Factorized linear discriminant analysis for phenotype-guided representation learning of neuronal gene expression data

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Abstract

A central goal in neurobiology is to relate the expression of genes to the structural and functional properties of neuronal types, collectively called their phenotypes. Single-cell RNA sequencing can measure the expression of thousands of genes in thousands of neurons. How to interpret the data in the context of neuronal phenotypes? We propose a supervised learning approach that factorizes the gene expression data into components corresponding to individual phenotypic characteristics and their interactions. This new method, which we call factorized linear discriminant analysis (FLDA), seeks a linear transformation of gene expressions that varies highly with only one phenotypic factor and minimally with the others. We further leverage our approach with a sparsity-based regularization algorithm, which selects a few genes important to a specific phenotypic feature or feature combination. We applied this approach to a single-cell RNA-Seq dataset of Drosophila T4/T5 neurons, focusing on their dendritic and axonal phenotypes. The analysis confirms results from the previous report but also points to new genes related to the phenotypes and an intriguing hierarchy in the genetic organization of these cells.

1 Introduction

The complexity of neural circuits is a result of many different types of neurons that specifically connect to each other. Each neuronal type has its own phenotypic traits, which together determine the role of the neuronal type in a neural circuit. Typical phenotypic descriptions of neurons include features such as dendritic and axonal laminations, electrophysiological properties, and connectivity [1–3]. However, the genetic programs behind these phenotypic characteristics are still poorly understood.

Recent progress in characterizing neuronal cell types and investigating their gene expression, especially with advances in high-throughput single-cell RNA-Seq [2], provides an opportunity to address this challenge. With massive data generated from single-cell RNA-Seq, we now face a computational problem: how to factorize the high-dimensional data into gene expression modules that are meaningful to neuronal phenotypes? Specifically, given phenotypic descriptions of neuronal types, such as their dendritic stratification and axonal termination, can one project the original data into a low-dimensional space corresponding to these phenotypic features and their interactions, and further extract genes critical to each of these components?

This problem can be approached by well-known methods with limitations. For instance, canonical correlation analysis (CCA) could find linear transformations of the data and the phenotypic features

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As a start, let us consider only two phenotypic features of neurons, dendritic stratification, and axonal lamination. We first asked whether we could factorize, for example, gene expressions and phenotypic features from autoencoders, as they are non-linear architectures. Here we propose a new analysis method named factorized linear discriminant analysis (FLDA). Inspired by multi-way analysis of variance (ANOVA) [9], this method factorizes data into components corresponding to phenotypic features and their interactions, and seeks a linear transformation that varies highly with one specific factor but not with the others. The linear nature of this approach makes it easy to interpret, as the weight coefficients directly inform the relative importance of each gene to each factor. We further introduce a sparse variant of the method, which constrains the number of genes contributing to each linear projection. We illustrate this approach by applying FLDA to a single-cell transcriptome dataset of T4/T5 neurons in Drosophila [10], focusing on two phenotypes: dendritic location and axonal lamination.

2 factorized linear discriminant analysis (FLDA)

Suppose that we are given gene expression data of single neurons which are typically very high-dimensional. These cells are classified into cell types, as a result of clustering in the high-dimensional space and annotations based on prior knowledge or verification outcome [11-13]. We know the phenotypic traits of each neuronal type, therefore each type can also be jointed defined by the phenotypic features. We want to find an interpretable low-dimensional embedding in which certain dimensions represent factors of phenotypic features or their interactions. This requires that variation along one of the axes in the embedding space causes the variation of only one factor. In reality, this is hard to satisfy due to noise in the data, and we relax the constraint by letting data projected along one axis vary largely with one factor while minimally with the others. Indeed, motivated by the linear factor models used in that maximally correlate with each other [4,5]. However, this approach cannot factorize gene expressions according to individual features, making the result hard to interpret. Another approach to this problem is autoencoders and variants [6,8]. It is challenging to understand the relationship between gene expressions and phenotypic features from autoencoders, as they are non-linear architectures.

As a start, let us consider only two phenotypic features of neurons, dendritic stratification, and axonal lamination, both of which can be described with discrete categories, such as different regions or layers in the brain [16,17,1,10]. Suppose that each cell type can be jointly represented by its dendritic location indexed as $i$ and axonal lamination indexed as $j$, with the number of cells within each cell type $n_{ij}$. This representation can be described using a contingency table (Figure 1A,B). Note here that we allow the table to be partially filled.

Let $x_{ijk}(k \in 1, 2, ..., n_{ijk})$ represent the expression values of $g$ genes in each cell ($x_{ijk} \in \mathbb{R}^{g}$). How to find linear projections $y_{ij} = u^T x_{ijk}$ and $z_{ijk} = v^T x_{ijk}$ that are aligned with features $i$ and $j$ respectively (Figure 1C)? We first asked whether we could factorize, for example, $y_{ijk}$, with respect to components depending on features $i$ and $j$. Indeed, motivated by the linear factor models used in

