Integration of spatial and single-cell data across modalities with weak linkage

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Abstract: single-cell sequencing methods have enabled the pro- 44 filing of multiple types of molecular readouts at cellular resolu-2 tion, and recent developments in spatial barcoding, in situ hy-3 bridization, and in situ sequencing allow such molecular readл outs to retain their spatial context. Since no technology can 5 provide complete characterization across all layers of biological modalities within the same cell, there is pervasive need for com-50 putational cross-modal integration (also called diagonal integration) of single-cell and spatial omics data. For current methods, the feasibility of cross-modal integration relies on the existence 52 10 of highly correlated, a priori "linked" features. When such 53 11 linked features are few or uninformative, a scenario that we call 54 12 13 "weak linkage", existing methods fail. We developed MaxFuse, 55 a cross-modal data integration method that, through iterative 56 14 co-embedding, data smoothing, and cell matching, leverages all 57 15 information in each modality to obtain high-quality integra-16 tion. MaxFuse is modality-agnostic and, through comprehen-17 sive benchmarks on single-cell and spatial ground-truth multi-18 ome datasets, demonstrates high robustness and accuracy in the 19 61 weak linkage scenario. A prototypical example of weak linkage 20 is the integration of spatial proteomic data with single-cell se-21 quencing data. On two example analyses of this type, we demon-22 strate how MaxFuse enables the spatial consolidation of pro- 64 23 teomic, transcriptomic and epigenomic information at single- 65 24 cell resolution on the same tissue section. 25

Single cell | Multi-modal integration | Diagonal integration | Matching | Multi omics | Spatial-omics | Protein | RNA | ATAC

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30 Introduction

Recent technological advances have enabled the profiling of 74 31 multiple biological modalities within individual cells, over 75 32 many cells in parallel. The growing list of modalities that 76 33 can now be profiled at the single-cell level include proteome 77 34 and metabolome (1, 2), transcriptome (3), and various as-78 35 pect of the epigenome such as methylation (4), histone mod- 79 ification (5-7), and chromatin accessibility (5, 8). In addi- 80 37 tion to technologies that operate on dissociated single cells, 81 38 rapid progress has been made on the in situ measurement of 82 39 transcriptome (9), proteome (10–14), epigenome (15), and 83 other modalities on histological tissue sections at single-cell 84 41 or close to single-cell resolution, retaining the spatial context. 85 42 These advances have spawned consortia-level efforts to con-86 43

struct multiomic single-cell and spatial atlases of each and every organ, across species, in healthy and diseased states.

To harness the new technologies and growing data resources for biological discovery, a primary challenge is the reliable integration of data across modalities. Cross-modal integration, also referred to as "diagonal integration" (16, 17), is the alignment of single cells or spatial spots across datasets where different features (or modalities) are profiled in each dataset. An example is the alignment of cells in a CODEX dataset, which measures protein abundance, to cells in a single-cell RNA sequencing (scRNA-seq) dataset, which measures RNA expression. This cross-modal integration step underpins many types of downstream analyses, and its importance is evident in the myriad methods that have already been developed to tackle it (18–24).

Despite the progress in this area, key limitations still hinder reliable cross-modal integration, as highlighted by recent surveys (16, 17, 25). A key factor limiting the accuracy of existing methods is the strength of linkage between modalities, as we define below. A feature is "linked" between two modalities if it can be measured in, or predicted by, both modalities. In the terminology of (16, 17), these linked features can serve as "anchors" for the integration. For example, to integrate single-cell or spatial ATAC sequencing (ATAC-seq) and single-cell or spatial RNA-seq data, most existing methods predict the "activity" for each gene in each cell/spot of the ATAC-seq data based on the accessibility of the gene's surrounding chromatin; then, each gene's ATAC activity can be linked to its RNA expression, mapping cells from the two datasets into the same feature space. Similarly, between RNA and protein assays, the abundance of each protein in the protein assay can be linked to the expression of its coding gene in the RNA assay. With the exception of bindSC (26), all existing methods, to our knowledge, rely crucially on the linked features and are designed for scenarios where there is a large number of linked features that exhibit strong cross-modality correlation, a situation that we refer to as "strong linkage". For example, between scRNA-seq and scATAC-seq, every gene in the genome can be linked, and the correlation between gene activity and RNA expression is often high enough for enough genes to allow for precise integration (18, 19, 22). To achieve strong linkage, some methods attempt to learn a mapping from the features of one modality to the features of

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the other modality through a "training set" consisting of data 144
where both modalities are simultaneously observed in each 145
cell/spot (23, 27). While this strategy may be applicable to- 146
wards the integration of data from biological systems that are 147

⁹¹ similar to the training set, it is questionable how well it can 148

⁹² generalize to unseen systems.

Cross-modality integration in scenarios of weak linkage, 150 93 where the number of linked features is small and/or the 151 94 between-modality correlation for the linked features is weak, 152 95 is especially challenging. A prototypical example of weak 153 96 linkage is between targeted protein assays (14, 28) and 154 97 transcriptome/epigenome assays such as scRNA-seq and 155 98 scATAC-seq. Such scenarios are becoming extremely com- 156 99 mon as spatial proteomic technologies are receiving wide- 157 100 spread adoption (10–14), complementing RNA and ATAC se- 158 101 quencing in achieving more complete tissue characterization 159 102 (see, for example, (29-32)). We will reveal, through compre- 160 103 hensive benchmarks, the limitations of existing state-of-the-161 104 art methods in such difficult cases. 105

Under both strong and weak linkage, the evaluation of ex- 162 106 isting methods have leaned heavily on systems with highly 107 distinct cell types whose separation only requires a crude 108 feature-level mapping between modalities. In fact, most ex-109 isting methods explicitly focus on the goal of "label transfer", 110 that is, the transfer of cell type labels from one modality to the 111 other. This goal only requires the integration to be accurate at 112 the resolution of the label. As we demonstrate in our bench-113 marks, even this seemingly modest goal of label transfer for $\frac{1}{170}$ 114 major cell types is unattainable in weak linkage scenarios by 174 115 current methods, much less the more challenging goal of in-116 tegration in continuously transitioning cell populations where 117 subtle distinctions need to be preserved between closely re-118 lated states. Yet, key biological discoveries often hinge on 175 119 the accurate preservation of fine cell state distinctions during 120 integration, 121 177

To address the above limitations, we developed MaxFuse 178 122 (MAtching X-modality via FUzzy Smoothed Embedding), a 179 123 model free, highly adaptive method that can accurately inte-124 grate data across weakly linked modalities. MaxFuse goes 181 125 beyond label transfer and attempts to match cells to pre-182 126 cise positions on a graph-smoothed low-dimensional embed-183 127 ding. MaxFuse starts by denoising the linked features in each 184 128 modality through borrowing information from all of the fea- 185 129 tures, and then performs an initial crude matching of cells 186 130 based on the denoised linked features. Then, MaxFuse iter-187 131 atively refines the matching step based on graph smoothing, 188 132 linear assignment, and CCA. These iterations use informa-133 tion from all features in both modalities to improve upon the 190 134 initial matching. The initial feature linkage may be derived 191 135 from domain knowledge or an existing integration, and thus, 192 136 MaxFuse can also be used to improve upon any existing inte-193 137 gration methods. 138 194 We systematically benchmarked the performance of Max- 195 139

We systematically benchmarked the performance of Max-195
 Fuse across protein, RNA, and chromatin accessibility 196
 single-cell multiome ground-truth datasets. Across a wide 197
 variety of datasets, MaxFuse has superior performance com-198
 pared to other state-of-the-art integration methods. Although 199

the largest improvements in accuracy are observed under weak linkage, under strong linkage MaxFuse is comparable to the current best method in integration performance with substantial improvement in speed.

We further demonstrate the analyses enabled by MaxFuse with two examples. First, in the integration of scRNA-seq and CODEX multiplexed in situ protein profiling data from the human tonsil, we show that MaxFuse can recover correct spatial gradients in the RNA expression of genes not included in the 46-marker protein panel. Next, MaxFuse is applied to an atlas-level integration of spatial proteomic and single-cell sequencing datasets, as part of a consortium-level effort to map cell organization and function across different regions of the human intestine (32). We demonstrate how to perform trimodal integration of CODEX, snRNA-seq, and snATAC-seq data to recover spatial patterns of RNA expression and transcription factor binding site accessibility at single-cell resolution.

Results

Cross-modality matching of single cells via iterative fuzzy smoothed embedding. Let data from the two modalities be represented by a pair of cell-by-feature matrices that contain all measured features in each modality. For convenience, call the two modalities Y and Z. In addition, we represent the initial knowledge about the linkage between the two modalities as another pair of cell-by-feature matrices whose columns have one-to-one correspondences. To distinguish between these two pairs of matrices, we call the former all-feature matrices and the latter linked-feature matrices. For example, when one modality is protein abundance over a small antibody panel and the other is RNA expression over the whole transcriptome, the two all-feature matrices have drastically different numbers of columns, one being the number of proteins in the panel and the other being the number of genes in the transcriptome; the linked feature matrices, on the other hand, have equal number of columns. where each column in the protein matrix is one protein and its corresponding column in the RNA linked-feature matrix is the gene that codes for the protein. When the number of cells is large, we recommend aggregating cells with similar features into meta-cells, as described in Materials & Methods, prior to applying MaxFuse. In that case, each row in the above matrices would represent a meta-cell. The procedure below does not depend on whether single- or meta-cells are used, and thus we will refer to each row as a "cell". The two pairs of matrices form the input of the MaxFuse pipeline in Figure 1A.

Stage 1 of MaxFuse aims to summarize cell-cell similarity within each modality and learn an initial cross-modal matching of cells. As shown in Figure 1A, this stage consists of three major steps. In step 1, for each modality, we use all features to compute a fuzzy nearest-neighbor graph connecting all cells measured in that modality. This graph, by utilizing the information in all features, provides the best possible summary of the cell-cell similarity for the given modality. In particular, cells that are close in this graph should

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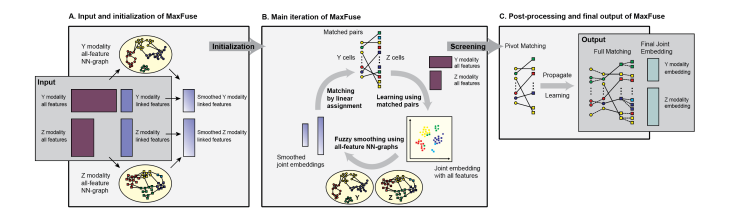


Figure 1: Overview of MaxFuse pipeline. (A) The input consists of two pairs of matrices. The first pair consists of all features from each modality, and the second pair consists of only the linked features. MaxFuse uses all features within each modality to create a nearest-neighbor graph (all-feature NN-graph) for cells in that modality. Fuzzy smoothing induced by the all-feature NN-graph is applied to the linked features in each modality. Cross-modal cell matching based on the smoothed linked features initializes the iterations in (B). (B) In each iteration, MaxFuse starts with a list of matched cell pairs. A cross-modal cell pair is called a pivot. MaxFuse learns CCA loadings over all features from both modalities based on these pivots. These CCA loadings allow the computation of CCA scores for each cell (including cells not in any pivot), which are used to obtain a joint embedding of all cells across both modalities. For each modality, the embedding coordinates then undergo fuzzy smoothing based on the modality-specific all-feature NN-graphs (obtained in (A)). The smoothed embedding coordinates are supplied to a linear assignment algorithm which produces an updated list of matched pairs to start the next iteration. (C) After iterations end, MaxFuse screens the final list of pivots to remove low-quality matches. The retained pairs are called refined pivots. Within each modality, any cell that is not part of a refined pivot is connected to its nearest neighbor that belongs to a refined pivot and is matched to the cell from the other modality in this pivot. This propagation step results in a full matching. MaxFuse further learns the final CCA loadings over all features from both modalities based on the refined pivots. The resulting CCA scores give the final joint embedding coordinates.

have comparable values for their linked features. Thus, in 232 200 step 2 of stage 1, MaxFuse boosts the signal-to-noise ratio in 233 201 the linked features within each modality by shrinking their 234 202 values, for each cell, towards the cell's graph-neighborhood 235 203 average. We call this step "fuzzy smoothing". After fuzzy 236 204 smoothing of linked features within each modality, MaxFuse 237 205 computes in step 3 distances between all cross-modal cell 238 206 pairs based on the smoothed linked features and applies linear 239 207 assignment (33) on the cross-modal pairwise distances to ob- 240 208 tain an initial matching of cells. The initial matching serves 241 209 as the starting point of stage 2 of MaxFuse. 210 242

