2	Solar simulated ultraviolet radiation inactivates HCoV-NL63
3	and SARS-CoV-2 coronaviruses at environmentally relevant doses
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21	Keywords: Coronavirus; HCoV-NL63; SARS-CoV-2; Solar simulated ultraviolet radiation; Viral
22	inactivation

### 23 ABSTRACT

24 The germicidal properties of short wavelength ultraviolet C (UVC) light are well established and used to inactivate many viruses and other microbes. However, much less is known about 25 26 germicidal effects of terrestrial solar UV light, confined exclusively to wavelengths in the UVA 27 and UVB regions. Here, we have explored the sensitivity of the human coronaviruses HCoV-28 NL63 and SARS-CoV-2 to solar-simulated full spectrum ultraviolet light (sUV) delivered at 29 environmentally relevant doses. First, HCoV-NL63 coronavirus inactivation by sUV-exposure 30 was confirmed employing (i) viral plaque assays, (ii) RT-qPCR detection of viral genome 31 replication, and (iii) infection-induced stress response gene expression array analysis. Next, a 32 detailed dose-response relationship of SARS-CoV-2 coronavirus inactivation by sUV was 33 elucidated, suggesting a half maximal suppression of viral infectivity at low sUV doses. Likewise, 34 extended sUV exposure of SARS-CoV-2 blocked cellular infection as revealed by plaque assay 35 and stress response gene expression array analysis. Moreover, comparative (HCoV-NL63 versus 36 SARS-CoV-2) single gene expression analysis by RT-qPCR confirmed that sUV exposure blocks 37 coronavirus-induced redox, inflammatory, and proteotoxic stress responses. Based on our findings, 38 we estimate that solar ground level full spectrum UV light impairs coronavirus infectivity at 39 environmentally relevant doses. Given the urgency and global scale of the unfolding SARS-CoV-40 2 pandemic, these prototype data suggest feasibility of solar UV-induced viral inactivation, an 41 observation deserving further molecular exploration in more relevant exposure models.

42

43 Abbreviations: MOI, multiplicity of infection; sUV, solar simulated ultraviolet light; UV,
44 ultraviolet.

### 46 **1. Introduction**

47 The germicidal properties of short wavelength ultraviolet C (UVC) light are well established 48 and widely used to inactivate many viruses and other microbes, and virucidal activity of solar UVC 49 targeting pathogenic coronaviruses has been explored in much detail before [1-3]. Given the 50 urgency and global scale of the unfolding SARS-CoV-2-caused COVID-19 pandemic, UV-51 induced inactivation of coronaviruses including SARS-CoV-2 has reemerged as a matter of much 52 contemporary research interest [2-8]. Indeed, recently, rapid and complete inactivation of SARS-53 CoV-2 by UVC has been substantiated experimentally, and virucidal UVC light sources (254 nm emission) are used for surface disinfection and decontamination [5,8]. Moreover, far UVC (222 54 55 nm) has attracted considerable attention due to its potent virucidal activity [2]. However, much 56 less is known about germicidal (and coronavirus-directed) effects of terrestrial (ground level) solar 57 UV light, a matter of much interest given the airborne spread of coronaviruses including SARS-58 CoV-2 [2,6]. UVC (< 290 nm) is not present in the solar spectrum reaching the Earth's surface, 59 and most of solar UV energy incident on the skin is from the UVA region (>95%; from 320-400 60 nm). Remarkably, the UVB (290-320 nm) proportion of total solar UV-flux received by skin can 61 be well below 2% depending on the solar angle, which determines the atmospheric light path length 62 and thereby the degree of ozone-filtering and preferential Rayleigh scattering of short wavelength 63 UV light [9].

Recently, the role of ground level (environmentally relevant) solar UV has been explored in the context of SARS-CoV-2 disinfection, and a role of solar UVB in human coronavirus inactivation has been substantiated based on atmospheric and geophysical simulations [2,6,10,11]. Specifically, inactivation times of SARS coronaviruses exposed to environmental photons with wavelengths between 290-315 nm have been calculated using OMI (ozone monitoring instrument)

69 satellite data for the sunlit earth [10]. Moreover, recent research has demonstrated that simulated 70 sunlight rapidly inactivates SARS-CoV-2 on surfaces including human saliva when exposed to 71 simulated sunlight representative of the summer solstice at 40 °N latitude at sea level on a clear 72 day [10]. Also, indirect effects of solar UVB exposure in reducing COVID-19 deaths have been 73 substantiated, potentially mediated by UVB-driven cutaneous vitamin D synthesis, among other 74 factors [12-14]. In addition, a role of solar UVA photons in the inactivation of coronaviruses has 75 been proposed [7].

76 Given the complexity of virucidal activity as a function of spectral composition from ultraviolet 77 to infrared, a topic recently reviewed by various authors, a more detailed knowledge and direct 78 evidence of solar UV-induced coronavirus inactivation (achievable at ground level and 79 environmentally relevant doses) would offer improved options that inform decisions at the basic 80 research, clinical care, and public health levels [2,6,8]. Here, for the first time, we have explored 81 the sensitivity of the human coronaviruses HCoV-NL63 and SARS-CoV-2 to solar simulated 82 ultraviolet light (sUV). Our findings suggest that solar UV delivered at environmentally relevant 83 dose levels inactivates HCoV-NL63 and SARS-CoV-2 coronaviruses with pronounced blockade 84 of infectivity protecting mammalian host cells.

