Cryopreservation of microglia enables single-cell RNA sequencing with minimal effects on disease-related gene expression patterns

Highlights

- High-quality scRNA-seq can be performed on cryopreserved microglia
- Cryopreservation of microglia does not affect disease-related gene expression
- Expression of immediate-early genes (IEG) is increased in cryopreserved microglia
- scRNA-seq of cryopreserved microglia reveals gene expression changes, including IEG

Morsey et al., iScience 24, 102357 April 23, 2021 © 2021 The Author(s).
https://doi.org/10.1016/j.isci.2021.102357
Cryopreservation of microglia enables single-cell RNA sequencing with minimal effects on disease-related gene expression patterns

Brenda Morsey, Meng Niu, Shetty Ravi Dyavar, Courtney V. Fletcher, Benjamin G. Lamberty, Katy Emanuel, Anna Fangmeier, and Howard S. Fox

SUMMARY
Microglia play a key role in brain development, normal homeostasis, and neurodegenerative disorders. Single-cell technologies have led to important findings about microglia, with many animal model studies using single-cell RNA sequencing (scRNA-seq), whereas most human specimen studies using archived frozen brains for single-nucleus RNA sequencing (snRNA-seq). However, microglia compose a small proportion of the total brain tissue; snRNA-seq depletes expression of microglia activation genes that characterize many diseases. Here we examine the use of purified, cryopreserved microglia for scRNA-seq. Comparison of scRNA-seq on paired fresh and cryopreserved microglia from rhesus monkeys revealed a high level of correlation of gene expression between the two conditions. Disease-related genes were relatively unaffected, but an increase in immediate-early gene expression was present in cryopreserved cells. Regardless, changes in immediate-early gene expression are still detectable. Cryopreservation of microglia is a suitable procedure for prospectively archiving samples.

INTRODUCTION
Microglia and central nervous system (CNS)-associated macrophages (CAM) make up the myeloid component of the CNS. In addition to their innate immune functions they are critical during brain development and for normal homeostasis. These cells also participate in the pathogenesis of CNS disorders, as well as protection and recovery from disease. The development and use of single-cell RNA sequencing (scRNA-seq) has led to significant advances in the study of many cell types, including microglia and CAM (Bonham et al., 2019; Geirsdottir et al., 2019; Gerrits et al., 2020; Goldmann et al., 2016; Hammond et al., 2019; Jordao et al., 2019; Keren-Shaul et al., 2017; Li et al., 2019; Masuda et al., 2019; Olah et al., 2018; Sousa et al., 2018; van der Poel et al., 2019). The study of microglia and CAM is complicated by their presence in the brain (and spinal cord), ruling out easy sampling such as for blood cells. Furthermore, in humans, access to biopsies and resections is quite limited.

Microglia and CAM originate from the yolk sac and populate the CNS during embryogenesis, whereas macrophages within the cerebrospinal fluid (CSF)-producing choroid plexus are derived from yolk sac precursors but are replaced during life by blood-derived cells (Goldmann et al., 2016). When fresh brain tissue is obtained, the tissue can be disassociated and cells purified for scRNA-seq; however, both these processes take time, and depending on when the tissue is procured (from experimental studies, the operating room, or the autopsy suite) and the specific protocol followed, it can be difficult to complete all processing in time to carry out the initial steps of scRNA-seq. Furthermore, microglia only make up approximately 10% of CNS cells (Lawson et al., 1992), limiting the number that can be studied.

To avoid the timing issue researchers have performed single nucleus RNA sequencing (snRNA-seq) using specimens stored frozen in tissue banks. For CNS studies, this technique has been frequently used to study specimens from those with Alzheimer disease (AD) (Del-Aguila et al., 2019; Grubman et al., 2019; Mathys et al., 2019; Zhou et al., 2020). Although archived samples still suffer from a low proportion of nuclei from microglial cells, the ability to use these archived samples represents a substantial advantage and may yield insight into disease pathogenesis. However, although snRNA-seq has been found to be applicable for many cell types, recently it was found to be not appropriate for the study of microglia. This is
due to lack of detection of activation genes, which typify many disease states such as AD, and may help explain the inconsistency found between single-cell expression studies on AD (Thrupp et al., 2020).

A number of procedures have been used to preserve cells after isolation for future use in scRNA-seq, including methanol, paraformaldehyde, dithio-bis(succinimidyl propionate), and a commercial proprietary reagent (Cell-Cover) fixation, as well as cryopreservation in dimethyl sulfoxide (DMSO) (Alles et al., 2017; Attar et al., 2018; Chen et al., 2018; Donlin et al., 2018; Gierahn et al., 2017; Guillaumet-Adkins et al., 2017; Thomsen et al., 2016; Zhou et al., 2020). A comparison of freshly isolated cells with cells cryopreserved in DMSO or fixed under two different conditions concluded that cryopreservation in DMSO was optimal for scRNA-seq (Wohnhaas et al., 2019). We have recently described the use of these cryopreserved preparations in an scRNA-seq study (Niu et al., 2020). In order to investigate the effect of cryopreservation on scRNA-seq of microglia, we compared scRNA-seq results from freshly isolated and cryopreserved cells from two rhesus monkeys. These monkeys had been infected with SIV and treated with combination antiretroviral therapy, which suppressed the blood plasma virus to below the limit of detection. Fresh, enriched brain macrophages and microglia were isolated from two SIV-infected rhesus monkeys and then further purified (Figure 1, top), followed by capture on the 10x Genomics platform, and cDNA library production. Microglia were also cryopreserved using DMSO, and two months later, the cells were thawed and purified (Figure 1, bottom), captured on the 10x Genomics platform, and cDNA libraries were similarly synthesized.

RESULTS

Fresh and cryopreserved microglia yield proper metrics for scRNA-seq studies

We routinely isolate enriched populations of microglia and macrophages from the brains of nonhuman primates in our studies on SIV as a model for the neurological effects of HIV and cryopreserve them for use in our future experiments (Bortell et al., 2018; Chaudhuri et al., 2013; Marcondes et al., 2001; Watry et al., 1995). Recently we reported the use of these cryopreserved preparations in an scRNA-seq study (Niu et al., 2020). In order to investigate the effect of cryopreservation on scRNA-seq of microglia, we compared scRNA-seq results from freshly isolated and cryopreserved cells from two rhesus monkeys. These monkeys had been infected with SIV and treated with combination antiretroviral therapy, which suppressed the blood plasma virus to below the limit of detection. Fresh, enriched brain macrophages and microglia were isolated from two SIV-infected rhesus monkeys and then further purified (Figure 1, top), followed by capture on the 10x Genomics platform, and cDNA library production. Microglia were also cryopreserved using DMSO, and two months later, the cells were thawed and purified (Figure 1, bottom), captured on the 10x Genomics platform, and cDNA libraries were similarly synthesized.

Figure 1. Sequence of flow cytometric purification of monkey microglia
Forward and side scatter area (Scatter, far left) were used to select the cell population, followed by side scatter area and height to eliminate doublets (Singlets, second from left). Viable cells, determined by dye exclusion from live cells, were selected to eliminate dead cells (Live, third from left) followed by selection of CD11b-positive cells (far right). The proportion of cells selected in each gate within each panel is indicated, as are the fluorochromes for the two rightmost panels. Data from animal B6T are shown for the fresh (top) and cryopreserved (bottom) specimens. The fresh cells had a live, CD11b-negative population (top right panel), whereas these were not present in the cryopreserved cells, instead a population of dead cells was present (bottom, third panel from left). Both these populations mapped to the low FSC-A/SSC-A cells present in the gate shown in the top and bottom left panels.
cDNA libraries from both sets of cells were subjected to sequencing, and reads mapped to the rhesus genome. The scRNA-seq data from both animals and both conditions resulted in high-quality sequencing statistics (Table 1), notable for a high number of genes detected with little difference between the fresh and cryopreserved cells and low variability between the four samples. However, although equal numbers of cells were loaded for capture, an average of 33% fewer cells were captured in the cryopreserved samples compared with the fresh. The data were subjected to quality assurance and quality control followed by graph-based clustering and projection in three dimensions using Uniform Manifold Approximation and Projection (UMAP) (Figure S1). Examination of maker genes (those significantly overexpressed in each cluster) revealed a small cluster of lymphoid cells (likely cytotoxic T cells and/or natural killer cells, with marker genes including CD3D, NKG7, GZMB, and LCK), which were then excluded from further analysis. The remaining cells expressed typical markers of microglia, such as P2RY12, CSF1R, AIF1, and CX3CR1 (Figure S1).

