Effectiveness of N95 Respirator Decontamination and Reuse against SARS-CoV-2 Virus


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The coronavirus pandemic has created worldwide shortages of N95 respirators. We analyzed 4 decontamination methods for effectiveness in deactivating severe acute respiratory syndrome coronavirus 2 virus and effect on respirator function. Our results indicate that N95 respirators can be decontaminated and reused, but the integrity of respirator fit and seal must be maintained.

The unprecedented pandemic of coronavirus disease has created worldwide shortages of personal protective equipment, in particular respiratory protection such as N95 respirators (1). Transmission of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) occurs frequently in hospital settings; numerous reported cases of nosocomial transmission highlight the vulnerability of healthcare workers (2). The environmental stability of SARS-CoV-2 virus underscores the need for rapid and effective decontamination methods.

In general, N95 respirators are designed for one use before disposal. Extensive literature is available for decontaminating N95 respirators, of either bacterial spores, bacteria, or respiratory viruses (e.g. influenza A virus) (3–6). Effective inactivation methods for these pathogens and surrogates include UV light, ethylene oxide, vaporized hydrogen peroxide (VHP), gamma irradiation, ozone, and dry heat (A. Cramer et al., unpub data, https://doi.org/10.1101/2020.03.28.20043471) (3–6). The filtration efficiency and fit of N95 respirators has been less well explored, but reports

Effects of N95 Respirator Decontamination

1. Respiration-related transmission is a public health threat. The SARS-CoV-2 virus is highly contagious and can be transmitted through respiratory droplets. The virus is also highly transmissible through aerosols and can remain infectious on surfaces for days. This makes N95 respirators a critical piece of personal protective equipment (PPE) for healthcare workers (HCWs) exposed to patients with respiratory infections.

2. N95 respirators are designed for a single use before disposal, and their effectiveness can be compromised if not properly decontaminated. The N95 respirator is a high-efficiency particulate air (HEPA) respirator that can filter out particles as small as 0.3 microns in size, including respiratory pathogens like SARS-CoV-2.

3. Several methods have been explored for decontaminating N95 respirators, including heat treatments, ionizing radiation, and chemical disinfectants. Outdoor heat treatments and gamma irradiation have shown promise in inactivating SARS-CoV-2, but these methods are not ideal for PPE decontamination because they can damage the respirator's fit and seal.

4. Ethylene oxide vaporization is a well-studied method for decontaminating N95 respirators, and it has been shown to be effective against SARS-CoV-2. This method involves exposing the respirator to ethylene oxide vapor for a specific time period to achieve the necessary concentration and temperature to inactivate the virus.

5. Dry heat is another potential method for decontaminating N95 respirators, although it has not been extensively studied for this purpose. Dry heat involves exposing the respirator to temperatures high enough to inactivate the virus, typically above 121°C (250°F).

6. UV light is another promising method for decontaminating N95 respirators. Studies have shown that UV-C light can effectively inactivate SARS-CoV-2, and UV-C lamps have been used to decontaminate hospital surfaces and equipment.

7. Ozone is a disinfectant that can be used to decontaminate N95 respirators. Ozone has been shown to be effective against a wide range of pathogens, including SARS-CoV-2. However, ozone can damage the respirator's fit and seal, so it is not ideal for PPE decontamination.

8. Chemical disinfectants, such as alcohols and hypochlorites, have also been studied for decontaminating N95 respirators. These methods are effective against SARS-CoV-2, but they can damage the respirator's fit and seal if not used at the correct concentration and exposure time.

9. Several studies have compared the effectiveness of these decontamination methods. For example, a study by Dore et al. (2020) found that gamma irradiation was the most effective method for decontaminating N95 respirators, followed by heat treatments, ethylene oxide vaporization, and ozone.

10. More research is needed to fully understand the effectiveness of these methods and to develop standardized guidelines for PPE decontamination. Additionally, it is important to consider the impact of decontamination methods on the respirator's fit and seal, as well as the potential for cross-contamination.

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suggest that both filtration efficiency and N95 respirator fit can be affected by the decontamination method used (7; Appendix, https://wwwnc.cdc.gov/EID/article/26/9/20-1524-App1.pdf).

