Construction of Its Evaluation System in Originally Designed Test-Chamber System and Sporicidal Activity of Aerosolized Hypochlorite Solution to \textit{Bacillus subtilis} Spores

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Effective spatial disinfection systems are required for human health care, public hygiene, and food and medicine manufacturing. Although some aerosolized disinfectants were already applied to its purpose, accurate evaluation systems were under-constructed. In this study, the spatial sporicidal activity of aerosolized hypochlorite solution (AHS) to dormant cells, \textit{Bacillus subtilis} spores, was evaluated by an originally designed chamber system. In the test-chamber, AHS was supplied and existed as micro-droplets, and environmental relative humidity (RH) could be controlled. Available chlorine (AC) exposure was also controlled with appropriate AC loading but was influenced by the acidity of AHS. Our results indicated that inactivation of spore was depend on AC exposure amount and time. On the other hand, unsaturated environmental RH markedly decreased spore inactivation. This study indicated that our test-chamber system can provide reproducible test data under a homogeneous air condition, and, thereby, that the data obtained by the chamber system should contribute to predicting the AC-required dose to disinfect a whole building.

Key words: Aerosolized hypochlorite solution / Sporicidal activity / Environmental relative humidity / Spatial antimicrobial activity evaluation chamber.

INTRODUCTION

As strong desire for disinfection and cleanness in a living environment and significance of countermeasures against infectious diseases and food contaminations, disinfection of facilities and sanitary control in living environment become of more importance and effective spatial disinfection or sterilization methods, greatly contributed to them, are expected. To solve these desires, several sanitary control systems, including UV irradiation method and directly air-filtration method, were invented, but it is considered that the gasified disinfectant is most effective even to high resistant dormant cells like a bacterial spore and has good prospects. Especially, bacterial spores generally are highly resistant to a variety of stresses, including heat, ultraviolet radiation and some disinfectants, and, therefore, spores survived after some kind of antimicrobial treatment against microbial contaminants often cause serious problems in food and pharmaceutical industries. Consequently, there is a need to develop effective sporicidal treatments to reduce hygienic risks (Nakagawa et al., 1998), and gasified sporicidal disinfectants are expected to be commercially available.

Several gaseous disinfectants, such as formaldehyde, ethylene oxide, ozone, hydrogen peroxide vapor and chlorine dioxide, have been reported to possess sporicidal activity and are utilized as an effective spatial disinfecting reagent in food and pharmaceutical industry.
Gaseous disinfectants are also advantageous for decontaminating atmosphere in large room and building and also for disinfecting device and equipment because of the easy dispersion and diffusional contact on large and complex surface. On the other hand, some gaseous disinfectants do not have enough sporicidal activities at a practical concentration and others have serious toxicity and metal corrosiveness. It is important and necessary to determine the treatment conditions, to monitor their residues and to introduce a ventilation system for keeping safety environment (Rogers et al., 2007). Therefore, the accurate modeling to molecular dispersion of gaseous disinfectant and death-kinetics of microbial cell are required. In the homogeneous disinfectant dispersion condition, it was proposed that the microbial cell death followed the Chick-Watson model (Chick 1908; Watson 1908). Mendes et al. (2007) were also propose that lethal modelling of gaseous disinfectant and its validation under practical conditions was necessary and important to use it in safety and reliability, as its microbicidal activity, like ethylene oxide gas, was extensively affected by environmental conditions.

It is also expected that aerosolized disinfectant solutions, including in aerosolized hypochlorite solution (AHS), are recently interesting for hygiene control of indoor environment in buildings and facility surface, like a gaseous disinfectant (Thorn et al., 2013; Martyny et al., 2005), but few practical uses of AHS were reported because of a shortage of basic experimental data for AHS disinfection. Oh et al. (2005) also suggested that aerosolized disinfectants diffuse in a manner similar to gaseous disinfectants, but it has also been considered experimentally to verify their micro-droplet behaviour (Burfoot et al., 1999). Therefore, it is important to understand the quantitative relationship between the dose of an aerosolized disinfectant and reduction of surviving microorganisms for hygienic control in industrial use. In this research, the test-chamber system, which able to supply aerosolized disinfectant with constant exposure rate, was newly designed and constructed. AHS antimicrobial activities were measured in the chamber under suitable treatment conditions, using Bacillus subtilis spore. Amount of available chlorine (AC) and environmental relative humidity (RH) were influenced to the sporicidal activity of AHS.