85

#### 86 2. Materials and Methods

87 *2.1. Chemicals* 

88 All chemicals were purchased from Sigma Aldrich (St. Louis, MO, USA).

89

### 90 2.2. Mammalian cell culture, viral propagation, and target cell infection

As established viral target cells infected by HCoV-NL63 and SARS-CoV-2, Calu-3 human
metastatic lung epithelial adenocarcinoma (HTB-55), Caco-2 human colorectal epithelial

93 adenocarcinoma (HTB-37) and Vero normal epithelial monkey kidney (CCL-81) cells (all from 94 ATCC, Manassas, VA, USA) were maintained according to published standard procedures [15-95 18]. In brief, all cells (Calu-3, Caco-2 and Vero) were cultured in Eagle's Minimum Essential 96 Medium (MEM) medium (Corning, Manassas, VA) supplemented with 10% bovine calf serum 97 (BCS, HyClone<sup>TM</sup> Laboratories, Logan, UT). Coronavirus HCoV-NL63 (NR-470) and its 98 genomic RNA (NR-44105) were obtained from BEI Resources (NIAID, NIH). SARS-CoV-2 99 strain WA1 (NR-52281; BEI Resources) was propagated in Vero cells unless specified otherwise 100 [6]. For viral stocks, cells were infected at a multiplicity of infection (MOI) of 0.01 and cultured 101 for 48 h. At that point, cells were harvested, homogenized, subjected to a single freeze-thaw cycle, 102 and then combined with the culture supernatant followed by centrifugation (3000 rpm, 10 min). 103 The viral titers of the final supernatant (after serial dilution) was determined by plaque forming 104 assay. All work with SARS-CoV-2 was performed under BSL3 conditions in a facility with 105 negative pressure and PPE that included Tyvek suits and N95 masks for respiratory protection.

106

### 107 2.3. Viral irradiation with solar simulated UV light (sUV)

108 A KW large area light source solar simulator, model 91293, from Oriel Corp. (Stratford, CT) 109 was used, equipped with a 1000W xenon arc lamp power supply, model 68920, and a VIS-IR band 110 pass blocking filter plus either an atmospheric attenuation filter (output 290-400 nm plus residual 111 650–800 nm for solar simulated light) [19,20]. For viral irradiation, viral stocks were diluted 112 >1:100 in PBS and irradiated in a sealed UV-transparent cuvette [BrandTech<sup>TM</sup> BRAND<sup>TM</sup> UV-113 Cuvets, providing transparency from 230 to 900 nm, widely used for DNA, RNA and protein 114 analysis (BrandTech<sup>™</sup> 759170, Fisher Scientific)]. The cuvette was inserted into a fully UV-115 transparent scintillation counter vial (Wheaton '180' low-potassium glass, SigmaAldrich

116 Z253081). The UV output was quantified using a dosimeter from International Light Inc. 117 (Newburyport, MA), model IL1700, with an SED240 detector for UVB (range 265–310 nm, peak 118 at 285 nm) or a SED033 detector for UVA (range 315–390 nm peak 365 nm) at a distance of 365 119 mm from the source, which was used for all experiments. In order to avoid artifactual thermal 120 effects of photon exposure on viral activity, cuvettes were placed on ice during irradiation. At 365 121 mm from the source, total solar UV intensity was 5.34 mJ/cm<sup>2</sup> s (UVA) and 0.28 mJ/cm<sup>2</sup> s (UVB).

122

### 123 2.4. HCoV-NL63 plaque forming assay and viral RNA quantification

124 A published standard procedure was followed [15,21]. For HCoV-NL63, target cells (CaCo-2 125 or Calu-3) were seeded in 6-well plates at approximately  $4 \times 10^5$  cells per well and incubated until 126 the monolayer was 80–90% confluent. Prior to infection, cells were washed with phosphate 127 buffered saline (PBS). Virus inoculum (MOI=0.01) in 500 µL of growth media supplemented with 128 2% horse serum (with standard penicillin/streptomycin and L-glutamine supplementation) was 129 added to each well. Viral entry was performed by incubation at 4°C for 30-60 min with gentle 130 agitation followed by 1 h incubation in 33°C, 5% CO<sub>2</sub>. Then, inoculum was removed and cells 131 were washed twice with PBS and replaced by 2 mL of normal growing media. After infection, 132 cells were washed twice with PBS and placed in the incubator and cultured in normal growth 133 media. Once plaques appeared (~5-7 d post infection), cells were fixed with 10% neutral buffered 134 formalin for 30 min at room temperature and stained with 1% crystal violet in 20% methanol for 135 20 min. Then, cells were washed several times with water, and plaques were counted and 136 representative pictures taken at 10x magnification using an inverted microscope (Nikon 137 Instruments, Melville, NY). In addition, viral RNA was extracted from cells and the respective 138 culture supernatant with the QIA amp Viral RNA Mini Kit (QIAGEN, Hilden, Germany). One step

- 139 RT-qPCR for HCoV-NL63 with absolute virus RNA quantification was performed using the
- 140 following primer/probe set as published before [22]:
- 141 forward primer 5'-ACGTACTTCTATTATGAAGCATGATATTAA-3'
- 142 reverse primer 5'-AGCAGATCTAATGTTATACTTAAAACTACG-3'
- 143 probe FAM-5'- ATTGCCAAGGCTCCTAAACGTACAGGTGTT -3'-NFQ-MGB

144 Briefly, RT-qPCR was carried out in a 20 µL reaction mixture with extracted RNA and One 145 step RT-qPCR 2x Master Mix containing ROX as a passive reference dye (Gold Biotechnology, 146 St. Louis, MO) and 300 nM forward and reverse primers and 200 nM MGB probe. Amplification 147 and detection were performed in ABI 7500 system (Applied Biosystems, Foster city, CA) under 148 the following conditions: first strand cDNA synthesis at 42°C for 30 min; initial denaturation/RT 149 inactivation at 95°C for 3 min; denaturation at 95°C for 10 sec and annealing/extension at 55°C 150 for 30 sec followed by 45 sec for data acquisition at 72°C. During amplification, the ABI PRISM 151 7500 sequence detector monitored real-time PCR amplification by quantitative analysis of the 152 fluorescence emissions. The reporter dye (FAM) signal was measured against the internal 153 reference dye (ROX) signal to normalize the signals for non-PCR-related fluorescence fluctuations 154 that occur from well to well. The cycle threshold (Ct) represented the refraction cycle number at 155 which a positive amplification was measured and was set at ten times the standard deviation of the 156 mean baseline emission calculated for PCR cycles 3 to 15. Genomic RNA from HCoV-NL63 was 157 used as a positive control.

