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Chlorhexidine mouthrinse as an adjunctive treatment for gingival



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[Intervention Review]

Chlorhexidine mouthrinse as an adjunctive treatment for gingival health

Patrice James¹, Helen V Worthington², Carmel Parnell³, Mairead Harding⁴, Thomas Lamont⁵, Andrea Cheung⁶, Helen Whelton⁷, Philip Riley²

¹Oral Health Services Research Centre, Cork University Dental School and Hospital, Wilton, Cork, Ireland. ²Cochrane Oral Health, Division of Dentistry, School of Medical Sciences, Faculty of Biology, Medicine and Health, The University of Manchester, Manchester, UK. ³HSE Louth Meath Dental Service, Navan, Ireland. ⁴Oral Health Services Research Centre, Cork University Dental School and Hospital (UCC), Wilton, Cork, and HSE South (CHO 4), Cork, Ireland. ⁵Dundee Dental School, University of Dundee, Dundee, UK. ⁶Cork University Dental School and Hospital (UCC), Wilton, Cork, Ireland. ⁷School of Dentistry, University of Leeds, Leeds, UK

Contact address: Patrice James, Oral Health Services Research Centre, Cork University Dental School and Hospital, Wilton, Cork, Ireland. p.james@ucc.ie.

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ABSTRACT

Background

Dental plaque associated gingivitis is a reversible inflammatory condition caused by accumulation and persistence of microbial biofilms (dental plaque) on the teeth. It is characterised by redness and swelling of the gingivae (gums) and a tendency for the gingivae to bleed easily. In susceptible individuals, gingivitis may lead to periodontitis and loss of the soft tissue and bony support for the tooth. It is thought that chlorhexidine mouthrinse may reduce the build-up of plaque thereby reducing gingivitis.

Objectives

To assess the effectiveness of chlorhexidine mouthrinse used as an adjunct to mechanical oral hygiene procedures for the control of gingivitis and plaque compared to mechanical oral hygiene procedures alone or mechanical oral hygiene procedures plus placebo/control mouthrinse. Mechanical oral hygiene procedures were toothbrushing with/without the use of dental floss or interdental cleaning aids and could include professional tooth cleaning/periodontal treatment.

To determine whether the effect of chlorhexidine mouthrinse is influenced by chlorhexidine concentration, or frequency of rinsing (once/day versus twice/day).

To report and describe any adverse effects associated with chlorhexidine mouthrinse use from included trials.

Search methods

Cochrane Oral Health's Information Specialist searched the following databases: Cochrane Oral Health's Trials Register (to 28 September 2016); the Cochrane Central Register of Controlled Trials (CENTRAL; 2016, Issue 8) in the Cochrane Library (searched 28 September 2016); MEDLINE Ovid (1946 to 28 September 2016); Embase Ovid (1980 to 28 September 2016); and CINAHL EBSCO (Cumulative Index to Nursing and Allied Health Literature; 1937 to 28 September 2016). We searched ClinicalTrials.gov and the World Health Organization International Clinical Trials Registry Platform for ongoing trials. No restrictions were placed on the language or date of publication when searching the electronic databases.



Selection criteria

We included randomised controlled trials assessing the effects of chlorhexidine mouthrinse used as an adjunct to mechanical oral hygiene procedures for at least 4 weeks on gingivitis in children and adults. Mechanical oral hygiene procedures were toothbrushing with/without use of dental floss or interdental cleaning aids and could include professional tooth cleaning/periodontal treatment. We included trials where participants had gingivitis or periodontitis, where participants were healthy and where some or all participants had medical conditions or special care needs.

Data collection and analysis

Two review authors independently screened the search results extracted data and assessed the risk of bias of the included studies. We attempted to contact study authors for missing data or clarification where feasible. For continuous outcomes, we used means and standard deviations to obtain the mean difference (MD) and 95% confidence interval (CI). We combined MDs where studies used the same scale and standardised mean differences (SMDs) where studies used different scales. For dichotomous outcomes, we reported risk ratios (RR) and 95% CIs. Due to anticipated heterogeneity we used random-effects models for all meta-analyses.

Main results

We included 51 studies that analysed a total of 5345 participants. One study was assessed as being at unclear risk of bias, with the remaining 50 being at high risk of bias, however, this did not affect the quality assessments for gingivitis and plaque as we believe that further research is very unlikely to change our confidence in the estimate of effect.

Gingivitis

After 4 to 6 weeks of use, chlorhexidine mouthrinse reduced gingivitis (Gingival Index (GI) 0 to 3 scale) by 0.21 (95% CI 0.11 to 0.31) compared to placebo, control or no mouthrinse (10 trials, 805 participants with mild gingival inflammation (mean score 1 on the GI scale) analysed, high-quality evidence). A similar effect size was found for reducing gingivitis at 6 months. There were insufficient data to determine the reduction in gingivitis associated with chlorhexidine mouthrinse use in individuals with mean GI scores of 1.1 to 3 (moderate or severe levels of gingival inflammation).

Plaque

Plaque was measured by different indices and the SMD at 4 to 6 weeks was 1.45 (95% CI 1.00 to 1.90) standard deviations lower in the chlorhexidine group (12 trials, 950 participants analysed, high-quality evidence), indicating a large reduction in plaque. A similar large reduction was found for chlorhexidine mouthrinse use at 6 months.

Extrinsic tooth staining

There was a large increase in extrinsic tooth staining in participants using chlorhexidine mouthrinse at 4 to 6 weeks. The SMD was 1.07 (95% CI 0.80 to 1.34) standard deviations higher (eight trials, 415 participants analysed, moderate-quality evidence) in the chlorhexidine mouthrinse group. There was also a large increase in extrinsic tooth staining in participants using chlorhexidine mouthrinse at 7 to 12 weeks and 6 months.

Calculus

Results for the effect of chlorhexidine mouthrinse on calculus formation were inconclusive.

Effect of concentration and frequency of rinsing

There were insufficient data to determine whether there was a difference in effect for either chlorhexidine concentration or frequency of rinsing.

Other adverse effects

The adverse effects most commonly reported in the included studies were taste disturbance/alteration (reported in 11 studies), effects on the oral mucosa including soreness, irritation, mild desquamation and mucosal ulceration/erosions (reported in 13 studies) and a general burning sensation or a burning tongue or both (reported in nine studies).

Authors' conclusions

There is high-quality evidence from studies that reported the Löe and Silness Gingival Index of a reduction in gingivitis in individuals with mild gingival inflammation on average (mean score of 1 on the 0 to 3 GI scale) that was not considered to be clinically relevant. There is high-quality evidence of a large reduction in dental plaque with chlorhexidine mouthrinse used as an adjunct to mechanical oral hygiene procedures for 4 to 6 weeks and 6 months. There is no evidence that one concentration of chlorhexidine rinse is more effective than another. There is insufficient evidence to determine the reduction in gingivitis associated with chlorhexidine mouthrinse use in individuals with mean GI scores of 1.1 to 3 indicating moderate or severe levels of gingival inflammation. Rinsing with chlorhexidine mouthrinse for 4 weeks



or longer causes extrinsic tooth staining. In addition, other adverse effects such as calculus build up, transient taste disturbance and effects on the oral mucosa were reported in the included studies.

PLAIN LANGUAGE SUMMARY

Chlorhexidine mouthrinse to reduce gingivitis and plaque build-up

Review question

Does the use of chlorhexidine mouthrinse (a broad spectrum antiseptic) in addition to other conventional tooth cleaning help to control and improve gingivitis (inflammation of the gums)? Does the frequency of rinsing or the concentration of the solution affect the result and are there any undesirable side effects?

Background

Gingivitis is a reversible condition when gums become red, swollen and can bleed easily. Gingivitis is also very common - studies suggest that as many as 50% to 90% of adults in the UK and USA suffer from it. In susceptible people gingivitis may lead to periodontitis, which is not reversible. In periodontitis inflammation is accompanied by loss of ligaments and bone supporting the teeth. If untreated it may eventually lead to tooth loss. Severe periodontitis is the sixth most widespread disease globally.

It is recognised that maintaining a high standard of oral hygiene is important for the prevention and treatment of gingivitis. Toothbrushing is the main method for maintaining good oral hygiene. Other cleaning methods commonly used include dental floss, interdental brushes and scaling and polishing carried out by a dental professional. Some people have difficulty controlling plaque build-up and preventing gingivitis using only conventional tooth cleaning. Therefore people sometimes use mouthrinses containing chlorhexidine in addition to conventional tooth cleaning. These mouthrinses are readily available over the counter; prescriptions generally not being required outside the USA.

Study characteristics

We included 51 studies that analysed a total of 5345 participants. The evidence in this review is up to date as of 28 September 2016. Generally study participants were children and adults who had gingivitis or periodontitis, were able to use usual tooth cleaning methods and were healthy. We did not exclude studies where some or all participants had medical conditions or special care needs as we considered the use of mouthrinses with chlorhexidine to be particularly relevant to them. The included studies assessed the effects of chlorhexidine mouthrinse used for at least 4 weeks in addition to conventional tooth cleaning on gingivitis in children and adults.

Key results

There is high-quality evidence that the use of mouthrinses containing chlorhexidine in addition to usual toothbrushing and cleaning for 4 to 6 weeks or 6 months leads to a large reduction in the build-up of plaque. There is also high-quality evidence of a moderate reduction in gingivitis in people with a mild level of it, although because the level of disease was already low this is not considered clinically important. The nature of the available evidence does not allow us to determine the level of reduction of gingivitis in people with moderate to severe levels of it.

There was no evidence that one concentration or strength of chlorhexidine rinse was more effective than another.

Rinsing for 4 weeks or longer causes tooth staining, which requires scaling and polishing carried out by a dental professional. Other side effects have been reported, including build-up of calculus (tartar), temporary taste disturbance and temporary shedding of/damage to the lining of the mouth.

Quality of the evidence

One study was assessed as being at unclear risk of bias, with the remaining 50 being at high risk of bias, however this did not affect the quality assessments for gingivitis and plaque as we believe that further research is very unlikely to change our confidence in the estimate of effect.

SUMMARY OF FINDINGS

Summary of findings for the main comparison.

Chlorhexidine mouthrinse compared with placebo/control mouthrinse/no mouthrinse for gingival health

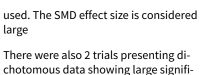
Patient or population: adults and children with gingivitis

Settings: any

Intervention: chlorhexidine mouthrinse

Comparison: placebo/control mouthrinse or no mouthrinse

Outcomes Illustrative comparative risks* (959		ive risks* (95% CI)	Relative ef- fect	Number of participants	Quality of the evidence	Comments		
	Assumed risk	Corresponding risk	(95% CI)	(studies)	(GRADE)			
	Control	Chlorhexidine						
Gingival Index 4 to 6 weeks (Löe and	The mean gingivitis scores ranged across	The mean gingivitis score in the chlorhexidine group was		805	⊕⊕⊕⊕ ^{2,3} high	The effect size at 6 months was similar		
Silness Gingival Index) (0 to 3 on an increasing scale)	control groups from 0.17 to 1.40 ¹	0.21 lower (0.11 to 0.31 low- er)		(10 trials)	-	Insufficient evidence for differences in effect size for different chlorhexidine concentration or frequency of use		
						Insufficient evidence to determine the effect size in individuals with moderate or severe levels of gingival inflammation on average (mean GI scores 1.1 to 3)		
Plaque 4 to 6 weeks	Plaque Index ranged from 0.75 to 1.06	The SMD was 1.45 lower in the chlorhexidine group in-		950	⊕⊕⊕⊕ ^{2,3} high	The effect for the Plaque Index (4 trials; 223 participants) was 0.58 (95% CI 0.39		
(various increas-	Turesky Modification	dicating a large reduction in plaque from 1.00 to 1.90		(12 trials)	-	to 0.78) lower		
ing scales including Plaque Index (0 to 3 scale) and Turesky Modification of the	of the Quigley and Hein Index ranged from 1.2 to 3.3	standard deviations				The effect for the Turesky Modification of the Quigley and Hein Index (5 trials; 546 participants) was 0.78 (95% CI 0.70 to 0.85) lower		
Quigley and Hein Index (0 to 5 scale))						There were also large effects for the plaque at 6 months		
Tooth staining 4 to 6 weeks	The mean tooth	The SMD for tooth staining in the chlorhexidine group		415	⊕⊕⊕⊝ ⁴	Data have not been converted to original scale as many different scales are		
o weeks	staining score was	was		(8 trials)	moderate	mai scale as many umerent scales are		



cant effect RR 5.41 (95% CI 2.03 to 14.47)

There was also a large effect for tooth staining for chlorhexidine at 7 to 12 weeks and 6 months

Other adverse effects

(various increasing

scales)

22 trials reported at least 1 adverse effect apart from extrinsic tooth staining and calculus formation in the chlorhexidine rinse arms. The adverse effects most commonly reported were taste disturbance/alteration (reported in 11 trials), effects on the oral mucosa including mucosal irritation, soreness, mild desquamation, mucosal ulceration/erosions, oral mucosal lesions (reported in 13 trials) and a general burning sensation and/or a burning tongue (reported in 9 trials)

CI: confidence interval; GI: Gingival Index; RR: risk ratio; SMD: standardised mean difference

1.07 (0.80 to 1.34) standard

deviations higher

GRADE Working Group grades of evidence

High quality: further research is very unlikely to change our confidence in the estimate of effect

Moderate quality: further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate Low quality: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate

Very low quality: we are very uncertain about the estimate

measured on differ-

ent scales

¹The mean gingivitis score for the control group was 0.93 (median is 1.0).

²Although most trials included in the meta-analyses were assessed as at high risk of bias we did not downgrade the GRADE assessments for this reason because we believe that further research is very unlikely to change our confidence in the estimate of effect.

³Not downgraded for high heterogeneity as results consistent.

⁴Downgraded as 8 trials at high risk of bias.

^{*}The basis for the assumed risk (e.g. the median control group risk across trials) is provided in footnotes. The corresponding risk (and its 95% CI) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI)



BACKGROUND

Description of the condition

Plaque-associated periodontal disease involves the inflammation of the gingivae (gums) and the supporting structures of the teeth. The first stage of the disease is plaque-associated gingivitis (inflammation of the gingivae), which is a reversible inflammatory condition characterised by redness and swelling of the gingivae and a tendency for the gingivae to bleed easily. It is caused by the accumulation and persistence of microbial biofilms (dental plaque) on the teeth (Löe 1965). Gingivitis is very common with studies suggesting that as many as 50% to 90% of adults in the UK and USA suffer from gingivitis (NICE 2012). In susceptible individuals, gingivitis may lead to the second stage of periodontal disease, periodontitis. Periodontitis is an irreversible condition, where inflammation of the gingivae is accompanied by connective tissue destruction and loss of the periodontal ligament and alveolar bone supporting the tooth. If untreated, periodontitis may ultimately lead to tooth loss. Severe periodontitis is the sixth most prevalent disease of man with a global age-standardised prevalence of 11.2% (Kassebaum 2014). It has a negative impact on oral-health-related quality of life (Al-Harthi 2013; Marcenes 2013; Needleman 2004; Tonetti 2015), affecting both physical function and social interaction (Cunha-Cruz 2007; Ng 2006). The response of the individual to gingival irritation by the microbial biofilm is modulated by local, systemic and genetic factors, and it is recognised that gingivitis will only progress to periodontitis in some individuals and at some sites. However as the microbial biofilm is a prerequisite for both conditions, gingival inflammation is considered a key risk factor in the initiation of periodontitis (Chapple 2015; Lang 2009). Indeed, consistent inflammation of the gingivae and gingival bleeding are predictive of future attachment loss and tooth loss (Schatzle 2004).

Although it is not possible to identify those individuals who will go on to develop severe periodontitis, the importance of maintaining a high standard of oral hygiene in the prevention and treatment of periodontal diseases is well recognised (Axelsson 2004; Ohrn 2009; van der Weijden 2011). Furthermore, the most important predictor of periodontal diseases, regardless of age is poor oral hygiene (Abdellatif 1987). Thus, the population approach to prevention of periodontal diseases should promote adequate plaque control and removal of the microbial biofilm for all individuals.

Description of the intervention

Mechanical disruption and removal of the microbial biofilm is commonly achieved by toothbrushing and interdental cleaning (Berchier 2008). However, adequate removal of the microbial biofilm with mechanical methods alone can be difficult for some, with many individuals finding it difficult to maintain an effective level of plaque control (Marsh 1992; Salzer 2015; Slot 2012; van der Weijden 2005; van der Weijden 2011; van der Weijden 2015). For these individuals, the level of plaque control that is achieved by regular mechanical cleaning alone may not be sufficient to prevent the onset or recurrence of periodontal diseases (Serrano 2015). Therefore chemical methods of removing the microbial biofilm such as antimicrobial mouthrinses have been recommended for use as either an adjunct to, or in some limited situations, a replacement for mechanical removal of the microbial biofilm

Chlorhexidine is a broad spectrum bisbiguanide antiseptic which was first introduced into clinical medicine in 1953 in the form of an antiseptic cream (Eley 1999; Foulkes 1973). It is a strong base and is practically insoluble in water (Karpinski 2015). In clinical medicine its water soluble salts, in particular chlorhexidine digluconate, are most commonly used (Foulkes 1973; Karpinski 2015). In dentistry, chlorhexidine is used in a variety of formulations and vehicles, such as mouthrinses, gels, sprays and varnishes. Problems can be encountered when delivering chlorhexidine as a gel or a toothpaste as the chlorhexidine may bind to ingredients in the products reducing its activity (Eley 1999). Although considerable advances have been made in recent years in the formulation of these and other chlorhexidine vehicles, mouthrinses have the advantage of having a long history of use for oral health. They are already accepted as part of a normal oral hygiene regimen, used following normal mechanical tooth cleaning (van der Weijden 2015). Chlorhexidine-containing mouthrinses are marketed worldwide under a number of trade names and are in widespread use for chemical plaque control. This family of rinses are mainly indicated for use as adjuncts to mechanical cleaning, in specific clinical situations where mechanical oral hygiene is difficult, such as postsurgery, in individuals with intermaxillary fixation, in fixed appliance orthodontic therapy and in individuals with intellectual and physical disabilities (Addy 1986). Chlorhexidine mouthrinse is mainly available in concentrations of 0.1%, 0.12% or 0.2% chlorhexidine digluconate as well as in low concentration (≤ 0.06%) rinse. The effect of chlorhexidine on the microbial biofilm is dosedependant (Keijser 2003). The optimum dose of chlorhexidine in a mouthrinse is considered to be 20 mg twice daily (Löe 1970) equivalent to 10 mL of 0.2% chlorhexidine mouthrinse (20 mg) or 15 mL of 0.12% chlorhexidine mouthrinse (18 mg) (Eley 1999; Keijser 2003). A rinse time of 30 seconds appears to be effective and acceptable although 60-second rinse times are also advocated (Bonesvoll 1974; Keijser 2003).

How the intervention might work

The action of chlorhexidine mouthrinse as a treatment adjunct in managing inflammation of the gingivae involves its antimicrobial properties, which reduce the bacterial load of the microbial biofilm and should therefore reduce levels of inflammation and help prevent primary and secondary plaque-associated periodontal disease. Chlorhexidine digluconate has the ability to kill a wide variety of micro-organisms including gram-positive and gram-negative bacteria (aerobes and anaerobes) (Emilson 1977) as well as fungi including yeasts (Greenstein 1986; Puig Silla 2008). When chlorhexidine mouthrinse is used, the positively charged (cationic) chlorhexidine molecule binds to the negatively charged microbial cell wall and interferes with the osmotic equilibrium of the micro-organism. Depending on the dose, chlorhexidine can have a bacteriostatic or bacteriocidal effect. At low concentrations, chlorhexidine is bacteriostatic causing leakage of low molecular weight substances from the microbial cell and inhibiting reproduction. At higher concentrations chlorhexidine is bacteriocidal and causes cell death by precipitating the cytoplasmic contents of the microbial cell (Greenstein 1986; Puig Silla 2008). This ability of chlorhexidine to adsorb and adhere to negatively charged surfaces is central to its success as an antimicrobial agent. The chlorhexidine molecule has the ability to adsorb to anionic (negatively charged) substrates such as hydroxyapatite, pellicle, salivary glycoproteins and mucous membranes exerting an immediate bacteriocidal effect. When



slowly released over time from the pellicle-coated enamel surface, it provides a prolonged bacteriostatic effect in vivo (Bonesvoll 1974; Bonesvoll 1974a; Bonesvoll 1978; Jenkins 1988; Rolla 1971). This property of chlorhexidine is known as its substantivity and is what sets chlorhexidine apart from many other antimicrobial agents.

Chlorhexidine mouthrinse use is associated with a number of local adverse effects the most common being the formation of brown staining on the teeth and oral tissues, particularly the tongue (Addy 1986). Several theories have been put forward to explain the cause of this characteristic staining, however most evidence points to the staining being a result of precipitation of anionic dietary chromogens (e.g. from tea, coffee, tannins from wine) onto adsorbed chlorhexidine cations (Addy 1985; Addy 1985a; Watts 2001). Other less common local adverse effects have also been reported including supragingival calculus accumulation (Eley 1999; Mandel 1994; Van Strydonck 2012), oral mucosal lesions (Addy 1986; Flotra 1971; Van Strydonck 2012), and altered taste perception (Addy 1986; Eley 1999; Marinone 2000; Van Strydonck 2012). The occurrence of side effects tends to be reduced with lower chlorhexidine concentrations (Addy 1986; Cumming and Löe 1973; Flotra 1971; Keijser 2003). With the exception of extrinsic staining of the teeth and calculus accumulation which require removal by professional tooth cleaning, the adverse effects are transient and resolve once chlorhexidine mouthrinse use has ceased (Flotra 1971; Greenstein 1986). These local adverse effects limit the use of chlorhexidine to short or moderate term use in specific clinical circumstances (van der Weijden 2015). Chlorhexidine is poorly absorbed by the oral tissues and the gastrointestinal tract and is considered to have very low toxicity (Foulkes 1973; Greenstein 1986). Parotid gland swelling has been reported following chlorhexidine mouthrinse use (Addy 1986; Eley 1999; van der Weijden 2010). There have been rare reports of type 1 hypersensitivity reactions to chlorhexidine used in the mouth or on the lips (Pemberton 2012) but severe reactions such as anaphylaxis do not appear to have occurred as a result of using chlorhexidine mouthrinse to maintain periodontal health.

Why it is important to do this review

Chlorhexidine mouthrinse is readily available and a prescription is generally not required outside the USA (Moran 2008). Its efficacy in controlling plaque and gingivitis in humans in the absence of active oral hygiene was first demonstrated over 40 years ago by Harald Löe and colleagues (Löe 1970). Nowadays chlorhexidine mouthrinse is mostly used as an adjunct to regular mechanical oral hygiene and is commonly regarded as the gold standard against which other antiplaque agents are measured (Jones 1997; Marsh 1992). The last decade has seen the publication of two meta-analyses (Gunsolley 2006; Gunsolley 2010) and a number of systematic reviews (including one metareview) of the effectiveness of chlorhexidine mouthrinse (Boyle 2014; Serrano 2015; van der Weijden 2015; Van Strydonck 2012). In most of these publications chlorhexidine mouthrinse was one of a number of antiplaque chemical agents under review. The one dedicated review of the effectiveness of chlorhexidine mouthrinse compared to placebo, control or regular oral hygiene employed slightly different inclusion criteria to this review but included tooth staining as an outcome in addition to gingivitis and plaque (Van Strydonck 2012). However, the search for this review was conducted in April 2011 and adverse effects other than extrinsic tooth staining were not prespecified outcomes in the review.

Given the numerous clinical indications for the use of chlorhexidine mouthrinse for gingivitis and plaque control, and the ready availability of chlorhexidine mouthrinse over-the-counter in many parts of the world, it is important to conduct a Cochrane systematic review of chlorhexidine mouthrinse as an adjunctive treatment for gingival health in order to provide, rigorous, up-to-date evidence for patients, oral health practitioners and policy makers. This should take into account the risk of bias of the studies that have been conducted, as well as assessing the possible adverse effects of the chlorhexidine mouthrinse.

OBJECTIVES

To assess the effectiveness of chlorhexidine mouthrinse used as an adjunct to mechanical oral hygiene procedures for the control of gingivitis and plaque compared to mechanical oral hygiene procedures alone or mechanical oral hygiene procedures plus placebo/control mouthrinse. Mechanical oral hygiene procedures were toothbrushing with/without the use of dental floss or interdental cleaning aids and could include professional tooth cleaning/periodontal treatment.

To determine whether the effect of chlorhexidine mouthrinse is influenced by chlorhexidine concentration, or frequency of rinsing (once/day versus twice/day).

To report and describe any adverse effects associated with chlorhexidine mouthrinse use (reporting of adverse effects is limited to those detailed in the included trials).

METHODS

Criteria for considering studies for this review

Types of studies

We included parallel-group randomised controlled trials (RCTs) irrespective of language or publication status where the duration of mouthrinsing was at least 4 weeks. The minimum duration of rinsing was selected to reflect the 'real life' use of chlorhexidine as a short-term adjunct to mechanical oral hygiene procedures. Studies where the duration of mouthrinsing was less than 4 weeks were excluded. Cross-over studies were excluded due to concerns that chlorhexidine could exert an effect beyond the washout period. It would not be possible to conduct a split-mouth study of mouthrinse use, however split-mouth studies comparing different scaling and root planing regimens and different periodontal surgical techniques often incorporate a chlorhexidine and placebo/control comparison. Such study designs were considered inappropriate to answer the question posed by this review and were excluded. We would have included properly designed cluster-RCTs if any such studies had met the inclusion criteria.

Types of participants

We included RCTs involving children or adults with gingivitis or periodontitis provided they were capable of performing normal mechanical oral hygiene procedures (in accordance with other Cochrane reviews, we classified all participants aged 16 years or less as children and those older than 16 years as adults). We included trials where some or all of the participants had medical conditions or special care needs as we considered the intervention to be particularly relevant to these individuals/groups.



Types of interventions

Experimental intervention: chlorhexidine mouthrinse (used at any concentration, volume, frequency or duration of rinsing) used in conjunction with mechanical oral hygiene procedures (toothbrushing with/without the use of dental floss or interdental cleaning aids and could include professional tooth cleaning/periodontal treatment).

Comparator interventions: mechanical oral hygiene alone (toothbrushing with/without the use of dental floss or interdental cleaning aids and could include professional tooth cleaning/periodontal treatment), or mechanical oral hygiene used in conjunction with placebo/control mouthrinse.

We included studies:

- where the chlorhexidine mouthrinse also contained fluoride;
- where gum care or antigingivitis dentifrices (that did not contain chlorhexidine) were used for mechanical oral hygiene provided that they were used in both experimental and comparator arms;
- with and without baseline prophylaxis (scale and polish) but all study arms had to have the same treatment.

We excluded studies:

- where chlorhexidine mouthrinse was used as a monotherapy in the absence of mechanical oral hygiene procedures;
- where the chlorhexidine mouthrinse formed part of a combined intervention with other agents (such as other chlorhexidine vehicles, e.g. dentifrice-containing chlorhexidine, or other antimicrobial agents (e.g. cetylpyridinium chloride (CPC)) that the comparator arm/s did not receive because we would have been unable to separate the effect of the chlorhexidine from the effect of the other active agents;
- where the mechanical oral hygiene procedures were not the same in both the chlorhexidine mouthrinse and the comparator arms e.g. studies or study arms where experimental or comparator arms received more intensive or more frequent professional mechanical cleaning or used additional mechanical cleaning aids e.g. dental floss as part of the intervention that was different from the other study arms;
- where chlorhexidine mouthrinse was applied locally e.g. with a brush or via subgingival irrigation.

Types of outcome measures

Primary outcomes

Gingivitis measured using any appropriate index/scale.

Only trials that reported gingivitis as an outcome were considered for inclusion in the review. The clinical features of gingivitis can be assessed non-invasively by visual assessment of colour, contour and gingival bleeding or invasively using an instrument to provoke bleeding or both (Lorenz 2009). The Gingival Index of Löe and Silness (GI) utilises both visual assessment and assessment of bleeding on provocation. It can be measured at specific sites using six index teeth (Löe and Silness 1963) or all teeth (Löe 1967). Each site is scored on a 0 to 3 increasing scale and an average score can be calculated for each tooth assessed. The mean score for an individual represents an average score for the areas examined. The Modified Gingival Index (MGI) (Lobene 1986) can be used as a full-mouth index or applied to selected teeth and is measured non-

invasively on a 0 to 4 increasing scale i.e. there is no attempt to use pressure to elicit bleeding. Many indices measure gingivitis solely by assessing bleeding on provocation but the techniques used to elicit bleeding can vary between indices. The Gingival Bleeding Index (Ainamo and Bay 1975) uses a blunt pocket probe to gently probe the orifice of the gingival crevice. If bleeding occurs within about 10 seconds after testing, a positive finding is recorded. The percentage of sites with gingival bleeding are calculated for the individual.

It is not uncommon for multiple measures of gingivitis to be presented within one study. Therefore the following hierarchy was developed to simplify data extraction.

- Where gingival inflammation and gingival bleeding were reported within the same study we extracted data on both outcomes.
- Where gingival inflammation was reported using more than one index within a study, the GI (Löe and Silness 1963; Löe 1967) was chosen over other indices and the MGI (Lobene 1986) was chosen next if the gingival index was not reported.
- Where gingival bleeding was reported within a study as bleeding on probing (BOP) (Ainamo and Bay 1975) and the proportion of sites with BOP using the GI (i.e. GI scores 2 or 3) (Löe and Silness 1963; Löe 1967), BOP was chosen over the proportion of sites with BOP (GI scores 2 or 3).

Within the gingival index, the severity (the average extent of the disease) was considered a measure of gingival inflammation and occurrence (the proportion of sites with BOP i.e. GI scores 2 or 3) was considered a measure of gingival bleeding.

Outcome data at 4 to 6 weeks, 7 to 12 weeks, 13 weeks to 6 months and greater than 6 months were collected and reported but the primary analysis of the gingivitis data was concerned with short-term use (4 to 6 weeks), along with a longer term measure at 6 months.

Secondary outcomes

Data on the following secondary outcomes were extracted from eligible trials that also reported gingivitis as an outcome.

- Dental plaque measured using any appropriate index/scale. Although dental plaque can be measured using a variety of different indices, the Plaque Index (Silness and Löe 1964) and the Turesky modification of the Quigley and Hein index (TQH) (Turesky 1970) are commonly used to measure dental plaque. Both use an increasing scale to quantify the amount of plaque present. The Plaque Index uses a 0 to 3 increasing scale to quantify the amount of plague on each surface (buccal, lingual, mesial and distal) of six index teeth. An average score for each tooth and for the individual can then be calculated. The TQH uses a 0 to 5 increasing scale to quantify the amount of plaque on the buccal and lingual surfaces of the teeth. An average score for the individual can then be calculated. Where dental plaque was measured using more than one index within a study; the Plaque Index (Silness and Löe 1964) was chosen over other indices and the TQH (Turesky 1970) was chosen next if the Plaque Index was not reported.
- Calculus measured using any appropriate index/scale.
- Tooth staining measured using any appropriate index/scale.



 Adverse effects (e.g. mucosal desquamation, taste disturbances, allergic reactions).

Search methods for identification of studies

Electronic searches

Cochrane Oral Health's Information Specialist conducted systematic searches in the following databases for randomised controlled trials and controlled clinical trials without language or publication status restrictions:

- Cochrane Oral Health's Trials Register (searched 28 September 2016) (Appendix 1);
- Cochrane Central Register of Controlled Trials (CENTRAL; 2016, Issue 8) in the Cochrane Library (searched 28 September 2016) (Appendix 2);
- MEDLINE Ovid (1946 to 28 September 2016) (Appendix 3);
- Embase Ovid (1980 to 28 September 2016) (Appendix 4);
- CINAHL EBSCO (Cumulative Index to Nursing and Allied Health Literature; 1937 to 28 September 2016) (Appendix 5).

Subject strategies were modelled on the search strategy designed for MEDLINE Ovid. Where appropriate, they were combined with subject strategy adaptations of the highly sensitive search strategy designed by Cochrane for identifying randomised controlled trials and controlled clinical trials as described in the *Cochrane Handbook for Systematic Reviews of Interventions* Chapter 6 (Lefebvre 2011).

Searching other resources

We searched the following trial registries for ongoing studies:

- US National Institutes of Health Ongoing Trials Register ClinicalTrials.gov (clinicaltrials.gov; searched 28 September 2016) (Appendix 6);
- World Health Organization International Clinical Trials Registry Platform (apps.who.int/trialsearch; searched 28 September 2016) (Appendix 7).

We searched the reference lists of included studies and relevant systematic reviews for further studies. We attempted to contact companies that manufacture chlorhexidine mouthrinse (Colgate Palmolive (Periogard), 3M (Peridex), GlaxoSmithKline (Corsodyl)) and companies that manufacture other types of mouthrinse (Johnson & Johnson (Listerine), P&G (Oral B brand)) to identify unpublished and ongoing studies.

We did not perform a separate search for adverse effects of interventions. Only the adverse effects that were described in the included studies were considered.

Data collection and analysis

Selection of studies

Two review authors independently and in duplicate screened the titles and abstracts of the list of studies identified by the searching process against the inclusion criteria for the review to identify eligible and potentially eligible studies. We obtained full-text copies of all the potentially eligible studies, and also of studies with insufficient information in the title/abstract to make a decision on eligibility. The review authors were not blinded to the journal name, authors, institution, results or sources of funding when assessing eligibility of the identified studies. Two review authors independently and in duplicate assessed the full-text copies to identify those that met the inclusion criteria. We contacted study authors for clarification or missing information relating to eligibility where necessary and feasible. We linked multiple reports of the same study together under one single study title. We resolved any disagreements on eligibility through discussion but, if this was not possible, an experienced member of the Cochrane Oral Health editorial team (H Worthington) was consulted to achieve consensus. We recorded any studies failing to meet the inclusion criteria, along with reasons for exclusion.

The eligibility of the non-English language reports were assessed by two review authors independently and in duplicate. Relevant sections of the reports were translated with the assistance of Cochrane Oral Health. Non-English language reports that met the inclusion criteria for the review were translated in full.

This process is summarised in the 'Study flow diagram' (Figure 1).



Figure 1. Study flow diagram.

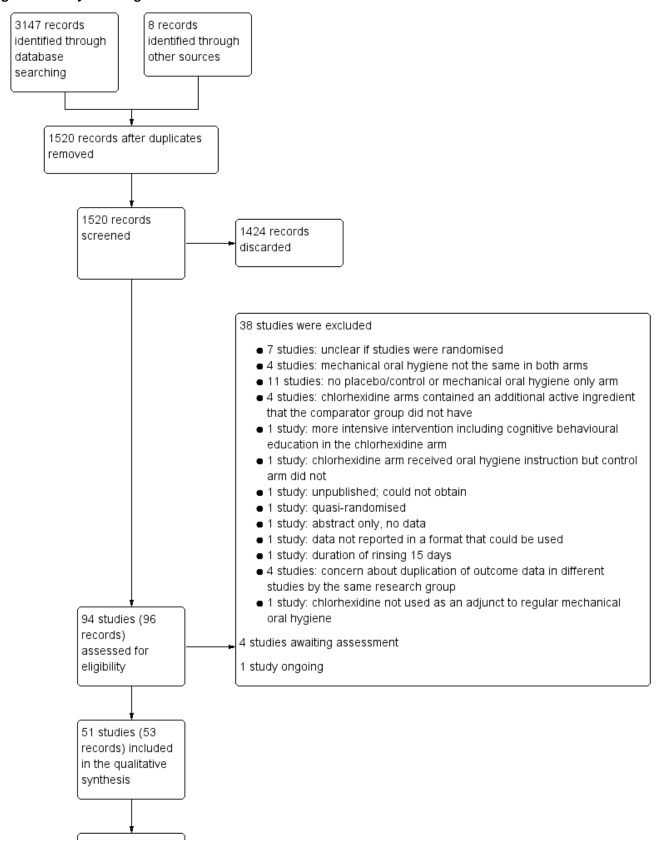




Figure 1. (Continued)

45 studies included in quantitative synthesis (meta-analysis)

Data extraction and management

Two review authors independently and in duplicate extracted data from the included studies using a customised Excel spreadsheet that was piloted on a small sample of studies. We contacted study authors for clarification or missing information where necessary and feasible. We resolved any disagreements through discussion and where agreement could not be reached, an experienced member of the Cochrane Oral Health editorial team (H Worthington) was consulted to achieve consensus.

We recorded the following data for each included study, which were then tabulated in the 'Characteristics of included studies' tables.

- Trial design, location/setting, number of centres, study duration.
- Details of the participants including demographic characteristics, criteria for inclusion and exclusion, and relevant information on gingivitis levels at baseline, numbers randomised to each study arm, and numbers analysed in each arm.
- Details of the type of experimental/comparator intervention, concentration and volume of mouthrinse, frequency and duration of rinsing, baseline prophylaxis (scale and polish), details of oral hygiene instruction (OHI), supervision of mouthrinsing, timing of mouthrinsing in relation to toothbrushing and postrinsing instructions.
- Details of the outcomes reported, including method and timing of assessment.
- Sample size calculations, source of study funding, information about adverse effects, proportion of smokers in the studies/ study arms and declarations/conflicts of interest.
- Outcome data: For gingivitis and plaque we extracted outcome data from the end point of each study with the exception of studies where the duration of rinsing was longer than 6 months. In this situation interim data for gingivitis and plaque at 6 months were extracted in addition to data at the end point of the study. We considered interim data on extrinsic tooth staining and calculus formation important in determining the time of onset of these adverse effects. Therefore, where they were reported, data on extrinsic tooth staining and calculus formation were extracted at interim time points as well as at the end point of each study. Outcome data were grouped according the following time intervals: 4 to 6 weeks, 7 to 12 weeks, 13 weeks to 6 months and greater than 6 months. Where data on extrinsic tooth staining or calculus were reported twice within one time interval, the later results were data extracted. Where an outcome was measured after rinsing has ceased, we considered the time point that rinsing ceased as the primary end point. Studies that measured gingivitis and plaque outcomes only after cessation of rinsing were included in a separate analysis of the long-term effects of rinsing with chlorhexidine mouthrinse on

- gingivitis and plaque. We did not extract extrinsic tooth staining and calculus data reported after rinsing had ceased.
- Where studies reported mean scores for the Gingival Index at 4 to 6 weeks or 6 months but did not report a measure of variance and a measure of variance could not be obtained from the authors of the studies, the variance was estimated from the standard deviations reported in similar trials that used the same index at the same time point as described in Chapter 16 of the Cochrane Handbook for Systematic Reviews of Interventions (Higgins 2011). Where mean scores and measures of variance were reported only in graphs and the data could not be obtained from the authors of the studies, data were estimated by reading the data off the graphs in the published report for inclusion in meta-analyses. The graphs were enlarged and the data were estimated independently and in duplicate. Any disagreements were resolved by discussion. A sensitivity analysis was conducted to gauge the effects of estimating these outcome data on the overall results.

Assessment of risk of bias in included studies

Two review authors assessed the risk of bias of all included studies, independently and in duplicate, using Cochrane's domain-based, two-part tool as described in Chapter 8 of the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). We contacted study authors for clarification or missing information where necessary and feasible. We tried to resolve any disagreements on risk of bias through discussion but, if this was not possible, an experienced member of the Cochrane Oral Health editorial team (H Worthington) was consulted to achieve consensus. A 'Risk of bias' table was completed for each included study. For each domain of risk of bias, we first described what was reported to have happened in the study in order to provide a rationale for the second part, which involved assigning a judgement of 'low risk' of bias, 'high risk' of bias, or 'unclear risk' of bias.

For each included study, we assessed the following seven domains of risk of bias.

Random sequence generation (selection bias): use of simple randomisation (e.g. random number table, computer-generated randomisation, central randomisation by a specialised unit), restricted randomisation (e.g. random permuted blocks), stratified randomisation and minimisation were assessed as at low risk of bias. Other forms of simple randomisation, such as repeated coin tossing, throwing dice or dealing cards, were also considered as at low risk of bias. If a study report used the phrase 'randomised' or 'random allocation' but with no further information, and if randomisation could not be confirmed, we assessed it as unclear for this domain.



- Allocation concealment (selection bias): use of centralised/ remote allocation, pharmacy-controlled randomisation (i.e. allocation of sequentially numbered mouthrinse containers of identical appearance) and sequentially numbered, sealed, opaque envelopes were assessed as at low risk of bias. If a study report did not mention allocation concealment, we assessed it as unclear for this domain.
- Blinding of participants and personnel (performance bias): if blinding was not mentioned, we assessed this domain as at unclear risk of bias. We only considered personnel blinding for studies with supervised mouthrinsing. If a study was described as double blind, we assumed that the intention was for participants and outcome assessors to be blinded to group allocation, however due to the propensity for chlorhexidine to stain the teeth and oral tissues and the association of its use with visible adverse effects such as mucosal desquamation and mucosal lesions, we considered the potential effect of these on blinding as follows.
 - Where tooth staining/adverse effects were reported narratively or measured using a scale (where relevant) and were higher in the chlorhexidine rinse arm/s, we considered that participants could have worked out which study arm they were in and this could have affected their oral health behaviours and hence the outcome and assessed this domain as at high risk of bias irrespective of whether participant blinding was indicated.
 - Where tooth staining/adverse effects were reported narratively or measured using a scale (where relevant) but were not higher in the chlorhexidine rinse arm/s and participant blinding was also indicated, we assessed this domain as at low risk of bias.
 - Where tooth staining/adverse effects were not reported, although we considered it likely in this situation that tooth staining or other visible adverse effects could have occurred and been noticeable in the chlorhexidine rinse arm/s after 4 weeks or more of rinsing, we assessed this domain as unclear as there was not enough information upon which to base a judgement.
 - Where it was not possible to blind participants and personnel due to differences in the experimental and comparator interventions this domain was assessed as at high risk of bias.
- Blinding of outcome assessment (detection bias): if blinding
 was not mentioned, we assessed this domain as at unclear
 risk of bias. If a study was described as double blind, we
 assumed that the intention was for participants and outcome
 assessors to be blinded to group allocation, however due to the
 propensity for chlorhexidine to stain the teeth and oral tissues
 and the association of its use with visible adverse effects such as
 mucosal desquamation and mucosal lesions, we considered the
 potential effect of these on blinding as follows.
 - * Where tooth staining/adverse effects such as mucosal desquamation and mucosal lesions that could be visible to the outcome assessor were reported narratively or measured using a scale (where relevant) and were higher in the chlorhexidine arm/s, we considered that outcome assessors could have worked out which study arm participants were in and therefore could not be adequately blinded. In this situation we assessed this domain as at high risk of bias irrespective of whether blinding of outcome assessment was indicated.

- * Where tooth staining/adverse effects such as mucosal desquamation and mucosal lesions that could be visible to the outcome assessor were reported narratively or measured using a scale (where relevant) but were not higher in the chlorhexidine arm/s and blinding of outcome assessment was also indicated, we assessed this domain as at low risk of bias.
- * Where tooth staining/adverse effects such as mucosal desquamation and mucosal lesions that could be visible to the outcome assessor were not reported, although we considered it likely that tooth staining or other visible adverse effects could have occurred and been noticeable in the chlorhexidine arm/s after 4 weeks of more of rinsing, we assessed this domain as unclear as there was not enough information upon which to base a judgement.
- Incomplete outcome data (attrition bias): if 10% or less of randomised participants were excluded from the analysis, we assessed this as at low risk of bias. However, when attrition was greater than 10%, assuming the missing participants in one group had a higher mean (e.g. gingivitis score) than those in the other group, as the attrition rate increased, so would the mean difference (MD) between groups, as described in Chapter 8 of the Cochrane Handbook for Systematic Reviews of Interventions (Higgins 2011). This situation led to a judgement of high risk of bias if we believed that the attrition was high enough to have resulted in a distortion of the true intervention effect, or if there was considerably greater attrition in one group than another. If attrition was greater than 10%, but with the additional factors of not being reported by group and insufficient reporting of reasons for attrition, this led to a judgement of unclear risk of bias. If it was not clear from the study report how many participants were randomised into each group, we assessed it as at unclear risk of bias for this domain.
- Selective reporting (reporting bias): if the study either reported outcomes not stated a priori in the methods section or the study protocol (if available) or did not report outcomes stated in the methods section, we assessed this as at high risk of bias. If gingivitis and plaque outcomes were reported with insufficient information to allow us to use the outcome data in a metaanalysis (e.g. no information on variance), we assessed it as at high risk of bias. We considered the reporting of adverse effects associated with the use of chlorhexidine mouthrinse to be important outcomes. For an assessment of low risk of bias we required either a statement that there were no adverse effects associated with the chlorhexidine rinse or if present, that adverse effects be reported by group. Where tooth staining was not reported or was measured using an index but insufficient data were reported to allow us to use the data in a meta-analysis we assessed it as at high risk of bias. Where tooth staining was not measured using an index but its presence/absence was reported we assessed this as at low risk of bias, provided that the information was reported for all participants by group.
- Other bias: any other potential source of bias that may feasibly alter the magnitude of the effect estimate (e.g. baseline imbalances in potentially important prognostic factors between intervention groups and differential diagnostic activity by outcome assessors).

We summarised the risk of bias as follows.



Risk of bias	Interpretation	In outcome	In included studies
Low risk of bias	Plausible bias unlikely to seriously alter the results	Low risk of bias for all key domains	Most information is from studies at low risk of bias
Unclear risk of bias	Plausible bias that raises some doubt about the results	Unclear risk of bias for one or more key domains	Most information is from studies at low or unclear risk of bias
High risk of bias	Plausible bias that seriously weakens confidence in the results	High risk of bias for one or more key domains	The proportion of information from studies at high risk of bias is sufficient to affect the interpretation of results

Unit of analysis issues

The participant was the unit of analysis. We were unable to use data from studies that reported number of sites rather than number of individuals in the meta-analyses, as failure to take the clustering into account could lead to an overestimate of effect, with narrower confidence intervals and smaller P values. Had we included cluster-RCTs we would have analysed the results taking account of the clustering present in the data, otherwise we would have used the methods outlined in Section 16.3.4 of the *Cochrane Handbook for Systematic Reviews of Interventions* in order to perform an approximately correct analysis (Higgins 2011).

Dealing with missing data

We attempted, where feasible, to contact the authors of studies to obtain missing data or for clarification. Where appropriate, we used the methods outlined in Chapter 16 of the *Cochrane Handbook for Systematic Reviews of Interventions* in order to estimate missing standard deviations for the main outcome only (gingivitis) (Higgins 2011). We did not use any further statistical methods or carry out any further imputation to account for missing data. Where mean scores and measures of variance were only reported in graphs and could not be obtained from the authors of the studies, data were estimated by reading the data off the graphs in the published report for inclusion in meta-analyses. The graphs were enlarged and the data were estimated independently and in duplicate. Any disagreements were resolved by discussion.

Assessment of heterogeneity

If meta-analyses were performed, we assessed the possible presence of heterogeneity visually by inspecting the point estimates and confidence intervals on the forest plots; if the confidence intervals had poor overlap then heterogeneity was considered to be present. We also assessed heterogeneity statistically using a Chi² test, where a P value < 0.1 indicated statistically significant heterogeneity. Furthermore, we quantified heterogeneity using the I² statistic. A guide to interpretation of the I² statistic given in Section 9.5.2 of the *Cochrane Handbook for Systematic Reviews of Interventions* is as follows (Higgins 2011):

- 0% to 40%: might not be important;
- 30% to 60%: may represent moderate heterogeneity;
- 50% to 90%: may represent substantial heterogeneity;
- 75% to 100%: considerable heterogeneity.

Assessment of reporting biases

Assessment of reporting bias within studies has already been described in the section Assessment of risk of bias in included studies.

Reporting biases can occur when reporting (or not reporting) research findings is related to the results of the research (e.g. a study that did not find a statistically significant difference/result may not be published). Reporting bias can also occur if ongoing studies are missed (but that may be published by the time the systematic review is published), or if multiple reports of the same study are published, or if studies are not included in a systematic review due to not being reported in the language of the review authors. We attempted to limit reporting bias in the first instance by conducting a detailed, sensitive search, including searching for ongoing studies, and any studies not reported in English were translated. If there were more than 10 studies included in a metaanalysis for the primary outcome (gingivitis), we assessed the possible presence of reporting bias by testing for asymmetry in a funnel plot. We carried out statistical analysis using the methods described by Egger 1997 for continuous outcomes.

Data synthesis

We carried out a meta-analysis only where studies of similar comparisons reported the same outcomes at the same time interval. We combined mean differences (MDs) where studies used the same scale and standardised mean differences (SMDs) where studies used different scales for continuous outcomes. For dichotomous outcomes, we reported risk ratios (RR) and 95% confidence intervals. Due to anticipated heterogeneity we used random-effects models for all meta-analyses.

We reported the results from studies not suitable for inclusion in a meta-analysis both in the results and in additional tables.

Subgroup analysis and investigation of heterogeneity

Where there were sufficient studies, we carried out the following subgroup analyses.

- Baseline prophylaxis (scale and polish) versus none.
- Children versus adults.
- Different chlorhexidine concentrations.
- Initial levels of gingivitis.
- Gingivitis only versus gingivitis + periodontitis.



Sensitivity analysis

In order to ensure our conclusions were robust, we carried out sensitivity analyses (where there were sufficient studies for each outcome) by excluding studies where outcome data were read from graphs, studies where standard deviations were estimated and studies where periodontal treatment was conducted during the study.

We would have carried out sensitivity analyses by excluding studies at high and unclear risk of bias, however this was not possible as apart from one study at unclear risk of bias, all of the included studies were at high risk of bias.

Presentation of main results

We produced a 'Summary of findings' table for chlorhexidine mouthrinse compared to placebo/control mouthrinse or no mouthrinse for the following outcomes: gingivitis measured using the Gingival Index at 4 to 6 weeks, plaque at 4 to 6 weeks, tooth staining at 4 to 6 weeks and other adverse effects. We used GRADE methods (GRADE 2004), and the GRADEpro GDT online tool for developing 'Summary of findings' tables (www.guidelinedevelopment.org). We assessed the quality of the body of evidence for each outcome by considering the overall risk of bias of the included studies, the directness of the evidence, the inconsistency of the results, the precision of the estimates, and the risk of publication bias. We categorised the quality of each body of evidence as high, moderate, low, or very low.

RESULTS

Description of studies

Results of the search

The searches resulted in 1520 references after de-duplication. Two review authors screened the titles against the inclusion criteria for this review, independently and in duplicate, discarding 1424 references in the process. We obtained full-text copies of the remaining 96 references (94 studies) and examined them independently and in duplicate, excluding 38 studies at this stage. A further four studies are awaiting assessment of their eligibility to be included in the review and one study is ongoing. Therefore, fiftyone studies (53 records) met the inclusion criteria for this review. This process is presented diagrammatically in Figure 1.

Included studies

Characteristics of the trial designs and settings

Fifty-one studies met the inclusion criteria for this review. All studies were of parallel-group design. Sixteen studies had two study arms and compared chlorhexidine rinse with placebo rinse. However, thirty-five studies had multiple study arms and not all study arms were relevant to this review. Twenty-two studies had three arms; 10 had four arms and two had five arms. One of the studies with four study arms (Flotra 1972) had three chlorhexidine rinse groups and one placebo rinse group but presented data only for two groups: the three chlorhexidine study arms combined and the placebo group. One additional study (Feres 2012) had three main therapeutic arms which were each divided into chlorhexidine rinse and placebo rinse groups using a 2 x 3 factorial design.

Sixteen studies were conducted in the USA, six in Brazil, five in the UK, four in Germany, three in India, three in Sweden, two

in each of the following countries: Canada, Mexico and Spain, and one in each of the following countries: Argentina, China, Iran, Italy, Norway, Switzerland, the Netherlands and Turkey. The settings of the studies were diverse and often poorly reported. Twenty-seven studies were conducted in a dental school/university setting and an additional three studies were assumed, based on the information provided, to have been conducted in a dental school/university setting (Emling 1992; Pereira 2011; Van Strydonck 2008). Two studies were conducted under "dental office conditions" (Grossman 1986; Grossman 1989), two in general dental practices (Eaton 1997; Fine 1985) and one in a private practice limited to periodontics (Sanz 1989). Three studies were conducted in a school setting (de la Rosa 1988; de la Rosa 1988b; Jayaprakash 2007). The remaining studies were conducted in military garrisons (Flotra 1972; Hase 1998); pensioners/nursing homes (Lopez-Jornet 2012; Weitz 1992); a bone marrow transplant unit (Ferretti 1987) and a knitting factory (Corbet 1997). Three studies were conducted at what we assumed were clinical research facilities (Charles 2004; Jose 2015; Stookey 2005) and in four studies the setting of the study was not clear (Flemmig 1990; Hase 1995; Sanz 1994; Taller 1993). The numbers of centres involved in the studies was often unclear but it appeared that all but two of the studies were conducted at a single centre. One study (Eaton 1997) was conducted at five suburban dental practices in Southern England and another study (Jose 2015) was conducted at two sites (Manchester and Wirral) in the UK.

In 19 studies direct support from industry was explicitly stated. Nine studies received support from P&G; three from GSK; two from Teledyne and one from each of the following companies: GABA, J&J, Oral B, Smithkline Beecham and Warner Lambert. Three studies that did not appear to be funded directly by industry explicitly stated that oral care products used in the study had been provided by industry (Anderson 1997; Sanz 1994; Stookey 2005). Three studies (Hase 1995; Hase 1998; Lang 1998) were associated with Biosurface Pharma AB through authorship and another study was associated with 'Dental Products Testing' and Pfizer through authorship (Charles 2004). Of the remaining studies that reported a source of support, two appeared to be funded by universities (Rahmani 2006; Taller 1993); six appeared to be funded by research grants (Anauate-Netto 2014; Bajaj 2011; Corbet 1997; Faveri 2006; Feres 2009; Feres 2012); and in 17 studies there was no statement regarding funding or support. Only one study (Zimmer 2015) stated that the conduct of the research and final decisions regarding the study report were independent of the funder (GSK).

Only 13 of the studies described a sample size calculation. For six of these studies (Faveri 2006; Feres 2009; Feres 2012; Graziani 2015; Turkoglu 2009; Van Strydonck 2008) the sample size required was reported and achieved but for three of these studies the sample size was based on pocket probing depth (PPD) (Faveri 2006; Feres 2012) and clinical attachment loss (CAL) (Feres 2009) rather than gingivitis. Two studies reported the sample size calculation and number of participants required in each group but the minimum sample size was not achieved at the end of the studies (Hase 1998; Lang 1998). Five of the studies (Axelsson 1987; Brightman 1991; Charles 2004; Eaton 1997; Zimmer 2015) described the sample size calculation but not the required sample size so we were unable to determine if the required sample size was achieved. In one study the sample size calculation was not described but the required sample size was reported and achieved (Bhat 2014).



Characteristics of the participants

A total of 6995 participants provided data for the included studies. The total number of participants providing data for the study arms relevant to this review (excluding the irrelevant study arms) was 5345 with the numbers analysed in each study ranging from 20 to 852.

Six studies involved children and adolescents (Anderson 1997; Bajaj 2011; Brightman 1991; de la Rosa 1988; de la Rosa 1988b; Jayaprakash 2007). The age range of the children was 8 to 16 and three of these studies also included some young adults aged 17 and 18 (Brightman 1991; de la Rosa 1988; de la Rosa 1988b). Four studies while predominantly focused on adults, included children and adolescents among the participants: in Axelsson 1987 participants were aged 16 to 50 years old; in Fine 1985 no participants under the age of 12 were included; and in Navarro 1998 the age range of participants was 14 to 35 years old. In Ferretti 1987, a study involving individuals undergoing bone marrow transplantation and chemoradiotherapy, the age range was 5 to 51 years old. The remaining forty-one studies involved adults. The age range of adult participants was 17 to 94 years with the mean age ranging from 20 to 83.

Twenty-two studies had a greater proportion of females than males and in four studies, the proportion of males was greater (Lang 1998; Southern 2006; Stookey 2005; Taller 1993). In five studies there was an equal balance of males to females (Bhat 2014; Graziani 2015; Pereira 2011; Turkoglu 2009; Zimmer 2006). Five studies involved male participants only (de la Rosa 1988; de la Rosa 1988b; Hase 1995; Hase 1998; Navarro 1998) and in the remaining 15 studies, the proportion of males to females was not reported. Eight studies included smokers among the participants (Charles 2004; Eaton 1997; Ernst 2005; Jose 2015; Joyston-Bechal 1993; Overholser 1990; Zimmer 2006; Zimmer 2015) with the proportion of smokers in each study ranging from 11% to 41%. 10 studies excluded smokers and in the remaining 33 studies, the smoking status of the participants was not reported.

Twenty-four studies reported gingivitis at baseline using the Gingival Index of Löe and Silness (Löe and Silness 1963; Löe 1967). In one study involving dental students (Lucas 1999) where normal gingiva were obtained in a pre-experimental phase by scaling and polishing and twice daily brushing, the mean baseline gingivitis score was zero. In two other studies (Navarro 1998; Jayaprakash 2007) gingivitis at baseline was particularly low, mean gingival index scores were 0.049 and 0.057 respectively. In the remaining 21 studies the mean gingival index ranged from 0.53 to 1.87 with an overall mean of 1.09.

In many of the included studies, the periodontal status of the participants was not explicitly described. In 15 studies, the participants appeared to have gingivitis but no periodontitis whereas in nine studies, the participants had periodontitis of varying severity in addition to gingivitis (Chaves 1994; Corbet 1997; Ernst 2005; Faveri 2006; Feres 2009; Feres 2012; Graziani 2015; Rahmani 2006; Sanz 1989). In three of these studies the periodontitis was untreated at baseline (Faveri 2006; Feres 2009; Feres 2012) and in one study the participants required osseous periodontal surgery (Sanz 1989). In 16 studies it was unclear whether the participants had periodontitis in addition to gingivitis but it was deemed likely that at least some of the participants in these studies had periodontitis; two studies involved older adults

(Lopez-Jornet 2012 (aged > 65 years) and Weitz 1992 (mean age 83)) and in the remaining 14 studies, the inclusion criteria allowed for the participants to have deep pockets and clinical attachment loss but the periodontal status of the people actually included in the study was not described. In 11 studies the participants clearly had gingivitis at baseline, but it was unclear whether the participants also had periodontitis.

Characteristics of the interventions

Nine studies compared chlorhexidine rinse and mechanical oral hygiene with mechanical oral hygiene alone. Two of these studies had additional chlorhexidine mouthrinse arms that were also included in the analysis (Jose 2015; Zimmer 2015). All of these studies except Jose 2015 also had additional arms that were not relevant to this review. Forty-two studies compared chlorhexidine rinse with placebo or control rinse. Six of these studies had additional chlorhexidine mouthrinse arms. Data from five of these six studies were included in the analysis (Flotra 1972; Graziani 2015; Hoffmann 2001; Jayaprakash 2007; Overholser 1990; Segreto 1986). We could not include data from the three separate chlorhexidine arms for Flotra 1972 because the authors combined the three arms into one experimental group. Twenty of these 42 studies comparing chlorhexidine rinse with placebo or control rinse also had additional arms that were not relevant to this review.

Of the studies with one concentration of chlorhexidine mouthrinse, 10 evaluated 0.2% chlorhexidine mouthrinse; 29 evaluated 0.12% chlorhexidine mouthrinse; two evaluated 0.1% chlorhexidine mouthrinse (Bajaj 2011; Ernst 2005) and four evaluated 0.05% or 0.06% chlorhexidine mouthrinse (Jayaprakash 2007; Joyston-Bechal 1993; Zimmer 2006; Zimmer 2015). Four studies had additional arms evaluating different chlorhexidine concentrations (Axelsson 1987; Flotra 1972; Hoffmann 2001; Segreto 1986) and in two studies, the concentration of the chlorhexidine rinse was not reported (Fine 1985; Turkoglu 2009).

Most of the studies evaluating 0.2% chlorhexidine mouthrinse used a volume of 10 mL twice daily for one minute whereas most of the studies evaluating 0.1% or 0.12% chlorhexidine mouthrinse used a volume of 15 mL twice daily. Duration of rinsing for the studies of 0.1% and 0.12% chlorhexidine mouthrinse varied from 30 seconds to 1 minute but most of the studies employed a 30-second rinse. The studies of 0.05% and 0.06% chlorhexidine mouthrinse predominantly used a volume of 10 mL once or twice a day for 30 seconds. In some studies with multiple arms different volumes of rinse were used in different arms: in three studies the placebo/control arms and other non-chlorhexidine active rinse groups rinsed with 20 mL while the chlorhexidine arms rinsed with 10 mL (Axelsson 1987) or 15 mL (Charles 2004; Overholser 1990); in another study the 0.1% chlorhexidine arm rinsed with 15 mL while the two 0.06% chlorhexidine arms used 10 mL of rinse (Hoffmann 2001).

In six studies the volume of rinse used by participants was not reported. In three of these studies participants rinsed twice daily but rinse volume and duration were not reported (Chaves 1994; Rahmani 2006; Sanz 1994). In two of the studies, participants rinsed once a day for 30 seconds with an unknown volume of rinse (Fine 1985; Zimmer 2006). In another study rinse volume, frequency and duration were not reported (Emling 1992).



Of the nine studies that compared chlorhexidine rinse and mechanical oral hygiene with mechanical oral hygiene alone, six involved regular mechanical cleaning using a toothbrush and toothpaste (Chaves 1994; Flemmig 1990; Jose 2015; Taller 1993; Zimmer 2006; Zimmer 2015). In one study floss and wood points were also used by participants (Fine 1985). In another study participants underwent conventional ultrasonic debridement (Rahmani 2006) and were given interdental cleaning aids, and participants in a further study used a toothbrush and no toothpaste (Van Strydonck 2008).

Thirty-five of the 51 studies compared chlorhexidine mouthrinse with placebo and seven studies compared chlorhexidine mouthrinse with control. The control rinses were 5% hydroalcohol solution (Charles 2004); coloured and flavoured 5% hydroalcohol solution (Overholser 1990); 0.5% benzene alcohol (Flotra 1972); saline (Bhat 2014; Graziani 2015) and water (Bajaj 2011; Hoffmann 2001). In one of these studies comparing 0.2% chlorhexidine mouthrinse with placebo rinse, participants were instructed to brush without toothpaste before using the mouthrinse (Hase 1995). In another study comparing 0.12% chlorhexidine mouthrinse with placebo rinse, all participants used a gum care dentifrice (Blend-amed) (Sanz 1994).

Thirty-six studies reported a baseline prophylaxis to remove plaque and thus assess the potential for chlorhexidine mouthrinse to inhibit plaque accumulation and its ability to reduce gingivitis. In two of the studies that reported a baseline prophylaxis, an additional prophylaxis was also provided 2 weeks before baseline (Brecx 1993; Hoffmann 2001). In another study the baseline prophylaxis was conducted during the first week of the study (Stookey 2005). Participants in eight of the nine studies in which some or all of the participants had periodontitis at baseline either had professional prophylaxis at baseline (Chaves 1994; Ernst 2005; Graziani 2015), or adjunctive professional non-surgical (Faveri 2006; Feres 2009; Feres 2012; Rahmani 2006) or surgical periodontal treatment (Sanz 1989) during the study. In the three studies where the participants with periodontitis had a professional prophylaxis at baseline only, scaling and polishing was supragingival (Graziani 2015) or the extent of the professional cleaning was not clear (Chaves 1994; Ernst 2005).

Participants in six studies did not have a baseline prophylaxis. These studies were thus designed to assess the potential for chlorhexidine mouthrinse to reduce plaque and gingivitis in individuals who do not obtain professional tooth cleaning or scaling prior to commencing mouthrinse use (Corbet 1997; Jayaprakash 2007; Jenkins 1993; Lopez-Jornet 2012; Southern 2006; Weitz 1992). One of these studies included participants with untreated periodontitis (Corbet 1997). Participants in this study received no professional prophylaxis and no oral hygiene instructions. In seven studies it was not clear whether a baseline prophylaxis had been carried out (Anauate-Netto 2014; Bajaj 2011; Bhat 2014; Emling 1992; Taller 1993; Turkoglu 2009; Zimmer 2015). In one study (Zimmer 2006), the screening examination was followed by calculus removal in the lower front teeth only. In another study, where it was not clear whether the participants had periodontitis in addition to gingivitis, there was no prophylaxis at baseline but supragingival scaling was performed at week 9 and subgingival scaling was performed after 13 weeks (Flotra 1972).

Twenty-two studies reported that participants received some form of oral hygiene instruction. Where details were reported

about the oral hygiene instruction, it usually involved instruction about toothbrushing technique. In two studies, oral hygiene instruction included instruction in the use of interdental cleaning aids (i.e. interdental brushes or dental floss or both) (Chaves 1994; Graziani 2015) and in another three studies participants were given interdental cleaning aids; presumably instruction in their use was included in the oral hygiene instruction but this not clear from the text (Fine 1985; Rahmani 2006; Taller 1993). In two studies participants were advised on duration and frequency of brushing but not technique (Pereira 2011; Van Strydonck 2008) and in another study, participants were given brief instruction about using the toothbrush they were given but were not given any instructions about duration or technique of toothbrushing (Zimmer 2006).

The participants in 27 studies did not receive any oral hygiene instruction. In 18 of these studies participants were advised to continue their usual oral hygiene practices throughout the duration of the studies. The participants in one of these 18 studies (Ferretti 1987) were undergoing bone marrow transplant and chemoradiotherapy and their usual oral hygiene regimen involved oral cleaning with a foam rubber toothbrush substitute and saline solution irrigations. In another of these studies where continuation of usual oral hygiene practices was advised, it was explicitly stated that use of interproximal cleaning devices was permitted if they were part of participants' usual oral hygiene routine (Zimmer 2015). In two studies it was not clear whether participants received oral hygiene instruction (Bajaj 2011; Stookey 2005). In one of these studies toothbrushing was supervised Monday to Friday in the mornings but it is not clear if oral hygiene instructions were provided to participants (Stookey 2005).

In 38 studies mouthrinsing was unsupervised. Mouthrinsing was fully supervised in only three studies, all conducted in a school setting (de la Rosa 1988; de la Rosa 1988b; Jayaprakash 2007). In 10 studies mouthrinsing was partially supervised for practical reasons. In five of the studies mouthrinsing was fully supervised on weekdays only (Axelsson 1987; Bajaj 2011; Bhat 2014; Lang 1998; Overholser 1990); in another two studies, mouthrinsing was supervised once a day on weekdays only (Charles 2004; Stookey 2005). In another three studies the mouthrinsing was supervised except for two weeks during the Chinese New Year (Corbet 1997); after participants were discharged from hospital (Ferretti 1987) and when participants were at military training (Hase 1998).

Thirty studies reported instructing participants to use the mouthrinse after toothbrushing. Three of these studies specified waiting 30 minutes after toothbrushing before using the mouthrinse (Feres 2012; Turkoglu 2009; Zimmer 2015). One of these studies specified waiting five timed minutes before mouthrinsing (Jose 2015). One study reported instructing participants to use mouthrinse at a separate time to toothbrushing (Charles 2004). In the remaining 20 studies the timing of mouthrinsing in relation to toothbrushing was not reported.

