1	Identification of neutralizing human monoclonal antibodies from Italian Covid-
2	19 convalescent patients
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27	One Sentence Summary: Neutralizing human monoclonal antibodies isolated from Covid-19
28	convalescent patients for therapeutic and prophylactic interventions.

30 ABSTRACT

31 In the absence of approved drugs or vaccines, there is a pressing need to develop tools for therapy and prevention of Covid-19. Human monoclonal antibodies have very good probability of being 32 safe and effective tools for therapy and prevention of SARS-CoV-2 infection and disease. Here we 33 describe the screening of PBMCs from seven people who survived Covid-19 infection to isolate 34 human monoclonal antibodies against SARS-CoV-2. Over 1,100 memory B cells were single-cell 35 sorted using the stabilized prefusion form of the spike protein and incubated for two weeks to allow 36 natural production of antibodies. Supernatants from each cell were tested by ELISA for spike 37 protein binding, and positive antibodies were further tested for neutralization of spike binding to 38 39 receptor(s) on Vero E6 cells and for virus neutralization *in vitro*. From the 1,167 memory B specific for SARS-CoV-2, we recovered 318 B lymphocytes expressing human monoclonals recognizing the 40 spike protein and 74 of these were able to inhibit the binding of the spike protein to the receptor. 41 42 Finally, 17 mAbs were able to neutralize the virus when assessed for neutralization *in vitro*. Lead candidates to progress into the drug development pipeline will be selected from the panel of 43 44 neutralizing antibodies identified with the procedure described in this study.

46 INTRODUCTION

The impact of the SARS-CoV-2 pandemic, with more than 3.5 million cases, 250,000 deaths and 47 more than 25 million people unemployed in the United States alone, is unprecedented. This first 48 49 wave of infection is likely to be followed by additional waves in the next few years, until herd immunity, acquired by vaccination or by infection, will constrain the circulation of the virus. It is 50 therefore imperative to develop therapeutic and preventive tools to face the next waves of SARS-51 52 CoV-2 infections as soon as possible. Among the many therapeutic options available, human monoclonal antibodies (mAbs) are the ones that can be developed in the shortest period of time. In 53 54 fact, the extensive clinical experience with the safety of more than 50 commercial mAbs approved 55 to treat cancer, inflammatory and autoimmune disorders provides high confidence on their safety. These advantages combined with the urgency of the SARS-CoV-2 pandemic, support and justify an 56 accelerated regulatory pathway. In addition, the long industrial experience in developing and 57 manufacturing mAbs decreases the risks usually associated with the technical development of 58 investigational products. Finally, the incredible technical progress in the field allows to shorten the 59 60 conventional timelines and go from discovery to proof of concept trials in 5-6 months (1). Indeed, in the case of Ebola, mAbs were the first therapeutic intervention recommended by the World 61 Health Organization (WHO) and they were developed faster than vaccines or other drugs (2). 62

63 The SARS-CoV-2 spike glycoprotein (S-protein) has a pivotal role in viral pathogenesis and it is considered the main target to elicit potent neutralizing antibodies and the focus for the development 64 of therapeutic and prophylactic tools against this virus (3, 4). Indeed, SARS-CoV-2 entry into host 65 cells is mediated by the interaction between S-protein and the human angiotensin converting 66 67 enzyme 2 (ACE2) (3, 5). The S-protein is a trimeric class I viral fusion protein which exists in a 68 metastable prefusion conformation and in a stable postfusion state. Each S-protein monomer is composed of two distinct regions, the S1 and S2 subunits. Structural rearrangement occurs when the 69 70 receptor binding domain (RBD) present in the S1 subunit binds to the host cell membrane. This

interaction destabilizes the prefusion state of the S-protein triggering the transition into the postfusion conformation which in turn results in the ingress of the virus particle into the host cell (6). Single-cell RNA-seq analyses to evaluate the expression levels of ACE2 in different human organs have shown that SARS-CoV-2, through the S-protein, can invade human cells in different major physiological systems including the respiratory, cardiovascular, digestive and urinary systems, thus enhancing the possibility of spreading and infection (7).

To identify potent mAbs against SARS-CoV-2 we isolated over a 1,100 S-protein specific-memory 77 B cells derived from seven Covid-19 convalescent donors. As the S-protein RBD domain is mainly 78 exposed when this glycoprotein is in its prefusion state (6), we screened naturally produced mAbs 79 against either the S1/S2 subunits and the S-protein trimer stabilized in its prefusion conformation 80 (6). This strategy allows us to identify mAbs able to recognize linear epitopes as well as highly 81 neutralizing trimer specific regions on the S-protein surface. The potent neutralizing effect of trimer 82 83 specific mAbs has already been shown for other pathogens including the respiratory syncytial virus (RSV) (8, 9). 84

In this paper we report the identification of a panel of 318 mAbs from which we plan to select leadcandidates for clinical development.

88 **RESULTS**

89 SARS-CoV2 induces a strong antibody response in patients contracting infection

Patients recovered from SARS-CoV-2 infection were enrolled in two ongoing clinical studies based 90 91 in Rome, Italy (National Institute for Infectious Diseases, IRCCS, Lazzaro Spallanzani) and Siena, Italy (Azienda Ospedaliera Universitaria Senese). We firstly examined whether these patients had 92 93 anti-SARS-CoV-2 S-protein antibodies. Plasma samples were evaluated by enzyme linked 94 immunosorbent assay (ELISA), to assess the polyclonal response to the S-protein trimer, and by their ability to neutralize the binding of the spike protein to Vero E6 cell (neutralization of binding 95 or NOB assay) and for their ability to neutralize the cytopathic effect caused by SARS-CoV-2 96 97 infection in vitro. Results shown in Table 1 and Figure 1 show that among the seven donors included in this study, six were able to produce high titers of SARS-CoV-2 S-protein specific 98 99 antibodies and in particular donors R-042, R-122 and R-188 showed the highest virus neutralizing titers. Only one patient (R-276) mounted low anti-Spike polyclonal response (Fig. 1A-B). 100 Interestingly, despite no statistically significant correlation was observed when Pearson correlation 101 102 analysis was performed, it is possible to observe a trend of correlation between S-protein binding, NOB titer and neutralization titer, suggesting that an abundant response against the S-protein trimer 103 in its prefusion conformation may be indicative of immunity against SARS-CoV-2 (Fig. 1C-D). A 104 105 bigger dataset may be needed to support this observation.

