A new chemical formulation for control of dental unit water line contamination: An 'in vitro' and clinical 'study'

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Abstract

Background: Water delivered by dental units during routine dental practice is highly contaminated. The aim of this study is to evaluate the efficacy of a new chemical solution flushed through Dental Unit Water Lines (DUWL) for the control of contamination inside dental units.

Materials and methods: Six old dental units equipped with a device designed to automatically flush disinfecting solutions through the water system (Castellini Autosteril) were selected. Water samples from DUWL effluents were collected in each dental unit for 10 randomly selected days, before and after a 5 minute DUWL disinfecting cycle with TetraAcetylEthileneDiamine (TAED) and persalt (Ster4spray produced by Farmec spa, and distributed by Castellini spa). Water samples were plated in R2A Agar and cultured at room temperature for 7 days, and the total number of heterotrophic microorganisms counted and expressed in Log10 CFU/mL. A general linear model was fitted and multiple regression ANOVA for repeated measures was used for the statistical analysis.

Results: The mean contamination in DUWL effluent at baseline was 5.45 ± 0.35 CFU/mL (range 4.79 to 5.93 CFU/mL). When water samples were tested "in vitro" against the chemical, no growth of heterotrophic bacteria was detected after a 5 minute contact in any of the water samples tested. After undergoing a 5 minute disinfecting cycle with the chemical, DUWL mean contamination in water effluents was 2.01 ± 0.32 CFU/mL (range 1.30 to 2.74 CFU/mL) (significant difference with respect to baseline).

Conclusions: An inbetween patient disinfecting procedure consisting of flushing DUWL with TAED and persalt equivalent to 0.26% peracetic acid could be useful in routine dental practice for cross-contamination control.
dental water to contain no more than 200 Colony Forming Units (CFU) /mL (2.3 in Log_{10} CFU/mL) of heterotrophic unfiltered output [4] Several methods have been suggested by which the DUWL contamination by heterotrophs might be kept under this limit and flushing protocols or chemical treatment are some of the options available to dentists [5,6].

Among chemicals now available, peracetic acid is one of the most powerful biocidal agent with a rapid and broad spectrum biocidal activity and could be a useful chemical for the purpose of controlling DUWL contamination [7,8], although, as delivered, it has a series of side-effects which have limited its use in dentistry [9–12].

In recent years, a new chemical formulation (TetraAcetylEthyleneDiamine in association with persalt) has been proposed as a non hazardous means of generating peracetic acid in situ in the absence of preformed peracetic acid side-effects [13,14].

The aim of this study was to evaluate the efficacy of the chemical to control DUWL bacterial contamination by heterotrophs both when tested "in vitro" and when flushed into DUWL.

Materials and methods

We selected for use in this study 6 dental units (Castellini Logos), all connected to municipal water and that had been in daily use for approximately 1 year. None of the selected units had ever been treated for removal of biofilm or reduction of planktonic bacteria. All dental units were equipped with a device designed to automatically flush disinfecting solutions through the water system (Castellini Autosteril).

a) DUWL contamination at baseline: a water sample (2 mls) was recovered in the morning before working for 10 randomly selected days from the high-speed handpiece line of each dental unit (60 samples). Samples were collected into a sterile tube added with filter-sterilized sodium thiosulphate at a final concentration of 18 µg/ml to oppose the growth-inhibiting effects of residual chlorine. Two split samples of each were made; the first split sample served to evaluate DUWL contamination at baseline.

b) DUWL contamination following "in vitro" contact with Ster4spray: the second split sample was used for the purpose. Ster4spray (produced by Farmed spa and distributed by Castellini spa Italy) is a fine powder containing a binary active system (TetraAcetylEthyleneDiamine and sodium perborate) which is activated by dissolving in water at a initial temperature of 35°C to form peracetyl ions at pH 8, equivalent to 0.26 % peracetic acid ensuring stable concentration up to 24 hours. The active is completely biodegradable and degrades to acetic acid, oxygen and water [14]. One part of each water sample was tested against nine parts of disinfectant. After five minute of contact time, one mL of the mixture was rapidly added to 9 mL of the recovery/neutralizer broth (3% polysorbate 80, 0.1% L-histidine, 0.3% lecithin, 0.5% sodium thiosulphate in phosphate buffer 25N) to prevent further inactivation taking place.

c) DUWL contamination after a disinfecting cycle with Ster4sprayflushed through DUWL by Autosteril: after collecting samples at baseline, each dental unit underwent (each morning for 10 days) a disinfecting cycle with Ster4spray consisting of flushing DUWLs with the disinfectant and washing it for 2 minutes after a 5 minute contact, and a further water sample was collected immediately after each cycle into a sterile tube added with filter-sterilized sodium thiosulphate.

Water samples (at baseline and after either "in vitro" disinfecting tests or "inside dental unit" disinfecting cycles) were plated in R2A Agar within 3 hours of collection and cultured at room temperature for 7 days, and the total number of heterotrophic microorganisms counted. All absolute counts were converted to Log_{10} values. This laboratory procedure has been recognized to be the best procedure to collect most of heterotrophic bacteria from DUWL [15–17].

