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1 The SARS-CoV-2 receptor-binding domain expressed in Pichia

2 pastoris as a candidate vaccine antigen

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30 Abbreviations: Receptor-binding domain (RBD), coronavirus disease 2019 (COVID-31 19), severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), angiotensin-32 converting enzyme 2 (ACE2), peripheral blood mononuclear cells (PBMC). Interleukin 2 33 (IL-2), Interleukin 4 (IL-4), Interleukin 6 (IL-6), tumor necrosis factor-alpha (TNF α), 34 interferon gamma (IFNy), non-human primates (NHP), Chinese hamster ovary cells 35 (CHO), baby hamster kidney cells (BHK21), human embryonic kidney cells (HEK293T), 36 immobilized metal ion affinity chromatography (IMAC), reversed-phase chromatography 37 (RP), circular dichroism (CD), optical density (OD), viral neutralization titer (VNT). 38 **Highlights**

- The RBD protein (C-RBD-H6 PP) is expressed with high purity in *P. pastoris*.
- Physico-chemical characterization confirms the right folding of the protein.
- The recombinant protein shows high antigenicity with sera from convalescents.
- The sera from animals inhibit the RBD-ACE2 binding and neutralize the virus.

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• The C-RBD-H6 protein stimulates IFNγ, IL-2, IL-6, IL-4, and TNFα in mice.

44 1. Abstract

45 The effort to develop vaccines based on economically accessible technological 46 platforms available by developing countries vaccine manufacturers is essential to 47 extend the immunization to the whole world population and to achieve the desired herd 48 immunity, necessary to end the COVID-19 pandemic. Here we report on the 49 development of a SARS-CoV-2 receptor-binding domain (RBD) protein, expressed in 50 yeast *Pichia pastoris*. The RBD was modified with addition of flexible N- and C-terminal 51 amino acid extensions aimed to modulate the protein/protein interactions and facilitate protein purification. Fermentation with yeast extract culture medium yielded 30-40 52 53 mg/L. After purification by immobilized metal ion affinity chromatography and 54 hydrophobic interaction chromatography, the RBD protein was characterized by mass-55 spectrometry, circular dichroism, and binding affinity to angiotensin-converting enzyme 56 2 (ACE2) receptor. The recombinant protein shows high antigenicity with convalescent 57 human sera and also with sera from individuals vaccinated with the Pfizer-BioNTech 58 mRNA or Sputnik V adenoviral-based vaccines. The RBD protein stimulates IFNy, IL-2, 59 IL-6, IL-4, and TNFα in mice secreting splenocytes from PBMC and lung CD3+ enriched 60 cells. Immunogenicity studies with 50 µg of the recombinant RBD formulated with alum, 61 induce high levels of binding antibodies in mice and non-human primates, assessed by 62 ELISA plates covered with RBD protein expressed in HEK293T cells. The mouse sera 63 inhibited the RBD binding to ACE2 receptor in an *in-vitro* test and show neutralization of 64 SARS-CoV-2 infection of Vero E6 cells. These data suggest that the RBD recombinant

protein expressed in yeast *P. pastoris* is suitable as a vaccine candidate against
COVID-19.

67 Keywords: SARS-CoV-2, *P. pastoris*, COVID-19, RBD, subunit vaccine

68 **1. Introduction**

The pandemic caused by SARS-CoV-2 has demonstrated the need to modify the way that public health, the governmental role, and international cooperation in science and health have been functioning up to now. It will not be possible to end the pandemic if the desired herd immunity is not achieved and for that it is necessary to vaccinate the vast majority of the world's population in the shortest possible time. It is not enough for industrialized countries to be vaccinated, so the contribution of the developing countries' vaccine manufacturers is more than welcome.

Cuba is an example of a global vision of public health and stands out for the strength of its biotechnology industry where one of the main achievements is the production of the Hepatitis B recombinant vaccine in yeast *P. pastoris* since 1991 [1-3] and the inclusion in the national immunization program since 1992. As a result, the incidence of acute hepatitis B in the country has decreased down to less than 50 cases and acute hepatitis B in children under 5 years old has not been reported since the year 2000 [4].

Therefore, it is feasible to select this technological platform for the development of a
vaccine candidate against SARS-CoV-2.

The mammalian cell expression systems as Chinese hamster ovary cells (CHO), baby hamster kidney cells (BHK21), human embryonic kidney cells (HEK293T), and mouse-

derived myeloma cell lines (NS0, SP2/0) are preferred for the expression of complex proteins for its capacity of proper post-translational modifications equivalent to that of humans, even though the selection of this recombinant protein production platform is limited by its low growth and productivity, and high production cost due to the relevant nutrient requirement [5]. The potential viral contamination of culture medium has also limited the use of the mammalian cell expression systems in large-scale production [6].

92 Pichia pastoris is one of the biotechnological platforms for the production of 93 recombinant proteins that overcome several limitations of bacteria as expression 94 protein aggregation and misfolding, svstem. such as production of toxic 95 lipopolysaccharides, lack of posttranslational modifications, and frequent protein 96 degradation due to the presence of proteases [7]. N-glycans in yeast secreted proteins 97 are highly mannosylated while mammalian proteins have hybrid sugar composition. P. 98 pastoris uses methanol, an inductor of the strong and tightly regulated AOX1 promoter 99 as an exclusive carbon source. Therefore methanol can be used to drive protein 100 expression altogether that it is reached an efficient secretion of the recombinant 101 proteins [8]. On the other hand, the advantages of protein production by *P. pastoris* lie 102 in the fact that the mechanism of protein expression in these microorganisms is close to 103 the ones in mammalian cells and include the right protein folding in the endoplasmic 104 reticulum, protein secretion by Kex2 as signal peptidase and low contamination with 105 host proteins due to its limited production of endogenous secretory proteins [9], which 106 simplifies and makes the purification process more feasible. Other significant 107 advantages of yeast include growth speed, and easy genetic manipulation allowing

108 linearized foreign DNA to be inserted at high efficiency in a chromosome via cross109 recombination to generate stable cell lines [10].

110 The receptor-binding domain of SARS-CoV-2 is a glycosylated 25 kDa protein domain 111 spanning residues N_{331} - K_{529} of the spike protein, including eight cysteine residues 112 forming four disulfide bonds. The domain contains two glycosylation sites (N₃₃₁ and 113 N_{343}) and a central twisted antiparallel beta-sheet formed by five strands with secondary 114 structure short helices and loops [11]. Although glycosylation plays a fundamental role 115 in the immunogenicity and stability of the RBD protein, the variability of the composition 116 and heterogeneity in size of the sugar chains may not influence the receptor binding 117 capacity [12]. Probably, for this reason the glycosylation introduced by *P. pastoris* more 118 distant from mammalian cells and therefore more distant to the humans could contribute 119 to the protein immunogenicity. RBD mediates cell entry through ACE2 host receptor and 120 the levels of RBD binding antibodies strongly correlate with neutralizing antibodies in 121 patients. Even more, the neutralizing antibody kinetics in patients mirrored the kinetics 122 of RBD antibody development [13]. Since RBD upon infection is the main target of 123 neutralizing antibodies, it has become the focus of vaccine design.

The SARS-CoV-2 RBD has been expressed at high levels in *P. pastoris* as a suitable vaccine candidate against COVID-19 [8;14;15]. The comparison between RBD obtained from *P. pastoris* and HEK293T mammalian cells by CD and tryptophan fluorescence shows that proteins were properly folded as well as that they have similar temperature stability despite differences in glycosylation of both expression platforms.

