

1 Long-term immune protection against SARS-CoV-2 escape variants upon a single 2 vaccination with murine cytomegalovirus expressing the spike protein

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18 Abstract

19 **Vaccines are central to controlling the coronavirus disease 2019 (COVID-19) pandemic but the durability**
20 **of protection is limited for currently approved COVID-19 vaccines. Further, the emergence of variants**
21 **of concern (VoCs) that evade immune recognition has reduced vaccine effectiveness, compounding the**
22 **problem. Here, we show that a single dose of a murine cytomegalovirus (MCMV)-based vaccine, which**
23 **expresses the spike (S) protein of the virus circulating early in the pandemic (MCMV^S), protects highly**
24 **susceptible K18-hACE2 mice from clinical symptoms and death upon challenge with a lethal dose of**
25 **D614G SARS-CoV-2. Moreover, MCMV^S vaccination controlled two immune-evading VoCs, the Beta**
26 **(B.1.135) and the Omicron (BA.1) variants in BALB/c mice, and S-specific immunity was maintained for**
27 **at least 5 months after immunization, where neutralizing titers against all tested VoCs were higher at**
28 **5-months than at 1-month post-vaccination. Thus, cytomegalovirus (CMV)-based vector vaccines might**
29 **allow for long-term protection against COVID-19.**

30 Introduction

31 The severe acute respiratory coronavirus-2 (SARS-CoV-2) emerged in Hubei province, China, in 2019 ¹.
32 Since then, the virus has spread worldwide and caused the ongoing COVID-19 pandemic. Although SARS-
33 CoV-2 was genetically largely stable during the initial months of the pandemic, several variants of concern
34 (VoCs) have emerged in the meantime and present an increased threat to global public health ² due to
35 their augmented transmissibility and/or antibody escape ^{3,4}.

36 In the European Union, six vaccines are currently approved for administration in humans ⁵, including two
37 mRNA-based vaccines, two vector-based vaccines, one adjuvanted protein vaccine, and an inactivated

38 virus vaccine. All authorized vaccines induce humoral immune responses against the spike (S) glycoprotein
39 of SARS-CoV-2, which drives viral entry into cells and is the key target for neutralizing antibodies. However,
40 vaccine-induced immunity wanes within months, resulting in reduced protection against infections and
41 disease⁶⁻⁸. Consequently, booster shots were recommended, but they also do not confer long-term
42 protection against infection and disease, especially in light of novel immune-evasive VoCs, such as the
43 Beta variant⁸, or the currently dominating Omicron VoC (Pango lineage B.1.1.529)⁹. Therefore, the need
44 for a vaccine that confers long-lasting and broad immune protection remains unmet.

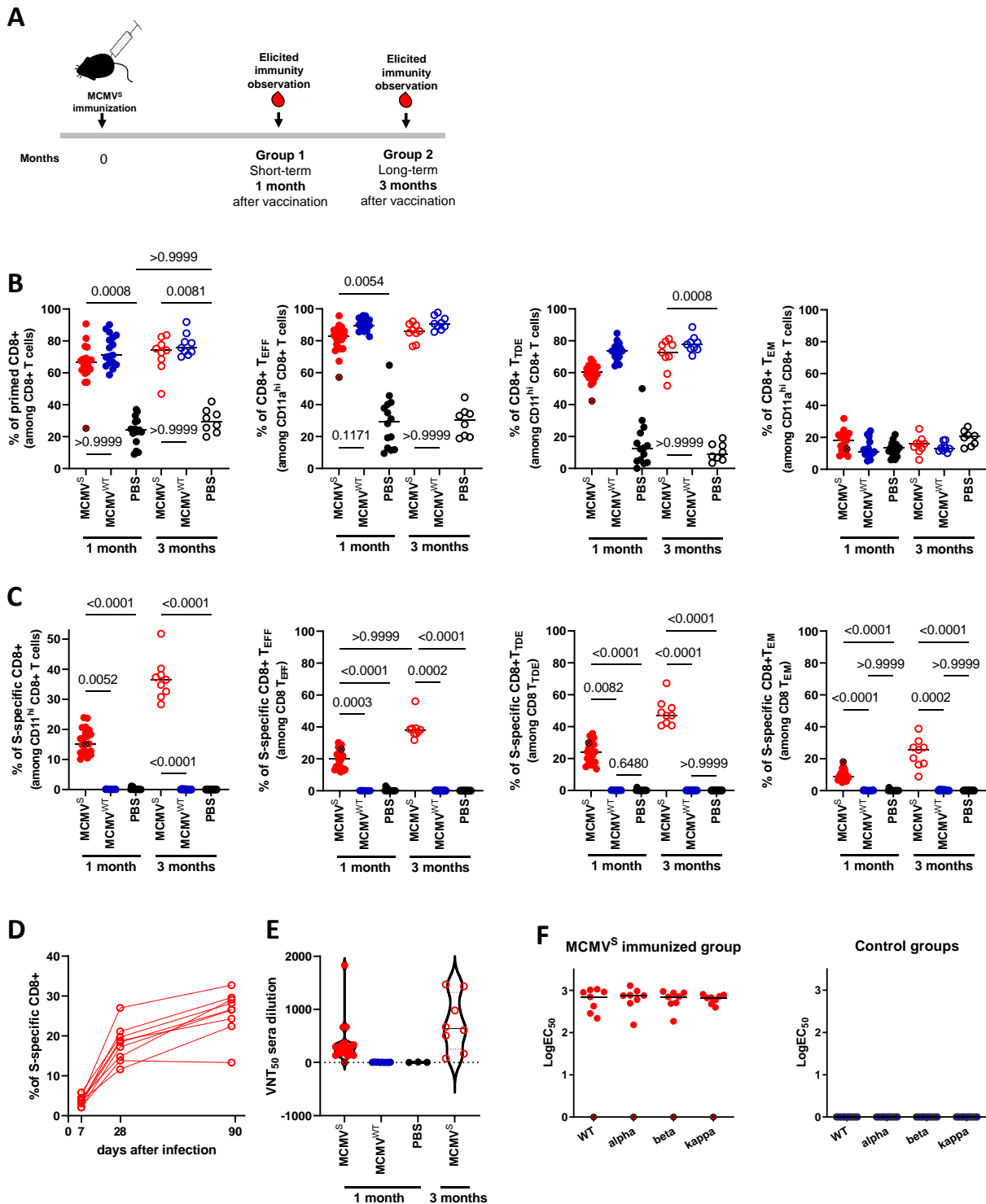
45 Murine cytomegalovirus (MCMV) belongs to the *betaherpesvirinae* subfamily and is a well-studied model
46 virus for human CMV (HCMV) infection¹⁰. CMVs induce uniquely robust anti-viral immunity, which persists
47 over a lifetime^{11,12}. This long-term maintenance of antigen-specific lymphocytes, known as memory
48 inflation^{13,14}, is characterized by an accumulation and persistence of CD8 T cells with an effector memory
49 (T_{EM}) phenotype, recognizing immunodominant viral antigens^{13,15}. Therefore, CMV has been explored as
50 a vaccine vector that provides long-term immune protection against many infectious targets, based on a
51 peculiarly strong CD8 T cell immunity¹⁶⁻²¹. MCMV infection also results in lasting humoral immune
52 responses toward viral antigens²², which can be exploited to retarget immune protection against other
53 viruses²³. Mouse models of SARS-CoV-2 infection for preclinical vaccine efficacy testing were initially
54 restricted to transgenic mice expressing the human ACE2 receptor (hACE2), or mouse-adapted SARS-CoV-
55 2 variants because the original strains of SARS-CoV-2 could not bind to mouse ACE2 (mACE2) receptors.
56 However, the N501Y mutation in the S protein observed in various VoCs, including Beta and Omicron
57 VoCs, allows mACE2 binding²⁴ and wild type (WT) mice infection, enabling the analysis of SARS-CoV-2
58 infection in the absence of exogenous receptor expression²⁵.

