Gene expression

**CellVGAE: an unsupervised scRNA-seq analysis workflow with graph attention networks**

David Buterez 1,*, Ioana Bica 2,3, Ifrah Tariq 4, Helena Andrés-Terré 1 and Pietro Liò 1

1Department of Computer Science and Technology, University of Cambridge, Cambridge CB3 0FD, UK, 2Department of Engineering Science, University of Oxford, Oxford OX1 3PJ, UK, 3The Alan Turing Institute, London NW1 2DB, UK and 4Computational and Systems Biology Program, Department of Biological Engineering, Massachusetts Institute of Technology, Cambridge, MA 02142, USA

*To whom correspondence should be addressed.

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**Abstract**

**Motivation:** Single-cell RNA sequencing allows high-resolution views of individual cells for libraries of up to millions of samples, thus motivating the use of deep learning for analysis. In this study, we introduce the use of graph neural networks for the unsupervised exploration of scRNA-seq data by developing a variational graph autoencoder architecture with graph attention layers that operates directly on the connectivity between cells, focusing on dimensionality reduction and clustering. With the help of several case studies, we show that our model, named CellVGAE, can be effectively used for exploratory analysis even on challenging datasets, by extracting meaningful features from the data and providing the means to visualize and interpret different aspects of the model.

**Results:** We show that CellVGAE is more interpretable than existing scRNA-seq variational architectures by analysing the graph attention coefficients. By drawing parallels with other scRNA-seq studies on interpretability, we assess the validity of the relationships modelled by attention, and furthermore, we show that CellVGAE can intrinsically capture information such as pseudotime and NF-kB activation dynamics, the latter being a property that is not generally shared by existing neural alternatives. We then evaluate the dimensionality reduction and clustering performance on 9 difficult and well-annotated datasets by comparing with three leading neural and non-neural techniques, concluding that CellVGAE outperforms competing methods. Finally, we report a decrease in training times of up to $\times$ 20 on a dataset of 1.3 million cells compared to existing deep learning architectures.

**Availability and implementation:** The CellVGAE code is available at https://github.com/davidbuterez/CellVGAE.

**Contact:** db804@cam.ac.uk

**Supplementary information:** Supplementary data are available at Bioinformatics online.

1 Introduction

scRNA-seq allows gene expression to be quantified at the level of individual cells; however, it introduces new challenges, as technical and biological limitations contribute to noisier and more complex data than previous sequencing techniques (Chen et al., 2019). A recent study found over 19 000 scRNA-seq studies as of October 2020 (Pasquini et al., 2021), and another identified over 1000 scRNA-seq tools as of September 2021 (Zappia and Theis, 2021), reflecting rapid iterative progress in the field. Non-neural tools are the most widespread, most likely for their simplicity and interpretability. However, it is typically beyond the capabilities of algorithms such as t-SNE (van der Maaten and Hinton, 2008) and UMAP (McInnes et al., 2020) to perform interpretable dimensionality reduction from the high-dimensional gene space. Consequently, this task, as well as denoising and imputation, are often handled by neural networks, especially Variational Autoencoders (VAEs) such as scVAE (Grombetch et al., 2020), Deep Count Autoencoder (Eraslan et al., 2019), scVI (Svensson et al., 2020), (Lopez et al., 2018) and DiffVAE (Bica et al., 2020). A popular alternative is graph-based clustering, popularized by Seurat (Stuart et al., 2019). Recent developments such as self-assembling manifolds (SAM) (Tarashansky et al., 2019) have since largely superseded Seurat and are competitive with neural networks.

In this work, we investigate a new machine learning approach with applications to dimensionality reduction and clustering. Based on the recent interest in both graph-based scRNA-seq clustering and graph neural networks, we propose a neural model that is built upon the variational graph autoencoder (VGAE) (Kipf and Welling, 2016b) with graph attention layers (GAT) (Velicković et al., 2018), named CellVGAE. Compared to other neural models which learn exclusively from the gene expression values, CellVGAE leverages
the connectivity between cells (represented as a graph) as an inductive bias to perform convolutions on a non-Euclidean structure, thus subscribing to the geometric deep learning paradigm. We use k-nearest neighbour (KNN) and Pearson correlation graphs (referred to as PKNN), based on their efficient implementations and widespread use. While methods based on Euclidean distance are popular, Pearson correlation has recently been argued to be preferable for scRNA-seq data (Kim et al., 2019).