158

### 159 2.5. SARS-CoV-2 plaque forming assay and viral RNA quantification

160 The quantification of infectious SARS-CoV-2 has been published before [18]. Target cells 161 (Vero or Calu-3) were infected in triplicates at an MOI of 0.005 (high titer) or 0.001 (low titer).

162 Briefly, cells were incubated with SARS-CoV-2 for 2 h and subsequently overlaid with 1%

163 methylcellulose in culture medium. After 3-4 days, the cells were fixed in 10% neutral buffered

- 164 formalin for 30 min, washed under tap water, and stained with 1% crystal violet. The number of
- 165 plaques was counted on a light table. Alternatively, infection of cells was determined by measuring
- 166 the amount of viral RNA. Cells were lysed in Trizol followed by RNA extraction with the
- 167 RNAeasy kit (Qiagen). After reverse transcription, cDNA corresponding to the gene encoding the
- 168 SARS-CoV-2 spike protein was quantified by qPCR with the Perfecta FastMix (QuantaBio) using:
- 169 forward primer (SARS-CoV-2) 5'-GCTGGTGCTGCAGCTTATTA-3'
- 170 reverse primer (SARS-CoV-2) 5'-AGGGTCAAGTGCACAGTCTA-3'
- at an annealing temperature of 60 °C. For normalization, *GAPDH* expression was measured using
  the following primers:
- 173 forward primer (*GAPDH*) 5'-TGGTGAAGGTCGGTGTGAAC-3'
- 174 reverse primer (GAPDH) 5'-CCATGTAGTTGAGGTCAATGAAGG-3'.
- 175

# 176 2.6. Human Stress & Toxicity PathwayFinder RT<sup>2</sup> Profiler<sup>TM</sup> gene expression array analysis of 177 infected host cells

Seven days post infection of Calu-3 host cells with either HCoV-NL63 (MOI=0.01) or HCoV-NL63 exposed to sUV (UVB portion: 706 mJ/cm<sup>2</sup>), total mRNA from host cells was isolated using the RNeasy Mini kit (Qiagen, Valencia, CA) following our published standard procedures. Reverse transcription was then performed using the RT<sup>2</sup> First Strand kit (Qiagen) from 500 ng total RNA. For gene expression array analysis, the human Stress & Toxicity PathwayFinder RT<sup>2</sup> Profiler<sup>TM</sup> technology (Qiagen), assessing expression of 84 stress response-related genes, was used as published before [23,24]. Quantitative PCR was run using the following conditions: 95 °C (10 min), followed by 40 cycles at 95 °C (15 s) alternating with 60 °C (1 min) (Applied Biosystems, Carlsbad, CA). Gene-specific products were normalized to a group of 5 housekeeping genes (*ACTB*, *B2M*, *GAPDH*, *HPRT1*, *RPLP0*) and quantified using the comparative  $\Delta\Delta$ Ct method (ABI PRISM 7500 sequence detection system user guide). Expression values were averaged across at least three independent array experiments, and standard deviation was calculated for graphing and statistical analysis as published before.

191

### 192 2.7. Individual RT-qPCR analysis

193 Total cellular mRNA was isolated using the Qiagen RNeasy Mini Kit (Qiagen, Gaithersburg, 194 MD) according to the manufacturer's protocol as published by us before [24]. Human primer 195 probes [CCL3 (Hs 00234142 m1), CSF2 (Hs 00929873 m1), HSPA6 (Hs 00275682 s1), IL1B 196 (Hs 00985639 m1), (Hs 00174097 m1), IL6 SOD2 (Hs 00167309 m1), TNF 197 (Hs 00174128 s1), and RSP18 (housekeeping gene; Hs 01375212 g1)], were obtained 198 from ThermoFisher Scientific (Waltham, MA). After cDNA synthesis, quantitative PCR reactions 199 were performed as follows: 10 min (95 °C) followed by 15 sec (95 °C), 1 min (60 °C), 40 cycles, 200 using the ABI7500 Real-Time PCR System (Applied Biosystems, Foster City, CA). Amplification 201 plots were generated, and Ct values were recorded as published before [24].

202

### 203 2.8. Statistical analysis

Unless stated differently, data sets were analyzed employing analysis of variance (ANOVA) with Tukey's posthoc test using the GraphPad Prism 9.1.0 software (Prism Software Corp., Irvine, CA); in respective bar graphs (analyzing more than two groups), means without a common letter differ (p < 0.05) as published before [24]. For bar graphs comparing two groups only, statistical

208 significance was calculated employing the Student's two-tailed t-test, utilizing Excel (Microsoft,

209 Redmond, WA). Experiments were performed in sets of at least three independent repeats. The

- level of statistical significance was marked as follows: \*p < 0.05; \*\*p < 0.01; \*\*\*p < 0.001.
- 211
- 212 **3. Results**

3.1. Solar simulated UV exposure of HCoV-NL63 blocks subsequent viral infection and
replication in Calu-3 human epithelial lung cells

215 First, we examined the feasibility of UV-inactivation of a pathologically relevant coronavirus 216 by employing a single dose of solar simulated UV light using a commercial xenon light source 217 with quantified spectral power distribution (Fig. 1A). To this end, we exposed human coronoavirus 218 NL63 (HCoV-NL63) in PBS to a high dose of sUV [equivalent to approximately 6 minimal 219 erythemal doses (MEDs; UVA: 13.46 J/cm<sup>2</sup>; UVB: 706 mJ/cm<sup>2</sup>)] and subsequently used it to infect 220 Calu-3 target cells for 7 days [2, 6, 8]. We used unexposed virus as controls. Strikingly, sUV pre-221 exposure strongly suppressed viral infectivity of target cells as demonstrated by quantitative 222 plaque assay analysis, indicating that sUV exposure caused a more than 8-fold decrease in viral 223 infectivity (Fig. 1B).

Next, we examined the dose-response relationship characterizing the inhibition of HCoV-NL63
viral replication (induced by sUV pre-exposure) by one step RT-qPCR analysis of the genomic
RNA copy number. We detected a significant inhibition at low sUV doses [UVA: 0.25 J/cm<sup>2</sup>; 13
mJ/cm<sup>2</sup> UVB]. Viral inactivation of more than 98 % occurred at doses equal and above 480 mJ/cm<sup>2</sup>
UVB (UVA: 9.04 J/cm<sup>2</sup>; Fig. 1C).

