COEM: Cross-Modal Embedding for MetaCell Identification

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Abstract

Metacells are disjoint and homogeneous groups of single-cell profiles, representing discrete and highly granular cell states. Existing metacell algorithms tend to use only one modality to infer metacells, even though single-cell multi-omics datasets profile multiple molecular modalities within the same cell. Here, we present Cross-Modal Embedding for MetaCell Identification (COEM), which utilizes an embedded space leveraging the information of both scATAC-seq and scRNA-seq to perform aggregation, balancing the trade-off between fine resolution and sufficient sequencing coverage. COEM outperforms the state-of-the-art method SEACells by efficiently identifying accurate and well-separated metacells across datasets with continuous and discrete cell types. Furthermore, COEM significantly improves peak-to-gene association analyses, and facilitates complex gene regulatory inference tasks.

1. Introduction

Recent advances in single-cell technologies have enabled measuring chromatin accessibility and gene expression levels simultaneously at single-cell resolution. Various sequencing technologies, such as sci-CAR (Cao et al., 2018), SNARE-seq (Chen et al.), SHARE-seq (Ma et al., 2020), and 10X Genomics Multiome, have been developed and applied to comprehensive studies of cell heterogeneity, developmental dynamics, and cis-regulatory elements (CREs) (Chu et al., 2022). However, extreme data sparsity is observed in Single-cell RNA sequencing (scRNA-seq) and Single-cell sequencing assay for transposase-accessible chromatin (scATAC-seq) data. This impedes cis-regulation inference at the single-cell level.

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Figure 1. Violin plot of Pearson correlation between metacell-aggregated gene expression and accessibility on (A) CD34+ bone marrow dataset and (B) T-cell depleted bone marrow dataset. There are a considerable number of negative peak-to-gene associations in metacells identified by SEACell with scATAC-seq, while use of COEM-identified metacells improves peak-to-gene correlation.

High-throughput single-cell profiling of biological samples typically leads to repetitive sampling of highly similar and statistically equivalent cells. The concept of metacells (Baran et al., 2019) has been proposed to maintain statistical confidence. Metacells are sets of scRNA-seq cell profiles representing distinct, highly granular cell states. Metacell aggregates, with their sufficient sequencing coverage, help reduce sparsity-derived problems in downstream analyses.

The MetaCells algorithm (Baran et al., 2019) infers metacells based on the partition of k-nearest neighbor (KNN) similarity graphs on scRNA-seq data only, but it fails on scATAC-seq data. The SEACells approach (Single-cell aggregation of cell-states) (Persad et al., 2022) is designed to identify metacells in RNA or ATAC modality. It significantly improves peak-to-gene associations, i.e. between chromatin accessibility peak and gene expression. However, SEACells identifies metacells based on only one modality: holding onto a strict biological assumption that accessible chromatin is associated with active transcription consistently, despite different data modalities possibly exhibiting a time lag be-
between chromatin remodeling of a gene and its transcription. The occurrence of this “time lag” manifests when:

1. Chromatin opens before transcription initiates
2. mRNA degrades follows chromatin closing (Li et al., 2021).

So, it is not surprising that we observe negative peak-to-gene associations across metacells, as Fig. 1 shows. We aim to investigate that integrated analysis of scATAC-seq and scRNA-seq data to improve our ability to discover and characterize cell states, while avoiding biased associations.

Our new algorithm, COEM, is designed to integratively analyze multi-omic sequencing single-cell data (e.g., gene expression and chromatin accessibility measured in the same cell). COEM addresses the fusion challenge (Baltrusaitis et al., 2018) by first learning joint low-dimensional latent representations for scRNA-seq and scATAC-seq data through a multi-view, multi-modal variational auto-encoder (VAE) model; further utilizing a spectral clustering approach, based on graph connectivity, to identify metacells representing highly granular, distinct cell states on both RNA and chromatin levels. We found COEM to be more accurate than SEACells in metacell identification across datasets with discrete cell types and continuous developmental trajectories. Also, COEM-identified metacells substantially improve the peak-to-gene association analysis and empower CREs prediction.

