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<th>MS2 virus inactivation by atmospheric-pressure cold plasma using different gas carriers and power levels</th>
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In this study, airborne MS2 bacteriophages were exposed for subsecond time intervals to atmospheric-pressure cold plasma (APCP) produced using different power levels (20, 24, and 28 W) and gas carriers (ambient air, Ar-O2 [2%, vol/vol], and He-O2 [2%, vol/vol]). In addition, waterborne MS2 viruses were directly subjected to the APCP treatment for up to 3 min. MS2 viruses with and without the APCP exposure were examined by scanning electron microscopy (SEM), reverse transcription-PCR (RT-PCR), and sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). Viral inactivation was shown to exhibit linear relationships with the APCP generation power and exposure time ($R^2 > 0.95$ for all energy levels tested) up to 95% inactivation (1.3-log reduction) after a subsecond airborne exposure at 28 W; about the same inactivation level was achieved for waterborne viruses with an exposure time of less than 1 min. A larger amount of reactive oxygen species (ROS), such as atomic oxygen, in APCP was detected for a higher generation power with Ar-O2 and He-O2 gas carriers. SEM images, SDS-PAGE, and agarose gel analysis of exposed waterborne viruses showed various levels of damage to both surface proteins and their related RNA genes after the APCP exposure, thus leading to the loss of their viability and infectivity.

This study was designed to investigate the effects of APCP on both airborne and waterborne MS2 viruses under different experimental conditions and, further, to elucidate the mechanisms by studying possible damage to RNA genes, the viral surface, and also surface proteins using reverse transcription (RT)-PCR and gel electrophoresis, as well as sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). The results provide guidance on the use of APCP as an alternative for combating viral threats.

MATERIALS AND METHODS

Preparation of MS2 suspensions. The aerosolized MS2 bacteriophage (ATCC 15597-B1) served as the challenge viral aerosol in this study. The double-top-agar-layer plaque technique described previously (37) was utilized to propagate MS2 virus using *Escherichia coli* (ATCC 15597) as the host organism. After culturing and washing, the sterile water containing the MS2 phages was decanted into a sterile 25-ml tube (Corning, Inc., Corning, NY, USA) and further centrifuged at 5,000 × g for 30 min to

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** Different Gas Carriers and Power Levels

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additional dry N2 flow rate (and the viral aerosols were delivered to the exposure test chamber by an nebulizer flow rate (S3 in the supplemental material. Current sets: 30 V and 0.68 A, 30 V and 0.8 A, and 30 V and 0.93 A. The and 28 W correspond, respectively, to the following three voltage and generation power levels and three different gas carriers, including ambient air, He-O2 (2%, vol/vol), and Ar-O2 (2%, vol/vol). On the basis of the dimensions of the plasma generation device and the exit flow rate (equal to the BioSampler’s sampling flow rate of 12.5 liters/min), the exposure time for the virosions inside the plasma device was calculated to be 0.12 s. Each of the exposed and control viral aerosol samples was continuously collected for 20 min. The BioSampler was charged with 20 ml of deionized (DI) water (Millipore Corporation, Billerica, MA, USA). The sampling flow rate was calibrated using a mini-Buck calibrator (A. P. Buck, Inc., Orlando, FL, USA) prior to each test. The viral aerosol samples collected with and without the APCP treatment were cultured for 1 to 2 days at 37°C using the double-top-agar-layer technique. The viruses were enumerated by manual counting of the number of PFU. Using the flow rate and the processing time, the cultivable viral aerosol concentration (number of PFU/m3) was determined. For each set of conditions, the viral aerosol inactivation experiment was conducted in three replicates. After sample collection, plasma discharge was sustained for 3 min at a generation power level of 28 W. A volume of 10 μl of virus suspensions (in number of PFU) with and without the APCP treatment, respectively.

