Cluster Tendency Assessment in Neuronal Spike Data

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Abstract

Sorting spikes from extracellular recording into clusters associated with distinct single units (putative neurons) is a fundamental step in analyzing neuronal populations. Such spike sorting is intrinsically unsupervised, as the number of neurons are not known a priori. Clustering can be performed using either the full dimensional waveforms or on low-dimensional representations obtained by dimensionality reduction or feature extraction methods. In this paper, we evaluate the utility of several methods for providing lower dimensional visualization of the cluster structure and on subsequent spike clustering. Experimental results are conducted on two datasets with ground truth labels. We also introduce a visualization technique called improved visual assessment of cluster tendency (iVAT) to estimate possible cluster structures in data without the need for dimensionality reduction. In data with a relatively small number of clusters, iVAT is beneficial in estimating the number of clusters to inform the initialization of clustering algorithms. With larger numbers of clusters, iVAT gives a useful estimate of the coarse cluster structure but sometimes fails to indicate the presumptive number of clusters. We show that noise associated with recording extracellular neuronal potentials can disrupt computational clustering schemes, highlighting the benefit of probabilistic clustering models. Results show that t-Distributed Stochastic Neighbor Embedding (t-SNE) feature extraction provides representations of the data that yield more accurate clustering than low-dimensional representations obtained with four other methods. Moreover, the clustering obtained using t-SNE representation outperforms clustering within the other four feature spaces and also with algorithms operating directly within the input dimensional space.
1 Introduction

Recording of extracellular signatures of action potentials, referred to as spikes, is a standard tool for revealing the activity of populations of individual neurons (single units). Single unit activity contains fundamental information for understanding brain microcircuit function in vivo and in vitro [Buzsáki, 2004]. Inferences about network activity can be made by identifying coincident activity and other temporal relationships among spiking patterns of different neurons [Brown et al., 2004]. However, the reliability of these inferences is strongly influenced by the quality of spike sorting, i.e., the detection and classification of spike events from the raw extracellular traces with the goal of identifying single-unit spike trains. Poor sorting quality results in biased cross-correlation estimates of the spiking activity of the different identified units [Ventura and Gerkin, 2012].

The typical workflow for spike sorting includes spike detection, feature extraction, and clustering. While detection is pretty straightforward and can be efficiently done with simple thresholding, the feature extraction and clustering procedures are far from being satisfactorily settled [Rossant et al., 2016]. It has been estimated that single or tetrode type electrodes (i.e. impedance < 100KΩ) can record neuronal activity within a spherical volume of 50 µm radius with amplitudes large enough to be detected (> 60µV). This volume of brain tissue constitutes about 100 neurons, however, it is expected that many of them will be silent [Buzsáki, 2004, Shoham et al., 2006]. To investigate the performance limits of spike sorting algorithms, different sets of ground truth data have been used. Pedreira et al. [2012] and Niediek et al. [2016] reported that it is possible to identify 8 to 10 out of 20 units with less than 50% false positive and false negative rates.

In practice, since we can’t physiologically verify how many neurons have been recorded, assigning the spikes within a recording to individual neurons remains a fundamental technical issue. The sorting is in essence an unsupervised learning challenge. Therefore, methods require one of two approaches: specification of a fixed value of the number of clusters to seek (c); or generation of candidate partitions for several possible values of c, followed by selection of a best candidate based on various post-clustering validation criteria. Moreover, improving spike classification to correctly identify cell types is a topic of interest highlighted by initiatives that aim to characterize and reconstruct different cell types in the brain and their role in health and disease [Jorgenson et al., 2015, Markram et al., 2015]. For that goal, Armananzas and Ascoli [2015] list the identification of the number of clusters as the first outstanding question in techniques for neuronal classification.

In summary, identifying the spiking trains of individual units within a recording is a three-faceted problem: (i) assessing the cluster tendency in the pre-clustering phase (before initializing the algorithm); (ii) clustering (iii) evaluation of the validity of the clusters that have been isolated, post-clustering [Bezdek, 2017]. Spike sorting algorithms usually start by projecting the data to a lower dimensional space. Although there are several reasons to do this such as reduction of computation time, the fundamental reason is that projections allow visualization of high dimensional input data. In turn, this facilitates the choice of a few
selected values of the integer c. Pre-specification of c is needed by almost all clustering algo-
risms as an input parameter (hyper parameter). In practice, since reduced dimensionality
embedding of the data often does not provide visually well separated clusters, it is common
to exclude large number of spikes and only take into account a small core portion of the
subsets that seems to have well-isolated clusters [Dehghani et al., 2016]. Omitting spikes to
obtain well-separated clusters may lead to single units with recognizable spike waveforms,
but it discards spikes that, as mentioned before, are fundamental for analyses of temporal
structure of spiking activity [Cohen and Kohn, 2011, Pazienti and Grün, 2006]. Therefore,
the bottleneck in spike sorting is at the pre-clustering stage: viz., inaccuracy of the assumed
data structure that is inferred by visualization of it in the lower dimensional space. If clus-
tering is to be done in a lower dimensional data space, errors here will affect both the initial
estimate of the cluster number and the performance of the clustering algorithm. Thus, this
study concerns itself with the pre-clustering stage.

The contributions of this manuscript can be summarized as follows:

• visualization of cluster structure: we compare the visualization of neuronal spike data
created using six methods (i) three well-known dimensionality reduction techniques:
principle component analysis (PCA), t-student stochastic neighborhood embedding
(t-SNE) and Sammon’s algorithm, (ii) two methods that extract features from the
waveforms: wavelet decomposition and features such as peak to valley amplitude and
Energy (PV-E), and (iii) a method that operates directly on waveforms in the input
space: improved visual assessment of tendency (iVAT). The analysis is performed on
two different type of ground truth data (labeled data): simulated spike sets and real
recorded spike sets, called dataset-1 and dataset-2, respectively. Our results indicate
that iVAT often shows the primary (or coarse) cluster structure, while t-SNE is often
capable of displaying finer cluster structure.

• Assessment of clustering quality using different projections of the data (i.e. PCA,
Waveform features, Wavelet features, t-SNE and Sammon)

  – by using ground truth data we evaluate quality of partitions obtained by clus-
tering in the upspace (input dimensional space; i.e., the waveforms) and also
in the five different two-dimensional representations obtained with the 5 projec-
tion methods. This test is performed by running c-means (also called k means
in the literature) and generating a number of clusters equal to the actual (i.e.
known) number of units. The quality of the partitions generated by each method
is evaluated by considering Dunn’s index (DI) (an internal index describing the
intrinsic quality of the generated clusters); the generalized Dunn’s index GDI_{33};
and the Adjusted Rand’s index (ARI) (an external measure of agreement between
computed partitions and the ground truth partition).

  – We also compare clustering in the downspace using c-means on the t-SNE em-
bedded data and Osort that is applied to the input data.
The outline of the paper is as follows. Section 2 provides a description of the datasets that we used (2.1), defines the problem (2.2), explains the methods used for data visualization (2.3 and 2.4), and lastly describes the measures used for evaluating clustering structure of the data (2.5). The results of the experiments on the datasets are given in section 3. Insights gained from the experiments are summarized in section 4.