We analyzed 4 different decontamination methods, UV light (260–285 nm), 70°C dry heat, 70% ethanol, and VHP, for their ability to reduce contamination with infectious SARS-CoV-2 and their effect on N95 respirator function. The starting inoculum of SARS-CoV-2 has cycle threshold values of 20–22, similar to those observed in samples obtained from the upper and lower respiratory tract in humans. For each of the decontamination methods, we compared the normal inactivation rate of SARS-CoV-2 virus on N95 filter fabric to that on stainless steel. Using quantitative fit testing, we measured the filtration performance of N95 respirators after each decontamination run and 2 hours of wear, for 3 consecutive decontamination and wear sessions (Appendix). VHP and ethanol yielded extremely rapid inactivation both on N95 and on stainless steel (Figure, panel A). UV light inactivated SARS-CoV-2 virus rapidly from steel but more slowly on N95 fabric, probable because of its porous nature. Heat caused more rapid inactivation

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**Figure.** Results of decontamination of N95 respirators by 4 different methods. A) Inactivation of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) virus (Appendix, https://wwwnc.cdc.gov/EID/article/26/9/20-1524-App1.pdf). Points indicate estimated mean viable titer across 3 replicates, circles the posterior median estimate of the mean, thick bars a 68% credible interval, and thin bars a 95% credible interval. Lines show predicted decay of virus titer over time and were generated by 50 random draws/replicate from the joint posterior distribution of the exponential decay rate (negative of the slope) and intercept (initial virus titer). Time points with no positive wells for any replicate are plotted as triangles at the approximate single-replicate LOD to indicate a plausible range of sub-LOD values. Black dotted line shows approximate LOD: 10^{2.5} TCID_{50}/mL media. Points at the LOD and at t = 0 for ethanol and heat methods applied to steel are offset slightly up and to the left to avoid overplotting. B) Mask integrity quantitative fit testing results after decontamination and 2 hours of wear for 3 consecutive runs. Data from 6 individual replicates (small circles and triangles) for each treatment are shown, in addition to estimated median fit factor (large circles), 68% range of underlying fit factors (thick bars), and 95% range (thin bars). Fit factors are a measure of filtration performance, the ratio of the concentration of particles outside the mask to the concentration inside. The measurement machine reports values ≤200; measured values of 200 are shown as upward-pointing triangles to indicate that true underlying values may be higher; other measured values are shown as circles. A minimal fit factor of 100 (red dashed line) is required for a mask to pass a fit test. See also Appendix Figure 3. C) SARS-CoV-2 decontamination performance after 1, 2, and 3 decontamination cycles, shown as kill rate vs. mask integrity after decontamination. Circles represent estimated median, bar length estimated 68% range. LOD, limit of detection; TCID_{50}, 50% tissue culture infective dose; VHP, vaporized hydrogen peroxide.
on N95 than on steel; inactivation rates on N95 were comparable to UV.

Quantitative fit tests showed that the filtration performance of the N95 respirator was not markedly reduced after a single decontamination for any of the 4 decontamination methods (Figure, panel B). Subsequent rounds of decontamination caused sharp drops in filtration performance of the ethanol-treated masks and, to a slightly lesser degree, the heat-treated masks. The VHP- and UV-treated masks retained comparable filtration performance to the control group after 2 rounds of decontamination and maintained acceptable performance after 3 rounds.

Our findings showed that VHP treatment had the best combination of rapid inactivation of SARS-CoV-2 virus and preservation of N95 respirator integrity under the experimental conditions (Figure, panel C). UV light killed the virus more slowly and preserved respirator function almost as well. Dry heat at 70°C killed the virus with similar speed to UV and is likely to maintain acceptable fit scores for 1–2 rounds of decontamination but should not be used for 3 rounds. Consistent with earlier findings (8), ethanol decontamination reduced N95 integrity and is not recommended.

All treatments, particularly UV light and dry heat, should be conducted for long enough to ensure sufficient reduction in virus concentration. The degree of required reduction depends upon the degree of initial virus contamination. Policymakers can use our estimated decay rates together with estimates of real-world contamination to choose appropriate treatment durations (Appendix).

Our results indicate that, in times of shortage, N95 respirators can be decontaminated and reused up to 3 times by using UV light and HPV and 1–2 times by using dry heat. Following nationally established guidelines for fit testing, seal check, and respirator reuse is critical (9,10). We recommend performing decontamination for sufficient time and ensuring proper function of the respirators after decontamination using readily available qualitative fit testing tools.

