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Single nucleotide detection using bilayer MoS₂ nanopores with high efficiency†

Payel Sen (Da and Manisha Gupta (D*b

Single nucleotide detection is important for early detection of diseases and for DNA sequencing. Monolayer (ML) MoS₂ nanopores have been used to identify and distinguish single nucleotides with good signal-tonoise ratio in the recent past. Here, we use a bilayer (BL) MoS₂ nanopore (~1.3 nm thick) to detect distinct single nucleotides with high spatial resolution and longer dwell time. In this study, the performance of similar sized (<3 nm) ML and BL MoS₂ nanopores for detection of a single nucleotide has been compared. Both single nucleotide and single stranded DNA translocations through them are studied. For single nucleotide detection, we observe that BL MoS₂ nanopores demonstrate twice the dwell time as compared to ML MoS₂ nanopores with 95% confidence. Single nucleotide detection rate for BL MoS₂ nanopores (50-60 nucleotides per s) is five-fold higher as compared to ML MoS₂ nanopores (10-15 nucleotides per s) in 10 pM analyte concentration. For single stranded DNA, we observe 89% (for 60 DNA molecules detected) single nucleotide detection efficiency with BL MoS₂ nanopores as compared to 85% for ML MoS2. The DNA sequencing efficiency through BL MoS2 nanopores is also found to be 8-10% better than through ML MoS₂ nanopores, irrespective of DNA sequencing orientation. Thus, owing to improved analyte/nanopore charge interaction BL MoS₂ nanopores can be used for single nucleotide detection with high resolution due to longer dwell time, detection rate and efficiency. This study demonstrates the improved ability of BL MoS₂ nanopores in sequencing DNA with 8-10% higher efficiency, two-times temporally resolved single-nucleotide current signatures and five-times higher detection rate, compared to ML MoS₂ nanopores.

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Introduction

DNA sequencing data can be used for detecting hereditary diseases based on genetic information and also determine the health effects of a microbial strain. DNA sequencing performed with nanopores is an important technique as it is label free and can be conducted in real-time with relatively low-cost and longread without amplification.1 Solid-state nanopores are used extensively for DNA sequencing due to nanopore stability and ease of fabrication.2-6 However, the sensing efficiency and resolution depend upon a lot of factors including the solid-state membrane thickness, nanopore size and material properties. A variety of materials like silicon, silicon oxide, silicon nitride, aluminium dioxide,10 and hafnium oxide11 have been used for sequencing DNA. In spite of being mechanically stable, sensing suffers from poor spatial resolution making them incapable of properly distinguishing four DNA bases. 2D material nanopores are rapidly emerging as a solution to improve spatial resolution

due to their ultra-thinness for identifying single nucleo-bases. ^{12–27} Molybdenum disulphide (MoS₂) is a type of 2D metal dichalcogenide that has been used in the last decade for nanopore fabrication. These MoS₂ nanopores have demonstrated good DNA sequencing with low sticking of the DNA and high signal-to-noise ratio. ^{12,16,20,21} High molybdenum (Mo) concentration around the MoS₂ nanopores is favorable for the analyte/nanopore wall surface charge interactions as this minimizes the noise. Thus, this is a promising material for nanopore fabrication.

van der Waals force separate the layers of 2D materials like MoS₂. Thus, the number of layers of MoS₂ separated can be controlled, which allows tunability of the membrane thickness. Single nucleotide and polynucleotide translocation through monolayer MoS₂ nanopores has been previously demonstrated experimentally.¹³ However, MoS₂ nanopores also suffer from high translocation velocity similar to other solid-state nanopores. Although different techniques have been employed to slow down DNA translocation through solid state nanopore, ²⁸⁻³¹ the control of translocation speed is still challenging. The thickness of monolayer (ML) MoS₂ (0.65 nm) is less than a single nucleotide (1.6–1.8 nm). Thus, the charge interaction along DNA/nanopore interface for ML MoS₂ gets compromised making DNA translocation fast. Molecular translocation

^aDepartment of Chemical and Materials Engineering, University of Alberta, Edmonton, Canada. E-mail: payel@ualberta.ca

^bDepartment of Electrical and Computer Engineering, University of Alberta, Edmonton, Canada. E-mail: mgupta1@ualberta.ca

