ORIGINAL ARTICLE

Plaque-left-behind after brushing: intra-oral reservoir for antibacterial toothpaste ingredients

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Abstract

Objectives Plaque is never fully removed by brushing and may act as a reservoir for antibacterial ingredients, contributing to their substantive action. This study investigates the contribution of plaque-left-behind and saliva towards substantivity of three antibacterial toothpastes versus a control paste without antibacterial claims.

Materials and methods First, volunteers brushed 2 weeks with a control or antibacterial toothpaste. Next, plaque and saliva samples were collected 6 and 12 h after brushing and bacterial concentrations and viabilities were measured. The contributions of plaque and saliva towards substantivity were determined by combining control plaques with experimental plaque or saliva samples and subsequently assessing their viabilities. Bacterial compositions in the various plaque and saliva samples were compared using denaturing gradient gel electrophoresis.

Results The viabilities of plaques after brushing with Colgate-Total® and Crest-Pro-Health® were smaller than of control plaques and up to 12 h after brushing with Crest-Pro-Health® plaques still contained effective, residual antibacterial activity against control plaques. No effective,

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residual antibacterial activity could be measured in saliva samples after brushing. There was no significant difference in bacterial composition of plaque or saliva after brushing with the different toothpastes.

Conclusions Plaque-left-behind after mechanical cleaning contributes to the substantive action of an antibacterial toothpaste containing stannous fluoride (Crest-Pro-Health®).

Clinical relevance The absorptive capacity of plaque-leftbehind after brushing is of utmost clinical importance, since plaque is predominantly left behind in places where its removal and effective killing matter most. Therewith this study demonstrates a clear and new beneficial effect of the use of antibacterial toothpastes.

Keywords Substantivity · Oral antibacterials · Toothpastes · Plaque · Saliva

Introduction

In the prevention of oral diseases, adequate oral home care is essential and the most applied method to attain this goal is the removal of dental plaque. Although mechanical removal usually yields over 80% removal, plaque is inevitably left behind in the majority of people [1]. Moreover, a plaque score of 0 can never be achieved [2]. Besides mechanical plaque removal by toothbrushing and interdental cleaning, a toothpaste may help to remove plaque and spread therapeutic ingredients, like fluorides and antibacterial agents, through the oral cavity. An effective oral antibacterial should have a broad spectrum of antibacterial activity, but at the same time it should not disturb the oral microbiome at



health [3]. Moreover, it should be compatible with other toothpaste ingredients [4]. There are several ways by which antibacterial agents can influence a biofilm, like interference with bacterial co-aggregation mechanisms or by affecting bacterial viability [5].

The efficacy of an antibacterial agent depends not only on its antibacterial power but also on its substantivity. Due to its capability to bind to oral surfaces and subsequent slow release in bio-active concentrations, a substantive antibacterial agent remains active in the oral cavity for a prolonged period of time [6-8]. A wellknown example of a substantive antibacterial agent is chlorhexidine. Effects on bacterial viability in saliva of a 0.2% chlorhexidine rinse could be perceived up to 7 h post-use [9]. Also substantive effects of an amine fluoride/stannous fluoride mouthrinse and toothpaste [10] and a triclosancontaining toothpaste [6] on biofilm vitality have been observed, respectively 7 and 24 h after application. However, it was never assessed whether absorption of antibacterial agents in plaque could substantially contribute to the observed substantive effects in addition to their adsorption to oral surfaces.

Recently, research implied that an in vitro biofilm can absorb antimicrobials, followed by substantive release in bio-active concentrations [11]. This yields the question whether antibacterial agents can be absorbed in plaque-left-behind after toothbrushing with an antibacterial toothpaste as well to inhibit further growth of oral biofilm. Absorption of fluoride in plaque-left-behind, a much smaller molecule than most antibacterial agents, has been studied and demonstrated to aid caries prevention [12, 13], but the absorption in plaque-left-behind and release of antibacterial agents from a toothpaste in a bio-active concentration has never been studied.

