The pre-exposure SARS-CoV-2-specific T cell repertoire determines the quality of the immune response to vaccination

**Highlights**

- SARS-CoV-2-specific T cell expansion rate indicates immune-response quality

- Naive and memory pre-exposure repertoires inversely predict vaccine-response quality

- Pre-existing memory T cells are excluded from high-quality vaccination responses

- Impaired antigen-specific pre-exposure T cell repertoire as a hallmark of immune aging

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**In brief**

Determinants of immune-response quality to SARS-CoV-2 remain poorly defined. Saggau et al. examine spike-specific naive and memory T cells pre- and post-vaccination and track pre-existing memory T cell receptors. They define T cell parameters of high-quality vaccine responses and identify high pre-existing memory and low naive T cell contributions as predictors of low-quality responses, particularly in the elderly.
The pre-exposure SARS-CoV-2-specific T cell repertoire determines the quality of the immune response to vaccination

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SUMMARY

SARS-CoV-2 infection and vaccination generates enormous host-response heterogeneity and an age-dependent loss of immune-response quality. How the pre-exposure T cell repertoire contributes to this heterogeneity is poorly understood. We combined analysis of SARS-CoV-2-specific CD4+ T cells pre- and post-vaccination with longitudinal T cell receptor tracking. We identified strong pre-exposure T cell variability that correlated with subsequent immune-response quality and age. High-quality responses, defined by strong expansion of high-avidity spike-specific T cells, high interleukin-21 production, and specific immunoglobulin G, depended on an intact naive repertoire and exclusion of pre-existing memory T cells. In the elderly, T cell expansion from both compartments was severely compromised. Our results reveal that an intrinsic defect of the CD4+ T cell repertoire causes the age-dependent decline of immune-response quality against SARS-CoV-2 and highlight the need for alternative strategies to induce high-quality T cell responses against newly arising pathogens in the elderly.

INTRODUCTION

Clinical symptoms of COVID-19 are highly diverse, ranging from no or mild symptoms to severe and often lethal disease, especially in the elderly. The rapid availability of highly effective SARS-CoV-2 vaccines is a cornerstone in the fight against the COVID-19 pandemic. However, vaccination efficacy is also variable and can be severely compromised in the elderly (Gustafson et al., 2020), as also observed following SARS-CoV-2 vaccination (Anderson et al., 2020; Collier et al., 2021; Li et al., 2021; Romero-Olmedo et al., 2022a). This is further indicated by more frequent break-through infections and an increased risk of developing severe symptoms in the elderly, even after full vaccination (Juthani et al., 2021; Wang et al., 2022; Yek et al., 2022). The
Figure 1. High- and low-quality CD4⁺ T cell responses develop upon SARS-CoV-2 vaccination
(A) Overview of the study design. Blood was drawn at the indicated time points and analyzed for SARS-CoV-2-specific CD4⁺ T cells and IgG. 34 individuals of our study received two doses of the BioNTech/Pfizer BNT162b2 mRNA vaccine, while 16 individuals received a heterologous immunization with AstraZeneca ChAdOx1 followed by Moderna mRNA-1273.
(B) Representative dot plot examples for the ex vivo detection of SARS-CoV-2 spike-reactive CD4⁺ T cells by ARTE pre- and post-vaccination. Absolute cell counts after magnetic CD154⁺ enrichment from 1x10⁷ PBMCs are indicated.
(C) Serum anti-spike IgG antibody concentrations pre- and post-second vaccination.
(D) Frequencies of SARS-CoV-2 spike-reactive Tmems pre- and post-vaccination (left) in comparison to convalescent COVID-19 patients with mild (non-hospitalized) or severe disease (based on WHO criteria WHO4–7) (right).
(E) Vaccination-induced expansion of spike-reactive CD4⁺ Tmems was calculated for each donor based on the frequency pre-vaccination and after the second vaccination. Individuals with the lowest (“bottom 10”) and highest (“top 10”) expansion rate are indicated in blue or orange, respectively.

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strong age association suggests that the immune status of the single individual is a critical component, which can also affect individuals from younger age groups. It is therefore a key question whether the antigen-specific pre-exposure status determines the quality of the response to SARS-CoV-2 vaccination, which would allow for identification of individuals at high risk and design of tailor-made protective measures.

CD4⁴ T cells orchestrate adaptive immune responses, and the T cell compartment is dramatically affected by aging. Decreasing thymic activity leads to a reduced production of new naive T cells in the elderly. This may affect mainly the response to newly encountered antigens, as the T cell receptor (TCR) repertoire diversity providing specific high-affinity TCRs is reduced (Cicin-Sain et al., 2010; Gustafson et al., 2020; Woodland and Blackman, 2006; Yager et al., 2008). On the contrary, a broad memory repertoire accumulating with age can also provide T cells with specificities reactive to newly arising antigens (Woodland and Blackman, 2006). Such pre-existing memory T cells have been shown to result from cross-reactivity to related, or even unrelated, antigenic peptides (Birnbaum et al., 2014; Sewell, 2012). However, exaggerated clonal expansions, or increased numbers of senescent or chronically stimulated T cells lacking full proliferative and effector capacity, may negatively affect the response of pre-existing memory T cells (Allen et al., 2020). Thus, the composition and status of the antigen-specific pre-exposure T cell repertoire, comprising both a diverse naive compartment and certain pre-existing memory T cell specificities, forms a key determinant for the quality of the resulting immune response.

Pre-existing memory T cells against SARS-CoV-2 have been demonstrated in unexposed donors (Bacher et al., 2020; Braun et al., 2020; Grifoni et al., 2020; Le Bert et al., 2020; Mateus et al., 2020; Nelde et al., 2020). T cell cross-reactivity of these pre-existing SARS-CoV-2-reactive memory cells to related “common cold” corona viruses (CCCoVs) (Bacher et al., 2020; Low et al., 2021; Loyal et al., 2021; Mateus et al., 2020; Nelde et al., 2020), but also other pathogens, has been described (Bacher et al., 2020; Bartolo et al., 2021; Stervbo et al., 2020; Tan et al., 2021). Importantly, pre-existing T cell memory against so far unexposed antigens is a common phenomenon for many antigens (Bacher et al., 2013; Campion et al., 2014; Kwok et al., 2012; Pan et al., 2021; Su et al., 2013). So far, it is still not clear to what extent pre-existing memory T cells contribute to the immune response against SARS-CoV-2 and how they influence the response quality. Several studies find correlations between the frequency of SARS-CoV-2-reactive pre-existing memory cells and mild or asymptomatic COVID-19 or improved immunity to vaccination (Loyal et al., 2021; Mallajosyula et al., 2021; Mateus et al., 2021; Swadling et al., 2022). We and others recently detected pre-existing memory T cells against SARS-CoV-2 in essentially all unexposed individuals (Bacher et al., 2020; Guerrero et al., 2021). These cells display rather low functional avidity to SARS-CoV-2 (Bacher et al., 2020), which was confirmed even for major conserved peptide epitopes between SARS-CoV-2 and CCCoVs (Dykema et al., 2021). Low-avidity CD4⁴ T cells are also a hallmark of severe COVID-19, suggesting their potential recruitment from the pre-existing memory compartment (Bacher et al., 2020). These data challenge the idea of a protective contribution of pre-existing memory CD4⁴ T cells to SARS-CoV-2 immunity and even suggest a potential negative impact on the immune-response quality. However, as of yet, the quantitative contribution of pre-existing memory T cells and their impact on the response quality has not been directly assessed.

Here, we employed antigen-reactive T cell enrichment (ARTE) (Bacher et al., 2013, 2016, 2019, 2020) in combination with clonal TCR tracking to follow the fate of pre-existing memory T cells and to characterize naive and memory SARS-CoV-2-specific T cells pre- and post-vaccination. We identified the pre-vaccination repertoire as a major determinant for the immune-response quality to vaccination. Strong clonal expansion, but exclusion of pre-existing memory T cells, was a hallmark of high-quality immune responses, which were characterized by high frequencies of high-avidity spike-specific T cells, strong interleukin-21 (IL-21) production, and higher concentration of anti-spike immunoglobulin G (IgG). Age-related changes in both the naive and memory pre-exposure T cell repertoire resulted in a CD4⁴ T cell-intrinsic inability to mount high-quality immune responses in the elderly (>80 years), emphasizing the need for novel intervention strategies.

