Improved integration of single-cell transcriptome and surface protein expression by LinQ-View

Highlights

- LinQ-View integrates mRNA and protein expression data to reveal cell heterogeneity

- LinQ-View prevents single dominant ADT features from affecting clustering

- LinQ-View presents a quantitative purity metric for CITE-seq data

- LinQ-View is specialized in handling CITE-seq data with fewer ADT features

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In brief

Li et al. present LinQ-View, a computational workflow that provides an effective solution for integrating multiple modalities of CITE-seq data and quantitative assessment of cluster purity. LinQ-View could be helpful for multimodal analysis and purity assessment of CITE-seq datasets that target specific cell populations.
Improved integration of single-cell transcriptome and surface protein expression by LinQ-View

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SUMMARY

Multimodal single-cell sequencing has enabled the simultaneous quantification of cell surface protein expression alongside unbiased transcriptional profiling. Here, we present LinQ-View, a toolkit designed for multimodal single-cell data visualization and analysis. LinQ-View integrates transcriptional and cell surface protein expression profiling data to reveal more accurate cell heterogeneity and proposes a quantitative metric for cluster purity assessment. Through comparison with existing multimodal methods on multiple public CITE-seq datasets, we demonstrate that LinQ-View efficiently generates accurate cell clusters, especially in CITE-seq data with routine numbers of surface protein features, by preventing variations in a single surface protein feature from affecting results. Finally, we utilized this method to integrate single-cell transcriptional and protein expression data from SARS-CoV-2-infected patients, revealing antigen-specific B cell subsets after infection. Our results suggest LinQ-View could be helpful for multimodal analysis and purity assessment of CITE-seq datasets that target specific cell populations (e.g., B cells).

INTRODUCTION

In the past decade, advances in single-cell sequencing have ushered in a new era of discovery in biology and medicine. Improved sensitivity, reliability, throughput, and sequencing depth have provided researchers immense power to study diverse biological systems at single-cell resolution (Cao et al., 2017; Hashimshony et al., 2012; Klein et al., 2015; Macosko et al., 2015; Picelli et al., 2013; Ramsköld et al., 2012; Rosenberg et al., 2018; Tang et al., 2009; Zheng et al., 2017). Recent technological advances have enabled the simultaneous measurement of multiple cell properties in combination with single-cell...
RNA sequencing (scRNA-seq), allowing for an unprecedented view of cell heterogeneity. Cellular indexing of transcriptomes and epitopes by sequencing (CITE-seq) and RNA expression and protein sequencing assay are capable of measuring the expression of surface protein markers alongside the transcriptome at single-cell resolution (Peterson et al., 2017; Stoeckius et al., 2017). Because much of foundational immunology has been elaborated through the use of specific lineage-restricted cell-surface proteins to define immune cell populations, direct estimation of protein expression might enable improved methods for delineating cell populations.

To gain a better understanding of immune functions at single-cell resolution, the integration of surface protein expression and transcriptional profiling data are critical. Existing single-cell analysis software packages are typically designed for the analysis of unimodal data and provide limited integration of additional profiling modalities, such as cell-surface protein expression data. In CITE-seq, protein expression is estimated through the use of antibody-derived tags (ADTs), which promise to better characterize immune cell heterogeneity (Peterson et al., 2017; Stoeckius et al., 2017; Stuart and Satija, 2019). Recently, a few methods have been published or pre-released for integrative analysis of CITE-seq data (Argelaguet et al., 2018, 2020; Hao et al., 2020; Kim et al., 2020; Shen et al., 2009, 2013; Wang et al., 2014). Although these tools are valuable, improvements to the workflow and imbalances between transcriptome and surface protein expression are warranted. In addition, there is a lack of a quantitative metric of cluster purity for CITE-seq data. Thus, there remains a need for efficient methods capable of integrating transcriptome and surface protein expression for single-cell CITE-seq data that allow downstream analyses with deeper resolution.

To overcome these challenges, we developed a single-cell analysis toolkit called “LinQ-View” for integrative analysis of CITE-seq data. LinQ-View integrates information from transcriptome and cell-surface protein expression levels by fusing pairwise distance matrices with an $L_\infty$ norm. On the basis of the integrative kernel, we developed a quantitative metric (purity score) to assess the purity of single-cell populations in both the transcriptome modality and the surface protein modality of CITE-seq data. Because the resulting matrix is also a distance matrix, subsequent downstream analyses can seamlessly utilize the protein- and transcriptome-linked views of single cells. By applying our integrated method to several public benchmark datasets, we demonstrated that LinQ-View is not only effective and efficient, but also superior on CITE-seq datasets with routine numbers of surface protein features, applicable to many research groups who focus on distinct areas of biology. We have made LinQ-View available on a public data repository for users worldwide at https://wilsonimmunologylab.github.io/LinQView.

RESULTS

LinQ-View combines single-cell mRNA and protein expression profiling data and enables multimodal downstream analysis

One possible representation of single-cell profiling data are based on pairwise comparisons of cells. Specifically, mRNA or protein expression profiles might be chosen as feature representations, and pairwise distances (or dissimilarities) can be calculated between all cells. The primary idea behind LinQ-View for CITE-seq is to calculate the dissimilarity representations for each single-cell profiling modality (i.e., mRNA and surface protein expression) and then combine the dissimilarities from each modality. We reason that combining dissimilarity representations might help to naturally emphasize different information from each modality and maximally capture and infer cell heterogeneity (Figure 1).

To calculate and integrate dissimilarity matrices, we first calculated pairwise Euclidean distances among cells from pre-processed gene or protein expression profiling data. To eliminate potential biases introduced from random noise in ADT signals, we applied a sigmoid filter to distances calculated from protein expression profiles. Second, because the feature scale and dimensions differ between gene and protein expression profiles, we rescaled each pairwise dissimilarity matrix by using a linear transformation to minimize the average sum of squared differences between the dissimilarity matrices. Finally, we integrated the dissimilarity matrices by using the $L_\infty$ norm. Broadly, this is equivalent to choosing the maximal distance between any pair of cells on the basis of either protein or gene expression. In conclusion, we simply maximize the cell-cell dissimilarity by using information from both single-cell mRNA and protein expression profiling data.

To quantitatively evaluate clustering quality, we developed a measurement to assess the cluster purity on both modalities of CITE-seq (Figure 1). Because the RNA modality is on the scale of the whole genome, whereas features of the ADT modality are highly variable and dataset dependent, we adopted an entropy-based metric (ROUGE score) for assessing the purity of single-cell populations by using transcriptome profiles from a previous study (Liu et al., 2020), and developed a quantitative metric for assessing the purity of surface protein profiles. In brief, our ADT scoring method defines the sum of the standard deviations of all ADT features as the diversity index of a group of cells, and quantifies the purity of a specific cluster by comparing its diversity index against the diversity index of the entire dataset (see STAR Methods for details). The value range of both RNA and ADT scores is between 0 and 1, and a higher score indicates higher purity. To quantify cluster purity for CITE-seq data with different numbers of ADT features, we designed an adjustable parameter termed the “rank” of the ADT score. The purpose of the rank is to avoid a few highly variable ADT features being masked by groups of much less variable ADT features. Further, we combined purity scores for multiple modalities using a geometric mean function for an overall assessment of clustering quality. The purity scoring for CITE-seq proposed in this study allows users to confidently evaluate the quality of their clusters and efficiently determine the best clustering algorithm for optimized resolution (Figure 1).

