Adoptive transfer of ex vivo expanded SARS-CoV-2-specific cytotoxic lymphocytes: A viable strategy for COVID-19 immunosuppressed patients?

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Abstract
Cellular and humoral response to acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infections is on focus of research. We evaluate herein the feasibility of expanding virus-specific T cells (VST) against SARS-CoV-2 ex vivo through a standard protocol proven effective for other viruses. The experiment was performed in three different donors’ scenarios: (a) SARS-CoV-2 asymptomatic infection/negative serology, (b) SARS-CoV-2 symptomatic infection/positive serology, and (c) no history of SARS-CoV-2 infection/negative serology. We were able to obtain an expanded VST product from donors 1 and 2 (1.6x and 1.8x increase of baseline VST count, respectively) consisting in CD3+ cells (80.3% and 62.7%, respectively) with CD4+ dominance (60% in both donors). Higher numbers of VST were obtained from the donor 2 as compared to donor 1. T-cell clonality test showed oligoclonal reproducible peaks on a polyclonal background for both donors. In contrast, VST could be neither expanded nor primed in a donor without evidence of prior infection. This proof-of-concept study supports the feasibility of expanding ex vivo SARS-CoV-2-specific VST.
from blood of convalescent donors. The results raise the question of whether the selection of seropositive donors may be a strategy to obtain cell lines enriched in their SARS-CoV-2-specificity for future adoptive transfer to immunosuppressed patients.

**KEYWORDS**
adoptive immunotherapy, COVID-19, lymphocyte expansion, respiratory virus, SARS-CoV-2, third-party donors, virus-specific T cells

1 | INTRODUCTION

The coronavirus disease 2019 (COVID-19) pandemic is causing an enormous health and economic global impact worldwide. Solid tumor or oncohematological patients who develop COVID-19 are at higher risk of mortality, surely due to their hampered humoral and cellular immune response against SARS-CoV-2. Adaptive T cell immune responses are increasingly recognized as key factor controlling viral clearance, severity of COVID-19, and humoral immune responses. A deep understanding of T cell responses to SARS-CoV-2 is critical to improve our assessment of the probability of SARS-CoV-2 reinfection, COVID-19 prognosis, the estimation of general population immunity, as well as for guiding vaccine development.

Although immunoglobulin G (IgG) and M (IgM) seroconversion rate after COVID-19 is common, it can vary from 80% to 100% in symptomatic and 15% to 95% in asymptomatic patients after SARS-CoV-2 infection. Moreover, the humoral response is not always achieved, and the presence of IgG can be as short as few days or weeks after recovery in nearly 40% of patients. In contrast, strong and long-lasting CD4 and CD8 T cell responses have been observed in nearly all infected cases, highlighting their predominant role in SARS-CoV-2 control and clearance.

Currenty, there are several ways to assess T cell response to SARS-CoV-2 infection such as ex vivo T cell flow cytometry and ELISPOT-based assays. However, limitations of these tests include expertise in interpretation, lack of consensus in the standardization of the methodology, and low burden of circulating SARS-CoV-2-specific T cells (<0.1% of total lymphocyte subset in peripheral blood). Furthermore, T cell cross-reactivity observed with other common seasonal human coronavirus could be regarded as a limitation since it may overestimate the real immunity in the general population. This issue is further aggravated by the current lack of evidence that such cross-reactivity could confer functional protection against SARS-CoV-2.

Innovative approaches, such as ex vivo virus-specific T cells (VST) expansion used for cytomegalovirus (CMV) and Epstein Barr virus (EBV), may be of value for SARS-CoV-2 since it could offer a great number of VST against SARS-CoV-2 for clinical or laboratory research purposes.

We present herein a proof-of-concept experiment to investigate the feasibility of expanding SARS-CoV-2 VST ex vivo. We report the results of extending the applicability of optimized VST expansion protocol against CMV and EBV to the SARS-CoV-2 and discuss about the potential implications of our findings.