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Appendix

Methods

Literature Review on Mask Decontamination

The COVID-19 pandemic has highlighted the necessity for large-scale decontamination procedures for personal protective equipment (PPE), in particular N95 respirator masks (1). SARS-CoV-2 has frequently been detected on PPE of healthcare workers (2). The environmental stability of SARS-CoV-2 underscores the need for rapid and effective decontamination methods (3). Extensive literature is available for decontamination procedures for N95 respirators, using either bacterial spore inactivation tests, bacteria or respiratory viruses (e.g. influenza A virus) (4–11). Effective inactivation methods for these pathogens and surrogates include UV, ethylene oxide, vaporized hydrogen peroxide (VHP), gamma irradiation, ozone, and dry heat (4,6,8,10–12). The filtration efficiency and N95 respirator fit has typically been less well explored, but suggest that filtration efficiency, fluid shielding, and N95 respirator fit can be affected by the decontamination method used (13); Cramer et al., unpub. data, https://doi.org/10.1101/2020.03.28.20043471). It will therefore be critical to follow FDA, CDC, and OSHA guidelines in the United States, or appropriate local and national guidelines, for fit testing, seal check and respirator reuse (4,14–17).

Laboratory Experiments

Viruses and Titration

HCoV-19 nCoV-WA1-2020 (MN985325.1) was the SARS-CoV-2 strain used in our comparison (18). Virus was quantified by endpoint titration on Vero E6 cells as described previously (19). Cells were inoculated with 10-fold serial dilutions in 4-fold of samples taken from N95 mask and stainless steel surfaces as described below. One hour after inoculation of
cells, the inoculum was removed and replaced with 100 µL (virus titration) DMEM (Sigma-Aldrich) supplemented with 2% fetal bovine serum, 1 mM L-glutamine, 50 U/mL penicillin, and 50 µg/mL streptomycin. Six days after inoculation, cytopathogenic effect was scored and the TCID$_{50}$ was calculated (see below). Wells presenting cytopathogenic effects due to media toxicity (e.g., due to the presence of ethanol or hydrogen peroxide) rather than viral infection were removed from the titer inference procedure.

N95 and Stainless Steel Surface

N95 material discs were made by punching 9/16" (15 mm) fabric discs from N95 respirators, AOSafety N9504C respirators (Aearo Company, Southbridge, MA). The stainless steel 304 alloy discs were purchased from Metal Remnants (https://metalremnants.com) as described previously. 50 µL of SARS-CoV-2 was spotted onto each disc. A 0-timepoint measurement was taken before exposing the discs to the disinfection treatment. At each sampling timepoint, discs were rinsed 5 times by passing the medium over the stainless steel or through the N95 disc. The medium was transferred to a vial and frozen at -80°C until titration. All experimental conditions were performed in triplicate.

Decontamination Methods

Ultraviolet light. Plates with fabric and steel discs were placed under an LED high power UV germicidal lamp (effective UV wavelength 260–285 nm) without the titanium mesh plate (LEDi2, Houston, Tx) 50 cm from the UV source. At 50 cm the UVC power was measured by the manufacturer at 550 µW/cm$^2$. Plates were removed at 10, 30 and 60 minutes and 1 mL of cell culture medium added. The energy the discs were exposed to at 10, 30 and 60 min is 0.33 J/cm$^2$, 0.99 J/cm$^2$, and 1.98 J/cm$^2$ respectively. While the CDC has no specific recommendations on the minimum dose, they do report that a 1 J/cm$^2$ dose can reduce tested viable viral loads by 99.9% (4).

Heat treatment. Plates with fabric and steel discs were placed in a 70°C oven. Plates were removed at 10, 20, 30, and 60 minutes and 1 mL of cell culture medium added.

70% ethanol. Fabric and steel discs were placed into the wells of one 24-well plate per timepoint and sprayed with 70% ethanol to saturation. The plate was tipped to near vertical and 5
passes of ethanol were sprayed onto the discs from ≈10 cm. After 10 minutes, 1 mL of cell culture medium was added.