There are several reasons motivating the introduction of a graph neural network (GNN) methodology. Firstly, the graph can support learning, acting as a valuable inductive bias and allowing the model to exploit relationships that are impossible or harder to model by the simpler dense layers. Secondly, graphs are generally more interpretable and visualizable; the GAT (Graph Attention Network) framework made important steps in bringing these desirable features to machine learning, a trait not shared by traditional (non-graph) methods. The attention mechanism is argued to (i) improve task performance, (ii) stabilize learning (reduce variance), and (iii) provide an extra layer of interpretability. Thirdly, by jointly using the variational autoencoder and graph neural networks, we allow future studies to exploit advances in both of these active research subjects, such as the newly published GATv2 (Brody et al., 2021) convolutional layer (already integrated into CellVGA).

For our evaluation of CellVGAE we compare with existing methods that: (i) achieve state-of-the-art performance on dimensionality reduction and clustering (unsupervised learning), (ii) are open source and (iii) are recognized in the scRNA-seq community. We select SAM and DiffVGA as recent advances in their respective fields and for their performance, and scVI for its popularity.

2 Materials and methods

In this work, we assume simple graphs (undirected, unweighted, without loops or multiple edges), defined as a tuple $G = (\mathcal{V}, \mathcal{E})$, where $\mathcal{V}$ is the set of vertices or nodes $\{v_1, v_2, \ldots \}$ and $\mathcal{E}$ is the set of edges between nodes, $\mathcal{E} \subseteq \mathcal{V} \times \mathcal{V}$. A common representation is given by a graph’s adjacency matrix $A$ (whose elements $a_{ij} = 1$ if $(v_i, v_j) \in \mathcal{E}$, with $a_{ij} = 0$ otherwise). For GNNs, we also assume that $D$-dimensional node features are represented by a $N \times D$ matrix $X$, where $N = |\mathcal{V}|$.

2.1 Variational graph autoencoder

The variational graph autoencoder (VGAE) is an unsupervised framework introduced in (Kipf and Welling, 2016b). Like the standard VAE, the VGAE has two components: an encoder and a decoder, which are trained to learn latent variables $z$, aggregated in an $N \times L$ matrix $Z$, where $L$ is the number of latent dimensions. The encoder or inference model is similar to the VAE, but does not depend only on the signal $X$, but also on the graph $A$:

$$q(Z|X, A) = \prod_{i=1}^{N} q(z_i|X, A) = N(z_i|\mu_i, \text{diag}(@^2)) \quad (1)$$

In the original formulation, the parameters $\mu, \sigma$ are learned by graph convolutional networks (GCN) (Kipf and Welling, 2016a); formally $\mu = \text{GCN}_{\mu}(X, A)$ and $\log \sigma = \text{GCN}_{\sigma}(X, A)$, where $X$ are features learnt by previous convolutional layers.

The decoder, or generative model, simply reconstructs an adjacency matrix using the inner product of latent variables:

$$p(A|Z) = \prod_{i=1}^{N} \prod_{j=1}^{N} p(A_{ij}|z_i, z_j) = p(A_{ij} = 1|z_i, z_j) = \sigma(z_i^T z_j) \quad (2)$$

Here, $\sigma$ is the sigmoidal function. Also, note that only the graph structure is reconstructed (not the node features). The loss function is of the form:

$$\mathcal{L} = \mathbb{E}_{q(Z|X, A)}[\log p(A|Z)] - D_{\text{KL}}[q(Z|X, A) \parallel p(Z)]$$

where the two components are the reconstruction loss $\mathcal{L}_{\text{RECON}}$ and a regularization term $\mathcal{L}_{\text{REG}}$. In the standard VGAE framework, the regularization is given by the Kullback-Leibler divergence $D_{\text{KL}}$ between $q(\cdot)$ and $p(\cdot)$.

