Splatters and Aerosols Contamination in Dental Aerosol Generating Procedures

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Abstract: Dental aerosol-generating procedures produce a large amount of splatters and aerosols that create a major concern for airborne disease transmission, such as COVID-19. This study established a method to visualise splatter and aerosol contamination by common dental instrumentation, namely ultrasonic scaling, air-water spray, high-speed and low-speed handpieces. Mock dental procedures were performed on a mannequin model, containing teeth in a typodont and a phantom head, using irrigation water containing fluorescein dye as a tracer. Filter papers were placed in 10 different locations to collect splatters and aerosols, at distances ranging from 20 to 120 cm from the source. All four types of dental equipment produced contamination from splatters and aerosols. At 120 cm away from the source, the high-speed handpiece generated the greatest amount and size (656 ± 551 µm) of splatter particles, while the triplex syringe generated the largest amount of aerosols (particle size: 1.73 ± 2.23 µm). Of note, the low-speed handpiece produced the least amount and size (260 ± 142 µm) of splatter particles and the least amount of aerosols (particle size: 4.47 ± 5.92 µm) at 120 cm. All four dental AGPs produce contamination from droplets and aerosols, with different patterns of distribution. This simple model provides a method to test various preventive strategies to reduce risks from splatter and aerosols.

Keywords: dental aerosol-generating procedures; COVID-19; splatter; droplets; aerosols; dental procedures

1. Introduction

The generation of aerosol and splatter creates a major risk for airborne contamination within the clinical environment [1–4]. Most routine dental treatments are aerosol-generating procedures (AGPs) that produce a mixture of splatter, droplets and aerosols that contain saliva, blood, irrigant water, and viable microorganisms (including bacteria and viruses) [1,5]. Commonly used dental instruments, including dental handpieces, air-water (triplex) syringes, and ultrasonic scalers all generate a large volume of splatter and aerosols, that are derived from patient fluids and coolant water, and that pose a risk to dental professionals and patients [1,5,6]. In order to assess the benefit of any protective methods or devices, it is first necessary to visualise very small amounts of contamination from splatter and aerosols during dental procedures. Such microdroplets are invisible and hence special methods are needed to visualise them and to map their spatial distribution within the working environment, and hence develop better ways to mitigate the risk of disease transmission.

There is an ongoing discussion in the literature on how to best define aerosols, droplets and splatter [7,8]. For the present study, splatter was defined as large liquid particles with the size ≥ 100 µm [9], whereas, droplets are medium size particles in the range of 5–100 µm [10], and aerosols are ≤ 5 µm [11,12]. The splattered fluid is large airborne
particles, and these fall onto surfaces in the dental operatory under the influence of gravity, following a ballistic trajectory from the point of origin. In contrast, droplets can remain suspended in the air until their water evaporates, and aerosols can remain suspended for several hours (depending on local air movement) and can travel for 1–3 m from their source [13].

Dental splatter, droplets and aerosols may contain pathogenic oral bacteria, nanoparticles [6,14–16], as well as respiratory viruses (including adenoviruses, influenza viruses, and the SARS-CoV-2 coronavirus [6,17]). Much greater attention has been focussed on dental aerosol-generating procedures (AGPs) because of the coronavirus disease 2019 outbreak (COVID-19), caused by the Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) virus. In some cases, especially when people are in close proximity to one another, COVID-19 has been proven to spread through airborne transmission [18,19]. The SARS-CoV-2 coronavirus can remain infectious in aerosols for extended periods, even when the water has evaporated, and particles that then settle onto surfaces can remain infectious for up to 72 h [20,21]. The airborne transmission of SARS-CoV-2, via salivary bioaerosols, poses a significant danger to healthcare workers that operate close to the face and oral cavities, such as dental practitioners, and oral and maxillofacial surgeons, especially when carrying out AGPs [5,19,22].

