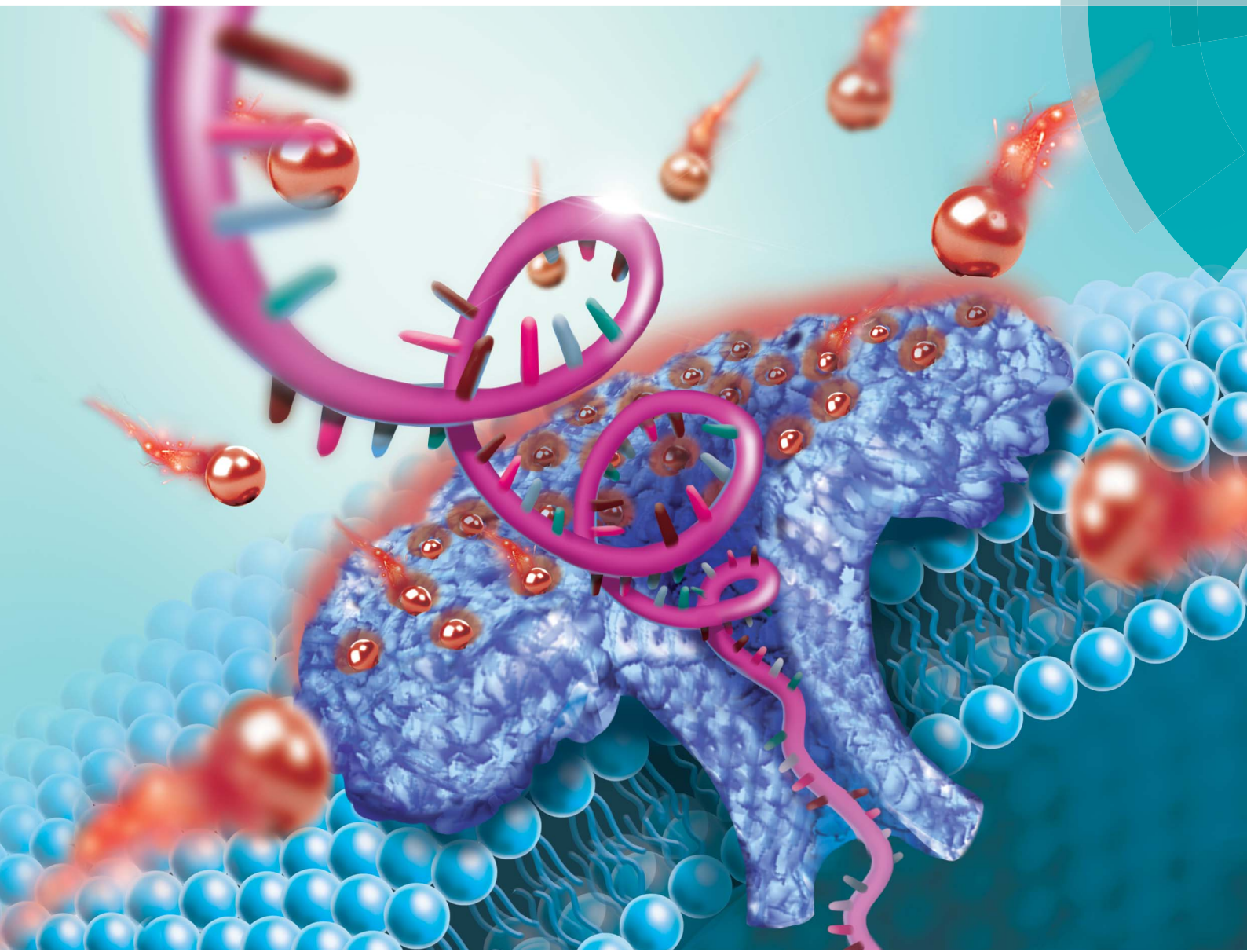


Chemical Science

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ISSN 2041-6539



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EDGE ARTICLE

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A lithium-ion-active aerolysin nanopore for effectively trapping long single-stranded DNA


 Cite this: *Chem. Sci.*, 2019, **10**, 354

All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 3rd September 2018
 Accepted 12th November 2018