In the meantime [14], we have demonstrated that plaqueleft-behind after brushing can absorb oral antibacterial agents from mouthrinses. By mixing control plaques from volunteers using a control toothpaste (without antibacterial claims) with plaques from volunteers using an antibacterial mouthrinse in addition to brushing with the control toothpaste, it could be demonstrated that these plaques contained residual, bio-available antibacterials that were released over a 2-h time period in sufficiently high concentrations to kill bacteria in control plaque [14].

In the present study, it was hypothesized that plaque-leftbehind after brushing may also act as a reservoir for antibacterial toothpaste components and therewith contribute towards their substantivity. Therefore we compare the contributions of plaque and of saliva to the prolonged substantivity of three antibacterial toothpastes versus a control toothpaste, without antibacterial claims as well as possible changes in bacterial composition of plaque and saliva after the use of the different toothpastes.



Toothpastes and recruitment of human volunteers

This unblinded clinical substantivity study was conducted in a parallel design and with exception of the Denaturing Gradient Gel Electropheresis (DGGE), entirely carried out by one professional dentist. For this study, four commercially purchased toothpastes, i.e., one control and three antibacterial toothpastes were selected, as listed in Table 1. The number of volunteers needed to complete the study was 74 (28 males, 46 females, age 19-29 years). They were all healthy dental and oral hygiene students recruited from the Center for Dentistry and Oral Hygiene at the University Medical Center Groningen. Six persons dropped out the study during use of the experimental toothpaste due to illness or not attending appointments. Each substantivity experiment consisted of two pairs of volunteers, one pair brushing for 2 weeks with the control toothpaste and the other pair brushing for 2 weeks with an antibacterial toothpaste and subsequently plaque and saliva samples were collected of all four volunteers (for more details see below). The order the volunteers came in was determinant for the allocation of the antibacterial toothpastes in alphabetical order over the volunteers (i.e., first Colgate Total® 5×, followed by Crest Pro Health® 5×, and Zendium Classic® 5×). The study was performed according to the guidelines of the Medical Ethics Committee of the University Medical Center Groningen, Groningen, The Netherlands, including the informed consent by the volunteers and the tenets of the Declaration of Helsinki.

Experimental protocol

The principle of the experimental set-up of the substantivity study for three antibacterial toothpastes versus one control toothpaste is schematically presented in Table 2. One experiment comprised two pairs of volunteers, one pair brushing for 2 weeks with the control toothpaste (viz. the control group) and the other pair for 2 weeks with an antibacterial toothpaste (viz. the experimental group). Volunteers were paired to minimize inter-individual differences. Mechanical cleaning, consisting of brushing and interdental cleaning, was done twice a day according to the habitual routine of the volunteers. Note that the control pair from one experiment acted as an experimental pair in the experiment 2 weeks later. After 2 weeks of brushing with the control or antibacterial toothpaste, plaque and saliva samples from each individual volunteer were collected 6 or 12 h after brushing in the morning by the volunteers themselves in the dental clinic. Before collection volunteers were carefully instructed by a professional dentist how to collect plaque from the buccal, lingual, palatal, and interproximal sides of the



Table 1 Toothpastes and their abbreviations as used in this study together with their main active components and manufacturer

Toothpaste	Abbreviation	Main active components	Manufacturer
Prodent Coolmint®	PC (control)	Sodium fluoride Sodium lauryl sulphate (SLS)	Sara Lee Household & Bodycare, Exton, USA.
Colgate Total®	CT	Triclosan Polyvinyl methylether maleic acid (PVM/MA)	Colgate-Palmolive Company, Piscataway, USA
		Sodium fluoride	
		SLS	
Crest Pro Health®	СРН	Stannous fluoride Sodium hexametaphosphate	Procter & Gamble, Cincinnati, USA
		SLS	
Zendium Classic®	ZC	Sodium fluoride Colostrum	Sara Lee Household & Bodycare, Exton, USA
		Lactoperoxidase	
		Lysozyme	
		Glucose oxidase	
		Amyloglucosidase	

dentition with a sterile cotton swab stick and a dental instrument [14]. This professional dentist also controlled compliance with the instructions during collection. Plaque was suspended in 2 ml sterile Reduced Transport Fluid [15]. In addition, approximately 2 ml unstimulated saliva was collected. All samples were vortexed and sonicated for

