INTRODUCTION

Filtering facepiece respirators (FFR) should be discarded after use for one work shift to control infection, especially if they come into contact with airborne pathogens, such as Mycobacterium tuberculosis, or influenza virus. During the severe acute respiratory syndrome (SARS) epidemic outbreak, consumers' demand for N95 respirators increased owing to their high collection efficiency. During the outbreak of Middle East respiratory syndrome (MERS) in Korea, pharmacies in South Korea sold many times more N95 FFRs than usual. However, these FFRs are sometimes reused, especially during a shortage or when their distribution is delayed. Economic considerations may also apply. The price of certified FFRs, such as National Institute for Occupational Safety and Health (NIOSH)-approved N95 FFRs, typically exceeds those of non-certified masks. Affordability considerations favor reuse.
identified. Storage containers should be disposed of or cleaned regularly. The FDA defines three kinds of reuse: between patients with adequate reprocessing, reuse by the same person with adequate reprocessing/decontamination, and repeated use by the same person over a period with or without reprocessing.

Before FFRs are reused, they may be decontaminated to control the growth of microorganisms on them. However, whether a decontaminated N95 FFR can be reused is an issue that requires detailed consideration. In some cases, the use of chemical disinfectants may require that an employer train workers on protecting themselves against chemical hazards and on complying with OSHA’s Hazard Communication, 29 CFR 1910.1200, and other standards. Contaminated objects with porous surfaces that cannot be disinfected may have to be disposed of. All personnel, clothing, equipment, and samples that leave a contaminated area (generally referred to as the Exclusion Zone) must be decontaminated to remove any harmful chemicals or infectious organisms that may be attached to them. Decontamination methods (i) physically remove contaminants, (ii) inactivate contaminants by chemical detoxification or disinfection/sterilization, or (iii) remove contaminants by a combination of both physical and chemical methods. NIOSH has published a series of research articles on mask decontamination.

In selecting decontamination methods, both decontamination and protective capability are considered. For example, ultraviolet germicidal irradiation (UVGI) and bleach reportedly do not significantly reduce the protective capability (penetration by contaminants) of FFRs. Bergman et al. tested many methods, involving UVGI and bleach, and found that FFRs that were treated with these two decontaminants and control samples exhibited expected filter aerosol penetration (<5%) and filter airflow resistance. Physical damage varied with treatment method. Further research is needed before any particular decontamination methods can be recommended. Other chemical and energetic methods also have potential for decontaminating FFRs, but few studies have addressed the elimination of viable microorganisms from FFRs. UVGI had been reported to be effective in eliminating H5N1 or MS2 coliphages from FFRs. UVGI had been reported to be effective in eliminating H5N1 or MS2 coliphages from FFRs.

The optimal dosages of decontamination methods are important for determining a comprehensive infection control strategy. Our work addresses the potential for cross-contamination of reused respirators with a view to overcoming FFR shortages and so to increase capacity for controlling future outbreaks.

### Practical Implications
- The survival of bacteria of reclaimed National Institute for Occupational Safety and Health-certified N95 filtering facepiece respirators (FFRs) after decontamination is important, especially for healthcare workers.
- Safe respirator usage after decontamination using various methods improves infection control and protection against biohazards.
- The optimal dosages of decontamination methods are important for determining a comprehensive infection control strategy.

### 2 | MATERIALS AND METHODS

#### 2.1 | Test system and decontamination methods

The main test variable in this study is the survival of bacteria that were loaded on N95 FFRs that were decontaminated by various methods under worst-case temperature and humidity, which prevail when an FFR is placed in a zipper bag in a healthcare worker’s pocket with the goal of preventing cross-contamination, and touching of the respirator. In the experiment, B. subtilis spores were the tested microbial strain; a six-jet Collison nebulizer (BGI, Waltham, MA) sprayed the spores into a test system, shown in Figure 1, where they were loaded on N95 FFRs by suction to simulate the respiratory
flow of workers during intensive activity. The experimental FFR was an N95 FFR (8210, 3M, St. Paul, MN), certified by NIOSH. It was divided into six pieces, to which five decontamination methods were applied; they involved ethanol, bleach, UV, an autoclave, and a traditional electric rice cooker (TERC), made in Taiwan, without steam. The treatment proceeded as follows.