2.2 Graph attention networks

Graph attention networks (GAT) are a powerful neural framework for graph-structured data, initially designed for supervised and semi-supervised learning, manifested through the graph attention layer, capable of performing self-attention on graph nodes. Given a set of node features $h \in \mathbb{R}^D$, a shared linear transformation parameterized by a weight matrix $W$ is applied to all nodes, followed by a learnable attention mechanism applied pairwise to the scaled node features: $e_v = d(W_h, W_{h'})$, where $W \in \mathbb{R}^{D \times D}$ and $d : \mathbb{R}^D \times \mathbb{R}^D \rightarrow \mathbb{R}$ assuming $D' > D$. Here, the attention mechanism can be applied independently $K$ times (multi-head attention), in two ways:

$$b'_i = \sum_{k=1}^{K} \sigma \left( \sum_{j \in \mathcal{N}_i} \alpha_{k}^{j} W^{k} h_{j} \right) \quad b'_i = \sigma \left( \sum_{k=1}^{K} \sum_{j \in \mathcal{N}_i} \alpha_{k}^{j} W^{k} h_{j} \right) \quad (3)$$

The left formulation is appropriate for the hidden (inner) layers as it outputs the concatenated hidden representations of dimension $K \cdot D'$. The variation on the right uses the mean of all the attention heads and is suitable for the last (output) layer.

2.3 CellVGAE architecture

CellVGAE’s encoder is based on graph attention layers. Using the established notation, where $X_0$ is the initial set of node features and $GAT_{1}^{(K)}$ represents the $r$th layer with $K$ attention heads, we define the first and subsequent inner neural layers for an architecture with $N$ inner layers:

$$X_i = \begin{cases} GAT^{(K)}_{1}(X_{0}, A) & \text{if } i = 1 \\ GAT^{(K)}_{i}(\text{ReLU}(X_{i-1}), A) & \text{if } 1 < i \leq N \end{cases} \quad (4)$$

The inner layers concatenate the representations learnt by multi-head attention, thus after the first layer, the number of output dimensions is $K \cdot D$, with $D_i$ the output dimension for each layer. The number of heads $K$ can be different for each layer. The final two layers learn the parameters $\mu, \sigma$ and follow the second branch of Equation 6:

$$\mu = \text{GAT}^{(K)}_{N}(\text{ReLU}(X_{N}), A) \ \log \sigma = \text{GAT}^{(K)}_{N}(\text{ReLU}(X_{N}), A) \quad (5)$$

with the exception that they use mean instead of concatenation. Apart from their demonstrated performance, GAT layers are appealing for several reasons. Since the coefficients indicate the contribution of nodes in a pairwise manner, they can be visualized as a weighted graph between the cells. We confirm that CellVGAE attention models non-spurious and accurate relationships in Supplementary Information SD and SE. Secondly, as GAT layers use
a linear transformation under the hood, a traditional layer weights analysis can be carried out, in a manner very similar to DiffVAE (detailed in Section 3). Currently, CellVGAE provides the users with the choice of GCN, GAT and GATv2 layers. A comparative study is provided in Supplementary Information SM.

We use the same inner product decoder as the original VGE, as well as the standard loss that combines regularization and reconstruction terms:

\[ \mathcal{L}(\mathbf{x}) = \mathcal{L}_{\text{REG}} + \mathbb{E}_{z \sim q(\mathbf{z} | \mathbf{x}, \mathbf{a})} \log p(\mathbf{a} | \mathbf{z}) \]  

with the mention that \( \mathcal{L}_{\text{REG}} \) can use any appropriate loss from the literature that minimizes the divergence between the learnt and prior distributions. CellVGAE can be used with the KL (Kingma and Welling, 2014) and MMD (Zhao et al., 2017) losses, normally implemented as:

\[ \mathcal{L}_{\text{REG}} = -D_{\text{KL}}[q(\mathbf{z} | \mathbf{x}, \mathbf{a}) \parallel p(\mathbf{z})] \]

\[ \mathcal{L}_{\text{REG}} = -D_{\text{MMD}}(q(\mathbf{z}) \parallel p(\mathbf{z})) \]  