Figure 1: Illustration of our approach. (A,B) In the example, cell types are jointly represented by two phenotypic features. If only some combinations of the two features are observed, one obtains a partial contingency table (B) instead of a complete one (A). (C) We seek linear projections of the data that separate the cell types in a factorized manner corresponding to the two features. Here $u$, $v$, and $w$ are aligned with Feature 1, Feature 2, and the interaction of both features, with the projected coordinates $y$, $z$, and $s$ respectively.
multi-way ANOVA and the idea of partitioning variance, we constructed an objective function as the following, and found $u^*$ that maximizes the objective (see detailed analysis in Appendix A):

$$u^* = \arg \max_{u \in \mathbb{R}^g} \frac{u^T N_A u}{u^T M_e u}$$

When we have a complete table, and there are $a$ levels for the feature $i$ and $b$ levels for the feature $j$, we have

$$N_A = M_A - \lambda_1 M_B - \lambda_2 M_{AB}$$

where $M_A$, $M_B$, and $M_{AB}$ are the covariance matrices explained by the feature $i$, the feature $j$, and the interaction of them. $\lambda_1$ and $\lambda_2$ are hyper-parameters controlling the relative weights of $M_B$ and $M_{AB}$ with respect to $M_A$. $M_e$ is the residual covariance matrix representing noise in gene expressions. Formal definitions of these terms are the following:

$$M_A = \frac{1}{a-1} \sum_{i=1}^{a} \sum_{j=1}^{b} (m_{i.} - m_.)(m_{i.} - m_.)^T$$

$$M_B = \frac{1}{b-1} \sum_{i=1}^{a} \sum_{j=1}^{b} (m_{.j} - m_.)(m_{.j} - m_.)^T$$

$$M_{AB} = \frac{1}{(a-1)(b-1)} \sum_{i=1}^{a} \sum_{j=1}^{b} (m_{ij} - m_{i.} - m_{.j} + m_.)(m_{ij} - m_{i.} - m_{.j} + m_.)^T$$

$$M_e = \frac{1}{N-ab} \sum_{i=1}^{a} \sum_{j=1}^{b} \left[ \frac{1}{n_{ij}} \sum_{k=1}^{n_{ij}} (x_{ijk} - m_{ij})(x_{ijk} - m_{ij})^T \right]$$

where

$$m_. = \frac{1}{ab} \sum_{i=1}^{a} \sum_{j=1}^{b} m_{ij}$$

$$m_{i.} = \frac{1}{b} \sum_{j=1}^{b} m_{ij}$$

$$m_{.j} = \frac{1}{a} \sum_{i=1}^{a} m_{ij}$$

in which

$$m_{ij} = \frac{1}{n_{ij}} \sum_{k=1}^{n_{ij}} x_{ijk}$$

An analogous expression provides the linear projection $v^*$ for the feature $j$, and $w^*$ for the interaction of both features $i$ and $j$. Similar arguments can be applied to the scenario of a partial table to find $u^*$ or $v^*$ as the linear projection for the feature $i$ or $j$ (see Appendix B for mathematical details).

Note that $N_A$ is symmetric and $M_e$ is positive definite. Therefore the optimization problem is a generalized eigenvalue problem [18]. When $M_e$ is invertible, $u^*$ is the eigenvector associated with the
largest eigenvalue of $M_e^{-1} N_A$. In general, if we want to embed $x_{ijk}$ into a $d$-dimensional subspace aligned with the feature $i$ ($d < a$), we can take the eigenvectors with the $d$ largest eigenvalues of $M_e^{-1} N_A$, which we call the top $d$ factorized linear discriminant components (FLDs). Since multi-way ANOVA can handle contingency tables with more than two dimensions, our analysis can be easily generalized to more than two features.

3 Sparsity-based regularization of FLDA

To select genes important to each phenotypic feature, we impose sparseness on the number of genes contributing to each axis. Briefly, we try to solve the following optimization problem:

$$ u^* = \arg \max_{u \in \mathbb{R}^g} \frac{u^T N_A u}{u^T M_e u} \quad \text{subject to} \quad ||u||_0 \leq l $$

(11)

from which the number of non-zero elements of $u^*$ is less or equal to $l$.

This is also known as a sparse generalized eigenvalue problem. To solve it, we used a recently developed statistical approach, the truncated Rayleigh flow method (Rifle). The algorithm of Rifle is composed of two steps [19]:

(1) obtain an initial vector $u_0$ that is close to $u^*$. We used the solution from the non-sparse FLDA as an initial estimate of $u_0$.

(2) iteratively, perform a gradient ascent step on the objective function, and then execute a truncation step that preserves the $l$ entries of $u$ with the largest values and sets the remaining entries to 0.

Pseudo-code for this step is presented below:

```plaintext
procedure RIFLE($N_A, M_e, u_0, l, \eta$)
    $\eta$ is the step size
    $t = 1$
    while not converge
        $\rho_{t-1} \leftarrow \frac{u_{t-1}^T N_A u_{t-1}}{u_{t-1}^T M_e u_{t-1}}$
        $C \leftarrow I + (\frac{\eta}{\rho_{t-1}})(N_A - \eta_{t-1} M_e)$
        $u_t \leftarrow \frac{Cu_{t-1}}{||Cu_{t-1}||_2}$
        Truncate $u_t$ by keeping the top $l$ entries of $u$ with the largest values and setting the rest entries to 0
        $u_t \leftarrow \frac{u_t}{||u_t||_2}$
        $t \leftarrow t + 1$
    end while
    return $u_t$
end procedure
```

4 Related method - linear discriminant analysis (LDA)

We name our method FLDA because its objective function has a similar format as that of linear discriminant analysis (LDA) [20,21]. LDA also models the difference among data organized in pre-determined classes. Formally, LDA solves the following optimization problem:

$$ u^* = \arg \max_{u \in \mathbb{R}^g} \frac{u^T \Sigma_b u}{u^T \Sigma_e u} $$

(12)

where $\Sigma_b$ and $\Sigma_e$ are estimates of the between-class and within-class covariance matrices respectively.