**MATERIALS AND METHODS**

**Bacterial strain and culture condition**

Bacillus subtilis ATCC6633 spore suspension (product code E-MN11, 10³ spores per mL) was purchased from Eiken Chemical Co., Ltd. (Tokyo, Japan). B. subtilis spores were cultivated at 37°C without shaking in NBg medium consisting of 8 g of Difco™ nutrient broth (product code 234000, Becton, Dickinson and Company, NJ, USA), 5 g of NaCl and 2 g of glucose per litre (pH 7.0). NBg agar medium supplementing with 1.5% agar was also used for colony formation.

**Chemicals**

Sodium hypochlorite solution with >5.0% AC (product code 197-02206; Wako Pure Chemical Industries, Ltd., Osaka, Japan) was used for aerosolized hypochlorite disinfectant in this study. Sodium hypochlorite pentahydrate with >39.0% AC (product code 195-17212; Wako Pure Chemical Ind., Ltd.) was also used as a standard substance for hypochlorite quantification. Weakly acidic (WA) -hypochlorite (pH 6.5) and basic (B) -hypochlorite (pH 9.0) solutions were prepared with an addition of 1 mol L⁻¹ of HCl solution. 2-[6-(4′-Amino)phenoxy-3H-xanthen-3-0-n-9-y]-benzoic acid (APF) and 2-[6-(4′-hydroxy)phenoxy-3H-xanthen-3-0-n-9-y]-benzoic acid (HPF) were purchased from Goryo Chemical Inc. (Sapporo, Japan) as a fluorescence probe to detect highly reactive oxygen species (ROS) such as hydroxyl radical and peroxynitrite. APF can also detect hypochlorite ion.

**Test-chamber system**

The test-chamber system shown in Fig. 1 was originally designed and constructed for this study (Ishikawa et al., 2016). The chamber was composed of an entrance box, air and AHS supply ports, a discharge port, a thermo-hygrometer, a humidity controller and a pair of gloves attached to a front panel. An HSP-SAN100 ultrasonic mist generator for AHS supply was provided by HSP Co. (Okayama, Japan). Highly purified air (Grade 2, Tomoe Shokai Co., Ltd., Tokyo, Japan) was introduced into the chamber via a filter unit of Millipore®-FH (pore size 0.45 μm, Millipore Co., MA, USA). A circulating aspirator (WJ-20, Shibata Scientific Technology Ltd., Tokyo, Japan) was connected in the discharge port. A humidity controller composed of TTM-004-2-R-AP-TE and TA503BL300V01-EP-BHAN-2M (Toplas Engineering Co., Ltd., Tokyo, Japan) was used to regulate RH in the chamber with controlling inflow rates of AHS and highly purified air. A thermo-hygrometer (TR-72wf, T&D Corporation, Nagano, Japan) monitored temperature and RH during an AHS treatment.

**Preparation of spore-attaching (SA) plates**

B. subtilis spore was collected with a centrifugation at 4,000×g for 10 min at 4°C in order to replace the buffer and to remove cell debris in the suspension (Eiken Chemical Co., Ltd.), subsequently washed twice with a sterilized distilled water (SDW), and resuspended in SDW. The resultant suspension was diluted stepwise,
and 5-µl aliquots of the suspension were subsequently poured onto grids of an inner surface of a lid of a poly-styrene 96-well microplate (product code 3870-096; AGC Techno Glass Co., Ltd., Shizuoka, Japan) and dried at room temperature for overnight in a vacuum desiccator. A series of 8 dilutions \((10^0 - 10^4)\) spores per grid\) was spotted in 6 replications (Fig. 2A).