DOI: 10.1039/c8sc03927e
rsc.li/chemical-science

Introduction

Nanopore techniques have been developed into powerful platforms for single molecule sensing in a rapid and low-cost way. Aerolysin (AeL), a rivet-shaped protein nanopore with an inner diameter of ~ 1.0 nm, was initially used to investigate the structural variation of α -helical peptides,¹ then applied to discriminate polymer sizes, study protein folding/unfolding, describe enzyme kinetics and monitor chemical bond making/breaking.²⁻⁶ In particular, AeL demonstrated its superior resolution in discrimination of very short oligonucleotides (≤ 10 nt).⁷⁻⁹ The high sensitivity and long reads are the two important parameters for DNA sensing and sequencing. Compared with other biological nanopores such as α -hemolysin (α -HL) and *Mycobacterium smegmatis* porin A (MspA),¹⁰⁻¹³ AeL has high sensitivity in direct discrimination of single nucleotides without chemical labelling, the incorporation of molecular motors or recognition probes.^{9,14} Previous work showed that AeL was able to recognise subtle differences between methylated cytosine and cytosine under serum conditions.¹⁴ Therefore, AeL shows great potential for direct DNA sensing and sequencing in a simple and cheap manner. However, all these achievements were based on very short oligonucleotides (≤ 10 nt). Few real applications have been achieved *via* AeL because it is difficult for long single-stranded DNA (ssDNA, > 10 nt) to enter

A lithium-ion-active aerolysin nanopore for effectively trapping long single-stranded DNA†

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Wild-type aerolysin (AeL) nanopores allow direct single nucleotide discrimination of very short oligonucleotides (≤ 10 nt) without labelling, which shows great potential for DNA sensing. To achieve real applications, one major obstacle of AeL is its poor capture ability of long single-stranded DNA (ssDNA, > 10 nt). Here, we have proposed a novel and robust strategy for the electrostatic focusing of long ssDNA into a lithium-chloride (LiCl)-active AeL. By using this method, for the first time we have demonstrated AeL detection of ssDNA longer than 100 nt. Due to screening more negative charges, LiCl improves AeL capture ability of long ssDNA (i.e. 60 nt) by 2.63- to 10.23-fold compared to KCl. Further calculations and molecular dynamics simulations revealed that strong binding between Li^+ and the negatively charged residue neutralized the AeL, leading to a reduction in the energy barrier for ssDNA capture. These findings facilitate the future high-throughput applications of AeL in genetic and epigenetic diagnostics.

into the AeL. For instance, AeL analysis of a 50 nt ssDNA shows an event frequency as low as $\sim 0.8 \text{ s}^{-1}$ at 100 mV at a concentration of $2.0 \mu\text{M}$.¹⁵ Most recently, our study demonstrated the discrimination of A, T, C and G in a 14 nt oligonucleotide but showed a low AeL capture rate of the ssDNA.¹⁶ This challenge largely restricts nanopore techniques from achieving direct sensing of long stranded DNA. To improve the capture ability of AeL for long stranded DNA, several strategies have been proposed. By increasing the applied voltage to a value as high as 70 mV, a 60 nt ssDNA showed an increased event frequency to 0.4 s^{-1} .¹⁷ In our previous work, by modulating the G-quadruplex DNA into an extended form using magnesium ions (Mg^{2+}), the subsequent capture of a 30 nt structured DNA was observed at an event frequency of 0.6 s^{-1} .¹⁸ By raising the temperature from 20 to 60 °C, the event frequency of a 50 nt ssDNA increased from 0.8 to 2.2 s^{-1} .¹⁵ By protonating AeL in an asymmetric pH environment, the translocation of a 16 nt ssDNA was improved to 5.6 s^{-1} .¹⁹ However, these methods increase the difficulty of the experiment and reduce the stability of the sensing system. The high voltage destabilizes the lipid bilayer membrane, and the high temperature causes variation in the salt concentration due to the non-negligible evaporation of water and might irreversibly denature the protein nanopores. The asymmetric pH environment induces spontaneous current blocks for AeL at applied voltages higher than 100 mV.¹⁹ Thus, a novel and robust approach is urgently needed to improve AeL's ability to capture long stranded DNA.