59 Recently, our group has demonstrated that an MCMV-based COVID-19 vaccine candidate encoding the S
60 protein from the Wuhan prototype (MCMV^S) induced long-lasting humoral and cellular immunity²⁶. Here,
61 we show that immunization with MCMV^S protects vaccinated K18-hACE2 mice against a lethal dose of the
62 SARS-CoV-2 D614G strain. Furthermore, the same vaccination protects BALB/c mice against viral challenge
63 with the immune-evasive Beta and Omicron strains, and the protection persists for at least 5 months with
64 no signs of immune waning because immunity, particularly virus neutralizing titer (VNT), at 5 months was
65 even stronger than at 5 weeks post-vaccination. Hence, our results argue that CMV-based SARS-CoV-2
66 vaccines may provide long-term and broad protection against multiple VOCs.

67 Results

68 Immune responses of MCMV^S-vaccinated K18-hACE2 mice

69 We have previously shown that MCMV^S immunization of BALB/c and C57BL/6 mice elicits a strong
70 humoral response as well as an inflationary CD8 T cell response towards the S antigen, respectively²⁶.
71 Here, we immunized K18-hACE2 mice intraperitoneally (i.p.) with either MCMV^S, MCMV^{WT}, or PBS and
72 challenged them one or three months later with a lethal dose of a SARS-CoV-2 D614G strain (**Fig. 1A**).
73 Mice were bled 7 days before the SARS-CoV-2 challenge, and blood leukocytes were analyzed by flow
74 cytometry (gating strategy shown in **Supplementary Fig. 1**). MCMV^S and MCMV^{WT} elicited comparable
75 levels of CD8 T cells in total, primed, effector (T_{EFF}), terminally differentiated effector (T_{TDE}), effector
76 memory (T_{EM}), and central memory (T_{CM}) compartments in short-term (1 month) and long-term (3 months)
77 groups (**Fig. 1B** and **supplementary Fig. 2**). We noticed that a single mouse in the short-term group
78 exhibited a very low frequency of primed CD8 T cells, possibly an indication of a technical failure. This
79 animal is shown for transparency and was marked with an X symbol (**Fig. 1B**, X -marked symbols).



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Figure 1. Immune status of MCMV^S-vaccinated K18-hACE2 mice

(A) Schematic image of the experimental setup. (B) CD3+CD8+CD4- T cells were gated for the primed subpopulation (CD11^{hi}CD44^{hi} – leftmost panel). Primed CD8 T cells were progressively gated into T_{EFF} (CD62L^{lo}), T_{TDE} (CD62L^{lo}KLRG1^{hi}), or T_{EM} (CD62L^{lo}KLRG1^{lo}) subpopulations. Frequencies of cells in each subset as a fraction of the parental population are shown. Horizontal lines show the median. (C) The percentages of S-specific cells in each subset shown in B are shown. (B,C) Each symbol is an individual animal; the MCMV^S-vaccinated animal with poor immune responses is marked with an X. Horizontal lines denote medians. Kruskal-Wallis tests were used for the statistical analyses (n=21 MCMV^S, n=15 MCMV^{WT}, PBS for 1-month; n=9 MCMV^S, MCMV^{WT}, n=8 PBS for 3-

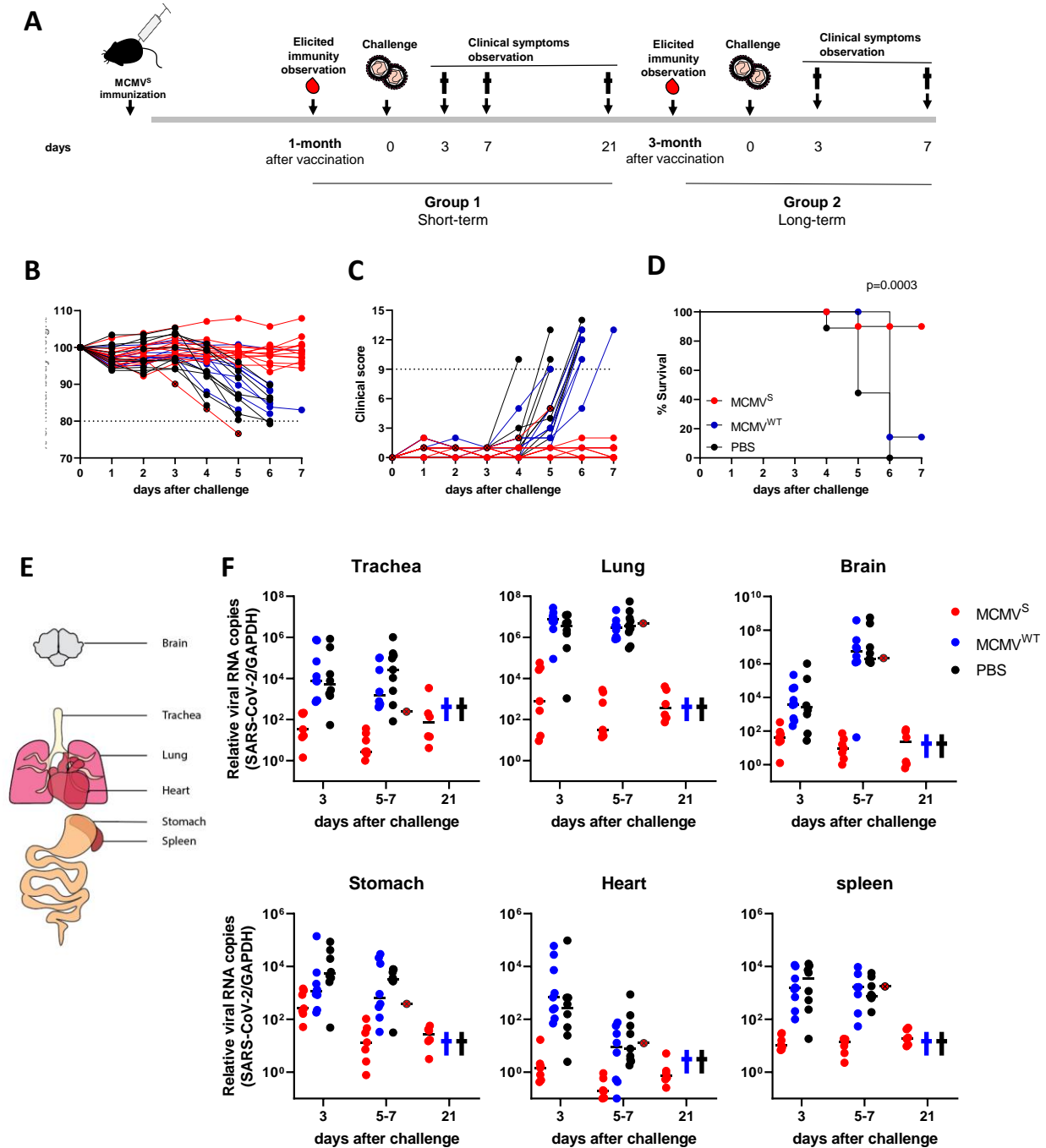
88 months). (D) Antigen-specific memory responses over time in blood shown as the percentage of antigen-specific cells within the
89 total CD8 compartment. Each line connects values from an individual mouse at indicated time points. (E) Violin plots of VNTs that
90 resulted in a 50% of reduction of SARS-CoV-2 infection (VNT_{50}). Each symbol indicates an individual mouse. (F) EC_{50} of the IgG
91 fraction specific for the respective S antigens from indicated variants measured by ELISA. Each symbol indicates an individual
92 mouse and each color represents vaccination with following vectors: Red=MCMV^S, Blue=MCMV^{WT}, and Black=PBS. Horizontal lines
93 show geometric means. One MCMV^S-vaccinated animal with poor immune responses is marked with an X throughout.