Different from FLDA, the representation of these classes is not explicitly formulated as a contingency table composed of multiple features. The consequence is that, when applied to the example problem in which neuronal types are organized into a two-dimensional contingency table with phenotypic features $i$ and $j$, in general, axes from LDA are not aligned with these two phenotypic features.
However, in the example above, we can perform two separate LDAs for the two features. This allows the axes from each LDA to align with its specific feature. We call this approach “2LDAs”. There are two limitations of this approach: first, it discards information about the component depending on the interaction of the two features which cannot be explained by a linear combination of them; second, it explicitly maximizes the segregation of cells with different feature levels which sometimes is not consistent with a good separation of cell type clusters.

5 Experiments

5.1 Datasets

In order to compare FLDA with LDA and the “2LDAs” approach and show the difference quantitatively, we created synthetic datasets. Four types of cells, each containing 25 examples, were generated from a Cartesian product of two features \( i \) and \( j \), organized in a 2x2 complete contingency table. Expressions of 10 genes were generated for these cells, in which the levels of Genes 1-8 were correlated with either the feature \( i \), the feature \( j \), or the interactions of them, and the levels of the remaining 2 genes were purely driven by noise (Figure 2A). Details of generating the data can be found in Appendix C, and analysis of these synthesized data is reported in Results.

To illustrate FLDA in analyzing single-cell RNA-Seq datasets for real problems of neurobiology, and demonstrate the merit of our approach in selecting a few important genes for each phenotype, we used a dataset of Drosophila T4/T5 neurons [10]. T4 and T5 neurons are very similar in terms of general morphology and physiological properties, but they differ by the location of their dendrites in the medulla and lobula, two distinct brain regions. T4 and T5 neurons each contain four subtypes, with each pair of the four laminating their axons in a specific layer in the lobula plate (Figure 3A). Therefore, we can use two phenotypic features to describe these neurons: the feature \( i \) indicates the dendritic location at the medulla or lobula; the feature \( j \) describes the axonal lamination at one of the four layers (a/b/c/d) (Figure 3B). In this experiment, we focused on the dataset containing expression data of 17492 genes from 3833 cells collected at a defined time during brain development.

5.2 Data preprocessing

The T4/T5 neuron dataset was preprocessed as previously reported [13, 15, 22]. Briefly, transcript counts within each column of the count matrix (genes \( \times \) cells) were normalized to sum to the median number of transcripts per cell, resulting in the normalized counts Transcripts-per-median or \( \text{TPM}_{gc} \) for Gene \( g \) in Cell \( c \). We used the log-transformed expression data \( E_{gc} = \ln (\text{TPM}_{gc} + 1) \) for further analysis. We adopted a common approach in single-cell RNA-Seq studies that is based on fitting a relationship between mean and coefficient of variation [23, 24] to select highly variable genes, and performed FLDA on the expression data with only these genes. As the number of cells was less than the number of genes, we first performed principal component analysis (PCA) and kept
Figure 3: FLDA on the dataset of T4/T5 neurons. (A) T4/T5 neuronal types and their dendritic and axonal phenotypes. (B) T4/T5 neurons can be organized in a complete contingency table. Here $i$ indicates the dendritic location and $j$ indicates the axonal termination. (C) SNR metric of each discriminant axis. (D) Projection of the data into the three-dimensional space consisting of the discriminant axis for the feature $i$ (FLD$_i$) and the first and second discriminant axes for the feature $j$ (FLD$_{j1}$ and FLD$_{j2}$). (E-G) Projection of the data into the two-dimensional space made of FLD$_i$ and FLD$_{j1}$ (E), FLD$_{j1}$ and FLD$_{j2}$ (F), or FLD$_{j2}$ and FLD$_{j3}$ (the third discriminant axis for the feature $j$) (G). Different cell types are indicated by different colors as in (A) and (D).

principal components (PCs) explaining $\sim 99\%$ of the total variance. In this experiment below, we set the hyper-parameters $\lambda$s in Equation (2) to 1.

5.3 Metrics

We included the following metrics to evaluate our method: We specified a metric of signal-to-noise ratio (SNR) to measure how well each discriminant axis separates cell types compared with noise estimated from the variance within cell type clusters; We defined explained variance (EV) for each discriminant axis to measure how much variance of the feature $i$ or $j$ is explained among the total variance explained by that axis; We calculated mutual information (MI) between each discriminant axis and each feature, to quantify how "informative" an axis is to a specific feature; Built on the calculation of MI, we included the modularity score which measures whether each discriminant axis depends on at most one feature [25]. The implementation details of these metrics can be found in Appendix D.

6 Results

To quantitatively compare the difference between FLDA and the alternative models of LDA and "2LDAs", we measured the proposed metrics from analyses of the synthesized datasets (Figure 2A). Given that the synthesized data were organized in a 2x2 contingency table, each LDA of the "2LDAs" approach could find only one dimension for the specific feature $i$ or $j$. Therefore, as a fair comparison, we only included the corresponding dimensions in FLDA (FLD$_i$ and FLD$_j$) and the top two linear discriminant components (LDs) in LDA. The overall SNR values normalized by that of LDA and the
We extracted a list of 20 genes each for the axis of FLDA to plot expression profiles of these genes in the eight neuronal types (Figure 4B,D). For both axonal termination and dendritic phenotype, we applied FLDA to the dataset of Drosophila T4/T5 neurons. The T4/T5 neurons could be organized into two groups, a/b vs c/d, corresponding to the upper or lower lobula plate, and FLD_{1j} divides them into another two, a/d vs b/c, indicating whether their axons laminate at the middle or lateral part of the lobula plate (Figure 3E,F). Unexpectedly, among these three dimensions, FLD_{1j} has a much higher SNR than FLD_{1i} and FLD_{3j}, whose SNR values are similar. This suggests a hierarchical structure in the genetic organization of T4/T5 neurons: they are first separated into either a/b or c/d types, and subsequently divided into each of the eight subtypes. In fact, this exactly matches the sequence of their cell fate determination during development, as revealed in a previous genetic study [26]. Finally, the last discriminant axis of the axonal feature FLD_{3j} separates the group a/c from b/d, suggesting its role in fine-tuning the axonal depth within the upper or lower lobula plate (Figure 3G). For this dataset, we also quantified the proposed metrics whose values are listed in Table 4.