Gene expression patterns are closely correlated between cryopreserved and fresh microglia, including disease-associated genes

We then explored single microglial cell gene expression from the two animals under the fresh and cryopreserved conditions. The gene expression levels from the different monkeys were highly correlated from both the fresh and cryopreserved conditions (Figure 2A). In comparing the cryopreserved with the fresh cells, a low proportion showed a 2-fold or greater change (all with a false discover rate <0.05), with 2.0% increased and 0.4% decreased in the cryopreserved cells compared with the fresh cells (Figure 2B). We then curated the gene list to those that are present in genomic DNA (as opposed to mitochondrial DNA), encoded proteins (versus regulatory RNAs), and excluded those encoding ribosomal proteins. A high level of correlation was again seen within the curated list, with only 1.8% of the gene expression values increased, and 0.1% decreased, in the cryopreserved cells compared with the fresh cells (Figure 2B).

snRNA-seq, which has been a key method in the analysis of brain cells and other cells from frozen archived specimens, has been found to be inappropriate for assessment of human microglia due to loss of ability to assess expression of key genes involved in disease pathogenesis when compared with scRNA-seq (Thrupp et al., 2020). As both monkeys and humans are primates, we assessed our findings on the effect of cryopreservation on microglial scRNA-seq in relation to that study. Using the criteria of expression change of 2-fold or greater change (as used by Thrupp et al., 2020), we found that 8.0% of the genes that were underrepresented in nuclei in snRNA-seq were increased in the cryopreserved compared with fresh cells we analyzed by scRNA-seq, whereas 0.9% were decreased (Figure 2C).

To obtain functional information on the expression changes resulting from cryopreservation, we analyzed a list of differentially expressed genes (DEGs) (538 genes showing a fold change of >|1.5| and false discovery rate of <0.05, Table S1) between the fresh and cryopreserved cells using Ingenuity Pathway Analysis (IPA). This revealed canonical pathway alterations (Table S2) including three signaling pathways: acute phase response, IL6, and TNFR2, which contained overlapping DEGs (Figure 2D). We next submitted the DEGs to Metascape to perform interactome analysis (using MCODE), which revealed densely connected protein networks, and subsequently performed gene set enrichment analysis. One network (Figure 2E, left) was highly enriched for cellular response to heat (gene ontology pathway 0034605, p = 10^{-7}), and a second (Figure 2E, right) was highly enriched for the AP1 pathway (protein interaction database pathway M167, p = 10^{-16.4}). The AP1 pathway is activated as part of the acute phase response (Hattori et al., 1993), linking these two independent analyses.

### Table 1. Sequencing statistics

| Sample   | Estimated number of cells | Mean reads per cell | Median genes per cell | Sequencing saturation | Reads mapped confidently to genome | Fraction reads in cells | Total genes detected | Median UMI counts per cell |
|----------|---------------------------|---------------------|-----------------------|-----------------------|-----------------------------------|------------------------|-----------------------|--------------------------|
| 85T fresh| 8,479                     | 48,148              | 1,429                 | 83.80%                | 86.30%                            | 93.50%                 | 19,535                | 2,551                    |
| 86T fresh| 6,338                     | 64,263              | 1,928                 | 72.50%                | 85.00%                            | 93.30%                 | 20,676                | 3,982                    |
| 85T cryo | 4,838                     | 47,217              | 1,927                 | 66.00%                | 86.70%                            | 93.40%                 | 19,864                | 3,772                    |
| 86T cryo | 4,921                     | 45,556              | 1,727                 | 72.30%                | 86.60%                            | 94.10%                 | 19,540                | 3,464                    |

The scRNA-seq statistics of the samples from animals 85T and 86T, with both fresh and cryopreserved preparations. Data were generated by the Cell Ranger software of 10x Genomics.
Figure 2. Evaluation of gene expression patterns between fresh and cryopreserved microglia, and comparison with other studies

In the panels with dot plots, the purple diagonal lines indicate ±2-fold change, open black circles < |2|-fold change, red up-pointing triangles ≥2-fold increased, and blue down-pointing triangles ≥2-fold decreased, in the cryopreserved compared with fresh condition.

(A) Dot plots comparing the expression of genes between the two monkeys in the fresh (left) and cryopreserved (right) conditions, revealing a high level of correlation using Pearson’s r statistic (both p < 0.001).

(B) Dot plots comparing the expression of genes between the two fresh and cryopreserved conditions, using all detected genes (left) and the curated protein-coding genes (right) as described in the text. Expression levels were highly correlated using Pearson’s r statistic (both p < 0.001).

(C) Expression of the genes decreased in snRNA-seq versus scRNA-seq from human microglia (Thrupp et al., 2020), in the genes (left) and the curated protein-coding genes (right) as described in the text.

In the snRNA-seq versus scRNA-seq comparison (Thrupp et al., 2020), the genes that were underrepresented in nuclei were highly represented in lists of genes whose expressions were increased in AD. Two of these lists were from scRNA-seq AD model studies in mice, denoting activation response microglia (ARM) and disease-associated microglia (DAM) (Keren-Shaul et al., 2017; Sala Frigerio et al., 2019), and a third is from a snRNA-seq study of human AD (Mathys et al., 2019). This led to the conclusion that snRNA-seq was not appropriate for the study of microglia. We found that only a small proportion of the genes listed had their expression changed by cryopreservation, with only 3.7% of the ARM, 3.0% of DAM, and 1.9% of the AD genes showing increased expression in cryopreserved cells; none showed decreased expression (Figure 2F).
We next evaluated our data using other relevant microglial gene lists (Table S3). Of the altered genes in an snRNA-seq examination of a neuroinflammatory demyelinating disease, human multiple sclerosis (MS) (Schirmer et al., 2019), the cryopreserved cells had 3.1% increased, and none decreased, gene expression. Furthermore, in a mouse scRNA-seq study of response to a demyelinating injury (denoted injury response microglia) caused by lysolecithin injection into the brain (Hammond et al., 2019), 4.2% of the genes found to have change were increased, and 0.8% decreased, in our cryopreserved microglia relative to fresh cells (Figure 2G).

Inflammatory conditions were examined in scRNA-seq studies in mice, using microglia isolated 3 (acute) or 24 h (subacute) after an intraperitoneal injection of lipopolysaccharide (LPS) (Gerrits et al., 2020; Sousa et al., 2018). It was found that 2.6% and 3.0%, respectively, of the genes with increased expression were also increased in cryopreserved versus fresh microglia. None of the genes in the LPS gene lists showed decreased expression in the cryopreserved cells relative to the fresh cells in our study (Figure 2H).

As the cells were subject to cold, we also examined genes responsive to cold stress (using Gene Ontology GO:0009409, response to cold, Table S3). Although a relatively small number of genes (42) are in this set, five (12%) of the genes were increased in the cryopreserved microglia relative to the fresh cells; none were decreased. Interestingly, cold stress in mice has been found to activate the nuclear factor (NF)–κB signaling pathway in microglia (Xu et al., 2020). Therefore, we examined genes known to be involved in activation of this pathway (using Gene Ontology GO:0051092, positive regulation of NF-κB transcription factor activity, Table S3). Here 3.4% were upregulated, and 1.4% downregulated, in cryopreserved microglia versus fresh cells (Figure 2I).

Immediate-early genes are increased in cryopreserved microglia

Previous studies revealed that induction of immediate-early genes (IEGs) can occur due to tissue processing for scRNA-seq (van den Brink et al., 2017), with microglia among the most prominent IEG responders (Wu et al., 2017). IEGs are genes whose transcripts are rapidly expressed after cellular stimulation, without the need for new protein synthesis to control this expression (Herschman, 1991). Using a curated list of IEGs (Wu et al., 2017) (Table S3), we found a high level of change in expression of these genes, with 55 of 128 (43%) homologous to the IEG list showing a greater than 2-fold increase in the cryopreserved cells relative to the fresh cells and none exhibiting a decreased expression (Figure 2J). This finding is consistent with the elevations in acute phase response and AP1 pathways (Figures 2D and 2E), as IEGs precede activation of these pathways.