2 | MATERIALS AND METHODS

2.1 | Donor selection

For the study purpose, we selected two healthcare workers with PCR-confirmed SARS-CoV-2 infection on 11 and 26 March 2020, respectively, and a volunteer healthy donor. All three volunteer donors were previously registered in REDOCEL, a database with high-resolution typed blood donors Human Leukocyte Antigen (HLA) system consenting to be contacted whether cellular product donations were required for adoptive therapies. This donor bank is enriched with young individuals carrying common Spanish HLA Class I/II alleles, which facilitates HLA matching for future patients. To carry out our proof-of-concept experiments, donors were selected in order to include three different immunologic profiles: two donors with PCR-confirmed history of SARS-CoV-2 infection, one of which was asymptomatic with no seroconversion (donor 1), while the other was mildly symptomatic and had detectable circulating IgG and IgM antibodies (donor 2), and a third donor without previous history of infection or seroconversion (donor 3).

Peripheral blood mononuclear cells (PBMCs) were obtained from 50 mL blood donations on 12 May 2020 after written informed consent. Cell processing and ex vivo expansion experiments were conducted at the GMP facility in Hospital Universitario y Politécnico La Fe. Summary of the entire protocol applied is described in Figure 1. This project has been approved by the ethical committee and institutional review board (registration number 2020-123-1).

2.2 | HLA typing

HLA-A, -B, -C, -DRB1, and -DQB1 loci typing of the blood donors included in the experiments was performed in the Histocompatibility Laboratory of the Valencia Transfusion Center by next generation sequencing (NGS) using commercially available reagents (GenDX, Utrecht, The Netherlands) and a MiniSeq platform (Illumina).

2.3 | Diagnostic and serologic testing for SARS-CoV-2

SARS-CoV-2 diagnostic was performed by real-time reverse transcriptase-polymerase chain reaction (rRT-PCR) by detection
gene E and gene N SARS-CoV-2 Realtime PCR Kit (Vircell, Spain). Serologic testing was performed by qualitative determination by chemiluminescence immunoassay (CLIA) Maglumi 2019-nCoV IgG, Maglumi 2019-nCoV IgM (SNIBE Diagnostic, Shenzhen, China), and COVID-19 VIRCLIA IgG, COVID-19 VIRCLIA (IgM and IgA) (Vircell).

### 2.4 Clinical laboratory tests

The analytical panel designed to evaluate the hematological changes related to SARS-CoV-2 infection included absolute lymphocyte and platelet counts, lymphocyte subpopulations analysis, as well as the serum levels of C-reactive protein, lactate dehydrogenase, D-dimers, fibrinogen, ferritin, cardiac troponin, and IL-6. Serum C-reactive protein (CRP) and Ferritin were measured by an immunoturbidimetric assay (Roche Hitachi), IL-6, and Procalcitonin by “ECLIA” method and LDH activity with the IFCC reference procedure.

### 2.5 Neutralization assay protocol

In the donors with PCR confirmed COVID-19 history, the neutralization capacity of circulating antibodies against the spike protein of SARS-CoV-2 was assessed using a vesicular stomatitis virus pseudotyped with the SARS-CoV-2 Spike protein. Experiments were performed as previously described with the exception that all tests were done in triplicate using fourfold antibody dilutions ranging from 1:20 to 1:20,480. The dose resulting in 50% neutralization was calculated using a three parameter logistic regression with the drc package in R using the LL3 function.

### 2.6 Generation of DCs

Donor-derived dendritic cells (DCs) were generated from freshly PBMCs isolated by gradient centrifugation with Ficoll-Paque (Lymphoprep StemCell Technologies) from 50 mL of anticoagulated (acid-citrate-dextrose) venous blood. After separating the
PBMCs, monocytes were isolated by immunomagnetic bead positive selection using the ClinMACS CD14 Reagent (Miltenyi Biotec, Germany) according to the manufacturer’s instructions. At this point, 50% of the CD14 negative fraction was cryopreserved with 10% DMSO + 90% human AB serum in polystyrene cryovials and kept at -80°C until its further usage.