It is of considerable interest to have a straightforward method to visualise and map the pattern of splatter/droplets/aerosols during AGPs. Past work has focussed mainly on the microbiological evaluation of bacteria contamination from aerosols and splatter following dental procedures, by counting aerobic bacteria colonies deposited onto agar plates positioned at various locations [23,24]. Other studies have used fluorescent tracers (such as sodium fluorescein) to show in broad terms the distribution of ejected material [13,25]. These methods have been used to show macro-spatial distributions of material produced by ultrasonic scalers [13], or other devices such as the high-speed air-turbine, or the triplex syringe [25]. To date, there is limited data that determined the distribution of aerosols and splatter at the microscale for the most commonly used devices, namely the ultrasonic scaler, triplex syringe, high-speed headpiece, and low-speed headpiece.

The aim of the current study was to map the microparticle distribution of splatter and aerosol contamination from routine dental procedures at defined distances using a fluorescent tracer, combined with fluorescence image analysis. The intention was to develop a model system for visualising the contamination from aerosols and splatter as a prerequisite to assessing potential mitigating measures that could reduce the extent of contamination during dental procedures.

2. Materials and Methods
2.1. Experimental Setup

The experiment was carried out in a 20 m² PC2 laboratory. The mock dental procedures were performed on the left mandibular incisor tooth (tooth 31, FDI World Dental Federation notation), on a Columbia phantom head mannequin containing typodont model teeth in both jaws. Sodium fluorescein (Sigma-Aldrich, St Louis, MO, USA) was added into the coolant water to give a final concentration of 10 mg/mL. The following types of dental equipment were used: ultrasonic piezoelectric dental scaler (Dentsply Cavitron, Charlotte, NC, USA), air-water spray triplex syringe in a portable dental cart (Forest™, Hillsboro, OR, USA), a high-speed air turbine handpiece (300,000 rpm) fitted with a tapered fissure bur (TiMax, NSK, Kanuma, Japan), and a low-speed air turbine handpiece (W&H implantMED™, Bürmoos, Austria) (1200 rpm) fitted with a round #5 dental bur. Representative images for each AGP are shown in Figure 1a. For the ultrasonic scaler and low-speed handpiece, the procedures were carried out for 15 s and 5 min, respectively. For the high-speed handpiece and triplex syringe, the procedures were performed for 5 s and 15 s, respectively. Water flow rates were adjusted as per the manufacturer’s recommendations. The irrigation flow rate was 40 mL/min for the ultrasonic scaler, 140 mL/min
for the triplex syringe, 15 mL/min for the high-speed handpiece, and 15 mL/min for the low-speed handpiece.

Figure 1. Representative images of liquid particles and experimental design for liquid particle collection. (a) Examples of the position of dental instruments. The fluorescein tracer is seen as green because of illumination with 405 nm violet light using an LED torch. The liquid generated by the ultrasonic scaler, triplex syringe, high-speed handpiece and low-speed handpiece can be seen. (b) A schematic diagram of the positions for splatter and aerosol collection at 10 different locations (No. 1–10). The angular positions in degrees were calculated relative to the sagittal plane of the mannequin.

Before each procedure, pieces of filter paper (retention size 2.00 µm, Filtech), with a size of 160 (length) × 100 mm (width), were placed in 10 different locations (refer to No. 1–10 in Figures 2–4), as illustrated in Figure 1b. To visualise the splatter pattern, the filter papers were visualised in a Gel Doc™ EZ Imager (Bio-Rad Laboratories Inc, USA). The aerosols retained in the filter paper fibres were imaged using a Leica DMi8 inverted fluorescent microscope (Leica Microsystems, Japan). Each experiment at each location was repeated four times. To prevent potential contamination between procedures, the whole area was cleaned between runs and 70% ethanol was used to wipe down surfaces.
Figure 2. (a) Representative splatter images; (b) mean fluorescence intensity across 10 locations (after images were converted into grayscale); (c) particle distributions at location 10 after 15 s use of the ultrasonic scaler, triplex syringe, high-speed handpiece and low-speed handpiece.
scaler and low-speed handpiece generated notable splatter contamination only at locations 4, 5 and 7 after 5 min of use (Figure 4).