Here, we report the design of an RBD protein vaccine candidate, its expression in *P. pastoris* yeast, protein purification, the physico-chemical characterization and the

capacity of the protein to elicit ACE2 binding inhibition antibodies, and neutralizing
response in mice and monkeys. Our approach differs from the previously reported
expression of RBD in *P. pastoris* [8] in the inclusion of the RBD glycosylation sites
maintaining the advantage of the glycosylation capacity of *P. pastoris*.

135 **2. Materials and Methods**

136 **2.1. Biological reagents, protein codification and serum panels**

137 Human and murine RBD and ACE2 receptor chimeric proteins (hFc-RBD, mFc-RBD, 138 hFc-ACE2, mFc-ACE2) were supplied by the Center of Molecular Immunology (CIM, 139 Havana, Cuba). All the chimeric proteins were purified by protein-A based purification of 140 the supernatant of stable transduce HEK293T cells and eluted in PBS. hFc-RBD was 141 conjugated with peroxidase (HRP) and henceforth named hFcRBD-HRP. H6-RBD was 142 produced as E. coli inclusion bodies [16], while RBD-H6 and C-RBD-H6 HEK were 143 secreted to the supernatant of stably transduced HEK293T. All three proteins were 144 purified by IMAC and the final buffer was exchanged to PBS. N-terminal end segment 145 (C), and C-terminal end six histidine tag (H6), added to the central RBD protein 146 sequence in a different protein constructions are codified as C-RBD-H6. Acronyms PP 147 and HEK next to the protein code refers to the expression system when *P. pastoris* or 148 HEK293T cells where used.

Different human sera used as controls include sera from volunteers vaccinated with Pfizer/BioNtech or Gamaleya's Sputnik V (Gam-COVID-Vac) vaccine, and sera from convalescent patients. All individuals gave their writing informed consent for the use of their serum.

153 2.2. Construction of Pichia pastoris strains expressing RBD from SARS-CoV-2 (C 154 RBD-H6 PP)

155 A sequence coding for residues 331-530 of the Spike protein of SARS-CoV-2 strain 156 Wuhan-Hu-1 (NCBI Acc. No. YP 009724390) with the appropriate N- and C-terminal 157 extensions was codon-optimized for S. cerevisiae using J-Cat [17] and cloned in-frame 158 with the KEX2 cleavage site of the pre-pro MATa sequence of pPICZaA (Invitrogen, 159 USA), placing it under transcriptional control of the *P. pastoris* AOX1 promoter. After 160 sequence verification, a representative clone was used to transform *P. pastoris* strain X-161 33 [18], selecting transformants by plating in YPD agar at Zeocin[™] concentrations of 162 100, 200, 400, and 800 µg/mL. Three transformants from the plate with the highest 163 concentration of Zeocin[™] were randomly picked, purified by further streaking on YPD-164 Zeocin[™], and used to prepare small seed banks. The seed banks were used in turn to 165 inoculate small-scale (50 mL) BMGY cultures [18] that were induced after 24 h of 166 growth at 28 °C by the addition of 0.5 % methanol. Culture supernatants were analyzed 167 at 96 h post-induction by SDS-PAGE and Western blotting with an anti-His6 antibody 168 (Promega, USA).

169 2.3. Fermentation

Fermentation was carried out in a 75-liters Chemap fermenter (Germany) with a working volume of 50 L of fermentation medium containing per liter of culture 8.8 mL of 85 % phosphoric acid, $6.92 \text{ g MgSO}_{4.7}\text{H}_2\text{O}$, $1.23 \text{ g (NH}_4)_2\text{SO}_4$, $16.77 \text{ g K}_2\text{HPO}_4$, 0.46 g CaCl₂.2H₂O, 32.3 mL of 98% glycerol, 4.61 g yeast extract, 4 mL histidine solution, 5 mL of 400X vitamin base solution (1 g L⁻¹ myo-inositol, 0.8 g L⁻¹ D-calcium

pantothenate, 0.8 g L^{-1} thiamine hydrochloride, 0.8 g L^{-1} pyridoxol hydrochloride, 0.2 g 175 176 L^{-1} nicotinic acid, 0.8 mg L^{-1} D (+) Biotin), and 1 mL of sterile-filtered trace 1000X element solution (6 g L⁻¹ CuSO₄.5H₂O, 0.415 g L⁻¹ KI, 3 g L⁻¹ MnSO₄.H₂O, 1 g L⁻¹ 177 Na₂MoO₄.2H₂O, 0.1 g L⁻¹ H₃BO₃, 20 g L⁻¹ ZnSO₄.7H₂O, 65 g L⁻¹ FeSO₄.7H₂O, 10 mL. 178 179 conc.H₂SO₄) was added after autoclaving [19]. After inoculation of the bioreactor with 4 180 L of pre-culture, the temperature was maintained at 30 °C and the pH at 4.75 by 181 pumping in a liquid ammonia solution. When a dissolved oxygen peak or a widening of 182 the pH peaks is observed, the fed-batch phase is started with glycerol 50 % at 540 mL/h 183 for 2-3 h. After one hour, the temperature is lowered to 25 ° C and raises the pH to 5.5. 184 When the 50 % glycerol increase is exhausted, 800 mL of methanol is added at 185 maximum flow of the peristaltic pump. The induction phase with methanol is started at 186 an initial flow of 240 mL/h, 4 h later the flow is increased to 380 mL/h, and then at 480 187 mL/h. This last flow is maintained until the end of fermentation (38-44 h).

188 2.4. Protein purification

189 After 48 h of fermentation in a yeast extract culture medium, the culture was harvested, 190 and cells were removed by centrifugation retention time between 5 -10 min at 15000 191 rpm at 4 °C. The fermentation supernatant was filtered in tandem conditions from 8 µm 192 - 3 µm - 0.45 µm using cellulose filters. The supernatant was concentrated with a 193 tangential flow filtration system with 30 kDa Hydrosart® membrane (Sartorius, 194 Germany), besides buffer exchanged was carried out against PBS buffer containing 5 195 mM of imidazole. The IMAC column (Chelating Sepharose[™] FF, Cytiva) was 196 equilibrated in the same buffer and the sample was loaded and sequentially washed by 197 using 30 column volumes of PBS containing 10 mM and 20 mM imidazole. Elution was

198 carried out by increasing the concentration of imidazole up to 250 mM in the equilibrium 199 buffer. The eluted protein was applied to a RP C4 column (Tosohaas, Japan) with 200 dimensions 50 mm in diameter by 250 mm long for a resin volume of 500 mL and with a 201 particle size of 15 to 20 µm. The column is coupled to a Shimadzu model LC-20AP 202 semi-preparative HPLC purification system. The column was equilibrated with 0.5 % 203 TFA solution, and protein was eluted by using a linear gradient of 1 % TFA solution (RP 204 solution A) and acetonitrile with 0.5 % TFA (RP solution B) from 32 to 45 % solution B in 205 40 minutes. The protein elutes between 35 and 37 % of solution B, approximately in one 206 column volume. The fraction collected with the purified protein was concentrated by 207 tangential flow filtration system with a 10 kDa Hydrosart® membrane and pooled 208 aseptically using a 0.22 μ m and stored at -20 °C.

209 2.5. ESI-MS analysis of the deglycosylated C-RBD-H6 PP protein

210 The volume equivalent to 100 µg of the protein dissolved in PBS (pH 7.2) was 211 deglycosylated with 1µL of PNGase-F (500 units, New England Biolabs) in presence of 212 0.5 M guanidine hydrochloride and 5 mM N-ethylmaleimide during 2 h at 37 °C. An 213 aliquot of 10 µg of deglycosylated protein was desalted by using ZipTip C18 (Millipore) 214 and the elution was loaded into the metal-coated nanocapillary for ESI-MS analysis. 215 The remaining deglycosylated protein was digested by using an in-solution buffer-free 216 trypsin digestion protocol previously reported [20] and adapted to the analysis of SARS-217 CoV-2 RBD proteins by introducing some modifications that provide full-sequence 218 coverage and detection of post-translational modifications in a single ESI-MS spectrum

219 [16]. Other experimental conditions for ESI-MS analysis are similar to those reported220 previously [16].