### AHS treatment

WA-hypochlorite and B-hypochlorite solutions adjusted to 9.0-4,000 mg L\(^{-1}\) of AC were filled into the ultrasonic mist generator. Micro-droplets (less than 10 µm average diameter) of AHS generated with it were supplied into the test-chamber at a rate of 73.8 mL h\(^{-1}\) and exhausted with an aspiration at 0.72 m\(^3\) h\(^{-1}\) to maintain the spatial AC concentration at a constant value. RH in the chamber was kept at saturated condition (95-100% RH) during the AHS treatment under this condition. As of keeping a RH of 70%, AHS and high purified air were supplied into the chamber intermittently, and AHS supply rate was, consequently, reduced to 37.6 mL h\(^{-1}\). The inner atmosphere of the chamber was stirred using water-resistant fans (SAN ACE 120WS, SANYO Electric Co., Ltd., Osaka, Japan) (Fig. 1), and the air flow in the chamber was also circulated along the inner wall surface. Micro-droplets of AHS were consequently supplied to the SA plates at a horizontal airflow of 0.6 m s\(^{-1}\) (Ishikawa et al., 2016).

The SA plates prepared were imported into the chamber through the entrance box and subsequently placed upside down on the exposure stage (Fig. 1 and Fig. 2C). Spores attached on the surface of an SA plate were exposed to AHS at room temperature. At the end of treatment, an SA plate was incubated at room temperature for 1 h in a dry environment. Finally, the resultant SA plate was mounted on a fresh 96-well microplate fulfilled with a NBg medium (370 µl per well), and cultivated at 37°C for 3-4 days (Fig. 2C). Spore viability on a grid was checked with a turbidity and/or a cell clot in the medium of its corresponding well.

### Most probable number (MPN) method for estimation of spore survival on an SA plate

The number of survived (germinable) spores on a grid of an SA plate was estimated (Ishikawa et al. 2015) using the most probable number (MPN) method described by Halvorson and Ziegler (1933). The number of survived spores per grid \((N)\) was calculated with the following equation:

\[
N = MPN \times D = \ln(n / q) \times D
\]

where \(D\) is the dilution factor, and \(n\) is the number of replicate grids (in this study \(n = 6\)) with same initial spore number \((10^0 - 10^4)\) intact spores in initial), and \(q\) is the number of grids showing no growth (no viable spore).
The log spore survival was calculated as the logarithm of the number of surviving spores \( N \) divided by initial spore number \( N_0 \):

\[
\text{Log spore survival} = \log_{10} \left( \frac{N}{N_0} \right) \quad (2)
\]

To confirm microbial contamination on an SA plate, a lid of the polystyrene 96-well microplate without spores was also exposed to AHS and cultivated in the same procedure.

**Colonify count method to detect viable spores**

To validate a viable spore count by the MPN method and an interfering activity of NBg medium to sporidial action of hypochlorite, survived spores were directly evaluated by colony-forming ability (colony count method). A portion of a spore suspension with or without treated by hypochlorite was diluted with SDW, spread on NBg agar plates, and then cultivated at 37°C for 16 h. Colonies were counted as survival spores.

**Quantification of AC**

Hypochlorite was quantified by the method described by Setsukinai et al. (2003) using sodium hypochlorite pentahydrate as a standard substance, and the hypochlorite concentrations of a purchased sodium hypochlorite solution and AHS during the treatment were converted to AC concentration as described by Dychdala (2001). Hypochlorite was evaluated specifically using APF and HPF in this method.

Micro-droplets of AHS during the treatment were captured with a Petri dish (diameter X height: 34.6 X 17.5 mm) filled with 14 mL of SDW and placed on the exposure stage of the test-chamber. After the AHS treatment, the solution collected in a dish was added to APF solution (10 µmol L\(^{-1}\) of APF dissolved in 50 mmol L\(^{-1}\) phosphate buffer saline, pH 7.4) at a volume of 1/50 to 1/10 to measure the amount of ROS and hypochlorite. The resultant cocktail was then incubated at 37°C for 1 h. The fluorescence of the reactant was measured using a FP-8200 spectrofluorometer (JASCO Corporation, Tokyo, Japan). The excitation wavelength was 490 nm, and the emission was filtered using a 515-nm barrier filter. The total amount of ROS excluding hypochlorite was quantified using HPF in the same protocol described above.

