Effect of Air Ions on Submicron T1 Bacteriophage Aerosols

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ABSTRACT

HAP, JOHN W. (Fort Detrick, Frederick, Md.), J. BRUCE HARSTAD, AND LEE M. BUCHANAN. Effect of air ions on submicron T1 bacteriophage aerosols. Appl. Microbiol. 14:888–891. 1966.—The effect of a high concentration of ionized air molecules on sampling T1 phage aerosols of submicron particle size was evaluated by comparing the phage recoveries of all-glass impingers (AGI-4) and type 6 filter papers. Sampler recoveries of all ionized aerosols were less than the recoveries of nonionized control aerosols. These reductions in recovery were greater with positive ions than with negative ions or ions of mixed polarity. The AGI-4 allowed considerable slippage, which was not affected by the air ions. Type 6 filter paper recoveries were less than AGI-4 recoveries. The air ions did not appear to affect the aerosol particle size as determined by an electron microscope.

Whitby (5) developed a sonic jet ionizer that produces a high concentration of ionized air molecules (air ions) and investigated their effect on inert aerosols. Whitby, Lundgren, and Peterson (6) found that aerosol particles produced by the atomization of dilute solutions of dyes carried a high electric charge that made the aerosol particles irregular in size and shape. Neutralization of the aerosol with a mixture of positive and negative air ions produced a more uniform aerosol and eliminated the migration that normally occurs with charged aerosols.

The action of air ions on bacteria suspended in small drops has been studied by Kreuger, Smith, and Go (3), who concluded that positive and negative air ions accelerated the rate of death of staphylococci, apparently by direct action on the cells and also by increasing the rate of evaporation.

Phillips, Harris, and Jones (4) found that a high concentration of air ions increased the exponential decay rates of aerosols of Serratia marcescens. The major part of the aerosol decay in the absence of air ions was due to biological decay; in the presence of air ions, most of the increase in decay was attributed to physical decay. Negative and positive ions caused similar increases in physical decay. In addition to the increase in physical decay, negative ions increased biological decay but positive ions did not.

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This paper reports on the effects of air ions of positive, negative, and mixed polarities on the sampling and particle size of submicron aerosols of T1 coliphage at a low relative humidity. The submicron aerosols were produced with a Dautrebande aerosol generator from concentrated aqueous suspensions of purified phage. The air ions were produced by the Whitby (5) sonic jet ionizer. The methods for phage purification and for generating, sampling, and particle-sizing submicron phage aerosols were those used by Harstad (2).

MATERIALS AND METHODS

Air ion apparatus. A sonic jet ionizer (Fig. 1), developed by Whitby, that is capable of producing a high concentration of air ions was used in these tests. The ionizer consists of a sharp needle located upstream from an orifice plate. A positive, negative, or alternating current is applied between the needle and the orifice plate to create a slightly visible corona at the tip of the needle. Air ions are formed as a stream of air passes through the corona. The ions are then forced through the plate orifice at sonic speeds, thus freeing the ions from the electric field around the corona. The particular specifications of the sonic jet ionizer are: voltage, 3,500 v; orifice diameter, 1.6 mm; needle spacing, 1.9 mm; ionized gas, dried and filtered air; air pressure, 29 psi; air flow, 87 liters per min; ions, positive, negative, or a mixture of both; ion concentration, 106 ions per cm3.

A Philco Ion Counter, model ICG-6, was used to estimate the number of unit ion charges emitted by the sonic jet ionizer. Ozone was not detected by the
stretched-rubber test, which is a simple method used by Crabtree and Erickson (1) for measuring approximate atmospheric ozone.

**Phage aerosol generation.** The virus used in the aerosol tests was T1 bacteriophage, one of the viruses parasitic to *Escherichia coli* B. Aqueous suspensions of T1 phage were prepared by concentrating and purifying large volumes of broth cultures by differential centrifugation and washing with distilled water. This thorough cleansing of the phage suspension to remove soluble and particulate contaminants was necessary to produce aerosols of minimal particle size. The concentration of the phage suspension used to fill the aerosol generator ranged from 0.5 \times 10^{11} to 1.5 \times 10^{11} phage particles per milliliter.

The Dautrebande De type aerosol generator (J. H. Emerson Co., Cambridge, Mass.) was used to produce the submicron phage aerosols. The generator was operated at an air pressure of 17.5 psi, which resulted in an air flow through the generator of 18.4 liters per min and a fluid atomization rate of 0.15 ml per min. After leaving the generator, the aerosol passed into a 4-liter cylindrical glass chamber, then entered a 45-liter glass carboy where the aerosol was mixed with 87 liters of air containing the air ions. After leaving the carboy, the aerosol entered a circular sampling manifold to which the aerosol samplers were connected. The excess aerosol was bled off through a filter located upstream from the sampling manifold. The relative humidity was controlled by drying the air before it passed through the air ionizer and was monitored by wet and dry bulb thermometers attached to the air. Air supplies for aerosol generation and ion production were filtered, which assured clean particle-free air. The chamber was grounded to eliminate the residual charge that remained on the chamber walls after ionization. A schematic diagram of the aerosol chamber and apparatus is shown in Fig. 2.

