Naïve B cells with low differentiation improve the immune reconstitution of HIV-infected patients

Jie Jia, Yu Zhao, Ji-Qun Yang, ..., Kun-Hua Wang, Jian-Hua Wang, Yi-Qun Kuang

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Highlights
Different immune responses show distinctive BCR repertoire
IGHV3 and IGHV4 are dominant genes expressed in naive and memory B cells of IRs
Naïve B cells mediate immune reconstitution in HIV-infected individuals on ART
Naïve B cells with low differentiation improve the immune reconstitution of HIV-infected patients

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SUMMARY
Incomplete immune reconstitution happens in some HIV-infected patients who have achieved persistent viral suppression under antiretroviral therapy (ART). We performed single-cell RNA sequencing for peripheral blood mononuclear cells to analyze B cell receptor (BCR) repertoire and B cell subtypes in health controls (non-HIV-infected, HCs), HIV-infected immunological responders (IRs), and immunological nonresponders (INRs). We found that the dominant usage of IGHV gene segments of naïve B cells and memory B cells were IGHV3 and IGHV4, and the diversity of BCR repertoire was decreased in INRs. Differentiation trajectory analysis showed that the low differentiation of naïve B cells was related to satisfactory immune status. The cell cycle of B cells with immune-specific genes of IgD+ B cells was degraded in INRs, which was mediated by the anaphase-promoting complex/cyclosome pathway in the phase of G2/M checkpoints. These findings provide significant insights to understand the function of B cell-mediated immune response in immune reconstitution after HIV infection.

INTRODUCTION
The introduction of antiretroviral treatment (ART) prevents HIV replication, thereby reducing the viral load to an undetectable level and gradually increasing CD4+ T cells to the normal level (>500 cells/µL) simultaneously. However, around 20%–40% HIV-infected patients failed to restore their CD4+ T cell counts (<350 cells/µL) after receiving ART; these patients are classified as immunological nonresponders (INRs). Compared with HIV-infected individuals whose CD4+ T cell counts have reached more than 350 cells/µL (immunological responders, IRs), INRs have severely compromised immunity, leaving them susceptible to AIDS and opportunistic infections.1,2 It was reported that deficient immune reconstitution may relate to reduced production of progenitor cells in bone marrow, reduced thymic output, and abnormal immune activation.3,4 However, these studies are not sufficient to reveal the mechanisms of failed immune reconstitution after HIV infection.

B cell-mediated immune response is sustained by HIV-specific memory B cells and plasma cells,6 suggesting an important role of B cells in immune reconstitution after HIV infection. HIV infection alters the memory B cell compartment and reduces naïve B cells by increasing cell apoptosis and exhaustion, which leads to a decrease in the quality of response to HIV.6 The B cell receptors (BCRs) can react to antigens, thus leading to B cell clone expansion and secretion of high-affinity antibodies.7 The variety of BCR is determined by the usage and diversity of V, D, and J genes in complementary determining region 3 (CDR3). BCR rearrangement corresponds to diversity of immunoglobulin (Ig) segments during B cell development.7 There is a linkage between the B cell clone and stereotypical immune response by analyzing BCR repertoires.8,9 Therefore, we intend to investigate the role of B cells in failed immune reconstitution after HIV infection.

Single-cell RNA sequencing (scRNA-seq) provides an effective way to explore cellular heterogeneity, differentiation state, and gene expression models of cell populations.10 In the present study, BCR repertoire and corresponding characteristics of B cell subtypes were detected. We showed that a high ratio of naïve B cells with low differentiation was related to satisfactory immune reconstitution. Furthermore, cell cycle of B cells was impaired and mediated by the anaphase-promoting complex/cyclosome (APC/C) pathway in G2/M checkpoints.
RESULTS

Three B cell subsets were identified by scRNA-Seq

We isolated peripheral blood mononuclear cells (PBMCs) of 13 participants to construct cDNA libraries and performed sc-RNA sequencing and sc-BCR sequencing using a 10x Genomics single-cell-based platform (Figure 1A). Three cluster (cluster 1, 2, and 3) cells were identified as B cells after data filtering (Figures 1B and S1). The B cell subsets corresponding to cluster 1 and cluster 2 were further annotated according to highly expressed CD19, CD79A, CD79B, and MS4A1 (Figure 1C). Furthermore, cluster 1 distinguishingly expressed high levels of TSL1A, IGHD, and IL4R (Figures 1D and 1E) and was defined as naïve B cells. Cluster 2 distinguishingly expressed IGHG1, IGHG2, AIM2, and TNFRSF13B and was defined as memory B cells (Figures 1D and 1E). Cluster 3 distinguishingly expressed IGHDA1, CD38, JCHAIN, and MZB1 and was defined as plasma cells (Figures 1C–1E). By analyzing the expression and distribution of selected marker genes, we identified and confirmed phenotypes of naïve B cells, memory B cells, and plasma cells.

