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Integrating Data Clustering and Visualization for the Analysis of 3D Gene Expression Data

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Abstract—The recent development of methods for extracting precise measurements of spatial gene expression patterns from three-dimensional (3D) image data opens the way for new analyses of the complex gene regulatory networks controlling animal development. We present an integrated visualization and analysis framework that supports user-guided data clustering to aid exploration of these new complex datasets. The interplay of data visualization and clustering-based data classification leads to improved visualization and enables a more detailed analysis than previously possible. We discuss (i) integration of data clustering and visualization into one framework; (ii) application of data clustering to 3D gene expression data; (iii) evaluation of the number of clusters k in the context of 3D gene expression clustering; and (iv) improvement of overall analysis quality via dedicated post-processing of clustering results based on visualization. We discuss the use of this framework to objectively define spatial pattern boundaries and temporal profiles of genes and to analyze how mRNA patterns are controlled by their regulatory transcription factors.

Index Terms—bioinformatics visualization, multimodal visualization, integrating Infovis/Scivis, visual data mining, three-dimensional gene expression, data clustering, cluster visualization, gene expression pattern, temporal expression variation, gene regulation, spatial expression pattern

I. INTRODUCTION

UNDERSTANDING the control of embryo development is a fundamental question in biology. A cell’s unique fate is determined by specific combinations of developmental regulatory factors that form part of complex genetic regulatory networks ultimately coordinating the expression of all genes. As a result, the developing embryo exhibits an extraordinarily complex set of spatial and temporal gene expression patterns. The basic structure of the genetic regulatory network is defined by the genome sequence. However, we currently cannot adequately decipher this information or correctly predict how patterns of gene expression evolve.

The Berkeley Drosophila Transcription Network Project (BD-NTP) is generating multiple complementary datasets to address these challenges using the early Drosophila developmental regulatory network as a model. These data sets include in vitro- and in vivo DNA binding data for key transcriptional regulators and, of particular relevance to this work, 3D gene expression data that describes the spatial output of the network at cellular resolution for multiple time points [1], [2].

A large variety of questions can be addressed using these new 3D gene expression datasets [2], [3]. For some analyses, such as logic-based network models, it is helpful to have an objective description of the pattern of a gene at a particular time point, i.e., to define which cells do or do not express a gene. Analysis of the temporal dynamics of gene expression, i.e., how patterns change over time, is essential for gaining a deeper understanding of complex network inter-relationships. Knowledge of the input and output of a network, i.e., the response of the gene expression network at time $t = t_{i+1}$ to the input of the expression levels of regulators at time $t = t_i$, is paramount to identifying regulatory interactions.

To address these and other challenges, we need a flexible visualization tool that allows for interactive exploration of the data. Since drosophila melanogaster has been used as a model for genetic research for decades, there exists a large accumulated body of knowledge about it. A tool designed for the analysis of 3D gene expression data must therefore allow researchers to incorporate this existing knowledge in the analysis, for example by providing ways to modify analysis results, and thus the visualization, accordingly. The tool must also capture the biological context of the embryo and allow different subsets of the data (cells or gene expression patterns) to be examined.

While visualization is a powerful approach to gain deeper insights into such complex data sets, it is limited in this case because the intricate and often subtle nature of 3D gene expression data makes visual detection of all existing features very difficult. For example, a typical feature of interest would be various groups of cells behaving similarly with respect to the expression of several genes. A human’s eye and mind, however, cannot readily...
compute relative concentrations of gene products. Data clustering has already proven to be very powerful at revealing details from conceptually simpler forms of expression data, such as that from microarray experiments, that are not easily detected visually in raw data. Appropriately defining clustering parameters, such as the number of clusters, as well as validation and interpretation of clustering results, is a non-trivial endeavor. To overcome these difficulties in both visual analysis and data clustering, we have adopted data clustering for 3D gene expression analysis by integrating it into PointCloudXplore (PCX). PCX is a visualization tool that features linked physical and information visualization views specifically developed for visualization of 3D gene expression data [4], [5].

Sections II and III present essential biological background necessary for understanding this work. After describing our integrated system in detail in Section IV, and evaluating the question of how to choose the number of clusters $k$ in Section V, we discuss, using a few example cases, how our integrated data clustering and visualization tool can be used in practice to address three relevant questions: (i) How can we usefully divide cells into distinct components of a gene’s expression pattern? (Section VI); (ii) What is the temporal variation of a gene expression pattern? (Section VII); and (iii) What components of a gene’s expression pattern are related to the expression patterns of the regulatory factors that control it? (Section VIII). In Section IX we present our conclusions and describe future plans.

II. BACKGROUND

All cells of living organisms contain DNA, which encodes the genetic information of the organism. Genes are functional subsequences of the DNA. Most genes code for the amino acid sequences of proteins and additional cis-regulatory elements that help to determine in which cells the gene’s product will be expressed. An important class of protein coding genes are developmental regulatory transcription factors that function by binding to cis-regulatory sequences in many genes and direct their patterns of gene expression. Complex genetic regulatory networks are built up where cascades of differently expressed transcription factors ultimately regulate all genes’ expression. These networks guide the development of all living organisms. The characteristic spatial and temporal patterns of regulatory transcription factors define the body plan of the developing animal (see Figure 1).

To provide a quantitative description of these patterns of gene expression in the early *Drosophila* embryo, the BDTNP has developed a data processing pipeline for extracting precise measurements of spatial gene expression patterns in 3D space. *Drosophila* embryos are first fluorescently stained and imaged using two-photon microscopy (see Figure 1, left). Each image is segmented to extract information, such as nuclear positions and volumes, as well as expression values in the neighborhood of each nucleus for the chosen genes [2]. The resulting *PointCloud* file contains information about either protein or mRNA expression of the genes. It is not practical to obtain the expression of more than a few genes in a single embryo, due to the limited number of different distinguishable fluorophores as well as the difficulty in adding multiple labels to embryos.

To allow relationships between multiple transcription factors and their target genes to be compared in a common coordinate framework, PointClouds are registered into a Virtual Embryo using both morphology and a common reference gene to determine cell correspondences [6], [7]. Because the spatial patterns of the genes change rapidly during stage 5, we stage the embryos based on invagination of cell membranes and group the PointClouds into six temporal cohorts [2].