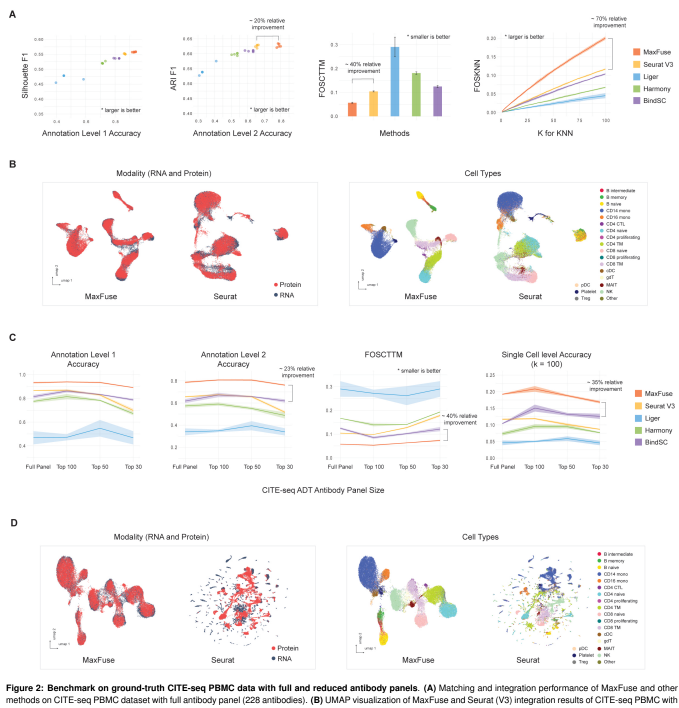
Stage 2 of MaxFuse, shown in Figure 1B, aims at improving 243 211 cross-modal cell matching quality by iterating the sequence $_{244}$ 212 of joint embedding, fuzzy smoothing, and linear assignment $_{_{245}}$ 213 steps. Starting with the initial matches obtained in stage 1, in $_{246}$ 214 each iteration, MaxFuse first learns a linear joint embedding 247 215 of cells across modalities by computing a canonical corre-248 216 lation based on all features of the cross-modal matched cell 249 217 pairs. Then, coordinates of this joint embedding are treated 250 218 as new linked features of each modality and fuzzy smoothing $_{251}$ 219 is applied on them based on the all-feature nearest-neighbor 220 graphs computed in stage 1. Finally, MaxFuse updates the 253 221 cell-matching across modalities by applying linear assign-254 222 ment on the pairwise distances of these fuzzy-smoothed joint 223 embedding coordinates. The resulting matching then starts $_{256}$ 224 the next iterate. Matching quality improves with each iter-225 ation until available information in all features, and not just 258 226 the linked features, have been used. 227 259

Stage 3 of MaxFuse aims at post-processing the last cross-260
modal cell matching from stage 2 and producing final out-261
puts. First, MaxFuse screens the matched pairs from the last 262
iterate in stage 2, retaining high quality matches as pivots. 263

The pivots are used in two complementary ways: (i) they are used one last time to compute a final joint embedding of all cells in both modalities; (ii) for any unmatched cell in either modality, its closest neighbor within the same modality that belongs to a pivot is identified and, as long as its distance to this neighbor is below a threshold, the match in the pivot is propagated to the cell. Thus, the final output of MaxFuse has two components: (i) a list of matched pairs across modalities, and (ii) a joint embedding of all cells in both modalities. More details on the MaxFuse algorithm are given in Materials & Methods.

Integration of transcriptome and targeted protein data with varying protein panel sizes. We benchmarked Max-Fuse on a CITE-seq dataset (34) containing simultaneous measurements of 228 protein markers and whole transcriptome on peripheral blood mononuclear cells. For comparison, we also applied four state-of-the-art integration methods: Seurat (V3) (24), Liger (22), Harmony (20), and BindSC (26) to this same dataset. Protein names were converted to RNA names manually to link the features between datasets. In each repetition of our experiment, we randomly subsampled 10,000 cells, applied all methods, and assessed using the benchmarking criteria to be described below. We performed 5 such repetitions and averaged the criteria across repetitions. We masked the known cell-cell matching between the protein and RNA modalities when applying all methods (treating Protein and RNA as two unpaired modalities), and then used the known matching for assessment.

Methods are assessed using six different criteria that measure both cell-type-level label transfer accuracy as well as cell-level matching accuracy. The first two criteria are based on label transfer accuracy. Cells are annotated at two levels



rigure 2: Bencimark on ground-truth CITE-seq PBMC data with full and reduced antibody panels. (A) Matching and integration performance of MaxFuse and other methods on CITE-seq PBMC dataset with full antibody panel (228 antibodies). (B) UMAP visualization of MaxFuse and Seurat (V3) integration results of CITE-seq PBMC dataset with reduced antibody panels. (D) UMAP visualization of MaxFuse and Seurat (V3) integration results of CITE-seq PBMC with the 30 most informative of the original 228 antibodies, colored by modality (left) or cell type (right).

of granularity: level-1, which differentiates between 8 ma- 274 264 jor cell types, and level-2, a finer classification which dif-275 265 ferentiates between 20 cell types. Label transfer accuracy is 276 266 expected to be higher for level-1 labels than for level-2 la- 277 267 bels. The proportion of matched pairs that share the same 278 268 label at both annotation levels are reported, with higher pro- 279 269 portions indicating higher matching quality. The next two 280 270 criteria measure the quality of the cross-modal joint embed-281 271 ding of cells. A high-quality joint embedding should preserve 282 272 biological signal, as reflected by the separation of known cell 283 273

types, while mixing the two modalities as uniformly as possible. Usually, there is a trade-off between biological signal preservation and uniformity of mixing. Thus, we report the F_1 scores computed based on average silhouette width (slt_f1) and adjusted Rand index (ari_f1), as proposed in Tran et al. (35). These scores aggregate quality assessments of biological signal preservation and modality mixing. For both criteria, higher F_1 indicates a better embedding. The fifth criterion is FOSCTTM, Fraction Of Samples Closer Than True Match (19, 36, 37), that quantifies the quality of joint embed-

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ding at single-cell resolution. For each cell, one can compute 341 284 the fraction of cells in the other modality that is closer than 342 285 its true match in the joint embedding. FOSCTTM is the av- 343 286 erage of this fraction over all cells in both modalities. The 344 287 lower this measure, the closer the true matches are in the joint 345 288 embedding, and hence, the better the joint embedding. The 346 289 last criterion is FOSKNN, Fraction Of Samples whose true 347 290 matches are among their <u>K-Nearest Neighbors</u> in the joint 348 291 embedding space. For any given $k \ge 1$, the higher this pro- 349 292 portion, the better the joint embedding. For precise defini- 350 293 tions and details of these criteria, see Materials & Methods. 351 294 Among all criteria described above, MaxFuse uniformly 352 295 dominates the methods by a sizable margin (Figure 2A). Im- 353 296 portantly, MaxFuse provides accurate cell matching across 354 297 weakly-linked modalities (level 1 accuracy 93.9%, $+ \sim 7\%$ 355 298 to the second best method, Figure S2B). The UMAP plots 356 299 calculated based on the post-integration embedding from re- 357 300 spective methods are shown in Figure 2B, colored by modal- 358 301 ity and by cell type. MaxFuse achieves both better mixing 350 302 of the two modalities (left panel) and better preservation of $_{360}$ 303 biological signals (right panel). For example, B cell subtypes 361 304 (B naive, intermediate, and memory cells) present a nicely 362 305 resolved developmental trajectory after MaxFuse integration, 363 306 but not after integration by other methods. 307 It is common to have an antibody panel that is of signif- $_{\scriptscriptstyle 365}$ 308 icantly smaller size than 228, especially for the emerging 366 309 spatial-proteomic datasets. To benchmark the performance 367 310 of MaxFuse against existing methods for smaller antibody 311 panels, we ordered the proteins according to their importance 312 for differentiating cell types (See Materials & Methods for ³⁶⁹ 313 370 details). We repeated the foregoing experiments when only 314 the top 100, 50, and 30 most important proteins are used ³⁷¹ 315 in the matching and integration process. At each antibody $^{\scriptscriptstyle 372}$ 316 panel size, we ran the experiment over five independent rep-373 317 etitions with randomly subsampled 10,000 cells, and aver-318 age the cell type annotation level matching accuracy, FOS-³⁷⁵ 319 CTTM and FOSKNN across repetitions (Figure 2C). Regard-³⁷⁶ 320 less of panel change, MaxFuse consistently outperformed 377 321 other methods. Additionally, MaxFuse successfully miti-378 322 gated the effect of reduced panel size on integration qual-379 323 ity: Even when the antibody panel size was reduced to 30, 380 324 MaxFuse maintained a > 90% annotation level 1 accuracy ³⁸¹ 325 while other methods produced variable and low quality cell 382 326 matching results ($\sim 10 - 70\%$, Figure S2B). Similarly, with ³⁸³ 327 a reduced antibody panel size (eg. 30 antibodies), the inte-³⁸⁴ 328 grated UMAP embedding (38) produced by other methods ³⁸⁵ 329 blurs the distinction between cell types, while MaxFuse em- 386 330 bedding still accurately captures the subtle structure of highly ³⁸⁷ 331 granular cell subtypes (e.g., the B cell subpopulations, Figure 388 332 389 2D and Figure S2A). 333

Systematic benchmark across multiple ground-truth ³⁹¹
 multiome modalities. We further benchmarked MaxFuse ³⁹²
 on four additional single-cell multiome datasets. The first ³⁹³
 is a CITE-seq dataset of human bone marrow mononuclear ³⁹⁴
 cells (BMMCs) that provides cell-matched measurements of ³⁹⁵
 the full transcriptome along with an antibody panel of size ³⁹⁶
 25 (34). The second is an ABseq dataset, also of BMMCs, ³⁹⁷

with an antibody panel of size 97 and the whole transcriptome (39). The third is an ASAP-seq PBMC dataset (40)with 227 antibodies and the whole epigenome measured in ATAC fragments. The fourth is a TEA-seq PBMC dataset (41) where we focused on the simultaneous measurements of 46 antibodies and the whole epigenome measured in ATAC fragments. Together, these datasets represent a diverse collection of measurement technologies over different modality pairs. We benchmarked the performance of MaxFuse against Seurat (V3), Liger, Harmony, and BindSC on these datasets. For datasets with simultaneous RNA and protein features, we linked each protein to its coding gene. For datasets with simultaneous ATAC and protein measurements, we linked each protein to the gene activity score (42) computed from the ATAC fragments mapping near its coding gene. As in the previous case, the known cell-cell correspondence across modalities were masked in the matching and integration stage for all methods, but used afterwards for evaluation.

We compared the performances of MaxFuse and the other four methods on these datasets using the collection of matching and integration quality measures described in the previous section (Figure 3A): cell type annotation matching accuracy, FOSCTTM, FOSKNN (*K* set as 1/200 dataset size), Silhouette F1 score, and ARI F1. Overall, MaxFuse outperformed other methods, often by a sizable margin (eg. ~ 20% relative improvement in terms of the metrics measured, Figure 3A and Figure S3.1A).

UMAPs of the MaxFuse cross-modal joint embeddings for each dataset are shown in Figure 3B, with the top row colored by modality and the bottom row colored by cell type annotation. Across the integration scenarios, MaxFuse mixed different modalities well in joint embeddings while retaining separation between cell types. Compared to the UMAPs of joint embeddings produced by other methods, MaxFuse consistently achieves substantial improvements (Figure 3B and Figure S3.2 A).

