

BIOPHYSICS

Aerolysin nanopores decode digital information stored in tailored macromolecular analytes

Chan Cao^{1,2}, Lucien F. Krapp^{1,2}, Abdelaziz Al Ouahabi³, Niklas F. König³, Nuria Cirauqui^{1,4,5}, Aleksandra Radenovic^{6*}, Jean-François Lutz^{3*}, Matteo Dal Peraro^{1,2*}

Digital data storage is a growing need for our society and finding alternative solutions than those based on silicon or magnetic tapes is a challenge in the era of “big data.” The recent development of polymers that can store information at the molecular level has opened up new opportunities for ultrahigh density data storage, long-term archival, anticounterfeiting systems, and molecular cryptography. However, synthetic informational polymers are so far only deciphered by tandem mass spectrometry. In comparison, nanopore technology can be faster, cheaper, non-destructive and provide detection at the single-molecule level; moreover, it can be massively parallelized and miniaturized in portable devices. Here, we demonstrate the ability of engineered aerolysin nanopores to accurately read, with single-bit resolution, the digital information encoded in tailored informational polymers alone and in mixed samples, without compromising information density. These findings open promising possibilities to develop writing-reading technologies to process digital data using a biological-inspired platform.

INTRODUCTION

DNA has evolved to store genetic information in living systems; therefore, it was naturally proposed to be similarly used as a support for data storage (1–3), given its high-information density and long-term storage (4) with respect to existing technologies based on silicon and magnetic tapes. Alternatively, synthetic informational polymers have also been described (5–9) as a promising approach allowing digital storage. In these polymers, information is stored in a controlled monomer sequence, a strategy that is also used by nature in genetic material. In both cases, single-molecule data writing is achieved mainly by stepwise chemical synthesis (3, 10, 11), although enzymatic approaches have also been reported (12). While most of the progress in this area has been made with DNA, which was an obvious starting choice, the molecular structure of DNA is set by biological function, and therefore, there is little space for optimization and innovation. Alternatively, precise synthesis of sequence-defined abiotic polymers has recently opened up the possibility to tune important parameters such as storage capacity, storage density, erasability, readability, and writing speed (5, 7, 8).

Decoding/reading these data can be achieved for DNA using a variety of sequencing techniques (13), whereas synthetic polymers are, so far, only deciphered by tandem mass spectrometry, a method that is efficient but requires large instruments (14, 15). Nanopores, the next generation of sequencing tool (13, 16), have been exploited for efficiently reading DNA based on the principle that translocation of different nucleobases through a nanopore generates a specific ionic current signature. More recently, nanopores have been ex-

plored for sensing digitally encoded DNA nanostructures (17, 18). Inspired by these recent advances, here, we encoded binary information in synthetic polymers tailor-made to be decoded using an engineered pore-forming toxin (PFT), aerolysin (19). By a rational and synergetic development of the pore readout (using specific aerolysin mutants) and the tailored digital analyte (using both DNA nucleotides and non-biological monomers), the translocation speed of the polymer was optimized to have a uniquely identifiable level-by-level signal, which delivered digital reading with single-bit resolution without compromising information density. Current DNA sequencing technologies based on biological nanopores (e.g., using *Mycobacterium smegmatis* porin A) are still limited by low base-calling accuracy as the maximal resolution is determined by the nearest four nucleotides translocating within the nanopore-sensing constriction (quadromers) (20, 21). Here, we show that aerolysin pores have the potential to achieve the molecular equivalent of single-base resolution for tailored digital analytes, which, in turn, allows for single-bit reading accuracy. Using deep learning, we were able to decode digital sequences encoding up to 4-bit information with a high accuracy while blindly detect the identity and relative concentration of polymer mixtures.

RESULTS AND DISCUSSION

Reading single-bit information encoded in polymers

Single-channel recording experiments were performed to analyze polymer translocation using the PFT aerolysin from *Aeromonas hydrophila* (Fig. 1A). Aerolysin is one of the best characterized among PFTs, it oligomerizes into a heptameric pore that features a novel and unique fold, constituted by two concentric β barrels held together by hydrophobic interactions (22–25). Aerolysin has been proposed to be a promising nanopore sensor, exhibiting high sensitivity for biomolecular detection and providing excellent current separation and a dwell time range suitable for accurate signal processing (26, 27). Using wild-type aerolysin, four types of nucleobase (28) and 13 natural amino acid have been identified directly (29), establishing this biological pore as a natural candidate for DNA and protein sequencing. Recently, aerolysin mutants have been rationally designed to further enhance the sensing properties of the wild-type

¹Institute of Bioengineering, School of Life Sciences, Ecole Polytechnique Fédérale de Lausanne (EPFL), 1015 Lausanne, Switzerland. ²Swiss Institute of Bioinformatics (SIB), 1015 Lausanne, Switzerland. ³Université de Strasbourg, Centre national de la recherche scientifique (CNRS), Institute Charles Sadron UPR22, 23 rue du Loess, 67034 Strasbourg Cedex 2, France. ⁴Department of Pharmaceutical Biotechnology, Universidade Federal do Rio de Janeiro, 21941-902 Rio de Janeiro, Brazil. ⁵CNRS, UMR5086, “Molecular Microbiology and Structural Biochemistry”, University of Lyon, 7 Passage du Vercors, 69367 Lyon, France. ⁶Institute of Bioengineering, School of Engineering, Ecole Polytechnique Fédérale de Lausanne (EPFL), 1015 Lausanne, Switzerland.