Proportions of unique BCR clonotypes of naïve B cells and memory B cells decreased in INRs

To reveal the BCR repertoire dynamic during immune reconstitution after HIV infection, we assessed the BCR repertoire among the HC, IR, and INR groups. Our results indicated that the abundance of naïve B cells and memory B cells with IGH was higher in the IR and INR groups than that in the HC group. The number of IGHV clonotypes of plasma cells was significantly higher in the IRs than that in the HC group (p < 0.01) (Figure 2A). Furthermore, we counted up the proportion of unique (detected in one group) and nonunique (detected in two groups) BCR clonotypes of naïve B cells and memory B cells in the comparison groups of HC vs. INR, HC vs. IR, and IR vs. INR. The results showed that the proportion of unique BCR clonotypes of naïve B cells and memory B cells was decreased in the INR group (p < 0.05) (Figure 2B), suggesting a decreased diversity of BCR repertoire in INRs. We were unable to analyze the BCR clonotypes of plasma cells due to their low percentage.

A high proportion of IGHV3 and IGHV4 gene usage in naïve B cells and memory B cells

We counted the frequency of BCR IGHV and IGHJ gene usage in naïve B cells and memory B cells among the HC, IR, and INR groups (Figure 3). The results showed that the dominant usage of the IGHV gene segment in naïve B cells (Figure 3A) and memory B cells (Figure 3B) were IGHV3 and IGHV4, and the dominant usage of the IGHJ gene segment was IGHJ4 (Figures 3C and 3D). The frequency of IGHJ4 in naïve B cells was higher in the IR and INR groups than that in the HC group (Figure 3C). Furthermore, the high frequency of IGHV3-11 and low frequency of IGHV4-34 and IGHV4-39 in naïve B cells were observed in the INR group compared with the IR and HC groups (Figure 3E). The high frequency of IGHV3-21, IGHV4-31, IGHV4-34, and IGHV4-59 and low frequency of IGHV3-23 and IGHV3-48 in memory B cells were observed in the INR group compared with the IR and HC groups (Figure 3F).

We further analyzed the CDR3 amino acids length distribution in three B cell subsets among the HC, IR, and INR groups. The results showed that the length of CDR3 was mainly distributed at 11–13 amino acids for most BCR clonotypes in naïve B cells, the dominant length of CDR3 was 10 amino acids for most BCR clonotypes in memory B cells, and the dominant length of CDR3 was 11 amino acids for most BCR clonotypes in plasma cells in the IR and INR groups (Figure 3G). There were no statistical differences in CDR3 length among the IR, INR, and HC groups.

Lower differentiation of naïve B cells in IRs

We further investigated the differentiation trajectories of naïve B cells, memory B cells, and plasma cells. The results showed that naïve B cells, memory B cells, and plasma cells formed into a relative process with different cell differentiation statuses in pseudo-time (Figure 4A). The diagrams showed a higher density of cell populations in high differentiation statuses in the IR and INR groups than that in the HC group, and the cells with low differentiation statuses were decreased in the IR and INR groups compared with the HC group (Figure 4A). By exploring the distribution of naïve B cells, memory B cells, and plasma cells in pseudo-time trajectories, we showed a higher density of memory B cells and plasma cells were distributed on the high differentiation statuses in the IR and INR groups than that in the HC group, but there was no significant difference in memory B cells and plasma cells distribution between the INR and IR groups (Figure 4A). For naïve B cells, the decreased density was observed on the low differentiation statuses in the INR group compared to the IR group, while most naïve B cells were distributed on low differentiation statuses.
Figure 1. Three B cell subsets were identified by scRNA-seq

(A) A schematic workflow of study design. PBMCs were collected and separated into three groups: healthy control (non-HIV-infected, HC), immunological responders (IRs), and immunological nonresponders (INRs).