2 Materials and Methods

2.1 Ground truth data

The importance of ground truth data for spike sorting validation is emphasized by Einevoll et al. [2012]. We use two labeled datasets, i.e., the label or membership of each spike for an individual neuron is known. Our first experiment uses simulations of extracellular traces (hereafter called dataset-1), and the second experiments uses data obtained from in-vivo experiments (hereafter called dataset-2).

**Dataset-1**: Pedreira et al. [2012] simulated extracelluar traces that contain the activity of 2 to 20 neurons with additive background noise. The single unit spike activity is generated by using average spike waveforms (templates) compiled from Basal Ganglia and Neocortical recordings. The background noise (i.e. lfp noise) is simulated by superimposition of thousands of spikes at random times which were then scaled down to a standard deviation of 0.1. Each simulated trace also contains multi-unit activity, which was generated by superimposing 20 waveforms with normalized amplitudes limited to 0.5. Dataset-1 thus provides us with simulated extracellular traces each containing 3 to 21 subsets of spikes. For each cluster number \( c \), five simulations using different sets of templates were generated (for a total of 95 datasets). The spikes were detected by voltage thresholding. The length of each spike waveform is 2 ms, with a sampling rate of 25 kHz, comprising 48 sample points. Thus, the upspace dimension for subsets in Dataset-1 was 48.

**Dataset-2**: We used the in vivo simultaneous intracellular and extracellular hippocampal recording datasets that are publicly available from the CRCNS website [Henze et al., 2009]. The dataset contains raw data of simultaneous intracellular and extracellular recordings from neurons in the CA1 region of the hippocampus of anesthetized rats. The experimental procedure consisted of inserting extracellular electrodes (either tetrodes, 13-\( \mu \)m polyimide-coated nichrome wires, or a single 60-\( \mu \)m wire) into the cell body layer of CA1, confirming the presence of unit activity in the recordings, and then inserting intracellular sharp-electrodes into the same region in close proximity to the extracellular electrode to impale a single neuron and record stable action potentials induced by current injections. With this method, it was possible to capture simultaneous spikes in the intracellular and extracellular recordings. We detected the intracellular spikes, and used those spike times to extract the extracellular spike train of that neuron (i.e. a labeled subset of spikes). Each spike waveform is 2 ms long which, with a sampling rate of 40 kHz, results in 80 sample points, so the upspace dimension
of each spike and subsequent sets in dataset-2 was 80. Dataset-2 is valuable because each 
spike waveform in the subset is a recorded signal from a physiological setting; hence, the 
variability in the probability distribution of each subset comes from either natural (e.g. the 
effect of other current sources in the extracellular medium) or experimental conditions (e.g. 
electrode drift). Each recorded trace has one labeled subset of spikes. Hence, to generate 
each mixture, we used the extracted spike subsets from different traces. In total, we obtained 
9 subsets of spikes from the database and then used combinations of 2,3,4,... to 9 of these 
subsets to create datasets containing spikes of 2 or more neurons (for a total of 502 datasets). 
In summary, the data for our experiments were mixtures of subsets of spikes each with different 
population size in either 80 or 48 dimensional input space.

2.2 Problem Definition

Let $X = \{x_1, \ldots, x_n\} \subset \mathbb{R}^p$ denote a set of vector data representing $n$ spikes generated by one 
or multiple neurons. The coordinates of $x_i$ are voltage samples that describe a spike event 
(they are always voltage samples in this article). The non-degenerate crisp $c$-partitions of 
the $n$ objects in a set $X$ can be represented by a $c \times n$ matrix $U$ in $M_{hcn}$, written in terms 
of the $c$ crisp subsets of it (the clusters $X_i$) as

$$
M_{hcn} = \{U \in \mathbb{R}^{cn} : u_{ik} \in \{0, 1\} \forall 1 \leq i \leq c, 1 \leq k \leq n; \sum_{i=1}^{c} u_{ik} = 1 \forall k; \sum_{i=1}^{c} u_{ik} > 0 \forall i\} \quad (1a)
$$

$$
U \in M_{hcn} \iff X = \bigcup_{i=1}^{c} X_i; X_i \cap X_j = \emptyset | i \neq j \quad (1b)
$$

Finding clusters in $X$ comprises three steps: deciding how many clusters ($c$) to look for; 
constructing a set of candidate partitions \{$U \in M_{hcn}$\} of $X$; and selecting a ”best” partition 
from CP (cf. equation (2) below) using a cluster validity index (CVI).

2.3 Dimensionality reduction and Feature extraction

Real data vectors in $\mathbb{R}^p$ usually have high dimensionality ($p > 3$) (e.g., images, videos, 
and multi-variate data streams). Feature selection Zhao et al. [2013] and dimensionality 
reduction algorithms van der Maaten et al. [2009] are used to (i) make a visual assessment 
of the structure of the data and (ii) to improve the performance of data-driven procedures, 
such as those for classification and clustering.

Here we focus on several well-known dimensionality reduction algorithms that have been 
used in a multitude of domains including neurosciences; we refer the interested reader to van 
der Maaten et al. [2009] and references therein for technical details.
Principal component analysis (PCA) is one of the most important and widely utilized linear
dimensionality reduction techniques [Theodoridis, 2009]. In order to find a low-dimensional
subspace suitable for projecting the input data, PCA projects the data along the directions
given by the leading eigenvectors of the data covariance, i.e., the directions associated with
the largest eigenvalues of the sample covariance matrix. In other words, PCA seeks a low-
dimensional representation that explains as much variance of the input data as possible.

In neuroscience research, another common approach is to extract features of the waveforms
that have a physical meaning: e.g. peak (P), valley (V), and energy (E) [Hattori et al., 2015,
Truccolo et al., 2011], hereby called PV-E. Another method based on wavelet transforms that
enables visualizing the data in the wavelet coefficient subspace has also been successfully
implemented in clustering packages such as Waveclus and Combinato [Niediek et al., 2016,
Quiroga et al., 2004].

Here, we will also consider two nonlinear dimensionality reduction techniques. The first of
these is t-SNE (t-student Stochastic Neighbor Embedding), developed by van der Maaten and
Hinton [2008]. It works by converting Euclidean distances between high-dimensional input
data into conditional probabilities. In doing so, t-SNE converts the geometric notion of
similarity into a statistical concept: if \( x_j \) is a neighbor of \( x_i \), then the conditional probability
\( p_{ji} \) is high. Then, t-SNE finds low-dimensional representations \( y_i \) and \( y_j \) of \( x_i \) and \( x_j \) by
minimizing the discrepancy between the upspace \( p_{ji} \) and downspace conditional probabilities
\( q_{ji} \), technically achieved by minimizing the Kullback-Leibler divergence between them. The
objective of t-SNE is to minimize the sum of the divergences over all the data points.

Two features of t-SNE should be noted. First, it is not a linear projection like PCA but rather
has a non-convex cost function, so its output may be different for different initializations.
Second, it is a parametric technique. Different settings of hyperparameters such as the
learning rate, the perplexity, and the iteration rate in the t-SNE algorithm generate different
maps in the scatterplots, and may cause misinterpretation of the data structure [van der
Maaten and Hinton, 2008].

The main parameter that affects the results of t-SNE is the perplexity, which is the limiting
condition for the entropy of the probability distribution of the similarities of datapoints
in the upspace. This means that the variance of the Gaussian that is centered over each
datapoint, i.e., the extent of the neighborhood around that point, is limited by the choice of
perplexity.