According to previous studies,¹⁹⁻²⁴ various possible effects could modulate DNA translocation dynamics, such as the geometry of the nanopore, the pH of the solution, the salt type and the electrolyte concentration gradient. These effects mainly

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† Electronic supplementary information (ESI) available: Protocol for aerolysin nanopore experiment, molecular dynamics simulations, histograms of the event frequencies and durations, and supporting figures. See DOI: [10.1039/c8sc03927e](https://doi.org/10.1039/c8sc03927e)



regulate the energy barrier for the entry of DNA, in particular they reduce the interaction between DNA and the entrance of the nanopores. As for aerolysin nanopores, they emerge as negatively-charged protein nanopores ($-52e$) with numerous aspartic (Asp) and glutamic (Glu) residues around the *cis* entrances, such as Asp216, Asp207, Glu307 and Glu415.²⁵ The strong electrostatic repulsion between the negatively charged nucleotides and the amino acids at the entrance contributes to a large energy barrier for DNA entering into the AeL. Therefore, a potential solution is to reduce the negative charges at the *cis* entrance of the AeL. The local charge distribution of a biological nanopore could be changed using site-directed mutagenesis and chemical modification.^{26–28} However, these methods are always complicated and time-consuming. Previous studies have demonstrated that lithium ions (Li^+) exhibit a specific preference over other alkali metal ions due to their high charge density and small size.^{23,29} As Li^+ shows a stronger binding affinity and a longer binding time with charged residues (*i.e.* Asp) compared with other alkali metal ions,³⁰ it could be used to screen the negative charges of a biological nanopore. Herein, we adopted lithium chloride (LiCl) to improve AeL's ability to capture long ssDNA. It has been reported for the first time that AeL detection of a 102 nt DNA (ssDNA_102) could be achieved in LiCl. Compared with KCl, LiCl enhanced the capture ability of AeL for ssDNA_102 by 3.89 to 9.07 fold.

Results and discussion

First, we demonstrated that the presented LiCl sensing system shows high stability in a wide voltage range (Fig. S1†). Fig. 1 illustrates a scheme of the nanopore experiment, depicting an individual ssDNA molecule captured in an AeL nanopore in M^+Cl^- ($\text{M}^+ = \text{Li}^+$ or K^+). For both electrolytes, we added the ssDNA in the *cis* side of the AeL, where the applied voltage is termed as grounded. Here, we adopted the event frequency as a surrogate of capture ability, which is defined as the number of blockades per second and evaluated through the inverse fitting of interevent time. We performed the assays of ssDNA_102 and obtained its real-time current traces in LiCl and KCl. The duration times of ssDNA_102 decrease as the applied voltage ranges from 100 to 160 mV, indicating the successful translocations of ssDNA_102 in M^+Cl^- ($\text{M}^+ = \text{Li}^+$ or K^+) (Fig. 1). Although the signals exhibit similar full blockades, the event frequencies of ssDNA_102 increase apparently when the electrolyte is changed from KCl to LiCl. The event frequency of ssDNA_102 in LiCl at 100 mV is $2.46 \pm 0.11 \text{ s}^{-1}$, producing a 9.07-fold increase compared to that in KCl ($0.27 \pm 0.04 \text{ s}^{-1}$) (Fig. 1 and S2†). When the applied voltage is increased to 160 mV, the event frequency of ssDNA_102 further increases from $2.26 \pm 0.23 \text{ s}^{-1}$ in KCl to $8.79 \pm 0.54 \text{ s}^{-1}$ in LiCl (Fig. 1 and S2†).

To figure out the enhancement mechanism in LiCl, we studied the behavior of polydeoxyadenines (dA_n) with different lengths in M^+Cl^- ($\text{M}^+ = \text{Li}^+$ or K^+). In LiCl, dA_{20} and dA_{40} exhibit a better linear frequency-voltage dependence, suggesting a more dominant diffusion-controlled capture process (Fig. 2a–c).^{19,31,32} However, dA_{60} shows an exponential frequency-voltage

