Rapid synchronous type 1 IFN and virus-specific T cell responses characterize first wave non-severe SARS-CoV-2 infections

Highlights

- Interferon response to non-severe SARS-CoV-2 can precede PCR detection of infection
- Virus-specific T cell responses are equally rapid but independent of interferon
- Virus-specific antibodies lagged 1–2 weeks behind T cells in first wave infections

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In brief

Non-severe SARS-CoV-2 infection in unvaccinated virus-naive individuals may reveal correlates of protection against future variants. In this context, Chandran et al. show rapid type 1 interferon and virus-specific T cell responses 1–2 weeks before virus-specific antibodies, supporting a role for T cell-mediated heterologous protection in next-generation vaccines.
Rapid synchronous type 1 IFN and virus-specific T cell responses characterize first wave non-severe SARS-CoV-2 infections

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SUMMARY

Effective control of SARS-CoV-2 infection on primary exposure may reveal correlates of protective immunity to future variants, but we lack insights into immune responses before or at the time virus is first detected. We use blood transcriptomics, multiparameter flow cytometry, and T cell receptor (TCR) sequencing spanning the time of incident non-severe infection in unvaccinated virus-naive individuals to identify rapid type 1 interferon (IFN) responses common to other acute respiratory viruses and cell proliferation responses that discriminate SARS-CoV-2 from other viruses. These peak by the time the virus is first detected and sometimes precede virus detection. Cell proliferation is most evident in CD8 T cells and associated with specific expansion of SARS-CoV-2-reactive TCRs, in contrast to virus-specific antibodies, which lag by 1–2 weeks. Our data support a protective role for early type 1 IFN and CD8 T cell responses, with implications for development of universal T cell vaccines.

INTRODUCTION

The host response in non-severe SARS-CoV-2 infection during the first epidemic wave, before vaccination, incorporates the mechanisms of effective host-defense in naive populations. These mechanisms remain a research priority because they may reveal determinants of protection against future variants of SARS-CoV-2 able to escape current vaccines, or future novel coronaviruses. To date, our knowledge has been limited to immune responses after the detection of the virus or onset of symptoms and to cross-sectional studies in which the time of infection was undefined. These data have provided strong evidence that germline encoded innate immunity mediated by type 1 interferon (IFN) responses contribute to protection against critical illness,1–3 although the extent to which they abrogate early symptomatic disease is not known. Perhaps more importantly, we still have very limited insights into the potential role of adaptive immune responses in early protection against disease in the first epidemic wave.4 These are of particular interest because they may inform the design of the next generation of vaccines with potential to provide cross-reactive immunity to future variants. We recently reported T cell responses in a subset of individuals associated with evidence for very early termination of infection before detectable viral replication or even seroconversion.5 The extent to which adaptive immune responses are a general feature of early responses in non-severe disease is not known. We sought to describe the temporal kinetics and relationships between the earliest immune responses to infection with an unbiased systems-level approach using genome-wide transcriptional profiling of weekly blood samples before, during, and after incident SARS-CoV-2 infections during the first epidemic wave in London and compared our findings with responses to other acute respiratory viruses using publicly available data from human challenge experiments.
RESULTS

Type 1 IFN and cell proliferation responses from 1 week before to 3 weeks after detection of asymptomatic and mild SARS-CoV-2 infection

We undertook a nested case-control study derived from a cohort of 400 healthcare workers at one London hospital recruited from March 23, 2020 to undergo weekly nasopharyngeal swab PCR tests and blood sampling when fit to attend work, as described previously. In this cohort, we detected 45 incident infections by PCR. Among these cases, we obtained 114 blood transcriptional profiles from 41 individuals spanning 3 weeks before to 3 weeks after the first PCR-positive result, including 12 individuals for whom samples were available before the first positive PCR. We also profiled convalescent samples from 16/41 individuals 5–6 months later. We compared these data to blood transcriptional profiles obtained from baseline samples in 55 consecutive uninfected controls who remained PCR- and seronegative for SARS-CoV-2 during follow-up (Figure S1; Table S1). None of the individuals who became infected required hospitalization. Among 38 individuals for whom blood transcriptomic data were available at the time of first positive PCR, 29 had no contemporary symptoms attributable to SARS-CoV-2 infection. Genome-wide transcriptional profiles from those who experienced an infection showed greatest perturbation compared with uninfected controls, both at the time of the first positive PCR test and independent of symptoms (Figure 1A; Figure S2D). Their profiles were significantly different from uninfected controls from the week before the first positive PCR to 3 weeks afterward. Six-month convalescent samples from a subset of these individuals were not significantly different from uninfected controls, indicating that the blood transcriptome had fully reverted to the baseline.