This limitation affects each datapoint separately based on the local density of the data. This
is the feature that enables t-SNE to avoid crowding points in the center of the map so that
cluster structure of the data in the upspace data is often seen in the tSNE downspace pro-
jection. This feature, however, comes at the cost of sacrificing the shape of the distribution
so that the distances between the clusters may not be meaningful. In other words, it is not
possible to infer reliable spatial information from the topology of the low-dimensional maps.
Fortunately, the topology is not relevant for our application: viz. revealing clusters in the
neuronal waveform data. The optimal choice of perplexity is dependent on the number of points (spikes) in the dataset. We found that for neuronal datasets with thousands of spikes (data points), as long as the extreme values in the parameter ranges are not selected, the t-SNE algorithm is not very sensitive to changes in perplexity. On the other hand, the reliability of t-SNE visualizations seems to decrease as the number of samples decreases. See [Mahallati et al., 2018] for an example.

We also consider another traditional nonlinear dimensionality reduction technique called the Sammon mapping [Sammon, 1969], which is one form of multidimensional scaling. Multidimensional scaling (MDS) seeks a low dimensional embedding of the input data while preserving all pairwise Euclidean distances (In a more general setting, t-SNE can be interpreted as a form of probabilistic MDS). However, high-dimensional data usually lies on a low-dimensional curved manifold, such as in the case of the Swiss roll [Tenenbaum et al., 2000]. In such cases, preserving pairwise Euclidean distances will not capture the actual neighboring relationships: the actual distance between two points over the manifold might be much larger than the distance measured by the length of a straight line connecting them, i.e., their Euclidean distance). Sammon mapping improves upon classic multidimensional scaling by directly modifying its original cost function, i.e., the distortion measure to be minimized. In particular, the Sammon mapping cost function weights the contribution of each pair of data points relative to the overall cost by taking into account the inverse of their pairwise distance in the original high-dimensional input space. In this way, Sammon mapping often preserves the local structure of the data better than classical multidimensional scaling.

While these five methods do not all produce lower dimensional data with an analytic projection function, we will call all downspace data sets projections.

2.4 iVAT

There are a number of imaging techniques that can be applied directly to the upspace data before clustering it. Here we describe the iVAT method described in [Havens and Bezdek, 2012], which is a generalization of the original VAT algorithm given by [Bezdek and Hathaway, 2002]. Improved Visual Assessment of Tendency (iVAT) is a visualization tool that uses the dissimilarity matrix, D, of the data to display potential cluster structure. The steps of the iVAT method are the following. The vectors in the dataset are represented as vertices in a complete graph, with the distances between them the weights of the graph. The algorithm first finds the longest edge in the graph. Then, starting at either end, it finds the minimal spanning tree (MST) of D based on Prim’s algorithm. Then, it reorders the rows (and columns) of D based on the order of edge insertion in the MST, creating D* (up to this point this is the original VAT algorithm). Then, iVAT transforms D* to D** by replacing each distance $d_{ij}$ in D* with the maximum edge length in the set of paths in the MST between vertices $i$ and $j$. When displayed as a gray-scale image, I(D**), possible
clusters are seen as dark blocks along the diagonal of the image. Images of this type are often called *cluster heat maps* in the neuroscience literature.

iVAT does not alter the physical meaning of the input data (even after the shortest path transformation), it just rearranges the objects in a way that emphasizes possible cluster substructure. The recursive computation of $D'^*$ given in Havens and Bezdek [2012] is $O(n^2)$. Appendix A.2 contains the pseudocode for iVAT. The iVAT algorithm requires no parameters to pick other than the dissimilarity function (d) used to convert X to D. This input matrix can actually be a bit more general than a true distance because its only requirements are that $D = D^T; d_{ij} \geq 0 \forall i, j; d_{ii} = 0 \forall i$. The most important points about this display technique are that it is applied directly to (a distance matrix of) the upspace data, so there is no distortion of the structural information introduced by a feature extraction function from the upspace to a chosen downspace, and iVAT preserves the physical meaning of the measured features. While any vector norm can be used to build an input matrix $D(X)$ from a set $X$ of feature vectors, the only distance used in this article is Euclidean distance. It is very important to understand that an iVAT image merely suggests that the input data has a certain number of clusters. Since iVAT can produce images from data of arbitrary dimensions, we can use it (or its scalable relative siVAT, Kumar et al. [2017]) to make a visual estimate of possible cluster structure in any upspace. While the iVAT algorithm is occasionally “wrong” (misleading), iVAT images usually provide some idea about the cluster structure of the input data [Bezdek, 2017].

Thus, iVAT provides clues about potential starting points for finding a useful partition of the input data. Mahallati et al. [2018] have shown the connection of VAT and iVAT to Dunn’s index and single linkage (SL) clustering. The intensity of the blocks in iVAT images are a (more or less) visual representation of the structure identified by single linkage clustering for labeled or unlabeled data. This suggests that iVAT might be regarded as a tool for “taking a peek” at a specific type of data structure in the input space.

### 2.5 Evaluating Cluster Quality

Cluster validity comprises computational models and algorithms that identify a “best” member amongst a set of candidate partitions (CP)

$$CP = \{U \in M_{hcn} : c_m \leq c \leq c_M\}$$

(2)

where $c_m$ and $c_M$ are the minimum and maximum specified values of the numbers of clusters sought.

The approach to identify a “best” partition $U$ (and concomitant value of $c$) in CP can be internal: using only information from the output results of the clustering algorithm, or external: using the internal information together with an outside reference matrix, usually the ground truth labels. Here, we use a classic internal scalar measure called Dunn’s index (DI)
Since we have labeled mixtures, we can calculate Dunn’s index on ground truth partitions in the upspace (input dimensional space) to give a measure of the compactness and isolation quality of the subsets in the original space. We have previously shown that this measure usually correlates with the quality of the visual assessment of potential cluster structure given by iVAT [Mahallati et al., 2018]. In the present work, we use a generalized version of Dunn’s index developed by Bezdek and Pal [1998] that uses the average distance from the mean as $\Delta$ and the average linkage clustering distance as $\delta$. It has been shown that this index is more robust with regards to sensitivity to outliers and hence produces more meaningful values for real life datasets with abundant aberrant points [Arbelaitz et al., 2013]. Equation 8 in Section A.3 gives the formula for GDI$_{33}$.

To evaluate the quality of the different clustering approaches we used adjusted Rand index (ARI) developed by Hubert and Arabie [1985], which is a well-known and fairly reliable criterion for performance assessment of the clustering results. The equation of the ARI is given in the appendix section A.3 (equation 9).