As a counterpoint to the above integration scenarios, we also considered the problem of integration of scRNA-seq and scATAC-seq data, on which multiple methods have demonstrated feasibility (18, 19, 22). The degree of overlap in the information contained in the RNA and ATAC modalities has been systematically measured in Lin and Zhang (43), where it was shown that, in terms of cell population structure, the information shared across RNA and ATAC is much higher than the information shared between RNA and protein for commonly used targeted protein panels. Thus, RNA and ATAC has stronger linkage and should be easier to integrate. We benchmarked MaxFuse against state-of-the-art methods for this problem on four public multiome datasets that simultaneously measure the chromatin accessibility and transcriptome expression for each cell: 10x mononuclear cells from peripheral blood (44), cells from embryonic mouse brain at day 18 postconception (44), cells from developing human cerebral cortex (45), and cells from human retina (46). The integration quality criteria described in the previous subsection are used to assess all methods, shown in Supplementary Materials. Across datasets and evaluation metrics, MaxFuse

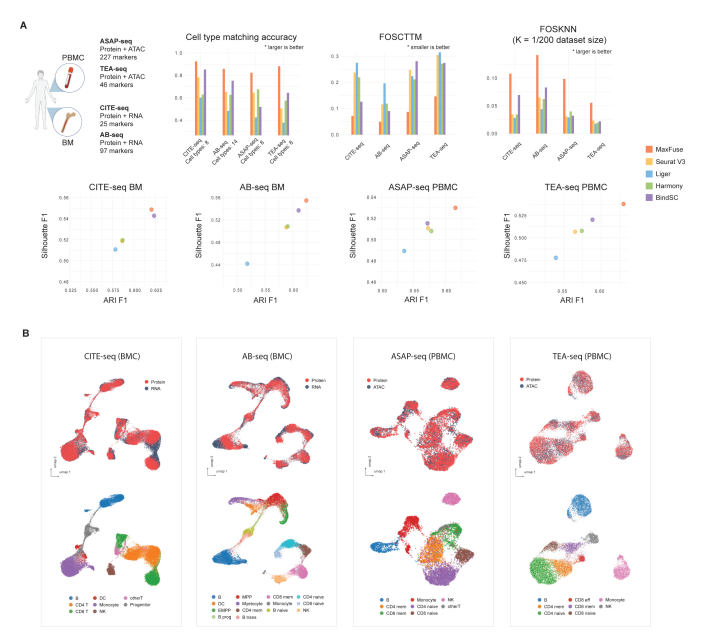


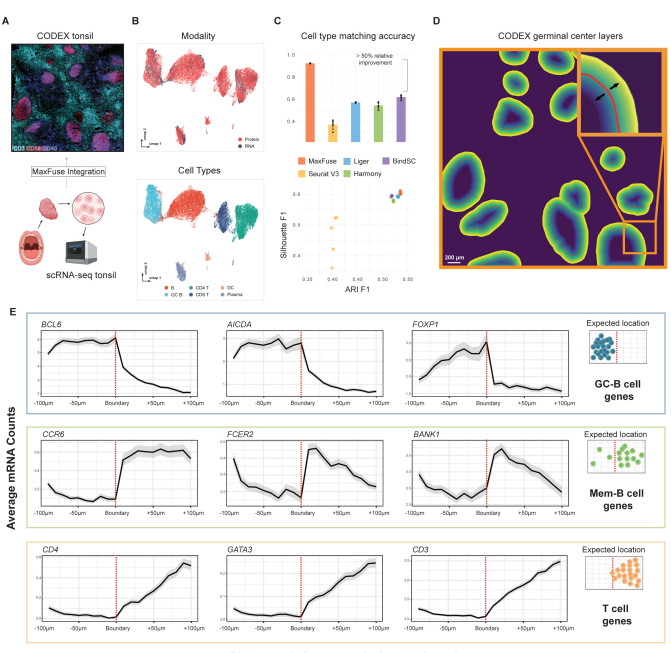
Figure 3: Systematic benchmark across multiple ground-truth data types with MaxFuse (A) Four different multiome datasets, generated by different technologies, were benchmarked. Cell type matching accuracy, FOSCTTM, FOSKNN (with K = 0.5% total cell counts of each dataset), and ARI and Silhouette F1 were evaluated across 5 methods. (B) UMAP visualization of MaxFuse integration results for the four ground-truth multiome datasets.

achieves best or close-to-best performance among methods, 413
and is comparable to scGLUE. However, MaxFuse is much 414
faster than scGLUE. For example, for the integration of a 415
dataset of 20,000 cells, MaxFuse took <5 minutes to finish 416
on a laptop with M1 Max chip while scGLUE took hours on 417
a comparable platform without CUDA acceleration. 418

Cross-modal integration of scRNA-seq and spatial 420 404 proteomic data enables information-rich spatial pat- 421 405 tern discovery. MaxFuse is particularly motivated by sce-422 406 narios where the signal-to-noise ratio in the cross-modal 423 407 linked features is low. Weak linkage is especially common 408 in spatial-omic data types due to technical limitations. For 424 409 example, high resolution spatial proteomic methods such as 425 410 CODEX, MIBI-TOF, IMC, and CosMx SMI can profile, at 426 411 sub-cellular resolution, a panel of 30-100 proteins (10-13). 427 412

Integration of such spatial proteomics datasets with singlecell transcriptomic and epigenomic datasets of the same tissue is often of interest, and particularly challenging due to the small number of markers in the spatial dataset and the weak linkage between modalities that is caused by both biological and technical differences. Thus, we demonstrated and benchmarked MaxFuse on the integration of CODEX multiplex imaging with 46 markers (47) with single-cell RNA-seq (48) of human tonsils from two separate studies (Figure 4A). Figure 4B shows the UMAPs of the MaxFuse integration colored by modality and by 6 major cell types.

Based on the pre-described benchmarking metrics, MaxFuse is the only method capable of integrating spatial proteomic and single-cell RNA-seq data. Existing state-of-the-art methods, Seurat (V3), Liger, Bindsc, and Harmony, failed to pro-



Distance relative to germinal center boundary

Figure 4: MaxFuse enables information-rich spatial pattern discovery (A) MaxFuse integrates human tonsil single-cell data: one dataset by CODEX from Kennedy-Darling et al (47) (upper panel), the other dataset by scRNA-seq from King et al (48) (lower panel). (B) UMAP visualization of MaxFuse integration of tonsil CODEX and scRNA-seq data, colored by modality (upper panel) and cell type (lower panel). (C) Metrics (cell type matching accuracy, Silhoutte F1 and ARI F1 score) evaluating performance of MaxFuse and other methods. Five batches of randomly sampled CODEX and scRNA-seq cells (total of 40k each batch) were sampled, and used for benchmarking for all methods. (D) Illustration of cell layers extending inwards/outwards from the germinal center boundary, with each layer consisting of 30 pixels ($\sim 11 \mu$ m). A total of 10 layers extending in each direction were examined. (E) For each of 9 genes, the average mRNA counts (linked by MaxFuse) across cells in each layer are plotted versus the position of the layer in reference to the germinal center boundary (inward on the left of boundary, outward on the right). For each group of 3 genes (row), their expected expression profile in reference to the germinal center boundary is shown on the right.

duce an embedding that integrates the two modalities while 485 428 preserving the cell population structure (Figure 4B and Fig-486 429 ure S4.1A). Evaluation results based on cell-type matching 487 430 accuracy is consistent with evaluation results based on the 488 431 joint embedding. At the level of the 6 major cell types shown 489 432 in Figure 4B, MaxFuse is able to achieve high label trans- 490 433 fer accuracy (93.3%), while the other methods fail to pre- $_{491}$ 434 serve cell type distinctions (40% - 60%, Figure 4B and Figure 492 435 S4.1B). 436 493

We further assessed whether MaxFuse can preserve, during 494 437 integration, the more subtle spatial variations within a cell 495 438 type that are captured by CODEX. We manually delineated 496 439 the boundaries of each individual germinal center (GC) from 497 440 the CODEX tonsil images based on CD19, CD21, Ki-67 pro- 498 441 tein expression patterns. From the boundaries, we then ex- 499 442 tended outward or inward, with each step covering roughly 500 443 one layer of cells (one step = 30 pixels erosion/dilation) (Fig- 501 444 ure 4C). Then, for each layer of cells, we calculated the av- 502 445 erage counts of specific genes, based on the scRNA-seq cells 503 446 that match to CODEX cells of that layer. We then asked if 504 447 known position-specific gene expression patterns relative to 505 448 the germinal center boundary are recovered in the integrated 506 449 scRNA-seq data. Indeed, MaxFuse was able to reconstruct 507 450 the spatial pattern of the GC from disassociated transcrip- 508 451 tomic data (Figure 4D): For GC-specific genes BCL6, AICDA 509 452 and FOXP1 (49-51) that relate to germinal center function- 510 453 ality, we observed high expression within the boundary and a 511 454 sharp drop in expression after passing the boundary layer; for 512 455 genes related to B cell memory CCR6, BANK1 and FCER2 513 456 (51–53) that should be enriched in B cells exiting from the 514 457 GC, we indeed saw a gradual increase outside of the GC and 515 458 then a quick decrease as the layer fully expands into the T cell 516 459 region; and finally for T cell related genes, for example CD4, 517 460 GATA3 and CD3 (54), we indeed saw a rapid increase out- 518 461 side of the GC boundary but no expression within. In com-519 462 parison, the integration with scRNA-seq produced by other 520 463 methods was incapable of accurately reconstructing the GC 521 464 spatial pattern (Figure S4.2A). 465

522 Tri-modal atlas-level integration of spatial and single-466 cell data with MaxFuse. In the consortium-level effort to 523 467 generate a comprehensive atlas across different regions of the 524 468 human intestine, colon and small bowel tissue from healthy 525 469 human donors were collected and systematically profiled by 526 470 CODEX, snRNA-seq, and snATAC-seq (32). We applied 527 471 MaxFuse to the integration of these three modalities (Fig- 528 472 ure 5A), with the goal of constructing high-resolution spatial 529 473 maps of full transcriptome RNA expression and transcrip- 530 474 tome factor binding accessibility. To perform tri-omic inte- 531 475 gration, we first conducted pairwise alignment of cells be- 532 476 tween protein (CODEX) and RNA (snRNA-seq), and cells 533 477 between RNA (snRNA-seq) and ATAC (snATAC-seq), as 534 478 previously described. The two sets of bi-modal cell-pairing 535 479 pivots were then "chained" together, with the pivot cells in 536 480 the RNA modality serving as the intermediary. This "chain- 537 481 ing" created a set of pivots linking all three modalities: Pro- 538 482 tein, RNA, and ATAC. Subsequently, we used these pivots to 539 483 calculate a tri-omic embedding via generalized CCA (gcca) 540 484

(21, 55). This allows for a joint UMAP embedding of the three modalities, shown in Figure 5B. We see that distinctions between major cell types are preserved and modalities are mixed within each cell type.

The MaxFuse integration produces, effectively, a joint profile of protein abundance, RNA expression, and chromatin accessibility at single-cell spatial resolution on the same tissue section. To confirm the post-integration consistency between the three modalities, we inspected whether CODEX's protein abundance aligns spatially with the expression and chromatin activity of the protein-coding gene, the spatial measurements of the latter two modalities imputed based on the MaxFuse integration. Figure 5C shows an example in CD163, a macrophage marker: The protein expression, RNA expression, and gene activity of CD163 are, as expected, uniquely enriched in the macrophage cell cluster (Figure 5C upper panel). Furthermore, protein, RNA, and ATAC activities of this gene all localize to the same spatial positions on the tissue section (Figure 5C lower panel). Other examples are shown in Supplementary Materials.

With the integration of the snATAC-seq and CODEX data, we can further map the spatial enrichment of transcription factor (TF) binding site accessibility. For each TF, this is achieved by first computing its motif enrichment score for each cell in the snATAC-seq data, and then the scores are transferred to the CODEX spatial positions based on the MaxFuse integration. Figure 5D shows such spatial profiles for 3 transcription factors: Binding motifs of IRF4, known to be a key regulator in immune cell differentiation (57), had increased accessibility in the immune-enriched compartments of the mucosa and submucosa layers (32). Binding motifs of KLF4, known to be required for the terminal differentiation of goblet cells (58), had heightened accessibility in the colonic crypts of the mucosa layer where goblet cells mature. Finally, binding motifs of SRF, a master regulator of smooth muscle gene expression. (59), had heightened accessibility in neighborhoods that are enriched for smooth muscle cells.