106

Isolation of naturally induced S-protein specific antibodies from SARS-CoV-2 convalescent patients

109 To retrieve mAbs specific for SARS-CoV-2 S-protein, peripheral blood mononuclear cells 110 (PBMCs) from the seven convalescent patients enrolled in this study were collected and stained 111 with fluorescent labeled S-protein trimer to identify antigen specific memory B cells (MBCs). The 112 gating strategy described in Fig. 2 was used to single cell sort into 384-well plates IgG⁺ and IgA⁺

MBCs binding to the SARS-CoV-2 S-protein trimer in its prefusion conformation. A total of 1,167 113 S-protein-binding MBCs were successfully retrieved with frequencies ranging from 0,17% to 114 1,41% (Table 2). Following the sorting procedure, S-protein⁺ MBCs were incubated over a layer of 115 3T3-CD40L feeder cells in the presence of IL-2 and IL-21 stimuli for two weeks to allow natural 116 production of immunoglobulins (10). Subsequently, MBC supernatants containing IgG or IgA were 117 tested for their ability to bind either the SARS-CoV-2 S-protein S1 + S2 subunits (Fig. 3A) or 118 trimer in its prefusion conformation (Fig. 3B) by enzyme linked immunosorbent assay (ELISA). A 119 panel of 318 mAbs specific for the SARS-CoV-2 S-protein were identified showing a broad range 120 of signal intensities (Table 2 and Fig. 3). 121

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123 Functional characterization of S-protein specific mAbs against SARS-CoV-2

Of the 318 supernatants containing S-protein specific mAbs, 265 were screened in vitro for their 124 125 ability to block the binding of the S-protein to Vero E6 cell receptors (Table 2 and Fig. 4A) and for their neutralization activity against the SARS-CoV-2 virus (Fig. 5A). In the NOB assay, 74 of the 126 127 265 tested (28%) S-protein specific mAbs were able to neutralize the antigen/receptor binding 128 showing a broad array of neutralization potency ranging from 52% to over 90% (Fig. 4B). In the viral neutralization assay, supernatants containing naturally produced IgG or IgA were tested for 129 their ability to protect the layer of Vero E6 cells from the cytopathic effect triggered by SARS-130 CoV-2 infection (Fig. 5A). When mAbs were able to protect Vero E6 from infection, i.e. showing 131 neutralization capacity against SARS-CoV-2, no cytopathic effect was observed (Fig. 5B, bottom-132 left box). On the contrary, when mAbs were not able to prevent infection, cytopathic effect on Vero 133 E6 was clearly visible (Fig. 5B, bottom-right box). Out of the 265 mAbs tested in this study, a panel 134 of 17 mAbs neutralized the virus and prevented infection of Vero E6 (Table 2). 135

137 **DISCUSSION**

Human monoclonal antibodies are an industrially mature technology with more than 50 products 138 already approved in the field of cancer, inflammation and autoimmunity. The well-established 139 140 safety profile and the large experience for their development, make mAbs ideal candidates for rapid development especially in epidemic and pandemic settings. So far mAbs have rarely been used in 141 the field of infectious diseases, mostly because the large quantities needed for therapy made them 142 143 not cost effective. However, in recent years, the incredible technological progress to isolate and screen memory B cells made possible to identify highly potent neutralizing mAbs and to further 144 improve their potency of several orders of magnitude through established engineering procedures. 145 146 This possibility resulted into a decreased amount of antibodies necessary for therapy thus making non-intravenous delivery of potent neutralizing mAbs possible. Several candidates are presently 147 under development in the field of HIV, pandemic influenza, RSV and many other infectious 148 diseases (11). Perhaps the most striking demonstration of the power of mAbs for emerging 149 infections came from the Ebola experience. In this case rapidly developed potent mAbs were among 150 151 the first drugs to be tested in the Ebola outbreak and showed remarkable efficacy in preventing mortality (2). Given the remarkable efficacy of this intervention, potent mAbs became the first, and 152 153 so far the only, drug to be recommended for Ebola by the WHO.

In the case of SARS-CoV-2, where so far we do not have any effective therapeutic nor prophylactic interventions, mAbs have the possibility to become one of the first drugs that can be used for immediate therapy of any patient testing positive for the virus, and even to provide immediate protection from infection in high risk populations. Preliminary evidences showed that plasma from infected people improves the outcome of patients with severe disease, therefore it is highly possible that a therapeutic and/or prophylactic mAb-based intervention can be highly effective (*12*). Furthermore, vaccination strategies inducing neutralizing antibodies have shown to protect non-

human primates from infection (13). These results further stress the importance of mAbs as a
measure to counterattack SARS-CoV-2 infection and to constrain its circulation.