For temporal comparisons, different cohorts are matched using the cellular flow fields that predict the positions of individual cells at each time point [3], [7]. This method enables us to follow gene expression levels within a particular cell over time using only data measured in fixed embryos. Hence, each cell in the Virtual Embryo contains gene expression levels for each of the six time steps. This cellular-level link between embryos of different ages makes it possible to study the development of gene expression patterns over time, as well as to use an mRNA expression pattern as an approximate substitute for a later protein expression pattern, when suitable protein data is not yet available [7].

Figure 1 (middle image) shows a 3D representation of a Virtual Embryo with an average expression pattern using the BDTNP’s visualization tool PCX. To provide an overview of all cells while preserving the relative spatial expression patterns, PCX offers a second physical view in which a cylindrical projection maps all cells onto a rectangular plane (see Figure 1, right). For simplicity, here we use this Unrolled View as our standard physical embryo view. A 3D view can equally be used to view embryos and developmental stages with more complex morphologies than in the early *Drosophila* embryo.

III. RELATED WORK

Data classification is the systematic grouping of data into categories according to some criteria. Data clustering is a class of techniques for unsupervised classification of data samples (here cells) into groups (clusters) of similar behavior. Data clustering provides means for the automatic discovery of data subclasses [8].

In some experimental contexts, such as expression microarrays, gene expression data is often represented as a data matrix, where each gene corresponds to a row and each data sample (cell, microarray, experiment, or condition) to a column. Each matrix entry describes the expression level of a gene in a specific experiment. In these applications, data clustering has proven very useful to classify expression data matrices and thereby identify characteristic sub-structures of each matrix.

Gene expression data clustering can roughly be subdivided into three applications: (i) clustering of genes to identify genes of similar function [9]; (ii) clustering of data samples to identify, for example, different tumor cell types [10]; and (iii) biclustering, i.e., clustering of genes and data samples at the same time to find subgroups of genes and data samples where highly similar activities are seen for the genes in the subset of data samples [11].
Clustering results are most commonly visualized using scatter-plots, plots of statistics, and color table views with columns and/or rows sorted with respect to the clustering. The broad applicability of clustering to gene expression has led to the development of several commercial and publicly available tools for clustering and visualization of gene expression data [12], [13], [14], [15], [16], [17]. However, these tools are limited to what essentially are one-dimensional analyses of gene expression in homogenized populations of cells. They do not take account of spatial position nor the complex relationship of expression across neighboring cells, and are consequently not suitable for interactive visualization and exploration of 3D gene expression data produced by the BDTNP.

Validation of clustering results and evaluation of an “optimal” number of clusters $k$ is an important problem in clustering of gene expression data. A survey of computational cluster validation techniques for gene expression data analysis is provided by Handl et al. [18]. Cluster evaluation functions are commonly subdivided in external and internal measures. External evaluation measures compare the result of a single clustering with a known set of class labels (the “gold standard” or “ground truth”). For our data a “gold standard” is not known, and consequently, we cannot consider external cluster evaluation functions. Internal evaluation measures do not rely on a “gold standard” but evaluate the clustering based on clustering results and the classified dataset.

The most common cluster evaluation measures consider the compactness, connectedness, and/or separation of a clustering. Such general measures, however, do not employ any specific characteristic of gene expression data. The Figure Of Merit (FOM) is an internal measure for gene clustering proposed by Yeung et al. [19] and extended by Datta and Datta [20] that employs explicitly the redundancies and correlations often present in gene expression data. In our application, the level of redundant information is generally low. As a result, FOM and analysis techniques such as the overabundance analysis proposed by Bendor et al. [21] are often not appropriate for our applications but may be interesting when the cells of the embryo are to be classified based on the information of a very large number of genes. To the best of our knowledge, none of these existing cluster quality measures directly employ the fact that genes are expressed in characteristic spatial patterns.

Internal cluster quality measures have been used to estimate the number of clusters $k$ in a dataset. Estimation of an “optimal” $k$ is usually done by computing a series of clustering results for an increasing number of clusters $k$. If a clustering algorithm and internal evaluation measure are adequate for the data to be classified, an “optimal” value of $k$ can often be identified as a “knee” (or elbow) of the resulting performance curve. Tibshirani et al. [22] introduced the gap statistic, a statistical procedure that formalizes this heuristic. Milligan and Cooper performed a Monte Carlo evaluation of 30 procedures for determining the number of clusters in a data set [23]. Existing cluster evaluation measures are designed to find “one perfect” $k$. As we show later in Section IV-D and V, when clustering cells in a 3D gene expression data set, we typically find a series of valid values for $k$, rather than the one “perfect” $k$.

To enable visualization of high-dimensional 3D gene expression data, PCX uses the established concept of linked multiple views [24]. Henze [25] proposed a system based on multiple views (termed portraits) for exploration of time-varying computational fluid dynamics data sets; advanced queries can be performed by selecting data subsets in these portraits. In the WEAVE system, a combination of Physical Views and Information Visualization Views (or abstract views as we refer to them in this paper) is used for exploration of cardiac simulation and measurement data [26]. Doleisch et al. [27] formalized the concept of using abstract views to define data queries.

It is often useful to interactively select data samples from a visual data representation, an operation generally referred to as brushing. A brush is an object that defines one specific selection of data samples. In PCX, brushing is used in a variety of views to select groups of cells with respect to associated quantities. To make this concept more intuitive to the biologist users, brushes are referred to in PCX as cell selectors and the operation of brushing as cell selection. Furthermore, cell selectors defined in one view are also highlighted in all other views, greatly aiding identification of further data properties. This process is termed linking.

PCX was also inspired by the work of Kosara et al. [28], Piringer et al. [29], and Fua et al. [30], who described several important extensions to standard scatter-plots and parallel-coordinates which are incorporated as abstract views in PCX and also used here [4], [5].

IV. DATA CLUSTERING AND VISUALIZATION PIPELINE

The PCX processing pipeline consists of two main interconnected components: visualization and data clustering. Visualization provides the ability to explore the data, to determine appropriate parameters for the clustering, to validate and analyze clustering results, and to modify clustering results using several dedicated cluster post-processing techniques (see Figure 2). Clustering provides ways for automatic identification of data features by classifying cells into groups (the clusters) based on similarity of their gene expression profiles. By highlighting clusters in the visualization, analysis and comparison of specific data features becomes possible, leading to a much more focused analysis of the data. Figure 2’s flowchart shows the basic structure of the data clustering and visualization pipeline as well as the connections between its main components, which are described in detail in the following subsections.