Table 2 General principle of the allocation of the volunteers to the control and three experimental toothpaste groups together with the brushing and collection time for the pairs of volunteers

Week	Volunteers brushing with	l			Plaque and saliva collection from volunteers		
	Control toothpaste Experimental toothpaste				Toothpaste		
	Prodent Coolmint (PC)	Colgate Total (CT)	Crest Pro Health (CPH)	Zendium Classic (ZC)	Control	Experimental	
1	AB						
2	CD						
3	EF	AB			AB		
4	GH	CD			CD		
5	IJ	EF			EF	AB	
6	KL	GH			GH	CD	
7	MN	IJ			IJ	EF	
8	OP		KL		KL	GH	
9	QR		MN		MN	IJ	
10	ST		OP		OP	KL	
11	UV		QR		QR	MN	
12	WX		ST		ST	OP	
13	YZ			UV	UV	QR	
14	ab			WX	WX	ST	
15	cd			YZ	YZ	UV	
16	ef			ab	ab	WX	
17	gh			cd	cd	YZ	
18					ef	ab	
19					gh	cd	

Each capital and small letter represents one volunteer



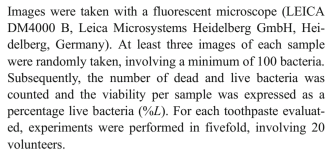
10 s at 30 W (Vibra Cell model 375, Sonics and Materials Inc., Danbury, CT, USA) in order to suspend bacterial clumps. Subsequently, both plaque and saliva samples of the individual volunteers in the control and experimental pairs were pooled. In total, four pooled samples were obtained in one experiment: a control (Pc) and an experimental plaque sample (Pe) and a control (Sc) and an experimental saliva sample (Se). These newly formed samples were studied with respect to their bacterial concentration and viability, immediately and 2 h after pooling of the pairs (for details, see below and Otten et al. [14]). For each antibacterial toothpaste, the experiment was done in fivefold, involving 5 pairs of volunteers per toothpaste (see Table 2) making a total of 20 volunteers.

In addition, control and experimental individual plaque samples from six volunteers were suspended after centrifugation in 0.5 ml TE buffer (10 mM Tris–HCl, pH 7.5, 1 mM EDTA) and were stored at -20° C for analysis of compositional similarity.

Analyses of plaque and saliva samples

The bacterial compositions of the plaques after use of the control or antibacterial toothpastes were compared using DGGE. After thawing, 0.5 ml plaque samples in TE buffer were centrifuged for 5 min at 13,000×g washed and vortexed with 200 µl TE. After DNA extraction, PCR was performed on 100 ng DNA with a T-gradient thermocycler. DGGE of PCR products generated with the F357GC/R518 primer set [16] was performed as described by Muyzer et al. [17]. The PCR products were applied on 8% (w/v) polyacrylamide gel in 0.5× TAE buffer (20 mM Tris acetate, 10 mM sodium acetate, 0.5 mM EDTA, pH 8.3). The denaturing gradient consisted of 30-80% denaturant (100% denaturant equals 7 M Urea and 37% formamide). A 10-ml stacking gel without denaturant was added on top. Electrophoresis was performed overnight at 120 V and 60°C. Gels were stained with silver nitrate [18]. DGGE gel images were converted and transferred into a microbial database with GelCompar II, version 6.1 (Applied Maths). The similarities in bacterial composition between control plaques and plaques collected after using an antibacterial toothpaste were analyzed using a ban based similarity coefficient (Dice). The clustering algorithm to calculate the dendograms was a non-weighted pair group method with arithmetic averages (UPGMA) [19].