Discussion

In this paper, we conceptually separated cross-modal integration of single-cell data into two different scenarios: across modalities with strong linkage (e.g., ATAC-RNA integration) and across modalities with weak linkage (e.g., RNAprotein integration for a targeted protein panel). Most existing methods are developed for integration across strongly linked modalities, and our ground-truth benchmark results suggest that their performances decay significantly as the strength of cross-modal linkage weakens. MaxFuse is motivated by and focuses on the challenging case of weak linkage, which has become increasingly common as many emerging study designs include spatial data with targeted marker panels to be collected jointly with single-cell sequencing data. MaxFuse relies on two key ideas to overcome weak linkage: The first is a "fuzzy smoothing" procedure that denoises the linked features by moving their values towards their graph-

linked features by moving their values towards their graphsmoothed values, with the graph determined by all features. The second is an iterative refinement procedure that improves

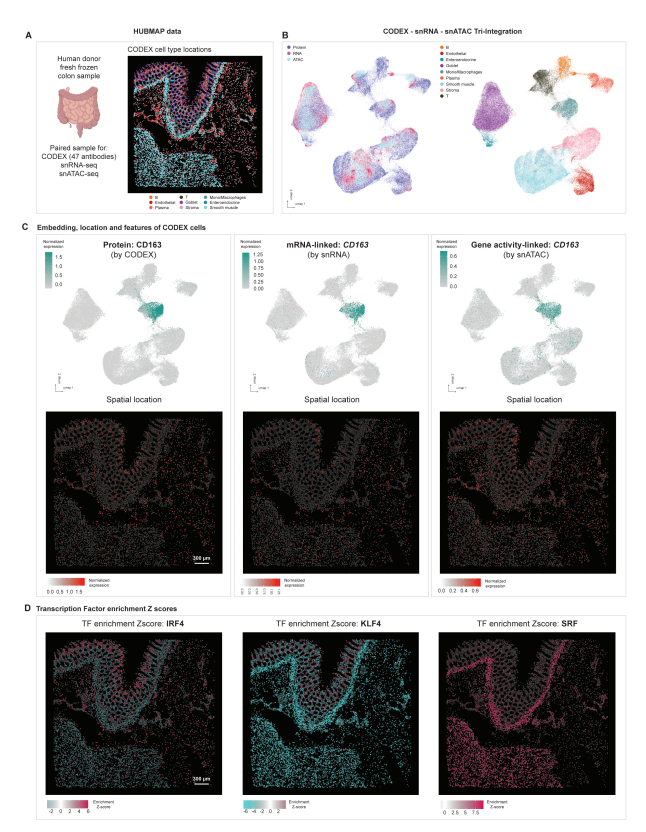


Figure 5: MaxFuse enables tri-modal integration with HUBMAP data (A) Overview of data: Patient paired CODEX, snRNA-seq, snATAC-seq single-cell human intestine data from HUBMAP consortium. Colon and small bowel data were integrated by MaxFuse respectively and this figure shows part of the colon data (CODEX data from one donor; snRNA-seq and snATAC-seq data from four donors). (B) UMAP visualization of the tri-modal integration embedding produced by MaxFuse, colored by modality: Protein, RNA and ATAC (left panel) and colored by cell type (right panel). (C) Upper panel: UMAP visualization of CODEX cells based on the integration embedding, overlaid with CD163 protein expression (from CODEX cells itself, left panel), *CD163* RNA expression (from matched snRNA-seq cells). Lower panel: Spatial location of CODEX cells based on their centroids' x-y position, overlaid with the same expression features as in the upper panel. (D) Spatial location of CODEX cells based on their centroids' x-y position factor motif enrichment score (Z-score, calculated by chromVAR (56)), based on their matched snATAC-seq cells.

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the cross-modal matching through an iterative cycle of co-596 541 embedding, graph-smoothing, and matching; this ensures 597 542 that information from all features, in both modalities, are 598 543 used to generate the final matching. We show that these key 599 544 ideas allow MaxFuse to substantially improve upon state-of- 600 545 the-art methods, achieving accurate integration of data from 601 546 targeted protein assays with data from transcriptome- and 602 547 epigenome-level assays. 548

While MaxFuse is motivated by the weak linkage scenario, 604 549 its applicability is universal. For strong linkage scenar-605 550 ios, methods based on deep learning, such as scGLUE, 606 551 achieve state-of-the-art integration performance but is hin-607 552 dered by high computational costs. In comparison, MaxFuse 608 553 achieves comparable performances as scGLUE on ground- 609 554 truth strong-linkage benchmark datasets at a considerably 610 555 lower computational cost. In addition, when joint embed-611 556 ding coordinates from other integration methods are avail-612 557 able, these coordinates could serve as linked features in Max-558 Fuse, which could then be further improved by the proce-559

dure. The light computation architecture and the flexibility ⁶¹⁴ in incorporating domain knowledge and existing integration ⁶¹⁵ results make the MaxFuse framework applicable to a wide ⁶¹⁶

range of cross-modal integration tasks.

564 Materials & Methods

565 The MaxFuse pipeline.

Input preparation Consider a pair of datasets $Y \in \mathbb{R}^{N_y \times p_y}$ 566 and $Z \in \mathbb{R}^{N_z \times p_z}$ from two modalities (termed Y-modality 567 and Z-modality for exposition convenience), with each row 568 corresponding to a cell and each column a feature. In the 569 ensuing discussion, we treat Y as the modality with a higher 570 signal-to-noise ratio. For concreteness, one can think of Y571 as a snRNAseq dataset and Z as a CODEX dataset. Suppose 572 625 there are two known functions $f_y: \mathbb{R}^{p_y} \to \mathbb{R}^s$ and $f_z: \mathbb{R}^{p_z} \to$ 573 \mathbb{R}^s such that $f_y(y)$ predicts the values of $f_z(z)$ in a cell if the 574 measured values under Y-modality are y in that cell and those 575 under Z-modality are z. For any matrix A with p_y columns, 628 576 let $f_u(A)$ denote the matrix with s columns and the same 629 577 number of rows as A, obtained from applying f_y on each row 630 578 of A and stacking the outputs as row vectors. For any matrix $_{631}$ 579 B with p_z columns, $f_z(B)$ is analogously defined. With $f_{y_{632}}$ 580 and f_z , we define $Y^\circ = f_y(Y) \in \mathbb{R}^{N_y \times s}$ and $Z^\circ = f_z(Z) \in \mathcal{I}_{fast}^{ooc}$ 581 $\mathbb{R}^{N_Z \times s}$. In the snRNAseq vs. CODEX example, if one has 582 a crude prediction for a subset S (with size |S| = s) of the 583 proteins then $f_z(z) = z_S$ returns the subvector indexed by S 584 while $f_y(y) = \hat{z}_S$ predicts the observed CODEX values for 585 these proteins based on transcriptomic information of a cell. 586 In summary, we start with a pair of original datasets (Y, Z)587 and a pair of datasets (Y°, Z°) with correspondence of 588 columns based on domain knowledge. 589

⁵⁹⁰ *Meta-cell construction* To alleviate sparsity and to scale to ⁶³⁵ ⁵⁹¹ large datasets, we start by constructing meta-cells. Take the ⁶³⁶ ⁵⁹² *Y*-modality for example. Let n_y be the desired number of ⁶³⁷ ⁵⁹³ meta-cells one aims for. We first construct a nearest-neighbor ⁶³⁸ ⁵⁹⁴ graph of the rows of *Y*, apply Leiden clustering with an ap- ⁶³⁹ ⁵⁹⁵ propriate resolution level to obtain n_y clusters, and average ⁶⁴⁰ over the rows within each cluster to obtain the features for each meta-cell that serves as the representative of the cluster. Consequently, we obtain $Y_m \in \mathbb{R}^{n_y \times p_y}$. Using this clustering structure (induced by Y as opposed to Y°), we can average feature vectors in Y° to obtain $Y^{\circ}_{m} \in \mathbb{R}^{n_{y} \times s}$. When desired, the same operation can be performed on the Z-modality to obtain $Z_{m} \in \mathbb{R}^{n_{z} \times p_{z}}$ and $Z_{m}^{\circ} \in \mathbb{R}^{n_{z} \times s}$. We recommend only constructing meta-cells for modalities with high signal-toratios. For example, if Y-modality contains snRNAseq data and Z-modality contains CODEX data, then we would construct meta-cells only in Y-modality. After this curation step, we have two pairs of datasets (Y_m, Z_m) and (Y_m°, Z_m°) . The former pair can have completely distinct feature sets, while the latter pair must have matching feature sets with corresponding columns. In Figure 1A, the former correspond to the pair of all feature matrices, and the latter correspond to the pair of linked feature matrices.

Fuzzy smoothing Let $G_Y \in \{0,1\}^{n_y \times n_y}$ be a nearest neighbor graph of Y_m where each row *i* is connected to k_i^Y rows that are closest in a chosen similarity measure, including itself. So row *i* of G_Y has k_i^Y entries equal to one and others zeros. In addition, all its diagonal entries are equal to one. Let $\mathcal{A}_Y(Y_m) = K_Y^{-1}G_YY_m$ and $\mathcal{A}_Y(Y_m^\circ) = K_Y^{-1}G_YY_m^\circ$ be locally averaged versions of Y_m and Y_m° over G_Y , respectively, where $K_Y = \text{diag}(k_1^Y, \ldots, k_{n_y}^Y)$. For a nearest neighbor graph G_Z , we define $\mathcal{A}_Z(Z_m)$ and $\mathcal{A}_Z(Z_m^\circ)$ in an analogous way. Finally, for any weight $w \in [0, 1)$ and any matrices A and B with n_y and n_z rows respectively, define

$$\widetilde{A} = S_Y(A; w) = wA + (1 - w)A_Y(A),$$

$$\widetilde{B} = S_Z(B; w) = wB + (1 - w)A_Z(B).$$
(1)

In this way, we define $\widetilde{Y}_{m}^{\circ} = S_{Y}(Y_{m}^{\circ}; w_{0})$ and $\widetilde{Z}_{m}^{\circ} = S_{Z}(Z_{m}^{\circ}; w_{0})$ with $w_{0} \in [0, 1)$. In Figure 1A, these are the smoothed Y-modality linked features and smoothed Z-modality linked features.

Initial matching via linear assignment As the columns in $\widetilde{Y}_{m}^{\circ}$ and in $\widetilde{Z}_{m}^{\circ}$ have correspondences, we can compute an $n_{y} \times n_{z}$ distance matrix D° where D_{ij}° measures the distance between the *i*-th row in $\widetilde{Y}_{m}^{\circ}$ and the *j*-th row in $\widetilde{Z}_{m}^{\circ}$ after projecting to respective leading singular subspaces. We obtain an initial matching $\widehat{\Pi}^{\circ}$ as the solution to the linear assignment problem (33, 60):

$$\begin{array}{ll} \text{minimize} & \langle \Pi, D^{\circ} \rangle \\ \text{subject to} & \Pi \in \{0, 1\}^{n_y \times n_z} \\ & \sum_i \Pi_{ij} \leq 1, \forall j, \ \sum_j \Pi_{ij} \leq 1, \forall i, \\ & \sum_{i,j} \Pi_{ij} = n_{\min}. \end{array}$$

Here, $n_{\min} = \min\{n_y, n_z\}$ and for two matrices A and B of the same size, $\langle A, B \rangle = \sum_{i,j} A_{ij} B_{ij}$ denotes the trace inner product. The estimator $\widehat{\Pi}^\circ$ provides a relatively crude matching using only the information provided by the prior knowledge encapsulated in f_y and f_z that link features in the two modalities. By definition, $\widehat{\Pi}^\circ$ gives n_{\min} pairs of matched

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 $_{641}$ rows between the two modalities. We call these matched $_{682}$

642 pairs *initial pivots*.

⁶⁴³ Cross-modality joint embedding and iterative refinement of ⁶⁸⁴ matching

From matched pairs to joint embedding An estimated match-⁶⁸⁶ ing $\widehat{\Pi}$ induces a cross-modality joint embedding of $Y_{\rm m}$ and ⁶⁸⁷ $Z_{\rm m}$. In particular, let $Y_{\rm m}^{r} \in \mathbb{R}^{n_y \times r_y}$ and $Z_{\rm m}^{r} \in \mathbb{R}^{n_z \times r_z}$ collect ⁶⁸⁸ the leading PCs of *all* features (i.e., $Y_{\rm m}$ and $Z_{\rm m}$) in the two ⁶⁸⁹ modalities, respectively. Here, the numbers of PCs to retain, ⁶⁹⁰ i.e., r_y and r_z , are chosen based on data. For any matrix A, let ⁶⁹¹ $[A]_{i}$. denote its *i*-th row. Suppose $\{(i_\ell, i_\ell') : \ell = 1, \dots, n_{\min}\}$ ⁶⁹² are the matched pairs specified by $\widehat{\Pi}$. We perform CCA on ⁶⁹³ data pairs

$$\{([Y_{\mathtt{m}}^{\mathtt{r}}]_{i_{\ell}}, [Z_{\mathtt{m}}^{\mathtt{r}}]_{i_{\ell}'}): \ell = 1, \dots, n_{\min}\}$$

to obtain the leading $r_{\rm cc}$ loading vectors for either modality,

⁶⁴⁶ collected as the columns of $\hat{C}_y = \hat{C}_y(\hat{\Pi})$ and $\hat{C}_z = \hat{C}_z(\hat{\Pi})$, ⁶⁴⁷ respectively. The cross-modal joint embedding induced by ⁶⁴⁸ $\hat{\Pi}$ is then $Y_m^{cc} = Y_m^r \hat{C}_y \in \mathbb{R}^{n_y \times r_{cc}}$ and $Z_m^{cc} = Z_m^r \hat{C}_z \in$ ⁶⁴⁹ $\mathbb{R}^{n_z \times r_{cc}}$, which are the predicted CC scores of Y_m^r and Z_m^r , ⁶⁵⁰ respectively.