RESULTS

SARS-CoV-2 vaccination generates qualitatively different CD4⁴ T cell responses

We analyzed the SARS-CoV-2 spike protein-specific CD4⁴ T cell response before and after vaccination in 50 SARS-CoV-2-naive individuals, which was confirmed by testing negative for SARS-CoV-2 IgG pre-vaccination (Figure S1A). Three donors became infected with SARS-CoV-2 during the course of our study, as indicated by an increase in nucleocapsid protein-specific IgG, and were excluded from further analysis (Figure S1A). Blood was drawn pre-vaccination, 3–5 weeks after the first vaccination, and 6–8 weeks after the second vaccination (Figure 1A).

To analyze the whole repertoire of spike-specific CD4⁴ T cells, including memory and naive cells, we used a previously developed highly sensitive technology, the ARTE assay (Bacher et al., 2013, 2016, 2019, 2020). Freshly isolated peripheral blood mononuclear cells (PBMCs) were stimulated ex vivo for 7 hours with a peptide pool spanning the whole spike-protein sequence.
Spike-reactive CD4+ T cells were detected based on up-regulation of CD154 (CD40L) and magnetically enriched from 1x10^7 PBMCs. This enabled a precise enumeration of rare spike-reactive T cells pre-vaccination (Figures 1B and S1B) with a minimum of 50 detected CD154+ cells (range 50–689, mean 166) after subtraction of background events (Figures S1C and S1D), which allowed detection of spike-reactive pre-vaccination responses in both the naive repertoire and from cross-reactive pre-existing memory cells (Figures S1B–S1D). Specificity of CD154+ cells pre- and post-vaccination was confirmed by expansion of sorted CD154+ memory T cells followed by restimulation with spike peptide pool or a negative control pool of influenza A proteins (Figure S1E), as reported previously (Bacher et al., 2020). Most donors of our study (n = 34) received two doses of the BioNTech/Pfizer BNT162b2 mRNA vaccine, while 16 individuals received a heterologous immunization with AstraZeneca ChAdOx1, followed by Moderna mRNA-1273 (Figure 1A and Tables S1 and S2). We observed no significant differences in the frequencies of spike-reactive CD4+ T cells or anti-spike IgG concentrations between these two vaccination regimens (Figures S1F and S1G).

Vaccination induced spike-specific IgG (Figure 1C) as well as spike-reactive memory CD4+ T cells (Tmems) at magnitudes comparable to those observed in convalescent COVID-19 patients with mild disease (Figure 1D). As expected, the post-vaccine response was characterized by a shift from naive to memory cells, especially of the effector memory phenotype (CD45RA–/CCR7–) (Figure S2A). Analysis of recent in vivo activation showed increased CD38 expression post-first vaccination that declined thereafter (Figure S2A). The post-vaccination response was further characterized by increased frequencies of IL-2-, IFN-γ-, IL-21-, and IL-10-producing cells (Figure S2B) that were in a similar range as in convalescent COVID-19 patients (Bacher et al., 2020).

To identify potential differences in the host-response quality, we calculated the SARS-CoV-2 spike-specific T cell expansion for each donor based on the frequencies pre-vaccination and post-second vaccination. The fold expansion displayed strong differences between individuals, ranging from a 2- to 312-fold increase in spike-reactive Tmems (Figure 1E). We next compared the bottom 10 expanders (≤15-fold expansion) and top 10 expanders (≥100-fold expansion) (Figure 1E). Donors contained in the top 10 group not only reached higher frequencies of spike-reactive Tmems (Figure 1F), but also had cells with a higher functional avidity (Figure 1G), as tested by restimulation of expanded spike-specific T cells with graded antigen concentrations (Figure S2C). Furthermore, top 10 expanders showed significantly increased IL-21 production (Figures 1H and 1I) but similar frequencies of IFN-γ, IL-2, or IL-10 producers (Figure S3D). Because IL-21 is an essential cytokine for B cell help, we next analyzed anti-spike IgG antibodies. Top 10 expanders displayed higher concentrations of anti-spike IgG (Figure 1J) as well as higher titers of SARS-CoV-2-neutralizing antibodies (Figure 1K). In line with these results, we observed a strong positive correlation between the fold expansion of spike-specific CD4+ T cells and anti-spike IgG concentrations (Figure 1L) or neutralizing antibody titers (Figure S3A) as well as the frequency of IL-21 producers (Figure S3B). The differences in anti-spike antibodies were also retained 6 months after the second vaccination (Figures 1J and 1K), although anti-spike IgG concentrations generally declined in all donors.

Overall, these data identify strong qualitative differences in the CD4+ T cell response to SARS-CoV-2 between individual vaccinees. A high-quality immune response is characterized by strong T cell expansion, resulting in high frequencies of high-avidity T cells and high IL-21 production, which correlates with high concentrations of anti-spike IgG antibodies.

High age predisposes individuals for poor T cell expansion despite increased pre-existing memory

We next asked which parameters might influence these qualitative differences between individuals. Vaccination efficacy is known to decline with age (Gustafson et al., 2020), and indeed the bottom 10 group contained a higher proportion of individuals >80 years of age (Figure 2A). Pre-existing T cell memory against SARS-CoV-2 in unexposed donors has been described, but its impact on infection and vaccination is controversially discussed (Moss, 2022). When we compared the frequencies of spike-specific memory cells before vaccination or the proportion of memory cells within the spike-specific T cells, we observed much higher frequencies of pre-existing memory T cells in the bottom 10 compared with the top 10 expander group (Figures 2B and 2C and Figure S3C). This is in line with our previous results, showing that pre-existing memory frequencies increase with age and correlate with the relative proportion of memory cells within the total CD4+ T cells, reflecting the cumulative immunological experience of an individual (Bacher et al., 2020).

To determine the potential impact of age and pre-existing memory in more detail, we compared different age groups within our cohort. Representative young (<25 years) and old (>80 years) individuals are depicted in Figure 2D, highlighting that the proportion of spike-reactive memory versus naive T cells recapitulates the memory-naive distribution in the total CD4+ T cell compartment (Figure 2D), as shown before (Bacher et al., 2020). Overall, the frequency of pre-existing memory T cells was variable between individuals but tended to increase with age and was particularly high in donors >80 years of age (Figure 2E). However, upon vaccination, spike-specific T cells poorly expanded in this age group, as shown for the representative young and old donor (Figure 2F) as well as for the whole cohort (Figure 2G). Consistent with this, the >80 years age group had the lowest frequency of spike-specific T cells post-vaccination (Figure 2H), as well as the lowest anti-spike IgG concentrations and SARS-CoV-2-neutralizing antibody titers (Figures 2I and 2J), despite having started with the highest frequencies of pre-existing memory cells pre-vaccination (Figure 2E). Spike-specific CD4+ T cells remained low even after the third vaccination in the >80 years age group, in contrast to anti-spike IgG antibodies, which reached similar concentrations as in the other age groups (Figures 2H and 2I). Thus, donor age has a strong negative influence on T cell response quality despite high frequencies of pre-existing memory T cells.

High pre-existing memory and a low proportion of naive T cells predict poor T cell responses

High pre-existing memory T cell frequencies were mainly found in the elderly, but not exclusively (Figure 2E). We therefore
analyzed whether pre-existing memory is an independent predictor of the T cell response quality. To dissect the role of pre-existing memory independently of the effects of high age, the >80 years group was depicted separately. Indeed, the Tmem frequency showed a strong negative correlation with the fold expansion of spike-specific T cells (Figure 3A), which was confirmed for each individual age group (Figure S3D). This was also obvious when donors were divided into several bins based on increasing numbers of pre-existing memory cells (Figure 3B). Accordingly, donors with the lowest frequencies of pre-existing memory showed the highest production of IL-21 (Figure 3C) and a tendency toward higher functional avidity, although not statistically significant (Figure 3D). These data indicate that pre-existing memory T cells expand poorly upon SARS-CoV-2
vaccination but also suggest that additional parameters may determine the selection of high-avidity T cells.

