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TOWARDS A STATISTICAL DESCRIPTION OF EXPERIMENTAL DATA FOR DETECTION-ESTIMATION PROBLEMS IN DNA TRANSLOCATIONS THROUGH NANOPORES

S. Michelet\(^1\), JP. Barbot\(^1\), O. François\(^1\), PY. Joubert\(^1\), P. Larzabal\(^1\), R. Kawano\(^2\), H. Sasaki\(^2\), T. Osaki\(^2\), S. Takeuchi\(^2\), B. Le Pioufle\(^1\)
\(^1\)SATIE, ENS Cachan, CNRS, UniverSud, 61 Avenue du Président Wilson, Cachan, France
\(^2\)KAST and IIS University of Tokyo, Japan

stephane.michelet@ens-cachan.fr, bruno.lepioufle@satie.ens-cachan.fr

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Abstract: This paper investigates the properties of DNA translocations signals in a stochastic framework. The considered signals are relative to the translocation of single strand DNA through natural nanopores, and are obtained using a planar patch clamp method. The stochastic signal analysis is carried out considering the statistical distribution of DNA translocation parameters, considered as random variables including the amplitude, the duration and the apparition of the DNA translocation events as well as the no-translocation signal features. For each of these variables, a distribution function is proposed and assessed using a Kolmogorov-Smirnov test, and their features are estimated. The DNA translocation signal stochastic analysis enables to characterize the detection and/or estimation performances of existing algorithms, such as a breakdown detection algorithm, in a stochastic framework. Moreover, it opens the way to the design of model based algorithms such as detection tests using a likelihood ratio or joint detection-estimation algorithms using a maximum likelihood approach, for an enhanced characterization of DNA translocations.

1 INTRODUCTION

In view of the DNA sequencing, a biochip dedicated to the DNA translocation through natural nanopores reconstituted on an artificial biomimetic membrane was designed in (Osaki et al., 2009). The biochip consists in a partition between a fluidic chamber and a channel, made with a thin film of parylen obtained by chemical vapor deposition, and micromachined through oxygen plama (see figure 1).

The biomimetic artificial membrane is built up on this partition, by the successive flow of lipids and buffers into the channel, as described in (Osaki et al., 2009) and the nanopore is created thanks to the insertion of an \(\alpha\)-hemolysin natural membrane protein. The application of a voltage on both sides of the membrane induces the movements of ions, and therefore the apparition of a current through the channel. The DNA strand crossing through the membrane induces a current blockade, measured thanks to a patch clamp amplifier. The amplitude and duration of this blockade characterizes the DNA composition and length. The blockade current constitutes the informative signal which is sampled and digitalized by the experimental setup. In order to avoid aliasing during the acquisition process, a so called anti-aliasing low pass filter is used to process the experimental data.

Getting DNA translocation signals is a delicate experiment, since the obtained signals depend on many parameters, such as temperature, humidity, sealing of the artificial membrane or surface conditions of the electrodes.

In this study, in order to avoid repetitive exper-
iments required to adjust the acquisition parameters left to the users and the dedicated data processing techniques, artificial signals are generated. The properties of these artificial signals are determined through the statistical investigations of actual biosignals. In section 2 the statistical properties of the signal are estimated, including the no-translocation current, the amplitude and duration of the DNA translocations events, and the delay between events. In section 3, corresponding artificial signals are generated and used to optimally design an amplitude-duration characterization algorithm based on a breakdown detection approach, and used to evaluate the amplitude-duration characterization performances. In section 4, thanks to the proposed statistical framework, the relevance of model based approaches is pointed out, in order to develop i) a detection test using likelihood ratio, and ii) a joint detection-estimation algorithm based on a maximum likelihood method.