221 **2.6.** Surface plasmon resonance experimental procedure

222 The interaction between mFc-ACE2 fusion protein and the recombinant C-RBD-H6 PP 223 was monitored by SPR using a BIACORE X (GE Health-care) at 25 °C in a multi-cycle 224 mode. Briefly, mFc-ACE2 was immobilized on a Protein A biosensor chip (GE Health-225 care) according to the manufacturer's protocol through the flow cell 1 (FC1). The FC2 226 was used as the reference cell. The real-time response of the C-RBD-H6 PP over the 227 immobilized mFc-ACE2 was recorded by duplicate in a concentration range from 15 to 228 2000 nM, at 10 µL/min flow rate for 120 s, while the dissociation took place for another 229 120 s. The running buffer was PBS (pH = 7.2). After each cycle the chip was 230 regenerated using pH = 2.0 glycine buffer. The equilibrium dissociation constant 231 (binding affinity, K_D) was estimated with the BIAevaluation® software (GE Healthcare) 232 using the Langmuir 1:1 interaction model. At least five curves were taken into account 233 for kinetics calculations.

234 **2.7. Structural Analysis by Circular Dichroism (CD) Spectroscopy**

CD spectra were acquired in a Jasco J-1500 CD spectrometer (Jasco, Japan). All measurements were carried out at 24 °C, the far UV CD spectra were studied at 100 μg/ml protein concentration (a 10 fold dilution in water of the stock solution) using a 1mm quartz cuvette. The near UV CD spectra were studied at the protein concentration of the stock solution in 20 mM pH 7.4 Tris buffer using a quartz cuvette of 10 mm path length. The spectra of the corresponding solution were subtracted. The far UV CD spectra were further analyzed by the BeStSel method [21;22] to estimate the secondary
structure content of the protein and the results were compared with the values derived
from the 3D coordinates of the crystallographic structure of the spike protein of SARSCoV-2 (PDB file 6yla) using DSSP method [23] implemented in Whatif program package
[24]

246 2.8. Animals and immunization schedules

247 Three different animal species were used for evaluation of immunogenicity of the C-248 RBD-H6 PP protein: BALB/c mice, Sprague–Dawley (SD) rats, and African green 249 monkeys (*Chlorocebus aethiops sabaeus*). Six- to eight-week-old female BALB/c mice, 250 and male and female SD rats were used for the study and housed in the animal facility. 251 The experimental protocols were approved by the Ethical Committee on Animal 252 Experimentation of the Center for Genetic Engineering and Biotechnology (CIGB, 253 Havana, Cuba) and the Center for Production of Laboratory Animals (CENPALAB, 254 Bejucal, Cuba).

The immunogen content per 500 μ L: 50 μ g of C-RBD-H6 PP protein adjuvated with 0.3 mg of aluminum hydroxide gel (Alhydrogel ®) in phosphate buffer (0.28 mg of disodium hydrogen phosphate, 0.31 mg of sodium dihydrogen phosphate dihydrate, 4.25 mg of sodium chloride).

BALB/c mice: Immunogenicity in mice was evaluated using a three-dose schedule with a 50 µg dose by intraperitoneal route with a 7 and 14 days interval before the second and third doses respectively. Blood was collected a week after the first boost and 7 and

14 days after the second boost. The number of animals per group, age, and gender ofthe animals are described in individual experiments.

Sprague Dawley rats: Immunogenicity of the C-RBD-H6 PP protein was tested in 20 SD rats (10 male and 10 female) during chronic toxicology study, with a 9 μg dose administered by the intramuscular route once a week, for 10 consecutive weeks for a total of 90 μg. Animals were bled three days after the last dose.

268 *Chlorocebus aethiops sabaeus* non-human primates: NHP ages between 3 to 6 years 269 and with 2-7 kg of weight were kept at the animal's facility at the CENPALAB. A total of 270 20 NHP were randomly assigned to 3 groups including the placebo (2 animals/gender, 271 total 4), the low dose (50 μ g, 3 animals/gender, total 6), and the high dose (100 μ g, 5 272 animals/gender, total 10). Seven days post-1st and 2nd boosting and after overnight 273 fasting, monkeys were sedated by intramuscular injection of ketamine hydrochloride (10 274 mg/kg) and bled from the femoral vein.

275 Specific anti-RBD titers and the inhibition of its interaction with ACE2 receptor were 276 evaluated using ELISA. Live SARS-CoV-2 neutralization was assessed using a 277 microneutralization assay.

278 2.9. Serum antibodies evaluation

279 Antibody detection by ELISA

The reactivity of sera from immunized animals was determined by ELISA. Briefly, 0.25
µg of RBD protein produced in HEK293T cells (Center for Molecular Immunology,
Havana) was used to coat 96-well microtiter plates (Corning Costar, Acton, MA) in 0.1
M sodium carbonate buffer (pH 9.6) at 4 °C overnight. After the plates were blocked

284 with 2 % skim milk, 0.05 % Tween 20, serially diluted mouse, rat or NHP sera or control 285 monoclonal antibodies SS-1, SS-4, SS-7 and SS-8 (Center for Genetic Engineering and 286 Biotechnology, Sancti Spiritus, Cuba), were added and incubated at 37 °C for 2 h in 0.2 287 % skim milk, 0.05 % Tween 20 in PBS, followed by six washes with PBS containing 288 0.05 % Tween 20. Bound antibodies were detected with horseradish peroxidase-289 conjugated goat anti-mouse IgG (SIGMA, USA), anti-Rat IgG (SIGMA, USA) anti-290 human IgG (1:10000, Jackson, USA) at 37 °C for 1 h, followed by washes. The reaction 291 was detected after the addition of 3.3-5.5-tetramethylbenzidine and quantified using a 292 microplate reader at 450 nm (BMG Labtech, Germany). This assay was also used for 293 the evaluation of RBD antigenicity using sera from immunized animals, and from 294 subjects that received Pfizer-BioNTech or Sputnik V vaccines, or are COVID-19 295 convalescents.

296 RBD to ACE2 plate-based binding assay

297 A competitive ELISA was performed to determine the inhibitory activity of the anti-RBD 298 polyclonal sera on the binding of the hFc-ACE2 coated plates to an hFc-RBD-HRP 299 conjugate. Briefly, the wells of ELISA plates were coated with 0.25 µg of recombinant 300 hFc-AEC2 as described above. A mixture containing an hFc-RBD-HRP conjugate and 301 serial dilutions of the sera were pre-incubated for 1h at 37 °C. A hundred microliters of 302 the mixture were added to hFc-ACE2 coated plates and further incubated for 90 min at 303 37 °C. The binding of the HRP tagged RBD to the receptor was detected after the 304 addition of 3,3-5,5-tetramethylbenzidine and reading at 450 nm. A similar assay was 305 used to characterize the ability of the C-RBD-H6 PP and C-RBD-H6 HEK proteins to 306 block the interaction of hFc-RBD-HRP with coated hFc-ACE2.