SARS-CoV-2-specific T cells within the naive pre-vaccination repertoire have so far not been analyzed, but the high sensitivity of the ARTE technology allows us to do this. In general, we observed a negative correlation between the frequencies of pre-existing memory cells and the percentage of naive cells within the spike-reactive pre-vaccination T cell pool (Figure S3E). Furthermore, the proportion of spike-reactive naive T cells showed a positive correlation with the expansion rate following vaccination (Figures 3E and 3F). The pre-vaccination naive pool size also correlated with IL-21 production (Figure 3G) as well as the functional avidity of spike-specific T cells post-vaccination (Figure 3H). Indeed, frequencies of IL-21 producers and high-avidity responses significantly dropped when the proportion of pre-vaccination spike-specific naive T cells was below 20% (Figures 3G and 3H). These data suggest that high-avidity T cells with high expansion potential are preferentially recruited from the naive compartment.

Overall, our data indicate that the vaccination success, defined by formation of a highly expanded, IL-21-producing, high-avidity T cell pool, is predicted by two interdependent, but not strictly linked, parameters: the frequency of pre-existing spike-specific memory T cells, mainly determined by the immunological experience of an individual, and the proportion of naive T cells within the pre-vaccination spike-specific T cell pool determined by thymic activity.

**Pre-existing memory T cell clonotypes minimally contribute to high-quality T cell responses**

Our results so far contradict the expectation that pre-existing memory T cells support immune responses against SARS-CoV-2 vaccination. To directly assess the contribution of pre-existing memory T cells to the post-vaccination repertoire, we performed clonal TCR tracking using TCR sequencing from ex vivo sorted spike-reactive memory T cells from 37 donors pre- and post-vaccination (Table S3). Consistent with their predicted low expansion capacity, pre-existing memory TCR-β chains accounted on average for 6.1% of the top 100 clonotypes post-vaccination, while most of the post-vaccination clonotypes were not detected at baseline (Figure 4A). Also, the cumulative abundance of pre-existing clonotypes contributing to the top 100 post-vaccination repertoire was on average only 10%. However, in two donors, pre-existing clonotypes made up more than 25% of the post-vaccination repertoire (Figure 4A). Analysis of the TCR-α repertoire showed similar results (Figure S4A). Next, we used the TCRdist algorithm to identify closely related clonotypes based on shared CDR3 sequence features (Dash et al., 2011).

**Figure 3. Pre-existing memory cells expand poorly upon vaccination**

(A) Spearman correlation between the frequencies of spike-reactive Tmems pre-vaccination (x axis) and the fold expansion post-second vaccination (y axis). (B–D) Donors were grouped by their pre-vaccination frequencies of spike-reactive Tmems (<1.5, n = 10; 1.5–2.5, n = 11; 2.5–5, n = 10; >5, n = 9). Donors >80 years of age (n = 6) are depicted as separate group. For each bin, the (B) fold expansion post-second vaccination, (C) percentage of IL-21 production within CD154+ Tmems, and (D) functional avidity of expanded spike-reactive T cells is indicated. (E) Spearman correlation between the proportion of spike-reactive naive cells pre-vaccination (x axis) and the fold expansion post-second vaccination (y axis). (F–H) Donors were grouped by their pre-vaccination proportion of spike-reactive naive cells (<20, n = 9; 20–30, n = 11; 30–45, n = 10; >45, n = 11). Donors >80 years of age (n = 6) are depicted as separate group. For each bin, the (F) fold expansion post-second vaccination, (G) percentage of IL-21 production within CD154+ Tmems, and (H) functional avidity of expanded spike-reactive T cells is indicated. Each symbol in (A–H) represents one donor; horizontal lines indicate mean in (B, C, F, G) and geometric mean in (D, H). Statistical differences: Kruskal-Wallis test with Dunn’s post hoc test in (B–D, F–H). To determine differences between the younger age groups, donors >80 years of age were excluded from statistical analysis. See also Figure S3.
Figure 4. TCR sequencing post-vaccination identifies primarily newly arising clonotypes
(A–C) Spike-reactive memory T cells were ex vivo sorted pre- and post-vaccination and analyzed by bulk TCR sequencing. Analysis of the top 100 expanded TCR−β spike-reactive clonotypes post-second vaccination is shown.
(A) The number of clonotypes (left graph) and their cumulative relative abundance (right graph) derived from the pre-existing, post-first or post-second vaccination repertoire is indicated. Each bar represents one donor (n = 37).
(B) Cumulative abundance of pre-existing or newly arising clonotypes found in the top 100 TCR repertoire post-second vaccination for different age groups (>25, n = 9; 26–39, n = 14; 40–60, n = 9, >80, n = 5).

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We identified several spike-reactive meta-clonotypes within our dataset (Figures S4B and S4C and Table S3) and also re-identified public immunodominant clusters previously reported in COVID-19 patients and SARS-CoV-2 vaccinees (Mudd et al., 2022; Rosati et al., 2021), confirming the spike specificity of our clonotypes. Also, within the biggest similarity clusters, most clonotypes were newly arising, while pre-existing clonotypes belonging to these clusters were only found in individual donors (Figure S4D).

Since the frequencies of pre-existing memory cells were particularly elevated in the elderly (Figure 2E), we next compared the recruitment of pre-existing clonotypes into the vaccine response in the different age groups (Figure 4B). There was no significant difference between age groups, and unexpectedly, the >80 years group showed a rather low contribution of pre-existing memory clonotypes to the post-vaccine response despite having the highest frequencies before vaccination (Figure 4B).

To better define the pre-existing memory dynamics, we analyzed the clonal fate in each individual donor. Clonal fate tracking is displayed as circos plots showing clonotypes pre-vaccination and the top 100 clonotypes post-second vaccination. Connecting lines show TCR-β sequences that are present at both time points, and the width of the connecting line indicates the relative size of each clone in the pre- or post-vaccination repertoire, respectively (Figures 4C and S5). Grouping donors according to the cumulative abundance of pre-existing clonotypes in the post-vaccination repertoire (<5%, 5–15%, >15%) (Figure S5A) correlated with three different response patterns (Figures 4D and 4E). In the “low contribution” group, the abundance of pre-existing clonotypes in the post-vaccination repertoire is negligible (<5%), whereas in the “hierarchy shift” group (5–15%), strongly expanded clonotypes pre-vaccination represent rather subdominant clonotypes post-vaccination. Thus, the T cell response in these groups is dominated by newly arising, strongly expanding clonotypes, most likely from the naïve compartment. In contrast, in the third, “high contribution” group (>15%), strongly expanded pre-existing memory clonotypes remain dominant post-vaccination, suggesting a low contribution of newly arising clonotypes. Donors >80 years of age mainly fell into the low contribution group, but to enable differentiation between age and pre-existing memory-mediated effects, they are further depicted separately.

Within these different response patterns, we next analyzed the independently assessed spike-specific vaccine-response quality parameters (Figures 4D and 4E). The low contribution group had the highest T cell fold expansion, as well as high IL-21 production after vaccination, while showing the highest proportion of spike-reactive naïve cells and the lowest frequencies of pre-existing memory cells pre-vaccination (Figures 4D and 4E). Furthermore, high and more stable concentrations of anti-spike IgG and SARS-CoV-2-neutralizing antibodies were found in this group (Figure 4E). Despite a tendency toward a slight quality loss, the hierarchy switch group was most similar to the low contribution group, indicating an overall high-quality vaccine response. In contrast, the high contribution group showed a significant decline in several quality parameters, such as low fold expansion, low IL-21, and a tendency toward reduced IgG stability while having a high frequency of pre-existing memory cells. The high contribution group was in many parameters not statistically different from the >80 years group, which displayed uniformly poor response quality (Figure 4E).

