The effects of an Ayurvedic medicinal toothpaste on clinical, microbiological and oral hygiene parameters in patients with chronic gingivitis: a double-blind, randomised, placebo-controlled, parallel allocation clinical trial

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(Index words: herbal medicinal toothpaste, chronic gingivitis, salivary anaerobe counts, clinical trial, gingival bleeding)

Abstract

Introduction Plant derived preparations have been essential components for maintenance of oral hygiene and the treatment of oral diseases globally since ancient times. Acacia chundra Willd, Adhatoda vasica Nees., Minusopsis elengi L., Piper nigrum L., Pongamia pinnate L., Pirerre, Quercus infectoria Olivier., Syzygium aromaticum L., Terminalia chebula Retz., Zingiber offici-nale Roscoe., individually or in combination, have been used for this purpose because of their beneficial effects.

Objectives To study the efficacy of an Ayurvedic toothpaste containing these herbs in patients with chronic gingivitis.

Methods Otherwise healthy males and non-pregnant females (n=80) aged 18-35 years with ≥20 teeth were randomly assigned to Group 1 (herbal toothpaste) and Group 2 (placebo toothpaste). Quigley Hein plaque index (PS), bleeding on probing (BOP) and probing pocket depth (PPD), were recorded for all teeth at six sites, and one ml of resting saliva was collected to ascertain anaerobic and aerobic bacterial counts at baseline, and at 4, 8, 12 and 24 weeks. Full-mouth prophylaxis was performed and instructions for brushing with the allocated toothpaste for 6 months were given at baseline. Sixty-six participants, 34 in Group 1 and 32 in Group 2 completed the study. Clinical examinations were performed by the same examiner blinded to group allocation.

Results Linear mixed model analysis revealed significant reductions of PS, BOP, PPD (p<0.0001) and total salivary anaerobic counts (p<0.05) in Group 1 at all prescribed visits compared to Group 2. Moreover the reduction increased overtime. No unpleasant effects of toothpaste use were reported.

Conclusions This study provides robust evidence of the beneficial antiplaque and antigingivitis effects of the test herbal toothpaste Sudantha\textsuperscript{6} on patients with chronic gingivitis.

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Introduction

Plaque induced chronic gingivitis is an oral health problem prevalent worldwide, affecting the dentition in people of all ages. Dental plaque, a well organized biofilm, has been established as the cause of chronic gingivitis [1]. Chronic gingivitis may progress to more destructive chronic and aggressive periodontitis, leading to loss of teeth [2, 3].

There is evidence that oral biofilm-associated diseases may affect systemic health by mechanisms such as spreading infections to adjacent tissues and spaces, hematogenous dissemination of oral biofilm organisms, or inflammatory mechanisms [4]. Elevated surrogate markers of inflammation in patients with periodontitis in cross-sectional studies substantiate this notion [5]. Further, evidence suggests that oral biofilm-associated chronic periodontitis enhances the risk of coronary heart disease and cerebrovascular disease, and that poor glycaemic control in diabetic patients with periodontitis is a concern for clinicians [6-8].

Hence the prevention and treatment of chronic gingivitis is important for maintenance of good oral health as well as general health. At present toothbrushes,
dental floss and toothpicks are employed for mechanical removal of the oral biofilm. However, this approach relies heavily on motivation and the ability of the individual to remove dental plaque efficiently on a daily basis. It has been reported that most people find brushing teeth and flossing difficult tasks, and are not able to remove plaque completely from all tooth surfaces [9]. So ineffective removal of plaque can lead to gingivitis, especially at difficult to reach sites of the dentition.

The use of chemotherapeutic agents is an accepted adjunctive mode for reducing plaque build-up, and incorporation of antiplaque and anti-gingivitis agents to the oral hygiene regimen has been found to be beneficial [10, 11]. These antimicrobial agents may be chemicals or herbal preparations. Various medicinal plants, individually or in combination, have been used in Sri Lanka for over 2000 years to maintain oral hygiene and to prevent periodontal diseases. Among these are *Acacia chundra* Willd., *Adhatoda vasica* Nees., *Mimusops elengi* L., *Piper nigrum* L., *Pongamia pinnata* (L.) Pirerre, *Quercus infectoria* Olivier., *Syzygium aromaticum* L., *Terminalia chebula* Retz., *Zingiber officinale* Rosce. Studies have shown beneficial therapeutic effects; antibacterial, antioxidant, antifungal, antiplasmodial and anti-inflammatory effects of these herbs.