A. Visualizing 3D Gene Expression Data

As described above, PCX is a visualization tool specifically developed for the analysis of 3D gene expression data [4],...
[5]. Physical and abstract views are integrated into a common framework using the established concept of brushing and linking. In physical views, color and height are used for visualizing spatial gene expression patterns (see Section II). In abstract views, physical cell positions are ignored and expression levels for multiple genes are plotted with respect to each other using scatter plots or parallel coordinates.

Selecting cells of interest can be executed in any view in PCX. Depending on the view, different data properties are employed to select cells. User-defined cell selections are then stored and managed in a central cell selector management system. Since all views have access to the same set of cell selectors, features of interest can be defined in any one view and then further analyzed in any other view (as will be shown later, in Figures 6, 13 and 14). The most common way to visualize cell selectors in PCX is to use a consistent color mapping. Depending on the current view, additional functions for highlighting cell selectors are available, such as cell selector-bands in 2D parallel coordinates (see, e.g., Figure 6 and 14).

B. Cluster Statistics

Analysis of statistical properties of clusters is essential for both the validation and analysis of clustering results. Cluster properties provided by PCX include the percentage of cells selected by a cluster, as well as the minimum, maximum, average, and standard deviation values for gene expression levels in a cluster. To compare these statistical properties for one gene in multiple clusters or multiple genes in one cluster, PCX provides box-plots and multi-dimensional color/transparency histogram plots. In histogram plots, we use both color and transparency to visualize the number of cells within a cluster that express the gene over a range of expression levels. Average curve plots (with optional error bars showing standard deviation values) aid in simultaneous analysis of multiple clusters in multiple genes. A simple example shown in Figure 3 illustrates the use of cluster statistics.

C. Data Selection

While it is possible to execute the clustering algorithms on an entire dataset, a more typical use pattern is to focus clustering on a data subset relevant to a specific line of scientific inquiry. The researcher, therefore, needs to define which parts of the data are relevant to address the current problem. In this section, we describe the different steps involved in the data selection process, as well as the effects of data selection on the cluster analysis, and describe how spatial information can be incorporated in the data analysis process. In the following two subsections, we will then describe clustering of 3D gene expression data and post-processing of clustering results.

3D gene expression data can be described as a matrix where each row represents one cell and each column one cell attribute, i.e., the expression of a gene at a specific time point or the $x$, $y$, or $z$ position of the cells in physical space. In order to define which parts of the expression data matrix are relevant, one needs to define: i) which rows (cells), and ii) which columns (gene+time point, $x$, $y$, or $z$) are of interest. Note that this form of data is quite different from that of gene expression microarray matrices, where each row represents a gene and each column represents expression under a different experimental condition, and spatial relationships are meaningless.

Cells of interest can be defined in PCX by using any cell selector or by using the results of a previous data clustering. Defining cells of interest focuses the analysis on a specific part of the data and also reduces the impact of surrounding noise on the analysis. By explicitly allowing data selection based on cell location, PCX overcomes one of the limitations of clustering methods designed for expression microarray data. By using an earlier clustering to define cell subsets of interest, one can first use PCX to group cells into a smaller number of clusters representing the predominant data features and then refine these clusters again using additional rounds of data clustering. In PCX, data clustering, as well as validation of clustering results, can in this way be performed in a step-by-step iterative process.

Defining which cell attributes are of interest is mandatory prior to clustering in PCX since these attributes define the actual biological context of the cells. To account for the complexities of 3D expression data, a variety of unique cell attribute data selection strategies is supported within PCX. First, genes of interest are generally identified based on visualization of the 3D gene expression data as well as based on input from other biological experiments, such as in vivo protein-DNA binding affinity data. Second, to account for spatial location in the clustering analysis, it is possible to directly use cell coordinates as input to the PCX clustering process. Adding this data enforces creation of spatially separated clusters along the AP ($x$) and/or the DV ($y$ and $z$) body axes. Individual weights can be defined for $x$, $y$, and $z$. These weights are then considered in the distance metric (see Section IV-D). However, in most cases the preferred way to incorporate spatial information in the analysis process is by splitting the newly computed clusters into their main independent spatial components. The main advantage of such a cluster post-processing technique over including cell coordinates...
in the clustering process is that cells with similar expression behavior in different parts of the embryo can be identified, and possible clustering artifacts due to the mixing of expression and spatial information can be prevented.

We observed an improved quality of analysis results by adding spatial information to the clustering process when classifying the static pattern of a single gene that has a wide spatial distribution. In the example shown in Figure 4, we classified the pattern of the gene giant (gt) using k-means clustering with and without using x cell positions in the clustering process. In the first case, three clusters were created, each selecting cells expressing gt at different levels, i.e., low, medium, and high expression (Figure 4a). By considering x cell positions, we create separate clusters for the different major spatial components of the gt expression pattern (see Figure 4b). In this case, each cluster includes only cells that express gt at specific levels, while the minimum and maximum expression level selected by each cluster also depends on its physical location. In this case, higher threshold levels were created in the anterior, and lower thresholds in the posterior region of the embryo (see Figure 4c). Creation of region-dependent threshold levels is often desirable when analyzing the static pattern of a single gene since each domain of a pattern may be regulated differently and, therefore, different thresholds may be appropriate. For gap genes with spatially distant, independent expression domains, such as gt, this simple strategy works well, whereas for patterns with shorter interdomain distances, such as eve, this strategy fails.

D. Clustering 3D Gene Expression Patterns

To implement clustering operations in PCX, we use portions of the open source clustering library “Cluster 3.0” [31]. We have integrated data clustering directly into PCX and created a dedicated GUI that provides access to data clustering and allows management of clustering results. Clustering algorithms currently available in PCX include the most commonly used methods for microarray gene expression data analysis, such as k-means, k-median, and k-medoid clustering, as well as several hierarchical clustering algorithms, and self-organizing maps (SOMs) [31], [9], [10], [32]. All these clustering algorithms require an appropriate distance function in order to define similarity between cells. In PCX we included the most common metrics for defining distances in gene expression space: Euclidean distance, city-block-distance, and several derivatives of the Pearson correlation [31].