A general linear model was fitted and multiple regression ANOVA for repeated measures was used to evaluate differences in CFU/mL between dental units, times (baseline and after disinfecting cycles) and the interaction of dental units x time; the Bonferroni t test was applied for significant values as a multiple-comparison t-test. The statistical analysis performed (multiple regression ANOVA for repeated measures by fitting a general linear model relating cfu to dental unit, disinfection and the interaction between dental unit x disinfection), not only allowed to evaluate any difference between cfu values before and after the disinfecting cycles, but also to evaluate any statistical difference in cfu between dental units and any difference between dental units in the cfu decreasing rate following the disinfecting cycles.

Results

a) DUWL mean contamination at baseline (in Log_{10}) was 5.45 ± .35 CFU/mL (range, 4.79 to 5.93 CFU/mL). Heterotrophic counts higher than 2.3 Log_{10} CFU/mL were found in all water samples. No significant difference in CFU/mL was found between dental units (F = 2.17; NS).

b) No growth of heterotrophic bacteria was detected in any of the water samples tested "in vitro" with Ster4spray.
c) DUWL mean contamination (in Log_{10}) in water effluents from dental units which had undergone disinfecting cycles with Ster4spray was 2.01 ± 0.32 CFU/ML (range 1.30 to 2.74 CFU/ML). Heterotrophic counts higher than 2.3 Log_{10} CFU/ML were only found in 10% of water samples. The difference in CFU/ML between values obtained after dental units had undergone disinfecting cycles and values at baseline was highly significant (F = 178.8; p < .01). No significant difference in CFU/ML decreasing rate after the disinfecting cycles was found between dental units (F = 2.18; NS).

Discussion

Water effluents from DUWL are highly populated in routine dental practice by heterotrophic bacteria principally originating from municipal water piped to the dental unit. Human pathogens, sucked back into the lines during dental procedures due to inadequate anti-retraction on dental units, have also been reported in some studies [3,18,19]. Both oral pathogens and heterotrophic bacteria can be responsible for severe diseases, and immune-compromised patients in particular may be at high risk [20].

As far as heterotrophic bacteria are concerned, the results of this study confirmed that DUWL are highly contaminated when dental units in use for several months receive no decontaminating treatment. According to our data, the great amount of DUWL contamination did not greatly differ from a dental unit to another and none of the water samples collected at the beginning of randomly selected working days reached CFU/ml values below the limit imposed by ADA for the year 2000 [4].

Interesting results have been obtained by testing the water samples against peracetic acid delivered by the chemical solution. After 5 minute of contact no growth of heterotrophic bacteria was found in any of 60 water samples tested. These results are in agreement with other reports showing that preformed peracetic acid possess a very rapid and broad spectrum microbicidal activity together with a very good activity against biofilm in waterlines used for haemodialysis [8–12].

However, despite these interesting properties, preformed peracetic acid has not been so far utilized in any study in dentistry with the aim to control DUWL contamination. Effectively, as delivered, preformed peracetic acid is unstable, potentially explosive, highly acidic and as a consequence highly corrosive. These properties make products containing preformed peracetic acid difficult to formulate for long term storage stability and difficult to handle and transport so limiting the use of this product in dentistry [9–11].

In the recent years, TAED with peroxygen source at near neutral pH has been clamed to provide a nonhazardous means of generating peracetic acid in situ, in the absence of the preformed peracetic acid side-effects [14].

Preliminary data from a previous study showed the relevant biocidal "in vitro" activity of the test formulation against human pathogens including spores, and data from the present study underline its great efficacy against heterotrophic bacteria both when tested "in vitro" and when flushed into DUWL [13].

In fact, DUWL flushing with the chemical solution left standing into DUWL for 5 minutes provided a good control of DUWL in all dental units studied, the mean CFU/ml values being well lower than the limit imposed from ADA, and values higher than 2.3 Log_{10} CFU/ML being only detected in 6 out of 60 samples.

These data are in agreement with other studies which have obtained similar results, but adopting different disinfecting procedure [21–23], consisting of introducing chemicals into water systems either continuously or intermittently during working pauses [24–28].

Although the mentioned treatments offer less potential for recolonization of waterlines since they keep into contact the chemicals with DUWL for longer periods of time, most of these agents can not probably ensure a rapid killing of viruses and bacteria eventually sucked back during dental procedures, since they are used at very low concentration in the continuous waterline supply or during working pauses [3].

Instead, we may speculate that intermittent between-patient treatment regimens using potentially biocidal concentrations of germicide, besides keeping low level of heterotrophic bacterial counts during dental procedures, could be also effective in eliminating oral pathogens eventually aspirated from patients under dental treatment and spread out during next procedures.

For this purpose the chemical formulation tested in this study might be very useful when used between patients in combination with dental units which incorporate the capacity to disinfect DUWL by the automatic flushing of lines.

Conclusions

i) Dental unit waterlines are highly contaminated during dental procedures and aerosols generated by dental instruments are a possible source of infection;

ii) The between-patient flushing of dental unit waterlines with disinfecting solutions with strong and rapid biocidal
activity could be very effective in controlling microbial contamination in water effluents during dental procedures.

**Competing interests**
None declared

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