\( \mathcal{L}_{\text{RECON}} \) is implemented as a binary cross entropy loss with negative sampling, where \( \hat{y}^{\text{pos}} \) is the decoder output for positive (real) edges, \( \hat{y}^{\text{neg}} \) is the decoder output for (randomly sampled) negative edges, and \( E \) is the total number of edges:

\[ \mathcal{L}_{\text{RECON}} = -\frac{1}{E} \left( \sum_i \log (\hat{y}_i^{\text{pos}}) + \sum_i \log (1 - \hat{y}_i^{\text{neg}}) \right) \]  

Optionally, CellVGAE can use a feature reconstruction neural network \( f_{\text{RECON}} \) in the decoder, addressing a limitation of the original VGE framework, such that the loss becomes:

\[ \mathcal{L}^*(\mathbf{x}) = \mathcal{L}(\mathbf{x}) + \text{MSE}(f_{\text{RECON}}(\mathbf{z}), \mathbf{x}) \]  

where MSE is the mean squared error.

2.4 CellVGAE workflow

The key idea is augmenting the gene expression matrix with cell connectivity information in the form of a graph. In this work, the used graphs take the form of KNN graphs based on Euclidean distance and PKNN graphs built using Pearson correlation, although the CellVGAE interface allows other metrics (such as Manhattan distance, cosine similarity, etc.) and completely custom graphs to be provided. In this formulation, the gene expression values associated with a cell are used as node features for the VGE. For this article, we log-normalized the expression matrix, a widespread practice in the scRNA-seq community (Boosehaghi and Pachter, 2021) and applied min-max scaling, which is also common in machine learning applications. Furthermore, for practical reasons such as training efficiency and balancing the number of features compared to the number of cells we generally select a number of top highly variable genes (HVGs) to use as node features, a standard practice for both neural and non-neural tools (Yip et al., 2019). The graph can be built from the original high-dimensional data, projections of the data to a lower number of dimensions (for example using Principal Component Analysis—PCA), or from a number of HVGs that can be different from the number of HVGs used as node features. Since we frequently use this last hyperparameter we refer to it as KHVG, the number of HVGs used to build the graph. As an unsupervised method, CellVGAE training is performed on the whole dataset. For evaluation purposes, a held-out test set is first used for each dataset to determine convergence and prevent overfitting, as well as suggest optimal graph settings (i.e. k and KHVG).

By propagating the transcriptomics information according to the cell connections, CellVGAE learns to reconstruct the original graph from the lower-dimensional latent space (Supplementary Information SM), producing high-quality compressed representations that can be used for downstream analysis, detailed in Section 3. The process is schematically represented in Figure 1.

3 Results

3.1 Accurate identification of clusters on challenging datasets

One of the motivating factors behind the SAM algorithm is the inability of the existing methods to analyse a novel scRNA-seq dataset of Schistosoma mansoni (including cell types such as \( \epsilon, \delta \) and \( \mu \)). More specifically, Tarashansky et al. (2019) showed that commonly used methods like PCA, Seurat and SIMLR (Wang et al., 2017) fail to distinguish any cluster formations. In turn, this renders tasks like cell (sub)type identification and finding marker genes difficult using current tools. To formalize this notion of difficulty, the authors introduce an unsupervised metric called network sensitivity, which measures the changes in cell-to-cell distances on randomly selected subsets of the gene expression matrix. An intuitive explanation of a high sensitivity score is that changes in the used features lead to a significantly different topological network. On the other hand, datasets robust to the said feature selection have many genes (signals) reinforcing the same structure, thus resulting in a low sensitivity score. The exact sensitivity ranking and numerical values for the S.mansoni, Macrophages and the other 9 datasets used in Results are available in (Tarashansky et al., 2019) and reproduced in Supplementary Figure S1. The S.mansoni dataset tops the network sensitivity chart by a significant lead, followed by the Macrophages dataset, which is in turn about twice as difficult as the next most challenging dataset.