We studied the therapeutic effects of a proprietary herbal toothpaste which has been formulated with an extract of these nine herbs. It is registered as Sudantha®, an Ayurvedic medicinal tooth-paste widely used in Sri Lanka (Reg.No.02/01/PV/08/143).

**Methods**

**Trial design**

Our study was a randomised, placebo-controlled, parallel allocation, 6-month clinical trial with blinding of the participants, clinical assessor and statistician. It was conducted at the Clinical Oral Microbiology Industry Collaborative Research Unit of the Faculty of Dental Sciences, University of Peradeniya, Sri Lanka. Approval for the trial was given by the Research and Ethics Review Committee of the Faculty of Dental Sciences of the University, and it was registered at the Sri Lanka Clinical Trials Registry (No.SLCTR/2010/012). All participants consented by voluntarily signing a witnessed agreement after receiving verbal and written information about the study. The trial was conducted in harmony with the guidelines for evaluating chemotherapeutic products for the control of gingivitis outlined by the American Dental Association [12, 13]. The outline of the study protocol is given in Figure 1

**Study population**

Based on a previous study of this herbal toothpaste, a sample of 35 was needed in each group to detect 30% reduction in bleeding on probing with 85% power. We recruited a sample of 80 in view of possible withdrawal of participants for non-compliance or reasons not related to the trial.

**Recruitment of participants**

Recruitment of participants commenced in November 2010 and the trial was completed in June 2012. Participants were selected from patients attending the diagnostic clinic of the Faculty of Dental Sciences, University of Peradeniya, and from the dental clinic of the Teaching Hospital, Kandy, Sri Lanka, after an oral and periodontal examination. Patients with generalised chronic gingivitis with over 30% sites with bleeding were selected as participants subject to the following inclusion criteria: non-smoking men and non-pregnant women between 18-35 years of age with at least 20 teeth, willingness to cooperate with the study protocol and attend all visits and agreeing to sign the consent form. The exclusion criteria were presence of gross oral pathology, presence of medical conditions that affect the integrity of periodontium or oral mucosa, any other chronic diseases that have an effect on periodontal disease, presence of teeth with defective or extensive restorations (including prosthetic crowns), pulpitis, caries, cracked enamel, use of any intra-oral appliances, perio-dontitis or history of treatment for periodontitis within the past twelve months, past history of known allergy to oral care products, regular use of anticonvulsants, anti-histamines, sedatives, tranquilizers, anti-inflammatory drugs or analgesics, need for antibiotic prophylaxis before the dental examination, or history of antibiotic use within the past three months.

Selected participants were randomly assigned to either the Group 1 or the Group 2 by a computer generated randomisation list (http://www.random.org). Investigator not involved in clinical examinations generated the random numbers and they were kept sealed until

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**Figure 1. Study Protocol**

- **Screening and selection**
- **Consent**
- **Random assignment**
  - **G 1 N=40**
  - **G 2 N=40**
- **Baseline examination + Prophylaxis**
- **Re-examination 4 WEEKS N=72**
- **Re-examination 8 WEEKS N=72**
- **Re-examination 12 WEEKS N=72**
- **Final examination 24 WEEKS N=66**
- **Data analysis**
- **BOP, PPD, PLAQUE**
- **Iml Saliva**
- **Adverse effects**
preliminary statistical analysis was over. Toothpaste tubes were in bags numbered 1-80. Participants were assigned the bags based on the number. Allocation ratio was 1:1.

Baseline examination was performed to assess the levels of plaque, bleeding on probing and pocket depth. All examinations were carried out by a single assessor, standardised with an experienced clinician and calibrated for intra-examiner variability (Kappa value 0.99 for PPD and 0.75 for plaque score). The examiner was blinded to the group allocation of the participant. At baseline, BOP and PPD using 15NC periodontal probe with a ball ended tip were recorded at 6 sites per tooth; mesio-facial, mid-facial, disto-facial, mesio-lingual, mid-lingual and disto-lingual. Then plaque score was recorded at 6 sites using Turesky modification of the Quigley and Hein plaque index (PS) after a plaque disclosing solution (Dent. Liquid plaque tester, Japan) was applied on all tooth surfaces [14, 15]. Thereafter, all participants were given brushing instructions and oral prophylaxis. No other instructions were given with regard to the use of toothpaste or brushing during the trial period. Participants were given the allocated toothpaste for home use and advised to brush twice daily for 24 weeks.