The CD14 + selected monocytes were matured for 10 days using a cocktail of cytokines into monocyte-derived DCs. Briefly, cells were seeded in petri dishes at 37°C and 5% CO2 in IMDM (Lonza Walkersville, USA) in a concentration of 1 x 10^6 cells/mL and differentiated for the first 5 days in the presence of 1000 IU/mL Granulocyte-Macrophage Colony-stimulating factor (GM-CSF) and interleukin (IL)-4, with the media changed and cytokines supplemented every other day. On day 5, cells were washed and reseeded at the same concentration in IMDM media supplemented with 1 µg/mL Prostaglandin E2 (Sigma Aldrich), 400 IU/mL tumor necrosis factor-alpha (TNFα), 1000 IU/mL IL-1B, and 1000 IU/mL IL-6 (all cytokines were manufactured by Miltenyi, unless otherwise stated). On day 7, IMDM was changed and cytokines supplemented at the same concentration as day 5. Finally, on day 10, DCs were harvested, counted, and gamma-irradiated with 30 Gy of 137Cs before being peptide-pulsed and used as antigen-presenting cells in the first culture stimulation.

2.7 | Generation of PHA blasts

The noncryopreserved CD14 negative fraction was used to generate PHA blasts required for repeated cycles of culture restimulation. Briefly, cells were culture for 7 days in T75 cm² flasks at concentration of 1 x 10^5 cells/mL in RPMI 1640 (Sigma) supplemented with 10% Fetal Bovine Serum (FBS, Thermo Fisher) and in the presence of IL-2. PHA blasts were added into the G-ReX at a ratio of 4:1 PBMC/PHA-blast. The previously transformed and irradiated autologous blasts were thawed and loaded for 2 hours in RPMI with 1 µg/mL of each PepTivator. After a washing step, incubated PHA-blasts were added into the G-ReX at a ratio of 4:1 PBMC/PHA-blast. From days 7 to 14 fresh media with 20 IU/mL IL-2 was added to the culture, and on day 14 a similar restimulation step with loaded PHA-blasts was performed. From days 14 to 21, the IL-2 concentration was increased to 50 IU/mL. On day 21, VST were harvested, aliquoted, and cryopreserved. Cellular product characterization was performed by culture sampling at days 14 and 21 of culture.

2.8 | Generation of SARS-CoV-2-specific T cell lines

After the DCs differentiation, the previously cryopreserved CD14 negative fraction was thawed and cells seeded in a gas-permeable rapid expansion cultureware (G-Rex) system (Wilson Wolf Manufacturing) at a density of 1 x 10^5/cm² in X-VIVO™ 15 Medium (Lonza), supplemented with 5% human AB serum (obtained from AB donors pooled plasma) and 2 mM L-glutamine (Lonza). The irradiated matured DCs were pulsed for 2 hours in RPMI 1640 with 1 µg/mL of proteins S, protein M, and protein N PepTivator SARS-CoV-2 (Miltenyi). These PepTivators are composed by a pool of lyophilized peptides, consisting mainly of 15-mer sequences with 11 amino acids overlap, covering the immunodominant sequence domains of the surface (or spike) glycoprotein (S), covering the complete sequence of the membrane glycoprotein (M) and the complete sequence of the nucleocapid phosphoprotein (N) of SARS-CoV-2 (GenBank MN908947.3, Protein QHD43416.1). DCs were then washed with RPMI and co-cultured in the G-ReX at 37°C, in 5% CO2 humidified atmosphere, at a ratio of 10:1 PBMC/DC for a period of 7 days. On day 7, cells were counted, fresh media supplemented with 20 IU/mL of IL-2 was added and culture underwent a first restimulation with peptide-pulsed PHA-blast. The previously transformed and irradiated autologous blasts were thawed and loaded for 2 hours in RPMI with 1 µg/mL of each PepTivator. After a washing step, incubated PHA-blasts were added into the G-ReX at a ratio of 4:1 PBMC/PHA-blast. From days 7 to 14 fresh media with 20 IU/mL IL-2 was added to the culture, and on day 14 a similar restimulation step with loaded PHA-blasts was performed. From days 14 to 21, the IL-2 concentration was increased to 50 IU/mL. On day 21, VST were harvested, aliquoted, and cryopreserved. Cellular product characterization was performed by culture sampling at days 14 and 21 of culture.