In most of the studies the duration of the intervention (mouthrinsing with chlorhexidine) was 6 months or less broken down as follows: 4 to 6 weeks (15 studies); 2 months (9 studies); 10 weeks (1 study); 3 months (9 studies); 4 months (2 studies) or 6 months (13 studies). In two studies the duration of the intervention was longer than 6 months; one study of 9 months' duration (Bajaj 2011) and another of 24 months' duration (Banting 1989).



In most studies the final follow-up assessment was at the end of the intervention phase. However in four studies participants ceased mouthrinsing before the final outcome assessment. In these studies the final assessment of the outcomes was carried out 4 weeks (Faveri 2006; Ferretti 1987), 4 months (Feres 2009) and 10 months (Feres 2012) after mouthrinsing had ceased. For this review, we considered the time that mouthrinsing ceased to be the primary end point for data extraction.

Characteristics of the outcomes

Gingivitis

Gingival inflammation

Thirty-eight studies included gingival inflammation as an outcome. The most commonly used index for measuring gingival inflammation was the Gingival Index of Löe and Silness (GI) (Löe and Silness 1963; Löe 1967). The following indices were also used to measure gingival inflammation: the Modified Gingival Index (MGI) (Lobene 1986), the Papillary Bleeding Score (PBS) (Loesche 1979), and the Papillary Marginal Gingival Index (PMGI) (de la Rosa and Sturzenberger 1976). Thirty-one studies measured gingival inflammation using the GI; two studies used the MGI (Eaton 1997; Overholser 1990); one study used the PBS (Anauate-Netto 2014); and four studies used the PMGI (de la Rosa 1988; de la Rosa 1988b; Ferretti 1987; Segreto 1986). In one of these studies (Segreto 1986), the PMGI was measured with visual inspection only (no probing) in addition to the GI. One study reported a total mean score for a non-specific gingival inflammation index (Fine 1985).

Gingival bleeding

Thirty-four studies included gingival bleeding as an outcome. The following indices were used to measure gingival bleeding: the GI (Löe and Silness 1963; Löe 1967) scores 2 and 3, the Gingival Bleeding Index of Ainamo and Bay (Ainamo and Bay 1975), the Eastman Interdental Bleeding Index (Caton and Polson 1985), the Bleeding Index/Bleeding on Marginal Probing (Saxton and van der Ouderaa 1989; van der Weijden 1994), a modification of the method described by Cowell et al (Cowell 1975), Papillary Bleeding Score (PBS) (proportion of sites with a PBS ≥ 2) (Loesche 1979) and the Papillary Bleeding Index (Muhlemann 1977; Saxer 1975).

Thirteen studies reported gingival bleeding using scores 2 and 3 from the GI. One of these studies (Southern 2006) also reported bleeding on probing after 15 seconds. Another two studies (Hase 1998; Lang 1998) measured bleeding on probing and the method used suggests that it was based on scores 2 and 3 of the GI. Three studies (Graziani 2015; Pereira 2011; Rahmani 2006) measured gingival bleeding using the Gingival Bleeding Index of Ainamo and Bay. Two studies (Brightman 1991; Overholser 1990) used the Eastman Interdental Bleeding Index. Two studies (Ernst 2005; Van Strydonck 2008) reported the Bleeding Index/Bleeding on Marginal Probing. One study (Joyston-Bechal 1993) used a modification of the method described by Cowell et al. One study (Anauate-Netto 2014) reported the proportion of sites with a PBS \geq 2 and three studies (Turkoglu 2009; Zimmer 2006; Zimmer 2015) used the Papillary Bleeding Index. A further seven studies measured the proportion of sites with bleeding on probing but were not specific about the index used (Chaves 1994; Faveri 2006; Feres 2009; Feres 2012; Flemmig 1990; Hase 1995; Taller 1993); three of these studies also reported the mean proportion of sites with gingival bleeding but were not specific about the index used (Faveri 2006; Feres 2009; Feres 2012).

Plaque

Forty-eight studies included plaque as an outcome. The majority of the studies measured dental plaque accumulation using the Plaque Index of Silness and Löe (Silness and Löe 1964) or the Turesky modification of the Quigley and Hein Index (TQH) (Turesky 1970). The following indices were also used to measure plaque: Soparker's modification of the TQH (Lobene 1982), the Modified Proximal Plaque Index (MPPI) (Zimmer 2005), the Quigley and Hein Plaque Index (Quigley 1962), the Approximal Plaque Index (API) (Lange 1977), a full-mouth plaque score (FMPS) (O'Leary 1972), and the method described by Greene and Vermillion 1964).

Twenty-one studies reported using the Plaque Index of Silness and Löe. Nineteen studies reported using TQH. One of these studies (Jose 2015) used Soparker's modification of the TQH, and one of these studies (Zimmer 2015) reported the MPPI in addition to TQH. Two studies in the same report that used the TQH (de la Rosa 1988; de la Rosa 1988b) did not report any outcome data for plaque. One study (Zimmer 2006) reported both the MPPI and the Quigley and Hein Plaque Index. The following indices were reported in one study each: the Quigley and Hein Plaque Index (Turkoglu 2009), the API (Ernst 2005), FMPS (Graziani 2015), and the method described by Greene and Vermilion (Lucas 1999). Three studies (Faveri 2006; Feres 2009; Feres 2012) measured the presence or absence of plaque accumulation but were not specific about the index used.

Adverse effects

Calculus

Twelve studies reported calculus as an outcome. Seven studies (Banting 1989; Charles 2004; Emling 1992; Flemmig 1990; Hase 1998; Overholser 1990; Sanz 1994) reported using the Volpe-Manhold Calculus Index (Manhold 1965; Volpe 1965). Two studies (Corbet 1997; Lang 1998) reported using the Calculus Surface Index (CSI) (Ennever 1961). One study (Anderson 1997) reported using the Retention Index (Björby and Löe 1966; Löe 1967). One study (Turkoglu 2009) used a calculus index described in the National Institutes of Health (NIH) report 'Oral health of United States adults' (report not available online) and in one study the index used was not specified (Grossman 1986). In two additional studies accumulation of calculus in the chlorhexidine rinse groups was noted but not measured using an index or scale (de la Rosa 1988; de la Rosa 1988b).

Extrinsic tooth staining

Twenty-one studies included extrinsic tooth staining as an outcome but this was measured and reported in a variety of different ways. In 16 studies, the following indices were used to measure extrinsic tooth staining: the Discolouration Index (Lobene 1968), the Discolouration Index of Lang and Räber (Lang and Räber 1981), the Shaw and Murray Index (Shaw 1977), the Gingival Modification of the Lobene Stain Index (GMSI) (Grundemann 2000; Lobene 1968), and Meckels stain grading method (Lang 1982). Seven studies (Axelsson 1987; Charles 2004; Emling 1992; Ernst 2005; Joyston-Bechal 1993; Navarro 1998; Overholser 1990) reported the Lobene Discolouration Index. Five studies (Anderson 1997; Brecx 1993; Corbet 1997; Hoffmann 2001; Lang 1998) reported the Discolouration Index of Lang and Räber. One study (Hase 1998) reported the Shaw and Murray Index and another (Jenkins 1993) a modification of the Shaw and Murray Index. One study (Van



Strydonck 2008) reported the GMSI and one study (Ferretti 1987) used Meckels stain grading method.

One study (Brightman 1991) reported using the Mean Stain Severity Index that was created at the Case Western Reserve University school of dentistry, to record intensity and area of stain. Another study (Graziani 2015) reported dividing the buccal surfaces of the eight central incisors into three areas: incisal, approximal and gingival according to Lobene 1968 and Grundemann 2000 and using a Staining Index (SI) to record the dichotomous presence or absence of staining in each area and to calculate the percentage of the total area showing staining. Three studies (Flemmig 1990; Grossman 1989; Sanz 1994) described their method of measuring tooth staining but did not specify a particular index.

Sixteen of the other included studies described information regarding tooth staining/discolouration/pigmentation reported by study participants (Anauate-Netto 2014; Bhat 2014; Feres 2009; Hase 1995; Pereira 2011); noted by the clinicians during the clinical examination but not measured using an index or scale (de la Rosa 1988; de la Rosa 1988b; Flotra 1972; Grossman 1986; Lopez-Jornet 2012; Turkoglu 2009; Zimmer 2006; Zimmer 2015); or listed as a reason for participant withdrawal from the studies (Banting 1989; Eaton 1997). In one study (Sanz 1989) the presence or absence of tooth staining was assessed from clinical photographs of the mandibular facial anterior teeth. We included information regarding tooth staining from these studies in meta-analyses where possible but were unable to include participant-reported data, data where staining of teeth were reported combined with tongue staining or denture staining or where tooth staining was reported grouped with other adverse effects.

Other adverse effects

Fourteen studies did not report any adverse effects apart from extrinsic staining of teeth or oral tissues alone or both (Brecx 1993; Brightman 1991; Ferretti 1987; Grossman 1989; Hoffmann 2001; Joyston-Bechal 1993; Lopez-Jornet 2012; Overholser 1990; Van Strydonck 2008; Zimmer 2015) or with calculus (Corbet 1997; Flemmig 1990; Grossman 1986; Sanz 1994). Twenty-two studies reported at least one adverse effect relating to the chlorhexidine mouthrinse use apart from extrinsic tooth staining/staining of the oral tissues and calculus. Three studies (Chaves 1994; Feres 2009; Segreto 1986) mentioned a bitter or unacceptable taste associated with the chlorhexidine mouthrinse and no other adverse effects, except for one of the studies (Feres 2009) which also mentioned tooth staining. Adverse effects were not mentioned in six studies (Bajaj 2011; Fine 1985; Jayaprakash 2007; Rahmani 2006; Southern 2006; Weitz 1992). The remaining six studies that did not report any adverse effects or issues with acceptability of the chlorhexidine rinse indicated that there were no oral mucosal lesions (Charles 2004); no severe/serious adverse effects/events (Faveri 2006; Stookey 2005); or adverse reaction/effect (Feres 2012); no irreversible pathology (Lucas 1999); or pathology related to the chlorhexidine mouthrinse use (Taller 1993).

Excluded studies

We excluded 38 studies from the review (see 'Characteristics of excluded studies' table). Below is a summary of the reasons for excluding these studies.

 In seven studies it was not clear whether the studies were randomised and attempts to contact the authors for clarification

- were unsuccessful (Bay 1975; Goutham 2013; Lang 1982; Ousehal 2011; Paknejad 2006; Todkar 2012; Vechis-Bon 1989).
- In four studies the mechanical oral hygiene procedures were not the same in the chlorhexidine rinse and comparator arms (Bouwsma 1992; Brown 2002; Madden 2008; Subhash 1985).
- In 11 studies the comparison was with another active rinse and there was no placebo/control or mechanical oral hygiene only group (Biswas 2014; CTRI 2014; Ernst 1998; Eshwar 2016; Haffajee 2009; McKenzie 1992; NCT01750801; NCT02546804; Persson 1991; Priya 2015; Radafshar 2017).
- Four studies were excluded because the chlorhexidine arm included another active agent that the comparator arms did not have. We would have been unable to separate the effect of the chlorhexidine rinse from the effect of the other active agent (Cortelli 2015; Luoma 1978; Quirynen 2005; Quirynen 2006).
- In one study the chlorhexidine arm received a more intensive intervention including cognitive behavioural education (2 hours) compared to the control arm who received standard care (Persson 1998). In another study (Caton 1993) the chlorhexidine arm received oral hygiene instructions but the control arm did not. The mechanical oral hygiene procedures were potentially not the same in both arms in these two studies.
- One study was unpublished and we were unable to obtain a copy of the study despite following up several lines of enquiry (Segreto 1993).
- In one study, it was not clear from the published report
 whether the study was a randomised trial. Additional
 information obtained from the contact author indicated quasirandomisation (participants were assigned to groups according
 to the order that they came to the University) (Leyes Borrajo
 2002).
- One study was reported in an abstract. The authors were contacted but they did not have access to the data (Saltini 1988).
- One study compared chlorhexidine, zinc and placebo mouthrinses but each rinse arm was further split into those who received a professional prophylaxis prior to the start of the study and those that did not. The study outcomes were not reported in a format that could be used in the review and the authors of the study were unable to provide any additional data (Fischman 1975).
- In one study (Nadkerny 2015) the duration of rinsing was only 15 days and this was not apparent until the full text of the paper was reviewed.
- In three studies we had concerns about the duplication of outcome data in different studies from the same research group (Gupta 2014; Gupta 2015; Karim 2014) and contact with the authors failed to provide an adequate explanation or reassurance. Another study from this research group was excluded due to these concerns (Gupta 2015a).
- One study was excluded because the chlorhexidine mouthrinse was not used as an adjunct to regular mechanical oral hygiene (Yates 2002).

Risk of bias in included studies

Allocation

Random sequence generation

Nineteen studies reported sufficient details of the method of random sequence generation to allow us to assign a judgement



of low risk of bias. The remaining 32 studies either stated that participants were randomised to groups, with no further details, or gave an inadequate description of the method. For these studies, we assigned a judgement of unclear risk of bias.

Allocation concealment

Eleven studies described various methods of allocation concealment which implied that the random sequence was adequately protected and administered as it was generated, thus we assigned a judgement of low risk of bias. Two studies were rated as being at high risk of bias because personnel involved in the studies used a random number table to assign participants to groups, and could have foreseen upcoming assignments (Faveri 2006; Fine 1985). The remaining 38 studies did not report sufficient information on the randomisation process to allow a judgement of either low or high risk of bias, so we assigned a judgement of unclear risk of bias to them.

Overall, seven studies had a rating of low risk of bias for the two above domains, and therefore were at low risk of selection bias. The two studies that had a rating of high risk of bias for allocation concealment were therefore at high risk of selection bias. The remaining 42 studies were at unclear risk of bias.

Blinding

The majority of studies contained a placebo arm with the intention of blinding participants and personnel. However, we considered blinding unlikely to be successful due to the propensity for chlorhexidine to stain the teeth and oral tissues and the association of its use with clinically apparent adverse effects such as mucosal desquamation and mucosal lesions.

Blinding of participants and personnel (performance bias)

Forty-one studies were assigned a judgement of high risk of bias either because staining or adverse events were reported at a higher rate in the chlorhexidine arms, or because a placebo comparison was not used and therefore no blinding was attempted. In these studies, participants could be aware of their allocated intervention and this could have affected their motivation and oral health behaviour. Two studies were assigned a judgement of low risk of bias as there was no difference in staining between the groups, so the blinding appeared to have been maintained (Ferretti 1987; Lopez-Jornet 2012). In the remaining eight studies, blinding was intended through the use of placebo but staining was not reported, so there was insufficient information on which to base a judgement. We rated these studies as being at unclear risk of bias.

Blinding of outcome assessment (detection bias)

Thirty-three studies were assigned a judgement of high risk of bias because the outcomes assessors could have noticed the higher rates of staining or adverse events or both in the chlorhexidine arms. Five studies were rated as having low risk of bias. Two of these studies showed no difference in staining between groups

(Axelsson 1987; Lopez-Jornet 2012). Two studies used masking strategies (whereby a subset of participants were assessed with and without plastic tooth covers and painting of the tongue with food dyes) which revealed that measurements of gingivitis were no different when an examiner thought the participant had received chlorhexidine or not (Grossman 1986; Grossman 1989). One study performed a sensitivity analysis on participants with no visible side effects, which showed no difference to the analysis including all participants, and we judged that bias had not affected the results (Zimmer 2006). We rated the remaining 13 studies at unclear risk of bias because extrinsic tooth staining was not reported. These studies either implied that the outcome assessors were blinded, or did not discuss it, but in any case it was not possible to judge whether or not any blinding could have been successful.

Incomplete outcome data

We did not have any concerns regarding attrition bias in 23 studies, so we rated them as at low risk of bias. Nineteen studies were assigned a judgement of high risk of bias due to either a high overall proportion of attrition, differences in the proportion of attrition between groups, or because the reasons for attrition differed between groups and were linked to the intervention. The remaining nine studies were rated as unclear because the numbers per group at either baseline or evaluation were not reported, thus preventing us from calculating the attrition.

Selective reporting

Fourteen studies adequately reported all expected and planned outcomes, so we rated them as having low risk of bias. The remaining 37 studies were rated at high risk of bias, mainly due to lack of reporting of measures of variance along with the means for each group, and no reporting of tooth staining and other adverse effects.

Other potential sources of bias

Eleven studies were assigned a judgement of low risk of bias. Three studies were assigned a judgement of high risk of bias due to baseline imbalances between groups in important prognostic factors or outcomes of the review or both (Bajaj 2011; Graziani 2015; Sanz 1989). The remaining 37 studies were rated as unclear due to a lack of reporting of details of either calibration of outcome assessment or baseline measurements, meaning that we were unable to assess the potential for differential measurement of outcomes or baseline imbalances respectively.

Overall risk of bias

Fifty studies were at high risk of bias overall as they had at least one domain rated at high risk of bias. One study was rated at unclear risk of bias overall as it had at least one domain rated at unclear risk of bias, but with no domains rated at high risk of bias (Lopez-Jornet 2012). A summary of the risk of bias of the included studies is presented in Figure 2.



Figure 2. Risk of bias summary: review authors' judgements about each risk of bias item for each included study.

	Random sequence generation (selection bias)	Allocation concealment (selection bias)	Blinding of participants and personnel (performance bias)	Blinding of outcome assessment (detection bias)	Incomplete outcome data (attrition bias)	Selective reporting (reporting bias)	Other bias
Anauate-Netto 2014	•	•		•	•	•	•
Anderson 1997	?	?		•		•	?
Axelsson 1987	?	?		•	•	•	?
Daiai 2044		ı					
Bajaj 2011	?	?		?	•	•	•
Bajaj 2011 Banting 1989	?	?	• •	?	•	•	?
	\vdash		• • •	?	•	•	?
Banting 1989	?	?		?	•	----	
Banting 1989 Bhat 2014	?	?	• • • • • • •	? •	•	••••	•
Banting 1989 Bhat 2014 Brecx 1993	?	?		? • •	•	•••••	?



Figure 2. (Continued)

•							
Chaves 1994	•	?	•	?	•	•	?
Corbet 1997	?	?		•			?
de la Rosa 1988	?	?		•	?		?
de la Rosa 1988b	?	?		•	?		?
Eaton 1997	•	?		•			•
Emling 1992	?	?			?	•	?
Ernst 2005	?	•			?	•	?
Faveri 2006	•		?	?	•	•	•
Feres 2009	•	•	?	?	•	•	?
Feres 2012	?	?	?	?	•	•	?
Ferretti 1987	?	?	•	•		•	?
Fine 1985	•		•	•	?		?
Flemmig 1990	?	?	•	•		•	?
Flotra 1972	•	•	•	•			?
Graziani 2015	•	•			•		
Grossman 1986	?	?		•			?
Grossman 1989	?	?		•	?		?
Hase 1995	•	?				•	?
Hase 1998	•	?	•	•	•	•	?
Hoffmann 2001	?	?	•	•	•	•	?
Javaprakash 2007	?		?	?	?		?



Figure 2. (Continued)

•							
Jayaprakash 2007	?	•	?	?	?		?
Jenkins 1993	?	•		•	•	•	?
Jose 2015	?	?		?	•		•
Joyston-Bechal 1993	?	?					•
Lang 1998	•	?				•	?
Lopez-Jornet 2012	?	?	•	•	•	•	?
Lucas 1999	?	?		•	•		?
Navarro 1998	?	?		•	?	•	?
Overholser 1990	•	•		•	•	•	•
Pereira 2011	?	?	•		•		?
Rahmani 2006	?	?	•	?	?		?
Sanz 1989	•	?		•	•		
Sanz 1994	?	?		•	•	•	?
Segreto 1986	?	?	?	?	•		?
Southern 2006	?	?	?	?	•	•	?
Stookey 2005	?	?	?	?			?
Taller 1993	?	?		?	•		?
Turkoglu 2009	?	•				•	•
Van Strydonck 2008	•	?			•	•	•
Weitz 1992	?	?	?	?	•		?
Zimmer 2006	•			•	•		•



Figure 2. (Continued)



Effects of interventions

See: Summary of findings for the main comparison

Chlorhexidine rinse versus placebo/control rinse or no intervention

Gingivitis

Gingival Index of Löe and Silness

At 4 to 6 weeks, eight studies analysing 466 participants (seven at high and one at unclear risk of bias) comparing chlorhexidine rinse with placebo or control rinse and two studies analysing 339 participants (both at high risk of bias) comparing chlorhexidine with no rinse, presented data using the Gingival Index (GI) of Löe and Silness (Löe 1967; Löe and Silness 1963) (Analysis 1.1). The mean GI score across the control groups ranged from 0.17 to 1.4 with a mean of 0.93 and a median of 1. There was no clear difference between the placebo/control and no rinse control subgroups (P = 0.33). The pooled effect estimate of the comparison for all trials showed a reduction in gingivitis in favour of chlorhexidine rinse (mean difference (MD) -0.21, 95% confidence interval (CI) -0.31 to -0.11, P < 0.0001) demonstrating a moderate effect (GI on 0 to 3 increasing scale). There was substantial heterogeneity between the studies (P = 0.001; $I^2 = 67\%$).

There was no clear difference between the different control group subgroups at 6 months (P = 0.11). The overall effect estimate showed a reduction in gingivitis in favour of the chlorhexidine rinse (MD -0.20, 95% CI -0.30 to -0.11, P < 0.0001, 13 high risk of bias studies, 2616 participants) (Analysis 1.2). The mean GI score across the control groups ranged from 0.1 to 1.25 with a mean of 0.8 and a median of 0.7. There was considerable heterogeneity (P < 0.00001; $I^2 = 96\%$).

There were insufficient data to determine the reduction in gingivitis associated with chlorhexidine mouthrinse use in individuals with higher mean GI scores of 1.1 to 3 (moderate or severe levels of gingival inflammation).

Investigation of heterogeneity

We undertook an investigation of the reasons for heterogeneity for the GI scores, at both 4 to 6 weeks and 6 months, for the following subgroups: adults compared to children; baseline gingivitis alone compared to gingivitis and periodontitis at baseline; prophylaxis/scale and polish at baseline compared to none; baseline mean gingivitis score <1 versus scores≥1, by undertaking metaregression in Stata version 14 (Stata 2015) (Additional Table 1; Table 2). The only subgroup analysis that was significant (P = 0.045) was for prophylaxis or not at 6 months, where the trials including a prophylaxis had a lesser effect for the chlorhexidine rinse than those without. This may explain some of the heterogeneity at 6 months.

Chlorhexidine concentration and frequency of rinsing

At 4 to 6 weeks there was no difference between the studies with concentrations of 0.2% and 0.1%/0.12% chlorhexidine, and there was an insufficient number of studies in the concentration subgroups at 6 months to undertake the analysis (Additional Table 3). Participants in one study included in the meta-analysis for 4 to 6 weeks for the GI rinsed once a day, participants in the other studies rinsing twice per day so we were unable to undertake an analysis. Participants in two studies at 6 months rinsed once per day compared with 11 rinsing twice but there was no difference between these subgroups (P = 0.56, Additional Table 4).

Other data

Meta-analyses of studies comparing chlorhexidine rinse with placebo or control rinse after 7 to 12 weeks and > 6 months of rinsing (all at high risk of bias), using the GI of Löe and Silness (Löe 1967; Löe and Silness 1963), showed an effect size similar to the 4 to 6 weeks and 6 month studies (MD -0.47, 95% CI -0.76 to -0.18 and MD -0.50, 95% CI -1.11 to 0.11 respectively). The uncertainty around the effect estimate at > 6 months (two studies of 9 and 24 months duration) is reflected in the 95% CI. Furthermore, there was considerable heterogeneity in both meta-analyses (P < 0.0001, I² = 86%; P < 0.0001, I² = 99% respectively) (Additional Table 5).

One 4-week study (Anauate-Netto 2014) at high risk of bias that was not included in the meta-analysis found no statistically significant differences in Papillary Bleeding Score (Loesche 1979) between the chlorhexidine (0.12%) and placebo group. Three studies (Fine 1985; Hoffmann 2001; Overholser 1990) at high risk of bias, reporting gingival inflammation at 4 to 6 months were not included in the meta-analysis. Two of the three studies found a reduction in the GI (P = 0.043) (Hoffmann 2001) and the modified GI (P < 0.001) (Overholser 1990) for the 0.1% and 0.12% chlorhexidine groups respectively compared to the control rinse (Additional Table 6). Results for 7 to 12 weeks studies not included in meta-analyses are supportive of a reduction in gingival inflammation associated with chlorhexidine mouthrinse use (Additional Table 6).

Gingival bleeding

A variety of different indices were used to measure gingival bleeding, therefore standardised mean differences (SMD) are presented. There was no clear difference between the placebo/control and the no rinse subgroups at either 4 to 6 weeks or 6 months (Analysis 1.3; Analysis 1.4). At 4 to 6 weeks, eight studies analysing 649 participants (all at high risk of bias) comparing chlorhexidine rinse with placebo/control or with no rinse that reported gingival bleeding were combined in a meta-analysis. The overall effect estimate for all studies, both placebo/control and no rinse controls, was moderate at 4 to 6 weeks (SMD -0.56, 95% CI -0.79 to -0.33) showing a reduction in gingival bleeding in favour



of the chlorhexidine rinse (P < 0.00001). There was evidence of moderate heterogeneity (P = 0.16; $l^2 = 34\%$).

At 6 months, eight studies analysing 1132 participants (all at high risk of bias) reported gingival bleeding and were combined in a meta-analysis. The overall effect estimate for all studies was also moderate at 6 months (SMD -0.72, 95% CI -1.02 to -0.42), showing a reduction in gingival bleeding in favour of the chlorhexidine rinse (P < 0.00001). There was evidence of considerable heterogeneity between the studies at 6 months (P < 0.00001; I² = 81%).

Chlorhexidine concentration and frequency of rinsing

No statistically significant differences were found between the subgroups with different chlorhexidine concentrations, at 4 to 6 weeks and 6 months (Additional Table 3). At both 4 to 6 weeks and 6 months all participants in the included studies for gingival bleeding were instructed to rinse twice per day, so no analysis for the effect of frequency of rinsing could be undertaken (Additional Table 4).

Other data

A meta-analysis of two studies (both at high risk of bias) analysing 196 participants comparing chlorhexidine rinse with no rinse controls using the Papillary Bleeding Index (Muhlemann 1977; Saxer 1975) after 8 weeks of rinsing showed no evidence of a difference in gingival bleeding (MD -0.07, 95% CI -0.16 to 0.02) (Additional Table 5).

Another meta-analysis of five studies (all at high risk of bias) analysing 182 participants comparing chlorhexidine rinse with placebo rinse using a variety of different gingival bleeding indices after 7 to 12 weeks of rinsing showed a strong effect in favour of the chlorhexidine rinse (SMD -1.29, 95% CI -1.85 to -0.72, P < 0.00001). There was evidence of substantial heterogeneity (P = 0.02; I^2 = 64%) (Additional Table 5).

Three studies analysing 99 participants (all at high risk of bias) comparing chlorhexidine rinse with placebo rinse, using bleeding on probing, presented longer-term data 1, 4 and 10 months after chlorhexidine mouthrinsing had ceased and were combined in a meta-analysis that showed a small retained effect in favour of chlorhexidine (MD -0.12, 95% CI -0.20 to -0.04) (Additional Table 5).

Three studies (Axelsson 1987; Graziani 2015; Sanz 1989) at high risk of bias reporting gingival bleeding at 4 to 6 weeks were not included in the meta-analysis (Additional Table 7). All three studies reported a reduction in gingival bleeding for 0.1%, 0.12% and 0.2% chlorhexidine rinses compared to placebo or control rinse.

Four studies (Banting 1989; Charles 2004; Lucas 1999; Sanz 1994) at high risk of bias reporting gingival bleeding at 6 months were not included in the meta-analysis (Additional Table 7). Three of these studies reported a reduction in gingival bleeding for 0.12% chlorhexidine rinse compared to placebo or control rinse. Banting 1989 demonstrated a reduction in gingival bleeding at 2 years for 0.12% chlorhexidine compared to placebo (P < 0.0001). Results for 7 to 12 weeks studies not included in meta-analyses are supportive of a reduction in gingival bleeding associated with chlorhexidine mouthrinse use (Additional Table 7).

Plaque

A variety of different indices were used to measure plaque, therefore SMDs are presented. There was no difference between the

placebo/control or no rinse control subgroups for plaque at 4 to 6 weeks (P = 0.93) (Analysis 1.5). The overall effect was based on 12 trials analysing 950 participants, 11 at high and one at unclear risk of bias, and showed a large effect in favour of chlorhexidine rinse (SMD -1.45, 95% CI -1.90 to -1.00). Four of these trials analysing 223 participants reported the Plaque Index (0 to 3 increasing scale) (Silness and Löe 1964) and showed a reduction in plaque in favour of chlorhexidine rinse (MD -0.58, 95% CI -0.78 to -0.39) (Analysis 1.6); five of these trials analysing 546 participants reported the Turesky modification of the Quigley Hein Index (0 to 5 increasing scale) (Turesky 1970) and also showed a reduction in plaque in favour of chlorhexidine rinse (MD -0.78, 95% CI -0.85 to -0.70) (Analysis 1.7).

At 6 months, although both subgroups showed a reduction in plaque favouring the chlorhexidine rinse (Analysis 1.8), there was a difference between the size of the effect estimates for the placebo/ control and no rinse subgroups (P = 0.02). The higher estimate for the chlorhexidine rinse compared to placebo/control rinse based on nine trials analysing 1933 participants (all at high risk of bias) with a large effect (SMD -1.59, 95% CI -1.89 to -1.29) is compared with a moderate effect for the chlorhexidine rinse compared to no rinse control groups based on 2 trials analysing 142 participants (both at high risk of bias) (SMD -0.68, 95% CI -1.35 to -0.01). There was substantial heterogeneity among the studies in each subgroup $(P < 0.00001; I^2 = 84\%; P = 0.06; I^2 = 72\% \text{ for placebo/control})$ and no rinse subgroups respectively). Five studies analysing 1108 participants reported data for the Plaque Index at 6 months. There was no clear difference between the placebo/control and no rinse subgroups, and the overall effect was -0.62 (95% CI -1.12 to -0.12) (Analysis 1.9). Six trials, all with placebo/no treatment controls, analysing 967 participants reported the Turesky modification of the Quigley Hein Index at 6 months and also showed a reduction in plaque in favour of chlorhexidine rinse (MD -0.73, 95% CI -0.88 to -0.57) (Analysis 1.10).

Chlorhexidine concentration and frequency of rinsing

No differences were found between the subgroups with different chlorhexidine concentrations (Additional Table 3). At 4 to 6 weeks the participants in one trial were instructed to rinse once per day compared with twice a day in the other 11 trials so no analysis for the effect of frequency of rinsing could be undertaken. Similarly at 6 months the participants in one trial were instructed to rinse once per day compared with twice a day in the other 10 trials so no analysis for the effect of frequency of rinsing could be undertaken (Additional Table 4).

Other data

Meta-analyses of studies that compared chlorhexidine mouthrinse with placebo (10 studies all at high risk of bias analysing 423 participants) or no rinse control (two studies both at high risk of bias analysing 196 participants) at 7 to 12 weeks showed a reduction in plaque (using various indices) in favour of chlorhexidine rinse (SMD -1.74, 95% CI -2.51 to -0.98 and SMD -0.77, 95% CI -1.07 to -0.47 respectively, Additional Table 5) both indicating a strong effect. There was considerable heterogeneity among the 10 placebo-controlled studies P < 0.00001; $I^2 = 91\%$.

One large 9-month study (at high risk of bias) (Bajaj 2011) analysing 852 participants comparing 0.1% chlorhexidine rinse with distilled water found a large reduction in plaque (0 to 3 increasing scale) in the chlorhexidine rinse group (MD -1.55, 95% CI -1.79 to -1.31, Additional Table 5).



Four studies analysing 132 participants (all at high risk of bias) comparing chlorhexidine rinse with placebo rinse presented data 1, 4 and 10 months after rinsing had ceased, using a variety of indices, and were combined in a meta-analysis that showed a large retained effect in favour of chlorhexidine (SMD -1.10, 95% CI -1.18 to -0.40, Additional Table 5). However, there was substantial heterogeneity between the studies (P = 0.02; P = 1.00).

One study (Sanz 1989) at high risk of bias reporting a plaque outcome at 6 weeks that was not included in the meta-analysis showed a reduction in plaque for 0.12% chlorhexidine rinse compared to placebo rinse (Additional Table 8).

Five studies at high risk of bias reporting a plaque outcome at 6 months that were not included in the meta-analysis also showed a reduction in plaque for 0.12%, 0.1%, 0.05% and 0.06% chlorhexidine rinse compared to placebo or control rinse (Banting 1989; Hoffmann 2001; Jayaprakash 2007; Lucas 1999; Sanz 1994) (Additional Table 8).

Banting 1989, which was not included in the meta-analysis demonstrated a reduction in plaque at 2 years for 0.12% chlorhexidine compared to placebo. Results for 7 to 12 weeks studies not included in meta-analyses were generally supportive of a reduction in plaque associated with chlorhexidine mouthrinse use (Additional Table 8).

Adverse effects

Calculus

Two studies analysing 102 participants, both at high risk of bias, and with placebo/control rinses, presented data on calculus at 4 to 6 weeks. There was insufficient evidence to determine whether using chlorhexidine rinse for 4 to 6 weeks was associated with increased calculus formation (SMD 0.02, 95% CI -0.09 to 0.14) (Analysis 1.11).

At 7 to 12 weeks, five studies (all high risk of bias) analysing 330 participants, with a placebo/control rinse presented calculus data, along with one study analysing 95 participants with a no rinse control also at high risk of bias (Analysis 1.12). There was a difference between the subgroups for control type, with there being insufficient evidence to determine an effect on calculus for chlorhexidine rinse compared to placebo/control rinse (SMD 0.14, 95% CI -0.08 to 0.36). The single study in the no rinse control group, however, showed more calculus in the chlorhexidine rinse group compared to the no rinse control group (SMD 1.02, 95% CI 0.59 to 1.45) demonstrating a large effect. There was evidence of substantial heterogeneity (P = 0.004; $I^2 = 71\%$).

At 6 months there was a difference between the subgroups for control type. There was only one study (at high risk of bias), analysing 91 participants, with a no rinse control group which showed an increase in calculus in the chlorhexidine rinse group (SMD 1.39, 95% CI 0.93 to 1.85) demonstrating a large effect (Analysis 1.13). The pooled result for the three studies (at high risk of bias) analysing 232 participants with placebo/control rinses showed an increase in calculus associated with chlorhexidine rinse (SMD 0.60, 95% CI 0.24 to 0.96) demonstrating a moderate effect. There was evidence of moderate heterogeneity (P = 0.16; I^2 = 46%).

Chlorhexidine concentration and frequency of rinsing

No statistically significant difference was found between the subgroups with different chlorhexidine concentrations at 7 to 12

weeks, however the data at 6 months indicated a larger effect (increased calculus formation in the chlorhexidine rinse group) for the lower concentration chlorhexidine mouthrinse (Additional Table 3). Participants in all the included trials with calculus data at 4 to 6, 7 to 12 weeks and 6 months were asked to use the mouthrinse twice a day, so no analysis of frequency could be undertaken (Additional Table 4).

Other data

Five studies at high risk of bias reporting interim and end scores for calculus at various time points from 4 weeks to 2 years (Anderson 1997; Banting 1989; Charles 2004; Grossman 1986; Sanz 1994) that were not included in the meta-analysis all found increased supragingival calculus in the 0.12% chlorhexidine rinse groups compared to the placebo and control rinse groups (Additional Table 9). Increased calculus was noted in two other studies but not measured using an index or scale (de la Rosa 1988; de la Rosa 1988b).

Extrinsic tooth staining

Different dichotomous and continuous indices were used to measure tooth staining. At 4 to 6 weeks both the dichotomous data (two studies at high risk of bias, 156 participants) and the continuous data (eight studies at high risk of bias, 415 participants) showed more tooth staining in the chlorhexidine rinse group. A large effect estimate was found for both the dichotomous data (risk ratio (RR) 5.41, 95% CI 2.03 to 14.47, Analysis 1.14) and continuous data (SMD 1.07, 95% CI 0.80 to 1.34, Analysis 1.16). There was no strong evidence of heterogeneity between the trials reporting continuous data (P = 0.14, P = 0.14, P = 0.14).

Only one study, analysing 118 participants, reported dichotomous data for tooth staining at 7 to 12 weeks, with more tooth staining in the chlorhexidine group (RR 2.50, 95% CI 1.29 to 4.83) (Analysis 1.15). Eleven studies, analysing 581 participants (at high risk of bias) reported data on different continuous scales showing an increase in tooth staining associated with chlorhexidine rinse at 7 to 12 weeks (SMD 1.19, 95% CI 0.98 to 1.40), demonstrating a large effect (Analysis 1.17).

Four studies analysing 323 participants, at high risk of bias, reported continuous data at 6 months using different tooth staining indices. The pooled effect estimate indicated a large effect for tooth staining due to the chlorhexidine rinse (SMD 1.54, 95% CI 1.22 to 1.86) (Analysis 1.18). There was no strong evidence of heterogeneity (P = 0.19; $I^2 = 36\%$).

Chlorhexidine concentration and frequency of rinsing

Subgroup comparisons for chlorhexidine concentration were not significant (Additional Table 3). In one study providing tooth staining data at both 4 to 6 weeks and 7 to 12 weeks, participants rinsed three times a day, and in one of the studies at 4 to 6 weeks participants rinsed only once per day, whereas in all the other studies participants rinsed twice per day (Additional Table 4). There were insufficient studies in the subgroups to make a meaningful comparison on frequency at any time point.

Other data

Thirteen studies reported information about tooth staining that could not be included in meta-analyses. Eight studies that reported data on extrinsic tooth staining could not be included in meta-



analyses because no measure of variance was reported (Charles 2004; Flotra 1972; Grossman 1989; Hoffmann 2001; Sanz 1994), no outcome data were reported (Grossman 1986) or because the mean value in the control groups was zero (Axelsson 1987; Graziani 2015). All of these studies reported higher levels of extrinsic tooth staining in the chlorhexidine compared to control arms at 4 to 6 weeks (Axelsson 1987; Flotra 1972; Graziani 2015); 7 to 12 weeks (Charles 2004; Grossman 1989); and at 6 months (Charles 2004; Grossman 1986; Grossman 1989; Hoffmann 2001; Sanz 1994). In five other studies tooth staining was reported combined with either denture staining (Lopez-Jornet 2012), discolouration of oral tissues (Hase 1995; Turkoglu 2009; Zimmer 2006) or was reported as 'discolouration' and it was not clear whether this related to teeth, oral tissues or both (Bhat 2014). In four of these five studies staining/discolouration was higher in the chlorhexidine rinse groups at 4 to 6 weeks (Bhat 2014; Hase 1995; Turkoglu 2009; Zimmer 2006) and 7 to 12 weeks (Zimmer 2006) (Additional Table

In five additional studies, increased tooth staining (de la Rosa 1988; de la Rosa 1988b; Feres 2009), 'pigmentation' (Pereira 2011) and 'yellow teeth' (Anauate-Netto 2014) were reported in relation to the chlorhexidine rinse arms. In two studies staining of teeth in the chlorhexidine rinse arm was listed as a reason for participant withdrawal from the studies (Banting 1989; Eaton 1997).

Other adverse effects

Twenty-two studies reported at least one adverse effect apart from extrinsic tooth staining and calculus formation in the chlorhexidine rinse arms. Apart from extrinsic tooth staining, the adverse effects most commonly reported in the chlorhexidine rinse arms of the included studies were taste disturbance/alteration (reported in 11 studies), effects on the oral mucosa (reported in 13 studies) and a general burning sensation or a burning tongue or both (reported in nine studies).

Taste disturbance/alteration was reported in 11 studies (Anauate-Netto 2014; Bhat 2014; Ernst 2005; Graziani 2015; Hase 1998; Jenkins 1993; Jose 2015; Lang 1998; Pereira 2011; Turkoglu 2009; Zimmer 2006). The proportion of participants experiencing taste disturbance/alteration was reported in eight of these studies and was consistently higher in the chlorhexidine rinse arms (3% to 30%) than in the placebo/control/oral hygiene (OH) only arms (0% to 7%) with one exception (Bhat 2014) where a higher proportion of participants in the saline control arm experienced taste disturbances (45.4% versus 63.6%).

A general burning sensation (Anderson 1997; Bhat 2014; Emling 1992; Hase 1995; Sanz 1989; Zimmer 2006) and/or a burning tongue (Anauate-Netto 2014; Jose 2015; Lang 1998) was reported in the chlorhexidine rinse arms in nine studies. The proportion of participants affected was reported in four of the nine studies (Anderson 1997; Bhat 2014; Jose 2015; Zimmer 2006) and ranged from 4.6% to 50% in the chlorhexidine arms compared to 0% in the placebo/control/OH only arms. Hase 1995 reported 'burning sensation' rated by participants on a visual analogue scale from none (0) to 100 (very much). The mean visual analogue scale score for the chlorhexidine rinse arm was approximately 18 compared to 8 for the placebo arm.

Effects on the oral mucosa of participants in the chlorhexidine rinse arms were reported in 13 studies. Specific effects were described as

mucosal irritation (Anauate-Netto 2014; Emling 1992), sensitivities of the oral mucosa (Ernst 2005), mucosal soreness (Jenkins 1993), mild desquamation (de la Rosa 1988; de la Rosa 1988b; Navarro 1998), mucosal ulceration (Hase 1995; Turkoglu 2009), mucosal erosions (Jenkins 1993), soft tissue conditions (Banting 1989), oral mucosal lesions (Axelsson 1987; Flotra 1972) and adverse mucosal reaction (Eaton 1997). Where it was reported, the proportion of participants experiencing effects on the oral mucosa was generally low. For example, clinically apparent mucosal ulceration was reported in 12% (n=3) of participants in the chlorhexidine rinse arm of one study (Turkoglu 2009) and mucosal erosions affected 4% (n=2) of participants in the chlorhexidine rinse arm of another study (Jenkins 1993). However one study reported a high proportion of oral mucosal lesions in participants in the chlorhexidine (0.2%) rinse arm (25%, n=6) (Axelsson 1987).

Discolouration/pigmentation of the oral soft tissues was reported in seven studies and was consistently higher in the chlorhexidine rinse arms ranging from 9% to 88% compared to 0% to 7% in the placebo/control/OH only arms (Graziani 2015; Pereira 2011; Hase 1995; Hase 1998; Lang 1998; Turkoglu 2009; Zimmer 2006). Coated tongue was reported as an adverse effect in one study (Jose 2015).

Less commonly reported adverse effects included transient anaesthetic sensation, hypoaesthesia and paraesthesia; affecting 5% to 18% of participants in the chlorhexidine rinse arms (Hase 1995; Hase 1998; Jenkins 1993; Jose 2015; Lang 1998; Zimmer 2006).

Lastly, in relation to acceptability, an unacceptable or unpleasant taste was reported by participants in the chlorhexidine rinse arms of eight studies (Anauate-Netto 2014; Chaves 1994; Eaton 1997; Feres 2009; Hase 1995; Jenkins 1993; Sanz 1989; Segreto 1986).

Sensitivity analysis

We undertook a sensitivity analysis for the GI at both 4 to 6 weeks and 6 months, excluding trials where we had to estimate the standard deviation from similar studies, or had to read the data from graphs. The pooled estimate for 4 to 6 weeks (excluding Jenkins 1993 and Sanz 1989) was: MD -0.23 (95% CI -0.36 to -0.10), and at 6 months (excluding Banting 1989; Chaves 1994; Jayaprakash 2007; Lucas 1999; Sanz 1994) MD -0.25 (95% CI -0.39 to -0.10). These estimates are similar to those for the overall pooling of the studies.

We undertook a sensitivity analysis for the GI at both 4 to 6 weeks and 6 months excluding trials where professional periodontal treatment was carried out during the study (i.e. Faveri 2006; Feres 2009; Feres 2012; Flotra 1972; Rahmani 2006; Sanz 1989). The pooled estimate for 4 to 6 weeks (excluding Rahmani 2006 and Sanz 1989) was: MD -0.21 (95% CI -0.32 to -0.10), and at 6 months (excluding Flotra 1972) MD -0.19 (95% CI -0.29 to -0.10). These estimates are similar to those for the overall pooling of the studies.

Assessment of reporting biases

The meta-analysis for GI at 4 to 6 weeks included 10 trials. Testing for asymmetry in a funnel plot using the methods described by Egger 1997, using "metabias" command in Stata (P = 0.019), although significant, the estimate of bias was negative indicating that the bias was lack of large studies (not small studies). The meta-analysis for GI at 6 months included 13 trials, and there was no evidence of publication bias from the funnel plot and this was



confirmed by testing for asymmetry in a funnel plot using the same methods (P = 0.95). Two of the plaque meta-analyses at 4 to 6 weeks and 6 months had more than 10 studies but funnel plots cannot be investigated when SMD models are used. All the other meta-analyses had less than 10 trials, so we were unable to assess reporting bias for these.

DISCUSSION

Summary of main results

There was high-quality evidence that using chlorhexidine mouthrinse as an adjunct to regular mechanical oral hygiene procedures for 4 to 6 weeks and 6 months results in a moderate reduction in gingivitis compared to placebo, control or mechanical oral hygiene alone. This reduction in gingivitis, in individuals with mild gingival inflammation on average (mean score of 1 on the 0 to 3 Gingival Index (GI) scale) is not considered to be clinically important. Heterogeneity was considerable in the meta-analyses at 4 to 6 weeks and 6 months and subgroup analysis was only able to explain some of the heterogeneity at 6 months, due to whether or not a prophylaxis or scale and polish was conducted at the start of the study. The standardised mean differences (SMDs) for gingival bleeding at 4 to 6 weeks and 6 months also indicate that chlorhexidine mouthrinse has a moderate effect on gingival bleeding compared to placebo, control or mechanical oral hygiene alone. There was insufficient evidence to determine the reduction in gingivitis associated with chlorhexidine mouthrinse use in individuals with mean GI scores of 1.1 to 3 indicating moderate or severe levels of gingival inflammation.

There was also high-quality evidence that chlorhexidine mouthrinse reduces plaque at 4 to 6 weeks and 6 months. SMDs for plaque for both control subgroups at 4 to 6 weeks and at 6 months indicate a large effect. The four studies that used the Plaque Index (0 to 3 increasing scale) and the four studies that used the Turesky Modification of the Quigley and Hein Index (0 to 5 increasing scale) at 4 to 6 weeks indicate a reduction in plaque in favour of the chlorhexidine mouthrinse.

There was no evidence that one concentration of chlorhexidine rinse was more effective than another in reducing gingivitis and plaque. Participants in most of the studies included in the meta-analyses used a rinse frequency of twice daily, limiting our ability to analyse the effect of rinse frequency on gingivitis and plaque. We were only able to evaluate the effect of rinse frequency for gingivitis measured at 6 months using the GI, however, no difference in effect was observed.

There was moderate-quality evidence that chlorhexidine rinse causes a large amount of extrinsic tooth staining when used for periods of 4 weeks or longer. Comparison of subgroups with different chlorhexidine concentrations was not statistically significant.

There was insufficient evidence to determine whether using chlorhexidine rinse for 1 to 3 months was associated with increased calculus formation. There was some evidence of increased calculus formation related to rinsing with chlorhexidine mouthrinse for a period of 6 months. However, overall evidence for the effect of chlorhexidine mouthrinse on calculus formation was inconclusive.

Twenty-two of the 51 studies included in this review reported at least one adverse effect apart from extrinsic tooth staining and calculus formation in the chlorhexidine rinse arms. The adverse effects most commonly reported in the chlorhexidine rinse arms of the included studies were taste disturbance/alteration (reported in 11 studies), effects on the oral mucosa including mucosal irritation, soreness, mild desquamation, mucosal ulceration/erosions and oral mucosal lesions (reported in 13 studies) and a general burning sensation or a burning tongue or both (reported in nine studies).

Overall completeness and applicability of evidence

Chlorhexidine mouthrinse is effective in reducing gingivitis and plaque when used as an adjunct to regular mechanical oral hygiene procedures for 4 weeks or longer. The consistency of effect for chlorhexidine mouthrinse on gingivitis and plaque across studies included in this review was remarkable though not surprising considering that chlorhexidine mouthrinse has long been regarded as the gold standard of antimicrobial mouthrinses. The primary outcome of this review was gingivitis which can be measured using a variety of different indices. The clinical features of gingivitis can be assessed non-invasively by visual assessment of colour, contour and gingival bleeding or invasively using an instrument to provoke bleeding or both (Lorenz 2009). Our review clearly demonstrates the many different indices that can be used to measure gingivitis. While these many indices were developed to provide a standardised reproducible method of quantifying gingivitis for descriptive or comparative purposes in epidemiology and clinical trials respectively, the subjectivity involved in their use remains problematic. The presence/absence of bleeding on provocation would appear to be the more objective approach (Ciancio 1986; Lorenz 2009; Polson and Caton 1985). However, the probing technique, the depth to which the instrument is inserted, the force with which it is applied, the time allowed to elapse between applying the stimulus and recording the outcome and even the angulation of the instrument can have an effect on the bleeding response (Lorenz 2009; Polson and Caton 1985; van der Weijden 1994). Moreover, calibration of examiners and demonstrating inter- and intraexaminer reliability are hampered by the inability to obtain the same response upon repeated application of the stimulus to provoke gingival bleeding at the same site (Lorenz 2009; Polson and Caton 1985). In synthesising the evidence from primary research studies evaluating chlorhexidine mouthrinse as an adjunctive treatment for gingival health, the many different approaches to measuring gingivitis, particularly gingival bleeding, presented difficulties in combining the study results to produce a summary of the effect of the mouthrinse.

To facilitate the analysis, we grouped the indices into those that measured principally gingival inflammation visually and invasively (e.g. the Gingival Index (Löe 1967; Löe and Silness 1963)) or visually only (e.g. the Modified Gingival Index (Lobene 1986)) and those that measured gingival bleeding (e.g. bleeding on probing (BOP) (Ainamo and Bay 1975) or the Eastman Interdental Bleeding Index (Caton and Polson 1985)). We endeavoured to include both types of index for measuring gingivitis in the review, therefore where both were reported within a study, both were data extracted. Our main analysis was based on the studies that reported gingival inflammation using the Gingival Index (Löe 1967; Löe and Silness 1963) at 4 to 6 weeks and 6 months to reflect both the short- and long-term effect of chlorhexidine mouthrinse. Our results showed that after 4 to 6 weeks of use, chlorhexidine mouthrinse reduced



gingivitis by a mean of 0.21 of a unit on the Gingival Index (GI) scale when compared to a placebo/control mouthrinse or mechanical oral hygiene alone. Based on these studies, a reduction in gingivitis of 0.11 to 0.31 of a unit on the GI scale could be expected with use of chlorhexidine mouthrinse for a period of 4 to 6 weeks in individuals with mild gingival inflammation on average (mean score of 1 on the 0 to 3 GI scale). A similar mean effect size of 0.20 of a unit reduction on the GI scale was found for gingivitis at 6 months also in individuals with mild gingival inflammation on average.

The GI is measured on a 0 to 3 increasing scale with 0 indicating normal gingivae. However, each one unit increase on the scale does not indicate the same proportionate deterioration in gingival condition and a low mean GI score could mask the presence of a small number of sites with moderate or severe levels of gingival inflammation. With this in mind, depending on the average gingivitis level in an individual, a one unit reduction could indicate a shift from severe inflammation and a tendency for spontaneous bleeding to moderate inflammation and BOP or from moderate inflammation and BOP to mild inflammation but no BOP. As previously mentioned, the participants who contributed data to the meta-analyses of the GI at 4 to 6 weeks had a low mean gingival index score at baseline. The clinical relevance of a mean reduction of one fifth of a unit on the GI scale is therefore not clear. Our results for gingival bleeding indicate that using chlorhexidine mouthrinse for 4 to 6 weeks or 6 months has a moderate effect on gingival bleeding. Although these meta-analyses included some studies where participants appeared to have a high proportion of sites with BOP at baseline, the variety of different gingival bleeding indices employed meant that we were unable to determine the mean reduction in gingivitis associated with chlorhexidine mouthrinse use in individuals with moderate or severe levels of gingival inflammation. When the SMDs for gingival bleeding were expressed in the units of the bleeding aspect of the GI (GI \geq 2) the effect size was equivalent to a 7% reduction in the percentage of sites with BOP at 4 to 6 weeks and a 4% reduction in the percentage of sites with BOP at 6 months. Use of chlorhexidine mouthrinse was associated with a large reduction in plaque at 4 to 6 weeks and 6 months. The mean reduction in plaque associated with chlorhexidine mouthrinse use at 4 to 6 weeks was equivalent to approximately half a unit on the 0 to 3 Plaque Index scale (Silness and Löe 1964) or three quarters of a unit on the 0 to 5 Turesky modification of the Quigley and Hein Index (TQH) scale (Turesky 1970).

Statistical heterogeneity was evident in many of our metaanalyses and was likely due to variation in the characteristics of the individual included studies. Investigation of the observed heterogeneity was conducted for our main outcome, measured using the GI at 4 to 6 weeks and 6 months using prespecified characteristics. There was a decrease in the effect estimate for the GI at 6 months when a prophylaxis or scale and polish was conducted at the start of the study compared to studies where no prophylaxis or scale and polish was undertaken. This is not consistent with our knowledge of the mechanism of action of chlorhexidine mouthrinse which would appear to be more effective in preventing plaque accumulation when applied to a clean tooth surface than reducing pre-existing plaque (Eley 1999). Nonetheless, the results indicate that some of the heterogeneity for the GI at 6 months was due to whether or not a prophylaxis or scale and polish was conducted at the beginning of the study. We have presented all of the meta-analyses despite the high heterogeneity due to the remarkable consistency of the direction of effect across the different analyses.

We included studies that evaluated chlorhexidine mouthrinse use in individuals with gingivitis only or gingivitis and periodontitis. Where it was reported, the level of gingivitis in the included studies at baseline was generally low. The majority of the studies that evaluated chlorhexidine mouthrinse use in individuals with periodontitis evaluated its effect as an adjunct to surgical or nonsurgical periodontal treatment. In a small number of the studies that evaluated chlorhexidine mouthrinse use in individuals with periodontitis, it was difficult to ascertain whether the professional tooth cleaning provided was appropriate for the level of disease present. For example, in two of these studies it was reported that professional cleaning was undertaken at baseline but no further details were provided (Chaves 1994; Ernst 2005). In another study where 17% to 50% of participants in the study groups were diagnosed with periodontitis, supragingival scaling and polishing only was provided at baseline (Graziani 2015). In one study a proportion of the participants clearly had periodontitis and abundant calculus, however no professional treatment or oral hygiene instructions were provided to the participants (Corbet 1997). The inclusion of these studies could be criticised because antimicrobials such as chlorhexidine have limited capacity to penetrate subgingivally and are therefore suggested for use as an adjunct to mechanical debridement rather than as an alternative therapy (Greenstein 1986; Ohrn 2009). Furthermore, self-medication with an effective antimicrobial agent such as chlorhexidine mouthrinse "may mask more serious underlying periodontal disease" (Tonetti 2015). It is well recognised that control of supragingival plaque alone will not be sufficient to treat periodontitis (Eley 1999; Hull 1980), disruption of the subgingival plaque by mechanical debridement will also be required (Kornman 1986; Ohrn 2009).

We would have liked to investigate how the effect of chlorhexidine mouthrinse on gingival health varies depending on the level of underlying periodontal disease. However, the periodontal status of the participants in the included studies often was not clearly reported at baseline. In many cases, the inclusion criteria allowed for individuals to have pocket probing depths and clinical attachment loss consistent with periodontitis but the periodontal status of the people actually enrolled in the study was not described. Furthermore, where participants were advised to continue their regular oral hygiene practices during the study, no information about their frequency of toothbrushing or their use of interdental cleaning aids was reported. Therefore, our ability to explore the effect of the adjunctive use of chlorhexidine mouthrinse on gingivitis in individuals with different levels of periodontal disease at baseline and in individuals with treated and untreated periodontitis was hampered both by the small number of studies available at each time point and by poor reporting of the relevant information in the included studies.

We were able to conduct a metaregression analysis of 4 to 6 week and 6-month studies where participants had gingivitis only compared to studies where participants had gingivitis and periodontitis (both treated and untreated during the study). The results did not disclose any statistically significant difference in the effect of the chlorhexidine mouthrinse on the gingival index at either time point, however only eight studies were included in the metaregression at 4 to 6 weeks and only nine studies were included



in the metaregression at 6 months. We cannot specifically comment on the effect of chlorhexidine used adjunctively during periodontal treatment on periodontal outcomes such as pocket probing depth and clinical attachment loss because the focus of this review was gingivitis and therefore periodontal outcomes were outside the scope of the review.

Retention of chlorhexidine in the oral cavity is key to its effectiveness as an antimicrobial agent (Bonesvoll 1974; Bonesvoll 1974a; Bonesvoll 1978; Jenkins 1988; Rolla 1971) and is dependant on a number of different factors including concentration and duration of application (Tomás 2010). In relation to concentration, the effect of chlorhexidine on plaque is dose-dependant and a similar effect can be achieved with larger volumes of lower concentration solutions providing the optimal dose of approximately 20 mg twice a day is delivered (Bonesvoll 1978). All but four of the included studies evaluated either 0.2%, 0.12% or 0.1% chlorhexidine mouthrinse and only two of these studies provided data for 0.2% compared to 0.1% (Axelsson 1987) or 0.12% (Segreto 1986) chlorhexidine mouthrinse within the same study. Most of the studies that evaluated 0.2% chlorhexidine mouthrinse employed a rinse volume of 10 mL twice a day while most of the studies that evaluated 0.12% or 0.1% chlorhexidine mouthrinse employed a rinse volume of 15 mL twice a day. Therefore regardless of the concentration of chlorhexidine mouthrinse used, the dose of chlorhexidine was effectively the same in the majority of the studies and corresponded to the optimal dose of 18 mg to 20 mg twice a day. It has been demonstrated that increasing the duration of rinsing with 0.12% or 0.2% chlorhexidine mouthrinse from 30 to 60 seconds results in greater substantivity of the rinse (Tomás 2010). The longer duration of rinsing employed by most of the studies in this review that evaluated 0.2% chlorhexidine mouthrinse could be expected to translate into greater retention of chlorhexidine in these studies and hence a greater effect on gingivitis and plaque. However, 75% of the chlorhexidine that is retained after rinsing for 60 seconds with 0.2% chlorhexidine rinse is retained after the first 30 seconds of rinsing (Bonesvoll 1974) and a rinse duration of 30 seconds for 0.12% chlorhexidine mouthrinse has been shown to be both effective and acceptable (Keijser 2003). Therefore, most of the studies included in this review used the optimal dose of chlorhexidine for at least 30 seconds which may partly explain why we did not find any difference in effect between the different concentration rinses. The two included studies that provided data for a direct comparison between 0.2% and 0.1%/0.12% chlorhexidine rinse found no difference in effect between the different concentration rinses used with the same volume and duration of rinsing (Axelsson 1987; Segreto 1986).

Studies that evaluated the effectiveness of chlorhexidine mouthrinse on gingivitis using a cross-over study design were excluded from this review. This was due to concerns that the duration of the washout phase between the trial periods might not be sufficient to rule out a carry-over effect due to chlorhexidine mouthrinse. Carry-over can threaten the internal validity of a cross-over study and is of particular concern in studies involving chlorhexidine mouthrinse due to the established substantivity or prolonged effect of chlorhexidine mouthrinse. Evidence-based guidance on the appropriate duration of washout phase is absent from the literature and the washout periods commonly used appear to range from 2 days to 3 weeks (Berchier 2010; Newcombe 1995). The very limited data we have in this review regarding gingivitis and plaque outcomes recorded after cessation

of chlorhexidine mouthrinsing suggest a residual effect on these outcomes weeks or months after rinsing with chlorhexidine has ceased. Although no firm conclusions can be drawn regarding the appropriateness of using a cross-over study design when the study includes a chlorhexidine mouthrinse arm, the uncertainty surrounding carry-over in relation to chlorhexidine mouthrinse warrants further investigation.

Low concentration (0.05%/0.06%) chlorhexidine mouthrinses are currently marketed for long-term daily use. Side-effects associated with chlorhexidine mouthrinse use are dose-dependant and tend to decrease when the concentration is lowered (Cumming and Löe 1973; Flotra 1971). Therefore in lowering the concentration of chlorhexidine, it is hoped that side effects will be reduced while maintaining the effect on gingivitis and plaque (Keijser 2003). Only five studies that evaluated chlorhexidine concentrations less than 0.1% met the inclusion criteria for this review. Two studies evaluated 0.05% (Jayaprakash 2007; Joyston-Bechal 1993) and three studies evaluated 0.06% chlorhexidine mouthrinse (Hoffmann 2001; Zimmer 2006; Zimmer 2015). Although all of these studies demonstrated a reduction in plaque, the effect on gingivitis was not clear. This might be explained by the lower concentration rinses having a mainly bacteriostatic action that may produce an effect below the threshold required to produce a corresponding reduction in gingivitis. Despite the reduced concentration of chlorhexidine, extrinsic tooth staining occurred in all four studies of low concentration chlorhexidine mouthrinse that reported a staining outcome (Hoffmann 2001; Joyston-Bechal 1993; Zimmer 2006; Zimmer 2015). There has been some debate about the mechanism by which chlorhexidine mouthrinse produces extrinsic tooth staining. However, it is currently understood to be a result of precipitation of anionic dietary chromogens onto adsorbed chlorhexidine cations (Watts 2001) and thus the mechanism by which chlorhexidine mouthrinse produces extrinsic tooth staining appears to be closely linked to its mechanism of action. Therefore lack of staining is likely to indicate lack of effectiveness (Addy 2005; Eley 1999). Some degree of tooth staining can therefore be expected to feature even with low chlorhexidine concentration rinses. However, with the exception of Zimmer 2006, there were no reports of other adverse effects relating to the low concentration chlorhexidine mouthrinse apart from staining of the teeth and oral

Use of chlorhexidine mouthrinse is associated with a number of adverse effects, the most common being extrinsic staining of the teeth and oral soft tissues (Addy 1986). In many of the studies included in this review, adverse effects were inadequately reported or not reported at all. It is important that adverse effects due to chlorhexidine mouthrinse use are reported alongside evidence for effectiveness with enough detail to allow health professionals and the public to make an informed decision about recommending, prescribing or using the mouthrinse.

From our included studies it was clear that rinsing with any concentration of chlorhexidine mouthrinse for 4 weeks or longer was associated with an increase in extrinsic tooth staining. A number of the included studies reported higher levels of calculus in the chlorhexidine rinse arms but the meta-analyses were inconclusive. Other adverse effects such as taste disturbance/ alteration, burning sensation and pigmentation of the oral soft tissues were also reported in a number of included studies. Effects on the oral mucosa such as mucosal ulceration, lesions



or erosions and a transient anaesthetic sensation, hypoaesthesia and paraesthesia were also reported in a number of included studies and can be considered more severe but the proportion of participants experiencing these adverse effects in any study tended to be low. While extrinsic tooth staining and calculus build-up due to chlorhexidine mouthrinse use must be removed by professional tooth cleaning incurring the costs associated with a visit to a dental professional, the other associated adverse effects are known to be transient in nature and to resolve once mouthrinsing ceases (Flotra 1971; Greenstein 1986). However, these adverse effects, although transient, can be severe and development of extrinsic tooth staining with short-term use of the mouthrinse is an obvious deterrent to using chlorhexidine mouthrinse.

Quality of the evidence

Although all the included studies were randomised controlled trials (RCTs), 50 of the 51 included studies were assessed as at high risk of bias overall, as they had at least one domain rated at high risk of bias. The reason many of the studies were assessed as at high risk of bias was due to problems with the blinding of participants, personnel, and outcome assessors. The majority of studies contained a placebo arm with the intention of blinding participants and personnel. However, we considered blinding unlikely to be successful due to the propensity for chlorhexidine to stain the teeth and oral tissues and the association of its use with clinically visible adverse effects such as mucosal desquamation and mucosal lesions. This also applied to the blinding of outcome assessors, in trials with either a placebo, control, or mechanical oral hygiene alone. This was not the fault of the investigators who in many studies went to great lengths to try to ensure blinding. Consequently we assessed 42 (82%) studies as at high risk of bias for either sort of blinding bias (performance bias or detection bias).

Thirty-seven studies were rated at high risk of selective reporting bias, mainly due to lack of reporting of measures of variance along with the means for each group, and no reporting of tooth staining and other adverse effects.

It is worth mentioning that 22 of the included studies were published prior to the first publication of the first iteration of the CONSORT Statement in 1996 (Begg 1996) and their quality was therefore partly judged according to reporting standards that did not exist at the time of their publication. An additional nine studies were published prior to the publication of the revised CONSORT Statement in 2001 (Moher 2001). However, disappointingly, instances of poor reporting were not confined to the studies that predated widespread awareness of the CONSORT Statement.

Despite most studies included in the meta-analyses being assessed as at high risk of bias we did not downgrade the GRADE assessments for gingivitis and plaque for this reason. This is because we believe that further research is very unlikely to change our confidence in the estimate of effect for these outcomes. The grade assessments for the body of evidence for chlorhexidine mouthrinse compared with placebo, control or mechanical oral hygiene alone for gingivitis measured using the Löe and Silness Gingival Index and for plaque measured using various indices were therefore rated as high quality. Although there was high heterogeneity for some of these meta-analyses we did not downgrade for this due to consistency of the direction of effect.

Over half of the included studies were either funded directly, received support for the study or were affiliated in some way with industry. Although a high level of industry funding is far from ideal, the reality is that without industry funding few trials evaluating the effectiveness of chlorhexidine mouthrinse would have been conducted. However, it is noteworthy that of the 19 included studies funded directly by industry, only one study (Zimmer 2015) stated that the conduct of the research and final decisions regarding the study report were independent of the company funding the research.

Potential biases in the review process

In order to prevent judgements about the eligibility criteria for studies being influenced by prior knowledge of the study results, the methods to be used in the review should be established and documented in advance of the review being undertaken (Higgins 2011). We made a number of changes to the protocol in relation to the types of studies and types of participants to be included in the review. We also clarified additional criteria for the inclusion of studies based on type of intervention that were implicit but not explicitly stated in the protocol. We amended the assessment of the blinding domains in the 'Risk of bias' tool to allow studies to obtain a judgement of 'low' or 'unclear' risk of bias in certain circumstances compared to the blanket judgement of high risk of bias that was stated in the published protocol.

A hierarchy to guide data extraction for gingivitis and plaque data was developed to facilitate data extraction and analysis. The main analysis however, was based on the main prespecified gingivitis index: the Gingival Index of Löe and Silness (Löe 1967; Löe and Silness 1963). Full details of the changes to the protocol are described in the Differences between protocol and review section. It is acknowledged that it may not always be appropriate to adhere to the published protocol (Higgins 2011). Therefore, while these changes to the protocol could introduce a risk of bias, the changes were made in the early stages of the review before any analyses of the results were undertaken and are therefore considered unlikely to have been influenced by knowledge of the study results or the results of the review. Furthermore, each decision was appropriately justified and was made to improve the scientific quality and clinical applicably of the review.