(ii) Exposure of waterborne MS2 viruses to APCP. In this part of the study, the same MS2 bacteriophage was treated with APCP while in suspension (not airborne); additionally, the testing examined longer exposure time intervals up to 3 min. The originally prepared MS2 bacteriophage was serially diluted; dilution factors were 10⁻¹ and 10⁻². In these experiments, 200 μl of viral suspensions with different concentrations was added into the wells of a 96-well plate with 3 replicates. Subsequently, the suspensions were exposed to the APCP produced using a microjet plasma generation device (38) at power levels of 20, 24, and 28 W and exposure times of 30, 60, 120, and 180 s. In the microjet plasma generator, two metal electrodes were separated from each other by a dielectric layer with a thickness of 0.5 mm. The openings in the two electrodes were 0.8 mm in diameter. One electrode was completely embedded in the device and powered by a direct current (DC) power supply (Matsusada Precision Inc., Shiga, Japan), and the other electrode was grounded for safety. Only negative high voltage was tested for this study. As described previously, the discharge sustaining voltage ranged from approximately 400 to 800 V with an operating current in the range of 20 to 35 mA (38). The flow rate of the gas carrier used for the liquid samples was 5 liters/min. Under these operating conditions, a plasma jet of 15 mm was generated in the liquid. Figure S4 in the supplemental material shows the temperature profile, measured using thermal imaging infrared cameras (FLIR Systems, Inc.), for the liquid suspension treated by the microjet plasma generation device in inactivating liquid-borne MS2 viruses. As observed from Fig. S4, the temperature in the liquid suspension ranged from 26.7°C to 36.2°C.

For each set of experimental conditions, the exposure control test was conducted in three replicates, and the viral survival rates were determined using equation 1, provided above, and used for airborne inactivation experiments. In addition, the damage to MS2 viral RNA genes was investigated using waterborne MS2 viruses with the APCP treatment. Agarose gel electrophoresis combined with RT-PCR was applied for analyzing the effects of the plasma treatment on these RNA genes. The extraction of viral RNA was performed following the procedure described in our previous study (37). The cDNA, the forward primers, and the reverse primers used in the previous work (37) were utilized here for amplifying viral surface protein genes: the A (maturation) protein gene (934 bp), capsid (coat) protein gene (160 bp), replicase protein gene (937 bp), and lysis protein gene (114 bp) (39, 40). The RT-PCR cycle conditions were set to the following: 30°C for 30 min (reverse transcription); 94°C for 2 min; 40 cycles in which each cycle included denaturation at 94°C for 1 min, annealing at 55°C for 1 min, and extension at 65°C for 2 min; and, finally, an extension at 65°C for 10 min (37). DI water served as the negative control in the RT-PCR experiments. Agarose gel electrophoresis was performed with the RT-PCR products and Sub-Cell GT instruments (Bio-Rad, Hercules, CA, USA) according to the manufacturer’s instructions. For the gel analysis, approximately 20 μl RT-PCR product was added to each designated well of 3% agarose gels. Gel electrophoresis was performed at a constant voltage of 60 V for 100 min, and then the gels were stained using GelRed solution (diluted 10⁴ times with DI water; Biotium, Hayward, CA, USA) and photographed using a Molecular Imager Gel Doc XR system (Bio-Rad, Hercules, CA, USA) at a wavelength of 254 nm under a UV lamp. In this effort, we did not use a real-time RT-quantitative PCR; thus, the results obtained here represent only those of a semiquantitative analysis of the damaged RNA genes for viruses inactivated by APCP.

To further investigate the mechanism of viral inactivation by APCP, we employed SDS-PAGE. The gel was prepared using a kit following the manufacturer’s instruction (Realtimees Beijing Biotechnology Co., Ltd., Beijing, China). For the electrophoresis buffer, 1X Tris-glycine in a total volume of 300 ml was used. The MS2 viruses were treated with APCP for 3 min at a generation power level of 28 W. A volume of 10 μl of suspension containing either control or treated MS2 virus and a protein ladder (10 to 170 kDa) was separately added to the prepared gel and subjected to SDS-PAGE analysis using vertical electrophoresis equipment (model number
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**RESULTS**