The bacterial concentrations in pooled control and experimental plaque and saliva samples (Pc, Pe, Sc, and Se) were determined using a Bürker-Türk counting chamber and a phase contrast microscope (Olympus BH-2, Japan). Bacterial viability was determined by staining 20 µl of a sample with 80 µl LIVE/DEAD stain (*Bac*LightTM, Molecular Probes Europe BV) to distinguish dead and live bacteria.



If plaque acts as a reservoir for antibacterials, these should be able to kill bacteria in unexposed plaque (Pc). In order to evaluate the antibacterial activity in the Pe sample, a novel mixture technique as described by Otten et al. [14] was used. In short, a 1:1 mixture of the Pc and Pe samples was made, denoted as PcPe of which the bacterial viability was assessed 2 h after combining. A comparable mixture, denoted as PcSe, was made to evaluate the antibacterial activity in saliva 2 h after combining to allow sufficient time for antibacterial action on the untreated bacteria in the mixtures. After 2 h, the bacterial viabilities of the mixtures were experimentally determined in combined plaque and plaque-saliva samples and compared with theoretically expected viabilities of the mixtures at t=0. Since both Pe and Se exert immediate antibacterial effects on Pc if any, theoretically expected viabilities of both PcPe and PcSe were calculated based on the viabilities of Pc, Pe, and Se at t=0 using the following equations

$$\%L_{\text{PcPe}} = \frac{(\%L_{\text{Pc}} \times C_{\text{Pc}}) + (\%L_{\text{Pe}} \times C_{\text{Pe}})}{C_{\text{Pc}} + C_{\text{Pe}}}$$
(1)

$$\%L_{\text{PcSe}} = \frac{(\%L_{\text{Pc}} \times C_{\text{Pc}}) + (\%L_{\text{Se}} \times C_{\text{Se}})}{C_{\text{Pc}} + C_{\text{Se}}}$$
(2)

where $\%L_{Pc}$, $\%L_{Pe}$, $\%L_{Se}$, C_{Pc} , C_{Pe} , and C_{Se} represent the experimental viabilities and bacterial concentrations of Pc, Pe, and Se.

Experimental viabilities, lower than theoretically expected, are an indication that the experimental plaque or saliva still contained residual antibacterial activity, 6 or 12 h after the last brushing.

Statistical analysis

The data were analyzed using one way ANOVA (analysis of variance) to indicate significant differences between bacterial viabilities and concentrations of plaque and saliva samples after brushing with the different toothpastes. The Student's t test was used for statistical comparison between the control and antibacterial toothpastes. Values of p < 0.05 were considered to indicate statistically significant differences.



Results

The bacterial composition of the experimental plaques was influenced by the toothpaste used and the largest dissimilarities in bacterial composition with respect to the control paste were observed 6 h after use of CPH, as can be seen in Table 3. Interestingly, the similarity after use of CPH increased between 6 and 12 h after last use, while for CT the similarity decreased over time.

In general, plaques collected after the use of CT and CPH had lower viabilities than the control plaque, while plaques collected after the use of ZC showed similar viability as the control plaque (Table 4). There was no statistically significant difference between viabilities of plagues collected 6 or 12 h after brushing with the control paste, CT or CPH, but plaques collected 12 h after brushing with ZC were more viable than when collected 6 h after brushing (p < 0.05). Moreover, the concentration of bacteria in plaque decreased significantly between 6 and 12 h after brushing with ZC and PC (p < 0.05). The differences in viability assessed immediately or 2 h after collection were not statistically significant.