To seek gene signatures for the discriminant components in FLDA, we applied the sparsity-based regularization to constrain the number of genes with non-zero weight coefficients. Here we set the number to 20, a reasonable number of candidate genes to be tested in a follow-up biological study. We extracted a list of 20 genes each for the axis of FLD_{1i} or FLD_{1j}. The relative importance of these genes to each axis is directly informed by their weight values (Figure 3A,C). Side-by-side, we plotted expression profiles of these genes in the eight neuronal types (Figure 3B,D). For both axes, the genes critical in separating cells with different feature levels are differentially expressed in corresponding cell types. On our gene lists, consistent with the original report [10], we found indicator genes for dendritic location such as TjAP-2, dpr2, CG34155, and CG12065, and those for axonal lamination such as klg, bi, pros. In addition, we found genes that were not reported in the previous study. For example, our results suggest that the genes Thor and pHCl-1 are important to the dendritic phenotype, and Lac and Mip are critical to the axonal phenotype. These are promising genetic targets to be tested in biological experiments.

Finally, FLDA allowed us to examine the component that depends on the interaction of both features and identify its gene signature, which provides clues to transcriptional regulation of gene expressions in the T4/T5 neuronal types (Figures 5 and 6). For instance, DIP_{ela} is exclusively expressed in T5c/d neurons, suggesting its expression is turned on only when the dendritic transcriptional program is set for T5 neurons and the axonal program is for c/d types. Oppositely, Neto is selectively enriched in T4a/b neurons, indicating activation of its expression under the condition that the dendritic transcriptional program is set for T4 neurons and the axonal program is for a/b types. These are known as the “AND” and “NOR” logics in genetic programs [27].
Selected genes for the dendritic phenotype

Selected genes for the axonal phenotype

Figure 4: Critical genes extracted from the sparse algorithm. (A) Weight vector of the 20 genes selected for the dendritic phenotype (FLD$_i$). The weight value is indicated in the color bar with color indicating direction (red: positive and green: negative) and saturation indicating magnitude. (B) Expression patterns of the 20 genes from (A) in eight types of T4/T5 neurons. Dot size indicates the percentage of cells in which the gene was expressed, and color represents average scaled expression. (C) Weight vector of the 20 genes selected for the axonal phenotype (FLD$_j$). Legend as in (A). (D) Expression patterns of the 20 genes from (C) in eight types of T4/T5 neurons. Dot size indicates the percentage of cells in which the gene was expressed, and color represents average scaled expression. Legend as in (B).

7 Discussion

Single-cell RNA-Seq has generated a large amount of neuronal transcriptomic data [11–15]. However, neurons display a great variety of phenotypes, making it a challenging task to understand transcriptional programs behind these phenotypic features [28]. Motivated by ANOVA, we developed FLDA to address this challenge. We illustrated FLDA focusing on two phenotypes of T4/T5 neurons in Drosophila, dendritic location and axonal termination. As multi-way ANOVA is applied to multiple factors, our approach can be easily generalized to more than two phenotypic features and applicable to additional characteristics such as electrophysiology and connectivity [3, 2].

FLDA factorizes gene expression data into features and their interactions, and finds a linear projection of the data that varies with only one factor but not the others. This provides a modular representation aligned with the factors [29]. Ridgeway and Mozer (2018) argued that modularity together with explicitness could define disentangled representations. Our approach is linear, which presents an explicit mapping between gene expressions and phenotypic features, therefore our approach can potentially serve as a supervised approach to disentanglement [30–32].

The linear nature of FLDA makes it extremely easy to interpret the representations, as the weight vector directly informs the relative importance of each gene. To allow the selection of a small set of critical genes, we leveraged our approach with sparse regularization. This makes FLDA especially useful to experimentalists who can take the list of genes and test them in follow-up genetic experiments. However, data other than gene expression may require non-linear representation learning. One way to deal with this is to use kernel tricks which are commonly used in machine learning [33]. As a future direction, we also hope to generalize ideas from this work to deep learning models.
8 Appendix

8.1 A. Objective functions

Here we derive the objective functions used in our analysis. Again if $x_{ijk} (k \in 1, 2, \ldots n_{ij})$ represents the expression values of $g$ genes in each cell ($x_{ijk} \in \mathbb{R}^g$), we seek to find a linear projection $y_{ijk} = u^T x_{ijk}$ that is aligned with the feature $i$.

8.1.1 Inspiration from ANOVA

We asked what is the best way to factorize $y_{ijk}$. Inspired by multi-way ANOVA [9], we identified three components: one depending on the feature $i$, another depending on the feature $j$, and the last one depending on the interaction of both features. We therefore followed the procedures of ANOVA to partition sums of squares and factorize $y_{ijk}$ into these three components.

Let us first assume that all cell types defined by $i$ and $j$ contain the same number of cells. With cell types represented by a complete contingency table (Figure 1A), $y_{ijk}$ can be linearly factorized using the model of two crossed factors. Formally, the linear factorization is the following:

$$ y_{ijk} = \mu + \alpha_i + \beta_j + (\alpha \beta)_{ij} + \epsilon_{ijk} \quad (13) $$

where $y_{ijk}$ represents the coordinate of the $k$th cell in the category defined by $i$ and $j$; $\mu$ is the average level of $y$; $\alpha_i$ is the component that depends on the feature $i$, and $\beta_j$ is the component that depends on the feature $j$; $(\alpha \beta)_{ij}$ describes the component that depends on the interaction of both features $i$ and $j$; $\epsilon_{ijk} \sim \mathcal{N}(0, \sigma^2)$ is the residual of this factorization.

