- CAFE: An Integrated Web App for High-Dimensional Analysis and Visualization in Spectral Flow
 Cytometry
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- 18 Running Title: Spectral Flow Cytometry Analysis Platform

19 Abstract

20 Spectral flow cytometry provides greater insights into cellular heterogeneity by simultaneous 21 measurement of up to 50 markers. However, analyzing such high-dimensional (HD) data is 22 complex through traditional manual gating strategy. To address this gap, we developed CAFE as 23 an open-source Python-based web application with a graphical user interface. Built with Streamlit, 24 CAFE incorporates libraries such as Scanpy for single-cell analysis, Pandas and PyArrow for 25 efficient data handling, and Matplotlib, Seaborn, Plotly for creating customizable figures. Its robust toolset includes density-based down-sampling, dimensionality reduction, batch correction, 26 27 Leiden-based clustering, cluster merging and annotation. Using CAFE, we demonstrated analysis 28 of a human PBMC dataset of 350,000 cells identifying 16 distinct cell clusters. CAFE can generate 29 publication-ready figures in real time via interactive slider controls and dropdown menus, 30 eliminating the need for coding expertise and making HD data analysis accessible to all. CAFE is 31 licensed under MIT and is freely available at https://github.com/mhbsiam/cafe.

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34 *Word count*: 4633

36 Introduction:

37 Flow cytometry is a widely used technique in immunology to identify and quantify immune cells based on specific surface markers¹. The development of spectral flow cytometry (SFCM) has 38 further expanded immunophenotyping capabilities allowing the simultaneous analysis of a greater 39 40 number of parameters through the complete emission spectra of fluorophores¹. Compared to conventional flow cytometry, SFCM uses spectral unmixing algorithms to deconvolute the 41 42 overlapping signals and achieves enhanced resolution and sensitivity to distinguish between different cell populations². SFCM can incorporate broader range of antibodies with up to 50 colors 43 in a single panel improving upon the conventional FCM where the number of parameters is limited 44 by the instrument constraints³. Incorporating more parameters substantially increases the 45 46 complexity in gating strategy which largely relies on established convention and prior knowledge^{4,5}. Additional gating steps and combinations of markers used to subset cells 47 48 complicate the interpretation of such high-dimensional data. Several clustering methods are 49 available to identify cell populations such as FlowSOM⁶, xShift⁷, SPADE⁸ and Phenograph⁹. 50 SPADE and FlowSOM utilize hierarchical clustering with the latter employing self-organizing maps (SOMs) to cluster cells, whereas xShift detects clusters based on shifts in local cell density⁶⁻⁸. 51 52 Phenograph, by contrast, constructs a K-nearest neighbor graph and applies the Louvain 53 algorithm to identify cell clusters, but Louvain can produce poorly connected or disconnected communities^{9,10}. 54

Recently, the Leiden clustering algorithm has emerged as a faster and more accurate alternative 55 to improve community detection in networks¹⁰. Single-cell RNA sequencing (scRNA-seq) tools: 56 Seurat¹¹ (R) and Scanpy¹² (Python) have integrated Leiden algorithms for community detection. 57 However, running Leiden within Seurat resulted in drawbacks including higher memory usage, 58 59 longer calculation time and random crashes in docker containers¹³. Scanpy resolves these issues, 60 and unlike Seurat, Scanpy improves visualization guality by using consistent KNN and SNN 61 graphs for both clustering and uniform manifold approximation and projection (UMAP) ^{13,14}. In February 2020, Phenograph version 1.5.3 was released, which incorporated an option to use 62 63 Leiden for clustering; however, the default parameter is set to Louvain through the latest release 64 (v.1.5.7). In our previous work, we showed that the use of Leiden algorithm in community detection for SFCM data provides superior result to Phenograph (Louvain), FlowSOM, and xShift¹⁵. 65 Currently, there is a scarcity of open-source tools to utilize Leiden algorithm for SCFM data 66 analysis¹⁶. 67

68 Here we present CAFE, Cell Analyzer for Flow Experiment, a user-friendly web application 69 developed in Python that works across Windows, MacOS, and Linux. The app is lightweight and 70 can perform high-dimensional SFCM data analysis using a standard computing machine (i.e., 71 Apple M1 chip with 16qb RAM), and it provides the flexibility to be deployed on HPC clusters for 72 enhanced scalability. Once installed, the tool runs entirely offline and does not require an active 73 internet connection to load files. This also enables users to maintain compliance with data 74 security, especially with protected health information (PHI) when analyzing patient samples. 75 CAFE can be used to process data, reduce dimension, batch correction, run Leiden clustering, 76 perform statistics and generate a wide range of figures. Figures can be adjusted and viewed within the tool in real time. Additionally, the tool offers Kernel Density Estimation (KDE)-based data 77 downsampling, advanced clustering with predefined markers, cluster quality evaluation, merging 78 79 subclusters into metaclusters, and cell type annotation. Designed as an open-source interactive data analysis platform, CAFE enables biologists with no-coding experience to analyze SFCM data 80 81 and create publication quality visualization with customizable parameters. CAFE is freely 82 available to download at: https://github.com/mhbsiam/cafe

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84 Methods:

85 *Implementation*

86 The CAFE webtool was developed using Python programming language due to its compatibility with Scanpy library¹². Figure 1 illustrates the components and workflows of CAFE. Streamlit 87 (streamlit.io) library was used to develop the web interface that provides dynamic updates based 88 89 on user inputs in the graphical user interface (GUI) without writing or editing code directly. 90 Streamlit was chosen due to its simplicity in development and compatibility with other Python 91 libraries across operating systems. Streamlit v1.39.0 is compatible with any modern HTML5 web browser. For data loading and processing, we relied on Pandas v2.2.3 with PyArrow v18.0.0 which 92 93 achieves faster data loading and processing compared to Pandas alone. We used NumPy v1.26.4 94 for data type and range selection, RGB array creation for color handling, and grid setup for 95 subplots. Seaborn v0.13.2, Matplotlib v3.9.2, and Plotly v4.24.1 libraries were used for data 96 visualization and users are provided with options to adjust parameters: plot size, color profile, and output formats in PNG, JPG, SVG, or PDF. CAFE integrates AnnData¹², a widely used framework 97 in single-cell RNA sequencing analysis that allows for efficient storage and manipulation of both 98

sparse and dense matrices along with metadata. CAFE outputs an AnnData object which can be
used outside of CAFE if users wish to deposit their data with analysis or perform custom analyses
using other tools.

102 The Scanpy library was used to perform key analyses including dimension reduction, batch 103 correction, and Leiden clustering. The user has the option to reduce dimension (sc.tl.pca) of the 104 data through Principal Component Analysis (PCA) or skip it. PCA is a linear dimensionality 105 reduction technique that retains the global structure of the data by capturing the variances across 106 all dimensions. Because the app performs PCA through Scanpy library, by default, the number of components retained is limited to the lesser of the two values: the number of cells or the number 107 108 of markers. Also, the Singular Value Decomposition (SVD) solver was set to "auto" that chooses 109 the most appropriate solver based on the size of the dataset; however, users have options to set 110 a percentage of variances they want to retain, and the type of solver used. The reduced dataset 111 is stored and can be further processed for batch correction (sc.pp.combat) using ComBat (Combined Batch)¹⁷. This is particularly useful if a user has collected samples in different batches 112 113 as the algorithm standardizes the data by making it comparable and removing unwanted 114 variability.

115 To group cells into distinct clusters based on marker expression profiles, Leiden clustering is run (sc.tl.leiden) and users can select either 'iGraph' or 'leidenalg' algorithm flavor^{10,18}. To define 116 clustering resolution, a user can choose from 0.01 to 2.0 where the lowest value provides the 117 118 lowest number of clusters. The user can fine tune Leiden calculation by altering the number of 119 neighbors and minimum distance values in Uniform Manifold Approximation and Projection 120 (UMAP) calculation within the app. CAFE generates AnnData object (H5AD) file, CSV outputs, 121 and visual outputs including UMAP plots, dot-heatmaps, expression pots, and barplots as high-122 resolution images and provides download buttons to save them to a desired folder. The app allows 123 various visualization settings, with changes made and displayed immediately within the app. The 124 generated AnnData object can be further used to perform a range of statistical analyses.

The app includes advanced functionalities for clustering and cluster evaluation. Because setting up appropriate values for Leiden resolution and UMAP parameters is central to obtaining quality clustering results, a user can leverage CAFE's Cluster Evaluation tab to generate multiple AnnData files with various combinations of these parameters and compare UMAP plots as well as Silhouette score, Calinski-Harabasz score, Davies-Bouldin score, and Elbow method to assess clustering results. Besides, CAFE provides clustering with pre-selected markers, merging 131 subclusters into metaclusters, and annotation of clusters directly within the app. The advanced 132 Downsampling tab offers to downsample (e.g. 20,000 events per sample) data using a PCA-KDE 133 based method. This approach combines PCA and KDE (Kernel Density Estimation) based 134 algorithm from scipy stats using Gaussian kernel function and silverman bandwidth¹⁹. KDE is 135 applied to the PCA-transformed data to estimate the density of data points. Based on these density estimates, the code probabilistically downsamples data, thus reducing sampling bias and 136 137 preserving original data distribution. This method offers an informed approach compared to simple 138 random downsampling, and it can be used to filter out noise while retaining meaningful biological 139 information in a smaller dataset.

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141 Statistical analysis

142 In the visualization tab under statistical analyses section, users can perform different statistical 143 tests and generate plots. The Shapiro-Wilk test from "scipy.stats" is used to determine if marker 144 expression within each group or cluster follows a normal distribution. Based on these results, the app recommends either parametric (T-test) or non-parametric (Mann-Whitney U test) tests using 145 146 "scipy.stats". For comparing multiple clusters, the app allows users to perform ANOVA 147 (scipy.stats.f oneway) or Kruskal-Wallis (scipy.stats.kruskal) tests. To reduce statistical artifacts, multiple testing correction is applied using the Benjamini-Hochberg False Discovery Rate (FDR) 148 149 through "statsmodels.stats.multitest". Additionally, effect size measures are computed to 150 complement statistical p-values, with users able to choose between parametric tests (Cohen's d) 151 or non-parametric tests (Cliff's Delta). Cohen's d is calculated using basic functions from Numpy, 152 while Cliff's Delta is computed with the "cliffs delta" package. To assess associations between 153 clusters and groups, we used Chi-square testing from "scipy.stats.chi2 contingency" and 154 contingency tables with "pandas.crosstab". Residual calculations were displayed in Streamlit as 155 tables to help users understand which clusters are more prevalent within certain groups.