3.1.1 The S.mansoni dataset

Figure 2 presents the UMAP plots of the evaluated dimensionality reduction techniques including multiple configurations of
For the second case study, we analyse the *Macrophages* dataset, where the authors examined the NF-κB activation dynamics and transcriptional profile of 823 macrophages treated with lipopolysaccharide (LPS) (Lane et al., 2017). We follow the methodology of Tarashansky et al. to remove cell cycle effects from 637 cells which were imaged for NF-κB response after 75, 150 and 300 min. The resulting expression matrix is used as input to CellVGAE and the other evaluated neural algorithms. Tarashansky et al. further argue that the cells can be grouped in two clusters by their activated signalling pathways: Myd88 and TRIF (MT) and only Myd88 (M).

As seen in Figure 3, the three illustrated CellVGAE models prefer to first and foremost cluster based on the NF-κB activation dynamics, such that cells belonging to the same chronological grouping are generally in close proximinity. At the same time, the CellVGAE representation respects the two pathway clusters found by SAM. The set of cells imaged after 300 min can be almost completely separated (fourth column in Fig. 3), although a small cluster of ‘300 mins’ cells is separated from the main agglomeration as it is part of the different MT cluster. However, when using less transcriptionomics information but otherwise identical settings (third column), or similarly for the settings of the second column, there is a continuum between the clusters emphasizing the preference to group by time since LPS induction. Thus, the continuous, non-interrupted topology is required to characterize the transcriptional profile and activation dynamics simultaneously.

### 3.1.3 The PBMC3k dataset

The PBMC3k dataset (10x Genomics) is difficult for SAM, DiffVAE and scVI in terms of cluster definition and separation, as illustrated and discussed in Figure 4. More specifically, none of the existing methods clearly separates the existing clusters, while CellVGAE is able to achieve both tight clusters and definitive separation, as shown by the unsupervised HDBSCAN clustering without any unlabelled cells. SAM is the only method with similarly tight clusters, but it is not capable of separating all cell populations even with its best-performing hyperparameters.

### 3.2 CellVGAE captures pseudo-temporal information in continuous blood cell differentiation

As CellVGAE is capable of representing complex biological signals in difficult scRNA-seq datasets (not exclusively based on transcriptionomics profiling), Fig. 3, we now evaluate CellVGAE on a continuous blood cell differentiation dataset consisting of 2730 myeloid progenitors with 3451 genes (Paul et al., 2015), investigating if the model can capture continuous phenotypes. To establish this, it is useful to consider the concept of pseudootime, defined as the ordered gradual change in gene expression and serving as a method to place cells along an inferred differentiation trajectory (Ji and Ji, 2019). It was previously shown that neural implementations such as the Deep Count Autoencoder (DCA) (Eraslan et al., 2019) can produce cell embeddings that respect the diffusion pseudo-time.

Like other neural algorithms and SAM, CellVGAE can successfully order the differentiating cells according to the pseudootime (Fig. 5). Differently, however, we notice that as the quantity of transcriptionomics information is increased, the clusters become
information, although this could be an interesting future direction.

For both figures and datasets, we chose the second layer of the visualization is enabled by plotting a graph of cells, where the attention coefficients give the edge weights. A simple strategy is to select the coefficients that can clearly separate the correct number of clusters, and with a high ARI (Fig. 4), we discover that all 14 Platelet cells are among the top 1% of all cells. In Supplementary Information SD and SE, the representation of this dynamic is particularly interesting since the Platelet cluster is by far the smallest, at just over 0.005% of all cells. In Supplementary Information SD and SE we examined whether the high attention coefficients for Platelets arise due to spurious connections in the KNN graph, and compared with PAGA (Wolf et al., 2019), a well-known algorithm for detecting and visualizing connections between cell clusters. The analyses revealed little contribution from the KNN graph structure alone, as well as similar, strong connections identified by PAGA. The other prevalent connections involve CD4 T, CD8 T, and FCGR3A cells. For this particular graph, we notice the existence of long paths and a relatively low number of connected components.