2 A STATISTICAL DESCRIPTION OF THE DNA TRANSLOCATION SIGNAL

In this study, the investigation is carried out in a stochastic framework for the current flowing through the nanopore. The considered experimental data are relative to the translocations of a 41mer ssDNA TTTTTTTTTTCATGAC-CTGGGGGAGTTGCGGAGGAAGGT, the concentration of which is 45 µM in a 1.0 M KCl, 10 mM PBS, 1 mM EDTA buffer featuring pH=7.4. The DNA translocations are conducted thanks to a 80 mV voltage applied between both sides of the lipid bilayer.

The stochastic characterization of DNA signals consists in the evaluation of the statistical distribution of the amplitude, denoted AMP, the duration (DUR), the delay between translocation (DBT), and the no-translocation signal (NTS), which are defined in figure 3.

2.1 Properties of the Current Through Nanopore in Absence of Translocation

Firstly we examine the statistical properties of the actual current flowing through an open α-hemolysin channel nanopore without any DNA stand translocation. An example of the current flowing through the nanopore is shown in figure 4.

In order to attest the assumed Gaussian distribution of the no-translocation signal, a Kolmogorov-Smirnov (KS) test was implemented. The KS test actually quantifies the distance between the cumulative distribution function (CDF) of the considered experimental data, denoted $F_n(x)$, and the CDF of a reference distribution denoted $F(x)$ (Kendall and Stuart, 1979). This KS will be prefered to the Chi-2 test which is sensitive to a lack of data in the experimental histogram. The KS distance is expressed by:

\[
\mu = \frac{1}{n} \sum_{k=1}^{n} x[k] \quad (1)
\]

\[
\sigma^2 = \frac{1}{n-1} \sum_{k=1}^{n} (x[k] - \mu)^2 \quad (2)
\]

where $x$ is the signal and $n$ the number of samples. Considering the available experimental data, the estimation using eq. (1) and eq. (2) leads to $\mu_{NTS} = 93.7$ pA and $\sigma_{NTS} = 1$ pA.

An histogram of this no-translocation signal is shown on figure 5, which seems to exhibit a Gaussian distribution. The mean and standard deviation parameters of the distribution, respectively denoted $\mu$ and $\sigma$, are estimated using:

![Figure 3: Features of the DNA translocation signal](image)

![Figure 2: Examples of current blockades relative to DNA translocation and translocation attempts.](image)

![Figure 4: Real current variations through α-hemolysin channel without any translocation event.](image)
\[ D_n = \sqrt{n} \times \sup_x |F(x) - F_n(x)| \]  \hspace{1cm} (3)

where \( n \) is the number of samples of the experimental data. If this distance \( D_n \) is greater than a predefined threshold, then the hypothesis according to which the experimental data distribution is close to the candidate reference distribution is rejected. The threshold is adjusted for a false reject rate of 1%.

Here, the KS test validates the normal distribution of the no-translocation current, as shown on figure 6, which exhibits the CDF \( F_n(x) \) and \( F(x) \).

2.2 Statistical Properties of Translocations Events

In this section, we characterize the translocation events through their duration and amplitude distribution. Indeed, the translocation event provokes a current blockade featured by a duration and an amplitude which give biological information on the ssDNA. The amplitude vs time duration graph permits to determine the length of the DNA, and provides information about its composition, such as the discrimination between polyU, polyC or polyA, (Akeson et al., 1999).

As usually admitted (Kasianowicz et al., 1996), only translocations with a current amplitude decreasing more than 80% of the initial value correspond to complete translocations. Others are translocation attempts which are not considered here.

2.2.1 Amplitude and Duration of the Translocation Events

Thanks to equations (1) and (2) the amplitude distribution mean value and standard deviation of the translocation amplitude \( \text{AMP} \) can be estimated: \( \mu_{\text{AMP}} = 89.2 \) pA and \( \sigma_{\text{AMP}} = 7.33 \) pA.

In this study, the translocation current amplitude \( \text{AMP} \) is assumed to be normally distributed, and a KS test implemented has validated this assumption.