(B) The UMAP plots of the B cell subsets distribution. A single dot represented a single cell.

(C) The bubble diagram of selected marker genes expression to distinguish the B cell subtypes from other PBMCs: naïve B cells (cluster 1), memory B cells (cluster 2), and plasma cells (cluster 3). The red bubble represented the higher expression of the marker genes; the large bubble represented a higher proportion of the B cell subset expressed marker genes.

(D) The heatmap of the selected marker genes expression to distinguish naïve B cells, memory B cells, and plasma cells. The red bubble represented the higher expression of the marker genes; the large bubble represented a higher proportion of the B cell subset expressed marker genes.

(E) The UMAP plots denote the distribution of subtype-specific marker genes in naïve B cells, memory B cells, and plasma cells. The single dot represents a single cell, and the red dots represent cells with a higher expression level of marker genes.

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The proportion of B cells with unique BCR clonotypes was increased in naïve B cells, memory B cells, and plasma cells between the IR and INR groups (Figure 4C). The diagram showed that the expression tendency of TCL1A and IGHM were increased in naïve B cells and there was a significant distinction of cell distribution in pseudo-time between the HC and IR groups, while cell distribution was mixed between the INR and IR groups. In addition, there was a high ratio of memory B cells with expression of AIM2 and TNFRSF13B and a high ratio of plasma cells with expression of XBP1 and MZB1 in the IR group compared with the the HC group, which suggested different characteristics of naïve B cells, memory B cells, and plasma cells among the HC, IR, and INR groups. The distribution of naïve B cells with TCL1A and IGHM, memory B cells with AIM2 and TNFRSF13B expression, and plasma cells with XBP1 and MZB1 expression were highly overlapped in the IR and INR groups rather than in the HC group, suggesting the similarities of naïve B cells, memory B cells, and plasma cells between the IR and INR groups (Figure 4C).

Taken together, naïve B cells had a lower degree of cell differentiation and expressed a higher level of IGHM in the HC group than that in the IR and INR groups. Naïve B cells had a higher degree of cell differentiation in IRs and INRs and had a differentiation relationship with memory B cells and plasma cells, suggesting a positive correlation between satisfactory immune status and the high abundance of naïve B cells with low differentiation.

**Abnormal function of B cells involved in failed immune reconstitution**

To analyze the function of B cells in immune reconstitution after HIV infection, we expanded the sample size to perform single-cell transcriptome sequencing. We detected the differential expressed genes (DEGs) of B cells among the 8 HC individuals, 10 IR individuals, and 8 INR individuals. A total of 75 significant DEGs were identified in B cells between the HC and INR groups, 81 significant DEGs between the HC and IR groups, and 18 significant DEGs between the IR and INR groups (Figures 5A and 5B). The top 10 DEGs were annotated in volcano plots (Figure 5A). A total of 50 significant DEGs were identified both in the comparison group of HCs vs. INRs and the comparison group of HCs vs. IRs, which may be related to HIV infection. A DEG of IGKV1-5 was identified in the comparison group of HCs vs. INRs and the comparison group of HCs vs. IRs, which may be related to HIV infection. A DEG of IGKV1-5 was identified in the comparison group of HCs vs. INRs and the comparison group of HCs vs. IRs (Figure 4C).

We further analyzed our data with gene set enrichment analysis (GSEA) (Table S1). Compared to the HC group, the increased gene sets in the INR groups were associated with immunological signatures for IgG+IgA+ memory B cells and naive B cells. Compared to the HC group, the increased genes in the IR group were associated with immunological signature for IgD+ B cells, APC/C-mediated degradation of cell cycle proteins, and G2/M checkpoints. Compared to the IR group, the increased genes in the INR group were associated with IL-4/IL-13 signaling and immunologic signature for PBMCs after exposure to Gag/Pol/Nef proteins of HIV (Figures 5D and 5E). The relative expression of immunologic signature genes for the above B cell subtypes was shown in Figure S2. Furthermore, the immunological signature genes for IgD+ B cells were mainly enriched in the process of cell cycle and involved in the early phase of the HIV life cycle, while these functions were not enriched in immunological signature genes for naïve B cells (Figure S3). Together, these results suggested an abnormal function of memory B cells and naive B cells, including IL-4 and IL-13 pathway and APC/C-mediated G2/M cell cycle, involved in failed immune reconstitution after HIV-1 infection.
We further analyzed DE-Gs in naïve B cells, memory B cells, and plasma cells that related to HIV infection under the criteria of significantly differential expressed in the comparison of HC vs. INR and comparison of HC vs. IR, but not in the comparison of IR vs. INR. The results showed that 147 DEGs in naïve B cells, 113 DEGs in memory B cells, and 22 DEGs in plasma cells were identified. Functional analysis demonstrated that DEGs that related to HIV infection in naïve B cells enriched the pathway of cytokine signaling in the immune system and DEGs that related to HIV infection in memory B cells enriched the pathway of interferon signaling (Figure 54).