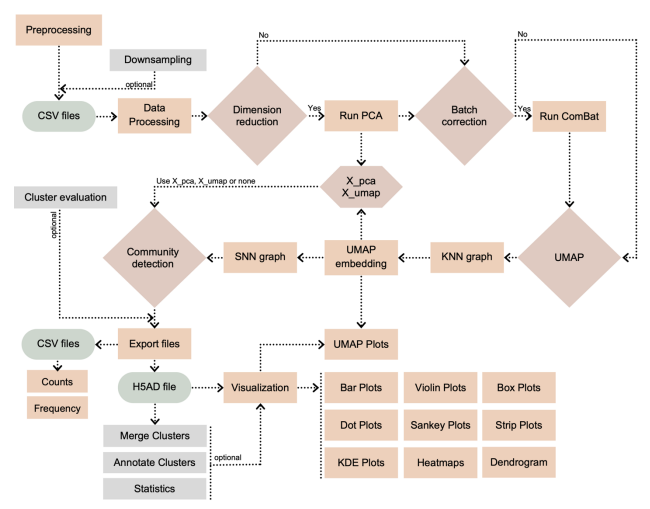
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157 **Performance and reproducibility**

We have set a global setting for Scanpy (sc.settings.n_jobs = -1) to use all available CPU cores. For advanced clustering, multi-threading was achieved using Python's joblib library. Two other libraries were used, watchdog v5.0.2 and iGraph v0.10.8¹⁸. Watchdog helps in monitoring file 161 change events and improves performance of Streamlit by providing real-time feedback. iGraph is 162 designed to handle complex networks and graph operations and is used by Scanpy as part of the 163 Leiden clustering to perform graph operations. We recommend 'iGraph' over 'leidenalg' as iGraph 164 is implemented in C and achieves advantages in performance compared to high-level interpreted languages such as Python. To export Plotly figures, we have used Kaleido engine v0.2.1. We 165 have tested the app with various datasets using an Apple M3 Pro System with 18GB of random-166 167 access memory (RAM). CAFE is primarily intended to be used using local computer; however, it can be scaled up using any High-Performance Computing (HPC) system that supports an HTML5 168 169 web browser. We also provided scripts in our GitHub page to generate AnnData with dimension 170 reduction and Leiden clustering through command-line interface (CLI) based HPC systems.



172



173 Figure 1: The flowchart outlines steps and components of CAFE's workflow.

174 Preprocessing includes compensation, data scaling/transformation using a standard FCM

software and scaled CSV files are then exported and renamed as Sample_Group.csv. Data

176 processing performs error checks and concatenation of CSV files into an AnnData object/H5AD

177 file. Major steps requiring user input include dimension reduction, batch correction, UMAP

(UMAP Uniform Manifold Approximation and Projection) and community detection. Outputs are
 downloadable as CSV, H5AD, PNG, JPG, SVG and PDF files.

181 **Results:**

182 To demonstrate the functionality of the app, we have analyzed 35-color spectral flow cytometry 183 data (Publicly available at FlowRepository: FR-FCM-Z3WR) of human peripheral blood 184 mononuclear cells (PBMC) obtained from COVID-19 hospitalized patients and healthy controls²⁰. 185 A total of 10 samples were analyzed with 5 from each group. For best practices, we installed and 186 ran CAFE through Pixi package manager. Users can also install and run the app using Anaconda 187 package manager as described in our Github documentation. Once initiated through a terminal 188 (pixi run cafe, or python cafe.py), a web browser opens with the CAFE app at localhost on port 189 8501. The default data loading limit is set to 3GB, but a user can change the value from the 190 cafe.py script if necessary.

191 Data processing

192 The uploaded public data were available as doublets-debris removed and CD45+ gated; so we 193 obtained the CSV files just by exporting scaled values from FlowJo v10.10.0. Data scaling is 194 generally recommended for high resolution clustering but there may be instances where users 195 may use raw values. Data can be similarly exported from other flow cytometry software such as 196 FCS Express. It is required that flow cytometry data have proper compensation. We recommend 197 manual inspection of flow cytometry data and removal of debris, dead cells, and doublets prior to 198 exporting the scaled files. A user can also gate on appropriate cell type and export the data to 199 obtain more focused clustering results. To streamline downstream analysis, we have implemented 200 a naming convention for the CSV files. Each CSV file name must begin with a unique 201 "SampleName" followed by "GroupName", separated by an underscore; for instance, "Sample01 Control.csv" and "Sample02 Treatment.csv". After loading the data, the app will 202 203 import the required libraries and perform initial checks for data structure and incorporate 204 SampleID and GroupName into the dataframe based on the CSV file names. Within the 205 dataframe, rows containing any missing values are skipped and anomalies in data structure are 206 reported. In this study, we used the advanced KDE-based downsampling option in CAFE to 207 downsample data to 35,000 cells per sample for a total of 350,000 cells and 12.25M data points (number of cells multiplied by number of markers). This is an optional step prior to data 208

¹⁸⁰

processing. After loading the files, the app processed (7.8 sec) and combined the expression dataand metadata without errors to create an AnnData object.