94 Importantly, only MCMV^S-vaccinated K18-hACE2 mice displayed high frequencies of CD8 cells recognizing
95 the antigenic epitope VNFNGL of the S protein, while control groups showed no S-specific T-cell
96 immunity (**Fig. 1B** and **C**). The long-term cohort showed a ~ two-fold increase of S-specific T-cells in all
97 examined subsets over the short-term cohort (**Fig. 1C**), and dynamic monitoring of S-specific CD8 T cells
98 in the peripheral blood confirmed a strong inflationary response at single animal level (**Fig. 1D**), in line
99 with our previous report in C57BL/6 mice²⁶. We also observed an increase in average neutralizing
100 antibody titers in the sera of long-term MCMV^S-vaccinated animals (**Fig. 1E**). Vaccinated mice showed high
101 titers of anti-S IgGs, which recognized various variants of interest and concern. Sera of mice mock-
102 immunized with MCMV^{WT} or PBS did not show any S-specific antibody response (**Fig. 1E** and **F**), arguing
103 strongly that the responses were not due to cross-reactions against potential MCMV antigens. Overall,
104 these results indicated that MCMV^S vaccination induces potent and lasting adaptive immunity towards
105 the S antigen.

106 Protection against SARS-CoV-2 challenge after a single vaccine dose

107 A week after the analysis shown in **Fig. 1**, short-term vaccinated K18-hACE2 mice were challenged with
108 2×10^3 PFU of SARS-CoV-2 (B.1 strain, D614G mutation), and daily monitored for weight loss and disease
109 indicators (**Fig. 2A**). All MCMV^S-immunized animals that robustly responded to vaccination (**Fig. 1B** and
110 **1F**) were protected from clinical signs and weight loss, while mock-vaccinated mice became severely ill,
111 reaching the humane end-point by day 5 post-infection or earlier (**Fig. 2B** and **2C**). MCMV^S-immunized
112 mice survived the lethal SARS-CoV-2 challenge, even when monitored for 3 weeks (**Fig. 2D** and
113 **Supplementary Fig. 3**).

114 SARS-CoV-2 can infect a broad range of tissues in K18-hACE2 mice as well as in humans^{27,28}. Therefore,
115 we tested whether MCMV^S vaccination decreases the SARS-CoV-2 viral load in the trachea, lungs,
116 stomach, brain, heart, and spleen (**Fig. 2E**). In all organs, MCMV^S-vaccinated animals showed reduced
117 SARS-CoV-2 viral RNA load for up to 3 weeks, while mock-vaccinated groups exhibited high viral RNA copy
118 numbers (**Fig. 2F** and **Supplementary Fig.4**). In particular, MCMV^S vaccination reduced viral load in the
119 trachea and lungs by roughly a 1,000-fold (**Fig. 2F**, trachea and lungs) and also markedly reduced viral
120 loads in the CNS (**Fig. 2F**, brain), which is known to be targeted by the virus in K18-hACE2 mice^{27,28}. The
121 viral burden in other organs showed a similar pattern with a strong decrease of detectable SARS-CoV-2
122 RNAs in MCMV^S-immunized animals at 7 days post-infection (dpi) in comparison with the control groups,
123 where viral RNA persisted until 7 dpi (**Fig. 2F**, stomach, heart, and spleen). To sum up, MCMV^S
124 immunization protected animals from developing clinical signs and high viral loads in all organs tested.



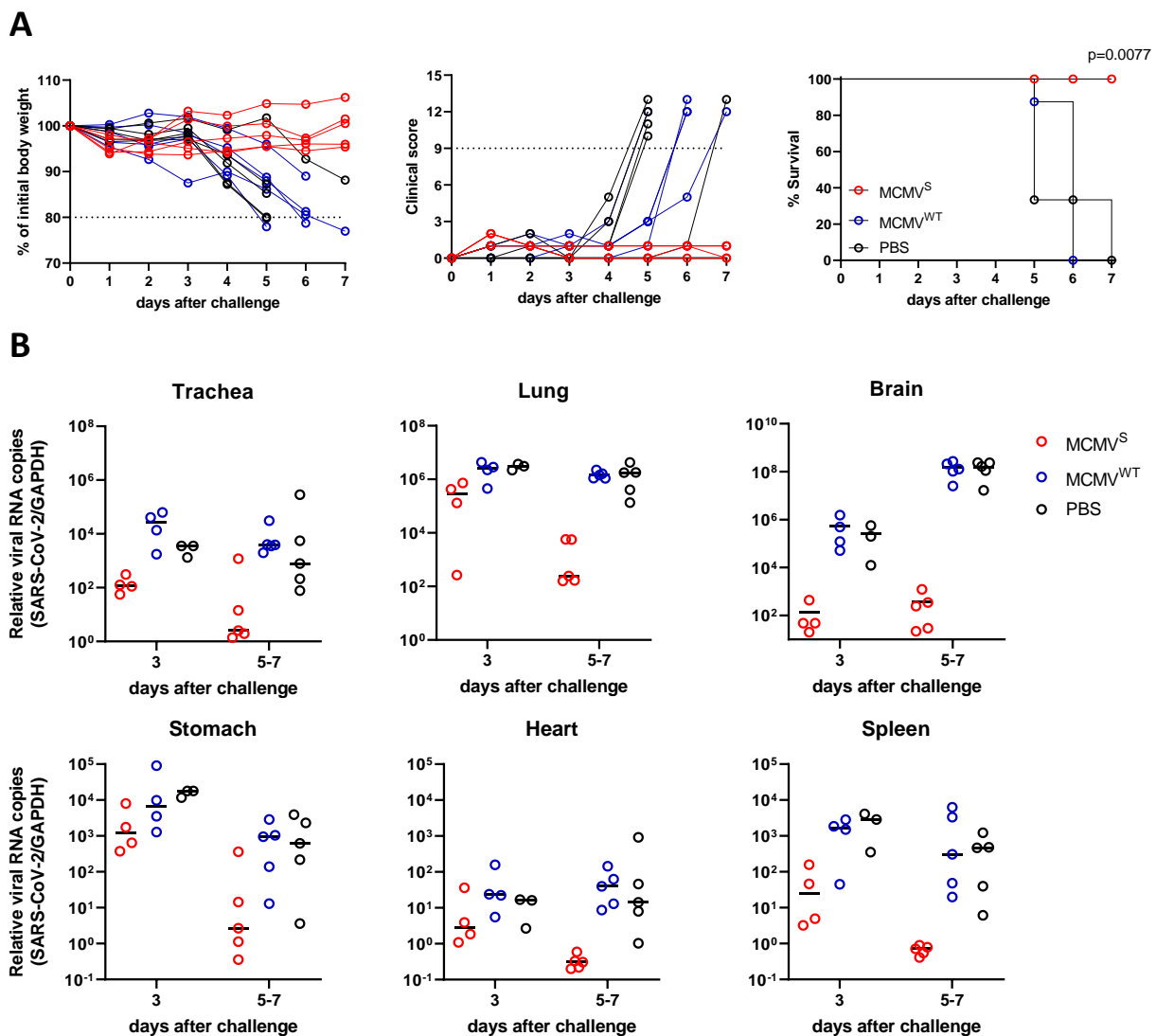
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126 **Figure 2. Short-term protection elicited by MCMV^S vaccination**

127 (A) Schematic image of experimental setup. (B) Percentage of body weight and (C) daily clinical scores upon challenge are shown.
 128 The dotted line indicates the threshold in score resulting in a humane end-point. (D) Mock-survival kinetics of challenged mice
 129 representing the % of mice reaching a humane end point. A log-rank (Mantel-cox) test was used for the statistical analysis. (E)
 130 Schematic illustration of the organs that were collected for virological analyses. (F) Relative viral RNA loads in the representative
 131 organs at the indicated time points normalized to the housekeeping gene *mGAPDH*. Cross symbols indicate no animals alive by
 132 the time of analyses. Horizontal lines indicate medians. The outlier animal with poor immunity is marked with an X-marked red
 133 dot throughout. Organs of mock-immunized mice that reached humane end-points before 7 days upon challenge were harvested
 134 on the day of euthanasia. Pooled data (n=6-8 per group) from two independent experiments are shown (except for the 21 days
 135 data set, which was generated once).