Only bacterial viabilities in saliva samples (Table 4) collected 12 h after brushing with CT (41% viability) were significantly different (p<0.05) compared to the control toothpaste (56% viability). Similarly, the bacterial viabilities of saliva collected 12 h after brushing with ZC (66% viability) were significantly different (p<0.05) from saliva collected 6 h after brushing with ZC (50% viability). Bacterial concentrations were not significantly different across the different saliva samples, with the exception of saliva collected 12 h after brushing with CPH, containing significantly (p < 0.05) less bacteria than the control saliva samples.

The experimental and theoretically expected viabilities of combined control and experimental plaques are compared in Table 5. For CT and ZC, experimental viabilities are smaller, respectively larger than theoretically expected, but these effects are not statistically significant. Experimental viabilities are significantly smaller (p < 0.05) than theoretically

Table 3 Similarities based on DGGE analysis in bacterial composition between control plagues, obtained during use of Prodent Coolmint® and experimental plaques, collected during the use of an antibacterial toothpaste

Collection	Similarity (%)		
time (h)	Colgate Total® (CT)	Crest Pro Health® (CPH)	Zendium Classic® (ZC)
6	77±7	60±14	75±7
12	67±6	67 ± 15	$71\!\pm\!8$

Volunteers (n=6) brushed for 2 weeks with a control toothpaste, followed by 2 weeks of brushing with an antibacterial toothpaste. Values are presented as averages±standard deviations for plaques collected 6 and 12 h after the last brushing

(C) in control (Pc) and experimental plaque (Pe) and control (Sc) and experimental saliva (Se) collected 6 and 12 h after brushing **Fable 4** Bacterial viability (%L) and concentration

Collection Evaluation	Evaluation	Control a	and experime	Collection Evaluation Control and experimental plaque						Control a	nd experim	Control and experimental saliva					
(II) arring	(III)	Prodent Cool (PC, control)	Coolmint® irol)	Colgate Tot	al® (CT)	Crest Pro Ha	ealth® (CPH)	Zendium C	lassic® (ZC)	Prodent Cooli (PC, control)	Coolmint® rol)	Colgate To	otal® (CT)	Crest Pro E	Prodent Coolmint® Colgate Total® (CT) Crest Pro Health® (CPH) Zendium Classic® (ZC) Prodent Coolmint® Colgate Total® (CT) Crest Pro Health® (CPH) Zendium Classic® (ZC) PC, control)	Zendium Cl	lassic® (ZC)
		$^{90}L_{ m Pc}$	$\frac{\text{%}^{-1}}{\text{%}^{-1}}$ $\frac{\text{%}^{-1}}{\text{(10^9/ml)}}$	$L_{ m Pe}$	C _{Pe} (10 ⁹ /ml)	$^{\prime\prime}$ CPe	C_{Pe} (10 ⁹ /ml)	$^{9/6}L_{ m Pe}$	$^{9/4}_{Pe} C_{Pe} $ (10 ⁹ /ml)	$^{\%}$	$\begin{array}{c c} \hline & & & & \\ \hline & \%L_{Sc} & & C_{Sc} & \\ \hline & & & & \\ \hline & & & & \\ \hline & & & & \\ \end{array}$	$^{9/6}L_{\mathrm{Se}}$ C_{Se} (10 $^{9}/\mathrm{ml}$)	C _{Se} (10 ⁹ /ml)	%T%	C _{Se} (10 ⁹ /ml)	%LSe	C _{Se} (10 ⁹ /ml)
9	0	51±16 45±14°	51±16 3.2±1.6 36±6 ^{a,b} 45±14 ^c 36±11 ^c	$36\pm6^{a,b}$ 36 ± 11^{c}	2.4±0.6	35±16 25±9ª,b,c	2.2±0.8 45±3 41±8°	45±3 41±8°	3.5±0.9	61±14 57±12	4.2±1.2	61±14 4.2±1.2 59±17 57±12 52±10	4.3±1.6 56±6 54±12	56±6 54±12	3.4±0.8	50±8 50±16	4.0±1.0
12	0 2	50 ± 15^{c} 48 ± 13	$2.5{\pm}1.1^{d}$	2.5 ± 1.1^{d} $29\pm19^{a,b,d}$ $29\pm13^{a,b}$	2.0±0.5	35±23° 37±14	2.0±0.8	$60\!\pm\!10^{\rm d,c} \\ 55\!\pm\!10^{\rm d}$	$1.6\pm0.8^{\mathrm{a,d}}$		4.3±1.8	$41\pm16^{a,c}$ 50 ± 17	4.7±2.5	60±9 c 58±6	$2.9\!\pm\!1.1^a$	66±11 ^{d,c} 62±7	5.3 ± 3.0