⁶⁵¹ Iterative refinement Let $\widehat{\Pi}^{(0)} = \widehat{\Pi}^{\circ}$ be the initial matching

obtained from Eq. (2). Fix a weight $w_1 \in [0,1)$ and the embedding dimension r^{cc} , we refine the estimated matching by iterating the following steps for t = 1, ..., T:

(i) Compute joint embedding $\{Y_{\rm m}^{\rm cc,(t)}, Z_{\rm m}^{\rm cc,(t)}\}$ induced by $_{709}^{708}$ $\widehat{\Pi}^{(t-1)};$

(ii) Apply fuzzy smoothing on joint embedding: $\widetilde{Y}_{m}^{cc,(t)} = \gamma_{11}$ $\mathcal{S}_{Y}(Y_{m}^{cc,(t)}, w_{1}), \widetilde{Z}_{m}^{cc,(t)} = \mathcal{S}_{Z}(Z_{m}^{cc,(t)}, w_{1});$ (12)

(iii) Calculate a distance matrix
$$D^{(t)} \in \mathbb{R}^{n_y \times n_z}$$
 where $D_{ij}^{(t)}$

measures the distance between $[Y_{m}^{CC, (t)}]_{i}$. and $[Z_{m}^{CC, (t)}]_{j}$. and obtain a refined matching $\widehat{\Pi}^{(t)}$ by solving Eq. (2) in ₇₁₆ which D° is replaced with $D^{(t)}$.

⁶⁶³ Figure 1B illustrates the foregoing refinement iteration.

Propagation of matching and post-processing For downtransformation for matching and post-processing For down-For down-For down-Transformation for each cell in Y a match in Z when possible, or *vice versa*, and sometimes both ways. In addition, we would like to have joint embedding of cells across different modalities in a common space. We now describe how MaxFuse achieves these goals. For down-For down-Fo

Filtering and final joint embedding Upon obtaining the 725 670 matched pairs $\{(i_{\ell}, i'_{\ell}) : \ell = 1, \dots, n_{\min}\}$ in $\widehat{\Pi}^{(T)}$, we rank ₇₂₆ 67 them in descending order of $D^{(T)}_{i_\ell i_\ell'}$ and only retain the top $_{^{727}}$ 672 $100 \times (1-\alpha)\%$ pairs, where α is a user-specified filtering ₇₂₈ 673 proportion (with a default $\alpha = 0$). The retained pairs are $\frac{1}{729}$ 674 called refined pivots. Then, we fit a CCA using the re-675 fined pivots and the corresponding rows in $Y_{\rm m}$ and $Z_{\rm m}$ to $_{_{731}}^{_{731}}$ get the associated CCA loading matrices $\hat{C}_y^{\rm e} \in \mathbb{R}^{p_y \times r^{\rm e}}$ and $_{_{732}}^{_{732}}$ 676 677 $\widehat{C}_z^e \in \mathbb{R}^{p_z \times r^e}$. Here the positive integer r^e is a user-specified $\frac{1}{733}$ 678 dimension for final joint embedding. Finally, the joint em-734 679 bedding of the full datasets is given by $Y^{e} = Y \widehat{C}_{y}^{e} \in \mathbb{R}^{N_{y} \times r^{e}}$ 680 and $Z^{e} = Z \widehat{C}_{z}^{e} \in \mathbb{R}^{N_{z} \times r^{e}}$, respectively. In Figure 1C, they 736 681

correspond to the Y-modality embedding and Z-modality embedding matrices.

Using pivots to propagate matching For each row index $i \in \{1, ..., n_y\}$ in Y-modality that does not have a match in Z-modality (i.e., *i* does not belong to any refined pivot), we search for the nearest neighbor of the *i*-th row in \widetilde{Y}_m (Y_m after fuzzy smoothing) that belongs to some refined pivot. Suppose the nearest neighbor is the j_i -th row with a match j'_i in Z-modality, then we call (i, j'_i) a matched pair obtained via propagation. We can optionally filter out any matched pair via propagation in which the nearest neighbor distance between $[\widetilde{Y}_m]_i$. and $[\widetilde{Y}_m]_{j_i}$ is above a user-specified threshold. The retained matched pairs composes the Y-to-Z propagated matching. We then repeat the above procedure with the roles of Y- and Z-modalities switched and obtain the Z-to-Y propagated matching.

Pooling all matched pairs from refined pivots and propagated matching together, we obtain a matching between meta-cells in Y-modality and those in Z-modality. Such a meta-cell level matching defines a single-cell level matching between the original datasets Y and Z by declaring (i, i') a matched pair for $1 \le i \le N_y, 1 \le i' \le N_z$ if the meta-cell that *i* belongs to is matched to the meta-cell that i' belongs to.

Scoring and directional pruning of matching For each singlecell level matched pair (i, i'), we compute Pearson correlation between the *i*-th row of Y^{e} and the *i'*-th row of Z^{e} (i.e., corresponding rows in final joint embedding) as its matching score. We use these matching scores to prune single-cell level matching, with the *direction* of pruning specified by user. Suppose the user wants to find for each cell in Z a match in Y (e.g., Z is a CODEX dataset and Y snRNAseq). Then for each cell index $1 \le i' \le N_z$, we first list all refined pivots and propagated matching pairs that contain i'. If the list is non-empty, we only retain the pair with the highest matching score. Otherwise, we declare no match for cell i' in Zmodality. If the direction is reversed, we apply the foregoing procedure with the roles of Y and Z switched. Furthermore, if no directional pruning is desired, we just keep all refined pivots and post-screening propagated matching pairs in the final single-cell matching.

After filtering, propagation, and potential pruning, the final list of matched pairs correspond to the final matching in Figure 1C.

A batched version of MaxFuse. Single-cell and spatial datasets can be large. To facilitate fast computation for large datasets, we developed a batched version of MaxFuse.

Batching Fix a desired pair of sample sizes (n_y, n_z) and meta-cell ratios $(N_y/n_y, N_z/n_z)$, we randomly partition the dataset under Y-modality (resp. Z-modality) into disjoint subsets of sizes roughly all equal to N_y (resp. N_z). Denote them as $Y^{[1]}, \ldots, Y^{[b_y]}$ and $Z^{[1]}, \ldots, Z^{[b_z]}$. We then apply the MaxFuse pipeline on each pair of data $\{Y^{[l]}, Z^{[m]}\}$, $1 \le l \le b_y$, $1 \le m \le b_z$ to get the refined pivots and the propagated matching, as well as their induced single-cell level matched pairs, for that pair of batches.

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Stitching After pooling all refined pivots from all batch 737 pairs, we obtain a multiple-to-multiple matching. For each 738 unique cell in Z-modality, we average all its matches in Y-739 modality, that is, we average matched cells in the modality 740 with a higher SNR. After this step, we get a pair of matrices 741 with rows paired. We then fit CCA on this pair of matrices 742 and get the loading matrices, which are then used to jointly 743 embed the whole datasets. Finally, with the joint embedding 744 of the whole datasets in Y- and Z-modalities, scoring and di- $_{789}$ 745 rectional pruning of matching are performed in the same way 700 746 as in MaxFuse without batching. 747 791

748 Systematic benchmarks on ground-truth datasets.

794 MaxFuse and other methods in comparison MaxFuse was 749 795 implemented in Python, and the four methods in comparison, 750 Seurat V3, Harmony, Liger, and BindSC, were implemented 751 in R. All benchmarking datasets were preprocessed in the 752 same way for all methods, including filtering of low-quality 753 cells, selection of highly variable genes and protein features 754 to be used in integration, feature linkage scheme (e.g., pro-755 tein to their corresponding gene names), and normalization of 756 raw observed values (except for Liger which required scaling 757 without centering). We used the default tuning parameters 758 in each method suggested by the respective tutorial except 759 for BindSC, for which we used the separate set of parame-760 ters suggested for the integration of protein-related data by 761 its method tutorial website. For MaxFuse, initial matching 762 used features that are weakly linked (e.g., protein CD4 and 763 RNA CD4) and are smoothed by all-feature nearest-neighbor 796 764 graphs. For refined matching, all features from both modal-797 765 ities were used (e.g., all proteins and RNAs that are highly 798 766 variable). For other methods in comparison, BindSC used 767 both the weakly linked features and all features, whereas oth-768 ers only used the weakly linked features by design. The full 801 769 detail (including preprocessing, implementation, and down-770 stream analysis and evaluation of MaxFuse and other meth-771 ods) is recorded and can be reproduced. 772 804

773 Evaluation metrics

1. Cell type matching accuracy: To evaluate the matching 774 performance for Seurat, Liger, Harmony, and BindSC, we 775 used the respective integration embedding vectors pro-776 duced by each method. For these methods, for each 777 cell in one modality, we regarded its nearest neighbor 778 from the other modality under Pearson correlation dis-779 tance in the embedding space as its match. For MaxFuse, 780 813 we directly used matched pairs produced in the final re-781 814 sult. For all methods, we use the same matching direc-782 tion (e.g., for each cell in CODEX data finding a matched⁸¹⁵ 783 cell in scRNAseq data) for fair comparison. Accuracy 784 of the matchings was measured by fraction of matched⁸¹⁷ 785 pairs with identical cell type annotations. Details on cell⁸¹⁸ 786 type annotation are given below in the description of each 819 787 benchmarking dataset. 788 820

2. FOSCTTM: Fraction of sample closer than true match 821 (FOSCTTM) was used to evaluate single-cell level align- 822 ment accuracy on datasets with ground-truth single-cell 823 level pairing. The measure has been used previously in cross-modality alignment benchmarking tasks (19, 36, 37). For such data, $N_y = N_z = N$, and FOSCTTM is defined as:

FOSCTTM =
$$\frac{1}{2N} \left(\sum_{i=1}^{N} \frac{n_y^{(i)}}{N} + \sum_{i=1}^{N} \frac{n_z^{(i)}}{N} \right),$$

where for each i, $n_y^{(i)} = |\{j|d(y_i, z_j) < d(y_i, z_i)\}|$ with da distance metric in the joint embedding space and for $l = 1, \ldots, N$, y_l and z_l the embedded vectors of the l-th cell with its measurements in Y and Z modality, respectively. The counts $n_z^{(i)}$, $i = 1, \ldots, N$, are defined analogously. A lower value of FOSCTTM indicates better integration performance.

3. FOSKNN: Fraction of sample with true match among k-nearest-neighbors (FOSKNN) was used to evaluate single-cell level alignment accuracy on datasets with ground-truth single-cell level pairing. For such data, $N_y = N_z = N$. For any method in comparison, let $\{y_i : i = 1, ..., N\}$ be the coordinates of cells in the joint embedding space from their Y modality information, and let $\{z_i : i = 1, ..., N\}$ be embedding coordiantes from their Z modality information. Then

FOSKNN =
$$\frac{1}{2N} \left(\sum_{i=1}^{N} \mathbf{1}_{E_{y,k}}^{(i)} + \sum_{i=1}^{N} \mathbf{1}_{E_{z,k}}^{(i)} \right)$$

where for i = 1, ..., N, $\mathbf{1}_{E_{y,k}}^{(i)}$ is the indicator of whether the k closest embedded vectors from Z modality to y_i includes z_i . The quantity $\mathbf{1}_{E_{z,k}}^{(i)}$ is defined analogously.