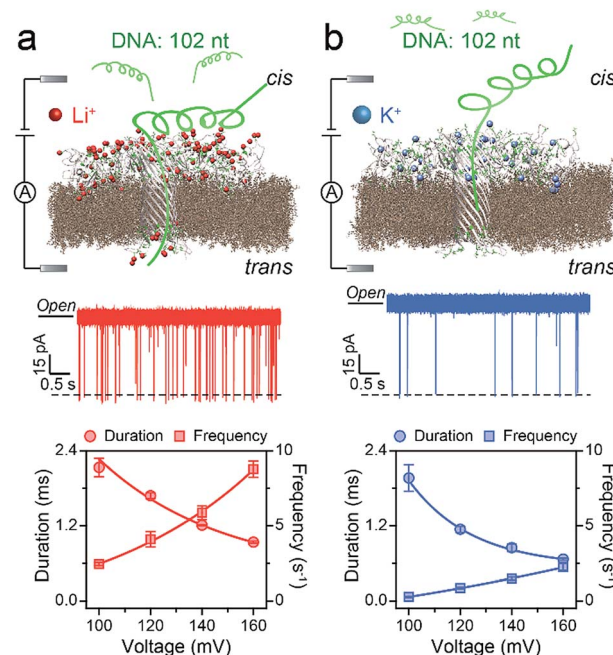


Fig. 1 Illustration of the detection of a 102 nt ssDNA (ssDNA_102) using an AeL nanopore in M^+Cl^- ($\text{M}^+ = \text{Li}^+$ or K^+). Top: the electrostatic attraction in LiCl and the relative electrostatic repulsion in KCl of a single ssDNA_102 molecule into AeL. The illustration is not to scale. Middle: raw current traces of ssDNA_102. More frequent blockade events were observed clearly in LiCl compared with KCl from the real-time current traces. Bottom: voltage dependence of ssDNA_102 translocation. Data in this work were obtained in 1 M M^+Cl^- ($\text{M}^+ = \text{Li}^+$ or K^+), 1 mM EDTA, 10 mM Tris buffer, and pH 8.0 with the presence of 2.0 μM ssDNA at the *cis* side of the AeL nanopore.

relationship, indicating a barrier-limited capture process (Fig. 2c). These results unravel the length-dependent capture of dA_n in LiCl: diffusion is predominantly responsible for short stranded dA_n and an energy barrier is primarily accountable for long stranded dA_n . As expected, similar results were obtained for dA_n in KCl (Fig. 2a–c).⁷ According to previous studies,^{33,34} the energy barrier for DNA capture mainly originates from DNA entropy and DNA-pore electrostatic interactions at the pore entrance. As the length of dA_n increases, entropy likely contributes to a portion of the energy barrier for the capture of long stranded dA_n . Therefore, the event frequencies reduce from 8.75 ± 0.06 to $5.81 \pm 0.31 \text{ s}^{-1}$ with dA_n elongating from 20 to 60 nt at 160 mV in LiCl due to an increased energy barrier. Interestingly, LiCl induces higher event frequencies for dA_n ($n = 20, 40, 60$) than KCl at the same applied voltage and the same DNA concentration. LiCl caused a 2.63- to 10.23-fold enhanced capture of dA_n ($n = 20, 40, 60$) compared to that obtained in KCl. The possible explanation could be a lower energy barrier at the *cis* entrance of the AeL for dA_n capture in LiCl than in KCl. As Li^+ screens more negative charges than K^+ , we deduced that the lower energy barrier in LiCl results from weaker DNA-AeL electrostatic repulsions at the *cis* entrance of the pore.

To demonstrate this speculation, we evaluated the electrolyte-dependent difference in energy barriers for dA_{60} , which shows the barrier-limited capture both in LiCl and KCl.