These data support the hypothesis that a diverse naïve compartment, allowing the selection of high-affinity T cells, is required for high-quality SARS-CoV-2 T cell responses rather than pre-existing memory T cells. To better quantify this effect, we analyzed the TCR diversity pre- and post-vaccination. The diversity of pre-existing memory before vaccination was not different between the groups. However, especially after the first, but also after the second, vaccination, a diverse SARS-CoV-2-specific repertoire developed in the low contribution and hierarchy switch groups, consistent with recruitment of multiple clonotypes from a diverse naïve repertoire. In contrast, the high contribution group, and in particular the “>80 years” donors, displayed a lower TCR diversity (Figure 4F). Also, when only analyzing the newly arising clonotypes, the >80 years group still had a reduced diversity, suggesting a limited and poorly expanding naïve pre-vaccination repertoire in the elderly. This is further supported by a strong reduction of the naïve T cell pool

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(C) Each circos plot represents the TCR-β sequences of the detected clonotypes pre-vaccination and the top 100 clonotypes post-second vaccination of one donor. Connecting lines show TCR-β sequences that are present at both time points. The fold expansion, the absolute number of clonotypes detected pre-vaccination, and the cumulative abundance of pre-existing clonotypes in the post-vaccine repertoire are indicated for each representative donor.

(D) Donors were grouped according to the cumulative abundance of pre-existing clonotypes in the top 100 post-vaccination repertoire (<5%, n = 11; 5–15%, n = 11; >15%, n = 9; donors >80 years of age, n = 5). Mean values for the indicated parameters were calculated, Z score normalized for each parameter and are shown as heatmap.

(E) Donors were grouped as described in (D) and the fold expansion, IL-21 production within CD154+ Tmems, the proportion of spike-reactive naïve T cells pre-vaccination, the frequencies of pre-existing Tmems, and anti-spike IgG concentrations or SARS-CoV-2 serum neutralization titers post-second vaccination are indicated.

(F) Renyi diversity profiles were calculated for mean values of the different pre-existing memory bins (<5%, n = 11; 5–15%, n = 11; >15%, n = 9; >80 y, n = 9). By varying the scaling parameter s, different diversity indices are calculated. At values of 0, 1, 2, and infinite richness, Shannon diversity, Simpson diversity, and Berger-Parker index are calculated (see STAR Methods). Thus, the sample with the highest value at s = infinite indicates a higher proportion of the most abundant TCRs, i.e., lower population diversity. A sample with a profile that is overall higher than the profiles of other samples is therefore more diverse. Diversity of the total spike-specific TCR-β repertoire pre-vaccination, post-first, and post-second vaccination as well as of the newly arising clonotypes post-second vaccination are shown.

(G) Representative CD45RA and CD31 staining of a young and old individual.

(H) Proportion of naive (CD45RA+CCR7+) within total CD4+ T cells in different age groups.

(I) Proportion of RTE (CD45RA+CD31+) within total CD4+ T cells in different age groups. Each symbol in (B, E, H, I) represents one donor. Box-and-whisker plots display quartiles and range in (B); truncated violin plots with quartiles and range are shown in (E). Symbols and error bars indicate mean and SEM in (F). Horizontal lines indicate mean in (H, I). Statistical differences: Kruskal-Wallis test with Dunn’s post hoc test in (E, F, H, I). Also see Figures S4 and S5.
and in particular of recent thymic emigrants (RTEs), especially in individuals >80 years of age (Figures 4G–4I), which is a well-known mechanism of immune aging (Mittelbrunn and Kroemer, 2021).

In summary, these data indicate how the contribution of pre-existing memory and naive T cells differentially affects the immune-response quality following vaccination and how this may be altered upon aging. High-quality immune responses were in general connected to a low contribution of pre-existing memory and a dominant contribution of newly arising clonotypes with the capacity to strongly expand. In contrast, pre-existing memory T cells seem to have low proliferative potential, leading to low-quality responses if not compensated, e.g., from the naive pool, like in the hierarchy shift group. This is particularly evident in the >80 years group, which showed a loss of proliferative capacity in the pre-existing memory, as well as reduced TCR diversity in the newly arising clonotypes, suggesting impaired recruitment from a reduced naive compartment.

The decline in T cell reactivity against the Omicron variant is variable between individuals

T cell responses are less affected by virus variants, such as Omicron B.1.1.529, compared with antibody responses due to the recognition of many different epitopes. However, low-quality T cell responses with restricted TCR diversity may be particularly at risk. We therefore analyzed the response against 80 selected spike peptides affected by Omicron B.1.1.529 BA.1 mutations (“Omicron peptides”) and their corresponding “wild-type peptides” (Figure 5A and Table S4). In vaccinated donors, the wild-type peptide pool accounted on average for 30% of the T cell response against the total spike protein and was further decreased by an average of 50% against the Omicron peptides (Figure 5B). The Omicron peptides stimulated a higher fraction of naive T cells (Figure 5C), as expected, due to sequence changes arising by the mutations, resulting in new T cell epitopes not being present in the wild-type spike used for vaccination (see Table S4).

In line with our previous results, T cell reactivity against the wild-type and Omicron peptides declined with donor age (Figure 5D). However, we did not observe a particular decline in the >80 years group, nor in individuals with a high contribution of pre-existing memory, but rather we observed high variability.
Figure 6. CCCoV cross-reactive T cells barely contribute to the SARS-CoV-2 vaccine response

(A–C) SARS-CoV-2 spike-reactive CD154⁺ Tmems were expanded post-second vaccination and restimulated in presence of autologous antigen-presenting cells with different antigens, including individual spike proteins from CCCoVs or a pool thereof.

[Legend continued on next page]
between individuals (Figures 5D and 5E). In summary, these data show that all age groups are similarly affected by the loss of reactivity against the Omicron variant. Larger cohorts are needed to clarify in the future whether the decline of reactivity against variants has stronger consequences in donors with low T cell response quality.

**CCCoV cross-reactive T cells are not major contributors to the anti-SARS-CoV-2 vaccine response**

Our data suggest that pre-existing memory does not strongly contribute to a high-quality vaccination response. However, pre-existing memory induced by related CCCoVs has been suggested to contribute to SARS-CoV-2 immunity upon infection and vaccination. To specifically determine the contribution of CCCoV-cross-reactive T cells to the SARS-CoV-2 spike-specific T cell response, we expanded SARS-CoV-2 spike-specific cells after the second vaccination. Restimulation proved high reactivity against the SARS-CoV-2 spike protein, but only low cross-reactivity to spike proteins from the CCCoV strains 229E, NL63, OC43, and HKU1, which was similar to other common viral antigens in individual donors (Figures 6A and 6B). Based on the reactivity to SARS-CoV-2 spike proteins and a pool of all four CCCoV spike proteins, we calculated the percentage of cross-reactivity for each donor. This revealed on average 5% cross-reactivity of the SARS-CoV-2 spike-specific T cells against CCCoVs upon vaccination, which was in a similar magnitude, as observed before, in COVID-19 patients (Figure 6C; Bacher et al., 2020).

Conversely, we analyzed whether vaccination affects the frequencies of T cells reactive against the spike proteins of CCCoVs, specifically the alpha coronavirus NL63 and the beta coronavirus OC43. As a high control for a cross-reactive response, we included the human adenovirus 5 hexon protein and restricted our analysis to the vaccinees receiving the first immunization with the adenovirus-vector-based ChAdOx1 vaccine, followed by Moderna mRNA-1273 (Figure 1A). Sequence alignment of the hexon protein from chimpanzee adenovirus Y25, which was used as a vector for ChAdOx1, showed high sequence homology (80% identity) with the human AdV5 hexon, whereas SARS-Cov-2 and CCCoV spike proteins had 30% sequence identity (Figure S6). Despite a 1–2 log increase of SARS-CoV-2 spike-specific T cells, CCCoV spike-reactive T cell frequencies remained stable post-vaccination (Figures 6D and 6E). In contrast, the response to the adenovirus hexon protein was increased after ChAdOx1 vaccination and remained stable afterward (Figures 6D and 6E). This indicates that cross-reactive T cells are increased upon vaccination when sequence homology is high enough, as it is the case for adenovirus-hexon-specific cells.

In summary, these data further indicate that CCCoV cross-reactive T cells are not the major drivers of the SARS-CoV-2 spike-specific CD4+ T cells upon vaccination, as previously also shown for SARS-CoV-2 infection (Bacher et al., 2020; Low et al., 2021; Mateus et al., 2020).