*Corresponding author. Email: matteo.dalperaro@epfl.ch (M.D.P.); aleksandra.radenovic@epfl.ch (A.R.); jflutz@unistra.fr (J.-F.L.)

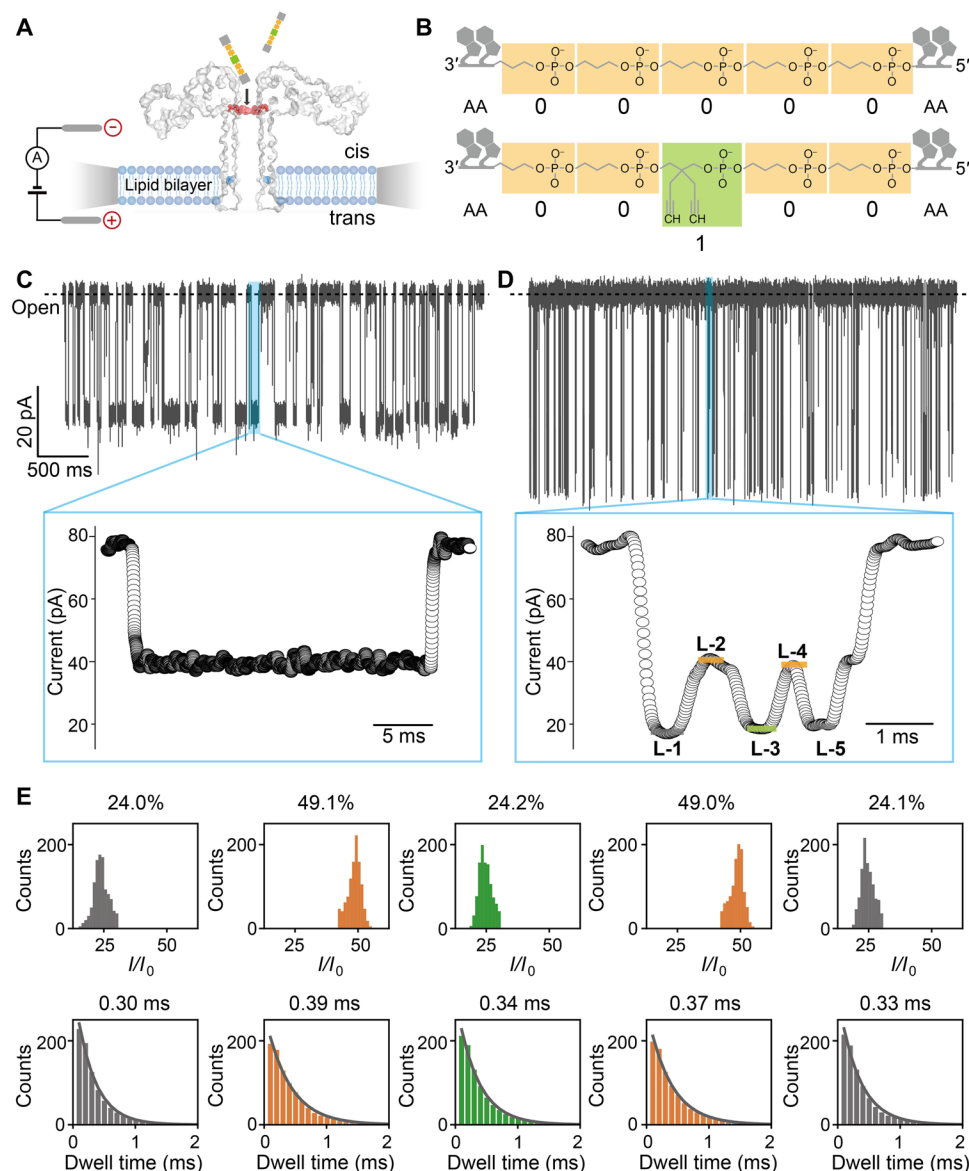


Fig. 1. Aerolysin reading of polymers encoding single-bit information. (A) Illustration of single-channel recording setup using an aerolysin pore; the *cis* and *trans* chambers are filled with 1.0 M KCl electrolyte buffer, and voltage is applied across the pore using two Ag/AgCl electrodes. Amino acid at 220 and 238 position are highlighted by red and blue, respectively. (B) Schematic structure of two representative polymers: AA00000AA and AA00100AA. (C) Raw current trace of AA00000AA during single-channel recording measurement (top). Magnification of one single event (bottom). (D) Raw current trace of AA00100AA measurement. Magnification of one single event (bottom) showing a multilevel signal: L-1 (gray), L-2 (orange), L-3 (green), L-4 (orange), and L-5 (gray). (E) I/I_0 histogram (top) and dwell time distribution (bottom) for L-1, L-2, L-3, L-4, and L-5, respectively. Relative fitted values are reported in each figure. All data were obtained using 1.0 M KCl, 10 mM tris, and 1.0 mM EDTA at pH 7.4, applying a bias potential of 100 mV.

pore (19). Notably, the K238A mutant has shown a significantly enhanced resolution for biomolecular recognition and turned out to be, likely because of a partial opening of one of the two main sensitive constrictions (Fig. 1A), the most suitable sensing system to detect the tailor-made molecular structure of the digitally encoded polymers ad hoc developed in this work.