In this work we addressed the question of whether mAbs recognizing SARS-CoV-2 can be 163 recovered from memory B cells of people who survived Covid-19 and whether some of them are 164 able to neutralize the virus. Our data show that SARS-CoV-2 specific mAbs can be successfully 165 isolated from most convalescent donors even if the frequency of S-protein specific memory B cells 166 is highly variable among them. In addition, approximately 28 % of isolated mAbs were able to 167 inhibit the binding of the S-protein to the receptor(s) on Vero E6cells. Finally a fraction of isolated 168 mAbs (N=17) were able to effectively neutralize SARS-CoV-2 with high potency when tested in 169 *vitro*. These data suggest that the method we implemented allows us to successfully retrieve mAbs 170 with potent neutralizing activity against SARS-CoV-2 and we plan to select the most promising 171 candidate(s) for drug development. Lead candidates will be further tested for the ability to generate 172 173 resistant viruses and for the ability to induce antibody-dependent disease enhancement in appropriate models. 174

176 MATERIALS & METHODS

177 Enrollment of SARS-COV-2 convalescent donors and human sample collection

This work results from a collaboration with the National Institute for Infectious Diseases, IRCCS – Lazzaro Spallanzani Rome (IT) and Azienda Ospedaliera Universitaria Senese, Siena (IT) that provided samples from SARS-CoV-2 convalescent donors who gave their written consent. The study was approved by local ethics committees (Parere 18_2020 in Rome and Parere 17065 in Siena) and conducted according to good clinical practice in accordance with the declaration of Helsinki (European Council 2001, US Code of Federal Regulations, ICH 1997). This study was unblinded and not randomized.

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Human peripheral blood mononuclear cells (PBMCs) isolation from SARS-CoV-2
 convalescent donors

Peripheral blood mononuclear cells (PBMCs) were isolated from heparin treated whole blood by density gradient centrifugation (Lympholyte-H; Cederlane). After separation, PBMC were: i) frozen in liquid nitrogen at concentration of 10 x 10⁶ PBMC/vial using 10% DMSO in heat-inactivated fetal bovine serum, (FBS) or ii) resuspended in RPMI 1640 (EuroClone) supplemented with 10% FBS (EuroClone), 2 mmol/L L-glutamine, 10 mmol and with 2 mmol/L penicillin, and 50 µg/mL streptomycin (EuroClone). Cells were cultured for 18 hour at 37°C with 5% CO₂. Blood samples were screened for SARS-CoV-2 RNA and for antibodies against HIV, HBV and HCV.

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196 Expression and purification of SARS-CoV-2 S-protein prefusion trimer

197 These expression vector coding for prefusion S ectodomain (kindy gift of Dr. Jason Mc Lellan) was 198 used to transiently transfect Expi293F cells (Thermo Fisher #A14527) using Expifectamine 199 (Thermo Fisher # A14525). Protein was purified from filtered cell supernatants using NiNTA resin 200 (GE Healtcare #11-0004-58), eluted with 250mM Imidazole (Sigma Aldrich #56750), dialyzed 201 against PBS, and then stored at 4°C prior to use.

202

203 Single cell sorting of SARS-CoV-2 S-protein⁺ memory B cells

Human peripheral blood mononuclear cells (PBMCs) from SARS-CoV-2 convalescent donors were 204 stained with Live/Dead Fixable Aqua (Invitrogen; Thermo Scientific) in 100 µL final volume 205 diluted 1:500 at room temperature (RT). After 20 min incubation cells were washed with phosphate 206 buffered saline (PBS) and unspecific bindings were saturated with 50 µL of 20% rabbit serum in 207 PBS. Following 20 min incubation at 4°C cells were washed with PBS and stained with SARS-208 209 CoV-2 S-protein labeled with Strep-Tactin®XT DY-488 (iba-lifesciences cat# 2-1562-050) for 30 min at 4°C. After incubation the following staining mix was used CD19 V421 (BD cat# 562440), 210 211 IgM PerCP-Cy5.5 (BD cat# 561285), CD27 PE (BD cat# 340425) IgD-A700 (BD cat# 561302), CD3 PE-Cy7 (BioLegend cat# 300420), CD14 PE-Cy7 (BioLegend cat# 301814), CD56 PE-Cy7 212 (BioLegend cat# 318318) and cells were incubated at 4°C for additional 30 min. Stained MBCs 213 214 were single cell-sorted with a BD FACSAria III (BD Biosciences) into 384 well plates containing 3T3-CD40L feeder cells and were incubated with IL-2 and IL-21 for 14 days as previously 215 216 described (10).

217

218 ELISA assay with S1 and S2 subunits of SARS-CoV-2 S-protein

219 The presence of S1- and S2-binding antibodies in culture supernatants of monoclonal S-proteinspecific memory B cells was assessed by means of an ELISA assay implemented with the use of a 220 commercial kit (ELISA Starter Accessory Kit, Catalogue No. E101; Bethyl Laboratories, 221 222 Montgomeri, TX, USA). Briefly, 384-well flat-bottom microtiter plates (Nunc MaxiSorp 384-well plates; Sigma-Aldrich) were coated with 25 µl/well of antigen (1:1 mix of S1 and S2 subunits, 1 223 µg/ml each; The Native Antigen Company, Oxford, United Kingdom) diluted in coating buffer 224 (0.05 M carbonate-bicarbonate solution, pH 9.6), and incubated overnight at 4°C. The plates were 225 then washed three times with 100 µl/well washing buffer (50mM Tris Buffered Saline (TBS) pH 226 8.0, 0.05% Tween-20) and saturated with 50 µl/well blocking buffer containing Bovine Serum 227

Albumin (BSA) (50mM TBS pH 8.0, 1% BSA, 0.05% Tween-20) for 1 hour (h) at 37°C. After 228 further washing, samples diluted 1:5 in blocking buffer were added to the plate. Blocking buffer 229 was used as a blank. After an incubation of 1 h at 37°C, the plates were washed and incubated with 230 25 µl/well secondary antibody (horseradish peroxidase (HRP)-conjugated goat anti-human IgG-Fc 231 Fragment polyclonal antibody, diluted 1:10,000 in blocking buffer, Catalogue No. A80-104P; 232 (Bethyl Laboratories, Montgomeri, TX, USA) for 1 h at 37°C. After three washes, 25 µl/well TMB 233 234 One Component HRP Microwell Substrate (Bethyl Laboratories, Montgomery, TX, USA) was added and incubated 10-15 minutes at RT in the dark. Color development was terminated by 235 addition of 25 µl/well 0.2 M H₂SO₄. Absorbance was measured at 450 nm in a Varioskan Lux 236 237 microplate reader (Thermo Fisher Scientific). The threshold for sample positivity was set at twice the OD of the blank. 238