**Phage aerosol sampling.** Two types of aerosol samplers, the all-glass impinger (AGI-4; Ace Glass Co., Vineland, N.J.) and Chemical Corps type 5 filter paper (Holingsworth and Vose Co., East Walpole, Mass.), were used to determine the phage aerosol concentration in the presence or absence of added air ions. The all-glass impinger was filled was 22 ml of Nutrient Broth (Difco) containing 0.1% Antifoam A (Dow Corning Corp., Midland, Mich.) and was operated at the maximal (near sonic) flow rate of 12.5 liters per min. The Chemical Corps type 6 filter paper, an ultra-high-efficiency paper composed of Bolivian or African Blue asbestos, esparto grass, and kraft fibers was cut into discs (2.5 cm in diameter) and sealed in in-line filter holders (7). The sampling flow rate through the holders was 1.0 liter per min. Immediately after sampling, the filter papers were placed in 100 ml of Nutrient Broth containing 0.1% Tween 20 (Atlas Powder Co., Wilmington, Del.) and then shaken for 15 min on a mechanical shaker to disintegrate the paper.

Slippage of the phage aerosol through the all-glass impinger was determined by backing up the impinger with a filter holder containing type 5 and type 5 Chemical Corps filter papers. The type 5 filter is a lower-efficiency paper of cellulose and asbestos fibers with a backing of cotton scrim. It was used only to support the more fragile type 6 paper and was assayed with the type 6 paper.

To compare the size and shape of the phage aerosol particles before and after the addition of air ions, an electrostatic precipitator was used. The precipitator, which was connected to the sampling manifold, collected aerosol samples on an electron microscope specimen grid. The grid was then examined in an electron microscope, and photographs of the aerosol particles were taken.

Sampler air flow rates were calibrated with a wet-test meter. The all-glass impingers (AGI-4) were also calibrated for clearance of orifice to flask bottom, and those with a clearance of 4 mm were selected for these tests.

Phage suspensions and aerosol samplers were assayed by making duplicate serial dilutions in Nutrient Broth and plating 1-ml samples of each dilution in triplicate, by use of an agar layer method described by Harstad (2).

**Design of experiments.** The aerosol trials were designed to compare the effects of a high concentration of air ions of positive, negative, and mixed polarities on the sampling and particle size of submicron T1 phage aerosols. Each aerosol trial consisted of three consecutive tests conducted on a single day: a control test in which no air ions were added to the aerosol, a...
TABLE 1. Effect of air ions on the sampling of submicron T1 phage aerosols

<table>
<thead>
<tr>
<th>Sampler</th>
<th>Aerosol condition</th>
<th>No. of comparisons</th>
<th>Mean recovery</th>
<th>Standard error</th>
<th>95% Confidence limits</th>
</tr>
</thead>
<tbody>
<tr>
<td>AGI-4</td>
<td>Mixed ions added</td>
<td>10</td>
<td>48.3</td>
<td>4.31</td>
<td>38.6-58.0</td>
</tr>
<tr>
<td></td>
<td>Negative ions added</td>
<td>10</td>
<td>43.6</td>
<td>5.79</td>
<td>30.5-56.7</td>
</tr>
<tr>
<td></td>
<td>Positive ions added</td>
<td>6</td>
<td>11.4</td>
<td>2.25</td>
<td>5.6-17.2</td>
</tr>
<tr>
<td>Type 6 filter paper</td>
<td>Mixed ions added</td>
<td>10</td>
<td>34.4</td>
<td>2.76</td>
<td>28.2-40.6</td>
</tr>
<tr>
<td></td>
<td>Negative ions added</td>
<td>10</td>
<td>30.0</td>
<td>4.36</td>
<td>20.1-39.9</td>
</tr>
<tr>
<td></td>
<td>Positive ions added</td>
<td>6</td>
<td>7.6</td>
<td>1.79</td>
<td>3.0-12.2</td>
</tr>
</tbody>
</table>

* Each trial yields two comparisons: ionized aerosol recovery/first control aerosol recovery and ionized aerosol recovery/second control aerosol recovery. Mean aerosol temperature, 22°C; range, 20 to 25°C; mean aerosol relative humidity, 29%; range, 24 to 32%.