**DISCUSSION**

Failed restoration of CD4+ T cell level leads to increased morbidity and mortality of patients with HIV infection. The abnormal B cell function is a critical factor for the progression of HIV infection, including excessive activation of immature/transitional and short-life plasma B cells, declining the resident memory B cells response, and increased expression of Fas to accelerate apoptosis progression of B cells. Moreover, higher expression of IL-10 of Breg or Treg cells could reduce or inhibit viral-specific effector T cells response in HIV-infected individuals. The results of the present study indicate a potential way in which B cells affect immune status after HIV infection.

Our study demonstrated that the BCR repertoire diversity decreased in INRs, and there was a similar feature of CDR3 amino acids length between memory B cells and naïve B cells among HCs, IRs, and INRs. Furthermore, the dominant IGHV gene usage of memory B cells and naïve B cells were IGHV3 and IGHV4, and the dominant IGHJ gene usage was IGHJ4. However, the frequency of IGHV3 and IGHV4 gene usage was changed obviously after virus infection. Previous studies provided evidence that COVID-19 infection resulted in changed IGH and IGHJ gene usage. Our data indicated a low level of IGHV4-34 and IGHV4-39 gene usage of naïve B cells and a low level of IGHV3-23 and IGHV3-48 gene usage of memory B cells in the INR group. HIV infection could reduce the response of memory B cells, which may correspond to the reduced frequency of usage of paired IGHJ4 to most IGKV genes in INRs compared to HCs. Overall, we assumed that B cells experienced obvious specific V(D)J rearrangements in INR individuals, which played an impaired role in HIV infection.

In our study, high levels of TCL1A and IGHD in naïve B cells, high levels of AIM2 and TNFRSF13B in memory B cells, and high levels of XBP1 and MZB1 in plasma cells were observed, which were consistent with previous studies. Furthermore, we identified naïve B cells as the starting point of differentiation to memory B cells by cellular trajectory analysis, which was coincidental with the similar distribution of BCR V/J gene usage between naïve B cells and memory B cells in the present study. In addition, we showed a reduced naïve B cell set with low differentiation in INRs compared with IRs. It has been shown that the excessive activation and apoptosis of immature B cells observed in advancing HIV-infected patients could explain the less differentiated degree of naïve B cells, and the correlation between declined immune response and a low percentage of naïve T cells was observed in HIV-infected individuals on ART. The high differentiation in memory B cells and plasma cells was associated with persistent or chronic HIV infection, even in HIV patients with restored CD4+ T cells after ART. The present study displayed a decreased percentage of memory B cells and low differentiation of naïve B cells in INRs, indicating a vital role of naïve B cells in immune reconstitution after HIV infection.

We further analyzed the changed function of B cells using GSEA. We revealed a gene set associated with immunological signatures for IgG/IgA memory B cells was highly expressed in INRs rather than in HCs. A recent study demonstrated that IgA and IgG antibodies of B cell lineages were related to the development of vulnerability to the HIV-1 site. The level of memory B cells was correlated with CD4+ T cell counts in...
patients, which could influence HIV-infected disease progression. Abnormal memory B cells could contribute to the ineffectiveness of the antibody response in HIV-infected individuals. The less differentiation of naive B cells in INRs than in IRs suggests naive B cells could be influenced by immune reconstruction. In addition, abnormal expression of the gene sets enriched in the function of APC/C-mediated degradation of cell cycle proteins and G2/M checkpoints were observed in IRs rather than in HCs. HIV-1 Vif degrades essential phosphorylation regulators to induce G2/M cell-cycle arrest, and HIV-1 Vpr promotes cell-cycle arrest through the depletion of CDC137. Combining the results of the present study, it was indicated that the abnormal cell cycle of B cells affected immune status after HIV-1 infection. Furthermore, a gene set associated with the signaling pathway of IL-4/IL-13 was highly expressed in INRs rather than in IRs. Our results indicated that low differentiation of naive B cells may be correlated with abnormal B cell (IgG+/IgA+ memory B cells and IgD+ B cells) response, and the altered cell cycle of these abnormal B cells may promote failed immune restoration after HIV infection.