2.9 | T cell clonality testing

T cell clonality was assessed in ex vivo blood samples and in the final expanded cellular product using a commercial kit (Master Diagnostica) via multiplexed amplification of the TCR γ locus using standardized primer sets and qualitative interpretation of fragment size distributions by capillary electrophoresis according to the BIOMED-2 protocol19,20.

2.10 | Flow cytometry analysis

Flow cytometry analyses were performed on days 0, 14, and 21 of cell expansion. Briefly, cells were resuspended in RPMI in Falcon round-bottom 5 mL FACS tubes at a concentration of 1 x 10^6 cells/mL and stimulated with 1 µg/mL of the specific peptides. About 10 ng/mL of PMA (phorbol 12-myristate 13-acetate) and 1 µM ionomycin (Sigma Aldrich) were used for a positive control, and DMSO at the concentration contained in the PepTivator was used as a negative control. After 2 hours of stimulation, 10 µg Brefeldin A (Sigma-Aldrich) was added to the tubes and kept overnight at 37°C in a 5% CO2 humidified atmosphere. Cells were then harvested, washed, permeabilized using Cytotox/Cytoperm (Becton Dickinson, San Jose, CA, USA), and stained with specific combinations of the following fluorochrome-conjugated antibodies: CD3-FITC, CD4-PECy7, CD4-BB700, CD8-APCCH7, CD19-PeCy7, CD56-APC, CD14-V450, interferon gamma (IFNγ)-PE and TNFα-APC (BD Biosciences). For the analysis of degranulation and assessing cytotoxic potential, day 21 expanded lymphocytes were restimulated in vitro with 1 mg/mL of the corresponding peptide pool in the presence of antihuman CD107a-BB700 antibody for 1 hour at 37°C in a humidified 5% CO2 atmosphere. DMSO was added as a negative control for spontaneous CD107a

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expression. After 1 hour, 1.2 µL/mL Monensin (2mM stock solution) and 10 µg Brefeldin A was added. Cells were then harvested, washed, and stained for additional surface molecules. In all staining, dead cells were excluded using the Fixable Viability Stain 510 (Becton Dickinson) and doublets were excluded by FSC-A versus FSC-H gating. Data were acquired on a FACSCanto II cytometer (Becton Dickinson) and analyzed using the FlowJo software (Tree Star, version 10, USA). A response was considered to be positive if the peptide stimulation was at least twice the background of the DMSO control sample from the same donor and higher than 0.15%. A donor was considered to have a detectable T cell memory to SARS-CoV-2, if a positive response was detectable to any of the SARS-CoV-2 PepTivators.

### 2.11 | Sterility testing and cryopreservation

Sterility of the expanded cells was assessed at the end of the culture by direct Gram staining, by testing for aerobic/anaerobic bacteria contamination using BacT/ALERT FA/FN Plus detection media (Biomerieux Diagnostics, USA) or fungal/mycobacterial contamination using BACTEC MYCO/F LYTIC (BD) medium and mycoplasma contamination was also ruled out by qPCR.

Expanded cells were aliquoted in polystyrene cryovials and cryopreserved with 10% DMSO + 90% human AB serum using a passive freezing container (Mr Frosty, Thermo Fischer Scientific) before being transfer into a liquid nitrogen tank.