3 Results and Discussion

3.1 Visual Assessment of Cluster Tendency

It is impossible to make a direct visual assessment of a set of recorded spike waveforms $X = \{x_1, \ldots, x_n\} \subset \mathbb{R}^p$, since each waveform has more than three voltage samples (i.e., dimensions), $p > 3$. The upspace dataset $X$ can be mapped to a downspace dataset $Y \subset \mathbb{R}^q$ by a feature extraction function $\phi : \mathbb{R}^p \mapsto \mathbb{R}^q$ in many different ways. Dimensionality reduction methods are commonly employed for visualization purposes to gain insights into the data structure; and to provide clustering algorithms with lower-dimensional data to increase the computational efficiency. Next, we will demonstrate that different dimensionality reduction methods provide different low-dimensional representations of the data, and hence suggest different numbers of clusters. Towards this end, we used spike subsets of the simulated dataset that includes 5 simulations for each combination of different number of spike subsets for $c$ from 3 to 21. Below we show some representative results: two cases of $c=3$ (one with low Dunn’s index, DI, and one with higher DI) and then one case each for $c=5$, $c=10$, $c=15$ and $c=20$. 
Figure 1 shows two cases from the dataset with c=3. The colors in Figure 1 correspond to the three data labels. Bear in mind, as you view this and subsequent figures, that in the real case, the data are always unlabeled, so the projected data will be just one color, and the apparent structure will be much less evident than it seems to be in these figures. Figure 1(a) is a ‘good’ case in which all the algorithms map the spikes to projections with visually well-separated clusters and iVAT agrees with them (the larger diagonal block contains two less apparent, sub-blocks). In 1(b) however, all 2D projections except t-SNE produce a single cluster (when plotted without colors), while t-SNE seems most successful in separating the three subsets (arguably, t-SNE shows c=2 clusters when colors are omitted). The iVAT image suggests c=2, conforming to the apparent (uncolored) pair of t-SNE clusters. The low value of DI is a warning that there is not much separation between these three subsets of waveforms.

| PCA | PV-E | Wavelet | t-SNE | Sammon | Waveforms | iVAT |
|-----|------|---------|-------|--------|-----------|------|
| ![PCA Image](image1.png) | ![PV-E Image](image2.png) | ![Wavelet Image](image3.png) | ![t-SNE Image](image4.png) | ![Sammon Image](image5.png) | ![Waveforms Image](image6.png) | ![iVAT Image](image7.png) |

(a) DI=0.168

| PCA | PV-E | Wavelet | t-SNE | Sammon | Waveforms | iVAT |
|-----|------|---------|-------|--------|-----------|------|
| ![PCA Image](image8.png) | ![PV-E Image](image9.png) | ![Wavelet Image](image10.png) | ![t-SNE Image](image11.png) | ![Sammon Image](image12.png) | ![Waveforms Image](image13.png) | ![iVAT Image](image14.png) |

(b) DI=0.068

**Figure 1:** Two different simulations at c=3

Figures 2 to 5 show representative mixtures of c=5, 10, 15 and 20 component mixtures. These examples, and many others not reported here, show that iVAT and t-SNE usually provide useful visual estimates of the number of clusters up to around c=15, but the other methods almost always fail with c = 4 or more subsets. We had 5 cases for each number of subsets (e.g. 5 different cases of mixture at c=10, etc.) and overall t-SNE provided the most consistent estimate of the presumptive numbers of mixture components. There were some cases for which iVAT failed to display the expected number of dark blocks in mixtures having fewer than 10 components. The block structure in some of the reproduced iVAT images is more apparent at higher resolutions than shown here. The block structure in some of the reproduced iVAT images is more apparent at higher resolutions than shown here. Our experiments suggest that iVAT is somewhat sensitive to noise in the waveforms, which often manifests itself as a falloff in intensity towards one end of the diagonal. See Figure 2 for an example.
Figure 2: Simulation at c=5. The iVAT image displays 4 clear blocks and some disconnected data due to noise in the lower right: the only projection that clearly shows c=5 is t-SNE.

(a) DI=0.158

Figure 3: Simulation at c=10. The iVAT image displays 10 blocks; the projection that clearly shows c=10 is t-SNE.

(a) DI=0.121

Figure 4: Simulation at c=15. The iVAT image displays 9 blocks and t-SNE shows 13 (colored), and 9 or 10 in black.

(a) DI=0.084

Figure 5: Simulation at c=20. The iVAT image displays 13 blocks and t-SNE shows 12
The next example in this section highlights the ability of iVAT to address two additional problems encountered in spike sorting, namely, anomaly detection and multistage clustering (aka “re-iteration” amongst subclusters). Figure 6(a) is a set called Z of n=4665 waveform vectors comprising a mixture of c=10 labeled subsets from simulated dataset-1. The 10 waveforms shown in Figure 6(b) are the average waveforms, \( \bar{Z}_{10} = \{ \bar{z}_i : 1 \leq i \leq 10 \} \), of the ten labeled subsets.

Visual inspection of Figure 6(b) suggests that \( \bar{z}_4 \), the average waveform of the 488 spikes for unit 4, here called Z_4, is an outlier (an anomaly) to the general shape of the other 9 graphs. This (easily seen) visual evidence suggests that Z_4 may form an anomalous cluster in the input or projection spaces. But this observation does not justify removal of all 488 unit 4 spikes from the input data. However, the iVAT image of Z_4 will corroborate our suspicion that Z_4 is an anomalous cluster in Z.

Figure 6(c) is the iVAT image of \( \bar{Z}_{10} \), and Figure 6(d) is the dendrogram of the clusters produced by extracting the single linkage hierarchy of clusters from the vectors in \( \bar{Z}_{10} \). The integers along the borders of the iVAT image of \( \bar{Z}_{10} \) show the identity of each pixel after iVAT reordering. The visualization in 6(c) is quite informative: it not only isolates \( \bar{z}_4 \) as an outlier (the single pixel at the lower right corner of the image), but it also depicts the other 9 graphs as members of a second large cluster. Moreover, this image suggests a hierarchical substructure within the 9x9 block. The intensities of \{5,7\} and \{6,10\} suggest that these pairs of subsets are closely related. The \{3,9\} block is next in intensity, followed by the 5x5 grouping of \{8, 5, 7, 9, 3\}, which are then coupled to \{6,10\}, and then this whole structure is embedded within the 9x9 block which includes \{1,2\}. We remark that the SL hierarchy is easily extracted by applying a back pass that cuts edges in the iVAT MST that reordered \( \bar{Z}_{10} \) (cf. Kumar et al. [2016]). Figures 6(c) and 6(d) make the relationship between iVAT and single linkage quite transparent. And Figure 6(c) illustrates how an iVAT image can suggest multicluster substructure in a data set.

Figures 6(e) and 6(f) are scatterplots of t-SNE projections of \( \bar{Z}_{10} \) corresponding to perplexity settings of 2 and 3. Both views show the labels of the 10 mean profiles, and both views seem to indicate that \( \bar{z}_4 \) is an outlier in the set \( \bar{Z}_{10} \). We show these two projections to emphasize that every run of t-SNE with different settings of its hyperparameters may produce different visualizations of its input data. On the other hand, the iVAT image is uniquely determined up to a choice of the distance measure used to construct D.

Figure 7(a) is the iVAT image of the data set Z shown in Figure 6(a). Comparing Figures 6(c) to 6(f) shows that iVAT very clearly suggests the same coarse cluster structure (c=2) in all of the upspace data that it sees in \( \bar{Z}_{10} \), the set of mean profiles. Neither image suggests that c=10; instead, both suggest that the best interpretation of the input data or its means is to first isolate the unit 4 waveform(s), and then regard the remaining spikes as a new cluster, which becomes a candidate for multistage clustering (reclustering, or re-iteration per Niediek et al. [2016]). Note that iVAT makes this information available whether the data are labeled or not.
Finally, Figures 7(b) and 7(c) are labeled and unlabeled t-SNE scatter plots of Z. Both views suggest that Z contains 5 clusters. Subset Z₄ is isolated in view 7(b), but not more isolated than subset Z₂, so t-SNE is less assertive about the anomalous nature of Z₄ than iVAT is. If the labels are available, reclustering might be applied to 9, 7, 5, 8 and/or 3,6,10 to make a finer distinction between spike subsets. If the labels are unavailable, it’s hard to see what can be inferred from the t-SNE projection about Z beyond the suggestion provided by view 7(c) that we should seek 5 clusters in Z.