In (Meller et al., 2000), the distribution of the translocation duration was approximated using a mixture of a Gaussian law and an exponentially decaying law. Here, for tractability purposes, a Rayleigh law (Eq. 4) seems to be more adequate to fit the \( \text{DUR} \) actual distribution law, and will therefore be preferred (Figure 7). The KS test validates this distribution law.

\[ f(DUR) = \frac{DUR}{r_{DUR}^2} \exp \left( -\frac{DUR^2}{2r_{DUR}^2} \right) \]  \hspace{1cm} (4)

For the duration distribution the Rayleigh parameter \( r \) is estimated according to equation (5).

\[ r_{DUR} = \frac{1}{2n} \sum_{k=1}^{n} (DUR[k])^2 = 924 \mu\text{sec} \]  \hspace{1cm} (5)

Figure 6: Cumulative distribution functions \( F_n(x) \) (Gaussian) \( F(x) \) (experimental data)

Figure 7: Cumulative distribution functions \( F_n \) and \( F \) relative to the Rayleigh distribution of \( \text{DUR} \)

2.2.2 Statistics of Delay Between Translocations Events

The distribution of the delay between translocations \( \text{DBT} \) is considered in this section and assumed to be a decreasing exponential, expressed by:

\[ f(DBT) = \alpha \exp(-\alpha DBT) \]  \hspace{1cm} (6)

where:

\[ \alpha = \frac{n}{\sum_{i=1}^{n} DBT(i)} \]  \hspace{1cm} (7)

\[ \alpha = (17.6 \text{ms})^{-1}. \]
2.2.3 Statistical Description of the DNA Translocation Signal

Finally, the distribution features of the random variables $AMP$, $DUR$, $DBT$ and $NTS$, estimated from experimental DNA translocation signals are gathered in table 1.

<table>
<thead>
<tr>
<th></th>
<th>Dist</th>
<th>param. 1</th>
<th>param. 2</th>
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<tr>
<td>$NTS$</td>
<td>Gaus.</td>
<td>$\mu = 93.7 \text{ pA}$</td>
<td>$\sigma = 1 \text{ pA}$</td>
</tr>
<tr>
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</tr>
<tr>
<td>$DUR$</td>
<td>Ray.</td>
<td>$r = 924 \mu$s</td>
<td></td>
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<tr>
<td>$DBT$</td>
<td>Exp.</td>
<td>$\alpha = (17.6 \text{ ms})^{-1}$</td>
<td></td>
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</table>

3 Performances of a no-parametric amplitude-duration estimation algorithm

In this section, the DNA translocation signal characterization results are used to evaluate, in a stochastic framework, the performances of an elementary translocation characterization algorithm. The considered algorithm is based on a breakdown detection technique, presented in (Osaki et al., 2010) which allows to estimate the amplitude and duration of translocation events, respectively denoted $\hat{AMP}$ and $\hat{DUR}$. To evaluate the performances of this characterization algorithm for various signal features, we build up artificial biomimetic signal considering the $AMP$, $DUR$, $DBT$ and $NTS$ distributions estimated in the previous section. Moreover, in order to take the possible experimental noise variance variations into account relative to the experimental set-up, we elaborate artificial signals featured by various signal to noise ratios ($SNR$), defined as:

$$SNR = 20 \log \left( \frac{|\hat{AMP}|}{\sigma_{NTS}} \right)$$  \hspace{1cm} (8)

An example of a 319 translocation signal sequence featured by a 30 dB $SNR$ is represented in figure 8, and figure 9 exhibits the detail of a single artificial translocation event. The implementation of the breakdown detection algorithm applied to this translocation sequence allows the $AMP$ and $DUR$ values of the 319 translocations to be estimated. The corresponding amplitude vs duration representation diagram is depicted in figure 10.

