1	Inactivation analysis of SARS-CoV-2 by specimen transport media, nucleic acid extraction
2	reagents, detergents and fixatives
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23	transport tubes; molecular extraction reagents; lysis buffers; clinical diagnostics;

24 Abstract

25 The COVID-19 pandemic has necessitated a rapid multi-faceted response by the scientific community, bringing researchers, health officials and industry together to address the 26 27 ongoing public health emergency. To meet this challenge, participants need an informed approach for working safely with the etiological agent, the novel human coronavirus SARS-28 CoV-2. Work with infectious SARS-CoV-2 is currently restricted to high-containment 29 laboratories, but material can be handled at a lower containment level after inactivation. Given 30 the wide array of inactivation reagents that are being used in laboratories during this pandemic, it 31 32 is vital that their effectiveness is thoroughly investigated. Here, we evaluated a total of 23 33 commercial reagents designed for clinical sample transportation, nucleic acid extraction and virus inactivation for their ability to inactivate SARS-CoV-2, as well as seven other common 34 35 chemicals including detergents and fixatives. As part of this study, we have also tested five filtration matrices for their effectiveness at removing the cytotoxic elements of each reagent, 36 permitting accurate determination of levels of infectious virus remaining following treatment. In 37 addition to providing critical data informing inactivation methods and risk assessments for 38 diagnostic and research laboratories working with SARS-CoV-2, these data provide a framework 39 40 for other laboratories to validate their inactivation processes and to guide similar studies for other pathogens. 41

43 1. Introduction

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Infection with the novel human betacoronavirus SARS-CoV-2 can cause a severe or fatal 44 respiratory disease, termed COVID-19 (1-3). As the COVID-19 pandemic has developed. 45 millions of clinical samples have been collected for diagnostic evaluation. SARS-CoV-2 has 46 been classified as a Hazard Group 3 pathogen in the UK, and as such, deliberate work with the 47 virus must be carried out in high containment laboratories (containment level 3 (CL3) in the UK) 48 with associated facility, equipment and staffing restrictions. Guidance from Public Health 49 England (PHE) and the World Health Organization (WHO) enable non-propagative testing of 50 51 clinical specimens to be carried out at the lower CL2, with the requirement that all non-52 inactivated material is handled within a microbiological safety cabinet (MSC) and that the process has been suitably and sufficiently risk assessed (4, 5). Guidance from the U.S. Centers 53 for Disease Control and Prevention requires that specimens must be inactivated (e.g. in nucleic 54 acid extraction buffer) before handling at biosafety level 2 (BSL-2) (6). To allow safe movement 55 of clinical samples from CL3/BSL-3 laboratories to CL2/BSL-2, virus inactivation procedures 56 should be validated, and formal validation of inactivation protocols are often an operational 57 requirement for clinical and research laboratories handling SARS-CoV-2. 58 59 Efficacy of virus inactivation depends on numerous factors, including the nature and concentration of pathogen, sample matrix, concentration of inactivation agent/s and contact time. 60 To date, there are limited data on efficacy of SARS-CoV-2-specific inactivation approaches in 61 62 the scientific literature and risk assessments have largely been based upon inactivation information for genetically related coronaviruses. Previous studies have found that treatment 63

65 are effective at inactivating SARS-CoV-1 and Middle East Respiratory Syndrome coronavirus

with heat, chemical inactivants, ultraviolet light, gamma irradiation and a variety of detergents

(MERS-CoV), other high-consequence human coronaviruses (7–13). However, limited 66 validation data exist for coronavirus inactivation by commercial sample transport media and 67 molecular extraction lysis buffers used in steps prior to nucleic acid extraction for diagnostic 68 69 testing. Furthermore, the precise composition of many commercial reagents is proprietary, preventing ingredient-based inference of inactivation efficacy between reagents. Some limited 70 71 preliminary data on SARS-CoV-2 inactivation are available (14–19), but given the current level of diagnostic and research activities, there is an urgent need to comprehensively investigate 72 SARS-CoV-2-specific inactivation efficacy of available methods to support safe virus handling. 73 74 An important consideration in inactivation assays is cytotoxicity, a typical effect of many chemical inactivants. To mitigate cytotoxic effects, the inactivation agent needs to be either 75 diluted out or removed from treated samples prior to testing for infectious virus. Each of these 76 77 methods for addressing cytotoxicity present their own challenges. Sample dilution requires the use of high titer stocks of virus (e.g. $>10^8$ PFU/mL) to be able to demonstrate a significant titer 78 reduction and reduces recovery of low level residual virus from treated samples, making it 79 80 difficult or impossible to distinguish complete from incomplete virus inactivation. In contrast, methods for purification of virus away from cytotoxic components in treated samples may also 81 82 remove virus or affect virus viability. Accurate quantification of remaining infectious virus ideally requires complete removal of cytotoxicity without compromising assay sensitivity, which 83 needs careful consideration of reagent and purification processes prior to performing inactivation 84 85 tests.

Here, we describe optimal methods for the removal of cytotoxicity from samples treated with commercial reagents, detergents and fixatives. These data were then used in evaluations of the effectiveness of these chemicals for inactivating SARS-CoV-2. This work, applicable to both

- 89 diagnostic and research laboratories, provides invaluable information for public health and basic
- 90 research responses to the COVID-19 pandemic by supporting safe approaches for collection,
- 91 transport, extraction and analysis of SARS-CoV-2 samples. Furthermore, our studies
- 92 investigating purification of a wide range of cytotoxic chemicals are highly applicable to
- 93 inactivation studies for other viruses, thereby supporting rapid generation of inactivation data for
- 94 known and novel viral pathogens.

96 2. Materials and Methods

97 2.1. Cells and virus

Vero E6 cells (Vero C1008; ATCC CRL-1586) were cultured in modified Eagle's 98 minimum essential medium (MEM) supplemented with 10% (v/v) fetal calf serum (FCS). Virus 99 used was SARS-CoV-2 strain hCOV-19/England/2/2020, isolated by PHE from the first patient 100 cluster in the UK on 29/01/2020. This virus was obtained at passage 1 and used for inactivation 101 studies at passage 2 or 3. All infectious work carried out using an MSCIII in a CL3 laboratory. 102 Working virus stocks were generated by infecting Vero E6 cells at a multiplicity of infection 103 104 (MOI) of 0.001, in the presence of 5% FCS. Cell culture supernatants were collected 72 hours post infection, clarified for 10 mins at $3000 \times g$, aliquoted and stored at -80°C until required. 105 Viral titers were calculated by either plaque assay or 50% tissue culture infectious dose 106 107 (TCID50). For plaque assays, 24-well plates were seeded the day before the assay (1.5×10^5) cells/well in MEM/10%FCS). Ten-fold dilutions of virus stock were inoculated onto plates 108 (100µL per well), inoculated at room temperature for 1 hour then overlaid with 1.5% medium 109 110 viscosity carboxymethylcellulose (Sigma-Aldrich) and incubated at 37°C/5% CO₂ for 3 days. For TCID50s, ten-fold dilutions of virus stock $(25\mu L)$ were plated onto 96-well plates containing 111 Vero E6 cell suspension (2.5×10^4 cells/well in 100µl MEM/5%FCS) and incubated at 37°C/5% 112 CO_2 for 5-7 days. Plates were fixed with 4% (v/v) formaldehyde/PBS, and stained with 0.2% 113 (v/v) crystal violet/water TCID50 titers were determined by the Spearman-Kärber method (20, 114 115 21).