To investigate the host response to infection, we identified differentially expressed transcripts by comparison of profiles from the time of first positive viral PCR to those of uninfected controls (Figure 1B). These were subjected to upstream regulator enrichment analysis to identify molecular pathways predicted to be activated at the level of cytokines, transmembrane receptors, kinases, and transcription factors that may be responsible for differential gene expression (Data S1). We filtered groups of target genes associated with each upstream regulator to include only those that had significantly greater co-correlated expression than would be expected at random in our blood transcriptomes to increase our confidence that these represent co-regulated genes in each molecular pathway (Figure S2E). Among those that were retained, the associated upstream regulators formed two clusters resulting from overlapping associations with target genes (Figure 1C), reflecting two predominant biological pathways. These were type 1 IFN responses and cell-cycle activity, as surrogates for innate immune activation and cellular proliferation, respectively (Figure S3A). We collated the differentially expressed genes linked to the most statistically enriched upstream regulator in each of the two clusters as a transcriptional module, resulting in a signal transducer and activator of transcription (STAT)1-regulated module to represent type 1 IFN responses and a Cyclin D1 (CCND1)-regulated module to...
represent the cell proliferation response. The validity of the functional annotation for each of these modules was confirmed by investigating their correlation and covariance with independently derived transcriptional signatures for type 1 and type 2 IFN responses and for cell proliferation. The STAT1 module correlated with both IFN response modules but showed much greater covariance with the type 1 IFN signatures (Figure S3B), consistent with our bioinformatic analysis of the functional pathway represented by this cluster of differentially expressed genes. Similarly, we found that the CCND1-regulated gene expression module showed good correlation and covariance with an independently derived cell proliferation module (Figure S3C). Type 1 IFN and cell proliferation responses both peaked with co-incident infection (Figures 2A and 2B), but significant increases in these responses were also evident in the week before the first positive PCR result. Type 1 IFN responses remained significantly elevated for 1 week after the first positive PCR, whereas the cell proliferation response remained elevated for 2 weeks after the first positive PCR. The peak of each of these responses over this time course discriminated infected individuals from non-infected controls with area under the receiver operating characteristic curve (AUROC) of 0.87 (95% CI, 0.78–0.94) and 0.92 (95% CI, 0.87–0.98) for the STAT1- and CCND1-regulated modules, respectively (Figures S4A and S4B), giving a measure of the consistency of both these responses in infected individuals. Despite this and the overlap in the temporal profiles of these two responses, the enrichment of STAT1- and CCND1-regulated modules representing each response at the individual participant level did not correlate, suggesting that they may be independently regulated or subject to idiosyncratic capacity for each of these responses at the level of individual participants (Figure 2C). The same observation was evident for differentially expressed genes combined as modules associated with each of the upstream regulators that reflected type 1 IFN or cell proliferation modules (Figure 2D).

Cell proliferation responses distinguish SARS-CoV-2 infection from other acute respiratory viruses and are predominantly attributable to T cell responses

Next, we compared the type 1 IFN and cell proliferation responses to incident SARS-CoV-2 infection with those of other acute respiratory viruses by comparing the peak expression of the STAT1- and CCND1-regulated modules in our cohort to that of publicly available samples of RSV, HRV, H3N2, and H1N1 infections (Figures 2E and 2F). Type 1 IFN responses were significantly higher in SARS-CoV-2 infection compared with all other groups. In contrast, the cell proliferation response was not significantly different between SARS-CoV-2 infection and other acute respiratory viruses, consistent with the predominant contribution of T cell responses to cell proliferation.
available longitudinal blood transcriptomic data derived from human challenge experiments with respiratory syncytial virus, human rhinovirus, and influenza virus (Figure S4D). Comparable enrichment of the type 1 IFN response was evident in each of these infections (Figure 2E), but the cell proliferation response was significantly greater to SARS-CoV-2 than the peak response to any of the other acute respiratory virus infections (Figure 2F). Of note, the peak cell proliferation response to SARS-CoV-2 infection did not correlate with the persistence of this response represented by CCND1 module expression at 2 weeks after the first positive PCR (Figure S4C). We tested the hypothesis that the cellular proliferation response may arise from rapid B cell or T cell expansion in response to infection by evaluating the correlation between the CCND1 module and expression of validated cell-type-specific signatures (Figure 3A). CCND1 module expression correlated with the transcriptional signature for T cells, but not B cells. The relationship between the cell proliferation response and T cell subsets was stronger for the CD8 T cell signature than for the CD4 T cell signature. To corroborate these findings, we undertook multiparameter flow cytometry of peripheral blood mononuclear cells (PBMCs) obtained in a subset of participants with contemporaneous PCR-positive infection and compared these to PBMCs from uninfected controls (Figure S5). Representation of the pooled multiparameter flow cytometry data by tSNE incorporating all T cells revealed new populations of CD4- and CD8-positive cells only in samples from infected participants, which also exhibited the highest levels of Ki67 staining as a marker of cell proliferation (Figure 3B), accompanied by HLA-DR expression as a marker of cell activation in exemplar cases (Figure 3C). In this subset of samples, only Ki67 staining of CD8 T cells was statistically enriched in infected individuals compared with controls (Figure 3D). Both CD4 and CD8 T cells showed statistically significant enrichment of Ki67 staining among selected memory T cell populations, but not naive T cells (Figure S5B). Of all other lymphocyte subsets, only NKT-like (CD3+CD56+) cells showed a significant increase in Ki67-positive staining (Figure S5C) but comprised on average 3% of circulating lymphocytes compared with T cells that comprised approximately 40% (Figure S5D).

Rapid clonal T cell expansion in response to SARS-CoV-2 infection associated with significant enrichment of SARS-CoV-2-reactive T cell receptors

To further evaluate the rapid T cell response to SARS-CoV-2 infection, we undertook sequencing of T cell receptor (TCR) \( \alpha \) and \( \beta \) chains in longitudinal samples to reflect dynamic changes in the T cell clonal repertoire. The abundance of \( \alpha \) and \( \beta \) chain

Figure 3. Cell proliferation response to SARS-CoV-2 infection in blood transcriptomic data is attributable to T cell proliferation

(A) Correlation of CCND1 module (representative of cell proliferation response) in all time points (−3 to +3 weeks) from individuals with SARS-CoV-2 infection with each of blood transcriptomic modules representative of B cells, pan-T cells, CD4 T cells, and CD8 T cells (regression lines shown in red, \( p \) values for Spearman rank correlations).

(B) tSNE plots of T cells from non-infected controls or individuals with co-incident PCR-positive SARS-CoV-2 infection. Contour plots are shown in the two left-hand panels, followed by dot plots colored by CD4/CD8 staining or relative KI67 staining as a proliferation marker. Red circles highlight a population of KI67 high CD4 and CD8 T cells exclusive to the PCR+ group (tSNE derived from flow cytometry data of CD4 and CD8 T cells, seven NICs, and nine PCR-positive).