239

240 ELISA assay with SARS-COV-2 S-protein prefusion trimer

ELISA assay was used to detect SARS-CoV-2 S-protein specific mAbs and to screen plasma from 241 242 SARS-CoV-2 convalescent donors. 384 well plates (Nunc MaxiSorp 384 well plates; Sigma Aldrich) were coated with $3\mu g/mL$ of streptavidin diluted in PBS and incubated at RT overnight. 243 Plates were then coated with SARS-CoV-2 S-protein at 3µg/mL and incubated for 1h at room 244 245 temperature. 50 μ L/well of saturation buffer (PBS/BSA 1%) was used to saturate unspecific binding and plates were incubated at 37°C for 1h without CO2. Supernatants were diluted 1:5 in PBS/BSA 246 1%/Tween20 0,05% in 25 µL/well final volume and incubated for 1h at 37°C without CO₂. 25 247 248 µL/well of alkaline phosphatase-conjugated goat anti-human IgG (Sigma-Aldrich) and IgA (Jackson Immuno Research) were used as secondary antibodies. In addition, twelve two-fold serial 249 250 dilutions of plasma from SARS-CoV-2 infected patients were analyzed in duplicate. Plasma 251 samples were diluted in PBS/BSA 1%/Tween20 0,05% (25 µL/well final volume; Starting Dilution 1:80) and incubated for 1h at 37°C without CO2. Next, 25 µL/well of alkaline phosphatase-252 conjugated goat anti-human IgG (Sigma-Aldrich) was added for 1h at 37°C without CO₂. Wells 253

were washed three times between each step with PBS/BSA 1%/Tween20 0,05%. PNPP (pnitrophenyl phosphate) (Thermo Fisher) was used as soluble substrate to detect SARS-CoV-2 Sprotein specific monoclonal antibodies and the final reaction was measured by using the Varioskan Lux Reader (Thermo Fisher Scientific) at a wavelength of 405 nm. Samples were considered as positive if optical density at 405 nm (OD₄₀₅) was two times the blank.

259

260 SARS-CoV-2 virus and cell infection

African green monkey kidney cell line Vero E6 cells (American Type Culture Collection [ATCC] #CRL-1586) were cultured in Dulbecco's Modified Eagle's Medium (DMEM) - High Glucose (Euroclone, Pero, Italy) supplemented with 2 mM L- Glutamine (Lonza, Milano, Italy), penicillin (100 U/mL) - streptomycin (100 μ g/mL) mixture (Lonza, Milano, Italy) and 10% Foetal Bovine Serum (FBS) (Euroclone, Pero, Italy). Cells were maintained at 37°C, in a 5% CO2 humidified environment and passaged every 3-4 days.

Wild type SARS CoV-2 2019 (2019-nCoV strain 2019-nCov/Italy-INMI1) virus was purchased from the European Virus Archive goes Global (EVAg, Spallanzani Institute, Rome). For virus propagation, sub-confluent Vero E6 cell monolayers were prepared in T175 flasks (Sarstedt) containing supplemented D-MEM high glucose medium. For titration and neutralization tests of SARS-CoV-2, Vero E6 were seeded in 96-well plates (Sarstedt) at a density of $1,5x10^4$ cells/well the day before the assay.

273

274 Neutralization of Binding (NOB) Assay

To study the binding of the Covid-19 Spike protein to cell-surface receptor(s) we developed an
assay to assess recombinant Spike protein specific binding to target cells and neutralization thereof.
To this aim the stabilized Spike protein was coupled to Streptavidin-PE (eBioscience # 12-4317-87,

278 ThermoFisher) for 30min at 4°C. and then incubated with VERO E6 cells and binding was assessed

by flow cytometry. The stabilized Spike protein bound VERO E6 cells with high affinity (data notshown).

To assess the content of neutralizing antibodies in sera or B-cell culture supernatants, two 281 microliters of nCoV19 Spike-Streptavidin-PE at 15-30µg/ml in PBS-1%FCS were mixed with two 282 microliters of various dilutions of sera or B-cell culture supernatants in U bottom 96well plates. 283 After incubation at 37°C for 1 hr, 25x103 Vero E6 cells suspended in two microliters of PBS 1% 284 285 FCS were added and incubated for additional 1 hr at 4°C. Non-bound protein and antibodies were removed and cell-bound PE-fluorescence was analyzed with a FACScantoII flow cytometer 286 (Becton Dickinson). Data were analyzed using the FlowJo data analysis software package 287 (TreeStar, USA). The specific neutralization was calculated as follows: NOB (%) = 1 - (Sample288 MFI value – background MFI value) / (Negative Control MFI value – background MFI value). 289