Finally, because the resulting cell-cell dissimilarity matrix is equivalent to a distance matrix, LinQ-View is compatible with most conventional clustering methods, for example, k-means, hierarchical clustering, Louvain, and Fuzzy c-means (Blondel et al., 2008; Butler et al., 2018; Dunn, 1973), and dimension reduction methods, for example, multidimensional scaling.
Sigmoid filtering prevents variations in a single dominant ADT feature from affecting clustering results in CITE-seq data with limited ADT features

Differing from measurements of RNA modality, which are on the whole-genome scale, ADT modalities are dependent upon the reagents, such as antibodies used in staining. Due to monetary considerations or a specific research focus, most CITE-seq datasets include only a few dataset-specific antibodies that researchers are interested in. Within datasets of this scale, the number of ADT features is usually much smaller than the number of RNA features. Therefore, variations in a single ADT feature have a much higher impact on the distance calculation than those in a single gene. This phenomenon could result in a failure to distinguish cells that have only a few genes differentially expressed when using both modalities and is particularly pronounced for CITE-seq datasets with fewer ADT features.

To demonstrate this phenomenon, we show that results generated directly from RNA- and ADT-distance matrices without a sigmoid filter failed to correctly distinguish memory and naive CD4⁺ T cell subsets (Figure 2A; Dataset 1). As there are only 10 ADT features measured in this dataset, variations in ADT features become dominant and mask the real pattern in RNA modality (a few differentially expressed genes). To eliminate such bias in ADT features, we introduced a sigmoid filter into the distance calculation of the ADT modality. In brief, we added a penalty to the difference of each ADT feature between any cell pair to reduce the influence of variation of each single ADT feature. The penalty is a sigmoid function, which keeps larger distances essentially constant while further reducing smaller distances. As a result, the sigmoid function filters out minor differences from ADT modalities. Results using the same dataset showed that the LinQ-View model with the sigmoid filter successfully distinguished two CD4⁺ T cell subsets (Figure 2B). We found that, without the sigmoid filter, the majority of the distance between these two CD4⁺ T cell subsets is contributed by the ADT modality (ADT, 71.2%; RNA, 28.8%), whereas with the sigmoid filter, the distance between these two CD4⁺ T subsets is more influenced by the RNA modality (ADT, 43.6%; RNA, 56.4%) (Figure 2C). These results demonstrate that the sigmoid filter is able to correctly distinguish the two CD4⁺ T subsets that have a minor difference in gene expression without being affected by variation in ADT features. Effectively, the sigmoid filter reduces the weights of small distances from the ADT modality, minimizing the interference from variations of a few ADT features.
CITE-seq purity scoring accurately quantifies the purity of cell clusters

We further tested the effectiveness of our purity scoring method in quantifying cluster purity on both real and simulated datasets. Given that the effectiveness of ROGUE scoring has been demonstrated in a previous study (Liu et al., 2020), we focused on a variance-model-based ADT scoring method. First, we investigated the performance of the ADT scoring method under different ranks by using a real CITE-seq dataset (Dataset 1). Results showed that ADT scores with a proper rank (rank = 2 or 3) can balance the highly variable groups (e.g., clusters 12, 15, 17) in the context of less variable groups, and accurately quantify the differences among groups with different variation levels; ADT scores with too high of a rank (rank = 4, 5) emphasize the difference between highly variable groups (e.g., cluster 12, 15, 17) and other groups, and ignores the difference among mid- to less variable groups (e.g., clusters 0–4, 7–10, 18–22); ADT scores with a rank that is too low (rank = 1) fail to emphasize the overall variation because of the lowest value range (Figures 3A and 3B).

Taken together, these results demonstrate that ADT scoring...
Figure 3. Performance of ADT scoring on both real and simulated datasets
(A) ADT scores under different ranks.
(B) Distributions of ADT scores under different ranks shown by boxplot.

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can highlight a few highly variable groups from other groups, whereas lower ranked ADT scores are less sensitive to variability and generate scores with only minor differences between maximum and minimum scores.

To further investigate ADT scoring, we generated a simulated dataset to demonstrate its performance on clusters with (1) different variation levels and (2) different feature sizes. The simulated dataset comprises six clusters and nine features (Figure 3C). We designed the six clusters with different variation levels to better test the effectiveness of ADT scoring on quantifying ADT variation. Briefly, cluster 1 has low variation on all features, cluster 3 has high variation on one feature; and clusters 2, 4, 5, and 6 have moderate variation on one, two, three, or four features, respectively (see STAR Methods for details). Using this approach, an effective ADT scoring system should be able to accurately quantify the different variation levels of all clusters. Our results showed that under all test ranks, our ADT scoring method was able to accurately quantify the variation level of six clusters: ADT scores decreased in order of the number of variation features per cluster: clusters 1, 2, 4, 5, and 6 (Figures 3D and 3E). The ADT scores of cluster 3 (high variation on one feature) were always lower than those of cluster 2 (mid variation on one feature). We also observed that the higher rank model preferred to highlight variation in a single feature (rank = 5, cluster 3), enabling sensitive detection from a large number of features (Figure 3D).

To demonstrate the ability of ADT scoring to capture variation in single or a few features among a large number of features, we added 9 and 18 negative features with low variation to the existing simulated dataset to generate two new simulated datasets (see STAR Methods for details). We then applied ADT scoring to the two new datasets and compared results among the three simulated datasets. Low-ranked ADT scores failed to accurately quantify the variations of clusters from a high number of features (Figure 3F). These results suggest a higher rank is required by datasets with more ADT features: rank = 2 and 3 performed best on the 9 feature dataset, rank = 3 and 4 performed best on the 18 feature dataset, and rank = 4 and 5 performed best on the 27 feature dataset. In conclusion, testing ADT scores on both real and simulated datasets suggested that ADT scoring is able to accurately quantify the variation level of clusters, and the rank of the ADT score should be increased as the number of feature numbers increase. Furthermore, LinQ-View uses a geometric mean to combine the purity scores of two modalities for an overall assessment of cluster purity.

**LinQ-view efficiently reveals NK and CD4+ T cell subpopulations**

Clustering and visualization of scRNA-seq data can help to identify rare and intermediate subpopulations. We sought to visualize and evaluate cell clustering by using LinQ-View on a public CITE-seq benchmark dataset (Dataset 1, see STAR Methods for details). Visualization of LinQ-View-processed samples revealed all modality-exclusive cell clusters compared with unimodal transcriptome- or surface protein-based analyses. Using unimodal analyses, we could distinguish two subgroups of natural killer (NK) cells (CD8− NK and CD8+ NK) only by ADT-based methods and two subgroups of CD4+ T cells (memory and naive CD4+ T cells) only by RNA-based methods (Figure 4A). In particular, the two ADT-exclusive NK cell subsets had identical expression levels of the CD8A gene but distinct expression levels of CD8 protein (Figures 4B and 4C), and therefore could be distinguished only by using information from protein expression. The two RNA-exclusive CD4+ T cells had slight differences in transcriptome expression and exhibited no difference on 10 tested surface proteins (Figure 4D), and could be distinguished only by using transcriptome expression. However, all four modality-exclusive subsets were resolved with LinQ-View (Figure 4A).

Using the CITE-seq purity score proposed in this study, we also evaluated the quality of LinQ-View clustering results, along with two unimodal clustering results. We found that both LinQ-View and RNA clustering achieved high RNA scores (ROGUE score), whereas ADT clustering achieved lower RNA scores (Figure 4E). Furthermore, several ADT clusters (ADT clusters 6, 9, 13, and 17) had a very low ROGUE score (<0.6), indicating that the gene expression profiles within those clusters were highly diverse. Comparison of ADT scores showed that ADT clustering achieved the highest ADT purity score, followed by LinQ-View clustering, and RNA clustering obtained the lowest score (Figure 4E). More specifically, purity scores of the NK cell subset identified by RNA clustering (RNA cluster 4) were 0.9598 (RNA) and 0.6903 (ADT), whereas purity scores of two NK cell subsets identified by LinQ-View (LinQ-View clusters 5 and 6) were 0.9553 and 0.9434 (RNA) and 0.7822 and 0.8279 (ADT). The purity scores of these NK cell subsets showed that LinQ-View distinguished NK cells into two subsets with high RNA purity scores and increased ADT scores at the same time. ADT clustering performed poorly in the purity test because there were several clusters with diverse gene expression (ADT clusters 6, 9, 13, and 17). Furthermore, we compared LinQ-View with unimodal methods on several public CITE-seq datasets (Figures S1 and S2).