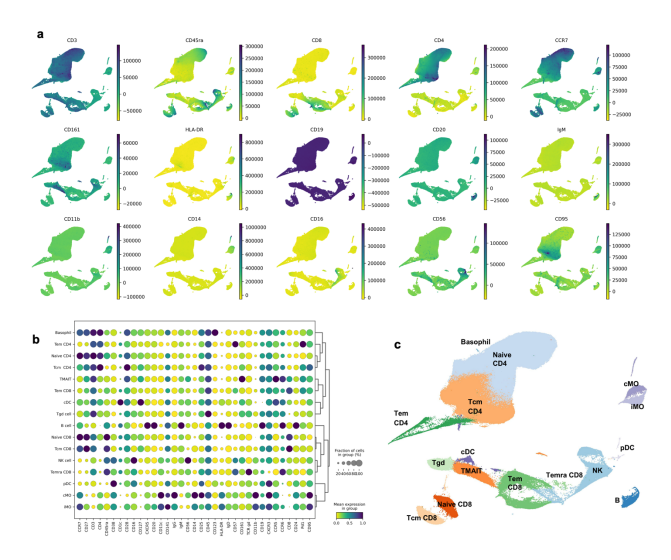
211 **Dimension reduction and batch effects**

212 After generating the AnnData object (H5AD file), we selected dimension reduction using PCA with

- SVD solver set to auto and retained components with 95% variance. The app ran PCA (2.16 sec),
- kept 12 components and generated (PC1 x PC2) graphs by Groups. Depending on the data size,
- a user can choose from auto, full, arpack, and randomized SVD solver. Randomized, for example,
- 216 is better suited for larger datasets as it provides a balance between speed and accuracy. For
- batch correction, we applied ComBat (1.06 sec) and proceeded to Leiden clustering.

218 Leiden clustering and metaclustering

219 For the dataset, we applied Leiden resolution of 1.0 with flavor set to iGraph, UMAP n neighbors 220 to 15, min dist to 0.1 and distance calculation method as Euclidean. A user has the option to use 221 a slider control to choose from resolution values 0.01 to 2.0. To find the optimal resolution, we 222 initially made use of Advanced Cluster Evaluation option in CAFE to generate a series of AnnData 223 files with varied Leiden resolution and n neighbor values and observed the UMAPs to find distinct 224 clusters that are biologically meaningful for the dataset. With Leiden resolution of 1.0, we initially 225 obtained a total of 30 clusters for the PBMC dataset which took 11.5 minutes for calculation. Once 226 clustering was completed, CAFE generated a frequency table of each sample by Leiden cluster 227 for the number of cells, frequency of cells, and median fluorescence intensity of each marker for 228 each cluster. Using these 3 tables, users can perform statistical analyses to compare cluster count 229 and frequency by groups and expression of marker proteins within clusters by group. Using the 230 Advanced Cluster Merging option, we merged the subclusters with similar profile into 231 corresponding metaclusters resulting in a new total of 16 clusters.



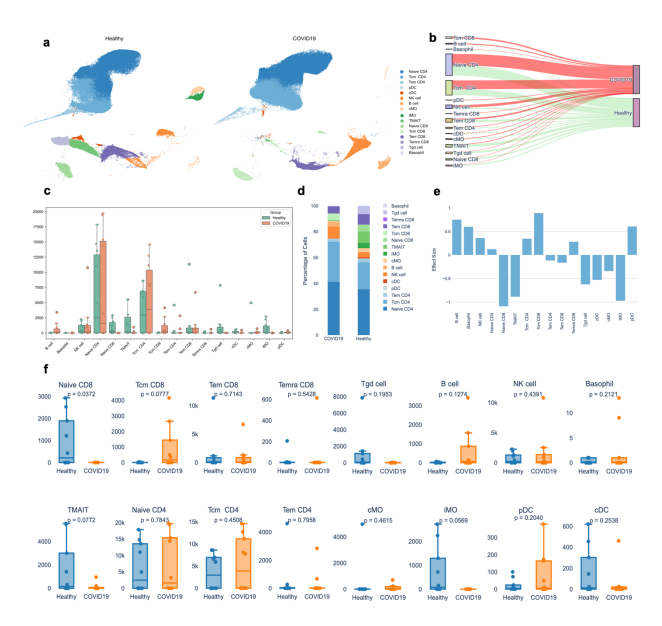
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234 Figure 2: Profiling of Human PBMCs Reveals Distinct Immune Suppopulations and Marker Expression Patterns. (a) UMAP plots showing selected marker expression intensities across all 235 236 cells in the UMAP space to highlight lineage-specific marker distribution. (b) Dot plot of all marker 237 expression across all identified PBMC cell types. Dendrogram highlighted distinct marker-based 238 groupings. (c) UMAP visualization showing 16 distinct clusters with annotated cell types including 239 Naive CD4 and CD8 T cells, central memory CD4 and CD8 T cells (Tcm), effector memory CD8 T cells (Tem), terminally differentiated effector memory CD8 T cells (Temra), mucosal-associated 240 invariant (MAIT) T cells, classical monocytes (cMO), intermediate monocytes (iMO), B cells, NK 241 cells, gamma delta ($y\delta$) T cells (Tgd), conventional dendritic cell (cDC) and plasmacytoid dendritic 242 cell (pDC). 243

244 Characterization of PBMC Subpopulations

To characterize the phenotypic properties, we examined the expression of surface markers for each identified cluster by protein expression UMAP plots (Figure 2a). We found high CD3 expression in T cells clusters with CD4 and CD8 expression showing corresponding T cell 248 subtypes. CD8+ effector memory (Tem) and central memory (Tcm) subsets were differentiated by 249 high CCR7 and CD27 expression in Tcm. We used CD45RA expression to identify terminally 250 differentiated Tem cells (Temra). Monocyte clusters were identified by CD14 and CD16 251 expression, distinguishing classical monocytes (cMO) from other monocyte subsets, while natural 252 killer (NK) cells showed high levels of CD56, corresponding with CD56^{Bright} NK cells. Based on 253 shared marker profiles and hierarchical ranking, T cell subsets (Tem, Tcm, and Temra) formed a 254 distinct grouping separate from B cells and myeloid-derived cells, reflecting the differential 255 expression of lineage-specific markers.