290

291 Viral propagation and titration

The SARS-CoV-2 virus was propagated in Vero E6 cells cultured in DMEM high Glucose 292 293 supplemented with 2% FBS, 100 U/mL penicillin, 100 µg/mL streptomycin. Cells were seeded at a density of 1x10⁶ cells/mL in T175 flasks and incubated at 37°C, 5% CO₂ for 18-20 hours. The sub-294 confluent cell monolayer was then washed twice with sterile Dulbecco's phosphate buffered saline 295 296 (DPBS). Cells were inoculated with 3.5 ml of the virus properly diluted in DMEM 2% FBS at a 297 multiplicity of infection (MOI) of 0.001, and incubated for 1h at 37°C in a humidified environment with 5% CO₂. At the end of the incubation, 50 mL of DMEM 2% FBS were added to the flasks. 298 The infected cultures were incubated at 37°C, 5% CO₂ and monitored daily until approximately 80-299 300 90% of the cells exhibited cytopathic effect (CPE). Culture supernatants were then collected, centrifuged at +4°C at 1,600 rpm for 8 minutes to allow removal of cell debris, aliquoted and 301 302 stored at -80°C as the harvested viral stock. Viral titers were determined in confluent monolayers of Vero E6 cells seeded in 96-well plates using a 50% tissue culture infectious dose assay (TCID50). 303 Cells were infected with serial 1:10 dilutions (from 10-1 to 10-11) of the virus and incubated at 304

 37° C, in a humidified atmosphere with 5% CO₂. Plates were monitored daily for the presence of SARS-CoV-2 induced CPE for 4 days using an inverted optical microscope. The virus titer was estimated according to Spearman-Karber formula (*14*) and defined as the reciprocal of the highest viral dilution leading to at least 50% CPE in inoculated wells.

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310 Semi-quantitative live SARS-CoV-2-based neutralization assay

311 To assess the neutralization titer of anti-SARS-CoV-2 plasma samples from Covid-19 convalescent donors, a semi-quantitative neutralization method was used. Plasma samples were heat-inactivated 312 for 30 minutes at 56°C and 2-fold serially diluted starting from 1:10 to 1:2,560 dilution, then mixed 313 314 with an equal volume of viral solution containing 100 TCID50 of SARS-CoV-2 diluted in D-MEM high Glucose 2% FBS. After 1 hour incubation at 37°C, 5% CO2, 100 µl of the virus-plasma 315 mixture at each dilution was passed to a cell plate containing a sub-confluent Vero E6 monolayer. 316 317 Plates were incubated for 3 days at 37°C in a humidified environment with 5% CO₂, then checked for development of CPE by means of an inverted optical microscope. The reciprocal of the highest 318 plasma dilution that resulted in more than 50% inhibition of CPE was defined as the neutralization 319 320 titer.

321

322 Qualitative live SARS-CoV-2-based neutralization assay

The neutralization activity of culture supernatants from monoclonal S-protein-specific memory B 323 cells was evaluated by means of a qualitative live-virus based neutralization assay against a one-324 325 point dilution of the samples. Supernatants were mixed in a 1:3 ratio with a SARS-CoV-2 viral solution containing 25 TCID50 of virus (final volume: 30 µl). After 1 hour incubation at 37°C, 5% 326 CO_2 , 25 µl of each virus-supernatant mixture was added to the wells of a 96-well plate containing a 327 sub-confluent Vero E6 monolayer. Following a 2-hours incubation at 37°C, the virus-serum mixture 328 was removed and 100 µl of DMEM 2% FBS were added to each well. The plates were incubated 329 for 3 days at 37°C in a humidified environment with 5% CO₂, then examined for CPE by means of 330

an inverted optical microscope. Absence or presence of CPE was defined by comparison of each
well with the positive control (plasma sample showing high neutralizing activity of SARS-CoV-2 in
infected Vero E6) and negative control (human serum sample negative for SARS-CoV-2 in ELISA
and neutralization assayl).

336 Author contribution

- 337 Emanuele Andreano, Ida Paciello isolated single memory B cells and identified S-protein specific
- 338 mAbs (cell sorting and ELISA).
- 339 Piero Pileri, Noemi Manganaro, Elisa Pantano performed NOB assay and produced recombinant
- 340 spike porotein.
- 341 Giulia Piccini, Alessandro Manenti and Emanuele Montomoli performed ELISA and viral
- 342 neutralization assay.
- 343 Marco Troisi, Fabiola Vacca, Concetta De Santi, Dario Cardamone, Anna Kabanova, contributed to
- 344 the characterization of positive memory B cells
- 345 Emanuele Nicastri, Chiara Agrati, Concetta Castilletti, Francesca Montagnani, Arianna Emiliozzi,
- 346 Massimiliano Fabbiani, Maria Rosaria Capobianchi, enrolled patients and isolated PBMCs.
- 347 Claudia Sala, Giuseppe Ippolito, Rino Rappuoli, Coordinated the project

348

- 349 Conflict of interest statement
- 350 RR is an employee of GSK group of companies.
- 351

352 ACKNWOLEDGMENT

We wish to thank Fondazione Toscana Life Sciences in the person of Dr. Fabrizio Landi and Dr. Andrea Paolini and all the administration for their incredible help and support. In particular we would like to thank Mr. Francesco Senatore, Mrs. Laura Canavacci and Mrs. Cinzia Giordano for their support in preparing all the documents needed for the ethical approval of the clinical studies started to carry out this project.

- 357 This work was possible thanks to the technology set up in our European Research Council (ERC) funded lab named vAMRes which
- allowed us to isolate mAbs from vaccinated and/or convalescent patients to tackle the global threat posed by antimicrobial resistance.
- 359 We wish to thank the National Institute for Infectious Diseases, IRCCS, Lazzaro Spallanzani Rome (IT) and the Azienda Ospedaliera
- 360 Universitaria Senese, Siena (IT), for providing blood samples from Covid-19 convalescent donors under studies approved by local
- 361 ethic committees. These studies are conducted under good clinical practice in accordance with the declaration of Helsinki. Donors
- 362 have given their written consent to participate to this study. We also wish to thank all the nursing staff that chose to cooperate for
- 363 blood withdrawal and all the donors that decided to participate to this study.