2. Method

2.1. Multi-Modal Joint Embedding

COEM learns a low dimensional representation from the joint scRNA-seq and scATAC-seq data, measured in the same cell. We adopt a modified multi-view VAE to extract a common latent space of the two data modalities. This is similar to previous work in VAEs representing the single-cell sequence data (Lopez et al., 2018; Li et al., 2022; Ding et al., 2018; Mao et al., 2020; Xu et al., 2020) in less noisy low dimensions. The generative model of the VAEs is derived as follows. Given a cell, the scRNA gene expression $x \in \mathcal{X}$, scATAC chromatin accessibility data $y \in \mathcal{Y}$, cell type $c$, and the common latent representation $z$, we have the following probability densities,

$$p_\theta(x, z | c) = p_\theta(x | z, c)p_\theta(z | c) \tag{1}$$

where $\theta$ represents the parameters of the generative models.

The likelihood $p_\theta$ of $x$ and $y$ depends on the common latent space $z$ and the cell type $c$. With the assumption that the distribution of observed scRNA-seq $x$, scATAC-seq $y$ are independent given the common latent representations $z$ and cell type $c$, $x \perp \perp y | z, c$. So, we have following joint generative model,

$$p_\theta(x, y, z | c) = p_\theta(x, y | z, c)p_\theta(z | c) \tag{2}$$

Moreover, we have the following Evidence Lower Bound (ELBO) for two modalities with conditional variational approximation $q_\phi$,

$$\log p_\theta(x | c) \geq -KL(q_\phi(z | x, c))|p_\theta(z | c)) + \mathbb{E}_{q_\phi(z | x, c)}(\log p_\theta(x | z, c)) \tag{3}$$

Normally, we use an inference network (encoder) to learn the variational approximation $q_\phi$ and a generative network (decoder) to learn $p_\theta$. Particularly, we assume $p_\theta(x | z, c)$ is a zero-inflated Poisson (ZIP) distribution, $p_\theta(y | z, c)$ is a negative binomial (NB) distribution, and $p_\theta(z | c)$ is a Gaussian Mixture Model $z \sim \mathcal{N}(\mu_c, \Sigma_c)$ (Li et al., 2022; Lopez et al., 2018). Given the proposed joint-embedding generative model, we adopt the scMVP (Li et al., 2022) architecture to model the generative network and inference network. Specifically, there is a two-channel attention-based encoder network for the scRNA-seq and scATAC-seq inputs, then concatenating them to derive the posterior distribution $p_\phi(z | x, y, c)$. Next, the imputed scRNA and scATAC profiles are reconstructed by an attention based two-channel decoder network to compute the likelihood $p_\theta(x, y | z, c)$.

2.2. Metacells Identification

Similar to previous work (Baran et al., 2019; Persad et al., 2022), COEM is a graph-based approach that employs spectral clustering to compute metacells. To be more specific, COEM consists of the following four steps:

1. Apply cross-modal joint embedding VAE to learn low common representations $z$ to encode the common information of scRNA-seq and scATAC-seq.
2. Build a $k$-nearest neighbors (KNN) graph on the low dimensional space $z$ based on Euclidean distances.
3. Then similarity matrix is constructed from the previous KNN graph by connectivity. The similarity matrix is based on radial basis function (RBF) kernel to encode the non-linear relations in the $z$.
4. Utilize spectral clustering to identify the cluster and to subsequently aggregate metacells as in (Persad et al., 2022).

This is one of the differences between COEM and SEACells (Persad et al., 2022), which uses archetypal analysis for metacell identification, instead of spectral clustering. While they both are graph based methods, instead of the
Figure 2. The main steps of COEM. 1. The joint profiled scRNA-seq and scATAC-seq are fed to the modified VAE to extract a common latent representation $z$. 2. Then a similarity matrix are constructed based on $z$ and KNN graphs. 3. Spectral clustering is applied to identify the higher granularity clusters where Metacells belongs to. 4. The Metacells features are aggregated for each clusters. 5. Metacells are available for downstreaming tasks.

| METHODS | CD34 | sciCAR | 10xPBMC | SNARE CELLLINE |
|---------|------|--------|---------|----------------|
| SPECTRAL | 23.7 | 6.3    | 45.2    | 1.7            |
| ARCHETYPAL | 191.5 | 103.6  | 277.4   | 28.0           |

Table 1. Running Time (seconds). We use the same low dimensional embeddings $z$ to compare archetypal analysis (used in SEACells) and spectral clustering (used in COEM).

convexity assumption of archetypal analysis of SEACells, spectral clustering of COEM purely utilizes the graphs’ connectivity information and enjoys a faster running speed. We empirically validate that spectral clustering can improve metacell identification and running time.

