ABSTRACT

Cryopreservation and thawing of PBMCs are inevitable processes in expanding the scale of experiments in human immunology. Here, we carried out a fundamental study to investigate the detailed effects of PBMC cryopreservation and thawing on transcriptomes. We sorted Tregs from fresh and cryopreserved/thawed PBMCs from an identical donor and performed single-cell RNA-sequencing (scRNA-seq). We found that the cryopreservation and thawing process minimally affects the key molecular features of Tregs, including \( \text{FOXP3} \). However, the cryopreserved and thawed sample had a specific cluster with up-regulation of genes for heat shock proteins. Caution may be warranted in interpreting the character of any cluster of cells with heat shock-related properties when cryopreserved and thawed samples are used for scRNA-seq.

Keywords: Cryopreservation; Single-cell analysis; Transcriptome; T-lymphocytes, regulatory

INTRODUCTION

No matter how sophisticated the experimental techniques of modern immunology, a batch effect originates from non-biological factors (1,2). The batch effect may corrupt the data produced by experiments, and it is difficult to correct later. A large-scale and simultaneous experiment under identical conditions is an effective approach to remove batch effects. In translational research using specimens originating from a defined cohort with human subjects, the cryopreservation and thawing process is inevitable in conducting a large-scale and simultaneous experiment (3,4). When PBMCs are tested in protein-based \textit{ex vivo} techniques, such as enzyme-linked immune absorbent spot and flow cytometry, the cryopreservation and thawing protocols provide stable and reproducible immunophenotype data (5,6). However, significant gene expression changes are occasionally detected after the cryopreservation and thawing process using comprehensive methods, such as transcriptome analysis (7).

Single-cell RNA-sequencing (scRNA-seq) is a powerful experimental technique for characterizing heterogeneous immune cell types at the molecular level (8). In human immunology studies, scRNA-seq is widely applied for various specimens, including normal
tissues, tumor tissues, bone marrow, and PBMCs (9-12). However, current single-cell sequencing using a water-in-oil microdroplet platform also has a significant batch effect among different sets of experiments, and it has been attempted to correct the batch effect by a bioinformatics approach (13). To minimize the batch effect, cryopreservation and thawing of specimens for scRNA-seq analysis has been introduced (14). Previous studies comparing fresh cells and cryopreserved/thawed cells showed that DMSO based cryopreservation is a reliable method without significant transcriptomic changes (15).

During immune landscape analysis of Tregs, we had a chance to compare the scRNA-seq data of Tregs sorted from fresh and cryopreserved/thawed PBMCs from an identical human subject. Our results suggest that unique transcriptomic features of Tregs are conserved after a cryopreservation and thawing process, but a specific cluster with heat shock protein gene expression was found only in the cryopreserved/thawed sample.

MATERIALS AND METHODS

Study subject
Peripheral blood was obtained from a single healthy donor twice at a 1-week interval for preparation of cryopreserved/thawed and fresh PBMCs. PBMCs were isolated from whole blood via standard Ficoll-Paque (GE Healthcare, Uppsala, Sweden) density gradient centrifugation. PBMCs were cryopreserved and thawed using a standard protocol (15). The freezing media comprised 70% FBS, 20% Roswell Park Memorial Institute (RPMI) 1640 media, and 10% DMSO. This study protocol was reviewed and approved by the Institutional Review Boards (KH2018-118). Written informed consent was obtained from the subject prior to the study.

Isolation of Tregs from PBMCs
Untouched CD4+CD25+CD127-Tregs were isolated from PBMCs using the EasySep Human CD4+CD127lowCD25+ Treg Isolation Kit (STEMCELL Technologies, Vancouver, Canada) according to the manufacturer’s instructions.

scRNA-seq
We generated scRNA-seq libraries using the Chromium Single Cell 3' Library & Gel Bead Kit v2 (10X Genomics, Pleasanton, CA, USA) according to the manufacturer’s instructions. Briefly, thousands of cells were separated into nanoliter-scale droplets. In each droplet, cDNA was produced by reverse transcription, adding a cell barcoding sequence and unique molecular identifier (UMI) to each cDNA molecule. Libraries were constructed and sequenced at a depth of approximately 50,000 reads per cell by the Novaseq 6000 platform (Illumina, San Diego, CA, USA).