**Batch effects correction for multimodal data by using LinQ-View**

One of the most challenging aspects of analyzing scRNA-seq data are the processing of multiple samples that have been produced in different batches. To evaluate the compatibility of LinQ-View with previously developed batch effects correction methods, we sought to demonstrate seamless integration with batch correction methods by performing LinQ-View analysis on two datasets of peripheral blood mononuclear cells (PBMCs) from healthy donors (Dataset 2, PBMC 1K, and Dataset 3, PBMC 10K). We performed multimodal analysis on two independent runs: one without batch effects correction as a negative control. Batch effects correction processed samples revealed all modality-exclusive cell clusters compared with unimodal transcriptome- or surface protein-based analyses. Using unimodal analyses, we could distinguish two subgroups of natural killer (NK) cells (CD8− NK and CD8+ NK) only by ADT-based methods and two subgroups of CD4+ T cells (memory and naive CD4+ T cells) only by RNA-based methods (Figure 4A). In particular, the two ADT-exclusive NK cell subsets had identical expression levels of the CD8A gene but distinct expression levels of CD8 protein (Figures 4B and 4C), and therefore could be distinguished only by using information from protein expression. The two RNA-exclusive CD4+ T cells had slight differences in transcriptome expression and exhibited no difference on 10 tested surface proteins (Figure 4D), and could be distinguished only by using transcriptome expression. However, all four modality-exclusive subsets were resolved with LinQ-View (Figure 4A).

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A

RNA cluster

Naive CD4+ T

Memory CD4+ T

NK

CD8- NK

CD8+ NK

CD4+ T

ADT cluster

LinQ-View cluster

Naive CD4+ T

Memory CD4+ T

NK

CD8- NK

CD8+ NK

CD4+ T

B

CD3

CD4

CD8

CD45RA

CD56

CD16

CD11c

CD14

CD19

CD34

RNA Cluster

LinQ-View Cluster

CD8+ NK

CD8- NK

C

CD8 (RNA)

NCAM1 (RNA)

CD56 (ADT)

CD3G (RNA)

CD8A (RNA)

CD3 (ADT)

D

Expression level

CD3

CD4

CD8

CD45RA

CD56

CD16

CD11c

CD14

CD19

CD34

E

Purity score

RNA cluster

ADT cluster

LinQ-View cluster

RNA score

ADT score

Clustering

RNA score (ROGUE)

ADT score

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Comparison of LinQ-View with existing multimodal methods in cell heterogeneity identification

To evaluate the effectiveness and robustness of LinQ-View, we compared the use of LinQ-View dissimilarity matrices with three representative published (or pre-released) methods: SNF, MOFA2, and Seurat 4. We compared two unimodal and four multimodal methods on our benchmark dataset. The results of the two unimodal methods, labeled as “RNA cluster” and “ADT cluster,” were used as baselines to reveal two CD4+ T cell subsets and two NK cell subsets on t-SNE embedding. The four multimodal clustering methods, LinQ-View, SNF, MOFA2, and Seurat 4, are labeled as “LinQ-View cluster,” “SNF cluster,” “MOFA2 cluster,” and “Seurat 4 cluster,” respectively. Using this approach, we demonstrated that LinQ-View and Seurat 4 correctly identified two NK cell subsets and the other two multimodal methods failed to separate any of these subsets at the same resolution (Figure S3A). Notably, MOFA2 successfully distinguished the two NK cell subsets at a higher resolution of 1.5 (Figure S3A). The SNF method failed to correctly separate any of these subgroups at increased resolution (Figure S3B).

We then assessed the cluster quality of different methods by using purity scores (Figures 6B and S3C). Clusters generated by LinQ-View, Seurat 4, and MOFA2 had a higher RNA score than those generated by SNF. MOFA2 (resolution = 0.9), Seurat 4, and SNF all generated one cluster with very low RNA scores, indicating improper clustering. For ADT purity, all methods achieved high ADT scores, and Seurat 4 and LinQ-View achieved the highest. Focusing on two NK cell subsets, all three methods achieved similar RNA scores, whereas LinQ-View and Seurat 4 achieved higher ADT scores than MOFA2 (Figure S3D and Table S1), indicating that LinQ-View, MOFA2, and Seurat 4 are capable of capturing unique patterns of ADT modality. Furthermore, we assessed the quality of subsets of CD4+ T cells generated by these methods. LinQ-View, Seurat 4, and MOFA2 (at resolution = 1.5) identified two, five, and four clusters from CD4+ T cells, respectively (Table S2). Purity test results showed that all methods obtained high overall RNA and ADT scores, whereas Seurat 4 generated a cluster with very low RNA and ADT scores (Seurat cluster 10, Figure 6C). In addition to cluster purity, a further investigation of ADT and RNA expression profiles for these CD4+ T cell subsets generated by three multimodal methods demonstrated that LinQ-View correctly identified naive and memory CD4+ T cell subsets, whereas MOFA2 and Seurat 4 generated a few clusters with similar RNA and ADT expression profiles (Figures 6D and 6E, Tables S3 and S4, see STAR Methods for details), suggesting LinQ-View reveals more accurate clusters for specific cell subsets. Furthermore, we also applied the comparison onto multiple public CITE-seq datasets to demonstrate the generalizability of the LinQ-View method (Figures S4 and S5, see STAR Methods for details).

By comparing the time efficiency of these methods, we demonstrated that LinQ-View and Seurat 4 had a much higher time efficiency than MOFA2, SNF, and CiteFuse (see STAR Methods for details). To better understand the limitations and application range of LinQ-View, we then tested LinQ-View on multiple single-cell datasets with different sizes (Tables S5 and S6). Results suggested that LinQ-View is capable of handling large but not massive single-cell datasets, and the best application range for LinQ-View is between 0 and 50K cells with fewer than 50 ADT features. This application is ideal for users who are specialized in studying distinct cell populations, such as B cells or T cells, which present limited numbers of biologically relevant ADT features.

LinQ-View identifies unique B cell subsets from SARS-CoV-2-infected patients

We have shown the effectiveness of LinQ-View in distinguishing major cell populations (e.g., T cells, B cells, and NK cells) on several public datasets (Figures 4, S1, S2, S4, and S5). To further demonstrate the utility of LinQ-View, we applied this pipeline to peripheral blood CD19+ B cells from 10 SARS-CoV-2-infected COVID-19 patients (Dataset 6). LinQ-View successfully identified several unique B cell subsets with similar transcriptional expression but distinct profiles for the five tested surface proteins. RNA clusters 0 and 1 (indicated by dashed circle with label “a”), two naive-like B cell subsets, were further divided into four subsets by LinQ-View: they are LinQ-View clusters 0, 3, 4, and 5 (Figure 7A). LinQ-View cluster 0 is negative to all tested ADTs (non-responder group), cluster 3 is positive to all tested ADTs (polyreactive group), cluster 4 represents an HLA-DR+ group, and cluster 5 is CD62L+ (Figure 7B). These four subsets are indistinguishable on RNA-based t-SNE because they have highly similar transcriptional expression. In addition, RNA cluster 2 (indicated by dashed circles with label “b”) was divided into two subsets: LinQ-View cluster 2, which is HLA-DR+, and cluster 10, which is HLA-DR0 (Figure 7B). These two subsets were also
indistinguishable by transcriptional expression (Figure 7A). Furthermore, LinQ-View identified three unique B cell subsets with distinct ADT profiles (LinQ-View clusters 1, 8, and 11) from RNA clusters 3, 5, and 6 (Figure 7A, indicated by dashed circles with label “c”). LinQ-View cluster 1 was negative to all tested ADTs, cluster 8 was positive for all tested ADTs, and cluster 11 was HLA-DR+ (Figure 7B). As indicated by the dashed circles in t-SNE embeddings, cells in group a (LinQ-View clusters 0, 3, 4, and 5), group b (LinQ-View clusters 2 and 10), and group c (LinQ-View clusters 1, 8, and 11) were mixed together on RNA t-SNE and could not be distinguished by using RNA only, even with higher resolution.
Figure 6. Comparison of cell heterogeneity identification among LinQ-View, MOFA2, Seurat 4, and SNF methods

We compared the LinQ-View method with three multimodal methods, MOFA2, Seurat 4, and SNF, on a CITE-seq dataset. The resolution for Louvain clustering was set to 0.9 if there is no indication.