In conclusion, our BCR repertoire and B cell subsets transcriptome analysis provides comparative information on the role of B cell-mediated immune response in immune reconstitution after HIV infection. We demonstrated that the BCR clonotypes were decreased in INRs, low differentiation of naive B cells with special immunological signature genes for IgD+ B cells was related to the immune status of HIV-1-infected patients, and the altered cell cycle of these B cells may promote failed immune restoration after HIV infection. Future studies are necessary to investigate the development and function of naive B cells after HIV infection.

Limitations of the study
We would like to unravel the potential function of B cells in HIV infection in the present study. Although, we performed function annotation for B cells among different groups, and an altered cell cycle of B cells was identified in the INR group, more specialized experiments need to be conducted to verify the altered functions of B cells in INRs. Also, it will be important to determine the relationship between heterogeneity of B cell differentiation and immune reconstitution.

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Supplemental information can be found online at https://doi.org/10.1016/j.isci.2022.105559.

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AUTHOR CONTRIBUTIONS

Conceptualization, Y.Q.K., K.H.W., and J.H.W.; Methodology, J.J. and Y.Z.; Resources, D.L., J.Q.Y., J.H.M., J.H.W., and X.L.Z.; Investigation, J.J., Y.Z., D.L., and J.H.M.; Formal Analysis, J.J., Y.Z., D.L., J.H.W., and Y.Q.K.; Writing – Original Draft, J.J. and Y.Z.; Writing – Review & Editing, Y.Q.K.; Supervision, Y.Q.K.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Chemicals, peptides, and recombinant proteins** | | |
| Chromium Single Cell 5′ Library Kit | 10X GENOMICS | PN-1000002 |
| Chromium Single Cell 5′ Gel Bead Kit | 10X GENOMICS | PN-1000003 |
| Chromium Single Cell 3′/5′ Library Construction Kit | 10X GENOMICS | PN-1000020 |
| Chromium Single Cell A Chip Kits | 10X GENOMICS | PN-1000009 |
| Chromium i7 Multiplex Kit | 10X GENOMICS | PN-120262 |
| Dynabeads MyOne™ SILANE | 10X GENOMICS | PN-2000048 |
| Ficoll-Paque™ PREMIUM density gradient media | GE Health | 17-5446-02 |
| Buffer EB | Qiagen | Cat#19086 |
| SPRselect Reagent Kit | Beckman Coulter | Cat#B23318 |
| High Sensitivity DNA assay Kit | Agilent | SD-U0000034 Rev. D |

| Deposited data | | |
| scRNA-seq for HIV-infected patients | This paper | BioProject: PRJNA783363 |
| scRNA-seq for healthy controls | Ref. [20] | Genome Sequence Archive: HRA001149 |

| Software and algorithms | | |
| R | R Development Core Team | cran.r-project.org |
| SingleR | R package | http://www.bioconductor.org/ |
| Seurat 3.1.1 | Ref.31 | http://satijalab.org/seurat |
| Cell Ranger | 10x Genomics | https://www.10xgenomics.com/support |
| Monocle | R package | http://cole-trapnell-lab.github.io/monocle-release/ |
| GSEA | GSEA and MSigDB Team | https://www.gseamsigdb.org/ |
| GraphPad Prism | GraphPad Software LLC | https://www.graphpad-prism.cn/ |

RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Yi-Qun Kuang (yq610433@hotmail.com).

Materials availability
This study did not generate new unique reagents.