Some clustering algorithms require additional parameters, such as the number of clusters k, to be specified by the user. In the context of 3D gene expression data there exists in general not a single “perfect” value for k, but we rather find a number of valid values, each representing a different level of detail. This behavior is due to the fact that quantitatively different expression levels of a gene may lead to multiple different outputs of the underlying genetic regulatory network. It is therefore valid to subdivide elongated structures formed in gene expression space into several sub-clusters.

For example, consider early-stage giant (gt) and Krüppel (Kr), which are expressed in spatially non-overlapping patterns, leading to formation of an L-shaped scatter plot (see Figure 5). Even though one could interpret this structure as one cluster – possibly indicating a NOT relationship between gt and Kr – it is also valid to subdivide this structure into, e.g., eight clusters, resulting in one cluster representing background expression, a three-level description of the pattern of Kr, and a four-level description of the gt pattern.

The choice of k depends on the level of detail required by the user. Therefore, PCX uses an interactive process to define k based on visualization. The spatial structure formed by the cells selected by clusters, cluster statistics, and standard data visualizations provides a way to decide if the number of clusters should be increased or reduced. Depending on the characteristic spatial patterns of genes, the cells included in a cluster often define some coherent spatial pattern. Thus, the presence of clusters that show high spatial scattering may be an indication that the chosen k was too large. To assist in this evaluation process, we have developed a dedicated cluster quality measure indicating the physical scattering of clustering results along with a function for suggesting a good initial k. These measures will be described in more detail in Section V. In Section VI, an example is provided where the pattern of eve is classified using different values of k.

Like a manually created cell selector, an automatically created cluster defines a subset of cells in the embryo and can, therefore, be stored and visualized in the same way as cell selectors. Thus, clustering can be used for highlighting data features in physical
or abstract views, enabling a much more focused analysis. In the visualization, PCX allows colors to be assigned to clusters either randomly, manually, or according to the average or ranked average expression of a selected gene in each cluster. Using physical views, the spatial pattern defined by a cluster can be analyzed, and abstract views allow for identification of cluster characteristics in gene expression space.

E. Cluster Post-processing

Cluster post-processing is essential to allow users to modify clustering results with respect to validation results or prior knowledge. There are four ways to post-process clusters in PCX. Manual correction and cluster filtering are two ways to correct small groups of misclassified cells. Cluster merging and splitting provide means to derive coarser or finer representations based on spatial information from the initial clustering.

Manual correction of clustering results can be performed in any physical view. By drawing on the embryo surface, one can interactively add and erase cells from the selection defined by a cluster. In contrast, filtering provides an automatic way to correct misclassified cells. Because genes are expressed in coherent spatial patterns, outliers in physical space tend to be also outliers in gene expression space. Therefore, we have developed a cluster filtering method that identifies and reassigns misclassified cells to the spatially neighboring cluster that is closest in expression space. First, all spatially independent components of a cluster that consist of less than $M$ cells are identified. To rule out false filtering, a minimum distance in physical space as well as a maximum error in expression space can be defined. In the example shown in Figure 6, it would be possible to exclude the cells shown in green from the filtering process either by increasing the minimum spatial distance or by reducing the maximum allowed error in expression space.

Merging clusters allows coarser representations to be created from an initial finer clustering. Such coarser descriptions often provide a clearer visualization that focuses on the main question being addressed (see, e.g., Section VI). Splitting clusters, on the other hand, provides means to derive finer representation from clusters based on spatial information. A cluster often consists of several spatially independent components (for example Figure 12), which may need to be treated differently in subsequent analysis. In general, however, one major component of a cluster may be defined by a number of small spatially independent components. PCX uses a modified single linkage clustering approach to split up such a cluster into a selected (often smaller) number of components.

The splitting algorithm works as follows. A cluster is first split into all its spatially independent components. The smallest components are subsequently merged with the spatially closest component. This approach is computationally more efficient and less sensitive to outliers than a classical single linkage clustering and also guarantees that the independent spatial components of a cluster are preserved while small, scattered components can still grow to define major cluster components. An example for cluster splitting will be described later in Section VI.

V. Identifying Good Values for $k$, the Number of Clusters

Many clustering algorithms, such as k-means, require the user to specify as an input parameter the target number of clusters, $k$. The quality of clustering results often depends on a proper choice of $k$. Unless users have a priori knowledge concerning the number of clusters present in the data, it is helpful for the user that the software offers a reasonable, initial value for $k$. Different approaches for finding an “optimal” $k$ have been proposed. Among them, those based on internal cluster measures appear to be more appropriate for our application [18], [23]. Our objective here, as described below, is to provide the user assistance in interactively searching for a good $k$ as opposed to trying to automatically compute the optimal value of $k$.

Even though internal cluster quality measures (see Section III) may be useful here, we are not aware of any such measure that takes the specific characteristics of 3D gene expression data into account. Since genes are often expressed in compact spatial patterns, we expect the derived clusters to be spatially compact. The presence of computed clusters with high spatial scattering typically suggest that the value of $k$ was too large. Because we do not use information about physical cell position in the clustering process, spatial compactness is a criterion available as an independent measure for clustering quality. As we will discuss below, spatial cluster scattering can also serve as a measure to indicate a series of adequate values of $k$. Combining spatial cluster scattering and the clustering error in expression space yields a method to identify a good initial value for $k$ that accurately reflects the structures present in the data, but with relatively low spatial scattering.

We propose to use

$$\varepsilon_{scatter}(k) = \frac{\sum_{i=1}^{k} R_s(i)}{\sum_{i=1}^{k} R_m(i)}$$

as an objective measure for the relative spatial scattering of a clustering result. $R_s(i)$ (with $s > 0$) is the number of spatially independent components of cluster $i$ consisting of at most $s$ cells. $R_m(i)$ thus defines all single cell regions in cluster $i$. $R_c(i)$ is the total number of spatially independent regions in cluster $i$. $\varepsilon_{scatter} \in [0,1]$ is independent of the clustering algorithm, usually has discontinuities, and shows a larger variation for smaller values.
that also accounts for larger regions as potential scatter, such as

One approach might be a less sensitive weighted cascade measure

indicate a series of appropriate values of

for

of

k

of

k

and a detailed description as shown in a) and a detailed description as shown in b).