In order to examine the statistical significance of changes in gene expression, we used gene set enrichment analysis (GSEA) to determine if one or more of the gene sets is significantly enriched or depleted in expression in the cryopreserved versus fresh cells. The 11 gene sets were assessed using the complete expression data from all groups (Table 2). The gene sets are the same as those assessed in Figure 2. The size is the number of genes within each gene set for which expression was measured in the current experiment. The normalized enrichment score (NES) indicates the degree to which a gene set is overrepresented for differences in gene set size, normalized for correlations between gene sets and the expression dataset. The positive numbers connote enrichment in the cryopreserved microglia. The nominal (NOM) p value reflects the significance of single set, whereas the false discovery rate (FDR) q-value adjusts the significance value for gene set size and multiple hypotheses testing.

| Gene list                                | Size | NES  | NOM p-val | FDR q-val |
|------------------------------------------|------|------|-----------|-----------|
| Immediate-early genes                    | 128  | 1.984| 0.0       | 0.0       |
| GO: Response to cold                     | 42   | 1.188| 0.145     | 0.319     |
| Multiple sclerosis microglia             | 64   | 1.101| 0.243     | 0.702     |
| Down-regulated in microglia nuclei       | 213  | 1.029| 0.377     | 1.0       |
| GO: Positive regulation of NF-κB         | 147  | 0.956| 0.654     | 1.0       |
| Acute LPS microglia                      | 152  | 0.941| 0.708     | 1.0       |
| Activation response microglia            | 108  | 0.935| 0.679     | 1.0       |
| Injury response microglia                | 119  | 0.830| 0.923     | 1.0       |
| Disease-associated microglia             | 202  | 0.817| 0.966     | 1.0       |
| Alzheimer disease microglia              | 53   | 0.701| 0.966     | 1.0       |
| Subacute LPS microglia                   | 367  | 0.680| 1.0       | 0.992     |

The gene sets are the same as those assessed in Figure 2. The size is the number of genes within each gene set for which expression was measured in the current experiment. The normalized enrichment score (NES) indicates the degree to which a gene set is overrepresented for differences in gene set size, normalized for correlations between gene sets and the expression dataset. The positive numbers connote enrichment in the cryopreserved microglia. The nominal (NOM) p value reflects the significance of single set, whereas the false discovery rate (FDR) q-value adjusts the significance value for gene set size and multiple hypotheses testing.

We next evaluated our data using other relevant microglial gene lists (Table S3). Of the altered genes in an snRNA-seq examination of a neuroinflammatory demyelinating disease, human multiple sclerosis (MS) (Schirmer et al., 2019), the cryopreserved cells had 3.1% increased, and none decreased, gene expression. Furthermore, in a mouse scRNA-seq study of response to a demyelinating injury (denoted injury response microglia) caused by lysolecithin injection into the brain (Hammond et al., 2019), 4.2% of the genes found to have change were increased, and 0.8% decreased, in our cryopreserved microglia relative to fresh cells (Figure 2G).

Inflammatory conditions were examined in scRNA-seq studies in mice, using microglia isolated 3 (acute) or 24 h (subacute) after an intraperitoneal injection of lipopolysaccharide (LPS) (Gerrits et al., 2020; Sousa et al., 2018). It was found that 2.6% and 3.0%, respectively, of the genes with increased expression were also increased in cryopreserved versus fresh microglia. None of the genes in the LPS gene lists showed decreased expression in the cryopreserved cells relative to the fresh cells in our study (Figure 2H).

As the cells were subject to cold, we also examined genes responsive to cold stress (using Gene Ontology GO:0009409, response to cold, Table S3). Although a relatively small number of genes (42) are in this set, five (12%) of the genes were increased in the cryopreserved microglia relative to the fresh cells; none were decreased. Interestingly, cold stress in mice has been found to activate the nuclear factor (NF)–κB signaling pathway in microglia (Xu et al., 2020). Therefore, we examined genes known to be involved in activation of this pathway (using Gene Ontology GO:0051092, positive regulation of NF-κB transcription factor activity, Table S3). Here 3.4% were upregulated, and 1.4% downregulated, in cryopreserved microglia versus fresh cells (Figure 2I).

Immediate-early genes are increased in cryopreserved microglia

Previous studies revealed that induction of immediate-early genes (IEGs) can occur due to tissue processing for scRNA-seq (van den Brink et al., 2017), with microglia among the most prominent IEG responders (Wu et al., 2017). IEGs are genes whose transcripts are rapidly expressed after cellular stimulation, without the need for new protein synthesis to control this expression (Herschman, 1991). Using a curated list of IEGs (Wu et al., 2017) (Table S3), we found a high level of change in expression of these genes, with 55 of 128 (43%) homologous to the IEG list showing a greater than 2-fold increase in the cryopreserved cells relative to the fresh cells and none exhibiting a decreased expression (Figure 2J). This finding is consistent with the elevations in acute phase response and AP1 pathways (Figures 2D and 2E), as IEGs precede activation of these pathways.

In order to examine the statistical significance of changes in gene expression, we used gene set enrichment analysis (GSEA) to determine if one or more of the gene sets is significantly enriched or depleted in expression in the cryopreserved versus fresh cells. The 11 gene sets were assessed using the complete expression data from all groups (Table 2). The gene sets are the same as those assessed in Figure 2. The size is the number of genes within each gene set for which expression was measured in the current experiment. The normalized enrichment score (NES) indicates the degree to which a gene set is overrepresented for differences in gene set size, normalized for correlations between gene sets and the expression dataset. The positive numbers connote enrichment in the cryopreserved microglia. The nominal (NOM) p value reflects the significance of single set, whereas the false discovery rate (FDR) q-value adjusts the significance value for gene set size and multiple hypotheses testing.
list. Although all show some degree of enrichment in the cryopreserved cells, the IEGs have the highest level, and only the IEGs were significantly enriched in expression (Table 2).

Changes in IEGs can still be assessed in cryopreserved microglia

Despite an increase in expression of IEGs due to cryopreservation, they still may be responsive to changes in experimental conditions. To assess this possibility, we examined IEG expression in our prior study of the effect of drug abuse on a model of neuroHIV in monkeys, which utilized cryopreserved microglia for scRNA-seq (Niu et al., 2020). In fact, a number of the IEGs were altered between the saline and methamphetamine conditions, with some IEGs increased (e.g., IL1B and DUSP2), and others decreased (e.g., CCL3 and CCL5), in expression by methamphetamine compared with saline treatment of the animals (Figure 3). Thus, there is no ceiling effect or blanket inability to detect differences of IEGs in response to different physiological conditions.

Trajectory analysis reveals change in state in the cryopreserved microglia

To understand the cells’ responses to the cryopreservation and subsequent recovery in scRNA-seq experiments, we conducted trajectory inference followed by pseudotime analysis on the monkey microglia. Trajectory analysis enables investigation of changes as cells progress through a dynamic process, here from fresh cells to cryopreserved, examining the transition from one condition to the other. This is then displayed as a pseudotime graph, with values corresponding to progress along the change between conditions. As there were different numbers of cells in the samples, to avoid bias between conditions the cell number was down-sampled in each to the amount in the smallest of the four samples (4,660 cells). To facilitate the analysis, the identified DEGs (Table S1) were used. As shown in Figure 4 (left and middle), the cells begin the trajectory from the presumed zero pseudotime point on the right. The vast majority (95.5%) of the fresh cells, as well 34.5% of the cryopreserved cells, lie in the initial stage. The trajectory then diverges into two separate paths at “1” with an upward-facing group containing 2.2% of the fresh cells and 40.1% of the cryopreserved cells and a downward-facing group containing 3.3% of the fresh cells and 25.3% of the cryopreserved cells. An example of the expression pattern of DEG is shown for the IEGs KLF2 (Figure 4, right) as well as FOS, PER1, and CCL3 (Figure S2). No one gene appears to drive the transition between fresh and cryopreserved; rather, as in other stressors, many of the IEGs show increased expression.

Analysis of tissue macrophage scRNA-seq data reveals IEGs are also increased by cryopreservation

A recent study compared freshly isolated cells with cryopreserved cells under different conditions and concluded that cryopreservation in DMSO, as performed here, was optimal (Wohnhaas et al., 2019). As part of this study, immune cells isolated from the livers of rats subjected to a choline-deficient diet (experimental) were compared with those fed a control diet (two rats per condition) using scRNA-seq on the 10x Genomics platform, again as performed here. To assess whether the effects of cryopreservation on IEGs are specific to microglia or present in other cell types, we downloaded the data from GEO (Series GSE127248) and filtered for the macrophages using graph-based clustering and marker genes (Figure S3). Combining the lists of differentially expressed genes found in either the experimental or control conditions between
fresh and DMSO cryopreserved cells revealed a low proportion of greater than 2-fold changes, 1.34% in the control condition and 0.91% in the experimental. Interestingly, between the two conditions in the rat macrophage experiment, only 16 genes were in common regarding increased expression due to cryopreservation. Thirteen (81%) of these are IEGs. These include the expression of the IEGs CCL3/Ccl3 (3.49-fold in monkey microglia, 4.59-fold and 3.89-fold in control and experimental rat liver macrophages, respectively) and FOS/Fos (6.51-, 10.45-, and 30.09-fold) (Figure 5A). Examining the list of curated IEGs, 16.2% of the genes showed increased expression in the cryopreserved cells from rats fed the control diet, whereas 13.0% were increased in those fed the experimental diet (Figure 5B). Trajectory analysis revealed that the rat liver macrophages also showed a progression from the fresh to cryopreserved cells, albeit with a more complex branching pattern (Figure S4).