These macromolecules are sequence-defined poly(phosphodiester)s prepared by automated phosphoramidite chemistry, as previously reported (30, 31) (Fig. 1B and fig. S1). The negatively charged backbone of these polymers ensures an efficient aerolysin capture and

the efficient translocation from the *cis* to *trans* compartment under an applied voltage (32). The polymers were digitally encoded using two monomers of different molecular structure. In the formed chains, the synthons *n*-propyl-phosphate and (2,2-dipropargyl)-propyl-phosphate represent bit-0 and bit-1, respectively. It shall be noted that this binary alphabet differs from the one that is usually used for mass spectrometry sequencing (33). Here, two monomers of markedly different bulkiness were selected to induce different pore current responses. Furthermore, automated phosphoramidite chemistry allows the use of both biological (i.e. natural DNA nucleotides) (10)

and nonbiological monomers (30). Thus, in the present work, bio-hybrid macromolecules, composed of the aforementioned nonnatural binary alphabet and natural nucleotides, were examined for optimal pore translocation. All the digitally encoded polymers examined in this study are listed in table S1, and their complete characterization is reported in figs. S40 to S85.

Digital decoding in aerolysin pores was first attempted with copolymers containing only bit-0 and bit-1 monomers. The negatively charged backbone of these polymers not only ensures efficient translocation but also speeds up the crossing time that would be too fast to allow decoding (32). As no signals were observed during the single-channel recording after addition of “00000” polymer in the *cis* side of the chamber, and the signal-to-noise-ratio was too low for “11111” (fig. S2), we introduced a di-deoxyadenosine at the polymer terminals (“AA0000AA,” Fig. 1B). This addition created highly detectable current blockade signals (39.0 ± 3.0 pA as mean residual current, with an open pore current of 76.0 ± 3.0 pA, and 18.9 ± 1.3 ms for dwell time), easily identifying the translocation of “AA0000AA” polymers (Fig. 1C). When a bit-1 monomer was inserted to create an “A00100AA” polymer, the additional moiety induced an obvious lowering of the current levels (Fig. 1D). Compared to AA0000AA, a clear decrease in the mean dwell time (2.8 ± 0.1 ms) was observed, and a fraction of the events ($\sim 28\%$) clearly showed a five-level signal (labeled L-1 to L-5 in Fig. 1D and fig. S3). To understand the relationship between the polymer chemical nature and current levels, we collected more than 10,000 blockade current events for statistical analysis (see Materials and Method and fig. S4). The relative current (I/I_0), fitted as Gaussian distribution, showed five distinct levels of values: $24.0 \pm 3.2\%$, $49.1 \pm 2.4\%$, $24.2 \pm 2.3\%$, $49.0 \pm 2.5\%$, and $24.1 \pm 2.4\%$, respectively (Fig. 1E; I_0 is the value of the open pore current, and I is the residual current value). The dwell times of each level, fitted with an exponential function, have values of 0.30 ± 0.01 , 0.39 ± 0.01 , 0.34 ± 0.01 , 0.37 ± 0.01 , and 0.33 ± 0.01 ms, respectively. The I/I_0 values of L-2 and L-4 are nearly

identical, while L-1, L-3, and L-5 are also quite similar, and all the current states share a similar characteristic dwell time (Fig. 1E). According to our previous studies, there is a strong correlation between the physical size of the translocated molecules and I/I_0 values (28); therefore, L-1 and L-5 can be likely interpreted as the blockade caused by the added volume of the di-deoxyadenosine moieties at the two terminals, while similarly L-3 to the bulky (2,2-dipropargyl)-propyl-phosphate (bit-1). The two lighter *n*-propyl-phosphates flanked between these bulkier groups (Fig. 1B) contribute instead to higher L-2 and L-4 current states. Therefore, the strategy of including nucleotides in our informational polymer not only prolonged the dwell time but also enhanced the potential resolution of the system to single-bit precision.