136 Protection against SARS-CoV-2 challenge three months after vaccination

137 The sera/plasma of convalescent COVID-19 patients contains anti-S antibodies, which begin to diminish
 138 around 28 days after infection^{29,30}. Similarly, clinically approved vaccines elicit immunity that wanes over
 139 time, and hence, durable immunity elicited by vaccines against COVID-19 is an unmet need. To define the
 140 long-term immune protection upon MCMV^S vaccination, K18-hACE2 mice were challenged with SARS-
 141 CoV-2 D614G strain three months after vaccination and daily monitored for clinical scores (**Fig. 2A**). All
 142 MCMV^S-vaccinated mice were protected against SARS-CoV-2 infection, uniformly surviving, maintaining
 143 their weights and showing no noticeable symptoms (**Fig. 3A**). Viral RNA burdens were reduced in the
 144 MCMV^S-immunized group in all examined organs. A few mice showed remaining viral RNA detected in the
 145 respiratory and gastrointestinal tracts, but only minor amounts of viral RNAs were detected in other
 146 organs (**Fig. 3B** and **Supplementary Fig. 5**), arguing that a single administration of the MCMV^S vaccine
 147 protects mice for at least 3 months.



148 **Figure 3. Long-term protection elicited by MCMV^S vaccination against SARS-CoV-2**

149 (A) Percentage of body weight (left), daily clinical scores (middle), and mock-survival kinetics of challenged mice representing the
 150 % of mice reaching a humane end point (right). Log-rank (Mantel-cox) test was used for the statistical analysis (n=5 each). (B)

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152 *Relative viral RNA loads in the representative organs at the indicated time points, normalized to GAPDH. In mice that reached*
153 *humane endpoints, organs were harvested on the day of euthanasia. Each symbol is an individual mouse, and horizontal lines*
154 *indicate the median of biological replicates.*

155 Sustained neutralizing antibody responses against Beta and the Omicron variants upon MCMV^S 156 immunization

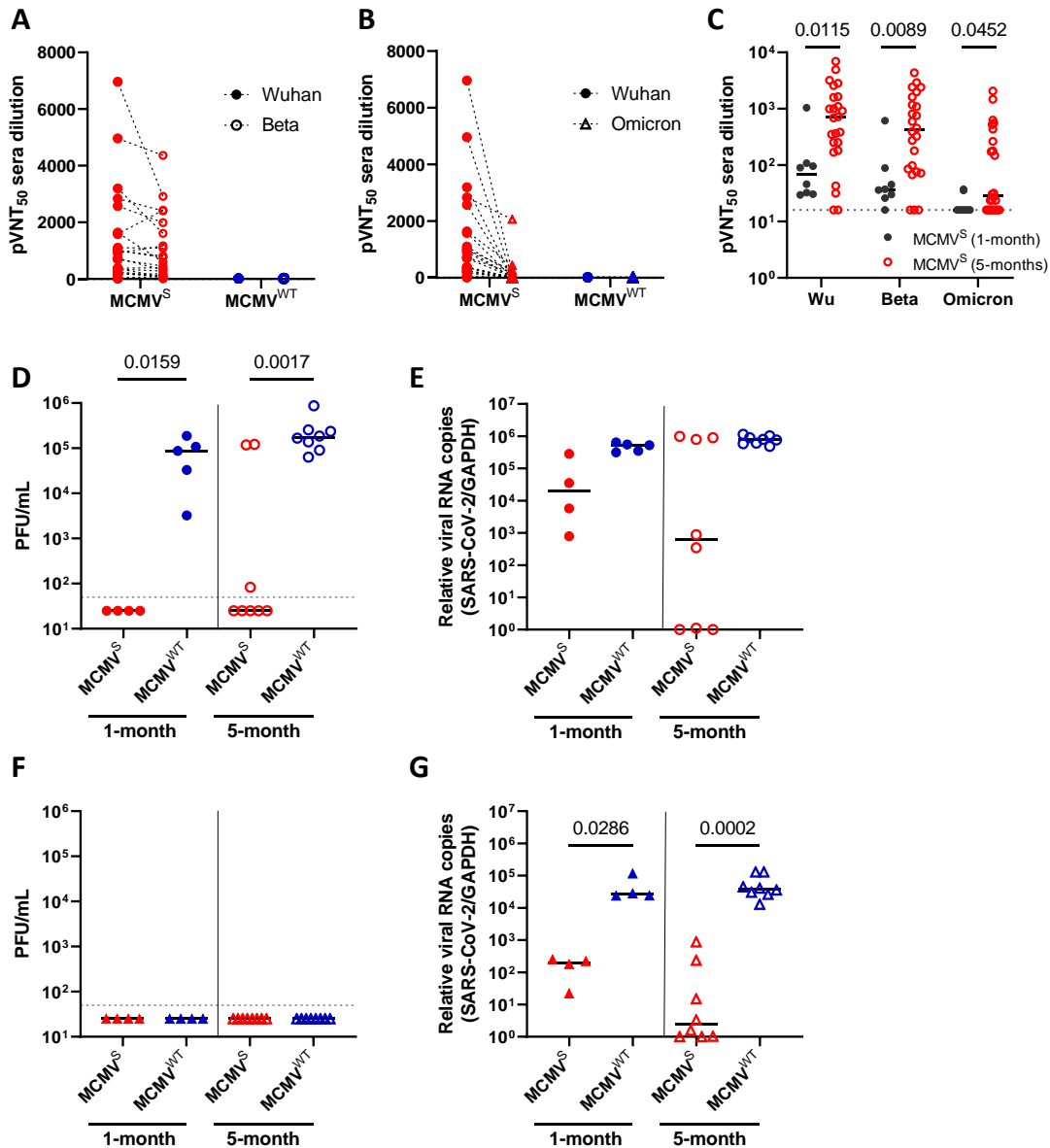
157 While our data on protection were encouraging, the emergence of immune-evading SARS-CoV-2 VoCs
158 compelled an examination of the protective ability of our MCMV-based vaccine against representative
159 VoCs. To this end, we conducted vaccine efficacy studies against the Beta (B.1.351) and the Omicron
160 variant (BA.1). Unlike the D614G strain, both VoCs contain the N501Y mutation in the S protein, which
161 allows SARS-CoV-2 infection in common inbred mice, such as BALB/c and C57BL/6²⁴. The prototype S
162 protein, present in MCMV^S, binds to hACE2 but does not bind to mACE2³¹. Hence, by immunizing BALB/c
163 mice, we evaluated the efficacy of a vaccine formulation that does not engage the mACE2 receptor and
164 has potentially fewer side effects. Moreover, to assess whether our vaccine can protect animals for more
165 than 3 months, we challenged the long-term cohort at 5-months post-vaccination. A short-term cohort
166 was challenged at 6 weeks after immunization as a point of reference. One week before the SARS-CoV-2
167 challenge, all mice were bled for immune status analyses.