**DISCUSSION**

The analysis of microglia by single-cell technologies has led to an increased understanding of various types of microglia in development, homeostasis, and changes due to aging and disease. Such studies have yielded important clues into the function of microglia in these processes. Single-cell transcriptomic studies from human specimens are often done with snRNA-seq; however expression of disease-related genes can be lost (Thrupp et al., 2020). Here we find that isolation and cryopreservation of microglia can be performed, and the cells can then be archived for later purification and use in scRNA-seq, with minimal changes in expression of disease-related genes. In both our study and a prior study on macrophages in rats (Wohnhaas et al., 2019), an increase in expression of IEGs was found in cryopreserved cells. This may be, in part, due to the freezing and/or recovery process used in cryopreservation, as previous studies have noted that IEGs are upregulated during tissue processing for scRNA-seq, especially notable in microglia (Wu et al., 2017).

IEGs were initially characterized as rapidly transcribed genes induced after growth factor stimulation of cells (Lau and Nathans, 1987). They are also induced in response to cellular differentiation as well as inflammation, neuronal activity, and stress (Bahrami and Drablos, 2016). Although we did not see enrichment of expression of genes associated with the response to cold, the IEG induction is likely due to cellular stress in the freezing and/or recovery process and has been noted by others in cryopreserved cells (Ruf-Zamojski et al., 2018).

Although IEGs are altered in the cryopreserved cells, this does not mean that these genes are not informative in studies. First, despite all precautions, isolation procedures are prone to lead to a level of IEG activation. Differences in IEG expression using cryopreserved microglia in scRNA-seq can still be detected between different in vivo conditions (Figure 3). Furthermore, when considering the genes whose expressions were suppressed in snRNA-seq relative to scRNA-seq (Thrupp et al., 2020), many of these were still significantly altered in studies on AD and MS (Mathys et al., 2019; Schirmer et al., 2019) (see Table S3). It is not clear which method best represents the status in vivo, as all have caveats. If one is not comparing results obtained by the same method, means of accounting for the known differences should be used.
Archiving of isolated microglia can lead to their productive use in subsequent studies without the loss of disease-related information. In addition, such cells can be used in a variety of additional studies, including other emerging single-cell technologies or long-established ones such as flow cytometry, in addition to nucleic acid, protein, or small molecule analyses. Furthermore, as cells are viable when recovered, cell culture experiments are also made possible. Cryopreserving cells will also allow the choice of specimens for future studies, instead of having to process for scRNA-seq before one knows the true utility of the specimen, such as not knowing the true diagnosis pending neuropathological or other evaluation. In addition to microglia, we have also reported the successful use of cryopreserved matured monocytes using scRNA-seq (Leon-Rivera et al., 2020). Other groups have reported the isolation and use of microglia from fresh autopsy human brain tissue from brain banks, in particular for studies on AD, and many have utilized microglia from fetal

Figure 5. Expression of selected IEGs in monkey microglia and rat macrophages
(A) Overlaid dot and violin plots indicate expression levels of the shown genes in cells from the current monkey microglia experiment (left) and the rat liver macrophages (control diet, center, and experimental diet, right), revealing increased expression in the cryopreserved compared with fresh condition.
(B) Dot plots of IEGs expression in rat liver macrophages in the control (left) and experimental diet (right) conditions; symbols as in Figure 2. Rat liver macrophage data from (Wohnhaas et al., 2019).
tissue as well as surgical resections for intractable epilepsy (Lue et al., 2019). We show here that cryopreserved microglia can be utilized for scRNA-seq in addition to other known uses.

Knowledge of the role of microglia in CNS function and dysfunction continues to grow, and single-cell technologies have contributed to this growth. In addition, techniques to perform single-cell procedure and analyses continue to expand. Cryopreservation will enable valuable specimens to be used for current and future studies to elucidate the role of microglia in CNS function and dysfunction.

Limitations of this study
Limitations include the use of only two biological replicates, and the examination on rhesus monkey microglia, as opposed to specimens from humans or other animal species. Furthermore, we only examined SIV-infected animals under suppressive antiretroviral treatment, and not those with other diseases or control animals. We do note the replicates were closely correlated, and that, among the different species examined by scRNA-seq, rhesus monkey microglia were found to show the highest similarity to human microglia (Geirsdottir et al., 2019). It has been found that performing tissue disruption and cellular isolation procedures in the cold was found to reduce or eliminate IEG upregulation (Adam et al., 2017; Wu et al., 2017). However, others have found that performing brain dissociation in the cold results in poor yield and lower viability and that the cells are prone to clump (Bordt et al., 2020). The reason for the decreased number of cryopreserved, relative to fresh, cells captured in the processing for scRNA-seq is not known, but may represent increased fragility of the cryopreserved cells. Cellular damage in cryopreserved cells used for scRNA-seq has been previously reported (Guillaumet-Adkins et al., 2017). Whether cryopreservation may introduce a bias in the cells analyzed, as with any isolation and purification procedure, remains to be determined, and future studies are needed to assess such a potential bias. We have successfully performed scRNA-seq from microglia samples that have been cryopreserved for over 7 years (Niu et al., 2020). Future studies can also address any limits on storage time on cryopreserved microglia, and changes that may occur due to different storage times. It would also be of interest if a similar study on brains frozen for different times was performed. Finally, direct comparison of different procedures (snRNA-seq, scRNA-seq from fresh and cryopreserved cells) can assist in comparing the methods and enabling meta-analyses to be done from studies performed with the different platforms.

Resource availability
Lead contact
Howard Fox, University of Nebraska Medical Center, hfox@unmc.edu.

Materials availability
This study did not generate new unique reagents.

Data and code availability
The scRNA-seq data from this study has been deposited in the NCBI GEO database, accession # GSE162663.

METHODS
All methods can be found in the accompanying transparent methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.102357.

ACKNOWLEDGMENTS
These studies were supported by NIH grants R01AI24965 (to C.V.F.), R01DA043164, and P30MH062261 (to H.S.F.) and a Nebraska Research Initiative Collaborative Grant (S.R.D.). We thank the DNA Sequencing (DNAS), Flow Cytometry Research (FCR), and Bioinformatics and Systems Biology (BSB) Cores at UNMC for assistance. They receive partial support from the NIH National Institute for General Medical Science IN-BRE P20GM103427 (DNAS, BSB), and COBRE P30GM110768 (DNAS) grants, NIH National Cancer Institute Cancer Support Grant P30CA036727 (FCR, BSB), and the Nebraska Research Initiative (FCR). Major instrumentation has been provided by the UNMC Office of the Vice Chancellor for Research, the University of
Nebraska Foundation, the Nebraska Banker’s Fund, and by the NIH NCRR/NIGMS Shared Instrument Program. We thank Dr. Jeffrey Lifson and staff of the Quantitative Molecular Diagnostics Core of the AIDS and Cancer Virus Program, Frederick National Laboratory, supported in part by contract HHSN261200800001E, for SIV viral load analyses. We thank Kelly Goff and Dr. Preston A. Marx of the Virus Characterization, Isolation and Production Core at Tulane National Primate Research Center, Covington, LA, supported in part by PS1RR00164 from the Office of Research Infrastructure Programs of the National Institutes of Health. We thank Robin Taylor for making the graphical abstract, which was created with BioRender.com. Finally, we thank Gilead Biosciences, USA, for the kind gift of emtricitabine and tenofovir alafenamide.

AUTHOR CONTRIBUTIONS
Conceptualization, B.M. and H.S.F.; data curation, N.M., B.G.L., and B.M.; formal analysis, B.M., N.M., and H.F.; funding acquisition, C.V.F. and H.S.F.; investigation, B.M., S.R.D., B.G.L., K.E., and A.F.; methodology, B.G.L., A.F., B.M., M.N., and H.S.F.; project administration, B.M. and B.G.L.; resources, S.R.D., C.V.F., and H.S.F.; supervision, C.V.F., and H.S.F.; writing – original draft, B.M., M.N., and H.S.F.; writing – review and editing, B.M., M.N., S.R.D., C.V.F., B.G.L., K.E., A.F., and H.S.F.

DECLARATION OF INTERESTS
The authors declare no competing interests.