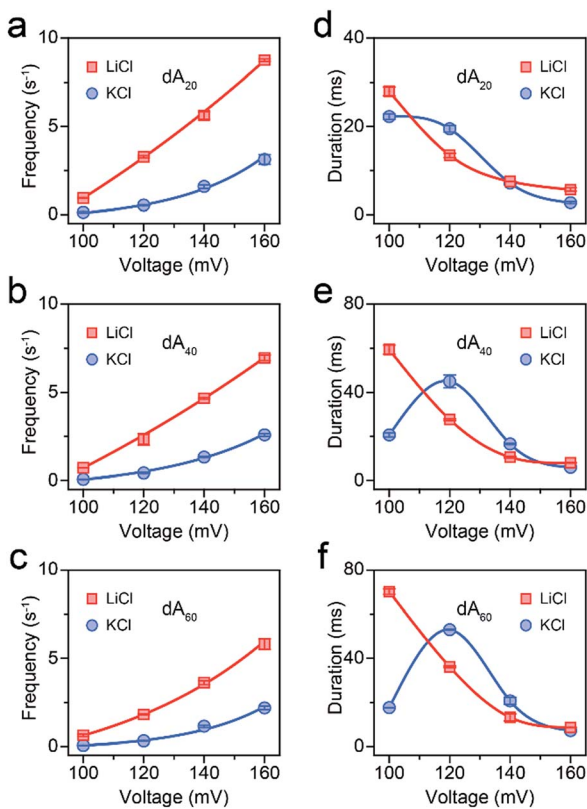


Fig. 2 AeL detection of dA_n with different lengths in M^+Cl^- ($M^+ = Li^+$ or K^+). Voltage-dependent (a–c) event frequencies and (d–e) blockade durations of dA_n ($n = 20, 40, 60$) in LiCl (red squares) and KCl (blue circles). The values of event frequencies and blockade durations were evaluated from transformed single-exponential functions (Fig. S3–S5†).

The event frequencies of dA_{60} shown in Fig. 2c are well-fitted with eqn (1):

$$f = f_0 e^{V/V_0} \quad (1)$$

where V is the applied voltage, V_0 is the applied voltage for a charged polymer to overcome Brownian motion, and f_0 is an extrapolated frequency in the absence of the applied voltage.^{33,34} According to the Van't Hoff–Arrhenius law, f_0 was calculated by eqn (2):

$$f_0 = \frac{CDA}{L} \times e^{\frac{-U}{k_B T}} \quad (2)$$

where U is the activation energy, k_B is the Boltzmann constant, T is the temperature, C is the concentration of the analyte in the bulk solution, D is the diffusion coefficient of the analyte, A is the sectional area of the pore and L is the pore length.^{33,34} The values of f_0 for dA_{60} are 5.29×10^{-3} in KCl and 4.65×10^{-1} in LiCl, respectively. Assuming $D(LiCl) \approx D(KCl)$, we obtained a reduction in the activation energy of $4.48 k_B T$ from LiCl to KCl for dA_{60} . Fig. 2d–f display hill-shaped duration-voltage curves and a threshold voltage of 120 mV for dA_n ($n = 20, 40, 60$) in KCl, indicating dA_n ($n = 20, 40, 60$) could not traverse across the AeL in KCl at a potential lower than 120 mV. The duration time for

a successful translocation is composed of the residence time of ssDNA inside the AeL and the translocation time of ssDNA through the AeL. Meanwhile for failed translocation, the duration time is only composed of the residence time of DNA inside the AeL. By increasing the applied voltage, we enhanced the electric driving force to overcome the energy barrier for the successful translocation of DNA.²⁹ As a result, the maximum duration times observed at the threshold voltage of 120 mV were 45.00 ± 2.95 ms for dA_{40} and 52.98 ± 0.26 ms for dA_{60} in KCl. Then, the duration times for dA_n ($n = 20, 40, 60$) decrease for applied potentials above 120 mV (Fig. 2a–c). In line with previous discussion, as Li^+ lowers the energy barrier by screening more negative charges than K^+ , the threshold voltage is greatly reduced for DNA capture in LiCl. Therefore, a monotonic exponential duration-voltage relation was obtained in LiCl for dA_n ($n = 20, 40, 60$) over the whole voltage range. The DNA translocation speed may be modulated by various factors. Note that the duration time of dA_n ($n = 20, 40, 60$) is determined by multiple non-covalent interactions between the pore and analyte. Our previous work revealed that the electrostatic interactions between AeL and ssDNA are not the only crucial factors in determining the DNA translocation speed.²⁸ Due to the small volume of the AeL, only a small amount of Li^+/K^+ can be accommodated inside the confined nanopore space. Therefore, the charge screening of ssDNA and/or AeL induced by counterions does not dominate DNA translocation speed. Synergistic effects including van der Waals forces, hydrogen bonding and geometrical structures will also lead to duration differences for ssDNA, and are under study in our future work.