168 Virus-neutralizing titers were examined by the pseudo-neutralization assay using pseudotyped VSV (pVSV)
169 that expressed S antigens from the Wuhan, the Beta, or the Omicron variant at 1 or 5 months post-
170 immunization. The neutralization capacity of sera was expressed as the titer resulting in 50% of
171 pseudovirus neutralization (pVNT₅₀), and a head-to-head comparison of pVNT₅₀ against Wuhan and Beta
172 (**Fig. 4A**) or Wuhan and Omicron VoCs (**Fig. 4B**) was performed at 5 months post-vaccination. While pVNT₅₀
173 values against both variants were reduced in most samples, the reduction of titers was less pronounced
174 in Beta (**Fig. 4A**) than in the Omicron variant (**Fig. 4B**), consistent with previous clinical observations³².
175 The neutralizing capacity was not due to cross-neutralizing effects against the vector, because sera from
176 animals immunized with the MCMV^{WT} did not neutralize the pseudoviruses (**Fig. 4A** and **4B**). Strikingly,
177 monitoring for immune waning by comparison of the neutralizing capacity at 5 weeks and 5 months post-
178 immunization revealed that pVNT₅₀ did not decrease but rather increased from 5 weeks to 5 months post-
179 immunization, and this was observed for all tested VoCs (**Fig. 4C**). Taken together, an MCMV-based
180 vaccine expressing a receptor-non-binding S antigen elicited strong and long-lasting neutralizing antibody
181 responses.

182 Reduced viral loads in animals at 1 and 5 months after MCMV^S immunization and challenge with Beta and 183 Omicron variants

184 A week after the bleeding, animals were challenged with 6x10⁴ PFU of the Beta or the Omicron variant of
185 SARS-CoV-2, and vaccine efficacy as determined by the reduction of viral burdens was measured. For this,
186 we analyzed viral RNA copies and infectivity in the lung at 3 days post-infection. We chose this setup since
187 infectious SARS-CoV-2 particles are only detectable for 3 days in the lungs and largely absent from
188 extrapulmonary organs^{27,33,34}. MCMV^S-immunized animals harbored fewer infectious particles in the lungs
189 as compared to the MCMV^{WT} control group (**Fig. 4D**). Parallel viral load analyses indicated that the short-
190 term cohort (1 month) had on average higher residual viral RNAs than the long-term cohort (5 months)
191 (**Fig. 4E**). However, three animals in the 5-months post-vaccination group showed high viral RNA loads
192 upon infection with the Beta VoC and two of the samples were also highly infectious (**Fig. 4D**). These
193 animals exhibited weak or undetectable neutralization against any tested variant (**Fig. 4C**), implying poor
194 vaccination response or vaccination failure in a subset of animals. MCMV^S-vaccinated animals also

195 controlled replication of the Omicron variant based on viral load, which was 100-fold reduced in the short-
 196 term cohort and 10,000-fold in the long-term challenge scenario (Fig. 4G). Isolation of infectious Omicron
 197 virus was not successful for unclear reasons at present. Taken together, our results are consistent with
 198 long-lasting and improving adaptive immunity against the S antigen over time upon vaccination with
 199 MCMV^S (Fig. 1C, 1E, and 4C).

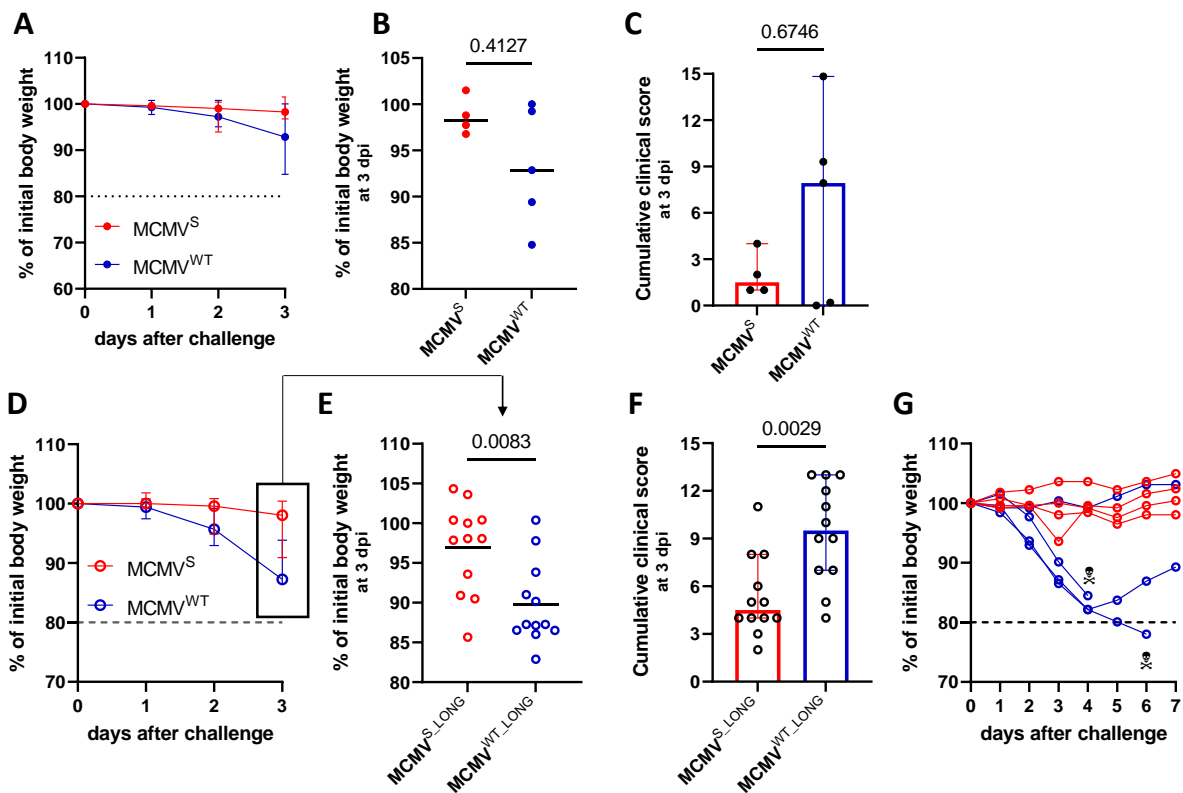


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201 **Figure 4. Beta- and Omicron-specific neutralizing antibodies and in vivo control at 1 and 5 months post-MCMV^S immunization**
 202 (A-B) pVNT₅₀ from the same mice at 5 months post-vaccination (A) against the Wuhan and Beta VoC or (B) Wuhan and Omicron
 203 VoC are connected by dashed lines (n=24). (C) Dot plots of pVNT₅₀ against each variant from the sera of the 1-month (n=8) and 5-
 204 months cohorts (n=24). Each symbol indicates an individual mouse. Solid horizontal lines show the median and the dotted line
 205 indicates the detection limit. (D) Infectious virus titers in lungs of SARS-CoV-2 Beta-challenged mice. (E) Relative viral RNA loads in
 206 lungs of SARS-CoV-2 Beta-challenged mice normalized to GAPDH. (F) Infectious virus titers in lungs of SARS-CoV-2 Omicron-
 207 challenged mice. (G) GAPDH normalized viral RNA loads in lungs of Omicron-challenged mice. Each symbol indicates an individual
 208 mouse, and solid horizontal lines indicate the median of biological replicates. Dashed lines indicate limits of detection. Two-tailed
 209 Mann-Whitney tests were used for statistical analysis (n=4 MCMV^S, n=5 MCMV^{WT} for 1-month; n=8 each for 5-months).

