Investigation of droplets released during digestive endoscopy using a high-speed camera (with video): a pilot study

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Abstract
Background A large release of droplets is often expected around the periphery of the digestive endoscope insertion site. Therefore, a sense of alarm over infection because of droplets that may be released during digestive endoscopy examination is increasing. This study aimed to investigate the droplets released during digestive endoscopy using a high-speed camera.

Methods We utilized a high-speed camera (FASTCAM SA-3, Photron Limited) capable of recording small, transparent droplets with a black background and high-brightness lighting. The obtained video files were analyzed using post-processing software. We divided the 20 models into the control (a spray bottle model and a cough model) and experimental groups (digestive endoscopy models). The sedative, proficiency of digestive endoscopy and the amount of gas injected were modulated to change the level of released droplets.

Results For the control groups, droplets were clearly observed using a high-speed camera. However, no droplet larger than 10 µm in size was observed in the experimental groups. Furthermore, the changes in the sedative, proficiency of digestive endoscopy and the amount of gas injected did not affect droplet formation.

Conclusions Based on high-speed camera photography, the risk of droplet generation during digestive endoscopy was not higher than that during violent expiratory events, such as coughing and sneezing.

Keywords Droplets · Endoscopy · COVID-19 · Coronavirus · High-speed camera

The coronavirus disease (COVID-19) outbreak, which began in November 2019, has rapidly spread worldwide [1]. Droplets released from the mouth of a COVID-19-infected patient are an important source of infection. Droplets released from an infected patient’s sneeze are capable of traveling a distance of over 2 m [2–4].

It is thought that a large release of droplets is expected around the periphery of the digestive endoscope insertion site. Therefore, a sense of alarm over infection through droplets that may be released during digestive endoscopy examination is increasing [5]. The epidemic prevention guidelines for digestive endoscopy against COVID-19 are set around a standard of the 2-m travel distance of droplets released from a normal sneezing patient and have no clear standard of the travel distance of droplets [6]. Therefore, most medical teams working in the digestive endoscopy room wear N95 masks and face shields, goggles, level-3 protective suits, and gloves, while minimizing the number of endoscopies performed. However, because of the excessive wearing of protective suiting, the accuracy of endoscopies is decreasing, while the procedure-related fatigue levels are increasing. Furthermore, as the duration of the procedure increases, the number of endoscopies performed decreases, which has a negative impact on patients.

Therefore, the aim of this study was to analyze the travel distance and range of droplets released during digestive endoscopy using a high-speed camera to aid in the development of epidemic prevention guidelines related to digestive endoscopy.
Materials and methods

Study population and ethical considerations

From May 2020 to June 2020 we investigated droplets from 20 models. The patients enrolled in the study and partaking in digestive endoscopy had no severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infection at 14 days prior to digestive endoscopy. Their age ranged from 21 to 75 years, with no history of traveling to or residing in countries at risk. The exclusion criteria were as follows: respiratory symptoms (coughing and difficulty breathing), fever, history of traveling to or residing in countries at risk without reporting SARS-CoV-2 infection at 14 days prior to the examination, swallowing impediments, the presence of aspiration pneumonia, and inability to undergo gastrointestinal endoscopic examinations. Regarding the droplet generated during digestive endoscopy, as our aim was to analyze only the droplets released through the gastrointestinal tract, patients with respiratory disease who could release sputum did not participate. This study was approved by the Boramae Hospital Institutional Review Board (IRB) (IRB number 07–2020-17). Informed consent was obtained from the patients.

Five spray bottle models, five normal cough models, and 10 digestive endoscopy examinees were analyzed in this study. The following subcases were constructed with 20 models: case A [spray bottles (n = 5)]; case B [cough models (n = 5)]; cases C, D, and E [gastroscopy (EGD) models (n = 5)]; and cases F, G, and H [colonoscopy (CFS) models (n = 5)] (Fig. 1).
Details of the measurement of the droplets

The high-speed camera’s photographs show in slow motion instantaneous phenomena, which cannot be photographed with a normal video or film camera, to aid in thoroughly confirming and understanding rapid forms with the naked eye. The movement of an arrow and the migration of a missile can be photographed, and even the evaporation of water and movement of precision explosives can be saved. As the droplets generated during digestive endoscopy are small, transparent, and move at a rapid pace, they are difficult to be confirmed with a naked eye or with normal cameras. Thus, we used a high-speed camera (FASTCAM SA-Z; Photron Limited, Tokyo, Japan), which can photograph 1000 frames/s, and it was calibrated to distinguish particle sizes of 20 µm (Fig. 2a). The measurement targets were the size of the droplet, travel distance, emission angle, and emission speed. For droplet determination, the examinee’s oral cavity and anal periphery were filmed.