We would like to thank the whole GSK Vaccines Pre-clinical Evidence Generation and Assay – Immunolgy function led by Dr. Oretta Finco for their availability and support as well as Mrs. Simona Tavarini, Mrs. Chiara Sammicheli, Dr. Monia Bardelli, Dr. Michela Brazzoli, Dr. Elisabetta Frigimelica, Dr. Erica Borgogni and Dr. Elisa Faenzi for sharing their expertise, extreme availability and technical support. We would also like to thank Dr. Mariagrazia Pizza and Dr. Simone Pecetta for initial insightful advices and discussions on this project.

369 We would like to thank Dr. Jason McLellan and his team for generously providing the SARS-CoV-2 S-protein stabilized in its 370 prefusion conformation used in this study. Furthermore, we would like to thank Dr. Daniel Wrapp and Dr. Nianshuang Wang for the 371 precious information and suggestions.

372 We gratefully acknowledge the Collaborators Members of INMI COVID-19 study group: Maria Alessandra Abbonizio, Amina 373 Abdeddaim, Fabrizio Albarello, Gioia Amadei, Alessandra Amendola, Mario Antonini, Tommaso Ascoli Bartoli, Francesco Baldini, 374 Raffaella Barbaro, Bardhi Dorian, Barbara Bartolini, Rita Bellagamba, Martina Benigni, Nazario Bevilacqua, Gianlugi Biava, 375 Michele Bibas, Licia Bordi, Veronica Bordoni, Evangelo Boumis, Marta Branca, Donatella Busso, Marta Camici, Paolo Campioni, 376 Alessandro Capone, Cinzia Caporale, Emanuela Caraffa, Ilaria Caravella, Fabrizio Carletti, Adriana Cataldo, Stefano Cerilli, Carlotta 377 Cerva, Roberta Chiappini, Pierangelo Chinello, Carmine Ciaralli, Stefania Cicalini, Francesca Colavita, Angela Corpolongo, 378 Massimo Cristofaro, Salvatore Curiale, Alessandra D'Abramo, Cristina Dantimi, Alessia De Angelis, Giada De Angelis, Maria 379 Grazia De Palo, Federico De Zottis, Virginia Di Bari, Rachele Di Lorenzo, Federica Di Stefano, Gianpiero D'Offizi, Davide Donno, 380 Francesca Faraglia, Federica Ferraro, Lorena Fiorentini, Andrea Frustaci, Matteo Fusetti, Vincenzo Galati, Roberta Gagliardini, 381 Paola Gallì, Gabriele Garotto, Saba Gebremeskel Tekle, Maria Letizia Giancola, Filippo Giansante, Emanuela Giombini, Guido 382 Granata, Maria Cristina Greci, Elisabetta Grilli, Susanna Grisetti, Gina Gualano, Fabio Iacomi, Giuseppina Iannicelli, Eleonora Lalle, 383 Simone Lanini, Daniele Lapa, Luciana Lepore, Raffaella Libertone, Raffaella Lionetti, Giuseppina Liuzzi, Laura Loiacono, Andrea 384 Lucia, Franco Lufrani, Manuela Macchione, Gaetano Maffongelli, Alessandra Marani, Luisa Marchioni, Raffaella Marconi, Andrea 385 Mariano, Maria Cristina Marini, Micaela Maritti, Alessandra Mastrobattista, Giulia Matusali, Valentina Mazzotta, Paola Mencarini, 386 Silvia Meschi, Francesco Messina, Annalisa Mondi, Marzia Montalbano, Chiara Montaldo, Silvia Mosti, Silvia Murachelli, Maria 387 Musso, Pasquale Noto, Roberto Noto, Alessandra Oliva, Sandrine Ottou, Claudia Palazzolo, Emanuele Pallini, Fabrizio Palmieri, 388 Carlo Pareo, Virgilio Passeri, Federico Pelliccioni, Antonella Petrecchia, Ada Petrone, Nicola Petrosillo, Elisa Pianura, Carmela 389 Pinnetti, Maria Pisciotta, Silvia Pittalis, Agostina Pontarelli, Costanza Proietti, Vincenzo Puro, Paolo Migliorisi Ramazzini, Alessia 390 Rianda, Gabriele Rinonapoli, Silvia Rosati, Martina Rueca, Alessandra Sacchi, Alessandro Sampaolesi, Francesco Sanasi, Carmen 391 Santagata, Alessandra Scarabello, Silvana Scarcia, Vincenzo Schininà, Paola Scognamiglio, Laura Scorzolini, Giulia Stazi, Fabrizio 392 Taglietti, Chiara Taibi, Roberto Tonnarini, Simone Topino, Francesco Vaia, Francesco Vairo, Maria Beatrice Valli, Alessandra 393 Vergori, Laura Vincenzi, Ubaldo Visco-Comandini, Pietro Vittozzi, Mauro Zaccarelli.

394 This publication was supported by the "Centro Regionale Medicina di Precisione" and all the people who answered the call to fight 395 with us the battle against SARS-CoV-2 and for their kind donations platform ForFunding on the 396 (https://www.forfunding.intesasanpaolo.com/DonationPlatform-ISP/nav/progetto/id/3380).

- 397 This publication was supported by the European Virus Archive goes Global (EVAg) project, which has received funding from the
- **398** European Union's Horizon 2020 research and innovation programme under grant agreement No 653316.
- 399

400 TABLE

401 Table 1. SARS-CoV-2 convalescent donors plasma analyses. Plasma S-protein binding titers for

402 each subject were measured by ELISA assays. Neutralization activity was detected by NOB and by

403 neutralization of SARS-CoV-2 infection of Vero cells.

404	Subject ID	Plasma S-protein Binding	Plasma Neutraliziation of binding (NOB)	Plasma Virus Neutralization
405	R-015	10240	>540	320
	R-042	5120	>540	640
406	R-122	10240	>540	640
	R-276	80	180	10
407	R-188	10240	>540	>2560
	S-4	2560	ND	ND
408	S-9	10240	ND	ND

410 Table 2. SARS-CoV-2 convalescent donors S-protein specific MBCs analyses. The Table 411 reports the number of S-protein-specific MBCs that were sorted and screened (for binding by 412 ELISA and for functionality by NOB and viral neutralization) for each subject enrolled in this 413 study.