Effects of terminal nucleobases on aerolysin capability for polymer reading

To optimize the polymer design and further understand the influence of the terminal nucleotides for decoding, a series of polymers that replaced the terminal di-deoxyadenosine groups with other types of dinucleotides were tested. According to our previous studies, due to backbone stereochemistry, single-stranded DNA prefers to enter the aerolysin pore from the 3'-end (34) (thus all polymers are oriented starting from the 3'-end; Fig. 1B). Translocation events of these polymers showed a qualitatively similar five-level signal (Fig. 2A). While L-1 values for AA00100AA and AA00100CC are nearly identical, in CC00100AA, the relative current is higher, demonstrating eventually that the first current blockade is associated to the 3' nucleobase and its chemical nature. Among all polymers, L-1 and L-5 of CC00100CC are the highest peaks, which further supports the hypothesis that the first and last current levels are induced by nucleobases at the terminals. Therefore, aerolysin nanopores are able to read not only tailor-made informational polymers but also different types of DNA bases and their order at the terminals. To systemically evaluate the enhancement of different dinucleotides for separation between bit-1 and bit-0 monomers, we compared the mean dwell time

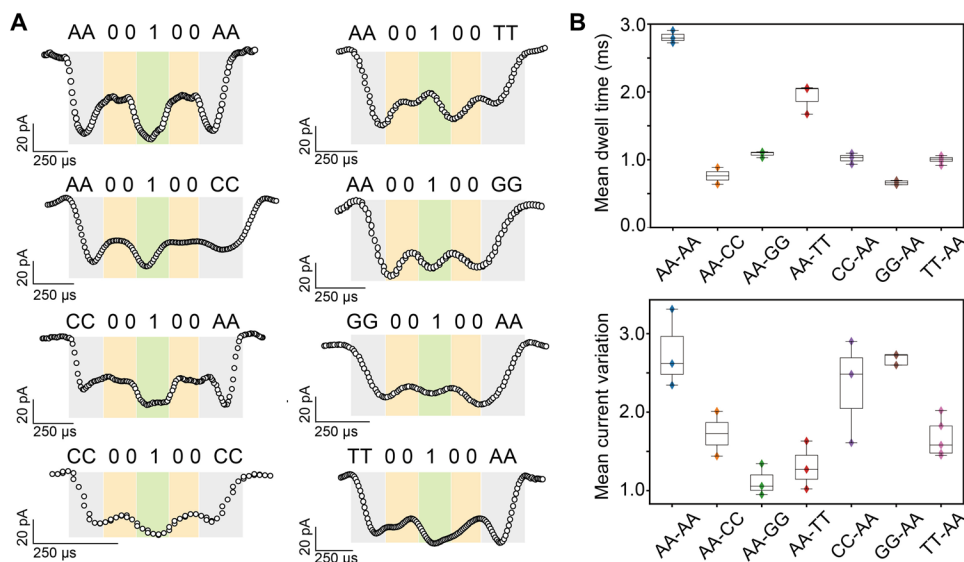


Fig. 2. Effects of terminal nucleobases on polymer reading. (A) Representative translocation events of the digital macromolecular analyte with different types of nucleotides at the chain termini. Colors scheme as in Fig. 1. (B) Mean dwell time and mean current variation for all polymers with different types of terminal dinucleotides. All data were obtained using 1.0 M KCl, 10 mM Tris, and 1.0 mM EDTA buffer at pH 7.4, applying a bias potential of 100 mV. Each value is an average obtained from at least three separate pore measurements.

(i.e., longer dwell time allows a more accurate determination of blockade current levels) and mean current variation (i.e., higher variation promotes a higher read accuracy for each bit, see Materials and Methods) of all polymers with different terminals (Fig. 2B). As di-deoxyadenosine at both terminals showed the longest dwell time and highest current variation among all polymers, it was chosen as the basic terminal block for the following design.

Decoding polymer sequences by engineered aerolysin pores using deep learning

We then tested the sensitivity of the pore for detecting bit-1 monomers when spanning the five available positions along the *n*-propyl-

phosphate backbone (i.e., AA10000AA, AA01000AA, AA00100AA, AA00010AA, and AA00001AA). For this task, we developed a deep learning approach to process the current signal, which was able to automatically classify a much larger fraction of events (~40%) with high accuracy (~84%; fig. S5). We used long short-term memory (LSTM) (35) recurrent neural network to read the events local extrema followed by a multilayer perceptron (MLP) to classify the polymers (Fig. 3A). These additional results showed that the detection of bit-1 monomers is difficult when flanking directly the terminal nucleobases (fig. S6). When only the innermost positions are considered (i.e., as in AA01000AA, AA00100AA, and AA00010AA), the resulting accuracy of the neural network is as high as 97.6% (Fig. 3B); therefore,