**DISCUSSION**

High immune-response variability is a critical feature of SARS-CoV-2 infection and vaccination. Understanding the individual intrinsic parameters determining the immune-response quality is key to identifying high-risk subgroups and developing protective measures. So far, it is poorly understood as to how the antigen-specific pre-exposure T cell repertoire impacts immune-response quality and how this is affected by age as the most relevant risk factor for poor protection. Our data showed that the response to SARS-CoV-2 vaccination is determined by the pre-exposure T cell repertoire, comprising two separate entities with opposing effects: SARS-CoV-2-specific naive and pre-existing memory T cells. Both are inversely affected by aging and together identify individuals at risk for impaired immunity.

Technical limitations in tracking rare antigen-specific T cells in unexposed individuals have restricted the study of the effect of the pre-vaccination repertoire on the immune-response quality. Rare cell-enrichment approaches allow identification of naive and memory T cells in the pre-exposure repertoire (Bacher et al., 2013, 2016, 2019, 2020; Moon et al., 2007; Obar et al., 2008; Pan et al., 2021; Su et al., 2013). Without enrichment, the detection is mainly restricted to high-frequency memory T cells and only in a variable fraction of unexposed donors (Bacher and Scheffold, 2013; Braun et al., 2020; Grifoni et al., 2008; Pan et al., 2021; Loyal et al., 2021; Mateus et al., 2021; Ogbe et al., 2021). By using ARTE, we identified the antigen-specific CD4+ T cell expansion rate as a critical parameter for vaccine-response quality, which was directly linked to the pre-vaccination repertoire. Rare T cell precursors with high expansion potential, likely from the naive repertoire, supported strong expansion upon vaccination, whereas pre-existing memory cells counteracted it.

Our data identified two interdependent, but not directly linked, parameters affecting immune-response quality: first, in the elderly, CD4+ T cell expansion of both the naive and memory compartment was severely compromised. This is in line with the current concept that aging has a strong effect on the T cell repertoire, with a decline in naive T cells and an impaired memory compartment (Mittelbrunn and Kroemer, 2021), which we confirmed here for SARS-CoV-2-specific T cells. Second, similar repertoire restrictions also occurred in individuals from younger age groups.
age groups with a high contribution of pre-existing memory cells, suggesting that this is a general parameter determining immune-response quality to SARS-CoV-2.

While the importance of the naive compartment is well accepted, the in vivo contribution of pre-existing memory cells is not clear (Lipsitch et al., 2020; Moss, 2022; Woodward and Blackman, 2006). Pre-existing memory T cells have been described for SARS-CoV-2 (Bacher et al., 2020; Bartolo et al., 2021; Braun et al., 2020; Le Bert et al., 2020; Loyal et al., 2021; Mateus et al., 2020; Neide et al., 2020), but also for many other pathogens (Bacher et al., 2013; Campion et al., 2014; Kwok et al., 2012; Pan et al., 2021; Su et al., 2013). Conflicting data exists regarding the age-associated prevalence of SARS-CoV-2-reactive pre-existing memory T cells (Bacher et al., 2020; Dowell et al., 2022; Loyal et al., 2021; Saletti et al., 2020). Recent correlative data suggested that pre-existing memory T cells may support an earlier and stronger anti-SARS-CoV-2 T and B cell re-action following infection or vaccination (Kundu et al., 2022; Loyal et al., 2021; Mateus et al., 2021; Meyer-Arndt et al., 2022; Swardling et al., 2022). A limitation of these studies is that they are based on technologies that only incompletely resolve the pre-vaccination repertoire, and the contribution of individual clonotypes to the in vivo response was not analyzed. By using clonal TCR tracking, we revealed a minimal contribution of pre-existing memory T cells in most individuals, which correlated with high T cell expansion. This indicates that in high-quality responders, pre-existing memory clones are surpassed by multiple strongly expanding low-frequency precursors, most likely from the naive pool. A similar clonal hierarchy shift was recently demonstrated for yellow fever vaccination (Pan et al., 2021). In contrast, individuals where pre-existing memory clonotypes remained dominant after vaccination displayed rather poor responses and a lower TCR diversity, as previously shown in mice (Cornberg et al., 2006). Low-avidity pre-existing memory cells may prevent naive T cell recruitment and high-affinity selection (Lanzer et al., 2014). Overall, this emphasizes the central importance of strong clonal expansions of diverse and high-avidity precursors that together form a high-quality T cell response.

A diverse TCR repertoire, rather than T cell abundance, was shown to be protective in pathogen-specific immunity (Wang et al., 2012). Furthermore, a limited TCR diversity has been associated with escape of viral mutants (Cornberg et al., 2006; Meyer-Olson et al., 2004). However, with our small number of analyzed individuals, we did not observe a particular reduction of the CD4+ T cell response to the B.1.1.529 Omicron variant in the elderly or in younger donors with low TCR diversity due to a high contribution of pre-existing memory. Instead, the decline against the Omicron variant was highly variable between the donors. This suggests that additional parameters such as precise epitope specificities or human leukocyte antigen (HLA) alleles might be important for successful adaption to viral variants. Nevertheless, given the already low frequencies and quality of spike-specific T cells in individual donors and especially the >80 years group, a further reduction by viral variants such as Omicron suggests a remaining vulnerability of these individuals despite vaccination. Clearly, larger cohorts are needed to address this point in more depth.

A protective role of pre-existing memory T cells has initially been proposed due to cross-reactivity to related CC-CoVs. However, CCCoV-reactive T cells only account for a variable part of the pre-existing memory to SARS-CoV-2 (Bacher et al., 2020; Bartolo et al., 2021). CCCoV cross-reactivity was reduced following vaccination, as previously demonstrated for SARS-CoV-2 infection (Bacher et al., 2020; Mateus et al., 2020), most likely due to low avidity of cross-reactive TCRs (Bacher et al., 2020; Dykema et al., 2021). In contrast, we found strong expansion of adenovirus hexon-specific memory T cells by the adenovector-based AstraZeneca ChAdOx1 vaccine. Similarly, clonal tracking after influenza vaccination shows strong involvement of memory T cells (Hill et al., 2021; Nienen et al., 2019). Thus, the role of pre-existing T cell memory is context dependent: effective cross-reactive T cell memory activation in vivo appears to depend on high affinity to homologous peptides, which seems to apply only to a minority of SARS-CoV-2 epitopes. Whether such rare pre-existing TCRs can directly contribute to an earlier response to SARS-CoV-2 in individual donors (Loyal et al., 2021; Mateus et al., 2021) remains to be further investigated.

High age is also a risk factor for severe COVID-19. SARS-CoV-2-specific CD4+ T cell frequencies are increased in severe COVID-19 (Moss, 2022), whereas decreased frequencies are found in old vaccinees (Anderson et al., 2020; Collier et al., 2021; Li et al., 2021; Romero-Olmedo et al., 2022a). However, in both situations, we identified similar low-avidity T cell responses (Bacher et al., 2020). This suggests that the recruitment of low-avidity T cells, presumably from an impaired pre-exposure repertoire, is a common denominator of low-avidity T cell responses. A higher dose, dissemination, and persistence of viral antigen or stronger costimulatory signals may account for the increased frequency of low-avidity SARS-CoV-2-specific T cells in severe COVID-19, as compared to vaccination.

In summary, a compromised pre-exposure repertoire determined low-quality CD4+ T cell responses, which is the common feature of poor vaccine responses, as well as severe COVID-19. Importantly, such a T cell repertoire-intrinsic defect may not be rescued by repeated vaccination (Romero-Olmedo et al., 2022a, 2022b), but vice versa: a low-avidity T cell response to vaccination may explain why the elderly remain at higher risk for developing severe disease (Yek et al., 2022). Thus, our results highlight the need to develop novel vaccination strategies targeting or diversifying restricted T cell repertoires in individuals at risk.