Figure 3. Splatter pattern by high-speed handpiece and triplex syringe after 5 s.

Figure 4. The splatter pattern by ultrasonic scaler and low-speed handpiece after 5 min surgeries.

3.1. Presence of Aerosols, 120 cm away from the Source

The aerosol particles that were retained on the filter paper fibres were imaged at location 10 (120 cm away from the source) after 15 s procedures (Figure 5a). Small aerosol particles (median particle diameter < 5 μm) were detected from all four equipment types (Figure 5a). The triplex syringe generated the aerosols with the smallest particle size (median diameter: 1.09 μm, versus 1.38 μm for both the ultrasonic scaler and the high-speed

2.2. Statistical Analysis

Images at each location were taken from four independent experiments and analysed using Fiji-ImageJ software to measure the diameter of the splatter, droplet and aerosol. For the calculation of the fluorescent value using Fiji (Figure 2b), each pixel of each fluorescent image was converted to grayscale (0–255; where 255 is the highest gray value) and the mean gray value was determined by the sum of the gray values of all the pixels divided by the number of
pixels. Using GraphPad Prism software (version 9.0.0), one-way ANOVA was used to assess differences between groups, with $p$ values $<0.05$ considered statistically different.

3. Results

3.1. Splatter Distribution Generated during 15 s Mock Procedures

After 15 s procedures on the mandibular central incisor tooth, all four types of equipment generated liquid contamination at the various locations in the vicinity of the equipment (Figure 2). The high-speed air turbine handpiece generated the most liquid particles and contaminated all 10 collection locations, including location 10 which was 120 cm away from the patient (Figure 2a). After quantification of fluorescent tracer intensity, the fluorescent images were converted to grayscale in Fiji and the data showed that the high-speed handpiece showed the greatest values, being significantly higher than the other three types of equipment, at all 10 locations (Figure 2b).

The distribution of splatter particles at the furthest location (No. 10) was assessed in greater detail, focussing on the size and frequency of the splatter particles. The data demonstrated that the ultrasonic scaler and low-speed handpiece generated fluid splashes with smaller size particles that were less than 300 $\mu$m (median diameter). In contrast, the triplex syringe and the high-speed handpiece generated larger particles ($>550 \mu$m median diameter). Most importantly, across four independent experiments, the high-speed handpiece gave the most abundant splatter contamination (>32,000 particles were analysed), while the ultrasonic scaler and low-speed handpiece caused less contamination (58 and 80 particles, respectively) (Figure 2c).

Furthermore, splatter particles were detected after using the high-speed handpiece and the triplex syringe for 5 s, especially for the high-speed handpiece, which contaminated all 10 locations even after 5 s of use (Figure 3). On the other hand, the ultrasonic scaler and low-speed handpiece generated notable splatter contamination only at locations 4, 5 and 7 after 5 min of use (Figure 4).

3.2. Presence of Aerosols, 120 cm away from the Source

The aerosol particles that were retained on the filter paper fibres were imaged at location 10 (120 cm away from the source) after 15 s procedures (Figure 5a). Small aerosol particles (median particle diameter $<5 \mu$m) were detected from all four equipment types (Figure 5a). The triplex syringe generated the aerosols with the smallest particle size (median diameter: 1.09 $\mu$m, versus 1.38 $\mu$m for both the ultrasonic scaler and the high-speed handpiece). The low-speed handpiece gave the largest aerosol particles (median diameter 1.75 $\mu$m) (Figure 5b). In all four independent experiments, the triplex syringe generated the most aerosol particles (>7100 particles) while the low-speed handpiece produced the least number of particles (311 particles) (Figure 5b).
handpiece). The low-speed handpiece gave the largest aerosol particles (median diameter 1.75 μm) (Figure 5b). In all four independent experiments, the triplex syringe generated the most aerosol particles (> 7100 particles) while the low-speed handpiece produced the least number of particles (311 particles) (Figure 5b).