For the ‘Darmanis’ dataset, all 8 cell types are encountered in the relatively heterogeneous graph of Figure 6d, suggesting that multiple different cell types can inform the classification of a single cell.

progressively more separated, starting from a single continuous cluster and reaching three clearly delimited clusters. The behaviours of DiffVAE under the same transcriptomics settings are illustrated in Supplementary Figures S3 and S4, respectively. DiffVAE successfully identifies the single, continuous cluster and can broadly discern two clusters, however not accurately and with no correlation to the amount of transcriptomics information. scVI successfully orders the cells by pseudotime but does not perform any separation. Thus, we recognize that CellVGAE excels at cleanly separating clusters (as in Fig. 2), especially when a large amount of gene expression information is available.

3.3 Graph attention coefficients enhance interpretability

We propose two alternative ways of visualizing the learnt attention coefficients (denoted by $a$). To quickly recapitulate, the attention coefficients measure the contribution of each cell to its neighbours in a pairwise manner (Equation 4). Thus, the numerical value associated with two nodes can be interpreted as an edge. For two nodes $i, j$ the coefficients $a_{ij}, a_{ji}$ are different, resulting in a graph that is weighted and directed. Each graph attention layer has its own independent set of coefficients.

First, we propose mapping the attention coefficients to a node-level representation by averaging over each node’s neighbourhood, acting as an overlay for two-dimensional projections like UMAP and where the highest resulting values indicate the nodes with the largest contribution in the neural model. Secondly, a finer-grained visualization is enabled by plotting a graph of cells, where the attention coefficients give the edge weights. A simple strategy is to select the top $n$ largest coefficients; alternatively and depending on the specific use case, the analysis can exclude certain cell types or focus on a different range of interactions (e.g. the smaller edge weights), providing a variable level of granularity.

We illustrate the two techniques in Figure 6, for two datasets. For both figures and datasets, we chose the second layer of the CellVGAE model and for simplicity have taken the mean across all the attention heads, but the same analysis applies per head and can, in fact, reveal the differences in what each one learns (the individual attention heads are illustrated in Supplementary Information 5N). In the standard GAT framework, the attention heads are not enforced to learn different representations, for example by minimizing mutual information, although this could be an interesting future direction.

For PBMC3k, a non-trivial finding from the UMAP plot in Figure 6a is that the Platelet cells are highly involved at the level of the second layer. In fact, by inspecting the graph representation (Fig. 6c), we discover that all 14 Platelet cells are among the top contributors, with some of the highest-weight edges (attention coefficients). The successful representation of this dynamic is particularly interesting since the Platelet cluster is by far the smallest, at just over 0.005% of all cells. In Supplementary Information SD and SE we examined whether the high attention coefficients for Platelets arise due to spurious connections in the KNN graph, and compared with PAGA (Wolf et al., 2019), a well-known algorithm for detecting and visualizing connections between cell clusters. The analyses revealed little contribution from the KNN graph structure alone, as well as similar, strong connections identified by PAGA. The other prevalent connections involve CD4 T, CD8 T, and FCGR3A cells. For this particular graph, we notice the existence of long paths and a relatively low number of connected components.

For the ‘Darmanis’ dataset, all 8 cell types are encountered in the relatively heterogeneous graph of Figure 6d, suggesting that multiple different cell types can inform the classification of a single cell.
Neurons, which are among the most activated, interact with an Endothelial cell, and another large group, of Fetal quiescent cells, is seen largely interacting with its kind, although it participates in other relationships, mostly with Fetal replicating cells. On this dataset, we notice the presence of self-loops, which might indicate that a cell’s own features are enough to distinguish itself. Overall, the graph is more fragmented (more small, connected components) indicating less cross-talk.

To further our understanding of the attention coefficients, we study if they encode redundant information. More specifically, we selected the top 80 cell pairs from Figure 6c and computed the pair-wise Euclidean distances in the 50-dimensional latent space (Supplementary File S2). We find that the overwhelming majority of cell pairs are distant. Thus, high attention coefficients do not correspond to highly similar cells, meaning that they capture an additional layer of information not present in standard VAEs.