(A) Clustering using two unimodal and four multimodal methods. Two CD4+ T cell subgroups and two NK cell subgroups were highlighted in the t-SNE embedding.

(B) Comparison of purity scores among four multimodal clustering methods.

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Among these unique B cell subsets identified by LinQ-View, clusters 4, 10, and 11 exhibited HLA-DR protein expression and cluster 5 exhibited CD62L protein expression, distinguishing them from transcriptionally similar subsets, which indicates significant phenotypic differences. Notably, some B cells expressed B cell receptors that non-specifically bound to multiple antigens. Clusters 3 and 8 exhibited strong binding to all tested ADTs and COVID-19 probes, suggesting that these two clusters might be composed of polyreactive B cells with non-specific binding features. Polyreactivity has been identified as a common feature of the antigen-specific B cell repertoire, and is defined by an antibody’s ability to bind multiple self and foreign antigens, likely because of increased conformational flexibility (Mouquet and Nussenzweig, 2012). We verified the polyreactive nature of a sampling of monoclonal antibodies (mAbs) cloned from cells that exhibited binding to multiple unrelated COVID-19 probes by performing a highly standardized polyreactivity ELISA (Guthmiller et al., 2020). We found that over half of all sampled mAbs were indeed polyreactive (Figure S6A). We next confirmed whether cells exhibiting non-specific binding were reactive to the streptavidin-PE-oligo (SA-PE-oligo) used to conjugate the COVID-19 probes. Ninety percent of sampled mAbs also reacted to SA-PE-oligo, confirming the non-specific nature of these cells (Figure S6B). Given that clusters 3 and 8 displayed normal percentages of mitochondrial gene expression and similar transcriptional expression profiles compared with other B cells, they were indistinguishable for conventional transcriptome-based unimodal methods (Figures 7C and 7D). However, due to the non-specific nature of these cells and our inability to assess whether and how they participate meaningfully in an immune response, we concluded that these clusters should be excluded from analysis. Future studies are warranted to investigate the role of polyreactive B cells in antibody responses to COVID-19.

By investigating immunoglobulin repertoires from these subsets, we identified additional differences on the basis of variable (V) gene locus, isotype, and degree of somatic hypermutation (SHM), which could further discern the identity of these B cell subsets (Figure 7E). Clusters 0, 2, 3, 4, 5, 10, 12, and 14 had the lowest median number of SHMs (0) and highest IgM/D proportion (>95%) and are thus most likely to be naive-like B cells. In contrast, clusters 6, 7, and 9 displayed an intermediate number of mutations and increased class switching to IgA/G (20%–70%) and so are most likely class-switched memory B cells from more recent immune responses to the SARS-CoV-2 proteins (Figure 7E). Clusters 1, 8, 11, and 13 displayed an increased median number of mutations and increased IgA/G proportion (30%–80%) and are most likely longer-lived memory B cells cross-reacting to the SARS-CoV-2 proteins or that had reacted early in the immune response and so have become more terminally differentiated. We also clustered B cells into six “probe hit” groups according to their COVID-19 antigen probe binding signals by using an existing approach (Dugan et al., 2021). These results revealed that spike-specific B cells were enriched in three memory-like B cell subsets (clusters 6, 7, and 9), with an increased number of SHMs, further supporting the notion that clusters 6, 7, and 9 are derived from a recent immune response to the SARS-CoV-2 proteins. Through this analysis, we were able to detect functional heterogeneity in the B cells that react to SARS-CoV-2 antigens after COVID-19 infection. Studies in our group are ongoing to better understand this functional heterogeneity (Dugan et al., 2021).

**DISCUSSION**

In this study, we use a simple and intuitive approach based on pairwise distances to represent variations between cells in multiple modalities. By applying LinQ-View to several CITE-seq datasets, we prove the effectiveness of harnessing variations in pairwise distances for maximally inferring cell heterogeneity. To assess the purity of single-cell populations using surface protein expression, we proposed a variance-based purity metric and demonstrated that this ADT scoring was capable of accurately quantifying the purity of single-cell populations on the basis of their ADT variability, and thus could be used to determine the best clustering algorithms for optimal resolution. We provided an empirical formula for users to choose the best rank for their data according to the number of ADT features found in their datasets. Notably, the geometric mean function utilized in our scoring method can be easily extended to datasets with more modalities in the future.

Compared with other multimodal methods, LinQ-View has unique advantages when analyzing CITE-seq data with a routine number (e.g., fewer than 50) of ADT features. Although the latest technology is now capable of measuring up to 228 ADT features in a single experiment, and it is likely that this number will grow in the near future, CITE-seq performed on less than 30,000 cells with a medium number of ADT (e.g., 20–50) will still be a valuable tool, especially for groups with smaller budgets or focusing on specific cell populations. As advances in CITE-seq technology significantly lower the barrier of entry, allowing more laboratories to adopt CITE-seq to investigate their research questions and verify their hypotheses, entry-level CITE-seq will be able to satisfy their research needs. Furthermore, many laboratories focus on only specific cell populations, for example, B cells or T cells, which are relatively small in number and have few associated ADT features of interest. Therefore, a small panel of ADT features of interest is more economical than a full panel of all ADT features for such research topics. Considering both performance and time efficiency, Seurat 4 is the best option for massive CITE-seq datasets with a high number of ADT features, and LinQ-View is the best option on large datasets with a limited number of ADT features (e.g., <50). In conclusion, LinQ-View provides an effective and efficient option, especially for CITE-seq data with limited ADT features for the R community.
The ultimate goal of single-cell methods is to elucidate both temporal and spatial aspects of cells, including spatial distribution, growth history, and interactions with other cells/molecules. In the future it is likely that more modalities will be accessed within a single experiment. To date, six modalities, including transcriptome, surface protein, T cell receptor (TCR) γδ, TCR αβ, sample identity from cell hashtag, and sample identity from single-guide RNA, can be identified by the expanded CRISPR-compatible CITE-seq for single immune cells (Mimitou et al., 2019). However, such multimodal profiling technologies pose new challenges for integrative computational analysis. A robust and extendable system is critical for handling the growing number of modalities. LinQ-View captures variations among cells by pairwise distances, which can be readily extended to multiple new modalities.

Limitations of study
The integration model in LinQ-View calculates distances between all pairs of cells, and then combines them into an integrated distance matrix. Because the memory usage and complexity of pairwise distance calculation grow quadratically as the size of datasets increases, the current design of LinQ-View has a limitation in time efficiency with increasing cell numbers. We have tested the time efficiency of LinQ-View on multiple datasets and have suggested the best application range of this method.

STAR★METHODS
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SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.crmeth.2021.100056.