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117 2.2. Reagents and chemicals used for SARS-CoV-2 inactivation

118	The commercial reagents evaluated in this study, along with their compositions (if
119	known) and manufacturers' instructions for use (if provided) are given in Supplementary Table
120	1. Specimen transport reagents tested were: Sigma Molecular Transport Medium (MM, Medical
121	Wire); eNAT (Copan); Primestore Molecular Transport Medium (MTM, Longhorn Vaccines and
122	Diagnostics); Cobas PCR Media (Roche); Aptima Specimen Transport Medium (Hologic);
123	DNA/RNA Shield, (Zymo Research); guanidine hydrochloride (GCHl) and guanidine
124	thiocyanate (GITC) buffers containing Triton X-100 (both Oxoid/Thermo Fisher); Virus
125	Transport and Preservation Medium Inactivated (BioComma). Molecular extraction reagents
126	tested were: AVL, RLT, ATL, and AL (all Qiagen); MagNA Pure external lysis buffer, and
127	Cobas Omni LYS used for on-board lysis by Cobas extraction platforms (Roche); Viral PCR
128	Sample Solution (VPSS) and Lysis Buffer (both E&O Laboratories); NeuMoDx Lysis Buffer
129	(NeuMoDx Molecular); Samba II SCoV lysis buffer (Diagnostics for the Real World);
130	NucliSENS lysis buffer (Biomerieux); Panther Fusion Specimen Lysis Tubes (Hologic); and an
131	in-house extraction buffer containing guanidine thiocyanate and Triton X-100 (PHE Media
132	Services). Detergents tested were: Tween 20, Triton X-100 and NP-40 Surfact-Amps Detergent
133	Solutions (all Thermo Scientific), and UltraPure SDS 10% solution (Invitrogen). Other reagents
134	assessed include: polyhexamethylene biguanide (PHMB, Blueberry Therapeutics);
135	Formaldehyde and Glutaraldehyde (TAAB); and Ethanol and Methanol (Fisher Scientific).
136	

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2.3. Removal of reagent cytotoxicity

Specimen transport tube reagents were assessed undiluted unless otherwise indicated. For
testing of molecular extraction reagents, mock samples were generated by diluting reagent in
PBS at ratios given in manufacturer's instructions. Detergents, fixatives and solvents were all

assessed at the indicated concentrations. All methods were evaluated in a spin column format, for 141 ease of sample processing within the high containment laboratory. Pierce Detergent Removal 142 Spin Columns (0.5mL, Thermo Scientific), Microspin Sephacryl S400HR (GE Healthcare), and 143 Amicon Ultra-0.5mL 50KDa centrifugal filters (Merck Millipore) were prepared according to 144 manufacturer's instructions. Sephadex LH-20 (GE Healthcare) and Bio-Beads SM2 resin (Bio-145 146 Rad) were suspended in PBS and poured into empty 0.8mL Pierce centrifuge columns (Thermo Scientific), and centrifuged for one min at $1000 \times g$ to remove PBS immediately before use. For 147 all matrices aside from the Amicon Ultra columns, 100µl of treated sample was added to each 148 149 spin column, incubated for two mins at room temperature, then eluted by centrifugation at 1,000 \times g for two mins. For Amicon Ultra filters, 500µl of sample was added, centrifuged at 14,000 \times g 150 for 10 mins, followed by three washes with 500µl PBS. Sample was then collected by 151 152 resuspending contents of the filtration device with 500µl PBS. To assess remaining cytotoxicity, a two-fold dilution series of treated filtered sample was prepared in PBS, and 6.5µl of each 153 dilution transferred in triplicate to 384-well plates containing Vero E6 cells (6.25×10^3 cells/well 154 in 25µl MEM/5%FCS) and incubated overnight. Cell viability was determined by CellTiter 155 Aqueous One Solution Cell Proliferation Assay (Promega) according to manufacturer's 156 157 instructions. Normalized values of absorbance (relative to untreated cells) were used to fit a 4parameter equation to semilog plots of the concentration-response data, and to interpolate the 158 concentration that resulted in 80% cell viability (CC20) in reagent treated cells. All analyses 159 160 were performed using GraphPad Prism 8 (v8.4.1, GraphPad Software). 161

162 2.4. SARS-CoV-2 inactivation

163 For commercial products, virus preparations (tissue culture fluid, titers ranging from $1 \times$ 10^6 to 1×10^8 PFU/ml) were treated in triplicate with reagents at concentrations and for contact 164 times recommended in the manufacturers' instructions for use, where available, or for 165 concentrations and times specifically requested by testing laboratories. Where a range of 166 concentrations was given by the manufacturer, the lowest ratio of product to sample was tested 167 168 (i.e. lowest recommended concentration of test product). Specimen transport tube reagents were tested using a ratio of one volume of tissue culture fluid to ten volumes of reagent, unless a 169 volume ratio of sample fluid to reagent was specified by the manufacturer. Detergents, fixatives 170 171 and solvents were tested at the indicated concentrations for the indicated times. For testing of 172 alternative sample types, virus was spiked into the indicated sample matrix at a ratio of 1:9, then treated with test reagents as above. All experiments included triplicate control mock-treated 173 174 samples with an equivalent volume of PBS in place of test reagent. Immediately following the required contact time, 1mL of treated sample was processed using the appropriately selected 175 filtration matrix. Reagent removal for inactivation testing was carried out in a larger spin column 176 177 format using Pierce 4mL Detergent Removal Spin Columns (Thermo Fisher), or by filling empty Pierce 10mL capacity centrifuge columns (Thermo Fisher) with SM2 Bio-Beads, Sephacryl S-178 179 400HR or Sephadex LH-20 to give 4mL packed beads/resin. For purification using Amicon filters, $2 \times 500 \mu$ l samples were purified using two centrifugal filters by the method previously 180 described, then pooled together. For formaldehyde and formaldehyde with glutaraldehyde 181 182 removal, one filter was used with $1 \times 500 \mu$ l sample volume, resuspended after processing in 500µl PBS, and added to 400ul MEM/5% FBS. For inactivation of infected monolayers, 12.5 183 cm^2 flasks of Vero E6 cells (2.5 × 10⁶ cells/flask in 2.5mL MEM/5% FBS) were infected at MOI 184 185 0.001 and incubated at 37°C/5% CO₂ for 24 hours. Supernatant was removed, and cells fixed

186	using 5mL of formaldehyde, or formaldehyde and glutaraldehyde at room temperature for 15 or										
187	60 mins. The fixative was removed, and monolayers washed three times with PBS before										
188	scraping cells into 1mL MEM/5% FBS and sonicated (3 \times 10 second on,10 seconds off at 100%										
189	power and amplitude) using a UP200St with VialTweeter attachment (Hielscher Ultrasound										
190	Technology). Supernatants were clarified by centrifuging at $3000 \times g$ for 10 mins.										
191											
192	2.5. SARS-CoV-2 quantification and titer reduction evaluation										
193	Virus present in treated and purified, or mock-treated and purified, samples was										
194	quantified by either TCID50 or plaque assay. As additional assay controls, unfiltered mock-										
195	treated sample was titrated to determine virus loss during filtration, and filtered test-reagent only										
196	(no virus) sample titrated to determine residual test buffer cytotoxicity. For TCID50 assays, neat										
197	to 10 ⁻⁷ ten-fold dilutions were prepared, and for plaque assays, neat to 10 ⁻⁵ ten-fold dilutions										
198	were prepared, both in MEM/5% FCS. TCID50 titers were determined by the Spearman-Kärber										
199	method (20, 21). Conditions where low levels of virus were detected such that TCID50 could not										
200	be calculated by Spearman-Kärber, TCID50 was calculated the Taylor method (22). Where no										
201	virus was detectable, values are given as less than or equal to the Taylor-derived TCID50 titer										
202	given by a single virus positive well at the lowest dilution where no cytotoxicity was observed.										
203	Titer reduction was calculated by subtracting the mean logarithmic virus titer for test-buffer-										
204	treated, purified conditions from the mean logarithmic virus titer for the PBS-treated, purified										
205	condition, with standard errors calculated according to (22).										
206											