(C) Representative flow cytometry data for HLA-DR and KI67 staining in either CD4 T cells or CD8 T cells from one non-infected control and one SARS-CoV-2 infected individual at the time of the first positive PCR infection. Numbers indicate percent positive for each marker including double-positives.

(D) Summary HLA-DR and KI67 staining data and median from seven uninfected controls and nine individuals with co-incident infection in either CD4 T cells or CD8 T cells. \( p \) value shown for Mann-Whitney test.
sequences in each sample were tightly co-correlated (Figure S6A). An expanded clone will increase or decrease in frequency depending on the sampling time point before and after the peak response. Therefore, we identified expanded TCR sequences as being statistically enriched at one time point compared with at least one other time point and summed the total number of expanded sequences for these TCRs at each time point per million of total TCR sequences (Figures S6B and S6C). These were compared with expanded sequences identified in the same way among a subset of six uninfected controls in whom we undertook TCR sequencing in samples from 5 successive weeks. By comparison to the pooled data from controls, a significant increase in expanded TCRs was evident in infected individuals by the time of the first positive PCR test up to a maximum abundance of >6% of total TCR sequences and persisted for at least 3 weeks for both \( \alpha \) and \( \beta \) chains (Figures 4A and S6). The abundance of expanded TCR sequences correlated significantly with the CCND1, but not the STAT1, regulated module, consistent with the hypothesis that the proliferation response reflected expansion of T cell clones (Figures 4B and 4C).

T cell clonal expansion was not explained by changes in MAIT cell- or NKT cell-associated TCR sequences (Figures S6E and S6F). A total of 102 expanded SARS-CoV-2-reactive \( \alpha \) or \( \beta \) chain TCR sequences from the contemporary VDJdb catalog\(^2\) were evident in 28 of the 41 individuals with SARS-CoV-2 infection, giving the odds ratio (OR±95% CI, Fisher’s exact test) for enrichment of antigen-specific TCR sequences in each case.

(F) Number of SARS-CoV-2-specific TCRs in VDJdb for which the reported HLA restriction (HLA A1 or HLA A2) matches the HLA haplotype of the individual in which the expansion was observed. The number is compared with the expected number of HLA matches if HLA allocation was random, and these numbers are used to derive the OR (±95% CI, Fisher’s exact test).

(G) Number of \( \text{ex vivo} \) SARS-CoV-2 peptide-reactive TCR sequences (\( \alpha \) and \( \beta \) genes) among expanded and unexpanded TCR sequences from individuals with SARS-CoV-2 infection, giving the OR (±95% CI, Fisher’s exact test) for enrichment of virus-specific TCR sequences.
(EBV)-reactive TCRs, which would represent non-specific bystander memory T cell proliferation. We also found a significant association between the reported HLA restriction of the TCRs in the database and the HLA type of the individuals in whom the matched expanded TCR was observed (Figure 4F).

The analysis above is dependent on accurate annotation of public SARS-CoV-2-reactive TCR sequences and is skewed toward anti-spike CD8 responses, reflecting the preponderance of MHC multimer sorted T cells in VDJdb. Therefore, it is likely to underestimate the proportion of SARS-CoV-2 expanded TCR sequences associated with incident infection in our participants because it will not include either private or CD4+ TCR specificities. To address this limitation and further confirm the significant enrichment of SARS-CoV-2-reactive T cell clones within our TCR analysis, we compared expanded $\alpha$ and $\beta$ TCR sequences in vivo with antigen-specific TCR sequences that expanded following ex vivo peptide stimulation in three individuals with convalescent PMBC samples available at 16 weeks after infection. Antigen-specific responses by CD4 and CD8 T cells to both structural and non-structural viral peptides in these samples were confirmed by flow cytometry (Figure S7B) and associated with stimulus-specific expansion of TCR sequences (Figure S7C). The ex vivo SARS-CoV-2 peptide-reactive TCRs were significantly enriched among in-vivo-expanded TCRs within the same individuals (Figure 4G).

Likewise, the in-vivo-expanded TCR sequences were significantly enriched among ex vivo SARS-CoV-2 peptide-reactive TCRs (Figure S7D). As predicted above, the overlap between in vivo and ex vivo SARS-CoV-2-associated TCR responses showed a higher OR than the overlap of in-vivo-expanded TCRs with the SARS-CoV-2 TCR catalog in VDJdb.

Circulating virus-specific antibodies lag 2 weeks behind transient increase in immunoglobulin gene expression in response to SARS-CoV-2 infection

The finding that the CCND1-regulated module did not correlate with our B cell signature does not exclude a B cell response. Emergence of antibodies to SARS-CoV-2 has been reported as early as 5 days after symptom onset.13 Flow cytometry of a subset of PBMCs available from those with contemporaneous PCR-positive infection showed no significant increase in frequency of selected B cell subsets, or among proliferating (Ki67-positive) B cells, compared with uninfected controls (Figures 5A and 5B). Interestingly, in blood transcriptional profiles we found increased expression of immunoglobulin (Ig) constant heavy- and light-chain transcripts, which peaked at the time of first PCR virus detection but was evident from 1 week before to 2 weeks after first PCR detection (Figures 5C and 5D). The increase in Ig gene expression in blood was less sustained than TCR expansion and returned to baseline by 3 weeks after the first positive PCR. In contrast, circulating antibodies to SARS-CoV-2 S1 spike protein that correlate with virus neutralization were not detectable until 1 week after the incident infection (Figure 5E) and continued to increase in this cohort for 8 weeks.8,9