Fig. 8. The patterns of

k

Fig. 8. The patterns of

gt

and

Kr,

with

w = 5

and

m = 36.

The suggested

k

is eight as shown in Figure 5.

Fig. 7. Cluster evaluation functions

ε_{\text{exp}}

(red) and

ε_{\text{scatter}}

(blue) for the clustering of

gt

and

Kr,

with

w = 5

and

m = 36. The suggested

k

is eight as shown in Figure 5.

of

k

than for large values of

k.

Local minima of

ε_{\text{scatter}}

indicate values of

k

for which clusters are relatively compact, and thus indicate a series of appropriate values of

k.

In the context of 3D gene expression data, clustering errors introduced by single cells isolated in physical space are quite common and our choice for

ε_{\text{scatter}}

performs well. An alternative approach might work better when these cluster outliers consist of small groups of cells. One approach might be a less sensitive weighted cascade measure that also accounts for larger regions as potential scatter, such as

ε_{\text{sc}}(k) = \frac{E_{\text{exp}}(k) + \sum_c \sum_i \kappa(i)}{\sum_c \sum_i \kappa(i)},

with

p > 0

being much smaller than the number of cells.

To evaluate the clustering error in expression space we use

ε_{\text{exp}},

the average distance, in expression space, of a cell to the center of the cluster it belongs to,

ε_{\text{exp}}(k) = \frac{1}{n} \sum_{i=1}^{n} \text{dist}(\text{center}(c_i), c_i),

where

n

is the number of clustered cells,

c_i

is the

i\text{th}

cell, \text{center}(c_i)

is the center of the cluster to which

c_i

belongs, and \text{dist}(\cdot, \cdot)

is the distance operator used in the clustering process.

We compute

ε_{\text{scatter}}

and

ε_{\text{exp}}

for

2 \leq k \leq m,

with

m

being the first value where

ε_{\text{scatter}} > 50%.

If the pattern of only one gene is used in the clustering, we use

ε_{\text{scatter}} > 60% as termination criterion instead because variations in background expression have a stronger impact on the cluster analysis, and because more complex structures are possible when multiple genes with spatially overlapping patterns are clustered. By using these thresholds for

ε_{\text{scatter}},

we ensure that we iterate over all potentially useful clusterings and do not terminate prematurely. We use

k = 2

as starting point because it represents the first potentially useful clustering. Furthermore, considering the relatively large value of

ε_{\text{exp}}(1),

starting at

k = 1

would result in a suggested value for

k

that is too small.

To identify a value

w

for

k

for which the error in expression space is sufficiently low to well characterize the data, we identify the first

k

for which the decrease in

ε_{\text{exp}}

is lower than the average decrease

\Delta \epsilon_{\text{exp}} = \frac{\epsilon_{\text{exp}}(2) - \epsilon_{\text{exp}}(m)}{m-2}.

Alternatively

w

could also be defined as the

k

that corresponds to the point of the

ε_{\text{exp}}

evaluation curve that is furthest from the line defined by

ε_{\text{exp}}(2)

and

ε_{\text{exp}}(m)

[33].

While the first approach tries to find the

k

for which the expression error has sufficiently decreased, the second approach tries to identify the so-called “knee” of the

ε_{\text{exp}}

evaluation curve. Both methods depend on

m,

but this dependency is well behaved, i.e., with increasing

m

the suggested

w

changes slowly and continuously. During the research and development of this work, both methods seem to work equally well.

We use the following algorithm to identify a good initial

k > w

that also results in a relatively low physical scattering:

\begin{align*}
    k &= w + 1 \\
    l &= k \\
    &\text{for } i \leftarrow 1 \text{ to } m \\
    &\quad \text{do } \{ \\
    &\quad \quad \text{if } (\epsilon_{\text{scatter}}(i) < \epsilon_{\text{scatter}}(l) + t) \\
    &\quad \quad \quad \text{then } k = i \\
    &\quad \quad \quad \text{if } (\epsilon_{\text{scatter}}(i) < \epsilon_{\text{scatter}}(l)) \\
    &\quad \quad \quad \text{then } l = i \\
\end{align*}

Initially,

k

is set to

w + 1,

which is the lowest value that results in a sufficiently low expression error. Then, the algorithm tries to optimize the expression error as well as the physical scattering by searching for a

k > w

that also results in a relatively low physical scattering.

Here we use a threshold of

\(t = 4\%\)

– determined through empirical testing – to restrict the maximal allowed increase in

ε_{\text{scatter}}

with respect to

l,

i.e., the

k

with the lowest relative physical scattering visited so far. Since

ε_{\text{exp}}

decreases with increasing values of

k,

the error in expression space for the suggested

k

is guaranteed to be smaller than

ε_{\text{exp}}(w).

Alternatively, one can also view the problem of finding a good initial

k

as an optimization problem by looking for the

k

that minimizes

w_{\text{total}}(k) = | \epsilon_{\text{exp}}(k) - \frac{1}{2} \epsilon_{\text{scatter}}(k) |,

where both

ε_{\text{exp}}

and

ε_{\text{scatter}}

are normalized. Conceptually, the first approach is more intuitive, does not require normalization of the evaluation functions, and will always suggest a minimum of

ε_{\text{scatter}}

if an adequate local minimum exist. Using

\(w_{\text{total}}\)

for finding a good initial

k

has the advantage that it does not rely on a threshold

r.

Furthermore, it may result in a more reliable suggestion in cases where

ε_{\text{scatter}}

is degenerate since

\(w_{\text{total}}\)

does not directly rely on the notion that the physical scattering increases with increasing values of

k.

In practice both approaches have shown to be useful.

Beginning with an initial, suggested value of

k,

the user can then determine the “best”

k

based on the information from the cluster evaluation and previews of the different evaluated clustering results using an Unrolled View. Even though the initial, suggested value of

k

may not always be optimal, our testing has revealed there is value in providing a “reasonable” value or range of values for

k.

Figure 7 shows the cluster evaluation functions for the clustering of

gt

and

Kr.

To provide an overview of both functions in one plot, we show

ε_{\text{scatter}}

along with

ε_{\text{exp}}(k) = \frac{\epsilon_{\text{exp}}(k)}{\epsilon_{\text{exp}}(2)}.

The suggested

k

is eight, which is also a strong local minimum of

ε_{\text{scatter}}.