We conclude this example with some general observations. First, the iVAT image is unique, while t-SNE plots are a function of three user-defined parameters. Second, single linkage clusters of the input data are available via clusiVAT [Kumar et al., 2016] once an iVAT image is built. Third, while Z has 10 labeled subsets of input spikes, neither iVAT nor t-SNE makes this prediction. This emphasizes the fact that labeled subsets may not necessarily be clusters with respect to any computational scheme designed to detect clusters.
Figure 6: iVAT and t-SNE visualizations of average waveforms of a mixture of 10 subsets of labeled simulated spikes
Figure 7: iVAT and t-SNE visualizations of a mixture of 10 subsets of labeled simulated spikes
We turn into dataset-2 to further investigate the limits of discernible spike subsets since as
mentioned previously, sets driven from dataset-2 are combinations of real spikes originated
from pyramidal cells in rat hippocampus (ref to [Henze et al., 2000]). We extracted the spike
subsets of nine individual neurons obtained from the different experimental trials. From
these nine subsets, we built 36 mixtures at c=2; 84 mixtures at c=3; 126 mixtures at c=4;
126 mixtures at c=5; 84 mixtures at c=6; 36 mixtures at c=7; 9 mixtures at c=8; and one
mixture of all nine subsets (c=1). This yields a total of 502 mixtures of labeled waveforms.

For the sake of brevity, we showcase four representative units and the various mixtures that
can be built from them at c=2, c=3, and c=4. Figure 8 shows the four representative subsets
(all nine subsets of the waveforms are shown in Figure 15).

Figure 8: The subsets of spikes generated by four representative units: X1, X2, X3, and
X4 containing 1173, 700, 779, and 382 spikes, respectively. Note that the waveforms in X4
are visually different than most of the waveforms in the other three subsets. This fuels an
expectation that mixtures with X4 as one component will be somewhat more separable than
mixtures without it.

Figure 9 shows all six views of pairs (Xi, Xj) made with 2D transformations of the 80D
(upspace, p=80) datasets for the mixtures of two representative single units. We will name
the mixtures (Xk, Xj)=Xkj and will follow this convention for all mixtures. For example,
the mixture of X1 and X2 is X12, and the mixture of X1, X2, and X4 becomes X124. The
waveforms comprising each mixture are also shown, with the average waveform for each
single unit in thick black. The colors of points in the 2D scatterplots correspond to class
labels of the mixed data. It is important to remember that in a real application, the data
are not labeled, so the associated 2D scatterplots will be mono-color dots in the plane. The
mixtures are ordered according to increasing values Dunn’s index. Observe that for each
mixture, different 2D projections may offer different interpretations of the cluster structure
in the upspace data. In 9(a), all five projections show one big cluster, far more evident if
the color labeling is missing, which is the case for real experiments in which we do not know
the membership of the waveforms. In cases like this, since the clusters are projected densely
side by side, human operators or algorithms tend to select only the core of the clusters. This
usually produces better values for cluster validity indices, but at the expense of unwarranted
confidence in subsequent analyses.
Figure 9: Mixture pairs of X1, X2, X3 and X4, ordered by increasing Dunn’s index

First, some general observations. Figures 9(a), 9(b), and 9(c) all have a DI value of around 0.09. This is a relatively low value that indicates a lack of separation between the two
components of the mixture. The iVAT images for these three cases are basically uniform (no
strongly visible dark blocks), which indicates that the upspace data are not well separated.
Separation emerges in Figures 9(d), 9(e), and 9(f), the three cases that have X4 as one
component. Dunn’s index is essentially doubled (0.18 up to 0.23), so upspace separation of
the pair of clusters has increased. The most visible separation is seen in the t-SNE downspace
scatterplot, which is mirrored in the upspace iVAT images: the strong dark block corresponds
to subset X4. Now we will discuss the six cases in more detail.

In 9(b), PCA, Wavelet, t-SNE, and Sammon show two clusters, while the PV-E plot shows
just one. In 9(c), all the projections except t-SNE point to one cluster; and in the other
images, which include X4, a less distorted and noisy set of spikes, all the projections do a
good job of mapping the clusters in a separable manner (for the wavelet projection in 9(e),
it is hard to see two clusters when there are no color labels). The iVAT image also follows
the same trend: the clarity of the two blocks generally becomes higher with a higher Dunn’s
index.

The Peak to Valley and Energy (PV-E), are the only real (physically meaningful) 2D features.
All the other 2D projections are dimensionless, i.e., they do not have physical meaning. It is
important to emphasize that neither the 2D projections nor iVAT produce clusters, all these
visual methods just suggest how many to look for.

The projections and the iVAT image of X23 with DI = 0.091 (Figure 9(b)) is a bit more
separable and clear than the mixture of X13 with DI = 0.097 (Figure 9(c)). Both values are
relatively small, and the difference between these two values (0.008) is negligible, indicat-
ing that these two cases are somewhat indistinguishable. The iVAT image for X14 clearly
suggests the c=2 at a Dunn Index of 0.228. This provides a much stronger indication of
reliability than the smaller DI values. Indeed, Dunn characterized a partition as being com-
 pact and separated if and only if DI > 1. DI values less than about 0.5 usually characterize
relatively poor cluster structure.

All the cases of mixtures of three subsets are portrayed in Figure 10, again ordered by
their Dunn’s index, which is quite low and nearly equal in all four views. The numerator
of DI is the minimum distance between any pair of subsets, and the denominator is the
largest distance between points in some clusters, so it is dominated by the smallest between-
subset distance and largest in-subset distance. Consequently, DI fails to recognize competing
clusters that cannot dominate either of the two factors in Dunn’s formulation. These non-
dominant clusters can often be seen in iVAT imagery. For example, in Figure 10(b), the
small yellow cluster seen in the t-SNE scatterplot of X124 appears as the small dark block in
the lower right corner of the corresponding iVAT image. In Figure 10(a) all the projections
except for t-SNE fail to point to c=3 and the iVAT image is not informative either. In Figure
10(c) for X234, the PV-E and Wavelet projections suggest that c=2, while PCA, t-SNE and
Sammon point to c=3. The t-SNE features provide the widest and most visible separation
between the three clusters. The iVAT image of X234 is weakly suggestive of c=3. Figure
10(d) for X134 provides a striking contrast in the ability of the visualization methods to
correctly portray the presumed structure in the data. PV-E and Wavelet suggest $c=1$, PCA
and Sammon imply $c=2$, and t-SNE points clearly to $c=3$. The iVAT image is pointing to
$c=2$, at a relatively low value of $DI=0.097$.

| PCA     | PV-E   | Wavelet | t-SNE  | Sammon | Waveforms | iVAT |
|---------|--------|---------|--------|--------|-----------|------|
| ![Plot](a) X123, $DI=0.074$ | ![Plot](b) X124, $DI=0.090$ | ![Plot](c) X234, $DI=0.091$ | ![Plot](d) X134, $DI=0.097$ |