Let us say that the features $i$ and $j$ fall into $a$ and $b$ discrete categories respectively. Then without loss of generality, we can require:

$$ \sum_{i=1}^{a} \alpha_i = 0 \quad (14) $$

$$ \sum_{j=1}^{b} \beta_j = 0 \quad (15) $$

$$ \sum_{i=1}^{a} (\alpha \beta)_{ij} = \sum_{j=1}^{b} (\alpha \beta)_{ij} = 0 \quad (16) $$

Corresponding to these, there are three null hypotheses:

$$ H_{01} : \alpha_i = 0 \quad (17) $$

$$ H_{02} : \beta_j = 0 \quad (18) $$

$$ H_{03} : (\alpha \beta)_{ij} = 0 \quad (19) $$

Here we want to reject $H_{01}$ while accepting $H_{02}$ and $H_{03}$ in order that $y_{ijk}$ is aligned with the feature $i$. 

Next, we partition the total sum of squares. If the number of cells within each cell type category is $n$, and the total number of cells is $N$, then we have

$$
\sum_{i=1}^{a} \sum_{j=1}^{b} \sum_{k=1}^{n} (y_{ijk} - \bar{y}(...) )^2 = bn \sum_{i=1}^{a} (\bar{y}_{i..} - \bar{y}(...))^2 + an \sum_{j=1}^{b} (\bar{y}_{.j} - \bar{y}(...))^2 \\
+ n \sum_{i=1}^{a} \sum_{j=1}^{b} (y_{ij} - \bar{y}_{i..} - \bar{y}_{.j} + \bar{y}(...))^2 + \sum_{j=1}^{b} \sum_{k=1}^{n} (y_{ijk} - \bar{y}_{ijk})^2
$$

(20)

where $\bar{y}$ is the average of $y_{ijk}$ over the indices indicated by the dots. Equation (20) can be written as

$$SS_T = SS_A + SS_B + SS_{AB} + SS_e$$

(21)

with each term having degrees of freedom $N - 1, a - 1, b - 1, (a - 1)(b - 1)$, and $N - ab$ respectively. Here $SS_A, SS_B, SS_{AB}$, and $SS_e$ are partitioned sum of squares for the factors $\alpha_i, \beta_j, (\alpha\beta)_{ij}$, and the residual.

ANOVA rejects or accepts a null hypothesis by comparing its mean square (the partitioned sum of squares normalized by the degree of freedom) to that of the residual. This is done by constructing F-statistics for each factor as shown below:

$$F_A = \frac{MS_A}{MS_e} = \frac{SS_A}{a - 1}$$

(22)

$$F_B = \frac{MS_B}{MS_e} = \frac{SS_B}{b - 1}$$

(23)

$$F_{AB} = \frac{MS_{AB}}{MS_e} = \frac{SS_{AB}}{(a - 1)(b - 1)}$$

(24)

Under the null hypotheses, the F-statistics follow the F-distribution. Therefore, a null hypothesis is rejected when we observe the value of a F-statistic above a certain threshold calculated from the F-distribution. Here we want $F_A$ to be large enough so that we can reject $H_{01}$, but $F_B$ and $F_{AB}$ to be small enough for us to accept $H_{02}$ and $H_{03}$. In other words, we want to maximize $F_A$ while minimizing $F_B$ and $F_{AB}$. Therefore, we propose maximizing an objective $L$:

$$L = F_A - \lambda_1 F_B - \lambda_2 F_{AB}$$

(25)

where $\lambda_1$ and $\lambda_2$ are hyper-parameters determining the relative weights of $F_B$ and $F_{AB}$ compared with $F_A$.

### 8.1.2 Objective functions under a complete contingency table

When the numbers of cells within categories defined by $i$ and $j$ ($n_{ij}$) are not all the same, the total sum of squares cannot be partitioned as in Equation (20). However, if we only care about distinctions between cell types instead of individual cells, we can use the mean value of each cell type cluster ($\bar{y}_{ij}$) to estimate the overall average value ($\bar{y}(...)$, and the average value of each category $i$ ($\bar{y}_{i..}$) or $j$ ($\bar{y}_{.j}$). Therefore, Equation (20) can be modified as the following:

$$\sum_{i=1}^{a} \sum_{j=1}^{b} \sum_{k=1}^{n_{ij}} (y_{ijk} - \bar{y}(...) )^2 = bn \sum_{i=1}^{a} (\bar{y}_{i..} - \bar{y}(...))^2 + an \sum_{j=1}^{b} (\bar{y}_{.j} - \bar{y}(...))^2 \\
+ \sum_{j=1}^{b} \sum_{k=1}^{n_{ij}} (\bar{y}_{ij} - \bar{y}_{i..} - \bar{y}_{.j} + \bar{y}(...))^2 + \sum_{j=1}^{b} \sum_{k=1}^{n_{ij}} (y_{ijk} - \bar{y}_{ijk})^2
$$