scRNA-seq data processing
The sequenced data were de-multiplexed using mkfastq (Cellranger, v3.0.2; 10X Genomics) to generate fastq files. After de-multiplexing, the reads were aligned to the human reference genome (GRCh38; 10X Cellranger reference GRCh38 v3.0.0) and feature-barcode matrices generated using the cellranger count, and then aggregated by cellranger aggr using default parameters. Subsequent analysis was performed using Seurat R package v3.1.5 (R Foundation, Vienna, Austria). Mitochondrial read counts were collected using PercentageFeatureSet function with the following option: pattern = "^MT-". In each cell, the gene expression was normalized based on the total read count and log-transformed. After
log-transformation, principal component analysis was performed and PC1 to PC6 applied following clustering and uniform manifold approximation and projection (UMAP).

**Identification of differentially expressed genes (DEGs)**
To identify DEGs for any 2 transcriptome profiles, we utilized the Wilcoxon rank sum test and receiver operating characteristics method in Seurat’s implementation based on a Bonferroni-adjusted p<0.05 and log2 fold change >0.25.

**Gene Ontology analysis for biological pathways (GO-BP)**
The Gene Ontology analysis was performed using Enrichr (https://amp.pharm.mssm.edu/Enrichr).

**RESULTS AND DISCUSSION**

**Single-cell transcriptomes of fresh and cryopreserved/thawed cells**
Applying a 10X Genomics scRNA-seq platform, we analyzed 4,990 cells from the fresh sample and 4,495 cells from the cryopreserved/thawed sample. The fresh sample had 18,449 median genes per cell, and the cryopreserved/thawed sample had 17,227 median genes per cell. Captured UMI, which represents the number of genes per cell, was relatively higher in the fresh sample (median 986; 1st–3rd quartile, 518–691) than the cryopreserved/thawed sample (median 600; 1st–3rd quartile, 858–1,131; *Fig. 1A*). The number of gene species was also relatively higher in the fresh sample (median, 3,503; 1st–3rd quartile, 3,011–4,019) than the cryopreserved/thawed sample (median 1,435; 1st–3rd quartile, 1,229–1,674; *Fig. 1A*). The proportion of mitochondrial transcripts, which represents contamination by dead or dying cells, was similar in the fresh sample (median 2.1; 1st–3rd quartile, 1.7–2.5) and the cryopreserved/thawed sample (median 2.3; 1st–3rd quartile, 1.8–2.5; *Fig. 1A*).

Next, we subjected all cells in each group to UMAP using the highly variable genes in each group and the Seurat package (16). We found six different clusters for the fresh sample (clusters 0–5) and seven different clusters for the cryopreserved/thawed sample (clusters 0–6; *Fig. 1B*). Cell cycle-associated gene expression was uniformly distributed among clusters (*Fig. 1C*). Moreover, the expression levels of key molecules defining Tregs, including \(CD3E\) (CD3), \(CD4\), \(IL2RA\) (CD25), and \(FOXP3\) were homogenous and not polarized to specific clusters in both the fresh and cryopreserved/thawed samples (*Fig. 1D–G*). This result indicates that the molecular identity of Tregs is still conserved across the different clusters after a cryopreservation and thawing process at the single-cell transcriptome level, even though the amount and variability of gene expression are reduced after cryopreservation and thawing.

**Cluster-specific up-regulated genes in fresh and cryopreserved/thawed cells**
To characterize each cluster identified in the fresh and cryopreserved/thawed samples, we investigated DEGs among different clusters in the fresh and cryopreserved/thawed samples. Representative up-regulated genes from each cluster were noted on a heatmap for the fresh sample (*Fig. 2A*) and the cryopreserved/thawed sample (*Fig. 2B*). A detailed list of genes up-regulated in each cluster is given in **Supplementary Tables 1 and 2**.