- **Ethanol**: Ethanol with various concentrations and volumes was added to the center of the surface of the N95 FFR using a pipette, the FFR was then dried in a petri dish that was placed in a biosafety cabinet (BSC) for 10 minutes.
- **Bleach**: A 0.4 mL volume of bleach with various concentrations (5.4% (w/w) as Cl₂: original; 2.7%: one part bleach to one part of deionized water; 0.54%: one part bleach to nine parts of deionized water) was added to the center of the surface of the N95 FFR using a pipette, the FFR was then dried in a petri dish in a BSC for 10 minutes.
- **UV**: An N95 FFR was placed 10 cm below a 6 W handheld UV lamp (model UVGL-58, VUP LLC, Upland, CA) that emitted a wavelength of 254 nm (UVC, 18.9 mW/cm²) or 365 nm (UVA, 31.2 mW/cm²). Both sides of each N95 FFR were exposed for different times - 1, 2, 5, 10 and 20 minutes - in a BSC. The UV intensity was measured using a handheld laser power and energy meter (OPHIR NOVAII, model Nova II PD300-UV) and was reported as a mean of five measurements over a 10 × 10 mm aperture with a swivel mount and a removable filter.
- **Autoclave**: The N95 FFR was heated for 15 minutes at 121°C and 103 kPa.
- **TERC**: The N95 FFR was placed in an electric rice cooker for dry heating for 3 minutes (149-164°C, without added water).

### 2.2 Sampling procedure

Each N95 FFR was placed into the system (Figure 1) for 30 minutes of bacterial bioaerosol sampling. The respiratory flow (85 L/min) of workers during high-intensity activities was used, and the face velocity for the whole N95 FFR was calculated as 8.3 cm/s. The N95 FFRs were cut into pieces with a diameter of 45 mm. Each had an effective diameter of 40 mm and a filtration area of 12.6 cm². The sampling flow rate of the pump was 6.3 L/min, which produced the desired face velocity.

*Bacillus subtilis* prototype strains (CCRC 12145, Taiwan Food Industry Research and Development Institute) were used to prepare an endospore suspension liquid for generating bacterial bioaerosols. The suspension was centrifuged at 1917 g for 5 minutes. The supernatant was discarded and the pellet was resuspended in sterile distilled water. This washing process was repeated twice, and spores were resuspended in approximately 55 mL of sterile distilled water to yield a uniform mixture which was poured into the Collison nebulizer. The spores were aerosolized at a pressure of 25 psi when the dilution air flow rate was 80 L/min, as presented in Figure 1. The stability of the bioaerosol concentration in the system was verified using an Andersen single-stage sampler (Andersen Inc., Atlanta, GA).

The aqueous packing density ($\alpha_{aq}$) of the retained liquid decontaminants was modified that in a previous report, in which $\alpha_{aq}$ was the volume fraction of the filter; it is calculated using the equation,

$$\alpha_{aq} = \frac{V_{aq}}{V_f}$$  \hspace{1cm} (1)

where $V_{aq}$ is the volume of liquid disinfectant that was spiked onto the test N95 FFR. The volume of the test N95 FFR ($V_f$) was 1.84 mL, which was estimated from the volume of water that was displaced by it. When 0.15, 0.4, 0.8, and 1.6 mL of ethanol were spiked onto the surface of the test N95, $\alpha_{aq}$ values of 0.082, 0.23, 0.44, and 0.87, respectively, were obtained.

After spores were loaded onto the FFRs for 30 minutes, the FFRs were decontaminated using one of the five aforementioned methods, and then placed in an incubator (Model: HONG-YU, HRM-80, Taichung, Taiwan) at the worst-case scenario temperature of 37°C (similar to body temperature) and 95% RH (the maximum feasible RH value), respectively, for 24 hours for another day of usage. Each batch test was conducted in triplicate. Figure 2 displays the sampling procedure.

After decontamination, elution was performed. It involved placing the test filter in a 50 mL centrifuge tube and then adding 20 mL of distilled water. The filter was vortexed for 2 minutes to ensure complete elution of all aerosolized bacterial spores. The eluent was then filtered through a 0.22 μm filter and subjected to colony counting using a Petri dish containing thallium acetate agar, as described in a previous study. Four sets of filter holders and pumps were used for each experiment, with a flow rate of 6.28 L/min for each pump and a face velocity of 8.3 cm/s.

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**FIGURE 1** Experimental system setup. A filter holder contains one piece of N95 FFR.
of sterilized water to soak the filters completely. A centrifuge was used to recover the loaded spores by desorption from the FFRs at a centrifugation speed of 3500 rpm for 10 minutes, followed by 1 minutes of vortexing. The centrifuged and vortexed suspension (0.1 mL) was uniformly applied to the TSA and then placed in an incubator for 24 hours. Colony forming units (CFUs) were counted and their relative survival (RS) was calculated as follows.

$$RS = \frac{C_f}{C_i} \times 100\%$$

where $C_f$ is the number of CFUs after decontamination and $C_i$ is the number of CFUs before decontamination (Figure 2).