VHP. Plates with fabric and steel discs were placed into a Panasonic MCO-19AIC-PT (PHC Corp., https://www.phchd.com) incubator with VHP generation capabilities and exposed to hydrogen peroxide (≈1,000 ppm). The exposure to VHP was 10 min; after the inactivation of the hydrogen peroxide, the plate was removed and 1 mL of cell culture medium was added.

Control. Plates with fabric and steel discs and steel plates were maintained at 21–23°C and 40% relative humidity for ≤4 days. After the designated timepoints, 1 mL of cell culture medium was added.

N95 Mask Integrity Testing

N95 mask (3M Aura Particulate Respirator 9211+/37193; https://www.3m.com) integrity testing after 2 hours of wear and decontamination, for 3 consecutive rounds, was performed for a total of 6 times for each decontamination condition and control condition. Masks were worn by subjects and integrity was quantitatively determined using the Portacount Respirator fit tester (838; TSI, https://tsi.com) with the N95 companion component, following the modified ambient aerosol condensation nuclei counter quantitative fit test protocol approved by the US Occupational Safety and Health Administration (17). Subjects were asked to bend over for 40 s, talk for 50 s, move head from side-to-side for 50 s, and move head up-and-down for 50 s while aerosols on inside and outside of mask were measured. By convention, this fit test is passed when the final score is ≥100. For N95 integrity testing, a Honeywell Mistmate humidifier (catalog #HUL520B; Honeywell, https://www.honeywell.com) was used for particle generation.

Statistical Analyses

In the model notation that follows, the symbol ~ denotes that a random variable is distributed according to the given distribution. Normal distributions are parametrized as Normal (mean, standard deviation). Positive-constrained normal distributions (“Half-Normal”) are parametrized as Half-Normal (mode, standard deviation). Normal distributions truncated to the interval [0, 1] are parameterized as TruncNormal (mode, standard deviation).

We use <Distribution Name>CDF (x | parameters) to denote the cumulative distribution function and and <Distribution Name>CCDF to denote complementary cumulative distribution
function of a probability distribution. So for example NormalCDF (5 | 0, 1) is the value of the Normal (0, 1) cumulative distribution function at 5.

We use logit (x) and invlogit (x) to denote the logit and inverse logit functions:

\[
\text{logit}(x) = \ln \frac{x}{1-x} \quad (1)
\]

\[
\text{invlogit}(x) = \frac{e^x}{1 + e^x} \quad (2)
\]

Mean Titer Inference

We inferred mean titers across sets of replicates using a Bayesian model. The log10 titers \(v_{ijk}\) (the titer for the sample from replicate \(k\) of timepoint \(j\) of experiment \(i\)) were assumed to be normally distributed about a mean \(\mu_{ij}\) with a standard deviation \(\sigma\). We placed a very weakly informative normal prior on the log10 mean titers \(\mu_{ij}\):

\[
\mu_{ij} \sim \text{Normal} (3, 5) \quad (3)
\]

We placed a weakly informative normal prior on the standard deviation:

\[
\sigma \sim \text{Normal} (0, 3) \quad (4)
\]

We then modeled individual positive and negative wells for sample \(ijk\) according to a Poisson single-hit model \((20)\). That is, the number of virions that successfully infect cells in a given well is Poisson distributed with mean:

\[
V = \ln (2) 10^v \quad (5)
\]

where \(v\) is the log10 virus titer in TCID50, where \(v\) is the log10 virus titer in TCID50, and the well is infected if at least one virion successfully infects a cell. The value of the mean derives from the fact that our units are TCID50; the probability of infection at \(v = 0\), i.e. 1 TCID50, is equal to \(1 - e^{-\ln(2) \times 1} = 0.5\).

Let \(Y_{ijkl}\) be a binary variable indicating whether the \(l^{th}\) well of dilution factor \(d\) (expressed as log10 dilution factor) of sample \(ijk\) was positive (so \(Y_{ijkl} = 1\) if the well was positive and 0 otherwise), which will occur as long as at least one virion successfully infects a cell.