**Evaluation of an AC load and an AC exposure**

The AC Loading rate \( \text{mg m}^{-3} \text{ h}^{-1} \) was calculated as follows.

\[
\text{AC loading rate} = C_{\text{AC}} \times Q_v \times V^{-1} \quad (3)
\]

\( C_{\text{AC}} \) (mg L\(^{-1}\)) is the AC concentration of the hypochlo-
Aerosolized hypochlorite solution (AHS) treatment condition in the test-chamber system. (A) Relative humidity (RH) in the test-chamber was illustrated with or without a humidity controlling. RH without a humidity controlling reached to 99.3 ± 1.0% (saturated humidity, bold line), and RH with 70% and 50% humidity controlling reached to 68.4 ± 2.2% (solid line) and 49.0 ± 3.0% (broken line), respectively. The treatment start time and finish time of the 1-hour AHS treatment were indicated with (↓) and (*), respectively. (B) Relationship between available chlorine (AC) loading rate introduced into the chamber and AC exposure rate to spore-attaching (SA) plates. The AHS treatment was performed at room temperature. Closed and opened circulars indicated the weakly acidic (WA-, pH 6.5) and the basic (B-, pH 9.0) AHS treatment in the saturated humidity, respectively, and triangles indicated the WA-AHS treatment in 70% humidity.

The test-chamber system was originally designed to expose an aerolized disinfectant like AHS to a microorganism contamination surface, such as an SA plate in this study. AHS was able to be kept at a constant concentration under a variety of RHs in the test-chamber because of continuously and constantly supplying and discharging of AHS micro-droplets. RH condition in the chamber during an AHS treatment was shown in Fig. 3A. RH was stable during AHS treatment for 1-3 hour, after the approximate 10-minute transition. RH at a stable stage were 99.3 ± 1.0% during AHS treatment without controlling by the humidity controller (saturated humidity). On the other hand, when the humidity controller was set at 70% and 50% RHs, average RHs were observed at 68.4 ± 2.2% and 49.0 ± 3.0%, respectively. Markedly temperature change was not observed during AHS treatments with a humidity controlling (data not shown).

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Performance of the test-chamber system for AHS treatment

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The relationship between AC loading rate introduced into the chamber and AC exposure rate to a target plate was determined, and a correlation was observed between them (Fig. 3B). According to comparison of AC exposure rates, approximately 4.6-fold higher AC exposure rate was observed in WA-AHS treatments, regardless of under saturated humidity or 70% RH, than in B-AHS treatments under saturated humidity. These results indicated that AC exposure rate could be controlled with an AC loading rate and pH of sodium hypochlorite solution in the mist generator.

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The AC exposure (mg m⁻²), indicated as a practical contact AC amount onto a target plate, was calculated as follows.

\[ \text{AC exposure} = C_{\text{PD}} \times \left( \frac{\text{Volume of SDW filled in a petri dish}}{\text{Area of petri dish}} \right)^{-1} \]

\( C_{\text{PD}} \) (mg L⁻¹) is the AC concentration in SDW collected from the petri dish. The AC exposure rate (mg m⁻² h⁻¹) was calculated by division of an AC exposure by a treatment time.

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5 times higher than the maximum AC concentration carried over in spore cultivations for MPN method using an SA plate after the AHS treatment. Spores were suspended in these solutions at a concentration of $2.0 \times 10^7$ spores mL$^{-1}$, and treated for 1 h at room temperature. Survived spores were evaluated by the colony count method. In present study, the highest AC load in the chamber system was at 824.3 mg m$^{-3}$, and concentration of AC carried over in spore cultivation was calculated to 16.9 mg L$^{-1}$, which is noticeably lower than 83 mg L$^{-1}$. The ratio of spore inactivation with the hypochlorite treatment in SDW was observed at >6.0-log reduction. No marked decrease of survived spores with the hypochlorite treatment in NBg medium was, however, observed. According to these results, NBg medium had enough inactivation activity for remaining AC on an SA plate after an AHS treatment.