307 2.10. Microneutralization of live SARS-CoV-2 virus in Vero E6

308 The neutralization antibody titers were detected by a traditional virus microneutralization 309 assay (MN50) using SARS-CoV-2 (CUT2010-2025/Cuba/2020 strain). Vero E6 cells 310 (2×10⁴ per well) were seeded in 96-well plates one night before use. Animals' sera were 311 inactivated at 56 °C for 30 min. The samples were prepared by two-fold serial sera 312 dilutions in the Eagle's Minimal Essential Medium (MEM, Gibco, UK) containing 2 % 313 (v/v) fetal bovine serum (Capricorn, Germany). SARS-CoV-2 strain at 100 TCID50 was 314 incubated in the absence or presence of diluted sera for 1 h at 37 °C. Afterward, Vero 315 E6 cell were overlaid with virus suspensions. At 96 h post-infection, the cells were 316 inspected for signs of cytopathogenic effects (CPE) by optical microscopy and stained 317 with neutral red (Sigma, USA). After three washes neutral red was dissolved in lysis 318 solution (50 % ethanol, 1 % acetic acid) for 15 min at 25 °C, and optical density (OD) 319 was detected at 540 nm. The highest serum dilution showing an OD value greater than 320 the cut-off was considered as the neutralization titer. The cut-off value is calculated as 321 the average of the OD of the cell control wells divided by two. The viral neutralizing 322 titers (VNT50) were calculated as the highest serum dilution at which 50 % of the cells 323 remain intact according to neutral red incorporation in the control wells (no virus added).

324 2.11. Cellular immune response

325 Long-term cellular immune response was evaluated in BALB/c mice using 326 subcutaneous administration of a formulation containing an equal antigen to alum ratio. 327 Animals receive 25 μ g of the C-RBD-H6 PP antigen by the subcutaneous route in a 100 328 μ L volume in a 0-14-35 days schedule. Blood samples were evaluated two weeks after

329 the last immunization and animals were euthanized 3 months later to assess for 330 systemic and lung resident cells response to in vitro antigen recall. Splenocytes were 331 flushed out by perfusion using gentamycin supplemented PBS and lung resident 332 lymphocytes were extracted after elimination of remnant blood by left auricular perfusion 333 with PBS. To detach leukocytes from lung tissue, a lung dissociation enzyme mix (130-334 095-927, Miltenyi, Germany) was used according to manufacturer instructions and using 335 a cell dissociator (GentleMACS Octo Dissociator, Miltenyi). CD3+ cells were further 336 selected from lung suspension using negative selection with magnetic beads (130-095-337 130, Miltenyi) and both lung CD3+ cells and splenocytes live cells were counted using a 338 flow cytometer (CyFlow, Sysmex, Germany).

339 For the re-stimulation assays, splenocyte or lung selected CD3+ suspensions were diluted to 10x10⁶ CD3+ live cells/mL and 50 µL of each sample was seeded in two 96-340 341 well U bottom tissue culture plates. Cells were re-stimulated with 50 µL of 20 µg/mL C-342 RBD-H6 PP, or just media (unstimulated) for 72 h, and the supernatant was analyzed at 343 1:2 dilution using Biolegend Deluxe cytokine kits for IL-2 (431004), IFNy (430804), IL-6 344 (431304), IL-4 (431104) and TNF α (430204) following manufacturer's instructions. Cells 345 were transferred at that point to anti IFNy coated ELISpot plates (Mabtech, Germany) 346 and results were analyzed 24 h later according to established procedures.

347 **2.12.** ELISpot assay with samples from previous naturally infected individuals

348 PBMCs from subjects with previous natural infection with SARS-CoV-2 were isolated 349 from 7 mL of whole blood collected using CPT tubes (Becton Dickinson, US), and store 350 in liquid nitrogen until analyzed. After overnight resting the cells in Optmizer media 351 (Gibco, Invitrogen, US) CD3+ live cells were counted using flow cytometry and seeded 352 on activation plates at 5×10^4 cells per well with 10 µg/mL of the recombinant C-RBD-H6 353 PP protein for 72 h. After transferring the cells to anti-IFNγ pre-coated plates (Mabtech, 354 GE). The amounts of T cells secreting the cytokine were detected after 20 h of 355 incubation as recommended by the manufacturer. All individuals gave their writing 356 informed consent for the use of their samples.

357 2.13. Statistical analysis

Prism 8.4.3 software was used to generate dose-response curves for ELISA test and to calculate EC50 values when needed. One way-ANOVA with Sidak's multiple comparisons tests-was used to determine the significance of differences and Spearman test was used to assess for parameters correlations. Wilcoxon matched paired test was used to perform paired analyzes for both stimulated/non-stimulated samples, the evolution of RBD specific titers and inhibition of ACE2 binding.

364 3. Results

365 3.1. Design of the C-RBD-H6 expression cassette

366 Recombinant C-RBD-H6 PP protein was designed as a potential subunit vaccine 367 candidate against SARS-CoV-2 displaying e modular structure consists of: a) a globular 368 central – RBD- domain comprising residues N₃₃₁-K₅₂₉ of the spike protein, b) additional 369 N- and C-terminal segments including polar and flexible linkers rich in Glycine and 370 Serine (Gly⁹-Ser¹⁵ and Gly²¹⁵-Ser²²⁹). The N- and C-terminal extensions are aimed to 371 modulate potential protein-protein interactions and facilitate protein purification through a high accessible hexa-Histidine tag (His²³²-His²³⁷). Topologically, the extensions are located in the opposite site of the protein respect to the receptor binding motif and its presence should sterically hinder potential aggregation problems associated to the presence of the exposed and disulfide bonded Cys⁷⁶ and Cys²¹⁰. (The C-RBD-H6 PP protein sequence is included as a supplemental material S1).

377 3.2. Expression and purification of the C-RBD-H6 PP

378 A construct for the expression in *P. pastoris* of the RBD from SARS-CoV-2 under 379 control of the AOX1 promoter, denominated pPICZa-CtagRBDH6, was prepared as 380 described in Materials and Methods and used to obtain RBD-expressing yeast clones. 381 Fig. 1 (lanes A-C) shows the result of the analysis of culture supernatants from 3 382 randomly selected clones after induction with methanol. There is a noticeable smear in 383 the supernatant from all three clones, ranging between 25 kDa and 50 kDa, that is 384 absent from the control transformed with the empty vector (lane V, Fig. 1), and resolves 385 into a ladder when probed by Western blotting with an anti-His6 antibody. Considering 386 the presence of three-potential N-glycosylation sites in the sequence of C-RBD-H6 PP 387 protein; and also that its molecular mass calculated from its cDNA sequence is 26 kDa 388 the presence of a broad and diffuse band detected on SDS-PAGE analysis suggest that 389 the C-RBD-H6 PP is being secreted in all three cases as a N-glycosylated protein. 390 Based on signal intensity in Western blotting, clone C was denominated X33-23 and 391 selected for further work (Fig. 1). The C-RBD-H6 PP protein was purified from the 392 supernatant following the procedures described in Materials and Methods, by IMAC, 393 followed by polishing using a RP to obtain a final preparation with high purity. After 394 purification, the C-RBD-H6 PP was obtained with a final yield between 30-40 mg/L of

culture medium, with more than 98% of purity at the end of the downstream process(Fig. 2 and 3).

397 **3.3. ESI-MS** analysis for determining the accurate molecular mass, the verification

398 *of the amino sequence and the assessment of disulfide bonds of the* 399 *deglycosylated C-RBD-H6 PP protein*

400 The C-RBD-H6 PP protein deglycosylated with PNGase-F was analyzed by ESI-MS 401 and the multiply charged spectrum is shown in Fig. 4.A. Deconvolution of this ESI-MS 402 spectrum (Fig. 4.B) showed a very intense signal at 26069.54 Da. This molecular mass 403 agrees very well with the calculated molecular mass (26069.87 Da) for the C-RBD-H6 404 PP considering the presence of four intramolecular disulfide bonds and the two out the 405 three potential N-glycosylation sites transformed into Asp residues due to the action of 406 PNGase-F. Also a low-abundance signal (see star in Fig. 4.B) indicates that a minor 407 fraction of this molecule is devoid of the extension of four N-terminal amino acids (-408 NSWF) not belonging to the RBD protein.