REFERENCES
Adam, M., Potter, A.S., and Potter, S.S. (2017). Psychrophilic proteases dramatically reduce single-cell RNA-seq artifacts: a molecular atlas of kidney development. Development 144, 3625–3632.
Alles, J., Karakios, N., Praktikno, S.D., Grosswendt, S., Wahrle, P., Ruffault, P.L., Ayoub, S., Schreyer, L., Boltengagen, A., Birchmeier, C., et al. (2017). Cell fixation and preservation for droplet-based single-cell transcriptomics. BMC Biol. 15, 44.
Attar, M., Sharma, E., Li, S., Bryer, C., Cubitt, L., Broxholme, J., Lockstone, H., Kinchen, J., Simmons, A., Piazza, P., et al. (2018). A practical solution for preserving single cells for RNA sequencing. Sci. Rep. 8, 2151.
Bahrami, S., and Drablos, F. (2016). Gene regulation in the immediate-early response process. Adv. Biol. Regul. 62, 37–49.
Bonham, L.W., Sinko, D.W., and Yokoyama, J.S. (2019). The transcriptional landscape of microglial genes in aging and neurodegenerative disease. Front. Immunol. 10, 1170.
Bordt, E.A., Block, C.L., Petrozzioello, T., Sadri-Vakili, G., Smith, C.J., Edlow, A.G., and Bilbo, S.D. (2020). Isolation of microglia from human tissue. STAR Protoc. 1, 100003.
Bortell, N., Basova, L., Najera, J.A., Morsey, B., Fox, H.S., and Marcondes, M.C.G. (2018). Sirtuin 1-chromatin-binding dynamics points to a common mechanism regulating inflammatory targets in SIV infection and in the aging brain. J. Neuroimmunome Pharmacol. 13, 163–178.
Chaudhuri, A.D., Yelamanchili, S.V., Marcondes, M.C., and Fox, H.S. (2013). Up-regulation of microRNA-142 in simian immunodeficiency virus encephalitis leads to repression of sirtuin1. FASEB J. 27, 3720–3729.
Chen, J., Cheung, F., Shi, R., Zhou, H., Lu, W., and Consortium, C.H.I. (2018). PBMC fixation and processing for Chromium single-cell RNA sequencing. J. Transl. Med. 16, 198.
Del-Aguila, J.L., Li, Z., Dubé, U., Mihindukulasuriya, K.A., Budde, J.P., Fernandez, M.V., Ibanez, L., Bradley, J., Wang, F., Bergmann, K., et al. (2019). A single-nucleus RNA sequencing study of Mendelian and sporadic AD in the human brain. Alzheimers Res. Ther. 11, 71.
Donlin, L.T., Rao, D.A., Wei, K., Slowkowski, K., McGeeachy, M.J., Turner, J.D., Meednu, N., Mizoguchi, F., Gutierrez-Arcelus, M., Lieb, D.J., et al. (2018). Methods for high-dimensional analysis of cells dissociated from cryopreserved synovial tissue. Arthritis Res. Ther. 20, 139.
Geirsdottir, L., David, E., Keren-Shaul, H., Weiner, A., Bohlen, S.C., Neuber, J., Balic, A., Giladi, A., Sheban, F., Dutertre, C.A., et al. (2019). Cross-species single-cell analysis reveals divergence of the primate microglia program. Cell 179, 1609–1622 e1616.
Gerrits, E., Heng, Y., Boddeke, E., and Eggen, B.J.L. (2020). Transcriptional profiling of microglia, current state of the art and future perspectives. Glia 68, 740–755.
Gierz, T.M., Wadsworth, M.H., 2nd, Hughes, T.K., Bryson, B.D., Butler, A., Satija, R., Fortune, S., Love, J.C., and Shalek, A.K. (2017). Seq-Well: portable, low-cost RNA sequencing of single cells at high throughput. Nat. Methods 14, 395–398.
Goldmann, T., Wieghofer, P., Jordao, M.J., Prutek, F., Hagemeyer, N., Frenzel, K., Amann, L., Staszewski, O., Kerdro, K., Krueger, M., et al. (2016). Origin, fate and dynamics of macrophages at central nervous system interfaces. Nat. Immunol. 17, 797–805.
Grubman, A., Chew, G., Ouyang, J.F., Sun, G., Choo, X.Y., McLean, C., Simmons, R.K., Buckbery, S., Vargas-Landin, D.B., Poppe, D., et al. (2019). A single-cell atlas of entorhinal cortex from individuals with Alzheimer’s disease reveals cell-type-specific gene expression regulation. Nat. Neurosci. 22, 2087–2097.
Guillaumet-Adkins, A., Rodriguez-Esteban, G., Mercu, E., Mendez-Lago, M., Jaitin, D.A., Villanueva, A., Vidal, A., Martinez-Marti, A., Felip, E., Vivancos, A., et al. (2017). Single-cell transcriptome conservation in cryopreserved cells and tissues. Genome Biol. 18, 45.
Hammond, T.R., Dufort, C., Dissing-Olesen, L., Giera, S., Young, A., Wysocker, A., Walker, A.J., Gergits, F., Sega, M., Nemesh, J., et al. (2019). Single-cell RNA sequencing of microglia throughout the mouse lifespan and in the injured brain reveals complex cell-state changes. Immunity 50, 253–271 e256.
Hattori, M., Tugores, A., Westwick, J.K., Veloz, L., Leffert, H.L., Karin, M., and Brenner, D.A. (1993). Activation of activating protein 1 during hepatic acute phase response. Am. J. Physiol. 264, G95–G103.
Herschman, H.R. (1991). Primary response genes induced by growth factors and tumor promoters. Annu. Rev. Biochem. 60, 281–319.
Jordao, M.J.C., Sankowski, R., Brendbeck, S.M., Sagar, Locatelli, G., Tai, Y.H., Tay, T.L., Schramm,
Alzheimer's disease. Cell. 2017.
Matcovitch-Natan, O., Dvir-Szternfeld, R., Ulland, T.K., David, E., Baruch, K., Lara-Astasio, D., Toth, B., et al. (2017). A unique microglia type associated with restricting development of alzheimer's disease. Cell. 769, 1276–1290 e1217.
Lau, L.F., and Nathans, D. (1987). Expression of a set of growth-related immediate early genes in BALB/c 3T3 cells: coordinate regulation with c-fos or c-myc. Proc. Natl. Acad. Sci. U S A 84, 1182–1186.
Lawson, L.J., Perry, V.H., and Gordon, S. (1992). Turnover of resident microglia in the normal adult mouse brain. Neuroscience 48, 405–415.
Leon-Rivera, R., Morsey, B., Niu, M., Fox, H.S., and Berman, J.W. (2020). Interactions of monocyties, HIV, and ART identified by an innovative scRNAseq pipeline: pathways to reservoirs and HIV-associated comorbidities. mBio 11, e01037–20.
Li, Q., Cheng, Z., Zhou, L., Darmanis, S., Neff, N.F., Okamoto, J., Gulati, G., Bennett, M.L., Sun, L.O., Clarke, L.E., et al. (2019). Developmental heterogeneity of microglia and brain myeloid cells revealed by single-cell RNA sequencing. Neuron 101, 207–223 e210.
Lue, L.F., Beach, T.G., and Walker, D.G. (2019). Alzheimer’s disease Research using human microglia. Cells 8, 838.
Marcondes, M.C., Burudi, E.M., Huhtonen-Resendiz, S., Sanchez-Alavez, M., Watry, D., Zandonatti, M., Henrikse, S.J., and Fox, H.S. (2001). Highly activated CD8(+) T cells in the brain correlate with early central nervous system dysfunction in simian immunodeficiency virus infection. J. Immunol. 167, 5429–5438.
Masuda, T., Sankowski, R., Staszewski, O., Bottcher, C., Amann, L., Sagar, Scheewe, C., Nessler, S., Kunz, P., van Loo, G., et al. (2019). Spatial and temporal heterogeneity of mouse and human microglia at single-cell resolution. Nature 566, 398–392.
Methys, H., Davila-Veldemar, J., Peng, Z., Gao, F., Mohammadi, S., Young, J.Z., Menon, M., He, L., Abdurrob, F., Jiang, X., et al. (2019). Single-cell transcriptomic analysis of Alzheimer’s disease. Nature 570, 332–337.
Niu, M., Morsey, B., Lamberty, B.G., Emanuel, K., Yu, F., Leon-Rivera, B., Raman, J.W., Gaskill, P.J., Matt, S.M., Ciborowski, P.S., et al. (2020). Methamphetamine increases the proportion of SIV-infected microglia/macrophages, alters metabolic pathways, and elevates cell death pathways: a single-cell analysis. Viruses 12, 1297.
Olah, M., Patrick, E., Villani, A.C., Xu, J., White, C.C., Ryan, K.J., Piehowski, P., Kapasi, A., Nejad, P., Cinpean, M., et al. (2018). A transcriptomic atlas of aged human microglia. Nat. Commun. 9, 539.
Ruf-Zamojski, F., Ge, Y., Nair, V., Zamojski, M., Pincas, H., Toufaily, C., Tome-Garcia, J., Stocekius, M., Stephenson, W., Smith, G.R., et al. (2018). Single-cell stabilization method identifies gonadotrope transcriptional dynamics and pituitary cell type heterogeneity. Nucleic Acids Res. 46, 11370–11380.
Sala Frigerio, C., Wolfs, L., Fattorelli, N., Thrupp, N., Voyerlyuk, I., Schmidt, I., Mancuso, R., Chen, W.T., Woodbury, M.E., Sivastava, G., et al. (2019). The major risk factors for Alzheimer’s disease: age, sex, and genes modulate the microglia response to abeta plaques. Cell Rep. 27, 1293–1306 e1296.
Schirmer, L., Velmeshev, D., Holmquist, S., Kaufmann, M., Werneburg, S., Jung, D., Vistnes, S., Stockley, J.H., Young, A., Stendel, M., et al. (2019). Neuronal vulnerability and multilineage diversity in multiple sclerosis. Nature 573, 75–82.
Sousa, C., Golebiewska, A., Povovathilingal, S.K., Kaoma, T., Pires-Afonso, Y., Martins, S., Coowar, D., Azuaje, F., Skupin, A., Balling, R., et al. (2018). Single-cell transcriptomics reveals distinct inflammation-induced microglia signatures. EMBO Rep. 19, e46171.
Thom森, E.R., Mich, J.K., Yao, Z., Hodge, R.D., Doyle, A.M., Jang, S., Shehata, S.I., Nelson, A.M., Shapovalova, N.V., Levi, B.P., et al. (2016). Fixed single-cell transcriptomic characterization of human radial glial diversity. Nat. Methods 13, 87–93.
Thrupp, N., Sala Frigerio, C., Wolfs, L., Skene, N.G., Fattorelli, N., Povovathilingal, S., Fourne, Y., Matthews, P.M., Theys, T., Mancuso, R., et al. (2020). Single-nucleus RNA-seq is not suitable for detection of microglial activation genes in humans. Cell Rep. 32, 108189.
van den Brink, S.C., Sage, F., Vertesy, A., Spanjaard, B., Peterson-Maduro, J., Baron, C.S., Robin, C., and van Oudenaarden, A. (2017). Single-cell sequencing reveals dissociation-induced gene expression in tissue subpopulations. Nat. Methods 14, 935–936.
van der Poel, M., Ulas, T., Mizee, M.R., Hisao, C.C., Miedema, S.S.M., Adelia, Schuurman, K.G., Helder, B., Tas, S.W., Schultz, J.L., et al. (2019). Transcriptional profiling of human microglia reveals grey-white matter heterogeneity and multiple sclerosis-associated changes. Nat. Commun. 10, 1139.
Watry, D., Lane, T.E., Streb, M., and Fox, H.S. (1995). Transfer of neuropathogenic simian immunodeficiency virus with naturally infected microglia. Am. J. Pathol. 146, 914–923.
Wohnhaas, C.T., Leparic, G.G., Fernandez-Albert, F., Kind, D., Gantner, F., Vollet, C., Hildebrandt, T., and Baum, P. (2019). DMSO cryopreservation is the method of choice to preserve cells for droplet-based single-cell RNA sequencing. Sci. Rep. 9, 10699.
Wu, Y.E., Pan, L., Zuo, Y., Li, X., and Hong, W. (2017). Detecting activated cell populations using single-cell RNA-seq. Neuron 96, 313–329 e316.
Xu, B., Lang, L.M., Lian, S., Guo, J.R., Wang, J.F., Liu, J., Yang, H.M., and Li, S.Z. (2020). Neurinflammation induced by secretion of acetylated HMGB1 from activated microglia in hippocampus of mice following chronic cold exposure. Brain Res. 1726, 146495.
Zhou, Y., Song, W.M., Andheya, P.S., Swain, A., Levy, T., Miller, K.R., Poliani, P.L., Cominelli, M., Grover, S., Gillillan, S., et al. (2020). Human and mouse single-nucleus transcriptomics reveal TREM2-dependent and TREM2-independent cellular responses in Alzheimer’s disease. Nat. Med. 26, 131–142.
Supplemental information