Figure 5. Detection of aerosols and droplets retained in filter paper fibres, at the furthest point from the source. (a) Representative fluorescent images of aerosols and droplets generated by four dental procedures. Yellow arrow: paper fibres; white arrow: aerosols/droplets. (b) The distribution of aerosols and droplets at location 10.

The aerosol particles were also imaged (Figure 6) at location 2 for the ultrasonic scaler and the triplex syringe (after 15 s), at location 4 for the low-speed (15 s, 5 min), and at location 10 for the low-speed handpiece and the triplex syringe (5 s). These images (Figure 6a,c,e) show that small aerosolised particles (<5 μm) were detected in these locations from various equipment, with similar median particle sizes (ranging from 1.01 to 1.38 μm, Figure 6b,d,f).
3.1. Comparison of Splatter and Aerosol Particle Size at Location 10

Figure 6. The visualisation (a,c,e) and histogram distributions (b,d,f) of aerosols particles on the filter paper fibres. White arrow: aerosol particles; yellow arrow: paper fibres.
3.3. Comparison of Splatter and Aerosol Particle Size at Location 10

The average diameter of splatter and aerosol particles at the furthest point was analysed (Figure 7). The high-speed handpiece generated significantly larger splatter particles (mean ± SD: 656 ± 551 µm, ranging from 257 to 3575 µm) than the other three types of dental equipment (Figure 7a). The splatter particles were comparable in size between the ultrasonic scaler (281 ± 188 µm, ranging from 200 to 1020 µm) and the low-speed handpiece (260 ± 142 µm, ranging from 257 to 3497 µm), while both the triplex syringe (566 ± 405 µm, ranging from 200 to 1240 µm) and the high-speed handpiece gave significantly larger splatter compared to the ultrasonic scaler (Figure 7a).

![Figure 7. Average diameter of splatter (a) and aerosols/droplets (b) particles by four aerosol-generating procedures (AGPs) at location 10. (a) Different dental AGPs generated different splatter particles at a distance 120 cm away from the source. ns: no significant difference. ***: p < 0.0002 vs. ultrasonic scaler; ****: p < 0.0002 vs. ultrasonic scaler. ###: p < 0.0002 vs. high-speed handpiece. (b) Aerosol particles difference between different AGPs. ****: p < 0.0002 vs. triplex syringe. ###: p < 0.0002 vs. low-speed handpiece.](image)

In terms of aerosols, the low-speed handpiece generated the largest aerosol particles that were retained on the filter paper fibres (mean ± SD: 4.47 ± 5.92 µm, ranging from 0.688 to 27.2 µm), while the triplex syringe produced the smallest aerosol particles (1.73 ± 2.23 µm, ranging from 0.688 to 29.15 µm) (Figure 7b).

4. Discussion

The present study provides a useful simple model for assessing the microscale distribution of liquid particles generated by four common dental devices. The high-speed handpiece generated the most and the largest splatter particles (>600 µm) at 1.2 m away from the source. Furthermore, aerosols (<5 µm) and droplets (<30 µm) were detected at 1.2 m away from the source, with the triplex syringe producing the greatest number of the smallest diameter aerosol particles.