409 To verify the amino acid sequence as well as the disulfide bonds the deglycosylated 410 protein was digested with trypsin using an in-solution buffer-free digestion protocol 411 developed in our group [16;20]. ESI-MS analysis of the tryptic peptides is shown in Fig. 412 4.C. Full-sequence coverage of C-RBD-H6 PP was verified and the assignment for 413 signals observed in this mass spectrum is summarized in Table 1. ESI-MS/MS analysis 414 of the signal detected at m/z 1399.64, 4+ confirmed that Asn³³¹ and Asn³⁴³ (two out the 415 three potential N-glycosylation sites) were transformed into Asp residues by the action 416 of PNGase F (see underlined residues in the Table). The four disulfide bonds (C₃₃₆-C₃₆₁, 417 C₃₇₉-C₄₃₂, C₃₉₁-C₅₂₅ and C₄₈₀-C₄₈₈) present in the native S protein of SARS-CoV-2 were 418 also detected in this ESI-MS spectrum (Fig. 4.C and Table 1). Tryptic peptides
419 containing free cysteine residues and S-S scrambling variants were not detected.

ESI-MS analysis of the deglycosylated C-RBD-H6 PP protein and its derived tryptic
peptides confirmed that N-glycans in its structure increased considerably its molecular
mass by SDS-PAGE analysis.

423 *3.4. Surface Plasmon Resonance characterization of the RBD-ACE2 binding* 424 *affinity*

425 As shown in Fig. 5, there was no appreciable signal in terms of response units (RU) 426 when measurements were performed a non-related protein used as negative control for 427 BIACORE experiments, immobilizing mFc-ACE2 ligand by Fc region on to a Prot A chip. In contrast, the association rate for C-RBD-H6 PP protein to mFc-ACE2 was 5.4×10⁵ M⁻ 428 1 ·s⁻¹, with a dissociation rate of 7.7×10⁻³ s⁻¹. The equilibrium was reached between 25 429 and 30 seconds, with an estimated dissociation constant of $K_D = 14.3 \times 10^{-9}$ M. The 430 431 association/dissociation rates as well as K_D were in the expected ranges, according to 432 the previously reported results for the RBD-ACE2 molecular interaction [25].

433 **3.5. Secondary structure analysis by CD Spectroscopy**

Fig. 6 shows the far UV CD spectrum of the C-RBD-H6 PP protein revealing characteristic bands similar to the observed in other recombinant RBD proteins reported previously [8], with maxima at 192 and 231 nm – due to the aromatic contribution - and minimum at 207 nm. Furthermore, as shown in Table 2 the secondary structure content of the protein estimated by BeStSel (7.9% helix, 28.7% beta - antiparallel, relaxed and right-handed-, 12.9% turn y 50.5% others) is very similar to the values assigned using
the 3D coordinates [11]. Moreover, as observed in Fig. 7 the near UV CD spectrum of
the protein is well structured, with bands at 263, 269, 277, 281 and 299 nm, indicative of
the presence of well-packed aromatic and cysteine residues as expected for a properly
folded protein.

444 3.6. Antigenicity analysis of C-RBD-H6 PP

445 To confirm the antigenicity of the yeast-derived protein, several ELISA tests were 446 performed with SS-1, SS-4, SS-7 and SS-8 anti-RBD monoclonal antibodies obtained 447 immunizing with RBD-H6 HEK protein produced in the mammalian cells HEK293T. One 448 of these monoclonal antibodies (SS-8) inhibits the protein binding to the ACE2 receptor 449 with an IC50 of 60 ng/mL. The binding to C-RBD-H6 PP protein was compared to the 450 binding to C-RBD-H6 HEK as shown in Fig. 8.A,E. The reactivity does not differ 451 between the two proteins. Human sera with high neutralization titers for live SARS-CoV-452 2 viral challenge of Vero E6 cells from eight COVID-19 naturally infected patients and 453 seven individuals that already received Pfizer-BioNTech or Sputnik V complete vaccine 454 schedules were incorporated into the testing panels. None properly folded C-RBD-H6 455 protein produced as inclusion body in BL-21 E coli strain was used as a negative 456 control. C-RBD-H6 PP protein obtained from yeast was able to maintain antigenicity 457 comparable to the protein purified from mammalian cells (Fig. 8.B,F and C,G). Minimal 458 signals were detected with the sera from animals immunized with RBD protein 459 expressed in *E. coli*, corroborating the need for a correctly folded protein to ensure the 460 exposition of immunodominant epitopes (data not shown). Further characterization was 461 conducted using polyclonal sera from mice and NHP immunized with C-RBD-H6 PP

and C-RBD-H6 HEK proteins, and with known capacity to neutralize SARS-CoV-2
infection. All polyclonal sera similarly detect the RBD protein expressed in both
heterologous expression system (Fig. 8.D,H).

The C-RBD-H6 PP protein was also used to recall *in vitro* a cellular response in terms of IFNγ secretion using ELISpot from PBMCs of COVID-19 naturally infected subjects with at least three months from hospital release. As shown in Fig. 8.I stimulation with 10 µg/mL of the yeast-produced protein induce IFNγ secretion from COVID-19 naturally infected subjects PBMCs.

470

3.7. Functionality of yeast-expressed RBD protein

471 ACE2 receptor binding competition assay

472 The functionality of the recombinant protein C-RBD-H6 PP was confirmed by the 473 assessment of its ability to bind to the chimeric hFc-ACE2 receptor bound to ELISA 474 plates and also to bind to the ACE2 receptor in Vero E6 cells both in solution or 475 attached to the plate. To evaluate the binding of the protein to hFcACE-2, a serial two-476 fold dilution curve of the test protein starting from 100 µg/mL was mixed with HEK293T 477 produced hFc-RBD-HRP conjugate at a fixed concentration and added to hFc-ACE2 478 coated plates. As shown in Fig. 9.A, two independent C-RBD-H6 PP protein 479 preparations effectively displaced the hFc-RBD HEK in the same magnitude that the 480 soluble hFc-ACE2 and the recombinant RBD-H6 HEK protein obtained in HEK293T 481 cells. Two non-related chimeric proteins P64K-VEGFH6 (expressed in *E. coli*), and hFc-482 VEGFR2 (produced in HEK293T cells) were used as negative controls

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483 A similar study was conducted on paraformaldehyde-fixed Vero E6 cells (Fig. 9.B) As 484 expected, no displacement of biotinylated hFc-RBD from the cell membrane was 485 observed in either case using as negative controls the non-related chimeric VEGF 486 protein fused to the N-terminal of *Neisseria meningitidis* (P64K-VEGFH6) or the human 487 Fc (VEGFR2-hFc), and 100 % of the signal was lost when the C-RBD-H6 PP protein, 488 soluble hFc-ACE2 or mFc-ACE2 were added (Fig. 9.C). These results suggest that the 489 yeast-expressed C-RBD-H6 PP protein can bind efficiently to the cell-surface ACE2 490 receptor.