**Limitations of the study**

Our study is restricted to only a small number of individuals within the different age groups due to the high experimental effort required for antigen-specific T cell enrichment and clonal tracking. Clonal tracking in the pre-immune repertoire is technically challenging and restricted to abundant precursor clones. Therefore, small precursor clones that might contribute to the post-vaccination response might be missed. Since we could not analyze the pre-vaccine naive TCR repertoire due to technical limitations in tracking low numbers of cells with low clonality, the direct origin of the rare, strongly proliferating clones in high-quality responders remains undefined, and a potential contribution of tissue-derived or stem cell memory cells cannot be excluded. Furthermore, the antigens driving the pre-existing memory response in different individuals remain unknown. Our analysis describes the average contribution of the pre-existing
memory population as well as the average of T cell quality parameters such as fold expansion and functional avidity. Individual T cell clones also may act differently in individual donors, and a positive contribution of individual pre-existing T cell clones to SARS-CoV-2 immunity remains to be further elucidated.

**STAR METHODS**

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**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at https://doi.org/10.1016/j.immuni.2022.08.003.

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**AUTHOR CONTRIBUTIONS**

Conceptualization: P.B., A.S.; investigation: C.Sa., G.R.M., E.R., S.M., B.M., A.K.K., N.B., J.G., R.R., M.V., U.M.G., H.M.R., P.B.; formal analysis: P.B., E.R.; resources: M.S., F.T., D.K.B., A.K., H.F., O.K., Y.K., M.J.G.T.V., O.C., P.K., A.F., C.Sch., S.S., B.H.; visualization: P.B., E.R.; supervision: A.K., H.F., A.F., S.S., B.H., A.S.; funding acquisition: P.B., A.S.; all authors provided discussion, participated in revising the manuscript, and agreed to the final version.

**DECLARATION OF INTERESTS**

E.R. is employee of Evotec SE. O.K. is employee of Peptides & elephants GmbH. O.C. reports grants or contracts from MSD, Pfizer; consulting fees from Biocon, Molecular Partners, Noxxon, PSI; honoraria for lectures from Abbott, Pfizer; participation on a data safety monitoring board or advisory board from Janssen, PSI, P.K. reports grants or contracts from BMBF, B-FAST, and NAPKON of the Network University Medicine and the State of North Rhine-Westphalia; consulting fees from Ambu GmbH, Gilead Sciences, Noxxon N.V., Pfizer Pharma; honoraria for lectures from Ambu GmbH, Gilead Sciences, MSD Sharp & Dohme GmbH, Pfizer Pharma GmbH, Scilink Comunicacion Cientifica SC; participation on an advisory board from Ambu GmbH, Gilead Sciences, Pfizer Pharma.

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### STAR METHODS

#### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| CD4-APC-Vio770 (clone: M-T466) | Miltenyi Biotec | Cat#130-113-251 RRID:AB_2726053 |
| CD8-VioGreen (clone: REA734) | Miltenyi Biotec | Cat#130-110-684, RRID:AB_2659241 |
| CD14-VioGreen (clone: REA599) | Miltenyi Biotec | Cat#130-110-525 RRID:AB_2655057 |
| CD20-VioGreen (clone: LT20) | Miltenyi Biotec | Cat#130-096-904 RRID:AB_2726147 |
| CD38-PE-Vio770 (clone: REA572) | Miltenyi Biotec | Cat#130-113-432 RRID:AB_2733228 |
| CD45RA-PE-Cy5 (clone: HI100) | BioLegend | Cat#304110 RRID:AB_314414 |
| CCR7-Brilliant Violet 785 (clone: G043H7) | BioLegend | Cat#353230 RRID:AB_2563630 |
| CD154-FITC (clone: REA238) | Miltenyi Biotec | Cat#130-109-469 RRID:AB_2751146 |
| IL-21-PE (clone: REA1039) | Miltenyi Biotec | Cat#130-117-421, RRID:AB_2727941 |
| TNFa-Brilliant Violet 650 (clone: MAb11) | BioLegend | Cat#502938 RRID:AB_2562741 |
| IL-10-PE-Dazzle (clone: JES3-9D7) | BioLegend | Cat#501426 RRID:AB_2566744 |
| IFN-γ-PerCP-Cy5.5 (clone: 4S.B3) | BioLegend | Cat#502526 RRID:AB_961355 |
| Ki-67 Alexa Fluor 700 (clone: B56) | BD Biosciences | Cat#561277 RRID:AB_10611571 |
| IL-2-BV711 (clone: 5344.111) | BD Biosciences | Cat#563946 RRID:AB_2738501 |
| CD31-APC-Vio770 (clone: REA730) | Miltenyi Biotec | Cat#130-110-810 RRID:AB_2657290 |
| CD69-PE (clone: REA824) | Miltenyi Biotec | Cat#130-112-613 RRID:AB_2659065 |
| CD4-VioBlue (clone: VIT4) | Miltenyi Biotec | Cat#130-113-219 RRID:AB_2726030 |
| CD154-APC (clone: 5C8) | Miltenyi Biotec | Cat#130-113-603, RRID:AB_2726191 |
| TNF-α-PE-Vio770 (clone: cA2) | Miltenyi Biotec | Cat#130-096-755 RRID:AB_2784483 |
| CD4-BV421 (clone: OKT4) | BioLegend | Cat#317434 RRID:AB_2562134 |
| CD154 MicroBead Kit, human | Miltenyi Biotec | Cat#130-092-658 |
| CD28 pure – functional grade, human (clone: 15E8) | Miltenyi Biotec | Cat#130-093-375 RRID:AB_1036134 |
| CD40 pure – functional grade, human (clone: HB14) | Miltenyi Biotec | Cat#130-094-133 RRID:AB_10839704 |
| CD14 MicroBeads, human | Miltenyi Biotec | Cat#130-050-201 RRID:AB_2665482 |

| **Biological samples** |        |            |
|-----------------------|--------|------------|
| Fresh human EDTA blood samples | N/A | N/A |

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| REAGENT or RESOURCE                      | SOURCE                      | IDENTIFIER |
|------------------------------------------|-----------------------------|------------|
| **Chemicals, peptides, and recombinant proteins** |                             |            |
| Human AB Serum                           | Sigma Aldrich               | Cat#H4522  |
| RPMI-1640 medium                         | Gibco, Life Technologies    | Cat#61870-044 |
| TexMACS medium                           | Miltenyi Biotec             | Cat#130-097-196 |
| X-Vivo15 medium                          | Lonza                       | Cat#BE02-060F |
| Human IL-2 (Proleukin)                   | Novartis                    | N/A        |
| Human IL-4                                | Miltenyi Biotec             | Cat#130-093-922 |
| Human GM-CSF                              | Miltenyi Biotec             | Cat#130-093-866 |
| Brefeldin A                              | Sigma Aldrich               | Cat#B7651  |
| **Critical commercial assays**            |                             |            |
| MS Columns                               | Miltenyi Biotec             | Cat#130-042-201 |
| LS Columns                               | Miltenyi Biotec             | Cat#130-042-401 |
| Viobility 405/520 Fixable Dye            | Miltenyi Biotec             | Cat#130-110-206 |
| Inside Stain Kit                         | Miltenyi Biotec             | Cat#130-090-477 |
| RNeasy Micro Kit                         | Qiagen                      | Cat#74004  |
| Human TCR Profiling Kit                  | Milaboratory                | Cat#TH-034-564252 |
| SMARTScribe Reverse Transcriptase        | Clontech                    | Cat#639538 |
| RNasin Ribonuclease Inhibitor            | Promega                     | Cat#N2511  |
| Uracil-DNA Glycosylase                   | New England Biolabs         | Cat#M0280S |
| Q5 Hot Start High-Fidelity DNA Polymerase| New England Biolabs         | Cat#M0493L |
| Anti-Spike IgG SARS-CoV-2-QuantiVac ELISA| Euroimmun                   | Cat#EI 2606-9601-10 G |
| SARS-CoV-2 NP IgG                        | AESKULISA                   | Cat#6122   |
| SARS-CoV-2 Neutralizing Ab ELISA Kit     | Invitrogen                  | Cat#BMS2326 |
| **Experimental models: Cell lines**       |                             |            |
| Vero cells                               | CLS Cell Lines Service GmbH | Cat#605372 |
| **Software and algorithms**              |                             |            |
| FlowJo                                   | Treestar                    |            |
| GraphPad Prism 9.3.1.                    | GraphPad                    |            |
| Genesis                                  | (Sturn et al., 2002)        |            |
| MIGEC MiGEC version 1.2.6                | (Shugay et al., 2014)       | N/A        |
| MiXCR version 3.0.14                     | (Bolotin et al., 2015)      | N/A        |
| R package “vegan”                        | (Oksanen et al., 2019)      | N/A        |
| R packages “ineq”                        | (Zeileis, 2014)             | N/A        |
| Ggplot2                                  | (Wickham, 2016)             | N/A        |
| **Other**                                |                             |            |
| PepMix™ SARS-CoV-2 (Spike Glycoprotein)  | JPT                         | Cat#PM-WCPV-S |
| SARS-CoV-2 B.1.1.529 BA.1 omicron peptides | Peptides& Elephants       | Cat#LB01999 |
| SARS-CoV-2 wildtype peptides             | Peptides& Elephants         | Cat#LB02004 |
| PepMix™ HCoV- HKU1 (Spike Glycoprotein)  | JPT                         | Cat#PM-HKU1-S-1 |
| PepMix™ HCoV-NL63 (Spike Glycoprotein)   | JPT                         | Cat#PM-NL63-S-1 |
| PepMix™ HCoV-OC43 (Spike Glycoprotein)   | JPT                         | Cat# PM-OC43-S-1 |
| PepMix™ HCoV-229E (Spike Glycoprotein)   | JPT                         | Cat# PM-229E-S-1 |
| PepTivator CMV pp65                      | Miltenyi Biotec             | Cat#130-093-438 |
| PepTivator CMV IE-1                      | Miltenyi Biotec             | Cat#130-093-493 |
| PepTivator Influenza A (H1N1) NA         | Miltenyi Biotec             | Cat#130-099-806 |
| PepTivator Influenza A (H1N1) MP1        | Miltenyi Biotec             | Cat#130-097-285 |

