RAPID FINITE STATE MACHINE CONTROL OF INDIVIDUAL DNA MOLECULES IN A NANOPORE

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Abstract: This paper demonstrates feedback voltage control of individual DNA hairpin molecules captured in a nanopore. A finite state machine is used to program voltage control logic, executed on a field-programmable gate array, for rapid detection and regulation of hundreds of DNA hairpins, one at a time. Prompt voltage reduction is used for extension of the dwell time of DNA hairpins in the nanopore. Then, voltage reversal after a preset dwell time is used for automated expulsion of molecules prior to hairpin unzipping. The demonstrated control authority of single molecular complexes captured in the nanopore device is an integral part of our ongoing research for direct monitoring and control of enzyme-bound biopolymers.

1 INTRODUCTION

Nanopore sequencing is based on electrophoretically driving a single-stranded DNA (ssDNA) or RNA molecule through a nano-scale pore (Deamer and Branton, 2002). The potential of this technology is high-speed, high throughout sequential identification of all nucleotides in any single DNA or RNA molecule. Many research groups are now exploring and developing biological and solid-state nanopores to achieve low-cost, high throughput nanopore-based sequencing (Rhee and Burns, 2006), in addition to other single molecule sensing applications (Dekker, 2007).

In the biological nanopore setup, a planar lipid bilayer is created across a 20 µm teflon aperture in a KCl solution. A single α-hemolysin protein channel is inserted into the planar lipid. The channel (pore) is 15 nm in length and varies in diameter. The cis-opening of the pore is 2.6 nm wide, opening to a 3.6 nm vestibule before narrowing to a limiting 1.5 nm width at the beginning of the stem. The remainder of the stem up to the trans-opening is 2 nm wide. The vestibule is large enough for double-stranded DNA (dsDNA) to enter, but the limiting stem is just wide enough for ssDNA to pass through. Across the bilayer, AgCl electrodes are used to apply a potential that produces an ionic current through the pore, passing from the cis side to the trans side of the pore with the trans-side voltage positive. As molecules translocate, the pore becomes partially blocked by the translocating molecule, causing a momentary drop in current. These translocation events can be characterized by the amplitude of the blockade current and the time the molecule spends in the pore, defined as the dwell time.

We use DNA oligomer that is 79 nucleotides total in length, with a 20 base pair hairpin (20 bphp). The hairpin is formed by the 3' end folding over and annealing on itself resulting in a 20 base pair region. The hairpin is thus the double-stranded segment, with the single-stranded segment 35 nucleotides long (4 unpaired bases in the doubled-stranded end loop). Upon capture of the ssDNA end, the hairpin enters the pore vestibule and remains until the hairpin is unzipped. A schematic of the nanopore system and an example 20 bphp translocation event is illustrated in Figure 1.

Regarding the resolution limits of ionic current measurements, homopolymers of ssDNA and block copolymers of RNA are distinguishable based on the measurable differences in the blockade current amplitude or kinetics (Akeson et al., 1999). However, translocation rates are too fast (up to 2 nucleotides/µsec, (Akeson et al., 1999)) to identify individual nucleotides in heterogeneous single-stranded polymers using existing biological nanopores (Dekker, 2007). In this paper and other...
The voltage field force exerted on the ssDNA causes it to dissociate from the enzyme after several milliseconds. The time-to-dissociation is captured in the nanopore, with the enzyme residing on top of the nanopore being too large to translocate through it. Binding of enzymes to DNA in this configuration has been shown to increase the dwell time of DNA in the nanopore by up to two orders of magnitude (up to 200 msec). Recently, kinetics of Escherichia coli exonuclease I binding to ssDNA has been quantified using voltage ramps for nanopore-based force spectroscopy (Hornblower et al., 2007). The voltage field force exerted on the ssDNA causes it to dissociate from the enzyme after several milliseconds before translocating. The time-to-dissociation in turn can be correlated to enzyme binding rate constants.

