1	Phage-like particle vaccines are highly immunogenic and protect against pathogenic
2	coronavirus infection and disease
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20	Running title: Phage vaccines against SARS-CoV-2 and MERS-CoV

21 ABSTRACT

22 The response by vaccine developers to the COVID-19 pandemic has been extraordinary with 23 effective vaccines authorized for emergency use in the U.S. within one year of the appearance of 24 the first COVID-19 cases. However, the emergence of SARS-CoV-2 variants and obstacles with 25 the global rollout of new vaccines highlight the need for platforms that are amenable to rapid 26 tuning and stable formulation to facilitate the logistics of vaccine delivery worldwide. We 27 developed a "designer nanoparticle" platform using phage-like particles (PLPs) derived from 28 bacteriophage lambda for multivalent display of antigens in rigorously defined ratios. Here, we 29 engineered PLPs that display the receptor binding domain (RBD) protein from SARS-CoV-2 and 30 MERS-CoV, alone (RBD_{SARS}-PLPs, RBD_{MERS}-PLPs) and in combination (hCoV-RBD PLPs). 31 Functionalized particles possess physiochemical properties compatible with pharmaceutical 32 standards and retain antigenicity. Following primary immunization, BALB/c mice immunized with RBD_{SARS}- or RBD_{MERS}-PLPs display serum RBD-specific IgG endpoint and live virus neutralization 33 34 titers that, in the case of SARS-CoV-2, were comparable to those detected in convalescent 35 plasma from infected patients. Further, these antibody levels remain elevated up to 6 months 36 post-prime. In dose response studies, immunization with as little as one microgram of RBD_{SARS}-37 PLPs elicited robust neutralizing antibody responses. Finally, animals immunized with RBD_{SARS}-38 PLPs, RBD_{MERS}-PLPs, and hCoV-RBD PLPs were protected against SARS-CoV-2 and/or MERS-39 CoV lung infection and disease. Collectively, these data suggest that the designer PLP system 40 provides a platform for facile and rapid generation of single and multi-target vaccines.

41 **INTRODUCTION**

42 Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), a positive-sense, 43 single-stranded RNA virus, was first isolated in late 2019 from patients with severe respiratory 44 illness in Wuhan, China (1, 2). This betacoronavirus is related to two other coronaviruses that are 45 highly pathogenic for humans – SARS-CoV and Middle East respiratory syndrome coronavirus 46 (MERS-CoV) (3). Although SARS-CoV no longer circulates in the human population, SARS-CoV-47 2 infection is the cause of the global pandemic of coronavirus disease 2019 (COVID-19), which 48 can progress to acute respiratory distress syndrome (ARDS) and death (4). The elderly, 49 immunocompromised, and those with certain co-morbidities (e.g., obesity, diabetes, and 50 hypertension) are at greatest risk of severe COVID-19 (5). In addition, MERS-CoV remains in 51 circulation and has infected over 2,500 people with lethal disease in ~834 (~34% case fatality 52 rate)(6). The virus has spread to 28 countries since emerging in the Kingdom of Saudi Arabia in 53 2012 (6, 7), and large outbreaks of human-to-human transmission have occurred in Riyadh and 54 Jeddah in 2014 and in South Korea in 2015 (8, 9).

The RNA genome of betacoronaviruses is approximately 30,000 nucleotides in length 55 56 (10). The 5' two-thirds encode nonstructural proteins that catalyze genome replication and viral 57 RNA synthesis, whereas the 3' one-third of the genome encodes the viral structural proteins, 58 including nucleoprotein and the spike (S), envelope, and membrane proteins. The S protein of 59 coronaviruses forms homotrimeric spikes on the virion (Fig 1A), which engages cell surface 60 attachment factors and entry receptors (11). Protein cleavage occurs sequentially during the entry 61 process to yield S1 and S2 fragments and is then followed by further processing of S2 to yield a 62 smaller S2' protein (12). The S1 fragment includes the receptor binding domain (RBD), the 63 predominant target of potently neutralizing monoclonal antibodies, while the S2 fragment 64 promotes membrane fusion (13-22).

65 A number of vaccine candidates against coronaviruses, targeting the S protein, have been 66 developed, including DNA plasmid, lipid nanoparticle encapsulated mRNA, inactivated virion, and

67 viral-vectored vaccines (23, 24). For instance, currently approved RNA-based vaccines that target the S protein of SARS-CoV-2 are safe and highly effective (25, 26); however, they suffer from 68 69 multiple-dose requirements and require strict "cold-chain" conditions that limit vaccine distribution 70 and pose constraints on the infrastructure and healthcare system. Conversely, adenovirus vector 71 vaccines against SARS-CoV-2 can have the advantage of single-dose administration (27), but 72 they are less effective than RNA vaccines and can lead to serious side effects, such as labored 73 breathing, transient neutropenia and lymphopenia, liver damage, and, in rare cases, neuropathies 74 like Bell's palsy and transverse myelitis (28-30). Furthermore, the success of platforms derived 75 from eukarvotic viruses (e.g., adenoviruses) has been tempered by safety concerns related to 76 pathogenicity, immunogenicity, and toxicity (31). Thus, there continues to be a need for 77 multifaceted, rapidly tunable vaccine platforms that can respond to novel emerging threats, 78 including natural variants, that provide long-term protection, and that are amenable to industrial 79 strategies to afford thermostable, single-shot formulations for efficient distribution in resource-80 limited settings.

81 Bacteriophage systems have been adapted as therapeutic and diagnostic (theranostic) 82 platforms. In these cases, phage-like particles (PLPs) derived from capsid proteins are utilized as 83 a scaffold for genetic and chemical modification (32, 33). For example, filamentous phages, such 84 as M13 and fd, and icosahedral phages, including P22, T4, AP205, MS2, and lambda, have been 85 used to display imaging reagents (34-36), synthetic polymers (37-39), bioactive peptides and 86 proteins (39-46). PLPs have also been employed as vaccine platforms to display antigens 87 including from Y. pestis (42) and P. falciparum (43). In these cases, the PLP decoration ligands are engineered using genetic and/or chemical modification of the capsid-associated proteins. 88

The Catalano laboratory has developed a "designer nanoparticle" platform adapted from phage lambda (47). Co-expression of the major capsid and "scaffolding" proteins in *Escherichia coli* affords icosahedral PLP shells that can be isolated in high yield. These can then be decorated *in vitro* with the lambda decoration protein (gpD) such that a single particle will display 140 trimeric

93 spikes of gpD (420 total copies) projecting from its surface (48, 49). The decoration protein can 94 be modified genetically to display heterologous peptides and proteins that can be displayed on 95 the PLP surface (39, 50, 51). Additionally, we have engineered a mutant gpD protein that contains 96 a Ser42->Cvs mutation (qpD(S42C)), which enables site-specific chemical modification of the 97 decoration protein using maleimide chemistry (Fig 1B) (39). Employing our in vitro system, PLPs 98 can be independently or simultaneously decorated with genetically and chemically modified 99 decoration proteins in rigorously defined surface densities (39, 46). Notably, the lambda PLP 100 platform has several advantages and unique features that can be harnessed for vaccine 101 development. For example, (i) PLPs can be decorated under defined *in vitro* conditions with 102 biological and synthetic molecules in varying surface densities using genetic and chemical 103 modification strategies; (ii) modification of the particle surface is fast and can be tuned in a user 104 defined manner, thereby streamlining the formulation process; (iii) intact phage and PLP shells 105 are natural adjuvants capable of stimulating the innate immune response (52, 53); (iv) the ability 106 to display antigens in high density can aid in regulating effector function (54); (v) decorated particles are monodisperse, stable, and possess physiochemical properties amenable for 107 108 pharmaceutical formulation and therapeutic applications (46). Therefore, the lambda system 109 described here allows for substantial flexibility and rigorous control in vaccine design and 110 development, setting the stage for the construction of an "all-in-one" vaccine platform.