256 Based on the expression profiles of marker proteins, we annotated the clusters using CAFE's 257 Advanced Annotation tab and classified them into 16 distinct cell types. We also used the dotplot 258 to confirm annotations of the cell types (Figure 2b). For instance, the B cell-specific marker CD19 259 and CD20 were used to identify the B cell cluster, the CD14 marker to identify monocytes, and 260 the CD16 marker to identify NK cells. High CD20 expression in B cell cluster indicated their mature 261 stage in immune response. Our annotated UMAP (Figure 2c) shows well-defined clusters that 262 correspond to PBMC lineages, including Naive CD4+ and CD8+ T cell and Tcm for both CD4+ 263 and CD8+ subsets. We also identified Tem and Temra cells, as well as mucosal associated 264 invariant CD8+ T cell (MAIT). We also identified cMO, intermediate monocytes (iMO), B cells, NK 265 cells, Gamma delta ($\gamma\delta$) T cells (Tqd) and dendritic cell (DC) types (cDC and pDC).



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Figure 3: Comparative Analysis of Immune Cell Subpopulations in Healthy and COVID-19 268 Individuals. (a) UMAP plots displaying distinct clustering patterns and differential distribution of 269 270 cell types in PBMC across healthy and COVID-19 group. (b) Sankey diagram illustrates the 271 distribution of cells across groups, with thicker flow indicating more cells. (c) Composite bar-strip plot summarizing cell count distribution across cell subpopulations. Dots represent each individual 272 samples colored by group. (d) Stacked bar chat showing distribution of cells in percentage across 273 two groups. (e) Effect size calculated using Cohen's d indicating changes in the number of cells 274 in COVID-19 compared to reference healthy control. (f) Comparison of individual cell type 275 frequencies between healthy and COVID-19 groups with p-values for statistical significance. 276 277 N=9/group

Distinct Cellular and Molecular Signatures Observed in COVID-19 Compared to Healthy Controls

UMAP analysis of COVID-19 hospitalized patients compared to healthy controls revealed distinct 281 282 clustering patterns between groups, particularly among monocytes, NK cells, and CD8 T cells (Figure 3a). To understand changes between the two groups, CAFE offers varied visualization 283 284 options, for instance, we used a Sankey diagram to demonstrate that MAIT cells and Tgd are 285 much less abundant in COVID-19 patients compared to healthy controls (Figure 3b). We also 286 found that CD8 Tcm and B cells were significantly expanded in COVID-19 patients. A composite 287 bar-strip plot also demonstrates the distribution of cells in frequency where each dot represented 288 each sample colored by specific group (Figure 3c). The total number of cells in iMO were largely 289 reduced in COVID-19 patients compared to healthy controls (Figure 3d). These data may indicate 290 a possible shift from an innate response towards an adaptive response. To quantify the effect size 291 of changes observed, we compared cell types within the COVID-19 group to healthy controls as 292 a reference and found changes in naive CD8, TMAIT, and iMO cells have a larger effect size, 293 demonstrating a bigger difference between the two groups (Figure 3e). We further compared 294 these cell types by plotting box plots for individual cell types (Figure 3f) which demonstrated a 295 non-statistically significant increase in Tcm CD8 (p=0.0777) and statistically significant decrease 296 in naive CD8 cells (p=0.0372) in COVID-19 patients compared to healthy controls.

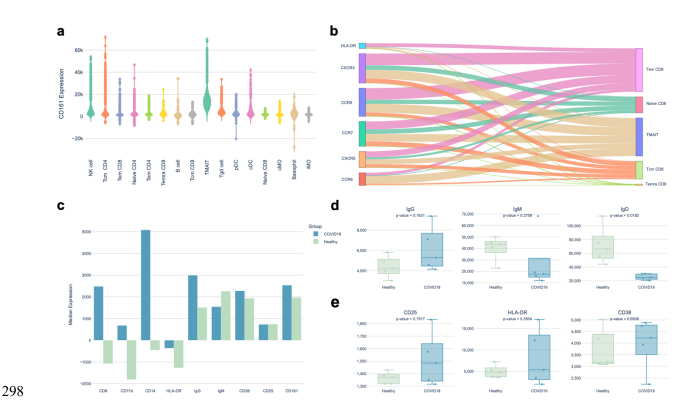


Figure 4: Marker expression and distribution differences between COVID-19 and healthy individuals. (a) Violin plot showed the median expression levels of CD161 across immune cell subtypes. (b) Sankey diagram illustrated marker expression in CD8+ T cells, with thicker flows indicating more cells expressing that marker. (c) Bar chart showed median expression of markers across all cell types between COVID-19 and healthy individuals where positive values indicated upregulation. Box plots displayed (d) the number of cells expressing IgG, IgM and IgD in B cells, and (e) the number of CD8+ T cells expressing CD25, HLA-DR, and CD38.