In (Benner et al., 2007), the interaction of DNA with the Klenow fragment (KF) of Escherichia coli DNA polymerase I was explored. In the absence of KF, capture and subsequent unzipping of 20 bphp at constant 180 mV reveals blockades with 20 pA mean amplitude and 4 msec median dwell time. Addition of KF and the dNTP complementary to the DNA template base in the KF catalytic site yielded a substantial increase in blockade dwell times (110 msec median lifetime for dGTP), attributable to ternary (DNA-KF-dGTP) complexes. Closer investigation of such blockades revealed a two-step pattern in greater than 97% of the blockades, the first step at 24 pA mean amplitude, and the second (terminal) step at 20 pA mean amplitude lasting 4ms consistent with the hairpin kinetics alone. It was demonstrated that the transition from step one to two resulted in dissociation of KF from DNA first, followed by hairpin dropping into the pore vestibule until unzipping occurred. As a initial effort at voltage control of enzyme-bound DNA, we demonstrated efficient automated detection of individual ternary complexes (< 3 msec), based on the characteristic 24 pA amplitude, and truncation of the blockade time by voltage reversal after 20 ms (Benner et al., 2007)).

This paper presents an extension of the control results presented in (Benner et al., 2007). Specifically, we demonstrate automated detection and manipulation of DNA hairpins. Rapid detection (< 2 msec) is based on computing a filtered mean amplitude of the ionic current in real time, and monitoring the mean relative to an amplitude range consistent with DNA hairpin blockades. Confirmation of our ability to detect DNA-enzyme blockades and control authority to aggregate the dwell times of hundreds of blockade events. Additionally, it complements our prior work (Benner et al., 2007), confirming our ability to detect DNA-enzyme blockades and DNA hairpin blockades. Confirmation of our ability to discern between each blockade type in real time is part of our ongoing work. Ultimately, nanopore-based characterization of enzyme dynamics will require direct detection and control of multiple DNA conformations relative to the enzyme, and direct control of enzyme-free DNA is a prerequisite toward developing

Figure 1: Schematic of nanopore and DNA, and plot of representative ionic current signal during a 20 bphp DNA translocation event under 180 mV applied potential. (I) At 180 mV, KCl ions pass through the open channel resulting in ~64 pA current. (II) Upon capture of the single-stranded end of the DNA molecule into the cis opening of the pore, the flow of ions is reduced to ~20 pA. (III) After ~5 msec, the voltage unzips the hairpin, causing ssDNA to pass through the pore into the trans chamber, completing the measured blockaded event. The duration of the event is referred to as dwell time.
this capability.

Direct control of ssDNA in a nanopore has been demonstrated (Bates et al., 2003), in which detection of DNA is based on monitoring the raw amplitude relative to a threshold level. Voltage level changes, comparable to those employed in this paper, were commanded to explore the zero and low voltage effects on ssDNA-pore interactions. In contrast to thresholding the raw ionic current amplitude, the windowed amplitude mean calculation we have used here filters the current noise. Additionally, detection depends on the mean remaining within a preset amplitude range (< 6 pA in spread) for multiple consecutive comparisons. Alternative methods for single molecule sensing and manipulation include optical tweezers and atomic force microscopy (Bustamante et al., 2003). For example, optical trapping has been used to sequence DNA by attaching a processive enzyme to a polystyrene bead (Abbondanzieri et al., 2005), (Greenleaf and Block, 2006). At present, greater spatial and temporal resolution of single DNA molecule polymerization has been achieved than with nanopores. However, these methods generally require more preparative steps, and far fewer molecules can be analyzed over a common time period.

2 CONTROL LOGIC SETUP

The nanopore system is setup in a 0.3 M KCl solution. A patch-clamp amplifier, Molecular Devices AxoPatch 200B, regulates the applied voltage and measures the ionic current through the channel. The data are recorded using the Molecular Devices Digidata 1440A digitizer, sampled at 50 kHz and low-pass filtered at 5 kHz with a four-pole Bessel filter. The voltage control logic is programmed using a finite state machine (FSM) within LabVIEW 8 software. The FSM logic is implemented on a field-programmable gate array (FPGA) hardware, National Instruments PCI-7831R. An FPGA is a reconfigurable hardware platform that permits fast measurement and voltage reaction times (1 μsec output sample time). An FSM is a logic construct where program execution is broken up into a series of individual states (Gill, 1962). Each state has a command associated with it, and transitions between states are a function of system measurements. Measurements of the pore current are processed and passed to the FSM as inputs. Changes in the FSM control logic are done as necessary, then re-compiled and re-routed to run on the FPGA. This achieves a balance between speed and flexibility, by enabling the system to react to events on the order of a microsecond, while also allowing for the control logic to be reconfigured as necessary between experiments.