The corresponding clustering result for

k = 8

is shown in Figure 5.

Figure 8 shows two additional example classifications
of $gt$ and $Kr$ using $k = 5$ and $k = 19$. $k = 5$ is the highest level for which $\varepsilon_{\text{scatter}} = 0$ and $k = 19$ is a local minimum of $\varepsilon_{\text{scatter}}$ ($\varepsilon_{\text{scatter}}(19) \approx 35.29\%$) close to the middle of the range. Here, we see that the suggested level of $k = 8$ provides a good compromise between high-level and low-level descriptions of the patterns. The value of $k$ that is best suited to investigate a biological question depends to a large degree on user requirements. Further example usages of $\varepsilon_{\text{scatter}}$ and $\varepsilon_{\text{exp}}$ are provided in Sections VI and VII.

$\varepsilon_{\text{exp}}$ and $\varepsilon_{\text{scatter}}$ are global cluster quality measurement functions in the sense that the clustering quality is evaluated based upon the entire dataset (in this case, all classified cells). Global error measures might not be appropriate if the user performs a clustering of a larger number of cells but is interested only in a small subset of clusters defining some local feature of interest.

VI. SINGLE PATTERN ANALYSIS

Genes are frequently expressed in complex patterns that show a wide range of quantitative changes in expression across the cells of an embryo. Although for some analyses, the data is best left unclassified in this form —simply using the expression values in all cells—it can also be revealing to divide a single pattern into one or more distinct regions. For example, on/off descriptions of expression have been useful in logical models of gene networks [34], [35].

However, discretizing a gene pattern via manual thresholding can be problematic—it may be very time-consuming, and the choice of thresholds is arbitrary and not fully data-dependent. To address this challenge, one can use, for example, k-means clustering and Euclidean distances to compute a number of data-dependent thresholds. Each of the $k$ clusters then represents a specific threshold range that can be interpreted as a different confidence level. Different components of a pattern may be regulated by different genes, so different thresholds may be appropriate for different regions. Cluster post-processing, such as splitting clusters into their main spatial components, allows different threshold levels to be selected for different components of a gene pattern. Alternatively, as described in Section IV-C, for genes with clearly distinct spatial expression domains, cell positions may be used in the initial clustering to enforce creation of separate clusters for spatially distant components of a pattern. Rather than choosing some arbitrary thresholds, clustering automatically suggests thresholds based on the histogram of the data. The k-means clustering algorithm seeks to minimize the mean squared distance from each data point (cell) to its nearest cluster center. To achieve this goal, the k-means algorithm will create $k$ cluster centers positioned according to the density distribution of the expression values of the selected gene.

Figure 9 shows three example classifications of the eve expression pattern using different numbers of clusters $k$. While $k = 2$ produces a threshold that is too high and does not capture all parts of each stripe, a clustering with $k = 3$ correctly identifies the seven stripes of the eve expression pattern. By increasing the number of clusters, additional details within the stripes along the dorsal-ventral axis can be seen, as well as an additional cluster selecting cells in the inter-stripe regions. This complex description illustrates that thinking of a gene as being either on or off is usually too simplistic. The fact that clustering automatically reveals differences along the dorsal-ventral axis demonstrates the usefulness of such analyses. The pair-rule genes, such as eve, are not typically thought of as dorsal-ventral regulators, but consistent with the clustering results, careful quantitation of the levels of eve and similar gene’s expression has shown they indeed show up to two-fold changes in expression along the dorsal-ventral axis suggesting a dorsal-ventral component in pair-rule regulation [3], [2]. Analyzing the actual meaning of these moderate changes requires computational tools, such as cluster analysis, to provide objective measures of their significance.

Figure 10 (left) shows the curves of the cluster evaluation functions $\varepsilon_{\text{scatter}}$ and $\varepsilon_{\text{exp}}$. In this case, $\varepsilon_{\text{scatter}}$ is rather smooth and monotonically increasing indicating that all $k$ with $\varepsilon_{\text{scatter}} > 0$ may result in valid clusterings of the eve pattern. This behavior can be explained by the very high signal-to-noise ratio of the eve expression data, which was averaged from dozens of embryos. The suggested $k$ is five, which is the largest $k$ for which only one cluster representing low eve expression is created (see Figure 10, right). A clustering with $k = 5$ provides a compromise between a high-level and low-level description of the eve expression pattern.

Binarized versions of the eve pattern (i.e., on/off descriptions) can be created by merging the different clusters, allowing one to easily compare the different classifications by defining their overlay (see Figure 11). While $k = 3$ and $k = 6$ result in similar classifications of the seven stripes, the clustering with $k = 2$ misses many cells of the pattern. Thus, first generating multiple clusters

Fig. 9. a) The expression pattern of eve at stage 5:9-25%; Classification of eve with b) $k = 2$; c) $k = 3$; and d) $k = 6$. While the $k = 2$ clustering produced a threshold that was too high, erasing too many cells from the pattern, the $k = 3$ clustering was better able to identify the seven stripes of the eve expression pattern. The $k = 6$ clustering identified additional characteristic variations within the stripes along the DV-axis as well as an additional cluster that selects some inter-stripe cells showing some higher expression of eve.

Fig. 10. Cluster evaluation functions $\varepsilon_{\text{exp}}$ (red) and $\varepsilon_{\text{scatter}}$ (blue) for the clustering of the eve expression pattern (left), with $w = 4$ and $m = 9$. The suggested $k$ is five, which is the largest $k$ for which only one cluster representing low background eve expression is created (right).
In Figure 14, the same clusters as in Figure 13 are shown to support simultaneous visualization of many more genes. PCX also provides 2D and 3D parallel coordinates due to the fact that only three gene dimensions can be visualized informatively in gene expression space visualization, but are limited to the regulatory relationships between transcription factors and their levels. Such plots can be very useful in identifying potential novel regulatory relationships.

Once derived, we use these individual stripe clusters to highlight the seven eve stripes via color in different abstract views. For example, the expression behavior of *gt*, *hb*, and *Kr*—three known transcriptional regulators of *eve*—can be revealingly analyzed within each of the eve stripes using a 3D scatter-plot (Figure 13). Here, large differences between stripes are visible, the seven stripes form very distinct point clusters within the scatter-plot. This behavior is consistent with current models suggesting that the expression pattern of *eve* does not simply consist of seven identical stripes, but that many stripes are regulated independently. The available data suggests that *gt*, *hb*, and *Kr* control some stripes, but the scatter-plot suggests that these factors have the potential to regulate all stripes by their unique combinations of expression levels. Such plots can be very useful in identifying potential novel regulatory relationships between transcription factors and their targets.