**Figure 10:** Three-subset mixtures of X1, X2, X3 and X4 at $c=3$ ordered by increasing values of Dunn’s index

Finally, a similar trend continues in the $c=4$ subset mixture of X1, X2, X3, and X4. The
PV-E and Wavelet features indicate only one big cluster, and PCA, Sammon, and the iVAT
image single out X4 while packing the other three sets of waveforms into a single cluster,
whereas, t-SNE maps the four subsets with arguably enough clarity to declare that X1234
has four clusters. It can be argued that while the input has $c=4$ labelled subsets, the primary
visual evidence does not support $c=4$, nor will there be a “best” set of clusters in the upspace
at this value of $c$. In other words, just because the subsets have 4 labels does not guarantee
that a cluster analysis of the data will agree. When you imagine the scatterplots in Figure
11 without colors there are not four distinguishable clusters present.
In summary, we note again that although the subsets of spikes were obtained from independent trials, they were all induced by intracellular current injections to hippocampal pyramidal cells and recorded from their close proximity in extracellular medium. This to some degree explains the similar average waveforms and high variability between spikes of one subset. We observed that, for example, from the thirty-six mixtures of two subsets that can be created from the nine spike sets, only 55% (20 cases) were mapped using t-SNE as well isolated clusters. Overall, visualizations of mixtures using any of the other projection methods and iVAT did not suggest discernible clusters. This points out the challenge in identifying neurons of the same class (e.g., pyramidal) from their spike waveforms, at least when they are induced by current injections.

3.2 Clustering and partition quality evaluation

3.2.1 Clustering in the input space and 2D spaces using c-means

So far, we have shown that the two-dimensional representations in our study may give different interpretations of the upspace data. This problem is highly dependent on the definition of similarity between spike waveforms of different units. Overall, iVAT and t-SNE were most helpful in assessing the pre-cluster presumptive structure of the waveform mixtures. In order to provide a more quantitative assessment of the effectiveness of the different low-dimensional representations in processing spike waveforms, we ran the c-means clustering algorithm on each of the 95 mixtures from dataset-1 and the 502 mixtures from dataset-2 (Recall that many authors refer to c-means as "k-means," where k is the notation chosen for the number of clusters: either notation is correct). We prefer to reserve (k) for the universally accepted description of the k Nearest Neighbor rule.

Dunn’s index and its generalizations provide measures of the intrinsic quality of the computed clusters (based on their distribution with respect to each other). Figure 12 shows the average Dunn’s index (DI) and generalized Dunn’s index (GDI\(_{33}\)) of the mixtures for the two datasets.
(a) The average(±SD) DI and GDI\(_{33}\) for the 95 mixtures in dataset-1

(b) The average(±SD) DI and GDI\(_{33}\) for the 502 mixtures in dataset-2

**Figure 12:** The average(±SD) Dunn’s and generalized Dunn’s indices of ground truth partitions for the mixtures in the two datasets

The two indices have the same trend: they decrease almost monotonically as the number of components (c) increases. However, the generalized version, GDI\(_{33}\), provides a much clearer idea of the trend than DI because it has higher values that reflect separation more clearly, and it avoids the bias of inliers and outliers that may affect Dunn’s index. On the other hand, Figure 12 also suggests that both indices tend to favor lower numbers of clusters. This is a different type of empirically observed bias that must be accounted for when relying on cluster validity indices. See Lei et al. [2017] for a discussion related to this point.

The DI and GDI\(_{33}\), as internal measures, were used to give a sense of the structure inherent in ground truth partitions of the data in the upspace. Then, to evaluate candidate partitions produced by c-means in the upspace and downspace data sets, we used the adjusted rand index (ARI), which compares the cluster structure of each c-means partition to its ground truth partner at every value of c.

The c-means clustering algorithm is executed on each low-dimensional representation ob-
tained with the five methods. The number of clusters to be generated (i.e., c) is set equal
to the number of labeled subsets in the ground truth partition (i.e., 2, 3, 4,...to 9 for cases
in dataset-2 and 3, 4, 5, ....to 21 for cases in dataset-1). For each computed partition, we
calculate the measure of agreement with ARI between the computed waveform memberships
and the memberships as given by the ground truth partition (recall that our data is labeled).

Figures 13(a) and 13(b) report the average ARI for mixtures in dataset-2 and dataset-1,
respectively. In each figure, the first column is the ARI of the clusters achieved by running
c-means on the input dimension space (the 48D waveforms for Dataset-1 and the 80D spike
waveforms for Dataset-2). The next columns show the average ARIs calculated for the
clusters achieved by c-means clustering on the 2D datasets produced by the five techniques.
The ARI maximizes at 1, so clustering in the 2D t-SNE downspace data provides c-means
clusters that, on average, slightly better match the ground truth partitions than c-means
clusters in the input space.

In order to highlight the importance of dimensionality reduction and feature extraction tech-
niques (the pre-clustering stage), this subsection presented a comparison between clustering
in the different downspaces and also the input space, using the same clustering algorithm
in all spaces. It is important, however, to recognize that the choice of clustering algorithm
also contributes to the accuracy of membership assignments. Given the good visualization
in t-SNE projections, the next subsection will investigate whether this better pre-clustering
technique improves the overall performance more than the difference in performance gained
by a change of the algorithm. In other words, how does clustering done on t-SNE projection
data compare to partitions produced by a different clustering algorithm on the input space?
(a) For dataset-1: The average ARIs of clusters obtained by c-means for the simulations of mixtures at each c value in the 48D upspace and the 5 2D-spaces

(b) For dataset-2: The average ARI for the clusters obtained by c-means for mixtures at each c value in the 80D upspace and the 5 2D-spaces

Figure 13: Average validity index (Adjusted Rand index) of the clusters obtained by c-means on the two datasets
3.2.2 Comparing c-means on t-SNE data with Osort clustering

In this section we compare the performance of c-means on t-SNE projections of the input data, which achieved the best overall ARI ranking among the 6 methods discussed in Section 3.2.1, to partitions on the input data obtained with a well-known spike sorting algorithm called Osort.

Osort is a greedy algorithm that clusters data in the upspace (input space). The idea is that the limit in discerning two waveforms from each other is the noise in the data. Therefore, the algorithm compares the squared Euclidean distances between spikes, with the square of the standard deviation of the raw signal as a measure of noise. If the distance is lower than noise, the two waveforms are paired in the same cluster; if the distance is higher than the noise, Osort creates a new cluster. At termination of Osort, a manual curation strategy is used to merge clusters or label spikes as anomalies. Although the results of Osort can be improved by this manual step, we wanted a fair comparison between the clusters created by Osort and c-means, and thus did not perform manual curation for Osort. The automated portion of the Osort algorithm retains clusters containing more than 50 spikes and merges the rest into one cluster.

Since the Osort algorithm requires the raw signal to measure the noise, we could only use dataset-1 for the experiments in this section. In dataset-2 the mixture of different units is made from extracted spikes from different raw traces, while in the simulation, i.e., dataset-1, a raw trace consisting of background noise is also simulated for each mixture.

