Heterogeneity in Neuronal Calcium Spike Trains based on Empirical Distance

Sathish Ande∗1, Jayanth R Regatti 1, Neha Pandey 1, Ajith Karunarathne 2, Lopamudra Giri 1, Soumya Jana 1
1Indian Institute of Technology Hyderabad, Telangana, India 2The University of Toledo, Ohio, USA
∗Email: ee15resch02003@iith.ac.in

Abstract—Statistical similarities between neuronal spike trains could reveal significant information on complex underlying processing. In general, the similarity between synchronous spike trains is somewhat easy to identify. However, the similar patterns also potentially appear in an asynchronous manner. However, existing methods for their identification tend to converge slowly, and cannot be applied to short sequences. In response, we propose Hellinger distance measure based on empirical probabilities, which we show to be as accurate as existing techniques, yet faster to converge for synthetic as well as experimental spike trains. Further, we cluster pairs of neuronal spike trains based on statistical similarities and found two non-overlapping classes, which could indicate functional similarities in neurons. Significantly, our technique detected functional heterogeneity in pairs of neuronal responses with the same performance as existing techniques, while exhibiting faster convergence. We expect the proposed method to facilitate large-scale studies of functional clustering, especially involving short sequences, which would in turn identify signatures of various diseases in terms of clustering patterns.

Index Terms—Calcium imaging; Neuronal spike trains; Empirical probability; Statistical dissimilarity; Heterogeneity.

I. INTRODUCTION

Neurons encode stimulus information in spike trains. In fact, heterogeneity in spike trains is a known manifestation of complex information processing, which enables diverse functions in the hippocampus, a brain region associated with memory and learning [1]. The said heterogeneity in spike trains has been investigated by clustering neuron pairs based on certain statistical similarities. An early attempt in this direction was based on a correlation-based similarity measure [2]. However, such a measure captures coincident firing, i.e., synchronicity in spike trains, but ignores time-delayed versions of similar patterns which are known to arise in complex neuronal networks. As a remedy, distance measures based on Lempel-Ziv (LZ) encoding have been suggested to identify the statistical similarities in synchronous or asynchronous spike trains [3], [4]. One such method was based on LZ-78 algorithm which needs long sequences for reliable performance. In the quest for a method that can be applied to short sequences, we consider LZ-76, a LZ-based fast method, but find it to be inaccurate. Against this backdrop, we propose a Hellinger distance measure based on empirical probabilities of patterns in each pair of spike trains [5]. Our method converges faster than LZ-78, and hence may be used on short sequences, while being comparably accurate. Further, we cluster pairs of neuronal spike trains and found two non-overlapping classes, and the clusters obtained using the proposed distance measure and the distance based on LZ-78 are found to behave similarly. This demonstrates the suitability of the proposed method as a fast-converging alternative to the existing slow technique.

The rest of this paper is organized as follows. Section II describes calcium imaging of hippocampal neurons and spike train inference, and introduces the notion of LZ distance and the proposed empirical distance measure. Further, Section III presents the results demonstrating suitability of the proposed method. Finally, Section IV concludes the paper.

II. MATERIALS AND METHODS

The workflow of the paper is schematically depicted in Fig. 2, and elaborated in the following.

A. Data Collection and Spike Train Inference

We performed time-lapse confocal imaging (using a Leica DMi6000B inverted microscope fitted with a Yokogawa CSU-X1 spinning-disk unit) on hippocampal neurons, cultured from 1 day postnatal Sprague-Dawley rats. In particular, we monitored intracellular calcium at 7-th day after plating using excitation at 488 nm and emission at 510 nm.
During imaging, neurons were kept in the attached incubation chamber maintained at 37°C and 5% CO2. The interval between successive images, while set at 1 s, was observed to vary between 0.8 s to 1 s. From the time-lapse image data (see Fig. 1 for a representative frame), the time course of spatially resolved Fluo-4 fluorescence intensity in neuron populations was obtained using Andor software. At present, out of a population of 28 neurons, we consider 8 neurons, indexed 1–8, for analysis (time course of calcium responses are shown in Fig. 3 for neurons 1–4 and heterogeneity in such responses is visually evident here). For each neuron, we inferred binary spike train from its time course using suitable normalization and a fast nonnegative deconvolution algorithm [5], [7]. Such spike sequences were used for further analysis.