### 3 | RESULTS

#### 3.1 | Donor SARS-CoV-2 infection history

Clinical characteristics of each donor are detailed in Table 1. Donor 1 is a 45-year-old male with a history of SARS-CoV-2 infection after close contact with a COVID-19 patient. He did not develop any clinical symptoms at any time. SARS-CoV-2 PCR positivity was confirmed at several time points in nasopharyngeal swab specimens, with the

| Parameter | Donor 1 | Donor 2 | Donor 3 |
|-----------|---------|---------|---------|
| Age (years) | 45 | 41 | 30 |
| Sex | Male | Male | Male |
| Ethnicity | Caucasian | Caucasian | Caucasian |
| HLA class I/II | | | |
| HLA-A | 01:01:01 | 02:01:01 | 11:01 |
| HLA-B | 18:01:01 | 49:01:01 | 08:01:01:01 |
| HLA-C | 04:01:01 | 06:02:01:02 | 03:03:01:01 |
| HLA-DRB1 | 03:01:01 | 11:01:01 | 03:01:01 |
| HLA-DQB1 | 02:01:01G | 02:01:01 | 02:01:01 |
| SARS-CoV-2 infection | | | |
| Symptoms | No | Mild | No |
| rRT-PCR swab test | Positive | Positive | NA |
| Hospitalization requirement | No | No | NA |
| Time from infectious contact to negative rRT-PCR (days) | 42 | 23 | NA |
| Status at blood donation | | | |
| Time since negative rRT-PCR (days) | 47 | 26 | NA |
| Symptoms | No | No | No |
| rRT-PCR swab test | Negative | Negative | Negative |
| Serologic testing | Negative | IgG+ | Negative |
| Neutralizing Abs | No | Yes | NT |
| T cell memory | Yes | Yes | No |

Note: Description of the relevant characteristics of each donor, their previous history in relation with SARS-CoV-2 infection and status at blood donation.

Abbreviations: NA, not applicable; NT, not tested.
first negative rRT-PCR occurring 42 days after the infectious contact. Serological testing at different time points (1, 2, and 4 months after the first positive PCR) was consistently negative.

Donor number 2 is a 41-year-old male with general mild symptoms, including myalgia and night-sweat but without pneumonia, with a disease duration of 23 days from the first positive PCR to negativity. This donor’s serological testing was consistently positive for anti-SARS-CoV-2 IgG at several time points (1, 2, and 4 months after the first positive PCR).

Donor 3 is a 30-year-old male with no prior history of close contact with infected individuals and no symptoms of SARS-CoV-2 infection since the outbreak of the COVID-19 pandemic in Spain, therefore representing individuals with no previous contact with this virus.

3.2 | Donor immune characterization at blood donation

At the time of blood donation (3 months after the first positive PCR for donors 1 and 2), all donors were asymptomatic, and the absence of RNA viral shedding was confirmed via a swab test. Results of the analytical panel assessing COVID-19 related laboratory changes at the time of donation were within the reference intervals. Additionally, the negativity of SARS-CoV-2 RNA in the blood samples was confirmed by rRT-PCR prior to the expansions. At the time of donation, neutralizing antibody testing showed no neutralization activity at a serum dilution of 1:20 for donor 1 and 50% neutralizing activity at a dilution of 1:48 (SE ± 5.01) for donor 2 (Figure 2A).

Assessment of T cell memory against SARS-CoV-2 was performed by flowcytometry in ex vivo fresh PBMCs samples measuring intracellular TNFα production in response to the different SARS-CoV-2 PepTivators (Figure 2B). CD8 + T cell responses were observed in both convalescent donors 1 and 2 for overlapping peptides of protein S (1.6% and 1.7%, respectively). However, both CD4 and CD8 responses for protein M and N could not reach sufficient levels to be deemed positive according to our defined threshold. In donor number 3, and in agreement with the absence of contact history with this new virus strain, no responses were detected to any of the SARS-CoV-2 peptides in neither the CD4 nor CD8 subpopulations.