- 229
- 230 >Figure 1<

### 231 3.2. Solar simulated UV exposure of HCoV-NL63 blocks subsequent infection of Caco-2 human

### 232 epithelial colorectal cells

233 In order to explore sUV effects on HCoV-NL63 infectivity in another human target cell, we 234 exposed the virus (in PBS) to a high dose of sUV [equivalent to approximately 6 MEDs (UVA: 235 13.46 J/cm<sup>2</sup>; UVB: 706 mJ/cm<sup>2</sup>)] and subsequently infected Caco-2 epithelial colon cells (Fig. 2). 236 As observed before with Calu-3 cells (Fig. 1), our quantitative plaque assay analysis showed that 237 the suppression of viral infectivity of Caco-2 target cells by sUV exposure caused a more than 4-238 fold decrease in plaque formation (Fig. 2A). Likewise, our dose response analysis by RT-qPCR of 239 genomic RNA copy numbers indicated that sUV exposure caused a pronounced suppression of 240 HCoV-NL63 viral replication at doses as low as 240 mJ/cm<sup>2</sup> UVB (UVA: 4.52 J/cm<sup>2</sup>; Fig. 2B). 241

242

### >Figure 2<

243

## 3.3. Stress response gene expression array analysis confirms solar UV-induced inhibition of HCoV-NL63 infectivity targeting Calu-3 human epithelial lung cells

246 Next, the cellular stress response of Calu-3 human epithelial lung cells, elicited by infection 247 with either mock-irradiated or sUV pre-exposed HCoV-NL63, was examined at the gene 248 expression level using the RT<sup>2</sup> Human Stress and Toxicity PathwayFinder<sup>TM</sup> PCR Array 249 technology. To this end, we infected Calu-3 target cells with sUV or mock-treated virus (doses as 250 in Figs. 1, 2) and profiled the gene expression at the end of the experiment. We observed global 251 HCoV-NL63-induced expression changes (antagonized by viral pre-exposure to sUV) as depicted 252 by Volcano plot (Fig. 3). As expected, HCoV-NL63 viral infection caused a pronounced 253 upregulation of stress response gene expression including genes encoding key regulators of

254 inflammatory signaling (such as CSF2, TNF, IL1B, IL1A, CCL3, CXCL10, NFKBIA, and IL6), 255 oxidative stress defense (such as SOD2), and heat shock response (such as HSPA6; Fig. 3). In 256 contrast, after viral sUV-exposure performed pre-infection, most of these infection-associated 257 expression changes were either attenuated or completely obliterated, an observation consistent 258 with pronounced suppression of HCoV-NL63 viral infectivity as a consequence of sUV-exposure. 259 Likewise, HCoV-NL63 viral infection-induced expression changes causing downregulation of 260 specific apoptotic modulators including BCL2L1, EGR1, CASP8, and CASP1, proliferation 261 markers such as PCNA, and heat shock response factors such as HSPA4, HSPH1, and HSP90AA2P 262 were completely absent in samples obtained from cells exposed to the pre-irradiated virus. 263 Strikingly, expression of seven specific genes (CDKN1A, CYP1A1, MDM2, HMOX1, RAD50, 264 HSPA1L, and E2F1) was modulated uniquely in response to exposure to sUV-preirradiated HCoV-265 NL63, a finding consistent with gene expression changes responsive to sUV-induced chemical 266 damage to viral components (including ribonucleic acids, proteins, and lipids) [1-3].

267

268

>Figure 3<

269

3.4. Dose-response relationship of solar simulated UV-induced inhibition of SARS-CoV-2
infectivity targeting Vero and Calu-3 mammalian cells

After demonstrating HCoV-NL63 coronavirus inactivation by sUV at an environmentally relevant dose level, we examined whether sUV-inactivation might also be applicable to SARS-CoV-2. To this end, we exposed the virus with a dose range of sUV, subsequently infected Vero monkey epithelial cells at two different multiplicities of infection (MOIs, high versus low titer), and measured the number of infectious virions three days later by plaque forming assay.

277 Strikingly, as observed with HCoV-NL63, sUV exposure caused a pronounced suppression of viral 278 infectivity. This antiviral effect, observable over a broad range of sUV doses, followed an 279 exponential decay curve with an effective ED<sub>50</sub> (sUV dose diminishing SARS-CoV-2 viral 280 infectivity by 50%) approximating 55 mJ/cm<sup>2</sup> (low titer) and 62 mJ/cm<sup>2</sup> (high titer) (Fig. 4A). 281 Next, we tested feasibility of achieving complete inhibition of SARS-CoV-2 replication by high 282 dose sUV [UVB portion: 1010 mJ/cm<sup>2</sup>, a maximum dose level similar to the one used in the HCoV-283 NL63-directed dose-response experiments (Fig. 1C)]. To this end, we pre-exposed SARS-CoV-2 284 to sUV and measured the amount of viral RNA (corresponding to the region of the viral genome 285 encoding the S protein) by RT-qPCR analysis. Indeed, complete inhibition was achieved at that 286 dose (Fig. 4B). We obtained similar results for sUV-exposed SARS-CoV-2 infections of Calu-3 287 human lung epithelial target cells with viral load in supernatants being monitored over three days 288 by RT-qPCR (Fig. 4C). Taken together, we conclude that SARS-CoV-2 is sensitive to sUV 289 suggesting viral inactivation at environmentally relevant exposure levels.