3.4 Retrieval of marker genes using neural weights analysis

To link the latent dimensions to the genes, the neural weights can be analysed, as initially proposed for DiffVAE. Since CellVGAE uses a simple inner product decoder, we focus on the encoder, a possibility analysed, as initially proposed for DiffVAE. Since CellVGAE uses a simple inner product decoder, we focus on the encoder, a possibility analysed, as initially proposed for DiffVAE.

That mark μ cells and finally an achaete-scute transcription factor, astf (Smp_142120) which is highly enriched in ε cells, a subpopulation of ν cells.

3.5 CellVGAE representations correspond to known biological signals

We make a first step towards visualizing what the model has learnt in terms of gene expression, starting with the learnt node embeddings, a matrix E of dimension N × L, where N is the number of cells and L is the latent (output) size, as well as Pν with dimension D × L. We select the vector g corresponding to a target gene of choice out of Pν and perform the multiplication E · g, which gives the CellVGAE expression value, used as the hue parameter when plotting. This type of analysis can suggest the contribution of a gene at the level of each cell, providing a granular view of expression within a cluster.

The proposed approach applies to all VAE-based methods and is not specific to CellVGAE. However, as illustrated in Figure 7, only CellVGAE is sensitive enough to visualize the learnt gene expression patterns, confirming that the model captures relevant biological information. Both CellVGAE and scVI successfully represent clusters enriched for some typical marker genes, such as NKG7, FCG3RA, and LYZ, although the tight cluster formations of CellVGAE aid the visualization. However, other genes lead to much more diffuse scVI representations, for example KLRG1, which should be expressed in NK cells (Wang et al., 2013), or CD74 which should be enriched in B cells (Naeim et al., 2008). DiffVAE does not convincingly capture gene-level information in any of the proposed examples.
3.6 Cell clustering performance on 10 well-annotated datasets

Next, we present an extensive evaluation of CellVGAE, SAM, DiffVAE and scVI on nine challenging and well-annotated scRNA-seq datasets (covering a wide range on the network sensitivity spectrum, Supplementary Information S4) where we show improved clustering performance, as well as on the RETINA dataset of 26.5K cells (Shekhar et al., 2016). Dataset availability and other characteristics are provided in Supplementary Table S6.

All 9 datasets were used to benchmark SAM alongside Seurat, SC3 (Kiselev et al., 2017) and SIMLR in (Tarashansky et al., 2019). Here, we do not reproduce the evaluation for the other algorithms. RETINA is included as it is the largest stand-alone dataset used to evaluate scVI in its publication. We run all four algorithms ourselves and report the results in Table 2. The main metric used to evaluate clustering performance is the adjusted rand index (ARI), a measure of the similarity between two data clusters. It is the standard choice for this type of evaluation and it is defined mathematically in Supplementary Information S5. We also include the silhouette coefficient (SC), similarly defined in Supplementary Information S6. Although we prioritize the ARI as a metric of clustering performance, as well as in RETINA dataset of 26.5K cells (Shekhar et al., 2016). Dataset availability and other characteristics are provided in Supplementary Table S6.

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3.7 Training time and scalability

With the introduction of CellVGAE, we diverge from existing VAE implementations by employing graph convolutions and raise new questions that exploit cell connectivity (CellVGAE and SAM) tend to form more complicated shapes when visualized in two dimensions using UMAP. In particular, one pattern common to both CellVGAE and SAM is the presence of clusters that are broadly linear in shape (SAM in Supplementary Fig. S7a, both in Supplementary Fig. S7c, CellVGAE in Supplementary Fig. S7g, SAM in Supplementary Fig. S7i and SAM in Fig. 5.). Differently from SAM, CellVGAE identifies clusters with topology involving holes and arcs, such as the Beta and Alpha cells from the Muraro dataset (Supplementary Fig. S7i), we observe that CellVGAE separates a cluster with mixed reference cell types, that differentially expresses Calm1, Scg2, Trpm1 and Scgn (Scanpy’s rank_genes_groups() with the ‘t-test’ method). Discarding this cluster (an additional 508 cells), we arrive at an ARI of 0.9775 and an SC of 0.7391, i.e. essentially matching DiffVAE’s ARI.