111 In this study, we report the development of a monovalent lambda PLP-based vaccine 112 against SARS-CoV-2 (RBD_{SARS}-PLP) or MERS-CoV (RBD_{MERS}-PLP) via decoration with the spike 113 RBD proteins from either virus. Additionally, we engineered bivalent PLPs that co-display spike 114 RBD proteins from both viruses (hCoV-RBDs-PLP) to serve as a bivalent vaccine candidate. 115 Intramuscular administration of RBD_{SARS}-PLPs, RBD_{MERS}-PLPs, and hCoV-RBDs PLPs in mice 116 induce robust and durable humoral immune responses, including the production of neutralizing 117 antibodies. Moreover, immunization also protected mice against lung infection, inflammation, and pathology after virulent SARS-CoV-2 and MERS-CoV challenge. 118

119

120 RESULTS

121 Design and construction of particles decorated with human coronavirus spike RBD 122 proteins. The lambda designer PLP platform was adapted to display CoV spike RBD proteins 123 using methods established in the Catalano laboratory (39, 46). Purified recombinant spike RBD 124 proteins derived from the SARS-CoV-2 and MERS-CoV isolates Wuhan-Hu-1 and EMC/2012 are 125 referred to as RBD_{SARS} and RBD_{MERS}, respectively, or collectively as hCoV-RBDs. These proteins 126 were crosslinked to the lambda decoration protein mutant, gpD(S42C), following a two-step 127 procedure. First, solvent accessible lysine residues in the hCoV-RBDs were modified with the N-128 hydroxysuccinimide ester groups of the SM(PEG)₂₄ heterobifunctional crosslinker, as depicted in 129 Fig 1C: SDS-PAGE analysis reveals an apparent single predominant product for both RBD_{SARS} 130 and RBD_{MERS} (Fig 2A,D, respectively, lanes 4). Next, the maleimide groups of the crosslinker 131 were reacted with the sole cysteine residue of gpD(S42C) to afford the gpD-RBD_{SARS} and gpD-132 RBD_{MERS} constructs (Fig 1C): SDS-PAGE analysis reveals multiple products in each case (Fig 133 2A,D, lanes 5). The reaction mixtures were fractionated by size exclusion chromatography (SEC) (Fig 2B,E), and the fraction containing a single predominant product for each hCoV RBD was 134 135 isolated and then concentrated for use in all subsequent experiments (fraction 32, Fig 2C,F).

136 The purified gpD-RBD constructs were the used to decorate PLPs, alone and in 137 combination at defined surface densities, as outlined in Fig 1D. Complete particle decoration 138 requires 420 copies of the decoration protein that assemble as trimeric spikes, projecting from 139 the shell surface (see Fig 1B); therefore, we define the number of gpD-binding sites occupied as 140 a surface density percentage. For instance, 60% RBD_{SARS}-PLPs denotes particles that are each 141 decorated with 252 copies of gpD-RBD_{SARS} with the remaining sites filled with wildtype (WT) gpD 142 (168 copies). Similarly, 40% hCoV-RBD PLPs are particles that are each decorated with 168 143 copies of gpD-RBD_{SARS}, 168 copies of gpD-RBD_{MERS}, and 84 copies of WT gpD. Analysis of the

reaction mixtures by agarose gel electrophoresis (AGE) demonstrated that particles can be decorated in a defined manner with either and both gpD-RBD constructs (**Fig 3A-C**).

146 Decorated particles were purified by SEC (Fig 3D) and characterized by multiple 147 approaches. Electron microscopy (EM) confirmed that decorated PLPs retain icosahedral 148 symmetry and are well dispersed. Close inspection of the micrographs revealed that particles 149 decorated with RBD densities greater than 20% are characterized by surface projections whose 150 number increases with increasing surface decoration (Fig 3E); we attribute these projections to 151 RBD proteins extending from the particle surface. Physiochemical characterization of PLP 152 preparations by dynamic and electrophoretic light scattering analyses reveal that the mean 153 hydrodynamic diameter of particles decorated with RBDs also increases as a function of RBD 154 surface density and have an overall surface charge that ranges from -12 to -29 mV (Table 1). The 155 polydispersity index (PDI) of the decorated particles ranges from (0.156 ± 0.05) to (0.285 ± 0.09) . 156 In sum, these data indicate that these particle preparations have a high degree of homogeneity 157 and are acceptable by pharmaceutical standards.

Having confirmed that PLPs can be decorated with varying surface densities of hCoV RBDs, we sought to determine whether the purified gpD-RBD constructs retain their antigenic characteristics following particle decoration. Thus, the antigenicity of RBD_{SARS}-PLPs, RBD_{MERS}-PLPs, and hCoV-RBD-PLPs was assessed using enzyme immunoassays. The data presented in **Figs 3F** and **3G** show that PLPs decorated with 20% gpD-RBD_{SARS} and/or 20% gpD-RBD_{MERS} retain antigenic properties, whereas no reactivity was noted for WT PLPs.

Particles decorated with gpD-RBD_{SARS} are immunogenic following one or two immunizations. To assess the immunogenicity of RBD_{SARS} PLPs, 6-week-old BALB/c mice were immunized by intramuscular (i.m.) inoculation with 10 ug of WT PLPs (control) or 60% RBD_{SARS}-PLPs, followed by administration of a booster dose three weeks later (**Fig 4A**). Serum samples were collected 14 days after the primary or booster immunization (day 35 post-prime) and again at days 63 and 174 post-prime. IgG responses against purified RBD protein were evaluated by

170 ELISA (Fig 4B). After a single immunization, 60% RBD_{SARS}-PLPs induced RBD_{SARS}-specific IgG 171 titers comparable to those detected in convalescent samples from SARS-CoV-2 recovered 172 patients (Fig 4B,C). At 14 days post-boost (day 35), these responses were elevated 16-fold (P <173 0.001). Importantly, mice immunized with 60% RBD_{SARS}-PLPs maintained high levels of RBD_{SARS}-174 specific IgG out to day 174 post-prime (Fig 4B), indicating that these particles elicit durable 175 humoral immune responses. In contrast, RBD_{SARS}-specific IgG was undetectable in mice 176 immunized with WT PLPs at all time points evaluated (Fig 4B). Furthermore, IgG subclass 177 analysis at 14 days post-prime and boost revealed that 60% RBD_{SARS}-PLPs elicit high levels of 178 RBD_{SARS}-specific lgG1, lgG2a, and lgG2b (Fig 4D).

179 Additionally, we evaluated serum from immunized animals for the capacity to neutralize 180 SARS-CoV-2 infection using a live virus focus-reduction neutralization test (FRNT). Serum from 181 mice immunized with WT PLPs did not display neutralizing activity (Fig S1), whereas serum from 182 mice immunized with 60% RBD_{SARS}-PLPs 14 days post-prime neutralized SARS-CoV-2 infection 183 comparable to the neutralizing activity detected in convalescent samples from SARS-CoV-2 184 recovered patients (Fig 4E-F). Neutralizing activity was enhanced 4.7-fold following the booster 185 immunization (P < 0.05), and high levels of SARS-CoV-2 neutralizing antibodies were maintained 186 out to day 174 post-prime (Fig 4E and Fig S1).

Vaccination with RBD_{SARS}-PLPs protects from virulent SARS-CoV-2 challenge. We 187 188 evaluated the protective activity of particles decorated with gpD-RBD_{SARS} using a recently 189 developed mouse-adapted strain of SARS-CoV-2 (SARS-CoV-2 MA10), which productively 190 replicates in the mouse lung and results in clinical manifestations of disease consistent with 191 severe COVID-19 in humans (55). Six-week-old BALB/c mice were immunized i.m. with WT PLPs 192 or 60% RBD_{SARS}-PLPs. At 184 days post-prime, mice were challenged intranasally with 193 10⁴ plaque-forming units (PFUs) of SARS-CoV-2 MA10 (Fig 5). Mice immunized with WT PLPs 194 rapidly lost weight following SARS-CoV-2 infection, whereas no weight loss was observed in mice 195 immunized with 60% RBD_{SARS}-PLPs (Fig 5A). At 4 days post-infection (dpi), mice were

euthanized, and lungs were collected for viral burden analysis and histopathology. While high
levels of infectious virus were detected in the lungs of mice immunized with WT PLPs, infectious
virus was undetectable in the lungs of mice immunized with 60% RBD_{SARS}-PLPs, as determined
by plaque assay (Fig 5B). Moreover, greatly reduced levels of genomic (Fig 5C) and subgenomic
(Fig 5D) viral RNA were detected in the lungs of mice vaccinated with 60% RBD_{SARS}-PLPs, as
compared to those immunized with WT PLPs. Collectively, these data indicate that i.m.
immunization with RBD_{SARS}-PLPs elicits durable protection against SARS-CoV-2 lung infection.

The effect of immunization with RBD_{SARS}-PLPs on lung inflammation and disease also was assessed by analyzing lung tissue for histopathological changes. Mice immunized with WT PLPs and challenged with SARS-CoV-2 MA10 had an abundant accumulation of immune cells in perivascular and alveolar locations, vascular congestion, and interstitial edema (**Fig 5E**). Conversely, immunization with 60% RBD_{SARS}-PLPs resulted in a marked reduction of lungassociated histopathological changes (**Fig 5E**), indicative of protection against SARS-CoV-2 induced lung inflammation and injury.