Generally, scatter-plots have proven to be a very intuitive and informative gene expression space visualization, but are limited due to the fact that only three gene dimensions can be visualized at once. PCX also provides 2D and 3D parallel coordinates to support simultaneous visualization of many more genes. In Figure 14, the same clusters as in Figure 13 are shown in a 2D parallel coordinate view of early stage *hb*, *gt*, *kni*, *Kr*, and *tll*. The average expression of the seven clusters in the different genes are shown via additional, thicker lines of darker colors and the associated standard deviations are shown via boxes placed on each parallel axis. Highly transparent color bands shown in front of the plot are used to further highlight the different clusters.

VII. TEMPORAL VARIATION ANALYSIS

Gene expression patterns are not static but highly dynamic. Understanding the temporal profile of a gene expression pattern is essential if we are to understand complex relationships between genes. Even though visual inspection of an expression pattern at different time steps provides an impression of the general temporal behavior of a gene, many important features, such as groups of cells with a similar temporal expression profile, are not easily detected and visual quantification of temporal change is not accurate. For example, the pattern of *giant* (*gt*) expression can be seen to change between six time cohorts within one hour, but it
Fig. 15. The expression pattern of *giant* (gt) shown at six different time cohorts of stage 5 of embryo development.

Fig. 16. Based on the patterns of gt shown in Figure 15, cells were classified into seventeen clusters as suggested by $\varepsilon_{\text{scatter}}$ and $\varepsilon_{\text{exp}}$. Two clusters selected cells showing only background expression of gt at all time steps and are therefore not shown here. Clusters 1, 8, and 16 were each split into their two main spatial components. The remaining clusters were not split, since no significant divergence in the temporal expression profile between their main spatial components could be identified. a) An unrolled view showing all eighteen clusters of interest. b-h) The user grouped the eighteen temporal clusters into seven main groups based on their average temporal expression profiles in gt. The six time steps are shown on the x-axis and the expression level on the y-axis of each plot. The spatial patterns defined by the different clusters are displayed in the accompanying unrolled view plots.

Fig. 17. Cluster evaluation functions $\tilde{\varepsilon}_{\text{exp}}$ (red) and $\varepsilon_{\text{scatter}}$ (blue) for the clustering of the six time steps of gt with $w = 10$ and $m = 54$. The suggested number of clusters $k$ is seventeen. $\varepsilon_{\text{scatter}}$ further indicates that seventeen is the highest $k$ for the particular level of detail with relatively low overall physical cluster scattering. It is not possible to rigorously describe how (see Figure 15). To show how PCX can assist in the analysis of the spatio-temporal expression pattern of genes, we have used clustering to classify cells into groups of similar temporal behavior.

In Figure 17, the curves of the cluster evaluation functions $\varepsilon_{\text{scatter}}$ and $\varepsilon_{\text{exp}}$ are shown. The suggested number of clusters $k$ is seventeen, which is also a local minimum of $\varepsilon_{\text{scatter}}$ with $\varepsilon_{\text{scatter}}(17) \approx 31.88\%$. The overall behavior of $\varepsilon_{\text{scatter}}$ indicates that $k = 17$ is the largest $k$ at the particular level of detail for which $\varepsilon_{\text{scatter}}$ is still relatively low. A comparison of $\varepsilon_{\text{scatter}}(17)$ to the next two lower local minima of $\varepsilon_{\text{scatter}}$ – with $\varepsilon_{\text{scatter}}(12) \approx 31.21\%$ and $\varepsilon_{\text{scatter}}(10) \approx 29.25\%$ – shows only a moderate increase in $\varepsilon_{\text{scatter}}$. When comparing $\varepsilon_{\text{scatter}}(17)$ to the $\varepsilon_{\text{scatter}}$ values of the next two larger local minima of $\varepsilon_{\text{scatter}}$ – with $\varepsilon_{\text{scatter}}(19) \approx 36.34\%$ and $\varepsilon_{\text{scatter}}(22) \approx 40.14\%$ – a significantly higher increase in relative physical cluster scattering is visible. This behavior can be interpreted as an indication that $k = 17$ may also provide a good compromise between a high-level and low-level description of the temporal variation of the gt expression pattern. A level of $k = 17$ was also confirmed to be appropriate by users of PCX.

Figure 16 shows as an example the result for gt, in which its expression patterns at six successive time cohorts were classified into seventeen clusters using k-means clustering and Euclidean distances. Two of the seventeen clusters selected cells showing only background expression at all time steps and are not shown. Each of the other fifteen clusters show distinct average expression profiles (the differently colored lines plotted in Figure 16), though some clusters show profiles that are closely related. In the figure, the user has grouped these clusters into seven main sub-groups based on their temporal average expression profiles, shown in panels b-h. In addition, clusters 1, 8 and 16 have each been split into two components to separate their anterior and posterior components.

Several trends can be readily seen from the different views of the analysis. The unrolled physical views show that clusters with similar average temporal expression profiles frequently, but not always, are adjacent to one another in the embryo. Expression within a set of clusters in the very anterior of the embryo increases particularly during the later time cohorts (visible, for example, in Figure 16b). Expression in the posterior margins of both of the major early gt stripes drops rapidly over the time series.
patterns. A complex combination of regulatory interactions drives these
than has been observed previously. These results suggest that
stripe moves anteriorly during this time series [36], [3], but the
(Figure 16f-h). It is known that the location of the posterior
eve stripe moves anteriorly during this time series [36], [3], but the
data show a much more complex pattern of temporal change than has been observed previously. These results suggest that
a complex combination of regulatory interactions drives these
patterns.

VIII. MULTIPLE PATTERN ANALYSIS
To dissect the complex regulatory interactions between genes, the expression patterns of different transcription factors that potentially act together as regulators may be used as input to cluster analysis. Cells are classified into clusters that have similar combinations of expression for the input set of regulators. Each cluster thus describes one potential sub-pattern that a regulatory
network composed of these factors could give rise to. The total
number of clusters then gives an approximation of the maximal
complexity of the output of the network. The results of such a
clustering can also be compared to the expression patterns of suspected target genes to assess possible regulatory relationships.