The test herbal toothpaste and placebo toothpaste were packed in identical tubes and dispensed by an investigator not involved with clinical examinations. The assessor performing clinical examination and salivary microbial analyses was blinded to the group allocation of the participants. The participants were also not aware of the type of toothpaste allocated to them. All measurements were repeated at 4, 8, 12 and 24 weeks. For clinical examinations, participants were instructed to refrain from brushing for about 12 hours before the clinic visit to standardize the time of collection of saliva for microbial culture. Saliva was collected and immediately cultured before the clinical examination.

Microbiological procedures

One ml of resting saliva was collected and centrifuged at 4000 rpm for 5 minutes and the pellet obtained was re-suspended in 1000µl of sterile isotonic saline and 10-fold dilutions of the re-suspended sample were prepared. Then 100µl of three dilutions 10⁻¹, 10⁻⁰ and 10⁻¹ dilutions were separately cultured in 5% sheep blood agar plates and incubated aerobically and anaerobically at 37°C. Total aerobic and anaerobic colony forming units (CFU) were counted after 24 and 48 hours. Total CFUs were calculated per one ml of saliva.

Statistical analysis

Analysis of data was performed using a statistical package Statistical Analysis Software SAS 9.1. Identity of each toothpaste tube was revealed at the end of the trial after the preliminary statistical analysis. Each participant was considered as an observational unit of analysis. For each variable i.e. PS, BOP and PPD values (six sites for each tooth was recorded and overall average was taken as the observation for analysis for each visit). Each parameter was recorded at baseline, 4, 8, 12 and 24 weeks. Likewise, each subject had 5 readings for salivary aerobic and anaerobic counts at baseline, 4, 8, 12 and 24 weeks examinations.

Since our data set includes multiple observations of clinical and microbiological parameters of same sampling unit (participant) over a six month period the analysis must address the issue of correlation between measures of the same sampling unit. Hence linear mixed model analysis was used as the statistical method in the data analysis.

Results

Study population

Sixty six participants (test n=34, placebo n=32) completed the 24-week trial. Mean age of the participants in Group 1 was 24.5±4.7 years and Group 2 it was 24.5±4.8 years. Six participants dropped out at 12 weeks for reasons not related to the trial, and 8 participants dropped out after the baseline examination. Hence 72 participants (31 male, 41 female) attended re-examination at 4 weeks, 8 weeks, 12 weeks (Group 1 Male – 17, Female – 19, Group 2 Male – 14, Female – 22). Another six participants were dropped out due to reasons not related to the trial. As a result, 66 participants completed all re-examination visits (Group 1 Male – 16, Female - 18, Group 2 Male – 12, Female – 20).

Baseline data

At the baseline there were no significant differences between mean values of Group 1 and Group 2 with

Table 1. Effect of different toothpaste use on plaque score during the trial

<table>
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B: Least Squares Means for Group Visit Interactions

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respect to all clinical parameters: PS (2.66, vs 2.50 with SED=0.1315, t=-1.21, p=0.23), BOP (0.59 vs 0.57 with SED=0.0293, t=-1.14, p=0.26), PPD (2.309 vs 2.305 with SED=0.0607, t=0.07, p=0.94), for all three variables.

Plaque scores, bleeding on probing, probing pocket depth during the trial

The results with respect to all three clinical variables PS, PPD and BOP were similar. The type 3 test of fixed effects for the three effects-Group (1 and 2), visit (baseline, 4,8,12 and 24 weeks) and interaction between group and visit, (Tables 1A & B, 2A & B and 3A & B) show that the type 3 test for all three effects are statistically significant (p<0.0001) in relation to all three clinical parameters. Figures 2, 3 and 4 show a general trend of a gradual reduction of all three parameters – plaque score, probing pocket depth and bleeding on probing, with time. It is also evident from the Figures 2, 3 and 4 that, at each visit the gap between values of two groups has increased and the decrease in all three parameters over time is greater in group 1 than the group 2. These differences between group 1 and group 2 were statistically significant at all visits – at 4, 8, 12 and 24 weeks (p<0.0001) in relation to all three clinical parameters – PS, PPD and BOP (Table 4).