- 4. Silhouette F1 score: Silhouette F1 score has been used to simultaneously measure modality mixing and information preservation post-integration process (21, 35). In brief, the F1 score was calculated by 2 · slt_mix · slt_clust/(slt_mix + slt_clust), where slt_mix is defined as one minus normalized Silhouette width with the label being modality index (two modalities); slt_clust is defined by the normalized Silhouette width with label being cell type annotations (e.g., "CD4 T", "CD8 T", "B", etc.). All Silhouette widths were computed using the silhouette() function from R package cluster.
- 5. ARI F1 score: Adjusted Random Index F1 score has been used to jointly measure modality mixing and information preservation post-integration process (21, 35). The score was calculated in a similar way to Silhouette F1 score, while the Adjusted Random Index was used instead of the Silhouette width. All ARI scores were computed using the function adjustedRandIndex() in R package mclust.

CITE-seq PBMC dataset The CITE-seq healthy human pbmc data with antibody panel of 228 markers was retrieved from Hao et al. (34). For benchmarking purposes, 5 batches of cells, each with 10k cells were randomly sampled from the original dataset, and selected for benchmarking. The first 15

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components of the embedding vector produced by all methvector produced by all meth-vector produced by all methvector produced by all meth-vector produced by all methvector produced by all meth-vector produced by all meth-vector

For antibody dropping, we ranked the importance of each in- 887 831 dividual antibody in the panel in terms of phenotyping con-888 832 tribution. The importance score was calculated by training a 889 833 random forest model (function randomForest in R pack-890 834 age randomForest, with default parameters) using all an-891 835 tibodies to predict cell type labels (annotation level 2 from 892 836 Hao et al.), then a permutation feature importance test (func- 893 837 tion varImp with default parameters in R package caret)894 838 was performed on the trained model to acquire the individ-895 839 ual importance scores. Then antibodies were ranked by the 896 840 importance scores, and 4 panels were used for antibody drop-841

ping test: (1) full 228 antibody panel; (2) top 100 most im-⁸⁹⁷
portant antibodies; (3) top 50 most important antibodies; (4) ⁸⁹⁸
top 30 most important antibodies.

CITE-seq BMC dataset The CITE-seq healthy human 901 845 BMC data with antibody panel of 25 markers was retrieved 902 846 from R package SeuratData 'bmcite', orignated from Hao 903 847 et al. (34). For benchmarking purpose, a total of 20k cells ₉₀₄ 848 were randomly sampled from the original dataset, and se-905 849 lected for benchmarking. The first 15 components of the and 850 embedding vectors produced by all methods were used for 907 851 benchmarking metric calculation. The UMAP visualization 908 852 of the integration process was also calculated with the first 15 $_{909}$ 853 components of the embedding vectors. The original cell type 910 854 annotation (lv2) from the R package was binned into 8 popu-855 lations: "DC", "progenitor", "monocyte", "NK", "B", "CD4 912 856 T", "CD8 T" and "Other T", and used for benchmarking. 857 913

Ab-seq BMC dataset The Ab-seq healthy human BMC 914 858 data with antibody panel of 97 markers, and whole transcrip-⁹¹⁵ 859 tome sequencing was retrieved from Triana et al. (39). All 916 860 cells in the dataset (\sim 13k), except cells belonging to cell⁹¹⁷ 861 types with insufficient amount of cells (< 50 cells, anno-⁹¹⁸ 862 tated as "Doublet and Triplets", "Early GMP", "Gamma delta 919 863 T cells", "Immature B cells", "Metaphase MPPs", "Neu-920 864 trophils" in Triana et al.) were excluded for integration, and 921 865 the remaining 14 cell types were used during benchmarking. 866

⁸⁶⁷ The first 15 components of the embedding vectors produced ⁹²²

⁸⁶⁸ by all methods were used for benchmarking metric calcula-

tion. The UMAP visualization of the integration process was
 also calculated with the first 15 components of the embedding
 vectors.

TEA-seq PBMC dataset The TEA-seq neutrophil-depleted 927 872 human PBMC dataset was retrieved from Swanson et al. 928 873 (**41**) (GSM4949911). This dataset is stained with 46 929 874 antibodies and contains chromatin accessibility informa-930 875 tion. Cell type annotation was performed using R package 931 876 Seurat(v4) WNN-multi-modal clustering pipeline: func- 932 877 tion FindMultiModalNeighbors was run on ADT 933 878 PCA (first 25 components) and ATAC LSI (2-50 com-934 879

ponents, calculated by R package Archr (42)). Subsequently, function FindClusters was used to generate unsupervised clustering (with parameter algorithm = 3, resolution = 0.2), followed by manual annotation. A total of 8 populations were identified ("Naive CD4", "Mem CD4", "Monocyte", "NK", "Naive CD8", "Mem CD8", "Effector CD8", "B", "NK"), and the total amount of cells was \sim 7.4k. ADT expressions and gene activity scores (calculated by R package Archr (42)) were used as input for Max-Fuse and other methods. Additionally, during matching refinement, MaxFuse used LSI reduction of the ATAC peaks (first 2-50 components) as features for the ATAC modality. The first 15 components of the embedding vectors produced by all methods were used for benchmarking metric calculation. The UMAP visualization of the integration process was also calculated with the first 15 components of the embedding vectors.

ASAP-seq PBMC dataset The ASAP-seq healthy human PBMC data (CD28 & CD3 stim PBMC control group) with an antibody panel of 227 markers, and chromatin accessibility information was retrieved from Mimitou et al. (40)(GSM4732109). Cell type annotation was performed using R package Seurat(v4) WNN-multi-modal clustering pipeline: function FindMultiModalNeighbors was run on ADT PCA (first 18 components) and ATAC LSI (2-40 components, calculated by R package Archr). Subsequently, function FindClusters was used to generate unsupervised clustering (with parameter algorithm = 3, resolution = 0.3), followed by manual annotation. A total of 9 populations were identified ("Naive CD4", "Mem CD4", "Monocyte", "NK", "Naive CD8", "Mem CD8", "B", "OtherT", "dirt"), and "dirt" was removed from subsequent usage, resulting in \sim 4.4 k cells used. ADT expressions and gene activity scores (calculated by R package Archr) were used as input for MaxFuse and other methods. Additionally, during matching refinement, MaxFuse used LSI reduction of the ATAC peaks (First 2-50 components) as features for the ATAC modality. The first 15 components of the embedding vectors produced by all methods were used for benchmarking metric calculation. The UMAP visualization of the integration process was also calculated with the first 15 components of the embedding vectors.

MaxFuse on Spatial-omics matching.

CODEX and scRNA-seq human tonsil CODEX multiplexed imaging data of human tonsil tissues with a panel of 46 antibodies were retrieved from Kennedy-Darling et al. (47). Images from tonsil-9338 (region X2-8, Y7-15) were used. Whole-cell segmentation was performed with a local implementation of Mesmer (61), with weights downloaded from: https://deepcell-data.s3-us-west-1.amazonaws.com/model-weights/Multiplex_Segmentation_20200908_2_head.h5. Inputs of segmentation were DAPI (nuclear) and CD45 (membrane). Signals from the images were capped at 99.7th percentile, with prediction parameter model_mpp = 0.8. Cells

smaller than 30 pixels or larger than 800 pixels were ex-992 935 cluded. Signals from individual cells were then extracted, 993 936 and scaled to the [0,1] interval, with percentile cutoffs $_{994}$ 937 of 0.5% (floor) and 99.5% (ceiling). Cell type annota-995 938 tion was performed using R package Seurat clustering 996 939 pipeline: function FindNeighbors was run on CODEX 997 940 protein PCA (first 15 components). Subsequently, function 998 941 FindClusters was used to generate unsupervised clus-999 942 tering (with parameter resolution = 1), followed by_{1000} 943 manual annotation. A total of 9 populations were identified₁₀₀₁ 944 ("B-CD22-CD40", "B-Ki67", "Plasma", "CD4 T", "CD81002 945 T", "DC", "Fibro/Epi", "Vessel", "Other", and "Dirt"), and 61003 946 populations (\sim 180k cells) were used in subsequent analysis₁₀₀₄ 947 ("B-CD22-CD40", "B-Ki67", "Plasma", "CD4 T", "CD81005 948 T", and "DC"). 949

Single-cell RNA-seq data of dissociated human tonsil cells1007 950 were retrieved from King et al. (48). The pre-processing and₁₀₀₈ 951 cell typing steps were done in R package Seurat, follow-1009 952 ing the description presented in King et al. In brief, tonsil₁₀₁₀ 953 cells ("t1", "t2" and "t3") were merged, then filtered by cri-1011 954 teria: nFeature_RNA > 200 & nFeature_RNA <_1012 955 7500 & percent.mt < 20, and subsequently value₁₀₁₃ 956 normalized by function SCTransform. Harmony batch₁₀₁₄ 957 correction was performed for different tonsils, with func-1015 958 tion RunHarmony. Unsupervised clustering was per-1016 959 formed by function FindNeighbors with harmony em-1017 960 bedding (1-27 dimensions) and function FindClusters₁₀₁₈ 961 with resolution = 0.5. A total of 8 population₁₀₁₉ 962 was defined ("B-CD22-CD40", "B-Ki67", "circulating B",1020 963 "Plasma", "CD4 T", "CD8 T", "DC", "Other"), and 6 pop-1021 964 ulations (\sim 13k cells) were used in subsequent analysis ("B-1022" 965 CD22-CD40", "B-Ki67", "Plasma", "CD4 T", "CD8 T", and₁₀₂₃ 966 "DC"). 967

Boundaries of germinal centers from the CODEX images₁₀₂₄
 were drawn manually, and dilation and erosion from the₁₀₂₅
 boundary was performed with python package skimage,

with function morphology.binary_dilation and 1026 971 morphology.disk. Ten layers inward or outward1027 972 from the boundary (each layer = 30 pixels, resolution:1028 973 376nm/pixel) was performed. Cells were assigned to each1029 974 layer by their centroids' locations. The RNA expression level1030 975 from each layer, based on the averaged CODEX matched1031 976 scRNA-seq cells, were plotted with R package ggplot2. 1032 977 The UMAP visualization of the integration process was cal-1033 978 culated with the first 15 components of the embedding vec-1034 979 tors. 1035 980

1036 HUBMAP atlas: CODEX, snRNA-seq and snATAC-seq hu-1037 981 man intestine CODEX multiplex imaging (48 markers), 1038 982 snRNA-seq and snATAC-seq of healthy human intestine cells 983 were acquired from Hickey et al. (32). For CODEX, samples1039 984 "B005 SB" and "B006 CL" were used, while for snRNA-1040 985 seq and snATAC-seq, single-ome sequencing data of four1041 986 donors ("B001", "B004", "B005", "B006") from the study1042 987 were used. Cells annotated as "B cells", "T cells", "Endothe-1043 988 lial", "Enteroendocrine", "Goblet", "Mono_Macrophages",1044 989 "Plasma", "Smooth muscle", and "Stroma" were selected1045 990

for the integration process. Cell counts for each modality₁₀₄₆

used for MaxFuse were: CODEX ~100k (small bowel) and ~70k (colon); snRNA-seq ~32k (small bowel) and ~16k (colon); snATAC-seq ~28k (small bowel) and ~21k (colon). CODEX protein expressions, snRNA-seq RNA expressions, snATAC-seq gene activity scores and LSI scores (calculated with R package Archr) were used as MaxFuse input (RNA expressions, gene activity scores and LSI scores were batch-corrected by Harmony (20), based on patient ID). The matching and integration process was done on colon and small bowel samples respectively.

Pairwise MaxFuse alignments of cells between protein (CODEX) and RNA (snRNA-seq), and cells between RNA (snRNA-seq) and ATAC (snATAC-seq) were performed. Refined pivots from the two bi-modal alignments were chained together by using the pivot cells in the RNA modality as the intermediary, resulting in a list of tri-modal pivots linking all three modalities. Subsequently, we used these pivots to calculate a tri-omic embedding via generalized CCA (gcca) (21, 55). In particular, we used the gcca formulation and algorithm described in (21).

The UMAP visualization of the tri-modal integration was calculated with the first 15 components of the embedding vectors (gcca scores in this case). Embeddings of CODEX cells were overlaid with their protein expressions, or their matched cells' RNA expressions, or gene activity scores. Spatial locations of these expression values and scores were plotted based on CODEX cells' x-y centroid locations. Additionally, we showed spatial locations of transcription factor motif enrichment scores (Z-score) of CODEX cells, based on their matched snRNA-seq cells, which were calculated by R package chromVAR (56). All values were capped between 5% - 95% quantiles for visualization purpose during plotting.