**Discussion**

To the best of our knowledge, we report the earliest in vivo immune responses to natural SARS-CoV-2 infection available to date, enabled by serial sampling of individuals at risk of infection during the peak of the first epidemic wave in London. The general paradigm for early antiviral host defense is dominated by induction of type 1 IFNs. Attenuated responses as a result of autoantibodies to type 1 IFNs and genetic polymorphisms associated with reduced expression of a type 1 IFN receptor subunit or with reduced expression of the IFN-inducible oligoadenylate synthetase gene cluster have all been associated with severe disease.2–4 These provide strong evidence that type 1 IFN responses contribute to effective protection against severe SARS-CoV-2 infection. We show that type 1 IFN responses can precede PCR detection of the virus and therefore may exert their protective effects in the earliest phases of infection, independent of symptoms. We propose that early detection of IFN-inducible genes in the blood transcriptome may arise from localized immune responses as a result of leukocyte trafficking through lymphoid tissues or the site of infection and may provide greater sensitivity than detection of circulating IFNs. As we have previously reported, an additional translational application of this finding is the detection of IFN-inducible transcripts in blood as a diagnostic biomarker of early viral infection that may precede PCR detection of the virus and symptoms.14

Alongside type 1 IFN responses, we detected an early cell proliferation response in the blood transcriptome, which we primarily attribute to CD8, and to a lesser extent CD4, T cell proliferation by correlation with cell-type-specific transcriptional modules, corroborated by flow cytometry to show a significant increase in Ki67-positive CD8 T cells and TCR sequencing to show expansion of T cell clones. While type 1 IFN responses were evident in a range of other acute respiratory virus infections modeled in human challenge experiments,1 the early T cell response to SARS-CoV-2 in our study was significantly greater than in other viral infections. By comparison with current databases of SARS-CoV-2-specific and HLA A1/A2 matched TCRs in VDJdb and of TCRs that expand specifically in response to structural and non-structural virus peptide pools, we were able to show that in-vivo-expanded T cell clones were highly enriched for SARS-CoV-2-reactive cells and that these were already evident by the time of first positive virus PCR. In individuals with COVID-19, T cell reactivity has been reported as early as 5–10 days after the onset of symptoms.15 Importantly, in one report, T cell proliferative responses to SARS-CoV-2 were evident in 92% of family contacts of COVID-19 cases independently of serostatus,16 and some people may have pre-existing cross-reactive T cells arising from previous seasonal coronavirus exposure.16–21 These may be expected to contribute to early viral clearance, as suggested in a subset of individuals with evidence of abortive SARS-CoV-2 infection2 and analogous to findings in influenza.22–24 We note that quantification of blood transcriptional perturbation revealed some outliers in the uninfected control group (Figure 1A). It is interesting to speculate whether these may also represent immune responses in an abortive infection.

If cross-reactivity were the primary driver of rapid T cell responses to SARS-CoV-2 infection, the fact that the early proliferative response discriminated infected and uninfected individuals with an AUROC of 0.92 would require pre-existing T cell priming to be a near ubiquitous feature of asymptomatic or non-severe
infection. Consistent with this hypothesis, among the largest studies of pre-pandemic blood samples, heterologous T cell reactivity to SARS-CoV-2 peptides with proven similarity to those of pre-existing seasonal coronaviruses has been reported in 81%.21 In this context, we hypothesize that the variation in T cell proliferative response and the lack of its correlation with type 1 IFN responses may be explained by differential levels of T cell priming in individual participants. We also identified a similarly rapid B cell response represented by transient enrichment of Ig gene expression in blood, but we were unable to corroborate any significant changes in the frequency of B cell subsets in a small sample available for flow cytometry. We speculate that the B cell response identified in the blood transcriptome may represent non-specific or sub-optimal activation of Ig gene expression, and because protective anti-S1 antibodies only became detectable after a 2-week lag, we hypothesize that the B cell response was unlikely to contribute significantly to rapid viral clearance in asymptomatic and non-severe infection.

**Figure 5.** Enriched immunoglobulin gene expression and antibody response to incident SARS-CoV-2 infection

(A and B) (A) Frequency of B cell (N, naive; CM, classical memory; AM, activated memory; PC, plasma cells; PB, plasmablasts) subsets among total CD19-positive cells and (B) Ki67-positive CD19 cells in PBMCs from six NICs and seven individuals with co-incident infection, showing individual data points and the median (bars). (C) Heatmap of immunoglobulin constant heavy- and light-chain gene expression in blood per individual (columns) stratified by time to first positive SARS-CoV-2 infection, compared with NIC and convalescent (Conv) samples 5–6 months after incident infection, presented as standardized (Z) scores of transcripts per million (TPM) using mean and SD of NIC. (D and E) Blood TPM of (D) IGHG1 and (E) relative IgG anti-S1 antibody levels stratified by time to first positive SARS-CoV-2 infection, compared with NIC and Conv samples. Individual data points shown with violin plots depicting median, IQR, and frequency distributions (*FDR < 0.05 by Kruskal-Wallis test for each group compared with NIC).