(26)
where

\[
\bar{y}_{ij} = \frac{\sum_{k=1}^{n_{ij}} y_{ijk}}{n_{ij}} \quad (27)
\]

\[
\bar{y}_{i..} = \frac{\sum_{j=1}^{b} \bar{y}_{ij.}}{b} \quad (28)
\]

\[
\bar{y}_{j..} = \frac{\sum_{i=1}^{a} \bar{y}_{ij.}}{a} \quad (29)
\]

\[
\bar{y}.. = \frac{\sum_{i=1}^{a} \sum_{j=1}^{b} \bar{y}_{ij.}}{ab} \quad (30)
\]

If we describe Equation (26) as:

\[
\tilde{SS}_T = \tilde{SS}_A + \tilde{SS}_B + \tilde{SS}_{AB} + \tilde{SS}_e \quad (31)
\]

then following the same arguments, we want to maximize an objective function in the following format:

\[
L = \frac{\tilde{SS}_A}{a-1} - \lambda_1 \frac{\tilde{SS}_B}{b-1} - \lambda_2 \frac{\tilde{SS}_{AB}}{(a-1)(b-1)} \frac{\tilde{SS}_e}{N-ab} \quad (32)
\]

### 8.1.3 Objective functions under a partial contingency table

When we have a representation of a partial table, we can no longer separate out the component that depends on the interaction of both features. Therefore, we use another model, a linear model of two nested factors, to factorize \( y_{ijk} \), which has the following format:

\[
y_{ijk} = \mu + \alpha_i + \beta_{j(i)} + \epsilon_{ijk} \quad (33)
\]

Note that we now have \( \beta_{j(i)} \) instead of \( \beta_j + \alpha\beta_{ij} \). In this model, we identify a primary factor, for instance, the feature denoted by \( i \) which falls into \( a \) categories, and the other (indexed by \( j \)) becomes a secondary factor, the number of whose levels \( b_i \) depends on the level of the primary factor. We merge the component depending on the interaction of both features into that of the secondary factor as \( \beta_{j(i)} \).

Similarly, we have

\[
\sum_{i=1}^{a} \sum_{j=1}^{b_i} \left[ \frac{1}{n_{ij}} \sum_{k=1}^{n_{ij}} (y_{ijk} - \bar{y}_{ij.} - \bar{y}_{i..} - \bar{y}..)^2 \right] = \sum_{i=1}^{a} \sum_{j=1}^{b_i} \left( \bar{y}_{ij.} - \bar{y}_{i..} - \bar{y}.. \right)^2 + \sum_{i=1}^{a} \sum_{j=1}^{b_i} \frac{1}{n_{ij}} \sum_{k=1}^{n_{ij}} (y_{ijk} - \bar{y}_{ij.})^2 \quad (34)
\]

which can be written as

\[
\tilde{SS}_T = \tilde{SS}_A + \tilde{SS}_B + \tilde{SS}_e \quad (35)
\]

with degrees of freedom \( N-1, a-1, M-a, \) and \( N-M \) for each of the terms, where \( M \) is:

\[
M = \sum_{i=1}^{a} b_i \quad (36)
\]
Therefore, we want to maximize the following objective:

\[
L = \frac{s_{SA}}{a-1} - \lambda \frac{s_{SB}}{M-a}
\]

(37)

\section*{8.2 B. FLDA with a partial contingency table}

Here we provide the mathematical details of FLDA under the representation of a partial table. When we have a partial table, if the feature \( i \) is the primary feature with \( a \) levels, and the feature \( j \) is the secondary feature with \( b_i \) levels, then \( N_A \) in Equation (1) is defined as follows:

\[
N_A = M_A - \lambda M_{B|A}
\]

(38)

where

\[
M_A = \frac{1}{a-1} \sum_{i=1}^{a} \sum_{j=1}^{b_i} (m_{ij} - m_{..}) (m_{ij} - m_{..})^T
\]

(39)

\[
M_{B|A} = \frac{1}{M-a} \sum_{i=1}^{a} \sum_{j=1}^{b_i} (m_{ij} - m_{..}) (m_{ij} - m_{..})^T
\]

(40)

and \( M \) is defined as in Equation (36). Correspondingly, \( M_e \) in Equation (1) is defined as:

\[
M_e = \frac{1}{N-M} \sum_{i=1}^{a} \sum_{j=1}^{b_i} \left[ \frac{1}{n_{ij}} \sum_{k=1}^{n_{ij}} (x_{ijk} - m_{ij}) (x_{ijk} - m_{ij})^T \right]
\]

(41)

and

\[
m_{..} = \frac{1}{M} \sum_{i=1}^{a} \sum_{j=1}^{b_i} m_{ij}
\]

(42)

\[
m_{ij} = \frac{1}{b_i} \sum_{j=1}^{b_i} m_{ij}
\]

(43)

The remaining mathematical arguments are the same as those for the complete table. In this scenario, because we don’t observe all possible combinations of features \( i \) and \( j \), we cannot find the linear projection for the interaction of both features.

\subsection*{8.3 C. Implementation details of data synthesis}

To quantitatively compare FLDA with LDA and the “2LDAs” approach, we synthesized data of four cell types, each of which contained 25 cells. The four cell types were generated from a Cartesian product of two features \( i \) and \( j \), where \( i \in \{0, 1\} \) and \( j \in \{0, 1\} \). Expressions of 10 genes were generated for each cell. The expression value of the \( h \)th gene in the \( k \)th cell of the cell type \( ij \), \( g_{ijk} \), was defined as the following:

\[
g_{ijk}^1 = i + \epsilon_{ijk}
\]

(44)

\[
g_{ijk}^2 = j + \epsilon_{ijk}
\]