B. Existing LZ-based Dissimilarity Measures

Versions (LZ-78, LZ-77, LZ-76) of LZ encoding scheme are based on suitable dictionaries that convert a given sequence \( X^n = (X_1, X_2, \ldots, X_n) \) into non-overlapping phrases [5], [8]. For instance, \( X^n = '001100110100111' \) is parsed as

\[
\text{LZ-78: 0-01-1-00-10-101-001-11} \\
\text{LZ-77: 0-01-0-10-010-101-010-00100-10011} - 111-11111-1111-111 \\
\text{LZ-76: 0-01-10-010100111} - 1-111-11111-1111-111.
\]

The complexity of a sequence \( X^n \) is defined as

\[
K(X^n) = \frac{c(X^n) \log(c(X^n))}{n},
\]

where \( c(X^n) \) is the number of phrases in the dictionary. For each version (LZ-78, LZ-77, LZ-76), the dictionary is different, and hence the complexity defined by (1) is different. However, each version of complexity \( K(X^n) \) is known to approach the entropy rate \( \frac{1}{2} H(X^n) \) as \( n \to \infty \), albeit at with a slow rate of convergence [5].

For two bit strings \( X^n \) and \( Y^n \) of equal length, the generic Lempel-Ziv distance is defined as [3]

\[
d_{LZ} = 1 - \min \left( \frac{K(X^n) - K(X^n|Y^n)}{K(X^n)}, \frac{K(Y^n) - K(Y^n|X^n)}{K(Y^n)} \right),
\]

where \( X^n|Y^n \) contains the phrases in \( X^n \) that are not in \( Y^n \). Specializing respectively to the versions LZ-78, LZ-77, L

\[\text{Fig. 3. Time course of normalized Fluo-4 intensity for four neurons indexed 1-4 in a neuron population.}\]

Fig. 4. Raster plots of synthetic spike trains for sequence length 200 (we use length of 500 for analysis) for synthetic trains 1-2 of pair (i), 3-4 of pair (ii), 5-6 of pair (iii) and 7-8 of pair (iv).

76, we define by (2) distances \( d_{LZ78}, d_{LZ77}, d_{LZ76} \), considering the corresponding suitable dictionaries.

C. Proposed Empirical Dissimilarity Measure

We computed the empirical contextual probabilities of symbols up to a pre-fixed maximum context length \( k \) in each given spike sequence \( X^n \) [5]. For instance, consider the sequence \( X^n = '01001111010100' \) and \( k = 2 \) (‘010’ - 4 - 4/5: indicates symbol ‘0’ occurs for four times when context ‘01’ appears for five times in \( X^n \)).

\[
\begin{align*}
&\text{• k=0 (no context): } '0' - 8 - 8/15; '1' - 7 - 7/15; \\
&\text{• k=1: '00' - 2 - 2/7; '01' - 5 - 5/7; '10' - 5 - 5/7; '11' - 2 - 2/7; } \\
&\text{• k=2: '000' - 0 - 0; '001' - 1 - 1; '010' - 4 - 4/5; '011' - 1 - 1/5; '100' - 2 - 2/5; '101' - 3 - 3/5; '110' - 1 - 1/2; '111' - 1 - 1/2. }
\end{align*}
\]

The empirical Hellinger distance between probability distributions \( P(x^n) \) and \( Q(x^n) \) of two spike sequences is defined in terms of Bhattacharyya coefficient \( BC(P, Q) \) as

\[
d_{EHD} = \sqrt{1 - BC(P, Q)},
\]

where

\[
BC(P, Q) = \sum_{x^n \in X^n} \sqrt{P(x^n)Q(x^n)}
\]

\[
= \sum_{x_1} \left[ \sum_{x_{n-k}} \left( \sum_{x_1} \sqrt{P(x_{n-k} | x_1)Q(x_{n-k} | x_1)} \right)^2 \right] \sqrt{P(x_1)Q(x_1)},
\]

and \( P(x_{n-k} | x_1) \) and \( Q(x_{n-k} | x_1) \) denote respective empirical conditional probabilities of the two sequences at context length \( k \).