3.3 | Results of SARS-CoV-2 VST expansion

T cell expansion and cell cultures were analyzed for cell number growth, cellular content, specificity according to cytokine production (day 14), and cytotoxic potential (day 21), as detailed in Table 2.

In terms of cellular growth, we observed an expected decrease in total cell number during the first 7 days of culture, corresponding to the death of non-SARS-CoV-2-specific (by-stander) cells. After the second stimulation with loaded autologous PHA-blasts, cell cultures from donors 1 and 2 expanded, reaching at day 14 a higher cell number than at the start of culture (1.6x and 1.8x increase, respectively). For donor 3, a marked decrease occurred from day 0 to day 7, with some cell number recovery until day 14 but never reaching the initial

![FIGURE 2](image-url) Donors characterization. (A) Neutralization capacity of circulating antibodies against the spike protein of SARS-CoV-2 assessed using a vesicular stomatitis virus pseudotyped with the SARS-CoV-2 Spike protein in both convalescent SARS-CoV-2 donors. (B) Example of the flow cytometry assessment of the anti-SARS-CoV-2 T cell memory in donor 2. TNFα was selected as the read-out, DMSO used as negative control and the CMV pp65 peptides and PMA/Ionomycin as positives controls.
TABLE 2 Expansion results

| Parameter | Donor 1 | Donor 2 | Donor 3 |
|-----------|---------|---------|---------|
| Day 7     | Fold expansion | 0.76x | 0.63x | 0.36x |
| Day 14    | Fold expansion | 1.6x | 1.8x | 0.96x |
| Specificity (%) TNFα/IFNγ | CD4+ Pep. M + N+S | 1.4 | 16.7 | 0 |
|           | CD8+ Pep. M + N+S | 0 | 27.0 | 0 |
| Day 21    | Fold expansion | 1.6x | 1.2x | - |
| Product identity (%) of total | CD3+ | 89.3 | 62.7 | - |
|           | CD4+ (% of CD3) | 60.1 | 58.6 | - |
|           | CD8+ (% of CD3) | 35.0 | 37.4 | - |
|           | CD56+ | 2.23 | 33.9 | - |
|           | CD19+ | 2.59 | 0.33 | - |
|           | CD14+ | 5.42 | 0.22 | - |
| Cytotoxic Potential (%) CD107a | CD4+ Pep. M | 3.3 | 5.0 | - |
|           | Pep. N | 0 | 0 | - |
|           | Pep. S | 12.8 | 20.4 | - |
|           | CD8+ Pep. M | 0 | 12.7 | - |
|           | Pep. N | 0 | 0 | - |
|           | Pep. S | 15.1 | 30.0 | - |
| Sterility | OK | OK | - |

Note: Summary of the cultures cell growth number, composition, and specificity. For donor number 3, culture was ended on day 14 because no SARS-CoV-2 specificity could be identified.

cell count and with a progressive decreased viability of the remaining cells.

On day 14, cultures were analyzed for their specificity to the SARS-CoV-2-peptides. Flow cytometry after stimulation with PepTivators M, N, and S, assessed TNFα and/or IFNγ-production in order to quantify the bulk responses to the viral peptides (Figure 3A). For both donors 1 and 2, CD4 + SARS-CoV-2-specific T cells could be identified (1.4% and 16.7%, respectively). For donor 2, 27% of the CD8 + T cells also responded to the viral antigens, representing the majority of the polyfunctional cells secreting TNFα and IFNγ simultaneously, suggesting their multifunctional effector potential. For donor number 3, besides the progressive decrease in cell numbers, T cells did not respond differently to the DMSO-negative control and the SARS-CoV-2 peptides, but only to the PMA/Ionomycin positive control, assuring the assay validity. For this reason, the cell culture was considered as not specific and was terminated at this time point.