Bacterial viability was evaluated immediately after plaque collection and 2 h afterwards. Values are presented as averages ±standard deviation over 18 experiments for Prodent Coolmint® collected 6 h after brushing, 19 experiments for Prodent Coolmint® collected 12 h after brushing, and 5 experiments for the experimental products. Every experiment involves two volunteers. After the and 2 pairs dropped out of the 12 h study experimental toothpaste, 1

Statistically different from 6 h at p<0.05, Student's t test



Statistically different from PC (control) at p < 0.05, Student's t test

Significant difference between the different toothpastes at p < 0.05, ANOVA Statistically different from ZC at p<0.05, Student's t test

Table 5 Experimentally measured and theoretically expected bacterial viabilities of combined control and experimental plaques (PcPe) and control plaque and experimental saliva samples (PcSe) collected 6 or 12 h after brushing

Control plaque combined with Pe and Sc, using	$%L_{PcPe}$				$\%L_{ m PcSe}$				
with re and Sc, using	6 h after collec	etion	12 h after colle	ection	6 h after collec	etion	12 h after colle	ection	
	Experimental	Expected	Experimental	Expected	Experimental	Expected	Experimental	Expected	
Colgate Total® (CT)	36±8	44±17	31±21	37±11	55±12	53±10	43±7	47±14	
Crest Pro Health® (CPH)	$22 \!\pm\! 9^a$	35 ± 5	31 ± 5^a	46±6	50 ± 12	51 ± 7	56±7	56±3	
Zendium Classic® (ZC)	42±9	38 ± 7	57 ± 13	54 ± 8	46 ± 6	$42\!\pm\!11$	65 ± 12	60 ± 7	

Values are presented as average percentage live bacteria±standard deviation over 5 experiments, involving 20 volunteers per comparison

expected ones for CPH, indicating residual antibacterial activity in plaques left behind 6 and 12 h after brushing with CPH. Combining control plaques with experimental saliva's (Table 5) did not yield any effect on bacterial viability, indicating that there is no residual antibacterial activity due to toothpaste components left in saliva 6 or 12 h after brushing.

Discussion

Our hypothesis that plaque-left-behind after brushing may act as a reservoir for antibacterial toothpaste components and therewith contribute towards their substantivity is confirmed by the present results for a stannous fluoride containing antibacterial toothpaste (CPH). Both antibacterial toothpastes, CT and CPH, reduced bacterial viability in plaque up to 12 h after brushing, but only plaque-left-behind after brushing with CPH still contained significant residual antibacterial activity up to 12 h after brushing to cause prolonged killing of bacteria in unexposed plaque. Whether or not this reservoir function of the plaque-left-behind will contribute to an clinically significant reduction in caries and gingivitis remains to be determined, however.

The effects of plaque-left-behind after brushing with an antibacterial toothpaste have been demonstrated by comparison with a control toothpaste (see Table 4). The control toothpaste chosen was a regular sodium fluoride—SLS containing paste, without any antibacterial claims, similar to most other sodium fluoride—SLS containing pastes. This is not withstanding the fact that both fluoride and SLS may have antibacterial efficacy [8, 20]. Note that the viabilities of plaque samples collected after use of the control paste are comparable with the ones measured in vivo after the use of Crest Regular®, another sodium fluoride—SLS containing toothpaste without antibacterial claims [21]. Viabilities of plaque samples obtained after brushing with triclosan/copolymer containing CT and stannous fluoride/

hexametaphosphate containing CPH toothpastes, however, show reduced viabilities with respect to the control, in line with their known antibacterial efficacies in vivo and in vitro [6, 21–23]. The differences in compositional similarity of the different experimental plaques with respect to the control plaque suggest that the antibacterial components in the different pastes act on different strains and species.