For the 95 mixtures in dataset-1 (5 at each c value), the t-SNE features were processed by c-means. For the same simulations, using the noise information from the raw trace, the Osort algorithm was also performed. The average ARI of the partitions obtained by the two methods is shown in Figure 14. Osort yields a smaller average ARI, indicating poorer matching to the ground truth partitions than c-means, in all cases except c=5. The clustering performance of Osort decreases dramatically as the number of spike subsets in the mixture increases. In contrast, c-means partitions of the t-SNE data are relatively constant and have somewhat higher ARI values, especially for higher values of c.

This provides further evidence that the space in which the clustering is performed plays an important role in the quality of the unit sorting. Figure 13 shows that clustering with c-means in the 2D t-SNE data works fairly well with both real and simulated data. Figure 13 also shows that c-means clustering in the input space is second best amongst the methods tested, and we think that upspace clustering should be a default option in all cases. We remind readers that other clustering algorithms might yield different results, and that almost every clustering algorithm will deliver clusters at any value of c.
Figure 14: The average clustering performance of Osort and c-means on t-SNE data. c-means on t-SNE provides a consistently high ARI even in mixtures with subsets of more than ten subsets.
Current sorting algorithms used for individual electrode recordings are limited in distinguishing more than a few neurons in mixtures of spikes obtained from each electrode [Niediek et al., 2016, Pedreira et al., 2012, Rutishauser et al., 2006]. Hence, what we consider to be the spike train of a single unit may, in fact, be the spike train of multiple units. This does not negate the usefulness of the findings which apply the results of these sorting algorithms to infer neural coding and brain function. Most of such research is performed based on the rate coding principle [Dayan and Abbott, 2001], which uses the spike rate of the sorted units to model the neuronal response. Rate coding models neglect sorting errors. As long as the spike rate changes according to the stimulus, the model will capture the response, whether the spike train consists of spikes of one or multiple neurons.

Nevertheless, there have been studies concerned with issues arising from sorting quality on the results of rate coding models. For example, Todorova et al. [2014] evaluated the quality of the off-line reconstruction of arm trajectories from electrode array recordings and showed that discarding spikes substantially degrades the decoding of the movement to the extent that decoding the unsorted recordings reached higher performance results. They also showed that adding the tuning model (temporal features) of the spiking to the sorting process does not always improve the sorting based on waveform features. We can use the analogy of a verbal fight or discussion among a few people. An observer can tell if the discussion is going smoothly or if it is heated based on the overall volume of the voices of the group, even if the words uttered by individuals is not discernible. This is why rate coding models are popular and successful in certain respects, but they cannot elucidate how neurons interact to give rise to brain functions [Akam and Kullmann, 2014, Huxter et al., 2003, Mehta et al., 2002, Rullen and Thorpe, 2001, Zuo et al., 2015]. We need to improve unit sorting if we want to model the temporal coding of neurons reliably.

In high-dimensional data, the role of visualization in gaining knowledge of the data structure is critical. There is no doubt, as in Plato’s allegory of the cave, that there is always a loss or distortion of structural information in any transformation from the upspace (aka: input space or input dimension) to any downspace. We investigated this issue using iVAT, a tool that enables direct visualization of cluster structure in the upspace as well as five dimensionality reduction methods. We showed that better sorting can be achieved by securing a visual assessment prior to clustering which affords an estimate of the cluster structure of the data (i.e., the number of clusters, c), or at least a small interval of integers that presumably bracket the true (meaning most distinguishable by some clustering algorithm) but unknown number of clusters. Our examples show that t-SNE is one of the best methods for projection of high dimensional data to the viewing plane. We note that t-SNE for the present analysis was parameterized with a perplexity of 30 and learning rate of 500. This was the empirically optimized setting for our data and we acknowledge that the need for parametrizing based on the data is a downside to using t-SNE to provide projected data for clustering.

In the first dataset, simulations were generated using average waveforms obtained from ex-
tracellular recordings in behavioral experiments. For the mixtures of spike subsets extracted from this dataset it was possible to estimate the presumptive cluster number in the data from the dark blocks in the iVAT images, even in some cases of mixtures of twelve subsets. Mixtures of higher subsets were sometimes displayed as compact and isolated clusters in the t-SNE projections. Our experiments confirm that when the data possess compact, well-separated clusters, visualization can be quite useful. We believe that dataset-1 represents mixtures of spike sets that are generated by different cell types, brain regions and brain states, and these can be distinguished based on their spike waveforms. In contrast, dataset-2 represents mixtures of spike sets that are induced from cells of the same class receiving intracellular current injections, hence providing spikes with similar waveforms. Therefore, classifying based on extracellular waveforms alone may not be feasible in the latter case (cells of the same type receiving the same input). It should be noted that in sorting of spikes for each electrode, different distances of the units from the electrode improves sorting, since the amplitude (energy) of the waveforms is different. However, these results indicate that a further subtype classification beyond the two main classes of inhibitory and pyramidal categories (i.e. subtypes of pyramidal cells) may not be feasible by considering only the spike waveforms.

Overall, from the visualizations of the data, we infer that there is a common waveform shape among many cells suggested by the existence of a large central region populated by most of the clusters. However, there are cells that have clearly distinct signatures. The individual clusters exhibit variability: for some neurons the spike waveforms define homogeneous and compact clusters, while others are elongated clusters in the nonlinear space. This suggests that the relation between waveform samples for different neurons is different (or the way the waveform samples interrelate is different in different neurons’ spikes).

There is an inherent variability in the waveforms imposed by the variations in the intracellular action potentials due to factors such as bursting [Henze et al., 2000]. The activity of neighboring neurons (that may sometimes overlap) and the background field potential also contribute to the variability in the waveforms [Buzsáki et al., 2012]. Slight electrode drift over the course of the experiment is also a possible source of variation in the waveform shape [Harris et al., 2016]. The extracellularly recorded potentials are already distorted signatures of intracellular actions potentials, which makes the dimensionality reduction stage even more critical. The problem of crowding in PCA maps is also well known. Analysis of the simultaneous extracellular and intracellular recordings shows that the probability distributions of spikes from different neurons in the PCA feature space have some degree of overlap [Harris et al., 2000]. Since the cluster visualization in PCA or other feature spaces usually exhibits overlapped or mixed clusters, it is common to leave a lot of non-outlier waveforms unsorted to get well-separated clusters. Our results show that the 2D t-SNE projection is the most reliable feature extraction scheme we tested. We believe that t-SNE works well since it is a probabilistic-based approach that is appropriate for neuronal data. In a nutshell, the variability caused by the noisy spikes can often be circumvented by converting the deterministic dissimilarity
measure between two waveforms into a probability of dissimilarities.

In this paper, we also demonstrated that the visual assessment of \( c \) from the iVAT images is often possible, highlighting that if clustering in the upspace is preferred, a visualization tool such as iVAT can be integrated into the package to inform the manual curation process. However, there were cases, in particular when \( c \) was high (> 10), that the iVAT image could not clearly indicate the number of clusters. In such cases relying on the visual assessment of a user makes the estimate of \( c \) subjective. We showed that extracellular neuronal waveforms generate noisy datasets that at times do not comprise well-separated clusters. So, the iVAT does not provide the definitive answer to the problem of spike sorting. Nevertheless, it provides insight into the coarse structure of the dataset. Moreover, we mentioned the relationship between iVAT and the single linkage clustering algorithm that is illustrated in Figures 6(c) and 6(d) (See Havens et al. [2009] and Mahallati et al. [2018] for further discussion). The majority of the edges in the MST that iVAT builds connect neighbor points and hence have very small values. The largest values in the MST usually correspond to edges that connect clusters (and the outlier points). The threshold between the small and large values reflects finer distinctions between clusters in the upspace. This threshold can be used in assigning spikes to clusters, in particular when the clustering is performed in the upspace (e.g. Osort).