210 Low doses of RBD_{SARS}-PLPs remain immunogenic and protect from virulent SARS-

211 CoV-2 challenge. We next evaluated the immunogenicity and protective efficacy of de-escalating 212 doses of particles decorated with gpD-RBD_{SARS}. Six-week-old BALB/c mice were immunized by 213 i.m. injection with 2.5, 1.0, or 0.25 µg of 60% RBD_{SARS}-PLPs and received a booster dose of the 214 same amount three weeks later. Control mice received two i.m. injections of 2.5 µg of WT PLPs, 215 following the same immunization schedule. Serum samples were collected on days 14, 35 (14 216 days post-boost), 63, and 84 post-prime, and IgG responses against purified RBD protein were 217 evaluated by ELISA. At 14 days, little response was observed in mice immunized with a single 218 injection of 0.25 µg of 60% RBD_{SARS}-PLPs, and no response was detected in control mice (Fig 219 6A). In contrast, mice immunized with 2.5 or 1.0 µg of 60% RBD_{SARS}-PLPs had detectable levels 220 of RBD_{SARS}-specific IgG which were comparable to mice immunized with a 10 µg dose (compare 221 Figs 4B and 6A), and to RBD_{SARS}-specific IgG detected in convalescent samples from SARS- CoV-2 recovered patients (Fig 4C). At 14 days post-boost (day 35 post-prime), RBD_{SARS}-specific
IgG responses were elevated in mice that received all doses of 60% RBD_{SARS}-PLPs, although
lower responses were observed for mice immunized with 0.25 µg (Fig 6A). Additionally, mice
immunized with all doses maintained high levels of RBD_{SARS}-specific IgG up to 84 days post-prime
(Fig 6A).

227 We also evaluated serum from this dose de-escalation study for the capacity to neutralize 228 SARS-CoV-2 infection (Fig 6B and Fig S2). Serum from mice immunized with WT PLPs did not 229 display neutralizing activity (Fig S2). While day 14 post-prime serum from mice immunized with 230 all doses of 60% RBD_{SARS}-PLPs displayed little to no neutralizing activity, day 35 (14 days post-231 boost) serum from mice immunized with 2.5 or 1.0 µg of 60% RBD_{SARS}-PLPs displayed potent 232 neutralizing activity that was maintained up to 84 days post-prime (Fig 6B). Despite detectable 233 levels of RBD_{SARS}-specific IgG, immunization with 0.25 µg of 60% RBD_{SARS} PLPs did not result in 234 neutralizing activity at any of the time points evaluated (Fig 6B).

235 Mice immunized with de-escalating doses of 60% RBD_{SARS} PLPs were challenged with 236 10⁴ PFU of SARS-CoV-2 MA10 at day 90 post-prime. Compared to control mice, mice immunized 237 with all doses (including 0.25 µg) of 60% RBD_{SARS} PLPs were protected from weight loss 238 associated with SARS-CoV-2 infection (Fig 6C). At 4 dpi, lungs were collected and assessed for viral burden. Immunization with 2.5 or 1.0 µg of 60% RBD_{SARS}-PLPs resulted in potent protection 239 240 against viral infection (Fig 6D) and reduced levels of viral genomic and subgenomic RNA (Fig 241 6E,F) in the lungs. Furthermore, histopathological analysis of lung tissues of mice immunized at 242 these doses showed minimal perivascular and alveolar infiltrates at 4 dpi, whereas those of 243 control mice were characterized by extensive inflammation (Fig 6G). Mice immunized with 0.25 244 µg of 60% RBD_{SARS} PLPs had levels of infectious virus and viral RNA in the lungs similar to those 245 in control mice (Fig 6D-F), consistent with an inability to neutralize SARS-CoV-2 infection (Fig 246 6B). Additionally, these mice displayed more severe lung inflammation and injury versus those 247 immunized with higher doses. Overall, the data indicate that neutralizing antibodies are essential for protection against SARS-CoV-2 infection and suggest that additional adaptive immune response elicited by vaccination with RBD_{SARS}-PLPs may contribute to protection against severe disease characterized by more extensive weight loss, such as the generation of memory T cells.

251 Vaccination with mosaic hCoV-PLPs protects from virulent SARS-CoV-2 and MERS-252 **CoV challenge.** As described above, PLPs can be decorated with multiple hCoV RBDs at varying 253 surface densities to generate mosaic PLPs. Thus, we evaluated the immunogenicity and 254 protective efficacy of particles simultaneously decorated with gpD-RBD_{SARS} and gpD-RBD_{MERS} in 255 comparison with their mono-RBD decorated counterparts (Fig 7). Six-week-old BALB/c mice were 256 immunized by i.m. injection with 7.5 µg of WT PLPs, 20% RBD_{SARS}-PLPs, 20% RBD_{MERS}-PLPs, 257 or 20% hCoV-RBD-PLPs (note, 20% of each of gpD-RBD_{SARS} and gpD-RBD_{MERS}). Mice received 258 a booster dose of the same amount three weeks later. Serum samples were collected on days 259 14, 35 (14 days post-boost), and 96 post-prime, and IgG responses against purified SARS-CoV-260 2 and MERS-CoV RBD protein were evaluated by ELISA. At each time point evaluated, mice 261 immunized with the bi-valent 20% hCoV-RBD-PLPs had similar levels of RBD_{SARS}- and RBD_{MERS}-262 specific IgG as mice immunized with the mono-valent 20% RBD_{SARS}-PLPs or 20% RBD_{MERS}-PLPs (Fig 7A, 7B). In addition, immunization with 20% hCoV-RBD-PLPs induced potent neutralizing 263 264 antibody responses against both SARS-CoV-2 (Fig 7C) and MERS-CoV (Fig 7D). We note, 265 however, that neutralizing titers against MERS-CoV were somewhat lower in mice immunized 266 with the bi-valent vaccine compare with the mono-valent MERS vaccine (Fig 7D).

267 Next, these mice were challenged with 10⁴ PFU of SARS-CoV-2 MA10 or 10⁵ PFU MERS-268 CoV at day 102 post-prime. Five days prior to MERS-CoV challenge, mice were i.n. inoculated 269 with Ad-hDPP4 to deliver the MERS-CoV cell entry receptor to lung cells (*56, 57*). Compared with 270 control mice, mice immunized with either RBD_{SARS}-PLPs or hCoV-PLPs were protected from 271 weight loss (**Fig 7E**) and lung viral burden (**Fig 7F**) associated with SARS-CoV-2 infection. 272 Similarly, mice immunized with either RBD_{MERS}-PLPs or hCoV-PLPs were protected from weight 273 loss (**Fig 7G**) and lung viral burden (**Fig 7H**) associated with MERS-CoV infection.

274

275 **DISCUSSION**

276 Numerous vaccine strategies and platforms have been developed for current and 277 emerging infectious diseases, and several studies support the use of CoV RBDs as vaccine 278 immunogens. For instance, the SARS-CoV-2 RBD is immunodominant, contains multiple 279 antigenic sites, and is the overwhelming target of neutralizing activity in COVID-19 convalescent 280 plasma (58). In addition, passive administration of monoclonal antibodies that target the RBD of 281 SARS-CoV-2 and other pathogenic CoVs protects rodents, ferrets, and non-human primates from 282 severe CoV infection (14, 59-68), indicating that the generation of RBD-specific antibodies is an 283 advantageous therapeutic strategy. Moreover, the RBD is a target of cross-neutralizing antibodies 284 (58, 69), and immunization with SARS-CoV-2 RBD functionalized nanoparticles can elicit cross-285 neutralizing antibodies against a number of SARS-related CoVs (70). Nevertheless and despite 286 significant advancements in diagnostic and therapeutic countermeasures, continued vaccine 287 development efforts are warranted, especially for viruses that pose a threat to global public health or to provide broad protection against multiple viruses. 288

289 In this study, we designed and constructed lambda PLPs decorated with the RBDs from 290 extant highly pathogenic human CoVs (i.e., SARS-CoV-2, MERS-CoV) as a tunable, multivalent 291 vaccine platform. The platform allows for symmetrical, high-density display of multiple target 292 antigens, thereby enhancing the stimulation of host immune responses. Properties of decorated 293 PLPs present several advantages for use as an effective vaccine delivery vehicle, including (i) a 294 particle size less than 200 nm, enabling entry into the lymphatic system (71) where adaptive 295 immune responses are initiated; (ii) additional physiochemical properties such as surface charge, 296 exposure of functional group, and hydrophobicity are easily manipulated by both genetic and 297 chemical approaches; and (iii) simultaneous delivery of foreign molecules can be achieved on a 298 single PLP. Further, we found no adverse effects due to immunization with phage lambda 299 particles by intramuscular routes in mice even after repeated doses or after challenge (i.e., no

evidence of vaccine-enhanced disease), indicating a favorable safety profile for future *in vivo*testing.