207 2.6. Serial passages of treated samples

in 2.5mL MEM/5% FBS) were inoculated with either 500 μ l or 50 μ l of treated filtered sample. Flasks were examined for cytopathic effect (CPE) and 500 μ l culture medium from each flask was used to inoculate new 12.5 cm ² flasks of Vero E6 cells after seven days. If no CPE was
Flasks were examined for cytopathic effect (CPE) and 500 μ l culture medium from each flask was used to inoculate new 12.5 cm ² flasks of Vero E6 cells after seven days. If no CPE was
was used to inoculate new 12.5 cm ² flasks of Vero E6 cells after seven days. If no CPE was
observed, this process was continued for up to four serial passages. For the duration of the
passage series, a flask of untreated cells was included as a control for cross-contamination
between flasks, and a SARS-CoV-2 infected control was included to ensure suitable conditions
for virus propagation. To distinguish CPE from any residual cytotoxicity associated with test
reagents, samples of cell culture medium were taken from each flask at the beginning and end of
each passage. Nucleic acid was extracted from cell culture media manually using a QIAamp
Viral RNA Mini Kit (QIAGEN) or using NucliSENS easyMAG or EMAG platforms (both
BioMérieux). Viral RNA levels were quantified by quantitative reverse-transcriptase PCR (qRT-
PCR) specific for the SARS-CoV-2 E gene (23) using TaqMan Fast 1-Step Master Mix (Applied
Biosystems) on a 7500 Fast Real-Time PCR System (Applied Biosystems). A positive result for
virus amplification was recorded if effects on the monolayer consistent with CPE and a decrease
in Ct across the course of a passage were observed.

225 **3. Results**

3.1. Reagent filtration optimization to minimize cytotoxicity and maximum virus recovery

Prior to evaluating their effectiveness at inactivating SARS-CoV-2, we investigated the 228 229 cytotoxicity of each reagent before and after filtration though one of five matrices: Sephadex 230 LH-20, Sephacryl S400HR, Amicon Ultra 50kDa molecular weight cut-off centrifugal filters, Pierce detergent removal spin columns (DRSC), and Bio-Beads SM2 nonpolar polystyrene 231 adsorbents. Reagents were diluted with PBS to the working concentrations recommended by the 232 233 manufacturer (for commercial sample transport and molecular extraction reagents), or the indicated concentrations (for all other chemicals), followed by a single reagent removal step with 234 each filtration matrix. Dilution series of filtered and unfiltered samples were generated to 235 236 determine concentration-dependent cytotoxicity, from which the CC20 value for each combination of reagent and filtration method were interpolated (Supplementary Figure 1). CC20 237 was chosen as, at this concentration, cells retain 80% viability and enable distinction of active 238 239 SARS-CoV-2 replication by visualisation of CPE in the monolayer. Table 1 shows the dilution factor of reagent-treated sample required to achieve the CC20 after filtration, with <1 indicating 240 241 complete removal of cytotoxicity. These data were used to determine the relative cytotoxicity removed by one filtration step for each combination of reagent and matrix (Figure 1A). 242

All unfiltered reagents tested here were cytotoxic, but the degree of cytotoxicity varied considerably as did the optimal filtration matrix for each reagent. The detergent Tween 20 used at 1% concentration was the least cytotoxic unfiltered, only requiring a dilution factor of 7.7 to reach the CC20, although only the Bio-Bead SM2 filters were effective at removing all cytotoxicity. The chemical fixative combination of 2% formaldehyde plus 1.5% glutaraldehyde

248	was the most cytotoxic unfiltered, requiring a dilution of over 4000 to reach the CC20, with only
249	the Amicon Ultra columns able to remove 100% of the cytotoxicity. However, for the majority
250	reagents (27/34) tested, filtration through at least one matrix type removed 100% of cytotoxicity
251	allowing neat eluate to be used directly in cell culture without further dilution. There were
252	several exceptions to this: DNA/RNA shield (maximum 99.4% cytotoxicity removal using
253	SM2); 40% GHCl (99.1% using Pierce DRSC); 4M GITC (99.7% using Pierce DRSC); MagNA
254	Pure (99.7% using SM2); AL buffer (87.4% using S400HR); Cobas Omni LYS (97.0% using
255	SM2); and NeuMoDx (93.4% using S400HR). For these reagents, filtered eluate was still
256	cytotoxic when used undiluted in cell culture. However, CC20 values indicated that this
257	remaining cytotoxicity would be removed by first or second $(10^{-1} - 10^{-2})$ dilutions in the TCID50
258	assay allowing evaluation of titer reduction using these reagents with the caveat that the effective
259	assay limit of detection (LOD) would be higher. Passing treated samples through more than one
260	column, or increasing the depth of the resin/bead bed within the spin column can also improve
261	cytotoxicity removal for some reagents (unpublished data).
262	In addition to cytotoxicity removal, a successful filtration method must also purify virus
263	without adversely affecting titer or integrity. We therefore assessed SARS-CoV-2 recovery after
264	each filtration method. Using an input titer of 1.35×10^6 TCID50/mL, triplicate purifications of
265	virus through Sephadex LH-20 or Pierce detergent removal spin columns resulted in recovery of

266 100% of input virus (Figure 1B). In contrast, the recoverable titer after one filtration through

Amicon Ultra filters was 2.13×10^5 TCID50/mL, an 85% reduction from input. Purification with

268 S400HR and Bio-Beads SM2 matrices resulted in recoverable titers of 1.08×10^6 TCID50/mL

and 8.99×10^5 TCID50/mL, a loss of 30% and 35% of input virus, respectively.

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271 **3.2.** SARS-CoV-2 inactivation by specimen transport and molecular extraction reagents

Specimen transport tubes are designed to inactivate microorganisms present in clinical 272 specimens prior to sample transport, while preserving the integrity of nucleic acids for molecular 273 274 testing. If effective, these products have the potential to streamline SARS-CoV-2 diagnostic processing in testing laboratories by eliminating the requirement for CL3 processing or, for 275 activities derogated to CL2, permitting processing outside an MSC. The BS EN 14476 standard 276 requires demonstration of a >4 \log_{10} titer reduction for virucidal suspension tests (22), and we 277 were able to demonstrate a \geq 4 log10 TCID50 titer reduction for all specimen transport media 278 evaluated in a tissue culture fluid matrix (Table 2). However, infectious virus remained 279 280 recoverable in treated samples after inactivation with most reagents tested (by either TCID50 or blind passage). The exceptions to this were PrimeStore MTM and 4M GITC, from which no 281 282 residual virus was detectable by either TCID50 or by the passaging of treated purified sample. While several contact times were evaluated for all these reagents, length of contact time had no 283 effect on either the level of virus titer reduction or whether virus remained detectable upon 284 285 passage.