Triclosan in toothpastes is often used in combination with a copolymer PVM/MA (polyvinyl methyl ether/maleic acid) [6, 24-25] and increases the permeability of the bacterial cell membrane, causing leakage of the cellular content, which finally leads to cell death [8]. Stannous fluoride (SnF₂) is recently used in toothpastes in combination with sodium hexametaphosphate to provide antiplaque [3, 26] and antigingivitis activity [3, 24]. Conflicting reports exists about the antibacterial efficacy of the enzyme containing toothpaste ZC, included in this study. Enzymes, like amyloglucosidase and glucose oxidase, are added to ZC toothpaste to enhance the host defense mechanisms [20]. Another ingredient added, supporting the host defense, is colostrum [27]. Whereas Tenovuo [27] concluded in a literature review that adding enzymes to a toothpaste improves the antibacterial capacity of saliva, in vivo the use of an enzyme containing toothpaste was unable to achieve differences in plaque and gingivitis scores as compared to a control toothpaste [7, 28]. Also in our study, we did not find an effect of the enzyme containing toothpaste on bacterial viabilities of plaques collected 6 or 12 h after brushing.

Despite the fact that both triclosan/copolymer containing CT and stannous fluoride/hexametaphosphate containing CPH toothpastes show reduced viabilities in plaque-left-behind after brushing, only plaques obtained after use of a stannous fluoride/hexamethaphosphate containing toothpaste still had sufficient residual antibacterial activity to cause significant bacterial killing (see Table 5). A triclosan/copolymer containing paste did not show such a contribution toward substantivity by plaque-left-behind in a statistically significant way, which probably suggests that the larger triclosan/copolymer molecules are



^a Statistically lower than theoretically expected values at p<0.05, Student's t test

not absorbed into plaque to the same extent as relatively small stannous fluoride molecules.

As might be expected from the lack of a direct effect of antibacterial agents on bacterial viability in saliva (Table 4), no contribution of saliva toward the prolonged substantivity of any of the antibacterial agents as delivered during toothbrushing was found (Table 5). Besides the antibacterial agents added to toothpastes, saliva exerts a continuous antibacterial effect [29]. The prolonged substantivity of chlorhexidine, accepted as the most effective oral antibacterial [8, 20], is generally attributed to weak adsorption followed by slow desorption to and from oral soft tissue surfaces to yield a prolonged effective antibacterial concentration in saliva [9]. Evidently, neither stannous fluoride nor the triclosan/copolymer combination adsorbs and/or desorbs in effective concentrations from the oral soft tissues into saliva.

In conclusion, plaque-left-behind after mechanical cleaning may contribute to the substantive action of an antibacterial toothpaste. Therewith, this study constitutes a pledge for the use of antibacterial toothpastes, since antibacterials can be absorbed in plaque-left-behind. The importance of antibacterials absorbed in plaque-left-behind is even amplified, when it is realized that plaque is predominantly left behind in places where its removal and effective killing matter most, i.e., in retention sites like fissures, interproximal spaces, gingival margins, or around orthodontic appliances that are hard to clean mechanically [2]. In this respect, it is worth mentioning that these conclusions are based on results obtained in a group of dental and oral hygiene students, who mostly have better skills to remove plague than the general public. In the general public, the amount of plaque-left-behind after brushing can be expected to be larger than in the current study, which may yield an even more pronounced contribution of plaque-left-behind toward the substantive action of antibacterial toothpaste components.

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Conflict of interest The authors declare that they have no conflict of interest.

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