414	Subject ID	Antigen Specific MBCs (%)	S-protein⁺ MBCs Sorted	S-protein⁺ mAbs (ELISA)	Neutraliziation of binding (NOB)	SARS-CoV-2 Neutralization
415	R-015	1,01	230	18	3	0
	R-042	0,27	39	3	1	0
416	R-122	0,61	5	2	0	0
410	R-276	0,27	66	4	2	0
	R-188	0,17	324	72	8	4
417	S-4	1,41	367	158	60	13
	S-9	0,87	136	53	ND	ND
418		Total	1,167	318	74	17

420 FIGURES

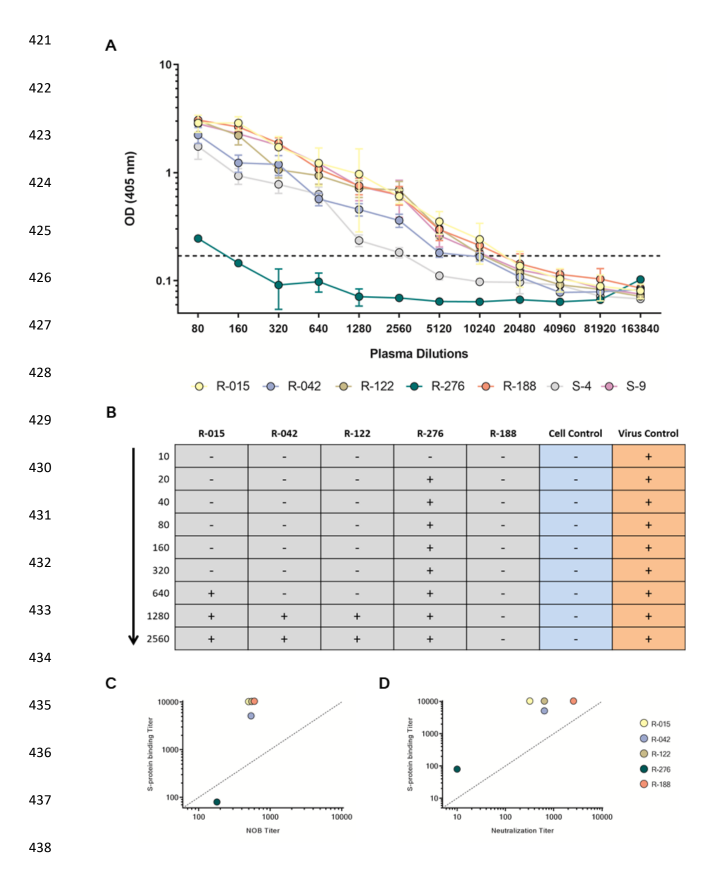


Fig. 1. S-protein binding and neutralization titration of SARS-CoV-2 convalescent donors 439 plasma. (A) Plasma samples were two-fold diluted starting at 1:80 to test their ability to bind the S-440 protein trimer in its prefusion state by ELISA. Results were considered as positive when the OD_{405} 441 value was at least two times higher than the blank. (B) Plasma samples were two-fold diluted 442 starting at 1:10 to test their ability to neutralize SARS-CoV-2 in vitro. Results were considered as 443 positive when no cytopathic effect (-) was observed on Vero E6 cells . (C) The graph shows on the 444 Y axis the Log_{10} S-protein binding titer and the X axis the Log_{10} NOB titer of plasma collected from 445 Covid-19 convalescent patients. Donors R-015 and R-122 are not visible in the graph as their data 446 overlap with those of donor R-188. (D) The graph shows on the Y axis the Log₁₀ S-protein binding 447 titer and the X axis the Log₁₀ neutralization titer of plasma collected from Covid-19 convalescent 448 patients. 449

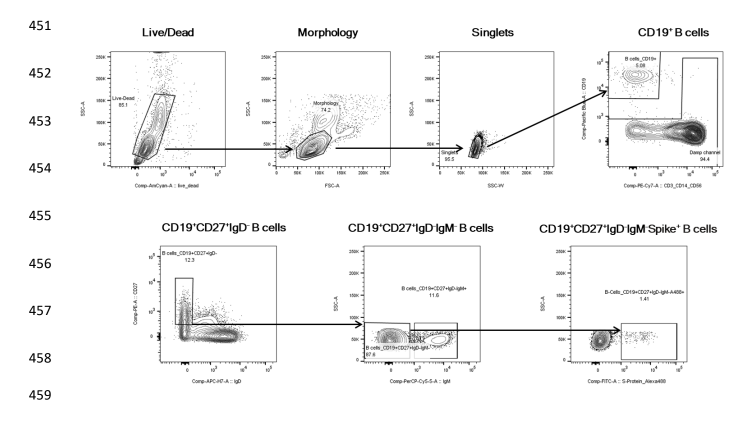


Fig. 2. Gating strategy for S-protein specific MBC single cell sorting. Starting from top left to
the right panel, the gating strategy shows: Live/Dead; Morphology; Singlets; CD19⁺ B cells;
CD19⁺CD27⁺IgD⁻; CD19⁺CD27⁺IgD⁻IgM⁻; CD19⁺CD27⁺IgD⁻IgM⁻S-protein⁺B cells.