We also sought to inform sample processing by examining inactivation by molecular 286 287 extraction lysis buffers used in several manual and automated extraction protocols within SARS-CoV-2 diagnostic and research laboratories. We could demonstrate a \geq 4 log10 reduction in 288 TCID50 titer for all but two molecular extraction reagents when evaluated using tissue culture 289 290 fluid (Table 3). The exceptions to this were AL and Cobas Omni LYS, where remaining cytotoxicity in the filtered eluate increased the TCID50 LOD to a level such that the maximum 291 calculable titer reductions were \geq 3.5 and \geq 3.9 log10 TCID50s, respectively. However, given no 292 293 virus was detected at any passage it is likely that infectious virus was effectively inactivated by

294 these two reagents. For reagents tested with multiple contact times (NucliSENS, Panther Fusion), shorter times (10 mins) were as effective at reducing virus titers as longer contact times. Most 295 reagents reduced viral titers to around the TCID50 assay LOD, indicating that any remaining 296 297 virus post treatment was present only at very low titers (<10 TCID50/mL), but higher levels of 298 virus were recoverable from samples treated with some extraction buffers. For NeuMoDx lysis buffer, although titers were reduced by $\geq 4 \log 10 \text{ TCID50s}$, an average of 91 (±38) TCID50/mL 299 remained detectable. Similarly, Buffer AVL reduced virus titers by 5.1 log10 TCID50s, but after 300 treatment virus was detectable in all treated samples replicates (average 54 (±18) TCID50/mL). 301 302 However, addition of four sample volumes of absolute ethanol following a 10 minute contact time with AVL (the next step in the QIAGEN Viral RNA Mini Kit manual), $a \ge 5.9 \log 10$ titer 303 reduction was recorded with no virus recoverable following passages in cell culture. 304 305 Panther Fusion lysis buffer was further tested against a relevant clinical sample matrix, pooled fluid from oropharyngeal (OP) and nasopharyngeal (NP) swab specimens, resulting in a 306 \geq 5.1 log10 titer with no remaining infectious virus detectable. We additionally evaluated the 307 tissue lysis buffer RLT using homogenised ferret lung as sample material, with treatment 308

resulting in a \geq 4.8 log10 titer reduction with no residual infectious virus detectable.

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311 3.3. SARS-CoV-2 inactivation by detergents

Detergents can be used to inactivate lipid enveloped viruses such as coronaviruses by disrupting the viral envelope, therefore rendering them unable to attach or enter cells (24–27).

- Here, we evaluated Triton X-100, SDS, NP40 and Tween 20 for their ability to inactivate SARS-
- CoV-2. SDS treatment at 0.1% or 0.5% reduced titers by \geq 5.7 and \geq 6.5 log10 TCID50s,
- respectively, while both concentrations of NP40 reduced titers by $\geq 6.5 \log 10 \text{ TCID50}$ with no

residual virus detectable following NP40 treatment. In contrast, up to 0.5% Tween 20 had no 317 effect on viral titers. Triton X-100 is commonly used in viral inactivation reagents, and here we 318 show that at both 0.1% and 0.5% v/v concentration, virus titers in tissue culture fluid were 319 320 reduced by \geq 4.9 log10 TCID50s, even with less than 2 min contact time (Table 4). Furthermore, we were unable to recover infectious virus from samples treated with 0.5% Triton X-100 for 10 321 mins or longer. We also saw effective inactivation of SARS-CoV-2 by SDS, NP40 and Triton X-322 100 in spiked NP and OP swab specimen fluid, but, importantly, we were not able to replicate 323 this in spiked serum; 1% Triton X-100 only reduced titers in human serum by a maximum of 2 324 325 log10 TCID50s with contact times of up to two hours. In addition to evaluating inactivation efficacy by detergents, we assessed the effects of 326 treatment on RNA integrity to determine their suitability for inactivation prior to nucleic acid 327 328 testing. Extracted RNA from treated samples was tested using a SARS-CoV-2-specific qRT-PCR, and the Ct difference between detergent-treated samples and mock-treated controls 329 determined (Table 4). A time-dependent increase in Ct value following treatment with 0.5% 330 Triton X-100 was observed, indicating a detrimental effect on RNA stability with increasing 331 treatment times. Treatment with NP40 had a marked effect, with a 30 minute treatment leading 332 to an increase in 9-10 Cts. While we saw no increase in Ct in tissue culture fluid samples treated 333 with 0.5% SDS, we observed an increase in Ct for SDS-treated swab fluid samples, likely due to 334 an increased concentration of RNases in clinical samples. 335 336

- 337

3.4. **SARS-CoV-2** inactivation by other chemical treatments

338 Fixation and inactivation of viruses by addition of formaldehyde, or a combination of 339 formaldehyde and glutaraldehyde, is a well-established protocol, particularly for diagnostic

340 electron microscopy (28, 29). 4% or 2% formaldehyde treatment for 15 or 60 mins reduced virus titers by \geq 4.8 log10 TCID50s when evaluated against a tissue culture fluid matrix, with no 341 remaining infectious virus detectable (Table 5). When infected monolayers were subjected to the 342 343 same treatment protocol, titer reductions were all $\geq 6.8 \log 10 \text{ TCID50s}$, with 60 min contact time moderately more effective than 15 min. However, in this format, a 60 min 4% formaldehyde 344 treatment was the only one from which no infectious virus was detectable. A mixture of 2% 345 formaldehyde with 1.5% glutaraldehyde tested on infected monolayers reduced virus titers by 346 \geq 6.7 log10 TCID50s with no remaining infectious virus detectable for both a 15 and 60 min 347 contact time. Polyhexanide biguanide (PHMB) is a polymer used as a disinfectant and antiseptic, 348 349 evaluated here as a potential lysis buffer, but it was only able to reduce viral titers by 1.6 log10 TCID50s at the highest concentration tested (2%). 350

352 4. Discussion

Samples containing infectious SARS-CoV-2 require an initial inactivation step before 353 downstream processing; given the rapid emergence of SARS-CoV-2, these inactivation protocols 354 355 have been guided by existing data for other coronaviruses and there is an urgent need to both confirm these historical data using the new virus and to validate new approaches for inactivating 356 357 SARS-CoV-2. We therefore analysed numerous commercially and commonly available reagents used by public health agencies and research laboratories around the world in their response to the 358 pandemic. In addition, to address challenges of reagent cytotoxicity in inactivation evaluation, 359 360 we provide data on the effectiveness of filtration methods for removing cytotoxicity from chemically treated samples. 361

Knowledge of the expected amount of infectious virus in clinical samples obtained from 362 363 COVID-19 patients is important when applying viral inactivation study data to diagnostic sample processing, allowing end users to interpret whether material they are handling is likely to 364 represent an infectious risk to themselves and others. These values are dependent on several 365 factors, including time post symptom onset, duration of symptoms, time elapsed between 366 sampling and testing, the presence of neutralizing antibody responses, and immunocompetency 367 368 of the individual (30). Data regarding quantitative infectious viral levels in typical clinical specimens are minimal, with most studies reporting viral loads determined by qRT-PCR only 369 (31-33). One study of 90 qRT-PCR positive NP or endotracheal (ETT) samples from COVID-19 370 371 patients estimated the median titer at 3.3 log10 TCID50/mL (30). Given we demonstrate >4 log₁₀ reduction in titer for all specimen transport reagents, this suggests that these reagents may 372 373 considerably decrease, even eliminate, the infectivity of a clinical sample. However, our 374 observation that residual virus could be recovered from most treated samples indicates that these

media cannot be assumed to completely inactivate SARS-CoV-2 in samples and that additional
precautionary measures should be taken in laboratories when it comes to sample handling and
transport.