302 We demonstrate that mono and mosaic hCoV-RBD decorated particles induce RBD-303 specific binding and virus neutralizing antibodies after one and two doses. Notably, these 304 responses remained highly elevated up to 6 months post-prime, indicating that the vaccine 305 platform can stimulate robust and durable immune responses. Additionally, immunized mice were 306 protected against virulent virus challenge (up to day 184 post-prime for SARS-CoV-2 and day 102 307 for MERS-CoV) as indicated by significantly reduced virus-induced weight loss, a marked 308 reduction in lung-associated inflammation and injury, and lower titers of infectious virus in the 309 lungs paired with diminished levels of viral genomic and subgenomic RNA. Consistent with 310 published studies (70). PLPs decorated with the SARS-CoV-2 RBD alone do not elicit IgG that 311 binds the MERS-CoV RBD or antibodies that neutralize MERS-CoV infection. In contrast, those 312 simultaneously decorated with the SARS-CoV-2 and MERS-CoV RBD proteins induce strong 313 immune responses against both pathogenic viruses. This supports that co-displaying multiple 314 antigens on the same particle is an effective means of broadening immune responses elicited by 315 our vaccine platform, and that the presence of multiple antigens do not interfere with each other.

316 A number of nanoparticles and antigen coupling strategies have been evaluated for vaccine design (72). These include nanoparticles based on the SpyCatcher-SpyTag system which 317 318 can be used to conjugate antigens to self-assembling protein nanoparticles (73-75), the Novavax 319 vaccine (76), and other phage systems (77). Most often, these immunization studies involve the 320 use of large amounts of antigen (e.g., 50-100 µg/dose), up to 4 immunizations, formulation with 321 an adjuvant (e.g., AddaVax), and vaccine administration routes not amenable to humans (e.g., 322 intraperitoneal) (75, 78). Our dose response studies show that immunization with as little as one 323 microgram of 60% RBD_{SARS}-PLPs elicited robust neutralizing antibody responses and conferred 324 protection against SARS-CoV-2 challenge. These data suggest that PLP-based vaccines 325 facilitate dose sparing immunization practices which, from both manufacturing and distribution perspectives, enhances their utility as a vaccine platform. Further studies will be necessary to test
 protective efficacy following immunization in a dose-dependent manner or through alternative
 routes.

While we did not define the epitopes targeted, the anti-SARS-CoV-2 monoclonal antibody 329 330 used in the enzyme immunoassays binds to a conserved epitope in the RBD that is only 331 accessible when the S protein is in conformation competent to bind the hACE2 receptor (79, 80). 332 Additionally, RBD-specific IgG subclass responses were assessed; however, further 333 characterization of the potential role of these in vaccine-mediated protection was not determined. 334 Considering the implications these may have to the development of safe vaccines (e.g., clinical 335 syndromes associated with vaccine-enhanced disease) (81), additional studies are needed to 336 examine the possible impacts to cellular effector function and evaluate mucosal and cellular 337 immunity in addition to immunity against variants of SARS-CoV-2 and MERS-CoV.

In summary, the versatility and robustness of the lambda system presented provides a promising vaccine candidate. Immunization with hCoV-RBD decorated PLPs induces antibodies with neutralizing activity, results in a durable immune response, and provides effective protection against virulent virus challenge. Continued efforts into the optimization of this platform could expand the diversity of protein-based vaccine technologies available and aid in the prevention of infectious diseases deleterious to global health.

344

345 METHODS

Purification of lambda PLPs. Lambda PLPs were expressed in *E. coli*BL21(DE3)[pNu3_E] cells and purified, as described (46). Briefly, self-assembled PLPs were
extracted from cell lysates in 20 mM Tris [pH 8.0, 4°C] buffer containing 0.4 mg/mL lysozyme and
0.04 mg/mL DNase. Following rate zonal centrifugation (10–40% sucrose density gradient), PLPs
were collected, concentrated, and exchanged into 20 mM Tris [pH 8.0, 4°C] buffer containing 15
mM MgCl₂, 1 mM EDTA, and 7 mM β-ME using centrifugal filter units (100k MWCO; Millipore

Amicon Ultra). Proteins were fractionated by anion exchange chromatography employing three 5 mL HiTrap Q HP columns connected in tandem and developed with a 30-column volume linear gradient to 1 M NaCl. The eluate was analyzed by SDS-PAGE, and PLP-containing fractions were pooled and exchanged into 50 mM HEPES [pH 7.4] buffer containing 100 mM NaCl and 10 mM MgCl₂ prior to storage at 4 °C.

357 **Purification of lambda decoration proteins.** WT and gpD(S42C) decoration proteins 358 were expressed in E. coli BL21(DE3)[pD] and BL21(DE3)[pDS542C] cells, respectively, and 359 purified, as described (46). Briefly, cell lysate supernatants were dialyzed overnight against 20 360 mM Tris [pH8.0, 4 °C] buffer containing 20 mM NaCl and 0.1 mM EDTA. Proteins were 361 fractionated employing three 5 mL HiTrap Q HP columns connected in tandem and developed 362 with a 30-column volume linear gradient to 1 M NaCl. Fractions containing decoration protein 363 were pooled, exchanged into 50 mM NaOAc [pH 4.8] buffer using centrifugal filter units (3k 364 MWCO; Millipore Amicon Ultra), and loaded onto three 5 mL HiTrap SP columns connected in 365 tandem. Bound proteins were eluted with a 30-column volume linear gradient to 0.5 M NaCl, and fractions containing decoration protein were pooled and dialyzed overnight against 20 mM Tris 366 367 [pH 8.0, 4 °C] buffer containing 20 mM NaCl and 0.1 mM EDTA for storage at 4°C. For long-term 368 storage at -80°C, aliquots of purified protein were supplemented with 20% glycerol.

369 Purification of recombinant hCoV RBD proteins. A pCAGGS expression vector 370 encoding the SARS-CoV-2 spike RBD was obtained from Dr. Florian Krammer, Icahn School of 371 Medicine at Mount Sinai (82). This construct contains sequences encoding the native signal 372 peptide (residues 1–14) followed by residues 319–541 from the SARS-CoV-2 Wuhan-Hu-1 spike 373 protein (GenBank MN908947.3) and a hexa-histidine tag at the C-terminus for purification. A 374 pTwist-CMV expression vector (Twist Biosciences Technology) encoding the MERS-CoV spike 375 RBD was kindly provided by Dr. Peter S. Kim, Stanford University (83). This construct contains 376 sequences encoding an N-terminal human IL-2 signal peptide following by residues 367-588 from 377 the MERS-CoV EMC/2012 spike RBD (GenBank JX86905.2) and dual C-terminal tags (octa378 histidine. AviTag) for purification. Recombinant protein expression and purification was performed 379 by the University of Colorado Cancer Center Cell Technologies Shared Resource. Due to the 380 involvement of glycans in the epitopes of some neutralizing antibodies (62, 84), RBDs were 381 expressed in Expi293F cells to retain normal glycosylation and antigenic properties. Expi293F 382 cells were transfected with plasmid DNA using ExpiFectamine transfection reagent 383 (ThermoFisher Scientific). At 72 h post-transfection, cell culture supernatants were centrifuged 384 (4,000 x g) for 20 min and filtered (0.2 micron). Recombinant proteins were purified by Ni-NTA 385 column chromatography using an ATKA purification system. Protein was eluted with 500 mM 386 imidazole and concentrated before assessing protein purity by SDS-PAGE and Coomassie 387 staining.