To film transparent droplets, a black background and high-brightness lighting were used in filming (Fig. 2b). To confirm the existence of the droplet on the high-speed camera, a coughing patient model in the state of not wearing the mouthpiece and the spray bottle model were used as the control groups. These groups (spray bottles and cough models) were compared with the droplet release photography of the experimental groups (digestive endoscopy models). Five patients undergoing EGD and five patients undergoing CSF participated in the control groups for the droplet release examination. To minimize stress of the participants,
blindfolds were used. The high-speed film footage was transferred to a computer, and the images were processed before analyzing the existence of droplets, which had an average duration of 1 h. Droplet generation was adjusted based on the use of sedatives, endoscopy procedure experience, and the amount of gas injected, and the analysis was performed, focusing on the insertion and removal points of the endoscope, which was expected to produce the most droplet generation.

Results

In the control groups (case A) of the spray bottle and cough models (case B), droplet particles were clearly observed. In the spray bottle model, the moving droplets were relatively uniform in size and direction. The droplet spread out within the radiation angle range of approximately 60 degrees, and the diameter of the droplet was 200–1000 µm. Droplets moving according to the movement of the tongue and lips were observed in the cough model. In the cough model, the droplets spread within the radial angle range of approximately 105 degrees, and the diameter of the droplets ranged from 50 to 1000 µm. However, in the EGD and CSF examinations, regardless of the use of sedatives, endoscopy proficiency, and amount of gas injected, no droplets moved parallel to the plane of the endoscope around the endoscope insertion site periphery (Fig. 3a–c). During EGD, saliva flowed down the sides of the mouth, and in CSF, intestinal fluids flowed down around the anus. Figure 1 and Video 1 demonstrate droplets from three representative models.

Discussion

Most of the previous studies on droplets reported to date have mostly focused on cough models for respiratory diseases [7, 8]. Recently, we introduced a brief report on droplets released during digestive endoscopy [9]. As the participants’ tongue and mouth were not restrained in the cough model, the saliva underneath the tongue moved freely. Furthermore, in the case of respiratory disease, phlegm and saliva production increased and, therefore, droplets could travel more than 2 m. However, during EGD, as the mouthpiece and endoscope are inserted into the oral cavity, secretions from inside the oral cavity only trickle down in the direction of gravity, and the probability of the secretions being emitted in droplet form along the plane of the endoscope was very low. During CSF, as the endoscope was sealed by the anus, no droplets were observed, and the intestinal fluids only flowed down around the anus.

Currently, preventative guidelines for digestive endoscopy have been developed based on cough models with droplet travel distances > 2 m. However, in high-speed photography during digestive endoscopy, no droplets moving parallel to the endoscope plane were confirmed. In our small-scale pilot study, the risk of droplet generation during digestive endoscopy was not higher than that during violent expiratory events, such as coughing and sneezing. Therefore, infection-related sections in the current COVID-19-related digestive endoscopy guidelines should be maintained, but it seems beneficial to revise the content related to droplets occurring at the entry point of the endoscope.
This study had some limitations. First, much time was spent on the high-speed camera post-processing procedure and, therefore, the number of study participants was low. Second, COVID-19-infected patients were excluded from the study. Therefore, further large-scale studies are required to overcome these limitations. Third, there was a limitation when measuring the exact particle size because the droplets move under the influence of the particle size, spray direction, gravity, and the flow of the surrounding atmosphere. In the case of the spray bottle model, the size of the nozzle is different for each manufacturer. Moreover, in the case of the cough model, the amount and size of droplets released may vary depending on the patient. In addition, as the movement direction of the particles may be different depending on the air circulation and gravity in the measurement room, it is necessary to record from three directions to measure the movement direction of the droplet. In a future research, we will consider shooting with special dyes for shooting to make accurate particle size measurements.

In conclusion, based on high-speed camera photography, no droplets were observed during digestive endoscopy. The mouthpiece and anus play a role in blocking the release of droplets at the entry site of the endoscope. We believe that the risk of droplet generation during digestive endoscopy was not higher than that of violent expiratory events, such as coughing and sneezing. This study’s findings can be effective for the development of preventative guidelines against COVID-19 in digestive endoscopy.

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Declarations

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