Limited SARS-CoV-2 inactivation data on molecular extraction reagents used in nucleic 378 379 acid detection assays are currently available. One study reported that Buffer AVL either alone or 380 in combination with ethanol was not effective at completely inactivating SARS-CoV-2 (15). By contrast, we could not recover any infectious virus from samples treated with AVL plus ethanol, 381 consistent with previous studies indicating that AVL and ethanol in combination is effective at 382 383 inactivating MERS and other enveloped viruses (10, 34), and indicating that both AVL and ethanol steps of manual extraction procedures should be performed before removal of samples 384 385 from primary containment for additional assurance. Our detergent inactivation data, indicating 386 that SDS, Triton X-100 and NP40, but not Tween 20, can effectively inactivate SARS-CoV-2 in tissue culture fluid and in pooled NP and OP swab fluid, corroborate findings of a recent study 387 (17); however, as has been demonstrated for other viruses (31), we observed an inhibitory effect 388 of serum on virus inactivation by detergent, highlighting the importance of validating 389 inactivation methods with different sample types. 390

Based on our findings comparing filtration matrices, we found that the optimum method for reagent removal for inactivation studies is determined by evaluating three factors: (i) effectiveness of cytotoxicity removal; (ii) efficiency of virus recovery; and (iii) the ease of performing these methods within a containment space. Methods permitting complete removal of cytotoxic reagent components with no or little effect on virus recovery give assurance that low levels of residual virus, if present, could be detected in virus inactivation studies. During reagent testing, there were several instances where we noted residual cytotoxicity in the neat eluate

contrary to what was expected based on the initial reagent removal data and is likely due to the
extended incubation period required for inactivation testing (up to 7 days, compared with
overnight for cytotoxicity evaluation). In all cases however, we were still able to enhance the
levels of titer reduction detectable when compared with what would have been achieved by
sample dilution alone.

In conclusion, we have evaluated methods for straightforward, rapid determination of 403 purification options for any reagent prior to inactivation testing, enabling establishment of 404 effective methods for sample purification while minimising virus loss. This is applicable to 405 406 inactivation studies for all viruses (known and novel), not only SARS-CoV-2. We have applied 407 these methods to obtain SARS-CoV-2 inactivation data for a wide range of reagents in use (or proposed for use) in SARS-CoV-2 diagnostic and research laboratories. In addition to guiding 408 409 laboratory risk assessments, this information enables laboratories to assess alternative reagents that may be used for virus inactivation and nucleic acid extraction, particularly considering 410 concerns about extraction reagent availability due to increased global demand caused by the 411 412 COVID-19 pandemic. Furthermore, chemical treatments evaluated here are commonly used for inactivation of a wide range of different viruses and other pathogens, and the results presented 413 414 may be used to directly inform and improve the design of future inactivation studies.

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- 422
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Table 1: Purification of reagents: Values [95% CI] represent the dilution factor required after one purification process to achieve the CC20 concentration [95% CI].

Туре	Reagent	Reagent:media	Post-filtration dilution factor of eluate needed for CC20							
		ratio or %v/v	Unpurified	Sephadex LH-20	Sephacryl S400HR	Amicon Ultra 50kDa	Pierce DRSC	Bio-Beads SM2		
	BioComma	10:1	36.2 [<i>30.1 – 44.0</i>]	<2 [n/a]	<2 [n/a]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]	12.1 [<i>9.2 – 16.4</i>]		
	Sigma MM	1.5:1	417 [<i>306 – 619</i>]	59.2 [<i>51.8 – 67.1</i>]	48.7 [<i>44.6 – 53.3</i>]	4.0 [<i>3.6 – 4.3</i>]	<1 [<i>n/a</i>]	7.6 [<i>6.5 – 8.9</i>]		
	eNAT	3:1	70.1 [<i>55.0 – 88.5</i>]	<1 [<i>n/a</i>]	2.8 [2.5 – 3.1]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]	24.4 [20.2 - 30.2]		
Specimen	Primestore MTM	3:1	56.2 [<i>47.2 – 66.3</i>]	<1 [<i>n/a</i>]	4.8 [<i>nc</i>]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]	18.3 [15.4 – 22.1]		
Transport	Cobas PCR Media	1:1	55.5 [<i>46.5 – 67.5</i>]	2.7 [<i>2.3 – 3.1</i>]	5.2 [<i>4.6 – 5.9</i>]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]	26.5 [<i>23.5 – 30.2</i>]		
Reagent	DNA/RNA Shield	10:1	1098 [<i>994 – 1231</i>]	1155 [<i>1076 – 1253</i>]	82.3 [<i><82.3 – 94.7</i>]	29.6 [<i>26.2 – 32.3</i>]	66.1 [<i>58.1</i> – 75.8]	7.1 [5.5 – 8.6]		
neugene	40% GHCI/Tx TM	10:1	245 [<i>205 – 288</i>]	24.5 [<24.5 – 31.5]	25.9 [<i><25.9 – 36.7</i>]	13.3 [<i><13.3 – 15.6</i>]	2.2 [nc]	119 [<i>103 – 135</i>]		
	2M GITC/Tx TM	10:1	245 [<i>215 – 277</i>]	19.4 [<i><</i> 19.4 – 23.9]	19.1 [<i>15.4 – 26.3</i>]	37.8 [<i>nc</i>]	<1 [<i>n/a</i>]	127 [<i>113 – 141</i>]		
	4M GITC/Tx TM	10:1	1054 [<i>889 - 1262</i>]	545 [487 - 613]	141 [<i>102 – 201</i>]	211 [<i>172 – 247</i>]	3.5 [<i>3.1 - 3.9</i>]	20.3 [<i>15.2 - 27.9</i>]		
	Buffer AVL	4:1	61.6 [50.8 – 75.1]	<1 [n/a]	3.2 [<i>2.9 – 3</i> .5]	<1 [n/a]	<1 [n/a]	26.1 [<i>21.5 – 32.3</i>]		
	MagNA Pure LB	1:1	1934 [<i>1348 – 2780</i>]	1391 [<i><1391–1654</i>]	474 [434 – 517]	346 [<i><346 – 382</i>]	59.1 [45.6 – 70.4]	5.8 [1.4 – 7.8]		
	NucliSENS	1:1	60.5 [<i>54.9 – 66.2</i>]	<1 [<i>n/a</i>]	4.3 [<i>4.0</i> – <i>4.9</i>]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]	4.6 [<4.6 - 6.7]		
	Panther Fusion	1.42:1	196 [<i><</i> 1 <i>96 – 214</i>]	<1 [<i>n/a</i>]	18.0 [<18.0 - 19.4]	15.9 [< <i>15.9</i> – <i>16.5</i>]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]		
	Buffer AL	1:1	61.9 [56.7 – 65.4]	37.4 [34.7 – 41.1]	7.8 [<i>6.6 – 9.3</i>]	30.5 [<i>25.5 – 36.3</i>]	29.5 [25.9 – 33.9]	16.5 [<i>14.6 – 18.9</i>]		
Molecular	Cobas Omni LYS	1:1	225 [<225 – 255]	142 [<i>nc</i>]	45.8 <i>[<45.8 – 55.6</i>]	117 [<i>nc</i>]	16.7 [<i>nc</i>]	6.7 [2.9 – 8.7]		
Reagents	PHE in-house LB	4:1	231 [<2 <i>31 – 310</i>]	26.2 [<i>22.0 - 31.8</i>]	11.4 [<i>9.9 - 13.2</i>]	2.7 [< <i>2.7 - 4.9</i>]	<1 [<i>n/a</i>]	12.9 [<i>9.8 - 17.9</i>]		
neugents	NeuMoDx LB	1:1	30.2 [<i>24.1 - 37.9</i>]	8.0 [<i>7.3 - 8.8</i>]	2.0 [1.7 – 2.4]	7.5 [6.6 - 8.1]	4.2 [0.4 - 6.9]	6.8 [<6.8 - 8.4]		
	E&O Labs VPSS	10:1	174 [<i>145 – 206</i>]	24.9 [22.1 - 28.4]	14.2 [11.7 - 17.5]	7.7 [<7.7 - 14.5]	<1 [<i>n/a</i>]	11.7 [8.5 – 16.4]		
	E&O Lab LB	10:1	69.0 [<i>62.7 – 76.9</i>]	9.5 [<9.5 – 11.0]	8.0 [7.4 – 8.7]	2.2 [<i>nc</i>]	<1 [<i>n/a</i>]	4.1 [3.5 – 4.7]		
	Samba SCoV LB	10:1	177 [<177 – 213]	68.2 [63.0 – 75.4]	27.3[24.2 - 30.1]	5.2 [<5.2 – 6.0]	<1 [<i>n/a</i>]	1.5 [1.0 – 1.8]		
	Buffer RLT	9:1	48.0 [<i>40.3 – 58.0</i>]	2.9 [2.3 – 4.3]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]	18.5 [15 <i>.3 – 22.8</i>]		
	Triton-X100	1%	185 [<i><185 – 211</i>]	48.4 [<48.4 - 58.4]	~17.22 [<i>nc</i>]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]		
Dotorgonts	Tween 20	1%	7.7 [6.9 – 8.6]	4.2 [<3.8 – 4.9]	1.3 [1.0 – 1.7]	4.4 [4.0 - 5.1]	4.9 [3.4 – 7.5]	<1 [<i>n/a</i>]		
Detergents	SDS	1%	69.6 [n/a]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]		
	NP40	1%	320 [<i><320 – 402</i>]	171 [<i><171 – 196</i>]	140 [<i>123 – 161</i>]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]	<1 [<i>n/a</i>]		
	Formaldehyde	4%	4207 [<i>3270 – 584</i> 4]	288 [<i>226 – 383</i>]	111 [<i>93 – 136</i>]	<1 [<i>n/a</i>]	51.6 [<i><</i> 5 <i>1.6 – 65.9</i>]	1309 [<i>1058 – 1685</i>]		
	Formaldehyde +	2% +	4227 [<i>3183 – 6027</i>]	39.8 [32.7 – 51.4]	97.9 [<i>82.9 -118</i>]	<1 [<i>n/a</i>]	22.6 [<22.6 – 27.2]	1545 [<i>1164 – 2203</i>]		
	Glutaraldenyde	1.5%		41 (1)	4.5. ()	4.5. ()				
Other	Ethanol	100%	63.3 [27.6 - 103]	<1 [n/a]	<1 [n/a]	<1 [n/a]	<1 [n/a]	8.8 [6.5 - 12.5]		
	Iviethanol	100%	108 [/9.5 – 155]	<1 [n/a]	<1 [n/a]	<1 [n/a]	<1 [n/a]	2.2 [1.9 - 2.5]		
	0.1% PHMB	10:1	30.1 [26.6 - 34.2]	9.5 [8.9 - 10.2]	<1 [n/a]	<1 [n/a]	<1 [n/a]	9.8 [<9.8 - 11.8]		
	1.0% PHMB	10:1	328 [304 – 356]	132 [111 – 154]	<1 [<i>n/a</i>]	<1 [n/a]	9.3 [<9.3 - 11.1]	203 [<203 – 299]		
	2.0% PHMB	10:1	837 [<837- 1141]	240 [<i>198 – 282</i>]	4.1 [3.7 - 4.5]	<1 [<i>n/a</i>]	25.0 <i>[<20.9 - 29.0</i>]	479 [<i><</i> 479 – 647]		