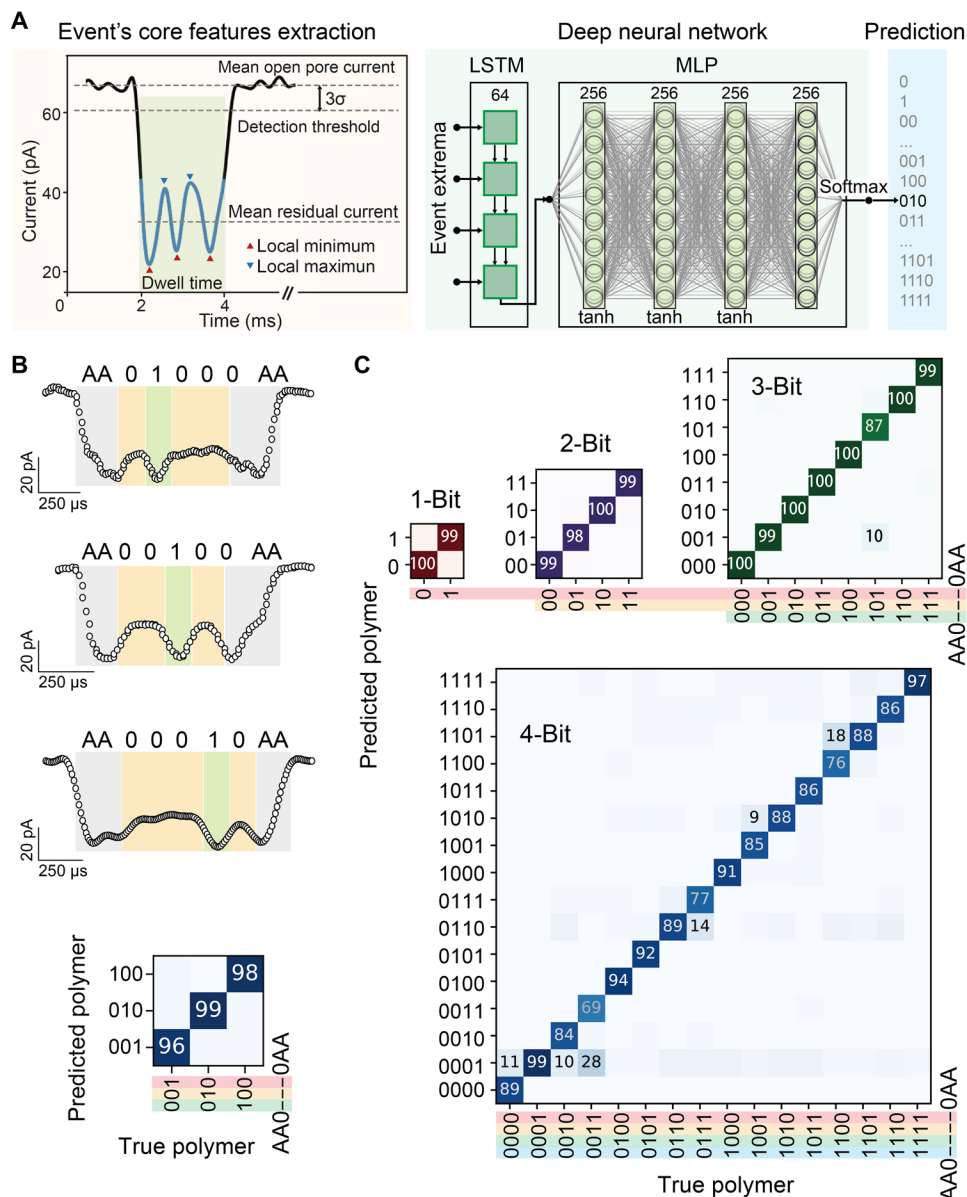


Fig. 3. Decoding polymer sequences using aerolysin pores and deep learning. (A) Details for nanopore signal processing and relative deep learning workflow. (B) Characteristic translocation events of polymers containing bit-1 at different positions, i.e., A01000AA, AA00100AA, and AA00010AA, and the corresponding confusion matrix results obtained by deep learning. (C) Confusion matrix of 1-, 2-, 3-, and 4-bit polymer sequences classification shown for a selection percentage of 10% (see figs. S35 to S38 for dependence on selection threshold). Columns represent true polymers from the test set, while rows are the polymers that deep learning assigned them to. All data were obtained using 1.0 M KCl, 10 mM tris, and 1.0 mM EDTA buffer at pH 7.4, applying a bias potential of 100 mV.