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AUTHOR CONTRIBUTIONS
L.L. designed the model, implemented the package, performed computational analyses of single-cell data, and wrote the manuscript. H.L.D. and C.S. collected samples, designed and performed experiments, and wrote the manuscript. L.Y-L.L. performed experiments and revised the manuscript. N.W.A. generated VDJ, 5’transcriptome, and feature libraries; performed Illumina sequencing; and revised the manuscript. M.K. helped in model design and revised the manuscript. N.-Y.Z. collected samples, expressed recombinant SARS-CoV-2 proteins, and generated mAbs. O.S. performed serum ELISAs and assisted in mAb generation. M.L.M., M.O.J., and K.S. coordinated the convalescent COVID-19 clinical study and collected patient samples. S.C. collected samples, performed ELISAs, and expressed recombinant SARS-CoV-2 proteins. H.A.U. collected samples and expressed recombinant SARS-CoV-2 proteins. C.H. revised the manuscript. C.N., R.P.J., D.F., A.J., and F.K. provided recombinant SARS-CoV-2 proteins. J.H. provided funding and resources for N.A. to perform sequencing and revised the manuscript. A.A.K. and P.C.W. supervised the work and wrote the manuscript.

DECLARATION OF INTERESTS
The authors declare no competing interests.
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KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Software and algorithms | R v3.6.3 | The R Foundation | https://www.r-project.org/ |
| RStudio v1.2.1335 | RStudio | https://www.rstudio.com/ |
| LinQ-View v0.99 | This paper | https://wilsonimmunologylab.github.io/LinQView/ |
| Seurat v3.9.9 | Hao et al., 2020 | https://satijalab.org/seurat/ |
| dplyr v1.0.2 | Hadley Wickham | https://dplyr.tidyverse.org/ |
| ggplot2 v3.3.2 | Hadley Wickham | https://ggplot2.tidyverse.org/ |
| Umap v0.2.6.0 | McInnes et al., 2018 | https://github.com/lmcinnes/umap |
| Rtsne v0.15 | Van Der Maaten, 2014 | https://github.com/jkrijthe/Rtsne |
| ROGUE | Liu et al., 2020 | https://github.com/PaulingLiu/ROGUE |

RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources should be directed to and will be fulfilled by the Lead Contact, Patrick C. Wilson (wilsonp@uchicago.edu).

Materials availability
This study did not generate new unique reagents.

Code availability
LinQ-View is implemented in R and freely available from the Wilson Lab GitHub (https://github.com/WilsonImmunologyLab/LinQView). Documents and tutorials are available from GitHub pages (https://wilsonimmunologylab.github.io/LinQView).

METHOD DETAILS

Datasets
Human multimodal datasets were obtained from public resources and previously published reports (Datasets 1-6).

Dataset 1: a dataset of 8,617 cord blood mononuclear cells (CBMCs), produced with CITE-seq (Stoeckius et al., 2017). NCBI GEO accession number is GSE100866. Notably, three ADT features were excluded in downstream analysis (CCR5, CCR7, CD10).

Dataset 2: a dataset of 713 peripheral blood mononuclear cells (PBMCs) from a healthy donor stained with TotalSeq-B antibodies. Data is available from 10x Genomics https://support.10xgenomics.com/single-cell-gene-expression/datasets/3.0.0/pbmc_1k_protein_v3.

Dataset 3: a dataset of 7,865 peripheral blood mononuclear cells (PBMCs) from a healthy donor stained with TotalSeq-B antibodies. Data is available from 10x Genomics https://support.10xgenomics.com/single-cell-gene-expression/datasets/3.0.0/pbmc_1k_protein_v3.

Dataset 4: a dataset of 8,412 cells from a dissociated Extranodal Marginal Zone B-Cell Tumor (MALT: Mucosa-Associated Lymphoid Tissue) stained with TotalSeq-B antibodies. Data is available from 10x Genomics https://support.10xgenomics.com/single-cell-gene-expression/datasets/3.0.0/malt_10k_protein_v3.

Dataset 5: a dataset of 5,247 peripheral blood mononuclear cells (PBMCs) from a healthy donor stained with TotalSeq-B antibodies. Data is available from 10x Genomics https://support.10xgenomics.com/single-cell-gene-expression/datasets/3.1.0/5k_pbmc_protein_v3.

Dataset 6: ten datasets of peripheral blood CD19+ B cells that were collected from SARS-CoV-2-infected subjects and stained with TotalSeq-C antibodies. The data are from clinical trial NCT04340050 and registered at ClinicalTrials.gov. Data has been published with a prior study (Dugan et al., 2021) and is available from Mendeley data: https://data.mendeley.com/datasets/3jdyw5jrjv/3. Subjects used in this study: S116, S166, S24, S376, S48, S414, S171, S305, S469, and S92. We measured the expression of five cell surface proteins and binding to seven COVID-19 protein probes by flow cytometry: spike protein, spike receptor binding domain (RBD), open reading frames 7a and 8 (ORF7a, ORF8), nucleoprotein (NP) and non-structural proteins (NSPs),
B cell immunoglobulin genes, as well as 5’ gene expression in the same immune cells simultaneously. Methodology for antigen probe generation and B cell sorting is described previously (Dugan et al., 2021). We used a pre-titrated cell staining panel that contained oligo-conjugated antibodies against CD62L, CD27, CD73, HLA-DR, and CD80.

Simulated datasets for ADT modality
To better assess the effectiveness of the ADT score proposed in this study, we generated three simulated datasets for ADT modality.

Simulated dataset 1
Simulated dataset 1 is composed of 9 features and 6 clusters. We generated random numbers by following a norm distribution to simulate normalized ADT data. We control the mean and standard deviation (SD) for each cluster and feature to generate data points for clusters with different variation levels. We define SD = 0.5 as low variance, SD = 1 as mid variance, and SD = 1.5 as high variance. We use absolute values in case of negative values. We set the number of cells (data points) for cluster 1–6 to 100, 50, 200, 50, 100, and 100 respectively. Means and SDs for each cluster and each feature have been listed in Table S7.

Simulated dataset 2
Simulated dataset 2 is based on simulated dataset 1. We simply added 9 negative features with low variance (Feature 1 of simulated dataset 1) to simulated dataset 1 to increase the number of features without changing the variance level of all clusters.

Simulated dataset 3
Simulated dataset 3 is based on simulated dataset 1. We simply added 18 negative features with low variance (Feature 1 of simulated dataset 1) to simulated dataset 1 to increase the number of features without changing the variance level of all clusters.

Pipeline design
The LinQ-View pipeline is composed of two major stages: pre-processing and joint analysis. The pre-process workflow was inspired by Seurat and other prior studies (Butler et al., 2018; Stuart et al., 2019; Wolf et al., 2018). As shown in Figure 1, pre-processing steps include cell quality control, unwanted gene removal (optional), data normalization (for both transcriptome and surface protein data), data scaling (for both transcriptome and surface protein data), highly variable gene (HVG) identification, linear dimension reduction (principal component analysis, PCA), and determination of the number of principal components (PC). Next, joint analysis was performed, such as calculating cell-cell pair-wise distances for each modality, joint distance calculation, cell population clustering, non-linear dimension reduction and visualization. The default cell population clustering method in LinQ-View is a community detection algorithm implemented in Seurat, and the LinQ-View toolkit is compatible with other clustering algorithms as well. Similarly, for non-linear dimension reduction, in addition to two default algorithms (t-SNE and UMAP), users may use any algorithm. Furthermore, other modalities, e.g. TCR, BCR and cell hashtags, can be applied to the identified cell populations to investigate feature (gene and protein) enrichment or feature-cluster correlation.