LB - lysis buffer; TM - transport Medium; nc - not able to be calculated.

Reagent	Virus matrix	Reagent: virus ratio	Contact time (mins)	Titer reduction Log10 (±SE)	Virus detectable in titration [†] (#replicates)	Virus detectable in culture (#replicates)
			10	≥ 4.8 (± 0.1)	Yes (2/3) [¢]	Yes (1/3)
Sigma MM	Tissue culture fluid	1.5:1	30	≥ 4.8 (± 0.1)	Yes (1/3) [¢]	Yes (1/3)
			60	≥ 4.8 (± 0.1)	No (0/3) [¢]	No (0/3)
			10	4.8 (± 0.2)	Yes (3/3)	Yes (3/3)
		1:3	30	5.1 (± 0.2)	Yes (3/3)	Yes (3/3)
ANAT	Tissue culture fluid		60	5.2 (± 0.2)	Yes (3/3)	Yes (3/3)
enal	lissue culture fiuld		10	≥ 5.1 (± 0.1)	No (0/3)*	Yes (1/3)
		3:1	30	≥ 5.1 (± 0.1)	No (0/3)*	Yes (1/3)
			60	≥ 5.1 (± 0.1)	No (0/3)*	No (0/3)
			10	≥ 5.1 (± 0.2)	No (0/3)*	No (0/3)
Primestore MTM	Tissue culture fluid	1:3	30	≥ 5.1 (± 0.2)	No (0/3)*	No (0/3)
			60	≥ 5.1 (± 0.2)	No (0/3)*	No (0/3)
		1:1.4	10	4.6 (± 0.1)	Yes (3/3)	Yes (3/3)
Cobas PCR Media	Tissue culture fluid		30	4.8 (± 0.1)	Yes (3/3)	Yes (3/3)
			60	4.8 (± 0.1)	Yes (3/3)	Yes (3/3)
			10	≥ 4.4 (± 0.1)	Yes (1/3)	No (0/3)
Aptima Specimen Transport Medium	Tissue culture fluid	5.8:1	30	≥ 4.4 (± 0.1)	No (0/3)	No (0/3)
			60	≥ 4.4 (± 0.1)	Yes (2/3)	Yes (1/3)
Virus Transport and			10	5.0 (± 0.2)	Yes (3/3)	Yes (3/3)
Preservation Medium	Tissue culture fluid	10:1	30	4.9 (± 0.2)	Yes (3/3)	Yes (3/3)
(Inactivated)			60	4.8 (± 0.2)	Yes (3/3)	Yes (3/3)
			10	≥ 4.8 (± 0.2)	No (0/3)**	tbc
DNA/RNA Shield	Tissue culture fluid	10:1	30	≥ 4.8 (± 0.2)	No (0/3)**	tbc
			60	≥ 4.8 (± 0.2)	No (0/3)**	tbc
2M GITC	Tissue culture fluid	10:1	30	≥ 4.6 (± 0.1)	No (0/3)*	Yes (1/3)
4M GITC	Tissue culture fluid	10:1	30	≥ 5.1 (± 0.2)	No (0/3)*	No (0/3)
40% GHCI	Tissue culture fluid	10:1	30	≥ 4.6 (± 0.1)	Yes (1/3)*	Yes (3/3)