To better understand the capture enhancement mechanism, we performed molecular dynamics (MD) simulations in M^+Cl^- ($M^+ = Li^+$ or K^+). The cations are defined as being bound within a cutoff distance of 3.1 Å from the heavy atoms of the pore.²³ Fig. 3a illustrates the distributions of the bound Li^+ , viewed from the *cis* side of the AeL (K^+ –AeL binding is shown in Fig. S6†). The binding number of Li^+/K^+ was averaged over a 10 ns MD simulation, fitting into a Gaussian distribution (Fig. 3b). The average binding numbers are 194 for Li^+ and 87 for K^+ (Fig. 3b). Although AeL exhibits an identical net charge in the two different electrolytes, the effective charges change due to the cation–AeL electrostatic interactions (Fig. 3a and S6†). The different Li^+/K^+ –AeL binding induced variation of the electrostatic potential distributions at the *cis* entrance of the pore. Since the Li^+/K^+ ions created a local concentration of positive ions in the pore cap, the pore vicinity is effectively polarized and the electrostatic potential increases.^{31,35} In KCl, AeL presents a highly negative electrostatic potential in a confined region at the *cis* entrance of the pore (Fig. 3c). However, a highly positive electrostatic potential occupies most areas of the pore cap in LiCl (Fig. 3c). As ssDNA is negatively charged, the high positive electrostatic potential facilitates the capture of ssDNA from the *cis* entrance of AeL in LiCl, in good accordance with our previous discussion.

Furthermore, we employed LiCl for different polynucleotides with the same length. The raw current traces of dA_{20} , dT_{20} and dC_{20} show more frequent blockades acquired in LiCl than KCl (Fig. 4a). The event frequencies are 8.75 ± 0.06 s⁻¹ for dA_{20} , 9.19

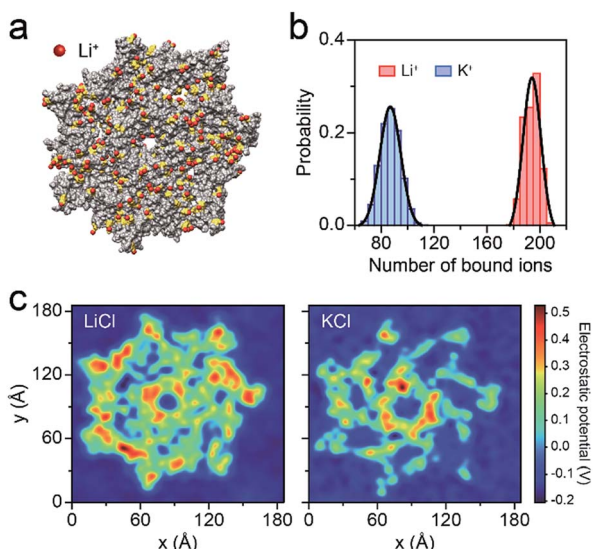


Fig. 3 Molecular dynamics (MD) simulations of AeL in M^+Cl^- ($M^+ = Li^+$ or K^+). (a) Li^+ bound to the AeL in MD simulations. Li^+ are represented in red, the negatively-charged amino acids are shown in yellow and the AeL is represented in gray. The illustration is not to scale. (b) The number of M^+ ($M^+ = Li^+, K^+$) bound to the AeL. The ions are regarded as being bound within 3.1 Å from the heavy atoms of the AeL. The binding numbers of Li^+/K^+ are fitted into a Gaussian function. (c) Average electrostatic potential around the *cis* entrance of the AeL.