The airborne spreading of splattered fluid, droplets and aerosols is a major concern in dental practice because of the risk of transmitting potentially pathogenic bacteria (such as *Legionella pneumophila* or *Pseudomonas aeruginosa*) and both oral and respiratory
viruses [2,3,26,27], including human influenza viruses and SARS-CoV-2. Visualising and mapping the distribution of these liquid particles is important to develop preventive strategies. There is limited data available on the microscale distribution of contamination induced by aerosol-generating devices. Previous studies used fluorescein tracers [13,25] or bacterial tracers [23] to detect “bio-aerosol” contamination at up to 2 m. However, different studies have used different definitions for aerosol and splatter particles: Allison et al. defined aerosol particles as ≤ 10 µm and splatter as > 10 µm in diameter [25], while Leung et al. used a cutoff value for droplets and aerosols of 5 µm (>5 µm—droplets; ≤5 µm—aerosols) [28], in the same manner as the present investigation.

The current study used a fluorescence gel imager and a microscope to capture images of splatter (≥100 µm), droplets (5–100 µm) and aerosol (≤5 µm). Such particles were found to be generated by the ultrasonic scaler, triplex syringe, high-speed handpiece, and low-speed handpiece. All four AGPs generated considerable contamination of the patient’s chest area (locations 4, 5, and 6), which is consistent with past studies with macroscale contamination pattern [23,25]. This patient chest contamination can be effectively dealt with using a disposable waterproof (plastic-backed) bib or apron for the patient.

The present data demonstrate that the high-speed handpiece generates the most splatter of large particles (>600 µm in diameter) and that this contamination extends a considerable distance, at least 1.2 m away from the mouth. This pattern is also consistent with past work with macroscale contamination [13,23,25,29].

Small aerosols particles (≤5 µm) are produced by several dental procedures, and these may pose an increased risk of transmitting respiratory infections, such as COVID-19 [6,30,31]. According to the World Health Organisation (WHO) definition, airborne aerosol particles that are < 5 µm in size and are generated by AGPs can remain in the air, and then travel over some distance to then cause infection if they are inhaled [12]. Our study utilised a fluorescent microscope to image the aerosol and droplet particles retained on the filter paper fibres in collection sites at 1.2 m away from the source. Both aerosol and droplet particles (0.6–30 µm in diameter) were generated by all four AGPs, with the triplex syringe generating the most aerosol and droplet particles, while the low-speed handpiece generated the least aerosol particles and the largest (4.47 ± 5.92 µm) aerosol particles. This indicates that all four dental AGPs need to be carefully considered in the context of planning risk-based additional precautions to prevent potential airborne transmission.

The limitation of our study is that splatter images taken by the Gel Doc imager used a relatively large size of filter paper (160 × 100 mm), and the paper mesh size gave a limited resolution (200 µm). Better resolution equipment is required to capture all the splatter and droplet particles. Another limitation of this study was that the water flow rate for high-speed and low-speed handpieces could be increased from 15 mL/min (in this study) to 50 mL/min (to mimic the real-world dental clinic) for the future studies. Furthermore, this study did not apply a high volume suction system during dental AGPs, which will be investigated as a preventative measure in future research. Notwithstanding the limitations of our study, it provides a quantitative method for visualising liquid particle contamination generated during dental AGPs. As the method demonstrates that the high-speed handpiece generated the most liquid contamination (even after 5 s), the use of this item of equipment should be minimised in the clinic during the COVID-19 pandemic at times when there is community transmission. However, further research in an actual clinical environment is required to validate potential preventive strategies to mitigate airborne transmission. Testing dental high volume suction would be useful as a direction for further work using this same model. In the same manner, the model could be used to test tracers such as coloured dyes or nonharmful bacteria to show their movement. Given that saliva is more viscous than water, the laboratory model could also be used to explore how changes in the fluid density and viscosity influence the production of aerosol and splatter.
5. Conclusions

This study established a method to visualise the contamination patterns of splatters, droplets and aerosols, generated by four dental AGPs. The data demonstrated that all four dental procedures generate splatter, droplets and aerosols contamination, at 120 cm away from the source. Among four dental AGPs, high-speed headpiece generated the most contamination. Precautions for reducing aerosol, droplet and splatter contamination from dental AGPs should be used in the treatment of all dental patients at all times.

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