532 Table 2: Virus inactivation by specimen transport tube reagents

 \dagger - samples titrated by TCID50, with a limit of detection of 5 TCID50/mL (0.7 Log10 TCID50/mL) unless stated * - limit of detection was 50 TCID50/mL (1.7 Log10 TCID50/mL) due to cytotoxicity in neat wells of TCID50 assay ** - limit of detection was 504 TCID50/mL (2.7 Log10 TCID50/mL) due to cytotoxicity in neat and -1 wells of TCID50 assay ϕ - titration by plaque assay; limit of detection was 3.3 PFU/mL (0.5 Log10 PFU/mL)

537	Table 3: Virus inactivation by molecular extraction re	agents
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Reagent	Virus matrix	Reagent: virus ratio	Contact time (mins)	Titer reduction Log10 (±SE)	Virus detectable in titration ⁺ (#replicates)	Virus detectable in culture (#replicates)
AVL	Tissue culture fluid	4:1	10	5.1 (± 0.1)	Yes (3/3)	Yes (3/3)
AVL + Ethanol	Tissue culture fluid	4:1:4 (AVL:virus: ethanol)	10 ^y	≥ 5.9 (± 0.2)	No (0/3)	No (0/3)
RLT (+BME)	Ferret lung homogenate	9:1	10	≥ 4.9 (± 0.2)	No (0/3)*	No (0/3)
MagNA Pure External LB	Tissue culture fluid	1:1	10	≥ 4.4 (± 0.2)	No (0/3)*	No (0/3)
AL	Tissue culture fluid	1:1	10	≥ 3.5 (± 0.2)	No (0/3)**	No (0/3)
Cobas Omni LYS	Tissue culture fluid	1:1	10	≥ 3.9 (± 0.1)	No (0/3)**	No (0/3)
PHE in-house LB	Tissue culture fluid	4:1	10	≥ 5.6 (± 0.1)	Yes (1/3)*	Yes (2/3)
	The second second second	10:1	30	≥ 5.2 (± 0.2)	No (0/3)*	Yes (2/3)
VPSS (E&U)	Tissue culture fiuld	1:1	10	≥ 5.1 (± 0.1)	No (0/3)*	Yes (1/3)
Lysis Buffer (E&O)	Tissue culture fluid	1:1	10	≥ 5.1 (± 0.1)	No (0/3)*	No (0/3)
NeuMoDx Lysis Buffer	Tissue culture fluid	1:1	10	4.3 (± 0.2)	Yes (3/3)*	Yes (3/3)
Samba SCoV LB	Tissue culture fluid	1:1	10	4.8 (± 0.1)	Yes (3/3)	Yes (3/3)
		1.1	10	≥ 5.0 (± 0.1)	Yes (2/3) [¢]	Yes (1/3)
NucliSENS LB	Tissue culture fluid	1:1	30	≥ 5.1 (± 0.0)	No (0/3) [¢]	Yes (1/3)
		2:1	10	≥ 4.9 (± 0.1)	No (0/3)*	No (0/3)
			10	≥ 4.4 (± 0.0)	No (0/3) [¢]	No (0/3)
Panther Fusion	Tissue culture fluid	1.42:1	30	≥ 4.4 (± 0.0)	No (0/3) [¢]	Yes (1/3)
Specimen Lysis Tubes			60	≥ 4.4 (± 0.0)	No (0/3) [¢]	Yes (1/3)
	Pooled swab material	1.42:1	30	≥ 5.1 (± 0.1)	No (0/3)	No (0/3)

 LB – Lysis buffer

 † - samples titrated by TCID50, with a limit of detection of 5 TCID50/mL (0.7 Log10 TCID50/mL) unless stated

 * - limit of detection was 50 TCID50/mL (1.7 Log10 TCID50/mL) due to cytotoxicity in neat wells of TCID50 assay

 ** - limit of detection was 504 TCID50/mL (2.7 Log10 TCID50/mL) due to cytotoxicity in neat and -1 wells of TCID50 assay

 \$\varphi\$ - titration by plaque assay; limit of detection was 3.3 PFU/mL (0.5 Log10 PFU/mL)

Table 4: Virus inactivation by detergents 543

Detergent	Virus matrix	Detergent: virus ratio	Contact time (mins)	Titer reduction Log10 (±SE)	Virus detectable in TCID50 [†] (#replicates)	Virus detectable in culture (#replicates)	RNA integrity‡ (Ct)
Tween 20	Tissue sulture fluid	0.1% v/v	30	0.0 (± 0.2)	Yes (3/3)	Yes (3/3)	n.d.
Tween 20	rissue culture fiuld	0.5% v/v	30	0.0 (± 0.2)	Yes (3/3)	Yes (3/3)	+0.2 (±0.0)
		0.1% v/v	30	≥ 4.9 (± 0.1)	Yes (3/3)	Yes (3/3)	n.d.
	Tierre eulture fluid		<2	5.9 (± 0.2)	Yes (3/3)	Yes (3/3)	+0.1 (±0.2)
	Tissue culture fiuld	0.5% v/v	10	≥ 6.2 (± 0.2)	No (0/3)	No (0/3)	+1.4 (±0.1)
Triton V 100			30	≥ 6.1 (± 0.2)	No (0/3)	No (0/3)	+3.6 (±0.1)
11101 X-100	Human sera	1.0% v/v	30	1.3 (± 0.2)	Yes (3/3)	Yes (3/3)	n.d.
			60	1.5 (± 0.2)	Yes (3/3)	Yes (3/3)	n.d.
			120	2.0 (± 0.2)	Yes (3/3)	Yes (3/3)	n.d.
	Pooled swab material	0.5% v/v	30	≥ 6.1 (± 0.2)	No (0/3)	tbc	+8.3 (±0.2)
	Tierre eulture fluid	0.1% v/v	30	5.7 (± 0.1)	Yes (3/3)	Yes (3/3)	+1.3 (±0.2)
SDS	rissue culture fiuld	0.5% v/v	30	≥ 6.5 (± 0.1)	Yes (1/3)	Yes (2/3)	-0.6 (±0.2)
	Pooled swab material	1.0% v/v	30	5.7 (± 0.2)	Yes (3/3)	tbc	+6.1 (±0.0)
	Tierre eulture fluid	0.1% v/v	30	≥ 6.5 (± 0.1)	No (0/3)	No (0/3)	+9.0 (±0.2)
NP40	rissue culture fiuld	0.5% v/v	30	≥ 6.5 (± 0.1)	No (0/3)	No (0/3)	+10.3 (±0.1)
	Pooled swab material	0.5% v/v	30	≥ 6.1 (± 0.2)	No (0/3)	tbc	+8.7 (±0.1)

544 545 546 n.d. - not done

† - limit of detection in TCID50 assay was 5 TCID50/mL (0.7 Log10 TCID50/mL)
 ‡ - difference in Ct in SARS-CoV-specific real-time RT-PCR compared to PBS-treated control, ± standard error