Benchmark on ground-truth strongly linked modalities.

MaxFuse and other methods specialized in ATAC-RNA integration in comparison We compared MaxFuse to three methods that specialize in ATAC-RNA integration: scGLUE (19), Maestro (62) and scJoint (63). For MaxFuse, the initial matching used the gene activity scores, while during refined matching the active RNA features and LSI embedding from ATAC were used. For other methods in comparison, we used their default settings. Metrics used for benchmarking were calculated similarly as described in previous sections. The full detail (including preprocessing, implementation, and downstream analysis and evaluation of MaxFuse and other methods specialized in ATAC-RNA integration) is recorded and can be reproduced.

Multiome scRNA - scATAC-seq human retina dataset Multiome (scRNA-seq & scATAC-seq) data of human retina cells was retrieved from Wang et al. (46). For input required by MaxFuse: gene activity and LSI scores of ATAC cells were calculated by R package Archr using the fragment files, while RNA counts were directly extracted. For other methods in comparison, we used their default settings. For benchmarking, a total of 20k cells were randomly sampled

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and used for testing. All cell types were used during integra-1109 1047 tion ("Rod", "OFF cone bipolar", "Mullerglia", "ON cone 1048 bipolar", "Rod bipolar", "Cone", "GABA amacrine", "Hor-1049 izontal", "Glyamacrine", "AII amacrine", "Retinal ganglion1112 1050 cell", "Astrocyte", "Microglia", annotated by Wang et al.).1114 1051 The first 15 components of the embedding vectors produced¹¹¹⁵ 1052 by all methods were used for benchmarking metric calcula-1053 1118 tion. 1054 1119

1120 10x Multiome peripheral blood mononuclear cells Multi-1121 1055 ome (scRNA-seq & scATAC-seq) data of human mononu-1122 1056 clear peripheral blood cells was retrieved from the 10x pub-1124 1057 lic data repository (44). For input required by MaxFuse: gene¹¹²⁵ 1058 activity and LSI scores of ATAC modality were calculated by1127 1059 R package Signac, the latter using the fragment files. RNA¹¹²⁸ 1060 counts were directly extracted from the cellranger out-1130 1061 put. Cell-type labels were transferred from CITE-seq PBMC¹¹³¹ 1062 1132 reference (34) using the method in (34). 1063 1133 1134

Multi-1135 10x Multiome day 18 embryonic mouse brain cells 1064 ome (scRNA-seq & scATAC-seq) data of developing mouse1137 1065 brain cells was retrieved from the 10x public data reposi-1066 tory (44). For input required by MaxFuse: gene activity and 1140 1067 LSI scores of ATAC modality were calculated by R package $_{1142}^{1141}$ 1068 Signac, the latter using the fragment files. RNA counts1143 1069 were directly extracted from the cellranger output. Cell-1144 1070 type labels were transferred from (64) using the method in¹¹⁴⁶ 1071 1147 (65). 1072 1148

10x Multiome developing human cerebral cortex cells1150 1073 Multiome (scRNA-seq & scATAC-seq) data of developing¹¹⁵¹ 1074 human cerebral cortex cells was retrieved from Trevino et1153 1075 al. (45). For input required by MaxFuse: gene activity and l_{1155}^{1154} 1076 LSI scores of ATAC modality were calculated by R package¹¹⁵⁶ 1077 Signac using the fragment files. RNA counts and $ATAC_{1158}^{1157}$ 1078 peak matrices were extracted from 10x cellranger out-1159 1079 put. The cell-type labels were taken from the original publi- $\frac{1160}{1161}$ 1080 cation. 1081 1162 1163

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AUTHOR CONTRIBUTIONS 1095

- Conceptualization: S.C., B.Z., G.P.N., N.R.Z., Z.M. 1096
- Algorithm Development and Implementation: S.C., N.R.Z., Z.M. 1097
- Analysis: S.C., B.Z., S.H., Z.M. 1098
- Contribution of Key Reagents and Tools: J.W.H., K.Z.L., M.S., W.J.G, G.P.N. 1099 1100 Supervision: G.P.N., N.R.Z., Z.M
- 1101
- Both S.C. and B.Z. contributed equally and have the right to list their name first in1186 1102 1103 their CV.

CONFLICT OF INTERESTS 1104

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Reference

- 1. Marlon Stoeckius, Christoph Hafemeister, William Stephenson, Brian Houck-Loomis, Pratip K Chattopadhyay, Harold Swerdlow, Rahul Satija, and Peter Smibert. Simultaneous epitope and transcriptome measurement in single cells. Nature methods, 14(9):865-868, 2017.
- 2. Payam Shahi, Samuel C Kim, John R Haliburton, Zev J Gartner, and Adam R Abate. Ab seq: Ultrahigh-throughput single cell protein profiling with droplet microfluidic barcoding. Scientific reports, 7(1):1-12, 2017.
- 3. Dominic Grün and Alexander van Oudenaarden. Design and analysis of single-cell se quencing experiments. Cell, 163(4):799-810, 2015.
- Ino D Karemaker and Michiel Vermeulen. Single-cell dna methylation profiling: technologies 4. and biological applications. Trends in biotechnology, 36(9):952-965, 2018.
- 5. Marek Bartosovic, Mukund Kabbe, and Goncalo Castelo-Branco. Single-cell cut&tag profiles histone modifications and transcription factors in complex tissues. Nature biotechnol oav. 39(7):825-835, 2021.
- 6. Sebastian Preissl, Kyle J Gaulton, and Bing Ren. Characterizing cis-regulatory elements using single-cell epigenomics. Nature Reviews Genetics, pages 1-23, 2022.
- 7. Wai Lim Ku, Kosuke Nakamura, Weiwu Gao, Kairong Cui, Gangging Hu, Qingsong Tang Bing Ni. and Keji Zhao. Single-cell chromatin immunocleavage sequencing (scchic-seq) to profile histone modification. Nature methods, 16(4):323-325, 2019.
- 8. Caleb A Lareau, Fabiana M Duarte, Jennifer G Chew, Vinay K Kartha, Zach D Burkett, Andrew S Kohlway, Dmitry Pokholok, Martin J Arvee, Frank J Steemers, Ronald Lebofsky, et al Droplet-based combinatorial indexing for massive-scale single-cell chromatin accessibility Nature Biotechnology, 37(8):916-924, 2019.
- Anjali Rao, Dalia Barkley, Gustavo S França, and Itai Yanai. Exploring tissue architecture 9. using spatial transcriptomics. Nature, 596(7871):211-220, 2021.
- 10. Yury Goltsev, Nikolay Samusik, Julia Kennedy-Darling, Salil Bhate, Matthew Hale, Gustavo Vazquez, Sarah Black, and Garry P Nolan. Deep profiling of mouse splenic architecture with codex multiplexed imaging. Cell, 174(4):968-981, 2018.
- 11. Michael Angelo, Sean C Bendall, Rachel Finck, Matthew B Hale, Chuck Hitzman, Alexan der D Borowsky, Richard M Levenson, John B Lowe, Scot D Liu, Shuchun Zhao, et al Multiplexed ion beam imaging of human breast tumors. Nature medicine, 20(4):436-442 2014
- 12. Charlotte Giesen, Hao AO Wang, Denis Schapiro, Nevena Zivanovic, Andrea Jacobs, Bodo Hattendorf, Peter J Schüffler, Daniel Grolimund, Joachim M Buhmann, Simone Brandt, et al Highly multiplexed imaging of tumor tissues with subcellular resolution by mass cytometry Nature methods, 11(4):417-422, 2014.
- Shanshan He, Ruchir Bhatt, Carl Brown, Emily A Brown, Derek L Buhr, Kan Chantranu 13 vatana, Patrick Danaher, Dwayne Dunaway, Ryan G Garrison, Gary Geiss, et al. High-plex imaging of rna and proteins at subcellular resolution in fixed tissue by spatial molecular imaging. Nature Biotechnology, pages 1-13, 2022.
- Emma Lundberg and Georg HH Borner. Spatial proteomics: a powerful discovery tool for 14. cell biology. Nature Reviews Molecular Cell Biology, 20(5):285-302, 2019.
- Yanxiang Deng, Marek Bartosovic, Sai Ma, Di Zhang, Petra Kukanja, Yang Xiao, Graham 15. Su, Yang Liu, Xiaoyu Qin, Gorazd B Rosoklija, et al. Spatial profiling of chromatin accessibility in mouse and human tissues. Nature, 609(7926):375-383, 2022.
- Ricard Argelaguet, Anna SE Cuomo, Oliver Stegle, and John C Marioni. Computational 16. principles and challenges in single-cell data integration. Nature biotechnology, 39(10): 1202-1215, 2021
- 17. Yang Xu and Rachel Patton McCord. Diagonal integration of multimodal single-cell data: potential pitfalls and paths forward. Nature Communications, 13(1):1-4, 2022
- 18. Jinzhuang Dou, Shaoheng Liang, Vakul Mohanty, Xuesen Cheng, Sangbae Kim, Jongsu Choi, Yumei Li, Katayoun Rezvani, Rui Chen, and Ken Chen. Unbiased integration of single cell multi-omics data. BioRxiv, 2020.
- 19. Zhi-Jie Cao and Ge Gao. Multi-omics single-cell data integration and regulatory inference with graph-linked embedding. Nature Biotechnology, pages 1-9, 2022.
- 20. Ilya Korsunsky, Nghia Millard, Jean Fan, Kamil Slowikowski, Fan Zhang, Kevin Wei, Yuriy Baglaenko, Michael Brenner, Po-ru Loh, and Soumya Raychaudhuri. Fast, sensitive and accurate integration of single-cell data with harmony. Nature methods, 16(12):1289-1296 2019
- 21. Bokai Zhu, Shuxiao Chen, Yunhao Bai, Han Chen, Nilanjan Mukherjee, Gustavo Vazquez, David R McIlwain, Alexandar Tzankov, Ivan T Lee, Matthias S Matter, et al. Robust single cell matching and multi-modal analysis using shared and distinct features reveals orches trated immune responses. bioRxiv, 2021.
- 22. Joshua D Welch, Velina Kozareva, Ashley Ferreira, Charles Vanderburg, Carly Martin, and Evan Z Macosko. Single-cell multi-omic integration compares and contrasts features of brain cell identity. Cell, 177(7):1873-1887, 2019.
- 23. Kevin E Wu, Kathryn E Yost, Howard Y Chang, and James Zou. Babel enables crossmodality translation between multiomic profiles at single-cell resolution. Proceedings of the National Academy of Sciences, 118(15):e2023070118, 2021.
- Tim Stuart, Andrew Butler, Paul Hoffman, Christoph Hafemeister, Efthymia Papalexi 24 William M Mauck III, Yuhan Hao, Marlon Stoeckius, Peter Smibert, and Rahul Satija. Comprehensive integration of single-cell data, Cell, 177(7):1888-1902, 2019.
- 25. Zhen Miao, Beniamin D Humphreys, Andrew P McMahon, and Junhvong Kim, Multi-omics integration in the age of million single-cell data. Nature Reviews Nephrology, 17(11):710-724. 2021.
- 26. Jinzhuang Dou, Shaoheng Liang, Vakul Mohanty, Qi Miao, Yuefan Huang, Qingnan Liang Xuesen Cheng, Sangbae Kim, Jongsu Choi, Yumei Li, et al. Bi-order multimodal integration of single-cell data. Genome biology, 23(1):1-25, 2022.
- 27. Zhana Duren, Xi Chen, Mahdi Zamanighomi, Wanwen Zeng, Ansuman T Satpathy Howard Y Chang, Yong Wang, and Wing Hung Wong. Integrative analysis of single-cell genomics data by coupled nonnegative matrix factorizations. Proceedings of the National Academy of Sciences, 115(30):7723-7728, 2018.
- 28 Vivien Marx A dream of single-cell proteomics Nature Methods 16(9):809-812 2019
- 29 Vidhya M. Ravi, Paulina Will, Jan Kueckelhaus, Na Sun, Kevin Joseph, Henrike Salié, Lea