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307 Altered Marker Expression Profiles in COVID-19 Patients

308 CAFE outputs a file with expression data for all markers on each cluster by sample for use in external plotting and statistical software. In addition, further exploration of specific markers within 309 310 clusters can be performed within CAFE. We used this approach to identify that MAIT cells have 311 the greatest expression of CD161, as shown by violin plot (Figure 4a). We also examined marker distribution in CD8 T cell populations and found that more cells within the Tem CD8 population 312 appeared activated based on greater HLA-DR, CXCR3, and CCR5 expression compared to other 313 CD8 T cell subsets (Figure 4b). We found that median expression of CD8, CD14, CD11b, IgG 314 315 were increased in COVID-19 patients compared to healthy controls across all clusters (Figure 316 4c). These reflect the overall differences in some of the cell populations we observed between groups. We examined marker expression within the B cell cluster and found that more B cells expressed IgG in COVID-19 samples compared to healthy controls although the difference was not statistically significant (p=0.1631), while IgD expression was statistically significantly reduced (p=0.0162) in COVID-19 samples compared to healthy controls (Figure 4d). We also examined activation markers in CD8+ T cells and found that COVID-19 patients had more CD8+ T cells that expressed CD25 and HLA-DR than healthy controls (Figure 4e).

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325 **Discussion:**

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As research in immunology increasingly relies on high-dimensional cytometric data, there is a growing need for a user-friendly analysis tools for everyday use. Here, we present CAFE as a free and open-source tool designed to address the analytical and accessibility issues posed by SFCM data. CAFE uses a GUI and interactive controls to enable immunologists to analyze complex data without needing specialized coding knowledge.

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333 Using a jupyter notebook, we have previously shown the ability of Scanpy's Leiden function 334 (scanpy.tl.leiden) to analyze a 50-color human PBMC dataset¹⁵. CAFE acts as a wrapper 335 combining packages within Streamlit to provide a web interface, offering more accessible and 336 extensive functionality than Jupyter notebooks' CLI. We demonstrated analysis of a 35-color 337 human PBMC dataset using CAFE in this study with 350,000 cells and 12.25M data points. Major 338 steps including data processing, dimension reduction, batch correction and Leiden clustering 339 were completed in under 12 minutes using an Apple M3 Pro laptop. In our analysis, we observed 340 COVID-19 patients with altered immune cell distributions and marker expression profiles. 341 consistent with prior findings, as well as MAIT cells expressing high CD161 and B cells expressing high CD20²⁰. 342

While developing CAFE, we have balanced compatibility and performance and included many options for customization of how the code processes and analyzes data while integrating default options and tooltips to help guide users. Our implementation of Pandas with PyArrow significantly improves processing speed over Pandas alone. However, transitioning to Polars' lazy evaluation framework could further speed up processing once compatibility issues between ARM and x86 machines are resolved. CAFE's web app design and functionality also revolved around simplicity as we de-emphasized features that are not commonly used in order to streamline user-

350 experience. For data dimension reduction, we adhered to a convention of using PCA as the 351 primary method and using PCA-reduced representations for constructing UMAP neighbor graphs, 352 as opposed to utilizing UMAP directly for primary dimensionality reduction. Although PCA is 353 designed for linear data, it effectively reduces noise and enhances clustering performance. Users 354 have the choice to skip PCA to perform Leiden clustering on the raw data or use UMAP 355 embeddings (i.e., X umap) to use UMAP reduced data for clustering. One viable alternative to 356 PCA for non-linear data structure is Kernel PCA, but we have skipped adding the kernel PCA 357 option in the CAFE workflow because it may not be practical since it is computationally taxing.

358 UMAP parameters and Leiden resolution largely influence the number of clusters for community 359 detection. Leiden clustering is performed on the graph structure, so evaluation of clustering guality 360 solely based on methods such as elbow or silhouette score is not ideal. Rather a combined approach with prior biological knowledge can inform the most correct clustering resolution. We 361 362 recommend using CAFE's advanced clustering evaluation tab to generate plots with a range of 363 varied UMAP parameters and Leiden resolutions for visual inspection. Using this approach, users 364 can select the most appropriate clustering resolution for each dataset. Since this is an 365 unsupervised algorithm, setting up an incorrect resolution can heavily skew the interpretation of 366 data.

367 Manual gating continues to be the gold standard in flow cytometry analysis, but it is limited by 368 sequentially drilling down into subsets of cells with 2-dimensional bi-axial gating. Thus, our goal 369 was to complement this hypothesis-driven approach with the unsupervised computational 370 algorithms. In this way, users can perform hypothesis-driven analysis with manual gating and 371 hypothesis-generating analysis with unsupervised clustering. Compared to other open-source 372 tools, CAFE provides a wide range of publication-ready visualization options. Due to its underlying 373 code in python, CAFE is highly scalable to datasets of millions of cells and takes advantage of 374 multi-threading to obtain higher performance. Previously, python-based Pytometry and CRUSTY integrated unbiased clustering algorithms within their tools^{21,22}. Pytometry incorporates the Leiden 375 algorithm¹⁰, which has been shown to be an improvement over the predecessor Louvain 376 algorithm²³: however, Pytometry requires coding using Python. CRUSTY incorporates an easy-377 378 to-use GUI but it does not offer the Leiden algorithm and relies on FlowSOM and Phenograph. 379 Another limitation of CRUSTY is that the cloud-based service limits users to analyzing 100,000 380 total events. There are also limited visualization and analysis options in CRUSTY and they rely on most of the Phenograph and FlowSOM default settings which cannot be customized. Cloud-381 382 based solutions may face limitations in availability, scalability, and data security. Users may be

383 prohibited from uploading data to cloud-based systems that have protected health information due to HIPAA. Among a few other GUI based tools, Cytoflow²⁴, Floreada (floreada.io), EasyFlow²⁵ 384 385 allow for flow cytometry data analysis but do not offer clustering. FlowPy (flowpy.wikidot.com) 386 allows for clustering but uses k-mean clustering rather than the most advanced algorithms 387 currently in use (i.e. Leiden). Additional tools like terraFlow²⁶ and CellEngine (CellCarta, Montreal, Canada) are for-profit spectral flow cytometry analysis softwares and the price of these may be 388 389 restrictive for many users. Finally, FlowJo is a staple for many immunologists and has some native 390 clustering capabilities. It also supports plugins for additional clustering algorithms, but these add-391 ons do not offer much customization in the clustering parameters.