3. Experiments

3.1. Evaluation Metrics

As in the SEACells paper (Persad et al., 2022), we use compactness, separation, and cell-type purity for the metacell benchmark. Compactness measures how homogeneous the cells within a metacell are. Separation assesses how metacells are distinct from each other. Cell-type purity, instead, measures the consistency of cell-types among cells that constitute a metacell. We computed diffusion components using principal components from RNA assay (RNA_PCA) or singular values singular values from ATAC assay (ATAC_SVD), and then quantified the compactness and separation of metacells. Since SEACells identifies metacells from one modality, we denote them as SeaC-ATAC and SeaC-RNA.

3.2. Results

We use five datasets for evaluation: sci-CAR cell line (Cao et al., 2018), SNARE-seq cell line (Chen et al.), 10X Genomics Multiome PBMC, 10X Multiome CD34+ bone marrow (hematopoietic stem and progenitor cells) and 10X Multiome T-cell depleted bone marrow (Persad et al., 2022). As showed in Table 2 and 3, and Table 4, 5, and 6 in appendix, COEM outperforms SEACells in most cases.

The sci-CAR dataset profiles well-labeled cells, including cell lines 293T, 3T3, a 293T/3T3 cell mixture, and A549 cells after 0, 1, or 3 hours of dexamethasone treatment. We evaluate cell-type purity in sci-CAR dataset, and metacells from COEM algorithm have greater purity (0.91) than SeaC-ATAC (0.67) and SeaC-RNA (0.74), which suggests that leveraging two modalities improves our ability to define cellular states, whereas archetypal analysis on joint embedding representations cannot guarantee high purity of metacells shown in Table 2.

Then, we measure compactness and separation across datasets with discrete cell types. Generally, COEM-based methods separate metacells better than SeaC-ATAC and SeaC-RNA shown in Tables 2, 3 and 4 to 6. But they may have higher compactness based on only one modality, since COEM balances cell variability from two modalities at the same time. The metacells identified by SEACells have lower compactness because metacell identification and compactness evaluation are based on the diffusion components from a single modality. But those metacells may not be well-
Table 2. Metacell Evaluation on sci-CAR celline dataset. Greater separation, lower compactness, and higher cell-type purity indicate better performance. COEM identified metacells with higher purity and separation.

| METHODS | RNA_PCA  | ATAC_SVD |
|---------|----------|----------|
|         | SEPARATION | COMPACTNESS | SEPARATION | COMPACTNESS |
| COEM    | 0.235 ±0.015 | 0.193 ±0.017 | 0.091 ±0.006 | 0.028 ±0.002 |
| SEAC-ATAC | 0.196 ±0.007 | 0.686 ±0.035 | 0.086 ±0.002 | 0.004 ±0.001 |
| SEAC-RNA | 0.177 ±0.051 | 0.150 ±0.062 | 0.041 ±0.015 | 0.032 ±0.014 |

Table 3. Metacell Evaluation on T-cell depleted bone marrow dataset. COEM metacells have higher separation than SEACell's.

| METHODS | RNA_PCA  | ATAC_SVD |
|---------|----------|----------|
|         | SEPARATION | COMPACTNESS | SEPARATION | COMPACTNESS |
| COEM    | 0.801 ±0.022 | 0.419 ±0.001 | 0.704 ±0.041 | 0.292 ±0.217 |
| SEAC-ATAC | 0.723 ±0.024 | 1.061 ±0.140 | 0.666 ±0.01 | 0.662 ±0.019 |
| SEAC-RNA | 0.704 ±0.056 | 0.640 ±0.153 | 0.566 ±0.070 | 0.547 ±0.291 |