We estimated the standard deviations for four studies that measured the Gingival Index at 6 months (Banting 1989; Jayaprakash 2007; Lucas 1999; Sanz 1994) and one study that measured the Gingival Index at 4 to 6 weeks (Sanz 1989) using data from the same outcomes measured at the same time point in other similar studies so that we could include these data in meta-analyses. Where mean scores or measures of variance of the mean scores or both were not reported in the text or in tabular form, and could not be obtained from the authors of the studies, data were estimated by reading them off the graphs in the published report for inclusion in meta-analyses for six studies (Brecx 1993; Brightman 1991; Chaves 1994; Ferretti 1987; Jenkins 1993; Joyston-Bechal 1993). The graphs were enlarged and the data were estimated independently and in duplicate. A sensitivity analysis for the Gingival Index at both 4 to 6 weeks and 6 months, excluding trials where we had to estimate the standard deviation from similar studies, or had to read the data from graphs, showed that the effect estimates were similar to those for the overall pooling of the studies.



Agreements and disagreements with other studies or reviews

Van Strydonck 2012 conducted a comprehensive review of the effect of chlorhexidine mouthrinse on plaque, gingival inflammation and tooth staining in studies of 4 weeks or longer. This review was very similar to our review but slight differences in the inclusion criteria and the dates of the literature searches meant that a slightly different group of studies were reviewed by Van Strydonck et al. The review focused on healthy adults only, considered chlorhexidine mouthrinse used as a monotherapy or as an adjunct, and excluded patients with periodontitis or dentures as well as orthodontic and implant patients. In addition, apart from tooth staining, adverse effects although reported in the review, were not prespecified outcomes. Of the 30 studies included in this review, 27 are common to our review. The results of the metaanalysis of studies reporting the GI at all time points included data from seven studies, six of which were included in our metaanalyses at either 4 to 6 weeks, 7 to 12 weeks or 6 months. The weighted mean difference for the GI in this review was -0.32 which is similar to the mean difference for the GI reported in our review. Extrinsic tooth staining, increased calculus formation and change of taste sensation were also noted to have frequently occurred in the included studies. Considerable heterogeneity in the design, study duration and rinse regimen were also noted to be present.

Boyle 2014 systematically reviewed the evidence relating to the efficacy of the most frequently used mouthrinses for the control of supragingival plaque and gingivitis. Both the scope of the literature search and the inclusion criteria were narrower than in our review. Of the 17 trials involving chlorhexidine mouthrinse included in this review 16 are common to our review. The reported 'summary relative differences' facilitated comparison between the different mouthrinse preparations being reviewed. Results for the chlorhexidine mouthrinse studies are supportive of the effectiveness of chlorhexidine mouthrinse primarily for the control of both gingivitis and plaque relative to control but also relative to other active mouthrinse preparations.

A systematic review by Serrano 2015 evaluated the effectiveness of adjunctive antiplaque chemical agents including chlorhexidine mouthrinse in RCTs of at least 6 months duration. The review included 14 studies evaluating chlorhexidine mouthrinse. Due to differences in inclusion criteria, 12 of these studies are common to our review. The authors reported a significant benefit for rinsing with 0.1% to 0.2% chlorhexidine mouthrinse based on a metanalysis of four studies (weighted mean difference on the GI -0.185) but a non-significant effect for chlorhexidine mouthrinse at concentrations of \leq 0.06%. In common with our review was the presence of high heterogeneity which made it difficult to pool the data and was an acknowledged limitation of the review.

Gunsolley 2006 conducted a meta-analysis of 6-month mouthrinse studies for the control of gingivitis and plaque. Six of the included studies evaluated the effect of 0.12% chlorhexidine mouthrinse using the GI and all six were common to our review. The analysis indicated a moderate effect for gingivitis and a large effect for plaque. A subsequent metareview also conducted by Gunsolley drew on the data from the earlier meta-analysis and found a weighted mean percentage reduction of 28.7% for the GI and 40.4% for the Plaque Index (Gunsolley 2010) based on the same six studies.

The efficacy of 0.12% chlorhexidine mouthrinse compared to 0.2% on plaque and periodontal parameters was systematically reviewed by Berchier 2010. The review included eight studies with rinse duration from 3 days to 3 months that directly compared 0.12% with 0.2% chlorhexidine mouthrinse. The review found no difference in the effect on gingivitis between the two concentrations of chlorhexidine. The small significant difference in plaque inhibition in favour of 0.2% chlorhexidine was thought to be of negligible clinical relevance. Our review included two studies that provided data for a direct comparison between 0.2% and 0.1%/0.12% chlorhexidine rinse (Axelsson 1987; Segreto 1986), one of which was also included in the Berchier review (Segreto 1986). There was no difference in effect between the different concentration rinses used with the same volume and duration of rinsing in these two studies. Similarly our subgroup analysis found no evidence that one concentration of chlorhexidine rinse was more effective than another in reducing gingivitis and plaque.

van der Weijden 2015 conducted a metareview of the evidence regarding the efficacy and safety of chemical agents in mouthrinses to manage gingivitis and plaque. The evidence for chlorhexidine mouthrinse was drawn from the systematic reviews (Serrano 2015; Van Strydonck 2012) and meta-analyses (Gunsolley 2006; Gunsolley 2010) described above. Berchier 2010 contributed data for the comparison of active ingredients. This review reaffirmed the effectiveness of chlorhexidine mouthrinse for reducing plaque and gingivitis. The body of evidence resulting from these three reviews was rated as 'strong'. Due to the heterogeneity in the meta-analyses, the authors advised caution in interpreting the difference of means as it "may not provide an exact measure of the results" (van der Weijden 2015). Although the authors concluded that chlorhexidine is the "first choice", they also acknowledged that the development of taste disturbance, tooth staining and calculus formation prohibit long-term adjunctive use of the mouthrinse.

AUTHORS' CONCLUSIONS

Implications for practice

This review found high-quality evidence of a large reduction in dental plaque with chlorhexidine mouthrinse used as an adjunct to mechanical oral hygiene procedures for 4 to 6 weeks and 6 months. We also found high-quality evidence from studies that reported the Löe and Silness Gingival Index of a reduction in gingivitis in individuals with mild gingival inflammation on average (mean score of 1 on the 0 to 3 Gingival Index (GI) scale), that was not considered to be clinically relevant. There was no evidence that one concentration of chlorhexidine mouthrinse was more effective than another. There was insufficient evidence to determine the reduction in gingivitis associated with chlorhexidine mouthrinse use in individuals with mean GI scores of 1.1 to 3 indicating moderate or severe levels of gingival inflammation. Rinsing with chlorhexidine mouthrinse for 4 weeks or longer causes extrinsic tooth staining. In addition, other adverse effects such as calculus build-up, transient taste disturbance and effects on the oral mucosa were reported in the included studies. Chlorhexidine mouthrinse is indicated in particular clinical situations for short periods of time. Using chlorhexidine mouthrinse for longer periods of time in individuals with special care needs who cannot maintain an adequate level of plaque control using mechanical cleaning methods alone must be carefully weighed against the adverse effects associated with its use.



Implications for research

The majority of the studies included in the main analysis of the Gingival Index at 4 to 6 weeks and 6 months involved healthy participants with low levels of gingivitis. There was insufficient evidence to determine the reduction in gingivitis associated with chlorhexidine mouthrinse use in individuals with mean GI scores of 1.1 to 3 indicating moderate or severe levels of gingival inflammation. Some of the studies included in the metaanalyses of gingival bleeding at 4 to 6 weeks and 6 months included participants with a high proportion of sites with bleeding on probing. However, due to the variety of different gingival bleeding indices employed, we were unable to determine the mean reduction in gingivitis associated with chlorhexidine mouthrinse use in individuals with moderate or severe levels of gingival inflammation. Further well-conducted randomised controlled trials are needed to investigate the effect size for adjunctive chlorhexidine mouthrinse use in individuals with moderate or severe levels of gingival inflammation. Agreement among the experts about which indices are most appropriate for measuring gingivitis in studies evaluating antimicrobial mouthrinses would facilitate future evidence syntheses. Evidence-based guidance regarding patient selection for chlorhexidine mouthrinse use would help to provide clarity regarding the 'particular clinical situations' in which short-term use of chlorhexidine mouthrinse is recommended.

Only five studies evaluating 0.05% or 0.06% chlorhexidine mouthrinse met the inclusion criteria for this review. These studies demonstrated a reduction in plaque but the effect of these low-concentration mouthrinses on gingivitis was not clear or consistent. As low-concentration chlorhexidine mouthrinse is currently marketed for long-term daily use, well-conducted randomised controlled trials are needed to investigate the effect of these mouthrinses on gingivitis and to record any adverse effects associated with their use.

Due to the unavoidable influence of extrinsic tooth staining associated with use of chlorhexidine mouthrinse on blinding

of participants, personnel and outcome assessors, particular attention must be given to robust study design and rigorous reporting of the other domains in the Cochrane 'Risk of bias' tool. In particular, estimates of effect should be fully reported with a measure of variance so that data can be included in meta-analyses. Evidence of adverse effects should also be fully reported for each study arm. The CONSORT Statement (Schulz 2010) should be used to optimise reporting in future studies. The appropriateness of using studies with cross-over design that include chlorhexidine mouthrinse warrants further investigation.

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CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

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Anauate-Netto 2014	
Methods	Trial design: parallel (3 arms - 2 arms relevant to this review)
	Location: dental clinics at University Bandeirante of Sao Paulo, Sao Paulo, Brazil
	Number of centres: 1
	Study duration: the recruitment period is not stated. Study duration and duration of rinsing was 4 weeks
Participants	Participants: healthy adults
	Inclusion criteria: at least 20 teeth, no clinical signs of periodontal disease, aged between 18 and 55 years, normal saliva secretion rate
	Exclusion criteria: smokers, pregnancy, taking oral topical or systemic medication
	Baseline gingivitis: (Papillary Bleeding Score) overall (3 groups) mean 1. Gp A: mean 1.1 (SD 0.5); Gp B: mean 1.0 (SD 0.5); Gp C: mean 0.9 (SD 0.4)
	Age at baseline (years): overall: 40. Gp A: mean 41.6 (SD 13.4); Gp B: mean 39.4 (SD 9.8); Gp C: mean 39 (SD 11.7)
	Gender: overall: male 24 (40%), female 36 (60%). Gp A: male 7 (35%), female 13 (65%); Gp B: male 8 (40%), female 12 (60%); Gp C: male 9 (45%), female 11 (55%)
	Number randomised: 60 (Gp A: 20; Gp B: 20; Gp C: 20)
	Number evaluated: 60 (Gp A: 20; Gp B: 20; Gp C: 20)
Interventions	Comparison: CHX rinse versus propolis rinse* versus placebo rinse
	Gp A (n = 20): CHX 0.12%, 15 mL, twice daily, duration not specified, 4 weeks
	*Gp B (n = 20): alcohol-free, 2% typified propolis (plus mint flavour, polioxyethelers, sorbitol, blue colour and water), 15 mL, twice daily, duration not specified, 4 weeks. We excluded this arm from the risk of bias and analysis

^{*} Indicates the major publication for the study



Anauate-Netto 2014 (Continued	Gp C (n = 20): placebo (same ingredients as Gp B without the propolis), 15 mL, twice daily, duration not specified, 4 weeks
	Prophylaxis at baseline: not reported
	OHI: none given. Subjects followed their usual oral hygiene procedures
	Non-supervised rinsing Timing of mouthrinsing in relation to toothbrushing: quote: "Rinsing was performed in the morning and before bedtime after ordinary oral hygiene procedures"
Outcomes	Gingivitis (gingivial inflammation: Papillary Bleeding Score, gingival bleeding: PBS ≥ 2); adverse reactions; assessed at 4 weeks
Funding	Quote: "This study was supported by a grant from FAPESP (Fundacao de amparo a pesquisado estado de Sao Paulo) protocol no. 2007/53047-3"
Notes	Sample size calculation: quote: "It was not possible to conduct a sample size and power analysis because of lack of randomized clinical trials of propolis on gingivitis"
	Adverse effects: Gp A (CHX): 23 reports (burning sensation, taste alteration, yellow teeth, breath alteration, tongue burning, mucosal irritation, bitter taste); Gp B (propolis): 7 reports (breath alteration, burning sensation, taste alteration, yellow teeth, bitter taste); Gp C (placebo): 9 reports (taste alteration)
	Non-smokers
	Declarations/conflicts of interest: none stated

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "a computer-generated list of random numbers was used"
		Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Low risk	Quote: "Rinses were prepared in dark-bottles, which were consecutively numbered according to the randomization schedule Study coordinator, examiners, and participants were unaware of group allocation. The group identity was generated and kept in Florianopolis, SC, Brazil while the study was conducted in Sao Paulo, SP, Brazil"
		Comment: remote/central randomisation should have ensured that the random sequence was implemented as it was generated, without any manipulation
Blinding of participants	High risk	Quote: "double-blind placebo-controlled" and "rinses
and personnel (perfor- mance bias) All outcomes		were prepared in dark-bottles." The placebo mouthrinse matched the propolis mouthrinse without the active ingredient
		Comment: the level of adverse reactions in the CHX group ($n=23$) was higher than in the propolis ($n=7$) and placebo groups ($n=9$) may have meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is not clear
Blinding of outcome assessment (detection bias) All outcomes	High risk	Quote: "double-blind". The incidence of adverse reactions in the CHX group (n = 23) was higher than in the propolis (n = 7) and placebo groups (n = 9)



Anauate-Netto 2014 (Continued	d)	
		Comment: the authors do not report precisely who was blinded, but even if this includes the outcome assessors the higher level of adverse effects including tooth staining in the CHX group could have allowed the outcome assessors to work out which individuals were in the CHX group and could have affected the outcome assessment. The direction of this potential bias is not clear
Incomplete outcome data (attrition bias) All outcomes	Low risk	All randomised participants were included in the analyses
Selective reporting (reporting bias)	High risk	All outcomes stated in the methods section are reported in the results section. Adverse effects including 'yellow teeth' are reported aggregated as the total number of participants affected by group. Insufficient information on tooth staining is reported for us to use the data in a meta-analysis
Other bias	Low risk	Quote: "PBS measurements were performed by an experienced examiner (AAA). Intra-examiner reliability exercises revealed a Kappa test score of 0.85, indicating adequate reproducibility of PBS measurements"
		Comment: the potential for differential diagnostic activity was minimised.
		Groups were balanced at baseline (NS differences) for age, gender and race. The groups appear balanced for PBS at baseline

Anderson 1997

Methods	Trial design: parallel (2 arms)		
	Location: Postgraduate Clinic of Orthodontics, University of Texas-Houston, USA		
	Number of centres: 1		
	Study duration: recruitment period is not reported. Study duration and duration of rinsing was 3 months		
Participants	Participants: adolescents undergoing orthodontic treatment		
	Inclusion criteria: quote: "The participants did not suffer from any systemic complications nor were they taking any medications that may have had an effect on the oral tissues. All the participants had banding of at least one molar per quadrant"		
	Exclusion criteria: not reported		
	Baseline gingivitis: (GI) overall 0.925. Gp A: mean 1.04 (no SD); Gp B: mean 0.81 (no SD)		
	Age at baseline (years): range 11-15		
	Gender: not reported		
	Number randomised: 32 (Gp A: 16; Gp B: 16)		
	Number evaluated: 28 (Gp A: 13; Gp B: 15)		
Interventions	Comparison: CHX rinse versus placebo rinse		
	Gp A (n = 16): CHX (Peridex) 0.12%,15 mL, twice daily, 30 seconds, 3 months		
	Gp B (n = 16): placebo (identical to the experimental rinse except for the active ingredient), 15 mL, twicdaily, 30 seconds, 3 months		



Anderson	1997	(Continued)
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Prophylaxis at baseline: quote: "they received a dental prophylaxis which included the removal of plaque, calculus and stains from the teeth by scaling and polishing" (8-10 days before baseline measurements)

OHI: they received oral and written toothbrushing instructions with the same toothpaste (Crest regular flavour) to be performed twice daily. The method used was the Modified Bass technique. Soft rounded toothbrushes were supplied to all patients

Non-supervised rinsing

Timing of mouthrinsing in relation to toothbrushing: not reported

Postrinse instructions: not reported

Outcomes

Gingivitis (gingival inflammation: GI Löe & Silness 1963); plaque (PI Silness & Löe 1964); calculus (Retention Index, Löe 1967); assessed at 1, 2 and 3 months follow-up

Probing depths (CP-12 Hu Friedy periodontal probe) - not relevant to this review and tooth staining (Discolouration Index, Lang 1981); assessed at 3 months only

Funding

Quote: "Partially supported by Procter & Gamble, by providing the chlorhexidine and placebo for the patients in this study"

Notes

Sample size calculation: not reported

Adverse effects: quote: "...no side effects were reported as a result of the use of either the experimental or the control solutions except for one patient in CHX group who acknowledged a burning sensation on the mucosa after the use of the experimental solution." No significant differences were found between the groups for discolouration or calculus accumulation, although tooth discolouration "was more evident in the experimental group" and for calculus accumulation "the higher mean values were always found in the experimental group"

Declarations/conflicts of interest: none stated

Bias	Authors' judgement	Support for judgement
Random sequence genera-	Unclear risk	Quote: "They were randomised to two groups"
tion (selection bias)		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "The control group received exactly the same instructions with a flavoured placebo solution which was identical to the experimental solution except for the active ingredient" and "the discoloration was more evident in the experimental group but it was not statistically significant"
		Comment: the study is not described as blind by the authors but the methods described indicate that it was intended for the participants to be unaware of the allocated intervention. However, the higher level of staining in the active group (although not statistically significant) meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is not clear
Blinding of outcome as-	High risk	No information is provided
sessment (detection bias) All outcomes		Comment: the study is not described as blind by the authors. The higher level of staining in the CHX group (although not statistically significant) meant that



Anderson 1997 (Continued)		the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear
Incomplete outcome data (attrition bias) All outcomes	High risk	12.5% (4/32) of randomised participants were not included in the analysis. Attrition by group: CHX 3/16 (18.8%), placebo 1/16 (6.3%) 2 individuals left the study after they received the baseline prophylaxis (CHX group), 1 before the 30 day evaluation (CHX group) and 1 was unable to return for the 60 and 90 day evaluations (control group)
		Comment: attrition was quite high in the CHX group as proportion of the total number in the group. Although the attrition is unlikely to be due to the intervention (CHX) (2 of the 3 participants who left the CHX group left the study before they started rinsing), if the missing participants had higher mean gingivitis scores in 1 group than the other, as the attrition rate increased, so would over/understatement of the mean difference
Selective reporting (reporting bias)	High risk	Mean scores and standard deviations for all outcomes were reported by tooth surface only which meant that with the exception of the tooth staining data (where we were able to combine data for buccal and lingual surfaces), we could not use the data in meta-analyses
Other bias	Unclear risk	No information is provided on intra- or inter-examiner reproducibility. There were no significant differences between the means of the plaque indices and gingival indices recorded on all surfaces for both experimental and control groups. The Retention Index, Discolouration Index and mean probing depth measurements were also balanced Mean age and gender are not reported

Axelsson 1987

Methods	Trial design: parallel (4 arms - 3 arms are relevant to this review)
	Location: dental school, Sweden
	Number of centres: 1
	Study duration: recruitment period is not reported. Study duration and duration of rinsing was 6 weeks
Participants	Participants: volunteers aged 16-50 years with gingivitis
	Inclusion criteria: not reported.
	Exclusion criteria: periodontal attachment loss or bone loss
	Baseline gingivitis (mean (SE): (GI Löe & Silness 1963) overall 1.21. Gp A: mean 1.18 (0.08); Gp B: mean 1.26 (0.07); Gp C: mean 1.19 (0.07); Gp D: mean 1.21 (0.08))
	Age at baseline (years): mean age not reported, range 16-50
	Gender: not reported
	Number randomised: 96, (Gp A: 24; Gp B: 24, Gp C: 24, Gp D: 24)
	Number evaluated: 88, (Gp A: 18; Gp B: 24, Gp C: 24; Gp D: 22)
Interventions	Comparison: CHX (Hibitane 0.2%) versus CHX (Hibitane 0.1%) versus Listerine* versus control
	Gp A (n = 24): CHX (Hibitane) 0.2%, 10 mL, twice daily, 60 seconds, 6 weeks
	Gp B (n = 24): CHX (Hibitane) 0.1%, 10 mL, twice daily, 60 seconds, 6 weeks



Rinsing was supervised Mon-Fri and unsupervised at weekends Timing of mouthrinsing in relation to toothbrushing is not reported Outcomes Gingivitis (gingival inflammation: GI Löe & Silness 1963, gingival bleeding: % GI scores 0, 1, 2 or 3) assessed at 6 weeks Plaque (TQH Turesky 1970); extrinsic tooth stain (Lobene 1968); assessed at 3 and 6 weeks A complete intraoral soft tissues examination was performed after 3 and 6 weeks Funding No information provided Notes Sample size calculation: quote: "The study was designed to provide a minimal power of 0.70 for detecting a clinically important difference to be statistically significant at the 0.05 probability level. The final sample size was based on the maximum determined among the separate requirements of the plaque and gingival index scores." The sample size required is not reported Adverse effects: mucosal lesions occurred in 6 subjects in the Hibitane group (0.2%), 1 subject in the control group complained of oral irritation which worsened during the study The order of examinations were a) soft tissue condition, b) stain index, c) gingival index, d) plaque index Declarations/conflicts of interest: none stated Groups A and B were combined for the main analysis	Axelsson 1987 (Continued)	*Gp C (n = 24): Listerine, 20 mL, twice daily, 30 seconds, 6 weeks. We excluded this arm from the risk of bias and analysis Gp D (n = 24): control rinse (composition not described), 20 mL, twice daily, 30 seconds, 6 weeks Prophylaxis at baseline: scaling and professional tooth cleaning after the baseline examination OHI: they were supplied with a soft nylon toothbrush and Colgate MFP toothpaste, were instructed to refrain from using commercial mouthrinses and continued to exercise their regular non-supervised self-performed plaque control programs
Outcomes Gingivitis (gingival inflammation: GI Löe & Silness 1963, gingival bleeding: % GI scores 0, 1, 2 or 3) assessed at 6 weeks Plaque (TQH Turesky 1970); extrinsic tooth stain (Lobene 1968); assessed at 3 and 6 weeks A complete intraoral soft tissues examination was performed after 3 and 6 weeks Funding No information provided Notes Sample size calculation: quote: "The study was designed to provide a minimal power of 0.70 for detecting a clinically important difference to be statistically significant at the 0.05 probability level. The final sample size was based on the maximum determined among the separate requirements of the plaque and gingival index scores." The sample size required is not reported Adverse effects: mucosal lesions occurred in 6 subjects in the Hibitane group (0.2%), 1 subject in the control group complained of oral irritation which worsened during the study The order of examinations were a) soft tissue condition, b) stain index, c) gingival index, d) plaque index Declarations/conflicts of interest: none stated Groups A and B were combined for the main analysis		Rinsing was supervised Mon-Fri and unsupervised at weekends
Sessed at 6 weeks Plaque (TQH Turesky 1970); extrinsic tooth stain (Lobene 1968); assessed at 3 and 6 weeks A complete intraoral soft tissues examination was performed after 3 and 6 weeks Funding No information provided Notes Sample size calculation: quote: "The study was designed to provide a minimal power of 0.70 for detecting a clinically important difference to be statistically significant at the 0.05 probability level. The final sample size was based on the maximum determined among the separate requirements of the plaque and gingival index scores." The sample size required is not reported Adverse effects: mucosal lesions occurred in 6 subjects in the Hibitane group (0.2%), 1 subject in the control group complained of oral irritation which worsened during the study The order of examinations were a) soft tissue condition, b) stain index, c) gingival index, d) plaque index Declarations/conflicts of interest: none stated Groups A and B were combined for the main analysis		Timing of mouthrinsing in relation to toothbrushing is not reported
A complete intraoral soft tissues examination was performed after 3 and 6 weeks No information provided Sample size calculation: quote: "The study was designed to provide a minimal power of 0.70 for detecting a clinically important difference to be statistically significant at the 0.05 probability level. The final sample size was based on the maximum determined among the separate requirements of the plaque and gingival index scores." The sample size required is not reported Adverse effects: mucosal lesions occurred in 6 subjects in the Hibitane group (0.2%), 1 subject in the control group complained of oral irritation which worsened during the study The order of examinations were a) soft tissue condition, b) stain index, c) gingival index, d) plaque index Declarations/conflicts of interest: none stated Groups A and B were combined for the main analysis	Outcomes	
Notes Sample size calculation: quote: "The study was designed to provide a minimal power of 0.70 for detecting a clinically important difference to be statistically significant at the 0.05 probability level. The final sample size was based on the maximum determined among the separate requirements of the plaque and gingival index scores." The sample size required is not reported Adverse effects: mucosal lesions occurred in 6 subjects in the Hibitane group (0.2%), 1 subject in the control group complained of oral irritation which worsened during the study The order of examinations were a) soft tissue condition, b) stain index, c) gingival index, d) plaque index Declarations/conflicts of interest: none stated Groups A and B were combined for the main analysis		Plaque (TQH Turesky 1970); extrinsic tooth stain (Lobene 1968); assessed at 3 and 6 weeks
Notes Sample size calculation: quote: "The study was designed to provide a minimal power of 0.70 for detecting a clinically important difference to be statistically significant at the 0.05 probability level. The final sample size was based on the maximum determined among the separate requirements of the plaque and gingival index scores." The sample size required is not reported Adverse effects: mucosal lesions occurred in 6 subjects in the Hibitane group (0.2%), 1 subject in the control group complained of oral irritation which worsened during the study The order of examinations were a) soft tissue condition, b) stain index, c) gingival index, d) plaque index Declarations/conflicts of interest: none stated Groups A and B were combined for the main analysis		A complete intraoral soft tissues examination was performed after 3 and 6 weeks
ing a clinically important difference to be statistically significant at the 0.05 probability level. The final sample size was based on the maximum determined among the separate requirements of the plaque and gingival index scores." The sample size required is not reported Adverse effects: mucosal lesions occurred in 6 subjects in the Hibitane group (0.2%), 1 subject in the control group complained of oral irritation which worsened during the study The order of examinations were a) soft tissue condition, b) stain index, c) gingival index, d) plaque index Declarations/conflicts of interest: none stated Groups A and B were combined for the main analysis	Funding	No information provided
control group complained of oral irritation which worsened during the study The order of examinations were a) soft tissue condition, b) stain index, c) gingival index, d) plaque index Declarations/conflicts of interest: none stated Groups A and B were combined for the main analysis	Notes	ing a clinically important difference to be statistically significant at the 0.05 probability level. The final sample size was based on the maximum determined among the separate requirements of the plaque
Groups A and B were combined for the main analysis		control group complained of oral irritation which worsened during the study
		Declarations/conflicts of interest: none stated
Risk of bias		Groups A and B were combined for the main analysis
	Risk of bias	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "Volunteers were assigned either to 1 of 3 different treatment groups or to a control group according to a randomised code by which double-blinding was maintained"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	Quote: "Volunteers were assigned either to 1 of 3 different treatment groups or to a control group according to a randomised code by which double-blinding was maintained"
		Comment: it is not clear who assigned the participants to the groups or whether participants or investigators could foresee assignment
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	The study is described as double-blind. The authors do not report precisely who was blinded. The composition of the control is not described. There were negligible differences in extrinsic tooth staining between the groups at 3 and 6 weeks
		Comment: neither participants nor personnel could be truly blind as different volumes and rinse times were used in different groups and rinsing was supervised on weekdays. Knowledge of which product was which could have influ-



Axelsson 1987 (Continued)		enced behaviour of participants and hence the outcome. The direction of this potential bias is not clear
Blinding of outcome assessment (detection bias) All outcomes	Low risk	The authors state that the study is "double-blind" but do not report precisely who was blinded. There were negligible differences in extrinsic tooth staining between the groups at 3 and 6 weeks. All intraoral examinations (including tooth staining) were performed by a single examiner
Incomplete outcome data (attrition bias) All outcomes	High risk	11% (8/72) of randomised participants were not included in the analysis Attrition by group: Hibitane 0.2% 6/24 (25%); Hibitane 0.1% (0%); control 2/24 (8.3%). Reasons for attrition: Hibitane 0.2%: mucosal lesions; control: oral irritation, personal reasons
		Comment: there were differential group losses with proportionately more losses from the Hibitane 0.2% group (due to the occurrence of mucosal lesions) which could have influenced the outcome
Selective reporting (reporting bias)	High risk	All outcomes stated in the methods section are reported in the results section. We calculated standard deviations for gingivitis and plaque from the standard errors reported in the tables. We also calculated the % of gingival units with a GI score of 2 or 3 however, no measure of variance was reported so we could not include the data in meta-analyses. Extrinsic tooth staining and adverse effects were adequately reported. We combined groups A and B for the main analysis
Other bias	Unclear risk	All intraoral examinations (including tooth staining) were performed by a single examiner. No information is provided on intraexaminer reproducibility. Mean gingivitis, plaque and stain scores at baseline appear to be reasonably balanced
		The age range is quite wide (16-50) but baseline data on age and gender balance are not reported

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Bajaj 2011	
Methods	Trial design: parallel (3 arms - 2 arms are relevant to this review)
	Location: Department of Pedodontics and Preventative Dentistry, Manipal College of Dental Sciences, Manipal, India
	Number of centres: 1
	Study duration: recruitment period not reported. Duration of study is 9 months; duration of rinsing is not explicitly stated, assumed to be 9 months
Participants	Participants: quote: "Children with similar socioeconomic status, dietary habits, oral hygiene status, and KAP (knowledge, attitude and practice) status were included. Further, only children who had a minimum of one to two established carious lesions were considered. The subjects were selected from residential schools"
	Exclusion criteria: not reported
	Baseline gingivitis: GI Löe & Silness 1967, Gp A (0.54 SD 0.22); Gp B (0.59 SD 0.73); Gp C (1.16 SD 0.21)
	Age at baseline (years): 8 to 12 years, not reported by group
	Gender: not reported
	Number randomised: 1431 (not reported by group)



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Number evaluated: 1309 (Gp A: 440; Gp B: 457; Gp C: 412)

Interventions

Comparison: CHX rinse versus Triphala rinse* versus control rinse

Gp A (440 analysed): CHX (Clohex 0.1% (diluted from 0.2%)) 10 mL, 1 min, once daily

*Gp B (457 analysed): Triphala (0.6%) 10 mL, not reported, once daily. We excluded this arm from the risk of bias and analysis

Gp C (412 analysed): control (distilled water) 10 mL, not reported, once daily

[Conflicting reports for duration of rinsing – 2 min rinse reported in methods for CHX rinse (duration of rinsing for Triphala and control not reported) and 1 min rinse reported in discussion for all 3 rinses. Rinse duration stated in the discussion is reported here]

Prophylaxis at baseline: not reported

OHI: not reported

Rinsing supervised during the week, non-supervised at weekends and during vacations

Timing of mouthrinsing in relation to toothbrushing: not reported

Postrinsing instructions: quote: "The children were advised not to eat or rinse for the next 30 min"

Outcomes

Gingivitis (gingival inflammation: GI Löe & Silness 1967) and plaque (PI Silness & Löe 1964) recorded at 3, 6 and 9 months follow-up. Microbiological analysis (*Streptococcus* and *Lactobacillus* counts) at baseline 6 and 9 months follow-up (not relevant to this review)

Funding

Quote: "This research project was funded by the Indian Council of Medical Research"

Notes

Sample size calculation: quote: "Large sample size was selected anticipating the possible attrition of the sample due to varying cultural background and migration of few students to other schools" (actual calculation not mentioned)

Adverse effects: not reported

Declarations/conflicts of interest: not reported

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "the subjects were allocated to the specific treatment by block randomization"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quotes: "it was a double-blind clinical trial Both solutions (chlorhexidine and Triphala) were made of identical colors. The bottles were then coded and then at the end of the study, the decoding was done" and "The schools were distributed in such a manner so that there was no intermingling within the students of different groups." The control solution was distilled water. Rinsing was supervised by teachers on school days
		Comments: the authors do not report precisely who was blinded. Blinding of the teachers is not explicitly stated. Higher levels of tooth staining are likely to have been noticeable in the CHX group, however, tooth staining is not reported and therefore not enough information is provided upon which to base



Bajaj 2011 (Continued)		a judgement on the likely effect of staining. The control group rinsed with distilled water and could have worked out which group they were in which could have affected their oral health behaviours
Blinding of outcome assessment (detection bias)	Unclear risk	Quote: "it was a double-blind clinical trial" No other information relating to blinding of outcome assessment is reported
All outcomes		Comment: the authors do not report precisely who was blinded. Higher levels of tooth staining are likely to have been noticeable in the CHX groups, however, tooth staining is not reported and therefore not enough information is provided upon which to base a judgement
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition after 9 months (all arms included) 122/1431 (8.53%). Numbers lost in each group are not reported. Authors report % lost by group as follows: CHX 7.36%, control 10.62%. Reasons for losses: quote: "the majority of times attrition occurred due to family reasons where the parents changed the school of their child"
		Comments: attrition is reasonably low (highest in the control group), reasonably balanced across groups and reasons for losses are unlikely to be related to the interventions
Selective reporting (reporting bias)	High risk	There are marked inconsistencies between data reported in text and data reported in figure 1 in relation to mean plaque scores. Tooth staining and adverse effects are not reported
Other bias	High risk	The number of examiners is not reported and there is no reference to training or calibration. Children are reported to have had similar SES, dietary habits, oral hygiene methods and KAP (knowledge, attitudes and practice) status. Children had a minimum of 1 to 2 established carious lesions. The control group (distilled water) had much higher mean plaque and gingivitis scores than the CHX group at baseline: plaque: CHX 0.76, control 1.76; gingivitis: CHX 0.54, control 1.16. (Data taken from the text rather than figures: see comment above about inconsistencies between the text and the figures)

Banting 1989

Danting 1909	
Methods	Trial design: parallel, 2 arms
	Location: not specified, but assumed to be the University of Western Ontario Dental School, Ontario, Canada
	Number of centres: assumed 1
	Study duration: recruitment period not stated. Duration of study and duration of rinsing was 24 months
Participants	Participants: employees from 3 hospitals, from the University of Western Ontario, and from the general public of London, Ontario
	Inclusion criteria: quote: "subjects had to be at least 18 years of age and with at least 16 natural teeth (including four molars) and some gingivitis"
	Exclusion criteria: quote: "Persons with advanced periodontal disease and/or gross dental neglect, those taking antibiotics regularly or within one week of the baseline examination, and those with a history of hepatitis, tuberculosis, heart disease, or other severe or debilitating medical conditions were excluded"
	Baseline gingivitis: not reported



Banting	1989	(Continued))
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Age at baseline: not reported but minimum age of 18 years

Gender: not reported

Number randomised: 456 (CHX 231, control 225)

Number evaluated at 6 months: 383 (CHX 190, control 193)

Number evaluated at 24 months: 272 (CHX 112, control 160)

Interventions

Comparison: CHX versus placebo rinse

CHX 0.12% (n = 231): 15 mL, twice daily, 30 seconds, 24 months

Control (placebo - composition not described) (n = 225): 15 mL, twice daily, 30 seconds, 24 months

Prophylaxis at baseline: quote: "All subjects were then (after baseline assessment) given a professional scaling and prophylaxis by a dental hygienist." Professional prophylaxis and scaling was repeated for all individuals every 6 months

OHI: none. Quote: "Toothbrushes and Crest dentifrice were also made available, but no additional oral care instruction was provided"

Non-supervised rinsing

Timing of mouthrinsing in relation to toothbrushing and postrinse instructions are not reported

Outcomes

Gingivitis (gingival inflammation: GI Löe & Silness 1967, gingival bleeding: GI scores 2 or 3), at 6, 12, and 24 months

Plaque (TQH Turesky 1970), at 6, 12, and 24 months

Calculus (supragingival calculus deposits (Volpe-Manhold Calculus Index, Volpe 1965) and subgingival calculus deposits) at 24 months only

Funding

Quote: "The support of this study by The Procter & Gamble Company, Cincinnati, Ohio, is gratefully acknowledged"

Notes

Sample size calculation: not reported

Adverse effects: quote: "Staining of the teeth was cited as the most common reason for subjects not continuing in the study." At the 24-month examination, subjects in the treatment group had higher mean supragingival calculus scores, however at the same time, more subjects in the CHX group were free of subgingival calculus. Soft tissue conditions were observed in 33% of the subjects at baseline, in 13% of the subjects at 1 year and in 12% of subjects at 2 years. The proportion and type of oral soft tissue conditions were similar for the treatment and control groups

Declarations/conflicts of interest: none stated. However, the second and third authors on the paper are affiliated to the Procter & Gamble Company

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "subjects were stratified by gender, age, and periodontal status and then randomly assigned to a control or treatment. Subjects residing in the same household were assigned to the same treatment group"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided



Banting 1989 (Continued)		
Blinding of participants	High risk	Tooth staining was noted in the CHX group
and personnel (perfor- mance bias) All outcomes		Comment: participant blinding is not mentioned but the use of a control 'placebo' (composition not described) indicate that it was intended for the participants to be unaware of the allocated intervention. However, the higher level of staining in the CHX group meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is not clear
Blinding of outcome assessment (detection bias) All outcomes	High risk	Quote: "Toothcover devices and tongue-masking were used at the one-year gingivitis examination to eliminate potential examiner bias related to extrinsic staining." Tooth staining was noted in the CHX group
		Comment: gingivitis and gingival bleeding results at 6 and 24 months and plaque and PPD results at 6, 12 and 24 months are at high risk of bias due to the higher level of tooth staining in the CHX group. The outcome assessors could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear. The impact of using tooth cover devices and tongue-masking at 12 months on blinding is not clear
Incomplete outcome data (attrition bias) All outcomes	High risk	6 months:
		16% (73/456) of randomised participants were not included in the analysis at 6 months. Attrition by group CHX 41/231 (18%), control 32/225 (14%)
		24 months:
		40% (184/456) of randomised participants were not included in the analysis at 24 months. Attrition by group CHX 119/231 (51.5%), control 65/225 (28.9%)
		Reasons for attrition: tooth staining was the most common reason. Other reasons were not reported
		Comment: attrition was high in both groups. There were proportionately more losses from the CHX group (due to tooth staining)
Selective reporting (reporting bias)	High risk	No baseline measure of gingival health was provided. No measures of variance were reported. Mean plaque scores were reported only in graphs. The mean supra- and subgingival calculus scores were not reported. We estimated the standard deviation for gingivitis (measured using the GI) as the median of the SDs in the placebo/control groups from similar studies that also used the GI at 6 months so that we could include the gingivitis data in the meta-analyses
Other bias	Unclear risk	Quotes: "One examiner was used for each of the soft-tissue, gingivitis and plaque measurements." "two examiners were used for the pocket depth assessments." No information is provided on intraexaminer reproducibility

and is not clear

Participant characteristics are not described

Subjects residing in the same household were assigned to the same treatment group. The number of participants who resided in the same household is not reported. The potential effect of clustering was not considered in the analysis

Bhat 2014

Methods Trial design: parallel (3 arms - 2 arms relevant to this review)



Bhat 2014 (Continued)	
	Location: Department of Public Health Dentistry, Darshan Dental College and Hospital, Udaipur, Rajasthan, India
	Number of centres: 1
	Study duration: recruitment period not reported. Study duration and duration of rinsing was 4 weeks
Participants	Participants: undergraduate dental students
	Inclusion criteria: quote: "The dental students with age group of 18–24 years, students willing to participate and with dentition of ≥20 teeth and a minimum of 5 teeth per quadrant, no relevant medical history and no periodontal treatment during past 3 months were included"
	Exclusion criteria: quote: "Students with orthodontic appliances or severe misaligned
	teeth, receiving antibiotic therapy or medication within past 6 months, presence of any systemic illness, subjects availing oral prophylaxis since past 6 months and students unable to comply with study appointment schedule were excluded"
	Baseline gingivitis: GI Löe & Silness 1963, Gp A: mean 0.77 (SD 0.43); Gp B: mean 0.62 (SD 0.52); Gp C: mean 0.91 (SD 0.84)
	Age at baseline (years): Gp A: mean 20.95 (SD 0.1); Gp B: mean 20.2 (SD 0.7); Gp C: 20.79 (SD 1.2). Range 18-24
	Gender: male 37 (51%), female 35 (49%), not broken down by group
	Number randomised: total 72 (Gp A: 24; Gp B: 24; Gp C: 24)
	Number evaluated: total 66 (Gp A: 22; Gp B: 22; Gp C: 22)
Interventions	Comparison: CHX (ClohexPlus) rinse versus herbal (Hiora) rinse* versus control (Nirlife) rinse
	Gp A (n = 24): CHX (0.20%) 10 mL, twice daily, 60 seconds, 4 weeks
	* Gp B (n = 24): herbal (Hiora) rinse 10 mL, twice daily, 60 seconds, 4 weeks. We excluded this arm from the risk of bias and analysis
	Gp C (n = 24): control (0.05% saline) 10 mL, twice daily, 60 seconds, 4 weeks
	Prophylaxis at baseline: not reported
	OHI: quote: "All the participants were instructed to follow their routine oral hygiene practices along with the assigned regimen and to maintain a reminder sheet on daily product use"
	Supervised and non-supervised rinsing: quote: "Each one of the daily rinses was supervised on each weekday and also supervised by a daily recall message for reminding them to use the assigned mouthwash"
	Timing of mouthrinsing in relation to toothbrushing: after toothbrushing
	Postrinsing instructions: not reported
Outcomes	Gingivitis: (gingival inflammation: GI Löe & Silness 1963); plaque (TQH Turesky 1970); adverse effects (burning sensation, dryness, taste disturbance, discolouration) assessed at 4 weeks
Funding	Quote: "Research funding: None declared"
Notes	Sample size calculation: quote: "From the pilot study, it was found that the overall prevalence of plaque and gingivitis was 80%. Considering for the drop outs logistic and technical problems the sample size was inflated by 20%, hence the sample size was 72 with 24 participants in each group"
	Adverse effects: a majority of the adverse effects resulted in Gp A (CHX group). It was observed that most of the study participants in Gp A complained of mild discolouration 18 (81.8%), moderate dis-



Bhat 2014 (Continued)

colouration 2 (9.1%) and dry mouth 18 (81.8%), whereas taste disturbance was higher in Gp C 14 (63.6%) as compared to Gp A 10 (45.4%). The least taste disturbance was seen in Gp B 2 (9.1%). The burning sensation was highest in Gp A 11 (50%) as compared to Gp B 9 (40.9%) and absent in Gp C

Declarations/conflicts of interest: quote: "Authors' conflict of interest disclosure: The authors stated that there are no conflicts of interest regarding the publication of this article"

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: " All the subjects were provided with their assigned mouthrinses and were divided into Group A, Group B and Group C randomly using simple lottery method with 24 participants in each group"
		Comment: if "simple lottery method" corresponds to drawing lots then we can consider risk of bias to be low
Allocation concealment (selection bias)	Unclear risk	Quote: "The assignment of the participants to the groups and the codes to the product was done by a person not involved in the examination"
		Comment: insufficient information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quotes: "All the mouthrinses were dispensed in the identical bottles and thereby ensuring a total subject masking. The examiner and the participants were also blinded with regard to the mouthrinse allocated to them thereby ensuring a double-blinded study" and "Normal saline was coloured to resemble the mouthwashes." A burning sensation was reported by almost half of the participants in the 2 active mouthrinse groups. 81.8% of study participants in the CHX group complained of mild discolouration and 9.1% of moderate discolouration
		Comment: higher levels of tooth staining were reported by most of the study participants in the CHX group. The higher level of staining in the CHX group meant that participants could have worked out which group they were in and this could have affected their oral health behaviours
Blinding of outcome assessment (detection bias) All outcomes	High risk	Quote: "The examiner and the participants were also blinded with regard to the mouthrinse allocated to them thereby ensuring a double-blinded study." A burning sensation was reported by almost half of the participants in the 2 active mouthrinse groups. 81.8% of study participants in the CHX group complained of mild discolouration and 9.1% of moderate discolouration
		Comment: the higher level of staining in the CHX group meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition after 4 weeks: 4/48 (8.3%). By group: CHX 2/24 (8.3%), control 2/24 (8.3%). Reasons for losses not broken down by group: quote: "Two students did not report after 1 month in spite of several recalls, three students were irregular in using mouthwash found during surprise visit and one of the students became ill and had to quit the study due to antibiotic coverage"
		Comments: losses are reasonably low and are balanced across groups
Selective reporting (reporting bias)	High risk	Standard deviations of mean scores for gingivitis (CHX group only) are not reported at 4 weeks. We used the SD for the CHX group at baseline so that we could include the gingivitis data (measured using the GI) in the meta-analyses. Adverse effects including 'discolouration' are reported (proportions with each adverse effect by group)



Bhat 2014 (Continued)

Other bias

Low risk

Quote: "The kappa statistical analysis for inter examiner variability for Turesky et al. Plaque index and Loe and Silness gingival index was 0.89 and 0.97, whereas the intra examiner was 0.90 and 0.94 respectively showing high degree of conformity in observational judgment. All recordings were made by a single examiner"

No statistically significant difference in mean age, mean baseline plaque scores and mean baseline gingival index scores

Brecx 1993

Methods

Trial design: parallel (3 arms - 2 arms relevant to this review)

Location: assumed to be at the Health Science Campus, University of Manitoba, Canada

Number of centres: 1

Study duration: recruitment period is not stated. Study duration and duration of rinsing was 3 months

Participants

Participants: quote: "Either medical or physiotherapy students, as well as non-dental laboratory staff, from the Health Science Campus, University of Manitoba"

Inclusion criteria: healthy adults

Exclusion criteria: not reported

Baseline gingivitis: (GI) Gp A: mean 0.6 (no SD); Gp B: mean 0.6 (no SD); Gp C: mean 0.6 (no SD)

Age at baseline (years): range 21-33

Gender: male 15 (42%), female 21 (58%), breakdown by group group data not reported, but gender distribution stated to be 'very similar' across treatment groups

Number randomised: 36 (assumed Gp A: 12; assumed Gp B: 12; assumed Gp C: 12)

Number evaluated: 31 (Gp A: 10; Gp B: 9; Gp C: 12)

Interventions

Comparison: CHX gluconate solution versus Meridol rinse* versus placebo rinse

Gp A (n = 12): CHX 0.2%,10 mL, twice daily, 1 min, 3 months

*Gp B (n = 12): Meridol, 10 mL, twice daily, 1 min, 3 months. We excluded this arm from the risk of bias and analysis

 ${\sf Gp\ C\ (n=12): placebo\ (quinine-hydrochloride\ solution),\ 10\ mL,\ twice\ daily,\ 1\ min,\ 3\ months}$

Prophylaxis at baseline: quote: "In the pretreatment phase, the individuals were subjected to a thorough prophylaxis, but not instructed in any form of mechanical plaque control." Subjects had teeth professionally cleaned and continued their usual oral hygiene for a period of 2 weeks until "month 0" at which time their teeth were polished again. They received a new toothbrush (Butler 411) for the 2 weeks preceding the beginning of month 0

OHI: none given. Quote: " ..but not instructed in any form of mechanical plaque control"

Non-supervised rinsing

Timing of mouthrisning in relation to toothbrushing: quote: "The participants were instructed to make sure to rinse twice daily regardless of their brushing habits. However, if they did brush, they were asked to use the mouthrinse after their toothbrushing"

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Brecx 1993 (Continued)	
Outcomes	Gingivitis (gingival inflammation: GI Löe & Silness 1967); plaque (PI Silness & Löe 1964), assessed at day 14, month 0, and after 1, 2, and 3 months of rinsing in addition to usual tooth cleaning
	Tooth stain (Modification of the Discolouration Index, Lang & Räber 1981). Time period not explicitly stated
	Dental plaque for vital fluorescence (VF) examination was collected at months 0, 1, 2, and 3 (not relevant to this review)
Funding	Quote: "This investigation was supported by a grant from GABA International Ltd, Therwil, Switzerland"
Notes	Sample size calculation: not reported
	Adverse effects: tooth staining i.e. higher DI scores in Meridol and CHX groups than placebo at end of study. At all time points the Meridol group showed less staining than the CHX group
	Declarations/conflicts of interest: none stated
Risk of bias	

Bias	Authors' judgement	Support for judgement
Random sequence genera-	Unclear risk	Quote: "The subjects were randomly assigned to three groups"
tion (selection bias)		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information provided
Blinding of participants and personnel (perfor- mance bias)	High risk	Quotes: "double-blind", "all solutions had the same color and were kept in the same kind of bottle (PET white, opaque)." A placebo containing 0.02% quinine-hydrochloride solution was used
All outcomes		Quote: "The DI increased with time in all groups At the end of the study, the Meridol and chlorhexidine groups exhibited higher scores (p<0.001) than the placebo group"
		Comment: the higher level of tooth staining in the CHX group meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is unclear
Blinding of outcome assessment (detection bias) All outcomes	High risk	Quotes: "double-blind", "All readings were noted by the same investigator" and "The DI increased with time in all groups At the end of the study, the Meridol and chlorhexidine groups exhibited higher scores (p<0.001) than the placebo group"
		Comment: the higher level of staining in the CHX group meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear
Incomplete outcome data (attrition bias) All outcomes	High risk	Assuming equal numbers in groups at the start (as this was not reported) attrition after 3 months was 2/24 (13.9%). Attrition by group was 2/12 (16.7%) CHX; no losses from placebo group. Reasons not broken down by group. Reasons for losses (all study arms): quote: "One subject decided to stop participation, and four others were dismissed for multiple evidence of non-compliance (noshow without several calls, no rinse during 1 min, too much solution left in the bottle, etc)"
		Comment: there were differential group losses with losses from the CHX group and none from the placebo group. The number of participants lost over the 3



Brecx 1993 (Continued)		months is small but because the groups were small at baseline the proportion ate losses are high
Selective reporting (reporting bias)	High risk	Mean scores for tooth staining at 4 weeks were not reported in the text and were read off graphs. Information on variance was only reported graphically as standard error bars on the graphs. Therefore, we estimated the standard error for gingivitis, plaque and tooth staining, from the graphs and used this information to calculate the standard deviations of the mean scores so that we could include the data in the meta-analyses. Apart from tooth staining, adverse effects were not mentioned
Other bias	Unclear risk	Quote: "All readings were noted by the same investigator, with an intra-individual variation below 5%"
		Comment: the examiner reproducibility is good
		Quote: "The periodontal status at time of selection (day-14) as well as age range and gender distribution were very similar"
		Comment: no supporting data are provided
		The mean GI scores for the 3 groups at baseline appear similar. The mean DI scores for the 3 groups at baseline also appear similar (on the graph). The mean plaque scores in the placebo group at baseline (on the graph) appear higher than in the CHX group. It is difficult to assess the baseline balance as the data are not reported or commented on in the text

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Methods	Trial design: parallel (2 arms)
	Location: assumed to be at Case Western Reserve University School of Dentistry, Cleveland, Ohio, USA
	Number of centres: 1

Study duration: recruitment period not stated. Study duration and duration of rinsing was 3 months

Participants

Participants: quote: "Subjects for this study were selected from the Department of Orthodontics at Case Western Reserve University School of Dentistry, Cleveland, Ohio"

Inclusion criteria: quote: "Forty-four subjects qualified on the basis of the following criteria: (1) They were under-going full-banded edgewise extraction treatment with brackets on their anterior teeth and bands on their molars; (2) four premolars had been extracted; (3) they were at least 11 years of age and no more than 17 years of age; (4) evidence of gingivitis was present, but there was no evidence of periodontitis; (5) there was no evidence of decalcification on their teeth; (6) there was no known hypersensitivity to chlorhexidine; (7) there were no known medical problems or evidence of current antibiotic therapy; and (8) no anterior composites were present"

Exclusion criteria: not specifically stated but are implicit in the inclusion criteria

Baseline gingivitis: not reported

Age at baseline (years): Gp A: mean 14.88 (SD 1.78); Gp B: mean 14.78 (SD1.52); range 11-17

Gender: overall: male (42%), female (58%). Group data not specified

Number randomised: 36 (Gp A: 18, Gp B: 18) Number evaluated: 34 (Gp A: 16, Gp B: 18)

Interventions

Comparison: CHX mouthrinse versus placebo mouthrinse



Brightman 1991 (Continued)

Gp A (n = 18): CHX (Peridex) 0.12%, 1/2 ounce, twice daily, 30 seconds, 3 months

Gp B (n = 18): placebo (identical to test mouthrinse but without the CHX), 1/2 ounce, twice daily, 30 seconds, 3 months

Prophylaxis at baseline: quote: "Each of the subjects was given a thorough prophylaxis and instructed in oral hygiene procedures"

OHI: quotes: "Participants were provided with 3 toothbrushes (one for each month of the study) and 3 tubes of toothpaste." "All subjects were instructed to use the Bass technique of toothbrushing and to brush with the Crest toothpaste that was provided." "The patients were all instructed to use one toothbrush every month. They were to brush only with Crest toothpaste with fluoride, once in the morning after breakfast and once in the evening before bedtime. They were all instructed to brush a minimum of 3 minutes to ensure thorough brushing"

Non-supervised rinsing

Timing of mouthrinsing in relation to toothbrushing: after brushing

Postrinsing instructions: no liquid or food to be taken after using mouthrinse for at least 30 minutes

Outcomes

Gingivitis (gingival inflammation: GI Silness & Löe 1964 and 1967; gingival bleeding: Eastman Interdental Bleeding Index, Caton and Polson 1985); plaque (PI Löe & Silness 1964 and 1967); staining intensity and area (method created at Case Western Reserve University School of Dentistry, Department of Periodontics); assessed at 6 weeks, and 3 months follow-up

Funding

Funding: not reported

Notes

Sample size calculation: quote: "This study was designed to provide a minimal power of 87% for detecting a clinically important difference to be statistically significant at the 0.05 probability level"

Adverse effects: quote: "Both the chlorhexidine group and the placebo group showed significant increases in the degree of stain, but the chlorhexidine group had a more significant amount of stain"

Declarations/conflict of interest: none stated

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Two tables of random numbers, one for the male population and one for the female population, were used, and the subjects were thus assigned to one of two treatment groups"
		Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "These measures (use of random number tables) ensured that neither the investigators nor the patients knew whether participants were in the experimental group or the placebo group"
		Comment: the authors indicated that the participants were blind to group allocation (although the use of a random number table would not have ensured blinding)
		Quotes: "The placebo mouthrinse was identical to the experimental mouthrinse except that the placebo mouthrinse did not contain chlorhexidine The appearance and the taste of both mouthrinses were similar. They were both colored blue, mint flavored, and contained in amber-colored bottles" and



Brightman 1991 (Continued)		
		"The mean stain severity index for the chlorhexidine group showed significant ly more stain at 12 weeks (p<0.001) for all of the sites measured"
		Comment: despite the efforts undertaken by the investigators the increased staining in the CHX group is likely to have compromised the attempted partici pant blinding
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Quote: "These measures (use of random number tables) ensured that neither the investigators nor the patients knew whether participants were in the experimental group or the placebo group"
		Comment: the authors indicated that the investigators were blind to group allocation (although the use of a random number table would not have ensured blinding)
		Quote: "The GI and the PI were scored by the primary investigator, and the stain index and the bleeding index were scored by a second investigator, who was a senior dental student. This was done to minimise bias, so that the investigator who was scoring the GI and PI was not influenced by the degree of extrinsic stain, which is a common side effect of chlorhexidine"
		Comment: despite the efforts undertaken by the investigators to blind outcome assessment, the primary investigator, although not scoring the stain, would likely still be aware of increased staining in some individuals and this meant that the outcome assessors could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition after 3 months was 2/36 (5.6%). Attrition by group: 2/18 (11.1%) CHX; no losses from placebo group. Reasons for losses from CHX group: "inability to cooperate" 2 lost from CHX mouthrinse group after less than 1 week
		Comment: it is not known whether the "inability to cooperate" of the 2 participants lost from the CHX group were related to use of the CHX mouthrinse. However, the number of participants lost over the 3 months is small and unlikely to have affected the results
Selective reporting (re- porting bias)	High risk	Mean scores for all outcomes are reported graphically with 95% CI bars in the graphs. We estimated the mean scores and 95% CIs for all outcomes. We used the information on the 95% CIs to calculate the SDs of the mean scores so that we could include the data in meta-analyses. Apart from tooth staining, adverse effects were not mentioned
Other bias	Unclear risk	The outcomes were measured by 2 investigators one of which was a senior dental student. No information is provided on intraexaminer reproducibility (Interexaminer not relevant as the 2 examiners recorded different variables)
		Comment: not enough information is provided
		Mean scores for gingivitis, plaque and tooth stain as represented on graphs ap pear quite balanced. However, the statistical significance of the differences is not reported

Charles 2004

Methods

Trial design: parallel (3 arms - 2 arms are relevant to this review)

Location: assumed research facility, USA

Number of centres: assumed to be 1



Charles 2004 (Continued)	
	Study duration: recruitment period is not stated. Duration of study and duration of rinsing was 6 months

Participants

Participants: adults aged 20-57

Inclusion criteria: quote: "a minimum of 20 sound, natural teeth; a mean plaque index (PI) (Turesky et al 1970) of at least 1.95; a mean GI (Loe & Silness 1963) of at least 0.95"

Exclusion criteria: quote: "Teeth that were grossly carious, fully crowned or extensively restored, orthodontically banded, abutments, or third molars were not included in the tooth count. Subjects with gross oral pathology or who were taking antibiotic or antiinflammatory drugs were excluded"

Baseline gingivitis: (GI), Gp A: mean 1.35 (SE 0.04); Gp B: mean 1.31 (SE 0.04); Gp C: mean 1.27 (SE 0.03)

Age at baseline (years): Gp A: mean 31.4 (SD 9.4); Gp B: mean 32.0 (SD 6.6); Gp C: mean 32.2 (SD 6.3). Range 20-57 (all arms of study)

Gender (reported for completers): overall; male 39 (36%); female 68 (64%). Gp A: male 13 (36%), female 23 (64%); Gp B: male 12 (35%), female 22 (65%); Gp C: male 14 (38%), female 23 (62%)

Number randomised: 108 (Gp A: 36, Gp B: 34, Gp C: 38) Number evaluated: 107 (Gp A: 36, Gp B: 34, Gp C: 37)

Interventions

Comparison: CHX 0.12% (Peridex) versus EO (essential oil mouthrinse, Listerine antiseptic)* versus control (5% hydroalcohol negative control)

Gp A (n = 36): CHX (Peridex) 0.12%,15 mL, twice daily, 30 seconds, 6 months

*Gp B (n = 34): Listerine, 20 mL, twice daily, 30 seconds, 6 months. We excluded this arm from the risk of bias and analysis

Gp C (n = 38): control (5% hydroalcohol), 20 mL, twice daily, 30 seconds, 6 months

Prophylaxis at baseline: quote: "Following the baseline examination, each subject received a complete dental prophylaxis to remove all plaque, calculus, and extrinsic stain"

OHI: none given. Quote: "....subjects followed their usual oral hygiene and dietary habits." Soft nylon toothbrushes and fluoridated toothpaste were provided to all subjects and replenished periodically as need for the duration of the study

Quote: "One of the daily rinses was supervised on each weekday"

Postrinsing instructions: not reported

Timing of mouthrinsing in relation to toothbrushing: quote: "Rinsings were not done at time of tooth-brushing but at separate times"

Outcomes

Gingivitis (gingival inflammation: GI Löe & Silness 1963; gingival bleeding: % bleeding sites (GI scores 2 or 3 indicating bleeding)); plaque (TQH, Turesky 1970); extrinsic tooth stain (Lobene Discolouration Index, Lobene 1968); supragingival calculus (Volpe-Manhold Calculus Index, Manhold 1965, Volpe 1965, 1967, Barnett 1989), assessed at 3 and 6 months follow-up

Funding

Funding: the authors are affiliated to Dental Products Testing and Pfizer

Notes

Sample size calculation: quote: "This study was designed to provide a minimal power of 0.80 for detecting a statistically significant difference in plaque and gingivitis scores at the 0.05 probability level"

Adverse effects: during the course of the study no oral mucosal lesions that could be attributed to any of the test mouthrinses were observed"

Smokers: by group Gp A: 8/36 (22%); Gp B: 8/34 (24%); Gp C: 6/38 (16%) (differences are not statistically significant)



Charles 2004 (Continued)

Declarations/conflicts of interest: the authors are affiliated with Dental Products Testing and Pfizer. No explicit conflicts of interest were declared

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Risk	c of	bı	as

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quotes: "Subjects were randomised into three groups", "assigned to either one of two test groups or a negative control group according to a computer generated random code"
		Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor-	High risk	Quote: "One of the daily rinses was supervised on each weekday." No mention of blinding of study participants
mance bias) All outcomes		Comment: the study compared CHX with essential oil and control (5% hydroal-cohol) and different volumes of the rinses were used making participant and personnel blinding difficult to achieve. Lack of blinding could have affected participants' oral health behaviours and hence the outcome but direction of this potential bias is unclear
Blinding of outcome assessment (detection bias) All outcomes	High risk	Quotes: "observer blind" and "At 3 and 6 months, there was significantly more gingival region stain in the essential oil group (p<0.05) and the chlorhexidine group (p<0.001) compared with the control group"
		Comment: the higher level of staining in the CHX group meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear
Incomplete outcome data (attrition bias) All outcomes	Low risk	Only 1 participant was lost over the 6 months. Attrition by group 1/38 (2.63%) placebo group. No losses from CHX group. Reasons for losses: personal reasons. Lost after the 3-month exam and results for this individual were included in the 3-month exam
		Comment: the reason the individual was lost was not related to the intervention
Selective reporting (reporting bias)	High risk	We calculated standard deviations for gingivitis and plaque from the standard errors reported in the tables. We could not include data on % bleeding sites in meta-analyses as the data were reported for total number of sites rather than individuals. Information on the variance of the mean calculus and stain scores was not reported and those outcomes could not be included in meta-analyses
Other bias	Low risk	Quotes: "All examinations were conducted by a single, experienced dental examiner. Reliability was established for the gingival index (GI) with a κ statistic of 0.75, which indicates satisfactory calibration."
		"There were no statistically significant differences among the treatment groups with respect to age, gender, smoking" There were no statistically significant differences among the treatment groups in relation to the gingival and calculus indices. The plaque index was significantly lower in the control group than in the CHX group at baseline
		Comment: all participants had a baseline prophylaxis to remove all plaque, calculus and extrinsic stain. It can be assumed that they were similar with respect to plaque and stain at the start of the study



Methods	Trial design: parallel (4 arms - 2 arms are relevant to this review) Location: USA, exact location not reported. Setting appears to be a university department of periodontics				
	Number of centres: 1 Study duration: recruitment period not stated. Study duration and duration of rinsing was 6 months				
Participants	Participants: adults with chronic generalised gingivitis				
	Inclusion criteria: quote: "presence of more than 20 teeth, not more than 4 sites with probing depth (PD) deeper than 6 mm, bleeding on probing (BOP) frequency of higher than 30% at screening visit with no systemic diseases or medications influencing gingival inflammation"				
	Exclusion criteria: opposite of the inclusion criteria				
	Baseline gingivitis: (GI) not reported Age at baseline (years): mean overall 35.1; range 19-62. No breakdown by group is reported Gender: male 32%; female 68%. No breakdown by group is reported Number randomised: 125 (not reported by group) Number evaluated: 104 (Gp A: 24; Gp B: 21; Gp C: 32; Gp D: 27)				
Interventions	Comparison: brush+CHX (CHX) versus brush+CHX irrigation (Irr+CHX) ^a versus brush+water irrigation (Irr+H ₂ O) ^b versus brush				
	Gp A (n = 24): CHX (Peridex) 0.12%, 1:3 dilution (with water), twice daily, no duration reported, 6 months				
	a Gp B (n = 21): CHX 0.04% Irr, 400 mL, once daily, no duration reported, 6 months. We excluded this arm from our risk of bias and analysis				
	b Gp C (n = 32): Irr+H $_{2}$ O, 400 mL, once daily, no duration reported, 6 months. We excluded this arm from our risk of bias and analysis				
	Gp D (n = 27): brush, N/A				
	Note: Numbers stated above relate to completers. No figures were reported by group for those who were randomised				
	Prophylaxis at baseline: yes, but no details reported OHI: quote: "After baseline data collection, all patients received instructions on the use of a toothbrush and dental floss by watching a standardized video tape." "[after prophylaxis] further oral hygiene instructions specific to the treatment group [were provided]" Postrinsing instructions: not reported				
	Non-supervised rinsing				
	Timing of mouthrinsing in relation to toothbrushing: not reported				
Outcomes	Gingivitis (gingival inflammation: GI Löe 1963); gingival bleeding: BOP (automated probe (Floride probe) using 20 grams of force, Gibbs et al 1988); plaque (PI Silness & Löe 1964); assessed at 3 and 6 months follow-up. Probing depth and subgingival microbial samples were taken from sites presenting probing depths > 4 mm and BOP at baseline visit (not relevant to this review)				
Funding	Funding: quote: "This study was supported by Teledyne Water Pik, Denver, CO" (USA)				
Notes	Sample size calculation: not reported Adverse effects: quote: "Several of the patients who did not return to the scheduled examination visits reported an unacceptable taste associated with the chlorhexidine. No patient reported adverse reaction or side effects to the mouthrinse"				



Chaves 1994 (Continued)

Declarations/conflicts of interest: not reported

Additional information was obtained from Chaves 1993

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Patients were randomly assigned to one of four groups based on a random number table that was computer generated"
		Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias)	High risk	Quote: "The clinical examiners were blinded to the treatment groups, however dental assistants and hygienists were responsible for providing information and support to their respective plaque control group"
Alloutcomes		Comment: blinding of participants was not possible due to the differences in the intervention groups being compared. The dental assistants and hygienists were aware of the group allocations of participants
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "The clinical examiners were blinded to the treatment groupsThe clinical examiners did not have access to any of the clinical or compliance records during the study"
		Comment: tooth staining is likely to have been noticeable in the CHX group after 6 months, and in this situation the outcome assessor could have worked out which individuals used CHX and not be adequately blinded. However, tooth staining is not reported and there is not enough information upon which to base a judgement
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition after 6 months (all study arms) was 21/125 (16.8%). Attrition by group is not reported. Reasons for loss to follow-up: several of the patients who did not return to the scheduled exam visits reported an unacceptable taste associated with CHX
		Comment: several participants were lost from the 2 CHX groups because they did not like the taste of the rinse. Exact numbers lost from each group are not reported but the number of participants in the 2 CHX groups completing the trial seemed to be appreciably lower than in the 2 groups not using CHX. Assuming the missing participants in 1 group had a higher mean (e.g. gingivitis score) than those in the other groups, as the attrition rate increased, so would the mean difference between groups
Selective reporting (reporting bias)	High risk	Mean scores for all outcomes are reported graphically with SD bars in the graphs. We estimated the mean scores and SDs for all outcomes so that we could include the data in meta-analyses. Information on adverse effects is reported but no specific information relating to tooth staining is reported
Other bias	Unclear risk	Quote Chaves 1993: "The clinical examinations were carried out by 6 different examiners with each patient being evaluated by the same examiner at all time periodsprior to the initiation of the study, all examiners participated in group seminars and clinical sessions on the assessment of clinical parameters and the use of the Florida Probe and were compared to a single standard examiner. Although no attempt was made to evaluate inter or intra-examiner reproducibility, no statistically significant differences were found between the examiners when PD, BOP and GI were evaluated"