To complete the analysis, we provide an extensive evaluation of CellVGAE on two additional values of k, k = 3 and k = 10 (Table 2 uses k = 5), using GAT (default for CellVGAE), GCN and GATv2 layers (Supplementary Table S4). We also experimented with an additional decoder component in the form of a dense neural network that reconstructs the gene expression values, as in existing VAE methods, with results reported in Supplementary Table S5. Overall, we find that the other graph settings perform slightly worse but still improve over existing methods. The addition of the decoder neural network does not lead to considerable differences.
questions about the scalability of the method for high-throughput sequencing datasets. An important difference is that CellVGAE uses a single graph that is fully loaded into memory throughout training (the impact in terms of memory utilization is minimal), avoiding the need to mini-batch the data.

We analysed the training time for eight random subsets of an open-source scRNA-seq dataset of 1.3 million mouse brain cells (from 10x Genomics). Combined with the fast GPU implementation of GNNs provided by PyTorch Geometric, CellVGAE offers between $\times 3.5$ to $\times 21$ faster training times than DiffVAE and scVI,
depending on the subset and hardware, as illustrated in Figure 8a. At the same time, CellVGAE GPU is between $18.5 \times 10^3$ to $33.5 \times 10^3$ faster than SAM, a CPU-exclusive algorithm. SAM failed to run for the ‘1M’ and ‘Full’ subsets, limiting its applicability to large-scale datasets.

We further study the training time and resource utilization of CellVGAE with several hyperparameters (Fig. 8b and c) on RETINA. The training time increases sublinearly in the number of graph edges and the number of attention heads (Fig. 8b). Similarly, doubling the hidden layer size produces only minimal increases in training time. On the other hand, the peak memory consumption grows more abruptly with the number of attention heads, increasing roughly linearly when doubling the hidden layer size (Fig. 8c). This is expected, as the parallel attention mechanism stores and updates separate copies of all the attention coefficients. We thus see that CellVGAE offers a trade-off between low training times and high memory usage.

As a finer-grained comparison, we also discovered that the training time of CellVGAE on a CPU is on the same order of magnitude as existing VAE models trained on GPUs (Fig. 8d, on RETINA), enabling the possibility to train large models when video memory is insufficient. Finally, we benchmarked the extremely efficient similarity search library Faiss (Johnson et al., 2017), an optional dependency of the CellVGAE implementation, for KNN graph generation on the same subsets of up to 1.3 million cells (Fig. 8e).

Our analysis using the PyTorch profiler reveals that the KNN graph itself occupies a negligible amount of memory, with occasional attention-related operations (e.g. concatenation, index select) greatly contributing to the peak consumption. As Faiss is GPU-enabled, it can perform exact KNN computations in under 5 min for all eight subsets, with the possibility to scale to billions of vectors with approximative algorithms.

4 Conclusion

We introduced CellVGAE, a machine learning architecture that integrates the benefits of graph-based clustering techniques with recent advancements in neural networks. We showed that CellVGAE performs consistently well in finding accurate, informative clusters, even when applied to complex datasets with subtle signals or when modelling properties not directly related to transcriptional data. The method produced excellent results on nine challenging datasets in terms of the most relevant clustering metrics. Furthermore, we delivered three strategies for interpretability: (i) high-weight gene identification, useful for gene markers and gene set enrichment analysis; (ii) visualization of learnt expression, per-gene; and (iii) a practical interpretation of the attention coefficients. Overall, we showed that the combined use of neural networks and graphs is effective for scRNA-seq data analysis. Lastly, by examining the training time and resource utilization, we concluded that CellVGAE is several times faster than existing VAE implementations on equivalent hardware.

Author contributions

D.B. conceived the presented ideas, designed and ran the experiments, wrote the manuscript text and prepared all the figures. I.B., I.T., H.A.-T. and P.L. helped supervise the project.

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Data Availability Statement

The data underlying this article are available in the article and in its online supplementary material.

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