**Airborne inactivation.** Overall, the APCP exposure significantly decreased the levels MS2 virus survival for both the airborne and liquid-borne states. Figure 2 shows the survival of the aerosolized viruses when exposed to the APCP produced using different power levels and gas carriers. Regardless of the gas carrier type, the survival of MS2 viruses was observed to decrease with increasing power levels (for all comparisons, \( P < 0.05 \) by ANOVA). On the other hand, for a given plasma generation power level, the loss of viability of aerosolized viruses strongly depended on the gas carrier type (for all comparisons, \( P < 0.0001 \) by ANOVA). The data from Fig. 2 suggest that the ambient air carrier produced the highest level of inactivation at power levels of 20 and 24 W, followed by the gas carriers Ar-\( \text{O}_2 \) (2%, vol/vol) and He-\( \text{O}_2 \) (2%, vol/vol). In contrast, at a power level of 28 W, the Ar-\( \text{O}_2 \) (2%, vol/vol) mixture was found to be the most efficient in inactivating aerosolized viruses, followed by ambient air and He-\( \text{O}_2 \) (2%, vol/vol). The APCP treatment performed using ambient air as the gas carrier at 28 W for 0.12 s inactivated more than 95% (1.3-log reduction) of the viruses in the air. At the same power level, APCP with Ar-\( \text{O}_2 \) (2%, vol/vol) was not as efficient, inactivating only \(<50\% \) (0.3-log reduction) of viruses. At 20 W, the inactivation produced by Ar-\( \text{O}_2 \) (2%, vol/vol) was as low as \(<10\% \) (0.05-log reduction); however, surprisingly, it increased to approximately 100% (negligibly low survival) when the power level increased from 20 to 28 W (Fig. 2). With Ar-\( \text{O}_2 \) (2%, vol/vol), the viral viability loss was even higher than the one achieved with ambient air at the same power level. From the linear regression analysis, we concluded that APCP with Ar-\( \text{O}_2 \) (2%, vol/vol) features the highest dependency of its viral aerosol inactivation on the power level, followed by plasma with He-\( \text{O}_2 \) (2%, vol/vol) and ambient air as gas carriers.

**Waterborne inactivation.** Constrained by the size of the plasma generation device available and the exit flow rate of 12.5 liters/min, the aerosol exposure time tested here was rather short (about 0.12 s). To extend the plasma exposure time, waterborne MS2 viruses were used for further tests. As seen in Fig. 3, the survival of waterborne MS2 viruses decreased as the exposure time increased from 30 to 180 s, when APCP was produced using the ambient air carrier at different power levels. Similar to the results of the experiments with aerosolized viruses, an increase in the power level was shown to decrease the survival of waterborne MS2 viruses (Fig. 3). Linear regression analysis indicated that there was a strong association between the exposure time and survival; i.e., a longer exposure time corresponded to lower survival of the waterborne MS2 viruses (\( R^2 > 0.95 \) for all energy levels). Similar results were observed with other gas carriers, such as Ar-\( \text{O}_2 \) (2%, vol/vol) and He-\( \text{O}_2 \) (2%, vol/vol), as shown in Fig. S5 and S6 in the supplemental material. The differences in survival and dependence on power level for different gas carriers can be attributed to the different rates of ROS generation arising from the different atomic chemistries; additionally, the discrepancy may be associated with the different die-off times of the ROS and the charged particles released from the APCP.

**Inactivation mechanisms.** Figure 4 presents the data obtained by analyzing the APCP chemistry produced under different gas carriers and power levels. Optical emission spectroscopy (OES) identified the ROS, such as O (atomic oxygen), in the emission peaks when testing was done with gas carriers such as Ar-\( \text{O}_2 \) (2%, vol/vol) and He-\( \text{O}_2 \) (2%, vol/vol). From the data presented in Fig. 4, it can be seen that the amount of O increased with increasing plasma generation power level. Typically, the plasma produced using Ar-\( \text{O}_2 \) (2%, vol/vol) had higher emission peaks of ROS in the emission spectra than the one generated with He-\( \text{O}_2 \) (2%, vol/vol). This, in turn, resulted in rates of virus survival with
He-O₂ (2%, vol/vol) higher than those found with Ar-O₂ (2%, vol/vol) (see Fig. S1 and S2 in the supplemental material). Among the ROS, the hydroxyl radicals (·HO) represent the most reactive form; accordingly, a lower rate of viral survival was observed with the APCP produced using ambient air than the APCP produced using the two other gas carriers. This result is consistent with those for the airborne and waterborne virus exposures seen in Fig. 2 and 3, respectively.