Table 5: Other reagent types 548

Reagent		Virus matrix	Reagent: virus ratio	Contact time (mins)	Titer reduction Log10 (±SE)	Virus detectable in TCID50 ⁺ (#replicates)	Virus detectable in culture (#replicates)	
Formaldehyde			4%	15	≥ 4.8 (± 0.2)	No (0/3)	No (0/3)	
		Tissue culture fluid		60	≥ 5.0 (± 0.2)	No (0/3)	No (0/3)	
			2%	15	≥ 4.8 (± 0.2)	No (0/3)	No (0/3)	
				60	≥ 5.0 (± 0.2)	No (0/3)	No (0/3)	
		Infected monolayer	4%	15	≥ 6.9 (± 0.2)	Yes (1/3)	Yes (1/3)	
				60	≥ 7.5 (± 0.2)	No (0/3)	No (0/3)	
			2%	15	≥ 6.8 (± 0.2)	Yes (2/3)	Yes (2/3)	
				60	≥ 7.3 (± 0.2)	Yes (2/3)	Yes (3/3)	
Formaldehyde + Glutaraldehyde		Tissue culture fluid	2% + 1.5%	60	≥ 5.0 (± 0.2)	No (0/3)	No (0/3)	
		Infacted monolower	29/ + 1 59/	15	≥ 6.7 (± 0.1)	No (0/3)	No (0/3)	
		infected monolayer	2/0 + 1.3/0	60	≥ 6.7 (± 0.1)	No (0/3)	No (0/3)	
Methanol‡		Infected monolayer	100%	15	≥ 6.7 (± 0.1)	No (0/3)	No (0/3)	
РНМВ	0.1%	Tissue culture fluid	10:1	30	1.4 (±0.2)	Yes (3/3)	Yes (3/3)	
	1.0%	Tissue culture fluid	10:1	30	1.5 (± 0.2)	Yes (3/3)	Yes (3/3)	
	2.0%	Tissue culture fluid	10:1	30	1.6 (± 0.2)	Yes (3/3)	Yes (3/3)	

549 550

† - limit of detection in TCID50 assay was 5 TCID50/mL (0.7 Log10 TCID50/mL)
 ‡ - ice cold methanol.

Reagent	Reagent	Manufacturer	Reagent composition	Recommended ratio of sample to	Recommended
Туре		Cat#		reagent	contact time
Specimen Transport Tube Reagents	Virus Transport and Preservation Medium (Inactivated)	BioComma Ltd. #YMJ-E	Not known	Swab placed directly into tube containing 3mL reagent	None given
	Sigma MM	Medical Wire #MWMM	Guanidine thiocyanate, Ethanol (concentrations unknown)	Up to 1 vol sample to 1.5 vols reagent (up to 0.67:1)	None given
	eNAT	Copan #608CS01R	42.5-45% guanidine thiocyanate, detergent, Tris- EDTA, HEPES.	Swab placed directly into tube containing 1 or 2mL reagent. For urine, 3:1	None given
	Primestore	Longhorn #PS-MTM-3	<50% guanidine thiocyanate, <23% ethanol	1:3	None given
	Cobas PCR	Roche #08042969001	≤40% guanidine hydrochloride, Tris-HCl	Swab placed directly into tube	None given
	Aptima Specimen Transport Medium	Hologic #PRD-03546	Not known	Swab OR 0.5mL VTM sample added to tube containing 2.9mL buffer	None given
	DNA/RNA Shield	Zymo Research #R1100	Not known	1:3	None given
	40% GHCL	Oxoid/Thermo Fisher #EB1351A	28.3% guanidine hydrochloride, 2.1% Triton X-100, Tris-EDTA	Swab placed directly into tube	None given
	2M GITC	Oxoid/Thermo Fisher #EB1349A	18.9% guanidine thiocyanate, 2.4% Triton X-100, Tris-EDTA	Swab placed directly into tube	None given
	4M GITC	Oxoid/Thermo Fisher #EB1350A	31.8% guanidine thiocyanate, 2.0% Triton X-100, Tris-EDTA	Swab placed directly into tube	None given
Molecular Extraction Reagents	NucliSENS Lysis Buffer	Biomerieux #200292	50% guanidine thiocyanate, <2% Triton X-100, <1% EDTA	1:2-1:200	10 mins
	Panther Fusion	Hologic #PRD-04339	Not known	1:1.42	
	Buffer AVL	QIAGEN #19073	50-70% guanidine thiocyanate	1:4	10 mins
	MagNA Pure 96 External Lysis Buffer	Roche #06374913001	30-50% guanidine thiocyanate, 20-25% Triton X- 100, <100mM Tris-HCl, 0.01% bromophenol blue.	1:1	None given
	Buffer AL	QIAGEN #19075	30-50% guanidine hydrochloride, 0.1-1% maleic acid	1:1	None given
	Cobas Omni LYS	Roche #06997538190	30-50% guanidine thiocyanate, 3-5% dodecyl alcohol, ethoxylated, 1-2.5% dithiothreitol	No instructions for use as off-board lysis buffer	None available
	L6 (Kingfisher formulation)	PHE Media Services	96.6% guanidine thiocyanate, 1.9% Triton X-100, Tris-EDTA	None available	None available
	Buffer RLT	QIAGEN #79216	30-50% guanidine thiocyanate	Tissue to be homogenized directly in undiluted buffer	None given
	NeuMoDx Viral Lysis Buffer	NeuMoDx Molecular,Inc. #401600	<50% guanidine hydrochloride, <5% Tween 20, <1% EDTA, <0.1% sodium azide	1:1	None given
	VPSS	E&O Laboratories #BM1675	Not known	Not known	Not known
	Lysis Buffer	E&O Laboratories #BM1676	Not known	Not known	Not known

551 Supplementary Table 1: Reagent Details

553



554

555 Figure 1: Effectiveness of five filtration matrices at removing cytotoxicity. (A) SARS-CoV-2

virus in clarified cell culture supernatant was treated with indicated reagent for 2mins at room 556 temperature before being purified through one of 5 filtration matrices: Sephadex LH-20 (blue); 557 Sephacryl S400HR (orange); Amicon Ultra 50kDa molecular weight cut off (red); Pierce 558 detergent removal spin columns (DRSC) (purple); or Bio-Bead SM2 (green). Values indicate the 559 percentage toxicity removal after one purification cycle relative to unpurified samples (based on 560 CC20 values – for more details see Table 1). (B) Percentage of input virus remaining in eluate 561 after one purification cycle through each filtration matrix. GHCl - guanidine hydrochloride; 562 GITC - guanidinium isothiocyanate; Tx - Triton X-100; PHMB - polyhexamethylene biguanide; 563 SDS - sodium dodecyl sulfate; NP40 - nonyl phenoxypolyethoxylethanol. LB - lysis buffer; TM 564 - transport medium 565

Percentage virus recovery after one filtration step



567

568 Supplementary Figure 1: Cytotoxicity of virus inactivation reagents after passing through

569 **purification matrices.** Concentration-response curves in Vero cells treated with a 2-fold serial

570 dilution of reagent. At 24 h post treatment cell viability was determined, with values normalized

- to mock treated cells. Each point represents the mean of triplicate wells, with error bars
- indicating standard deviation. Graphs are representative of at least 2 independent experiments.
- 573 Matrices used: Sephadex LH-20 (blue); Sephacryl S400HR (orange); Amicon Ultra 50kDa
- 574 molecular weight cut off (red); Pierce detergent removal spin columns (DRSC) (purple); or Bio-
- 575 Bead SM2 (green). (A) Reagents used in specimen transport tubes: GHCl guanidine
- 576 hydrochloride; GITC guanidinium isothiocyanate; Tx Triton X-100; TM Transport Medium
- 577 (B) Reagents used in molecular extraction protocols: PHMB polyhexamethylene biguanide. (C)
- 578 Detergents commonly used for virus inactivation: SDS sodium dodecyl sulfate; NP40 nonyl
- 579 phenoxypolyethoxylethanol. (D) Other reagents commonly used for virus inactivation.
- 580

Dilution Factor (1:10^x)