**MPN method using an SA plate was useful to determine number of survival spores**

Comparison of the numbers of viable spores which counted by MPN method with an SA plate and by colony counting method was shown in Fig. 4. In this experiment, intact spores without any disinfection treatments were used. No marked difference was observed between viable spore counts obtained both with these methods. Consequently, this result proposed that estimation of the survived spore number with a MPN method using an SA plate was enough quality for evaluation of disinfection activity in AHS treatment.

**NBg medium possessed an interfered activity against the killing reaction of hypochlorite**

The TABLE 1 shows the sporicidal activities of WA-hypochlorite added at 83 mg of AC per 1 liter of NBg medium and sterilized distilled water. Eighty three milligrams per liter of AC concentration is approximately

![FIG. 4](image)

**Comparison of survival spore counts measured by the colony counting method and the MPN method. Each plot was an average value of independent duplicated assays.**

**TABLE 1.** Inactivation of *Bacillus subtilis* spores dipped into a weakly acidic (WA-) hypochlorite solution with or without addition of NBg medium.$^a$

| Medium | Added amount of WA-hypochlorite (mg-AC per 1 liter) | Survived *Bacillus subtilis* spores |
|--------|---------------------------------------------------|-----------------------------------|
|        |                                                   | Colony count (CFU mL$^{-1}$)      | Log reduction ratio |
| SDW    | None                                              | $2.1 \times 10^7$                 | -                   |
| SDW    | 83                                                | $<1.0 \times 10^7$                | >6.3                |
| NBg    | None                                              | $2.4 \times 10^7$                 | -0.1                |
| NBg    | 83                                                | $1.9 \times 10^7$                 | 0.0                 |

$^a$ Spores were dipped into a WA-hypochlorite solution at room temperature for 1 h.

![FIG. 5](image)

**Reduction of survived spores during the aerosolized hypochlorite solution (AHS) treatment at room temperature. WA-AHS was introduced into the chamber at an available chlorine (AC) loading rate of 15.0 mg m$^{-3}$ h$^{-1}$ under the condition of saturated humidity.**

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SPORICIDAL HYPOCHLORITE MIST

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**TABLE 2.** Required available chlorine (AC) to reduce a survived spore to approximately 5 logs.

| Hypochlorite solution | RH          | Log reduction of survival spores | Required amount of available chlorine |
|-----------------------|-------------|----------------------------------|--------------------------------------|
|                       |             | AC exposure                       | AC load                              |
|                       |             | mg m⁻² | fold | mg m⁻³ | fold |
| Weakly acidic (pH 6.5) | Saturated   | 5.4   |     | 4.7    | 1.0  |
| Basic (pH 9.0)        | Saturated   | 4.7   |     | 6.4    | 1.4  |
| Weakly acidic (pH 6.5) | 70% RH    | 5.0   |     | 162.7  | 34.6 |

* Saturated indicates 99.3 ± 1.0% RH.

* The measured value of humidity set at 70% RH was 68.4 ± 2.2% RH (mean ± SD).

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**Characterization of the sporicidal activity of the AHS treatment**

The time course in spore survival during WA-AHS treatment under saturated humidity at room temperature was shown in Fig. 5. WA-AHS was introduced into the chamber at an AC loading rate of 15.0 mg m⁻³ h⁻¹. Values of the spore survival were decreased gradually as the treatment time increasing.

Relationship between the log spore survival and AC exposure was presented in Fig. 6A. Despite pH difference of AHS, the rate of spore inactivation and AC exposure were similar between the WA- and B-AHS treatments under saturated humidity (Fig. 6A). Whereas, decreased sporicidal activity in the WA-AHS treatment under 70% RH was observed, compared with that under the saturated humidity. The required AC exposure to reduce the spore survival to approximately 5 logs were described in TABLE 2. In 70% RH, required AC exposure was observed at 162.7 mg m⁻² and this value was approximately 35-fold higher than those obtained in saturated humidity.