- Vollmer, Ugne Kuliesiute, Jasmin von Ehr, Jasim K. Benotmane, Nicolas Neidert, Mariet280
 Follo, Florian Scherer, Jonathan M. Goeldner, Simon P. Behringer, Pamela Franco, Mo-1281
- 1196
 hammed Khiat, Junyi Zhang, Ulrich G. Hofmann, Christian Fung, Franz L. Ricklefs, Ka-1282

 1197
 trin Lamszus, Melanie Boerries, Manching Ku, Jürgen Beck, Roman Sankowski, Mariusteas

 1198
 Schwabenland, Marco Prinz, Ulrich Schüller, Saskia Killmer, Bertram Bengsch, Axel K.1284

 1199
 Walch, Daniel Delev, Oliver Schnell, and Dieter Henrik Heiland. Spatially resolved multi-1285

 1200
 omics deciphers bidirectional tumor-host interdependence in glioblastoma. Cancer Cell, 401286
- (6):639–655.e13, 2022. ISSN 1535-6108. doi: https://doi.org/10.1016/j.ccell.2022.05.009.
 Amin Abedini, Ziyuan Ma, Julia Frederick, Poonam Dhillon, Michael S Balzer, Rojeshtzes
 Shrestha, Hongbo Liu, Steven Vitale, Kishor Devalaraja-Narashimha, Paola Grandi, et al.1289
 Spatially resolved human kidney disease progression. *bioRxiv*, 2022.
 1201
- 1206
 31. Anuja Sathe, Kaishu Mason, Susan M Grimes, Zilu Zhou, Billy T Lau, Xiangqi Bai, Andrewi 292

 1207
 Su, Xiao Tan, H Lee, Carlos J Suarez, et al. Colorectal cancer metastases in the liver estab-1293

 1208
 lish immunosuppressive spatial networking between tumor associated spp1+ macrophages1294

 1209
 and fibroblasts. *Clinical Cancer Research: an Official Journal of the American Association*295

 1210
 for *Cancer Research*, pages CCR–22, 2022.
- 1211
 32. John W Hickey, Winston R Becker, Stephanie A Nevins, Aaron Horning, Almudena Espint297

 1212
 Perez, Roxanne Chiu, Derek C Chen, Daniel Cotter, Edward D Esplin, Annika K Weimer,1298

 1213
 et al. High resolution single cell maps reveals distinct cell organization and function across1299

 1214
 different regions of the human intestine. *bioRxiv*, 2021.
 1300
- Rainer Burkard, Mauro Dell'Amico, and Silvano Martello. Assignment problems: revisedr301 reprint. SIAM, 2012. 1302
- Yuhan Hao, Stephanie Hao, Erica Andersen-Nissen, William M Mauck III, Shiwei Zheng,1303
 Andrew Butler, Maddie J Lee, Aaron J Wilk, Charlotte Darby, Michael Zager, et al. Integrated:304
 analysis of multimodal single-cell data. *Cell*, 184(13):3573–3587, 2021.
- Hoa Thi Nhu Tran, Kok Siong Ang, Marion Chevrier, Xiaomeng Zhang, Nicole Yee Shin1306
 Lee, Michelle Goh, and Jinmiao Chen. A benchmark of batch-effect correction methods for1307
 single-cell rna sequencing data. *Genome biology*, 21(1):1–32, 2020. 1308
- Jie Liu, Yuanhao Huang, Ritambhara Singh, Jean-Philippe Vert, and William Stafford Noble.1309
 Jointly embedding multiple single-cell omics measurements. In Algorithms in bioinformat-1310
 ics.... International Workshop, WABI..., proceedings. WABI (Workshop), volume 143. NIH1311
 Public Access, 2019. 1312
- April R Kriebel and Joshua D Welch. Uinmf performs mosaic integration of single-cell multi-1313
 omic datasets using nonnegative matrix factorization. *Nature communications*, 13(1):1–17,1314
 2022. 1315
- Etienne Becht, Leland McInnes, John Healy, Charles-Antoine Dutertre, Immanuel WH316
 Kwok, Lai Guan Ng, Florent Ginhoux, and Evan W Newell. Dimensionality reduction for1317
 visualizing single-cell data using umap. *Nature biotechnology*, 37(1):38–44, 2019.
- Sergio Triana, Dominik Vonficht, Lea Jopp-Saile, Simon Raffel, Raphael Lutz, Danieli319
 Leonce, Magdalena Antes, Pablo Hernández-Malmierca, Diana Ordoñez-Rueda, Beáta Ra-1320
 masz, et al. Single-cell proteo-genomic reference maps of the hematopoietic system enable1321
 the purification and massive profiling of precisely defined cell states. *Nature immunology*,1322
 22(12):1577–1589, 2021.
- Eleni P Mimitou, Caleb A Lareau, Kelvin Y Chen, Andre L Zorzetto-Fernandes, Yuhan Hao,
 Yusuke Takeshima, Wendy Luo, Tse-Shun Huang, Bertrand Z Yeung, Efthymia Papalexi,
 et al. Scalable, multimodal profiling of chromatin accessibility, gene expression and protein
 levels in single cells. *Nature biotechnology*, 39(10):1246–1258, 2021.
- Elliott Swanson, Cara Lord, Julian Reading, Alexander T Heubeck, Palak C Genge, Zachary Thomson, Morgan DA Weiss, Xiao-jun Li, Adam K Savage, Richard R Green, et al. Simultaneous trimodal single-cell measurement of transcripts, epitopes, and chromatin accessibility using tea-seq. *Ellie*, 10:e63632, 2021.
- Jeffrey M Granja, M Ryan Corces, Sarah E Pierce, S Tansu Bagdatli, Hani Choudhry, Howard Y Chang, and William J Greenleaf. Archr is a scalable software package for integrative single-cell chromatin accessibility analysis. *Nature genetics*, 53(3):403–411, 2021.
- Kevin Z. Lin and Nancy R. Zhang. Quantifying common and distinct information in single-cell multimodal data with tilted-cca. *bioRxiv*, 2022. doi: 10.1101/2022.10.07.511320.
- 1251 44. 10X Genomics. 10x genomics datasets, 2022.
- Alexandro E Trevino, Fabian Müller, Jimena Andersen, Laksshman Sundaram, Arwa Kathiria, Anna Shcherbina, Kyle Farh, Howard Y Chang, Anca M Paşca, Anshul Kundaje, et al. Chromatin and gene-regulatory dynamics of the developing human cerebral cortex at single-cell resolution. *Cell*, 184(19):5053–5069, 2021.
- 46. Sean K Wang, Surag Nair, Rui Li, Katerina Kraft, Anusri Pampari, Aman Patel, Joyce B
 Kang, Christy Luong, Anshul Kundaje, and Howard Y Chang. Single-cell multiome of the
 human retina and deep learning nominate causal variants in complex eye diseases. *bioRxiv*,
 2022.
- Julia Kennedy-Darling, Salil S Bhate, John W Hickey, Sarah Black, Graham L Barlow,
 Gustavo Vazquez, Vishal G Venkataraaman, Nikolay Samusik, Yury Goltsev, Christian M
 Schürch, et al. Highly multiplexed tissue imaging using repeated oligonucleotide exchange
 reaction. European Journal of Immunology, 51(5):1262–1277, 2021.
- Hamish W King, Kristen L Wells, Zohar Shipony, Arwa S Kathiria, Lisa E Wagar, Caleb Lareau, Nara Orban, Robson Capasso, Mark M Davis, Lars M Steinmetz, et al. Integrated single-cell transcriptomics and epigenomics reveals strong germinal center–associated etiology of autoimmune risk loci. *Science Immunology*, 6(64):eabh3768, 2021.
- Stella Maris Ranuncolo, Jose M Polo, Jamil Dierov, Michael Singer, Tracy Kuo, John Greally, Roland Green, Martin Carroll, and Ari Melnick. Bcl-6 mediates the germinal center b cell phenotype and lymphomagenesis through transcriptional repression of the dna-damage sensor atr. *Nature immunology*, 8(7):705–714, 2007.
- Masayuki Kuraoka, T Matt Holl, Dongmei Liao, Mandy Womble, Derek W Cain, Alexander E Reynolds, and Garnett Kelsoe. Activation-induced cytidine deaminase mediates central tolerance in b cells. *Proceedings of the National Academy of Sciences*, 108(28):11560– 11565, 2011.
- Antony B Holmes, Clarissa Corinaldesi, Qiong Shen, Rahul Kumar, Nicolo Compagno, Zhong Wang, Mor Nitzan, Eli Grunstein, Laura Pasqualucci, Riccardo Dalla-Favera, et al. Single-cell analysis of germinal-center b cells informs on lymphoma cell of origin and outcome. Journal of Experimental Medicine, 217(10), 2020.

- 52. Dan Suan, Nike J Kräutler, Jesper LV Maag, Danyal Butt, Katherine Bourne, Jana R Hermes, Danielle T Avery, Clara Young, Aaron Statham, Michael Elliott, et al. Ccr6 defines memory b cell precursors in mouse and human germinal centers, revealing light-zone location and predominant low antigen affinity. *Immunity*, 47(6):1142–1153, 2017.
- Sean P Saunders, Erica GM Ma, Carlos J Aranda, and Maria A Curotto de Lafaille. Nonclassical b cell memory of allergic ige responses. *Frontiers in immunology*, 10:715, 2019.
- Lyssia Belarif, Caroline Mary, Lola Jacquemont, Hoa Le Mai, Richard Danger, Jeremy Hervouet, David Minault, Virginie Thepenier, Veronique Nerrière-Daguin, Elisabeth Nguyen, et al. II-7 receptor blockade blunts antigen-specific memory t cell responses and chronic inflammation in primates. *Nature communications*, 9(1):1–13, 2018.
- Jon R Kettenring. Canonical analysis of several sets of variables. *Biometrika*, 58(3):433– 451, 1971.
- Alicia N Schep, Beijing Wu, Jason D Buenrostro, and William J Greenleaf. chromVAR: inferring transcription-factor-associated accessibility from single-cell epigenomic data. *Nature methods*, 14(10):975–978, 2017.
- Sorim Nam and Jong-Seok Lim. Essential role of interferon regulatory factor 4 (irf4) in immune cell development. Archives of pharmacal research, 39(11):1548–1555, 2016.
- Jonathan P Katz, Nathalie Perreault, Bree G Goldstein, Catherine S Lee, Patricia A Labosky, Vincent W Yang, and Klaus H Kaestner. The zinc-finger transcription factor klf4 is required for terminal differentiation of goblet cells in the colon. 2002.
- Zhigao Wang, Da-Zhi Wang, GC Teg Pipes, and Eric N Olson. Myocardin is a master regulator of smooth muscle gene expression. *Proceedings of the National Academy of Sciences*, 100(12):7129–7134, 2003.
- Shuxiao Chen, Sizun Jiang, Zongming Ma, Garry P Nolan, and Bokai Zhu. One-way matching of datasets with low rank signals. arXiv preprint arXiv:2204.13858, 2022.
- Noah F Greenwald, Geneva Miller, Erick Moen, Alex Kong, Adam Kagel, Thomas Dougherty, Christine Camacho Fullaway, Brianna J McIntosh, Ke Xuan Leow, Morgan Sarah Schwartz, et al. Whole-cell segmentation of tissue images with human-level performance using large-scale data annotation and deep learning. *Nature biotechnology*, 40(4):555–565, 2022.
- Chenfei Wang, Dongqing Sun, Xin Huang, Changxin Wan, Ziyi Li, Ya Han, Qian Qin, Jingyu Fan, Xintao Qiu, Yingtian Xie, et al. Integrative analyses of single-cell transcriptome and regulome using maestro. *Genome biology*, 21(1):1–28, 2020.
- Yingxin Lin, Tung-Yu Wu, Sheng Wan, Jean YH Yang, Wing H Wong, and YX Wang. scjoint integrates atlas-scale single-cell rna-seq and atac-seq data with transfer learning. *Nature Biotechnology*, 40(5):703–710, 2022.
- Gioele La Manno, Kimberly Siletti, Alessandro Furlan, Daniel Gyllborg, Elin Vinsland, Alejandro Mossi Albiach, Christoffer Mattsson Langseth, Irina Khven, Alex R Lederer, Lisa M Dratva, et al. Molecular architecture of the developing mouse brain. *Nature*, 596(7870): 92–96, 2021.
- Mo Huang, Zhaojun Zhang, and Nancy R Zhang. Dimension reduction and denoising of single-cell rna sequencing data in the presence of observed confounding variables. *bioRxiv*, 2020.