210 Protection against SARS-CoV-2-associated illness against immune-evasive Beta VoC

211 Next, we assessed protection against disease development upon challenge with the Beta or Omicron VoC.
 212 In groups challenged 1-month post-vaccination with the Beta variant, MCMV^S-immunized animals
 213 retained their weights and displayed few clinical symptoms, while MCMV^{WT} mock-immunized animals lost
 214 around 10% of their initial weights (Fig. 5A-C). Similarly, MCMV^S-vaccinated animals were overall
 215 protected from weight loss in comparison with MCMV^{WT}-controls at 5-months post-vaccination (Fig. 5D)
 216 and displayed lower clinical scores (Fig. 5E and 5F). A randomly pre-selected subset of animals was
 217 monitored for 7 days, where 50% of animals in the mock-immunized group reached the humane end-
 218 point, while body weights were maintained in all MCMV^S-vaccinated animals (Fig. 5G). Animals that were
 219 challenged with the Omicron variant showed no weight loss and very mild clinical symptoms
 220 (Supplementary Fig. 6), in line with reports by other studies. Animals vaccinated with MCMV^S for 5
 221 months were nevertheless protected against the mild symptoms elicited by the Omicron infection,
 222 because the cumulative clinical scores at dpi 3 were on average 2-fold lower in this than in the MCMV^{WT}
 223 control group. Altogether, our data strongly indicate that a single dose of MCMV^S vaccination provides
 224 long-lasting protection against the immune evasive Beta variant of SARS-CoV-2.



225

226 **Figure 5. Protection against the Beta variant-associated illness after 1-month or 5-months after vaccination**

227 (A, D) Percentage of body weight monitored for 3 days after upon infection with the Beta variant. Medians with 95% confidence
 228 interval are shown. (B, E) Percentages of initial body weight in individual animals at 3 dpi. Each symbol indicates an individual
 229 mouse. (C, F) Cumulative clinical scores at dpi 3. Each symbol shows summed-up clinical scores up to 3 dpi. Medians (bar graphs)
 230 with 95% confidence interval are shown. (G) Body weight loss was monitored for a week after the SARS-CoV-2 infection. Each line
 231 connecting dots indicates an individual mouse, and skull symbols indicate animals that reached a humane end-point. Closed
 232 symbols show one month (A-C) and open symbols present five months (D-G) after vaccination. Dashed lines indicate the humane-
 233 end point. Solid horizontal lines show the median of biological replicates (n=4 MCMV^S, n=5 MCMV^{WT} for A-C; n=12 each for D-F).
 234 Statistical significance was analyzed using two-tailed Mann-Whitney tests.

235 Discussion

236 Several vaccines have been developed to combat the COVID-19 pandemic. However, immune waning
237 remains an unresolved problem^{6,7,9,35}, and vaccines that result in lasting and broad protection against
238 SARS-CoV-2 variants remain unavailable. The emergence of the Omicron variant has emphasized the
239 urgent need for such a vaccine, as the currently authorized vaccine regimens showed heterogeneous
240 protection levels against this variant^{36,37}, resulting in 30-88% of vaccine effectiveness against
241 hospitalization and frequent breakthrough infections³⁸. Viral vectors that were authorized early in the
242 COVID-19 pandemic, such as Vaxzevria (ChAdOx1-S) and the COVID-19 vaccine Janssen (Ad26.COVS.2.S)³⁹,
243 exhibit the same restrictions as the mRNA-based vaccines³⁶. CMV has received attention as a new vaccine
244 vector tool due to the induction of a strong and lasting T-cell immunity²¹. While HCMV seroprevalence is
245 estimated to exceed 90% in some geographic areas, the pre-existing immunity to CMV does not hinder
246 vaccine responses and protection, as demonstrated in studies with rhesus monkeys^{17,40}. MCMV shares
247 structural and functional homology with HCMV and allows *in vivo* analyses in the natural host. Hence, we
248 used this model virus to test whether a CMV-based vector might in principle induce long-lasting protection
249 against SARS-CoV-2 infection and disease development. Our vaccine induced strong and durable S-specific
250 CD8 T cells (**Fig. 1C-D**) and neutralizing antibody responses (**Fig. 1E** and **4C**). Importantly, immune
251 protection against lethal challenge at 3 months post-immunization (**Fig. 3**) was just as robust as at 1-month
252 post-infection (**Fig. 2**) and protection against immune evasive VoCs was maintained at 5 months post-
253 vaccination (**Fig. 4** and **5**). Hence, our data indicate that our approach may provide durable immune
254 protection against COVID-19 upon vaccination.

255 Neutralizing antibody responses correlate with vaccine effectiveness⁴¹ and are therefore considered an
256 important predictor of vaccine potency. We have previously shown that MCMV-based vaccines may elicit
257 strong antibody responses whose affinity and neutralizing capacity increase over time²⁶. Here, we show
258 that the MCMV^S vaccine induces antibodies that neutralize the Beta and Omicron variants and protect
259 against disease upon infection with these viruses, with neutralization titers increasing over time (**Fig. 4A-**
260 **C**). This feature is unique for our vaccine approach and in stark contrast with other vaccine formulations,
261 which require prime/boost administration for optimal performance^{39,42,43}. Considering that neutralizing
262 antibodies against SARS-CoV-2 wane at similar rates upon infection or vaccination with currently used
263 vaccines^{30,35,37,44}, our vaccine provides an improvement over both of these scenarios.

264 Interestingly, even the vaccinated animals with weak neutralizing antibody responses against SARS-CoV-
265 2 variants were protected against illness and showed lowered viral loads in the lungs (**Fig. 4**). This
266 phenomenon was also demonstrated in a preclinical study with rhesus macaques and in clinical studies,
267 where study subjects were protected although no neutralizing antibodies were observed⁴⁵⁻⁴⁷. One may
268 speculate that T-cell immunity elicited by MCMV^S vaccination protected these animals, but we cannot
269 exclude the possibility that memory B lymphocytes in MCMV^S-immunized mice responded to the
270 challenge by rapidly generating neutralizing antibodies upon challenge, thus limiting virus replication and
271 protecting the host. It is finally also possible that the concerted action of these two lymphocyte lineages
272 protected the vaccinated mice against challenge. To differentiate between these scenarios, one may use
273 MCMV vectors eliciting T-cell responses against S epitopes only, or mice lacking T-cells, but these
274 experiments go beyond the scope of our study.

275 In this study, we showed that an S antigen that cannot bind to the target ACE2 receptor might still provide
276 robust immune protection. Namely, we cloned the S antigen from the prototypical Wuhan strain of SARS-

277 CoV-2 into our vaccine vector, but the binding affinity of the Wuhan S protein to murine ACE2 receptors
278 is low³¹. Consequently, the S antigen from our vaccine could not bind to the mACE2 receptor. On the
279 other hand, the N501Y mutation in the S protein enables infection of common laboratory inbred mice,
280 such as BALB/c or C57BL/6²⁴, and this mutation is present in the Beta and the Omicron VoCs. Therefore,
281 MCMV^S elicited protective immunity against variants that engaged the mACE2, although the S expressed
282 by the vaccine vector did not. This is interesting because some studies reported that the binding of the S1
283 subunit of the S protein to ACE2 on endothelial cells may affect endothelial barrier integrity and cardiac
284 activity⁴⁸⁻⁵⁰. While we cannot formally exclude that long-term S antigen expression may drive adverse
285 effects unrelated to ACE2 binding, our data demonstrate that our vaccine formulation protects mice against
286 SARS-CoV-2 variants in absence of ACE2 receptor engagement, which is especially important if low levels
287 of S antigen persist in the host.

288 The data presented here demonstrate that an MCMV-based vaccine candidate expressing the full-length
289 prototype S protein is highly immunogenic and protective in mice, with robust activation of both arms of
290 the adaptive immune system, cellular and humoral responses. We demonstrated robust protection
291 against a lethal dose of the SARS-CoV-2 D614G variant in K18-hACE2 mice and the immune-evasive Beta
292 and Omicron variants in BALB/c mice by a single dose of our MCMV^S vaccine. Future research needs to
293 focus on a head to head comparison of CMV-based vaccines with other COVID-19 vaccine formulations,
294 on the length of protection and administration routes, paving the way towards clinical trials with
295 appropriate CMV vectors. However, by demonstrating in proof of concept that a CMV vector can provide
296 long-term protection against COVID-19 upon a single immunization shot, we provide here the first and
297 crucial step in this direction.