Further characterization of ACE2 binding was conducted in solution using yeast and mammalian-derived proteins coated in different concentrations in ELISA plates. ACE2 EC-50 on RBD coated plates result in values ranging between 100 and 300 ng/mL in agreement with the values reported for commercial sources of RBD and with our data using RBD variants purified from HEK293T supernatant. Using the same system stability of the protein was verified by incubating the protein at 4 °C, 37 °C, and 50 °C for 2 h without significant loss in ACE2 binding activity (data not shown)

These results confirm the specific binding of the yeast-expressed RBD with SARS-CoV-2 receptor ACE2 either cell-bound or soluble, indicating that the antigen herein obtained is functional.

501 Yeast-expressed RBD protein elicited RBD-ACE2 receptor binding inhibition and SARS502 CoV-2 neutralizing antibodies in rodents and NHP

503 To evaluate the immunogenicity of C-RBD-H6 PP protein, BALB/c mice were 504 immunized thrice at 7 days intervals by the intraperitoneal route with 50µg of the 505 protein. The IgG antibody response in mice at days 21 and 35 shows that the C-RBD-506 H6 PP protein was able to induce antibody responses in ELISA coated with the RBD 507 protein produced in mammalian HEK293T cells. Seroconversion was achieved a week 508 after the first boost (Fig. 10.A). The administration of a second booster at day 21 509 significantly increases both RBD-specific IgG titers and the inhibitory potential of the 510 sera to reduce the protein binding to its cognate receptor ACE2 a fact that correlates 511 with live SARS-CoV-2 neutralization (Fig. 10.B,C).

The C-RBD-H6 PP protein was also tested in SD rats using intramuscular administration of 9 µg every 7 days for 10 weeks. Assessment of IgG and RBD-ACE2 receptor binding inhibition three days after the last immunization indicates high RBDspecific antibody titers that correlate with the inhibitory titer. The former also display a high correlation with the neutralization titer of live SARS-CoV-2 virus in Vero E6 cells (Fig. 10.D,F).

518 The C-RBD-H6 PP protein evaluation in NHP using short intramuscular administration 519 schedule on days 0-14-28, and two dose levels indicate a dose-response effect with 520 seroconversion of 83 % (5 of 6 animals) and 100% (the 10 animals) for the 50 µg and 521 100 µg dose respectively after the first booster and a 100 % seroconversion in both 522 schedules after the second booster. Total IgG titers increased to 44,240 StdU/mL and 523 62,435 StdU/mL respectively for monkeys included in the low and high-dose groups. In 524 both cases, the titers were significantly higher than those detected in the convalescent 525 panel of sera. The geometric median ACE2 binding inhibition titer for 50 µg dose was 526 1:230, while a significantly higher value of 1:705 was detected for the animals receiving 527 the 100 µg dose. Both values were higher than those detected in a panel of COVID-19

528 convalescent subjects (p<0.001, Kruskal-Walis). The higher titers for the 100 µg dose 529 correlate with and enhancement in the inhibition of RBD-ACE2 binding for this group. 530 The analysis of the neutralization titer of live SARS-CoV-2 virus in Vero E6 cells 531 corroborates these findings, indicating that 50µg dose is sufficient to induce ACE2 532 binding inhibition titers with an EC50 geometric mean of 1:66 serum dilution, a value 6 533 times higher than the reported for the convalescent panel. A significant increase to 534 1:141 in this parameter was detected for the 100 µg dose pointing to a dose-related 535 effect (Fig. 10.G,J).

536 The specific IgG antibody and the inhibition of RBD-ACE2 binding were only detected in 537 C-RBD-H6 PP protein-inoculated animals but not in control animals. The immune 538 responses evidenced the boosting effect and the specific dose-dependent effect, as 539 compare to negative results for control animals.

540 *Cellular response*

541 Cellular response recall was evaluated three months after the last immunization in 542 BALB/c mice receiving subcutaneous doses of 25 µg in alum in a 0-14-35 days 543 schedule. The evaluation of the presence of a memory response in the animal's spleens 544 indicates a significant induction of clones secreting IFNy in response to incubation with 545 the C-RBD-H6 PP antigen. Furthermore, the analyses of the supernatant of the recall 546 reaction point to the predominant induction of IFNy followed by IL-2, IL-6 and to a lower 547 extend TNF α and IL-4 (Fig. 11). Our findings in the systemic compartment (Fig. 11.A) 548 were similar to those found for CD3+ cells enriched from mice lungs (Fig. 11.B),

549 indicating that the response can also be recalled in the organ primarily affected by 550 SARS-CoV-2 infection.

551 **4. Discussion**

552 Despite the impressive development of prophylactic vaccines against COVID-19 and 553 that several vaccines have reached the Emergency Use Authorization or 554 Pharmaceutical Registry, there is not a sufficient supply of vaccines. In addition, the 555 challenge of the continuous emergence of new mutant strains of the virus will require 556 updating the antigen sequences included in the vaccines and administer booster doses 557 to maintain the immunity of the population.

558 Results from this study demonstrate that it is possible to develop a vaccine candidate 559 expressing RBD protein of SARS-CoV-2 using the yeast *P. pastoris*. The RBD was 560 selected base on the state of the art evidence of the main contribution of the epitopes of 561 RBD to the neutralization activity of the sera [13;26-28]

562 Moreover neutralizing antibody responses generated after immunization with Pfizer-563 BioNtech or Moderna vaccines of previously infected subjects, are mainly due to anti-564 RBD antibodies. The sera depletion of antibodies targeting the RBD abrogates sera 565 neutralization capacity [29]

566 Optimal conditions for the production of a recombinant protein in *P. pastoris* expression 567 system differ according to the target protein. Indeed, we were able to obtain 30-40 mg/L 568 of the RBD with more than 98% of purity, close to the yield obtained in previous report 569 by Arbeitman C.R., et al [8], an essential condition to develop a vaccine candidate. In 570 addition to the protein yield, it is important the sugar composition. In *P. pastoris* the 571 glycosylation pattern is characterized by the high mannose content. Mannosylation 572 enhances activation of antigen-presenting cells like macrophages and dendritic cells, 573 functioning as immunopotentiator while increases the antigen immunogenicity 574 compared with its nonglycosylated counterparts [30]. The ionic interactions dependent 575 on sugar composition also could help to stabilize RBD structure.

576 Our results on the affinity of the *P. pastoris* recombinant protein C-RBD-H6 PP by the 577 receptor comply with those of other authors performing binding assays in BIACORE for 578 the interaction pair RBD-ACE2 [31-33]. It means that probably the subunit vaccine 579 based on C-RBD-H6 PP may elicit an effective antibody response against SARS-CoV-580 2, avoiding the virus entry and replication by blocking RBD-ACE2 interaction in real 581 SARS-CoV-2 infection context.

The correct glycosylated protein during the transit in the endoplasmic reticulum and Golgi apparatus, go to the secretory pathway. If the protein is unable to fold properly is degraded in the cytosol. This mechanism guarantees that only properly folded proteins are secreted [8;34]. Our results demonstrated the presence of four intramolecular disulfide bonds identical to those present in the native RBD of SARS-CoV-2. These results as well as the secondary structure determined by CD spectroscopy confirm the right folding of the C-RBD-H6 PP.

Remarkably despite the known difference in the sugar composition of the protein expressed in *P. pastoris* compared to the expression systems in mammals, the antigenicity of the protein is similar both with polyclonal sera from mice and monkeys immunized with the recombinant protein, such as with human sera from individuals with natural coronavirus infection or vaccinated with the Pfizer-BioNTech or Sputnik V

594 vaccines against SARS-CoV-2. The C-RBD-H6 PP protein also stimulated cellular 595 response mediated by IFNy secretion in lymphocytes isolated from convalescent 596 subjects. The C-RBD-H6 PP protein also inhibited the RBD binding to the ACE2 597 receptor in a competitive ELISA regardless of whether the RBD receptor was obtained 598 recombinant or it is found directly in the membrane of Vero E6 cells. In the same 599 manner, the sera from mice, rats and NHP immunized with C-RBD-H6 PP protein 600 inhibited the RBD-ACE2 receptor binding and neutralize the SARS-CoV-2 in 601 microneutralization tests.