3 | RESULTS

Figure 3 presents the decontaminating effect of 0.4 mL ($\alpha_{aq} = 0.23$) ethanol at various concentrations on the RS of $B. \text{subtilis}$ spores that were loaded on the N95 FFR. An RS of $89 \pm 6\%$ was obtained after spiking with 50% ethanol, and $73 \pm 5\%$ was obtained after spiking with 70% ethanol. The lowest RS of $68 \pm 3\%$ was obtained when the concentration of ethanol was 80%. The result that was obtained using 95% ethanol (RS = $73 \pm 7\%$) was close to that obtained using 70% ethanol although the samples that were spiked with 95% ethanol sometimes yielded slightly higher values of RS than were obtained using the 80% ethanol samples. An RS of $59 \pm 8\%$ was obtained in 24 hours without decontamination. The 50%, 70%, 80%, and 95% ethanol-treated samples had RS values of $33 \pm 8\%$, $22 \pm 8\%$, $20 \pm 2\%$ and $26 \pm 7\%$ after 24 hours, respectively.

Figure 4 shows the effect of 70% ethanol on the RS of $B. \text{subtilis}$ spores. Just after spiking with ethanol, the RS was found to have declined from 100% to 68%-75%. When 0.4 mL ($\alpha_{aq} = 0.23$) of 70% ethanol was applied, the RS fell to 22% in 24 hours. The RS fell to 20% when 80% ethanol was used.

In the bleach decontamination test, no colony was recovered after 5.4%, 2.7% or 0.54% NaOCl was used, constituting no dilution, twofold, and 10-fold dilution, respectively (Figure 5). This study found that NaOCl, even when diluted 10-fold from standard bleach, had a strong decontamination effect, with a 100% bactericidal effect.

Similar results were achieved using UVC. No colony was recovered after exposure to UVC for as little as 5 minutes (Figure 6). However, RS remained above 20% after 20 minutes of irradiation by UVA, exponentially decaying with increased exposure time (Figure 6).

Figure 7 presents the RS values that were achieved using the five decontamination methods. Four of the methods - involving 0.54% NaOCl, UVC, an autoclave, and a TERC - effectively sterilized almost 100% of the bacteria. Decontamination with 70% ethanol yielded an initial RS of approximately 75% and an RS that remained above 20% after 24 hours of storage.

4 | DISCUSSION

Decontamination using ethanol yielded higher RS of spores than the other four decontamination methods (Figure 7). The other four
methods yielded RS values of close to zero, indicating effective sterilization. The biosafety manual that was published by the World Health Organization (WHO) notes that alcohols are effective against vegetative bacteria but not spores. The US CDC also states that alcohols can eliminate many or even all pathogenic microorganisms except for bacterial spores. Our findings are consistent with the recommendations of the WHO and the US CDC.

In this study, 59 ± 8% of the loaded spores survived on N95 FFRs for 24 hours without decontamination. This result is consistent with some previous studies of the survival rates of *B. subtilis*.
spores at 37°C and an RH of 85%\textsuperscript{21} or 95%\textsuperscript{21}. Moreover, the results of Sagripanti and Bonifacino\textsuperscript{32} suggest that the nature of the challenged surface may affect the sporicidal activity of some chemical agents. For example, a study by Li et al revealed that the average values of RS (%) that were obtained by elution from a Nuclepore filter for hardy \textit{B. subtilis} were 64% and 48% at sampling times of 1 and 30 minutes, respectively. The average RS (%) values of \textit{B. subtilis} from a gelatin filter were 128% and 108% at sampling times of 1 and 30 minutes, respectively\textsuperscript{26}. Our results were comparable to those obtained for Nuclepore filter samples.

The WHO biosafety manual mentions that alcohols should be used at concentrations of approximately 70% (v/v) in water to
maximize their germicidal effectiveness. The WHO and the US CDC both recommend the use of 70% alcohol.\textsuperscript{15,30} The US CDC also notes that the biocidal activity of alcohol diminishes sharply at dilutions of weaker than 50% (v/v), and the optimal bactericidal concentration is 60%-90% (v/v).\textsuperscript{15} From the result of quadratic polynomial regression, the lowest RS occurred at the 83% and 76% alcohol for the initial and 24 hours samples, respectively (Figure 3). The RS values of approximately 70%-95% (v/v) (Figure 3) that were obtained in this study support the US CDC’s recommendation.