Cryopreservation of microglia enables single-cell RNA sequencing with minimal effects on disease-related gene expression patterns

Brenda Morsey, Meng Niu, Shetty Ravi Dyavar, Courtney V. Fletcher, Benjamin G. Lamberty, Katy Emanuel, Anna Fangmeier, and Howard S. Fox
Figure S1. Quality assurance and quality control (QA/QC) of monkey microglia (Related to Table 1 and Results). QA/QC was performed using overlaid dot and violin plots (with the Y-axis in log-scale) of the indicated measures. (A) Counts were filtered to include cells with between 800-15,000 total counts (left), Detected Features to include cells with between 300-5,000 genes detected (middle), and the percent of counts mapping to the Mitochondrial genome filtered to cells with 15% or less (right). (B) To remove non-myeloid cells, graph-based clustering was performed followed by projection in three dimensions using UMAP. Cluster 6 (double-headed arrow) was enriched in cells expressing genes found in lymphocytes, specifically cytotoxic T cells and/or natural killer cells, and cells in this cluster were excluded from further analysis. (C) Dot plots showing expression of microglia marker genes from the 23,637 cells remaining cells after elimination of cluster 6. The number of cells expressing both indicated markers is indicated in the upper right, those with only one of the markers at the upper left and right (along axes), and neither marker lower left (near origin).
Figure S2. IEG expression along trajectory analysis (Related to Figure 4). Trajectory analysis of the fresh to cryopreserved state change, with cells color-coded by expression of the indicated IEG.
Figure S3. Identification of macrophages in the rat liver immune cell experiment (Related to Figure 5). (A) Graph-based clustering revealed groups of cells with marker genes enriched in the clusters, enabling identification of cell types from the liver immune cell experiment. Figure shows results projected in three dimensions using UMAP. B cells were identified by the expression Cd19, Cd79a, and Cd79b; Macrophages by Cd14, Spi1, and Csf1r; Mito High are cells with high expression of genes encoded by the mitochondrial genome; Neutrophils by Mmp8 and Ccl6; NK cells by Nkg7; plasmacytoid Dendritic Cells (pDC) by Siglech; and T cells by Cd3d, Cd3e, and Cd3g. (B) Cells in the clusters with Macrophage markers (red, other cell types and colors dimmed) were selected for further analysis.
Figure S4. Trajectory analysis of fresh and cryopreserved rat liver macrophages (Related to Figures 5 and Results). Trajectory analysis for the two conditions of rat macrophage analyzed by scRNA-seq. The transition from fresh to the cryopreserved condition is indicated, as on the left, with pseudotime indicated on the right.
Isolation of total brain microglia/macrophages

Microglia/macrophages-enriched brain isolation was performed using a modification of our previously described procedure (Marcondes et al., 2001). The brain was sectioned and meninges were removed in a 150 mm dish on ice with 4°C Hank’s balanced salt solution (HBSS; Invitrogen, Carlsbad, CA). The remaining roughly minced tissue was homogenized with a Dounce homogenizer using additional 4°C HBSS as necessary. Homogenate was washed twice with 4°C HBSS and centrifuged at 4°C for 7 minutes at 550 x g. The brain tissue was then digested at in 37°C in HBSS containing 28 U/ml DNase I and 8 U/ml papain (approximately 2.5 U/gram of tissue) on a nutating mixing platform at 37°C for 30 minutes (Sigma, St. Louis, MO). While digesting, the enzymes were inactivated by addition of 2.5% FBS (volume/volume). Digested tissue was centrifuged then resuspended in 25 mL of 4°C HBSS. The cell suspension was mixed with 7 mL of 90% Percoll (GE HealthCare, Pittsburg, PA) and centrifuged at 4°C for 15 minutes at 550 x g, with the acceleration set to 5 and the brake set to 1. The resulting fatty upper layer and HBSS were removed in a 150 mm dish on ice with 4°C Hank’s balanced salt solution (HBSS; Invitrogen, Carlsbad, CA). The remaining tissue was gently washed twice with 4°C HBSS and centrifuged at 550 x g, with the acceleration set to 5 and the brake set to 1.

Animals

Two male, five-year-old Indian origin rhesus macaques were purchased from New Iberia Research Center (New Iberia, LA). The monkeys tested negative for the indicated viral pathogens: SIV, SRV, STLV-1, B-virus, and measles; and bacterial pathogens: salmonella, shigella, campylobacter, yersinia, and vibrio. Macaques were housed in compliance with the Animal Welfare Act and the Guide for the Care and Use of Laboratory Animals in the nonhuman primate facilities at Department of Comparative Medicine, University of Nebraska Medical Center (UNMC). The primate facility at UNMC has been accredited by the AAALAC international. This study was reviewed and approved by the UNMC Institutional Animal Care and Use Committee under protocol 16-001-02-FC titled “Assessment of Antiretroviral Pharmacology in Lymphoid Tissues using the SIV macaque model” approved 2/25/2016. Animals were maintained in a temperature-controlled (23 ± 2°C) indoor climate with 12-hour light/dark cycle. They were fed Teklad Global 25% protein primate diet (#2055, Envigo) supplemented with fresh fruit or vegetables, and water ad libitum. The monkeys were observed twice daily for health status by the animal care and veterinary personnel.