**Limitations of the study**

Our study has some important limitations. The precise time of exposure to SARS-CoV-2 or transmission of infection was not possible to determine. This was offset by including longitudinal samples in 12 subjects obtained before detection of incident infection by PCR, providing enough statistical power to show that both type 1 IFN and cell proliferation responses were statistically enriched in the week before the first positive PCR result. We had limited access to PBMCs to assess frequency, phenotypic, and functional characteristics of SARS-CoV-2-reactive T cells, preventing further validation of a predominant virus-specific CD8 T cell response to early infection. The accumulating database of SARS-CoV-2-specific TCR sequences allowed us to relate clonal T cell expansion with HLA-class-1-restricted antigen specificity. This only accounted for a small fraction of expanded sequences and does not exclude proliferation of bystander T cells. However, the lack of significant enrichment of CMV- or EBV-specific TCRs among expanded clones, typically a major component of bystander responses to early viral infection, and the lack of enrichment for IFN-γ activity or other signatures of T cell activation in the blood transcriptome argue against generalized bystander T cell activation. We also confirmed that early *in vivo* TCR expansion in infected individuals was enriched for SARS-CoV-2 TCR expansion in response to *ex vivo* stimulation with viral peptides, but future studies will be required to undertake this analysis at single-cell level to establish the relative contribution of CD4 and CD8 T cell clones. Our focus on the blood compartment means that we do not have direct...
measurements of responses at the site of host-pathogen interactions. Analysis of bulk RNA samples for transcriptional profiling and TCR sequencing restricted our ability to evaluate transcriptional heterogeneity at the cellular level, further characterized expanded T cell clones, or undertake TCR analysis with transcriptional heterogeneity at the cellular level, further characterizing expanded T cell clones, or undertake TCR analysis with paired I/D chains. Most importantly, since less than 5% of infections lead to hospitalization,26 our study design precluded comparison of severe and non-severe outcomes that would require substantially greater sample size. Nonetheless, our data reflect immune responses in asymptomatic and non-severe infection, which incorporate correlates of effective host defense to natural infection in a SARS-CoV-2 naive population, providing further evidence for the importance of early type 1 IFN and T cell responses. Human challenge experiments that control for variation in time and dose of exposure will offer the best opportunities to acquire the granular detail of early immune responses. Larger-scale studies will be required to assess the frequency of SARS-CoV-2 T cell reactivity in naive populations and determine whether early type 1 IFN or T cell responses predict outcomes.

Although vaccine rollout is likely to be the primary immunological strategy to control the pandemic,27 understanding the determinants of effective natural immunity will remain a critical objective to enable risk stratification and novel vaccine design as the virus evolves. In particular, identification of the antigenic determinants of the earliest T cell responses in asymptomatic SARS-CoV-2 infection is a priority to inform development of potential universal coronavirus vaccines.

CONSORTIA

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SUPPLEMENTAL INFORMATION

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AUTHOR CONTRIBUTIONS

Study conception, design, and oversight, G.C., C.M., T.A.T., J.C.M., and M.N.; data acquisition, A.C., J.R., G.N., L.S., A.R.B., C.P., J.M.G., and M.W.; data analysis and interpretation, A.C., J.R., G.N., L.S., R.K.G., A.R.B., A.W., T.R., M.R.d.M., B.S.-M.D.D.E., J.A.G.-A., A.S., T.B., D.M.A., R.J.B., A.M., G.S.T., M.K.M., B.M.C., and M.N.
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### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| BV421-conjugated anti 4-1BB, clone 4B4-1 | Biolegend | Cat no: 309820; RRID:AB_2563830 |
| PE-conjugated anti Blimp-1, clone 6D3 | BD biosciences | Cat no: 564702; RRID:AB_2738901 |
| FITC-conjugated anti CCR7, clone 150503 | BD biosciences | Cat no: 561271; RRID:AB_10561679 |
| BV711-conjugated anti CD11c, clone 3.9 | Biolegend | Cat no: 301630; RRID:AB_2562192 |
| APC-conjugated anti CD138, clone DL-101 | Biolegend | Cat no: 352308; RRID:AB_10869646 |
| V500-conjugated anti CD14, clone M5E2 | BD biosciences | Cat no: 561391; RRID:AB_10611856 |
| Pe-conjugated anti CD161, clone 191B8 | Miltenyi | Cat no: 130-092-677; RRID:AB_571632 |
| BV786-conjugated anti CD19, clone SJ25C1 | BD biosciences | Cat no: 740968; RRID:AB_2740593 |
| AF700-conjugated anti CD20, clone 2H7 | BD biosciences | Cat no: 560631; RRID:AB_1727447 |
| BV421-conjugated anti CD21, clone B-Ly4 | BD biosciences | Cat no: 562966; RRID:AB_2737921 |
| PE/Cy7-conjugated anti CD24, clone MLS | Biolegend | Cat no: 311120; RRID:AB_2259843 |
| BUV395-conjugated anti CD27, clone L128 | BD biosciences | Cat no: 563815; RRID:AB_2744349 |
| BUV805-conjugated anti CD3, clone UCHT1 | BD biosciences | Cat no: 612895; RRID:AB_2870183 |
| BV510-conjugated anti CD3, clone OKT3 | Biolegend | Cat no: 317332; RRID:AB_2561943 |
| PE-Dazzle-conjugated anti CD38, clone HIT2 | Biolegend | Cat no: 303538; RRID:AB_2564105 |
| BUV395-conjugated anti CD4, clone SK3 | BD biosciences | Cat no: 565550; RRID:AB_2738273 |
| BV711-conjugated anti CD45RA, clone H100 | BD biosciences | Cat no: 56373; RRID:AB_2738392 |
| Pe-Dazzle594-conjugated anti CD56, clone QA17A16 | Biolegend | Cat no: 392410; RRID:AB_2728406 |
| BV605-conjugated anti CD69, clone FN50 | Biolegend | Cat no: 310938; RRID:AB_2562307 |
| APC-Cy7-conjugated anti CD71, clone CY1G4 | Biolegend | Cat no: 334110; RRID:AB_2563117 |
| PerCP/Cy5.5-conjugated anti CD71, clone Cy1G4 | Biolegend | Cat no: 334114; RRID:AB_25631175 |
| AlexaFluor700-conjugated anti CD8a, clone RPA-T8 | Biolegend | Cat no: 301028; RRID:AB_493745 |
| BV605-conjugated anti CXCR4, clone I2G5 | Biolegend | Cat no: 306521; RRID:AB_2562443 |
| V500-conjugated anti HLA-DR, clone G46-6 | BD biosciences | Cat no: 561224; RRID:AB_10563765 |
| APC-conjugated anti IFNg, clone 4S.B3 | Biolegend | Cat no: 502512; RRID:AB_315237 |
| BUV805-conjugated anti IgD, clone IA6-2 | BD biosciences | Cat no: 742039; RRID:AB_2871332 |
| APC/Cy7-conjugated anti IgM, clone MHM88 | Biolegend | Cat no: 314520; RRID:AB_10900422 |
| Pe-Cy7-conjugated anti Ki67, clone 20Raj1 | ThermoFisher | Cat no: 25-5699-42; RRID:AB_2573462 |
| AF488-conjugated anti Ki67, clone B56 | BD biosciences | Cat no: 561165; RRID:AB_10611866 |
| AlexFluor647-conjugated anti OX40, clone Ber-ACT35 | Biolegend | Cat no: 350018; RRID:AB_2571938 |
| BB700-conjugated anti PD-1, clone EH12.1 | BD biosciences | Cat no: 566460; RRID:AB_2744348 |
| BV785-conjugated anti TCR Va7.2, clone 3C10 | Biolegend | Cat no: 351722; RRID:AB_2566042 |
| Functional grade anti-CD28, clone CD28.2 | ThermoFisher | Cat no: 16-0289-81; RRID:AB_468926 |