D. Clustering using Gaussian mixture models

We adopted the Gaussian mixture model for clustering the dissimilarities between the spike trains of different neuron pairs in population. In particular, the mixture probability density function (pdf) is assumed to follow [9]

\[
p(x; \theta) = \sum_{i=1}^{K} w_i \phi(x; \mu_i, \sigma_i^2),
\]

where \( \phi(\cdot) \) indicates the Gaussian pdf, and \( \mu_i, \sigma_i^2 \) and \( w_i \) respectively denote the mean, variance and mixing weight of the \( i \)-th Gaussian component. Further, the parameter vector \( \theta = (\mu_i, \sigma_i^2, w_i)_{i=1}^{K} \), assuming \( K \) Gaussian components.
III. RESULTS

We begin by demonstrating our tools on synthetic spike trains. Subsequently, such tools are applied to experimentally observed calcium spike trains.

A. Synthetic Spike Trains

We generated four pairs of spike trains as follows: (i) a pair following independently identically distributed (iid) Poisson model with firing rate $\lambda = 50$ Hz; (ii) an independent pair following Poisson model with firing rates 50 Hz and 300 Hz; (iii) an iid pair, each following a Poisson model with a random $\lambda$ picked for each time window of length 50, and uniformly distributed in the interval [50 300] Hz; (iv) a phase-shifted pair of periodic spike trains with period 20 ms. In (i)-(iii), a spike in each $dt = 1$ ms interval is generated when a uniform random number in [0,1] turns out to be less than $\lambda dt$. The raster plot of all 8 spike trains, described above, is shown in Fig. 4. We next compare empirical Hellinger distance measure $d_{EH}$ for $k = 0, 2, 4, 6, 8, 10$ with existing LZ based distance measures $d_{LZ78}$, $d_{LZ77}$, as shown in Figs. 5(a)-5(d) respectively. We computed $d_{EH}$ using (3) and $d_{LZ78}$, $d_{LZ77}$, $d_{LZ76}$ using (2). In pair (i) (as well as in pair (iii)), the computed distances $d_{LZ78}$ and $d_{EH}$ are small as spike trains are iid (see Figs. 5(a) -5(c)). However, the distances $d_{LZ77}$ and $d_{LZ76}$ are large. A similar behavior is observed in case of non-random periodic pair (iv) as well. So, $d_{LZ77}$ and $d_{LZ76}$ appear unsuitable as measures of distance, and will not be considered for further analysis. Revisiting case (iv), $d_{EH}$ quickly converges to zero, whereas $d_{LZ78}$ converges slowly. In case (ii), where rival spike trains have different distribution, $d_{LZ78}$ and $d_{EH}$ both capture the dissimilarity. However, $d_{EH}$ behaves similarly to $d_{LZ78}$ for suitable $k$ (= 6), albeit with faster convergence.

B. Calcium Spike Trains

As mentioned earlier, we consider eight neurons, indexed 1-8, whose spike trains are inferred. The corresponding raster plot is shown in Figure 6. Taking successive neurons as pairs (1-2, 3-4, 5-6, 7-8), we next compared $d_{EH}$ for $k = 0, 2, 4, 6, 8, 10$ with $d_{LZ78}$ for different sequence lengths in respective Figs. 7(a)-7(d). We observe that the proposed distance measure $d_{EH}$ converges faster than $d_{LZ78}$ for all $k.$
Further, \( d_{\text{EHD}} \) approximates \( d_{\text{LZ78}} \) for \( k = 4 \) beyond sequence length 200. However, this analysis needs to be performed for more neurons to obtain optimum \( k \). Next, we considered sequence length 200, and clustered neuronal spike train pairs using Gaussian mixture models based on distance measures \( d_{\text{LZ78}} \) and \( d_{\text{EHD}} \) with \( k = 4 \). Varying number \( K \) of clusters, we observed two significant clusters, which oscillated with temporal shift. Specifically, refer to Fig. 8 for such behaviour for both distance measures, when the starting frame is shifted by 0, 1, 2 and 3 samples. Interestingly, the variations in mean values of those clusters (for each of \( d_{\text{LZ78}} \) and \( d_{\text{EHD}} \)), when plotted against temporal shifts ranging 0–140 (refer Figs. 9(a) and 9(b), respectively), show oscillatory fluctuations that are small compared to the difference in mean. Further, we performed k-means clustering on means of the two clusters (labeled L, low-mean, and H, high-mean), each pair generated for temporal shifts 0–140, and similarly assigned a second label L or H. Next, we plotted relative histogram for classes LL, LH, HL and HH in Fig. 10 for both distance measures at hand. We observed that the relative count in LL in case of \( d_{\text{LZ78}} \) is less compared to \( d_{\text{EHD}} \). It potentially indicates that \( d_{\text{LZ78}} \) shows some neuron pairs as statistically dissimilar, while those are actually similar.