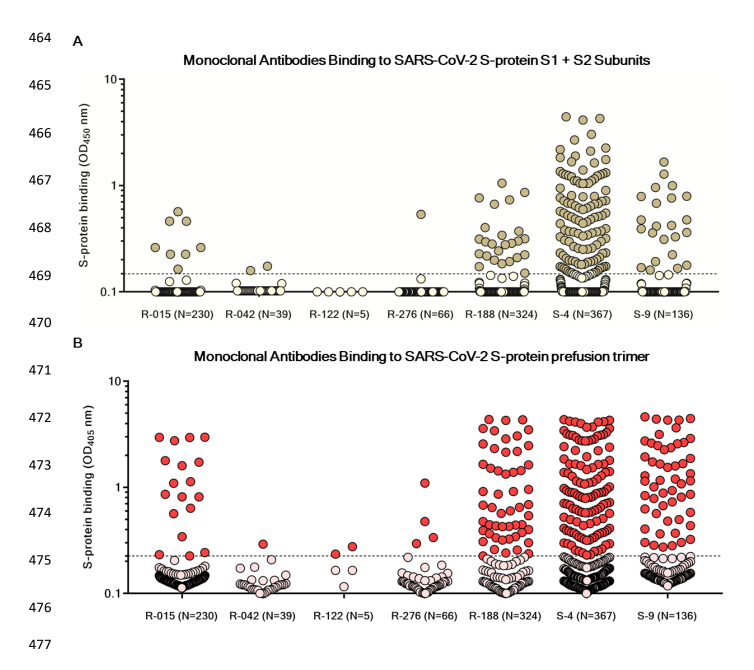


Fig. 3. Identification of SARS-CoV-2 S-protein specific mAbs isolated from convalescent 478 **donor.** (A) The graph shows supernatant tested for binding to the SARS-CoV-2 S-protein S1 + S2479 subunits. Threshold of positivity has been set as two times the value of the blank (dotted line) 480 Darker dots represent mAbs which bind to the S1 + S2 while light yellow dots represent mAbs 481 482 which do not bind- Total number (N) of single cell sorted B cell supernatants screened for binding is also shown for each donor. (B) The graph shows supernatant tested for binding to the SARS-483 CoV-2 S-protein stabilized in its prefusion conformation. Threshold of positivity has been set as 484 two times the value of the blank (dotted line). Red dots represent mAbs which bind to the S-protein 485

- 486 while pink dots represent mAbs which do not bind. Total number (N) of single cell sorted B cell
- 487 supernatants screened for binding is also shown for each donor.

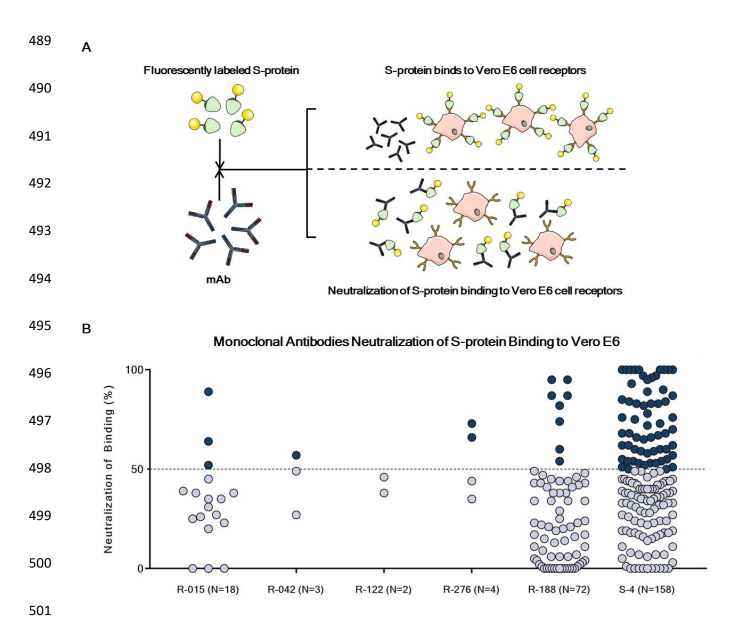


Fig. 4. Neutralization of S-protein binding to Vero E6 cell receptors by S-protein specific 502 mAbs. (A) Schematic representation of the neutralization of binding (NOB) assay used to screen 503 isolated S-protein specific mAbs for their ability to abrogate the interaction between SARS-CoV-504 2/Vero E6 receptors. (B) The graph shows supernatant tested by NOB assay. Threshold of positivity 505 has been set as 50% of binding neutralization (dotted line). Dark blue dots represent mAbs able to 506 507 neutralize the binding between SARS-CoV-2 and receptors on Vero E6 cells, while light blue dots represent non-neutralizing mAbs. The total number (N) of S-protein specific mAbs supernatants 508 screened by NOB assay is shown for each donor 509

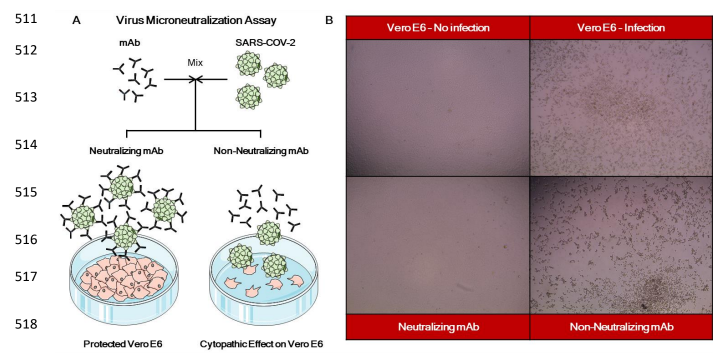




Fig. 5. SARS-CoV-2 neutralization assay for S-protein specific mAbs. (A) Schematic representation of the virus neutralization assay used in this study to assess functional activities of Sprotein specific mAbs. (B) Representative microscope images that showing the cytopathic effect of SARS-CoV-2 or the protection protective efficacy of the screened supernatants on Vero E6 cells by screened mAbs.

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