290

291

>Figure 4<

292

## 3.5. Solar simulated UV exposure of SARS-CoV-2 prevents stress response gene expression elicited by viral infection of Calu-3 human epithelial lung cells as detected by array analysis

Next, to determine Calu-3 human epithelial lung cell stress response gene expression elicited by SARS-CoV-2 as a function of viral pre-exposure to sUV, we employed expression analysis using the Human Stress and Toxicity PathwayFinder<sup>TM</sup> PCR Array technology. To this end, we infected Calu-3 target cells with sUV or mock-treated virus as outlined before, followed by comparative gene expression profiling at the end of the experiment. We observed multiple SARS-

300	CoV-2-induced expression changes (antagonized by viral pre-exposure to sUV) as shown in the
301	Volcano plot depiction [displaying statistical significance (P value) versus magnitude of change
302	(fold change)] (Fig. 5). SARS-CoV-2 infection caused a pronounced upregulation of stress
303	response gene expression including genes encoding key regulators of inflammatory signaling
304	including IL1A, IL1B, IL6, TNF, CCL3, CXCL10, CSF2, and NFKBIA, oxidative stress defense
305	such as SOD2, and heat shock response such as HSPA6 (Fig. 5). In contrast, after infection with
306	sUV-exposed virus, most of these infection-associated expression changes were either attenuated
307	or completely obliterated, an observation consistent with pronounced suppression of SARS-CoV-
308	2 infectivity as a consequence of sUV-exposure. Remarkably, these expression changes closely
309	mirrored those observed in response to HCoV-NL63 infection that occurred with or without viral
310	exposure to sUV (Fig. 3).
211	

- 311
- 312

#### >Figure 5<

313

314 Likewise, we observed a striking similarity between the gene expression changes elicited by 315 HCoV-NL63 and SARS-CoV-2 (and blocked by viral sUV pre-exposure), modulating redox, 316 inflammatory, and proteotoxic stress responses in Calu-3 human epithelial lung cells (Fig. 6). 317 Specifically, sUV-induced (UVB portion: 706 mJ/cm<sup>2</sup>) viral inactivation was apparent from independent RT-qPCR assessment of mRNA levels ('no sUV' versus 'sUV') interrogating genes 318 319 encoding key regulators of redox (SOD2), inflammatory (IL1B, TNF, CCL3, IL6, CSF2), and 320 proteotoxic ('heat shock'; HSPA6) stress responses in Calu-3 target cells as detailed above. Thus, 321 our data suggest that similar to HCoV-NL63, sUV exposure of SARS-CoV-2 interrupts the viral

life cycle causing suppression of viral replication and virus-induced inflammatory and cellular
 stress responses in mammalian target cells.

- 324
- 325 >Figure 6<
- 326

### 327 **4. Discussion**

328 Identification and mechanistic exploration of environmental factors that might determine 329 coronavirus infectivity are of significant interest with relevance to both basic molecular research 330 and public health-related preventive and interventional investigations [2]. Here, we have explored 331 for the first time the effects of full spectrum (UVA + UVB) solar ultraviolet radiation on 332 coronavirus infectivity and demonstrate that sUV inactivates HCoV-NL63 and SARS-CoV-2 333 coronaviruses at environmentally relevant doses. First, we observed that exposure of HCoV-NL63 334 and SARS-CoV-2 to sUV (performed at acute dose levels relevant to human populations 335 worldwide) blocks subsequent viral infection and replication in relevant primate target cells 336 [human: Calu-3 lung epithelial, Caco-2 colorectal epithelial; monkey: Vero kidney epithelial (Figs. 337 1, 2, 4)]. Blockade of viral infectivity in response to sUV pre-exposure was also confirmed using 338 stress response gene expression profiling in array (Figs. 3, 5) and independent RT-qPCR format 339 (Fig. 6) elicited in Calu-3 target cells by coronavirus infection (HCoV-NL63 and SARS-CoV-2). 340 Remarkably, dose levels used throughout this pilot study are representative of terrestrial ground 341 level exposure suggesting environmental relevance, and significant coronavirus inactivation was 342 detectable even at low exposure levels expected to be beneath the cutaneous sunburn-inducing 343 threshold (Figs. 1, 2, 4) [2,6,8]. In this context, it is remarkable that recent research has already

344 indicated that ground level solar UV displays significant virucidal effects targeting coronaviruses

345 including SARS-CoV-2 [2,6,11,13]. However, the complexity of human exposure levels to solar 346 UV as a function of solar zenith angle, seasonality, spectral distribution, and latitude remain to be 347 addressed before any firm conclusions relevant to human populations can be drawn. Specifically, 348 the anti-viral activity of specific spectral components of sUV remains to be determined since the 349 light source employed in our prototype studies emitted full spectrum simulated solar UV, and the 350 action spectrum of virus inactivation by solar UV remains largely undefined. For example, it is 351 possible that the UVA portion of ground level sUV significantly contributes to the coronavirus-352 directed effects described by us [7]. It therefore remains to be seen if indirect impairment of viral 353 structure and infectivity occurs by alternative mechanisms, such as UVA-driven photosensitization 354 and oxidative stress (mediated by formation of reactive oxygen species including singlet oxygen), 355 that might be operative in addition to direct inactivation of viral genomic RNA through nucleic 356 acid base photodamage. It will also be interesting to explore potential mechanistic synergisms 357 underlying virucidal effects that occur upon combined UVA and UVB as compared to separate 358 spectral exposure. Likewise, experimental conditions used throughout our studies (including viral 359 irradiation in PBS and exposure performed in cell culture medium) might limit the applicability of 360 our conclusions in the context of relevant coronavirus transmission situations that involve more 361 complex determinants of infectivity including the role air-borne and aerosol transmission and 362 intermediate surface retention [6].

Addressing urgency and global scale of the unfolding SARS-CoV-2 pandemic requires an improved understanding of environmental factors that modify viral infectivity [2,6,8]. Taken together, our data suggest feasibility of sUV-induced viral inactivation targeting HCoV-NL63 and SARS-CoV-2 coronaviruses, a finding to be substantiated by future mechanistic exploration performed in more relevant *in vivo* exposure models.

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375

- **Declaration of competing interest**
- 377 The authors state no conflict of interest.
- 378

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### 495 Figure legends

496 Figure 1. Solar simulated UV pre-exposure antagonizes HCoV-NL63 viral infectivity 497 targeting Calu-3 human epithelial lung cells. Virus in PBS was exposed to sUV or left 498 unexposed followed by Calu-3 target cell infection (0.01 MOI) and post infection culture over 7 499 days followed by analysis. (A) Spectral power distribution (irradiance) of the solar simulator light 500 source equipped with appropriate cut-off filter (sUV: UVB + UVA, solid black line). (B) Plaque 501 assay after viral exposure to sUV (UVB portion: 706 mJ/cm<sup>2</sup>) as visualized by light microscopy 502 (10 x magnification); bar graph summarizes numerical data. (C) RT-qPCR of viral genome 503 replication in target cells [left panel: amplification curves as a function of sUV dose (UVB portion 504 as indicated); right panel: bar graph summarizing numerical data].