Salivary microbial counts

Table 5A and 5B show the type 3 test of fixed effects for the three effects (group, visit and group*visit) in relation to the total salivary aerobic and anaerobic counts. According to Table 5A, type 3 test for visit is highly significant (p<0.0001) indicating that there is a reduction of total aerobic counts with time in both groups but there is no significant group effect (p>0.05) indicating herbal toothpaste had no significant effect on aerobic counts. There is also no significant effect on the counts due to the combined effect of visit and group indicating the effect did not vary due to the interaction between the test toothpaste use and the duration of its use (p>0.05)

Table 5B shows the type 3 test of fixed effects for dependent variable total anaerobic counts in saliva. It indicates that all three effects – visit, group and interaction between group and visit are statistically significant (p<0.001, p<0.05 and p<0.05 respectively). It also shows that there was a reduction of anaerobic counts during the trial irrespective of group, the group which used the herbal toothpaste had significantly lower counts compared to the group which used the placebo toothpaste (p=0.04).
Table 5A. Effect of different toothpaste use on the salivary aerobic counts during the trial

Null Model Likelihood Ratio Test

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Type 3 Tests of Fixed Effects

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Table 5B. Effect of different toothpaste use on the salivary anaerobic counts during the trial

Null Model Likelihood Ratio Test

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Type 3 Tests of Fixed Effects

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Discussion

We investigated the efficacy of an Ayurvedic medicinal toothpaste formulated with nine medicinal herbs in the management of plaque induced chronic generalised gingivitis. The clinical variables, plaque score, gingival bleeding on probing, probing pocket depth, and salivary anaerobic and aerobic bacterial counts were evaluated during the six-month study period. At all re-examinations the test group had reduced plaque score ($p=0.0001$), mean bleeding on probing ($p=0.0001$) and mean probing pocket depth $p=0.04$. There was a
statistically significant reduction of total salivary aerobic counts in both groups over time when compared with baseline counts. However, these reductions were not statistically significant between groups. The salivary anaerobic counts were also reduced in both groups with observed reduction in the Group1 which used herbal toothpaste being greater compared to that of the placebo group ($p=0.0001$). The reduction of anaerobic counts in the group which used herbal toothpaste was also increased with the duration of its use ($p=0.04$ Table 5B). Initial oral prophylaxis and the Hawthorn effect may have contributed partly to the general reduction of all clinical parameters and aerobic and anaerobic bacterial counts in this clinical trial. However, the finding that the use of herbal toothpaste resulted in statistically significantly greater reduction of all clinical parameters and anaerobic bacterial counts could be due to the antibacterial and antiinflammatory properties of the herbal ingredients used in the toothpaste. This is the first study reporting a six-month clinical trial using a toothpaste formulated with a mixture of herbal extracts, in which the clinical, microbial and oral hygiene parameters have reduced significantly when compared with a placebo toothpaste which had the same ingredients in the herbal toothpaste except the herbal extract. Another finding of this study is the absence of any significant difference in aerobic bacterial counts between the two groups suggesting that this herbal toothpaste does not significantly alter the aerobic bacterial counts. This is a favourable finding as the aerobic bacteria in the oral microbiome are associated with gingival health.

The reduction of anaerobes during the trial period explains the clinical improvement of gingivitis as resolution of gingival inflammation is associated with concomitant reduction of anaerobes in the oral biofilm. Further, the improvements obtained in BOP and plaque score in this study with the use of the test herbal toothpaste was better than in studies which used either the twice daily antimicrobial mouth rinses in addition to routine toothbrushing [16, 17, 18], or the combined effect of antiseptic mouth rinse and antigingivitis and antiplaque dentifrice in another study [19]. Hence the use of antimicrobial rinses as an adjunct to tooth brushing would appear to be superfluous. An added advantage of using the test toothpaste is the complete absence of tooth staining and taste alterations related to its use, when compared to other agents such as chlorhexidine [20].

This trial was a sequel to our previous study with volunteers [21] which showed that three months’ use of this particular herbal toothpaste conferred significant improvement in PS, BOP and salivary anaerobic counts, with no significant effect on the total salivary aerobic counts [20]. It is noteworthy that the beneficial effects observed in this study were with twice a day, unsupervised home use of this toothpaste, with no reinforcement of oral health or brushing instructions during the trial.

In conclusion, this trial provides evidence for the therapeutic benefits of the Ayurvedic medicinal test toothpaste through its anti-inflammatory effects and confirms the previously documented anti-inflammatory and antibacterial properties of the herbs used in this formula.

Acknowledgements

We are grateful to the staff of the Veterinary Research Institute, Gannoruwa, Sri Lanka for providing sheep blood for this study, Mrs. M R D M Senanayake, for technical assistance, the staff of the Division of Microbiology for helping in many ways for this project, and Link Natural Products (Pvt) Ltd., Sri Lanka, for funding this study.

Conflicts of interest

This study was sponsored by Link Natural Products Private Limited which produces the test herbal toothpaste Sudantha. Study was conducted, data collected and analysed at University of Peradeniya. PSR had full access to all the data in this study and takes complete responsibility for the integrity of the data, and the accuracy of the data analysis.

References