To provide an example of such multi-gene clustering, we exam-
ined the relationship between the three transcriptional regulators
giant (gt), hunchback (hb), and KRüppel (Kr) and the second
stripe of the eve gene. These three factors are well-characterized
regulators of this expression stripe; hb is an activator and Kr and
gt are repressors [37]. As discussed in Section VI, the seven
stripes of eve form characteristic clusters in gene expression space
with respect to gt, hb, and Kr expression. By using these three
factors’ expression patterns as input to a clustering analysis, we
can identify the potential expression pattern components that can be defined based on these regulators (see Figure 18). We used their
mRNA expression values from the first temporal cohort (0%-3%
invagination) to simulate their protein expression values at the
third temporal cohort (9%-25% invagination) – the stage of the
eve comparison target. We have found this lag, on average, to be
optimal for all regulators [7]. In the example, cells are classified
into twenty-two clusters that map to locations throughout the
embryo. Eight of these clusters are of interest to the control of eve
stripe 2, five that lie within the stripe and three in the flanking
inter-stripe regions. The five clusters within stripe 2 define the
center, the anterior and posterior borders, as well as a ventral
portion of the stripe, suggesting that these characteristic parts of stripe 2 may be different (see Figure 18b).

To validate the structure formed by the clusters against the
target pattern, cluster colors are mapped onto an expression
surface of eve, in which height shows the level of expression
(Figure 19). It can be seen that the five clusters fit closely to the
expression pattern of the target stripe 2.

Based on the average expression curves, the characteristic
expression pattern of the potential regulators in the eight clusters
that are within and flanking stripe 2 are easily visible (see
Figure 18c, d). Here, hb is expressed at high levels in all clusters
except those posterior of stripe 2, consistent with its known role
as an activator of stripe 2. Kr is expressed at high levels only
posterior of stripe 2 and gt is expressed at high levels only anterior
of stripe 2, consistent with their known roles as repressors that
define the posterior and anterior borders of stripe 2, respectively.

Interestingly, the two clusters that form short ventral patches on
eve stripe 2 (yellow and blue) show significantly lower expression
of hb than the two clusters that lie dorsally to them (red and
green) (see Figure 18c, d). This correlates with a lower level of
eve expression in this ventral margin (Figure 19) and suggests
that this reduced expression may be the result of lower activation
by hb. hb is typically thought of as regulating gene expression
only along the anterior/posterior axis of the embryo. The cluster
analysis suggests that it may also be able to mediate differential
transcription along the dorsal-ventral axis. However, if we were
to add a dorsal-ventral gene, such as snail (sna) (see Figure 1)
into the analysis, it would be difficult to distinguish if the ventral
gap in eve stripe 2 resulted from direct inhibition by sna, if
sna acted via inhibiting ventral hb expression, or if all three
expression patterns are parallel manifestations of dorsal-ventral
patterning systems, each acting separately. Thus, cluster analysis
can be used for identifying interesting correlations that might
result from novel biological interactions or phenomena, but the
analyses should be confirmed by experimental data.

This case study illustrates that clustering the expression patterns
of multiple regulators can provide confirmation and additional
insights into known regulatory interactions. It is likely that the
extension of this strategy to less well-characterized systems will
suggest potential regulatory interactions that can then be tested.
IX. Conclusions

Our overall objective for this work has been to provide important new capabilities to accelerate scientific knowledge discovery. Our work helps biologists, who aim to discover potentially new, experimentally verifiable biological interactions, by providing the ability to to define, analyze, and iteratively refine clusters in multiple, linked views. For computational biologists, we have presented objective methods for classifying quantitative data-points in spatial datasets.

We have shown how data clustering and visualization can be integrated into one framework and how our system can be used effectively to explore and analyze 3D spatial expression data. A system of linked multiple views is used for data exploration and for steering the analysis process, helping bridge spatial patterns of expression with abstract views of quantitative expression information.

Data clustering then provides means for automatically defining cell selections, depicting characteristic data features and in this way improving the visualization. We have shown how dedicated post-processing of clustering results based on visualization and user knowledge improves the analysis. We have demonstrated how the combination of \( \varepsilon_{\text{scatter}} \) as measure to describe the relative physical scattering of clustering results and \( \varepsilon_{\text{exp}} \) to suggest a good initial value for \( k \) in combination with visual validation of clustering results can be used to determine appropriate values for \( k \).

Analysis of 3D spatial gene expression data is a challenging task requiring unique strategies not encountered in studies of 1D non-spatial data, such as microarray expression data. Using our integrated data visualization and clustering approach, we have shown how the pattern of a gene and its temporal variation can be defined and analyzed. We have shown how suspected relationships between genes can be analyzed to address the question of how the pattern of a gene is created by the action of multiple regulators.

Along with the first release of the BDTNP 3D gene expression database, we have also made a version of PCX freely available to the public [1]. Data clustering and 3D parallel coordinates are currently in active use by BDTNP members and will soon also be included in the public version of PCX.

In PCX, spatial information is incorporated in the analysis process mainly by using cluster post-processing techniques such as splitting of clusters. Alternatively, \( x \), \( y \), and \( z \) cell positions can directly be added to the cluster analysis. However, this may result in clusters defined by a complex mix of spatial and expression influences which may not be easy to interpret.

Development of additional analysis techniques that effectively integrate spatial and gene expression information is one focus of future work. Adaptation of spatial clustering methods, such as the dual clustering approach proposed by Cheng-Ru et al. [38], is only one promising approach. Alternatively, one could perform clustering based on gene expression information only, then split the resulting clusters into spatially distinct sub-clusters, and then perform a re-clustering based on the centers of the detected sub-clusters. In PCX we currently use hierarchical clustering only for partitioning of the data. By traversing the data hierarchy created in a hierarchical clustering, exploration of the data at multiple levels of detail becomes possible. In addition to clustering with cells, clustering of genes as well as biclustering promises to provide further insights into the data. In addition, matrix decomposition techniques, such as principal component analysis (PCA) and singular value decomposition (SVD) [39], [40], have successfully been applied to other types of gene expression data. Integration of these and other analysis techniques into PCX should further increase its value for practical use and impact.

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References

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