Chaves 1994 (Continued)

The groups appear reasonably balanced (on graph) with respect to GI, BOP and PI at baseline. The actual data at baseline are not reported

Comment: not enough information is provided

Corbet 1997

Methods

Trial design: parallel (2 arms)

Location: knitting factory assumed to be in Ho Tung, Guangdong, China

Number of centres: assumed to be 1

Study duration: recruitment period not stated. Duration of study 6 months, but data only reported for 3

months

Participants

Participants: quotes: "All were employed in a knitting factory in the area of Ho Tung, Province of Guangdong, People's Republic of China. However, most of the participants were migrant workers and originated from other Provinces such as Henan, Hebei and Sichuan" and "..a population with established gingivitis and abundant supra- and subgingival plaque and calculus"

Inclusion criteria: quote: "The subjects were admitted to the study if they were in good general health and had not been exposed to antibiotic therapy for at least 6 months. Furthermore, at least 20 teeth had to be present. No probing depth of pockets exceeded 6 mm and no loss of attachment of more than 2 mm was encountered"

Exclusion criteria: quote: "Subjects with known allergies, haematological disorders or diabetes mellitus were not allowed to participate. Also, pregnant women were excluded"

Gingivitis (at baseline for the completers): GI: overall mean 1.43 (SE 0.01); Gp A: mean 1.40 (SE 0.02); Gp B: mean 1.46 (SE 0.01)

Age at baseline (years): range 17-39. Overall mean 22.5 (4.95). Gp A: mean 23.0 (4.34); Gp B: mean 22.3 (5.26)

Gender: overall: male 13 (21.7%), female 47 (78.3%). Gp A: male 5 (25%), female 15 (75%); Gp B: male 8 (20%), female 32 (80%)

Number randomised: 60 (Gp A: 20; Gp B: 40)

Number evaluated: At 3-month measurement: 36 (Gp A: 13; Gp B: 23)

Interventions

Comparison: CHX versus placebo control

Gp A (n = 20): CHX (Peridex) 0.12%, 15 mL, twice daily, 45 seconds, 6 days per week for 6 months

Gp B (n = 40): placebo (identical in composition to the test rinse only without the active ingredient), 15 mL, twice daily, 45 seconds, 6 days per week for 6 months

Prophylaxis at baseline: quote: "A unique feature of the present study was that no pre-experimental prophylaxis was performed prior to the commencement of the supervised rinsing"

OHI: none given. Quote: "No attempts were made to influence the sparsely performed oral hygiene practices of the participants"

Postrinsing instructions: not reported

Supervision of rinsing: quote: "The supervision of the daily rinses by factory staff assured an almost 100% compliance and hence, it may be assumed that optimal conditions for a test of the therapeutical principle of chlorhexidine mouthrinses were provided... All rinsings were supervised by one of the supervisors in the factory. The rinsings were performed at 8.00 am and at 7.00 pm, and were timed for 45s." Note: there was a 2-week period during the study when participants were not supervised as they returned home during the Chinese New Year holiday. Quote: "During the 2 weeks of the Chinese New Year holiday (January 1995), mouthrinses were provided for the subjects to take to their homes and instructions on their usage was given by the supervisor"

Timing of mouthrinsing in relation to toothbrushing: not reported

Outcomes

Gingivitis (gingival inflammation: GI Löe & Silness 1963, Löe 1967; gingival bleeding: sites with scores 2 or 3 indicating bleeding); plaque (PI Silness & Löe 1964); stain (Discolouration Index Lang & Räber



Corbet 1997 (Continued)		us Surface Severity Index system (CSSI), Ennever et al 1961); assessed at 3 and 6 te: Quote: "this report is limited to the 3-month data"
Funding	Quote: "This study was supported by a grant for collaborative research of the Clinical Research Foundation (CRF) for the Promotion of Oral Health, University of Berne, Switzerland"	
Notes	Sample size calculation: not reported Adverse effects: not reported except for tooth staining and calculus Declarations/conflicts of interest: none stated	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quotes: "Randomised controlled clinical trial", "subjects were divided into two groups matched according to the mean gingival index"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "double-masked." The placebo rinse was identical in composition to the test rinse only without the active ingredient (CHX). Rinsings were supervised and blinding of the factory supervisors is not explicitly stated but it is considered unlikely that they were aware of the content of the rinses
		Quotes: "The overall increase in mean discoloration index was significantly higher in the Test than the Control group" and "There were significantly (p < 0.05) more surfaces scoring DI = 1, 2 or 3 in the Test group (39%) than in the control group (28%)"
		Comment: the participants in the CHX group may have noticed the increased staining and worked out which group they were in and this may have affected their oral health behaviours
Blinding of outcome assessment (detection bias) All outcomes	High risk	Quotes: "double-masked", "The same examiners calibrated for one particular index system scored the same clinical index in all subjects at baseline and after 3 months", "The overall increase in mean discoloration index was significantly higher in the Test than the Control group" and "There were significantly (p < 0.05) more surfaces scoring DI = 1, 2 or 3 in the Test group (39%) than in the control group (28%)"
		Comment: it is not clear if the "double-masked" included the examiners. Even if the masking refers to the examiners, it is likely that the examiners were aware of which participants were using CHX due to the increased tooth staining associated with CHX rinse
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition at 3 months was 24/60 (40%). Attrition by group: 7/20 (35%) CHX, 17/40 (42.5%) placebo. Reasons for losses (not broken down by group): the participants were mostly migrant workers from other provinces within China. 19 subjects failed to return from their homes after the Chinese New Year holiday and 5 were excluded due to pregnancy
		Quote: "The mean baseline scores for those subjects completing the study were not significantly different for the test and control group, respectively, although they were slightly different when compared with the baseline data of the original study population"



Corbet 1997 (Continued)		Comment: losses were very high and although the reasons for losses are not related to the intervention the proportion of those lost is so high that a distortion of the true intervention effect cannot be ruled out
Selective reporting (reporting bias)	High risk	We calculated standard deviations for gingivitis (GI), plaque, calculus and staining from the standard errors reported in the text. No information was reported on the variance for gingival bleeding (% sites scoring 0, 1, 2, or 3). No information regarding adverse effects apart from tooth staining and calculus were reported. This is a 6-month study but only 3-month data are reported
Other bias	Unclear risk	No information is provided on intra- and interexaminer reproducibility.
		Baseline characteristics of the participants were balanced for GI, PI, CSSI, DI, age and gender

de la Rosa 1988

Trial design: parallel (2 arms) Location: a boarding school in Monterrey, Mexico Number of centres: 1 Study duration: recruitment period not stated. Duration of study and duration of rinsing was 12 weeks			
Participants: boarding school boys Inclusion criteria: quote: "For acceptance into the study, they had to have some degree of plaque-induced gingivitis" Exclusion criteria: quote: "Those with gross oral neglect or pathoses that needed prompt care were not accepted" Baseline gingivitis: (Papillary Marginal Gingivitis Index; mean baseline score of all sites graded for individuals examined at 6 weeks) Gp A: mean 0.6933; Gp B: mean 0.6513 Age at baseline (years): range 8-18 Gender: 100% male Number randomised: not reported Number evaluated: 92 (Gp A: 46; Gp B: 46)			
Comparison: CHX versus placebo			
Gp A (completers n = 46): CHX 0.12%, 15 mL, twice daily, 30 seconds, 12 weeks			
Gp B (completers n = 46): placebo (composition not described), 15 mL, twice daily, 30 seconds, 12 weeks			
Prophylaxis at baseline: quote: "all subjects received a dental prophylaxis to remove any dental accretions"			
OHI: quote: "At the beginning of the studies, the subjects received soft toothbrushes and brushing instructions"			
Postrinsing instructions: not reported Supervised rinsing			
Timing of mouthrinsing in relation to toothbrushing: quote: "The evening rinsing was preceded by a one-minute toothbrushing with a sodium fluoride dentifrice, Crest"			
Gingivitis (gingival inflammation: Papillary Marginal Gingivitis Index, de la Rosa and Sturzenberger 1976, severity (mean score all sites) and occurrence (proportion of sites with a PMGI score ≥1)); plac (TQH, Turesky 1970); assessed at 6 and 12 weeks follow-up			
Funding: quote: "The work was supported by a grant of the Procter & Gamble Company, Cincinnati, OH"			
Sample size calculation: not reported			



de la Rosa 1988 (Continued)

Adverse effects: quotes: "Some mild epithelial desquamations were observed in some chlorhexidine subjects... There was no discomfort reported in association with these transient incidences in our study. Neither was there a consequence on oral health... there was an increase in extrinsic tooth stain and supragingival calculus in the chlorhexidine groups.." and "There was no lasting side effect on the oral soft tissue or the health of the users"

Declarations/conflicts of interest: potential conflict of interest: the second author is a Senior Dentist at the Procter and Gamble Company and the third author is an Associate Director, Peridex Research, Procter and Gamble Company

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "subjects were stratified by age intervals (<12 and >/= 12) and stratified by seven intervals of gingivitis scores ranging from <0.25 to >1.50. Within these strata, the subjects were distributed into the chlorhexidine or the place-bo groups at random"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	The authors report that the study is "double-blind". The comparison group rinsed with placebo rinse, no details are given regarding the formulation or presentation of the placebo
		Comment: it is not clear whether the 'supervisory personnel' who dispensed the toothpaste and mouthrinse and supervised the rinsing were blind. The authors report an increase in extrinsic tooth stain and supragingival calculus in the CHX group. The higher level of tooth staining and calculus in the CHX group meant that participants could have worked out which group they were in (or that they were in the active mouthrinse group) and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is unclear
Blinding of outcome assessment (detection bias) All outcomes	High risk	The authors report that the study is "double-blind". Quote: "Previous records were not available at subsequent examinations nor did examiners have any knowledge as to which treatment group a given subject belonged"
		Comment: the authors report an increase in extrinsic tooth stain and supragingival calculus in the CHX group. The higher level of staining and calculus in the CHX group meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Numbers at baseline not reported, so attrition from baseline cannot be calculated. Between 6 weeks and 3 months 4 (8%) participants were lost from the CHX group and 6 (11.5%) participants were lost from the placebo group
Selective reporting (reporting bias)	High risk	The SDs for the main outcome (mean gingivitis occurrence and severity) are not reported. The ancillary outcome (plaque) is mentioned but not fully reported possibly because the result was not statistically significant
Other bias	Unclear risk	No information is provided on intra- and interexaminer reproducibility.
		The groups appear balanced for gingivitis at baseline. The authors state that the baseline scores demonstrate comparability of the test and control groups. However, the statistical significance of the differences is not reported



Methods	Trial design: parallel (2 Location: a boarding so	arms) Chool in Monterrey, Mexico		
	Number of centres: 1			
	Study duration: recruit	ment period is not stated. Study duration and duration of rinsing was 10 weeks		
Participants	Participants: boarding school boys			
	Inclusion criteria: quote: "For acceptance into the study, they had to have some degree of plaque-induced gingivitis"			
	Exclusion criteria: quote: "Those with gross oral neglect or pathoses that needed prompt care were not accepted"			
	Baseline gingivitis: (Papillary Marginal Gingivitis Index; mean score of all sites graded at baseline for those examined at 10 weeks) Gp A: mean 0.4544; Gp B: mean 0.4263			
	Age at baseline (years): range 8-18 Gender: 100% male			
	Number randomised: r	not reported		
	Number evaluated: 99			
Interventions	Comparison: CHX versus placebo			
	Gp A (completers n = 49): CHX 0.12% (Peridex), 15 mL, twice daily, 30 seconds, 10 weeks			
	Gp B (completers n = 50): placebo (composition not described), 15 mL, twice daily, 30 seconds, 10 weeks			
	Prophylaxis at baseline: quote: "all subjects received a dental prophylaxis to remove any dental accretions"			
	OHI: quote: "At the beginning of the studies, the subjects received soft toothbrushes and brushing in-			
	structions" Postrinsing instructions: not reported			
	Supervised rinsing			
	Timing of mouthrinsing in relation to toothbrushing: quote: "The evening rinsing was preceded by a one-minute toothbrushing with a sodium fluoride dentifrice, Crest"			
Outcomes	Gingivitis (gingival inflammation: Papillary Marginal Gingivitis Index, de la Rosa and Sturzenberger 1976; severity (mean score all sites) and occurrence (proportion of sites with a PMGI score ≥1); assessed at 10 weeks follow-up			
	Plaque (TQH, Turesky 1970)			
Funding	Funding: quote: "The work was supported by a grant of the Procter & Gamble Company, Cincinnati, OH"			
Notes	Sample size calculation			
	Adverse effects: quotes: "Some mild epithelial desquamations were observed in some chlorhexidine subjects There was no discomfort reported in association with these transient incidences in our study. Neither was there a consequence on oral health there was an increase in extrinsic tooth stain and supragingival calculus in the chlorhexidine groups" and "There was no lasting side effect on the oral soft tissue or the health of the users"			
	Declarations/conflicts of interest: potential conflict of interest: the second author is a Senior Dentist at the Procter and Gamble Company and the third author is an Associate Director, Peridex Research, Procter and Gamble Company			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Unclear risk	Quote: "subjects were stratified by age intervals (<12 and >/= 12) and stratified by seven intervals of gingivitis scores ranging from <0.25 to >1.50. Within		



de la Rosa 1988b (Continued)		these strata, the subjects were distributed into the chlorhexidine or the place- bo groups at random"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	The authors report that the study is "double-blind". The comparison group rinsed with placebo rinse, no details are given regarding the formulation or presentation of the placebo
		Comment: it is not clear whether the 'supervisory personnel' who dispensed the toothpaste and mouthrinse and supervised the rinsing were blind. The authors report an increase in extrinsic tooth stain and supragingival calculus in the CHX group. The higher level of tooth staining and calculus in the CHX group meant that participants could have worked out which group they were in (or that they were in the active mouthrinse group) and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is unclear
Blinding of outcome assessment (detection bias) All outcomes	High risk	The authors report that the study is "double-blind". Quote: "Previous records were not available at subsequent examinations nor did examiners have any knowledge as to which treatment group a given subject belonged"
		Comment: the authors report an increase in extrinsic tooth stain and supragingival calculus in the CHX group. The higher level of staining and calculus in the CHX group meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Numbers at baseline not reported, so attrition from baseline cannot be calculated
Selective reporting (reporting bias)	High risk	The SDs for the main outcome (mean gingivitis occurrence and severity) are not reported. The ancillary outcome (plaque) is mentioned but not fully reported possibly because the result was not statistically significant
Other bias	Unclear risk	No information is provided on intra- and interexaminer reproducibility.
		The groups appear balanced for gingivitis at baseline. The authors state that the baseline scores demonstrate comparability of the test and control groups. However, the statistical significance of the differences is not reported

Eaton 1997

Methods	Trial design: parallel (2 arms) Location: suburban general dental practices in the South of England at Staines, Ripley, Lydd, Leigh on Sea, and West Malling, UK Number of centres: 5 Study duration: recruitment period is not stated. Study duration and duration of rinsing was 3 months
Participants	Participants: healthy subjects, existing patients of the 5 general dental practices Inclusion criteria: quote: "Inclusion criteria were: a willingness to participate in the study, a minimum of 16 natural teeth, the presence of chronic gingivitis but no pocket probing depths in excess of 4 mm or radiographic evidence of bone loss, no need for urgent treatment, between the ages of 18 and 65



Eaton 1997 (Continued)

years, in good health and a willingness to refrain from using any mouthrinses other than those provided in the study"

Exclusion criteria: quote: "Exclusion criteria were: acute gingivitis or other periodontal conditions, existing lesions of the oral mucosa, the receipt of antibiotics within 14 days of the baseline examination, history of diabetes, hepatitis B, tuberculosis or other infective or metabolic diseases, allergy to oral care products, the use of chlorhexidine or other antibacterial mouthrinses within three months of the start of the study, pregnancy or childbirth within the previous year, partial dentures, clinically unacceptable restorations, smoking more than 10 cigarettes per day and the receipt of any medication with a history of potential to alter gingival response"

Baseline gingivitis: (Modified GI) Gp A: mean CHX 0.56 (SD 0.4); Gp B: mean 0.54 (SD 0.4) Age at baseline (years): Gp A: mean 34.8 (SD 11.4); Gp B: mean 34.9 (SD 12.7); range of 18-65

Gender: overall: male 41 (34%), female 80 (66%).

Number randomised: 121 (Gp A: 60; Gp B: 61). 24 subjects were at 4 practices each and 25 subjects were

at the fifth practice

Number evaluated: 98 (Gp A: 48; Gp B: 50)

Interventions

Comparison: CHX versus control (placebo in aqueous solution)

Gp A (n = 60): CHX (Peridex) 0.12%, 15 mL, twice daily, 30 seconds, 3 months

Gp B (n = 61): control, 15 mL, twice daily, 30 seconds, 3 months

Prophylaxis at baseline: quote: "All were then (after baseline examination) given a dental prophylaxis... After initial baseline prophylaxis, patients were given no further prophylaxes during the study" OHI: quote: "Patients were instructed to use only these oral hygiene items [medium, multitufted toothbrush and sodium fluoride containing toothpaste] throughout the study. They were asked to brush their teeth as normally"

Postrinsing instructions: not reported

Non-supervised rinsing

Timing of mouthrinsing in relation to toothbrushing: mouthrinsing after toothbrushing, after breakfast and last thing at night

Outcomes

Gingivitis (Modified Gingival Index, Eaton 1997); plaque (PI Silness & Löe 1964); assessed at 6 and 12 weeks follow-up

Funding

Funding: Procter and Gamble (Health & Beauty Care) Europe Ltd sponsored the study

Notes

Sample size calculation: quote: "Sample size calculations were based on detecting a difference of 20% in gingivitis between placebo and ChD group if it existed at a 2-tailed significance level of 5% with 95% power"

Participants included smokers (smoking up to and including 10 cigarettes a day)

Adverse effects: participants in the CHX group left the study due to staining of teeth (4), taste (1) and adverse mucosal reaction (1)

Declarations/conflicts of interest: potential conflict of interest with author E Zak (Procter and Gamble Health & Beauty Care Europe). No other declarations other than that of funding (above)

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "mouthrinseswere dispensed on a randomised double-blind basis, according to a random code"
		Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided



Eaton 1997 (Continued)		
Blinding of participants and personnel (perfor- mance bias)	High risk	The authors state that the study is double-blind. The CHX and placebo mouthrinses were distributed in coded but otherwise indistinguishable containers
All outcomes		Comment: tooth staining from the CHX was noted as a reason for loss to follow-up in the study and may have been visible to the participants in the test group. This meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is unclear
Blinding of outcome as-	High risk	The authors state that the study is double-blind
sessment (detection bias) All outcomes		Comment: although the authors indicate that the clinical examiners were blinded to the treatment groups, tooth staining from the CHX was noted as a reason for loss to follow-up in the study. This meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition at 12 weeks 23/121 (19%). Attrition by group: 12/60 (20%) CHX, 11/61 (18%) placebo. Reasons for loss to follow-up: CHX: 5 withdrawn because they had taken antibiotics during the study for non-oral reasons, a further 6 for factors relating directly to the mouthrinses (4 toothstaining, 1 taste, 1 adverse mucosal reaction), 1 was unwilling/unable to continue in the study. Placebo: 8 withdrawn because they had taken antibiotics during the study for non-oral reasons, 3 for miscellaneous reasons including using a different mouthrinse, heavy smoking and unspecified unwillingness to continue
		Comment: half of those lost from the CHX group were lost due to reasons directly related to using the CHX mouthrinse. Most losses in the placebo group were protocol violations. Although numerically the dropouts are similar, the reasons are different and may bias in favour of CHX
Selective reporting (reporting bias)	High risk	Adverse effects including tooth staining were reported in relation to with- drawals from the study only but are considered important outcomes and should have been reported for all participants where relevant
Other bias	Low risk	Interexaminer κ scores 0.78-0.85 for PI and 0.73-0.94 for the Modified Gingival Index. Examiners maintained κ scores 0.51-0.90 for PI and 0.73-1.00 for the Modified Gingival Index during the 12 months required to complete the study. No mention of intraexaminer calibration

Emling 1992

Methods	Trial design: parallel (5 arms - 4 arms relevant to this review) Location: assumed to be at School of Dental Medicine, University of Pennsylvania, Philadelphia, PA, USA Number of centres: 1 Study duration: recruitment period is not stated. Study duration and duration of rinsing was 2 months
Participants	Participants: healthy subjects regardless of race or sex
	Inclusion criteria: minimum of 20 crowned natural teeth; quote: "All subjects reported that they were free from any known reaction or sensitivity to any dentifrices, mouthrinses, or oral hygiene products. The female subjects entered into the study gave verbal assurance that they were not pregnant All subjects agreed to refrain from any routine dental treatment or prophylaxis other than emergency pro-

Groups are balanced for age, sex, gingivitis and plaque at baseline



Emling 1992 (Continued)

cedures. They also agreed to use only the assigned treatments, and to comply with daily use as directed by the instructions given to them at the time of the product distribution"

Exclusion criteria: quote: "No subjects had orthodontic bands or diseases present of the hard tissues (osteomyelitis, fractures, cysts, odontomas, etc). There were no neoplastic growths in the oral cavity, advanced periodontal diseases, or diseases of the buccal mucosa, tongue, or floor of the mouth such as abscesses, ulcerations, hyperplasia, hypertrophy, or granulomas. None of the subjects entered into the study were currently using antibiotics or had used any antibiotics at least two weeks prior to the onset of the study"

Baseline gingivitis is reported separately for buccal and lingual surfaces

Baseline gingivitis for buccal surfaces: Gp A: mean 1.04 (0.59); Gp B: mean 0.92 (0.48); Gp C: mean 0.95 (0.62); Gp D: mean 1.24 (0.68); Gp E: mean 1.00 (0.48)

Baseline gingivitis for lingual surfaces: Gp A: mean 1.26 (0.56); Gp B: mean 1.18 (0.61); Gp C: mean 1.18 (0.55); Gp D: mean 1.26 (0.55); Gp E: mean1.07 (0.51)

Age at baseline (range): 18-60

Gender: not stated. Subjects were accepted without reference to race or sex

Number randomised: 65 (assumed to be 13 in each group)

Number evaluated: not reported

Interventions

Comparison: (Crest + CHX) versus (Rembrandt + CHX) versus (Crest + CHX until week 4 then Rembrandt + CHX from week 4 to week 8)* versus (Crest + placebo) versus (Rembrandt + placebo)

Gp A (n = 13): CHX (Peridex) 0.12% + Crest; amount not reported; frequency not reported; rinse time not reported, 8 weeks

Gp B (n = 13): CHX (Peridex) 0.12% + Rembrandt; amount not reported; frequency not reported; rinse time not reported, duration not reported

*Gp C (n = 13): CHX (Peridex) 0.12% + Crest (baseline to week 4); CHX (Peridex) 0.12% + Rembrandt; amount not reported; frequency not reported; rinse time not reported; duration not reported. We excluded this arm from our data extraction, risk of bias and analysis

Gp D (n = 13): placebo + Crest; amount N/A; frequency not reported; rinse not reported; rinse time not reported; duration not reported

Gp E (n = 13): placebo + Rembrandt; amount N/A; frequency not reported; rinse not reported; rinse time not reported; duration not reported

Prophylaxis: not reported

OHI: quote: "At each examination time, subjects reported to the study having not brushed their teeth

for a period of 10-12 hours"

Postrinsing instructions: not reported

Non-supervised rinsing

Timing of mouthrinsing in relation to toothbrushing: after toothbrushing

Outcomes

Gingivitis (gingival inflammation: GI Löe & Silness 1963); plaque (TQH,Turesky 1970); extrinsic tooth stain (area and intensity) (DI, Lobene 1968); supragingival calculus (Volpe-Manhold Calculus Index, Volpe-Manhold 1965); assessed at 2, 4, 6, and 8 weeks follow-up

Funding

Funding: not reported

Notes

Sample size calculation: not reported

Adverse effects: quote: "Eight subjects using Peridex complained of burning and/or irritation of the gingiva or palatal mucosa. No other changes of the gingiva were observed during the study." Increases in stain intensity and area were noted in all CHX rinse groups during the study. Groups C and D had significant increases in mean calculus scores

Declarations/conflicts of interest: none stated. First author works for International Dental Research Ltd, Moorestown, NJ, USA



Emling 1992 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence genera-	Unclear risk	Quote: "Subjects were randomly assigned to one of 5 groups"
tion (selection bias)		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quotes: "The study was conducted in a double-blind manner. Neither the subjects nor the clinical examiners were aware of the product assignments at any time during the study", and "The dentifrices and the mouthrinses were all in similar packages labelled only with a subject's assigned number and the investigators emergency day and night phone numbers"
		Increases in stain intensity and area were noted in all CHX rinse groups during the study
		Comment: The higher level of tooth staining in the CHX groups meant that participants could have worked out which group they were in (or that they were in 1 of the active mouthrinse groups) and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is unclear
Blinding of outcome assessment (detection bias) All outcomes	High risk	Quotes: "The study was conducted in a double-blind manner. Neither the subjects nor the clinical examiners were aware of the product assignments at any time during the study", and "The dentifrices and the mouthrinses were all in similar packages labelled only with a subject's assigned number and the investigators emergency day and night phone numbers"
		Increases in stain intensity and area were noted in all CHX rinse groups during the study
		Comment: the higher level of tooth staining in the CHX groups meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	No information is provided
Selective reporting (reporting bias)	Low risk	All outcomes stated in the methods section are adequately reported in the results section. For gingivitis, plaque, stain intensity and stain area, means and SDs for buccal and lingual surfaces were combined for each group. Then, CHX + Crest and CHX + Rembrandt groups were combined into a CHX + toothpaste group and placebo + Crest and placebo + Rembrandt were combined into a placebo + toothpaste group to be included in meta-analyses
Other bias	Unclear risk	No information is provided on number of examiners or intra- and interexaminer reproducibility
		Baseline GI, plaque, calculus and stain indices were not statistically significantly different between the groups at baseline



Methods	Trial design: parallel (3 arms - 2 arms are relevant to this review)
	Location: Department for Operative Dentistry, Johannes Gutenberg University of Mainz, Germany
	Number of centres: 1
	Study duration: recruitment period not stated. Study duration and duration of rinsing was 4 weeks
Participants	Participants: quotes: "soldiers stationed in Mainz, Germany" and "students from the dental clinic of Johannes Gutenberg University Mainz, Germany"
	Inclusion criteria: quote: "Each volunteer had gingivitis or slight periodontitis"
	Exclusion criteria: quote: "General exclusion criteria were systemic diseases and long-term medications"
	Baseline gingivitis (GI, Löe & Silness): Gp A: mean 1.21 (SD 0.68); Gp B: mean 1.11 (SD 0.88); Gp C: mean 1.09 (SD 0.71)
	Age of completers (years): Gp A: mean 28.4 (SD 8.5); Gp B: mean 32.2 (SD 12.2); Gp C: mean 31.3 years (SD 9.5). Overall age range 18-58. Gp A: range 18-50; Gp B: range 22-58; Gp C: range 22-54
	Gender: not reported
	Number randomised: 101 (not reported by group)
	Number evaluated: 90 (Gp A: 33; Gp B: 29; Gp C: 28)
Interventions	Comparison: CHX versus Hexetidine* versus placebo
	Gp A (completers n = 33): CHX 0.1%, 15 mL, twice daily, 30 seconds, 4 weeks
	* Gp B (completers n = 29): Hexetidine 0.1%, 15 mL, twice daily, 30 seconds, 4 weeks. We excluded this arm from the risk of bias and analysis
	Gp C (completers n = 28): placebo, 15 mL, twice daily, 30 seconds, 4 weeks
	Prophylaxis at baseline: quote: "Each subject's teeth were professionally cleaned afterwards"
	OHI: quote: "The subjects were given detailed instructions on oral hygiene. Each subject received a new toothbrush (Oral B, P35, Gilette/Oral B) and several trial packages of a standardized toothpaste (Emex, GABA). The subjects were asked to brush their teeth with the new toothbrush and toothpaste a ter breakfast and dinner each day"
	Non-supervised rinsing
	Timing of mouthrinising in relation to toothbrushing: after toothbrushing
Outcomes	Gingivitis (gingival inflammation: GI Löe & Silness 1967; gingival bleeding: Bleeding Index, van der Wei jen 1994); plaque (Approximal Plaque Index, Lange 1977); extrinsic tooth stain (Discolouration Index, no reference provided but as described it appears to be the severity aspect of the Discolouration Index (Lobene 1968)); assessed at 2 weeks (GI and DI) and at 4 weeks (all indices). Volunteers were questioned about changes to taste, oral sensitivity, taste of mouthrinse, and about their opinion of the mouthrinse (the latter only asked after 4 weeks); assessed at 4 weeks follow-up
Funding	Not reported
Notes	Sample size calculation: not reported
	Adverse effects: quote: "A total of 28 of 90 subjects reported obvious changes in their sense of taste after using the mouthrinse. The number of subjects using mouthrinse B reported the most incidences of a change in their sense of taste. A total of 18 of 29 subjects (62.1%) complained about changes in their sense of taste. Only 8 of 33 subjects (24.2%) using mouthrinse A reported a change in the sense of taste.



Ernst 2005 (Continued)

In the group using mouthrinse C, 2 out of 28 subjects (7.1%) complained about obvious changes in their sense of taste... A relatively high number of subjects using mouthrinse B complained about oral mucosa sensitivities. Two subjects out of this groups showed such severe changes in terms of intensive red, inflamed areas at the intermediate check-up that mouthrinse use was discontinued immediately... In group 1 (using mouthrinse A), 5 of 33 subjects (15%) showed sensitivities of the oral mucosa... A total of 24% of the subjects in group 1 suffered from changes in their sense of taste." Tooth staining was significantly higher in the CHX group compared to the 2 other groups

Smokers: Gp A 17/33 (51.5%); Gp B 12/29 (41.4%); Gp C 8/28 (28.6%)

Declarations/conflicts of interest: none stated

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quotes: "divided randomly into two groups" and "the solutions were handed out randomly to each subject"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Low risk	Quote: "All mouthrinses were coded and placed into identical bottles. The bottles were labelled A, B, or C by the pharmacy at the Universtiy Clinic Mainz, Mainz, Germany. The solutions were handed out randomly to each subject"
		Comment: the details suggest that concealment of allocation sequence was adequate
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quotes: "Double-blind" and "All mouthrinses were coded and placed into identical bottles. The bottles were labelled A, B or C by the pharmacy at the University Clinic Mainz, Mainz, Germany each solution contained the same application and dosage instructions. The bottles were not decoded until after all the follow-up examinations and final statistical analyses had been completed." The placebo contained propyleneglycol, glycerol, gentian-tincture, pigment E 124 and purified water
		Comment: the CHX group developed more tooth staining than the placebo group wwhich meant that participants could have worked out which group they were in (or that they were in 1 of the active mouthrinse groups) and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is unclear
Blinding of outcome assessment (detection bias) All outcomes	High risk	Quotes: "Double-blind" and "All mouthrinses were coded and placed into identical bottles. The bottles were labelled A, B or C by the pharmacy at the University Clinic Mainz, Germany each solution contained the same application and dosage instructions. The bottles were not decoded until after all the follow-up examinations and final statistical analyses had been completed All exams were done by clinicians at the Department of Operative Dentistry, Johannes Gutenberg University"
		Comment: the CHX group developed more tooth staining than the placebo group which meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear
Incomplete outcome data (attrition bias)	Unclear risk	Attrition at 5 weeks 11/101 (10.9%). Not broken down by group
All outcomes		Reasons for losses: non-compliance



Ernst 2005 (Continued)		
Selective reporting (reporting bias)	Low risk	All outcomes stated in the methods section are adequately reported in the results section. Adverse effects including tooth staining are reported in detail
Other bias	Unclear risk	No information is provided on number of examiners, intra- and interexaminer reproducibility or training of examiners
		The mean scores for GI, Bleeding Index, Approximal Plaque Index and DI appear similar at baseline. Mean age of completers and age range appear similar. There were a higher proportion of cigarette smokers in Gp A compared to Gp C. Gender is not reported

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Interventions	Comparison: (CHX + SRP) versus (placebo control + SRP)
	Number evaluated: 29 (Gp A: 14; Gp B: 15)
	Number randomised: 30 (Gp A: 15; Gp B: 15)
	Gender: overall: male 13 (45%); female 16 (55%). Gp A: male 5 (36%), female 9 (64%); Gp B: male 8 (53%), female 7 (47%)
	Age at baseline (years): range not stated. Gp A: mean 48.5 (+/-8.8); Gp B: mean 42.1 (+/-6.5)
	Baseline gingivitis: (gingival gleeding yes/no), Gp A: mean 33.02 (+/-11.37); Gp B: mean 40.17 (+/-19.46)
	Exclusion criteria: quote: "The exclusion criteria were as follows: previous periodontal therapy, pregnancy, nursing, smokers, any systemic condition that could affect the progression of periodontal disease or that required antibiotic coverage for routine dental therapy, allergy to CHX and antibiotic therapy in the previous 6 months"
	Inclusion criteria: quote: "The study included subjects >30 years of age with at least 15 teeth and a minimum of six teeth with at least one site with PD between 5 and 7 mm and CAL between 5 and 10 mm"
Participants	Participants: adults referred to the periodontal clinic of Guarulhos University, Brazil
	Study duration: the recruitment period is not stated. Study duration is 84 days (-21 days to +63 days post-therapy. Duration of rinsing was 63 days (-21 days to +42 days post-therapy)
	Number of centres: assumed to be 1
	Location: Periodontal Clinic of Guarulhos University, Guarulhos, Sao Paulo, Brazil
Methods	Trial design: parallel (2 arms)

Gp A (n = 14): CHX 0.12% + SRP, 15 mL, twice daily, 1 min, 63 days

Gp B (n = 15): placebo control (composition not described) + SRP, 15 mL, twice daily, 1 min, 63 days

Prophylaxis at baseline: quote: "During the initial phase, subjects received... full-mouth supragingival scaling"

OHI: quote: "During the initial phase, subjects received... instruction in proper home-care techniques." All participants received scaling and root planing "completed in a maximum of six appointments lasting approximately 1h each." The treatment was completed in 21 days. The CHX rinsing began with the SRP and continued for 42 days after the end of this therapy

Non-supervised rinsing

Timing of mouthrinsing in relation to toothbrushing: after toothbrushing



Faveri 2006 (Continued)		
Outcomes	Gingivitis (gingival blee sessed at 42 and 63 day	eding 0/1, BOP 0/1); plaque (plaque accumulation 0/1); suppuration (0/1); as- ys postSRP
	Probing depth (mm); C to this review)	AL (mm) assessed and microbiological monitoring also carried out (not relevant
Funding	Quote: "This study was	supported by Guarulhos University, Guarulhos, SP-Brazil and FAPESP"
Notes	would provide an 80% probing pocket depth	n: quote: "The sample size calculation determined that 14 subjects per group power to detect a true difference of 1.0 mm between test and placebo using reduction in pockets $>$ /6 mm as the primary outcome variable. This calculation illed comparison of α = 0.05"
	Adverse effects: quote:	"No severe adverse effects were reported by any of the subjects"
	Non-smokers	
	Declarations/conflicts	of interest: none stated
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "During the enrollment visit, each subject was given a code number, and a computer-generated table was used to have them randomly assigned to receive one of the two proposed treatments"
		Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	High risk	Quote: "During the enrollment visit, each subject was given a code number, and a computer-generated table was used to have them randomly assigned to receive one of the two proposed treatments. The coordinator of the study assigned participants to their groups"
		Comment: it is likely that the study co-ordinator was aware of the upcoming assignments when randomising the subjects
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	The study is described as single-blind. The university pharmacy prepared the placebo and CHX rinses and packaged them in opaque plastic tubes. The study co-ordinator marked the code number of each subject on a set of 9 tubes. The co-ordinator gave the coded tubes to the 2 examiners who at no time during the study had access to information about the contents of the tubes or the assignment of subjects to groups. Quote: "All study personnel, including the biostatistician and participants, were blinded to treatment assignment for the duration of the study"
		Comment: blinding was unlikely to be maintained if there was more staining and other minor adverse effects in the CHX group (it is stated that no serious adverse effects were reported by any of the subjects). However, as these outcomes were not reported, it is unclear whether blinding was maintained
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	The study is described as single-blind. The university pharmacy prepared the placebo and CHX rinses and packaged them in opaque plastic tubes. The study co-ordinator marked the code number of each subject on a set of 9 tubes. The co-ordinator gave the coded tubes to the 2 examiners who at no time during the study had access to information about the contents of the tubes or the assignment of subjects to groups. Quotes: "All study personnel, including the biostatistician and participants, were blinded to treatment assignment for the duration of the study" and "One examiner performed all clinical measurements and treatment was performed by the second examiner"



Faveri 2006 (Continued)		
		Comment: blinding was unlikely to be maintained if there was more staining and other adverse effects in the CHX group (it is stated that no serious adverse effects were reported by any of the subjects). However, as these outcomes were not reported, it is unclear whether blinding was maintained
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition at 12 weeks: 1/30 (3.3%). 1 subject lost from the CHX group because they had taken an antibiotic during the course of the study
Selective reporting (reporting bias)	High risk	All outcomes are reported graphically with bars representing a measure of variance in the graphs. We obtained additional outcome data from the first author. Toothstaining is considered to be an important side effect of CHX mouthrinsing but this is not reported
Other bias	Low risk	Quote: "The clinical monitoring was performed by two trained and calibrated examiners One examiner performed all clinical measurements and treatment was performed by the second examiner"
		The examiners were calibrated to provide reproducible measurements for probing depth and CAL. Inter- and intraexaminer variability is reported for probing depth and CAL measurements
		"The demographic and clinical baseline characteristics were similar between the two groups (P > 0.05)" $$

		n	

Methods

Trial design: parallel (3 arms - 2 arms relevant to this review)

Location: Periodontal Clinic of Guarulhos University, Guarulhos, Sao Paulo, Brazil

Number of centres: 1

Study duration: recruitment period is not stated. Study duration was 6 months and 21 days. Duration of rinsing was 63 days

Participants

Participants: quote: "population referred to the Periodontal Clinic of Guarulhos University"

Inclusion criteria: quote: "subjects with untreated previously periodontal disease... good general health and were diagnosed with generalized chronic periodontitis based on the current classification of the American Academy of Periodontology. The inclusion criteria were as follows: >30 years of age, at least 15 teeth, minimum of six teeth with at least one interproximal site with PD between 5 and 7 mm and CAL between 5 and 10 mm, at least 30% of the sites with PD and CAL >/5 mm and presence of bleeding on probing (BOP)"

Exclusion criteria: quote: "The exclusion criteria were as follows: previous subgingival periodontal therapy, smoking, pregnancy, nursing, systemic diseases that could affect the progression of periodontal disease (e.g. diabetes and immunological disorders), long-term administration of anti-inflammatory medication, need for antibiotic coverage for routine dental therapy, antibiotic therapy in the previous 6 months and allergy to CHX"

Baseline gingivitis: (gingival bleeding) Gp A: mean 29.6 (+/- 13.8); Gp B: mean 34.9 (+/- 19.4); Gp C: mean 40.3 (+/- 22.5)

Age at baseline (years): range not stated. Gp A: mean 45.1 (+/- 9.6); Gp B: mean 39.6 (+/- 6.1); Gp C: mean 42.5 (+/- 7.3)

Gender: overall: male 24 (40%), female 36 (60%). Gp A: male 8 (40%), female 12 (60%); Gp B: male 7 (35%), female 13 (65%); Gp C: male 9 (45%), female 11 (55%)



Fere	s 2009	(Continued)
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Number randomised: 60 (Gp A: 20; Gp B: 20; Gp C: 20)

Number evaluated: 60 (Gp A: 20; Gp B: 20; Gp C: 20)

Interventions

Comparison: (SRP + CHX) versus (SRP + placebo) versus (SRP, professional plaque control and placebo)*

All participants received full-mouth SRP under local anaesthesia in 4-6 appointments of 1 hour each. SRP was completed 21 days after baseline. Participants started rinsing at the beginning of the study and continued rinsing for 42 days after the end of the SRP

Gp A (n = 20): SRP + CHX 0.12%, 15 mL, twice daily, 1 min, 63 days

Gp B (n = 20): SRP + placebo, 15 mL, twice daily, 1 min, 63 days

*Gp C (n = 20): SRP, professional plaque control and placebo, 15 mL, twice daily, 1 min, 63 days (data not included in the review). We excluded this arm from our risk of bias and analysis

Prophylaxis at baseline: quote: "During the initial phase, all subjects received... full-mouth supragingival scaling"

OHI: quote: "During the initial phase, all subjects received instruction on proper homecare techniques"

Non-supervised rinsing

Timing of mouthrinsing in relation to toothbrushing: after toothbrushing

Outcomes

Outcomes measured at 2 and 6 months post-therapy (i.e. first follow-up examination occurred approx 20 days after rinsing had ceased at approx 83 days after baseline)

Gingivitis: (gingival bleeding (0/1); BOP (0/1)); visible plaque (0/1) measured at 2 and 6 months post-therapy

Suppuration (0/1); PD (mm); CAL (mm); also microbiological monitoring; assessed at 2 and 6 months post-therapy (not relevant to this review)

Funding

This study was supported in part by research grants from the state of Sao Paulo research foundation and FOGARTY from the National Institutes of Health (NIH), USA

Notes

Sample size calculation: quote: "The ideal sample size to assure adequate power for this clinical trial was calculated considering differences of at least 1 mm between groups for clinical attachment level (CAL) in initially deep periodontal sites [probing depth (PD) > 7 mm]... Based on these calculations, it was defined that 16 subjects per group would be necessary to provide an 80% power with an [alpha] of 0.05"

Adverse effects: quote: "No severe adverse effects were reported by any of the subjects. Two subjects from the CHX group reported adverse events during the study, such as tooth staining and an unpleasant taste"

Non-smokers

Declarations/conflicts of interest: none stated

Notes: intention-to treat analysis. The last available recordings were carried forward to represent all subsequent time points of evaluation. Data from professional plaque control group are not included in the review

Risk of bias

Bias

Authors' judgement Support for judgement



Feres 2009	(Continued)
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Random sequence generation (selection bias)

Low risk

Quote: "Each subject was given a code number at the enrolment visit and the study coordinators used a computer-generated table (in blocks of three) to randomly allocate them to one of three therapeutic groups"

Comment: adequate method of random sequence generation

Allocation concealment (selection bias)

Low risk

Quotes: "Each subject was given a code number at the enrolment visit and the study coordinators used a computer-generated table (in blocks of three) to randomly allocate them to one of three therapeutic groups." The mouthrinses were prepared and dispensed in opaque plastic tubes by the Guarulhos university pharmacy. "The study coordinators marked the code number of each subject on a set of 9 tubes according to the therapy assigned"

Additional information provided by the study authors: "Allocation concealment was assured by placing the numbered bottles in indistinguishable plastic bags with the same numbering. The numbered bags with the medication/placebo were given to the therapists by a clinical assistant, following the sequence of numbers from 1 to 60. This assistant and the clinicians had no access to the randomization list, assuring the allocation concealment. The study coordinators had access to the list but did not participate on the clinical treatment/examination of the patients"

Comment: the details suggest that concealment of allocation sequence was adequate

Blinding of participants and personnel (performance bias) All outcomes Unclear risk

CHX and placebo rinse was packaged in opaque plastic tubes with the patients code number written on the tubes. "The examining researchers had no access to information about the contents of the tubes or assignment of subjects to the three therapies. All study personnel, including the biostatistician and participants were not aware of the treatment assignments for the duration of the study"

2 subjects from the CHX group reported adverse events during the study, such as tooth staining and an unpleasant taste

Comment: blinding of all participants is not possible in this study because of differences between the interventions (subjects in the professional plaque control group attended the clinic twice a week and the others did not) although the CHX and placebo groups could be considered blind. The presence of noticeable tooth staining was noted by 2 of the participants. It is not clear if other participants in the CHX group noticed increased tooth staining

Blinding of outcome assessment (detection bias) All outcomes

Unclear risk

Quotes: "The examining researchers had no access to information about the contents of the tubes or assignment of subjects to the three therapies. All study personnel, including the biostatistician and participants were not aware of the treatment assignments for the duration of the study" and "One examiner carried out all clinical measurements in a given subject and treatment was performed by the second clinician. Thus, the monitoring clinician was masked to the treatment protocol"

Comment: the presence of noticeable tooth staining was noted by 2 of the participants and would also have been visible to the outcome assessors. As tooth staining was not measured in the course of the study, it is not clear if other participants in the CHX group had increased tooth staining that would have been visible to the outcome assessors

Incomplete outcome data (attrition bias)
All outcomes

Low risk

Intention-to-treat analysis was carried out using last available recordings for those lost to follow-up. Therefore all subjects who entered the study were included in the analysis at all time points



eres 2009 (Continued)				
		Attrition (all study arms) 11/60 (18.3%). Dropouts by group: CHX 5/20 (25%), placebo 3/20 (15%). Reasons for loss to follow-up: CHX: 2 moved to another city, 1 got pregnant and 2 could not be contacted; placebo: 3 could not be contacted		
		Comment: dropouts are high particularly in the CHX group. Reasons for losses do not appear to be related to the intervention		
Selective reporting (reporting bias)	High risk	All outcomes stated in the methods section are adequately reported in the results section. However, tooth staining is considered to be an important side effect of CHX mouthrinsing and apart from being self-reported by 2 participants this is not otherwise reported		
Other bias	Unclear risk	The 2 examining researchers were calibrated for the periodontal PD and CAL measurements only		
		Quote: "No statistically significant differences were observed among groups for any parameter at baseline"		
Feres 2012				
Methods	Trial design: parallel (6 arms) (2 x 3 factorial design; 2 arms relevant to this review are described here)			
	Location: Periodontal Clinic of Guarulhos University, Brazil			
	Number of centres: 1			
	Study duration: the recruitment period is not stated, however, the study duration was 1 year and the duration of rinsing was 2 months			
Participants	Participants: healthy subjects with untreated generalized chronic periodontitis			
	Inclusion criteria: at least 30 years old, at least 6 teeth, at least 1 site with PD and CAL 4 mm or more, BOP			
	Exclusion criteria: previous subgingival periodontal therapy, pregnant or breastfeeding, smoking, systemic disease that could affect progression of periodontal disease, long-term use of anti-inflammatory medication, need for antibiotic premedication for routine dental therapy, use of antibiotic therapy in previous 6 months, allergy to metronidazole, amoxicillin or CHX			
	Baseline gingivitis: (% of sites with BOP) overall mean for SRP group (1): 70.6 (SD 23.8); within group (1) CHX + SRP 69.2 \pm 26.5, SRP 72.0 \pm 21.4			

Gender: overall: male 12 (30%), female 28 (70%); CHX: male: 7, female: 13; placebo: male: 5, female: 15

Number randomised: 40 (Gp A: 20; Gp B: 20)

Number evaluated: 40 (Gp A: 20; Gp B: 20) at 3 months; ITT analysis (last observation carried forward) used to account for dropouts at 6 and 12 months (not reported by CHX/placebo)

Interventions Comparison: (1) SRP versus (2) SRP + metronidazole versus (3) root planing + metronidazole + amoxicillin

Each arm was further randomised to receive CHX rinse or placebo rinse. Only the first arm (1) was eligible for inclusion in this review

Gp A (n = 20): CHX 0.12% + SRP, 15 mL, twice daily, 1 min, 2 months



Feres 2012 (Continued)	
, ,	Gp B (n = 20): placebo + SRP as above
	Prophylaxis at baseline: SRP (4 to 6 1-hour sessions under local anaesthesia over 14 days) and "all subjects received periodontal maintenance at the three post-treatment monitoring visits"
	OHI: "During the initial phase, all subjects received instruction on proper home-care techniques and were given the same dentifrice (Colgate Total; Colgate Palmolive Co, São Bernardo do Campo, SP, Brazil) to use during the study period"
	Non-supervised rinsing Timing of mouthrinsing in relation to toothbrushing: "Subjects were instructed to rinse in the morning, 30 min after breakfast and toothbrushing, and also at night 30 min after toothbrushing"
Outcomes	Gingivitis (BOP yes/no, gingival bleeding yes/no); plaque (% sites with accumulation); periodontitis (probing depth, CAL); and suppuration assessed at 3, 6 and 12 months
	Because participants received periodontal maintenance at 3, 6 and 12 months, the 3-month data are considered to be the only data relevant to this review
Funding	Quote: "This study was supported by Research Grant #07/55291-9 from Fundação de Amparo à Pesquisa do Estado de São Paulo (FAPESP, Brazil)"
Notes	Sample size calculation: accounting for 15% attrition, 39 participants were needed per group (the initial 3 groups before further division into CHX/placebo) in order to detect a difference of 4 sites with probing depth 5 mm or greater at 90% power and 5% significance (unclear whether the study is powered to detect a difference between CHX/placebo within each of the 3 arms)
	Adverse effects: adverse effects reported appear to be related to the metronidazole and amoxicillin, but there were no statistically significant differences between any of the 3 groups
	Declarations/conflicts of interest: quote: "The authors declare that they have no conflict of interests"
	Data for the SRP+CHX and SRP groups within group (1) were obtained from the study authors

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quotes: "each of the selected subjects was given a code number during the enrolment visit and the study coordinators (M.Fe. and L.C.F.) used a computer-generated table to allocate them to one of the three therapeutic groups" and "A second randomization was performed to allocate half of the subjects in each group to rinse withCHX or placebo"
		Comment: second randomisation not adequately described
Allocation concealment (selection bias)	Unclear risk	Comment: not described
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Quotes: "double-blind" and "Masking of CHX rinsing may have been hampered by tooth staining"
		Comment: tooth staining is likely to have been noticeable in the CHX group rinsing for 2 months, therefore participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. However, tooth staining is not reported and there is not enough information upon which to base a judgement
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quotes: "double-blind" and "Masking of CHX rinsing may have been hampered by tooth staining"



Feres 2012 (Continued)		
		Comment: tooth staining is likely to have been noticeable in the CHX groups at 3 months follow-up and in this situation the outcome assessor could have worked out which individuals used CHX and not be adequately blinded. However, tooth staining is not reported and there is not enough information upon which to base a judgement
Incomplete outcome data (attrition bias) All outcomes	Low risk	All randomised participants were included in the analyses. ITT analysis (last observation carried forward) used to account for dropouts at 6 and 12 months (not reported by CHX/placebo)
Selective reporting (reporting bias)	High risk	Tooth staining is considered to be an important side effect of CHX mouthrinsing but this is not reported. Only the primary outcome (PD) was reported by CHX/placebo in the published article
Other bias	Unclear risk	Calibration carried out with high Kappa scores for inter- and intraexaminer agreement (> 92%), but this may have only been for the primary outcomes of the study, which were for periodontitis (PD and CAL)

Ferretti 1987

Methods	Trial design: parallel (2 arms)		
	Location: assumed to be University of Kentucky Medical Center, USA		
	Number of centres: 1		
	Study duration: recruitment period not stated. Study duration is 90 days. Duration of rinsing was 60 days		
Participants	Participants: quote: "patients undergoing assessment for bone marrow transplantation"		
	Inclusion criteria: quote: "patients undergoing assessment for bone marrow transplantation patients with dentition"		
	Exclusion criteria: quote: "Two patients were not included in evaluation because, for medical reasons, they did not undergo bone marrow transplantation. These excluded patients were similar in all other aspects to the 33 patients who were evaluated"		
	Baseline gingivitis: not reported		
	Age at baseline (years): Gp A: mean 25.3 ±3.3; Gp B: mean 27.1±3.9. Gp A: range 5-48; Gp B: range 5-51		
	Gender: not reported		
	Number randomised: 35 (Gp A: 18; Gp B: 17)		
	Number evaluated: assumed 33 (Gp A: 16; Gp B: 17) but this is unclear. See note under 'Attrition bias'		
Interventions	Comparison: CHX versus placebo control		
	Gp A (n = 16): CHX 0.12%, 15 mL, 3 times daily, 30 seconds, 60 days		
	Gp B (n = 17): placebo, 15 mL, 3 times daily, 30 seconds, 60 days		
	Prophylaxis at baseline: quote: "Necessary dental care, including dental cleaning was implemented before the initiation of chemoradiotherapy"		
	OHI: none, usual but with foam rubber toothbrush substitute and saline solution irrigations		



Ferretti 1987 (Continued)	Partially supervised rinsing. Quote: "Supervised by the nursing staff of the transplant unit for the duration of the patients' hospitalization (usually 35-45 days). After discharge, patients continued this regimen for the remainder of the 60-day treatment period" Timing of mouthrinsing in relation to toothbrushing: after toothbrushing (with a foam rubber toothbrush)
Outcomes	Gingivitis (gingival inflammation: PMGI, de la Rosa 1976); plaque (TQH, Turesky 1970); extrinsic tooth stain (Meckel's stain grading method, Lang 1977) assessed at 33, 60, and 90 days follow-up Mucositis, oral streptococci and candida are also reported but are not relevant to this review
Funding	Quote: "This study was supported by funds from the University of Kentucky, the Procter & Gamble Co, and the Veterans Administration"
Notes	Sample size calculation: not reported Adverse effects: quote: "Extrinsic dental staining was similar throughout the study for both the chlorhexidine and control groups. There was a modest, although not significant, increase in dental staining noted in both groups during the treatment" Declarations/conflicts of interest: none stated

Bias	Authors' judgement	Support for judgement	
Random sequence genera-	Unclear risk	Quote: "Through prospective randomisation"	
tion (selection bias)		Comment: no description of the method of random sequence generation	
Allocation concealment (selection bias)	Unclear risk	No information is provided	
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "in a double-blind fashion." The placebo was identical in composition to the CHX only without the CHX. Rinsing was supervised by the nursing staff. Blinding of the nursing staff although likely is not explicitly stated	
Alloutcomes		Comment: there was a modest although not significant increase in dental staining noted in both groups during the treatment	
Blinding of outcome assessment (detection bias)	High risk	Quotes: "in a double-blind fashion" and "All clinical indexes were scored by the same examiner"	
All outcomes		Comment: there was a modest although not significant increase in dental staining noted in both groups during the treatment. The incidence of oral candida and mucositis is markedly lower in the CHX group which may have affected blinding by potentially alerting the examiner to the group assignment of the subjects	
Incomplete outcome data (attrition bias) All outcomes	High risk	According to Table 2: patient characteristics, 35 patients were randomised (18 CHX and 17 control) and 33 were evaluated (16 CHX and 17 control). 2 participants were lost from the CHX group because they did not undergo bone marrow transplantation for medical reasons. We assume that the remaining patients were evaluated for inclusion in the study rather than evaluated at the end of the study. This is because data presented within the bar charts for the CHX group indicate substantial losses to follow-up throughout the trial in the CHX group (44% by day 33). Losses from the control groups are not reported but include 2 deaths in the control group early on post-transplant	



Ferretti 1987 (Continued)				
Selective reporting (reporting bias)	Low risk	All outcomes are reported graphically with bars representing a measure of variance (assumed to be SD) in the graphs. We estimated the mean scores and SDs for each outcome so that we could include the data in meta-analyses		
Other bias	Unclear risk	Quote: "All clinical indexes were scored by the same examiner"		
		Comment: there is no mention of training for the clinical examiner.		
		The groups were comparable with regard to age, pretransplant disease status, type of transplant, graft-versus-host disease prophylaxis and graft-versus-host disease. The baseline plaque and gingivitis scores appear balanced between the groups. The control group appears to have had less extrinsic staining at baseline but the statistical significance of the difference is not reported		

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Methods	Trial design: parallel (3 arms - 2 arms are relevant to this review)		
	Location: dental practice in the UK		
	Number of centres: 1		
	Study duration: the recruitment period is not stated. Study duration and duration of rinsing was 4 months		
Participants	Participants: quote: "patients coming to the surgery"		
	Inclusion criteria: quote: "patients had to have some degree of gingival inflammation. A gingival score system was chosen to measure the degree of inflammation present, and patients selected were required to have a total score of greater than 36 points"		
	Exclusion criteria: quote: "No patient under the age of 12 years was included"		
	Baseline gingivitis: not reported		
	Age at baseline (years): range not stated but minimum age was 12		
	Gender: not reported		
	Number randomised: 133 (Gp A: 45, Gp B: 50, Gp C: 38)		
	Number evaluated: 116 - breakdown not specified		
Interventions	Comparison: (CHX rinse + OH) versus (PVP rinse + OH)* versus OH		
	Gp A (n = 45): CHX (concentration not reported) + OH (toothbrushing plus floss and/or wood points),		

volume not reported, once daily, 30 seconds, 4 months

*Gp B (n = 50): PVP + OH (toothbrushing plus floss and/or wood points), volume not reported, once daily, 30 seconds, 4 months. We excluded this arm from our risk of bias and analysis

Gp C (n = 38): OH (toothbrushing plus floss and/or wood points), 4 months

Prophylaxis at baseline: quote: "At the first visit thorough scaling and polishing...were given"

OHI: quote: "At the first visit thorough... instruction in oral hygiene techniques were given"

Non-supervised rinsing

Timing of toothbrushing in relation to mouthrinsing: not reported



Fine 1985 (Continued)	Duration of treatment: 4 months
Outcomes	Gingivitis ('a gingival inflammation index'). Total score was recorded and presented in results; assessed at 1 week, 1 month, and 4 months follow-up
Funding	Not reported but Napp Laboratories helped with the statistical analysis
Notes	Sample size calculation: not reported Adverse effects: quote: "Several patients having been on chlorhexidine-gluconate mouthwash previ-
	ously preferred the PVP-I as it did not stain the teeth or fillings There was also marked differences in the individual tendency to develop extrinsic dental staining after using chlorhexidine as a mouth rinse Patients also reported a drop in the number of aphthous ulcers whilst using PVP-I"
	Declarations/conflicts of interest: none stated
	There appears to be an error in Figure 1. Outcome data were estimated from the graph at 3 months but as the outcome was not measured at 3 months we assumed that these data related to the outcome at 4 months

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Once the patient had been selected he/she was then put into one of three groups by completely random allocation according to a random allocation code"
		Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	High risk	Quote: "Once the patient had been selected he/she was then put into one of three groups by completely random allocation according to a random allocation code." This was done by the dental nurse
		Comment: the use of an open random allocation schedule means that the practice dental nurse could likely foresee assignment and there is a possibility that randomisation could have been subverted
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: participant blinding was not possible for CHX rinse plus oral hygiene versus oral hygiene alone
Blinding of outcome assessment (detection bias)	High risk	Quote: "During the visit neither the dentist nor the hygienist knew which group the patient was in"
All outcomes		Comment: since this study was conducted in a single practice, and randomisation was done by the dental nurse, it is unlikely that blinding was maintained. In addition, tooth staining was noted to have occurred in the CHX group
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Attrition at 3 months (all study arms): 17/133 (12.78%). Losses not broken down by group. Reasons for losses not reported
Selective reporting (re-	High risk	Outcome measured at 1 and 4 months and reported in graphs
porting bias)		Gingivitis: quote: "a gingival inflammation index" reported as 'total score'. No information on the variance of the total score is provided. The data show an unusual increase in mean gingivitis score for CHX at 3 months which is not discussed



Fine 1985 (Continued)

Other bias

Unclear risk

There were 2 examiners: a dentist and hygienist. There is no reference to cali-

bration

Baseline demographic data are not reported. Mean total gingivitis scores ap-

pear similar in Figure 1

Flemmig 1990

Methods

Trial design: parallel (4 arms - 2 arms relevant to this review)

Location: USA. Setting not reported

Number of centres: assumed to be 1

Study duration: recruitment period not stated. Study duration and duration of rinsing was 6 months

Participants

Participants: quote: "patients with naturally-occurring gingivitis"

Inclusion criteria: quote: "1. Minimum of 16 natural teeth including at least 4 molars, not including third molars. 2. Demonstrated Bleeding on Probing (BOP) using a standard periodontal probe at screening examination and an electronic pressure sensitive probe* (25p) at baseline examination at a minimum of 6 sites on a minimum of 4 teeth. These sites must have been equal to or less than 4 mm pocket probing depth and were considered sites with moderate to severe gingivitis"

Exclusion criteria: quote: "1. Systemic disorders or medications which might influence the appearance of the oral tissue or the nature of the plaque. 2. History of adverse reactions which might affect the participation in this study. 3. Used systemic antibiotics for a period of more than 7 days 30 days prior to the baseline examination, or 10 days prior to either the 3 month or 6 month examinations. 4. Diabetics, pregnant or lactating women, or patients on long-term anti-inflammatory medications"

Baseline gingivitis: (GI) Gp A: mean 0.52 (+/-0.04); Gp B: mean 0.56 (+/-0.05); Gp C: mean 0.51 (+/-0.04); Gp D: mean 0.54 (+/-0.04)

Age at baseline (years): mean age 36.6 years. Range not stated

Gender: not reported

Number randomised: 222 (Gp A: 58, Gp B: 55, Gp C: 54, Gp D: 55) Number evaluated: 175 (Gp A: 43, Gp B: 40, Gp C: 44, Gp D: 48)

Interventions

Comparison: CHX rinse versus CHX irrigationa versus water irrigationb versus toothbrushing

Gp A (n = 58): CHX (Peridex) 0.12%, 15 mL, twice daily, 30 seconds, 6 months

^aGp B (n = 55): CHX irrigation 0.06%, 200 mL, once daily, time not reported, 6 months. We excluded this arm from our risk of bias and analysis

^bGp C (n = 54): water irrigation, 500 mL, repeat not reported, time not reported, 6 months. We excluded this arm from our risk of bias and analysis

Gp D (n = 55): toothbrushing, volume N/A, frequency N/A, time not reported, duration N/A

Prophylaxis at baseline: quote: "Patients received a full mouth professional removal of supra- and subgingival plaque and calculus and a prophylaxis after the baseline and 6-month examinations"

OHI: quote: "During the entire course of the study subjects were encouraged to continue to practice their usual oral hygiene and no additional oral hygiene instructions were given, except that all patients used the same dentifrice"

Non-supervised rinsing



lemmig 1990 (Continued)			
(continued)	Timing of mouthrinsing	g in relation to toothbrushing: not reported	
Outcomes	probe 25 p); plaque (Pl faces of the 12 anterior each anterior surface, culus (Volpe-Manhold	ammation: GI, Löe 1967; gingival bleeding: BOP yes/no using pressure sensitive, , Silness & Löe 1964); extrinsic tooth stain (Staining Index, photo of facial surmaxillary and mandibular teeth were graded for (1) overall stain, (2) stain of (3) % coverage against 2 sets of established photo standards); supragingival cal Calculus Index, Volpe 1965); PPD (using pressure sensitive probe 25 p) - not relevessed at 3 and 6 months follow-up	
Funding	Quote: "This study was supported by the Teledyne Research Assistance Program"		
Notes	Sample size calculation: not reported		
	frequent reasons were soft tissue reactions co ed in the exclusion of c	s: "No predominant reasons for not completing the study was found. The most patient not available for examination, tooth staining, and pregnancy. Adverse ensisting of epithelial desquamation of the ventral surface of the tongue resultione patient from the study who irrigated with chlorhexidine" and "Both CHX irried in significant (P ≤ 0.05) increase in the calculus index and stain compared to ter irrigation"	
	Declarations/conflicts of interest: none stated		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	Unclear risk	Quote (from report): "After the baseline examination patients were assigned to one of the following treatment groups after balancing according to sex and	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote (from report): "After the baseline examination patients were assigned to one of the following treatment groups after balancing according to sex and marginal gingival bleeding"
		Quote (from correspondence): "Treatment was assigned randomly the study protocol is not available"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	Quote (from correspondence): "allocation was concealed the study protocol is not available"
		Comment: not enough information is provided
Blinding of participants and personnel (perfor- mance bias)	High risk	The authors report that this is a single-blind study which must apply to the examiner, since participants could not be blinded (irrigation versus rinse versus toothbrushing alone)
All outcomes		Quote: "At 6 months calculus was increased by 273.2% in the CHX rinse group compared to the toothbrushing group. In the CHX rinsing group stain at the facial surfaces of anterior teeth was increased by 74.2% compared to toothbrushing"
		Comment: the higher level of tooth staining in the CHX group meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is unclear
Blinding of outcome assessment (detection bias) All outcomes	High risk	The authors report that this is a single-blind study which must apply to the examiner, since participants could not be blinded (irrigation versus rinse versus toothbrushing alone). Examiner 1 assessed soft tissues and the plaque index. The other examiners each assessed 1 of the remaining parameters (GI, pocket probing depth and BOP and Calculus Index). Each clinical parameter was assessed by the same examiner throughout the study. Tooth discolouration and staining was assessed in a single-blind manner. Quote: "At 6 months calculus



Flemmig 1990 (Continued)		
		was increased by 273.2% in the CHX rinse group compared to the toothbrushing group. In the CHX rinsing group stain at the facial surfaces of anterior teeth was increased by 74.2% compared to toothbrushing"
		Comment: the higher level of staining in the CHX group meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition 22/113 (19.5%). Attrition by group: CHX 15/58 (25.86%), toothbrushing (normal oral hygiene) 7/55 (12.73%). Losses higher in CHX group. Reasons for loss to follow-up (not broken down by group): patient not available for examination, tooth staining and pregnancy. Insufficient information to determine if losses are due to the intervention but authors report a significant increase in stain in the rinse group compared to the toothbrushing group at 6 months and staining was one of the reasons for loss to follow-up
Selective reporting (reporting bias)	Low risk	All outcomes stated in the methods section are adequately reported in the results section. Bacteriological results are presented in another paper
		SD was estimated as sqrt(MSE) so that these data could be included in meta- analyses
Other bias	Unclear risk	There were 4 examiners, each measuring a different outcome. No measure of intraexaminer reliability for each index is given
		Comment: there is no mention of training or calibration of the clinical examiners
		Baseline demographic data by group are not reported

Flotra 1972

Age at baseline (years): mean age: 27; range: 19-39 Gender: not reported Number randomised: 50 (Gp A: 40, Gp B: 10) Number evaluated: 28 (Gp A: 22, Gp B: 6)
Gender: not reported
Age at baseline (years): mean age: 27; range: 19-39
Baseline gingivitis: (GI): Gp A: mean 1.11; Gp B: 1.22
Exclusion criteria: not reported
Inclusion criteria: not reported
Participants: Norweigan Air Force soldiers
Study duration: recruitment period not stated. Study duration and duration of rinsing was 17 weeks
Number of centres: 1
Trial design: parallel (4 arms). *Note: data are reported for the 3 treatment groups combined. Therefore, there are 2 overall groups in the analysis: control and experimental Location: examinations were conducted at the military garrison at Rygge, approximately 50 kilometre south of Oslo, Norway
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Flotra 1972 (Continued)	
(continued)	Gp A (n = 40): 0.2% CHX rinse, 0.1% CHX rinse, 0.1% CHX acetate rinse, 10 mL, twice daily, 1 min, 17 weeks
	Gp B (n = 10): control (benzene alcohol),10 mL, twice daily, 1 min, 17 weeks
	Prophylaxis at baseline: none at baseline. Supra- and subgingival scaling was performed during the study starting during week 9
	OHI: none given. Quote: "The established oral hygiene routine was not otherwise changed"
	Non-supervised mouthrinsing
	No information provided on timing of rinsing in relation to toothbrushing or postrinse instructions
Outcomes	Gingivitis (gingival inflammation: GI Löe & Silness 1963); plaque (PI Silness & Löe 1964); calculus (Retention Index, Bjorby & Löe 1966); and pocket depth (mm) - not relevant for this review; assessed at 8, 11, and 17 weeks follow-up
Funding	Not reported
Notes	Sample size calculation: not reported
	Adverse effects: quote: "The taste of the control solution, like the test solution, was some-what unpleasant"
	Side effects are reported in a separate publication. During the experimental period, 11 participants exhibited lesions in the oral mucosa (10 CHX and 1 control). 3 of the 10 participants developed more severe lesions (2 in the 0.2% CHX gluconate group and 1 in the 0.1% CHX acetate group). 1 individual (0.1% CHX gluconate) developed what appeared to be a viral stomatitis and another (0.1% CHX acetate) a viral parotitis. 1 individual developed small ulcers under the upper lip - probably due to an aphthous lesion. In the test groups at 4 weeks (n = 48), 12% of the tooth surfaces without fillings and 62% of the silicate fillings became discoloured while 36% of the test persons developed discoloured tongues during the experimental period
	Declarations/conflicts of interest: none stated