505

Figure 2. Solar simulated UV pre-exposure antagonizes HCoV-NL63 viral infectivity targeting Caco-2 human epithelial colorectal cells. Virus in PBS was exposed to sUV or left unexposed followed by Caco-2 target cell infection (0.01 MOI) and post infection culture (7 days) followed by analysis. (A) Plaque assay after viral exposure to sUV (UVB portion: 706 mJ/cm<sup>2</sup>) as visualized by light microscopy (10 x magnification); bar graph summarizes numerical data. (B) RT-qPCR detection of viral genome replication in target cells; left panel: amplification curves (as a function of sUV-dose); right panel: bar graph summarized numerical data.

Figure 3. Solar simulated UV pre-exposure of HCoV-NL63 prevents stress response gene expression elicited in Calu-3 human epithelial lung target cells. Treatments were performed as detailed in Fig. 1. (A) Target cell stress response [control (HCoV-NL63) versus sUV (UVB portion: 706 mJ/cm<sup>2</sup>) pre-exposed virus] assessed by  $RT^2$  Profiler<sup>TM</sup> Stress and Toxicity Pathway

gene expression array analysis [volcano plot depiction: p value over log2 (fold expression change)]. (B) Scatter plot depiction comparing expression changes elicited by untreated control virus (top panel) or sUV pre-exposed virus (bottom panel). (C) Venn diagram depicting expression changes induced by mock-irradiated virus (control) versus sUV pre-irradiated virus. (D) Tabular summary of numerical values specifying gene expression changes at the mRNA level (p<0.05).</p>

523

524 Figure 4. Solar simulated UV exposure of SARS-CoV-2 antagonizes subsequent viral 525 infection and replication in African green monkey Vero and Calu-3 human epithelial lung cells. (A) SARS-CoV-2 was sUV-irradiated (UVB portion: up to 480 mJ/cm<sup>2</sup>; or remained 526 527 unirradiated) in PBS and subsequently used to infect Vero cells at two different MOIs (high versus 528 low titer). Dose response of plaque formation as a function of sUV pre-exposure dose was 529 assessed; a representative experiment (left panel, top and bottom rows) and quantification (right 530 panels) are depicted. (B) Detection of viral genome replication in Vero cells with quantification 531 of viral RNA after infection using mock or sUV pre-irradiated virus (UVB portion: 1010 mJ/cm<sup>2</sup>) 532 as assessed by RT-qPCR after 24 h. (C) Infection of Calu-3 cells with SARS-CoV-2 [sUV pre-533 exposed (UVB portion: 706 mJ/cm<sup>2</sup>) versus unirradiated virus]. The presence of infectious virions 534 in the supernatants was quantified over the course of three days post infection by RT-qPCR (nd: 535 not detectable).

536

Figure 5. Solar simulated UV pre-exposure of SARS-CoV-2 prevents stress response gene expression elicited in Calu-3 human epithelial lung target cells. Treatment and analysis were performed as detailed in Fig. 3. (A) Target cell stress response [control (SARS-CoV-2) versus sUV (UVB portion: 706 mJ/cm<sup>2</sup>)-preirradiated virus] assessed by  $RT^2$  Profiler<sup>TM</sup> Stress and

*Toxicity Pathway* gene expression array analysis [volcano plot depiction: p value over log2 (fold
expression change)]. (B) Scatter plot depiction comparing expression changes elicited by untreated
control virus (top panel) or sUV pre-exposed virus (bottom panel). (C) Venn diagram depicting
expression changes induced by mock-irradiated virus (control) versus sUV pre-irradiated virus.
(D) Tabular summary of numerical values of gene expression changes at the mRNA level (p<0.05).</li>

Figure 6. Comparative analysis of redox, inflammatory, and proteotoxic stress response gene expression in Calu-3 human epithelial lung cells elicited by HCoV-NL63 and SARS-CoV-2 (with and without viral sUV pre-exposure). Gene expression as assessed by single RT-qPCR quantification in virus-exposed target cells as a function of viral pre-exposure ['no sUV' versus 'sUV' (UVB portion: 706 mJ/cm<sup>2</sup>)]. Bar graphs depict fold change ('sUV' versus 'no sUV') normalized to housekeeping gene expression (*RPS18*; gray bar: no sUV pretreatment; black bar: sUV-pretreatment).