Cell quality control
Cell quality control removes all unwanted cells, such as dying cells, cycling cells and cell doublets from the current analysis. Two commonly used cell quality indicators are the number of detected genes and proportion of mitochondrial genes. Too few detected genes may indicate poor library or sequencing quality, whereas too many detected genes may indicate cycling cells or multiplets. In this study, all cells that have less than 200 genes detected or more than 2500 genes detected have been excluded. In addition, a high proportion of mitochondrial genes may indicate poor library or sequencing quality, whereas too many detected genes may indicate cycling cells or multiplets. In this study, all cells that have less than 200 genes detected or more than 2500 genes detected have been excluded. In addition, a high proportion of mitochondrial genes may indicate dying cells. Determining an appropriate threshold for number of detected genes is easy, but determining the threshold for proportion of mitochondrial genes is relatively difficult. The threshold for the number of detected genes will vary for different single cell techniques due to differences in sequencing depth and sensitivity; as a result, previous studies have summarized empirical values for different techniques (Wang et al., 2019). For the threshold proportion of mitochondrial genes, it is standard to set an arbitrary low threshold to filter out all unwanted cells (e.g. 5% for 10X data). The fixed threshold works for most cases but may remove too many cells in some specific cases. Here, we propose a strategy called soft threshold to determine the threshold of the proportion of mitochondrial genes. A soft threshold was set to the 95th percentile of the current dataset distribution, and the soft threshold was subject to a sealing point as the maximum threshold in the case of particularly poor cell quality. Of note, a recent study suggested that the proportion of mitochondrial genes may vary for different single-cell techniques (Wang et al., 2019). Its results showed that proportions of mitochondrial genes from Smart-seq/Smart-seq2 data (TPM) are much higher (in average 4-fold) than those from 10x data. In this study, a 10% sealing was applied to all datasets.

Integrating transcriptome and cell surface proteins
In this study, we performed joint analysis by integrating cell-cell pairwise distance matrixes from transcriptome expression and surface protein signals. The cell-cell pairwise distance matrices were calculated separately for each modality (in this instance, transcriptome expression and surface protein signals). Because the number of ADT features (usually from 10–50) is typically much less than the number of RNA features (usually 1000–2000 highly variable genes), variants in a single ADT feature have a much higher impact on the distance calculation than those in a single gene. This phenomenon may result in a failure to distinguish some cell groups that only have a few genes differentially expressed (e.g. memory and naïve CD4+ T cells) when using both modalities, because the random noise in ADT features can mask the signal pattern in so few differentially expressed genes. To eliminate the potential bias in ADT features, we introduced a sigmoid filter into the distance calculation of the ADT modality.
We define the sigmoid filter as:

\[ f(x) = \frac{1}{1 + e^{-n(x-k)}} \]

where \( x \) denotes the original distances, \( n \) and \( k \) are hyperparameters of sigmoid function, and \( e \) is Euler’s number (we set \( e = 2.72 \) in practice). Then the filtered distances \( F(x) \) will be

\[ F(x) = xf(x) = \frac{x}{1 + e^{-n(x-k)}} \]

and the original equation for Euclidian distance is defined as:

\[ D_e(P, Q) = \|P - Q\|_F = \sqrt{\sum_{i=1}^{N} (p_i - q_i)^2} \]

Here, \( P = (p_1, p_2, p_3, \ldots, p_N) \) and \( Q = (q_1, q_2, q_3, \ldots, q_N) \) are vectors of two cells \( P \) and \( Q \); \( p_i \) and \( q_i \) are the normalized value of the \( i \)-th ADT feature of cell \( P \) and \( Q \); and \( N \) is the number of ADT features; and \( \| \cdot \|_F \) denotes the Frobenius Norm. The sigmoid filtered Euclidian distance \( D_{e(\text{sigmoid})}(P, Q) \) is defined as:

\[ D_{e(\text{sigmoid})}(P, Q) = \|F(P - Q)\|_F = \|P - Q\|f(P - Q)\|_F \]

\[ = \sqrt{\sum_{i=1}^{N} (f(p_i) - f(q_i))^2} = \sqrt{\sum_{i=1}^{N} \left( \frac{|p_i - q_i|}{1 + e^{-n(p_i-q_i)}} \right)^2} \]

After testing with real data and simulated data, the suggested values of parameters \( n \) and \( k \) were set to \( n = 10 \) and \( k = 0.5 \). The sigmoid filter is able to eliminate the potential bias from random noise in ADT data. Some sub-cell groups with only a few differentially expressed genes, e.g. memory and naïve CD4+ T cells, could be identified by joint analysis after ADT data was scaled by the sigmoid filter (Figure 2).

In addition, after investigating the distribution of each ADT signal, we proposed a more precise way to distinguish small distances between negative and positive signals (most likely to be accurate distances) from those within negative/positive (most likely due to noise). Similar to the way users can define positive/negative in flow cytometry, users can define a threshold between negative and positive signals (most likely to be accurate distances) from those within negative/positive (most likely due to noise). In conclusion, we applied a sigmoid filter in our distance calculations to reduce the impacts of minor differences of ADT in the joint model, avoiding potential bias from ADT noise.

After sigmoid filtered Euclidian distances are constructed for ADT data, then the scaled distances for RNA (\( D_{RNA}' \)) and ADT (\( D_{ADT}' \)) were calculated by the following equations:

\[ D_{RNA}' = \alpha D_{RNA} \]

\[ D_{ADT}' = (1 - \alpha) D_{ADT} \]

where \( D_{RNA} \) and \( D_{ADT} \) are Euclidian distance matrices calculated from transcriptome expression (RNA) and surface protein signal (ADT). We determine an optimized value of linear operator \( \alpha \) by solving the following convex problem:

\[ \min_{\alpha} \frac{1}{2m} \|\alpha D_{RNA} - (1 - \alpha) D_{ADT} \|_2 \quad \text{subject to } \alpha \in [0, 1] \]

where \( m = N(N-1)/2 \), and \( N \) is the number of cells. This convex optimization problem can be solved by a gradient descent algorithm.

After scaling two distances into the same level, we introduced an \( L_\infty \) norm model to calculate the joint distance:

\[ D_{joint,ij} = \|D_{RNA}'_{ij} \cdot D_{ADT}'_{ij}\|_\infty = \max \left( D_{RNA}'_{ij}, D_{ADT}'_{ij} \right), \quad i,j \subset C \]

where \( D_{RNA}'_{ij} \cdot D_{ADT}'_{ij} \cdot D_{ADT}'_{ij} \cdot D_{ADT}'_{ij} \cdot D_{ADT}'_{ij} \cdot D_{ADT}'_{ij} \cdot D_{ADT}'_{ij} \cdot D_{ADT}'_{ij} \) are joint distance, scaled RNA distance, scaled ADT distance between cell \( i \) and \( j \), respectively. \( \| \cdot \|_\infty \) denotes \( L_\infty \) norm. \( C \) is the set of all cells.

In addition, the contributions of each modality can be quantified using an \( L_\infty \) norm model. The contribution of each modality was roughly quantified by the ratio of the number of distances from this modality to the total number of distances.
ContributionRNA = \frac{m - \| D_{\text{joint}} - D'_{\text{RNA}} \|_0}{m} × 100\%

ContributionADT = \frac{m - \| D_{\text{joint}} - D'_{\text{ADT}} \|_0}{m} × 100\%

Here \( D_{\text{joint}}, D'_{\text{RNA}}, D'_{\text{ADT}} \) are the lower triangles of the \( N \times N \) matrix, so the total number of distances is \( m = N(N - 1)/2 \). \( \| \cdot \|_0 \) denotes \( L_0 \) norm, which is equal to the number of non-zero elements.