In order to quantify the characterization performances of the algorithm, we compute the true positive rate and the false positive rate of the characterization algorithm, considering $SNRs$ ranging from 6 to 46 dB. The true positive rate (TPR) is computed as the rate of the estimated $AMP$-$DUR$ values of each considered translocation event which are close to the actual values within a predefined distance $\nu$. On the other hand, the false positive rate (FPR) is defined as the rate of the estimated $AMP$-$DUR$ values which are wrongly positioned at a distance smaller than a varying distance $\nu$. Then, the receiver operational characteristic (ROC) which plots the TPR as a function of FPR for various values of $\nu$ can be considered to quantify the characterization performances (Bradley, 1997). ROC curves obtained for the considered translocations data are presented in figure 11. One can note for example that a 90 % TPR is reached at the cost of a 0.01 % FPR considering a translocation sequence with $SNR = 28$ dB, and that the same 90 % TPR is reached at the cost of a 1 % FPR when the $SNR$ falls down to 18 dB. An other means of quantifying the performance of the amplitude-duration estimation algorithm is to evaluate the mean square error ($MSE$) defined in equation (9), of the characterization as a function of the $SNR$ of the translocation signal.

$$MSE = \frac{1}{n} \sum_{i=1}^{n} \left( \frac{\hat{AMP}_i - AMP_i}{\mu_{AMP}} \right)^2 + \left( \frac{\hat{DUR}_i - DUR_i}{\mu_{DUR}} \right)^2$$  \hspace{1cm} (9)

where $\hat{AMP}$ and $\hat{DUR}$ are the estimated values of $AMP$ and $DUR$ respectively, $n$ is the number of translocations equal to 319, and where the contribution of the amplitude and duration errors are normalized by their mean values in order to give them the same weight in the computation of the $MSE$. The $MSE$ computed according to equation (9) and expressed in percent is represented in figure 12. One can note that the $MSE$ falls from 30 % to 0.02 % when the $SNR$ rises from 6 dB up to 46 dB, respectively.

![Figure 8: Generated artificial signal.](image_url)
Figure 9: Single translocation event in the artificial signal.

Figure 10: Diagram duration vs amplitude for the artificial translocations.

Figure 11: ROC of the duration-amplitude characterization.

4 DISCUSSION

We have proposed a statistical characterization of nanopore DNA translocation current allowing well known methods of amplitude/duration characterizations (Basseville and Nikiforov, 1993) to be implemented and evaluated using intensive computer simulations.

More challenging now is the use of this statistical characterization to optimally build new model based approaches to improve the characterization performances. The proposed modelling of translocation signals opens this way. As a complete statistical characterisation of a steplike signal is now available, several ways of investigation are opened. Let us briefly point out two of them for further works:

i) a model based segmentation procedure which detects multiple change points in a steplike signal can be built on a generalized likelihood ratio test or on information theoretic criterion such as Akaike information criterion like tests. Moreover, since this segmentation technique considers the DNA translocation signal as a whole sequence, it avoids the well known drawbacks relative to sliding window data processing approaches.

ii) a regularized maximum likelihood method can be built, looking for the unknown parameters $\theta$ as:

$$\hat{\theta} = \arg \min_{\theta} \{ \| s \ast (t) - s(\theta) \|_2^2 + \lambda \| \nabla s(\theta) \|_1 \}$$  \hspace{1cm} (10)

with $\theta = [t_1, t_2, ..., t_N, a_1, a_2, ..., a_N]^T$ where $t_i$ are the step location parameters and $a_i$ are the step amplitude parameters.

$\nabla s(\theta)$ is the gradient of the solution. As we are looking for a steplike signal, for regularization purposes a $l_1$ norm will be used for the gradient.

$s \ast (t)$ is the actual recorded signal and $s(\theta)$ is a candidate signal. Recent developments in convex constraint optimisation open the way to an efficient optimisation of the criterium expressed in equation (10). $\lambda$ is a parameter used to adjust the contribution of each terms of the regularization criterion.

This provides a statistical framework for DNA translocation characterisation.

ACKNOWLEDGEMENTS

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