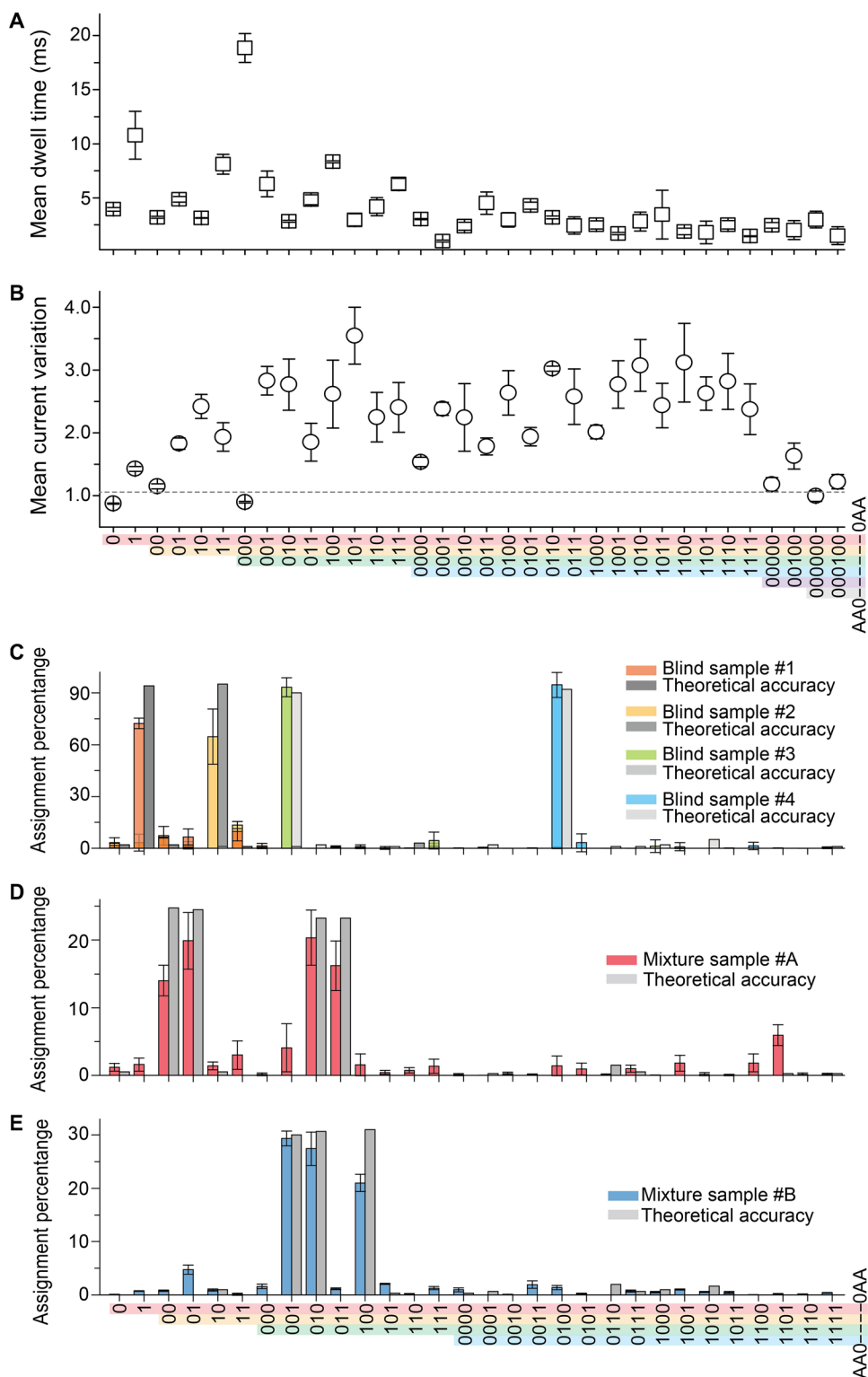


Fig. 4. Statistical analysis and assignment of specific polymer's identity and relative concentration in a mixture. Statistical analysis of (A) mean dwell time and (B) current variation of all polymers. (C) Assignment percentage of blind polymer samples #1 (orange), #2 (yellow), #3 (green), and #4 (blue), respectively. Assignment percentage of mixture sample #A (D) and mixture sample #B (E). The theoretical accuracy is shown by the gray columns. All data were obtained using 1.0 M KCl, 10 mM tris, and 1.0 mM EDTA buffer at pH 7.4, applying a bias potential of 100 mV.

these positions were only used for single-bit reading, hereafter indicated in bold (e.g., AA0**0000**AA). We tested between AA0- and -0AA flanking terminals the polymers encoding for 1 to 4 bits of information generating a library of 30 different polymer sequences (i.e., $2^1 + 2^2 + 2^3 + 2^4$; Fig. 3C and figs. S7 to S34). While each of these 30 polymers were measured by a single nanopore independently, we report a reading accuracy of 99.0% for 1-bit polymers, 99.0% for 2-bit, 98.0% for 3-bit, and 86.5% for 4-bit; the accuracy is slightly decreasing for longer polymers that host a larger number of combinations and with higher event selection thresholds (Fig. 3C and figs. S35 to S38).

To better evaluate the reading capability of aerolysin nanopores, we performed a statistical analysis across $N = 546$ separate nanopore measurements of all 30 polymers (~ 6.6 M events in total; Fig. 4A). Notably, few 5-bit and 6-bit polymers were also included to expand the polymer landscape. In general, the averaged dwell time decreased as polymer length increased, tending nonetheless to a value of 2.5 ± 0.8 ms. In particular, if the total coding length is shorter than 3 bits, the dwell time is longer, likely due to the presence of a less abundant negatively charge density per bit, which is steered slower by the applied voltage; while dwell times tend to converge to higher value when the polymer length is longer than 4 bits, and charge density is less affected by the terminal groups (Fig. 4A). On the other hand, although 4-bit polymer dwell times decreased, their current variation increased (Fig. 4B), indicating that more information is encoded in the longer polymers.