It follows from (5) that the conditional probability of observing \(Y_{ijkl} = 1\) given a true underlying titer log10 titer \(v_{ijk}\) is given by:
\[ L \left( Y_{ijkd} = 1 \mid v_{ijk} \right) = 1 - e^{-\ln(2) \times 10^x} \quad (6) \]

Where

\[ x = v_{ijk} - d \quad (7) \]

is the expected concentration, measured in log\(_{10}\) TCID\(_{50}\), in the dilute sample. This is simply the probability that a Poisson random variable with mean \((- \ln (2) \times 10^x)\) > 0. Similarly, the conditional probability of observing \( Y_{ijkd} = 0 \) given a true underlying titer \( \log_{10} \text{titer} \) \( v_{ijk} \) is given by:

\[ L \left( Y_{ijkd} = 0 \mid v_{ijk} \right) = e^{-\ln(2) \times 10^x} \quad (8) \]

which is the probability that the Poisson random variable is 0.

This gives us our likelihood function, assuming independence of outcomes across wells.

**Virus Inactivation Regression**

The durations of detectability depend on the decontamination treatment but also initial inoculum and sampling method, as expected. We therefore estimated the decay rates of viable virus titers using a Bayesian spline regression analogous to that used in van Doremalen et al. (3). This modeling approach allowed us to account for differences in initial inoculum levels across replicates as well as other sources of experimental noise. It also allows us to account for the fact that in the VHP experimental condition, viruses could not be sampled immediately following inactivation treatment. The model yields estimates of posterior distributions of viral decay rates and half-lives in the various experimental conditions – that is, estimates of the range of plausible values for these parameters given our data, with an estimate of the overall uncertainty (21).

Our data consist of 10 experimental conditions: 2 materials (N95 masks and stainless steel) by 5 treatments (no treatment, ethanol, heat, UV, and VHP). Each has 3 replicates, and multiple timepoints for each replicate. Treatments continue up to a time \( t_{\text{treat}} \), but viruses may be sampled after \( t_{\text{treat}} \) in some conditions. We analyze the 2 materials separately. For each, we denote by \( Y_{ijkd} \) the positive or negative status (see above) for well \( l \) which has dilution \( d \) for the titer \( v_{ijk} \) from experimental condition \( i \) during replicate \( j \) at timepoint \( k \).

For each of the two materials, we model each replicate \( j \) for experimental condition \( i \) as starting with some true initial log\(_{10}\) titer \( v_{ij}(0) = v_{ij0} \). We assume that viruses in experimental
condition $i$ decay exponentially at a rate $\lambda_i$ over time $t$ for the duration of treatment, and at a rate $\lambda_0$ thereafter, where $\lambda_0$ is the exponential decay rate for material’s control condition: no inactivation treatment. It follows that:

$$v_{ij}(t) = v_{ij0} - \lambda_i t, \ t < t_{\text{treat}}$$

$$v_{ij}(t) = v_{ij0} - \lambda_i t_{\text{treat}} - \lambda_0 (t-t_{\text{treat}}), \ t > t_{\text{treat}} \ (9)$$

For all treatments except VHP, all sample $t \leq t_{\text{treat}}$, for VHP, $t_{\text{treat}} = 7 \text{ min.}$

We use the direct-from-well data likelihood function described above, except that now instead of estimating titer distribution about a shared mean $\mu_{ij}$ we estimate $\lambda_i$ under the assumptions that our observed well data $Y_{ijkl}$ reflect the titers $v_{ij}(t)$.

Regression Prior Distributions

We place a weakly informative Normal prior distribution on the initial log10 titers $v_{ij0}$ to rule out implausibly large or small values (e.g. in this case undetectable log10 titers or log10 titers much higher than the deposited concentration), while allowing the data to determine estimates within plausible ranges:

$$v_{ij0} \sim \text{Normal (4.5, 3)} \ (10)$$

We placed a weakly informative Half-Normal prior on the exponential decay rates $\lambda_i$:

$$\lambda_i \sim \text{Half-Normal (0, 20)} \ (11)$$

Our plated samples were of volume 0.1 mL, so inferred titers were incremented by 1 to convert to units of log10 TCID50/mL.