388 Crosslinking of lambda decoration protein (gpD(S42C)) to hCoV RBDs. SARS-CoV-389 2 or MERS-CoV RBD protein was exchanged into 0.01 M PBS [pH 7.2] buffer using centrifugal 390 filter units (10k MWCO; Millipore Amicon Ultra). A 5-fold molar excess of SM(PEG)₂₄ (PEGylated, 391 long-chain sulfosuccinimidyl 4-(N-maleimidomethyl)cyclohexane-1-carboxylate crosslinker; 392 ThermoFisher Scientific) was added, and the mixture was incubated (30 min, 25°C) for 393 modification of solvent accessible primary amines. Reduced gpD(S42C) was added to the mixture 394 (1:1, molar equivalent) and incubated (1 h, 25°C) to yield the crosslinked products (gpD-RBD_{SARS}, 395 gpD-RBD_{MERS}). The reaction was guenched with the addition of 0.2% β -ME (30 min, 25°C), and 396 crosslinked proteins were purified by size-exclusion chromatography employing a 24 mL 397 Superose 6 Increase 10/300 GL column equilibrated and developed with 40 mM HEPES [pH 7.4] 398 buffer containing 150 mM NaCl, 200 mM arginine, 0.1 mM EDTA, and 2 mM β-ME. Fractions 399 containing cross-linked proteins were identified by SDS-PAGE and were pooled. The purified 400 proteins were and exchanged into 20 mM Tris [pH 8.0, 4°C] buffer containing 20 mM NaCl and 401 0.1 mM EDTA for storage at 4°C. Protein concentration was quantified spectroscopically.

402 *In vitro* PLP expansion and decoration. PLPs were expanded and subsequently 403 decorated *in vitro*, as described (46). Briefly, purified PLPs were expanded in 10 mM HEPES [pH

404 7.4] buffer containing 2.5 M urea for 30 min on ice and then exchanged into 10 mM HEPES [pH 405 7.4] buffer containing 200 mM urea using centrifugal filter units (100k MWCO; Millipore Amicon 406 Ultra). Expanded shells (30 nM) were decorated in a stepwise fashion with modified and WT 407 decoration protein at 25°C in 10 mM HEPES [pH 7.4] buffer containing 50 mM urea, 10 mM 408 arginine, and 0.1% Tween 20. Proteins were added in the following order: (1) gpD-RBD_{SARS} (0.69-409 8.33 µM final concentration, 30 min incubation); (2) gpD-RBD_{MERS} (1.39-8.33 µM, 30 min 410 incubation); (3) wildtype gpD (5.55-13.19 µM, 60 min incubation). Decorated PLPs were purified 411 by size-exclusion chromatography employing a 24 mL Superose 6 Increase 10/300 GL column 412 equilibrated and developed with 100 mM HEPES [pH 6.6] buffer containing 200 mM NaCl, 200 413 mM arginine, 1 mM EDTA, and 2 mM β -ME at a flow rate of 0.3 mL/min. Fractions containing 414 decorated PLPs were pooled and exchanged into 50 mM HEPES [pH 7.4] buffer containing 100 415 mM NaCl and 10 mM MgCl₂ for storage at 4°C.

416 Transmission Electron Microscopy. Carbon-coated copper grids (400 mesh; CF400-417 CU) were glow-discharged using a Pelco EasiGlo system (Ted Pella, Inc.; Redding, CA, USA) 418 with a plasma current of 15 mA, negative glow discharge head polarity, glow discharge duration 419 of one minute, and held under vacuum for 15 seconds. PLP preparations were diluted to 10 nM 420 using water (sterile, double distilled) and spotted twice onto grids. Following sample adsorption 421 (20 s), excess liquid was wicked off using a Whatman #1 filter paper. Grids were washed using 422 water, and excess liquid was wicked off. Samples were negatively stained (20-25 s) with filtered 423 2% (w/v) methylamine tungstate [pH 6.7] and 50 µg/mL bacitracin (0.2 µm Nucleopore 424 polycarbonate syringe filters). Excess stain was wicked off, and grids were allowed to air-dry for 425 at least one hour before transferring to a grid storage box. Samples were maintained covered 426 throughout this procedure and stored at room temperature until imaged. Images were acquired 427 on a FEI Tecnai G2 transmission electron microscope at an accelerating voltage of 80 kV and 428 equipped with a 2k × 2k CCD camera. Images were processed in Fiji (85) and measurements 429 were based on 100 particles with values reported as mean \pm standard deviation.

430 Particle size and charge measurements. Decorated particles were diluted to 2 nM (55-431 700 µL) using 10 mM HEPES [pH 7.4] for dynamic light scattering (DLS) and electrophoretic light 432 scattering (ELS) analyses. Particle size and charge measurements were acquired on a Malvern 433 Panalytical Zetasizer Nano ZS (He-Ne laser 633 nm light source; 5 mW maximum power). Size 434 information obtained from the correlation function was based on hydrodynamic diameter 435 (weighted mean reported as Z-average, Z-Ave) and particle size distributions (intensity, volume 436 PSD) with polydispersity index (PDI) values provided to serve as an estimate of the width of the 437 distributions. Overall surface charge was reported as zeta potential [mV]. Successive sample 438 measurements were done in quadruplicate with values reported as mean ± standard deviation.

439 Virus and cells. Vero E6 (CRL-1586, American Type Culture Collection (ATCC)) were 440 cultured at 37°C in Dulbecco's Modified Eagle medium (DMEM, HyClone 11965-084) 441 supplemented with 10% fetal bovine serum (FBS), 10 mM HEPES [pH 7.3], 1 mM sodium 442 pyruvate, 1X non-essential amino acids, and 100 U/ml of penicillin-streptomycin. SARS-CoV-2 443 strain 2019 n-CoV/USA WA1/2020 was obtained from BEI Resources. The virus was passaged once in Vero E6 cells and titrated by focus formation assay (FFA) on Vero E6 cells. The mouse-444 445 adapted SARS-CoV-2 strain MA10 (kindly provided by R. Baric, University of North Carolina at 446 Chapel Hill) elicits disease signs in laboratory mice similar to severe COVID-19 (55) was used for 447 all challenge studies. All work with infectious SARS-CoV-2 and MERS-CoV was performed in 448 Institutional Biosafety Committee approved BSL3 and A-BSL3 facilities at the University of 449 Colorado School of Medicine and University of Maryland School of Medicine using positive 450 pressure air respirators and other personal protective equipment.

451 **Mouse experiments.** Animal studies were carried out in accordance with the 452 recommendations in the Guide for the Care and Use of Laboratory Animals of the National 453 Institutes of Health. The protocols were approved by the Institutional Animal Care and Use 454 Committees at the University of Colorado School of Medicine (Assurance Number A3269-01) and 455 the University of Maryland School of Medicine (Assurance number D16-00125 (A3200-01)).

456 Female BALB/c mice were purchased from The Jackson Laboratory. 6-8-week-old mice were 457 immunized with WT PLPs (control), RBD_{SARS}-PLPs, RBD_{MERS}-PLPs, or hCoV-RBD PLPs 458 preparations in 50 µl PBS via i.m. injection in the hind leg. Mice were boosted three weeks after 459 the primary immunization using the same route. For virulent virus challenge, mice were 460 anaesthetized by intraperitoneal injection with 50 µL of a mix of xylazine (0.38 mg/mouse) and 461 ketamine hydrochloride (1.3 mg/mouse) diluted in PBS. Immunized BALB/c mice were inoculated intranasally (i.n.) with 10⁴ PFU of SARS-CoV-2 MA10. For MERS-CoV challenge, immunized 462 mice were first inoculated i.n. with 2.5 x 10⁸ PFU of a replication incompetent adenovirus vector 463 464 to deliver human DPP4 (Ad5-hDPP4) into the lungs of mice (57). Five days later, mice were 465 inoculated i.n. with 10⁵ PFU of MERS-CoV (Jordan strain). All mice were weighed and monitored 466 for additional disease signs daily. At 4 days post-inoculation (dpi), animals were euthanized by 467 bilateral thoracotomy and tissues were collected for virological, immunological, and pathological 468 analyses.