At the start of study, both macaques were intravenously inoculated with 200 μl of a SIVmac251 stock, which contained 2.48 x 10⁹ copies/ml of SIV RNA. Virus stocks were provided by the Virus Characterization, Isolation and Production Core at Tulane National Primate Research Center (Covington, LA) as described previously (Del Prete et al., 2013). Five-weeks post-inoculation, combined antiretroviral therapy (cART) was initiated. One animal received cART consisting of two drugs: 4 mg/ml dolutegravir (DTG), 40 mg/ml emtricitabine (FTC) in a vehicle of 15% Kleptose and 0.1 N NaOH at a pH of 7.4, while the other was treated with four drug cART, consisting of 4 mg/ml DTG, 1mg/ml tenofovir alafenamide fumarate (TAF), 40 mg/ml FTC, 12 mg/ml maraviroc (MVC) in a vehicle of 15% Kleptose and 0.1 N NaOH at a pH of 7.4. Both animals were treated once daily with a subcutaneous injection of 1 ml cART/kg body weight.

Animals remained on the cART drug regimen for six months following suppression of virus in the plasma (viral load of <50 SIV copies/ml plasma) and were sacrificed as per experimental design for the above-named approved study (for animal 85T a total of 302 days of SIV infection, 277 days of cART, and documented suppression for 181 days, for 86T seven additional days for each measure). Plasma viral load was determined from EDTA-anticoagulated plasma separated from blood by centrifugation. Plasma was stored at -80°C until assessed for viral load. SIV RNA viral load was measured using a PCR-based assay by the Quantitative Molecular Diagnostics Core at the National Cancer Institute (Frederick, MD) essentially as described previously (Li et al., 2016). At necropsy, deeply anesthetized animals were perfused intracardially with sterile PBS containing 1 U/ml heparin to clear blood including blood-borne cells from the brain. Brains were harvested for histopathology and molecular studies, and approximately half of the brain then taken for single cell microglia/macrophages isolations and further experimental studies.
both hemocytometer and Coulter Counter Z1.

The brain microglia/macrophages isolates were either used for scRNA-seq as described below, or cryopreserved for later use. To cryopreserve, cells were centrifuged at 4°C at 550 x g for 5 minutes and supernatant was removed. The pellet was dissociated by tapping and then resuspended by the dropwise addition of a solution of 4°C 10% DMSO in FBS at a concentration of 10 million cells per milliliter. Cells were transferred to cryopreservation tubes and placed in a Mr. Frosty for slow controlled freezing at -80°C. After 24 hours, cryotubes were transferred to liquid nitrogen for long-term storage.

**scRNAseq preparation of fresh brain isolates**

All procedures were completed using RNAase-free tubes and filter tips. Brain microglia/macrophages were washed in PBS, and stained with UV-blue live/dead assay for 30 minutes at 4°C. Cells were centrifuged at 300 x g for 8 minutes at 4°C and resuspended in e-bioscience flow cytometry staining buffer for 5 minutes at 4°C. Cells were again centrifuged at 300 x g for 8 minutes at 4°C and resuspended in MACS separation buffer with 0.1% BSA (Miltenyi, Gladbach, Germany). Cells were counted and volume was adjusted for staining with non-human primate CD11b microbeads (Miltenyi). Twenty million cells were reconstituted in 80 μL of MACS buffer and reacted with 40 μL of CD11b microbeads at 4°C for 15 minutes. After incubation, cells were washed with MACS buffer with 0.1% BSA, centrifuged at 300 x g for 8 minutes at 4°C, and the resulting pellet was resuspended in 1 ml of MACS buffer and loaded onto two MACS Separator LS columns. The positive CD11b cells were collected and isolated were counted on Coulter Counter Z1 and assessed for viability and concentration using a hemocytometer and trypan blue exclusion. CD11b-enriched isolates were then stained with Brilliant Violet 605-labeled anti-mouse/human CD11b antibody clone M1/70 (Biolegend, San Diego, CA) for 45 minutes at 4°C. Cells were washed with e-bioscience flow cytometry staining buffer and cell concentration was set to 5 million cells per ml for fluorescence activated cell sorting (FACS). Machine standards were set with unstained and single stained cells. Live/dead standards were also set using the Amine reactive compensation Bead kit (Invitrogen, Carlsbad, CA). Cells were sorted based on size, singlets, live, CD11b+ events using an Aria2 flow cytometer (BD Biosciences). Sorted cells were adjusted to a concentration of 1,000 cells per µl in DMEM/F12 with 10% FBS. FACS files were analyzed using FlowJo software (FlowJo, Ashland, OR).

**scRNAseq preparation of cryopreserved brain isolates**

Samples of cryopreserved enriched brain microglia/macrophages isolates, stored in liquid nitrogen described above, were rapidly thawed in a 37°C water-bath. Once thawed, 1 ml of media consisting of DMEM/F12 with 2.5mM L-glutamine and 15mM HEPES with 10% FBS, supplemented with 1% DNase (Sigma Aldrich) was slowly added to cells using filter tips. Cells were incubated at room temperature for 10 minutes, gently mixed at 5 minutes using a pipet. After 10 minutes, the cells were transferred dropwise into a 15 ml tube and 10 ml of the above media with 1% DNase containing media. Cells were incubated at room temperature for 5 minutes and centrifuged at 25°C 300 x g for 8 minutes and resuspended in 1 ml of DMEM/F12 with 20% FBS, and incubated at 37°C in a 5% CO₂ incubator for 15 minutes to enable recovery. Media was removed and cells were washed and counted using a Coulter Counter for use. Once cell concentration was known, cells were transferred to ice-cold PBS, and sequentially stained with the Live/Dead Assay, enriched immunomagnetically by positive selection by MACS using CD11b magnetic beads, and then stained with CD11b-BV605 and processed for FACS purification, all as described above for the fresh brain microglia/macrophage isolates.

**scRNAseq capture and library preparation**

Post-sort, fresh and cryopreserved isolates were concentrated to approximately 1,000 cells per µl, assessed by trypan blue for viability and concentration. Based on 10x Genomics parameters targeting 8,000 cells, the ideal volume of cells was loaded onto the 10x Genomics (Pleasanton, CA) Chromium GEM Chip and placed into Chromium Controller for cell capturing and library preparation. We used the 10x Genomics Single Cell 3’ GEM, Library, & Gel bead kit v3. Briefly, the microfluidics combines the Single Cell 3’ Gel Beads containing unique barcoded primers with a unique molecular identifier (UMI), followed by lysis of cells and barcoded reverse transcription of RNA, amplification of barcoded cDNA, fragmentation of cDNA (with peak consistently in the range of 470-500 bp), 5’ adapter attachment and sample indexing as the manufacturer instructed with version 3 reagent kits. The prepared libraries were then sequenced using an Illumina Nextseq550 sequencer with the NSQ 500 hi- Output KT v2 (150 CYS)
Bioinformatic Analysis
Rhesus monkey samples

Bioinformatic analysis was performed basically as described in Niu et al., 2020. Demultiplexing and filtered feature barcode matrices generation was performed by Cell Ranger (version 3.1, 10x Genomics) pipeline. The raw base call (BCL) files were converted into FASTQ files, and the reads aligned to custom combined genomes of Macaca mulatta (Mmul 10) and the SIV genome (using NCBI reference sequence M33262.1, note no SIV sequences were identified). Cell Ranger also performed filtering, barcode counting, UMI counting, and sample aggregation. As a result, a gene expression matrix was generated containing the raw UMI counts for each cell for each sample, along with a matrix consisting of matrices of all samples.

Partek (St. Louis, MO, USA) Flow version 9.0 was used for in-depth analysis, with the gene expression matrices for each sample as the input. Single cell QA/QC was performed to filter out low-quality cells based on the Total UMI count (<800 or >15,000), Detected Gene count (<300 or >5,000), and Mitochondrial UMI proportion (>15%), resulting in the removal of 784 cells, leaving 23,792. This was followed by a noise reduction step, in which the genes that had zero expression in all cells were removed from further analysis. Normalization was performed by converting the UMI to CPM (counts per million reads), then adding 1 and converting to the log2 value.

Initial principle component analysis (PCA) was performed with a setting of 50 PCs for calculation. Graph-based clustering was performed using the first 15 PCs from the PCA (according to the PC scree plot), and Louvain as the clustering algorithm. Data visualization was performed using Uniform Manifold Approximation and Projection (UMAP). A cluster of contaminating lymphocytes was identified as described in the text, and those cells were excluded from further examination. Differentially expressed genes (DEGs) analysis to assess differences between the fresh and cryopreserved cells was performed using Gene Specific Analysis (GSA) in Partek Flow software. GSA functions to select the best statistic model for each gene’s differential expression from its pool to produce p-values. The five response distributions in the pool are Normal, Lognormal, Lognormal with shrinkage, Negative Binomial, and Poisson distributions. Trajectory analysis and pseudotime calculation was performed (also using Partek Flow, using Monocle 2 (Qiu et al., 2017)) on the set of 538 DEGs, chosen with a fold-change of > |1.5| and a false discovery rate of < 0.05 (Table S1).