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393 CAFE, while addressing many of these limitations as an open-source alternative, has its own 394 practical considerations. CAFE is intended to be run locally which requires installing a package 395 manager such as Pixi or Anaconda/Miniconda3 through terminal. Performance is also dependent 396 on the user's machine. For larger datasets, we recommend utilizing our provided scripts in Github 397 to run data processing step through an HPC cluster by allocating more RAM. Once the user has 398 Anndata file generated with cluster information, all the downstream analysis and figure generation 399 steps become significantly less computationally demanding. Ultimately, CAFE's aim is to become 400 a secure, scalable, and open-source platform accessible to a broad range of researchers to run 401 complex analyses through a simple intuitive graphical user interface.

402

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408

409 **Data availability:** The raw data are publicly available at FlowRepository: FR-FCM-Z3WR. The 410 downsampled .csv files are available from Figshare: 27940719. The processed Anndata object 411 after dimensionality reduction and clustering is also available from Figshare: 27940752.

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413 **Code availability:** CAFE is freely available at https://github.com/mhbsiam/cafe.

415	Co	ntributions: M.H.B.S. and D.J.T. conceived the software development. M.H.B.S. created the
416	sof	tware and web application; M.H.B.S. analyzed the data. M.H.B.S. and D.J.T. wrote the paper.
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420		
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479 **Figure Legends**:

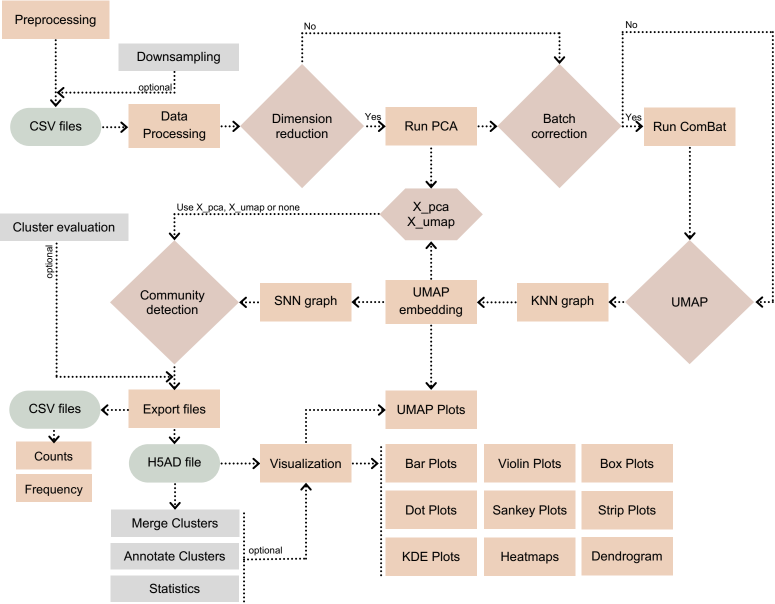
Figure 1: The flowchart outlines steps and components of CAFE's workflow. Preprocessing
includes compensation, data scaling/transformation using a standard FCM software and scaled
CSV files are then exported and renamed as Sample_Group.csv. Data processing performs error
checks and concatenation of CSV files into an AnnData object/H5AD file. Major steps requiring
user input include dimension reduction, batch correction, UMAP (UMAP Uniform Manifold
Approximation and Projection) and community detection. Outputs are downloadable as CSV,
H5AD, PNG, JPG, SVG and PDF files.

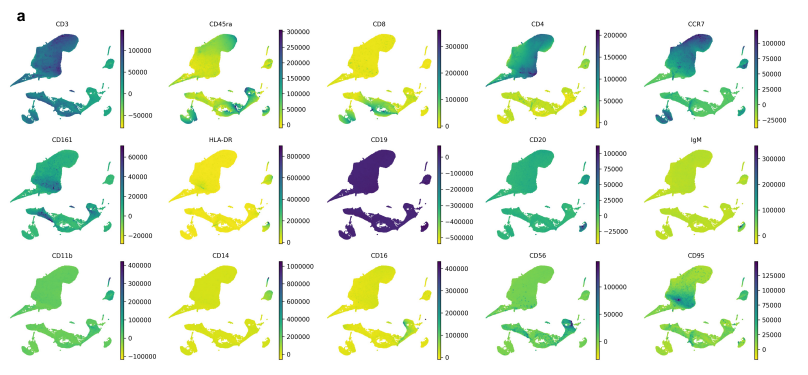
487 Figure 2: Profiling of Human PBMCs Reveals Distinct Immune Subpopulations and Marker Expression Patterns. (a) UMAP plots showing selected marker expression intensities across all 488 cells in the UMAP space to highlight lineage-specific marker distribution. (b) Dot plot of all marker 489 490 expression across all identified PBMC cell types. Dendrogram highlighted distinct marker-based 491 groupings. (c) UMAP visualization showing 16 distinct clusters with annotated cell types including 492 Naive CD4 and CD8 T cells, central memory CD4 and CD8 T cells (Tcm), effector memory CD8 493 T cells (Tem), terminally differentiated effector memory CD8 T cells (Temra), mucosal-associated 494 invariant (MAIT) T cells, classical monocytes (cMO), intermediate monocytes (iMO), B cells, NK 495 cells, gamma delta ($\gamma\delta$) T cells (Tgd), conventional dendritic cell (cDC) and plasmacytoid dendritic 496 cell (pDC).