To further develop a mechanistic understanding of the inactivation of MS2 viruses due to exposure to the APCP, we examined the control (untreated) and plasma-treated viral suspension samples using a scanning electron microscope. The MS2 virion size is about 27 nm (39), while MS2 RNA is tightly compacted within the virion and confined to a radius of 8.3 ± 1 nm (40). In contrast, the MS2 coat protein shell is approximately 13.6 ± 2 nm in size and has a thickness of 2.1 ± 1 nm (40). As seen from Fig. 5, the images of virions exposed to the plasma treatment show more fragments than nonexposed samples, which show spherical particles. Nonetheless, due to the low resolution, the detailed surface damage might not be immediately identified from the images. In contrast, damage to the viral RNA genes that code for four major viral surface proteins, the maturation protein (A protein), the capsid protein, the replicase protein, and the lysis protein, was confirmed (Fig. 6). Figure 6 shows the results of agarose gel electrophoresis of viral RNA genes coding for surface proteins of waterborne MS2 viruses upon plasma exposure over 30, 60, 120, and 180 s at a power level of 24 W when ambient air served as the gas carrier. The RT-PCR products were detected in the target bands, and the gel band intensity data generally suggest a positive association between the time of exposure to APCP and the degree of gene damage. One exception was the reappearance of the replicase protein gene with 120 s exposure, which was unexpected and might have
been caused by the nonuniform plasma exposure to the viral suspension due to the airflow used to produce the plasma. In general, damage to the maturation and replicase genes made the virus unable to express the related proteins, which impeded its ability to seek the host and replicate. For example, even if all the surface proteins are intact after the APCP treatment, damage solely to the RNA genes may prevent the virus from replicating inside the host. In addition to damage to RNA genes, the plasma exposure could also have caused damage to surface proteins.

For MS2 viruses, the following proteins are known to exist on its surface: the maturation protein (40 kDa), the coat protein (13.86 kDa), the replicase protein (60 kDa), and the lysis protein (8.87 kDa) (39, 40). To study the effects of APCP on these surface proteins, SDS-PAGE of the APCP-treated MS2 viruses was conducted. Replicase protein (red arrow) and maturation protein (blue arrow) were observed on the SDS-polyacrylamide gel image (Fig. 7). Overall, we have found from the SDS-polyacrylamide gel image (Fig. 7) that with exposure to APCP the band intensity for both the replicase and maturation proteins decreased; e.g., it was barely seen for the maturation protein after the 180-s exposure. Also observed from Fig. 7, some bands disappeared after the APCP exposure. These data suggest that the APCP exposure can significantly degrade the MS2 viral surface proteins. Because the capsid and lysis proteins have smaller sizes, they were not clearly separated by SDS-PAGE under the conditions applied in this work.

DISCUSSION
The differences in viral survival levels observed in this study for different gas carriers can be linked mainly to the differences in plasma chemistry. It has recently been reported that an increase of the oxygen concentration in the mixture of H₂ and O₂ from 0.25% to 0.75% generated a >1.5-fold increase in the level of inactivation of surface-borne MS2 viruses (36). A >99% (>2-log reduction) inactivation of waterborne MS2 viruses was achieved in this study as a result of their 3-min exposure to ACPC with He-O₂ (2%, vol/vol) (see Fig. S6 in the supplemental material), and such an efficiency may be further enhanced by optimizing the oxygen concentration. For example, a selected oxygen percentage of between 0.5% and 1.5% in the He-O₂ gas carrier was reported to produce high levels of atomic oxygen (41). Ambient air with an oxygen content of approximately 21% has a composition vastly different from that of the two other carriers utilized in this study. Accordingly, different reactive products are released. In a previous work (34), hydroxyl radicals rather than atomic oxygen were observed earlier when generating APCP with the ambient air carrier. As shown in Fig. S4 in the supplemental material, the highest temperature generated in the liquid suspension was about 36.2°C, and such a temperature does not affect the viability of the viruses.