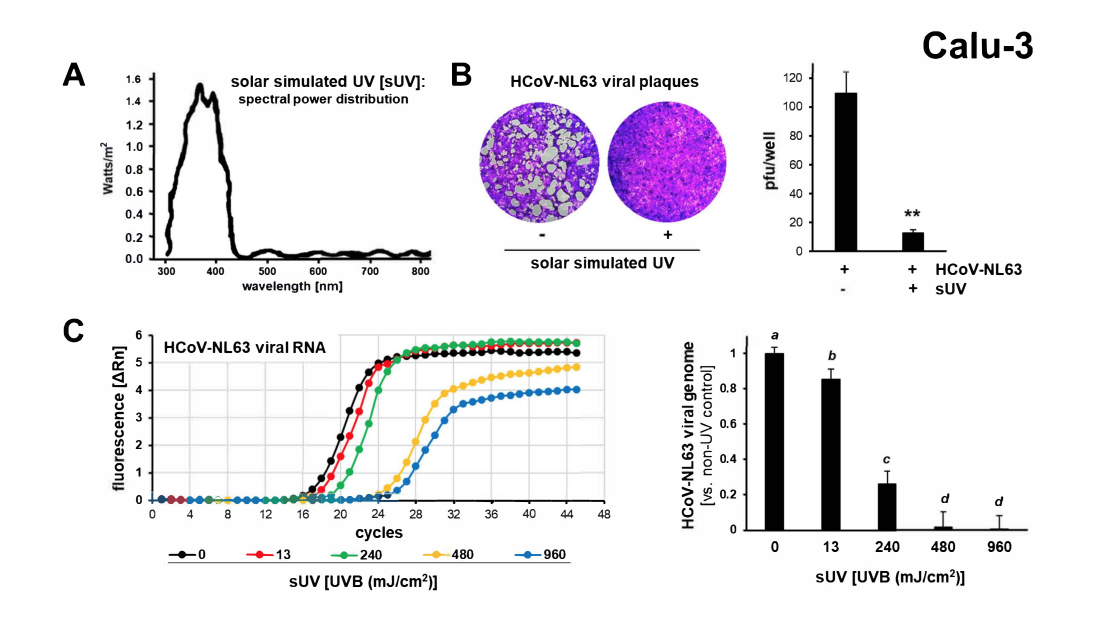
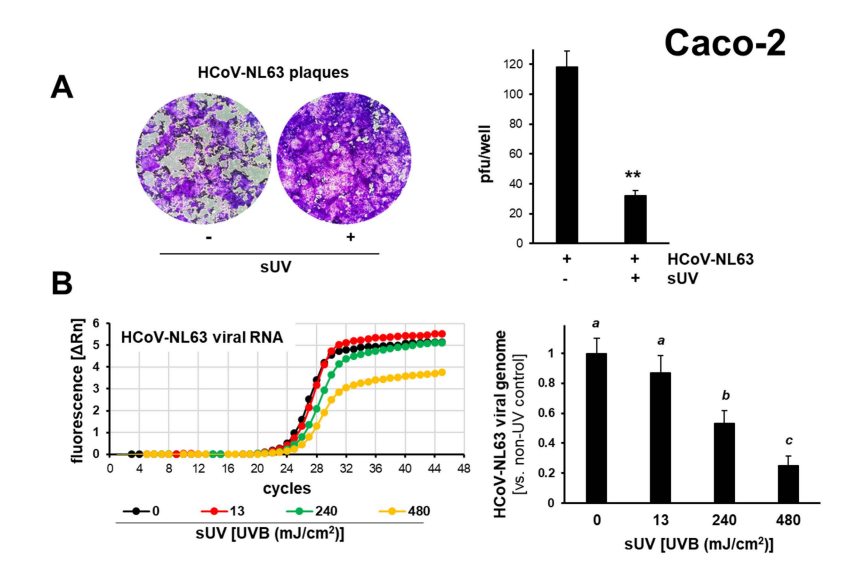
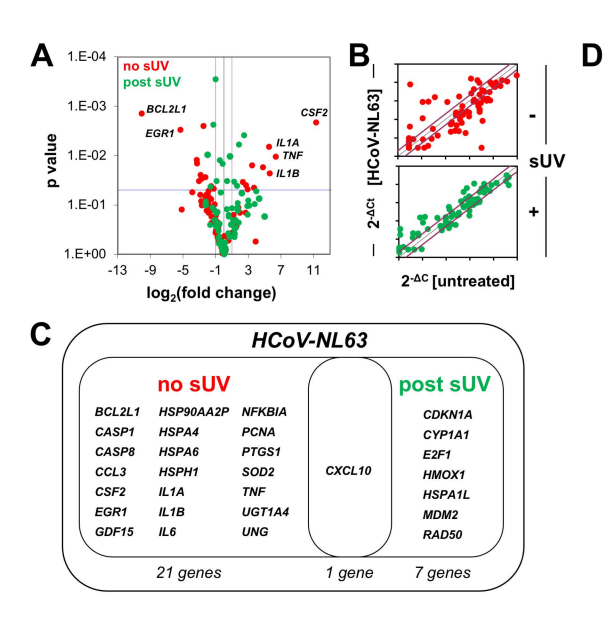