The end culture products (day 21) for both donors 1 and 2 mainly consisted of CD3 + cells (80.3% and 62.7%, respectively) and with a CD4 + dominance. Interestingly, in the culture from our serology positive donor 2, an important NK cell population expanded (CD56 + CD3 −; 33.9% of cells), in contrast with the culture from donor 1, where only 2.2% of CD56 + CD3 − could be detected (Figure 3B). At this time point of the culture, the cytotoxic potential of the expanded VST was evaluated by measuring cell degranulation through CD107a staining. The responses were measured separately for each viral PepTivator in order to discern the magnitude of response and expansion elicited to the epitopes of the M, N and S proteins (Figure 3C). For both donors 1 and 2, strong degranulation responses were observed to overlapping peptide-pool spanning the protein S from SARS-CoV-2 and in both CD4 + (12.8% and 20.4%) and CD8 + (15.1% and 30%) sub-populations, respectively. Additional CD4 responses to PepTivator M were identified for both donors 1 and 2 (3.3% and 5%). In terms of CD8 + T cell population responding to the membrane glycoprotein for SARS-CoV-2, responses could be detected in donor 1, yet these did not reach greater than twice the signal to the background (our set criteria to be deemed positive, DMSO 3.23%, pep. M 5.77%). In contrast, a clear positive population of 12.7% CD107a + CD8 + T cells responded to this peptide after subtracting the background.

An additional T-cell clonality study comparing samples from day 0 and day 21 supported cell expansion numbers and flowcytometry data. After expansion, multiple reproducible peaks on a polyclonal background could be detected in both donor 1 and donor 2, suggesting that the expanded cellular product is enriched for oligoclonal antigen-specific T cells. The intensity of these peaks was more prominent for donor 2, which is also in accordance with the higher percentage of SARS-CoV-2 VSTs observed by flowcytometry (Figure 4).

4 | DISCUSSION

This proof-of-concept experiment shows that VST expansion against SARS-CoV-2 is feasible under GMP conditions and can achieve a nearly twofold increase in the number of this VST from patients with symptomatic and asymptomatic PCR-confirmed SARS-CoV-2 infection. We observed a predominance of CD4 + T cells in the final VST product. The SARS-CoV-2 specificity of the expanded product was

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**Note:** Summary of the cultures cell growth number, composition, and specificity. For donor number 3, culture was ended on day 14 because no SARS-CoV-2 specificity could be identified.
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analyzed by flow cytometry showing a robust TNFα and IFNγ polarization after exposure to protein S, M, and N. Clonality studies of the final product also confirmed the presence of clonal VST against SARS-CoV-2 in both cases.

The two SARS-CoV-2 infected donors presented herein exhibited different patterns of VST expansion. We observed a higher number of expanded SARS-CoV-2 VST in donor 2 (symptomatic) in comparison to donor 1 (asymptomatic), suggesting that symptomatic donors could be more suitable for expansion. In contrast, we did not observe SARS-CoV-2 VST expansion in donor 3 (no prior history or contact with COVID-19). The fact that donor 2 had neutralizing antibodies suggests a stronger T cell response and consequently a high number of expanded VST. Our observation is supported by prior findings, where the neutralizing antibody titer significantly correlated with the numbers of SARS-CoV-2 VST. However, we found a stronger cytokine response to protein S in CD8+ T cells in comparison to CD4+ T cells in both donors, as measured by the fraction of CD107a staining. This is very similar to influenza virus infection, where viral surface hemagglutinin elicits mostly CD4+ T cell responses, whereas the majority of CD8+ T cell responses are specific to viral internal proteins. But a clear understanding of epitope sensitivity and HLA-restrictions from the CD4+ and CD8+ T cells providing a protective immunity against SARS-CoV-2 is still lacking and merits further research.

Another intriguing observation was the high percentage (33%) of NK cells obtained from donor 2 in the expanded VST product. Adaptive NK cells could also play a protective role against this new emerging virus.