FIG 6 Image of gel after agarose gel electrophoresis of viral RNA genes coding for the surface proteins A protein, capsid protein, replicase protein, and lysis protein of waterborne MS2 viruses. The gas carrier was ambient air, the plasma generation power was 24 W, and the time of exposure to plasma was 30, 60, 120, or 180 s. Lanes: 1, D2000 marker (numbers on the left are molecular sizes [in base pairs]; 2, MS2-positive control; 3, time of exposure of 30 s; 4, time of exposure of 60 s; 5, time of exposure of 120 s; 6, time of exposure of 180 s; 7, negative control (DI water).

FIG 7 SDS-polyacrylamide gel image of the APCP-treated and untreated (control) MS2 viral surface proteins. The plasma was generated with the device used to test waterborne viruses, the plasma generation power was 28 W, and the time of exposure was 180 s. Lanes: 1, protein ladder; 2, untreated MS2 virus control; 3, plasma-treated MS2 viral proteins; 4, negative control (DI water). Red arrow, replicase protein (60 kDa); blue arrow, maturation protein (40 kDa).
Therefore, the inactivation of MS2 viruses by APCP that we documented here was primarily associated with the generated ROS, such as hydroxyl radicals and atomic oxygen, and not the heat generated from the plasma device. These species could have reacted with RNA genes, resulting in their damage (Fig. 6). The ROS generated from the plasma is described to be able to break structurally important bonds of peptidoglycan (i.e., C—O, C—N, or C—C bonds) and to lead to the destruction of the bacterial cell wall (42). Alternative causes may also be present. For example, one study pointed out that the plasma inactivation of lambda phage occurred mainly due to the damage to the coat protein, while its DNA damage at early stages was negligible (43).

Here, we observed damage to the surface proteins; i.e., the band intensity was significantly decreased compared to that for the control or disappeared, as seen in Fig. 7. A previous study has shown that the A protein is exposed to the capsid surface and directly binds the MS2 genomic RNA at its 5′ and 3′ ends (40). In addition to RNA packaging, the main function of the A protein of an MS2 virion is to identify the host. In contrast, the viral replicase and lysis proteins are involved in the replication and lysis of E. coli bacteria, respectively (40). Therefore, any damage to the A (maturation) and replicase proteins from the external assault would result in the loss of virus infectivity as well as RNA packaging. On the other hand, the MS2 coat protein, the primary structural protein shell, has a total of 90 homodimers arranged in a quasi-equivalent T=3 lattice on the capsid surface (44). The MS2 coat protein was shown to bind a stem-loop structure in viral RNA for both encapsidation of the genome and translational repression of replicase synthesis (44). Besides, the coat protein is reported to interact with MS2 RNA at 20 sites along the RNA backbone (44). Unfortunately, in this study, the bands for lysis and coat proteins did not show up in the gel due to their small sizes. Nonetheless, damage to the coat proteins would affect the infection ability of the MS2 viruses.

In this study, we have observed a significant inactivation of MS2 viruses resulting from their exposure to APCP. The data suggest that the inactivation of the MS2 viruses for both the airborne and waterborne states depends on the power level, exposure time, and gas carrier. The viral inactivation process is primarily attributed to the ROS released from APCP, including damage to the virus surface proteins and RNA genes. Although the magnitude of the effect of APCP is generally higher than that present in the atmosphere, results presented here can facilitate a better understanding of virus decay in ambient air. Future research efforts should be directed toward determining the efficiency of APCP against human viral pathogens, such as H1N1 and H7N9 influenza viruses and even Ebola virus. Owing to its complex contents, e.g., an assembly of photons, electrons, positive and negative ions, free radicals, and excited or nonexcited molecules and atoms (34, 45), the currently achieved understanding of the biological mechanism of APCP-induced viral inactivation is rather limited, and this work contributes to the development of a better understanding of viral inactivation when viruses are in contact with APCP. Clearly, the APCP technology has advantages over chemical methods with respect to secondary environmental pollution and cost (12), and the information obtained here will also serve as a foundation for developing efficient plasma-based technologies and strategies to control viral agents.

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