Immunologically well-known genes were found among representative up-regulated genes in each Treg cluster. For example, \(CCR7\), a marker of naïve or central memory T cells, was significantly up-regulated in clusters 0 and 2 in the fresh sample and in clusters 0 and 3 in
the cryopreserved/thawed sample (Fig. 2A and B, red arrows). KLRB1 and TNFRSF18 were commonly up-regulated in cluster 5 in the fresh sample and cluster 6 in the cryopreserved/thawed sample (Fig. 2A and B, blue arrows), which are the genes for surface protein CD161 and glucocorticoid-induced TNF receptor, respectively. Intriguingly, the genes associated with heat shock proteins, such as HSPA1A, HSPA1B, HSPA6, HSPB1, HSPE1, and HSP90AA1, were exclusively up-regulated in cluster 4 in the cryopreserved/thawed sample (Fig. 2B, pink box). When we plotted the variable genes in the fresh and cryopreserved/thawed samples, genes for heat shock proteins, including HSPA1A, were specifically variable in the cryopreserved/thawed samples (Fig. 2C). This cryopreservation/thawing-specific gene expression pattern was exemplified at single-cell resolution by HSPA1A and confirmed that cluster 4 was specifically found in Tregs from the cryopreserved/thawed sample (Fig. 2D).
Figure 2. Cluster-specific up-regulated genes in fresh and cryopreserved/thawed cells. (A) Heatmaps presenting up-regulated genes from each cluster for the fresh sample and (B) the cryopreserved/thawed sample. The arrows indicate the commonly up-regulated genes in the clusters from both fresh and cryopreserved/thawed samples. The pink box indicates cluster 4 in the cryopreserved/thawed sample, with multiple up-regulated heat shock protein genes. (C) Plots of the variable genes from the fresh or cryopreserved/thawed samples, or both. (D) UMAP plot of the representative expression pattern for HSPA1A.
Identification of unique transcriptional signatures in cryopreserved/thawed cells

To further evaluate the gene expression changes associated with the cryopreservation and thawing process, we investigated the variably expressed genes in fresh and cryopreserved/thawed samples. Among 910 genes up-regulated in any cluster from the cryopreserved/thawed sample, 561 (61.6%) genes overlapped with the genes up-regulated in the clusters in the fresh sample (Fig. 3A). When we analyzed the gene lists for each cluster in GO-BP, 228 of 391 (58.3%) pathways in the cryopreserved/thawed sample overlapped with the pathways up-regulated in the clusters in the fresh sample (Fig. 3A; Supplementary Tables 3 and 4).

Next, we analyzed the similarity of each cluster between the fresh and cryopreserved/thawed samples based on GO-BP. Cluster 0 in the fresh sample had 27 enriched pathways and was most similar to cluster 5 in the cryopreserved/thawed sample, with 26 overlapping pathways (96.3%; Fig. 3B). Cluster 3 in the fresh sample had 188 enriched pathways and was most similar to cluster 1 in the cryopreserved/thawed sample, with 130 overlapping pathways (69.1%; Fig. 3B).

We evaluated the DEGs specifically up-regulated in cluster 4 in the cryopreserved/thawed sample, which demonstrated up-regulation of heat shock protein genes. T-cell activation-related genes, including CD69, JUN, DUSP1, and DUSP2, were commonly up-regulated in cluster 6 in the cryopreserved/thawed sample (Fig. 3C). Commonly up-regulated genes between clusters 4 and 6 (n=24) were analyzed by GO-BP, finding that pathways related to ER stress, apoptosis, cell cycle arrest, and DNA damage were significantly enriched (Fig. 3D; Supplementary Table 5). These data indicate that cluster 4 resembles cluster 6.
in gene expression signatures, but acquires expression of heat shock protein genes in the cryopreservation and thawing process.

In conclusion, we show that cryopreservation and thawing minimally affect the key molecular features of Tregs although cell clustering results were not identical. However, the cells with up-regulated heat shock protein genes were clearly clustered in Tregs after cryopreservation and thawing, and this cluster may represent the signature of recent heat exposure applied during a cryopreservation and thawing process. Although cryopreserved and thawed samples can be used for scRNA-seq, caution might be warranted in interpreting the character of any cluster of cells with heat shock-related properties.

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SUPPLEMENTARY MATERIALS

Supplementary Table 1
Marker genes for each cluster from the fresh sample
Click here to view

Supplementary Table 2
Marker genes for each cluster from the cryopreserved/thawed sample
Click here to view

Supplementary Table 3
DEGs and associated biological pathways for each cluster from the fresh sample
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Supplementary Table 4
DEGs and associated biological pathways for each cluster from the cryopreserved/thawed sample
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Supplementary Table 5
DEGs and associated biological pathways for commonly up-regulated genes (n=24) from both cluster 4 and 6 in the cryopreserved/thawed cells
Click here to view
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