(Continued on next page)
**Continued**

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Biological samples** | | |
| Blood RNA samples | This study | N/A |
| Peripheral blood mononuclear cell samples | This study | N/A |
| Bacillus Calmette Guerin (BCG) | NIBSC (UK) | Cat no: 07/274 |
| **Chemicals, peptides, and recombinant proteins** | | |
| Recombinant human IL-2 | Peprotech | Cat no: 200–02 |
| SARS-CoV-2 overlapping peptide pools covering structural spike, nucleoprotein, membrane protein, NSP7, NSP12, NSP13 | GL Biochem Shanghai | Custom |
| RLT buffer | Qiagen | Cat no: 79216 |
| Ficoll-Hypaque Plus | GE healthcare | Cat no: 17-1440-03 |
| Dimethyl sulfoxide solution | Sigma Aldrich | Cat no: 67-68-5 |
| Fetal bovine serum | Sigma Aldrich | Cat no: F7524 |
| Phosphate buffered saline | ThermoFisher | Cat no: 10010023 |
| Brilliant violet buffer | BD biosciences | Cat no: 563794 |
| **Critical commercial assays** | | |
| Tempus™ Blood RNA tubes | ThermoFisher | Cat no: 4342792 |
| Tempus Spin RNA Isolation Kit | ThermoFisher | Cat no: 4380204 |
| GlobinClear kit | ThermoFisher | Cat no: AM1980 |
| TURBO DNA-free kit | ThermoFisher | Cat no: AM2238 |
| Kappa Hyperprep kit | Roche | Cat no: 07962363001 |
| Nextseq 500/550 High Output 75 cycle kit | Illumina | Cat no: 20024906 |
| LIVE/DEAD™ Fixable Blue Dead Cell Stain Kit | ThermoFisher | Cat no: L34962 |
| Foxp3 / Transcription Factor staining buffer | eBioscience | Cat no: 00-5523-00 |
| RNEasy kit | Qiagen | Cat no: 74004 |
| **Deposited data** | | |
| Blood RNAseq data from the present study cohort | ArrayExpress | ArrayExpress: E-MTAB-10022 |
| Microarray transcriptional profiles from PPD stimulated PBMC | ArrayExpress | ArrayExpress: E-MTAB-11345 |
| TCRseq data | NCBI Short Read Archive | NCBI Short Read Archive: SUB9362448 |
| Supplemental Data S1 and Data S2 | Mendeley Data | https://data.mendeley.com/datasets/vm4dvpxxdy/2 |
| | | https://doi.org/10.17632/vm4dvpxxdy.2 |
| **Software and algorithms** | | |
| Kallisto | Bray et al., 2016 | https://github.com/pachterlab/kallisto |
| Tximport | Bioconductor | https://bioconductor.org/packages/release/bioc/html/tximport.html |
| BioMart | Bioconductor | https://bioconductor.org/packages/release/bioc/html/biomaRt.html |
| sva package | Bioconductor | https://bioconductor.org/packages/release/bioc/html/sva.html |
| Ingenuity Pathway Analysis | Qiagen | https://digitalinsights.qiagen.com/products-overview/discovery-insights-portfolio/analysis-and-visualization/qiagen-ipa/ |
| Gephi v0.9.2 | Jacomy et al., 2014 | https://gephi.org/ |
| XGR | Fang et al., 2016 | https://xgr.r-forge.r-project.org/ |

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RESOURCES AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Mahdad Noursadeghi (m.noursadeghi@ucl.ac.uk).

Materials availability
Requests for access to samples will be considered (subject to a material transfer agreement) by an access committee via an online application (https://covid-consortium.com/application-for-samples/), on the basis of availability and scientific merit within the scope of research ethics approvals, participant consent, and data governance. Responses to applications will be made within 4 weeks.