ELISA. SARS-CoV-2 and MERS-CoV RBD-specific antibody responses in mouse and 469 470 human sera were measured by ELISA. Immunol 4HBX plates were coated with 0.2 µg of 471 recombinant SARS-CoV-2 RBD protein (Wuhan-Hu-1, GenPept QHD43416) or MERS-CoV RBD protein overnight at 4°C. Coating antigen was removed, and plates were blocked with 200 µl of 472 473 3% non-fat milk in PBS/0.1% Tween-20 (PBS-T) for one hour at room temperature. After blocking, 474 wells were washed once with PBS-T and then probed with serum samples diluted in PBS-T 475 supplemented with 1% non-fat milk for 1.5 h. Samples were removed, and wells were washed three times with PBS-T and probed with secondary antibodies diluted at 1:4000 in PBS-T; goat 476 anti-mouse IgG-HRP (Southern Biotech, 1030-05), goat anti-human IgG Fc-HRP (Southern 477 478 Biotech, 2014-05), goat anti-mouse IgG1-BIOT (Southern Biotech, 1071-08), goat anti-mouse IgG_{2a} Human ads-BIOT (Southern Biotech, 1080-08), goat anti-mouse IgG_{2b}-BIOT (Southern 479 480 Biotech, 1091-08), goat anti-mouse IgG_3 human ads-BIOT (Southern Biotech, 1100-08). 481 Biotinylated antibodies were subsequently probed with streptavidin-HRP (Southern Biotech,

482 7100-05). Detection antibody was removed, and wells were washed three times with PBS-T. 483 Plates were developed using 3,3',5,5'-tetramethylbenzidine (TMB; Sigma, T0440), followed by 484 the addition of 0.3 M H₂SO₄. Plate absorbance was read at 450 nm on a Tecan Infinite M plex 485 plate reader. Endpoint titers are reported as the reciprocal of the final dilution, in which 486 absorbance at 450 nm was above the blank average plus three times the standard deviation of 487 all blanks in the assay. The immunoreactivity of RBD_{SARS}-PLPs, RBD_{MERS}-PLPs, and hCoV-RBD PLPs was measured in a similar manner. In this case, plates were coated with 0.2 µg of WT PLPs 488 or RBD_{SARS}-PLPs, RBD_{MERS}-PLPs, and hCoV-RBD PLPs overnight at 4°C. Following blocking and 489 490 washing, wells were incubated for 1.5 h at room temperature with either chimeric human anti-491 SARS-CoV spike antibody clone CR3022 (Absolute Antibody, Ab01680) or mouse anti-MERS-492 CoV spike antibody clone D12 (Absolute Antibody, Ab00696) prepared in a 2-fold dilution series 493 in PBS-T with a starting dilution of 1:200 and signal was developed as described above.

494 SARS-CoV-2 focus reduction neutralization test (FRNT). Vero E6 cells were seeded in 96-well plates at 10⁴ cells/well. The next day serum samples were heat-inactivated at 56°C for 495 496 30 minutes and then serially diluted (2-fold, starting at 1:10) in DMEM (HyClone, 11965-084) plus 497 1% FBS in 96-well plates. Approximately 100 focus-forming units (FFU) of SARS-CoV-2 USA-498 WA1/2020 were added to each well and the serum plus virus mixture was incubated for 1 h at 499 37°C. Following co-incubation, medium was removed from cells and the serum sample plus virus 500 mixture was added to the cells for 1 h 37°C. Samples were then removed and cells overlaid with 501 1% methylcellulose (MilliporeSigma, M0512) in minimum essential media (MEM) (Gibco, 12000-502 063) plus 2% FBS and incubated for 24 hours at 37°C. Cells were fixed with 1% paraformaldehyde 503 (PFA; Acros Organics, 416780030) and probed with 1 µg/mL of chimeric human anti-SARS-CoV spike antibody (CR3022, Absolute Antibody, Ab01680) in Perm Wash (1X PBS/0.1% 504 saponin/0.1% BSA) for 2 h at room temperature. After three washes with PBS-T, cells were 505 506 incubated with goat anti-human IgG Fc-HRP (Southern Biotech, 2014-05) diluted at 1:1000 in

Perm Wash for 1.5 h at room temperature. SARS-CoV-2-positive foci were visualized with TrueBlue substrate (SeraCare, 5510-0030) and counted using a CTL Biospot analyzer and Biospot software (Cellular Technology Ltd, Shaker Heights, OH). The FRNT₅₀ titers were calculated relative to a virus only control (no serum) set at 100%, using GraphPad Prism 9.1.2 (La Jolla, CA) default nonlinear curve fit constrained between 0 and 100%.

512 **MERS-CoV neutralization assay.** To determine the inhibitory activity of mouse sera 513 against MERS-CoV, 3,950 TCID₅₀/ml of MERS-COV-Jordan was incubated with diluted sera for 514 30 minutes at room temperature and the inhibitory capacity of each serum dilution was assessed 515 by TCID₅₀ assay as previously described (*56*, *86*).

516 Quantification of SARS-CoV-2 genomic and subgenomic RNA. To quantify viral 517 genomic and subgenomic RNA, lung tissue of mice challenged with SARS-CoV-2 was 518 homogenized in TRIzol reagent (Life Technologies, 15596018) and total RNA was isolated using 519 a PureLink RNA Mini kit (Life Technologies, 12183025). Single-stranded cDNA was generated 520 using random primers and SuperScript IV reverse transcriptase (Life Technologies, 18091050). SARS-CoV-2 genomic or subgenomic RNA copies were measured by qPCR using the primer and 521 522 probe combinations listed in **Table S1** (Integrated DNA Technologies). To quantify SARS-CoV-2 523 genomic RNA, we extrapolated viral RNA levels from a standard curve generated from known 524 FFU of SARS-CoV-2 from which RNA was isolated and cDNA generated as previously described 525 (87). To quantify SARS-CoV-2 subgenomic RNA, we extrapolated viral RNA levels from a 526 standard curve using defined concentrations of a plasmid containing an amplified SARS-CoV-2 527 subgenomic fragment (pCR-sgN TOPO). Briefly, a subgenomic RNA fragment was amplified by 528 RT-PCR using RNA isolated from SARS-CoV-2-infected Vero E6 cells. The 125 bp amplicon is 529 composed of the joining region between the 3' UTR and the N gene, with inclusion of a 530 transcription-regulatory sequence (TRS) of the SARS-CoV-2 Wuhan-Hu-1 (NC 045512.2). This 531 amplicon was cloned into the pCR4 Blunt TOPO vector (Invitrogen, K2875J10), sequence confirmed, and used in a dilution series of defined gene copies. All qPCR reactions were prepared 532

with Taqman Universal MasterMix II (Applied Biosystems, 4440038) and were analyzed with
Applied Biosystems QuantStudio ViiA 7 analyzer.

535 Plague assay. Vero E6 cells were seeded in 12-well plates one day prior to virus 536 inoculation. Lung homogenates were serially diluted in DMEM supplemented with 2% FBS, 537 HEPES, and penicillin-streptomycin and incubated on cells for one hour at 37°C. Afterwards, cells 538 were overlaid with 1% (w/v) methylcellulose in MEM plus 2% FBS and incubated at 37°C. At 3 539 dpi, overlays were removed, and plates were fixed with 4% PFA for 20 minutes at room 540 temperature. After removal of PFA, plates were stained with 0.05% (w/v) crystal violet in 20% 541 methanol (10-20 min). Crystal violet was removed, and plates were rinsed with water or PBS. 542 Plagues were counted manually to determine infectious virus titer.

Histopathology. Mice were euthanized at 4 days following SARS-CoV-2 MA10 or MERSCoV challenge. The lungs were removed and fixed with 10% formalin. 5-micron sections were
stained with H&E for histological examination. Slides were examined in a blinded fashion for total
inflammation, periarteriolar, and peribronchiolar inflammation and epithelial cell denuding.

547 **Statistical Analysis.** Statistical significance was assigned when *P values* were < 0.05 548 using Prism Version 9.1.2 (GraphPad). Tests, number of animals (n), median values, and 549 statistical comparison groups are indicated in the Figure Legends.