On the basis of the model generated by deep learning, the expectation is that any item in the library of polymer sequences can be identified directly with high confidence. To test this hypothesis, we performed blind tests to identify the given polymers and their relative concentration when mixed. Following this blind procedure, we were able to correctly detect polymer “AA0**10**AA” among all 30 polymers, with a percentage of $72.0 \pm 3.0\%$ (Fig. 4C, orange), which is close to the predicted accuracy of the deep learning model (94.0%, fig. S39). Similarly, we correctly assigned polymer “AA0**100**AA” with an accuracy of $64.0 \pm 5.0\%$ (yellow), “AA0**0010**AA” with an accuracy of $93.2 \pm 5.2\%$ (green), and “AA0**01000**AA” with an accuracy of $94.5 \pm 4.2\%$ (blue). The high-assignment percentages are consistent with the prediction accuracy predicted by deep learning (i.e., 95.0, 90.0, and 92.0%, respectively). We then measured a mixture sample (#A) in which polymers AA0**000**AA, AA0**10**AA, AA0**100**AA, and AA0**110**AA were blindly mixed at equimolar ratio, recapitulating the composition with an accuracy of $14.0 \pm 2.0\%$, $20.0 \pm 4.0\%$, $20.0 \pm 4\%$, and $16.0 \pm 4\%$, respectively (Fig. 4D). This is similar to the accuracy for a 30-polymer classification given an equimolar concentration ratio of the four polymers (i.e., 24.7, 24.5, 23.2, and 23.2%). Furthermore, a second mixture (#B) containing AA0**0010**AA, AA0**0100**AA, and AA0**1000**AA in an equimolar ratio was tested, and an approximately equal assignment was observed (i.e., $29.4 \pm 1.4\%$, $27.4 \pm 3.1\%$, and $21.0 \pm 1.6\%$), similar to the predicted accuracy obtained by deep learning (30.0, 30.6, and 31.0%; Fig. 4E).

Conclusion

In conclusion, we have demonstrated that tailor-made informational polymers can be efficiently decoded by using a specific variant of the aerolysin pore (K238A). In particular, the design of an optimal bio-inspired writing-reading framework allowed for single-bit resolution, which is unprecedented in analytical chemistry. The aerolysin pore structure can, in principle, be further tuned by engineering

its main two sensitive constrictions to optimize the translocation for efficient reading. On the other hand, the vast chemical space accessible to informational polymers can be further explored to enhance optimal decoding by biological nanopore. Informational polymers hybridized with DNA nucleobases keep some of the advantages of synthetic DNA used as support for data storage (36). For instance, different terminal nucleobases, which allow for more efficient capture by the nanopore, can be readily discriminated (Fig. 2), opening the possibility to use canonical DNA bases to define data structure in a format that can enable random access (37–39).

Writing-reading digital data using this biological-inspired nanopore-based platform can offer numerous advantages. First, single-bit resolution on the proposed informational polymer theoretically provides the opportunity to increase the information density of existing DNA-based solutions. For instance, the best resolution in solid-state nanopore platform for digital reading of dsDNA is 8-bps difference for an average mass of ~ 5 kDa (17), whereas the mass difference between different bit units in our informational polymers is ~ 70 -fold lower (76 Da). Similarly, the minimum distance between two bits in solid-state nanopore solutions is 76 bps, while in the aerolysin framework, we can achieve the molecular equivalent of single-base resolution (140 Da, Fig. 2). Second, nanopore sensing does not require additional labeling, and there is no theoretical upper limit for the reading length (40), further reducing the overall cost and workflow time. Nanopore sensing, which relies on an electrical readout, naturally enables large-scale parallelization based on already established technologies (41), thus allowing the construction of more affordable and portable devices for data management.

Toward practical polymer-based data storage, the next important challenge will entail sequencing longer polymers for which the translocation mechanism has to be optimized to preserve as much as possible single-bit resolution and information density. Possible solutions include (i) reducing negative charge density to obtain longer dwell times for precise reading, (ii) optimizing interbit separation to eliminate the influence from neighboring subunit, and (iii) increasing the rigidity of the polymer backbone to reduce the signal noise. One benefit of informational polymers is that they can be better tailored on the properties of the reader; therefore, new definitions of fundamental bit units and blocks, as well as their sequential assembly or controlled fragmentation, can be designed ad hoc to achieve efficient data storage and retrieval (33, 42, 43). In summary, this bio-inspired platform based on hybrid DNA-polymer analytes that encode digital information read by a biological nanopore sensor opens up new promising perspectives for polymer-based memories, with important advantages for ultrahigh density, long-term storage, and device portability.

MATERIALS AND METHODS

Synthesis of the macromolecular analytes

The polymers used in the nanopore experiments were synthesized by automated phosphoramidite chemistry on an Expedite DNA synthesizer (Perceptive Biosystem 8900), as previously described (30, 31). All polymers were characterized by electrospray ionization–high resolution mass spectrometry (HRMS) (table S1 and figs. S40 to S82), and their purity was controlled by anion-exchange high-performance liquid chromatography (figs. S83 to S85), on an Agilent Apparatus equipped with a column Dionex BioLC DNAPac-PA100 and ultraviolet detectors (260 and 280 nm).