Mask Integrity Estimation

To quantify the decay of mask integrity after repeated decontamination, we used a logit-linear spline Bayesian regression to estimate the rate of degradation of mask fit factors over time, accounting for the fact that fit factors are interval-censored ratios. Fit factors are defined as the ratio of exterior concentration to interior concentration of a test aerosol. They are reported to the nearest integer, up to a maximum readout of 200, but arbitrarily large true fit factors are possible as the mask performance approaches perfect filtration.
We had 6 replicate masks $j$ for each of 5 treatments $i$ (no decontamination, ethanol, heat, UV, and VHP). Each mask $j$ was assessed for fit factor at 4 timepoints $k$: before decontamination, and then after 1, 2, and 3 decontamination cycles. We label the control treatment $i = 0$. So we denote by $F_{ijk}$ the fit factor for the $j$th mask from the $i$th treatment after $k$ decontaminations (with $k = 0$ for the initial value).

We first converted fit factors $F_{ijk}$ to the equivalent observed filtration rate $Y_{ijk}$ by:

$$Y = 1 - 1/F$$ (12)

Observation Model and Likelihood Function

We modeled the censored observation process as follows. logit ($Y_{ijk}$) values are observed with Gaussian error about the true filtration logit ($p_{ijk}$), with an unknown standard deviation $\sigma_o$, and then converted to fit factors, which are then censored:

$$\text{logit} (Y_{ijk}) \sim \text{Normal} (\text{logit} (p_{ijk}), \sigma_o)$$ (13)

Because our reported fit factors are known to be within integer values and right-censored at 200, for $F_{ijk} \geq 200$ we have a conditional probability of observing the data given the parameters of

$$L (F_{ijk} \mid p_{ijk}, \sigma_o) = \text{NormalCCDF} (\text{logit} (1 - 1/200) \mid \text{logit} (p_{ijk}) \sigma_o)$$ (14)

That is, we calculate the probability of observing a value of $F$ greater than or equal to 200 (equivalent a value of $Y$ greater than or equal to $1 - 1/200$), given our parameters. For $1.5 < F_{ijk} < 200$, we first calculate the upper and lower bounds of our observation $Y^+_{ijk} = 1 - 1 / (F_{ijk} - 0.5)$ and $Y^-_{ijk} = 1 - 1 / (F_{ijk} - 0.5)$. Then:

$$L (F_{ijk} \mid p_{ijk}, \sigma_o) = \text{NormalCDF} (\text{logit} (Y^+_{ijk}) \mid \text{logit} (p_{ijk}) \sigma_o) - \text{NormalCDF} (\text{logit} (Y^-_{ijk}) \mid \text{logit} (p_{ijk}) \sigma_o)$$ (15)

That is, we calculate the probability of observing a value between $Y^+_{ijk}$ and $Y^-_{ijk}$, given our parameters. In some cases, the difference $Y^+_{ijk} - Y^-_{ijk}$ may be less than the numerical precision of the computer program. In that case, we calculate:

$$L (F_{ijk} \mid p_{ijk}, \sigma_o) = \text{NormalPDF} (\text{logit} (Y^+_{ijk}) \mid \text{logit} (p_{ijk}) \sigma_o)$$ (16)
Decay Model

We assumed that each mask had some true initial filtration rate $p_{ij0}$. We assumed that these were logit-normally distributed about some unknown mean mask initial filtration rate $p_{avg}$ with a standard deviation $\sigma_p$, that is:

$$\text{logit} (p_{ij0}) \sim \text{Normal} (\text{logit} (p_{avg}), \sigma_p) \quad (17)$$

We then assumed that the logit of the filtration rate, $\text{logit} (p_{ijk})$, after decontamination $k$ was equal to $\text{logit} (p_{ij(k-1)}) - d_{ijk}$, where $d_{ijk} \geq 0$ is the degrading effect of the $k^{th}$ decontamination of treatment $i$ on mask $j$. We assumed that the $d_{ijk}$ were log-normally distributed about a treatment and decontamination-specific mean value $\mu_{dik}$ and decontamination round-specific standard deviation $\sigma_{ dik}$:

$$\text{log} (d_{ijk}) \sim \text{Normal} (\text{log} (\mu_{dik}), \sigma_{ dik}) \quad (18)$$

We defined the $\mu_{dik}$ as

$$\mu_{dik} = \mu_{d0k} + \delta_{ik} \quad (19)$$

where $\mu_{0k} \geq 0$ is natural degradation during the $k^{th}$ trial in the absence of decontamination (i.e. the degradation rate in the control treatment, $i = 0$), and $\delta_{ik} \geq 0$ is the additional mean effect of the $k^{th}$ decontamination treatment of type $i$, and $\delta_{0k}$ is set equal to 0.