In our first experiment, the objective was to efficiently detect individual DNA hairpin events, and increase the blockade dwell time by lowering the applied voltage from 180 mV to 150 mV upon detection. This is referred to as dwell time extension control. Next, we sought to aggregate the extended blockade dwell times, by expelling the DNA using voltage reversal of -50 mV after 10 ms at 150 mV. This is referred to as dwell time aggregation control. The motivation was to increase the nominal hairpin dwell time, and expel the molecule before unzipping the hairpin. A typical 20 bphp event at constant 180 mV voltage is shown in Figures 1 and 2a. The probability histogram of the base 10 logarithm of dwell time (Figure 2aIII, blue) is unimodal, with median dwell time of 2.8 ms. The median amplitude of the event plot in Figure 2aII is 20.9 pA with an interquartile range (IQR) of 1.7 pA. Only 6% of events are in the subset range of 13-18 pA (2aIII, yellow). For the same experiment at constant 150 mV voltage (data not shown), the events cluster around a median amplitude of 15 pA and 87% of 150 events are in the 13-18 pA range. Thus, under extension and aggregation control for which the voltage is reduced to 150 mV for all detected events, a larger

Certain inefficiencies in FPGA signal routing into the sampling loop caused the additional 0.65 ms of delay in the reaction time. By bringing global signals inside the sampling loop, the delay has recently been eliminated, reducing detection time to 1.35 ms.

3 EXPERIMENTS AND RESULTS
percentage of blockades should have a mean amplitude within the 13-18 pA range.

3.1 Dwell Time Extension (Figure 2b)

Upon diagnosis of a DNA hairpin event using the mean filtered current, the command voltage is reduced to 150 mV until the hairpin unzips and the DNA translocates through the pore. Using 180 mV for capture results in more events than 150 mV, while reducing to 150 mV extends the life of the hairpin. Dwell time extension is useful for terminal base-pair sequencing by machine learning methods (Vercoutere et al., 2003). After each translocation, the FPGA resets the voltage to 180 mV. A representative event is shown in Figure 2bI. The event plot (Figure 2bII) pattern shows that events faster than the nominal diagnosis time of 2.0 ms are unaffected by extension control, and events with longer dwell times converge to the ∼15 pA mean amplitude as expected. The concave trend is also consistent with an equation for event’s mean amplitude vs. dwell time. In particular, for an event at 21 pA (median amplitude at 180 mV) for 2.4 ms, and at 15 pA (median amplitude at 150 mV) for 2.0 ms of detection time and 0.4 ms of transient time. While

\footnote{Step changes in voltage induce a capacitive transient, and the transient at the end of each event is ∼0.4 ms for changing from 150 mV to 180 mV. Thus, 2.4 ms at 21 pA is 2.0 ms of detection time and 0.4 ms of transient time. While}
x ms, an approximate mean amplitude \( \bar{I} \) is
\[
\bar{I} = \frac{2.4 \times 21 + 15 \times x}{2.4 + x}.
\] (1)

When \( x \approx 24 \) ms, as in Figure 2bI, \( \bar{I} = 16 \) pA. Equation (1) closely matches the mean amplitude vs. dwell time data (Fig. 2bII). Also, the fraction of events within the subset range 13-18 pA increased to 41%, as shown in the yellow histogram overlaid on the blue probability histogram (Fig. 2bIII).

### 3.2 Dwell Time Aggregation (Figure 2c)

The objective was to aggregate the dwell times of the extended events by applying 150 mV for 10 ms upon diagnosis of a hairpin event, followed by voltage reversal of -50 mV for 5 ms. The reversal time of 5 ms is known to be sufficient to clear the DNA from the channel, prepping the pore for the next event. Aggregation control would imply a measure of control over the distribution of the events, in addition to temporal control of individual molecular events. A representative event is shown in Figure 2cI. As before, the event plot (Fig. 2cII) pattern shows that events faster than the nominal diagnosis time of 2.0 ms are unaffected by aggregation control. Within the subset range of 13-18 pA, the median amplitude is 16 pA with 0.7 pA IQR (amplitude histogram shown in Fig. 2cII). The 16 pA median is consistent with (1), since for \( x = 10.0 \) ms, \( \bar{I} = 16 \) pA. Also in the subset range, and the median dwell time is 12.4 ms with 0.1 ms IQR. The low IQR indicates a high degree of control over the distribution of events that extend to at least 10 ms at 150 mV. The median dwell time of 12.4 ms is commensurate with 2.0 ms of detection time, 10 ms at 150 mV, and 0.4 ms due to a transient that is included at the end of each event resulting from voltage reversal\(^3\).