Data and code availability
- Single-cell RNA-seq data have been deposited at Sequence Read Archive (SRA) database in National Center for Biotechnology Information (NCBI) and are publicly available as of the date of publication. This paper does not report original code. The accession number for the scRNA-seq data reported in this paper is PRJNA783363 (BioProject).
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.
- Codes of preprocessing, normalization, clustering and plotting of single-cell datasets are available on Zenodo: https://doi.org/10.5281/zenodo.7262476.
EXPERIMENTAL MODEL AND SUBJECT DETAILS

Human subjects
HIV-infected IR and INR were enrolled under the following criteria: (1) HIV-1 positive; (2) aged 30–50 years old; (3) male; (4) Han ethnicity; (5) sustainable viral load suppression (<50 copies/mL plasma) over 2 years after ART; (6) no HIV-1 viral load blipped during ART; (7) no immune reconstitution inflammatory syndrome (IRIS) occurred during ART; and (8) no hepatitis C virus (HCV)/HBV/tuberculosis (TB) co-infection. Ultimately, 10 HIV-infected IR (mean age of 39.40 ± 9.94, CD4+ T cells >500 cells/μL) and 8 HIV-infected INR (mean age of 43.67 ± 8.74, CD4+ T cells <350 cells/μL) were enrolled at the Third People’s Hospital of Kunming City, the Zhengzhou Sixth People’s Hospital, and the Guangzhou Eighth People’s Hospital. In addition, 8 male Han ethnicity healthy controls (HC) with a mean age of 38.00 ± 7.12 were used as the control group, and the transcriptome and BCR sequencing data of the 5 HC subjects were obtained from selected samples under a 10x Genomics single-cell platform by Illumina sequencing in previously published work.20

Ethics statement
This study was approved by the Ethics Committee of the First Affiliated Hospital of Kunming Medical University (No. 2018-L-42).

METHOD DETAILS

Preparation for PBMC suspension
Approximately 5 mL of anti-coagulated venous blood of 26 participants (8 HCs, 10 IRs and 8 INRs) was obtained and processed within 1 hour of collection. PBMCs were prepared by using Ficoll Paque™ PREMIUM density gradient media (GE Health, 17-5446-02) according to the manufacturer’s instructions.

BCR sequencing and data processing
The construction and sequencing of the BCR library were performed on platforms of Chromium™ Single Cell V(D)J Solution (Gene Denovo), which used GemCode to package barcoded single-cell 5’ gel beads (10x genomics, PN-1000020, PN-1000009, PN-1000003), a master mix with cells, and partitioning oil on a microfluidic chip to generate single-cell Gel Bead-In-EMulsion (GEMs). Then GEMs were recycled and purified by Dynabeads MyOne™ SILANE (10x genomics, PN-2000048), the single-cell 5’ gel bead was dissolved and any co-partitioned cell was lysed by Buffer EB (Qiagen, Cat#19086) to release mRNAs and performed reverse transcription to produce barcoded, full-length cDNA from poly-adenylated mRNAs. After incubation, the GEMs were broken and pooled post-GEM-RT were recovered, barcoded cDNAs were extracted and purified by SPRSelect Reagent Kit (Beckman Coulter, Cat#B23318), then full-length V(D)J segments can then be enriched and amplified by PCR to generate sufficient mass for library construction.

The amplification products were enzymatic fragmentation and the sizes were selected by using SPRSelect Reagent Kit (Beckman Coulter, Cat#B23318) to generate variable length fragments that collectively span the V(D)J segments of the enriched BCR transcripts. By adding primer sequences and sample index, the enriched transcripts were used to construct the library via end repair, A-tailing, adaptor ligation, and PCR amplification. The single-cell V(D)J reagent kit protocol produces V(D)J enriched and 5’ gene expression Illumina-ready sequencing libraries, a library comprising standard Illumina paired-end constructs which begin and end with P5 and P7, the libraries were quality controlled by Agilent 2100 Bioanalyzer using High Sensitivity DNA assay Kit (Agilent, Cat# 02100-00034 Rev. D). The final libraries contained the P5 and P7 priming sites were used in Illumina bridge amplification. After sequencing, we used Cell Ranger (https://support.10xgenomics.com/single-cell-vdj/software/overview/welcome) to assemble, quantify, and annotate paired V(D)J transcript sequences.