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RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Petra Bacher (p.bacher@ikmb.uni-kiel.de).

Materials availability
This study did not generate new unique reagents.

Data and code availability
- TCR sequencing data generated during this study are provided in Table S3.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Blood donors
Vaccine blood donors were recruited following approvals by the Institutional Review Board of UKSH Kiel (identifier D409/21) and Lübeck (identifier IRON AZ 15–304). SARS-CoV-2 naïve vaccinees were enrolled between December 2020 and July 2021. Convalescent COVID-19 patients were enrolled between April and September 2020 (ethics committee of the UKSH Kiel (Identifier D474/20); University Hospital Frankfurt (Identifier 11/17); University of Cologne (Identifier 20–1295; 08–160) (Bacher et al., 2020). All blood donors signed informed consents. See Table S2 for demographic information on vaccine blood donors.

Vero cells
A Vero cell-based virus-neutralization test (cVNT) was used for measurement of SARS-CoV-2-neutralizing antibodies (see below). Vero cells (CLS Cell Lines Service GmbH, Eppelheim, Germany, order no. 605372) were incubated at 37 °C, 5% CO2 and 90% humidity in Dulbecco’s modified Eagle’s medium supplemented with 10% (v/v) fetal calf serum, 3.7 g/L NaHCO3, 4.5 g/L glucose, 2 mM L-glutamine, and 1% (v/v) of Pen-Strep-Fungi mix containing 10,000 U/ml penicillin, 10 mg/mL streptomycin, and 25 µg/mL amphotericin B (all reagents from Bio&SELL GmbH, Feucht, Germany).

METHOD DETAILS

Antigens
Pools of lyophilized 15-mer peptides with 11–amino acid overlap, covering the complete protein sequence were purchased from JPT (Berlin, Germany): SARS-CoV-2 spike, 229E spike, OC43 spike, HKU1 spike, NL63 spike and Miltenyi Biotec (Bergisch Gladbach, Germany): Influenza A H1N1 (HA, MP1, MP2, NP and NA), CMV (pp65, IE-1), EBV (EBNA1, BZLF1, LMP2A, LMP1), AdV (Hexon). Peptides covering the sequence regions of the SARS-CoV-2 omicron variant B.1.1.529 BA.1 and the corresponding wildtype peptides were obtained from peptides&elephants (Hennigsdorf, Germany).

Peptides were resuspended according to manufacturers’ instructions and cells were stimulated at a concentration of 0.5 µg/peptide/ml until otherwise indicated in the figures and figure legends. Tetanus-toxoid was purchased from Statens Serum Institute and used at a concentration of 10 µg/mL.

Antigen-reactive T cell enrichment (ARTE)
Peripheral blood mononuclear cells were freshly isolated from EDTA blood on the day of blood donation by density gradient centrifugation (Biocoll; Biochrom, Berlin, Germany). Antigen-reactive T cell enrichment (ARTE) was performed as previously described.
(Bacher et al., 2013, 2016, 2019, 2020). In brief, 1-2×10^7 PBMCs were plated in RPMI-1640 medium ( Gibco), supplemented with 5% (v/v) human AB-serum ( Sigma Aldrich, Schnelldorf, Germany) at a cell density of 1 × 10^6 PBMCs/2 cm² in cell culture plates and stimulated for 7 h in presence of 1 µg/ml CD40 and 1 µg/ml CD28 pure antibody (both Miltenyi Biotec, Bergisch Gladbach, Germany). 1 µg/ml Brefeldin A ( Sigma Aldrich) was added for the last 2 h.

Cells were labeled with CD154-Biotin followed by anti-Biotin (CD154 MicroBead Kit, Miltenyi Biotec) and magnetically enriched by two sequential MS columns (Miltenyi Biotec). Surface staining was performed on the first column, followed by fixation and intracellular staining on the second column. Frequencies of antigen-specific T cells were determined based on the cell count of CD154+ T cells after enrichment, normalized to the total number of CD4+ T cells applied on the column. For each stimulation, CD154+ background cells enriched from the non-stimulated control were subtracted.

Flow cytometry
Cells were stained in different combinations of fluorochrome-conjugated antibodies (see Key resources table). Viability 405/520 Fixable Dye (Miltenyi Biotec) was used to exclude dead cells. For intracellular staining cells were fixed and permeabilized with the Inside stain Kit (Miltenyi Biotec). Data were acquired on a LSR Fortessa ( BD Bioscience, San Jose, CA, USA). Screening of expanded T cell lines on 384-well plates was performed on a MACSQuantX Analyzer (Miltenyi Biotec). FlowJo (Treestar, Ashland, OR, USA) software was used for analysis.

Quantification of SARS-CoV-2 spike-specific antibodies
Serum antibodies were measured as described previously (Geisen et al., 2021a, 2021b). Briefly, SARS-CoV-2 S1 IgG antibodies were measured according to the manufacturer’s instructions ( QuantiVac, Euroimmun, Lübeck, Germany). Samples with antibody values above the highest standard (384 BAU/ml) were diluted (1:10 or 1:20) and re-analyzed. SARS-CoV-2 Nucleocapsid-protein IgG antibodies were measured using the AESKULISA SARS-CoV-2 NP IgG test.

Measurement of SARS-CoV-2-neutralizing antibodies
Dilutions of each serum were tested in duplicate in an in-house 96-well format Vero cell-based virus-neutralization test (cVNT) ( Neumann et al., 2021; Rose et al., 2022a, 2022b). In brief, 2.5 × 10^4 Vero cells (order no. 605372, CLS Cell Lines Service GmbH, Eppelheim, Germany) were seeded per well and incubated at 37°C under standard conditions. On the next day, sera were heat-inactivated (56°C for 30 min) and diluted (1:10 to 1:1280) in supplemented Dulbecco’s modified Eagle’s medium. Then, 25 µl of the serum dilution were mixed with 25 µl of virus suspension containing 50 plaque-forming units (pfu) of an own pre-VOC B.1 strain. This virus was isolated from the upper respiratory tract of a 2020 COVID-19 patient, as previously described ( Stromer et al., 2020b). The resulting 50 µl were incubated for 1 h at 37°C. Vero cells were washed with PBS, combined with 50 µl of the virus-serum dilutions and incubated for one hour on a shaker at room temperature. Thereafter, 50 µl of fresh cell culture medium supplemented with 20% fetal calf serum (v/v) was added per well and plates were incubated for four days under standard conditions. Then, cells were fixed with paraformaldehyde and stained with crystal violet. The cytopathic effect formation was compared with an untreated cell control (medium only, negative) and a viral control (50 pfu, positive) ( Neumann et al., 2021; Stromer et al., 2020a). The geometric mean of the duplicates was calculated as titers.