The need for the accelerated development of the vaccine candidate during the pandemic, led to the neutralizing activity of the NHP sera being evaluated just one week after the third dose, reaching titers comparable to the panel of convalescent sera used as a control. It is known that a long time for bleeding would lead to the maturation and selection of B cell clones producing high avidity antibodies and consequently higher neutralizing titers.

608 Conclusions

Optimal conditions for the production of a recombinant protein in *P. pastoris* expression
system differ according to the target protein. Indeed, we were able to obtain 30-40 mg/L
of the RBD with more than 98% of purity.

Our ESI-MS results demonstrated the presence of four intramolecular disulfide bonds identical to those present in the native RBD of SARS-CoV-2, and the CD spectrum of the protein indicates the presence of well-packed aromatic and cysteine residues. These results together with the high receptor binding affinity in BIACORE assays confirm the right folding of the C-RBD-H6 PP protein.

The C-RBD-H6 PP antigenicity and the capacity of the sera from immunized mice, rats and NHP to inhibited the RBD-ACE2 receptor binding and neutralize the live virus infection to Vero E6 cells, points to the feasibility of the protein as a vaccine candidate.

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627 Authors Contributions

628 GGN: provided original ideas and study concept and design, data curation, analysis and 629 interpretation of data, and drafting, review and editing of the final version of the 630 manuscript and studies supervision. MLF: contributed to analysis and interpretation of 631 data, study design, review of the manuscript and studies supervision. LJGL: contributed 632 to ESI-MS study design, acquisition of data, analysis and interpretation of data, and 633 drafting and review of the manuscript. LAER, IAM and YRG: performed the ESI-MS 634 studies, acquisition of data, analysis and interpretation of data. GCH: performed 635 alycosylation studies. ACR: performed drafting and execution of Biacore studies. GMP: 636 performed the protein purification studies. MPI and JZS: performed fermentation 637 studies. GCS: provided original idea and drafting of the genetic construction, and 638 performed structural analysis experiments by CD spectroscopy. AMMD: provided 639 original ideas and performed the genetic construction and review of the manuscript, and

640 DGR: performed the genetic construction and protein expression experiments. MBR: 641 provided original ideas, study designs, analysis and interpretation of data, drafting and 642 review of the manuscript, graphic and statistical processing and execution of 643 antigenicity, immunogenicity and cellular response studies. IGM and CCHA: performed 644 antigenicity, immunogenicity studies and analysis of cellular response in mouse, rat, 645 and NHP. OCS: study design and supervision of the microneutralization experiments. 646 GLP: contributed to analytical procedures and donor patients selection and evaluation 647 of the immunological and functional response. JVH, EMD, EPV and MAA: study 648 concept and supervision.

649 Disclosure

- 650 MLF, MBR, AMMD, DGR, ACR, GCS, GMP, EPV, MAA and GGN are co-authors of the
- 651 Center for Genetic Engineering and Biotechnology patent application comprising the C-
- 652 RBD-H6 PP protein as a vaccine antigen against SASR-CoV-2.
- All authors approved integrally the final article.

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mFc-RBD and hFc-RBD chimeric proteins.

659

660 Figure Legends

Fig. 1. Analysis of culture supernatants from randomly selected RBD-secreting *P. pastoris* clones. In all cases 10 μL of each sample were analyzed under reducing conditions. Panel A: Coomassie Blue-stained 12.5 % SDS-PAGE; panel B: Western blotting of an identical gel with an anti-His tail antibody. Lane V: Strain transformed with the empty vector. Lanes A, B and C: clones transformed with the RBD plasmid.

Fig. 2. RP-HPLC. Analysis of C-RBD-H6 PP protein produced in *Pichia pastoris*, with
98.6 % of purity; analyzed on a reversed phase C8 Vydac analytical column. The
gradient is shown with a blue line.

Fig. 3. Protein electrophoresis. Coomassie Blue stained 12,5 % SDS-PAGE gel of 10
µg of the purified C-RBD-H6 PP under reducing conditions. Lane 1: C-RBD-H6 PP
protein; Lane 2 Molecular weight markers.

672 Fig. 4. ESI-MS spectra. (A) Multiply-charged and (B) deconvoluted ESI-MS spectra of 673 C-RBD-H6 PP protein previously N-deglycosylated with PNGase-F. The parenthesis in 674 (B) shows the expected molecular mass considering the presence of four intramolecular disulfide bonds and the Asn³³¹ and Asn³⁴³ transformed into Asp residues by the 675 676 PNGase F during the deglycosylation step. (C) ESI-MS spectrum of the tryptic peptides 677 generated by an in-solution buffer-free trypsin digestion protocol of the N-678 deglycosylated C-RBD-H6 PP protein. The star in (B) and (C) indicates the low-679 abundance species that do not contain the four residues (-NWSF) located at the N-680 terminal end. Nt and Ct-His₆ represents the N-terminal end and the His₆-tag C-terminal 681 end peptides of C-RBD-H6 PP protein, respectively.

682 Fig. 5. SPR analysis from C-RBD-H6 PP protein obtained in *P. pastoris* interacting 683 with mFc-ACE2 receptor in a single-cycle BIACORE experiment. (A) Sensorgrams 684 corresponding to one of the replicates of the protein dissolved in PBS, pH = 7.2, (B) 685 Non-related protein (also expressed in yeast), used as negative control for the 686 interaction with immobilized mFc-ACE2 cell receptor. As expected, there were no 687 significant signals for the protein used as negative control; compared to the curves 688 obtained for C-RBD-H6 PP protein in the same experimental conditions (similar axis scale is shown). 689

690 Fig. 6. Far UV CD spectrum of the C-RBD-H6 PP protein.

Fig. 7. Near UV CD spectrum of the C-RBD-H6 PP protein. Bands at 263, 269, 277,
281 and 299 nm, indicate the presence of well packed aromatic and cystine residues.

693 Fig. 8. Antigenicity of the RBD protein produced in P. pastoris yeast. C-RBD-H6 694 PP (upper panel) or RBD-H6 HEK (lower panel) proteins were used for coating ELISA 695 plates and for animal immunization. Sera and monoclonal antibodies were used in serial 696 two-fold dilutions. (A,E). SS-1, SS4, SS-7 and SS-8 monoclonal antibodies; (B,F). 697 COVID-19 sera from convalescents; (C,G) Sera from subjects immunized with Pfizer-698 BioNTech (red squares) and Sputnik vaccines (black triangles); (D,H). Mice (grey) and 699 NHP (blue) sera; (I). C-RBD-H6 PP stimulates INFy secretion in CD3+ cells from 700 naturally infected individuals. Wilcoxon matched paired test.

Fig. 9. Binding inhibition capacity of two independent protein preparations of the
C-RBD-H6 PP (L04 and L07). (A). The C-RBD-H6 PP protein displaced the hFc-RBDHRP from the hFc-ACE2 coated ELISA plates, and (B) from the ACE2 receptor in Vero
E6 cells. (C). The C-RBD-H6 PP protein was recognized by the soluble ACE2 receptor.

P64K-VEGFH6 and hFc-VEGFR2 non-related chimeric proteins were used as negativecontrols.