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation is not mentioned in the study report
		Quote (from correspondence): "The allocation of participants to the various groups was performed in this way: First we decided how many we wanted in each group, the appropriate no of lots for each group were made and drawn for each participant from a jar"
		Comment: adequate method of random sequence generation
Allocation concealment	Low risk	Allocation concealment is not mentioned in the study report
(selection bias)		Quote (from correspondence): "The drawing of lots was performed by Dr Waerhaug and myself (Prof Gjermo), with particular attention that the group allocation was not revealed for Dr Flötra who performed the examinations. This was done at the Department of Periodontology at the University of Oslo, whereas the actual examinations were performed at a military garrison at Rygge, approximately 50 km south of Oslo. Dr Flötra brought the results to Oslo before the seal for the allocation was broken and data analyzed"
		Comment: we consider it unlikely that participants or investigators enrolling participants could foresee assignments



Flotra 1972 (Continued)

Blinding of participants and personnel (performance bias) All outcomes High risk

Quote: "The experiment was a double-blind study." Additional information was obtained from the second author of the paper: "neither the examiners nor the participants were aware of which group they belonged to." The control mouthrinse contained 0.5% benzene alcohol

Quotes: "The test mouth washes were distributed in coded plastic bottles.." and "The taste of the control solution, like the test solution was somewhat unpleasant"

In the test groups, 12% of the tooth surfaces and 62% of the silicate fillings became discoloured while 36% of the test persons developed discoloured tongues during the experimental period

Comment: the higher level of tooth staining in the CHX group meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is unclear

Blinding of outcome assessment (detection bias) All outcomes High risk

Quote: "The experiment was a double-blind study." Additional information was obtained from the second author of the paper: "neither the examiners nor the participants were aware of which group they belonged to" and "One investigator recorded the Plaque Index throughout the study and another recorded the gingival index" and "Dr Flötra brought the results to Oslo before the seal for the allocation was broken and data analyzed"

In the test groups, 12% of the tooth surfaces and 62% of the silicate fillings became discoloured while 36% of the test persons developed discoloured tongues during the experimental period

Comment: despite the efforts taken to maintain blinding, the higher level of staining in the CHX group meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear

Incomplete outcome data (attrition bias)
All outcomes

High risk

From Table 1 and 2 n = 28 at 17 weeks. 22/50 (44%) lost to follow-up at 17 weeks. By group: Gp A 18/40 (45%); Gp B 4/10 (40%). Reasons for losses given for 11/22: 2 excluded because of stomatitis and parotitis, 9 for various reasons such as discharge from service, transfers, etc

Comment: losses to follow-up although proportionately balanced between the groups were unacceptably high

Selective reporting (reporting bias)

High risk

The outcome data for the 3 CHX rinse groups are not reported by individual rinse group; combined data for the 3 CHX groups are presented in the results. The results for the Retention Index are not reported fully. Pocket probing depth is measured at baseline but not reported. Side effects including oral lesions, discolouration of the teeth and tongue are reported in detail in a separate publication although not in a format that would allow their inclusion in meta- analyses

Other bias

Unclear risk

2 examiners, 1 for each index. No measurement of intraexaminer reliability

Baseline demographic data are not reported. Mean GI and PI in the 2 groups at baseline appear similar (not significantly different for GI)

Graziani 2015

Methods

Trial design: parallel (4 arms)



Graziani 2015 (Co	ontinued)
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Location: Unit of Dentistry and Oral Surgery, University Hospital of Pisa, Italy

Number of centres: 1

Study duration: the recruitment period is not stated. Study duration and duration of rinsing was 5 weeks

Participants

Participants: healthy adults

Inclusion criteria: at least 20 teeth, the 8 incisors free of restorations or prosthetic reconstructions, general good health

Exclusion criteria: systemic illness (including cardiovascular, renal of liver diseases), smokers, periodontal/dental treatment within the 35 days preceding the study, pregnant or lactating females, undergoing orthodontic treatment, inability to sign consent form

Baseline gingivitis: (Löe & Silness GI) Gp A: mean 1.8 (95% CI 1.4 to 2.2); Gp B: mean 1.6 (95% CI 1.3 to 1.9); Gp C: mean 1.8 (95% CI 1.5 to 2.2); Gp D: mean 1.5 (95% CI 1.3 to 1.8)

Age at baseline (years): Gp A: mean 29.7 (95% CI 23.4 to 35.9); Gp B: mean 39.5 (95% CI 31.6 to 35.9); Gp C: mean 36 (95% CI 30.3 to 37.2); Gp D: mean 29.7 (95% CI 24 to 35.5)

Gender: overall: male 36 (51%), female 34 (49%). Gp A: male 6 (40%), female 9 (60%); Gp B: male 11 (52%), female 10 (48%); Gp C: male 10 (53%), female 9 (47%); Gp D: male 9 (60%), female 6 (40%)

Number randomised: 70 (Gp A: 15; Gp B: 21; Gp C: 19; Gp D: 15)

Number evaluated: 70 (Gp A: 15; Gp B: 21; Gp C: 19; Gp D: 15)

Interventions

Comparison: CHX rinse 1 versus CHX rinse 2 versus CHX rinse 3 versus control rinse

Gp A (n = 15): CHX 0.2% (with alcohol), 10 mL, twice daily, 1 min, 5 weeks

Gp B (n = 21): CHX 0.2% (alcohol-free), 10 mL, twice daily, 1 min, 5 weeks

Gp C (n = 19): CHX 0.2% with antidiscolouration system (alcohol-free), 10 mL, twice daily, 1 min, 5 weeks

Gp D (n = 15): H₂O/NaCl control, 10 mL, twice daily, 1 min, 5 weeks

Prophylaxis at baseline: supragingival scaling and polishing

OHI: quote: "Oral hygiene instructions were provided... as follows: interdental brushing... and/or dental floss when interdental embrasures did not allow interdental brushing. Interdental cleaning was followed by electric toothbrushing... A sodium lauryl sulfate-free toothpaste was provided to each participant"

Non-supervised rinsing

Timing of mouthrinsing in relation to toothbrushing: not reported

Postrinsing instructions: participants were instructed to refrain from tea, coffee and red wine consumption for at least 1 hour after rinsing

Outcomes

Gingivitis (gingival inflammation: GI Löe & Silness 1963; gingival bleeding: full-mouth bleeding score, Ainamo and Bay 1975); plaque (full-mouth plaque score, O'Leary 1972); tooth discolouration/staining (staining index, modification of Lobene 1968, Grundemann 2000) using digital photographic measurements of the buccal surfaces of the 8 incisors, tooth colour was measured using spectrophotometric analysis; other adverse effects; assessed at 1, 3 and 5 weeks follow-up

Funding

Quote: "Johnson & Johnson supported this study with a grant to purchase the Vita EasyShade spectrophotometer and to perform the data analysis (Dr D'Aiuto). UCLH/UCL received a proportion of funding from the Department of Health's NIHR Biomedical Research Centres funding scheme"



Graziani 2015 (Continued)

Notes

Sample size calculation: sample size was geared towards comparing each treatment mean with the control mean. The calculation was reported and achieved. Quote: "A sample size of 12 individuals per group, with at least three subjects to compensate for dropouts, with 3 treatment arms and one control group would provide any-pair power of 80% (α = 0.05) (Kruskal-Wallis Test) for comparing each treatment mean with the control mean"

Adverse effects: "At 35 days, staining was frequent, reported in well above 60% of all cases rinsing with the three CHX formulations." Tongue staining incidence: Gp A: 1 at 35 days; Gp B: 2 at 21 days; Gp C: 2 at 7 and 21 days. Taste alteration: Gp A: 2 at 7 days; Gp B: 2 at 7 days and 2 at 21 days; Gp C: 1 at 35 days. Gastric acidity: Gp B: 1 at 7 days. No serious side effects reported

Non-smokers

Declarations/conflicts of interest: quote: "Dr Graziani has received lecture fees from Johnson & Johnson and from Curaden healthcare" (Johnson & Johnson and Curaden were the manufacturers of 2 of the 3 CHX rinses)

Participants in the study were described as having "mild periodontal involvement and periodontal inflammation limited to approximately one-fifth of the entire dentition"

We combined groups A, B and C for the main analysis

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "a computer-generated random sequence was used to assign participants to one of the four treatment groups"
		Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Low risk	Quote: "random sequence conducted by a research fellow not directly involved in the experiments was used to assign participants Allocation to the treatment was concealed as a code number identifying the allocated group. It was sealed in an opaque envelope which was opened at the baseline visit after completion of clinical measurements, photographs, supragingival scaling and polishing"
		Comment: numerous steps were taken to ensure that the random sequence was applied exactly as it was generated without any manipulation
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quotes: "double-blind" and "the mouthrinse bottles were dispensed to the participants according to the allocated group by a research fellow not directly involved in the research. The control rinse was saline"
		Quote: "At 35 days, staining was frequent, reported in well above 60% of all cases rinsing with the three CHX formulations"
		Comment: the higher level of tooth staining in the CHX groups meant that participants could have worked out which group they were in (or that they were in 1 of the active mouthrinse groups) and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is unclear
Blinding of outcome assessment (detection bias)	High risk	Quotes: "double-blind" and "Clinical examinations were performed by an examiner unaware of the group allocation"
All outcomes		Comment: the higher level of staining in the CHX groups meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear



Graziani 2015 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Low risk	All randomised participants were included in the analyses (ITT analysis) Attrition at 35 days was 17/70 (24.3%). By group: Gp A 4/15 (26.7%); Gp B 3/21 (14.3%); Gp C 4/19 (21.1%); Gp D 6/15 (40%)
Selective reporting (reporting bias)	High risk	Percentage reduction in full-mouth bleeding score was not reported for Gp A. Due to incomplete reporting of this outcome, data on full-mouth bleeding score could not be included in a meta-analysis
Other bias	High risk	Full-mouth bleeding score measurement was carried out by a "blinded, calibrated and experienced examiner." Apart from that statement, training and calibration of the 1 outcome assessor are not mentioned. The 4 groups do not appear to be balanced with respect to mean age, % periodontitis diagnosed, mean number of pockets and % of pockets ≥5 mm. The 95% confidence intervals indicate some statistically significant differences between the groups but the P values for the differences are not reported

Grossman 1986

Methods	Trial design: parallel (2 arms)		
	Location: USA. Quote: "Dental office conditions"		
	Number of centres: assumed to be 1		
	Study duration: recruitment period not stated. Study duration and duration of rinsing was 6 months		
Participants	Participants: adults		
	Inclusion criteria: quote: "To be accepted into the study, subjects had to have at least 16 natural teeth including 4 molars"		
	Exclusion criteria: quote: "Subjects were not enrolled if they had existing soft-tissue damage due to ill fitting appliances, evidence of overt periodontal disease or gross dental neglect, an unfavorable medical history, or a history of routine antibiotic use"		
	Baseline gingivitis: (GI) examiner A (Gp A: mean 0.5530, Gp B: mean 0.5388); examiner B (Gp A: mean 0.7472, Gp B: mean 0.7470)		
	Age at baseline (years): Gp A: 33.02; Gp B: 32.25. Range Gp A: 18-58; Gp B: 18-60		
	Gender: Gp A: male 72 (33%), female 147 (67%); Gp B: male 71 (34%), female 140 (66%)		
	Number randomised: 430 (Gp A: 219, Gp B: 211)		
	Number evaluated: 380 (Gp A: 187, Gp B: 193)		
Interventions	Comparison: CHX versus placebo		
	Gp A (n = 187): CHX 0.12%,15 mL, twice daily, 30 seconds, 6 months		
	Gp B (n = 193); placebo, 15 mL, twice daily, 30 seconds, 6 months		

Gp B (n = 193): placebo, 15 mL, twice daily, 30 seconds, 6 months

Prophylaxis at baseline: quote: "All subjects received a thorough prophylaxis after the baseline examination for plaque, gingivitis, and oral soft tissue health"

OHI: quote: "A sodium fluoride dentifrice (Crest) and soft toothbrushes were given to each subject to be used according to their own individual habits"

Non-supervised mouthrinsing

No information is provided on timing of rinsing in relation to toothbrushing or postrinse instructions



Grossman 1986 (Continued)	Duration of treatment: 6 months		
Outcomes	Gingivitis (gingival inflammation (GI, Löe 1967); gingival bleeding (GI sites with scores 2 or 3 indicating bleeding)); plaque (TQH, Turesky 1970); oral soft tissue effects; calculus (index not reported); and tooth stain (index not reported) and tongue stain assessed at 3 and 6 months follow-up		
Funding	Quote: "The support of this study by the Procter & Gamble Company, Cincinnati, Ohio, is gratefully acknowledged"		
Notes	Sample size calculation: not reported		
	Adverse effects: quotes: "As expected with the use of any oral antimicrobial, some extrinsic tooth staining was observed in the chlorhexidine group" and "Supragingival calculus was higher in the group using chlorhexidine" and "The systematic evaluation of the oral structures did not yield any indication of chlorhexidine-related changes in the oral mucosa"		
	Declarations/conflicts of interest: none stated		
	Gingivitis data were extracted for examiner A only		
Risk of bias			

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "subjects were stratified by sex, age and gingivitis severity. Within strata they were assigned to one of two treatment groups by random permutations of 2"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	Comment: no information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quotes: " double-blind clinical trial" and "Neither the subject nor the dental examiners knew which treatment the subject was assigned to at any time during the study." The control mouthrinse was identical to the CHX mouthrinse but without the CHX gluconate
		Quote: "more extrinsic tooth staining was observed in the chlorhexidine group"
		Comment: the higher level of tooth staining in the active group meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is not clear
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quotes: " double-blind clinical trial" and "Neither the subject nor the dental examiners knew which treatment the subject was assigned to at any time during the study" and "gingivitis was measured independently by two examiners" and "Prior to the 6-month examination, a subset of subjects was fitted with custom-made tooth covers fabricated from opaque plastic. This procedure was included to eliminate bias due to chlorhexidine related tooth stain. The subjects' tongues were also coated with a paste mixture of food colors to eliminate the chance of bias which could result from stained tongues during the gingivitis examinations" and "The GI and bleeding scores recorded by each examiner were consistent regardless of whether subjects were graded with or without tooth covers"
		Comment: the presence of tooth staining in the CHX group did not appear to have an effect on the measurement of gingivitis. The effect of the tooth staining in the CHX group on plaque is less clear and the measurement of plaque



Grossman 1986 (Continued)		could have been affected if the examiners noticed the increased tooth staining in the active group. It is also unclear how the tooth staining may have influenced recording of outcomes at 3 months when no tooth covers were used
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition at 6 months was 50/430 (11.6%). Attrition by group: CHX 32/219 (14.6%); control: 18/211 (9%). Reasons: quote: "Most of the drop outs were people who had moved or could not meet examination schedules" Comment: assuming the missing participants in 1 group had a higher mean (e.g. gingivitis score) than those in the other group, as the attrition rate increased, so would the mean difference between groups
Selective reporting (reporting bias)	High risk	Outcome data for gingivitis, occurrence, severity and bleeding are reported separately for examiner A and B. We extracted gingivitis data for examiner A only. Information on the variance of the mean scores for gingivitis and plaque were not reported. We used the SDs for each clinical parameter at baseline so that we could include the data in meta-analyses. Outcome data for tooth and tongue staining and calculus (measured at 3 and 6 months) are mentioned in the methods and discussion but are not reported in the results
Other bias	Unclear risk	The demographics of the study population were balanced at baseline. Gingivitis was measured independently by 2 examiners. There is no mention of training or calibration for either examiner

Grossman 1989

Methods	Trial design: parallel (4 arms - 2 arms relevant to this review)
metrious.	Location: assumed to be Northfield, New Jersey, USA. Quote: "Dental office conditions"
	, , , , ,
	Number of centres: 1
	Study duration: recruitment period is not stated. Study duration and duration of rinsing was 6 months
Participants	Participants: quote: "adult males and females from the Northfield, New Jersey area"
	Inclusion criteria: quote: "To be accepted into the study, subjects had to sign an informed consent, have at least 16 natural teeth (including four molars), and have at least one gingival site that bled on probing"
	Exclusion criteria: quote: "Subjects were not enrolled if they had evidence of overt periodontal disease or gross dental neglect, an unfavorable medical history, or a history of routine antibiotic use"
	Baseline gingivitis: (GI) Gp A: mean 0.5332, Gp B: mean 0.5227, Gp C: mean 0.5457, Gp D: mean 0.4994
	Age at baseline (years): Gp A: mean 36.8, Gp B: mean 37.1, Gp C: mean 35.9, Gp D: mean 36.5. Range no reported
	Gender: overall: male 166 (33%), female 336 (67%). Gp A: male 35 (31%), female 78 (69%); Gp B: male 4 (36%), female 83 (64%); Gp C: male 42 (33%), female 85 (67%); Gp D: male 43 (32%), female 90 (68%)
	Number randomised: 502 (Gp A: 113, Gp B: 129, Gp C: 127, Gp D: 133). *Note: unclear if this is the number initially randomised
	Number evaluated: 481
	Note: n by group taken from Table 1 showing initial balance for subjects completing the study
Interventions	Comparison: CHX versus phenolic essential oils ^a versus Sanguinarine ^b versus placebo



Grossman 1989 (Continued)

Gp A (n = 113): CHX 0.12%, 15 mL, twice daily, 30 seconds, 6 months

^aGp B (n = 129): phenolic essential oils, 20 mL, twice daily, 30 seconds, 6 months. We excluded this arm from our risk of bias and analysis

^bGp C (n = 127): Sanguinarine, 15 mL, twice daily, 15 seconds (2 consecutive rinses of 15 seconds twice a day), 6 months. We excluded this arm from our risk of bias and analysis

Gp D (n = 133): placebo, 15 mL, twice daily, 30 seconds, 6 months

Prophylaxis at baseline: quote: "Following baseline examinations... all subjects received a thorough prophylaxis"

OHI: none given. Quote: "A sodium fluoride dentifrice (Crest, Proctor and Gamble) and soft toothbrushes were given to all subjects to be used according to individual habits"

Non-supervised rinsing

Timing of mouthrinsing in relation to toothbrushing: not reported

Outcomes

Gingivitis (gingival inflammation: GI Löe & Silness 1967; gingival bleeding: GI grades 2 or 3); plaque (PI, TQH, Turesky 1970); extrinsic tooth stain (photographs of the facial surfaces of the 12 anterior teeth in the maxilla and mandible were graded for dental stain intensity (0-4 scale none-very dark) and coverage (0-6 scale none-heavy)); oral soft tissue health assessed at 3 and 6 months follow-up

Funding

Quote: "The support of this study by the Procter & Gamble Company, Cincinnati, Ohio is gratefully acknowledged"

Notes

Sample size calculation: not reported

Adverse effects: quote: "As would be expected, the extended use of chlorhexidine resulted in more extrinsic dental stain than in other groups. In contrast to previously published reports using different methods of evaluation, use of the phenolic rinse produced a significant increase in dental stain as well. After 3 and 6 months, those subjects using either chlorhexidine or the rinse containing phenolic compounds experienced a significant increase in stain when compared to those on placebo or the sanguinarine rinse"

Declarations/conflicts of interest: none reported, however some of the authors worked for the Procter & Gamble Company

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Subjects were stratified by age and GI score at baseline. Quote: "Within strata, subjects were randomly assigned to one of four treatments"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	The authors state that the study is double-blind. Quotes: "Neither the subjects nor the examiners knew which treatments were assigned at any time during the study" and "All products were supplied to the subjects in identical amber pint bottles." The placebo rinse was Peridex without the CHX
		Comment: there was a significantly higher level of staining in the CHX group compared to the placebo group and this meant that participants could have worked out which group they were in, and this could have affected their oral



irossman 1989 (Continued)		health behaviours, and hence the outcome. The direction of this potential bias is not clear
Blinding of outcome assessment (detection bias) All outcomes	Low risk	At the 6-month examination each subject was graded twice, first without a custom-made plastic tooth cover in place and then with the tooth cover in place. Quote: "In addition the dorsum of the tongue was stained brown with food color to imitate the antimicrobial tongue stain." Analysis of gingivitis scores recorded with and without masking devices revealed that the reductions for each treatment group were comparable
		Comment: the presence of tooth staining in the CHX group did not appear to have an effect on the measurement of gingivitis (our primary outcome). The effect of the tooth staining in the CHX group on plaque is less clear and the measurement of plaque could have been affected if the examiners noticed the increased tooth staining in the active group. It is also unclear how the tooth staining may have influenced recording of outcomes at 3 months when no tooth covers were used
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Total number randomised is not specified, only number completing. Baseline data on completers is provided in Table 1, but number does not tally with number of completers reported in the abstract. No details are provided on number of losses to follow-up
Selective reporting (reporting bias)	High risk	The covariance adjusted mean scores for gingivitis, bleeding and plaque are presented with the covariance analysis mean squared error term for the 3 and 6-month exams. SD was calculated as the sqrt of the covariance analysis mean squared error so that these data could be included in meta-analysis. Data on mean extrinsic tooth stain accumulation is reported without any information on variance of the mean scores
Other bias	Unclear risk	No information is provided on intra- and interexaminer reproducibility
		Quote: "All groups were well-balanced at the beginning of the study with respect to age, gender, initial plaque, stain, gingivitis or bleeding grades"

Hase 1995

Methods	Trial design: parallel, (3 arms - 2 are relevant to this review)
	Location: not explicitly stated and authors from Sweden and Finland (setting not reported)
	Number of centres: 1
	Study duration: recruitment period is not stated. Study duration and duration of rinsing was 4 weeks
Participants	Participants: adult males with gingivitis
	Inclusion criteria: males aged 18 to 60 years; at least 25% of all gingival sites (6 sites per tooth) having a minimum Löe & Silness GI score of 2; minimum 10 teeth per jaw; no pocket probing depth greater than 5 mm
	Exclusion criteria: people with buccal or extensive interproximal restorations reaching beyond the axial angle of the teeth assigned for the gingival fluid flow measurement; people with crowns or bridgework; previous hypersensitivity or reactions to drugs; alcohol or drug addiction; severe liver or renal disease; psychiatric disorders; severe disability with multiple drug therapy; current antibiotics or antiphlogistics (including sprays with corticosteroids) or anticholinergic antiparkinson or antidepressant treatments
	Baseline gingivitis: (% sites BOP) Gp A: mean 16 (SE unclear); Gp B: mean 14 (SE unclear); Gp C: mean 19 (SE unclear)



Library	Better health.	Cochrane Database of Systematic Reviews		
Hase 1995 (Continued)				
,	Age at baseline (years): Gp A: mean 41 (SE 2); Gp B: mean 4 18-60	41 (SE 2); Gp C: mean 39 (SE 2). Age range		
	Gender: males only			
	Number randomised: 62 (Gp A: 21; Gp B: 20; Gp C: 21)			
	Number evaluated: 57 (Gp A: 21; Gp B: 18; Gp C: 18)			
Interventions	Comparison: CHX digluconate rinse versus delmopinol hydrochloride rinse* versus placebo rinse			
	Gp A (n = 21): CHX (0.2%), 10 mL, 1 min, twice daily (morni	ing and evening)		
	*Gp B (n = 20): delmopinol hydrochloride (0.2%), same roubias and analysis	utine. We excluded this arm from our risk of		
	Gp C (n = 21): same routine using placebo (flavoured in ordactive solutions)	der to imitate the sensation of rinsing with the		
	Prophylaxis at baseline: supra- and subgingival scaling			

OHI: participants were instructed to brush their teeth without toothpaste

Timing of rinsing in relation to toothbrushing: after brushing without toothpaste

Outcomes

Quote: "The split-mouth technique was used in the study in order to avoid interference between various assessments." Plaque measurements were made on the left halves of the jaws and gingivitis measurements on the right

Gingivitis (gingival BOP, no reference), measured at 2 and 4 weeks; plaque (PI, Silness & Löe 1964; plaque wet weight, Scheinin & Makinen 1971 - not relevant to this review) measured at 4 weeks

Subjective opinion of participants regarding taste, staining (teeth or tongue or both), clean mouth, vesicles or ulcerations or both, burning sensation, transient anaesthetic sensation, all measured at 4 weeks on a 1 to 100 mm VAS

Gingival fluid flow (Löe & Holm Pedersen 1965) measured at 2 and 4 weeks

Safety (physical examinations including blood tests and ECG), measured before starting treatment and within 3 weeks after the termination of the study - not relevant to this review

Funding

Not reported

Non-supervised rinsing

Postrinsing instructions: not reported

Duration of treatment: 4 weeks

Notes

Sample size calculation: not reported. Quote: ".. the number of patients in this study turned out to be too small to yield conclusive results on the effect of either delmopinol or chlorhexidine on pre-established gingivitis"

Adverse effects: results from the safety assessment were focussed on delmopinol rather than CHX. Others were reported on a VAS scale. Treatment with CHX resulted according to patients in more staining of teeth and tongue than the other 2 mouthrinses. An anaesthetic sensation transient anaesthesia at the tip of the tongue in the oral mucosa was experienced more clearly by the patients in the delmopinol group than by those using placebo or CHX

Results from the VAS (estimated from Figure 3):

- -Taste: CHX 58 (SE 8), placebo 23 (SE 5), delmopinol 60 (SE 8)
- -Staining of teeth/tongue: CHX 38 (SE 7), placebo 9 (SE 1), delmopinol 10 (SE 5)



Hase 1995 (Continued)

- -Clean mouth: CHX 29 (SE 9), placebo 33 (SE 7), delmopinol 30 (SE 8)
- -Vesicles and/or ulcerations: CHX 9 (SE 1), placebo 8 (SE 1), delmopinol 7 (SE 0.5)
- -Burning sensation: CHX 18 (SE 5), placebo 8 (SE 0.5), delmopinol 17 (SE 6)
- -Transient anaesthetic sensation: CHX 21 (SE 9), placebo 10 (SE 4), delmopinol 38 (SE 9)

Declarations/conflicts of interest: none explicitly stated but first and last authors were employees of Biosurface AB, Malmo, Sweden

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Patients were allocated to treatment groups according to a computer-generated randomisation list, where treatment was balanced within blocks of 6 patients"
		Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	Quote: "Packaging and labelling were carried out at an independent Clinical Service Department"
		Comment: it is not clear whether the assignment of subjects to groups was carried out at the independent Clinical Service Department
Blinding of participants and personnel (perfor- mance bias)	High risk	Quotes: "double-blind" and "The various test solutions were dispensed in identical 200 ml amber glass bottles" and "The placebo was flavoured in order to imitate the sensation of rinsing with the active solutions"
All outcomes		Comment: the higher incidence of staining in the CHX group (plus worse taste, greater transient anaesthetic sensation, and greater burning sensation in the active groups) meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcomes. The direction of this potential bias is not clear
Blinding of outcome assessment (detection bias) All outcomes	High risk	The study is reported to be "double-blind". There was 1 clinical examiner. Quote: "The split-mouth technique was used in the study in order to avoid interference between various assessments"
		Comment: the authors do not report precisely who was blinded, but even if this includes the outcomes assessor he higher incidence of staining in the CHX group meant that they could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is not clear
Incomplete outcome data (attrition bias) All outcomes	High risk	Losses to follow-up at 4 weeks: 3/42 (7%). By group: placebo 3/21 (14.29%). There were no losses from the CHX group. Reasons for losses by group: placebo, 1 did not meet the eligibility criteria and 2 took antibiotics during the trial
		Comment: although reasons for attrition do not appear to be related to the particular interventions, the differential loss introduces the possibility that the results may be distorted
Selective reporting (reporting bias)	Low risk	Information on means and SEs are presented graphically. From correspondence with the author/research team, additional information was obtained on means and SEs which enabled calculation of SDs from SEs and inclusion of these data in the meta-analyses
Other bias	Unclear risk	No information is provided on intra- and interexaminer reproducibility.



Hase 1995 (Continued)

Mean age was balanced across groups at baseline. No statistically significant demographic differences between the groups were recorded. Baseline BOP and mean plaque scores appear balanced but the statistical significance of the differences is not reported

Hase 1998

Methods

Trial design: parallel (3 arms - 2 arms relevant to this review)

Location: military regiment, Halmstad, Sweden

Number of centres: 1

Study duration: recruitment period is not stated but the duration of rinsing was 6 months

Participants

Participants: military conscripts and military staff (adult males) with gingivitis

Inclusion criteria: males aged at least 18 years; at least 25% bleeding sites of all gingival sites (6 sites per tooth) on BOP using a Florida-probe with a pressure of 0.25 N to the bottom of the pocket; minimum 16 healthy teeth without crowns, bridge-work, or ill-fitting dental restorations

Exclusion criteria: 4 or more teeth with cavities or pocket probing depth 5 mm or more; previous hypersensitive reactions to drugs; alcohol or drug addiction; severe liver or renal disease; psychiatric disorders; severe disability with multiple drug therapy; receiving antibiotics within 6 weeks of the prestudy visit; antiphlogistics (including sprays with corticosteroids) or anticholinergic drugs such as antiparkinson drugs or antidepressants

Baseline gingivitis: (% sites BOP) Gp A: mean 42.7 (SE 2.7); Gp B: mean 47.3 (SE 2.6); Gp C: mean 43.9 (SE 2.7) (unpublished data)

Age at baseline (years): Gp A: mean 24 (SD 10); Gp B: mean 26 (SD 7); Gp C: mean 23 (SD 7)

Gender: males only

Number randomised: 140 (Table 2 page 748) (Gp A: 47; Gp B: 48; Gp C: 45)

Number evaluated (per protocol analysis): 100 (Gp A: 30; Gp B: 37; Gp C: 33)

Interventions

Comparison: CHX digluconate rinse versus delmopinol hydrochloride rinse* versus placebo rinse

Gp A (n = 47): CHX (0.2% Hibitane Dental), 10 mL, 1 min, twice daily

*Gp B (n = 48): delmopinol hydrochloride (0.2%), same routine. We excluded this arm from our risk of bias and analysis

Gp C (n = 45): same routine using placebo (same as Gp B but without delmopinol and sodium hydroxide)

Prophylaxis at baseline: supra- and subgingival scaling

OHI: participants were instructed not to change their normal mechanical tooth cleaning methods during the study

Partly supervised rinsing (mostly supervised but occasionally unsupervised due to military training)

Timing of rinsing in relation to toothbrushing: after toothbrushing

Postrinsing instructions: not reported

Outcomes

Gingivitis (gingival BOP using a Florida-probe with a pressure of 0.25 N to the bottom of the pocket (Gibbs 1988, Löe 1967)), measured at 3 and 6 months



Hase 1998	(Continued)
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Plaque (PI, TQH, Turesky 1970) measured at 3 and 6 months

Calculus (Volpe-Manhold Calculus Index, Volpe 1967), measured at 3 and 6 months

Extrinsic tooth staining (Staining Index of Shaw & Murray 1977), measured at 3 and 6 months

Adverse events (transient anaesthetic sensation, taste modification, staining (teeth or tongue or both)) were recorded

Microbiological monitoring of plaque and salivary microflora was performed during the treatment period and also 3 months after the end of treatment for 65 patients - results (excluding BOP which is presented here) presented separately - not relevant to this review

Funding

Not reported

Notes

Sample size calculation: based on BOP. The standard deviation was estimated to 14 and a difference of 10% units between placebo and delmopinol was considered worth detecting. Based on a power of 95% at significance level of 5%, 50 participants per group were required. Therefore this was not achieved

Adverse effects: transient anaesthetic sensation in the oral mucosa especially at the tip of the tongue in delmopinol and CHX groups reached the same level of reporting at 6 months. Taste modification was commonly reported in both the delmopinol and CHX groups. Staining of teeth or tongue was common in the CHX group

Adverse effects at 6 months (Table 4) (% participants experiencing the adverse effect):

local anaesthesia/hypoasthesia/parasthesia CHX 18%, placebo 0%, delmopinol

22%; taste loss/taste perversion: CHX 16%, placebo 0%, delmopinol 24%; discolouration teeth/discolouration tongue: CHX 13%, placebo 7%, delmopinol 8%

Declarations/conflicts of interest: nothing explicitly stated but first author was employee of Biosurface AB, Malmo, Sweden

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Patients were allocated to treatment groups according to a computer-generated randomisation list, where treatment was balanced within blocks of 6 patients"
		Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	Quote: "Packaging and labelling were carried out at an independent Clinical Service Department"
		Comment: it is not clear whether the assignment of subjects to groups was carried out at the independent Clinical Service Department
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	The study is reported to be "double-blind" and "The various test solutions were dispensed in identical 200 ml amber glass bottles." The placebo was the same composition as the delmopinol rinse but without the delmopinol and sodium hydroxide. It is unclear who supervised the rinsing and if they were blinded to allocation of the rinse
		Comment: the higher level of staining of the teeth and tongue in the CHX group (plus greater taste modification and transient anaesthetic sensation in the active groups) meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcomes. The direction of this potential bias is not clear



Hase 1998	(Continued)
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Blinding of outcome assessment (detection bias) All outcomes	High risk	The study is reported to be "double-blind". Assessments were made by 5 trained dental hygienists who followed the same subjects throughout the treatment period	
		Comment: the authors do not report precisely who was blinded, but even if this includes the outcomes assessors the higher level of staining of the teeth and tongue in the CHX group meant that they could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is not clear	
Incomplete outcome data (attrition bias) All outcomes	High risk	At 6 months 29 participants are excluded from the analysis in the study arms of interest to this review but only 16 are reported as withdrawn from the study with reasons for withdrawal. Data presented here for loss to follow-up are based on the total lost to follow-up (n = 29)	
		Attrition at 6 months 29/92 (31.5%). By group: CHX 17/47 (36.2%), placebo 12/45 (26.7%). Reasons for loss to follow-up: CHX: adverse events/lack of cooperation, release from military service; placebo: adverse events/lack of cooperation, release from military service, moved away	
		Comment: very high attrition which may be related to type of mouthrinse and could feasibly introduce bias in the results	
Selective reporting (reporting bias)	Low risk	Information on means and SEs are presented graphically. From correspondence with the author, additional information was obtained on means and SEs which enabled calculation of SDs from SEs and inclusion of these data in the meta-analyses	
Other bias	Unclear risk	Assessments were made by 5 trained dental hygienists who followed the same subjects throughout the treatment period. A calibration session for all assessments was performed before the start of treatment but the results of the calibration (Inter- and intraexaminer reliability) are not reported	
		Mean age was balanced across groups at baseline. No statistically significant demographic differences between the groups were recorded. Baseline BOP and mean plaque scores appear balanced but the statistical significance of the differences is not reported	

Hoffmann 2001

Hoffmann 2001	
Methods	Trial design: parallel (5 arms - 4 arms relevant to this review)
	Location: subjects were recruited from the University of Technology, Dresden. The setting was the University Periodontal Clinic
	Number of centres: 1
	Study duration: the period of recruitment is not stated. Study duration and duration of rinsing was 6 months after a run in period of 2 weeks
Participants	Participants: medical students
	Inclusion criteria: quote: "exhibiting good oral hygiene Inclusion criteria were at least 20 teeth, ginigival index (GI) <1.5, and no periodontitis"
	Exclusion criteria: quote: "Exclusion criteria were allergic reactions to any of the components of any mouth rinse, antibiotic/antiphlogistic medication in the previous 8 weeks, dental-treatment needs (e.g. due to acute carious lesions), pregnancy or any kind of systemic diseases"



Hoffmann 2001 (Continued)

Baseline gingivitis: not reported but had to be <1.5 for inclusion in the study

Age at baseline (years): mean age is not reported. Age range 18-36 years

Gender: males 28/81 (34.6%), females 53/81 (65.4%)

Number randomised: 81 (CHX 0.10% 16, CHX 0.06% 17, CHX 0.06%/F 17, ASF 16, control 15)

Number evaluated: 73 (CHX 0.10% 14, CHX 0.06% 16, CHX 0.06%/F 17, ASF 15, control 11)

Interventions

Comparison: CHX 0.10% versus CHX 0.06% versus CHX 0.06%/F versus ASF* versus control

- CHX 0.10% (Chlorhexamed) (n = 16): 15 mL, twice daily, 30 seconds, 6 months
- CHX 0.06% (Corsodyl Zahnfleisch Fluid) (n = 17): 10 mL, twice daily, 30 seconds, 6 months
- CHX 0.06%/F (Odol med3 Depot) (n = 17): 10 mL, twice daily, 30 seconds, 6 months
- *ASF (Meridol) (n = 16): 10 mL, once daily, 30 seconds, 6 months. We excluded this arm from our risk of bias and analysis
- Control (water) (n = 15): 10 mL, twice daily, 30 seconds, 6 months

Prophylaxis at baseline: professional oral hygiene was performed 2 weeks before baseline and at baseline teeth were polished again

OHI: quote: "At the time of selection (day -14) all participants received oral-hygiene instructions." However, they were then advised to follow their habitual tooth cleaning procedure using the new toothbrush provided and their usual toothpaste for 2 weeks. At each follow-up no attempt was made to influence participants' other cleaning habits

Assumed to be non-supervised

Timing of mouthrinsing in relation to toothbrushing: after brushing

Outcomes

Gingivitis (gingival inflammation: GI, Löe & Silness 1967); plaque (PI, Silness & Löe 1964); tooth staining (Discolouration Index (Lang & Räber 1981) assessed at day -14, at baseline and at 1, 2, 3, and 6 months

Funding

Not reported

Notes

Sample size calculation: not reported

Adverse effects: Quotes: "One subject in the control group, however, had to be taken out because of leukoplakia..., no dropout occurred due to side effects of any of the mouthwash preparations" and in relation to tooth staining "Except the 0.06% CHX, all test solutions showed significantly higher DI scores when compared to the control... However from a clinical point of view, regarding the DI, no important differences between the active groups were seen" and "the 0.1% resulted in stronger staining than the ASF and the 0.06% CHX.."

Declarations/conflicts of interest: none stated

Bias	Authors' judgement	t Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Quote: "According to their clinical parameters and their sex, the 85 panellists were distributed at random and stratified to the 5 test groups"	
		Comment: no description of the method of random sequence generation	
Allocation concealment (selection bias)	Unclear risk	No information is provided	



Hoffmann 2001 (Continued)

Blinding of participants and personnel (performance bias) All outcomes High risk

The authors report this to be a "blind clinical study"

Quote: "The different test solutions were presented in similar bottles. However each participant got his/her specific information regarding the use of mouthwashes from an independent person not involved in the examinations"

The control rinse was tap water which the authors recognise as an issue (page 93). Quote: "After investigation it was revealed that some of the participants of the control group did not understand the scientific sense of rinsing with tap water"

The 0.1% CHX group rinsed with 15 mL and the other groups rinsed with 10 mL

In relation to tooth staining, quotes: "Except the 0.06% CHX, all test solutions showed significantly higher DI scores when compared to the control.. However from a clinical point of view, regarding the DI, no important differences between the active groups were seen" and "the 0.1% resulted in stronger staining than the ASF and the 0.06% CHX.."

Comment: the control group cannot be considered blind. The higher level of staining in the active groups (ASF and CHX) meant that participants could have worked out that they were in 1 of the active mouthrinse groups and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is unclear

Blinding of outcome assessment (detection bias) All outcomes

High risk

The authors report this to be a "blind clinical study"

Quote: "One investigator assessed PII and GI, while another estimated DI during the study"

In relation to tooth staining, quotes: "Except the 0.06% CHX, all test solutions showed significantly higher DI scores when compared to the control... However from a clinical point of view, regarding the DI, no important differences between the active groups were seen" and "the 0.1% resulted in stronger staining than the ASF and the 0.06% CHX.."

Comment: the higher level of staining in the active groups meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. Even though a separate examiner was used to record PI and GI and the DI the examiners recording the PI and GI would still be aware of the increased tooth staining. The direction of this potential bias is not clear

Incomplete outcome data (attrition bias)
All outcomes

High risk

85 individuals were randomised to groups but only 81 were "included in the study". The total number originally allocated to each group (from the 85) is not reported therefore 85 is used as the denominator for the calculation of overall attrition and 81 is used as the denominator for calculating the attrition in each group

Attrition at 6 months 8/85 (9.4%). Attrition by group: CHX 0.1% 2/16 (12.5%), CHX 0.06% 1/17 (5.9%), CHX 0.06%/F 0%, ASF 1/16 (6.3%), control 4/15 (26.7%). Reasons for loss to follow-up: quote: "One subject in the control group.. had to be taken out because of leukoplakia; two others did not show up at the last examination. Another two losses were in the 0.1% CHX group, but these cases were not due to any adverse effects. Furthermore, two losses occurred because of antibiotic medication during this long experimental time"

Comment: there were differential losses between the groups with a notably higher loss in the control group. This is conjunction with the loss of 4 participants before the study started indicate an increased risk of bias



Hoffmann 2001 (Continued)			
Selective reporting (reporting bias)	High risk	Results for all outcomes were reported in text at 3 and 6 months and at all tir points in box plot diagrams. Median values only were reported and no information on the variance of the scores was provided	
Other bias	Unclear risk	There is no mention of training and calibration of the 2 examiners	
		There were more females than males in each group at baseline ranging from 66% in the control group to 70% in both 0.06% CHX groups. It is reported that there were no differences between the groups at baseline for the 3 clinical parameters	
		No other baseline data are reported by group	

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Interventions

Trial design: parallel 3 arms

Location: Sri Laksmi Narasimha High School, Bangalore, India

Number of centres: 1

Study duration: the recruitment period is not reported. Study duration and duration of rinsing was 6

months

Participants Participants: school children (from Sri Laksmi Narasimha High School) in Bangalore city

Inclusion criteria: quote: "Inclusion criteria involved written informed consent from parents, good general health of children, a minimum of 12 gradable teeth, agreement to delay any elective dental treatment, including oral prophylaxis during the course of the study, and the agreement to comply with the study visits and procedures"

Exclusion criteria: quote: "Exclusion criteria involved antibiotic therapy, history of early-onset periodontitis, acute necrotizing ulcerative gingivitis, gross oral pathology, treatment for cancer or seizure disorders and conditions that interfered with the examination procedures"

Baseline gingivitis: (GI, Löe & Silness 1963) CHX 0.0536, CHX/NaF 0.0562, placebo 0.0609. SD not reported

Age at baseline (years): mean age not reported. Age range 13-16 years

Gender: not reported

Number randomised: 150 (numbers allocated to each group not reported, assumed 50 in each group at baseline)

Number evaluated: not reported

Comparison: CHX rinse versus CHX/NaF rinse versus placebo rinse

Gp A (n = assumed to be 50) CHX 0.05%: 10 mL, once daily, 6 months

Gp B (n = assumed to be 50) CHX/NaF 0.05%: 10 mL, once daily, 6 months

Gp C (n = assumed to be 50) placebo: 10 mL, once daily, 6 months

Rinse duration not reported

Prophylaxis at baseline: quote: "...no oral prophylaxis ... were given"

OHI: quote: "no ...oral hygiene instructions were given"



Jayaprakash 2007 (Continued)	Supervised rinsing Timing of mouthrinsing in relation to toothbrushing: not stated, however rinsing was done in school likely at a different time to toothbrushing Postrinsing instructions: instructed not to eat or drink anything for an hour after using the mouthrins	
Outcomes	Gingivitis (gingival inflammation: GI, Löe & Silness 1963) and plaque (PI, Silness & Löe 1964), assessed after 1, 3, and 6 months	
Funding	Not reported	
Notes	Sample size calculation: not reported Adverse effects: not reported Declarations/conflicts of interest: not reported Gingivitis data for Gp A and Gp B were combined for the main analysis	
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Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "The students were randomly numbered 1-150 and the mouthwash samples were numbered randomly 1-150 by the mouthrinse manufacturer the students who were assigned with particular numbers were provided with the mouthwash with the same number"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Low risk	Quote: "The students were randomly numbered 1-150 and the mouthwash samples were numbered randomly 1-150 by the mouthrinse manufacturer the students who were assigned with particular numbers were provided with the mouthwash with the same number"
		Comment: coding of mouthrinse samples done by manufacturer. We consider it unlikely that participants or investigators enrolling participants could foresee assignments
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	The study is reported to be "double-blind". The investigator supervised the mouthrinsing performed by the students daily. No information is provided on the composition of the placebo. Tooth staining is not mentioned
		Comment: tooth staining is likely to have been noticeable in both CHX groups compared to placebo during this 6-month study and in this situation participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. However, tooth staining is not reported and there is not enough information upon which to base a judgement
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	The study is reported to be "double-blind". The investigator supervised the mouthrinsing performed by the students daily. Tooth staining is not mentioned
		Comment: tooth staining is likely to have been noticeable in both CHX groups compared to placebo during this 6-month study and in this situation the outcome assessor could have worked out which individuals were in the placebo group and not be adequately blinded. However, tooth staining is not reported and there is not enough information upon which to base a judgement



Jayaprakash 2007 (Continued)			
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The number in each group at the start and end of the study is not reported	
Selective reporting (reporting bias)	High risk	Mean scores for gingivitis and plaque are reported without any measure of variance of the mean scores. We estimated the SD for gingivitis (measured using the GI) as the median of the SDs in the placebo/control groups from similar studies that also used the GI at 6 months so that we could include the gingivitis data in the meta-analyses. Furthermore, adverse effects including tooth staining are considered to be important outcomes but are not reported	
Other bias	Unclear risk	Outcomes were assessed by 1 individual. Details of training and intraexaminer reliability are not reported Baseline demographic data for the participants are not reported	

Jenkins 1993

Methods	Trial design: parallel, 2 arms
	Location: dental clinic, University of Wales College of Medicine, Cardiff, Wales, UK
	Number of centres: 1
	Study duration: recruitment period not stated. Study duration and the duration of rinsing was 6 weeks
Participants	Participants: adults with gingivitis (employees of the University of Wales College of Medicine and South Glamorgan Area Health Authority)
	Inclusion criteria: between 18 and 60 years old; minimum 16 natural teeth; minimum Löe & Silness GI score of 0.5
	Exclusion criteria: wearing dental prosthesis or appliance; advanced restorative dental work; medical or pharmacotherapy history affecting the study
	Baseline gingivitis: (Löe & Silness GI) Gp A: mean 0.89 (SD 0.24); Gp B mean 0.88 (SD 0.21) - means read from graph (Figure 3)
	Age at baseline (years): 18 to 60 (not reported by group)
	Gender: not reported
	Number randomised: 102 (Gp A: 51; Gp B: 51)
	Number evaluated: 99 (Gp A: 49; Gp B: 50)
Interventions	Comparison: (CHX gluconate + sodium fluoride (NaF) rinse) versus (placebo rinse (without CHX or fluoride))
	Gp A (n = 51): CHX (0.12%) plus NaF (100 ppm F), 15 mL, 1 min, twice daily (morning and evening)
	Gp B (n = 51): same routine using placebo
	Prophylaxis at baseline: none but could request and receive at any time a polishing of the anterior teeth if they perceived a cosmetic problem
	OHI: none, but all participants received same toothbrush and fluoride toothpaste but the use of any oral hygiene products other than those supplied was not permitted
	Non-supervised rinsing



Jenkins 1993 (Continued)	
,	Timing of mouthrinsing in relation to toothbrushing: not reported
	Postrinsing instructions: not reported
Outcomes	Gingivitis (gingival inflammation: GI Löe & Silness 1963 "modified to standardize the probing force, to elicit bleeding, using a constant pressure probe set at 25 g"); plaque (PI, TQH, Turesky 1970); stain area (modification of Shaw & Murray stain index 1977, stain severity graded according to colour (0 to 5 scale)), all measured at 6 weeks Problems associated with the use of rinses, particularly due to staining were elicited by questionnaire
Funding	Quote: "The authors are grateful to Smith Kline Beecham for supporting this study"
Notes	Sample size calculation: not reported
	Adverse effects: quotes: "Mucosal erosions were clinically apparent in two subjects using the active rinse and a further 2 subjects in this group reported mucosal soreness" and "soreness tingling and taste disturbance were more commonly reported for the active rinse and the difference between the groups reached statistical significance for taste disturbance ($P < 0.05$). Significantly more subjects (49%) using the active rinse were noted to have staining than using the control rinse (4%) ($P < 0.001$). Also 82% of the active group reported staining compared with 14% in the control group ($P < 0.001$). 2/3 of those reporting staining on active treatment classed it as quite bad or unacceptable, compared to none on the control rinse ($P < 0.001$). In response to the question of liking the taste of the mouthrinse coded as yes = 1, equivocal = 1.5 and no = 2 there was a clear preference for the control rinse ($P < 0.001$)"
	Declarations/conflicts of interest: not reported

Bias	Authors' judgement	Support for judgement
Random sequence genera-	Unclear risk	Quote: "volunteers randomly allocated to the active or control rinse"
tion (selection bias)		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Low risk	Quote: "volunteers were dispensed the allocated rinse through the Dental Hospital Pharmacy where a sealed code-breaker was kept"
		Comment: we consider it unlikely that participants and investigators enrolling participants could foresee assignment
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	The study is reported to be "double-blind"
		The control rinse was identical to the test rinse "except that it contained neither chlorhexidine or fluoride"
		Quote: "Significantly more subjects (49%) using the active rinse were noted to have staining than using the control rinse (4%) (P < 0.001)"
		Comment: the higher level of staining in the CHX group meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is not clear
Blinding of outcome assessment (detection bias) All outcomes	High risk	The study is reported to be "double-blind". Quote: "Significantly more subjects (49%) using the active rinse were noted to have staining than using the control rinse (4%) $(P < 0.001)$ "
		Comment: the authors do not report precisely who was blinded, but even if this includes the outcomes assessors the higher incidence of staining in the active group meant that they could not be adequately blinded and this could



Jenkins 1993 (Continued)		have affected the outcome assessment. The direction of this potential bias is not clear
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition after 6 weeks was 3/102 (2.9%). Attrition by group: 2/51 (3.9%) CHX 0.12%, 1/51 (2.0%) control. Reasons for losses not broken down by group: quote: "three subjects withdrew, 1 for reasons of taste of the active rinse and 2 for reasons unrelated to treatment"
		Comment: losses are very low and not considered sufficient to have any effect on the results
Selective reporting (reporting bias)	Low risk	Mean gingival and plaque data, stain area and intensity are reported graphically with SD (as a number on the graph). Mean scores were estimated from graphs. Stain area only was used in the meta-analysis
Other bias	Unclear risk	No information is provided on number of examiners, examiner training or intra- or interexaminer reproducibility so it is unclear whether or not there was a risk of differential diagnostic activity. Baseline participant characteristics (age, gender, etc) are not reported. There was no significant difference between the CHX and control groups for plaque, gingivitis, proportion of smokers and tooth staining at baseline

Jose 2015

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Trial design: parallel group (3 arms)

Location: Manchester and Wirral, UK, Clinical Research Facility

Number of centres: 2

Study duration: the recruitment period is not stated. Study duration and duration of rinsing was 6 weeks

Participants

Participants: healthy adults recruited from the sites' databases and via advertisements

Inclusion criteria: at the screening visit, subjects were asked to brush as they normally would for 1 timed minute and to expectorate into a white cup. Evidence of blood in the expectorant or bleeding while brushing were the initial inclusion criteria. Additional inclusion/exclusion criteria used were: ≥16 permanent, gradable teeth, including 4 molars (protocol amendment 3: changed from 20 gradable teeth); and ≥20 or more bleeding sites at the baseline examination

Exclusion criteria: women who were pregnant, lactating or those of child bearing potential not practising a reliable method of contraception were excluded from the study. Subjects who were taking antibiotics within 2 weeks before the screening visit or throughout the study were excluded, as were those on concomitant medication that, in the opinion of the investigator, might interfere with the outcome of the study. Other than having mild to moderate gingivitis, subjects were to be in good oral health with no active caries, no heavy calculus deposits and no more than 5 periodontal pockets measuring ≥5 mm in depth (protocol amendment 3: changed from more than 3 pockets >/- 5 mm in depth)

Baseline gingivitis: (GI, Löe & Silness) Gp A: mean 1.24 (SE 0.009); Gp B: mean 1.24 (SE 0.009); Gp C: mean 1.24 (SE 0.01)

Age (years): Gp A: mean 37.8; Gp B: mean 36.2; Gp C: 36.5. Range 18-63 years. Gp A: range 18-62; Gp B: range 18-63; Gp C: range: 18-56

Gender: overall: male 85 (26%), female 239 (74%). Gp A: male 23 (21%), female 85 (79%); Gp B: male 30 (28%), female 79 (73%); Gp C: male 32 (30%), female 75 (70%)

Number randomised: 324 (Gp A: 108; Gp B: 109; Gp C: 107)



Jose 2015 (Continued)

Number evaluated (ITT): 319 (Gp A: 105; Gp B: 109; Gp C: 105)

Interventions

Comparison: (CHX rinse with alcohol + standard fluoride toothpaste) versus (CHX rinse alcohol-free + standard fluoride toothpaste) versus (brushing alone with standard fluoride toothpaste)

Gp A (n = 108): CHX with alcohol 0.2%, 10 mL, twice daily, 1 min, 6 weeks

Gp B (n = 109): CHX alcohol-free 0.2%, 10 mL, twice daily, 1 min, 6 weeks

Gp C (n = 107): brushing alone (standard toothbrush and toothpaste)

All 3 groups brushed twice daily with full brush head of toothpaste for 1 timed minute

Prophylaxis at baseline: quote: "A complete dental prophylaxis, which included flossing to ensure removal of all plaque, was performed by an appropriately trained professional at the baseline visit or on a separate visit within 7 days of baseline"

OHI: the first use of the product/toothbrushing (1 timed minute) was supervised but it is not clear whether the participants received instructions on brushing technique. We assumed not

Rinsing was unsupervised (apart from the first use of the rinse)

Timing of mouthrinsing in relation to toothbrushing: quote: "subjects brushed their teeth with a full brush head of toothpaste for one timed minute, rinsed their mouth with water then waited for five timed minutes. After five minutes, they swished with 10 ml of their assigned mouthwash (mouthwash groups only) for a timed one minute followed by expectorating"

Duration of rinsing: 6 weeks

Outcomes

Gingivitis (gingival inflammation: GI, Löe & Silness 1963; gingival bleeding: Gingival Severity Index); plaque (Soparkar's modification of the TQH, Turesky 1970, Lobene 1982); oral soft tissue examination and CHX pharmacokinetic variables (from saliva samples) - not relevant to this review at 6 weeks follow-up

Funding

The study and writing support for the manuscript were funded by GSK Consumer Healthcare

Notes

The study protocol was registered at Clinicaltrials.gov (NCT01751178) and at www.gsk-clinicalstudyregister.com/study/RH01561#ps

Sample size calculation: not reported

Adverse effects: quote: "A total of 338 TEAEs (treatment emergent adverse events) were recorded among 174 (53.7%) of subjects (Table 4a). A large number of TRAEs (treatment related adverse events) were recorded in the two chlorhexidine groups where 27.8% of subjects in the 0.2% CHX-alcohol group and 24.8% in the 0.2% CHX-alcohol-free group reported at least one TRAE compared to only 3.7% in the brushing alone group. The most commonly reported TRAEs were coated tongue, glossodynia, oral paraesthesia, ageusia, dry mouth, oral hypoaesthesia and dysgeusia (Table 4b). There was one serious adverse event, not related to study product and one severe event (oral paresthesia) in the 0.2% CHX-alcohol group which did resolve. The subject withdrew from the study due to the event. A second subject withdrew from the study due to a non-TRAE"

Smoking status: overall smoker 43 (13%), non-smoker 281 (87%). Gp A: smoker 14 (13%), non-smoker 94 (87%); Gp B: smoker 15 (14%), non-smoker 94 (86%); Gp C: smoker 14 (13%), non-smoker 93 (87%)

Declarations/conflicts of interest: none stated, however the authors of the study are either employees of GSK Consumer Healthcare or employees of Intertek Clinical Research Services, Wirral and Manchester, UK which has received funding from GSK Consumer Healthcare

Groups A and B were combined in the main analysis



Jose 2015 (Continued)

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "Subjects were then stratified, according to the study site (Manchester or Wirral, UK), baseline number of bleeding sites and smoking status, and randomised into one of three treatment groups"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	The study is described as "examiner-blinded". All participants could not be blinded due to the design of the study which included a 'brushing alone' group
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	The study is described as "examiner-blinded". Adverse events are described but tooth staining was not measured or mentioned
		Comment: tooth staining is likely to have been noticeable in the CHX groups after 4 weeks, and in this situation the outcome assessor could have worked out which individuals used CHX and not be adequately blinded. However, tooth staining is not reported and there is not enough information upon which to base a judgement
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition (ITT rather than PP): 5/324 (1.5%). Gp A: 3/108 (2.8%), Gp B: no losses, Gp C: 2/107 (1.9%). Reasons for loss to follow-up: Gp A: 2 subjects with adverse events and 1 withdrew consent; Gp C: 2 subjects withdrew consent
		Comment: we do not believe the losses described could pose a risk of bias
Selective reporting (reporting bias)	High risk	All outcome stated in the methods section are described fully in the results. Tooth staining is considered to be an important side effect of CHX mouthrinsing but is not reported
Other bias	Low risk	Quote: "All assessments of GI were performed by a single, experienced examiner." Repeat assessment of PI were performed and showed excellent agreement (kappa value for PI 0.891 (95% CI 0.884 to -0.889).
		Baseline demographics and % bleeding sites at baseline appear balanced (Table 1). Mean Gingival Severity Index, GI and PI appear similar at baseline

Joyston-Bechal 1993

Methods	Trial design: parallel, 2 arms
	Location: Department of Oral Medicine and Periodontology, The London Hospital Medical College, UK
	Number of centres: 1
	Study duration: recruitment period is not stated. Study duration and duration of rinsing was 8 weeks
Participants	Participants: adults with gingivitis
	Inclusion criteria: over 18 years old; minimum 20 natural teeth; minimum Community Periodontal Index of Treatment Needs (CPITN) score of greater than 1 in at least 2 sextants but less than 4 in any sextant. This method of selection ensured that all subjects showed signs of gingivitis but no pockets greater than 5.5 mm in depth were present



Joyston-Bechal 1993 (Continued)

Exclusion criteria: medical history affecting the study; pregnancy or lactation; open carious lesions; partial dentures; orthodontic banding; current periodontal treatment; mentally or physically handicapped with impaired dexterity

Baseline gingivitis: not reported

Age at baseline (years): not reported. Mean age of completers at baseline (years): Gp A: mean 30.5 (SE 2.5); Gp B: mean 32.8 (SE 2.9)

Gender (completers): overall: male 38.46%; female 61.54%. Gp A: male 6 (31.6%), female 13 (68.4%); Gp B: male 9 (45.0%), female 11 (55.0%)

Number randomised: 47 (not reported by group)

Number evaluated: 39 (Gp A: 19; Gp B: 20)

(The text of the paper indicates that there were 20 individuals in the test group and 19 in the control group but Table 1 reports 19 individuals in the test group and 20 in control group. We have extracted the data presented in Table 1)

Interventions

Comparison: (CHX + sodium fluoride (NaF) rinse) versus (placebo rinse (similar colour and flavour but without CHX or fluoride))

Gp A (n = 19 analysed): CHX (0.05%) + 0.05% NaF (500 \pm 25 ppm NaF), 10 mL, 30 seconds, twice daily

Gp B (n = 20 analysed): same routine using placebo

Note: all subjects received the control mouthrinse for a run in period of 10 days to allow familiarisation with the requirements of the study

Prophylaxis at baseline: all teeth were scaled and polished on the first visit after assessments

OHI: subjects were asked to follow their routine dental hygiene practices, no oral hygiene instructions were given. All participants received the same toothbrush and fluoride toothpaste

Non-supervised rinsing

Timing of mouthrinsing in relation to toothbrushing: after brushing with toothpaste and rinsing with water

Postrinsing instructions: not reported

Outcomes

Gingivitis (gingival bleeding - Cowell 1975 with the following modification: the periodontal probe was inserted into the periodontal pockets to a depth of only 1-2 mm and not to the base of the pocket); plaque (PI, Silness & Löe 1964); degree of staining (Lobene Stain Index 1968), all measured at 8 weeks

Funding

Quote: "We are most grateful to Johnson and Johnson Limited for financial support for this study"

Notes

Sample size calculation: not reported

Adverse effects: not reported apart from tooth staining. The active rinse group had more staining than the control group at baseline before the scale and polish and more staining than the control at the end of the study

Smokers: by group: Gp A: 5/19 (26% of completers); Gp B: 7/20 (35% of completers) (differences between the groups are not statistically significant)

Declarations/conflicts of interest: not reported

|--|



Joyston-Bechal 1993 (Continue	ed)	
Random sequence generation (selection bias)	Unclear risk	Quote: "randomised into test and control groups"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	Quote: "The bottles were randomly numbered so that neither examiners or subjects were aware of the distribution of test or placebo solutions"
		Comment: not enough information is provided. It is not clear how the allocation sequence was concealed before participants were randomised to groups
Blinding of participants	High risk	The authors describe the study as "double-blind"
and personnel (perfor- mance bias) All outcomes		Quotes: "The bottles were randomly numbered so that neither examiners or subjects were aware of the distribution of the test or placebo solutions" and "The placebo solution was of similar colour and flavour but with no active components" and "Test and placebo solutions were supplied in identical bottles" Baseline assessments of staining were completed before the scale and polish and there was a statistically significant difference between the groups at baseline with the test group having a higher mean staining score at baseline and at the end of the study. The authors state that "subjects and examiners were unaware of any staining" but it is not clear how this information was obtained
		Comment: the active rinse group had more staining than the control group at baseline before the scale and polish and more staining than the control group at the end of the study. The higher incidence of staining in the active group meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is not clear
Blinding of outcome assessment (detection bias) All outcomes	High risk	Quotes: "double-blind study" and "The bottles were randomly numbered so that neither examiners or subjects were aware of the distribution of the test or placebo solutions" and "Test and placebo solutions were supplied in identical bottles" Baseline assessments of staining were completed before the scale and polish and there was a statistically significant difference between the groups at baseline with the test group having a higher mean staining score at baseline and at the end of the study. The authors state that "subjects and examiners were unaware of any staining" but it is not clear how this information was obtained
		Comment: the authors do not report precisely who was blinded, but even if this includes the outcomes assessors the higher level of staining in the active group meant that the outcome assessors could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is not clear
Incomplete outcome data (attrition bias)	High risk	Attrition after 8 weeks was 8/47 (17.0%). Losses not broken down by group and reasons for losses not reported
All outcomes		Comment: assuming the missing participants in 1 group had a higher mean (e.g. gingivitis score) than those in the other group, as the attrition rate increased, so would the mean difference between groups. Also, although attrition was not reported by group, it could be associated with the outcomes
Selective reporting (reporting bias)	High risk	Mean scores are reported graphically with a measure of variance (assumed to be SD) as a bar on the graphs. Mean scores and SDs were estimated from graphs so that these data could be included in meta-analyses. Apart from tooth staining, adverse effects are not mentioned
Other bias	Low risk	Outcome assessment at the initial visit and the final visit were by different examiners but their interexaminer reproducibility is reported and is adequate.