Data and code availability
- Open access to new transcriptional profiling data and essential anonymised metadata is available online through ArrayExpress: E-MTAB-10022 and ArrayExpress: E-MTAB-11345. TCR sequencing data are available at NCBI Short Read Archive: SUB9362448. Requests for access to individual de-identified participant level data will be considered (subject to a data transfer agreement) by an access committee via an online application (https://covid-consortium.com/application-for-samples/), on the basis of availability and scientific merit within the scope of research ethics approvals, participant consent, and data governance. Responses to applications will be made within 4 weeks.
- No custom code was generated for the present analysis.
- Any additional information required to reanalyze the data reported in this work paper is available from the Lead contact upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Study design
We undertook a case control study nested within our COVIDsortium health care worker cohort. Participant screening, study design, sample collection, and sample processing have been described in detail previously.6,7,9,10 Healthcare workers were recruited at St Bartholomew’s Hospital, London, UK in the week of lockdown in the United Kingdom (between 23rd and 31st March 2020). Participants underwent weekly evaluation using a questionnaire and biological sample collection (including serological assays) for up to 16 weeks when fit to attend work at each visit, with further follow up samples collected at 6 months. Participants with available blood RNA samples who had PCR-confirmed SARS-CoV-2 infection (Roche cobas diagnostic test platform) at any time point were included as ‘cases’. A subset of consecutively recruited participants without evidence of SARS-CoV-2 infection on nasopharyngeal swabs and who remained seronegative by both Euroimmun antiS1 spike protein and Roche anti-nucleocapsid protein throughout follow-up were included as uninfected controls. The baseline characteristics of the study participants are provided in Table S1.

Ethical approval
The study was approved by a UK Research Ethics Committee (South Central - Oxford A Research Ethics Committee, ref. 20/SC/ 0149). All participants provided written informed consent.

METHOD DETAILS

Blood RNA sequencing
Blood samples for RNA sequencing were collected in Tempus Blood RNA tubes. For ‘cases’, we included all available RNA samples, including convalescent samples at week 24 of follow-up for a subset of participants. For uninfected controls, we included baseline

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| FlowJo v10.7.1      | BD biosciences | N/A |
| Decombinator v4      | Peacock et al., 2021 | https://github.com/innate2adaptive/Decombinator |
| Other               | VDJdb database (accessed 1st November 2021) | Bagaev et al., 2020 | https://vdjdb.cdr3.net/ |
|                     | Blood transcriptional profiles from selected human respiratory virus challenge studies | Gene Expression Omnibus | https://www.ncbi.nlm.nih.gov/geo/ (accession no: GSE73072) |
For multiparametric flow cytometry cells were plated in 96-well round-bottomed plates (0.5-1x10^6 per sample) and washed with PBS.

Flow cytometry

For multiparametric flow cytometry cells were plated in 96-well round-bottomed plates (0.5-1x10^6 per sample) and washed once in PBS (PBS; ThermoFisher) then stained with Blue fixable live/dead dye (ThermoFisher) for 20 min at 4°C in PBS, followed by selected antibodies (see key resources table and Table S2). Cells were washed again in PBS and incubated with saturating concentrations of monoclonal antibodies against markers to be stained on the cell surface, diluted in 50% Brilliant violet buffer (BD biosciences) and 50% PBS for 30 min at 4°C unless stated. After surface Ab staining cells were resuspended in fix/perm buffer (eBiosciences, Foxp3/Transcription Factor staining buffer kit, fix perm concentrate diluted 1:3 in fix/perm diluent) for 45-60 min at 4°C. Cells were then washed in 1x perm buffer (10x perm buffer Foxp3/Transcription Factor staining buffer kit diluted to 1X in ddH2O) and saturating concentrations of intranuclear targets (Ki67) were stained in 1X perm buffer for 30-45 min 4°C. Cells were washed twice in PBS then analyzed by flow cytometry using the LSR II flow cytometer (BD biosciences).

Ex vivo T cell stimulation

5x10^5 PBMC were cultured with anti-CD28 (0.5 μg/mL) and rIL-2 (20 IU/mL) + pools of overlapping peptides (2 μg/mL per peptide) covering structural proteins (spike, nucleoprotein and membrane protein) and replication transcription complex (RTC) proteins (NSP7, NSP12, NSP13), as detailed previously, comprising pools of 15-mer peptides overlapping by 10 amino acids >80% purity (GL Biochem Shanghai). Intracellular IFNγ cytokine staining post-expansion was performed as detailed previously, with brefeldin A being added on day 7 and PBMC harvested and stained 16-18 h later. In parallel, for T cell receptor sequencing, PBMC were collected on day 8, washed twice in sterile PBS, frozen at -80°C in RLT buffer (Qiagen), before thawing and extracting RNA using the RNEasy kit (Qiagen) according to manufacturer’s instructions.

QUANTIFICATION AND STATISTICAL ANALYSIS

Blood RNA sequencing data analysis

Sample processing batch effects were evaluated by principal component analysis at genome wide level (Figure S2A) and among the interstretch of the 10% genes with least variable expression in each sample processing batch (Figure S2B). A batch effect evident in the least variant gene expression analysis was corrected using the ComBat function in the sva package in R, allocating samples with PC2 score <0 and >0 (in Figure S2B) to separate batches. Principal component analysis (PCA) of the least low variance gene expression after batch correction showed no further separation of samples by processing batch (Figure S2C). Molecular degree of perturbation (MDP) was calculated as previously described. Transcripts were included only if more than one sample had a TPM count above the limit of detection, and the SD (SD) of TPM among uninfected controls was >0.5. The TPM values for each individual dataset were then transformed to a Z score using the mean and SD for each transcript among uninfected controls used as a standard reference. The MDP expression of each sample/dataset was then represented as the sum of all Z scores>2 divided by the total number of transcripts. Differential gene expression between datasets from individuals with co-incident infection and non-infection controls was identified using a Mann-Whitney test with false discovery rate <0.05 and absolute fold difference >1.5 (or Log2 0.585). Analysis of upstream transcriptional regulation of the differentially expressed genes was performed using Ingenuity Pathway Analysis (Qiagen, Venlo, The Netherlands) and visualised as network diagram using the Force Atlas two algorithm in Gephi v0.9.2. We depicted all statistically over-represented molecules (false discovery rate <0.05), predicted to be upstream of >2 target genes, and annotated with one of the following molecular functions: cytokine, transmembrane receptor, kinase and transcriptional regulator, representing the canonical components of molecular pathways responsible for transcriptional reprogramming in immune responses. The biological pathways represented by the upstream regulators were identified by Reactome pathway enrichment analysis using XGR as previously described. For visualization, 20 pathway groups were identified by hierarchical clustering of Jaccard indices to quantify similarity between the gene compositions of each pathway. For each group, the pathway with the largest total number of genes was then selected to provide a representative annotation.
Transcriptional modules
To identify co-regulated gene networks used as transcriptional modules, we calculated the average correlation coefficient for pairwise correlations of the expression levels of each group of target genes associated with predicted upstream regulators in our transcriptomic dataset, and compared this to the distribution of average correlation coefficients obtained from random selection of equivalent sized groups of genes repeated 100 times. Groups of target genes with average correlation coefficients that exceeded the mean of the distribution of equivalent sized randomly selected groups by \( \geq 2 \) SD (Z score \( \geq 2 \)) with false discovery rate <0.05 were identified as transcriptional modules representing the functional activity of the associated upstream regulator (Figure S2E).