TABLE 2. Comparison of type 6 filter paper and all-glass impingers (AGI-4) for the recovery of ionized and nonionized submicron T1 phage aerosols

<table>
<thead>
<tr>
<th>Aerosol condition</th>
<th>No. of tests</th>
<th>Mean relative efficiency (%)</th>
<th>Standard error</th>
<th>95% Confidence limits</th>
</tr>
</thead>
<tbody>
<tr>
<td>No ions added (control)</td>
<td>26</td>
<td>63.5</td>
<td>2.94</td>
<td>57.4-69.6</td>
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<tr>
<td>Mixed ions added</td>
<td>5</td>
<td>43.2</td>
<td>5.57</td>
<td>27.7-58.6</td>
</tr>
<tr>
<td>Negative ions added</td>
<td>5</td>
<td>39.8</td>
<td>7.31</td>
<td>19.5-60.1</td>
</tr>
<tr>
<td>Positive ions added</td>
<td>3</td>
<td>40.8</td>
<td>0.47</td>
<td>38.8-42.8</td>
</tr>
</tbody>
</table>


† Two control aerosol tests for each ionized aerosol test.

Aerosol test in which air ions were added, and a final control test in which no air ions were added. The conditions of all tests were the same except for the addition of air ions. A series of 13 trials (days), consisting of 39 aerosol tests, was conducted. Five trials were conducted with negative ions and ions of mixed charge, and three trials were conducted with positive ions. The schedule followed for each aerosol test consisted of filling the Dautrebande aerosol generator with 6 ml of the phage suspension and disseminating the phage aerosol continuously for 15 min, 10 min to establish equilibrium inside the aerosol mixing chamber followed by a 5-min aerosol-sampling period, and then 10 min to flush the aerosol chamber with clean, filtered, dry air. In the tests in which air ions were added, the Whitby sonic jet ionizer was turned on and off simultaneously with the Dautrebande aerosol generator. Except for the electrostatic precipitator, two samplers of each type were used and the samples were pooled for phage assays.

RESULTS

Preliminary experiments. In the first few aerosol tests conducted with air ions of mixed charge, the

FIG. 3. Electron micrograph of particles from a T1 phage aerosol of mixed charge.
chamber was not grounded and sampler recoveries were noticeably higher during the second control test than during the first. To find the reasons for this phenomenon, an electrometer was used to determine whether there was a charge on the chamber walls prior to the first and second control tests. No charge was present prior to ionization, and a positive charge did remain on the walls of the chamber after ionization was stopped and the aerosol chamber was flushed. When the chamber was grounded, the residual charge was eliminated and sampler recoveries during the first and second control tests matched. Apparently, the positive charge remaining on the chamber walls after ionization caused a repulsion of the positively charged aerosol particles during the second control test, thereby permitting a greater number of organisms to be available for sampling. All data reported here were obtained when the chamber was grounded.

**Effect of added air ions on sampler recovery.** In Table 1, sampler recovery of ionized aerosols is compared with the recovery obtained for control (no ions added) aerosols as given by: recovery = ionized aerosol recovery/control aerosol recovery. AGI-4 and type 6 filter paper recovery of all ionized aerosols was less than the recoveries of control aerosols. This loss in sampler recovery was more pronounced with positively charged aerosols than with negatively charged aerosols or aerosols of mixed charge. The recovery with negatively charged aerosols was somewhat less than that with aerosols of mixed charge, but the differences were not considered significant.

In Table 2, type 6 filter paper recoveries are compared with AGI-4 recoveries as given by: relative recovery = type 6 filter paper/AGI-4. Type 6 filter paper recoveries were less than AGI-4 recoveries for control aerosols and ionized aerosols. Air ions reduced recoveries with the type 6 filter paper sampler to a greater extent than with the AGI-4 sampler (Table 1). Type 6 filter paper recovery was approximately 40% of AGI-4 recovery for each of the three ion treatments; this value was 63.5% for control aerosols.

Table 3 shows the slippage of the phage aerosols through the AGI-4, defined as: slippage = backup filter/backup filter + AGI-4. Slippage was slightly higher for ionized aerosols than for control aerosols, but none of the differences was considered significant.

**Effect of air ions on aerosol particle size.** Electron micrographs of phage aerosols showed no significant changes in size or shape between ionized and control aerosols. Figure 3 is an electron micrograph of a phage aerosol exposed to air ions of mixed charge.

**Discussion**

Previous studies on the sampling of submicron T1 phage aerosols in the absence of air ions have shown that biological loss, such as death of the phage in the aerosol and killing of phage by the samplers, is much larger than physical loss of the aerosol (2). The present study revealed that air ions affected the stability of submicron T1 phage aerosols, resulting in a reduction of sampler recoveries. The reduced phage recoveries could be attributed to (i) aerosol death, which is a function of biological stability; (ii) physical aerosol loss from fallout or migration to the chamber walls; or (iii) killing of phage by the samplers, which is also a function of biological stability. The data do not reveal which of these factors was predominant. Slippage of the aerosol through the sampler was not a factor, because AGI-4 slippage was not affected by the ion treatments and the previous study revealed that type 6 filter paper is virtually a complete collector of submicron particles.

**Acknowledgments**

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**Literature Cited**