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418 Materials and Methods

419 Cell culture and viruses

420 Vero E6 (CRL-1586) and M2-10B4 cells (ATCC CRL-1972) were cultured as described previously²⁶. Caco-2
421 cells (ACC 169) were purchased from DSMZ (Braunschweig, Germany) and were cultured in DMEM (Gibco,
422 NY, USA) supplemented with 20% fetal bovine serum (FBS), 2 mM L-glutamine, 100 IU/mL penicillin and
423 100 µg/mL streptomycin. The FI strain of SARS-CoV-2 (GISAID database ID: EPI_ISL_463008) was described
424 previously as a D614G variant⁵¹ and was passaged on Caco-2 cells in the biosafety level 3 (BSL3) laboratory
425 at HZI.

426 SARS-CoV-2 genome sequences are available on GISAID and GenBank under the following accession
427 numbers: SARS-CoV-2 B.1.351 (Beta) FFM-ZAF1/2021 (GenBank ID: MW822592)⁵² and SARS-CoV-2
428 B.1.1.529 (BA.1) FFM-ZAF0396/2021 (EPI_ISL_6959868; GenBank ID: OL800703)⁵³. MCMV^{WT} refers to the
429 BAC-derived molecular clone (pSM3fr-MCK-2fl clone 3.3)⁵⁴. MCMV^S was generated by *en passant*
430 mutagenesis, as described previously²⁶. Briefly, the codon-optimized S protein of the Wuhan variant
431 replaced the viral ie2 protein.

432 The expression vector for SARS-CoV-2 S protein of Omicron (BA.1) (based on isolate hCoV-
433 19/Botswana/R40B58_BHP_3321001245/2021; GISAID Accession ID: EPI_ISL_6640919) was generated by
434 Gibson assembly as described previously⁵⁵ and then subsequently introduced in pseudotype VSV
435 backbone that lacks VSV glycoprotein G (VSV-G)⁵⁶. A plasmid encoding the S protein of SARS-CoV-2 Beta
436 (B.1.351) has been previously reported²⁶.

437 Virus stock generation and plaque assay

438 BAC-derived mutant MCMVs were propagated on M2-10B4 cells and concentrated by sucrose density
439 gradient centrifugation. BAC-derived MCMV was reconstituted by transfection of BAC DNA into NIH-3T3
440 cells (ATCC CRL-1658) using FuGENE HD transfection reagent (Promega, WI, USA) according to the
441 manufacturer's instructions. Transfected cells were cultured until viral plaques appeared and passaged 5
442 times in M2-10B4 cells before virus stock production. Virus stocks were prepared on ice. First,
443 supernatants of infected M2-10B4 cells were collected and infected cells were pelleted (5,000 × g for 15
444 min). The resulting cell pellets were homogenized in DMEM supplemented with 5% FBS and cell debris
445 was removed by centrifugation (12,000 × g for 10 min). Collected supernatants were resuspended in VSB
446 buffer (0.05 M Tris-HCl, 0.012 M KCl, and 0.005 M EDTA, adjusted to pH 7.8) and then concentrated by
447 centrifugation through a 15% sucrose cushion in VSB buffer (23,000 × g for 1.5 h). The resulting pellet was
448 re-suspended in 1-1.5 mL VSB buffer, briefly spun down, and supernatants were aliquoted and kept at -
449 80°C.

450 SARS-CoV-2 D614G was generated and viruses were quantified by plaque assays as described before²⁶
451 with a minor modification that Caco-2 cells were used for virus production. SARS-CoV-2 Beta and Omicron
452 BA.1 stocks were generated as described previously and titers were determined by the median tissue
453 culture infective dose (TCID₅₀) method⁵³.

454 Pseudotyped viruses were harvested as described before ^{26,55}. In brief, 293T cells were transfected with
455 expression plasmids (pCG1) encoding different S proteins of SARS-CoV-2 variants by using the calcium-
456 phosphate method. At 24h post-transfection, the medium was removed and cells were inoculated with a
457 replication-deficient VSV vector lacking its glycoprotein and coding instead for an enhanced green
458 fluorescent protein (GFP) (kindly provided by Gert Zimmer, Institute of Virology and Immunology,
459 Mittelhäusern, Switzerland). Following 1 h incubation at 37°C, the cells were washed with PBS, and culture
460 media containing anti-VSV-G antibody (culture supernatant from I1-hybridoma cells; ATCC CRL-2700)
461 were added. The pseudotype virus was harvested at 16-18 h post-infection.

462 *Virus in vivo* infection

463 K18-hACE2 mice were obtained from Jackson Laboratories and bred in the core animal facility of
464 Helmholtz Center for Infection Research, Braunschweig. BALB/c mice were purchased from Envigo (IN,
465 USA). All animals were housed under Specific Pathogen Free (SPF) conditions at HZI during breeding and
466 infection. All animal experiments were approved by the Lower Saxony State Office of Consumer Protection
467 and Food Safety.

468 K18-hACE2 mice (2-6 months old) were intraperitoneally (i.p.) immunized with 10⁶ PFU of recombinant
469 MCMV^S or MCMV^{WT} diluted in PBS or treated with PBS (200 µL per animal). BALB/c mice (4-7 months old)
470 were i.p. immunized with 2x10⁵ PFU of recombinant MCMV^S or MCMV^{WT}. All mice were weekly or bi-
471 weekly monitored and scored for their health status after vaccination. Blood was analyzed at indicated
472 time points.

473 SARS-CoV-2 challenge experiments were performed in the HZI BSL3 laboratory essentially as described ⁵⁷
474 with the following modifications: K18-hACE2 mice were intranasally (i.n.) infected with 2x10³ PFU of
475 D614G SARS-CoV-2, while BALB/c mice were challenged with 6x10⁴ PFU of the VoCs. SARS-CoV-2-infected
476 mice were monitored for weight loss and clinical status daily, according to the animal permit.

477 SARS-CoV-2-challenged animals were scored daily to monitor any signs of disease development. Animals
478 were scored based on five criteria: spontaneous/social behavior, fur, fleeing behavior, posture, and
479 weight loss. Each score indicates the following: no signs of symptoms (score=0), mild and/or sporadic
480 symptoms (score=1), moderate and/or frequent symptoms (score=2), and severe symptoms with a clear
481 sign of heavy suffering (score=3). Weight loss criterion is scored as follows: ≤1 % (score=0), 1-10 %
482 (score=1), 10-20% (score=2), and >20% (score=3). Mice with a score of 3 in one criterion, or an overall
483 score of ≥9, were removed from the experiments.

484 *Organ harvest*

485 The trachea, lungs, heart, spleen, stomach, and brain were harvested at 3 or 7 days post-SARS-CoV-2
486 challenge, homogenized in 500 or 1,000 µL PBS with an MP Biomedical FastPrep 24 Tissue Homogenizer
487 (MP Biomedicals, CA, USA) and stored at -80°C until the usage for plaque assay or qRT-PCR analysis. In
488 animals that reached humane endpoints before 7 days post-challenge, organs were harvested on the day
489 of euthanasia.

490 *RNA isolation and viral loads analyses*

491 RNA was isolated according to the manufacturer's protocol (Rneasy RNA isolation kit, Qiagen). Shortly,
492 250 µL of organ homogenates in 750 µL Trizol were centrifuged at 16,000 x g for 3 min. The resulting
493 supernatants were carefully collected and washed with the same volume of 70% Ethanol. The mixed
494 solution was transferred into a collection tube and centrifuged at 10,000 x g for 30 sec. After decanting

495 the flow-through, the column was washed once with 700 μ L of RW1 wash buffer and twice with 500 μ L
496 RPE buffer. Lastly, 40 μ L of nuclease-free water was added to the column for RNA elution, and isolated
497 RNAs were kept at -80°C .