Of note, by comparing with other conventional norm models (e.g. an \( L_1 \) norm model and \( L_2 \) norm model), we find that the \( L_\infty \) norm model generates more accurate and robust results than any other norm models. In addition, since distances from multiple modalities were scaled into the same level, too few ADT numbers can cause high individual feature weights, resulting in potential errors from ADT random noise. The \( L_\infty \) model is able to overcome this challenge, which is difficult for \( L_1 \) or \( L_2 \) norm model. A comparison between the \( L_\infty \) model and the \( L_1 \) model demonstrated that the \( L_\infty \) model correctly distinguished memory and naïve CD4 T cell subsets whereas the \( L_1 \) model failed. After that, cell clustering and non-linear dimension reduction based on joint distances will be able to represent variations in both transcriptome expression and surface protein expression. In summary, the \( L_\infty \) norm model takes valuable information for cell heterogeneity identification from both modalities, and is able to avoid potential bias in distance integration caused by too few ADT numbers.

**PURITY SCORE OF CELL CLUSTERS FOR CITE-SEQ DATA**

To comprehensively assess the purity of multi-modal cell subsets, we developed two purity scores, and , for both modalities in CITE-seq data. In detail, we adopted ROGUE score from a previous study to assess the purity of single cell populations on transcriptome modality, and developed a quantitive metric to assess the purity of single cell populations on surface protein modality. As demonstrated in a previous study, the ROGUE score is able to accurately quantify the purity of single cell populations using transcriptome data. For a given cell population, the ROGUE score calculates the entropy reduction (ds) of each gene using an expression entropy model, and then quantifies the purity of this cluster by a statistic measurement:

\[ S_{\text{RNA}} = \text{ROGUE} = 1 - \frac{\sum_{\text{sig}} ds}{\sum_{\text{sig}} ds + K} \]

\( ds \) denotes entropy reduction of each gene; \( \sum_{\text{sig}} ds \) denotes the summary of significant \( ds \). \( K \) is a pre-defined parameter and is dependent on single cell sequencing technology.

Distinct from transcriptome modality, which is at the whole genome scale level, the number of features in the surface protein modality of CITE-seq is determined by users and is usually limited, disabling the application of the ROGUE score on the surface protein modality of CITE-seq. We developed a score to access the purity of single cell populations on the surface protein modality by evaluating the variation level among all ADT features of given populations. We firstly calculate the summary of standard deviations among all ADT features of entire dataset as a benchmark. Then we assess the purity of each clusters in this dataset by calculating:

\[ S_{\text{ADT}} = 1 - \frac{\sum SD}{\sum SD^2 + K_{\text{ADT}}} \]

\( SD \) denotes standard deviation of each ADT feature for a given group; \( n \) denotes a non-negative (\( n = 1,2,3,\ldots \)) rank that usually goes higher along the raise of number of ADT features and can be chosen by users. An empirical formula (floor of natural logarithm of ADT feature number could be helpful in choosing rank: rank=\( \text{floor}(\ln(N)) \), \( N \) is the number of ADT features. \( K_{\text{ADT}} \) denotes the summary of standard deviations among all ADT features of entire dataset using the same rank. It is the benchmark variation level of the entire dataset. In this way, \( S_{\text{ADT}} \) for a population will be between 0 and 1. Similar to \( S_{\text{RNA}} \), a higher score indicates better purity. \( S_{\text{ADT}} \) equals 0.5 if the variation level of a cluster is the same as that of the entire dataset.

We also developed a geometric mean function to combine the two purity scores for an overall assessment of cluster purity. Notably, this metric is also expandable to datasets with multiple modalities in the future by simply increase the rank to number of modalities of new datasets.

\[ S = \sqrt{S_{\text{ADT}}S_{\text{RNA}}} \]

In practice, users can calculate and compare purity scores for multiple clustering results on the same dataset in order to determine the best clustering algorithm and optimized resolution.

**Parameter setting in model comparison**

We performed clustering analysis using the same algorithm (Louvain clustering, implemented in the Seurat package) under the same resolution (0.9) for all 4 multimodal methods. For easier comparison and identification of cell subsets across different methods, we visualized the results of all six methods on the same t-SNE embedding (LinQ-View t-SNE). Method-specific clustering and
visualization of three multimodal methods can be found in the supplemental information (Figure S7). This dataset was pre-processed by following conventional pre-processing procedures: we performed log normalization for RNA data and CLR normalization for ADT data, selected 2000 highly variable genes for RNA data, performed PCA analysis for RNA data and selected the top 20 PCs for further analysis. Notably, three ADT features were excluded from this analysis (CCR5, CCR7, CD10). More specifically, for the LinQ-View method, Louvain clustering was performed on a joint distance matrix. The joint distance matrix was generated under a default parameter setting. For the SNF model, we set the number of neighbors (K) to 20, a hyperparameter alpha to 0.5 and the number of iterations (T) to 10 as suggested in SNF tutorial. For MOFA2, we adopted default model parameter settings using a “get_default_model_options” function and set the number of factors to 15. The mofa2 model was trained on 2000 highly variable genes and all ten ADT features. Then we ran the MOFA2 model using a “run_mofa” function and the model converged in 155 iterations. For Seurat 4, the number of PCs for the RNA modality was set to 20 and the number of PCs for the ADT modality was set to 5.

**Time efficiency test**
All time efficiency tests were performed on a 2019 version iMac with 3.1 GHz Intel Core i5 processor and 64 GB 2400 MHz DDR4 memory, with operating system version macOS Mojave (10.14.6). R environment version is 3.6.3 (for LinQ-View and SNF) and 4.0.3 (for MOFA2 and Seurat 4).

**Data pre-processing**
Pre-processing steps were adapted from Seurat (Butler et al., 2018; Stuart et al., 2019). Transcriptome data was processed by cell quality control, unwanted gene removal (optional), data normalization, data scaling, highly variable gene (HVG) identification, linear dimension reduction (PCA) and determination of the number of PC. Surface protein data was processed by data normalization and data scaling.

**ADT and RNA expression for CD4+ T cell subsets generated by three multimodal methods on Dataset 1**
We investigated the ADT and RNA expression profiles for the CD4+ T cell subsets generated by three multimodal methods. Differences among subsets generated by MOFA2 and Seurat4 on CD3, CD4, CD45RA and CD14 were not significant or had a low fold change in expression difference, and these subsets showed no difference in corresponding genes (Figure 6D). This indicates that the CD4+ T cell subsets generated by MOFA2 and Seurat 4 are most likely being compromised by small variability in the expression of a few ADT features rather than real gene or protein expression patterns. An investigation of the expression of 10 differently expressed genes between naïve and memory CD4+ T cells showed that the subsets identified by MOFA2 and Seurat4 show no difference in these gene markers (Figure 6E). Furthermore, the gene markers identified from 5 Seurat clusters were highly similar, suggesting that there is no significant difference among these subsets in gene expression (Tables S3 and S4). Taken together, single dominant ADT features are commonly found in existing CITE-seq data which have a limited number of ADT features (i.e., <50). Interference from the ADT modality having an adverse effect on the results is common with existing methods, because these methods generate ADT variances directly from normalized ADT counts or PCA results of ADT modality without any further processing. Our results show that the sigmoid filter and model we proposed in this paper works well on CITE-seq data with limited ADT features by preventing variations in a single dominant ADT feature from impacting clustering results.

**Comparison of LinQ-View with existing multi-omics methods on multiple CITE-seq datasets**
To demonstrate the generalization of LinQ-View method on CITE-seq data, especially CITE-seq data with routine numbers of surface protein features, we further compared LinQ-View with two commonly used multimodal methods: MOFA2 and Seurat 4. In this comparison, we applied all three methods onto two public benchmark CITE-seq datasets generated by 10X genomics (Dataset 2 and 3). All clustering was performed using the Louvain algorithm under the same resolution (0.6 for dataset 2, 0.8 for dataset 3). Purity score rank for ADT modality was set to 3 for both datasets. For easier comparison and locating cell subsets across different methods, we visualized the results of all three methods on the same t-SNE embedding (LinQ-View t-SNE). Parameter settings for all three methods are as same as described in “Parameter setting in model comparison” section.