(45)
\[ g_{ijk}^3 = i \land j + \epsilon_{ijk} \] (46)

\[ g_{ijk}^4 = i \lor j + \epsilon_{ijk} \] (47)

\[ g_{ijk}^5 = 2i + \epsilon_{ijk} \] (48)

\[ g_{ijk}^6 = 2j + \epsilon_{ijk} \] (49)

\[ g_{ijk}^7 = 2i \land j + \epsilon_{ijk} \] (50)

\[ g_{ijk}^8 = 2i \lor j + \epsilon_{ijk} \] (51)

\[ g_{ijk}^9 = \epsilon_{ijk} \] (52)

\[ g_{ijk}^{10} = 2 + \epsilon_{ijk} \] (53)

where

\[ i \land j = \begin{cases} 1, & \text{if } i = 1, j = 1 \\ 0, & \text{otherwise} \end{cases} \] (54)

and

\[ i \lor j = \begin{cases} 0, & \text{if } i = 0, j = 0 \\ 1, & \text{otherwise} \end{cases} \] (55)

were interactions of the two features. Here \( \epsilon_{ijk} \) was driven by Gaussian noise, namely,

\[ \epsilon_{ijk} \sim \mathcal{N}(0, \sigma^2) \] (56)

We synthesized datasets of 5 different \( \sigma \) values (\( \sigma \in \{0.2, 0.4, 0.6, 0.8, 1.0\} \)). This was repeated 10 times and metrics for each \( \sigma \) value were calculated as the average across the 10 repeats.

8.4 D. Implementation details of the metrics used in the study

We measured the following metrics in our experiments:

8.4.1 Signal-to-Noise Ratio (SNR)

Because we care about the separation of cell types, we define the SNR metric as the ratio of the variance between cell types over the variance of the noise, which is estimated from within-cluster variance. For the entire embedding space, given \( q \) cell types, if the coordinate of each cell is indicated by \( c \), then we define the overall SNR metric as the following:

\[
SNR_{overall} = \frac{\text{tr}(\Sigma_{p=1}^q n_p (\bar{c}_p - \bar{c}_.)(\bar{c}_p - \bar{c}_.)^T))}{\text{tr}(\Sigma_{p=1}^q \Sigma_{k=1}^{n_p} (c_{pk} - \bar{c}_p)(c_{pk} - \bar{c}_p)^T)}
\] (57)

where \( \bar{c}_p \) is the center of each cell type cluster, and \( \bar{c}_. \) is the center of all data points.
Let \( c \) denote the embedded coordinate along a specific dimension. The SNR metric for that axis is therefore:

\[
SNR = \frac{\sum_{p=1}^{n_p} n_p (\bar{c}_p - \bar{c}_{..})^2}{\sum_{p=1}^{n_p} \sum_{k=1}^{n_p} (c_{pk} - \bar{c}_p)^2}
\]  

(58)

8.4.2 Explained Variance (EV)

We want to know whether the variation of a specific dimension is strongly explained by that of a specific feature. Therefore, we measure, for each axis, how much of the total explained variance is explained by the variance of the feature \( i \) or \( j \). Formally, given the embedded coordinate \( y_{ijk} \), we calculate the EV as the following:

\[
EV_i = \frac{\sum_{a=1}^{a} \sum_{b=1}^{b} n_{ij} (\bar{y}_{i..} - \bar{y}_{...})^2}{\sum_{a=1}^{a} \sum_{b=1}^{b} \sum_{k=1}^{n_{ij}} (y_{ijk} - \bar{y}_{...})^2}
\]  

(59)

\[
EV_j = \frac{\sum_{a=1}^{a} \sum_{b=1}^{b} n_{ij} (\bar{y}_{j..} - \bar{y}_{...})^2}{\sum_{a=1}^{a} \sum_{b=1}^{b} \sum_{k=1}^{n_{ij}} (y_{ijk} - \bar{y}_{...})^2}
\]  

(60)

where \( \bar{y} \) is the average of \( y_{ijk} \) over the indices indicated by the dots.

8.4.3 Mutual Information (MI)

The MI between a discriminant axis \( u \) and a feature quantifies how much information of the feature is obtained by observing data projected along that axis. It is calculated as the MI between data representations along the axis \( y = u^T X \) and feature labels of the data \( f \), where \( X \) is the original gene expression matrix:

\[
I(y, f) = H(y) + H(f) - H(y, f) = -\sum_{y \in Y} p(y) \log_2 p(y) - \sum_{f \in F} p(f) \log_2 p(f) - \sum_{y \in Y} \sum_{f \in F} p(y, f) \log_2 p(y, f)
\]  

(61)

Here \( H \) indicates entropy. To calculate \( H(y) \) and \( H(y, f) \), we discretize \( y \) into 10 bins.

8.4.4 Modularity

Ridgeway and Mozer (2018) argued that in a modular representation, each axis should depend on at most a single feature. Following the arguments in their paper, the modularity score is computed as follows: we first calculate the MI between each feature and each axis \( (m_{if} \text{ denotes the MI between one axis } i \text{ and one feature } f) \). If an axis is perfectly modular, it will have high mutual information for only one feature and zeros for the others, we therefore compute a template \( t_{if} \) as the following:

\[
t_{if} = \begin{cases} 
\theta_i, & \text{if } f = \arg \max_g m_{ig} \\
0, & \text{otherwise}
\end{cases}
\]  

(62)

where \( \theta_i = \max_g m_{ig} \). We then calculate the deviation from the template as:

\[
\delta_i = \frac{\sum_j (m_{if} - t_{if})^2}{\theta_i^2 (N - 1)}
\]  