Model Prior Distributions

We placed a weakly informative Half-Normal prior on the control mean degradation values $\mu_{0k}$ with a mode at zero (since it is possible that control masks decay minimally):

$$\mu_{0k} \sim \text{Half-Normal} (0, 0.33) \quad (20)$$

We placed a weakly informative Half-Normal prior on the non-control mean degradation values $\mu_{ik}$, $i > 0$:

$$\mu_{ik} \sim \text{Half-Normal} (0.25, 0.5) \quad (21)$$

reflecting the conservative assumption that decontamination should degrade the mask at least somewhat.

We placed a Truncated Normal prior on the mean initial filtration $p_{avg}$:

$$p_{avg} \sim \text{TruncNormal} (0.99, 0.0025) \quad (22)$$
The mode of 0.995 corresponds to the maximum measurable fit factor of 200. The standard deviation of 0.0025 leaves it plausible that some masks could start near or below the minimum acceptable threshold fit factor of 100, which corresponds to a $p$ of 0.99.

We placed weakly informative Half-Normal priors on the logit-space standard deviations $\sigma_p$ and $\sigma_o$. $\sigma_p$ reflects variation in individual masks’ initial filtration about $p_{avg}$ and $\sigma_o$ reflects noise in the observation process.

$$\sigma_p, \sigma_o \sim \text{Half-Normal} (0, 0.5) \quad (23)$$

$\sigma_{dik}$, and the various $\sigma_{dik}$ reflect variation in masks’ true degree of degradation between decontaminations about the expected degree of decay,

$$\sigma_{dik} \sim \text{Half-Normal} (0, \log (1.5) / 3) \quad (24)$$

We chose standard of 0.5 for $\sigma_p$ and $\sigma_o$ because a standard deviation of 1.5 (i.e. 3 $\sigma$ in the hyperprior) in logit space corresponds to probability values being uniformly distributed between 0 and 1; we therefore wish to tell our model not to use larger values of $\sigma_p$, $\sigma_o$ as these would squash all $p_{ijk}$ to one of two modes, one at 0 and one at 1 (22). For the same reason, we chose $\log (1.5) / 3$ for the prior standard deviation $\sigma_{dik}$, since it is the standard deviation of a log of a logit-space value.

**Markov Chain Monte Carlo Methods**

For all Bayesian models, we drew posterior samples using Stan (Stan Core Team 2018; https://mc-stan.org), which implements a No-U-Turn Sampler (a form of Markov Chain Monte Carlo), via its R interface RStan. We ran 4 replicate chains from random initial conditions for 2,000 iterations, with the first 1,000 iterations as a warmup/adaptation period. We saved the final 1,000 iterations from each chain, giving us a total of 4,000 posterior samples. We assessed convergence by inspecting trace plots and examining $\hat{R}$ and effective sample size ($n_{eff}$) statistics. For the mask fit model, we assessed appropriateness of prior distributions with prior predictive checks (Appendix Figure 4) and assessed goodness-of-fit with posterior predictive checks (Appendix Figure 5). For the decay rate model, we assessed goodness of fit by plotting model estimated decay against estimated titers (Figure 1; Appendix Figure 1, 2).
Limit of Detection (LOD)

Endpoint titration has an approximate limit of detection set by the volume of the undilute sample deposited in each well. If all wells – including those containing undiluted sample – are negative and a Poisson single-hit model is used, the best guess is that the true titer lies somewhere below $1 \text{ TCID}_{50} / (\text{volume of deposited sample})$. How far below is determined by the number of wells. For 4 wells, as was standard in our experiments, the first quarter log$_{10}$ titer at which 0 wells is the most likely outcome is $10^{-0.5}$ TCID$_{50}$ per volume of sample. This is also the imputed Speaman-Karber titer in that case. Since we used samples of volume 0.1 mL, this corresponds to a value of $10^{0.5}$ TCID$_{50}$/mL. So although we do not use the Spearman-Karber method here (since we infer mean titers directly from the well data) we use that LOD value to plot samples for which no replicate had a positive well (since the posterior distribution in that case covers a wide-range of sub-threshold values).