Identification of clonotyping and diversity
Cell barcodes were grouped into clonotypes if they share the same set of productive CDR3 nucleotide sequences. A three-layer donut chart was used to visualize the repertoire clonality. Rarefaction analysis was used to compare the repertoire diversity, rarefaction curves were interpolated from 0 to the current sample size and then extrapolated up to the size of the largest of the samples. Repertoire information of all samples was exhibited using a set of basic features: CDR3 abundance, CDR3 length, Variable (V) and Joining (J) segment usage, the length of V/J gene in the CDR3 region, and V-J gene paired frequency in CDR3 junctions.
Single-cell RNA sequencing and data processing

The PBMC suspensions of 26 participants were loaded on a 10x Genomics GemCode Single-cell instrument that generates single-cell Gel Bead-In-EMulsion (GEMs). Single-cell capturing and downstream library constructions were performed using Chromium i7 Multiplex Kit (10x GENOMICS, PN-120262) and Chromium Single Cell 5’ Library Kit (10x GENOMICS, PN-1000002). After dissolution of the Gel Bead in a GEM, primers containing Illumina® R1 sequence, a 16-nt 10 x barcode, a 10-nt Unique Molecular Identifier (UMI), and a poly-dT primer sequence were released and mixed with cell lysate and Master Mix. Barcoded full-length cDNAs were then reverse-transcribed from poly-adenylated mRNAs. Residual biochemical reagents and primers were removed by using Dynabeads MyOne™ SILANE (10x GENOMICS, PN-2000048) in a post-GEM reaction mixture. Then, the barcoded cDNAs were amplified by PCR to construct the library. Read primers R1 and R2, sample index P5 and P7 were then added during library construction via End Repair, A-tailing, Adaptor Ligation, and PCR. The final libraries containing the P5 and P7 primers were used in Illumina bridge amplification.

ScRNA-seq was performed on the Illumina platform by Gene Denovo Biotechnology Co. 10x Genomics Cell Ranger was used to convert raw data files to FASTQ files. Reads with low-quality barcodes and UMIs were filtered out and then clean reads mapped to reference genome, and reads mapped to transcriptome uniquely and intersecting an exon at least 50% were considered for UMI counting. Quantification was performed based on corrected UMI sequences and valid barcodes, the cells by gene matrices were produced via UMI counting and cell barcode calling. The cells with gene matrices for each sample were individually imported to Seurat version 3.1.1. Cells with a high number of UMIs (≥ 8000), a high proportion of mitochondrial genes (≥ 10%), and abnormal gene numbers <500 or >4000 were filtered out. Then, we used “LogNormalize” to normalize gene expression and used Seurat to minimize the effects of batch effect and behavioral conditions on clustering. We performed canonical correspondence analysis, and used Mutual Nearest Neighbors (MNN) to construct correspondence relationships among cells, the correspondence relationship was used as anchors to complete data integration and batch normalization. Furthermore, we measured the correlation between each cell subset, then used Uni-form Mani-fold Approximation and Projection for Dimension Reduction (UMAP) to visualize cell clusters. Cell type annotation was performed by the SingleR R packages.

Trajectory analysis of cell differentiation

Cell trajectory was analyzed using the matrix of cells and gene expressions by Monocle. Monocle reduced the space down to one with two dimensions and ordered the cells (sigma = 0.001, lambda = NULL, param. gamma = 10, tol = 0.001).32 The trajectory has a tree-like structure, including tips and branches. The branches occur because cells execute alternative gene expression programs. Monocle develops BEAM to test for branch-dependent gene expression by formulating the problem as a contrast between two negative binomial GLMs.33 In addition, monocle can find genes that are differentially expressed between groups of cells and assess the statistical significance of those changes, we identified key genes related to the development and differentiation process with FDR <10^{-5}.

Differential gene expression and function annotation

Differentially expressed genes (DEGs) were analyzed between HCs and IRs, HCs and INRs, IRs and INRs. DE-Gs were selected under the condition of log2 fold change (FC) ≥ 1 and adjusted p < 0.05. For functional annotation, gene set enrichment analysis (GSEA) was performed with 1000 permutations for each test. Nom p < 0.05 and |NES| > 1 was statistically significant for GSEA.

QUANTIFICATION AND STATISTICAL ANALYSIS

The statistical analysis was performed using GraphPad Prism 8.0 (GraphPad Software LLC) with an unpaired Student’s t test in the comparison groups. p < 0.05 was considered statistically significant, values denoted with * reflected significance levels as follows: *p < 0.05; **p < 0.01; ***p < 0.001. Results were shown as the mean ± standard error of the mean (SEM).