Another reason why having a reliable dimensionality reduction stage is important is revealed by our results on Dunn’s index, which showed that, DI, in common with many other internal cluster validity indices, is usually monotonic in \( c \). So, the common practice of running the clustering algorithm for several values of \( c \) and then choosing the best partition based on the optimal value of any cluster validity index may not be very effective. Moreover, by computing both DI and GDI\textsubscript{33} for the same data, we demonstrated that there is no agreement about a generic CVI, a fact that has been shown before in previous experiments on internal CVIs [Vendramin et al., 2010]. Indeed, in the real (unlabeled data) case, it is wise to compute a number of different internal CVIs, with a view towards ensemble aggregation of the results. To appreciate the disparity that different CVIs can cause, see Arbelaitz et al. [2013] for an extensive survey of 30 internal CVIs tested on 20 real data sets. See Vega-Pons and Ruiz-Shulcloper [2011] for a survey of ensemble approaches to clustering. Fournier et al. [2016] have applied this method to aggregation of partitions obtained by different clustering methods used for sorting spike waveforms. Here we suggest using an ensemble approach on the votes cast by different internal cluster validity indices - DI and its 18 GDIs are just a few of the ones available in Arbelaitz et al. [2013] - for each partition in CP. We think this approach will greatly improve the final interpretation of structure in unlabeled data. This will be the objective of our next foray into spike sorting clustering algorithms.
Figure 15: The subsets of spikes of the 9 individual neurons used in the study. Each subplot title displays the label of the experiment in [Henze et al., 2009] dataset and the number of spikes in each subset: e.g., #745 means there are 745 waveforms in the sample.
A.2 iVAT algorithm

| Algorithm: VAT |
|----------------|
| **In** | $D$, an $n \times n$ matrix of dissimilarities: $D = D^T$; $d_{ij} \geq 0 \forall i, j$; $d_{ii} = 0 \forall i$ |
| **Set** | $K = \{1, 2, \ldots, n\}$; $I = J = \emptyset$ |
| | Select $(i, j) \in \arg\max\{D_{st} : s \in K, t \in K\}$ |
| | $P(1) = i$; $I = \{i\}$; $j = K - \{i\}$ |
| | Initialize MST at either end of edge with largest weight in $D$ |
| **For** | $m = 2, \ldots, n$ do: select $(i, j) \in \arg\min\{D_{st} : s \in I, t \in J\}$ |
| | Select $(i, j) \in \arg\min\{D_{st} : s \in I, t \in J\}$ |
| | $P(m) = j$; $I = I \cup \{j\}$; $J = J - \{i\}$; $d_{m-1} = d_{ij}$ |
| **For** | $1 \leq i, j \leq n$ do: |
| | $[D^*]_{ij} = [D]_{P(i)P(j)}$ |
| **Out** | VAT reordered dissimilarities $D^*$: arrays $P, d$ |
| **% Create VAT RDI $I(D^*)$ using $D^*$** |
| Algorithm: iVAT |
| **In** | $D^*$ = VAT reordered dissimilarity matrix: $D^* = [0]$ |
| **For** | $k = 2$ to $n$ do: |
| | $j = \arg\min\{D^*_r : r = 1, \ldots, k-1\}$ |
| | $D^*_{kc} = D^*_{k} ; c = j$ |
| | $D^*_{kc} = \max\{D^*_{kj} : D^*_{jc} \}; c = 1, \ldots, k-1; c \neq j$ |
| **For** | $j = 2, \ldots, n; i \neq j$ do: |
| | $D^*_{ji} = D^*_{ij}$ |
| **Out** | iVAT reordered dissimilarities $D^*$ |
| **% Create iVAT RDI $I(D^*)$ using $D^*$** |

A.3 Cluster Validity Indices

A.3.1 Dunn’s Index

Let $X_i$ and $X_j$ be non empty subsets of $\mathbb{R}^p$, and let $d : \mathbb{R}^p \times \mathbb{R}^p \mapsto \mathbb{R}^+$ be any metric on $\mathbb{R}^p \times \mathbb{R}^p$. Define the diameter $\Delta$ of $X_k$ and the set distance $\delta$ between $X_i$ and $X_j$ as:

$$
\Delta(X_k|d) = \max_{x, y \in X_k} \{d(x, y)\},
$$

$$
\delta(X_i, X_j|d) = \min_{x \in X_i, y \in X_j} \{d(x, y)\} = \delta_{SL}(X_i, X_j|d).
$$
Then for any partition $U \leftrightarrow X = X_1 \cup \ldots \cup X_i \cup \ldots X_c$, Dunn’s separation index of $U$ is:

$$DI(U|d) = \frac{\min_{1 \leq i \leq c} \{ \min_{1 \leq j \neq i \leq c} \{ \delta(X_i, X_j|d) \} \}}{\max_{1 \leq k \leq c} \{ \Delta(X_k|d) \}}$$  \hfill (5)

### A.3.2 Generalized Dunn’s Index

$$\Delta_3(X_k|d) = 2 \left( \frac{\sum_{x \in X_k} d(x, \bar{v}_k)}{|X_k|} \right)$$  \hfill (6)

$$\delta_3(X_i, X_j|d) = \frac{1}{|X_i||X_j|} \sum_{\substack{x \in X_i \\ y \in X_j}} d(x, y)$$  \hfill (7)

$$GDI_{33}(U|d) = \frac{\min_{1 \leq i \leq c} \{ \min_{1 \leq j \neq i \leq c} \{ \delta_3(X_i, X_j|d) \} \}}{\max_{1 \leq k \leq c} \{ \Delta_3(X_k|d) \}}$$  \hfill (8)

where $\bar{v}_k = \frac{\sum_{x \in X_k} x}{|X_k|}$ is the mean or centroid of the cluster. The notation $|d$ in equations 3, 4, 5, 6, and 7 for $\Delta$ and $\delta$ indicate that these formulas are valid for any metric $d$ on the input space.

### A.3.3 Adjusted Rand Index

Let $V \in M_{hrn}$ be the crisp partition of the $n$ objects possessing $r$ clusters, according to ground truth labels. Let $U \in M_{hcn}$ be any crisp partition of $n$ objects with the $c$ clusters generated by any clustering algorithm. Note that $r$ does not necessarily equal $c$. The ARI is a measure of similarity between $U$ and $V$, computed as:

$$ARI(U|V) = \frac{2(ae - bc)}{(a + b)(e + b) + (a + c)(e + c)}$$  \hfill (9)

where,

- $a =$ Number of pairs of data objects belonging to the same subset in $U$ and $V$.
- $b =$ Number of pairs belonging to the same subset in $V$ but to different subsets in $U$.
- $c =$ Number of pairs belonging to the same subset in $U$ but to different subsets in $V$. 

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e = Number of pairs not in the same subset in V nor the same subset in U.

Hubert and Arabie [1985] developed this correction to eliminate bias due to chance from Rand’s index.
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