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558 AUTHOR CONTRIBUTIONS

- 559 B.J.D., A.C., M.B.F., C.E.C., and T.E.M. designed the experiments. B.J.D., A.C., S.W.,
- and R.J. performed the experiments. B.J.D., A.C., S.W., R.J., M.B.F., C.E.C., and T.E.M.
- 561 performed data analysis. B.J.D., A.C., C.E.C., and T.E.M. wrote the initial draft of the manuscript.
- 562 All authors provided comments and edits to the final version.
- 563

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758 FIGURE LEGENDS

759 Figure 1. Design and construction of hCoV RBD decorated particles. (A) Atomistic 760 model of the SARS-CoV-2 virion with homotrimeric spikes of the S protein shown in shades of 761 pink (Nanographics GmbH). (B) Cryo-EM reconstruction of the phage lambda capsid with only 762 the density from the trimeric spikes of the decoration protein, gpD, displayed (Lander et al., 2008). 763 Shown to the right in cartoon representation is the crystal structure of an isolated gpD trimer (PDB: 764 1C5E) modified to contain the serine to cysteine mutation (red spheres) of the gpD variant. 765 gpD(S42C). (C) Reaction schematic for the chemical crosslinking of human coronavirus RBD 766 proteins (hCOV RBDs) to qpD(S42C) via lysine amide coupling and thiol-maleimide chemistry. 767 The products (gpD-RBD_{SARS}, gpD-RBD_{MERS}) contain a PEG crosslinker, SM(PEG)₂₄, with a spacer arm length of 95.2 Å. (D) PLPs are decorated with RBDs at the desired surface densities either 768 769 independently (top) or in combination (bottom) with remaining gpD-binding sites filled using wildtype gpD such that all 420 binding sites are occupied. 770

771 Figure 2. Construction of hCoV RBDs. (A) Proteins at various steps in the chemical 772 modification of gpD(S42C) for conjugation to RBD_{SARS} were fractionated by SDS-PAGE and visualized by Coomassie Blue staining. Lanes: (1) gpD(S42C); (2) RBD_{SARS}; (3) gpD(S42C) and 773 774 RBD_{SARS} in the absence of crosslinker; (4) RBD_{SARS} modified with the crosslinker; (5) gpD-775 RBD_{SARS} crosslinked proteins (red arrows). (B) SEC chromatogram of the gpD-RBD_{SARS} reaction 776 mixture (blue); collected fractions numbers are indicated at the top. SEC chromatograms of 777 unmodified RBD_{SARS} (red) and unmodified gpD(S42C) (purple) are shown for comparison. (C) 778 SDS-PAGE of fractions eluting from the SEC column in panel B. The migration of purified gpD-779 RBD_{SARS}, unmodified RBD_{SARS}, and gpD(S42C) are denoted as D-R_s, R_s, and D, respectively. 780 Unless otherwise stated, gpD-RBD_{SARS} fraction 32 (red star) was used for all subsequent 781 experiments. (D) Proteins at various steps in the chemical modification of gpD(S42C) for 782 conjugation to RBD_{MERS} were fractionated by SDS-PAGE and visualized by Coomassie Blue 783 staining. Lanes: (1) gpD(S42C); (2) RBD_{MERS}; (3) gpD(S42C) and RBD_{MERS} in the absence of 784 crosslinker; (4) RBD_{MERS} modified with the crosslinker; (5) gpD-RBD_{MERS} crosslinked proteins (red 785 arrows). (E) SEC chromatogram of the gpD-RBD_{MERS} reaction mixture (yellow); collected fractions 786 are indicated at the top. (F) SDS-PAGE of fractions eluting from the SEC column in panel E. The 787 migration of purified gpD-RBD_{MERS}, unmodified RBD_{MERS}, and gpD(S42C) are denoted as D-R_M, R_M, and D, respectively. Unless otherwise stated, gpD-RBD_{MERS} fraction 32 (red star) was used 788 789 for all subsequent experiments.

790 Figure 3. PLP decoration with hCoV RBDs. PLPs were decorated with (A) gpD-RBD_{SARS} 791 (D-R_s, blue), (**B**) gpD-RBD_{MERs} (D-R_M, yellow) or (**C**) with both constructs at the indicated surface 792 densities as outline in Fig 1D and the reaction mixtures were fractionated by AGE. In all gels, 793 naked PLPs (no decoration protein, lanes 1) and WT PLPs (decorated with 100% wildtype gpD 794 (grey), lanes 2) served as controls for the decoration reactions. (D) Representative SEC 795 chromatograms showing WT PLPs (grey; retention time (RT), 8.2 mL), 20% RBD_{SARS} PLPs (blue; 796 RT, 8.1 mL), 20% RBD_{MERS} PLPs (yellow; RT, 8.1 mL), and 20% hCOV RBD PLPs (black; RT, 797 8.1 mL). (E) Electron micrographs of PLPs decorated with hCOV RBDs (magnification, 98,000x; 798 scale bars, 100 nm). White arrows indicate the surface density attributed to the RBD proteins 799 projecting from the PLP surface. (F-G) Antigenicity of the purified hCoV-PLPs. ELISA plates were 800 coated with 0.2 µg WT PLPs, 20% RBD_{SARS} PLPs, 20% RBD_{MERS} PLPs, 20% hCoV PLPs, and 801 recombinant RBD protein as indicated. Plates were then probed with 2-fold serial dilutions of a 802 detection antibody, and signals were developed following standard ELISA protocols. For (F), 803 recombinant SARS-CoV-2 RBD protein is denoted as rRBD_{SARS}, and plates were probed with an 804 anti-SARS-CoV-2 RBD monoclonal antibody. For (G), recombinant MERS-CoV RBD protein is 805 denoted as rRBD_{MERS}, and plates were probed with an anti-MERS-CoV RBD detection antibody. 806 Figure 4. RBD_{SARS}-PLPs are immunogenic. (A) Schematic of immunization protocol and

sample collection. (B) WT BALB/c mice (n = 5 mice/group) were immunized with 10 μ g of WT

808 PLPs (control) or 60% RBD_{SARS}-PLPs by intramuscular (i.m.) injection on days 0 and 21. Animals 809 were bled on days 14, 35, 63, and 174 and RBD_{SARS}-specific IgG endpoint titers were determined 810 by ELISA. P values were determined by one-way ANOVA with Tukey's multiple comparisons test. *P < 0.05. ***P < 0.001. (**C**) RBD_{SARS}-specific IgG ELISA endpoint titers in plasma from pre-811 812 pandemic controls (n = 10) and convalescent SARS-CoV-2 PCR positive patients (n = 20). (D) 813 RBD_{SARS}-specific IgG subclass responses on days 14 and 35 were determined by ELISA. P 814 values were determined by unpaired student's t test. *P<0.05. (E-F) SARS-CoV-2 neutralizing 815 activity was determined by a focus reduction neutralization test (FRNT). (E) The dilution of serum 816 that inhibited 50% of infectivity (FRNT50 titer) was calculated for each sample by nonlinear 817 regression analysis, as described in Materials and Methods. P values were determined by oneway ANOVA with Tukey's multiple comparisons test. *P<0.05, ***P<0.001. (F) SARS-CoV-2 818 FRNT50 titers in plasma from pre-pandemic controls (n = 10) and convalescent SARS-CoV-2 819 820 PCR positive patients (n = 20).

821 Figure 5. Immunization with RBD_{SARS}-PLPs protects against SARS-CoV-2 infection and disease. (A) WT BALB/c mice (n = 5/group) were immunized with 10 µg of WT PLPs 60% 822 823 RBD_{SARS}-PLPs by intramuscular (i.m.) injection on days 0 and 21. At day 184, mice were 824 challenged intranasally (i.n.) with 10⁴ PFU of mouse-adapted SARS-CoV-2 MA10. Mice were 825 monitored daily for weight changes. P values were determined by two-way ANOVA with Tukey's 826 multiple comparisons test. **P<0.01, ****P<0.0001. (B-D) At 4 dpi, viral burden in the lung was 827 quantified by plaque assay (B), RT-qPCR for viral genomic RNA (C), and RT-qPCR for N 828 subgenomic RNA (D). (E) Histopathology in lung tissue sections was evaluated by hematoxylin 829 and eosin staining. P values were determined by Mann-Whitney test in **B**, **D**; or by unpaired 830 student's t-test in **C**. ***P*<0.01, ****P*<0.001.

Figure 6. Immunization with RBD_{SARS}-PLPs at low doses elicits potent and durable
SARS-CoV-2 antibody responses and protective immunity. (A-B) WT BALB/c mice (n =
5/group) were immunized with 2.5 μg of WT PLPs (control) or variable doses (2.5, 1.0, or 0.25)

µg) of 60% RBD_{SARS}-PLPs by i.m. injection on days 0 and 21. Animals were bled on days 14, 35, 834 835 63, and 84. (A) RBD_{SARS}-specific IgG endpoint titers were determined by ELISA. P values were 836 determined by Mann-Whitney test. **P<0.01. (B) SARS-CoV-2 neutralizing activity in serum 837 samples was determined by FRNT. FRNT50 titer was calculated for each sample by nonlinear 838 regression analysis. P values were determined by Mann-Whitney test. **P<0.01. (**C**) At day 90 839 post-prime, mice were challenged i.n. with 10⁴ PFU of SARS-CoV-2 MA10 and monitored daily 840 for changes in weight. P values were determined by two-way ANOVA with Tukey's multiple comparisons test. *P<0.05, **P<0.01. (D-G) At 4 dpi, viral burden in the lung was quantified by 841 842 plaque assav (D), RT-qPCR for viral genomic RNA (E), and RT-qPCR for N subgenomic mRNA 843 (F), and histopathology in lung tissue sections was evaluated by hematoxylin and eosin staining (G). P values were determined by Mann-Whitney test. **P<0.01, ***P<0.001. 844