Relationship between the log spore survival and AC loads was also presented in Fig. 6B. Approximately 3.7-fold higher AC load in B-AHS treatment under saturated humidity was required for the similar spore inactivation, compared with in WA-AHS treatment (Fig. 6B and Table 2). This result indicated that AC provided in B-AHS treatment was extremely lower than that in WA-AHS treatment. Approximately 20-fold higher AC load in the WA-AHS treatment under 70% RH was required for similar spore inactivation in that under saturated humidity. These results indicated that environmental humidity was also affected to the AHS treatment.
DISCUSSION

Hypochlorite solution have been investigated as an aerosolized disinfectant (Thornton et al., 2013; Martyny et al., 2005) because of high microbicidal activity, high safety (low toxicity) for human and easy affordability, especially when its pH was adjusted to be 5.5-6.5 (Ono et al., 2012). In this study, weakly acidic hypochlorite solution (pH6.5) was used for an aerosolized disinfectant and evaluated its microbicidal activity in our newly designed test-chamber system.

Microbicidal activity of AHS was evaluated by AC exposure and AC load, which were indicated as the amount of AC loaded onto a spore-attaching (SA) plate and the amount of AC loaded into the chamber, respectively. Generally, AC load is easily and exactly controlled during an AHS treatment in practical indoor space, and relationship between AC exposure and AC load become useful information. In our study, high correlation between AC exposure rate and AC loading rate was confirmed under WA-AHS treatment regardless of relative humidity (Fig. 3B). On the other hand, AC exposure rate in WA-AHS treatment (pH6.5) was markedly higher than those in B-AHS treatment (pH9.0). This result might indicate that AC in micro-droplets of WA-AHS was more retainable than B-AHS during the generation of AHS by ultrasonication and/or the AHS treatment. The retentivity of dissolved chlorine in aqueous solution was considered to be high in the following order: ClO > HClO > Cl2 (Fukuzaki 2006; Urano and Fukuzaki 2010), however our results were seemed inconsistent to that. Aerosolizing of hypochlorite solution increased the surface area of micro-droplet so that some material transfer was promoted, and might consequently promoted changes in composition within micro-droplet including volatilization of dissolved chlorine (Urano and Fukuzaki 2010). In other words, aerosolizing methods might be influenced dissolved hypochlorite concentration. However, further detailed investigation is necessary to confirm this hypothesis.

Generally, microbicidal activity of hypochlorite solution was higher under moderately acidic condition than alkaline condition, since the concentration of HClO with microbicidal activity was increased. It was, therefore, predicted that WA-AHS treatment was more effective than B-AHS treatment. In fact, however, B-AHS treatment was also effective likewise to WA-AHS on same AC exposure (Fig. 6A). It was proposed that long-time treatment of B-AHS might wipe the pH effect of hypochlorite microbicidal action out under the saturated humidity.

On the other hand, although AC exposure rate in WA-AHS treatment at 70% RH was equivalent to those in saturated humidity when the AC loading rate was same (Fig. 3B), the sporicidal activity of the former was remarkably weakened (Fig. 6). Permeability of AC into spores was influenced by or depend on moisture on the SA plate. The effect of moisture on antimicrobial activity of the agents has been also investigated by Santiesteban-López et al. (2007). Antibacterial susceptibilities of food-borne bacteria to individual and binary mixtures of potassium sorbate with natural phenolic compounds were influenced by pH and water activity. It was also predicted that lifetime of micro-droplet of AHS in low RH might be shortened and easily volatilized before attacking spores. Furthermore, if AHS treatment will be planned to achieve the spore reduction of 5 logs under more low humidity conditions, for example 50% RH, it is necessary to expose AHS with very high AC Load. Such a high AC load must be avoided in our test-chamber system, because of safety and damage of the test-chamber. Currently, a new test-chamber system is being designed to carry out AHS treatments under high AC load in safety.

Our study indicated that AHS have enough sporicidal activity under the saturated humidity, but, in fact, it will be difficult to supply micro-droplets of AHS to the entire space of the target room(s) homogeneously, since air quality and airflow in the room are temporally and spatially heterogeneous. Whereas, the test-chamber system used in this study was designed to reproduce an ideal space with a homogeneous air condition. Thereby, the data obtained using the chamber should contribute to predicting the required AC dose to treat a whole building by integrating the data temporally and spatially.

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