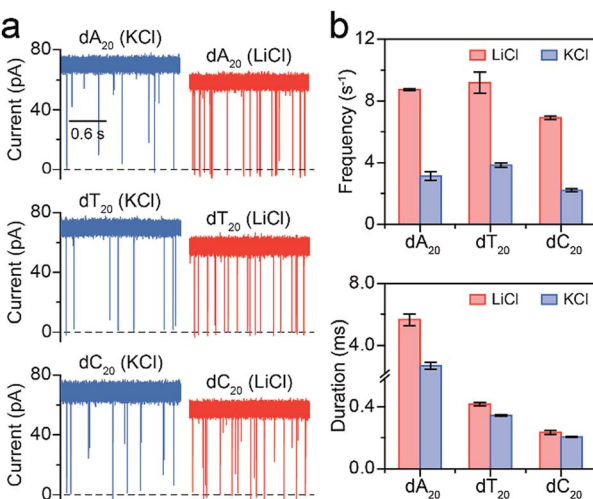


Fig. 4 Analysis of different polynucleotides with the same length in M^+Cl^- ($M^+ = Li^+$ or K^+). (a) Raw current traces of dA_{20} , dT_{20} and dC_{20} . (b) The event frequencies and durations of dA_{20} , dT_{20} and dC_{20} . Event frequencies reveal obvious enhancements for these ssDNA in LiCl. The values of event frequencies and durations were extracted from transformed single-exponential functions (Fig. S7†). Data were obtained at +160 mV in 1 M M^+Cl^- ($M^+ = Li^+$ or K^+), 1 mM EDTA, 10 mM Tris buffer, and pH 8.0 with the presence of 2.0 μ M ssDNA at the *cis* side of the AeL nanopore.

± 0.68 s^{-1} for dT_{20} , and 6.91 ± 0.11 s^{-1} for dC_{20} at 160 mV in LiCl, providing a 2.39-to-3.15-fold increase compared to those in KCl, respectively (Fig. 4a). The difference in event frequencies

might result from the structural variations of dA_{20} , dT_{20} and dC_{20} induced by the cation-ssDNA interactions. The duration times are 5.65 ± 0.38 ms for dA_{20} , 0.42 ± 0.01 ms for dT_{20} , and 0.23 ± 0.01 ms for dC_{20} at 160 mV in LiCl (Fig. 4b). The duration differences in LiCl between dA_{20} and dT_{20} , dA_{20} and dC_{20} , and dT_{20} and dC_{20} are 5.23 ± 0.39 , 5.42 ± 0.39 and 0.19 ± 0.02 ms, respectively. In KCl, the differences between dA_{20} and dT_{20} , dA_{20} and dC_{20} , and dT_{20} and dC_{20} are 2.34 ± 0.24 , 2.48 ± 0.24 and 0.14 ± 0.02 ms, respectively (Fig. 4b). Obviously, these values in KCl are smaller than those gained in LiCl, implying that LiCl provides a higher discrimination ability in the duration time for AeL with polynucleotides. Moreover, we found that LiCl leads to a 1.65-fold increase in event frequency for dA_{40} at 160 mV using an α -HL (Fig. S8†), which is smaller than the 2.69-fold increase obtained by the AeL. It is reasonable as more negative charges were screened at the *cis* entrance of the AeL due to the different charge distributions of the two pores.^{11,25}

Conclusion

In conclusion, we have presented a novel and robust method based on LiCl to enhance AeL capture of long ssDNA. By introducing LiCl, the AeL achieved the detection of a ssDNA longer than 100 nt for the first time. LiCl increased the ssDNA capture ability by 2.63 to 10.23 folds (*i.e.* 60 nt) compared with KCl at the same potential and DNA concentration. The calculations demonstrated that LiCl leads to an experimental reduction in activation energy of $4.48 k_B T$ for dA_{60} in LiCl. Further MD simulations revealed that the strong Li^+ -AeL binding induces a great screen of negative charges on the pore surface, resulting in electrostatic focusing of long ssDNA into the AeL. Our previous work has shown that the event frequency of a 30 nt ssDNA is relatively low ($\sim 0.6 s^{-1}$) in 1 M $MgCl_2$, and the effect of divalent cations might be less favorable than monovalent cations for DNA capture. This general strategy is simple and practical without exposing the analyte and nanopore to extreme conditions (*i.e.* high temperatures and low pH). This work would facilitate various real applications of AeL in life science, such as genetic and epigenetic detection, enzyme activity evaluation and signaling pathway description. These findings also provide new insight into the construction of highly-sensitive AeL arrays for drug separation and screening, and direct DNA sensing and sequencing at high throughput and without labelling.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This research was supported by the “Chen Guang” Project from the Shanghai Municipal Education Commission and Shanghai Education Development Foundation (17CG27), National Natural Science Foundation of China (21421004 and 61871183) and the Fundamental Research Funds for the Central Universities (222201718001, 222201717003, 222201714012).

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