Additional bioinformatic analyses were performed using human gene IDs of the rhesus monkey orthologs. Gene Set Enrichment Analysis (GSEA) (Subramanian et al., 2005) was performed on the expression data (using GSEA (version 4.1.0)). For the analysis, 1000 permutations were used to calculate the p values with permutation type set to phenotype. To prepare the expression data, the cellranger aggr function of Cell Ranger was used to aggregate the matrices of all samples, followed by Loupe Browser (version 4.2.0, 10x Genomics) to calculate the average UMI for each gene per sample considering the size factor. The source of the gene sets are referenced within the text, the Gene Ontology (GO) lists were obtained from the Molecular Signatures Database (Liberzon et al., 2015). To facilitate pathway analysis the gene set was filtered to include a curated list containing only rhesus monkey genes with human homologues, and then removing non-protein coding genes, such as ribosomal RNA, miRNA, snRNA, lncRNA, as well as genes that coded for ribosomal proteins and mitochondrial-encoded proteins, as described (Niu et al., 2020). A further noise reduction step was taken to exclude genes that were not expressed in at least 99.9% of the remaining cells. Ingenuity Pathway Analysis (IPA) (QIAGEN, Germantown, MD, USA) was used to examine canonical pathway alterations. Metascape (Zhou et al., 2019) was used to perform interactome analysis using the molecular complex detection (MCODE) clustering algorithm (Bader and Hogue, 2003).

Rat immune cell data

A total of eight rat immune scRNA-seq samples (Wohnhaas et al., 2019) were downloaded from GSE127248 in the MEX format. Sample data contain the unfiltered feature-barcode matrices of all barcodes. The data were uploaded to Partek Flow, and cell barcode QA/QC was performed to determine whether a given cell barcode is associated with a cell for all 8 samples, followed by leaving a total of 35,349 cells for further analysis. Single cell QA/QC was then performed for Total UMI count and Detected Gene count, leaving a total of 31,604 cells. Noise reduction filtering was used to exclude genes that were not expressed in at least 99.9% of the remaining cells, followed by PCA, graph-based clustering, and
visualization using UMAP. Clusters containing macrophages (as shown in Figure S3) were selected, resulting in 10,405 cells. DEGs analysis, trajectory analysis and pseudotime calculation were performed similarly to the above analysis of the rhesus monkey samples.

**Statistics**

Alpha was set to < 0.05 for all analyses reported. Differentially expressed genes were determined using gene-specific analysis as performed by Partek Flow version 9.0. Pearson correlation analysis was performed using Prism version 9.0.0 (GraphPad Software, San Diego, CA).

**KEY RESOURCES TABLE**

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Anti-CD11b antibody clone M1/70, BV605 | BioLegend | Cat# 101257 |
| CD11b microbeads, non-human primate | Miltenyi Biotec | Cat# 130-091-100 |
| **Bacterial and Virus Strains** | | |
| SIVmac251 | Tulane National Primate Research Center | Del Prete et al., 2013 |
| **Biological Samples** | | |
| Indian origin rhesus macaques | New Iberia Research Center | N/A |
| **Chemicals, Peptides, and Recombinant Proteins** | | |
| Kleptose HPB Parenteral GradeEP-USP/NF | Roquette | Cat# 346111107 |
| Dolutegravir Sodium | Adooq Biosciences | Cat# A16162 |
| Maraviroc | Adooq Biosciences | Cat# A10556 |
| Emtricitabine | Gilead Biosciences | N/A |
| Tenofovir alafenamide | Gilead Biosciences | N/A |
| DNase I (for cryopreserved cell recovery) | Sigma Aldrich | Cat# 10104159001 |
| DNase (for microglia isolation from brain) | Sigma Aldrich | Cat# D4527-200KU |
| Papain | Sigma Aldrich | Cat# P5306 |
| DMSO | Sigma Aldrich | Cat# D2650 |
| **Critical Commercial Assays** | | |
| Live Dead assay UV blue | Invitrogen | Cat# L23105 |
| Arc Amine reactive compensation Bead kit | Invitrogen | Cat# A10346 |
| Single Cell 3’ GEM, Library, & Gel bead kit v3 | 10x Genomics | Cat# 1000075 |
| Single Cell 3’ Chip B Kit | 10x Genomics | Cat# 1000074 |
| NSQ 500 hi-Output KT v2 (150 CYS) | Illumina | Cat# FC-404-2002 |
| NSQ 500 hi-Output RGT CART v2 (150 CYS) | Illumina | Cat# 15057931 |
| NSQ 500 hi-Output FC CART v2 | Illumina | Cat# 15065973 |
| **Deposited Data** | | |
| NCBI GEO database, accession # GSE162663 | This paper | [https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE162663](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE162663) |
| NCBI GEO database, accession # GSE127248 (rat liver immune cells) | Wohnhass, CT, et al., Sci Rep. 2019 9:10699. | [https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE127248](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE127248) |
| Software and Algorithms |
|-------------------------|
| **Cell Ranger version 3.1** | 10x Genomics | [https://support.10xgenomics.com/single-cell-gene-expression/software/pipelines/latest/what-is-cell-ranger](https://support.10xgenomics.com/single-cell-gene-expression/software/pipelines/latest/what-is-cell-ranger) |
| **Loupe Browser version 4.2.0** | 10x Genomics | [https://support.10xgenomics.com/single-cell-gene-expression/software/visualization/latest/installation](https://support.10xgenomics.com/single-cell-gene-expression/software/visualization/latest/installation) |
| **Partek Flow version 9.0** | Partek | [https://www.partek.com/partek-flow](https://www.partek.com/partek-flow) |
| **Prism version 9.0.0** | GraphPad Software | [https://www.graphpad.com](https://www.graphpad.com) |
| **Metascape update 09-16-2020** | Zhou Y, et al., Nature Communication 2019, 10:1523 | [https://metascape.org/gp/index.html#main](https://metascape.org/gp/index.html#main) |
| **GSEA version 4.1.0** | Subramanian A, et al, PNAS 2005, 102:15545 | [https://www.gsea-msigdb.org/gsea/msigdb/index.jsp](https://www.gsea-msigdb.org/gsea/msigdb/index.jsp) |
| **Molecular Signatures Database version 7.2** | Subramanian A, et al, PNAS 2005, 102:15545 | [http://www.gsea-msigdb.org/gsea/msigdb/index.jsp](http://www.gsea-msigdb.org/gsea/msigdb/index.jsp) |

**Supplemental References**

Bader, G.D., and Hogue, C.W. (2003). An automated method for finding molecular complexes in large protein interaction networks. BMC Bioinformatics 4, 2.

Del Prete, G.Q., Scarlotta, M., Newman, L., Reid, C., Parodi, L.M., Roser, J.D., Oswald, K., Marx, P.A., Miller, C.J., Desrosiers, R.C., et al. (2013). Comparative characterization of transfection- and infection-derived simian immunodeficiency virus challenge stocks for in vivo nonhuman primate studies. J Virol 87, 4584-4595.

Li, H., Wang, S., Kong, R., Ding, W., Lee, F.H., Parker, Z., Kim, E., Learn, G.H., Hahn, P., Policicchio, B., et al. (2016). Envelope residue 375 substitutions in simian-human immunodeficiency viruses enhance CD4 binding and replication in rhesus macaques. Proc Natl Acad Sci U S A 113, E3413-3422.

Liberzon, A., Birger, C., Thorvaldsdottir, H., Ghandi, M., Mesirov, J.P., and Tamayo, P. (2015). The Molecular Signatures Database (MSigDB) hallmark gene set collection. Cell Syst 1, 417-425.

Qiu, X., Mao, Q., Tang, Y., Wang, L., Chawla, R., Pliner, H.A., and Trapnell, C. (2017). Reversed graph embedding resolves complex single-cell trajectories. Nat Methods 14, 979-982.
Subramanian, A., Tamayo, P., Mootha, V.K., Mukherjee, S., Ebert, B.L., Gillette, M.A., Paulovich, A., Pomeroy, S.L., Golub, T.R., Lander, E.S., et al. (2005). Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles. Proc Natl Acad Sci U S A 102, 15545-15550.

Wohnhaas, C.T., Leparc, G.G., Fernandez-Albert, F., Kind, D., Gantner, F., Viollet, C., Hildebrandt, T., and Baum, P. (2019). DMSO cryopreservation is the method of choice to preserve cells for droplet-based single-cell RNA sequencing. Sci Rep 9, 10699.

Zhou, Y., Zhou, B., Pache, L., Chang, M., Khodabakhshi, A.H., Tanaseichuk, O., Benner, C., and Chanda, S.K. (2019). Metascape provides a biologist-oriented resource for the analysis of systems-level datasets. Nat Commun 10, 1523.