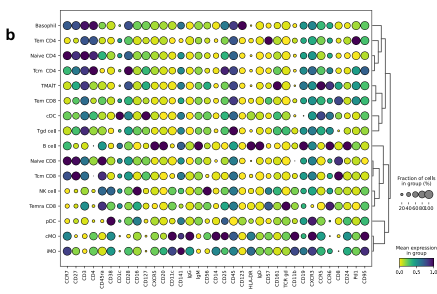
497 Figure 3: Comparative Analysis of Immune Cell Subpopulations in Healthy and COVID-19 498 **Individuals.** (a) UMAP plots displaying distinct clustering patterns and differential distribution of 499 cell types in PBMC across healthy and COVID-19 group. (b) Sankey diagram illustrates the 500 distribution of cells across groups, with thicker flow indicating more cells. (c) Composite bar-strip 501 plot summarizing cell count distribution across cell subpopulations. Dots represent each individual 502 samples colored by group. (d) Stacked bar chat showing distribution of cells in percentage across 503 two groups. (e) Effect size calculated using Cohen's d indicating changes in the number of cells 504 in COVID-19 compared to reference healthy control. (f) Comparison of individual cell type 505 frequencies between healthy and COVID-19 groups with p-values for statistical significance. 506 N=9/group.

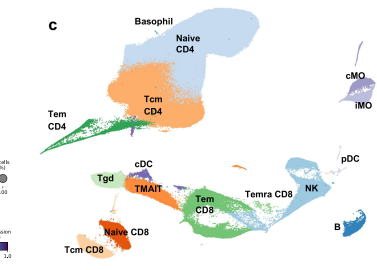
507 Figure 4: Marker expression and distribution differences between COVID-19 and healthy

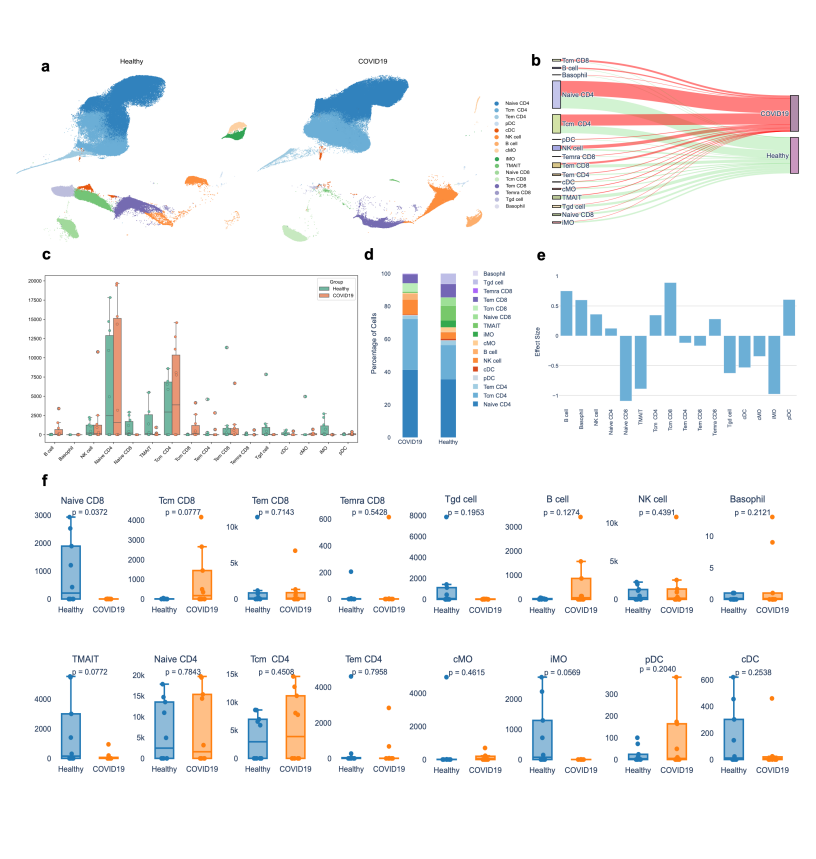
individuals. (a) Violin plot showed the median expression levels of CD161 across immune cell
 subtypes. (b) Sankey diagram illustrated marker expression in CD8+ T cells, with thicker flows
 indicating more cells expressing that marker. (c) Bar chart showed median expression of markers
 across all cell types between COVID-19 and healthy individuals where positive values indicated
 upregulation. Box plots displayed (d) the number of cells expressing IgG, IgM and IgD in B cells,
 and (e) the number of CD8+ T cells expressing CD25, HLA-DR, and CD38.

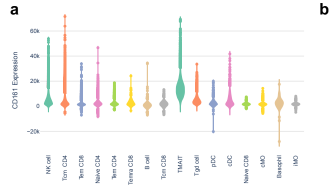




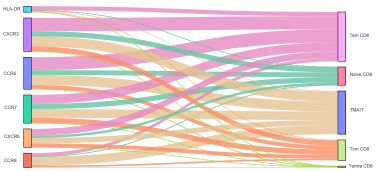








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