(63)

where \( N \) is the number of features. The modularity score for the axis \( i \) is \( 1 - \delta_i \). The mean of \( 1 - \delta_i \) over \( i \) is defined as the overall modularity score.
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9 Additional Information

| Sigma | 0.2   | 0.4   | 0.6   | 0.8   | 1      |
|-------|-------|-------|-------|-------|--------|
| EV_j of FLD_i | 0.94915015 | 0.89798334 | 0.83949665 | 0.75799516 | 0.66325434 |
| EV_j of FLD_j | 0.03684044 | 0.04535285 | 0.02913673 | 0.01936836 | 0.01216407 |
| EV_j of FLD_i | 0.03665295 | 0.04499388 | 0.02981225 | 0.02231451 | 0.01190785 |
| EV_j of FLD_j | 0.94874927 | 0.89794858 | 0.82187076 | 0.74982447 | 0.65690059 |
| I(i, FLD_i) | 0.99360769 | 0.99308033 | 0.95913323 | 0.86075507 | 0.71272067 |
| I(i, FLD_j) | 0.49493103 | 0.28223236 | 0.15676406 | 0.090115 | 0.07770875 |
| I(j, FLD_i) | 0.4643901 | 0.28223236 | 0.15676406 | 0.090115 | 0.07770875 |
| I(j, FLD_j) | 0.99298066 | 0.98740331 | 0.94179684 | 0.85538379 | 0.73796543 |
Table 2: MI and EV metrics measured from LDA on the synthesized data

| Sigma | EV_i of LD1   | EV_i of LD2   | EV_j of LD1   | EV_j of LD2   |
|-------|--------------|--------------|--------------|--------------|
| 0.2   | 0.48776682   | 0.4769204    | 0.51329241   | 0.6015657    |
| 0.4   | 0.45116523   | 0.44081294   | 0.51243319   | 0.46221964   |
| 0.6   | 0.52330355   | 0.36777595   | 0.36741921   | 0.47571376   |
| 0.8   | 0.42727731   | 0.44296763   | 0.3870769    | 0.37019437   |
| 1     | 0.36337134   | 0.32343567   | 0.33454171   | 0.32053017   |

Table 3: MI and EV metrics measured from “2LDAs” on the synthesized data

| Sigma | EV_i of LD_i | EV_i of LD_j | EV_j of LD_i | EV_j of LD_j |
|-------|--------------|--------------|--------------|--------------|
| 0.2   | 0.97822626   | 0.00520168   | 0.97847676   | 0.9780679    |
| 0.4   | 0.92099596   | 0.0094905    | 0.92212988   | 0.99508033   |
| 0.6   | 0.85186621   | 0.00939646   | 0.83392262   | 0.96055224   |
| 0.8   | 0.76552256   | 0.00952869   | 0.75588205   | 0.87531047   |
| 1     | 0.66916737   | 0.00696314   | 0.66316636   | 0.73704612   |

Table 4: Metrics measured from FLDA, LDA and “2LDAs” on the dataset of T4/T5 neurons

| Model | Axis         | SNR     | EV_i   | EV_j   | I(i, u) | I(j, u) | Modularity |
|-------|--------------|---------|--------|--------|---------|---------|------------|
| FLDA  | FLD_i        | 47.4564008 | 0.94162107 | 2.09E-05 | 0.99569955 | 0.07323188 | 0.99459067 |
|       | FLD_j_1      | 17.7680626 | 0.004905 | 0.97893759 | 0.00952869 | 0.99800712 | 0.99972387 |
|       | FLD_j_2      | 13.6348569 | 0.00949837 | 0.9284645 | 0.00946416 | 0.99806253 | 0.99972387 |
|       | FLD_j_3      | 4.56306359 | 8.00E-07 | 0.81863597 | 0.005781 | 0.94929582 | 0.99996294 |
| LDA   | LD1          | 48.1795978 | 0.94162107 | 2.09E-05 | 0.99569955 | 0.07323188 | 0.99459067 |
|       | LD2          | 18.2363587 | 0.004905 | 0.97893759 | 0.00952869 | 0.99800712 | 0.99972387 |
|       | LD3          | 14.016217 | 0.00949837 | 0.9284645 | 0.00946416 | 0.99806253 | 0.99972387 |
|       | LD4          | 4.64133888 | 8.00E-07 | 0.81863597 | 0.005781 | 0.94929582 | 0.99996294 |
| 2LDAs | LD_i         | 47.3372616 | 0.94162107 | 2.09E-05 | 0.99569955 | 0.07323188 | 0.99459067 |
|       | LD_j_1       | 17.4834949 | 0.004905 | 0.97893759 | 0.00952869 | 0.99800712 | 0.99972387 |
|       | LD_j_2       | 13.7035177 | 0.00949837 | 0.9284645 | 0.00946416 | 0.99806253 | 0.99972387 |
|       | LD_j_3       | 4.57167846 | 8.00E-07 | 0.81863597 | 0.005781 | 0.94929582 | 0.99996294 |

Figure 5: Additional plots for FLDA on the dataset of T4/T5 neurons. (A, B) Projection of the original gene expression data into the two-dimensional space made of the first and second (FLD_{ij}^1 and FLD_{ij}^2) (A) or the second and third (FLD_{ij}^2 and FLD_{ij}^3) (B) discriminant axes for the component that depends on the combination of both features i and j. Different cell types are indicated in different colors as in (B).
Figure 6: Additional plots for critical genes extracted from the sparse algorithm. (A) Weight vector of the 20 genes selected for the interaction of both dendritic and axonal features ($\text{FLD}_{ij}$). The weight value is indicated in the color bar with color indicating direction (red: positive and green: negative) and saturation indicating magnitude. (B) Expression patterns of the 20 genes from (A) in eight types of T4/T5 neurons. Dot size indicates the percentage of cells in which the gene was expressed, and color represents average scaled expression.