Independently derived Type 1 and Type 2 interferon inducible modules and cell-type specific transcriptional modules were described previously.\(^{36,38,39}\) To derive an independent cell proliferation module, PBMC were isolated from BCG-vaccinated individuals and stimulated \( in \ ames / 2x10^5 \) colony forming units of Bacillus Calmette–Guérin Russia (NIBSC) for 6 days to drive proliferation of antigen specific T cells. Stimulated and unstimulated PBMC were subjected to transcriptional profiling, differential gene expression and Reactome pathway enrichment analysis as previously described.\(^{38}\) Differentially enriched transcripts annotated to the “Cell Cycle” Reactome pathway (Table S3) were used to derive a transcriptional signature for T cell proliferation. The expression of each module was represented by the geometric mean log\(_2\) TPM value of its constituent genes.

Data from human challenge studies
Publicly available data from previously published human viral challenge studies were downloaded from Gene Expression Omnibus (accession GSE73072). We calculated module scores for the STAT1 and CCND1 modules as the mean expression across all constituent genes, using log\(_2\)-transformed microarray data. Only participants who developed evidence of infection following inoculation were included, as per the original study definitions.\(^{11}\) The peak enrichment of STAT1 and CCND1-regulated modules for each infected individual was calculated was represented by the highest log\(_2\) TPM ratio to the mean of uninfected controls, across the time course of each dataset (Figure S4).

Flow cytometry data analysis
Flow cytometry data were analyzed using FlowJo (version 10.7.1 for mac, Tree Star). Single stain controls were prepared with cells or anti-mouse IgG beads (BD biosciences). Fluorescence minus one (FMOs) were used for gating. Selected cell populations were quantified using established combinations of antibody stains (Table S2). For tSNE an equal number (2223 cells) of CD4 and CD8 T cells from each of the seven control and nine PCR + samples were concatenated and tSNE was calculated on single cells expression values for the following markers: CD4, CD8, HLA-DR, Ki67, CD45RA, CCR7 (Iterations 1000, perplexity 30, eta 4979; KNN algorithm, Exact. Gradient algorithm, Barnes-Hut).

T cell receptor sequencing and analysis
The alpha and beta genes of the TCR were sequenced from all time points for which RNA was available within the first 4 weeks of the study for all participants who were PCR + at any time point, and for six randomly selected individuals who remained PCR- and seronegative throughout the study. The pipeline introduces unique molecular identifiers attached to individual cDNA molecules which allows correction for sequencing error PCR bias, and provides a quantitative and reproducible method of library preparation. Full details for both the experimental TCRseq library preparation and the subsequent TCR annotation (V, J and CDR3 annotation) using Decombinator V4 have been described previously.\(^{40,41,42}\) Expanded TCRs were defined as any TCR which changed significantly between any two time points (Figure S6B-S6C). The boundaries (shown as blue dotted lines) were defined as the maximum TCR abundance which might be observed at time 2, given its abundance at time 1, assuming Poisson distribution of counts with \( p < 0.0001 \), to give a false discovery rate of \( <1 \) in 1000. TCR abundances are normalised for total number of TCRs sequenced in each sample, and expressed as counts/million. MAIT TCRs were defined as any TCR alpha containing TRAV1-2 paired with TRAJ12, TRA20 or TRAJ33. INKT TCRs were defined as TCRs containing TRAV10 paired with TRAJ18. The VDJdb database\(^{43}\) (https://vdjdb.cdr3.net/), accessed on 1\(^{st}\) November 2021, was searched for any TCR annotated for CMV, EBV or SARS-Cov-2. TCRs annotated for multiple antigens were excluded.

The number of in vivo expanded TCRs defined as described above which matched the antigen annotated TCRs for each set of antigen was then calculated. As a control, we calculated the average overlap observed using 10 same size random sets of TCRs from the same individuals, instead of the expanded TCRs. The VDJdb included the HLA restriction of each TCR determined experimentally. We measured the association between the VDJdb HLA of each matched TCR, with the known HLA haplotype of the individual in which the expansion was observed. We considered a match “correct” if the individual had at least one allele which matched that of the annotated matched TCR. The statistical association was calculated by comparing the number of “correct” and “incorrect” HLA matches with the number expected if HLA of all individuals was randomly shuffled.

The in vitro peptide stimulated T cells from three individuals were TCR sequenced after in vitro expansion, and compared to cultures from the same pool of T cells cultured without antigen but under identical culture conditions. T cell receptors were considered expanded if the frequency of the TCR was at least eight-fold higher in the antigen stimulated than the control cultures.