Joyston-Bechal 1993 (Continued)

There was a statistically significant difference between the groups at baseline with the test group having a higher mean staining score at baseline and at the end of the study. No other apparent sources of bias were identified

Lang 1998

Methods

Trial design: parallel (3 arms - 2 arms are relevant to this review)

Location: Berne, Switzerland. University Department of Periodontology and Fixed Prosthodontics

Number of centres: 1

Study duration: the recruitment period is not stated but the duration of rinsing is 6 months

Participants

Participants: males and females from a telecommunication company in Berne, Switzerland

Inclusion criteria: quote: "Inclusion criteria were: 18 years old and above, with ≥25% of all gingival sites (six sites around each tooth) with bleeding upon probing (BOP) using the Florida probe with a standardized force of 0.25N... at least 16 sound natural teeth without crowns, bridgework or ill-fitting dental restorations"

Exclusion criteria: quote: "Exclusion criteria were: more than four pockets ≥5 mm - third molars and the distal sites of the second molars not included, caries with cavities, known previous hypersensitivity reactions to drugs, alcohol or drug addiction, severe liver or renal disease, psychiatric disorders, severe disability with multiple drug therapy, pregnancy or planned pregnancy and breast feeding. Neither should patients have been on treatment with antibiotics within the last 6 weeks prior to the prestudy visit, nor should they have treatment with anti-inflammatory or anti-cholinergic drugs"

Baseline gingivitis: not reported

Age at baseline (years): mean ages (SD): delmopinol 39 (8) years; CHX 40 (12) years; placebo 43 (11) years. Range not stated

Gender: males: 85.8%, females: 14.2%

Number randomised: 162 (CHX 50, delmopinol 53, placebo 53) (Note: discrepancy between the total number entering the study and the number included in per protocol analysis in each group at baseline)

Number evaluated: 133 (CHX 41, delmopinol 47, placebo 45)

Interventions

Comparison: CHX rinse versus delmopinol rinse* versus placebo rinse

- CHX 0.2% (Hibitane) (n = 50): 10 mL, twice daily, 60 seconds, 6 months
- *Delmopinol (Decapinol) (n = 53): 10 mL, twice daily, 60 seconds, 6 months. We excluded this arm from our risk of bias and analysis
- Placebo (n = 53): 10 mL, twice daily, 60 seconds, 6 months

Prophylaxis at baseline: quote: "After baseline, but before starting to rinse, the patients underwent professional cleaning of the teeth including supra- and subgingival scaling"

OHI: none. Quote: "..patients were instructed to brush their teeth as usual... No oral hygiene instructions were given before or during the study"

Partially supervised rinsing (supervised on weekdays, unsupervised at weekends)

Timing of mouthrinsing in relation to toothbrushing: if rinsing took place in connection with toothbrushing the rinsing should always be performed after the mechanical cleaning of the teeth



Lang 1998 (Continued)

Outcomes

Gingivitis (gingival inflammation: GI, Löe 1963; BOP, bleeding yes/no Florida probe to bottom pocket, Löe 1967, Gibbs 1988); plaque (PI, Silness & Löe 1964); calculus (Calculus Surface Index / Calculus Surface Severity Index (CSI/CSSI), Ennever 1961, Conroy 1968); stain (DI, Lang & Räber 1981) assessed at baseline, 3 months and 6 months

Oral soft tissue examined and adverse effects reported during the study

Funding

Not reported

Notes

Sample size calculation: quote: "The sample size calculation was based on BOP. The standard deviation was estimated to 14 and a difference of 10 percentage units between placebo and delmopinol was considered worth detecting. The calculation which was based on a power of 95% and a significance level of P = 0.05, resulted in a required number of 50 subjects in each treatment group"

Adverse effects: quote: "No serious adverse events were reported in this study... one of the most frequent adverse events when rinsing with delmopinol was a transient anaesthetic sensation in various parts of the oral mucosa. This event was reported as 'anaesthesia local' or 'paraesthesia'... Taste affection ('taste loss', 'taste perversion') was reported by 35% and 20% of the patients in the delmopinol group after 3 and 6 months, respectively...The patients in the chlorhexidine group.. reported this event to the same extent, namely 35% after 3 months and 14% after 6 months.. Staining of the teeth and tongue was reported more often in the chlorhexidine group than the delmopinol group both at the 3-month (69% and 20% of patients in each respective group) and 6-month examination (88% and 16% in each respective group)... The most common adverse events causing withdrawal from treatment by the 10 patients rinsing with chlorhexidine were, staining of teeth, especially black staining (four patients), altered or bad taste (four patients), burning tongue (three patients) and black or brown staining of the tongue (three patients). The patient rinsing with delmopinol who was withdrawn from treatment due to adverse events reported anaesthesia of mucous membrane, burning tongue, dry mouth and taste alteration. The withdrawal patient in the placebo group due to adverse events reported bad taste, stomach problems, and tendency to vomit"

Declarations/conflicts of interest: none declared, however, 2 of the authors are associated with Biosurface Pharma AB

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Patients were allocated to treatment groups, according to a computer-generated randomization list, where treatment was balanced within blocks of six patients"
		Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	Quote: "Packaging and labelling were carried out at an independent Clinical Service Department in Uppsala, Sweden. The various test solutions were dis- pensed in identical 200 ml amber glass bottles"
		Comment: it is not clear whether the assignment of subjects to groups was carried out at the independent Clinical Service Department
Blinding of participants	High risk	The authors report the study to be "double-blind"
and personnel (performance bias) All outcomes		Quote: "The various test solutions were dispensed in identical 200 ml amber glass bottles." The placebo consisted "of the vehicle of active delmopinol solution except delmopinol and sodium hydroxide"
		Quote: "rinsing with chlorhexidine resulted in statistically significantly more staining than both placebo and delmopinol"
		Comment: while every effort was made to ensure blinding of participants and personnel, the higher level of staining of the tongue and teeth in the



Lang 1998 (Continued)		CHX group relative to the placebo group meant that participants could have worked out which group they were in and this could have affected their oral
		health behaviours and hence the outcomes. The direction of this potential bias is not clear
Blinding of outcome assessment (detection bias)	High risk	The authors report the study to be "double-blind"
All outcomes		Quotes: "The various test solutions were dispensed in identical 200ml amber glass bottles" and "Assessments were performed by trained examinersE ach examiner scored one type of parameter for all patients throughout the whole study" and "Rinsing with chlorhexidine resulted in statistically significantly more staining than both placebo and delmopinol" and "The decisions to exclude patients/visits were taken before breaking the blind"
		Comment: while every effort was made to ensure blinding of outcome assessment, the higher level of staining of the tongue and teeth in the CHX group relative to the placebo group meant that outcome assessors could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is not clear
Incomplete outcome data (attrition bias) All outcomes	High risk	Different data are reported for the number of patients excluded from the per protocol analysis (Table 2) and the number of patients withdrawn from treatment and reasons for withdrawal (Table 3). After 6 months (Table 2) 17/103 (16.5%) randomised participants were not included in the final analysis. Atrition by group (Table 2): CHX 9/50 (18%), placebo 8/53 (15.1%). Total number of patients withdrawn from treatment and reasons for withdrawals by group: CHX (n = 13, 24%) adverse events (n = 10) and refusal to cooperate (n = 3); placebo (n = 2, 4%), adverse events (n = 1), refusal to cooperate (n = 1)
		Comment: there are a high proportion of losses between the groups whichever source of information (Table 2/Table 3) is used and reasons for losses are unbalanced between the groups with a high proportion of the withdrawals in the CHX group due to adverse effects
Selective reporting (reporting bias)	Low risk	All outcomes stated in the methods section are adequately reported in the results section
		Information on means and SEs are presented graphically. From correspondence with the author, additional information was obtained on means and SEs which enabled calculation of SDs from SEs and inclusion of these data in the meta-analyses
Other bias	Unclear risk	Although this is an experienced research team and the examiners were trained there is no mention of calibration or reliability of outcome measurements
		Mean age appears similar across groups. 86% of participants were male but the breakdown for gender across groups is not reported
		16 and 10 patients in the CHX and placebo groups were using medication at the time of the premedical screening (P < 0.001)
		Assessments of GI were performed directly after BOP. Quote: "thus the possible traumatising effect on the gingival crevice may have affected the results of the GI"

Lopez-Jornet 2012

Methods Trial design: parallel (2 arms)



opez-Jornet 2012 (Continued)		
	Location: pensioners home in Murcia, Spain	
	Number of centres:1	
	Study duration: recruitment period not stated. Study duration and duration of rinsing was 30 days (with a 7-day whitening period before rinsing commenced)	
Participants	Participants: volunteers from a pensioners home in Murcia, Spain of either sex over 65 years of age	
	Inclusion criteria: subjects of either sex over 65 years of age, with removable dentures who upon clinical examination did not present manifestations of oral candidiasis in any of its forms (pseudomembra nous, erythematous, hyperplastic or angular cheilitis). None of the subjects had used adhesives for	
	their dentures, and all of them were in good general health	
	Exclusion criteria: smokers, patients with hypersensitivity or allergy to the study medication, individuals with oral mucosal disease and antibiotic treatment in the month prior to inclusion in the study	
	Baseline gingivitis: (GI Löe & Silness 1963) Gp A: CHX 1.51 \pm 0.98, Gp B: placebo 1.33 \pm 0.69	
	Age at baseline (years): mean 75 years (range 65-94)	
	Gender: total: 30 males (42.86%), 40 females (57.14%); CHX: 12 males (34.29%) and 23 females (65.71%); placebo: 18 males (51.43%) and 17 females (48.57)	
	Number randomised: 70 (Gp A: 35; Gp B: 35)	
	Number evaluated: 70 (Gp A: 35; Gp B: 35)	
Interventions	Comparison: CHX rinse versus placebo control rinse	
	Gp A: CHX 0.2%, 10 mL, 30 seconds, twice daily for 30 days	
	Gp B: placebo, same formulation without CHX 10 mL, 30 seconds, twice daily for 30 days	
	Prophylaxis at baseline: none	
	OHI: both groups were supplied with a whitening rinse and toothbrush with 0.05% fluoridated tooth-paste, and an instruction sheet and were advised to use for 7 days prior to recording of Silness & Löe P and GI. They also received instructions for accurate oral hygiene	
	Non-supervised rinsing	
	Timing of mouthrinsing in relation to toothbrushing: not reported	
	Postrinsing instructions: quote: "the subjects avoided rinsing with water, eating and drinking for 30 min following use of the mouthwash. They also received instructions for accurate oral hygiene, and were required to avoid all products other than those included in the study"	
Outcomes	Gingivitis (ginginal inflammation: GI Löe & Silness 1963) and plaque (Silness & Löe 1964) measured ter 30 days of rinsing	
	Microbiological evaluation of <i>Candida albicans</i> (colony forming units) measured at baseline and after 30 days of rinsing - not relevant to this review	
	Adverse effects: teeth/dentures/tongue staining, mucosal desquamation, alteration in taste sensation after 15 and 30 days of rinsing	
Funding	Not reported	
Notes	Sample size calculation: not reported	

Adverse effects: quote: "In reference to the presence of adverse effects, examination of the oral cavity in both groups revealed statistically significant differences after 15 days and at the end of the study (30



Lopez-Jornet 2012 (Continued)

days) in terms of staining of the teeth/dentures and tongue. No patient showed mucosal desquamation or alterations in taste sensation"

Number and % of participants with observed adverse effects at 30 days (Table 4): tongue staining: CHX: 11, 31.43%; placebo 8, 22.86. Denture/dental staining: CHX 2, 5.71%; placebo: 3, 8.57%

Non-smokers

Declarations/conflicts of interest: not reported

The GI data appear to be transposed (Table 2 page 295). These were entered in the opposite order in RevMan for analysis

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Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quotes: "randomized" and "the randomization list in the form of envelopes was generated using specifically designed software"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	Quotes: "randomized" and "the randomization list in the form of envelopes was generated using specifically designed software" and "both products were supplied in identical opaque bottles"
		Comment: mentioning the use of envelopes does not provide enough information to make a judgement. It is not clear how the allocation sequence was concealed before participants were randomised to groups
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quotes: "double-blind" and "both products were supplied in identical opaque bottles" and "both groups revealed statistically significant differences after 15 days and at the end of the study (30 days) in terms of staining of the teeth/dentures and tongue." The placebo was the same formulation as the test rinse without the CHX
		Comment: there does not appear to be significantly more staining in the CHX group compared to the placebo group
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quotes: "double-blind" and "All examinations were carried out by a single experienced examiner" and "both groups revealed statistically significant differences after 15 days and at the end of the study (30 days) in terms of staining of the teeth/dentures and tongue"
		Commet: there does not appear to be significantly more staining in the CHX group compared to the placebo group
Incomplete outcome data (attrition bias) All outcomes	Low risk	There were no losses to follow-up
Selective reporting (reporting bias)	Low risk	All outcomes stated in the methods section are reported in the results section
Other bias	Unclear risk	The groups were comparable at baseline in relation to age, gender, gingivitis, plaque and colony forming units of <i>Candida albicans</i>
		Denture/dental staining was not reported at baseline only at 15 and 30 days
		Quote: "All examinations were carried out by a single experienced examiner"



Lopez-Jornet 2012 (Continued)

Comment: intraexaminer reliability is not reported

Lucas 1999	
Methods	Trial design: parallel (4 arms - 2 arms are relevant to this review)
	Location: Faculty of Dentistry, National University of Nordeste, Corrientes, Argentina
	Number of centres: 1
	Study duration: the duration of recruitment is not stated. 2 studies are reported; the first was a 21-day study with participants abstaining from all oral hygiene practices, the second involved the use of CHX or placebo rinse as an adjunct to oral hygiene for 180 days. The second study is relevant to this review and is described here
Participants	Participants: dental students
	Inclusion criteria: quote: "All had at least sixteen natural teeth including four molars"
	Exclusion criteria: quote: "Patients affected with systemic diseases or on regular medication were excluded"
	Baseline gingivitis: mean CHX: 0, mean placebo: 0
	Age at baseline (years): mean: 20.9 years. Range not reported
	Gender: male: 40%, female: 60%
	Number randomised: 20 (Gp A: 10; Gp B: 10)
	Number evaluated: not explicitly stated but appears to be 20 (Gp A: 10; Gp B: 10)
Interventions	Comparison: CHX mouthrinse versus placebo mouthrinse
	Gp A (n = 10): CHX mouthrinse (0.12%), 10 mL, twice daily, 180 days
	Gp B (n = 10): placebo mouthrinse, 10 mL, twice daily, 180 days
	The rinse duration was not reported
	Prophylaxis at baseline: quote: "Clinically normal gingiva was obtained by scaling, polishing and twice daily tooth brushing"
	OHI: quote: "Each subject participating in this study received regular instruction and reinforcement in personal plaque control procedures throughout the entire pre-experimental phase"
	Assumed to be non-supervised rinsing Timing of mouthrinsing in relation to toothbrushing: participants were advised to rinse 30 min after toothbrushing
Outcomes	Gingivitis (gingival inflammation: GI, Löe & Silness 1967, severity (mean GI score of entire mouth) and occurrence (proportion of surfaces with GI ≥ 1), gingival bleeding: GI, Löe & Silness 1967 (proportion of surfaces with GI 2 or 3)); plaque (PI, Greene & Vermillion 1964); effect on oral tissues assessed at days 0, 90 and 180 days
Funding	Not reported
Notes	Sample size calculation: not reported



Lucas 1999 (Continued)

Adverse effects: quotes: "The clinical and tactile evaluation of the oral tissues throughout the experimental period did not reveal any type of irreversible pathology associated with the use of chlorhexidine" and "some extrinsic staining was observed in the chlorhexidine group"

Non-smokers

Declarations/conflicts of interest: none stated. CHX and placebo were kindly supplied by the Microsules Bernabo Laboratories, Buenos Aires, Argentina

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Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence genera-	Unclear risk	Quote: "The subjects were divided at random into four groups"
tion (selection bias)		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "Neither the experimental subjects nor the examiner responsible for the clinical examinations had previous knowledge of what group was rinsing with chlorhexidine or placebo. (Chlorhexidine and placebo rinses were kindly supplied with a code number which was revealed after all data collection was completed and tabulated.)" The composition of the placebo is not reported. Although tooth staining and calculus were not measured the authors note that "some extrinsic tooth staining was observed in the chlorhexidine group" and that "Supraginigival calculi were more numerous in the group using chlorhexidine"
		Comment: the higher level of tooth staining and calculus in the CHX group meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is unclear
Blinding of outcome assessment (detection bias) All outcomes	High risk	Quote: "Neither the experimental subjects nor the examiner responsible for the clinical examinations had previous knowledge of what group was rinsing with chlorhexidine or placebo. (Chlorhexidine and placebo rinses were kindly supplied with a code number which was revealed after all data collection was completed and tabulated)"
		Although tooth staining and calculus were not measured the authors note that "some extrinsic tooth staining was observed in the chlorhexidine group" and that "Supraginigival calculi were more numerous in the group using chlorhexidine"
		Comment: the higher level of staining in the CHX group meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear
Incomplete outcome data (attrition bias) All outcomes	Low risk	The number of participants at 180 days is not explicitly stated and the results are presented by number of surfaces not number of participants. However it would seem that all the participants were included in the data at 180 days
Selective reporting (reporting bias)	High risk	Reported numbers represent number of surfaces of teeth evaluated rather than the number of individuals. No measures of variance were reported for either outcome. We estimated the SD for gingivitis (measured using the GI) as the median of the SDs in the placebo/control groups from similar studies that also used the GI at 6 months so that we could include the gingivitis data in the



Lucas 1999 (Continued)		meta-analyses. Tooth staining is considered to be an important outcome but this is not adequately reported			
Other bias	Unclear risk	Training of the single examiner and intraexaminer reliability are not mentioned. The baseline demographics of the groups are not reported. There is noway to determine if the groups were similar at the start of the study			
dayawa 1000					
Methods	Trial design: paral	llel, (3 arms - 2 arms relevant to this review)			
		ental clinic, State University of Ponta Grossa, Brazil			
	Number of centres				
	Study duration: re days	ecruitment period is not stated but the study duration and duration of rinsing was 42			
Participants	Participants: males aged between 14 and 35 years				
	Inclusion criteria: minimum 16 permanent teeth (including 4 molars, but excluding teeth with excessive decay, crowns, bridges, fixed orthodontic appliances, and third molars)				
	Exclusion criteria: active periodontal disease; deposits of calculus; use of antibiotics, anti-inflammatory, and antibacterial or other drugs that could influence the study				
	Baseline gingivitis (completers): (GI, Löe & Silness 1963) Gp A: mean 0.043 (SE 0.021); Gp B: mean 0.065 (SE 0.026); Gp C: mean 0.040 (SE 0.022)				
	Age at baseline (years): range 14 to 35 (not reported by group)				
	Gender: males only				
	Number randomised: not reported				
	Number evaluated: 31 (Gp A: 9; Gp B: 11; Gp C: 11)				
Interventions	Comparison: CHX	K digluconate rinse versus Plantago major rinse* versus placebo rinse			
	Gp A (n = 9 analyse	ed): CHX (0.12%), 10 mL, once daily before bedtime			
	* Gp B (n = 11 analysed): same routine using Plantago major. We excluded this arm from our risk of bias and analysis				
	Gp B (n = 11 analysed): same routine using placebo				
	Prophylaxis at baseline: participants had a thorough prophylaxis to remove plaque				
	OHI: yes. Quote: "Patients were subjected to a regular program of plaque control not supervised and instructed to swish 10 mls of solution daily before bedtime"				
	Non-supervised rinsing				
	Timing of mouthri	insing in relation to toothbrushing: not reported			
	De atripais a inatro	Destriction in the state and was not considered			

Gingivitis (gingival inflammation: GI, Löe & Silness 1963); plaque (PI, TQH, Turesky 1970); tooth staining

(Lobene Stain Index 1968); adverse events (soft tissues), measured at 3 and 6 weeks follow-up

Not reported

Outcomes

Funding

Postrinsing instructions: not reported



Navarro 1998 (Continued)

Notes Sample size calculation: not reported

Adverse effects: 1 person reported epithelial desquamation in the CHX group

Declarations/conflicts of interest: unclear from translation

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "Patients were randomized into treatment group, positive control or placebo"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "Patients were randomised into treatment group, positive control or placebo following a double-blind scheme." A placebo solution containing distilled water, essence, colorant and methylparaben 0.1% was used. The CHX group showed an increase in tooth pigmentation at 6 months
		Comment: the higher incidence of staining in the CHX group meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is not clear
Blinding of outcome assessment (detection bias)	High risk	Quote: "Patients were randomised into treatment group, positive control or placebo following a double-blind scheme." No further information is given
All outcomes		Comment: the authors do not report precisely who was blinded, but even if this includes the outcomes assessors the higher level of staining in the active group meant that they may not have been adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is not clear
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The number of patients at baseline is not reported. Results are presented for the 31 subjects who completed the study. The authors report that 1 person in the CHX group left the study between 0 and 21 days due to epithelial desquamation and that others withdrew from the study citing personal reasons
		Comment: not enough information is provided
Selective reporting (reporting bias)	Low risk	All outcomes stated in the methods section are adequately reported in the results section. We calculated SDs from the SEs reported so that the data could be included in meta-analyses
Other bias	Unclear risk	No information is provided on intra- and interexaminer reproducibility. At baseline the groups had similar mean gingivitis, plaque and staining scores

Overholser 1990

Methods Trial design: parallel (3 arms - 2 arms are relevant to this review)

Location: assumed to be at the Baltimore College of Dental Surgery, Dental School, University of Mary-

land at Baltimore, USA



Overholser 1990	(Continued)
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Number of centres: 1

Study duration: the recruitment period is not stated. Study duration and duration of rinsing was 6 months

Participants

Participants: healthy adults

Inclusion criteria: quote: "...with pre-existing plaque and gingivitis but without clinical evidence of periodontitis... minimum of 20 sound natural teeth, and plaque and gingival index means ≥1.95 as determined by the Turesky modification of the Quigley-Hein index (Turesky et al 1970) and the modified gingival index (Lobene et al 1986), respectively"

Exclusion criteria: quote: "Grossly carious, fully crowned, orthodontically banded, abutment and third molar teeth were not included. Subjects with gross oral pathology or on antibiotic, antibacterial or anti-inflammatory therapy were excluded from the study"

Baseline gingivitis: (modified GI) Gp A: mean 2.28 (SE 0.03); Gp B: mean 2.23 (SE 0.02); Gp C: mean 2.22 (SE 0.02)

Age of completers (years): Gp A: mean 29.24; Gp B: mean 29.17; Gp C: mean 28.62. Range 21-62 years. Gp A: range 23-62; Gp B: range 21-47; Gp C: range 22-42

Gender (completers): overall: male 44 (36%), female 80 (65%). Gp A: male 22 (54%), female 19 (46%); Gp B: male 10 (24%), female 31 (76%); Gp C: male 12 (29%), female 30 (71%)

Number randomised: 128 (not reported by group)

Number evaluated: 124 (Gp A: 41; Gp B: 41; Gp C: 42)

Interventions

Comparison: CHX rinse versus Listerine rinse* versus control

Gp A (n completers = 41): CHX (Peridex) 0.12%,15 mL, twice daily, 30 seconds, 6 months

*Gp B (n completers = 41): Listerine, 20 mL, twice daily, 30 seconds, 6 months. We excluded this arm from our risk of bias and analysis

Gp C (n completers = 42): control (flavoured coloured 5% hydroalcohol solution), 20 mL, twice daily, 30 seconds, 6 months

Prophylaxis at baseline: quote: "Following the baseline examination, patients were given a complete prophylaxis to remove all plaque, calculus and extrinsic stain"

OHI: none given. Subjects followed their usual oral hygiene and dietary habits, but were instructed to refrain from using other mouthrinses

Rinsings were supervised twice daily on weekdays

Timing of mouthrinsing in relation to toothbrushing: not reported

Postrinsing instructions: quote: "Subjects were instructed not to rinse, eat or drink for 30 mins following use of the test rinses"

Outcomes

Gingivitis (gingival inflammation: modified GI, no reference; gingival bleeding: Interdental Bleeding Index, Caton & Polson 1985); plaque (TQH, Turesky 1970); extrinsic tooth stain (Lobene Stain Index, Lobene 1968); supragingival calculus (Volpe 1965, Manhold 1965, Volpe 1967); assessed at 3 and 6 months follow-up. Plaque was collected for microbiological study (reported separately)

Funding

Quote: "This study was supported by a grant from the Warner-Lambert Company"

Notes

Sample size calculation: not reported

Adverse effects: no abnormal soft tissue findings were noted in any group. Quote: "The PX group, showed significant increases (P<0.001) in stain at 3 and 6 months, compared to its baseline"

Smokers: by group: Gp A: 5/41 (12%); Gp B: 2/41 (5%); Gp C: 6/42 (14%)



Overholser 1990 (Continued)

Declarations/conflicts of interest: none stated

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RISK OF DIAS		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Subjects were assigned to groups according to a computer-generated random code"
		Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Low risk	Product was dispensed in coded amber bottles. Product code was not disclosed to the examiners or recorded on case report forms
		Comment: we consider it unlikely that participants and investigators enrolling participants could foresee assignment
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	The authors state that the study is double-blind. The products were dispensed in coded amber bottles. The placebo control was a flavoured, coloured 5% hydroalcohol solution. Personnel dispensing rinses did not participate in any other aspect of the study. Quote: "Significantly greater (P<0.01) extrinsic stain was observed in the PX group when compared to LA and the control at both 3 and 6 months"
		Comment: the higher level of tooth staining in the CHX group compared to the other 2 groups meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is not clear
Blinding of outcome assessment (detection bias) All outcomes	High risk	The authors state that the study is double-blind. The products were dispensed in coded amber bottles. Quotes: "Product code was not disclosed to the exam iners or recorded on case report forms" and "Subjects refrained from all oral hygiene and use of experimental products on examination days until after the examination were completed in order to eliminate possible bias due to product odor" Separate forms were used to record the outcome data at each exam ination interval and the examiner did not have access to case report forms during the study. To assure maximum objectivity and reproducibility, 1 examiner was used for gingivitis, bleeding and plaque indices and another for the stain and calculus indices. Quote: "Significantly greater (P<0.01) extrinsic stain was observed in the PX group when compared to LA and the control at both 3 and months"
		Comment: despite the considerable efforts made to ensure blinding of outcome assessment, the higher level of staining in the CHX group meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is not clear
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition at 6 months 4/128 (3.1%). Losses not broken down by group. 3 subjects were lost because they were unable to comply with the supervised rinsing schedule. 1 subject in the Peridex group dropped out when stain began to form
		Comment: attrition is very low over the 6 months of the study
Selective reporting (reporting bias)	Low risk	All outcomes stated in the methods section were addressed in the results with the exception of microbiological plaque analysis which is reported in a separate paper. We calculated SDs from the SEs reported so that the data could be included in meta-analyses
Other bias	Low risk	Efforts were made to standardise examiners in a series of sessions in which the examiners reviewed the clinical criteria in each index prior to the initiation of



Overholser 1990 (Continued)

the study. There is no indication of inter- or intraexaminer reliability. To assure maximum objectivity and reproducibility, 1 examiner was used for gingivitis, bleeding and plaque indices and another for the stain and calculus indices

Comment: the standardisation sessions and use of 1 examiner to record each index would reduce inter- and intraexaminer variability.

Quote: "The treatment groups were well-balanced with respect to age and smoking status." The authors report a statistically significant imbalance in the distribution of sexes within groups. The treatment-by-sex interaction was investigated and was not significant. The groups appear balanced for gingival index, plaque scores, bleeding index, supragingival calculus and tooth staining at baseline (although no P values for the differences are reported)

Pereira 2011

Methods

Trial design: parallel (3 arms - 2 are relevant to this review)

Location: Department of Periodontics, Dental School, University of Fortaleza, Brazil

Number of centres: 1 centre

Study duration: recruitment period not stated. Study duration and duration of rinsing was 90 days

Participants

Participants: adult subjects from the University of Fortaleza

Inclusion criteria: quote: "Bleeding index > 20%, presence of at least 20 natural teeth and absence of supragingival calculus and other plaque retentive factors, such as carious cavity and restoration excess"

Exclusion criteria: quote: "Participants with medical disorders and under antimicrobial therapy, as well as smokers, pregnant women and individuals presenting a probing depth > 3 mm were excluded from the trial"

Baseline gingivitis: bleeding index, Gp A (0.27 \pm 0.02); Gp B (0.22 \pm 0.04); Gp C (0.23 \pm 0.07)

Age at baseline (years): mean age not reported. Age range 27 to 42 years (not reported by group)

Gender: male: 15 (50%), female: 15 (50%) (not reported by group)

Number randomised: 30 (Gp A: 10; Gp B: 10 and Gp C: 10)

Number evaluated: 30 (Gp A: 10; Gp B: 10 and Gp C: 10)

Interventions

Comparison: CHX rinse versus Ocimum gratissimum rinse* versus placebo control rinse

Gp A (n = 10): CHX (0.12%) 10 mL, 1 min, 3 times daily

*Gp B (n = 10): Ocimum gratissimum (Og) 10 mL, 1 min, 3 times daily. We excluded this arm from our risk of bias and analysis

Gp C (n = 10): triethanolamine (qsp), alcohol, water (qsp), nipagin (0.2%), glycerine (2.5%), aspartame (qsp); 10 mL, 1 min, 3 times daily

Prophylaxis at baseline: quote: "After the initial examination all teeth of each subject were polished with a pumice and flossed to eliminate plaque remnants"

OHI: quote: "A personal 'kit' containing a new toothbrush, a commercial dentifrice with no anti-inflammatory properties and tests or control mouth rinses was given to all participants. They were instructed to brush their teeth for 1 minute, three times a day, using their habitual technique"

Non-supervised rinsing



Pereira 2011 (Continued)			
	Timing of mouthrinsin rinsed with one of the	g in relation to toothbrushing: quote: "After each brushing the participants formulations"	
	Postrinsing instruction	ns: not reported	
	Duration of rinsing: 90	days	
Outcomes	Gingivitis (bleeding index (BOP), Ainamo & Bay 1975); plaque (TQH, Turesky 1970); hard and soft oral tissues for the presence of any adverse reaction. All at 90 days		
Funding	Quote: "Source of support – Nil"		
Notes	Sample size calculation: not reported		
	Adverse effects: quote: "The tests mouth rinses had a good acceptance and did not show formations of abscess, ulceration or allergic reactions. However, three volunteers from CLX (CHX) group reported pigmentation and temporary taste disturbance"		
	Non-smokers		
	Declarations/conflicts of interest: quote: "Conflict of interest: none declared"		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera-	Unclear risk	The participants were assigned to groups by random permutations of 3	
tion (selection bias)		Comment: no description of the method of random sequence generation	
Allocation concealment (selection bias)	Unclear risk	No information is provided	
Blinding of participants and personnel (performance bias)	High risk	Quotes: "double-blind" and "The bottles were previously coded to warrant that neither the examiner nor the participants knew their content, which was revealed by the pharmacist only after the study was completed" and " three valuntoers from CLY (CHY) group reported pigmentation and temporary taste	

All outcomes volunteers from CLX (CHX) group reported pigmentation and temporary taste disturbance" The control mouthrinse had the same formulation as the test rinses but without CHX or Og Comment: the higher incidence of pigmentation and temporary taste disturbance in the CHX group meant that participants could have worked out which group they were in (or that they were in 1 of the active mouthrinse groups) and this could have affected their oral health behaviours High risk Quotes: "double-blind" and "The bottles were previously coded to warrant Blinding of outcome assessment (detection bias) that neither the examiner nor the participants knew their content, which was All outcomes revealed by the pharmacist only after the study was completed" and ".. three volunteers from CLX (CHX) group (30%) reported pigmentation and temporary taste disturbance" Comment: the higher incidence of pigmentation in the CHX group meant that the outcome assessor could have worked out that these 3 participants were in the CHX group and could not be adequately blinded Incomplete outcome data Low risk Quote: "All participants completed the trial" (attrition bias)

All outcomes



Pereira 2011 (Continued)				
Selective reporting (reporting bias)	High risk	All outcomes stated in the methods section are adequately reported in the results section. Tooth staining is considered to be an important outcome but is not adequately reported		
Other bias	Unclear risk	Quote: "At the beginning, there was no statistically significant difference between the control and tests groups with respect to PLI and BI (P>0.05) means"		
		No information is provided on the balance of demographic factors (age, gender etc) between groups		
		Quote: "A single, previously calibrated examiner scored the BI and the plaque Index (PLI)." No information on intra- and inteexaminer reliability (from the previous calibration) are reported		

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Methods	Trial design: parallel (3 arms - 2 arms are relevant to this review)
	Location: assumed university dental school, Iran
	Number of centres: 1
	Study duration: quote: "This study was conducted in 2003 for 10 months." Duration of rinsing and end point for outcome assessment was 6 weeks
Participants	Participants: those attending for treatment in the periodontal clinic of the University of Mashhad, Dental school, Iran
	Inclusion criteria: non-pregnant, no risk factors for increased susceptibility to bacterial endocarditis, no current medications or antibiotics within past 2 months, moderate adult periodontitis, at least 1 site with 4 mm or more pocket depth among their 6 lower anterior teeth and without a history of professional periodontal treatment within the past 6 months
	Exclusion criteria: not reported
	Baseline gingivitis: (GI Löe & Silness 1963) Gp A: mean 1.97 (SD 0.19); Gp B: mean 1.78 (SD 0.39); Group C mean 1.85 (SD 0.14)
	Age at baseline (years): mean age and age range are not reported
	Gender: not reported
	Number randomised: 30 (Gp A: 10; Gp B: 10; Gp C: 10)
	Number evaluated: not reported
Interventions	Comparison: (ultrasonic SRP with 0.2% CHX delivered through the tip of ultrasonically activated scaler)* versus (CHX 0.2% + ultrasonic SRP with distilled water) versus (ultrasonic SRP using distilled water alone)
	*Gp A (n = 10): ultrasonic SRP with 0.2% CHX delivered through the tip of ultrasonically activated scale after baseline examination and 1 week later (data from this group were not extracted). We excluded this arm from our risk of bias and analysis

Gp B (n = 10): CHX (0.2%) + ultrasonic SRP with distilled water after baseline examination and 1 week

Gp C (n = 10): ultrasonic SRP using distilled water alone after baseline examination and 1 week later Prophylaxis at baseline: quote: "Following the baseline measurements, the entire dentition was ultrasonically scaled and irrigated according to the treatment protocol for each group. After one week the

later: CHX twice daily, 6 weeks



Rahman	i 2006	(Continued)
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same procedure was repeated for all groups until all supra- and subgingival calculus was removed according to both visual and tactile inspection"

OHI: quote: "At the first visit all patients were instructed tooth brushing using the Bass technique and interdental cleaning aids were given and given similar toothbrushes and toothpastes." OHI was reinforced 1 week after the baseline measurements and throughout the study

Non-supervised rinsing

Timing of mouthrinsing in relation to toothbrushing: not reported

Postrinsing instructions: not reported

Outcomes

Gingivitis (gingival inflammation: GI Löe & Silness 1963, gingival bleeding: Gingival Bleeding Index, Ainamo & Bay 1975); plaque (PI Silness & Löe 1964) measured at 6 weeks

Probing pocket depth (6 sites on each tooth using a William's periodontal probe)

CAL (6 sites on each tooth using a William's periodontal probe) also measured at 6 weeks - not relevant to this review

Funding

Quote: "Research Vice-Chancellor of Mashhad University of Medical Sciences who supported the study by a grant"

Notes

Sample size calculation: not mentioned

Adverse effects: not reported

Declarations/conflicts of interest: not stated

Bias	Authors' judgement	Support for judgement
Random sequence genera-	Unclear risk	Quote: "randomly divided into three treatment groups"
tion (selection bias)		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: participants were not blinded. The different interventions meant that participants would know which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is not clear
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Blinding of outcome assessment not mentioned but, even if there was outcome assessor blinding, it is likely that there would be more staining in the CHX group thus compromising any blinding. However, tooth staining is not mentioned in the report so it remains unclear
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Losses to follow-up are not reported. Number of participants analysed at 6 weeks is not reported
Selective reporting (reporting bias)	High risk	Adverse effects are not mentioned. Tooth staining is considered to be an important side effect of CHX mouthrinsing but this is not reported
Other bias	Unclear risk	Number of examiners and training of examiners not mentioned



Rahmani 2006 (Continued)

Quote: "There were no significant differences in the clinical parameters between test and control groups at the baseline examination"

Methods	Trial design: parallel, 2 arms
	Location: private practice limited to periodontics, Los Angeles, USA
	Number of centres: 1
	Study duration: recruitment period not stated. Study duration and duration of rinsing was 6 weeks
Participants	Participants: patients with periodontitis who required osseous periodontal surgery
	Inclusion criteria: diagnosis of AAP Class III periodontitis; requiring periodontal osseous surgery; able to perform adequate mechanical plaque control
	Exclusion criteria: patients receiving osseous grafts, pregnancy; diabetes or other metabolic diseases; receiving systemic antibiotics within 3 months of the start of the study; receiving anti-inflammatory or other systemic drugs capable of altering clinical response
	Baseline gingivitis: not reported
	Age at baseline (years): not reported
	Gender: not stated
	Number randomised: 40 (Gp A: 19; Gp B: 21)
	Number evaluated: 38 (Gp A: 17; Gp B: 21)
Interventions	Comparison: CHX gluconate rinse versus placebo rinse used postperiodontal surgery
	Gp A (n = 19): CHX (0.12%), 15 mL, 30 seconds, twice daily
	Gp B (n = 21): same routine using placebo
	Prophylaxis at baseline: full-mouth SRP followed by osseous periodontal surgery in 1 quadrant (periodontal dressing then placed over entire surgical area). All participants received prescription for Tylenol II as pain relief
	OHI: prior to surgery patients received instructions in oral hygiene
	Non-supervised rinsing
	Timing of mouthrinsing in relation to toothbrushing: after usual oral hygiene practices
	Postrinsing instructions: not reported
Outcomes	Gingivitis (gingival inflammation: modified Löe & Silness GI 1963 using a pressure sensitive probe standardised at 30 g inserted to the bottom of the sulcus - reported as GI severity (overall mean scores) and GI bleeding (scores 2 and 3), measured at 4 and 6 weeks (not measured at 1 and 2 weeks due to early healing of wound masking the inflammatory status of the gingiva))
	Plaque (Silness & Löe PI 1964 - reported as plaque accumulation and visible plaque), measured at 1, 2, 4 and 6 weeks
	Extrinsic stain (stain/no stain scored from oral photographs of mandibular facial anterior teeth), measured at 6 weeks



Sanz 1989	(Continued)
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Participants' subjective assessment (questionnaire on whether or not they liked the assigned rinse including taste and compliance), unclear when measured (reported at 1 and 2 weeks)

Periodontal status (pocket probing depth and CAL), measured at 4 and 6 weeks - not relevant to this review

Postsurgical soft tissue health (epithelialisation - poor/moderate/good), measured at 1, 2, 4 and 6 weeks - not relevant to this review

Pain (participants' own perception on 0 to 10 scale and by medication consumption pain on soft palpation with periodontal probe - none/mild/severe), measured at 1, 2, 4 and 6 weeks - not relevant to this review

Funding

Quote: "This study was supported in part by a grant from the Procter & Gamble Company, Cincinnati, OH"

Notes

Sample size calculation: not reported

Adverse effects: no case of desquamation was seen with the use of 0.12% CHX rinse in this study. 24% of participants (n = 5) in the CHX group disliked the taste of the mouthrinse mainly because of a 'burning sensation' and 'a too strong taste'. Quote: "In the CHX group, 47.1% of the patients who did not have stain at the time of surgery developed light extrinsic dental stain" compared to 4.7% of patients in the placebo group

Declarations/conflicts of interest: not stated

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "they were randomly assigned to receive either chlorhexidine or place- bo mouthrinse via a computer-generated random list"
		Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	The authors describe the study as double-blind. The control mouthrinse was identical to the test mouthrinse but without CHx gluconate. Quote: "at 6 weeks, 47.1% (8/17) of the patients in the CHX group and 4.7% (1/21) in the placebo group showed light stain" (on the lower anterior teeth)
		Comment: the higher level of tooth staining in the CHX group meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is not clear
Blinding of outcome assessment (detection bias) All outcomes	High risk	The authors describe the study as double-blind. Clinical examinations were carried out. Quotes: "by an independent clinical examiner (not the periodontist) at pre-surgery, 1, 2, 4, and 6 weeks post-surgically" and " at 6 weeks, 47.1% (8/17) of the patients in the CHX group and 4.7% (1/21) in the placebo group showed light stain" (on the lower anterior teeth)
		Comment: the higher level of staining in the CHX group meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is not clear
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition at 6 weeks overall 2/40 (5%). Attrition by group: CHX 2/19 (11%), all participants in the control group contributed data for the main outcomes of gingivitis, plaque and staining.



Sanz 1989 (Continued)		
		2 subjects were lost from the CHX group due to concomitant systemic antibiotic use. In the placebo group, 1 subject did not have a PD and PAL measurements taken and was excluded from analysis of these measurements
		Comment: attrition is low (although all losses were from the CHX group) and do not appear to be related to the intervention
Selective reporting (reporting bias)	High risk	No measures of variance were reported. We estimated the SD for gingivitis (measured using the GI) as the median of the SDs in the placebo/control groups from similar studies that also used the GI at 4-6 weeks so that we could include the gingivitis data in the meta-analyses
Other bias	High risk	Training and calibration of the 1 (independent) clinical examiner is not mentioned. Baseline demographics of the subjects are not described. With the exception of visible plaque and stain, the baseline balance of the outcomes is not reported
		Staining was poorly reported but appears that 94.2% of CHX group had staining at baseline compared to 33.3% in placebo

Sanz 1994	
Methods	Trial design: parallel (3 arms - 2 arms are relevant to this review)
	Location: unclear
	Number of centres: 1
	Study duration: recruitment period not stated but the duration of the intervention was 6 months
Participants	Participants: adults with a history of established gingivitis
	Inclusion criteria: aged 18 to 65 years; minimum 16 natural teeth (including 4 anterior teeth in each jaw); GI Löe & Silness 1967 score greater than 0.7
	Exclusion criteria: evidence of periodontitis (probing pockets greater than 4 mm); medical history that would interfere with participation; evidence of gross oral neglect
	Baseline gingivitis: (GI Löe & Silness 1967) Gp A: mean 1.57; Gp B: mean 1.53
	Age at baseline (years): Gp A: mean 33.5 (range 18 to 60); Gp B: mean 33.6 (range 18 to 65)
	Gender: Gp A: male 29 (41%), female 42 (59%); Gp B: male 29 (43%), female 39 (57%)
	Number randomised: 139 (Gp A: 71; Gp B: 68) (208 including ineligible arm)
	Number evaluated: 130 (Gp A: 66; Gp B: 64) (191 including ineligible arm)
Interventions	Comparison: (CHX digluconate rinse + gum care dentifrice) versus (placebo rinse + gum care dentifrice) versus (placebo rinse + CHX digluconate and zinc lactate dentifrice)*
	*We excluded this arm from our risk of bias and analysis
	Gp A (n = 71): CHX (0.12%) rinse + gum care dentifrice containing allantoin, urea, azulene, medical soat calcium carbonate, and sodium monofluorophosphate 1100 ppm (Blend-a-med Parodontoseschutz), quantities and time not specified, twice daily
	Gp B (n = 68): as above but the rinse was a placebo
	Prophylaxis at baseline: quote: "After this examination all subjects received a dental prophylaxis in order to remove completely existing plaque and calculus"



Sanz 1994 (Continued)	O.U		
		s were available to all participants but no OHI was provided	
	Non-supervised rinsing		
	Timing of rinsing in rela	ation to toothbrushing: immediately after toothbrushing	
	Postrinsing instruction	s: none	
	Duration of treatment:	6 months	
Outcomes	Gingivitis: gingival inflammation (using a modification of GI Löe & Silness 1967. An electronic periodontal pressure sensitive probe standardised at 25 g was used and mean scores were reported); gingival bleeding (GI = 2 was recorded when bleeding appeared on probing, measured at 3 and 6 months)		
	Plaque (PI Silness & Lö	e 1964), measured at 3 and 6 months	
	Calculus (Volpe-Manho	old Calculus Index (Volpe 1965)), measured at 3 and 6 months	
	dard; overall stain mea	urfaces of the 12 anterior teeth. Comparisons made against a photographic stansured on a 0 to 6 scale; each tooth measured on a 0 to 4 scale for stain intensity; on a 0 to 6 scale for stain coverage), measured at 3 and 6 months	
	Oral soft tissue health a	attributable to product use, measured at 3 and 6 months	
Funding	Rinse and dentifrice su	pplied by Procter & Gamble	
Notes	Sample size calculation: not reported		
	Adverse effects: quote: "No adverse effects on oral soft tissues were seen and no deviations from clinically normal were recorded which would indicate possible side effects which are related to any of the products used in this study." There was a statistically significant increase in calculus for the positive control group compared with the control group. Quote: "Statistically significant more overall staining, more intense staining and stain coverage per tooth were detected for the positive control group and the experimental group compared with the control group"		
	Declarations/conflicts of interest: nothing explicitly stated but 2 of the 5 authors were employees of Procter & Gamble (manufacturers of the products being tested)		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Quote: "Subjects were then stratified by plaque Index (PII) and gingival index (GI) and gender and they were randomly assigned to one of the three treatment groups"	
		Comment: no description of the method of random sequence generation	
Allocation concealment (selection bias)	Unclear risk	No information is provided	
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	The study is described as double-blind. The control mouthrinse was Peridex without the CHX gluconate. Quotes: "All products were supplied in identical tubes and bottles, respectively" and "Statistically significant more overall staining, more intense staining and stain coverage per tooth were detected for the positive control group and the experimental group compared with the control group"	

Comment: the higher level of staining in the CHX group than the placebo group meant that participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the out-

come. The direction of this potential bias is unclear



Sanz 1994	(Continued)
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Blinding of outcome assessment (detection bias) All outcomes	High risk	The study is described as double-blind. Quotes: "The oral examinations were carried out by 2 examiners each carrying out the same examinations throughout the study" and "Statistically significant more overall staining, more intense staining and stain coverage per tooth were detected for the positive control group and the experimental group compared with the control group"
		Comment: the authors do not report precisely who was blinded, but even if this includes the outcomes assessors the higher level of staining in the CHX group meant that the outcome assessor could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition at 6 months overall 17/208 (8.2%). Attrition by group: CHX + gum care dentifrice 5/71 (7%); placebo + gum care dentifrice 4/68 (6%); placebo + experimental dentifrice 8/69 (11.6%)
		Reason for losses: quote: "Reasons not associated with product use"
		Comments: losses are low and although exact reasons for losses are not described and not broken down by group they are thought unlikely to have affected the outcome
Selective reporting (reporting bias)	High risk	The main outcome data presented in tables were broken down into subgroups with low and high initial gingivitis. The overall outcome data were presented only in graphs. No measures of variance were reported. We estimated the mean scores for gingivitis from the graphs. We estimated the SD for gingivitis (measured using the GI) as the median of the SDs in the placebo/control groups from similar studies that also used the GI at 6 months so that we could include the gingivitis data in the meta-analyses
Other bias	Unclear risk	Training and calibration of the 2 examiners is not mentioned. Quote: "All treatment groups were well balanced at the beginning of the study with regard to age, gender, initial plaque and initial gingivitis index scores as summarized in table 1"

Segreto 1986

Methods	Trial design: parallel, 3 arms
	Location: University of Texas Health Science Centre, San Antonio, Texas, USA
	Number of centres: 1
	Study duration: recruitment period not stated. Study duration and duration of rinsing was 3 months
Participants	Participants: adult volunteers from the University of Texas Health Science Centre at San Antonio Texas and towns people from San Antonio
	Inclusion criteria: minimum 16 natural teeth (including 4 molars)
	Exclusion criteria: existing soft tissue damage caused by ill-fitting appliances; advanced periodontal disease; dental neglect needing prompt professional attention; unfavourable medical history; regular use of antibiotics; pregnancy
	Baseline gingivitis (from Table 8): (GI Löe 1967) Gp A: mean 0.6745 (SD not reported); Gp B: mean 0.6468 (SD not reported); Gp C: mean 0.6750 (SD not reported)
	Age at baseline (years) (from Table 7): Gp A: 31.29 (range 18 to 61); Gp B: mean 32.33 (range 18 to 60); Gp C: mean 30.74 (range 18 to 60)



Segreto	1986	(Continued)
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Gender: overall: male 234 (39%), female 363 (61%). Gp A: male 80 (40%), female 122 (60%); Gp B: male 81 (41%), female 118 (59%); Gp C: male 73 (37%), female 123 (63%)

Number randomised: 597 (Gp A: 202; Gp B: 199; Gp C: 196)

Number evaluated: 454 (Gp A: 144; Gp B: 157; Gp C: 153)

Interventions

Comparison: 0.2% CHX gluconate rinse versus 0.12% CHX gluconate rinse versus placebo rinse

Gp A (n = 202): CHX (0.2%), 15 mL, 30 seconds, twice daily

Gp B (n = 199): CHX (0.12%), 15 mL, 30 seconds, twice daily

Gp C (n = 196): same routine using placebo

Prophylaxis at baseline: after the baseline examination all subjects received a thorough prophylaxis

OHI: participants followed usual habits but were given fluoride toothpaste and soft toothbrush

Non-supervised rinsing

Timing of mouthrinsing in relation to toothbrushing: not reported

Postrinsing instructions: none reported

Outcomes

Gingivitis: gingival inflammation measured using GI (Löe 1967) and Papillary Marginal Gingivitis Index (PMGI) (de la Rosa & Sturzenberger 197, visual inspection only - no probing); severity (mean extent of disease) and occurrence (meaning mean % gingival sites with GI or PMGI score ≥ 1). Gingival bleeding measured using GI (Löe 1967) mean % of gingival sites with a GI score ≥ 2)

Plaque (plaque disclosed using standard red dye and quantity on facial and lingual surfaces assessed using TQH PI (Turesky 1970))

Oral soft tissue effects (visual-tactile examination of the oral mucosa), measured at 6 weeks and 3 months

All measured at 6 weeks and 3 months

Funding

Quote: "The support of this study by the Procter & Gamble Company, Cincinnati, Ohio, is gratefully acknowledged"

Notes

Sample size calculation: not reported

Adverse effects: quote: "With the exception of some subjective comments, there were no differences in the systematic examination of oral soft tissues which would indicate that adverse effects were produced by either chlorhexidine mouthrinse in comparison to the placebo mouthrinse." 12% of subjects in the 0.2% group dropped out because they did not like the mouthrinse, compared to 4% of the 0.12% group and 1% of the placebo subjects

The results are reported separately for examiner A and examiner B. Data for examiner A only were extracted and used in the analysis

Declarations/conflicts of interest: not stated

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "they were stratified by age, sex, and gingivitis severity. Within strata they were assigned to one of three treatment groups by random permutations of 3"
		Comment: no description of the method of random sequence generation



Segreto 1986 (Continued)		
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	The study is described as double-blind. Quotes: "Neither the subject nor the dental examiners knew to which treatment the subject was assigned at any time during the study" and " the mouthrinses were supplied in pint bottles with child-resistant closures." The placebo mouthrinse was identical in composition to the 2 active mouthrinses but without the CHX
		Comment: tooth staining is likely to have been noticeable in the CHX groups after 3 months, therefore participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. However, tooth staining is not reported and there is not enough information upon which to base a judgement
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	The study is described as double-blind. Quote: "Neither the subject nor the dental examiners knew to which treatment the subject was assigned at any time during the study"
		Comment: tooth staining is likely to have been noticeable in the CHX groups after 3 months, and in this situation the outcome assessor could have worked out which individuals used CHX and not be adequately blinded. However, tooth staining is not reported and there is not enough information upon which to base a judgement
Incomplete outcome data (attrition bias)	High risk	Attrition at 3 months overall 143/597 (24%) and by group: CHX (0.2%) 58/202 (29%), CHX (0.12%) 42/199 (21%), control 43/196 (22%)
All outcomes		Reasons for losses: most were people who had moved or could not meet examination schedules, 12% of subjects in the 0.2% group dropped out because they did not like the mouthrinse, compared to 4% of the 0.12% group and 1% of the placebo subjects
Selective reporting (reporting bias)	High risk	No measures of variance are reported so the data could not be included in meta-analyses. Furthermore, tooth staining is considered to be an important side effect of CHX mouthrinsing but this is not reported
Other bias	Unclear risk	Training and calibration of the 2 examiners is not mentioned. We used data for examiner A only. Intraexaminer reliability is not reported for this examiner
		Baseline demographic and clinical data for the 3 groups appear balanced

Southern 2006

Methods	Trial design: parallel (3 arms - 2 arms are relevant to this review)
	Location: Virginia, USA (type of setting not reported assumed to be a university dental school)
	Number of centres: not reported
	Study duration: recruitment period is not reported. Study duration and duration of rinsing was 3 months
Participants	Participants: healthy adults with gingivitis
	Inclusion criteria: minimum age of 18 years; Löe & Silness 1967 GI score of 2 or 3; general good health
	Exclusion criteria: advanced periodontal disease (AAP IV or greater); use of antibiotics in the 90 days before the start of the study; need for antibiotics premedication; anterior facial restorations; use of a



Southern 2006 (Continued)

daily antigingivitis rinse in the 3 months before the start of the study; pregnancy; smoker; poor general

Baseline gingivitis: not reported. Quote: "Randomization produced similar equivalent baseline groups that exhibited overall GI scores of 2 or 3"

Age at baseline (years): mean 25 (SD 7.67) (not reported by group but "Randomization produced similar equivalent baseline groups")

Gender: male n = 35 (57%), female n = 28 (43%) (not reported by group but "Randomization produced similar equivalent baseline groups")

Number randomised: 63 (not reported by group)

Number evaluated: 60 (not reported by group)*

*Text states: "final sample size of 60 (30 in each group)" - which does not make sense as there were 3 groups. Assumed final sample size 20 in each group (21 at baseline)

Interventions

Comparison: CHX rinse versus herbal rinse* versus placebo rinse

Gp A (n = not reported, assumed 21): CHX (0.12%), half an ounce (15 mL), length of time not specified, twice daily (morning and evening)

*Gp B (n = not reported, assumed 21): same as above with herbal rinse (filtered spring water, vegetable glycerin, echinacea, goldenseal, calendula, aloe, bloodroot, grapefruit seed extract, citric acid, spearmint oil, peppermint oil and cinnamon). This arm is not included in our risk of bias or analysis

Gp C (n = not reported, assumed 21): same as above with placebo rinse

Prophylaxis at baseline: none. Quote: "no prophylaxis was conducted prior to study initiation"

OHI: all groups given same soft toothbrush and whitening fluoride toothpaste, and instructed to follow their usual oral hygiene routine (but not to use any other oral rinse during the study)

Partially supervised rinsing: as described in methods it appears to be non-supervised rinsing but the authors state that "A research assistant was responsible for product allocation and supervising rinsing procedures"

Timing of mouthrinsing in relation to toothbrushing: after usual mechanical oral hygiene routine

Postrinsing instructions: not reported

Outcomes

Gingivitis (gingival inflammation: GI Löe 1967 and BOP), measured at 1, 2 and 3 months

Plaque (PI Silness & Löe 1964), measured at 1, 2 and 3 months

Oral soft tissue health, measured at baseline, 1, 2 and 3 months

Funding

Not reported

Notes

Sample size calculation: not reported

Adverse effects: not reported

Non-smokers

Declarations/conflicts of interest: not stated

Risk of bias

Bias

Authors' judgement Support for judgement



Southern 2006 (Continued)	Unalograiak	Queto, "participants were randomized to an affiliate and distance."
Random sequence generation (selection bias)	Unclear risk	Quote: "participants were randomized to one of three product groups"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	Quotes: "participants were randomized to one of three product groups" and "Individuals were identified by code numbers throughout the study" and "A research assistant was responsible for product allocation and supervising rinsing procedures"
		Comment: not enough information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Quotes: "product identifiers were removed from all containers" and "The examiner and participants were blinded to product allocation" and "A research assistant was responsible for product allocation and supervising rinsing procedures." It is not clear if the research assistant was blinded to allocation of mouthrinses
		Comment: the participants rinsed for 3 months. No information is provided about the ingredients in the placebo rinse. Tooth staining, although likely in the CHX group, is not mentioned in the report. The research assistant who was responsible for product allocation appears to have also supervised the mouthrinsing which may have compromised personnel blinding (although the methods imply that the rinsing was unsupervised so whether the rinsing was supervised/unsupervised is not clear)
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quotes: "The examiner and participants were blinded to product allocation." Individuals were identified by code numbers throughout the study. Participants were given fluoride whitening toothpaste "to decrease the possible side effects of staining and lessen examiner bias"
		Comment: tooth staining is likely to have been noticeable in the CHX groups after 3 months, and in this situation the outcome assessor could have worked out which individuals used CHX and not be adequately blinded. However, tooth staining is not reported and there is not enough information upon which to base a judgement
Incomplete outcome data (attrition bias)	Low risk	Attrition at 3 months overall 2/42 (4.8%), 1 participant from each group. Reasons for loss to follow-up: personal reasons
All outcomes		Comment: attrition is low and there are no differential group losses
Selective reporting (reporting bias)	High risk	The GI was recorded but the mean GI was not reported. Instead the change in proportion of GI scores of 2 or 3 compared to placebo was reported. No data are reported for probing pocket depth. Quote: "statistical analysis showed no statistically significant change in PPD in any of the groups." Tooth staining is considered to be an important side effect of CHX mouthrinsing but this is not reported
Other bias	Unclear risk	Quote: "A single calibrated dental hygienist was used to collect all data"
		Comment: no information provided on intraexaminer reproducibility
		Quotes: "Randomization produced similar equivalent baseline groups" and "Randomization produced similar equivalent baseline groups that exhibited overall GI scores of 2 or 3"
		Insufficient information is provided upon which to judge the baseline balance of demographic factors



Stookey 2005

Methods Trial design: parallel (4 arms - 2 arms are relevant to this review)

Location: USA, setting "clinical site"

Number of centres: 1

Study duration: the recruitment period is not reported but the duration of rinsing was 6 months

Participants

Participants: healthy adults

Inclusion criteria: at least 18 years of age with minimum 16 gradable natural teeth with 4 molars and in good general health. Subjects also had to have a minimum baseline whole mouth average GI score 0.50, at least 10 bleeding sites and a maximum baseline plaque score of 3

Exclusion criteria: quote: "Patients were unable to continue in the study for the following reasons: participation in any other dental study; use of other oral care products; evidence of rampant caries, obvious periodontal disease, chronic neglect requiring urgent treatment, history of any medical diseases that may interfere with the study (e.g. bleeding tendencies, infectious diseases) wearing removable or fixed orthodontic devices; use of antibiotics or immunosuppressives within 1 week prior to baseline, 3- or 6-month exams; use of antiinflammatory drugs or analgesics within 48 hours of baseline, 3- or 6-month exam; known hypersensitivity to chlorhexidine or tartrazine; oral prophylaxis outside of study; use of oral chlorhexidine products or participation in an oral rinse study within 3 months prior to baseline examination; pregnancy/nursing; noncompliance by missing more than five consecutive supervised rinses and/or more than 15% of all supervised rinses

Baseline gingivitis: (Löe & Silness GI) Gp A: mean 0.794 (SE 0.025); Gp B: mean 0.792 (SE 0.018); Gp C mean 0.800 (SE 0.018); Gp D mean 0.814 (SE 0.020)

Age at baseline (years): Gp A: mean 32.9; Gp B: mean 33.7; Gp C: mean 33.8; Gp D mean 34.3. Gp A: range 18-53; Gp B: range 18-59; Gp C: range 18-57; Gp D range 18-66

Gender: number in each group not reported. Male: Gp A: 60%; Gp B: 63%; Gp C: 59%; Gp D: 62%. Female: Gp A: 40%; Gp B: 37%; Gp C: 41%; Gp D: 38%

Number randomised: 366 (Gp A: 49; Gp B: 97; Gp C: 103; Gp D: 102)

Number evaluated (6 months): 298 (Gp A: 40; Gp B: 82; Gp C: 90, Gp D: 86)

Interventions

Comparison: CHX rinse versus cetylpyridinium chloride rinse (CPC 0.075%)^a versus cetylpyridinium chloride rinse (CPC 0.1%)^b versus placebo

Gp A (n = 49): CHX (Peridex) 0.12%: 15 mL, twice daily, 30 seconds, 6 months

 a Gp B (n = 97): CPC 0.075%: 15 mL, twice daily, 30 seconds, 6 months. This arm was excluded from our risk of bias and analysis

 $^{
m b}$ Gp C (n = 103): CPC 0.10%: 15 mL, twice daily, 30 seconds, 6 months. This arm was excluded from our risk of bias and analysis

Gp D (n = 102): placebo (of CPC): 15 mL, twice daily, 30 seconds, 6 months

Prophylaxis at baseline: quote: "approximately 1 week following the baseline examination, qualifying subjects received a thorough dental prophylaxis"

OHI: supervised toothbrushing but unclear if there was OHI

Quote: "From Monday to Friday, subjects came to the clinical site each morning for supervised brushing and rinsing.. Subjects were instructed to follow the same instructions in the evening and on the weekends"



Stookey 2005 (Continued)

,,	a 0.243% sodium fluor	ng in relation to toothbrushing: quote: "Subjects were instructed to brush with ide toothpaste (Crest Cavity Protection) using a disposable Anchor toothbrush, rinse with 15 ml of product for 30 seconds"	
Outcomes	Gingivitis (GI Löe & Silness 1963 to measure gingival inflammation and bleeding (GI scores 2 or 3)); plaque (TQH PI, Turesky 1970); oral soft tissue examinations; assessed at 3 and 6 months follow-up		
Funding	The test formulations (CPC) were provided by the Procter & Gamble Company		
Notes	Adverse effects: quote: "There were no serious adverse events reported during the study that were deemed related to the test products. OST examinations showed that subjects rinsing with the chlorhexidine treatment had significantly more "tongue lesion" comments at month 3 than those rinsing with either the CPC rinse or the placebo rinse. There were no significant differences between either of the CPC rinse groups and the placebo group in the number of subjects that had OST comments at 3 or 6 months"		
	Declarations/conflicts of interest: nothing explicit stated, however 2 of the co-authors (Dr Witt and Dr Gibb) were employees of the Procter & Gamble Company at the time the study was published		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Quotes: "Subjects were randomly assigned to four treatments, balancing for gender and baseline mean GI score" and "The randomization was performed such that the sample size per group ratio was 2:2:2:1 with 1 representing the chlorhexidine rinse group"	
		Comment: no description of the method of random sequence generation	
Allocation concealment (selection bias)	Unclear risk	No information is provided	
Blinding of participants and personnel (perfor-	Unclear risk	Quotes: "double-blind" and "all mouthrinses were packed in identical amber bottles." The control rinse was a placebo of CPC rinse.	
mance bias) All outcomes		Quote: "From Monday to Friday, subjects came to the clinical site each morning for supervised brushing and rinsing Subjects were instructed to follow the same instructions in the evening and on the weekends"	
		Comment: it is not clear if the personnel supervised the rinsing were blinded to product allocation. Tooth staining is likely to have been noticeable in the CHX group after 6 months, therefore participants could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. However, tooth staining is not reported and there is not enough information upon which to base a judgement	
Blinding of outcome as-	Unclear risk	Quote: "double-blind"	
sessment (detection bias) All outcomes		Comment: tooth staining is likely to have been noticeable in the CHX group after 6 months, and in this situation the outcome assessor could have worked out which individuals used CHX and not be adequately blinded. However, tooth staining is not reported and there is not enough information upon which to base a judgement	
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition at 6 months all arms: $68/366$ (18.6%). Attrition by group: CHX $9/49$ (18%), control $16/102$ (15.7%). Reasons for attrition: (not broken down by group) protocol violation (NSAIDS) ($n=15$); medication use outside study protocol ($n=41$ at 3 months, 17 at 6 months); not available ($n=10$ at 3 months, 17	



Stookey 2005 (Continued)		at 6 months); non-compliance (n = 5 at 3 months and 5 at 6 months); adverse event (n = 5 at 3 months and 8 at 6 months); non-study related medical reasons (n = 2 at 3 months and 6 at 6 months) Comment: attrition was high and some losses were possibly related to the intervention
Selective reporting (reporting bias)	High risk	We calculated SDs from the SEs reported so that the data could be included in meta-analyses. Tooth staining is considered to be an important side effect of CHX mouthrinsing but this is not reported
Other bias	Unclear risk	Number of examiners not mentioned. Training and calibration of examiners not mentioned
		Subjects balanced at baseline with regard to gender, age, gingival health and plaque

Taller 1993	
Methods	Trial design: parallel (3 arms - 2 arms are relevant to this review)
	Location: New Jersey, USA (type of setting not reported)
	Number of centres: not reported
	Study duration: recruitment period is not reported. Study duration and duration of rinsing was 5 weeks
Participants	Participants: adults with gingival bleeding
	Inclusion criteria: adults with gingival bleeding; non-contributory medical history; not taking antibiotics; minimum of 20 teeth; minimum total gingival bleeding score of 10 (bleeding points recorded on 4 areas of 6 teeth (0 = no bleeding; 1 = bleeding) so that maximum score would be 24)
	Exclusion criteria: not reported
	Baseline gingivitis: (gingival bleeding score on 0 to 24 scale as described above) Gp A: mean 14.46 (SD 3.13); Gp B: mean 13.57 (SD 3.84); Gp C: mean 13.77 (SD 1.64)
	Age at baseline (years): mean 46 (range 21 to 71) (not reported by group but "no statistical differences between the ages of the groups ($P = 0.23$ for one way and over)")
	Gender: (reported for completers only) males 23 (57.5%), females 17 (42.5%) (not reported by group but "sex was sufficiently randomized ($P = 0.92$ by chi square test)")
	Number randomised: 42 (not reported by group)
	Number evaluated: 40 (Gp A: 13; Gp B: 14; Gp C: 13)
Interventions	Comparison: CHX gluconate rinse versus baking soda/hydrogen peroxide toothpaste* versus control (usual care)
	Gp A (n = 13 analysed): CHX (0.12%), 15 mL, 30 seconds, twice daily
	*Gp B (n = 14 analysed): baking soda/hydrogen peroxide toothpaste used instead of usual toothpaste. We excluded this arm from our risk of bias and analysis
	Gp C (n = 13 analysed): usual toothpaste
	Prophylaxis at baseline: not reported, assume none



OHI: all groups given OHI, same toothbrush and dental floss, and instructed to brush and floss 3 times per day	
Non-supervised rinsing	
Timing of mouthrinsing in relation to toothbrushing: twice per day after brushing and flossing	
Postrinsing instructions: not reported	
Gingivitis (gingival bleeding on probing on a 0 to 24 scale - a score of 12 would mean that 50% of sites bled), measured at 5 weeks	
Signs of pathology of oral tissues	
Not clear. The authors acknowledged the support of the University of Medicine & Dentistry of New Jersey on the project	
Sample size calculation: not reported	
Adverse effects: quote: "There was no evidence of pathology of the oral tissues of any subject"	
Declarations/conflicts of interest: not stated	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "Subjects were randomly assigned to each group"
		Comment: no description of the method of random sequence generation
Allocation concealment (selection bias)	Unclear risk	No information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Blinding of participants is not stated and not possible with the interventions being tested and this could have affected participants' oral health behaviours and hence the outcome. The direction of this potential bias is unclear
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "the same examiner scored all subjects and did not know the group each subject was assigned to at any time"
		Comment: blinding attempted but, as there is no information on tooth staining, it is not clear whether or not it would be possible for the outcome assessor to remain blind
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition at 5 weeks 2/42 (4.8%). Losses not broken down by group. Reason for losses: did not return at 5 weeks
		Comment: it is not clear which groups the participants were lost from, however losses are low and are considered unlikely to have affected the outcome
Selective reporting (reporting bias)	High risk	Tooth staining is considered to be an important side effect of CHX mouthrinsing but this is not reported
Other bias	Unclear risk	1 examiner scored all subjects which eliminates interexaminer variability as a source of bias. However, there is no mention of training or intraexaminer reproducibility
		The groups are balanced at baseline for bleeding points, age and sex



Turkoglu 2009

Methods

Trial design: parallel, 2 arms

Location: Department of Periodontology, Ege University, Izmir, Turkey

Number of centres: 1

Study duration: quote: "subjects were recruited...over a period of 1 year between 2006 and 2007."