The optimal bactericidal concentrations for various microorganisms may vary.\textsuperscript{15} Pseudomonas aeruginosa was destroyed by ethanol at concentrations of 30%-100%, and Serratia marcescens, E. coli, and Salmonella typhosa were destroyed by ethanol at concentrations of 40%-100%. Gram-positive organisms, such as Staphylococcus aureus and Streptococcus pyogenes, were slightly more resistant to ethanol, being destroyed by ethanol concentrations of 60% to 95%. When the effect of ethanol against M. tuberculosis was evaluated, 95% ethanol was found to kills the tubercle bacilli in water or sputum suspension within 15 seconds. As the challenge bioaerosol in the current study is B. subtilis spore, the RS remained at 75 ± 17% (Figure 7) indicating that the B. subtilis spores were more resistant than those mentioned above, including tubercle bacilli. Ethanol does not affect the viability of Bacillus spores in current guidance,\textsuperscript{1,10,15,30} and a comprehensive examination of disinfection against different target microorganisms should therefore be performed.

The average RS (%) values for hardy B. subtilis that were obtained by elution from FFRs decayed to 23 ± 8% in 24 hours of treatment with 70% ethanol but to 59 ± 8% without ethanol (Figure 7). The decay to 59% was caused mainly by the nature of the FFR surface and effects of storage. The combined effect of FFR surface, storage, and ethanol treatment might have been expected to yield an RS of 44% (59% × 75%), but was only 22% in this study. As ethanol should not have this much of an effect on spores,\textsuperscript{1,10,15,30} the nature of the FFR surface, treatment with ethanol, and the storage conditions may have an interaction effect. Therefore, the mechanism by which ethanol affects the amounts of spores that survive on FFRs should be further investigated.

Figure 4 presents the effect of the $\alpha_{aq}$ of 70% ethanol on the RS of B. subtilis spores. When $\alpha_{aq}$ exceeded 0.23, the initial RS was around 70% and that after 24 hours was about 20%. Lin et al\textsuperscript{21} found that when 1.5 mL of artificial saliva was dropped onto an N95, its surface tension caused it to form a sphere-like droplet that was attached to the hydrophobic first layer, before it slowly penetrated the second and third layers. However, in the present study, which is based on observation, 70% ethanol penetrated the filter rapidly and quickly evaporated to the air. Accordingly, increasing $\alpha_{aq}$ had little effect on the RS of B. subtilis spores and may have been responsible for the rapid evaporation of 70% ethanol.

About 60%-70% of the B. subtilis spores that were loaded on N95 FFRs survived after 24 hours of storage without decontamination, whereas only approximately 20% of spores retained their culturability after 20 minutes of irradiation by UVA (Figure 6), whose disinfection effect was comparable to that of ethanol (Figure 4). From the result of exponential decay regression, the half-life (the value of RS reduces to 50%) was 10 and 0.17 minutes for the UVA and UVC irradiation, respectively (Figure 6). Although UVA could not decontaminate as effectively as UVC, it did have some decontaminating effect. This finding warrants further study.

The results in our study verify the biocidal efficacy of bleach (0.54% NaOCl), UVC, and an autoclave, which are well known means of sterilization.\textsuperscript{15,30} Interestingly, the TERC exhibited biocidal efficacy as a sterilizing device. In the WHO biosafety manual,\textsuperscript{30} heat is
regarded as one of the most commonly used physical agents for decontamination against pathogens. “Dry” heat, which is non-corrosive, is applied to many items of laboratory-ware, which can withstand temperatures of at least 160°C for 2-4 hours. In this study, the TERC is used as a dry heating device and was found to exhibit a biocidal efficacy that reaches effective sterilization in 3 minutes. The results achieved using the TERC provide useful information regarding effective means for decontaminating and reusing FFRs.

Notably, when an N95 FFR is reused, the biocidal efficacy of the decontamination treatment, filter quality, fit factor (which is affected by physical damage to the frame or rubber strap), and toxic residual chemicals on FFR must all be considered. For example, bleach can harm the wearer if not properly used to decontaminate an N95 FFR before reuse. Safe disposal of spent bleach is important, and users may decide to neutralize the microbial activity of the bleach before disposal. Solutions can be neutralized by reaction with chemicals such as sodium bisulfite, or glycine. Considering the potential health risks, the method of decontamination using bleach must be modified such as by the use of chemical methods for neutralizing residuals.

The RS is a function of decontamination and the biological characteristics of pathogens. The filter quality combines penetration and pressure drop and is affected by the physical characteristics of the FFR. However, this study focused on RS because it is a useful metric for quantifying sterilization or degree of disinfection. In summary, bleach, UVC, the autoclave, and the TERC provide effective sterilization. However, ethanol and UVA are ineffective and not cleared as active means for decontaminating and reusing FFRs.

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