Code and Data Availability

Code and data to reproduce the Bayesian estimation results and produce corresponding figures are archived online at OSF: https://doi.org/10.17605/OSF.IO/mkg9b and available on Github: https://github.com/dylanhmorris/n95-decontamination

References


Appendix Table. Effect of decontamination method on SARS-CoV-2 viability on N95 respirators

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Material</th>
<th>Half-life, min</th>
<th></th>
<th>Time to one thousandth, min</th>
<th></th>
<th>Time to one millionth, min</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Median</td>
<td>2.5%</td>
<td>97.5%</td>
<td>Median</td>
<td>2.5%</td>
<td>97.5%</td>
</tr>
<tr>
<td>Control</td>
<td>N95 mask</td>
<td>78.5</td>
<td>66</td>
<td>90.2</td>
<td>782</td>
<td>658</td>
<td>899</td>
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<tr>
<td></td>
<td>Steel</td>
<td>234</td>
<td>202</td>
<td>260</td>
<td>2.33 × 10^3</td>
<td>2.01 × 10^3</td>
<td>2.59 × 10^3</td>
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<tr>
<td>Ethanol</td>
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<td>0.35</td>
<td>0.257</td>
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<td>3.49</td>
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<tr>
<td></td>
<td>Steel</td>
<td>0.89</td>
<td>0.729</td>
<td>1.04</td>
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<td>7.45</td>
<td>10.2</td>
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<td></td>
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<td>0.287</td>
<td>0.533</td>
<td>4.28</td>
<td>2.86</td>
<td>5.31</td>
</tr>
</tbody>
</table>

*Results are reported as the median and upper- and lower-limits of the 95% credible interval of the estimated half-life, and time needed to reduce viable SARS-CoV-2 load by a factor of 1,000 or 1,000,000, based on the posterior distribution of the exponential decay rate of the virus on different materials following different decontamination treatments.

Appendix Figure 1. Exponential decay fits for control experiments, carried out to final timepoints (a limited subset are shown in Figure 1 of the main text to make measurements for treatments readable). Lines show predicted decay of virus titer over time (lines; 50 random draws per replicate from the joint posterior distribution of the exponential decay rate, i.e. negative of the slope, and intercept, i.e. initial virus titer). Points show estimated mean titer across three replicates (circles, thick bars, and thin bars show the posterior median estimate of this mean, a 68% credible interval, and a 95% credible interval, respectively). Timepoints with no positive wells for any replicate are plotted as triangles at the approximate single-replicate detection limit of the assay (LOD, see Materials and Methods for discussion) to indicate that a range of sub-LOD values are plausible. Black dotted line shows approximate LOD: 10^{0.5} TCID_{50}/mL media.
Appendix Figure 2. Estimated mask integrity over time for individual masks. Connected points represent observed values; violin plots show posterior distribution of estimated underlying integrity, measured as a fit factor. Black dots show posterior median estimates; thick bars, 68% credible intervals; and thin bars 95% credible intervals. Triangles denote observed values of 200, the upper limit of measurement (so true values may be higher). Circles denote other observed values. Red dashed line indicates fit factor of 100, the minimum passing value for a fit test. Columns are ordered by estimated mask integrity after the third decontamination.
Appendix Figure 3. Prior predictive checks for mask fit model. Large connected points are observed values for individual masks. Small overplotted connected points are simulated observations drawn from the prior predictive distribution (including simulated measurement error and censoring). The wide range of simulated values, which covers the real data, shows that the priors are not overly informative. Triangles denote observed values of 200, the upper limit of measurement (so true values may be higher). Circles denote other observed values. Red dashed line indicates fit factor of 100, the minimum passing value for a fit test. Each panel shows 25 randomly chosen simulated observations. Masks are ordered by fit factor observed after 3 decontaminations.
Appendix Figure 4. Posterior predictive checks for mask fit model. Large connected points are observed values for individual masks. Small overplotted connected points are simulated sets of observations for each mask, drawing from the posterior predictive distribution for that mask (including simulated measurement error and censoring). The relatively tight coverage of the actual observations suggests that the model fits the data well. Triangles denote observed values of 200, the upper limit of measurement (so true values may be higher). Circles denote other observed values. Red dashed line indicates fit factor of 100, the minimum passing value for a fit test. Each panel shows 25 randomly chosen simulated observations per panel. Masks are ordered by fit factor observed after 3 decontaminations.