Fig. 2





## Calu-3

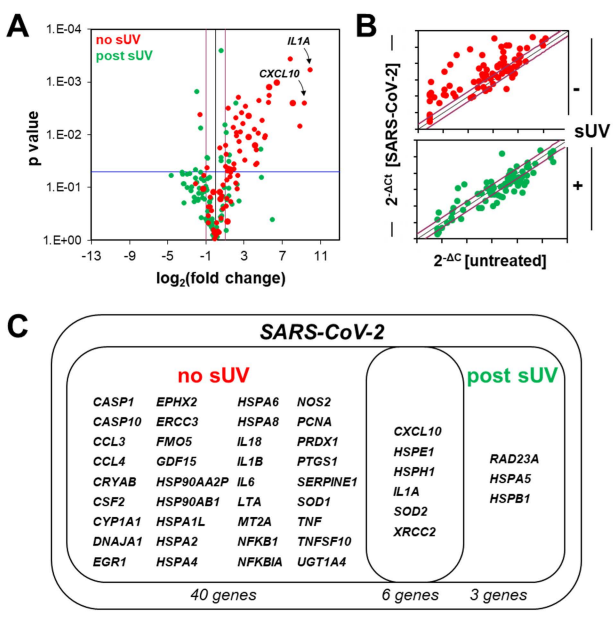
### HCoV-NL63

Gene	Description	fold change (p≤0.05)	
Gene		no sUV	post sUV
CSF2	Colony stimulating factor 2 (granulocyte-macrophage)	2452.4	
TNF	Tumor necrosis factor	81.6	-
IL1B	Interleukin 1, beta	48.3	-
IL1A	Interleukin 1, alpha	45.9	
SOD2	Superoxide dismutase 2, mitochondrial	27.3	-
CCL3	Chemokine (C-C motif) ligand 3	12.9	-
HSPA6	Heat shock 70kDa protein 6 (HSP70B')	10.8	-
CXCL10	Chemokine (C-X-C motif) ligand 10	7.4	3.6
NFKBIA	NFKB inhibitor alpha	6.5	-
CYP1A1	Cytochrome P450, family 1, subfamily A, polypeptide 1	-	5.5
IL6	Interleukin 6 (interferon, beta 2)	4.9	
CDKN1A	Cyclin-dependent kinase inhibitor 1A (p21, Cip1)	-	3.0
MDM2	Mdm2 p53 binding protein homolog	-	2.4
HMOX1	Heme oxygenase (decycling) 1	-	-2.1
PTGS1	Prostaglandin-endoperoxide synthase 1	-2.4	-
RAD50	RAD50 homolog	-	-2.4
CASP1	Caspase 1, apoptosis-related cysteine peptidase	-3.0	-
HSPA1L	Heat shock 70kDa protein 1-like	-	-3.1
E2F1	E2F transcription factor 1	-	-4.0
UGT1A4	UDP glucuronosyltransferase 1 family, polypeptide A4	-4.9	
HSP90AA2P	Heat shock protein 90kDa alpha, class A member 2	-5.8	1 <b>-</b> 1
PCNA	Proliferating cell nuclear antigen	-6.3	-
GDF15	Growth differentiation factor 15	-6.7	-
UNG	Uracil-DNA glycosylase	-7.3	-
HSPH1	Heat shock 105kDa/110kDa protein 1	-8.4	-
CASP8	Caspase 8, apoptosis-related cysteine peptidase	-9.8	-
HSPA4	Heat shock 70kDa protein 4	-10.3	-
EGR1	Early growth response 1	-40.5	-
BCL2L1	BCL2-like 1	-1078.6	-

Fig. 4

Α Vero high titer low titer 350 120 300 100  $ED_{50} = 54.9$  $ED_{50} = 62.4$ high titer 250 pfu/well pfu/well (95% CI 54.2 - 73.3) (95% CI 45.2 - 69.8) 80 200  $R^2 = 0.99$  $R^2 = 0.97$ 60 150 low titer 40 100 20 50 13 240 480 0 60 120 0 0 sUV [UVB (mJ/cm<sup>2</sup>)] **6** 510 Ó Ō 510 170 340 170 340 sUV [UVB (mJ/cm<sup>2</sup>)] sUV [UVB (mJ/cm<sup>2</sup>)] B С Calu-3 Vero SARS-CoV-2 viral genome infectious SARS-CoV-2 in supernatant [pfu/well] 5x10<sup>5</sup> 1.5 [relative abundance (fold)] \*\*\* \*\*\* 4x10<sup>5</sup> 1.0 3x10<sup>5</sup> 2x10<sup>5</sup> \*\*\* 0.5 1x10<sup>5</sup> nd nd nd 0 0 sUV sUV ÷ ÷ + t day 3 day 2 day 1

Fig. 5



		SARS-CoV-2		
•	B	fold change (p≤0.05)		
Gene	Description	no sUV	post sUV	
IL1A	Interleukin 1, alpha	982.3	26.9	
CXCL10	Chemokine (C-X-C motif) ligand 10	654.8	2.0	
IL1B	Interleukin 1, beta	469.5	-	
TNF	Tumor necrosis factor	284.0	-	
CCL3	Chemokine (C-C motif) ligand 3	230.7	-	
IL6	Interleukin 6 (interferon, beta 2)	89.6	-	
CSF2	Colony stimulating factor 2 (granulocyte-macrophage)	51.1	-	
NFKBIA	NFKB inhibitor alpha	50.6	-	
CCL4	Chemokine (C-C motif) ligand 4	48.2	-	
EPHX2	Epoxide hydrolase 2, cytoplasmic	37.4	-	
TNFSF10	Tumor necrosis factor (ligand) superfamily, member 10	36.1	-	
SOD2	Superoxide dismutase 2, mitochondrial	29.9	2.7	
UGT1A4	UDP glucuronosyltransferase 1 family, polypeptide A4	23.4	-	
CYP1A1	Cytochrome P450, family 1, subfamily A, polypeptide 1	23.2	-	
MT2A	Metallothionein 2A	20.5	-	
LTA	Lymphotoxin alpha (TNF superfamily, member 1)	19.6	-	
HSP90AA2P	Heat shock protein 90kDa alpha, class A member 2	19.0		
XRCC2	X-ray repair cross complementing 2	18.2	4.4	
NOS2	Nitric oxide synthase 2, inducible	17.1	-	
HSPA6	Heat shock 70kDa protein 6 (HSP70B')	11.5	-	
HSPE1	Heat shock 10kDa protein 1 (chaperonin 10)	11.2	2.9	
CRYAB	Crystallin, alpha B	9.3	-	
PRDX1	Peroxiredoxin 1	9.1	-	
HSPH1	Heat shock 105kDa/110kDa protein 1	8.5	2.2	
CASP1	Caspase 1, apoptosis-related cysteine peptidase	7.5	-	
SOD1	Superoxide dismutase 1, soluble	6.9	-	
HSPA1L	Heat shock 70kDa protein 1-like	6.3 6.0	-	
SERPINE1	Serpin peptidase inhibitor, clade E, member 1		-	
PTGS1	Prostaglandin-endoperoxide synthase 1	5.5	-	
FMO5 HSPA4	Flavin containing monooxygenase 5	5.5 4.6	2	
CASP10	Heat shock 70kDa protein 4	4.6	-	
PCNA	Caspase 10, apoptosis-related cysteine peptidase Proliferating cell nuclear antigen	4.6	-	
HSPA2	Heat shock 70kDa protein 2	4.5	-	
NFKB1	Nuclear factor of kappa 1	3.9	-	
DNAJA1	DnaJ (Hsp40) homolog, subfamily A, member 1	3.8	-	
HSP90AB1	Heat shock protein 90kDa alpha, class B member 1	2.7		
HSP90AB1 HSPA8	Heat shock 70kDa protein 8	2.3	-	
IL18	Interleukin 18 (interferon-gamma-inducing factor)	2.3	-	
GDF15	Growth differentiation factor 15	-3.1	-	
EGR1	Early growth response 1	-12.4	-	
RAD23A	RAD23 homolog A (S. cerevisiae)	-12.4	-2.8	
HSPA5	Heat shock 70kDa protein 5		-3.8	
HSPB1	Heat shock 27kDa protein 1		-4.3	
	How show Zhibu protoin 1	-	4.0	

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