Different approaches exist to obtain VST. One consists of isolating these T-cells from whole blood or leukapheresis products with virus reactive cells using an automated device capturing IFNγ-secreting cells. This approach has been already assayed using...
overlapping peptides of SARS-CoV-2 in COVID-19 convalescent donors, but the reduced specific T cell number along with the presence of other nonspecific T cells in the final product represents important limitations. To assure their clinical efficacy, HLA testing of donors and recipients is required in order to match HLA restrictions of the selected cells or even confirming the specificity of the IFNγ + cells. As an alternative, the application to SARS-CoV-2 of expansion protocols initially designed for CMV, EBV, and adenovirus (16) is feasible and leads to a highly specific VST product against SARS-CoV-2.

Donor 2 donation yielded a higher amount of SARS-COV-2 VST and could be considered as the most suitable donor. A starting bulk PBMC product with no detectable (by cytometry) CD4 memory and only 1% of CD8 + anti-SARS-CoV-2 T cells could be enriched into a cellular product with a specificity of around 25% CD4 + and 42% CD8 + protein S and M specific VSTs. Although the PepTivator used herein did not include all SARS-CoV-2 proteins, we were able to expand enough specific SARS-CoV-2 VST. Although it is not clear which is the optimal dose of VST, it has been shown that an adoptive transfer of doses as low as $10^4$ VST/Kg can expand in vivo and control viral replication in a high proportion of cases with a low incidence of graft versus host disease. For clinical purposes, we can speculate that with a standard donation of 450 mL of whole blood could give rise to sufficient cells to treat five (70 Kg) COVID-19 patients, assuming a starting cautious infusion dose of $0.5 \times 10^6$ VST/Kg.

Finally, we would like to highlight that the ex vivo expanded SARS-CoV-2 VST product has several potential utilities for COVID-19 research. First, it offers a higher number of VST. Second, it could help to identify the SARS-CoV-2 relevant antigens for T cell recognition for vaccine development. Third, it could be used to elucidate the immune profile of a protective T cell immunity against SARS-CoV-2. And at last, it could be useful to study the HLA restrictions from T cell receptors recognizing SARS-CoV-2 infected cells in order to optimize adoptive cellular therapies.

The limitations of the current experiments comprise the low number of donors included, the low blood volume used (50 mL) for expansion, and the fact that the peptides used during the expansion do not cover all SARS-CoV-2 proteins. It remains to be determined if the use of overlapping peptides spanning all SARS-CoV-2 proteins can improve protocol performance, increasing the number of expanded VST.

### 5 CONCLUSIONS

In this proof-of-concept study, we conclude that it is possible to expand ex vivo SARS-CoV-2 VSTs from convalescent donors of
SARS-CoV-2 infection, using a standard protocol to obtain virus-specific VSTs. Contrariwise, this approach was not successful to prime de novo responses to SARS-CoV-2 peptides in an uninfected healthy donor.

CONFLICT OF INTEREST
The author(s) declare that they have no conflict of interests.

AUTHOR CONTRIBUTIONS
Conceptualization and design, MG, C.A.-G., and JLP; Methodology C.A.-G., JM, C.F.-G., and VL; Formal Analysis, DP, MPC, MDG., EMG-B., and CA; Investigation, AS, PS, I.G.-S., A.B.-R., AL, and I.L; Resources, AP, LL, JS, and CA; Review & Editing, JDLR, RG, and MAS; Funding Acquisition, GS; Project Administration, GS, JLP. All authors conducted the experiments in their field of expertise and participated in the revision of the manuscript. All authors read and approved the final manuscript.

INSTITUTIONAL REVIEW BOARD STATEMENT
The study was conducted according to the guidelines of the Declaration of Helsinki, and approved by the Institutional Review Board (or Ethics Committee) of Hospital Universitario y Politécnico La Fe (protocol code: 2020-123-1 and date of approval: March 27, 2020)

INFORMED CONSENT STATEMENT
Informed consent was obtained from all subjects involved in the study.

DATA AVAILABILITY STATEMENT
The data that support the findings of this study are available on request from the corresponding author. The data are not publicly available due to privacy or ethical restrictions.

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