Summary statistics for the histograms in Figure 2cIII are reported in Table 1. In (Mathe et al., 2004), the authors characterize hairpin unzipping at a set of constant voltages by fitting a curve to an unzipping probability data profile. Specifically, for an unzipping time \( t \) at voltage \( V \), the unzipping probability is the fraction of events with dwell time less than \( t \), divided by the total number of events\(^4\). For a set of \( t \) values, the unzipping probability data profile is shown in Fig. 3 for our experiments.

At 180 mV constant, under extension control, and at 150 mV constant. As in (Mathe et al., 2004), we fit a line to the data, revealing a characteristic unzipping time \( t_u^V \) for constant voltage \( V \). For amplitude range 10-30 pA and dwell time range 0.3-500 ms, it is revealing to compare the median dwell times with the fitted \( t_u^V \) constants. For \( V = 180 \) mV, the median is 2.6 ms and \( t_u^{180} = 4.2 \) ms. For \( V = 150 \) mV, the median is 25.2 ms and \( t_u^{150} = 39.4 \) ms. We observe that data trimming has a significant affect over the quality of the fit to the data, and consequently over the value for \( t_u^V \). In contrast, the median does not vary as much, suggesting a sensitivity of \( t_u^V \) to outliers, in addition to the fitting method used. For example, for a dwell time range of 0.3-4000 ms at \( V = 150 \) mV, the median is

### Table 1: Summary statistics for Figure 2cIII.

<table>
<thead>
<tr>
<th>Figure No.</th>
<th>No. of Events</th>
<th>Median Dwell Time (ms)</th>
<th>IQR (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2aIII(^b)</td>
<td>472(^b)</td>
<td>2.8</td>
<td>4.2</td>
</tr>
<tr>
<td>2bIII(^c)</td>
<td>76(^d)</td>
<td>31.6</td>
<td>62.0</td>
</tr>
<tr>
<td>2cIII(^e)</td>
<td>256(^e)</td>
<td>12.4</td>
<td>0.1</td>
</tr>
</tbody>
</table>

\(^a\)Blue histogram, for events within 10 to 30 pA range. 
\(^b\)6% (27 events) within subset 13-18 pA range. 
\(^c\)Yellow subset histogram, for events within 13-18 pA range. 
\(^d\)41% of the 187 events within 10 to 30 pA range. 
\(^e\)55% of the 466 events within 10 to 30 pA range.

Figure 3: Unzipping probability data profile, defined as fraction of events with dwell time less than each unzipping time \( t \), and line \( 1 - \exp(-t/t_u^V) \) fit to profile for constant voltages \( V = 180 \) mV (red) and \( V = 150 \) mV (blue). Characteristic unzipping time constant \( t_u^V \) at constant voltage \( V \) is generated by fit. Symbols: ◦ for \( V = 180 \) mV, × for extension control (transitions from 180 mV to 150 mV), and * for \( V = 150 \) mV.
34.9 ms (+ 9.7) and \( \tau_1 = 55.7 \text{ ms (} + 16.3 \text{)}. \) Careful selection and analysis of statistical models appropriate for our data (with outliers always present) is part of our ongoing work.

### 4 CONCLUSIONS

We have shown that single DNA hairpin molecules captured in a biological nanopore can be detected and reacted to using a finite state machine implemented on a field-programmable gate array. The dwell time of such translocation events can be extended to gain more signal, which can in turn be analyzed offline using machine learning methods to yield terminal base-pair specific signatures. The signatures can then be used for real-time identification of terminal base pairs. Additionally, the finite state machine is capable of ejecting a molecule from the pore after it has been detected but prior to unzipping the hairpin. Rapid DNA hairpin detection (< 2 msec) relied on a mean filtered amplitude, which was required to remain within a preset amplitude range (< 6 pA in spread) for multiple consecutive threshold comparisons. The method will be tuned to differentiate DNA-enzyme blockades from DNA alone blockades in real time as part of our ongoing work. Ultimately, nanopore-based characterization of enzyme dynamics will require direct detection and control of multiple DNA conformations relative to the enzyme, and direct control of enzyme-free DNA is a prerequisite toward developing this capability.

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### REFERENCES