Aerolysin productions

The aerolysin full-length sequence was cloned in the pET22b vector with a C-terminal hexahistidine tag to aid purification as described in our previous work (19). The QuikChange II XL Kit from Agilent Technologies was used for performing site-directed mutagenesis on the aerolysin gene, following manufacturer's instructions. The recombinant protein K238A was expressed and purified from BL21 DE3 pLys *Escherichia coli* cells. Cells were grown to an optical density of 0.6 to 0.7 in Luria-Bertani media. Protein expression was induced by the addition of 1 mM isopropyl β -D-1-thiogalactopyranoside (IPTG) and subsequent growth over night at 20°C. Cell pellets were resuspended in lysis buffer [20 mM sodium phosphate (pH 7.4) and 500 mM NaCl], mixed with cOmplete Protease Inhibitor Cocktail (Roche), and then lysed by sonication. The resulting suspensions were centrifuged (12,000 rpm for 35 min at 4°C), and the supernatants were applied to an HisTrap HP column (GE Healthcare) previously equilibrated with lysis buffer. The protein was eluted with a gradient more than 40 column volumes of elution buffer [20 mM sodium phosphate (pH 7.4), 500 mM NaCl, and 500 mM imidazole], and buffer was exchanged into final buffer (20 mM tris and 500 mM NaCl at pH 7.4) using a HiPrep Desalting column (GE Healthcare). The purified protein was flash-frozen in liquid nitrogen and stored at -20°C.

Single-channel recording experiments

Phospholipid of 1,2-diphytanoyl-*sn*-glycero-3-phosphocholine powder (Avanti Polar Lipids Inc., Alabaster, AL, USA) was dissolved in octane (Sigma-Aldrich Chemie GmbH, Buchs, Switzerland) for a final concentration of 1.0 mg per 100 μ l. Purified K238A aerolysin mutant was diluted to the concentration of 0.2 μ g/ml and then incubated with Trypsin-agarose (Sigma-Aldrich Chemie GmbH, Buchs, SG Switzerland) for 2 hours under 4°C temperature to activate the toxin for oligomerization. The solution was finally centrifuged to remove trypsin.

Nanopore single-channel recording experiments were performed by Orbit Mini equipment (Nanonion, Munich, Germany). Phospholipid membranes were formed across a MECA 4 recording chip that contains a 2 \times 2 array of circular microcavities in a highly inert polymer. Each cavity contains an individual integrated Ag/AgCl microelectrode and is able to record four artificial lipid bilayers in parallel. The measurement chamber temperature was set to 25°C for all experiments.

Polymers in powder form were prediluted in water, to a stock concentration of 2.0 mg/ml, and added to the *cis* side of the chamber in 1.0 M KCl solution buffered with 10 mM tris and 1.0 mM EDTA (pH 7.4) to the final concentration of 20 μ mol. All experiments shown here were repeated in 3 to 15 different pores.

Current signal processing

The raw signals are segmented on the basis of voltage discontinuities and large time-scale discontinuities to separate the signals segments where the pore is blocked or where a second pore is inserted into the membrane. For each segment, the open pore current distribution is measured by fitting a Gaussian function on the peak distribution of current with the highest mean current. The signal segments with an open pore current distribution of mean between 67 to 98 pA and SD between 1.5 to 4.2 pA are kept.

The events are extracted using a current threshold at 3σ from the open-pore current distribution (Fig. 3A). The relative current $I_{rel} = \frac{I}{I_0}$

is computed from the mean open pore current (I_0). The cores of the events are extracted by removing the current drop at the beginning and end of the events using an adaptive current threshold. The dwell time, average relative current, current variation $\sigma_{rel} = \frac{\sigma}{\sigma_0}$ (σ_0 is the value of the open pore current SD, and σ is the residual current SD), and local extrema are computed. The events are selected on the basis of the dwell time (0.4 to 30.0 ms) and the average relative current (15 to 60%) discarding the events that are too short or too long and removing the outliers. In average, this initial filtering procedure discards ~10% of the events (fig. S4).

To detect and label different level in the signal, the local relative current extrema are used to generate a Gaussian mixture model (GMM) with three components: low, high, and transition level. The low and high Gaussian models correspond to the two main modes of the relative current extrema distribution. The transition level describes possible change of state between high and low level. Each event is segmented into low, high, and transition levels of based on the level type with the highest probability predicted by the GMM. Last, the transition levels, which are not transition between high and low such as high-transition-high and low-transition-low, are merged into a single high and low level, respectively.

Last, to classify the current events, a machine learning approach was devised including two steps. The first one is the classification of every events, and the second is the assessment of the quality of the prediction of the classifier (Fig. 3A). The neural network architecture for both the classification and the assessment is an LSTM (35) neural network followed by an MLP using the position in time and relative current of the local extrema for each event as input features. The features are rescaled by a fixed factor to decrease the training time. The classifier is composed of a LSTM with state size of 64 without any activation function, followed by a four fully connected hidden layers of size 256 with hyperbolic tangent as activation functions, and, last, an output layer of size 30 with softmax activation function. The neural networks for the classification and assessment are trained together using a three-part loss functions. The first part is the full-classification cross-entropy loss of the predictions from the classifier and the polymers label. The second part is the assessment cross-entropy loss between the predicted and actual prediction validity from the classifier. The third part is the reinforcement classification loss, which is the full classification cross-entropy loss scaled by the assessment prediction.

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at <http://advances.sciencemag.org/cgi/content/full/6/50/eabc2661/DC1>

[View/request a protocol for this paper from Bio-protocol.](#)

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