Study duration and duration of rinsing was 4 weeks

Participants

Participants: adults with gingivitis attending the Department of Periodontology, Ege University

Inclusion criteria: 18 to 45 years old; gingivitis associated with dental plaque; no sign of destructive periodontal disease; clinical attachment loss less than 3 mm; minimum 20 teeth (teeth that were grossly carious fully crowned or extensively restored, orthodontic banded, abutments, or third molars were not included in the tooth count)

Exclusion criteria: use of tobacco products; current or historic serious systemic disease affecting immune response (e.g. diabetes, immune disorders, hepatitis, HIV); use of antibiotics, anti-inflammatory, or immunosuppressive drugs within 3 months of the start of the study; periodontal therapy within 3 months of the start of the study; pregnancy or lactation; use of oral contraceptives

Baseline gingivitis: (Papillary Bleeding Index) Gp A: mean 1.47 (SD 0.5); Gp B: mean 1.44 (SD 0.7)

Age at baseline (years): Gp A: mean 27.56 (SD 8.3) (range 18 to 45); Gp B: mean 25.44 (SD 5.6) (range 19 to 35)

Gender: Gp A: males 13 (52%), females 12 (48%); Gp B: males 12 (48%), females 13 (52%)

Number randomised: 87 (Gp A: 45; Gp B: 42)

Number evaluated: 50 (Gp A: 25; Gp B: 25)

Interventions

Comparison: CHX digluconate rinse versus placebo rinse

Gp A (n = 45): CHX (concentration not reported), 10 mL, 1 min, twice daily (morning and before bedtime)

Gp B (n = 42): same procedure as above using placebo rinse

Prophylaxis at baseline: not reported. Assumed none

OHI: all participants given the same fluoride toothpaste and toothbrush, and given toothbrush instruction (Modified Bass technique), instructed to brush twice daily

Non-supervised rinsing

Timing of mouthrinsing in relation to toothbrushing: 30 min after toothbrushing

Postrinsing instructions: no rinsing with water or eating or drinking for 30 min after using their allocated rinse

Outcomes

Gingivitis (gingival bleeding: Papillary Bleeding Index, Saxer 1975)

Plaque (PI, Quigley & Hein 1962)

Calculus (scale not specified; references Oral Health of United States Adults. Epidemiology and oral disease prevention program. Bethesda, MD: National Institute of Dental Research; 1987; 159-67. NIH publication no 87-2868); probing depth* and gingival crevicular fluid cytokine levels*, all measured at 4 weeks using a Williams periodontal probe

Adverse effects, recorded weekly throughout the study



Turkoglu 2009 (Continued)	*Not relevant to this re	view		
Funding	Quote: "This work was supported by an unrestricted grant from Procter & Gamble. We would like to thank Drogsan Company for supplying the CHX and placebo mouthrinses"			
Notes	Sample size calculation: based on detecting a 0.5 difference in plaque index and papillary bleeding dex at 5% significance and 80% power, required 15 participants per group, which was achieved Adverse effects: of the 25 who rinsed with CHX; 5 subjects experienced taste disturbances and 14 showed discolouration of teeth and/or tongue and 3 subjects had mucosal ulcerations			
	Non-smokers			
	Declarations/conflicts	of interest: not stated		
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Unclear risk	Quote: "patients were randomly assigned to CHX or placebo groups by taking into account the gender, age and the extent of the gingivitis"		
		Comment: no description of the method of random sequence generation		
Allocation concealment (selection bias)	Low risk	Quote: "patients were randomly assigned to CHX or placebo groups by taking into account the gender, age and the extent of the gingivitis by an independent periodontist who kept the allocation information confident until the data collection and biochemical analysis were completed"		
		Comment: we consider it unlikely that participants and investigators could foresee assignment of the intervention		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Double-blind. Quotes: "The subjects were unaware of the type of mouthrinse provided to the patient" and "The placebo mouthrinse was composed of CHX mouthrinse ingredients except that it lacked the active ingredients (Chlorhexidine digluconate). Both CHX and placebo bottles were similar in appearance"		
		Comment: many of the participants in the CHX group experienced taste disturbances and/or discolouration of the teeth and/or tongue. Subjects in this group could have worked out which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is not clear		
Blinding of outcome assessment (detection bias) All outcomes	High risk	Double-blind. Quote: "The same examiner recorded all clinical measurements over the course of the study that was unaware of the type of mouthrinse provided to the patient." In addition an independent periodontist performed the randomisation of participants to groups and kept the allocation information confidential until the analyses were completed		
		Comment: more than half of the subjects in the CHX showed discolouration of the teeth and/or tongue. It is unlikely that the clinical examiner could remain unaware of the group allocation of these subjects and this could have affected the outcome assessment. The direction of this potential bias is not clear		
Incomplete outcome data (attrition bias) All outcomes	High risk	Losses to follow-up at 4 weeks 37/87 (42.5%). By group: CHX 20/45 (44.4%),placebo 17/42 (40.5%). Reasons for loss to follow-up: CHX: 3 discontinued the intervention and 17 were lost follow-up due to antibiotic usage, mucosal ulcerations, irregular mouthrinse usage. Placebo: 3 discontinued the intervention and 14 were lost follow-up due to antibiotic usage, mucosal ulcerations, irregular mouthrinse usage		



Turkoglu 2009 (Continued)		Comment: the numbers lost in each group for each reason are not fully broken down. Losses are very high especially considering the short study duration. With such a high rate of attrition, if the missing participants in 1 group had a higher mean (e.g. gingivitis score) than those in the other group, as the attrition rate increased, so would the MD between groups
Selective reporting (reporting bias)	Low risk	All outcomes stated in the methods section are adequately reported in the results section. Adverse effects were only reported for the CHX group (presumably there were no adverse effects in the control group)
Other bias	Low risk	Quotes: "All measurements were performed by a single-blinded and calibrated examiner the intra-examiner reliability was high as was revealed by intraclass correlation coefficient 0.87 for PD measurements and 0.85 for plaque measurements" and "There were no differences between CHX and placebo groups in the distribution of the extent of gingivitis, gender and age. The groups were balanced at baseline for PD, PBI, PI and CI values (P>0.05)"

Van Strydonck 2008

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Study design: parallel (3 arms 2 arms are relevant to this review)

Location: the Netherlands. Department of Periodontology, Academic Centre for Dentistry (ACTA)

Number of centres:1

Study duration: Recruitment period is not reported. Study duration and duration of rinsing was 6 weeks

Participants

Participants: non-dental students at the University of Amsterdam

Inclusion criteria: quotes: "Good general health, no medical or dental history or medication which might interfere with the outcome or the progress of the study" and "a minimum of 18 scorable natural teeth excluding third molars or crowned teeth with porcelain or gold restorations. To be enrolled in the study, the subjects were required to have a minimum of 40% bleeding sites as determined by the Bleeding on Marginal Probing Index (BOMP)"

Exclusion criteria: quote: "Subjects were excluded if they had any physical limitations or restrictions which might preclude normal toothbrushing skills. They were also excluded if they had used an oral CHX product or had taken a systemic antibiotic or antiinflammatory drug for 3 consecutive days within the previous 3 months. Subjects with removable prostheses or orthodontic appliances were not allowed to participate"

Baseline gingivitis: Bleeding on Marginal Probing Index (Van der Weijden 1994) Gp A: 1.21 (SD 0.24), Grp B: 1.22 (SD 0.25), Gp C: 1.26 (SD 0.26)

Age at baseline: mean Gp A: 21, Gp B: 22, Gp C: 23. Age range 18-65 years

Gender: Gp A: male 21% female 79%, Gp B: male 26% female 74%, Gp C: male 37% female 63%

Number randomised: 150 (Gp A 50, Gp B 50, Gp C 50) Number evaluated: 140 (Gp A 47, Gp B 47, Gp C 46)

Interventions

Comparison: (CHX mouthrinse and template control brush) versus (template control brush) versus (template test brush (with slow-releasing delivery of 124 mg CHX digluconate))*

Gp A (n = 50): template control brush and CHX rinse 0.2% CHX digluconate: 10 mL, twice daily, 60 seconds and brush twice daily for 1 min without dentifrice

Gp B (n = 50): template control brush: brush twice daily for 1 min without dentifrice



Van Strydonck 2008 (Continued)

*GP C (n = 50): test brush (with slow-releasing delivery of 124 mg CHX digluconate): brush twice daily for 1 min without dentifrice. We excluded this arm from our risk of bias and analysis

Prophylaxis at baseline: quote: "At baseline subjects received a supragingival prophylaxis to render them plaque and stain free"

OHI: quote "..instructed to brush twice daily without a dentifrice for 1 min (in the morning and in the evening)"

Non-supervised rinsing

Timing of mouthrinsing in relation to toothbrushing: rinsed twice daily after toothbrushing without a dentifrice

Postrinsing instructions: quote: "subjects were asked to refrain from rinsing eating or drinking for 30 mins after using their assigned product"

Outcomes

Gingivitis (gingival bleeding: Bleeding on Marginal Probing Index (Saxton & Van der Ouderaa, Van der Weijden 1994); plaque (Silness & Löe 1964, Danser 2003); stain (Gründermann Modified Stain Index (Gründermann 2000) all measured at 3 and 6 weeks

Funding

Quote: "This study was supported by a grant from Oral-B Laboratories, Belmont, CA, USA." GSK provided the Corsodyl mouthrinse

Notes

No dentifrice was used when toothbrushing

Sample size calculation: very detailed description provided. Quote: ".... a sample size of 45 subjects per treatment group was needed to ensure an 80% (power = $1-\beta$) or greater chance of detecting differences of ≥ 0.11 whole-mouth BOMP units"

Adverse effects: apart from tooth staining, quotes: "no adverse events were reported" and "No differences were detected in the proportion of oral tissue abnormalities among the groups, with the exception of the tongue. Changes noted were the presence of stain or discoloration on the tongue. Treament Ctb+R (Gp A) yielded a statistically significantly (P=0.0001) greater proportion of abnormal observations than treatments Ttb (Gp C, excluded from this review) and Ctb (Gp B)"

Declarations/conflicts if interest: quote: "The authors declare that they have no conflict of interests"

Risk of bias

Bias Authors' judgement Support for judgement		Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Randomisation was performed using a computer-generated list of random-numbers"
		Comment: adequate method of random sequence generation
Allocation concealment	Unclear risk	The study co-ordinator was responsible for allocation concealment
(selection bias)		Comment: not enough information is provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Blinding of participants is not possible due to the interventions being tested: template toothbrush versus template toothbrush and CHX mouthrinse and this could have affected participants' oral health behaviours and hence the outcome. The direction of this potential bias is unclear
Blinding of outcome assessment (detection bias) All outcomes	High risk	Quote: "Study products were coded and distributed to the subjects in a location away from the examiners to ensure and maintain blinding the examiners were blind to treatment randomization and records of earlier examinations were not available at the time of re-examinations One examiner assessed all plaque scores and performed all stain evaluations Another examiner assessed all the bleeding scores using BOMP and all safety evaluations"



Van Strydonck 2008 (Continue	d)	
		Quote: "The mean stain scores for treatment Ctb+R (CHX) were statistically greater (P=0.0001) than for treatments Ttb (CHX template toothbrush) and Ctb (control toothbrush)"
		Comment: despite the efforts made to ensure blinding of outcome assessment, the higher level of tooth staining in the CHX rinse group meant that examining clinicians could not be adequately blinded and this could have affected the outcome assessment. The direction of this potential bias is unclear
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition: all arms: 10/150 (6.7%). By group: CHX rinse 3/50 (6.0%), template control brush 3/50 (6%). Reasons for losses not broken down by group. Subjects lost after randomisation and before baseline examination: 4 were disqualified, 4 refused to participate and 2 were excluded for other reasons Comment: all losses were after randomisation before the start of the trial. They were balanced across groups
Selective reporting (reporting bias)	Low risk	All outcomes stated in the methods sections were reported in the results section
Other bias	Low risk	Quote: "Both examiners were well-trained and had been involved in previous studies." Calibration is not mentioned
		There was no statistically significant difference in mean age, or whole mouth plaque and gingival bleeding levels

Weitz 1992

Methods	Trial design: parallel, 2 arms				
	Locaton: geriatric facility, USA				
	Number of centres: 1				
	Study duration: recruitment period not stated. Study duration and duration of rinsing was 60 days				
Participants	Participants: residents in a geriatric facility				
	Inclusion criteria: residents in the geriatric facility receiving "emergency only" care. Quote: " at least 60 years old and in good health"				
	Exclusion criteria: quote: "Patients presenting with a history of antibiotic therapy within the last 3 months were not accepted into the study sample"				
	Baseline gingivitis: GI (Löe & Silness 1963) Gp A: 1.88, Gp B: 1.80. SD not reported				
	Age at baseline: at least 60 years old at recruitment, mean 83 years. Not reported by group				
	Gender: overall: male 12 (33.33%), females 24 (66.66%). Not reported by group				
	Number randomised: 36 (number in each group is not reported). 12 with removable dentures, 12 with overdentures and 12 with no prosthesis				
	Number evaluated: 36 (number in each group is not reported)				
Interventions	Comparison: CHX rinse versus placebo rinse				
	Gp A: CHX (0.12%) (n = not reported): 15 mL, twice daily, 30 seconds				
	Gp B: placebo (n = not reported): 15 mL, twice daily, 30 seconds				



Weitz 1992 (Continued)					
(Prophylaxis at baseline: none, prophylaxis performed after final measurements at 60 days				
	OHI: none. Quote: " instructed to continue their normal oral hygiene routine"				
	Non-supervised rinsing				
	Timing of mouthrinsing in relation to toothbrushing: not reported				
	Postrinsing instructions: not reported				
Outcomes	Gingivitis (gingival inflammation: GI (Löe & Silness 1963), measured at 60 days				
	Plaque (PI Silness & Löe 1964), measured at 60 days				
Funding	Not reported				
Notes	Sample size calculation: not reported				
	Adverse effects: not reported				
	Declarations/conflicts of interest: not reported				

Risk of bias

Bias	Authors' judgement	Support for judgement	
Random sequence genera-	Unclear risk	Quote: "patients were randomly divided into two groups"	
tion (selection bias)		Comment: no description of the method of random sequence generation	
Allocation concealment (selection bias)	Unclear risk	No information is provided	
Blinding of participants and personnel (performance bias)	Unclear risk	The study is not described as a single-blind study but the control group rinsed with a quote "matched placebo". Information on tooth staining is not reported	
mance bias) All outcomes		Comment: not enough information is provided	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Blinding of outcome assessor is not mentioned and is unlikely. Information tooth staining is not reported	
Incomplete outcome data (attrition bias) All outcomes	Low risk	There were no losses to follow-up	
Selective reporting (reporting bias)	High risk	No measure of variance was reported. We were unable to estimate the SD and therefore could not include these data in meta-analyses. Furthermore, tooth staining is considered to be an important side effect of CHX mouthrinsing but this is not reported. Adverse effects are not reported	
Other bias	Unclear risk	No information is provided on number of examiners and examiner training and calibration	
		Quote: "At baseline, the active and control groups were not statistically different (independent t-test, P<0.05) with respect to age, gingival index or plaque index"	



Zimmer 2006

Methods

Trial design: parallel (4 arms - 2 arms relevant to this review)

Location: Department of Operative and Preventive Dentistry and Endodontics, Heinrich Heine University of Düsseldorf, Germany

Number of centres: 1

Study duration: recruitment period is not stated. Study duration and duration of rinsing was 8 weeks

Participants

Participants: adult blood donors who visited the blood bank of the University Hospital, Düsseldorf, Germany in July and August 2004

Inclusion criteria: minimum Papillary Bleeding Index and Modified Proximal Plaque Index scores per tooth of 0.5 and 1.5 respectively

Exclusion criteria: wearing fixed orthodontic appliances; severe periodontitis (defined as CAL 5 mm or greater in a minimum of 3 teeth); long-term use of drugs with anti-inflammatory effects within 1 month prior and/or during the study; removable dentures; less than 20 natural teeth; regular use of dental floss or antimicrobial mouthrinses during past 3 months; furcation involvement; pathological tooth mobility

Baseline gingivitis: (Papillary Bleeding Index) Gp A: mean 1.23 (SD 0.44); Gp B: mean 1.25 (SD 0.45); Gp C: mean 1.19 (SD 0.44); Gp D: mean 1.27 (SD 0.45)

Age at baseline (years): mean 31.7 (range 20 to 64.4) (not reported by group but authors state that there were no statistically significant differences between groups)

Gender: 78 (50%) males, 78 (50%) females (not reported by group)

Number randomised: 156 (Gp A: 39; Gp B: 39; Gp C: 39; Gp D: 39)

Number evaluated: 156 (Gp A: 39; Gp B: 39; Gp C: 39; Gp D: 39)

Interventions

Comparison: (CHX + sodium fluoride (NaF) rinse and toothbrushing) versus (cetylpyridinium chloride (CPC) + sodium fluoride (NaF) rinse and toothbrushing)^a versus (flossing and toothbrushing)^b versus toothbrushing

Gp A (n = 39): CHX (0.06%) + 0.025% NaF, amount not specified, 30 seconds, once daily (after tooth-brushing at bedtime)

*Gp B (n = 39): CPC (0.1%) + 0.025% NaF, amount not specified, 30 seconds, once daily (after tooth-brushing at bedtime). We excluded this arm from our risk of bias and analysis

 * Gp C (n = 39): flossing once daily (after toothbrushing at bedtime). We excluded this arm from our risk of bias and analysis

Gp D (n = 39): usual toothbrushing routine

Prophylaxis at baseline: quote: "The screening examination was followed by calculus removal in the lower front teeth"

OHI: quote: "Participants received brief instructions for the procedure they had to perform in addition to using the toothbrush, dental floss and oral rinsing. The dental floss instruction was demonstrated with a plastic tooth model did not exceed 2 minutes. The subjects were advised to brush their teeth in the usual manner. No instructions concerning brushing technique and brushing time were given." All participants received same toothbrush and fluoride toothpaste

Non-supervised rinsing

Timing of mouthrinsing in relation to toothbrushing: after brushing with toothpaste

Postrinsing instructions: not reported



Zimmer 2006 (Continued)

Outcomes

Gingivitis (gingival bleeding: Papillary Bleeding Index, Saxer & Muhlemann 1975); plaque (Modified Proximal Plaque Index, Lange 1977 and Zimmer 2005 and Quigley and Hein Index, Quigley and Hein 1962); adverse effects ("discomfort in taste, discomfort in sensibility, gingival damage, gingival bleeding, staining of teeth and tongue, mouth burning during application, and white plaque on tongue immediately after use"), all measured at 4 and 8 weeks

Funding

Quote: "This study was supported by GlaxoSmithKline"

Notes

Sample size calculation: not reported

Adverse effects: reported with no double counting - if any participant experienced more than 1 adverse effect, only the most 'relevant' one was reported

- At 4 weeks: no side effects: Gp A: 21/39, Gp B: 15/39, Gp C: 36/39, Gp D: 37/39. Discomfort in taste: Gp A: 5/39, GP B: 7/39, Gp C: 0, Gp D: 1/39. Discomfort in sensibility: Gp A: 2/39, Gp B: 2/39, Gp C: 0, Gp D: 0. Gingival damage: Gp A: 0, Gp B: 0, Gp C: 3/39, Gp D: 0. Stain on teeth/tongue: Gp A: 4/39, Gp B: 9/39, Gp C: 0, Gp D: 0. Mouth burning during application: Gp A: 7/39, Gp B: 5/39, Gp C: 0, Gp D: 0. Bleeding of the gingiva: Gp A: 0, Gp B: 0, Gp C: 0, Gp D: 1. White plaque on tongue immediately after use: Gp A: 0, Gp B: 1/39, Gp C: 0, Gp D: 0
- At 8 weeks: no side effects: Gp A: 19/39, Gp B: 10/39, Gp C: 38/39, Gp D: 39/39. Discomfort in taste: Gp A: 6/39, GP B: 4/39, Gp C: 0, Gp D: 0. Discomfort in sensibility: Gp A: 2/39, Gp B: 1/39, Gp C: 0, Gp D: 0. Gingival damage: Gp A: 0, Gp B: 0, Gp C: 1/39, Gp D: 0. Stain on teeth/tongue: Gp A: 6/39, Gp B: 19/39, Gp C: 0, Gp D: 0. Mouth burning during application: Gp A: 6/39, Gp B: 4/39, Gp C: 0, Gp D: 0. Bleeding of the gingiva: Gp A: 0, Gp B: 0, Gp C: 0, Gp D: 0. White plaque on tongue immediately after use: Gp A: 0, Gp B: 1/39, Gp C: 0, Gp D: 0

Smokers: 33 (21.15%) overall. By group: Gp A: 10%; Gp B: 31%; Gp C: 28%; Gp D: 15%

Declarations/conflicts of interest: not reported

Risk of bias

Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Low risk	Quote: " the 156 participants were randomly assigned to four groups with 39 subjects in each group In a box containing 156 envelopes in four strata (three strata with 40 and one with 36), each participant had to draw one envelope containing the number of the attributed product"		
		Comment: drawing of lots is an adequate method of random sequence generation		
Allocation concealment (selection bias)	Low risk	Quote: " the 156 participants were randomly assigned to four groups with 39 subjects in each group In a box containing 156 envelopes in four strata (three strata with 40 and one with 36), each participant had to draw one envelope containing the number of the attributed product"		
		Quote: "The assignment of subjects to groups was performed by a person not involved in the examination"		
		Comment: allocation of subjects to groups was concealed up to the point that participants drew an envelope and were allocated to a group		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Comment: participants were not blinded. The different interventions meant that participants would know which group they were in and this could have affected their oral health behaviours and hence the outcome. The direction of this potential bias is not clear		
Blinding of outcome assessment (detection bias)	Low risk	Quotes: "The study was conducted by a blinded operator" and "Because clinically visible side effects might have influenced examiner blinding, an addi-		



Zimmer 2006 (Continued) All outcomes		tional statistical analysis was performed for subjects without visible side effects. Again, compared to all subjects, only small discrepancies of this subgroup and no changes in statistical significance were found. This finding indicates that clinically visible side effects did not affect examiner accuracy" Comment: aAlthough a risk of bias of outcome assessment due to staining of teeth/tongue in both active mouthrinse groups was likely, it did not appear to influence the results
Incomplete outcome data (attrition bias) All outcomes	Low risk	All randomised participants were included in the analyses
Selective reporting (reporting bias)	High risk	Adverse effects were reported with no double counting - if any participant experienced more than 1 adverse effect, only the 'most relevant' one was reported. No information is provided on how the 'most relevant' side effect was chosen to be reported and information about participants who experienced more than 1 adverse effect is not reported
Other bias	Low risk	Quote: "Before starting the study the intraexaminer reliability was tested by repeated measurements of MPPI, QHI, and PBI and analysed by Cohen's κ test. the results were as follows: MPPI=0.86, QHI=0.82 and PBI=0.88." No other apparent bias identified
		Quote: "All examinations were conducted by one single examiner (GK)"
		Smokers: CHX NaF (4) 10.26%, control (6) 15.38%. The authors state that this difference failed to reach statistical significance (P = 0.076)

Zimmer 2015

Methods	Trial design: parallel (4 arms - 3 arms are relevant to this review)
	Location: Dental School of the Witten/Herdecke University, Witten, Germany
	Number of centres: 1
	Study duration: quote: "Subjects were recruited between April 20th, 2010 and June16th, 2010." Study duration and duration of rinsing was 8 weeks
Participants	Participants: healthy adults
	Inclusion criteria: quotes: "subjects aged 18–65 years and having a mean PBI per tooth >0.5 were en-

Inclusion criteria: quotes: "subjects aged 18–65 years and having a mean PBI per tooth ≥0.5 were enrolled in the study" and "... Informed consent was obtained by each subject. Each subject had a minimum of 20 permanent natural teeth. Third molars, orthodontically banded, fully crowned or extensively restored or abutment teeth were not included in the tooth count"

Exclusion criteria: quote: "Subjects were excluded from the study if they were pregnant or breast-feeding, had diabetes type I or II, or severe periodontal disease, or wore removable dentures. Severe periodontal disease was defined as the presence of clinical attachment loss of more than 5 mm according to the periodontal disease classification of the American Academy of Periodontology.... However, a subject was excluded only if this applied for a minimum of three teeth. Subjects were also excluded if any of the following conditions was present: use of antibiotics within 2 weeks prior the first examination or use of any systemic medication which would have an effect on gingival conditions within 30 days prior to the screening visit, or recent history (within the last year) of alcohol or other substance abuse. Dental professionals, dental students and employees of the sponsor or members of their immediate families were also excluded from the study"

Baseline gingivitis: (Papillary Bleeding Index) Gp A: mean 1.06 (SD 0.4); Gp B: mean 0.99 (SD 0.37); Gp C: mean 0.94 (SD 0.31); Gp D: mean 1.02 (SD 0.39).



Zimmer 2015 (Continued)

Age (years): overall: mean 33.9. Gp A: mean 33.8; Gp B: mean 34.0; Gp C: mean 33.9; Gp D: mean 33.8. Range 18-65 years

Gender: nt reported

Number randomised: 160 (Gp A: 40; Gp B: 40; Gp C: 40; Gp D: 40) Number evaluated: 155 (Gp A: 39; Gp B: 39; Gp C: 40; Gp D: 37)

Interventions

Comparison: (twice daily toothbrushing and 0.06% CHX digluconate + 0.025% F as sodium fluoride mouthrinse) versus (twice daily toothbrushing and an experimental alcohol-free mouthrinse with 0.06% CHX digluconate + 0.025% F as sodium fluoride) versus (twice daily toothbrushing alone) versus (twice daily toothbrushing and an experimental alcohol-free mouthrinse with 0.06% CHX digluconate + 0.03% cetylpyridinium chloride (CPC) + 0.025% F as sodium fluoride)*

Gp A (n = 40): twice daily toothbrushing and rinsing twice a day for 30 seconds with 10 mL of an alcohol-containing mouthrinse with 0.06% CHX digluconate + 0.025% fluoride as sodium fluoride for 8 weeks

Gp B (n = 40): twice daily toothbrushing and rinsing twice a day for 30 seconds with 10 mL of an experimental alcohol-free mouthrinse with 0.06% CHX digluconate + 0.025% fluoride as sodium fluoride for 8 weeks

Gp C (n = 40): twice daily toothbrushing alone

*Gp D (n = 40): twice daily toothbrushing and rinsing twice a day for 30 seconds with 10 mL of an experimental alcohol-free mouthrinse with 0.06% CHX digluconate + 0.03% cetylpyridinium chloride (CPC) + 0.025% fluoride as sodium fluoride. We excluded this arm from our risk of bias and analysis

Prophylaxis at baseline: not reported

OHI: quotes: "All participants received a short instruction on how to conduct the attributed oral hygiene measurements. The subjects were advised to brush their teeth in the morning and in the evening postprandial in the usual manner. No instructions concerning brushing technique and brushing duration were given. After using the brush, the subjects rinsed their mouth with tap water" and "During the study period, the use of other than the attributed oral hygiene tools strictly was prohibited... Interproximal cleaning devices had been permitted if they were part of the usual oral hygiene routine of the included subjects"

Timing of toothbrushing in relation to mouthrinsing: quote: "The subjects of the rinsing groups waited for 30 min. Thereafter, they rinsed for 30 s with 10 ml of the assigned rinse. Afterwards, the subjects refrained from drinking, eating and rinsing for at least 30 min. To control the waiting and the rinsing time, each subject was provided with a digital stop watch"

Outcomes

Gingivitis (gingival bleeding: Papillary Bleeding Index, Saxer 1975); plaque (TQH, Turesky 1970; Modified Proximal Plaque Index, Zimmer 2005) assessed at 4 and 8 weeks follow-up. All adverse events occurring during the study period were recorded using a questionnaire and by clinical examination. Occurrence of discomfort in taste, discomfort in sensibility, gingival damage, gingival bleeding, staining of teeth and tongue, mouth burning and white plaque on tongue immediately after use was registered at baseline and after 4 and 8 weeks. Oral soft tissue examination was performed at each visit

Funding

Quote: "This study was funded by GlaxoSmithKline Consumer Healthcare (Bühl, Germany). GlaxoSmithKline was involved in the design and conduct of the study and provided logistical support during the trial. Data management and statistical analysis were performed by an independent institute at the Heinrich-Heine-University Düsseldorf. The manuscript was prepared by Prof. Zimmer, Dr. Naumova and Dr. Jordan. GlaxoSmithKline was permitted to review the manuscript and suggest changes, but the final decision on content was exclusively retained by the authors"

Notes

Sample size calculation: quote: "... was calculated on the basis of the following assumptions: power 0.8; alpha error: 0.05; delta-QHI between groups after 8 weeks: 0.3 (SD 0.5); one tailed analysis (Axum 7.0)"

Adverse effects: tooth staining at 8 weeks: Gp A: 18/39, Gp B: 21/39 and Gp C: 8/40. Tongue staining at 8 weeks: Gp A: 18/39, Gp B: 22/39 and Gp C: 14/40



Zimmer 2015 (Continued)

Gastrointestinal adverse events were found in all groups (Gp 1 (A): 1 case, Gp 2 (B): 2 cases, Gp 3 (no data extracted): 7 cases, Gp 4 (C): 1 case); in 5 cases (1 in Gp 2 (B) and 4 in Gp 3 (no data extracted)), gastrointestinal infection was given as reason; in 4 cases, diarrhoea; and in 1 case, 'stomach burning' and meteorism, respectively. No serious adverse events were observed in the study

Smokers by group: Gp A: 10/40 (25%); Gp B: 9/40 (23%); Gp C: 10/40 (25%) (differences between the groups were not statistically significant)

The trial was registered at the Clinical Trials Register of the National Institutes of Health (NCT01811615, www.clinicaltrials.gov)

Declarations/conflicts of interest: none stated

Tooth staining data for groups A and B (at 4 weeks) were combined in the main analysis

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Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Using the stratification by sex and PBI (PBI/tooth≥0.5 and<1.0 or PBI/tooth≥1.0), the 160 eligible participants were allocated by block randomization to one of four groups with 40 subjects each. Block randomization was performed by a statistician not involved in the study"
		Comment: adequate method of random sequence generation
Allocation concealment (selection bias)	Low risk	Quotes: "Subjects within each group were randomly assigned to one of the following groups" and "Block randomization was performed by a statistician not involved in the study"
		Comment: we consider it unlikely that participants and investigators enrolling participants could foresee assignment
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quotes: "All personal instructions and delivering of the study products were performed by a study nurse not involved (A.P.) in the study examinations. The subjects were forbidden to tell the examiner their regimen during the study visits" and "With respect to tooth staining, chi square-test revealed significant more occurrences in the three rinsing groups when compared to the negative control (P<0.001)"
		Comment: due to the design of the study the participants in toothbrushing only group could not be blinded. The higher incidence of staining in the CHX groups means that blinding may not be possible in a comparison against the control
Blinding of outcome assessment (detection bias) All outcomes	High risk	The study was designed to be examined blind. Quotes: "All assessments of an individual subject in the course of the study have been conducted by the same investigator (P.K.) who had experience from previous clinical studies" and "with respect to tooth staining, chi square-test revealed significantly more occurrences in the rinsing groups when compared to the negative control (P<0.001)"
		Comment: the higher incidence of staining in the CHX groups means that blinding may not be possible in a comparison against the control
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition 2/120 (1.7%). By group: Gp A: 1/40 (0.03%); Gp B: 1/40 (0.03%); Gp C: no losses. Reasons for losses: Gp A: informed consent withdrawn; Gp B: protocol violation
		Comment: attrition is minimal and unlikely to affect the outcome



Zimmer 2015 (Continued)		
Selective reporting (reporting bias)	Low risk	All outcomes stated in the methods sections were reported in the results section $ \label{eq:control} % \begin{center} \begin$
Other bias	Unclear risk	No mention of intraexaminer reproducibility
		Groups were balanced at baseline for age and smoking status, gingivitis and plaque
		Tooth staining at baseline is not reported

ASF = amine fluoride and stannous fluoride; BOP = bleeding on probing; CAL = clinical attachment level; CHX = chlorhexidine; CI = Calculus Index; CI = confidence interval; CSSI = Calculus Surface Severity Index; DI = Discolouration Index; F = fluoride; Gp = group; GI = Gingival Index; ITT = intention-to-treat analysis; MSE = mean squared error; OH = oral hygiene; OHI = oral hygiene instruction; PAL = probing attachment level; PBS = Papillary Bleeding Score; PD = pocket depth; PI = Plaque Index; PMGI = Papillary Marginal Gingival Index; PPD = probing pocket depth; ppm = parts per million; PVP = polyvinyl pyrrolidone; SD = standard deviation; SE = standard error; SES = socioeconomic status; SRP = scaling and root planing; TQH = Turesky modification of the Quigley and Hein Index; VAS = visual analogue scale.

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Bay 1975	It is unclear from the published report whether the study is a randomised trial. Attempts to contact the first author for clarification were unsuccessful
Biswas 2014	Comparison is with another active rinse and there is no placebo/control or mechanical OH only group
Bouwsma 1992	The test group used a triangular wooden interdental cleaner once daily after toothbrushing which the CHX group did not use. The mechanical OH procedures were not the same in both groups
Brown 2002	The study compares CHX plus mechanical OH and 2-monthly professional prophylaxis versus mechanical OH and 6-monthly check up and professional prophylaxis (standard care). The mechanical OH procedures were not the same in both groups
Caton 1993	The CHX group received OH instructions but the control group did not. The mechanical OH procedures were potentially not the same in both groups
Cortelli 2015	The CHX arms used CHX gel (tongue brushing for 1 min after rinsing) in addition to CHX rinse. We were unable to separate the effect of the CHX rinse from the effect of the other active agent
CTRI 2014	Comparison is with another active rinse and there is no placebo/control or mechanical OH only group
Ernst 1998	Comparison is with another active rinse and there is no placebo/control or mechanical OH only group
Eshwar 2016	Comparison is with another active rinse and there is no placebo/control or mechanical OH only group
Fischman 1975	This study compared CHX, zinc and placebo mouthrinses but each rinse group was further split into those who received a professional prophylaxis prior to the start of the study and those that did not. The study outcomes were not reported in a format that could be used in the review and data were not available from the authors of the study
Goutham 2013	It is unclear from the published report whether the study is a randomised trial. Attempts to contact the first author for clarification were unsuccessful



Study	Reason for exclusion
Gupta 2014	Concern about inclusion of duplicated outcome data from another study by the same authors
Gupta 2015	Concern about inclusion of duplicated outcome data from another study by the same authors
Gupta 2015a	Concern about inclusion of duplicated outcome data from other studies in other published trials from these researchers
Haffajee 2009	Comparison is with another active rinse and there is no placebo/control or mechanical OH only group
Karim 2014	Concern about inclusion of duplicated outcome data from another study by the same authors
Lang 1982	It is unclear from the published report whether the study is a randomised trial. Attempts to contact the first author for clarification were unsuccessful
Leyes Borrajo 2002	It is unclear from the published report whether the study is a randomised trial. Additional information obtained from the contact author indicate quasi-randomisation (participants were assigned to groups according to the order that they came to the university)
Luoma 1978	CHX, F and placebo groups also used a toothpaste at evenings and weekend with the same composition as their respective rinse solution. Therefore the CHX rinse group also used a CHX-containing toothpaste. We were unable to separate the effect of the CHX rinse from the effect of the other active agent
Madden 2008	The CHX group received a more intensive intervention including 2-monthly OH instruction compared to the control group who received 6-monthly OH alone. The mechanical OH procedures were (potentially) not the same in both groups
McKenzie 1992	Comparison is with another active rinse and there is no placebo/control or mechanical OH only group
Nadkerny 2015	Duration of rinsing was 15 days
NCT01750801	Comparison is with another active rinse and there is no placebo/control or mechanical OH only group
NCT02546804	Comparison is with another active rinse and hot salt mouthwash and there is no placebo/control or mechanical OH only group
Ousehal 2011	It is unclear from the published report whether the study is a randomised trial. Attempts to contact the first author for clarification were unsuccessful
Paknejad 2006	It is unclear from the published report whether the study is a randomised trial. Attempts to contact the first author for clarification were unsuccessful
Persson 1991	Comparison is with another active rinse and there is no placebo/control or mechanical OH only group
Persson 1998	The CHX group received a more intensive intervention including cognitive behavioural education (2 hours) compared to the control group who received standard care
Priya 2015	Comparison is with another active rinse and there is no placebo/control or mechanical OH only group



Study	Reason for exclusion
Quirynen 2005	The study involves 1-stage full-mouth disinfection involving use of CHX gel at baseline in addition to the mouthrinses. We were unable to separate the effect of the CHX rinse from the effect of the other active agent
Quirynen 2006	The study involves 1-stage full-mouth disinfection involving use of CHX gel at baseline and 1 week in addition to the mouthrinses. We were unable to separate the effect of the CHX rinse from the effect of the other active agent
Radafshar 2017	Comparison is with another active rinse and there is no placebo/control or mechanical OH only group
Saltini 1988	This is an abstract. It was not possible to contact the first author, therefore co-authors were contacted but they did not have access to the data
Segreto 1993	We were unable to obtain a copy of this unpublished study despite following up several lines of enquiry
Subhash 1985	The brushing method differs between the groups. The mechanical OH procedures are not considered to be the same in both groups
Todkar 2012	It is unclear from the published report whether the study is a randomised trial. Attempts to contact the first author for clarification were unsuccessful
Vechis-Bon 1989	It is unclear from the published report whether the study is a randomised trial. Attempts to contact the first author for clarification were unsuccessful
Yates 2002	Experimental gingivitis study, the CHX mouthrinse is used in the absence of any mechanical tooth cleaning

CHX = chlorhexidine; OH = oral hygiene; F = fluoride.

Characteristics of studies awaiting assessment [ordered by study ID]

CTRI/2011/05/001774

Methods	Study design: parallel group 3 arms
	Location: Apollo Hospital, Dental and Facial Surgical Centre, RV Dental College, Santosh Hospital, Bangalore, Karnataka, India
	Number of centres: appears to be 4
	Study duration: 7/8 months. Duration of rinsing appears to be 6 months
Participants	Participants: minimum 12 years old with mild to moderate gingivitis
	Inclusion criteria: with a reasonable standard of oral hygiene with no severe gingivitis (score of not 2 on the GI). Subjects with mild to moderate gingivitis, with probing depths of not more than 3 mm
	Age 12-99 eligible for inclusion
	Total sample size 120; 96 evaluable subjects required. Not clear how many subjects actually participated in trial
Interventions	Chlorhexidine 0.12% (Orocleanse) with alcohol versus chlorhexidine 0.12% (Oroclear) without alcohol versus placebo in conjunction with regular self-performed oral hygiene measures



CTRI/2011/05/001774 (Continued)	Rinsing with 15 mL twice daily for 30 seconds for 6 months
Outcomes	Gingival inflammation: GI, also stain indices
	Brief summary of results mentions PI, PBI and CI not specified in methods
Notes	Study completed. Brief summary of results reported but no outcome data are reported. We contacted the author for further information but did not receive a reply

NCT01898000

Methods	Study design: parallel, (3 arms - 2 arms relevant to this review)
	Location: Government Dental College and Research Institute, Bangalore, India
	Number of centres: not clear
	Study duration: duration of rinsing 60 days
Participants	Participants: subjects with chronic generalised gingivitis reporting to outpatient Department of Periodontics, Government Dental College and Research Institute, Bangalore from January 2013 to April 2013
	Quote: "instructed not to rinse/eat anything for 30 min after mouthwash use. Subjects were also asked to refrain from all other unassigned forms of oral hygiene aids, including dental floss and chewing gum during the study. No oral hygiene instructions like brushing and flossing were given to the subjects to exclude the influence of improved oral hygiene practices on the results"
Interventions	Group I: placebo mouthwash. Group II: Triphala (TRP) mouthwash. Group III: chlorhexidine mouthwash. Subjects were instructed to use 15 mL of mouthwash twice daily
Outcomes	PI, GI and OHI-S, and microbiological colony counts were recorded on baseline and 7, 30 and 60 days respectively
Notes	Study completed. Brief summary of results reported but no outcome data are reported. We contacted the author for further information but did not receive a reply

NCT02065414

Methods	Study design: parallel, (3 arms - 2 arms are relevant to this review)
	Location: not clear "Clinical Centre"
	Number of centres: 1
	Study duration: 4 weeks
Participants	Participants: aged 18 years and older with gingivitis, but without moderate/advanced periodontitis
Interventions	Chlorhexidine rinse (Corsodyl) 0.12% versus experimental mouthrinse (Listerine Advance Gum Defense) versus placebo
	Chlorhexidine: rinse with 10 mL for 60 seconds twice daily for 4 weeks (5 min after brushing and rinsing mouth with water)
	Experimental rinse: rinse with 20 mL for 30 seconds twice daily for 4 weeks (immediately after brushing and rinsing mouth with water)



NCT02065414 (Continued)	
	Placebo: rinse with 20 mL for 30 seconds twice daily for 4 weeks (immediately after brushing and rinsing mouth with water)
Outcomes	Gingivitis: gingival inflammation, Modified Gingival Index at 4 weeks; gingival bleeding, Gingival Bleeding Index at 1 and 4 weeks. Plaque: Turesky Modification of the Quigley and Hein Plaque Index at 4 weeks. Extrinsic tooth stain: Macpherson Modification of the Lobene Stain Index at 4 week
Notes	Study completed. Contacted investigator for further details but did not receive a reply
Radvar 2016	
Methods	Trial design: parallel, (3 arms - 2 arms relevant to this review) (study number 2 only, study number 1 does not meet the inclusion criteria for the review)
	Location: Periodontology Department, Mashad Dental School, Iran
	Number of centres: assume 1 centre
	Study duration: recruitment period not stated. Duration of rinsing was 4 weeks and the outcome was assessed 6 weeks after scaling and root planing
Participants	Participants: outpatients with chronic periodontitis referred to the Periodontology Department, Mashad Dental School, Iran
	Inclusion criteria: chronic periodontitis with pocketing and attachment loss in all quadrants (over 45 years old)
	Exclusion criteria: "Patients on antibiotic therapy or anti-inflammatory drugs, those with any history of systemic disease, or allergy to components of the mouthwash, those who had undergone any form of non-surgical or surgical periodontal therapy in the last 6 months, pregnant and lactating mothers and smokers were excluded"
	Baseline gingivitis: BOP, Group A (66.8 \pm 14.85); Group B (71.08 \pm 10.23); Group C (57.18 \pm 19.47)
	Age at baseline (years): not reported. Over 45 years old
	Gender: not reported
	Number randomised: 30 (number randomised to each group not reported, assumed 10 in each group at baseline)
	Number evaluated: not reported
Interventions	Chlorhexidine rinse versus herbal rinse* versus control rinse
	Group A (assume n = 10): CHX (concentration not reported) 10 mL, 1 min, twice daily
	*Group B (assume n = 10): herbal mouthwash (<i>S alba, M sylvestris and A officinalis</i>) 10 mL, 1 min, twice daily. (We excluded this arm from our risk of bias and analysis)
	Group C (assume n = 10): control (normal saline) 10 mL, 1 min, twice daily
	Prophylaxis at baseline: quote: "Scaling and root planing was accomplished in two visits for all patients and local anaesthesia was used when required"
	OHI: not reported
	Non-supervised rinsing
	Timing of mouthrinsing in relation to toothbrushing: not reported



Radvar 2016 (Continued)	Postrinsing instructions: quote: "Subsequent rinsing with water was not allowed" Duration of rinsing: 4 weeks
Outcomes	Gingivitis: BOP (no reference), PD, CAL were measured using a periodontal probe at baseline and then 6 weeks after scaling and root planing
Notes	Sample size calculation: not reported Adverse effects: quote: "No adverse reaction was seen in the herbal mouthwash group, however, in the CHX group there were some adverse reactions such as dental and tongue staining" Declarations/conflicts of interest: none reported It is not clear exactly when the outcome was assessed. Outcome assessment appears to have been completed 2 weeks after rinsing ceased We are awaiting clarification from the authors about various aspects of this study

BOP = bleeding on probing; CAL = clinical attachment level; CHX = chlorhexidine; CI = Calculus Index; GI = Gingival Index; OHI = oral hygiene instruction; OHI-S = Oral Hygiene Index-Simplified; PBI = Papillary Bleeding Index; PD = pocket depth; PI = Plaque Index.

Characteristics of ongoing studies [ordered by study ID]

NCT02709785

Trial name or title	SmartMouth Advanced Clinical Formula clinical research design protocol
Methods	Trial design: parallel, (3 arms - 2 arms relevant to this review)
	Location: Saint Louis University Center for Advanced Dental Education, Saint Louis, Missouri, USA and Southern Illinois University School of Dental Medicine, Alton, Illinois, USA
	Number of centres: 2
	Study duration: March 2016 to June 2017. Study duration 10 weeks. Duration of rinsing 6 weeks
Participants	Participants: adults with gingivitis or chronic periodontitis
	Inclusion criteria: subjects between the ages of 18 to 80, with a diagnosis of gingivitis or chronic periodontitis, mean GI \geq 0.4 and PI \geq 1.0, subjects that are in good medical health at time of the study, at least 20 permanent natural teeth
	Exclusion criteria: use of local or systemic antibiotics during the course of the study, subjects that are pregnant or nursing mothers, undergoing orthodontic therapy, subjects wearing removable prostheses, subjects taking medication which is altering the gingiva or causing inflammation, gingival overgrowth, history of sensitivity or suspected allergies following the use of oral hygiene products, subjects taking anti-inflammatory or anticoagulant medications that would alter the gingiva and promote bleeding, subjects that require antibiotic prophylaxis prior to dental treatment, subjects that have acute dental problems requiring immediate treatment
	Baseline gingivitis: GI ≥ 0.4 (Löe & Silness 1963)
	Age at baseline: 18-80 years are eligible
	Prophylaxis: all subjects will be given a complete dental prophylaxis to remove plaque, calculus, and stain
	OHI: written and verbal OHI will be given that include a regular regimen of brushing twice daily and flossing daily. All subjects will be assigned a commercially available ADA-accepted toothbrush and dental floss



NCT02709785 (Continued)	Number to be randomised: 85 (each group will have at least 25 subjects)
Interventions	0.12% chlorhexidine rinse versus SmartMouth Advanced Clinical Formula (ACF) rinse versus place- bo rinse
Outcomes	Gingivitis: gingival inflammation: GI Löe & Silness 1963; gingival bleeding (GI scores 2 and 3); Turesky modification of the Quigley and Hein Plaque Index, Turesky 1970. Extrinsic tooth staining: modification of the Lobene Tooth Stain Index, Lobene 1968. Supragingival calculus on the lingual surfaces of the 6 mandibular anterior teeth will be scored using the Volpe-Manhold Calculus Index, Volpe 1965. Safety assessments and assessment of the oral soft tissues will be conducted at each visit by visual examination of the oral cavity. Evaluation of taste perception, malodour via questionnaire
Starting date	March 2016
Contact information	Contact: David D Miley, Saint Louis University Center for Advanced Dental Education, Saint Louis, Missouri, USA
Notes	Sponsor: Saint Louis University
	Collaborator: Triumph Pharmaceuticals
	NCT02709785

ADA = American Dental Association; GI = Gingival Index; PI = Plaque Index; OHI = oral hygiene instruction.

DATA AND ANALYSES

Comparison 1. CHX versus placebo/control mouthrinse or no mouthrinse

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Gingival Index (0-3) 4-6 weeks	10	805	Mean Difference (IV, Random, 95% CI)	-0.21 [-0.31, -0.11]
1.1 CHX versus no rinse	2	339	Mean Difference (IV, Random, 95% CI)	-0.13 [-0.31, 0.05]
1.2 CHX versus placebo/control rinse	8	466	Mean Difference (IV, Random, 95% CI)	-0.23 [-0.34, -0.13]
2 Gingival Index (0-3) 6 months	13	2616	Mean Difference (IV, Random, 95% CI)	-0.20 [-0.30, -0.11]
2.1 CHX versus no rinse	2	142	Mean Difference (IV, Random, 95% CI)	-0.12 [-0.18, -0.05]
2.2 CHX versus placebo/control rinse	11	2474	Mean Difference (IV, Random, 95% CI)	-0.22 [-0.33, -0.11]
3 Gingival bleeding 4-6 weeks	8	649	Std. Mean Difference (IV, Random, 95% CI)	-0.56 [-0.79, -0.33]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
3.1 CHX versus no rinse	4	459	Std. Mean Difference (IV, Random, 95% CI)	-0.69 [-0.89, -0.50]
3.2 CHX versus placebo/control rinse	4	190	Std. Mean Difference (IV, Random, 95% CI)	-0.36 [-0.77, 0.06]
4 Gingival bleeding 6 months	8	1132	Std. Mean Difference (IV, Random, 95% CI)	-0.72 [-1.02, -0.42]
4.1 CHX versus no rinse	2	142	Std. Mean Difference (IV, Random, 95% CI)	-0.49 [-0.83, -0.16]
4.2 CHX versus placebo/control rinse	6	990	Std. Mean Difference (IV, Random, 95% CI)	-0.79 [-1.16, -0.41]
5 Plaque 4-6 weeks	12	950	Std. Mean Difference (IV, Random, 95% CI)	-1.45 [-1.90, 1.00]
5.1 CHX versus no rinse	3	433	Std. Mean Difference (IV, Random, 95% CI)	-1.43 [-2.39, -0.47]
5.2 CHX versus placebo/control rinse	9	517	Std. Mean Difference (IV, Random, 95% CI)	-1.48 [-2.07, -0.89]
6 Plaque 4-6 weeks PI (0-3)	4	223	Mean Difference (IV, Random, 95% CI)	-0.58 [-0.78, -0.39]
6.1 CHX versus no rinse	2	114	Mean Difference (IV, Random, 95% CI)	-0.59 [-0.94, -0.24]
6.2 CHX versus placebo/control rinse	2	109	Mean Difference (IV, Random, 95% CI)	-0.50 [-0.97, -0.04]
7 Plaque 4-6 weeks TQH (0-5)	5	546	Mean Difference (IV, Random, 95% CI)	-0.78 [-0.85, -0.70]
7.1 CHX versus no rinse	1	319	Mean Difference (IV, Random, 95% CI)	-0.83 [1.00, -0.66]
7.2 CHX versus placebo/control rinse	4	227	Mean Difference (IV, Random, 95% CI)	-0.76 [-0.85, -0.68]
8 Plaque 6 months	11	2075	Std. Mean Difference (IV, Random, 95% CI)	-1.43 [-1.76, -1.10]
8.1 CHX versus no rinse	2	142	Std. Mean Difference (IV, Random, 95% CI)	-0.68 [-1.35, -0.01]
8.2 CHX versus placebo/control rinse	9	1933	Std. Mean Difference (IV, Random, 95% CI)	-1.59 [-1.89, -1.29]
9 Plaque 6 months PI (0-3)	5	1108	Mean Difference (IV, Random, 95% CI)	-0.62 [-1.12, -0.12]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
9.1 CHX versus no rinse	2	142	Mean Difference (IV, Random, 95% CI)	-0.30 [-0.42, -0.18]
9.2 CHX versus placebo/control rinse	3	966	Mean Difference (IV, Random, 95% CI)	-0.86 [-1.46, -0.25]
10 Plaque 6 months TQH (0-5)	6	967	Mean Difference (IV, Random, 95% CI)	-0.73 [-0.88, -0.57]
10.1 CHX versus placebo/control rinse	6	967	Mean Difference (IV, Random, 95% CI)	-0.73 [-0.88, -0.57]
11 Calculus 4-6 weeks	2		Mean Difference (IV, Random, 95% CI)	Subtotals only
11.1 CHX versus placebo/control rinse	2	102	Mean Difference (IV, Random, 95% CI)	0.02 [-0.09, 0.14]
12 Calculus 7-12 weeks	6	425	Std. Mean Difference (IV, Random, 95% CI)	0.32 [-0.04, 0.69]
12.1 CHX versus no rinse	1	95	Std. Mean Difference (IV, Random, 95% CI)	1.02 [0.59, 1.45]
12.2 CHX versus placebo/control rinse	5	330	Std. Mean Difference (IV, Random, 95% CI)	0.14 [-0.08, 0.36]
13 Calculus 6 months	4	323	Std. Mean Difference (IV, Random, 95% CI)	0.80 [0.33, 1.26]
13.1 CHX versus no rinse	1	91	Std. Mean Difference (IV, Random, 95% CI)	1.39 [0.93, 1.85]
13.2 CHX versus placebo/control rinse	3	232	Std. Mean Difference (IV, Random, 95% CI)	0.60 [0.24, 0.96]
14 Tooth staining 4-6 weeks dichoto- mous	2	156	Risk Ratio (M-H, Random, 95% CI)	5.41 [2.03, 14.47]
14.1 CHX versus no rinse	1	118	Risk Ratio (M-H, Random, 95% CI)	4.44 [1.43, 13.80]
14.2 CHX versus placebo/control rinse	1	38	Risk Ratio (M-H, Random, 95% CI)	9.88 [1.37, 71.44]
15 Tooth staining 7-12 weeks dichoto- mous	1		Risk Ratio (M-H, Random, 95% CI)	Subtotals only
15.1 CHX versus no rinse	1	118	Risk Ratio (M-H, Random, 95% CI)	2.5 [1.29, 4.83]
16 Tooth staining 4-6 weeks	8	415	Std. Mean Difference (IV, Random, 95% CI)	1.07 [0.80, 1.34]
16.1 CHX versus no rinse	1	94	Std. Mean Difference (IV, Random, 95% CI)	1.54 [1.08, 2.00]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
16.2 CHX versus placebo/control rinse	7	321	Std. Mean Difference (IV, Random, 95% CI)	0.97 [0.73, 1.22]
17 Tooth staining 7-12 weeks	11	581	Std. Mean Difference (IV, Random, 95% CI)	1.19 [0.98, 1.40]
17.1 CHX versus no rinse	1	95	Std. Mean Difference (IV, Random, 95% CI)	1.32 [0.88, 1.77]
17.2 CHX versus placebo/control rinse	10	486	Std. Mean Difference (IV, Random, 95% CI)	1.17 [0.93, 1.41]
18 Tooth staining 6 months	4	323	Std. Mean Difference (IV, Random, 95% CI)	1.54 [1.22, 1.86]
18.1 CHX versus no rinse	1	91	Std. Mean Difference (IV, Random, 95% CI)	1.18 [0.73, 1.62]
18.2 CHX versus placebo/control rinse	3	232	Std. Mean Difference (IV, Random, 95% CI)	1.69 [1.38, 1.99]

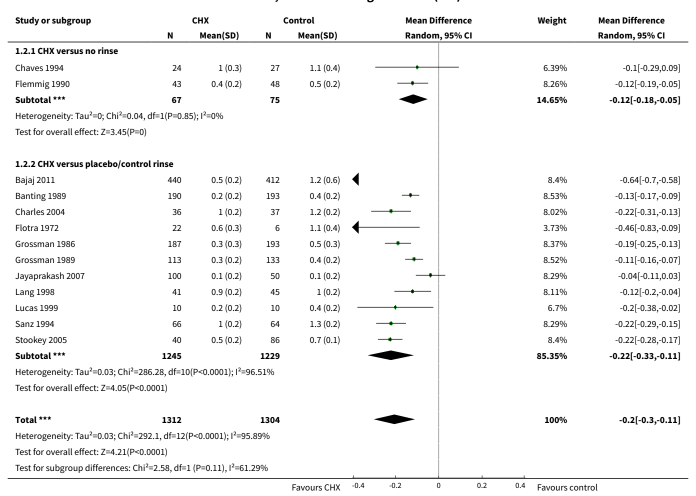
Analysis 1.1. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 1 Gingival Index (0-3) 4-6 weeks.

Study or subgroup		СНХ	C	Control	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
1.1.1 CHX versus no rinse							
Jose 2015	214	1.1 (0.1)	105	1.2 (0.1)	+	20.28%	-0.08[-0.11,-0.04]
Rahmani 2006	10	1.1 (0.3)	10	1.4 (0.3)		8.05%	-0.28[-0.55,-0.01]
Subtotal ***	224		115			28.33%	-0.13[-0.31,0.05]
Heterogeneity: Tau ² =0.01; Chi ² =2	2.16, df=1(P=	0.14); I ² =53.61%					
Test for overall effect: Z=1.44(P=0	0.15)						
1.1.2 CHX versus placebo/contr	rol rinse						
Axelsson 1987	42	0.6 (0.3)	22	1 (0.3)		13.75%	-0.37[-0.53,-0.22]
Bhat 2014	22	0.4 (0.4)	22	1 (0.8)		5.17%	-0.6[-0.98,-0.22]
Ernst 2005	33	0.5 (0.5)	28	0.6 (0.5)		9.4%	-0.11[-0.35,0.13]
Graziani 2015	55	0.7 (0.7)	15	0.8 (0.7)		4.6%	-0.12[-0.52,0.29]
Jenkins 1993	49	0.5 (0.3)	50	0.7 (0.3)	 + │	16.8%	-0.18[-0.28,-0.08]
Lopez-Jornet 2012	35	0.8 (0.8)	35	1.2 (0.9)		4.82%	-0.4[-0.79,-0.01]
Navarro 1998	9	0 (0.1)	11	0.2 (0.3)		12.12%	-0.13[-0.31,0.06]
Sanz 1989	17	1.1 (0.6)	21	1.2 (0.6)		5.01%	-0.13[-0.51,0.26]
Subtotal ***	262		204		•	71.67%	-0.23[-0.34,-0.13]
Heterogeneity: Tau ² =0.01; Chi ² =1	11.37, df=7(P	=0.12); I ² =38.44%	6				
Test for overall effect: Z=4.44(P<0	0.0001)						
Total ***	486		319		•	100%	-0.21[-0.31,-0.11]
Heterogeneity: Tau ² =0.01; Chi ² =2	27.51, df=9(P	=0); I ² =67.28%					
Test for overall effect: Z=4.11(P<0	0.0001)						
<u>·</u>	•			Favours CHX	-0.5 -0.25 0 0.25 0.5	Favours cor	ntrol



Study or subgroup		снх		Control Mean Difference Weight		Weight Mean Difference				
	N	Mean(SD)	N	Mean(SD)		Rand	lom, 9	5% CI		Random, 95% CI
Test for subgroup differences: Chi ² =0.94, df=1 (P=0.33), l ² =0%										
				Favours CHX	-0.5	-0.25	0	0.25	0.5	Favours control

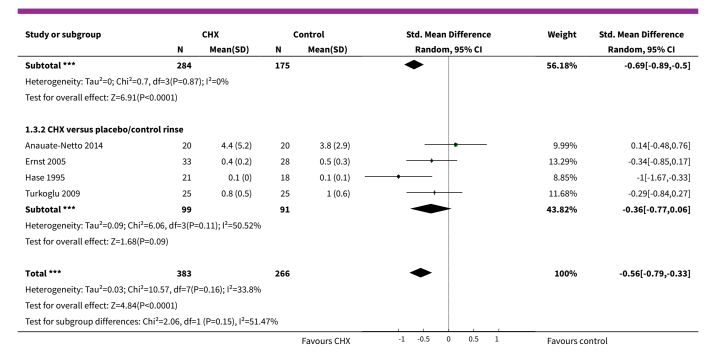
Analysis 1.2. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 2 Gingival Index (0-3) 6 months.



Analysis 1.3. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 3 Gingival bleeding 4-6 weeks.

Study or subgroup	СНХ		c	ontrol	Std. Mean Difference	Weight	Std. Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
1.3.1 CHX versus no rinse							
Jose 2015	214	0.1 (0.1)	105	0.2 (0.1)		27.62%	-0.66[-0.9,-0.42]
Rahmani 2006	10	0.3 (0.2)	10	0.5 (0.3)		5.09%	-0.97[-1.91,-0.03]
Taller 1993	13	0.4 (0.2)	13	0.5 (0.1)	 	6.47%	-0.92[-1.73,-0.1]
Van Strydonck 2008	47	0.7 (0.3)	47	1 (0.3)		17.01%	-0.67[-1.09,-0.26]
				Favours CHX	-1 -0.5 0 0.5 1	Favours co	ntrol





Analysis 1.4. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 4 Gingival bleeding 6 months.

Study or subgroup		СНХ	c	ontrol	Std. Mean Difference	Weight	Std. Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
1.4.1 CHX versus no rinse							
Chaves 1994	24	0.2 (0.1)	27	0.3 (0.2)	-+-	10.38%	-0.58[-1.15,-0.02]
Flemmig 1990	43	0.3 (0.1)	48	0.3 (0.1)		12.43%	-0.44[-0.86,-0.03]
Subtotal ***	67		75		•	22.81%	-0.49[-0.83,-0.16]
Heterogeneity: Tau ² =0; Chi ² =0.15, df	=1(P=0.7); I ² =0%					
Test for overall effect: Z=2.89(P=0)							
1.4.2 CHX versus placebo/control	rinse						
Grossman 1986	187	0.1 (0.1)	193	0.1 (0.1)	+	15.2%	-0.48[-0.68,-0.28]
Grossman 1989	113	0 (0.1)	133	0.1 (0.1)		14.61%	-0.57[-0.83,-0.31]
Hase 1998	30	0.3 (0.2)	33	0.4 (0.2)	-+-	11.18%	-0.55[-1.05,-0.04]
Lang 1998	45	0.1 (0.1)	47	0.3 (0.1)		11.36%	-1.85[-2.34,-1.36]
Overholser 1990	41	0.3 (0.3)	42	0.3 (0.4)	-+ 	12.21%	-0.24[-0.67,0.19]
Stookey 2005	40	0.1 (0.1)	86	0.2 (0.1)		12.64%	-1.16[-1.56,-0.75]
Subtotal ***	456		534		•	77.19%	-0.79[-1.16,-0.41]
Heterogeneity: Tau ² =0.18; Chi ² =35.6	6, df=5(P	<0.0001); I ² =85.9	8%				
Test for overall effect: Z=4.08(P<0.00	01)						
Total ***	523		609		•	100%	-0.72[-1.02,-0.42]
Heterogeneity: Tau ² =0.14; Chi ² =36.5	7, df=7(P	<0.0001); I ² =80.8	6%				
Test for overall effect: Z=4.7(P<0.000	1)						
Test for subgroup differences: Chi ² =:	1.28, df=1	L (P=0.26), I ² =22.	13%				
				Favours CHX	-2 -1 0 1 2	Favours co	ontrol



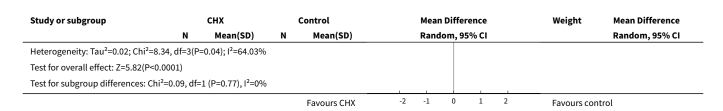
Analysis 1.5. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 5 Plaque 4-6 weeks.

Study or subgroup		СНХ	Control		Std. Mean Difference	Weight	Std. Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
1.5.1 CHX versus no rinse							
Jose 2015	214	2.5 (0.9)	105	3.3 (0.6)	•	9.92%	-1.05[-1.29,-0.8]
Rahmani 2006	10	0.5 (0.5)	10	0.9 (0.4)	+	7.17%	-0.79[-1.71,0.13]
Van Strydonck 2008	47	0.3 (0.3)	47	1 (0.3)	+	8.94%	-2.37[-2.9,-1.84]
Subtotal ***	271		162		•	26.04%	-1.43[-2.39,-0.47]
Heterogeneity: Tau ² =0.63; Chi ² =2	20.6, df=2(P<	0.0001); I ² =90.29	%				
Test for overall effect: Z=2.91(P=0	0)						
1.5.2 CHX versus placebo/cont	rol rinse						
Axelsson 1987	42	0.4 (0.5)	22	1.2 (0.5)	+	8.67%	-1.68[-2.27,-1.08]
Bhat 2014	22	0.7 (0.1)	22	1.5 (0.2)		6.06%	-4.59[-5.76,-3.42]
Ernst 2005	33	0.4 (0.2)	28	0.5 (0.2)	+	9%	-0.68[-1.2,-0.16
Graziani 2015	55	0.1 (0.1)	15	0.2 (0.1)	+	8.67%	-1.02[-1.61,-0.42]
Hase 1995	21	0 (0)	18	0.8 (0.4)	-	7.41%	-2.55[-3.41,-1.68]
Jenkins 1993	49	1.2 (0.8)	50	1.8 (0.5)	+	9.39%	-1[-1.42,-0.58]
Lopez-Jornet 2012	35	0.8 (0.8)	35	1.1 (0.9)	*	9.2%	-0.27[-0.74,0.2]
Navarro 1998	9	1.7 (0.8)	11	2.3 (0.4)	+	7.09%	-0.92[-1.86,0.02]
Turkoglu 2009	25	1.7 (0.6)	25	2.6 (0.5)	+	8.48%	-1.55[-2.19,-0.91]
Subtotal ***	291		226		♦	73.96%	-1.48[-2.07,-0.89]
Heterogeneity: Tau²=0.69; Chi²=6	55.44, df=8(P	<0.0001); I ² =87.7	7%				
Test for overall effect: Z=4.9(P<0.	0001)						
Total ***	562		388		•	100%	-1.45[-1.9,-1
Heterogeneity: Tau²=0.52; Chi²=8	86.66, df=11(P<0.0001); I ² =87.	31%				
Test for overall effect: Z=6.31(P<0	0.0001)						
Test for subgroup differences: Ch	ni ² =0.01, df=1	L (P=0.93), I ² =0%					

Analysis 1.6. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 6 Plaque 4-6 weeks PI (0-3).

Study or subgroup		СНХ	(Control	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
1.6.1 CHX versus no rinse							
Rahmani 2006	10	0.5 (0.5)	10	0.9 (0.4)	-+-	16.37%	-0.36[-0.74,0.02]
Van Strydonck 2008	47	0.3 (0.3)	47	1 (0.3)	=	36.8%	-0.73[-0.85,-0.61]
Subtotal ***	57		57		•	53.17%	-0.59[-0.94,-0.24]
Heterogeneity: Tau ² =0.05; Chi ² =3	3.24, df=1(P=	0.07); I ² =69.16%					
Test for overall effect: Z=3.3(P=0)							
1.6.2 CHX versus placebo/contr	rol rinse						
Hase 1995	21	0 (0)	18	0.8 (0.4)	+	31.11%	-0.71[-0.9,-0.52]
Lopez-Jornet 2012	35	0.8 (0.8)	35	1.1 (0.9)	-+ 	15.72%	-0.23[-0.63,0.17]
Subtotal ***	56		53		•	46.83%	-0.5[-0.97,-0.04]
Heterogeneity: Tau ² =0.09; Chi ² =4	1.63, df=1(P=	0.03); I ² =78.39%					
Test for overall effect: Z=2.12(P=0	0.03)						
Total ***	113		110		•	100%	-0.58[-0.78,-0.39]
				Favours CHX	-2 -1 0 1 2	Favours cor	ntrol





Analysis 1.7. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 7 Plaque 4-6 weeks TQH (0-5).

Study or subgroup		СНХ	c	Control	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
1.7.1 CHX versus no rinse							
Jose 2015	214	2.5 (0.9)	105	3.3 (0.6)	+	20.51%	-0.83[-1,-0.66]
Subtotal ***	214		105		◆	20.51%	-0.83[-1,-0.66]
Heterogeneity: Not applicable							
Test for overall effect: Z=9.73(P<0	0.0001)						
1.7.2 CHX versus placebo/contr	ol rinse						
Axelsson 1987	42	0.4 (0.5)	22	1.2 (0.5)	-	9.94%	-0.79[-1.03,-0.55]
Bhat 2014	22	0.7 (0.1)	22	1.5 (0.2)	H	58.99%	-0.78[-0.88,-0.68]
Jenkins 1993	49	1.2 (0.8)	50	1.8 (0.5)	-+-	8.79%	-0.65[-0.91,-0.39]
Navarro 1998	9	1.7 (0.8)	11	2.3 (0.4)		1.77%	-0.58[-1.15,-0.02]
Subtotal ***	122		105		•	79.49%	-0.76[-0.85,-0.68]
Heterogeneity: Tau ² =0; Chi ² =1.28	, df=3(P=0.7	3); I ² =0%					
Test for overall effect: Z=17.58(P<	(0.0001)						
Total ***	336		210		•	100%	-0.78[-0.85,-0.7]
Heterogeneity: Tau ² =0; Chi ² =1.79	, df=4(P=0.7	7); I ² =0%					
Test for overall effect: Z=20.08(P<	(0.0001)						
Test for subgroup differences: Ch	i ² =0.51, df=1	. (P=0.47), I ² =0%					
				Favours CHX	-2 -1 0 1	2 Favours cor	ntrol

Analysis 1.8. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 8 Plaque 6 months.