845 Figure 7. Immunization with mosaic hCoV-RBD-PLPs protects against both SARS-CoV-2 and MERS-CoV challenge. (A-B) WT BALB/c mice (n = 5-10 mice/group) were 846 847 immunized with 7.5 µg of WT PLPs (control), 20% RBD_{SARS}-PLPs, 20% RBD_{MERS}-PLPs, or mosaic 848 20% hCoV-RBD-PLPs by intramuscular (i.m.) injection on days 0 and 21. At the times indicated, 849 RBD_{SARS}-specific (A) and RBD_{MERS}-specific (B) IgG endpoint titers were determined by ELISA. 850 Mann-Whitney test. *P<0.05, ***P<0.001. (C) SARS-CoV-2 neutralizing activity was determined 851 by a focus reduction neutralization test (FRNT). The dilution of serum that inhibited 50% of infectivity (FRNT50 titer) was calculated. Mann-Whitney test. *P<0.05, **P<0.01, ***P<0.001, 852 853 ****P<0.0001. (D) MERS-CoV neutralizing activity was determined by a plaque reduction 854 neutralization test (PRNT). The dilution of serum that inhibited 50% of infectivity (PRNT50 titer) 855 was calculated. Mann-Whitney test. **P<0.01, ***P<0.001, ****P<0.0001. (E-H) At day 102 postprime, mice were challenged intranasally (i.n.) with 10⁴ PFU of SARS-CoV-2 MA10 (E-F) or 10⁵ 856 PFU MERS-CoV (G-H). (E and G) Mice were monitored daily for weight changes. Two-way 857 858 ANOVA with Tukey's multiple comparisons test. **P<0.01, **P<0.01. (F and H) At 4 dpi, the

- 859 burden of infectious SARS-CoV-2 (F) and MERS-CoV (H) in the lung was quantified by plaque
- 860 assay. Mann-Whitney test. **P*<0.05, ***P*<0.01.

TABLE 1 | CHARACTERIZATION OF PLP PREPARATIONS

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	Zetasizer*						
Sample	Z-Ave (d.nm)	PDI	Intensity PSD (d.nm)	Volume PSD (d.nm)	ZP (mV)		
Naked PLPs	66.2 ± 1.4	0.073 ± 0.01	71.8 ± 2.1	58.6 ± 1.1	-25.1 ± 0.8		
WT PLPs	73.8 ± 1.3	0.081 ± 0.04	80.1 ± 3.0	65.6 ± 1.6	-17.4 ± 2.4		
5% RBD _{SARS} PLPs	82.4 ± 3.0	0.156 ± 0.05	90.8 ± 2.4	68.5 ± 0.7	-18.8 ± 2.4		
20% RBD _{SARS} PLPs	111.5 ± 6.5	0.285 ± 0.09	121.3 ± 6.0	86.8 ± 5.2	-28.5 ± 1.6		
60% RBD _{SARS} PLPs	110.7 ± 7.1	0.223 ± 0.06	131.9 ± 17	94.4 ± 6.7	-11.5 ± 2.6		
10% RBD _{MERS} PLPs	84.3 ± 3.3	0.164 ± 0.05	89.7 ± 1.5	70.7 ± 0.6	-11.8 ± 1.4		
20% RBD _{MERS} PLPs	94.9 ± 3.4	0.212 ± 0.06	101.7 ± 4.2	76.6 ± 1.4	-22.2 ± 2.1		
40% RBD _{MERS} PLPs	93.1 ± 3.3	0.176 ± 0.04	102.3 ± 4.8	77.7 ± 1.7	-15.2 ± 2.1		
20% hCOV RBD PLPs	115.3 ± 16	0.277 ± 0.11	112.9 ± 9.1	85.7 ± 5.2	-23.3 ± 2.8		

*PLP preparations were diluted in 10 mM HEPES [pH 7.4], and measurements acquired in quadruplicate at 25°C. Values shown are mean ± standard deviation.

Particle sizing data is based on particle diameter (d.nm) as measured by dynamic light scattering from two to three independent analyses. These are reported as Z-average (Z-Ave; weighted mean hydrodynamic size), polydispersity index (PDI; measure of sample heterogeneity) and particle size distributions (PSD; intensity, volume).

Particle charge data is based on overall surface charge (Zeta potential, ZP) as measured by electrophoretic light scattering.

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TABLE S1: SARS-CoV-2 qPCR primer and probe combinations

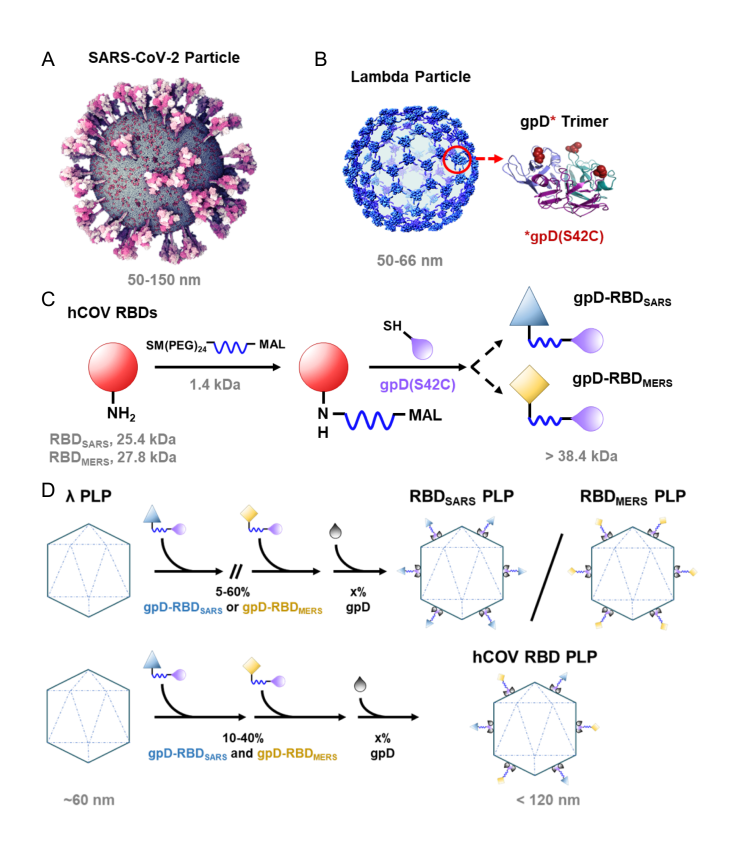
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SARS-CoV-2 E gene qPCR primer and probe

E Sarbeco fwd 1	5'	ACAGGTACGTTAATAGTTAATAGCGT	3'
E Sarbeco rev 2	5'	ATATTGCAGCAGTACGCACACA	3'
E Sarbeco probe 1	5'	FAM-ACACTAGCCATCCTTACTGCGCTTCG-BHQ	3'

SARS-CoV-2 N (sgRNA) qPCR primer and probe

sgN SARS2 fwd	5'	ACCTTCCCAGGTAACAAACCAACCA	3'
sgN SARS2 rev	5'	CACCAAACGTAATGCGGGGTGC	3'
sgN SARS2 probe	5'	FAM-CTGATAATGGACCCCAAAATCAGCGA-BHQ	3'



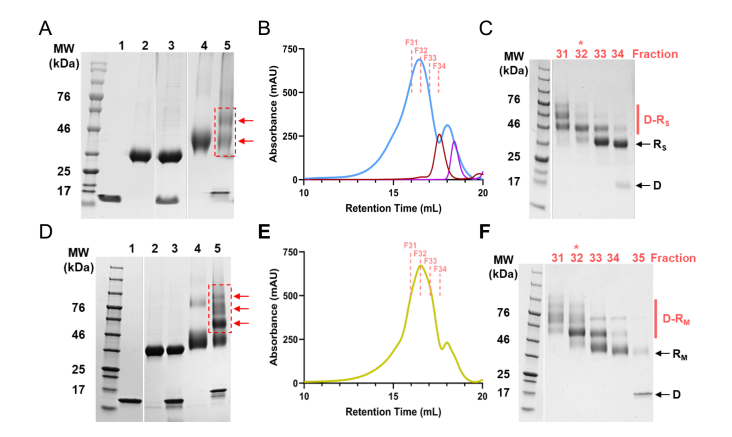


Figure 2