IV. CONCLUSION

In this paper, we proposed an empirical Hellinger distance measure to quantitatively statistical dissimilarity in neuronal calcium spike trains. We illustrated our method with synthetic as well as experimental spike trains, and showed fast convergence. At the same time, our method exhibited similar (or, slightly superior) clustering behavior. The quantitative analysis of how fast is our method compared to existing methods would be considered as a future work. Further, we would like to extend our analysis for spike trains recorded with micro-electrode array [10]. We expect the proposed method to facilitate large-scale studies of functional clustering, especially involving short sequences, which would in turn identify signatures of various diseases in terms of clustering patterns. In particular, the proposed scheme could assume importance in functional clustering of neurons in dissociated cultures.

V. ACKNOWLEDGMENT

We thank Drs. Mennerick and Gautam for providing materials and equipment. Sathish Ande thanks the Ministry of Electronics and Information Technology (MeitY), the Government of India, for fellowship grant under Visvesvaraya PhD Scheme.

REFERENCES

[1] I. Soltész and A. Losonczy, “Cal pyramidal cell diversity enabling parallel information processing in the hippocampus,” Nature neuroscience, vol. 21, no. 4, pp. 484–493, 2018.

[2] S. Schreiber, J.-M. Fellous, D. Whitmer, P. Tiesinga, and T. J. Sejnowski, “A new correlation-based measure of spike timing reliability,” Neurocomputing, vol. 52, pp. 925–931, 2003.

[3] M. Christen, T. Ott, and R. Stoop, “Spike train clustering using a lempel-ziv distance measure,” in Proceedings of the international symposium on nonlinear theory and its applications (NOLTA). Citeseer, 2004, pp. 379–82.

[4] M. Christen, A. Koh, T. Ott, and R. Stoop, “Measuring spike pattern reliability with the lempel-ziv-distance,” Journal of neuroscience methods, vol. 156, no. 1-2, pp. 342–350, 2006.

[5] S. Ande, J. Regatti, N. Pandey, S. Avasarala, S. Swain, A. Karunarathne, L. Giri, and S. Jana, “Information content in neuronal calcium spike trains: Entropy rate estimation based on empirical probabilities,” in 6th International IEEE-EMBS Conference on Biomedical Engineering and Sciences (IECBES 2020). IEEE, 2020 (Accepted). [Online]. Available: https://arxiv.org/abs/2102.00723

[6] L. Giri, A. K. Patel, W. A. Karunarathne, V. Kalyanaraman, K. Venkatesh, and N. Gautam, “A g-protein subunit translocation embedded network motif underlies gpcr regulation of calcium oscillations,” Biophysical journal, vol. 107, no. 1, pp. 242–254, 2014.

[7] J. T. Vogelstein, A. M. Packer, T. A. Machado, T. Sippy, B. Babadi, R. Yuste, and L. Paninski, “Fast nonnegative deconvolution for spike train inference from population calcium imaging,” Journal of neurophysiology, vol. 104, no. 6, pp. 3691–3704, 2010.

[8] J. M. Amigo, J. Szczepanski, E. Wajnryb, and M. V. Sanchez-Vives, “Estimating the entropy rate of spike trains via lempel-ziv complexity,” Neural Computation, vol. 16, no. 4, pp. 717–736, 2004.

[9] M. Geoffrey J and P. David, Finite Mixture Models. Wiley Series in Probability and Statistics, 2000.

[10] M. Mahmud and S. Vassanelli, “Processing and analysis of multichannel extracellular neuronal signals: State-of-the-art and challenges,” Frontiers in neuroscience, vol. 10, p. 248, 2016.