Surrogate neutralizing antibodies (Figure S1G) were measured using a surrogate virus neutralization test (cPASS, Medac, Wedel, Germany, or BMS2326, Invitrogen, Vienna, Austria). For the comparison of the tests, samples were measured in parallel on both ELISAs and the incubation time of the thermo test was adjusted accordingly. IgG values above 35,6 BAU/ml and neutralization above 30% were considered as positive, according to the manufacturer’s recommendation.

Expansion and re-stimulation of antigen-reactive T cells
For expansion of antigen-specific T cell lines, PBMCs were stimulated for 6 h, CD154+ cells were isolated by MACS and further purified by FACS sorting on a FACS Aria Fusion ( BD Bioscience, San Jose, CA, USA) based on dual expression of CD154 and CD69. Purified CD154+ T cells were expanded in presence of 1:100 autologous antigen-loaded irradiated feeder cells in TexMACS medium ( Miltenyi Biotec), supplemented with 5% (v/v) human AB-serum (GemCell), 200 U/ml IL-2 (Proleukin; Novartis, Nürnberg, Germany), and 100 IU/ml penicillin, 100 µg/ml streptomycin, 0.25 µg/ml amphotericin B (Antibiotic Antimycotic Solution, Sigma Aldrich) at a density of 2.5 × 10^6 cells/cm². During expansion for 2–3 weeks, medium was replenished and cells were split as needed.

For re-stimulation, fastDCs were generated from autologous CD14+ MACS isolated monocytes (CD14 MicroBeads; Miltenyi Biotec) by cultivation in X-Vivo™15 medium ( BioWhittaker/Lonza), supplemented with 1000 IU/ml GM-CSF and 400 IU/ml IL-4 (both Miltenyi Biotec). Before re-stimulation expanded T cells were rested in RPMI-1640 supplemented with 5% human AB-serum without IL-2 for 2 days. 0.5–1×10^6 expanded T cells were plated with fastDCs in a ratio of 1:1 of in 384-well flat bottom plates and re-stimulated for 6 h, with 1 µg/ml Brefeldin A ( Sigma Aldrich) added for the last 4 h.

Analysis of functional avidity
For determining the functional avidity, in vitro expanded SARS-CoV-2-specific T cells were re-challenged with decreasing antigen concentrations (0.5, 0.25, 0.01, 0.025, 0.0025, 0.001, 0.00025 µg/peptide/ml) and analyzed for re-expression of CD154 and cytokines. Antigen concentrations required for half-maximal response (EC50 values) were calculated from dose-response curves using GraphPad PRISM. These curves were plotted as a semi-logarithmic plot, where the amount of antigen is plotted (on the X axis) as the...
log of antigen concentration and the response is plotted (on the Y axis) using a linear scale. To compare the EC50 values of different donors, the bottom and top of the curve were defined as 0 and 100%, respectively.

**T cell receptor (TCR) sequencing**

For TCR sequencing, 5x10^7 PBMCs pre-vaccination and 1-2x10^7 PBMCs post-vaccination were stimulated with the spike peptide pool for 6h. CD154<sup>+</sup> cells were magnetically enriched and further purified by FACS sorting on a FACS Aria Fusion (BD Bioscience, San Jose, CA, USA) based on dual expression of CD154 and CD69. A maximum of 1x10^6 CD154<sup>+</sup> cells were analyzed. To increase the sensitivity of the used TCR profiling kit, based on 5<sup>0</sup>RACE and switch oligo approaches, Jurkat cells were spiked into samples with less than 1x10^4 CD154<sup>+</sup>CD69<sup>+</sup> cells. RNA was isolated using the RNeasy Micro Kit (Qiagen). TCR libraries were prepared using human TCR profiling kit (MiLaboratory), according to the manufacturer’s protocol. In detail, 21 PCR cycles were used for the 1<sup>st</sup> PCR and 15 cycles for the 2<sup>nd</sup> PCR amplification.

Libraries were produced for n = 37 individuals for a total of n = 111 samples, i.e. 3 time points (baseline, post 1<sup>st</sup> vaccination, post 2<sup>nd</sup> vaccination) for each individual. Libraries were pooled equimolarly and sequenced on an Illumina NovaSeq6000 machine, SP v1.5 flow cell and 300 cycles. Data of both TCR<sub>α</sub> and β chains were obtained.

**TCR repertoire analyses**

PCR and sequencing error correction were performed through identification and selection of unique molecular identifiers using the software MiGEC, version 1.2.6 (Shugay et al., 2014). Filtered sequences were aligned on a TCR gene reference, clonotypes were identified, grouped and CDR3 sequences were identified using the software MiXCR version 3.0.14 (Bolotin et al., 2015). Clonotype tables containing clonotype counts, frequencies, CDR3 nucleotide and amino acid sequences and V(D)J genes were obtained and used for further analysis.

The Jurkat TCRs were then excluded from further analysis. The presence and abundance of baseline clonotypes was evaluated at 1<sup>st</sup> and 2<sup>nd</sup> post-vaccination time points. Pre-existing clonotypes were defined as clonotypes present both at baseline and at post-vaccination. Newly arising clonotypes post 1<sup>st</sup> vaccination were defined as clonotypes present in both post-vaccination time points. Newly arising clonotypes post 2<sup>nd</sup> vaccination were defined as clonotypes detected only after 2<sup>nd</sup> vaccination. Overlap among baseline clonotypes and the most abundant 100 clonotypes after 2<sup>nd</sup> vaccination was analyzed.

Diversity profiles of the TCR<sub>α</sub> repertoire were evaluated for all time points using Rényi diversity profiles (Rényi, 1961). By varying the α parameter, different diversity indices are calculated: at α = 0, 1, 2 and infinite, the profiles provide the logarithm of richness, the Shannon diversity, the reciprocal Simpson diversity, and the Berger-Parker index, respectively. This means that the sample with the highest value at α = 0 has the highest richness, but that the lower value at α = infinite indicates higher proportion of the most abundant sequence. A sample with a profile that is overall higher than the profiles of other samples is more diverse, i.e. has more different clonotypes.

**TCR similarity analyses**

TCR similarity clusters were identified using the TCRdist algorithm of the CONGA software v0.1.1 (Schattgen et al., 2022) on the single TCR<sub>α</sub> and TCR<sub>β</sub> chains separately. The TCRdist threshold for cluster identification for beta chain was set to 20 and for alpha to 15. An igraph object was created with the igraph package (Csardi and Nepusz, 2006), and greedy clustering was applied. Network plots were created with Gephi (Bastian et al., 2009) and Biorender showing the biggest TCR clusters, defined as clusters including a minimum of 20 unique TCRs.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Statistical parameters including the exact value of n, the definition of center, dispersion and precision measure, and statistical significance are reported in the Figures and the Figure Legends. Statistical tests were performed with GraphPad PRISM software 9.3 (GraphPad Software, La Jolla, CA, USA). Statistical tests were selected based on appropriate assumptions with respect to data distribution and variance characteristics, p values <0.05 were considered statistically significant.

**Software**

Flow cytometry data were analyzed using FlowJo (Treestar, Ashland, OR, USA) software. Graphics and statistics were created with GraphPad PRISM software version 9.3.1. (GraphPad Software, La Jolla, CA, USA). Heatmaps were generated using Genesis software, version 1.7.7 (Sturn et al., 2002). For TCR sequence analyses the software MiGEC version 1.2.6 (Shugay et al., 2014), and MiXCR version 3.0.14 (Bolotin et al., 2015) were used to obtain clonotype tables. For downstream analyses the R software v4.1.2 was used. To calculate the diversity measures the R packages vegan (Oksanen et al., 2019) and ineq (Zeileis, 2014) were used. Ggplot2 (Wickham, 2016) was used for visualization.