4. Discussion

Our experiments show that integrating information from both scRNA-seq and scATAC-seq is beneficial for identifying accurate and robust metacells from single-cell data. Especially joint-embedding, low-dimensional representations can enable the identification of cellular states with high cell-type purity and well-separated metacells. On top of that, the proposed COEM algorithm enjoys a faster running time. Moreover, it avoids high numbers of negative peak-to-gene associations that are false. Hence, the joint profiled sequence data are able to depict a comprehensive landscape of cells’ stages. Furthermore, (Bilous et al., 2021) has demonstrated that metacells inferred from scRNA-seq are compatible with RNA velocity model. Therefore, COEM is also able to be applied to multi-omic velocity model for estimating epigenomic and transcriptomic dynamics.

In future work, we would explore more machine learning techniques for multi-modal sequence data integration and metacell identification. For example, scATAC-seq and scRNA-seq are naturally causally related. It is interesting to utilize this unique relation to learn more robust, causally sufficient, and efficient representations (Blei, 2020; Wang & Jordan, 2021; Veitch et al., 2021; Dou et al., 2022; Tabib et al., 2021). Besides, the results are promising for use of an optimal transport approach to model cellular stage changes across different modalities on the metacell level (Schiebinger et al., 2019).
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A. Appendix

In this section, we show the results of additional experiments on datasets including Snare Cellline, 10x_PBMC, and 10X Multiome CD34+ bone marrow.

Table 4. Metacell Evaluation on SNARE-seq celline dataset

| RNA_PCA | METHODS | SEPARATION | COMPACTNESS |
|---------|---------|------------|-------------|
|         | COEM    | 0.280 ± 0.015 | 0.856 ± 0.185 |
|         | SEAC-ATAC | 0.191 ± 0.013 | 0.617 ± 0.112 |
|         | SEAC-RNA  | 0.067 ± 0.011 | 0.067 ± 0.004 |

Table 5. Metacell Evaluation on CD34+ bone marrow dataset

| RNA_PCA | METHODS | SEPARATION | COMPACTNESS |
|---------|---------|------------|-------------|
|         | COEM    | 0.266 ± 0.011 | 0.035 ± 0.006 |
|         | SEAC-ATAC | 0.203 ± 0.013 | 0.037 ± 0.007 |
|         | SEAC-RNA  | 0.256 ± 0.007 | 0.026 ± 0.003 |

Table 6. Metacell Evaluation on PBMC dataset

| RNA_PCA | METHODS | SEPARATION | COMPACTNESS |
|---------|---------|------------|-------------|
|         | COEM    | 0.515 ± 0.127 | 0.349 ± 0.115 |
|         | SEAC-ATAC | 0.406 ± 0.129 | 3.315 ± 1.21 |
|         | SEAC-RNA  | 0.428 ± 0.133 | 0.294 ± 0.046 |

| ATAC_SVD | METHODS | SEPARATION | COMPACTNESS |
|---------|---------|------------|-------------|
|         | COEM    | 0.159 ± 0.021 | 0.124 ± 0.008 |
|         | SEAC-ATAC | 0.097 ± 0.010 | 0.004 ± 0.001 |
|         | SEAC-RNA  | 0.095 ± 0.0188 | 0.037 ± 0.012 |

| ATAC_SVD | METHODS | SEPARATION | COMPACTNESS |
|---------|---------|------------|-------------|
|         | COEM    | 0.318 ± 0.014 | 0.136 ± 0.010 |
|         | SEAC-ATAC | 0.284 ± 0.015 | 0.025 ± 0.003 |
|         | SEAC-RNA  | 0.244 ± 0.123 | 0.115 ± 0.021 |

| ATAC_SVD | METHODS | SEPARATION | COMPACTNESS |
|---------|---------|------------|-------------|
|         | COEM    | 0.181 ± 0.038 | 0.030 ± 0.013 |
|         | SEAC-ATAC | 0.169 ± 0.024 | 0.022 ± 0.008 |
|         | SEAC-RNA  | 0.034 ± 0.009 | 0.141 ± 0.005 |