498 Eluted RNAs were analyzed further to assess viral RNAs in the given organs by quantitative reverse
499 transcription polymerase chain reaction (RT-qPCR). The reaction was performed with a total volume of 20
500 μ L containing 2 μ L of sample RNAs or positive control RNAs, 5 μ L TaqPath 1-step RT-qPCR Master Mix with
501 ROX reference dye, and 1.5 μ L probe/primer sets. 2019-nCoV RUO kit was used to detect SARS-CoV-2
502 RNAs (Integrated DNA Technologies (IDT), USA), and Taqman Rodent GAPDH control reagents
503 (ThermoFischer Scientific, USA) were used for endogenous GAPDH RNAs. For absolute viral RNA
504 quantification, a standard curve was generated by serially diluting a SARS-CoV2 plasmid with the known
505 copy numbers 200,000 copies/ μ L (2019-nCoV_N_Positive Control, #10006625, IDT, USA) at 1:2 ratio in all
506 PCR analyses, with a quantitation limit of 20 copies of the plasmid standard in a single qPCR reaction. The
507 viral RNA of each sample was quantified in triplicate and the mean viral RNA was calculated by the
508 standard. RT-qPCR was performed using the StepOnePlus™ Real-Time PCR system (ThermoFischer
509 Scientific, USA) according to the manufacturer's instructions.

510 Detection of infectious SARS-CoV-2 in the lungs

511 Lung organ homogenates were serially diluted in DMEM (Gibco, NY, USA) supplemented with 5% FBS and
512 100 IU/mL penicillin, and 100 μ g/mL streptomycin. 100 μ L of sample dilutions were transferred onto
513 confluent VeroE6 cells in a 96-well format. After inoculation for 1 h at 37°C , the inoculum was removed
514 and 1.5% methylcellulose in MEM supplemented with 5% FBS, 2 mM L-glutamine, 100 IU/mL penicillin,
515 and 100 μ g/mL streptomycin was added to the cells. The infected cells were incubated at 37°C for 48 h
516 before inactivation with a 4% formalin solution in PBS for 10 min at RT. The fixed cells were subjected to
517 immunofluorescent staining against the SARS-CoV-2 N protein. Briefly, fixed cells were permeabilized with
518 0.1% Triton X-100 (Sigma-Aldrich, MA, USA) for 10 min at RT and blocked with 1% BSA (Sigma-Aldrich, MA,
519 USA) in PBS for 30 min at RT. Thereupon, cells were incubated with a monoclonal anti-SARS-CoV-2 N
520 protein antibody (Abcalis, AB84-E02, 10 μ g/mL) for 30 min at RT. After washing three times with PBS with
521 0.05% Tween-20 (PBS-T), a secondary antibody anti-mouse IgG conjugated with Alexa488 (Cell Signaling
522 Technology, #4408, 1:500) was added for 30 min at RT. After washing three times with PBS-T, the stained
523 cells were visualized using Incucyte S3 (Sartorius, Goettingen, Germany).

524 Flow cytometry quantification of S-specific T cells

525 Peripheral blood was harvested and red blood cells were removed by short osmotic shock. Thereupon,
526 lymphocytes were stained with S-derived VNFNFNGL-specific tetramers (Kindly provided by Ramon Arens,
527 Leiden University) for 30 min at RT. Subsequently, cells were stained with fluorescent-labeled antibodies
528 against CD3 (17A2, eBiosciences, CA, USA), CD4 (GK1.5, BioLegend, CA, USA), CD8a (53-6.7, BD Bioscience,
529 CA, USA), CD44 (IM7, BioLegend, CA, USA), CD11a (M17/4, BioLegend, CA, USA), CD62L (MEL-14,
530 BioLegend, CA, USA), and KLRG1 (2F1, BioLegend, CA, USA) for 30 min at 4°C . Dead cells identified by 7-
531 AAD viability staining solution (BioLegend, CA, USA) were excluded from all analyses. The labeled cells
532 were analyzed by flow cytometry (BD LSRFortessa™ Cell Analyzer) and subsequent analyses were done in
533 detail in FlowJo Software v10.

534 Detection of anti-spike antibodies in mouse sera

535 ELISA (Enzyme-Linked ImmunoSorbent Assay) was used to detect anti-S IgGs in vaccinated animal sera
536 and performed as described previously⁵⁸. Briefly, antigens were immobilized on 384-well plates and

537 blocked with 2% milk powder in PBS-T. Mouse sera were serially diluted starting at a concentration of
538 1:100. S1-S2-His of different S variants were used to identify the antigen-specific binding and BSA or cell
539 lysates were used as control for unspecific binding. EC₅₀ was analyzed by a statistical analysis tool in
540 GraphPad Prism 9.

541 *In vitro* live virus neutralization titer (VNT) assay

542 The serum neutralization assay was performed as described before ²⁶. Briefly, heat-inactivated sera were
543 serially diluted and incubated with 100 PFU/100 µL of SARS-CoV-2 for an hour at RT. Thereupon, they
544 were transferred to 96-well plates seeded with Vero-E6 cells and inoculated with serum and virus for 1 h.
545 After the inoculum removal, the cells were overlaid with 1.5% methylcellulose and incubated at 37°C and
546 5% CO₂ for 3 days. The cells were fixed with 4% formaldehyde, followed by crystal violet staining and
547 plaque counting. Serum-neutralizing titer that results in a 50% reduction of virus plaques (VNT₅₀) was
548 analyzed by GraphPad Prism 9 nonlinear regression analysis.

549 Pseudovirus neutralization assay

550 Pseudovirus neutralization assays were performed as described in the previous publications ^{4,26}. For
551 neutralization experiments, pseudotyped particles and heat-inactivated serum dilution were mixed at a
552 1:1 ratio and incubated for 60 min at 37°C before being inoculated onto VeroE6 cells grown in 96-well
553 plates. At 24 h post-infection, GFP expression was measured by using Incucyte S3 (Sartorius, Goettingen,
554 Germany).

555 Statistics

556 One-way ANOVA with Kruskal-Wallis correction was performed for multiple-group analysis. Two-tailed
557 Mann-Whitney tests were used to compare the difference between two independent groups. Log-rank
558 (Mantel-cox) tests were performed to compare the survival distributions of groups. Statistical analysis was
559 calculated by GraphPad Prism 9.

560 Data availability

561 All SARS-CoV-2 genome sequences are available as mentioned above. The D614 SARS-CoV-2 variant
562 (GISAID database ID: EPI_ISL_463008), SARS-CoV-2 B.1.351 (Beta) FFM-ZAF1/2021 (GenBank ID:
563 MW822592), and SARS-CoV-2 B.1.1.529 (BA.1) FFM-ZAF0396/2021 (EPI_ISL_6959868; GenBank ID:
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597 Author Contributions

598 Y.K. and L.C.S. conceived and designed the study; Y.K. H.J. and B.F. performed *in vivo* experiments; F.B.
599 and M.Hu. performed and analyzed ELISA serology; Y.K., H.J., and F.B. analyzed the data; Y.K. and L.C.S.
600 interpreted the data; Y.K. and L.C.S. wrote the manuscript; K.E., M.Z.C., M.W., S.C., M.Ho., and S.P.
601 provided recombinant virus or clinical isolates; L.C.S. supervised the study.
602 All authors reviewed the results and approved the final version of the manuscript. Correspondence and
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604 Competing interests

605 The authors LCS and YK are applicants for a patent based on MCMV as a vaccine vector. The authors
606 declare no other competing interests.