707 Fig. 10. Immunogenicity of C-RBD-H6 PP protein in three animal species using 708 different immunization routes and schedules. (A) Evaluation of RBD-specific IgG in 709 BALB/c mice 7 days after second and third intra-peritoneal immunization (n=10); (B). 710 Evaluation in BALB/c mice of EC50 for the ACE2 binding inhibition and PRNT in the 711 microneutralization assay (n=10): C. Correlation analyses of the ACE2 binding inhibition 712 and microneutralization tests in mice (C). Sperman, r=0.6482, p=0.0478); (D). 713 Evaluation of RBD-specific IgG in SD rats 3 days after 10 weakly subcutaneous 714 immunization (n=10); (E). Evaluation in SD rats of EC50 for the ACE2 binding inhibition 715 and PRNT in the microneutralization assay (n=20); (F). Correlation analyses of the 716 ACE2 binding inhibition and microneutralization tests in rats (Spearman, r=0.9233, p=<717 0.0001); (G). Evaluation of RBD specific IgG in NHP with 50 µg and 100 µg dose 14 718 days after three subcutaneous immunizations every second week with 6 and 10 animals 719 respectively; (H). Evaluation in NHP of EC50 for the ACE2 binding inhibition; (I). 720 Evaluation in NHP of EC50 for the PRNT in the microneutralization assay. (J). 721 Association Correlation analyses of the Inhibition and microneutralization tests in NHP 722 (Spearman, r=0.8994, p<0.0001).

723 Fig. 11. Heatmap of the cytokine response. Splenocytes (A) and Lung CD3+

enriched cells (B), after restimulation with C-RBD-H6 PP. Cells were obtained three
months after the last immunization, from 4 to 5 mice per group that received three
subcutaneous 25 µg doses. Non-stimulated controls were subtracted from re-stimulated
samples.

- 728 **Table 1.** Summary for the sequence verification of the N-deglycosylated C-RBD-H6 PP
- considering the ESI-MS analysis of tryptic peptides generated by the in-solution buffer-
- 730 free digestion.
- 731

Code ^{a)}	<i>m/z</i> _{Theor}	z	<i>m/z</i> _{Exp}	Assignment
C ³³⁶ - C ³⁶¹ , Nt	1399.64	4	1399.63	NWSFFSNIGGSSGGS- ³³¹ DITNLCPFGEVFDATR ^{346 b)}
				/
				³⁵⁸ ISNCVADYSVLYNSASFSTFK ³⁷⁸
				(Native C^{336} - C^{361} , N-terminal end, Asn ³³¹ and Asn ³⁴³ \rightarrow Asp)
	1266.09	4	1266.07	FSNIGGSSGGS- ³³¹ DITNLCPFGEVFDATR ^{346 b)}
C ³³⁶ -				/
C ³⁶¹ , Nt- NWSF				³⁵⁸ ISNCVADYSVLYNSASFSTFK ³⁷⁸
				(Native C^{336} - C^{361} , N-terminal end -NWSF, Asn ³³¹ and Asn ³⁴³ \rightarrow Asp corresponding to the two N-glycosylation sites)
F ³⁴⁷ -R ³⁵⁵	557.28	2	557.26	³⁴⁷ FASVYAWNR ³⁵⁵
K ³⁵⁶ - R ³⁵⁷	303.21	1	303.20	³⁵⁶ KR ³⁵⁷
				³⁷⁹ CYGVSPTK ³⁸⁶
C ³⁷⁹ -	1020.81	3	1020.79	
C ⁴³²	765.86	4	765.85	⁴²⁵ LPDDFTGCVIAWNSNNLDSK ⁴⁴⁴
				(Native C ³⁷⁹ -C ⁴³²)
C ³⁷⁹ *- C ⁴³²	982.71	4	982.69	⁶⁴ CYGVSPTK ⁷¹
				I
				⁴¹⁸ IADYNYKLPDDFTGCVIAWNSNNLDSK ⁴⁴⁴
				(Native C ³⁷⁹ -C ⁴³² , 1 missed cleavage)
C ³⁹¹ - C ⁵²⁵	992.52	4	992.47	³⁸⁷ LNDLCFTNVYADSFVIR ⁴⁰³
				I
				⁵¹⁰ VVVLSFELLHAPATVCGPK ⁵²⁸
				(Native C ³⁹¹ -C ⁵²⁵)
G ⁴⁰⁴ - R ⁴⁰⁸	575.28	1	575.27	⁴⁰⁴ GDEVR ⁴⁰⁸
Q ⁴⁰⁹ - K ⁴¹⁷	450.25	2	450.24	⁴⁰⁹ QIAPGQTGK ⁴¹⁷
I ⁴¹⁸ -K ⁴²⁴	443.72	2	443.71	⁴¹⁸ IADYNYK ⁴²⁴

V ⁴⁴⁵ - R ⁴⁵⁴	609.80	2	609.78	⁴⁴⁵ VGGNYNYLYR ⁴⁵⁴
L ⁴⁵⁵ -R ⁴⁵⁷	435.27	1	435.26	⁴⁵⁵ LFR ⁴⁵⁷
S ⁴⁵⁹ - R ⁴⁶⁶	495.77	2	495.76	⁴⁵⁹ SNLKPFER ⁴⁶⁶
C ⁴⁸⁰ - C ⁴⁸⁸	1589.38	3	1589.35	⁴⁶⁷ DISTEIYQAGSTPCNGVEGFNCYFPLQSYGFQPTNGVGYQP
	1192.29	4	1192.26	(Native C ⁴⁸⁰ -C ⁴⁸⁸)
Ct*-His ₆	799.68	3	799.67	⁵²⁹ K-GGSGGSSSSSSSSSSSIEHHHHHH ^{b)}
	600.01	4	600.00	(C-terminal end, 1 missed cleavage)
Ct-His ₆	756.98	3	756.97	GGSGGSSSSSSSSSIEHHHHHH ^{b)}
	567.99	4	567.98	(C-terminal end)

732

733 Nt: N-terminal end, Ct-His₆: His-tag C-terminal end. $C^{\#}-C^{\#}$ corresponds to tryptic 734 peptides linked either by intermolecular disulfide bonds or a tryptic peptide that contains 735 an intramolecular disulfide bond in its structures.

736 m/z_{calc} correspond to the calculated m/z values for all tryptic peptides generated by the

in-solution buffer-free digestion of the N-deglycosylated protein.

738 m/z_{exp} correspond to the experimental m/z values for all tryptic peptides observed in the

739 ESI-MS analysis shown in Fig. 4.C.

Regions of the sequence written in italic do not correspond to the RBD of SRAS-CoV-2 and were inserted in the cloning stage, while underlined residues indicate the conversion of N-glycosylated asparagines (Asn^{331} and Asn^{343}) into aspartic acid residues by the action of PNGase-F (Asn^{331} and Asn^{343} ->Asp).

744

745 **Table 2.** Secondary structure content of the protein estimated by CD (BeStSel) and 3D

746 coordinates (DSSP)

747

	Secondary Structure Content, %			
Secondary Structure Element	Method			
	BeStSel	DSSP		
Helix	7.9	9.3		
Beta antiparallel	28.7	22.4		
Beta parallel	0.0	0.0		
Turn	12.9	22.4		
others	50.5	45.9		
Helix1 (regular)	3.3	-		
Helix2 (distorted)	4.6	-		
Beta Antiparallel_1 (left-handed)	0.9	-		
Beta Antiparallel_2 (relaxed)	12.0	-		
Beta Antiparallel_3 (right-handed)	15.8	-		

748

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866

Α

V A B C



V A B C



20 kDa \rightarrow

В











-O-C-RBD-H6 PP













3.0