Using this approach, LinQ-View, MOFA2, and Seurat 4 identified 7, 7, and 8 cell clusters from dataset 2, respectively (Figure S4A). Purity scores of cell clusters generated by three methods showed that all three methods achieved high overall purity scores (Figure S4B). MOFA2 achieved highest RNA score but lowest ADT score, and Seurat 4 achieved highest ADT score but lowest RNA score, indicating the imbalance weighting between ADT and RNA modalities. More specifically, MOFA2 weights RNA modality higher than ADT modality, and Seurat 4 weights ADT modality higher than RNA modality on tested dataset. Notably, besides these statistical differences, there are non-trivial differences among the clustering results of the three methods. For easier demonstration, we labeled two groups of cells using red and blue dashed line and label “a” and “b” (Figure S4A). For group “a” cells, LinQ-View divided them into four clusters (cluster 1, 2, 4, and 5), MOFA2 divided them into three clusters (cluster 0, 2, and 5), and Seurat 4 divided them into five clusters (cluster 1, 2, 4, 6, and 7). A further investigation of surface protein marker expression demonstrated that LinQ-View clusters have distinct surface protein expression patterns: cluster 1 is CD3+CD4+CD45RO+, cluster 2 is CD3+CD45RO+, cluster 4 is CD3+CD4+CD45RA+, and cluster 5 is CD16+CD56+CD45RA+ (Figure S4C). Seurat 4 generated similar results as LinQ-View among cell group “a”. However, there is a minor difference between LinQ-View clusters 1 and 4 and Seurat 4 clusters 2 and 4. Surface protein expression of these clusters showed that LinQ-View clustering is better because LinQ-View cluster 1 and 4 has much more
distinct expression of CD45RA than Seurat 4 cluster 2 and 4. Furthermore, MOFA2 failed to detect the difference of surface protein expression among these cells and generated three clusters with similar surface protein expression patterns. For group “b” cells, Seurat 4 grouped those cells as cluster 6 whereas LinQ-View identified the difference from those cells and clustered them into clusters 5 and 6 (Figure S4A). A further investigation of surface protein and gene marker expression demonstrated that LinQ-View clusters 5 and 6 have distinct protein expression (e.g. CD56, and CD4) and gene expression (e.g. GNLY, NKG7, LST1, and AIF1) (Figure S4D). Notably, MOFA2 also clustered these cells into two individual clusters (clusters 5 and 6), despite that MOFA2 cluster 5 also improperly included some cells from other clusters.

On dataset 3, LinQ-View, MOFA2, and Seurat 4 identified 26, 19, and 19 cell clusters, respectively (Figure S5A). Purity scores of cell clusters generated by three methods showed that all three methods achieved high overall purity scores (Figure S5B). Similarly to dataset 2, MOFA2 achieved highest RNA score but lowest ADT score, and Seurat 4 achieved highest ADT score but lowest RNA score, indicating the imbalance weighting between ADT and RNA modalities. Similarly, LinQ-View, MOFA2, and Seurat 4 also generated different cell clusters on some groups of cells. For easier demonstration, we labeled two groups of cells using red and blue dashed line and label “a” and “b” (Figure S5A). For group “a” cells, LinQ-View and Seurat 4 identified two clusters (LinQ-View cluster 8 and 9, Seurat 4 cluster 14 and 15). A further investigation of surface protein and gene marker expression demonstrated that LinQ-View clusters 8 and 9 have distinct expression profiles on surface protein CD25 and a few gene markers (e.g. TCL1A, IGHG1, YBX3, IGHG2, and S100A6) (Figure S5C). However, MOFA2 failed to detect those gene expression differences and clustered these cells into one cluster (MOFA2 cluster 5). For group “b” cells, LinQ-View clustered these cells into two clusters (LinQ-View cluster 1 and 11), MOFA2 clustered these cells into two clusters (MOFA2 cluster 3 and 8), and Seurat 4 clustered these cells into three clusters (Seurat 4 cluster 8, 9, and 11) (Figure S5D). A further investigation of surface protein and gene marker expression demonstrated that the two LinQ-View clusters (clusters 1 and 11) have distinct expression profiles on two surface protein markers (CD25 and CD127) and a few gene markers (e.g. ANXA1, IL7R, SELL, FOS). However, clusters of group “b” cells identified by Seurat 4 and MOFA2 have similar expression profiles on these surface protein and gene markers, indicating these two methods failed to properly integrate information from two modalities.

Taken together, comparison on datasets 2 and 3 demonstrated that all three methods are able to integrate multiple modalities of CITE-seq data and correctly identify major cell populations, moreover, LinQ-View is able to balance contribution between two modalities and reveal more accurate clusters with distinct protein and gene expression profiles compared to other two methods. Combined with the comparison on dataset 1 in the main text, we demonstrated that LinQ-View identified cell subsets accurately and consistently on multiple CITE-seq datasets, and proved the generalizability and applicability of LinQ-View method.

**Time efficiency test of existing multimodal methods**

We assessed the time efficiency of these multimodal methods. Using the same test dataset (8067 cells after quality control), SNF required 3–4 hours whereas the LinQ-View model ran in approximately 30 seconds. CitFuse (using the SNF model as kernel) took more than 8 hours, most likely due to the lack of dimension reduction before modality integration. Thus, in addition to poor performance in multimodal integration, time efficiency test results demonstrate that SNF and CitFuse are not compatible with large-scale single cell CITE-seq datasets. In addition, MOFA2 required 6 minutes to train the MOFA2 model, and Seurat 4 only took 20 seconds on finding multi-modal neighbors using the same dataset. These results demonstrated that LinQ-View and Seurat 4 not only performed better in integrating RNA and ADT modalities, but also had a much better time efficiency than MOFA2. However, because distances were computed between all pairs of cells, the model design of LinQ-View has a limitation on time efficiency with increasing cell number. For example, Seurat 4 is able to handle a massive single cell data set of 170K cells with 228 ADT features whereas all of the other existing methods (including LinQ-View) failed. Taken together, time efficiency test results suggested Seurat 4 has the highest efficiency, especially with massive datasets (>50K cells); LinQ-View is efficient on large size datasets (<50K cells); MOFA2 is about 10 times slower than the previous two methods and is able to handle medium size datasets (<20K cells) in a reasonable timeframe; and the SNF model and CitFuse are too time-consuming to process even medium size datasets.

Notably, we observed that the time efficiency of MOFA2 is highly variable and dataset-dependent when applied MOFA2 on multiple CITE-seq datasets. Runtime of model training step of MOFA2 on datasets 1, 2, 3, 4, and 5 are 385.48, 12.59, 49.17, 59.22, and 72.67 seconds, respectively. Considering dataset 1 (8067 cells, 10 ADTs), dataset 3 (7865 cells, 17 ADTs) and dataset 4 (8412 cells, 17 ADTs) are similarly sized, our results suggest that time efficiency of MOFA2 is highly dataset-dependent.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

The boxplots and medians were generated using an R package “ggplot2” (Figures 3B, 3E, 4E, 6B, 6C, and 7E). The upper and bottom edges of the box indicate the 25th percentile (Q1) and 75th percentile (Q3), respectively. The range between Q1 and Q3 is defined as the interquartile range (IQR). The horizontal bar in the box indicates the median (Q2). The top edge of the upper vertical line indicates the maximum (Q3 + 1.5IQR) and the bottom edge of the lower vertical line indicates the minimum (Q1-1.5IQR). The violin plots were generated using an R package “Seurat” (Figures 4C, 4D, 6D, and 6E). We identified differentially expressed genes and proteins between two groups of cells using a Wilcoxon Rank Sum test implemented in “FindAllMarkers” function in an R package “Seurat”.