Study or subgroup	CHX Control Std. Mean Difference		Std. Mean Difference	Weight	Std. Mean Difference		
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
1.8.1 CHX versus no rinse							
Chaves 1994	24	0.8 (0.3)	27	1 (0.8)		8.6%	-0.32[-0.87,0.23]
Flemmig 1990	43	0.4 (0.3)	48	0.8 (0.3)		9.45%	-1[-1.44,-0.57]
Subtotal ***	67		75		•	18.05%	-0.68[-1.35,-0.01]
Heterogeneity: Tau ² =0.17; Chi ² =3.62	, df=1(P=	0.06); I ² =72.34%					
Test for overall effect: Z=2(P=0.05)							
1.8.2 CHX versus placebo/control r	inse						
Bajaj 2011	440	0.7 (0.3)	412	2 (0.8)	+	11%	-2.1[-2.27,-1.93]
Charles 2004	36	1.7 (0.5)	37	2.2 (0.4)	→	9.07%	-1.03[-1.52,-0.54]
Flotra 1972	22	0.2 (0.2)	6	0.9 (0.4)	←	4.14%	-3.3[-4.59,-2]
Grossman 1986	188	0.6 (0.5)	189	1.6 (0.5)	+	10.67%	-1.9[-2.14,-1.65]
				Favours CHX	-2 -1 0 1 2	Favours co	ontrol



Study or subgroup		СНХ	C	ontrol	Std. Mean Difference	Weight	Std. Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Grossman 1989	113	0.8 (0.5)	133	1.5 (0.5)	-+-	10.47%	-1.43[-1.71,-1.15]
Hase 1998	30	1 (0.7)	33	1.7 (0.6)		8.77%	-1.07[-1.6,-0.54]
Lang 1998	41	0.3 (0.3)	45	0.8 (0.4)		9.19%	-1.4[-1.88,-0.93]
Overholser 1990	41	0.8 (0.5)	42	1.6 (0.5)	+-	9.01%	-1.61[-2.11,-1.11]
Stookey 2005	40	1.4 (0.5)	85	2 (0.5)		9.64%	-1.33[-1.74,-0.92]
Subtotal ***	951		982		•	81.95%	-1.59[-1.89,-1.29]
Heterogeneity: Tau ² =0.16; Cl	hi²=49.48, df=8(P	<0.0001); I ² =83.8	3%				
Test for overall effect: Z=10.2	24(P<0.0001)						
Total ***	1018		1057		•	100%	-1.43[-1.76,-1.1]
Heterogeneity: Tau ² =0.25; Ch	hi ² =84.73, df=10(P<0.0001); I ² =88.	2%				
Test for overall effect: Z=8.47	7(P<0.0001)						
Test for subgroup difference	s: Chi ² =5.85, df=1	L (P=0.02), I ² =82.	89%				
				Favours CHX	-2 -1 0 1 2	Favours co	ontrol

Analysis 1.9. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 9 Plaque 6 months PI (0-3).

	СНХ	(Control	Mean Difference	Weight	Mean Difference	
N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI	
43	0.4 (0.3)	48	0.8 (0.3)	*	20.54%	-0.32[-0.45,-0.19]	
24	0.8 (0.3)	27	1 (0.8)	-	19.14%	-0.2[-0.52,0.12]	
67		75		♦	39.68%	-0.3[-0.42,-0.18]	
df=1(P=0.5); I ² =0%						
0001)							
l rinse							
440	0.7 (0.3)	412	2 (0.8)	•	20.71%	-1.29[-1.37,-1.21]	
22	0.2 (0.2)	6	0.9 (0.4)	-+-	19.11%	-0.79[-1.12,-0.46]	
41	0.3 (0.3)	45	0.8 (0.4)	*	20.49%	-0.48[-0.62,-0.34]	
503		463		•	60.32%	-0.86[-1.46,-0.25]	
.2, df=2(P<	0.0001); I ² =97.92	2%					
01)							
570		538		•	100%	-0.62[-1.12,-0.12]	
9.21, df=4(P<0.0001); I ² =98.	.09%					
01)				į			
=3.06, df=1	1 (P=0.08), I ² =67.	34%					
֡֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜֜	43 24 67 df=1(P=0.5 0001) I rinse 440 22 41 503 .2, df=2(P<01) 570 9.21, df=4(01)	N Mean(SD) 43 0.4 (0.3) 24 0.8 (0.3) 67 df=1(P=0.5); l²=0% 0001) I rinse 440 0.7 (0.3) 22 0.2 (0.2) 41 0.3 (0.3) 503 .2, df=2(P<0.0001); l²=97.92 01) 570 9.21, df=4(P<0.0001); l²=98	N Mean(SD) N 43 0.4 (0.3) 48 24 0.8 (0.3) 27 67 75 df=1(P=0.5); I²=0% 0001) I rinse 440 0.7 (0.3) 412 22 0.2 (0.2) 6 41 0.3 (0.3) 45 503 463 .2, df=2(P<0.0001); I²=97.92% 01) 570 538 9.21, df=4(P<0.0001); I²=98.09%	N Mean(SD) N Mean(SD) 43 0.4 (0.3) 48 0.8 (0.3) 24 0.8 (0.3) 27 1 (0.8) 67 75 df=1(P=0.5); I²=0% 0001) I rinse 440 0.7 (0.3) 412 2 (0.8) 22 0.2 (0.2) 6 0.9 (0.4) 41 0.3 (0.3) 45 0.8 (0.4) 503 463 .2, df=2(P<0.0001); I²=97.92% 01) 570 538 9.21, df=4(P<0.0001); I²=98.09% 01)	N Mean(SD) N Mean(SD) 43 0.4 (0.3) 48 0.8 (0.3) 24 0.8 (0.3) 27 1 (0.8) 67 75 df=1(P=0.5); l²=0% 0001) 1 rinse 440 0.7 (0.3) 412 2 (0.8) 22 0.2 (0.2) 6 0.9 (0.4) 41 0.3 (0.3) 45 0.8 (0.4) 503 463 .2, df=2(P<0.0001); l²=97.92% 01) 570 538 9.21, df=4(P<0.0001); l²=98.09% 01)	N Mean(SD) N Mean(SD) Random, 95% CI 43	

Analysis 1.10. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 10 Plaque 6 months TQH (0-5).

Study or subgroup	СНХ			Control	Mean Difference		Weight	Mean Difference			
	N	Mean(SD)	N	Mean(SD)		Rand	dom, 95	% CI			Random, 95% CI
1.10.1 CHX versus placebo/contro	l rinse										
				Favours CHX	-5	-2.5	0	2.5	5	Favours contr	ol



Study or subgroup		СНХ	C	ontrol	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
Grossman 1986	188	0.6 (0.5)	189	1.6 (0.5)	•	20.56%	-0.95[-1.05,-0.85]
Overholser 1990	41	0.8 (0.5)	42	1.6 (0.5)	+	15.42%	-0.82[-1.04,-0.61]
Grossman 1989	113	0.8 (0.5)	133	1.5 (0.5)	*	19.5%	-0.73[-0.86,-0.6]
Hase 1998	30	1 (0.7)	33	1.7 (0.6)	+	11.29%	-0.7[-1.02,-0.38]
Stookey 2005	40	1.4 (0.5)	85	2 (0.5)	*	17.38%	-0.62[-0.8,-0.44]
Charles 2004	36	1.7 (0.5)	37	2.2 (0.4)	+	15.86%	-0.47[-0.68,-0.26]
Subtotal ***	448		519		•	100%	-0.73[-0.88,-0.57]
Heterogeneity: Tau ² =0.03; Ch	ni²=23.59, df=5(P	=0); I ² =78.81%					
Test for overall effect: Z=9.39	(P<0.0001)						
Total ***	448		519		•	100%	-0.73[-0.88,-0.57]
Heterogeneity: Tau ² =0.03; Ch	ni²=23.59, df=5(P	=0); I ² =78.81%					
Test for overall effect: Z=9.39	(P<0.0001)						
				Favours CHX -5	-2.5 0 2.5	5 Favours con	rol

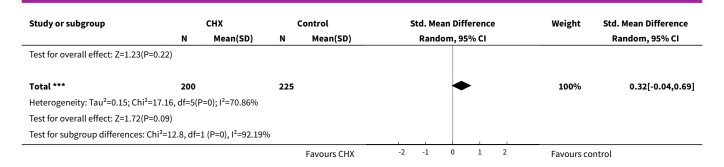
Analysis 1.11. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 11 Calculus 4-6 weeks.

Study or subgroup		СНХ	(Control		Mean Difference Weight		Weight	Mean Difference		
	N	Mean(SD)	N	Mean(SD)		Rai	ndom, 95% (:1			Random, 95% CI
1.11.1 CHX versus placebo/c	ontrol rinse										
Emling 1992	26	0.5 (0.3)	26	0.5 (0.5)			•			28.62%	0.01[-0.21,0.23]
Turkoglu 2009	25	0.3 (0.2)	25	0.3 (0.3)						71.38%	0.03[-0.11,0.17]
Subtotal ***	51		51							100%	0.02[-0.09,0.14]
Heterogeneity: Tau ² =0; Chi ² =0	.02, df=1(P=0.9)); I ² =0%									
Test for overall effect: Z=0.37(I	P=0.71)										
				Favours CHX	-100	-50	0	50	100	Favours contro	 [

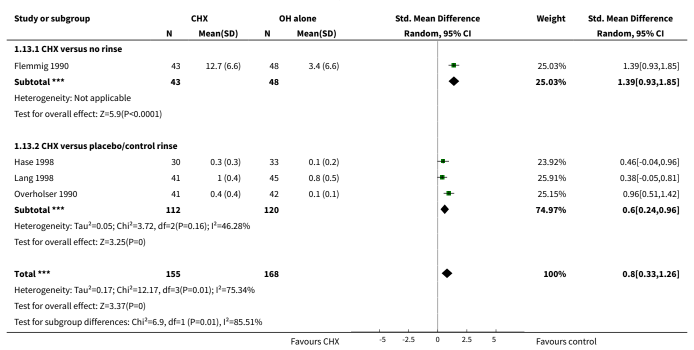
Analysis 1.12. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 12 Calculus 7-12 weeks.

Study or subgroup		СНХ	c	ontrol	Std. Mean Difference	Weight	Std. Mean Difference
	N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
1.12.1 CHX versus no rinse							
Flemmig 1990	44	8 (4.9)	51	3 (4.9)		18.01%	1.02[0.59,1.45]
Subtotal ***	44		51		•	18.01%	1.02[0.59,1.45]
Heterogeneity: Not applicable							
Test for overall effect: Z=4.67(P<0.0	0001)						
1.12.2 CHX versus placebo/contr	ol rinse						
Corbet 1997	13	2.7 (0.5)	23	2.4 (0.9)	+	13.03%	0.35[-0.33,1.04]
Emling 1992	26	0.5 (0.3)	26	0.4 (0.5)		15.62%	0.28[-0.27,0.82]
Hase 1998	35	0.2 (0.3)	34	0.1 (0.2)	-	17.11%	0.18[-0.29,0.66]
Lang 1998	41	0.8 (0.4)	49	0.9 (0.6)	-+ 	18.29%	-0.22[-0.64,0.2]
Overholser 1990	41	0.2 (0.2)	42	0.1 (0.1)	+-	17.94%	0.32[-0.12,0.75]
Subtotal ***	156		174		•	81.99%	0.14[-0.08,0.36]
Heterogeneity: Tau ² =0; Chi ² =4.18,	df=4(P=0.38	3); I ² =4.24%					
				Favours CHX	-2 -1 0 1 2	Favours co	ontrol





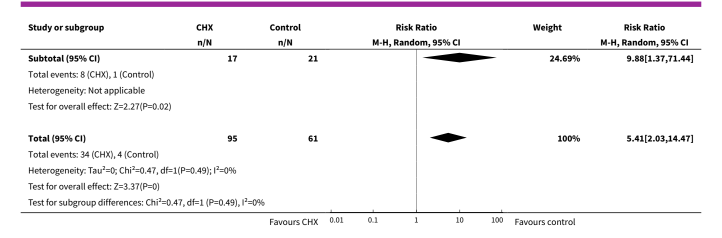
Analysis 1.13. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 13 Calculus 6 months.



Analysis 1.14. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 14 Tooth staining 4-6 weeks dichotomous.

Study or subgroup	СНХ	Control	Risk Ratio			Weight	Risk Ratio		
	n/N	n/N	M-H, Random, 95% CI					M-H, Random, 95% CI	
1.14.1 CHX versus no rinse									
Zimmer 2015	26/78	3/40					75.31%	4.44[1.43,13.8]	
Subtotal (95% CI)	78	40					75.31%	4.44[1.43,13.8]	
Total events: 26 (CHX), 3 (Control)									
Heterogeneity: Not applicable									
Test for overall effect: Z=2.58(P=0.01)									
1.14.2 CHX versus placebo/control rins	e								
Sanz 1989	8/17	1/21			-	— .	24.69%	9.88[1.37,71.44]	
		Favours CHX	0.01	0.1	10	100	Favours control		





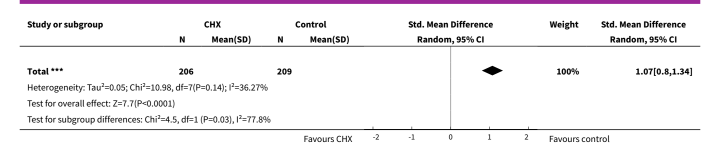
Analysis 1.15. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 15 Tooth staining 7-12 weeks dichotomous.

Study or subgroup	снх	Control			Risk Ratio			Weight	Risk Ratio
	n/N	n/N M-H, Random, 95% CI			M-H, Random, 95% CI				
1.15.1 CHX versus no rinse									
Zimmer 2015	39/78	8/40				-		100%	2.5[1.29,4.83]
Subtotal (95% CI)	78	40			•	•		100%	2.5[1.29,4.83]
Total events: 39 (CHX), 8 (Control)									
Heterogeneity: Not applicable									
Test for overall effect: Z=2.73(P=0.01)									
		Favours CHX	0.01	0.1	1	10	100	Favours control	

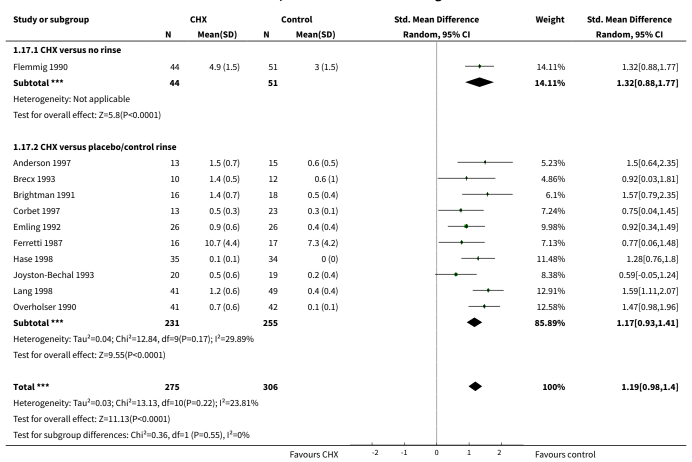
Analysis 1.16. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 16 Tooth staining 4-6 weeks.

	СНХ	,	Control	Std. Mean Difference	Weight	Std. Mean Difference
N	Mean(SD)	N	Mean(SD)	Random, 95% CI		Random, 95% CI
47	1 (0.5)	47	0.4 (0.3)		17.65%	1.54[1.08,2]
47		47		•	17.65%	1.54[1.08,2]
1)						
rinse						
10	1.1 (0.4)	12	0.7 (0.3)		7.14%	1.1[0.19,2.02]
16	0.9 (0.6)	18	0.5 (0.4)		10.64%	0.78[0.07,1.48]
26	0.9 (0.6)	26	0.5 (0.5)		14.32%	0.64[0.08,1.2]
33	0.7 (0.4)	28	0.3 (0.3)		14.49%	1.25[0.7,1.8]
16	4.2 (2.9)	17	0.9 (0.7)		8.94%	1.55[0.76,2.34]
49	0.1 (0.1)	50	0 (0)		19.39%	1.02[0.6,1.44]
9	0.1 (0.4)	11	0 (0.1)		7.43%	0.39[-0.5,1.28]
159		162		•	82.35%	0.97[0.73,1.22]
df=6(P=	0.38); I ² =6.42%					
.)						
	47 47 47 10 11) 16 26 33 16 49 9	47 1 (0.5) 47 10 1.1 (0.4) 16 0.9 (0.6) 26 0.9 (0.6) 33 0.7 (0.4) 16 4.2 (2.9) 49 0.1 (0.1) 9 0.1 (0.4) 159 df=6(P=0.38); l²=6.42%	47 1 (0.5) 47 47 47 10) rinse 10 1.1 (0.4) 12 16 0.9 (0.6) 18 26 0.9 (0.6) 26 33 0.7 (0.4) 28 16 4.2 (2.9) 17 49 0.1 (0.1) 50 9 0.1 (0.4) 11 159 162 df=6(P=0.38); l²=6.42%	47 1 (0.5) 47 0.4 (0.3) 47 47 47 10 1.1 (0.4) 12 0.7 (0.3) 16 0.9 (0.6) 18 0.5 (0.4) 26 0.9 (0.6) 26 0.5 (0.5) 33 0.7 (0.4) 28 0.3 (0.3) 16 4.2 (2.9) 17 0.9 (0.7) 49 0.1 (0.1) 50 0 (0) 9 0.1 (0.4) 11 0 (0.1) 159 162 df=6(P=0.38); l²=6.42%	47 1 (0.5) 47 0.4 (0.3) 47 47 48 10 1.1 (0.4) 12 0.7 (0.3) 16 0.9 (0.6) 18 0.5 (0.4) 26 0.9 (0.6) 26 0.5 (0.5) 33 0.7 (0.4) 28 0.3 (0.3) 16 4.2 (2.9) 17 0.9 (0.7) 49 0.1 (0.1) 50 0 (0) 9 0.1 (0.4) 11 0 (0.1) 159 162 df=6(P=0.38); l²=6.42%	47 1 (0.5) 47 0.4 (0.3) 47 47 47 17.65% 17.65% 10 1.1 (0.4) 12 0.7 (0.3) 16 0.9 (0.6) 18 0.5 (0.4) 26 0.9 (0.6) 26 0.5 (0.5) 33 0.7 (0.4) 28 0.3 (0.3) 16 4.2 (2.9) 17 0.9 (0.7) 49 0.1 (0.1) 50 0 (0) 9 0.1 (0.4) 11 0 (0.1) 159 162 47 17.65% 17.65% 17.65% 17.65% 17.65% 17.65% 17.65% 17.65% 17.65% 17.43% 10.44% 10.64% 1





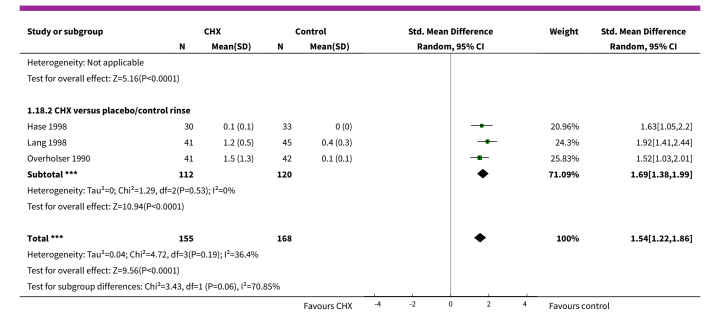
Analysis 1.17. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 17 Tooth staining 7-12 weeks.



Analysis 1.18. Comparison 1 CHX versus placebo/control mouthrinse or no mouthrinse, Outcome 18 Tooth staining 6 months.

Study or subgroup	СНХ		(Control		Std. Mean Difference We			Weight	Std. Mean Difference	
	N	Mean(SD)	N	Mean(SD)		Ran	dom, 95%	6 CI			Random, 95% CI
1.18.1 CHX versus no rinse											
Flemmig 1990	43	5.3 (1.9)	48	3 (1.9)			-	-		28.91%	1.18[0.73,1.62]
Subtotal ***	43		48				- •	•		28.91%	1.18[0.73,1.62]
				Favours CHX	-4	-2	0	2	4	Favours conti	rol





ADDITIONAL TABLES

Table 1. Random-effects metaregression analyses of Gingival Index (GI) at 4 to 6 weeks

Characteristic	Number of studies	Slope es- timate	95% CI	Slope interpretation	P value
Adults versus children	10 - no stud- ies with just children				
Gingivitis alone versus gingivitis with perio	8	0.12	-0.14 to 0.38	Increase in GI effect estimate for gingivitis and perio	0.30
Prophylaxis or not	9	0.05	-0.22 to 0.32	Increase in GI effect estimate for prophylaxis	0.66
Baseline gingivitis < 1 versus ≥ 1	9	0.02	-0.25 to 0.30	Increase in GI effect estimate for higher baseline score	0.84

CI = confidence interval.

Table 2. Random-effects metaregression analyses of Gingival Index (GI) at 6 months

Characteristic	Number of studies	Slope es- timate	95% CI	Slope interpretation	P value
Adults versus children	13	-0.17	-0.42 to 0.09	Increase in GI effect estimate for adults	0.185
Gingivitis alone versus gingivitis with perio	9	0.15	-0.14 to 0.44	Increase in GI effect estimate for gingivitis and perio	0.25
Prophylaxis or not	11	-0.13	-0.25 to -0.004	Increase in GI effect estimate for no prophylaxis	0.045



Table 2. Random-effects metaregression analyses of Gingival Index (GI) at 6 months (Continued)

Baseline gingivitis 9 -0.05 -0.39 to 0.30 Decrease in GI effect estimate for high- 0.75 er baseline score

< 1 versus ≥ 1

CI = confidence interval.

Table 3. Chlorhexidine concentration: all trials

Outcome (index)	Time	Chlorhex- idine con- centra- tion	Studies (partici- pants)	MD/SMD 95% CI	Effect P value	Subgroup P value	
Gingival inflamma-	4 to 6 weeks	0.2	6 (552)	MD -0.27 (-0.46 to -0.09)	0.003 favours CHX	0.41	
tion (Gingi- val Index)	weeks	0.1 and 0.12	5 (253)	MD -0.19 (-0.27 to -0.10)	< 0.0001 favours CHX	_	
Gingival inflamma-	6 months	0.2	1 (86)	MD -0.12 (-0.20 to -0.04)	0.005	Too few	
tion (Gingi- val Index)		0.1 and 0.12	10 (2352)	MD -0.22 (-0.33 to -0.11)	< 0.00001 favours CHX	studies in subgroup	
		0.05	1 (150)	MD -0.04 (-0.11 to 0.03)	0.28	_	
Gingival bleeding	4 to 6 weeks	0.2	4 (472)	SMD -0.71 (-0.90 to -0.51)	< 0.00001 favours CHX	0.18	
bleeding w	Weeks	0.1 and 0.12	3 (127)	SMD -0.32 (-0.85 to 0.21)	0.23	_	
Gingival bleeding	6 months	0.2	2 (155)	SMD -1.20 (-2.48 to 0.08)	0.07	0.34	
		0.12	6 (977)	SMD -0.57 (-0.79 to -0.36)	< 0.00001 favours CHX	_	
Plaque	4 to 6 weeks	0.2	8 (685)	SMD -1.75 (-2.45 to -1.04)	< 0.00001 favours CHX	0.04	
		0.1 and 0.12	4 (215)	SMD -0.95 (-1.23 to -0.66)	< 0.00001 favours CHX	_	
Plaque	6 months	0.2	2 (149)	SMD -1.26 (-1.61 to -0.90)	< 0.00001 favours Chx	0.65	
		0.1 and 0.12	8 (1898)	SMD -1.38 (-1.75 to -1.00)	< 0.00001 favours CHX	_	
Calculus	4 to 6 weeks	0.12	1 (52)	MD 0.01 (-0.21 to 0.23)	0.93		
Calculus	7 to 12	0.2	2 (159)	SMD -0.03 (-0.43 to 0.36)	0.86	0.05	
	weeks	0.12	4 (266)	SMD 0.52 (0.13 to 0.91)	0.10	_	
Calculus	6 months	0.2	2 (149)	SMD 0.41 (0.09 to 0.74)	0.01	0.005	
		0.12	2 (174)	SMD 1.17 (0.76 to 1.59)	< 0.00001 favours control	_	

0.08

< 0.00001 favours control

< 0.00001 favours control



Tooth

staining

Table 3. Chlorhexidine concentration: all trials (Continued)							
Tooth 4 to 6 staining weeks		0.2 2 (116)		SMD 1.45 (1.04 to 1.87)	< 0.00001 favours control	0.05	
J		0.1 and 0.12	6 (299)	SMD 0.96 (0.68 to 1.24)	< 0.00001 favours control		
Tooth staining	7 to 12 weeks	0.2	3 (181)	SMD 1.38 (1.05 to 1.71)	< 0.00001 favours control	0.10	
		0.12	7 (361)	SMD 1.2 (0.96 to 1.45)	< 0.00001 favours control		

SMD 0.59 (-0.05 to 1.24)

SMD 1.79 (1.41 to 2.17)

SMD 1.33 (1.00 to 1.66)

CHX = chlorhexidine; CI = confidence interval; MD = mean difference; SMD = standardised mean difference. Studies where the concentration of the chlorhexidine mouthrinse was not reported (Turkoglu 2009) and where data relate to a combination of different chlorhexidine concentrations (Flotra 1972) were excluded from this analysis. Therefore, in certain analyses, the number of studies and participants presented in the table are different from the data presented in the main analysis.

Table 4. Chlorhexidine rinse frequency of use: all trials

0.05

0.2

0.12

6 months

1 (39)

2 (149)

2 (174)

Outcome (index)	Time	Chlorhexidine fre- quency	Studies (partici- pants)	MD/SMD 95% CI	Effect P value	Subgroup P value
Gingival in- flammation	4 to 6 weeks	Twice per day	9 (785)	MD -0.22 (-0.33 to -0.11)	< 0.0001	Too few — studies in
(Gingival In- dex)	weeks	Once per day	1 (20)	MD -0.13 (-0.31 to 0.06)	0.18	subgroup
Gingival in- flammation	6 months	Twice per day	11 (1614)	MD -0.17 (-0.20 to -0.13)	< 0.0001	0.56
(Gingival In- dex)		Once per day	2 (1002)	MD -0.34 (-0.93 to 0.25)	0.26	_
Gingival bleeding	4 to 6 weeks	Twice per day	8 (649)	SMD -0.56 (-0.79 to -0.33)	< 0.0001	-
Gingival bleeding	6 months	Twice per day	8 (1132)	SMD -0.72 (-1.02 to -0.42)	< 0.0001	-
Plaque	4 to 6 weeks	Twice per day	11 (930)	SMD -1.49 (-1.97 to -1.02)	< 0.0001	Too few – studies in
	WEEK2	Once per day	1 (20)	SMD -0.92 (-1.86 to 0.02)	0.05	subgroup
Plaque	6 months	Twice per day	10 (1223)	SMD -1.34 (-1.66 to -1.03)	< 0.0001	Too few – studies in
		Once a day	1 (852)	SMD -2.10 (-2.27 to -1.93)	< 0.0001	subgroup
Calculus	4 to 6 weeks	Twice per day	1 (50)	MD 0.03 (-0.11 to 0.17)	0.70	-
Calculus	7 to 12 weeks	Twice per day	5 (373)	SMD 0.33 (-0.11 to 0.77)	0.14	-



 Table 4. Chlorhexidine rinse frequency of use: all trials (Continued)

Calculus	6 months	Twice per day	4 (323)	SMD 0.80 (0.33 to 1.26)	0.0007	-
Tooth stain- 4 to 6 ing weeks		3 times per day	1 (33)	SMD 1.55 (0.76 to 2.34)	< 0.0001	Too few - studies in
5	mg weeks	Twice per day	5 (310)	SMD 1.18 (0.93 to 1.44)		subgroup
		Once per day	1(20)	SMD 0.39 (-0.5 to 1.28)	0.39	
Tooth stain-	7 to 12 weeks	3 times per day	1 (33)	SMD 0.77 (0.06 to 1.48)	0.03	Too few - studies in
5	Weeks	Twice per day	9 (496)	SMD 1.26 (1.04 to 1.49)	< 0.0001	subgroup
Tooth stain- ing	6 months	Twice per day	4 (323)	SMD 1.54 (1.22 to 1.86)	< 0.0001	-

CI = confidence interval; MD = mean difference; SMD = standardised mean difference.

Table 5. Results for gingivitis and plaque at 7 to 12 weeks, > 6 months and long term (after cessation of mouthrinsing)

Outcome (index)	Time	Chlorhex- idine conc (%)	Studies (partici- pants)	MD/SMD 95% CI	Effect P value	Heterogeneity
Results for studies	with no rinse	control arms	5			
Gingival bleeding	7 to 12 weeks	< 0.1	2 (196)	MD -0.07 (-0.16 to 0.02)	0.13 favours CHX	P = 0.19, I ² = 40%
Plaque	7 to 12 weeks	< 0.1	2 (196)	SMD -0.77 (-1.07 to -0.47)	< 0.00001 favours CHX	P = 0.61, I ² = 0%
Results for studies	with placebo	/control rins	e arms			
Gingival inflam- mation (Gingival Index)	7 to 12 weeks	0.2 and 0.12	4 (144)	MD -0.47 (-0.76 to -0.18)	0.001 favours CHX	P < 0.0001, I ² = 86%
Gingival inflam- mation (Gingival Index)	> 6 months	0.1 and 0.12	2 (1124)	MD -0.50 (-1.11 to 0.11)	0.11 favours CHX	P < 0.0001, I ² = 99%
Gingival bleeding	7 to 12 weeks	0.12 and < 0.1	5 (182)	SMD -1.29 (-1.85 to -0.72)	< 0.00001 favours CHX	P = 0.02, I ² = 64%
Gingival bleeding	Long term	0.12	3 (99)	MD -0.12 (-0.2 to -0.04)	0.003 favours CHX	P = 0.33, I ² = 11%
Plaque	7 to 12 weeks	0.2, 0.12 and < 0.1	10 (423)	SMD -1.74 (-2.51 to -0.98)	< 0.00001 favours CHX	P < 0.00001, I ² = 91%
Plaque	> 6 months	0.1	1 (852)	MD -1.55 (-1.79 to -1.31)	< 0.00001 favours CHX	N/A
Plaque	Long term	0.12	4 (132)	SMD -1.10 (-1.18 to -0.40)	< 0.002 favours CHX	P = 0.02, I ² = 71%



CHX = chlorhexidine; CI = confidence interval; conc = concentration; MD = mean difference; N/A = not applicable; SMD = standardised mean difference

There were no subgroup differences between the different chlorhexidine concentrations, therefore the overall effect for all concentrations combined is reported.

Table 6. Studies with a gingival inflammation outcome not included in meta-analyses

Time	Study ID	Compari- son	Index	CHX mean (SD)	Control mean (SD)	Total n	Notes
4 to 6 weeks	Anau- ate-Netto 2014	CHX 0.12% vs placebo	PBS	0.9 (0.6)	0.7 (0.4)	40	Did not report GI. Quote: "no statistically significant differences were detected among groups"
7 to 12 Anderson weeks 1997	Anderson 1997	CHX 0.12% vs placebo	GI	0.345	0.895	28	Reported mean GI by surface + SD. No overall SD. We calculated overall mean. Quote: "The means of thegingival indices did not show any significant differences (P<0.05) 1 or 2 months after baseline. However there were significant differences (P<0.05) in the changes recorded at 30 and 60 days at all sites inthe experimental group"
	de la Rosa 1888b	CHX 0.12% vs placebo	PMGI sever- ity (mean score of all sites grad- ed)	0.1413	0.2902	92	Did not report GI or a SD. Quote: "the effect of the chlorhexidine rinse on the occurrence and severity of gingivitis amounted to a 51% reduction of the disease compared to the placebo rinsedifferences were statistically significant"
	de la Rosa 1988a	CHX 0.12% vs placebo	PMGI sever- ity (mean score of all sites grad- ed)	0.2892	0.4526	99	Did not report GI or a SD. Quote: "the gingivitis reductions were 34% and 36% for occurrence and severity respectivelydifferences were statistically significant"
	Eaton 1997	CHX 0.12% vs placebo	mGI	0.42 (0.383)	0.55 (0.382)	98	Did not report GI. Quote: "the pooled mean mGI score improved by 25% from 0.56 at baseline to 0.42 at 12 weeks in the ChD (CHX) group but showed no change (0.54 to 0.55) in the placebo group"
	Ferretti 1987	CHX 0.12% vs placebo	PMGI	0.8	1.94	33	Did not report GI and SD depicted in graph but not reported. Quote: "Significant reductions ingingivitis scores were seen on days 33 (P<0.0001) and 60 (P<0.001) for those patients using chlorhexidine rinse"
	Segreto 1986	0.2% vs 0.12% vs placebo	GI	0.4112/0.3640	0.5039	454	Did not report a SD. Quote: "Gingivitis severity by the GI method was significantly lower at 3 months for both chlorhexidine groups compared to the placebo group. Differences ranged from 28-46% and averaged 37% for the 0.12% group. Differences ranged from 18-40% for the 0.20% group and averaged 29%"

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	Weitz 1992	CHX 0.12% vs placebo	GI	1.69	1.86	36	Did not report a SD. Quote: "the active (CHX) groups had significantly lower gingivitis scores than the respective control groups. Overall, the active group had a 10.27% reduction in the gingival indexcompared to insignificant changes in the control groups"
6 months	Fine 1985	CHX+OH vs OH	Not clear. Quote: "a gingival in- flammation Index"	NR	NR	83	Outcomes reported in graphs which are difficult to decipher. Quote: "Whilst there was a general improvement in all groups of patients, no one group was statistically significantly different from the other"
	Hoffmann 2001	CHX 0.1%, CHX 0.06%, CHX 0.06% +F vs con- trol	GI	Median 0.15/0.29/0.34	Median 0.45	58	No SD. At 3 months "only the 0.1% CHX was different from the control". At 6 months "the 0.1% CHX showed significant differencesin the GIwhen compared to the 0.06% CHX/F" (P = 0.043)
	Overholser 1990	CHX 0.12% vs control	mGl	0.81 (SE 0.065)	1.166 (SE 0.063)	83	Did not report a SD. Quote: "PX (CHX) inhibited gingivitis development by 26.8% (P<0.001) at 3 months and by 30.5% (P<0.001) at 6 months, compared to the control"

CHX = chlorhexidine; F = fluoride; NR = not reported; OH = oral hygiene; SD = standard deviation; SE = standard error.

Total n is the number of participants analysed in the study arms relevant to the review.

PBS (Papillary Bleeding Score, Loesche 1979) is measured on a 0-5 increasing scale.

GI (Gingival Index, Löe 1967; Löe and Silness 1963) is measured on a 0-3 increasing scale.

PMGI (Papillary Marginal Gingivitis Index, de la Rosa and Sturzenberger 1976) is measured on a 0-3 increasing scale.

mGI (modified Gingival Index, Lobene 1986) is measured on a 0-4 increasing scale.

Table 7. Studies with a gingival bleeding outcome not included in meta-analyses

Time	Study ID	Compari- son	Index	CHX mean (SD)	Control mean (SD)	Total n	Notes
4 to 6 weeks	Axelsson 1987	CHX 0.2% vs CHX 0.1% vs placebo	Mean % of gingival units with GI score 2 or 3	12%/11%	25%	64	Did not report a SD. Quote: "In allstudy groups, the % of gingival units scored GI 2+3 was reduced between baselineand end of trial"
	Graziani 2015	CHX 0.2% +Alc vs CHX 0.2% no Alc vs CHX+ADS	FMBS/BOP	Not clear	13.47%	70	Partial reporting of outcome data. Quote: "Statistically significant decreases in mean full-mouth scores of gingival inflammation were noted for all experimental study groups at day 35 compared to baseline. Between-group changes in

	a g	vs control rinse	5 • • • • • • • • • • • • • • • • • • •	utcome not included in meta-analyses (Continued)			FMBSwere statistically significant only when CHX2 (mean difference of 43.4 ± 22.4 , P = 0.05) and CHX 3 (mean difference of 46.1 ± 23.1 , P = 0.05) were compared to the CTRL group" (note: CHX 2 = CHX no Alc, CHX 3 = CHX + ADS)
	Sanz 1989	CHX 0.12% vs placebo	Mean % sites GI score 2 and 3	18.78%	31.31%	38	Did not report a SD. Quote: "Beginning at 4 weeks of rinsing gingival bleeding was significantly lower in the CHX group compared to the placebo group by an average of 41.6% (P < 0.05). At 6 weeks that reduction was 40%, (P < 0.05)"
7 to 12 weeks	Corbet 1997	CHX 0.12% vs placebo	Mean % sites GI score 2 and 3	27%(anterior) 42%(posterior)	52%(anterior) 75%(posterior)	36	Did not report a SD. Quote: "The difference between the mean percentage of GB of the test and control groups at 3 months was highly significant (P < 0.001)"
	Segreto 1986	CHX 0.2% vs CHX 0.12% vs placebo	Mean % sites GI score 2 and 3	Examiner A: 3.4%/2% Examiner B: 7.2%/6.2%	Examiner A: 3.81% Examiner B: 14.9%	454	Did not report a SD. O.2% Quote: "Bleeding was 31% lower (range 11-52%) compared to the placebo group" (P > 0.05) O.12% Quote: "gingival bleeding was significantly lower by an average of 53% for both examiners (range 48-59%)" (P ≤ 0.05)
6 months	Banting 1989	CHX 0.12% vs placebo	Mean % sites GI score 2 and 3	2.41%	4.12%	383	Did not report a SD. Regarding outcomes at 6 months to 2 years: Quote: "Subjects in the treatment groupdisplayed between 42% and 51% fewer sites with moderate to severe gingivitis (GI scores of 2 or 3) compared with subjects in the control group" (P < 0.0001)
	Charles 2004	CHX 0.12% vs control	Mean % sites GI score 2 and 3	11.01%	20.65%	1156 sites	% of bleeding sites in each group is presented. There is a ur of analysis error (number of sites rather than number of subjects). Quote: "There was a considerable reduction in perce bleeding sites in the chlorhexidinegroups at 6 months copared with both control and baseline"

4.2%

Graph

20

130

Did not report a SD. Quote: "Although the percentage of bleeding surfaces in the chlorhexidine group was less than in

ences were not significant" (P = 0.07)

graph. Did not report a SD.

the placebo group on days 90 (33%) and 180 (76%), the differ-

% bleeding sites reported incompletely in text and also in a

Lucas 1999

Sanz 1994

CHX 0.12%

vs placebo

CHX 0.12%

vs placebo

Mean %

sites GI

3

3

score 2 and

% sites GI

score 2 and

1%

Graph

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ADS = antidiscolouration system; Alc = alcohol; BOP = bleeding on probing; CHX = chlorhexidine; FMBS = Full-Mouth Bleeding Score; GI = Gingival Index; SD = standard deviation. Total n is the number of participants analysed in the study arms relevant to the review.

Table 8. Studies with a plaque outcome not included in meta-analyses

Time	Study ID	Comparison	Index	CHX mean (SD)	Control mean (SD)	Total n	Notes
4 to 6 weeks	Sanz 1989	CHX 0.12% vs placebo	PI	0.452	0.9907	38	Did not report a SD
7 to 12 weeks	Anderson 1997	CHX 0.12% vs placebo	PI	0.3175*	0.8425*	28	Mean GI by surface + SD reported. We calculated the overall mean. No overall SD
	de la Rosa 1888b	CHX 0.12% vs placebo	TQH	NR	NR	92	Quote: "The reductions in dental plaque were not statistically significant"
	de la Rosa 1988a	CHX 0.12% vs placebo	TQH	NR	NR	99	Quote: "The reductions in dental plaque were not statistically significant"
	Segreto 1986	0.2% vs 0.12% vs placebo	TQH	1.14/1.01	1.58	451	Did not report a SD
	Weitz 1992	CHX 0.12% vs placebo	PI	1.84	2.21	36	Did not report a SD
6 months	Banting 1989	CHX 0.12% vs placebo	ТQН	Graph	Graph	383	Data presented in a graph. Did not report a SD. Quote: "Subjects in the treatment group had significantly lower mean plaque scores than those in the control group at six months, and at one and two years. The difference between the groups ranged from 35% to 46%"

Table 8. Studies with a plaque outcome not included in meta-analyses (Continued)

	Hoffmann 2001	CHX 0.1%/CHX 0.06%/ CHX 0.06%+F vs con- trol rinse	PI	Median 0.13/0.25/0.27	Median 0.72	58	Median only. Did not report a SD
•	Jayaprakash 2007	CHX 0.05%/ CHX 0.05% +F vs placebo	PI	0.0813/0.0459	0.1189	100	Did not report a SD
•	Lucas 1999	CHX 0.12% vs placebo	OHI-S	0.33	0.59	20	Did not report a SD
	Sanz 1994	CHX 0.12% vs placebo	PI	Graph	Graph	130	Data presented in a graph. Did not report a SD. Quote: "reductions in Plaque Indexwere statistically significant for the positive control group compared with the reduction obtained with the control group These reductions were 41% (positive control)after 3 months and 35%after 6 months"
Plaque > 6 months	Banting 1989	CHX 0.12% vs placebo	TQH	Graph	Graph	272	Data presented in a graph. Did not report a SD. Quote: "Subjects in the treatment group had significantly lower mean plaque scores than those in the control group at six months, and at one and two years. The difference between the groups ranged from 35% to 46%"

CHX = chlorhexidine; F = Fluoride; NR = not reported; SD = standard deviation.

Total n is the number of participants analysed in the study arms relevant to the review.

PI (Plaque Index, Silness and Löe 1964) is measured on a 0-3 increasing scale.

TQH (Turesky modification of the Quigley and Hein Index, Turesky 1970) is measured on a 0-5 increasing scale.

OHI-S (Simplified Oral Hygiene Index, Greene and Vermillion 1964) is measured on a 0-3 increasing scale.

Table 9. Studies with a calculus outcome not included in meta-analyses

Time	Study ID	Comparison	Index	CHX mean (SD)	Control mean (SD)	Total n	Notes
4 to 6 weeks	Anderson 1997	CHX 0.12% vs placebo	RI	0.1075	0.0475	29	Mean RI by surface + SD reported. We calculated overall mean. No overall SD
7 to 12 weeks	Anderson 1997	CHX 0.12% vs placebo	RI	0.0875	0.0525	28	Mean RI by surface + SD reported. We calculated overall mean. No overall SD

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	Charles 2004	CHX 0.12% vs control	VM	0.37 (SD NR)	0.11 (SD NR)	74	Did not report a SD
	Sanz 1994	CHX 0.12% vs placebo	VM	Graph	Graph	130	Data presented in graph. Did not report a SD. Quote: "All groups developed calculus after the initial cleaning at baseline. This increase was only statistically significant for the positive control group compared with the control group at 6 months"
Calculus 6 months	Banting 1989	CHX 0.12% vs placebo	VM	NR	NR	383	Results reported at 24 months only. See results at > 6 months below
	Charles 2004	CHX 0.12% vs control	VM	0.45 (SD NR)	0.21 (SD NR)	73	Did not report a SD
	Grossman 1986	CHX 0.12% vs placebo	Not speci- fied	NR	NR	380	Quote: "Supragingival calculus was higher in the group using chlorhexidine but this increase in calculus did not diminish the therapeutic effects of chlorhexidine since subjects with significant increases in calculus also had significant decreases in gingivitis"
	Sanz 1994	CHX 0.12% vs placebo	VM	Graph	Graph	130	Data presented in a graph. Did not report a SD. Quote: "All groups developed calculus after the initial cleaning at baseline. This increase was only statistically significant for the positive control group compared with the control group at 6 months"
Calculus > 6 months	Banting 1989	CHX 0.12% vs placebo	VM	NR	NR	272	At 24 months subjects in the treatment group had higher mean supragingival calculus scores, but at the same time more subjects were free of subgingival calculus

CHX = chlorhexidine; NR = not reported; SD = standard deviation.

Total n is the number of participants analysed in the study arms relevant to the review.

RI (Retention Index, Björby and Löe 1966) is measured on a 0-3 increasing scale.

VM (Volpe-Manhold Calculus Index, Manhold 1965; Volpe 1965) measures calculus present on the lingual surface of the lower 6 anterior teeth. Calculus is measured in 3 planes using a standard periodontal probe. The greatest value allowed for any 1 plane is 3 units, therefore the maximum score per tooth is 9 units or 54 units per subject. The mean per subject score is obtained by dividing the total calculus score by the number of lower anterior teeth. A mean calculus score for the group is then calculated.

Table 10. Studies with a staining outcome not included in meta-analyses

Time	Study ID	Comparison	Index	CHX mean (SD/SE) or n (%)	Control mean (SD)	Total n	Notes
4 to 6 weeks	Axelsson 1987	CHX 0.2%/0.1% vs control	DI (Lobene)	0.1171 (0.297)	0	64	Not included in meta- analysis as control mean is 0. We combined data from the 0.2% and 0.1% CHX groups
	Bhat 2014	CHX 0.2% vs control	Reported number and proportion of participants with mild or moderate 'discolouration' in text	20 (91%)	0	44	Not clear whether discolouration relates to teeth or oral tissues or both
	Flotra 1972 & 1971 (4- month study that reports this outcome at 4 weeks)	CHX 0.2%/0.1%/0.1% (acetate) vs control	No index Reported narratively	12% of tooth sur- faces with- out fillings 62% of sili- cate fillings	NR	48 (at 4 weeks)	Quote: "12% of the tooth surfaces without fillings became discolored within the first 4 weeks of the experimentthis happened more frequently on the interproximal surfaces than on the labial surfaces (ratio2:1). Sixty-two per cent of the silicate fillings in these areas were discol-
	Graziani 2015	CHX 0.2% vs control	SI	0.2228 (0.18)	0	70	ored" Not included in meta- analysis as control mean is 0. We combined data from the 3 CHX groups
	Hase 1995	CHX 0.2% vs placebo	Subjective of participants regarding staining of teeth <u>and/or</u> tongue using VAS	38 (SE 7)	9 (SE 1)	39	Teeth and/or tongue stain- ing reported together. Da- ta estimated from a graph
	López-Jor- net 2012	CHX 0.2% vs placebo	Registration of side effects including denture/dental staining (n and %)	2 (5.71%)	3 (8.57%)	70	Denture and dental stain- ing reported together
	Turkoglu 2009	CHX (conc not reported) vs placebo	No index Reported narratively	14 (56%)	Assumed 0	50	Quote: "Of the 25 subjects who rinsed their mouth with CHX mouthrinse14

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		J	e not included in meta-analyses (Continued)				showed discolouration of teeth and/or tongue"
	Zimmer 2006	CHX 0.06%+F +OH vs OH	Staining of teeth and tongue registered at final examination (n)	4	0	78	Stain on teeth/tongue re- ported together. If more than 1 side effect was present, only the most rel- evant was listed i.e. side effects were reported with no double counting
7 to 12 weeks	Charles 2004	CHX 0.12% vs control	DI (Lobene)	1.61 (SD NR)	0.01 (SD NR)	74	Did not report a SD
	Grossman CHX 0.12% vs Not specified Quote: "Photographs of fa- 1989 placebo cial surfaces of the 12 anterior teeth (maxil- lary and mandibular, cuspid to cuspid) were graded for stain intensity and coverage"		4.66 (SD NR)	2.59 (SD NR)	246	Did not report a SD	
	Sanz 1994	CHX 0.12% vs placebo	Not specified	NR	NR	130	Data presented in a graph at 6 months only See 6- month results below
	Zimmer 2006	CHX 0.06% +F +OH vs OH	Staining of teeth and tongue registered at final examination (n)	6	0	78	Stain on teeth/tongue re- ported together. If more than 1 side effect was present, only the most rel- evant was listed i.e. side effects were reported with no double counting
6 months	Charles 2004	CHX 0.12% vs control	DI (Lobene)	2.08 (SD NR)	0.01 (SD NR)	73	Did not report a SD
	Grossman CHX 0.12% vs Not specified 1986 placebo		NR	NR	380	Outcome data not reported. Quote: "Some extrinsic tooth staining was observed in the chlorhexidine group"	
	Grossman 1989	CHX 0.12% vs placebo	Not specified Quote: "Photographs of facial surfaces of the 12 anterior teeth (maxillary and mandibular, cuspid to cuspid) were graded for stain intensity and coverage"	5.15 (SD NR)	2.75 (SD NR)	246	Did not report a SD

Table 10. Studies with a staining outcome not included in meta-analyses (Continued)

Hoffmann 2001	CHX 0.1%/ 0.06%/0.06%+F vs control	DI (Lang and Räber)	1.13/1.02/1.06 (SD NR)	0.38 (SD NR)	58	Median only. Did not report a SD
Sanz 1994	CHX 0.12% vs placebo	Not specified Quote: "Photographs of facial surfaces of the 12 anterior teeth (maxillary and mandibular, cuspid to cuspid) were graded for stain intensity and coverage"	Graph	Graph	130	Data presented in graph Did not report a SD Quote: "Statistically signif- icant more overall stain- ing, more intense stain- ing and stain coverage per tooth were detected for the positive control groupcompared with the control group"

CHX = chlorhexidine; conc = concentration; F = fluoride; NR = not reported; OH = oral hygiene; SD = standard deviation; SE = standard error; VAS = visual analogue scale. Total n is the number of participants analysed in the study arms relevant to the review.

DI (Discolouration Index, Lobene 1968): gingival and body regions of the tooth are scored for intensity (0-3 increasing scale) and severity (0-3 increasing scale).

DI (Discolouration Index, Lang and Räber 1981) is measured on a 0-3 increasing scale.

SI (Staining Index): the buccal surfaces of the 8 central incisors were divided into 3 areas: incisal, approximal and gingival according to Lobene 1968 and Grundemann 2000 and a SI was used to record the dichotomous presence or absence of staining in each area and to calculate the percentage of the total area showing staining.



APPENDICES

Appendix 1. Cochrane Oral Health's Trials Register search strategy

From July 2014, searches of Cochrane Oral Health's Trials Register were conducted using the Cochrane Register of Studies using the search strategy below:

- 1 ((plaque or gingivitis or gingiva* or periodont* or "gum disease*"):ti,ab) AND (INREGISTER)
- 2 ((mouthwash* or mouthrinse* or wash* or rins*):ti,ab) AND (INREGISTER)
- 3 ((chlorhexidine* or MK-412a or "MK 412a" or MK412a or CHX or Corsodyl or Peridex or Hibident or Prexidine or Parodex or Chlorexil or Peridont or Eludril or Perioxidin or Chlorohex or Savacol or Periogard or Chlorhexamed or Nolvasan or Sebidin or Tubulicid or hibitane):ti,ab) AND (INREGISTER)
- 4 (#1 and #2 and #3) AND (INREGISTER)

Previous searches of Cochrane Oral Health's Trials Register were undertaken in April 2010 and February 2012 using the Procite software and the search strategy below:

((plaque or gingivitis or gingiva* or periodont* or "gum disease*") AND (mouthwash* or mouthrinse* or wash* or rins*) AND (chlorhexidine* or MK-412a or "MK 412a" or MK412a or CHX or Corsodyl or Peridex or Hibident or Prexidine or Parodex or Chlorexil or Peridont or Eludril or Perioxidin or Chlorohex or Savacol or Periogard or Chlorhexamed or Nolvasan or Sebidin or Tubulicid or hibitane))

Appendix 2. Cochrane Central Register of Controlled Trials (CENTRAL) search strategy

- #1 MeSH descriptor Dental Plaque, this term only
- #2 ((dental or oral or tooth or teeth) and plaque)
- #3 MeSH descriptor Gingivitis explode all trees
- #4 gingivitis
- #5 ((gingiva* near/3 inflamm*) or (gingiva* near/3 bleed*) or (gingiva* near/3 pocket*) or (gingiva* near/3 attachment*))
- #6 MeSH descriptor Periodontitis explode all trees
- #7 periodont*
- #8 ((haemorrhag* near/5 gum*) or (hemorrhag* near/5 gum*) or (hemorrag* near/5 gum*) or (haemorrhag* near/5 gingiva*) or (hemorrhag* near/5 gingiva*) or (hemorrhag* near/5 gingiva*))
- #9 ((bleed* near/5 gum*) or (bleed* near/5 gingiva*))
- #10 ((inflamm* near/5 gum*) or (inflamm* near/5 gingiva*))
- #11 ((gum next disease) or (disease* near/5 gum*) or (disease near/5 gingiva*))
- #12 (#1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7 OR #8 OR #9 OR #10 OR #11)
- #13 MeSH descriptor Mouthwashes explode all trees
- #14 (mouthrins* or "mouth rins*" or mouth-rins* or rinse* or rinsing* or mouthwash* or "mouth wash*" or mouth-wash*)
- #15 ((oral near/3 wash*) or (dental near/3 wash*) or (mouth* near/3 wash*) or (oral near/3 rins*) or (dental near/3 rins*) or (mouth* near/3 rins*))
- #16 (#13 OR #14 OR #15)
- #17 MeSH descriptor Chlorhexidine, this term only
- #18 chlorhexidine
- #19 (MK-412a or "MK 412a" or MK412a)
- #20 (CHX):ti,ab,kw
- #21 (Corsodyl or Peridex or Hibident or Prexidine or Parodex or Chlorexil or Perident or Eludril or Perioxidin or Chlorohex or Savacol or Periogard or Chlorhexamed or Nolvasan or Sebidin or Tubulicid or hibitane)
- #22 (#17 OR #18 OR #19 OR #20 OR #21)
- #23 (#12 AND #16 AND #22)

Appendix 3. MEDLINE Ovid search strategy

- 1. Dental plaque/
- 2. ((dental or oral or tooth or teeth) and plaque).mp.
- 3. exp Gingivitis/
- 4. gingivitis.mp.
- 5. ((gingiva\$ adj3 inflamm\$) or (gingiva\$ adj3 bleed\$) or (gingiva\$ adj3 pocket\$) or (gingiva\$ adj3 attachment\$)).mp.
- 6. exp Periodontitis/
- 7. periodont\$.mp.
- 8. ((haemorrhag\$ adj5 gum\$) or (hemorrhag\$ adj5 gum\$) or (hemorrag\$ adj5 gum\$) or (haemorrhag\$ adj5 gingiva\$) or (hemorrhag\$ adj5 gingiva\$)).mp.



- 9. ((bleed\$ adj5 gum\$) or (bleed\$ adj5 gingiva\$)).mp.
- 10.((inflamm\$ adj5 gum\$) or (inflamm\$ adj5 gingiva\$)).mp.
- 11.((gum adj disease) or (disease\$ adj5 gum\$) or (disease adj5 gingiva\$)).mp.

12.or/1-11

- 13.exp Mouthwashes/
- 14. (mouthrins or "mouth rins or mouth-rins or rinse or rinsing or mouthwash or "mouth wash or "mouth wash or "mouth wash or "mouth wash or mouth wash or mo
- 15.((oral or dental or mouth) adj3 (wash\$ or rins\$)).mp.

16.or/13-15

- 17.Chlorhexidine/
- 18.chlorhexidine.mp.
- 19.(MK-412a or "MK 412a" or MK412a).mp.
- 20.CHX.ti,ab.
- 21.(Corsodyl or Peridex or Hibident or Prexidine or Parodex or Chlorexil or Perident or Eludril or Perioxidin or Chlorohex or Savacol or Periogard or Chlorhexamed or Nolvasan or Sebidin or Tubulicid or hibitane).mp.
- 22.or/17-21
- 23.12 and 16 and 22

This subject search was linked to the Cochrane Highly Sensitive Search Strategy (CHSSS) for identifying randomised trials (RCTs) in MEDLINE: sensitivity-maximising version (2008 revision) as referenced in Chapter 6.4.11.1 and detailed in box 6.4.c of the *Cochrane Handbook for Systematic Reviews of Interventions*, Version 5.1.0 (updated March 2011) (Lefebvre 2011).

- 1. randomized controlled trial.pt.
- 2. controlled clinical trial.pt.
- 3. randomized.ab.
- 4. placebo.ab.
- 5. drug therapy.fs.
- 6. randomly.ab.
- 7. trial.ab.
- 8. groups.ab.
- 9. or/1-8
- 10. exp animals/ not humans.sh.
- 11. 9 not 10

Appendix 4. Embase Ovid search strategy

- 1. Dental plaque/
- 2. ((dental or oral or tooth or teeth) and plaque).mp.
- 3. exp Gingivitis/
- 4. gingivitis.mp.
- 5. ((gingiva\$ adj3 inflamm\$) or (gingiva\$ adj3 bleed\$) or (gingiva\$ adj3 pocket\$) or (gingiva\$ adj3 attachment\$)).mp.
- 6. exp Periodontitis/
- 7. periodont\$.mp.
- 8. ((haemorrhag\$ adj5 gum\$) or (hemorrhag\$ adj5 gum\$) or (hemorrag\$ adj5 gum\$) or (haemorrhag\$ adj5 gingiva\$) or (hemorrhag\$ adj5 gingiva\$)).mp.
- 9. ((bleed\$ adj5 gum\$) or (bleed\$ adj5 gingiva\$)).mp.
- 10.((inflamm\$ adj5 gum\$) or (inflamm\$ adj5 gingiva\$)).mp.
- 11.((gum adj disease) or (disease\$ adj5 gum\$) or (disease adj5 gingiva\$)).mp. [mp=title, original title, abstract, name of substance word, subject heading word, unique identifier]
- 12.or/1-11
- 13.exp Mouthwashes/
- 14. (mouthrins or "mouth rins or mouth-rins or rinse or rinsing or mouthwash or "mouth wash or "mouth wash or mouth-wash or mouth-wash.) .mp.
- 15.((oral or dental or mouth) adj3 (wash\$ or rins\$)).mp.
- 16.or/13-15
- 17.Chlorhexidine/
- 18.chlorhexidine.mp.
- 19.(MK-412a or "MK 412a" or MK412a).mp.
- 20.CHX.ti,ab.



21.(Corsodyl or Peridex or Hibident or Prexidine or Parodex or Chlorexil or Perident or Eludril or Perioxidin or Chlorekex or Savacol or Periogard or Chlorhexamed or Nolvasan or Sebidin or Tubulicid or hibitane).mp.

22.or/17-21

23.12 and 16 and 22

The above subject search was linked to adapted version of the Cochrane Embase Project filter for identifying RCTs in Embase Ovid (see http://www.cochranelibrary.com/help/central-creation-details.html for information):

- 1. Randomized controlled trial/
- 2. Controlled clinical study/
- 3. Random\$.ti,ab.
- 4. randomization/
- 5. intermethod comparison/
- 6. placebo.ti,ab.
- 7. (compare or compared or comparison).ti.
- 8. ((evaluated or evaluate or evaluating or assessed or assess) and (compare or compared or comparing or comparison)).ab.
- 9. (open adj label).ti,ab.
- 10. ((double or single or doubly or singly) adj (blind or blinded or blindly)).ti,ab.
- 11. double blind procedure/
- 12. parallel group\$1.ti,ab.
- 13. (crossover or cross over).ti,ab.
- 14. ((assign\$ or match or matched or allocation) adj5 (alternate or group\$1 or intervention\$1 or patient\$1 or subject\$1 or participant \$1)).ti,ab.
- 15. (assigned or allocated).ti,ab.
- 16. (controlled adj7 (study or design or trial)).ti,ab.
- 17. (volunteer or volunteers).ti,ab.
- 18. trial.ti.
- 19. or/1-18
- 20. (exp animal/ or animal.hw. or nonhuman/) not (exp human/ or human cell/ or (human or humans).ti.)
- 21. 19 not 20

Appendix 5. CINAHL EBSCO search strategy

- S1 MH "Dental Plaque+"
- S2 ((dental or oral or tooth or teeth) and plaque)
- S3 MH "Gingivitis+"
- S4 gingivitis
- S5 (gingiva* N3 inflamm*) or (gingiva* N3 bleed*) or (gingiva* N3 pocket*) or (gingiva* N3 attachment*)
- S6 MH "Periodontitis+"
- S7 periodont*
- S8 (haemorrhag* N5 gum*) or (hemorrhag* N5 gum*) or (hemorrag* N5 gum*) or (haemorrhag* N5 gingiva*) or (hemorrhag* N5 gingiva*) or (hemorrag* N5 gingiva*)
- S9 (bleed* N5 gum*) or (bleed* N5 gingiva*)
- S10 (inflamm* N5 gum*) or (inflamm* N5 gingiva*)
- S11 "gum disease" or (disease* N5 gum*) or (disease* N5 gingiva*)
- S12 S1 or S2 or S3 or S4 or S5 or S6 or S7 or S8 or S9 or S10 or S11
- S13 MH "Mouthwashes+"
- S14 mouthrins* or "mouth rins*" or mouth-rins* or rinse* or rinsing* or mouthwash* or "mouth wash*" or mouth-wash*
- S15 (oral N3 wash*) or (dental N3 wash*) or (mouth* N3 wash*) or (oral N3 rins*) or (dental N3 rins*) or (mouth* N3 rins*)
- S16 S13 or S14 or S15
- S17 MH "Chlorhexidine+"
- S18 chlorhexidine
- S19 MK-412a or "MK 412a" or MK412a
- S20 TI CHX or AB CHX
- S21 (Corsodyl or Peridex or Hibident or Prexidine or Parodex or Chlorexil or Peridont or Eludril or Perioxidin or Chlorohex or Savacol or Periogard or Chlorhexamed or Nolvasan or Sebidin or Tubulicid or hibitane)
- S22 S17 or S18 or S19 or S20 or S21
- S23 S12 and S16 and S22

The above subject search was linked to Cochrane Oral Health's filter for identifying RCTs in CINAHL:

S1 MH Random Assignment or MH Single-blind Studies or MH Double-blind Studies or MH Triple-blind Studies or MH Crossover design or MH Factorial Design



S2 TI ("multicentre study" or "multicenter study" or "multi-centre study") or AB ("multicentre study" or "multicenter study") or BU ("multicentre study" or "multi-centre stud

S3 TI random* or AB random*

S4 AB "latin square" or TI "latin square"

S5 TI (crossover or cross-over) or AB (crossover or cross-over) or SU (crossover or cross-over)

S6 MH Placebos

S7 AB (singl* or doubl* or trebl* or tripl*) or TI (singl* or doubl* or trebl* or tripl*)

S8 TI blind* or AB mask* or AB blind* or TI mask*

S9 S7 and S8

S10 TI Placebo* or AB Placebo* or SU Placebo*

S11 MH Clinical Trials

S12 TI (Clinical AND Trial) or AB (Clinical AND Trial) or SU (Clinical AND Trial)

S13 S1 or S2 or S3 or S4 or S5 or S6 or S9 or S10 or S11 or S12

Appendix 6. US National Institutes of Health Ongoing Trials Register (ClinicalTrials.gov) search strategy

chlorhexidine AND mouthrinse AND gingivitis

chlorhexidine AND mouthwash AND gingivitis

Appendix 7. World Health Organization International Clinical Trials Registry Platform search strategy

chlorhexidine AND mouthrinse AND gingivitis

chlorhexidine AND mouthwash AND gingivitis

HISTORY

Protocol first published: Issue 9, 2010 Review first published: Issue 3, 2017

Date	Event	Description
16 August 2010	Amended	Amended declaration of interest statement.

CONTRIBUTIONS OF AUTHORS

Helen Whelton (HW) conceived the idea for the review. Patrice James (PJ), Carmel Parnell (CP), Philip Riley (PR), Thomas Lamont (TL) and Mairead Harding (MH) identified the studies to be included in the review. PJ, CP, PR, TL, MH and Andrea Cheung (AC) carried out data extraction. PJ, AC, PR and TL compiled the characteristics of included studies tables. PJ, CP, PR, MH and TL assessed the risk of bias for the included studies. Helen Worthington (HVW) statistically analysed the data. PJ, TL and AC drafted the background section. PJ, HVW and PR drafted the body of the review. HW, MH, CP and AC commented on and edited the draft review. HVW and PR provided advice and guidance throughout the review process.

DECLARATIONS OF INTEREST

As former Director of the Oral Health Services Research Centre, Cork, Helen Whelton has conducted clinical trials of products for GlaxoSmithKline (GSK) and she received consultancy fees in relation to this research. Mairead Harding (Deputy Director of the Oral Health Services Research Centre) was involved in the development of educational materials with regard to tooth wear/dental erosion and has conducted epidemiological studies and presented the findings on tooth wear and dental erosion at conferences. Fees received from GlaxoSmithKline for this work were paid to the Oral Health Services Research Centre, Cork and not to Mairead Harding. Philip Riley is a salaried member of the Cochrane Oral Health editorial team. Editorial control of the review and final decisions about the content of the review were maintained by Helen Worthington who is Co-ordinating Editor of Cochrane Oral Health. The remaining co-authors declare no conflict of interest.



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• Cochrane Oral Health Global Alliance, Other.

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DIFFERENCES BETWEEN PROTOCOL AND REVIEW

In Spring 2016, in conjunction with the editorial board of Cochrane Oral Health, the decision was made to address the comparison of chlorhexidine mouthrinse with placebo, control or mechanical oral hygiene alone in this review and to report the comparison of chlorhexidine mouthrinse with other active mouthrinses in a subsequent review. This decision will enable us to comprehensively report all of the results for the objectives set out in the published protocol (James 2010) across two reviews.

A number of additional changes were made in relation to the criteria for including studies in the review after the protocol was published but early in the review process.

- · Types of studies
 - * It was decided to exclude cross-over trials due to concerns that chlorhexidine could exert an effect beyond the washout period.
 - * When we wrote the protocol, we did not anticipate encountering split-mouth studies meeting the inclusion criteria because it is not possible to conduct a split-mouth study when using mouthrinse as the intervention. However, we encountered split-mouth studies comparing different scaling and root planing regimens and different periodontal surgical techniques that incorporated a chlorhexidine and placebo/control comparison. Such study designs were considered inappropriate to answer the question posed by this review and were excluded.
- Types of participants
 - * Individuals no longer need to have undergone periodontal treatment and be in the maintenance phase to be included. This change was made to allow us to include studies where chlorhexidine was used as an adjunct to surgical and non-surgical periodontal therapy. Individuals of any age, gender or race with periodontal disease (gingivitis or periodontitis) provided they are capable of performing mechanical oral hygiene procedures are included.
- Risk of bias
 - * Assessment of blinding for participants, personnel and outcome assessors was changed to allow for low and unclear risk of bias in these domains in certain situations.

The following were not explicitly addressed in the protocol and required clarification.

- Types of intervention
 - * We clarified that the mechanical oral hygiene procedures must be the same in both the chlorhexidine mouthrinse and the comparison arms so that the chlorhexidine mouthrinse is the only difference between the arms to ensure that the groups are truly comparable.
 - * Studies where the chlorhexidine mouthrinse also contained fluoride were included.
 - * Studies where gum care or antigingivitis dentifrices (that do not contain chlorhexidine) are used for mechanical oral hygiene in both test and comparator arms were included.
 - * Studies where chlorhexidine mouthrinse formed part of a combined intervention with other agents (such as other chlorhexidine vehicles, dentifrice containing chlorhexidine, or other antigingivitis agents (e.g. cetylpyridinium chloride (CPC)) that the comparator arm/s did not receive were excluded because the effect of the chlorhexidine could not be separated from the effect of the other active agents.



- Types of outcome
 - * A hierarchy to guide data extraction of gingivitis and plaque data was developed to facilitate data extraction and analysis. The main analysis however, was based on the main prespecified gingivitis index: the Gingival Index of Löe and Silness (Löe 1967; Löe and Silness 1963).

INDEX TERMS

Medical Subject Headings (MeSH)

Chemotherapy, Adjuvant; Chlorhexidine [adverse effects] [*therapeutic use]; Dental Plaque [complications] [*drug therapy]; Dental Plaque Index; Dental Prophylaxis; Gingivitis [*drug therapy] [etiology]; Mouthwashes [adverse effects] [*therapeutic use]; Oral Hygiene; Publication Bias; Randomized Controlled Trials as Topic; Time Factors; Tooth Discoloration [chemically induced]

MeSH check words

Adolescent; Adult; Aged; Aged, 80 and over; Child; Female; Humans; Male; Middle Aged