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through ML MoS₂ nanopore has been slowed down by using transmembrane viscosity gradient for producing temporally resolved current blockades capable of distinguishing different types of DNA nucleotides.¹³

In this work, we experimentally investigated the role of 2D material membrane properties on the translocation speed of DNA. It is found that the behavior of monolayer (ML) and bilayer (BL) MoS₂ is different under a vertical electrical field applied using the cis and trans electrodes.32 The strong interlayer coupling in the BL MoS2 leads to a gradation in potential throughout the thickness as each layer will experience a different electric potential.³² Due to this potential gradient the phosphate groups are trapped and detrapped by the Molybdenum atoms in the membrane. An additional pull is created because of the increased negative (analyte DNA)-positive (Mo atoms lining the nanopore surface) charge interaction. This can help immobilize molecules at the nanopore for longer time producing highly resolved signals. Molecular dynamics simulation study conducted previously concluded that BL MoS2 can slow down peptide translocation producing stepwise distinguishable current blockades for single nucleotide sensing.22 Thus, we have conducted detailed single and poly nucleotide translocation experimental studies through ML and BL MoS2 nanopores. The translocations were conducted using picomolar concentration and the dwell times were then extracted from these measurements for further analysis.

Experimental

Standard microfabrication techniques were used to fabricate free-standing solid-state silicon nitride (SiN_x) membranes on silicon support. Electron Beam Lithography (EBL) was used along with Reactive Ion Etching (RIE) to fabricate a 50 nm pore at the centre of the previously fabricated SiN_x membrane. Then, monolayer (ML) and bilayer (BL) MoS2 flakes were transferred on the pore via exfoliation using scotch tape. A dual-stage microscope was used to ensure a properly aligned and centred MoS₂ transfer. ESI Fig. S1a[†] presents the detailed fabrication flowsheet of MoS2 nanopores. 20 µm sized MoS2 flakes were obtained repeatedly by this method. The membranes were first characterized to determine and ensure the number of MoS₂ layers using JEOL JEM-ARM200CF Scanning/Transmission Electron Microscope (STEM). The nanopore was STEM drilled on the free standing MoS₂ membrane. Nanopores with diameters between 2.5-3 nm were obtained repeatedly and were suitable for our application. To prevent nanopore expansion or material redeposition due to prolonged beam exposure, intense imaging on the same nanopore was avoided.

300 mM filtered KCl solution at pH = 7 was prepared by adding 3 mM Tris-HCl buffer. The solution was then degassed in vacuum for 90 minutes and used for the experiments after attaining room temperature. 0.5 mM solutions of single nucleotides (deoxyadenosine triphosphate: dA, deoxythymidine triphosphate: dT, deoxycytidine triphosphate: dC and deoxyguanosine triphosphate: dG) were procured from Integrated DNA Technologies for nanopore translocation and dwell-time analysis thereof. 5 mM solution of single stranded (ss)

Custom-made Teflon half-cells were used to contain the electrolyte (KCl). As Teflon or poly-tetra-fluoro-ethylene are both hydrophobic polymers, they help in maintaining a clean environment. They also have high chemical resistance which ensures a well-insulated sensing environment. The half cells were sealed with polydimethylsiloxane (PDMS) to ensure that the electrolytic exchange between the chambers is only via the nanopore. PDMS, also reduces capacitive noise which helps in obtaining improved signals. The half-cells were thoroughly cleaned by ultra-sonication before each experiment. The nanopores/membranes were dipped in acetone and kept in vacuum for 30 min. The nanopore was cleaned by gradually pulling the acetone through it. The cleaning steps include three acetone-cleaning repeats followed by four IPA washes in vacuum (20 min each) to remove acetone residue. Fresh solvent is used for every cycle of acetone and IPA treatment. The membrane with the embedded nanopore was then mounted between the half cells. Two Ag/AgCl electrodes were cleaned using ethanol and DI water and re-chlorinated by treating them with bleach for an hour. The electrodes were then dipped in each half-cell to apply a voltage across the nanopore bearing membrane. The entire cell-assembly was enclosed in a Faraday cage (Warner instruments) to prevent electrical interference. Filtered and buffered 300 mM KCl solution (previously prepared) was introduced in both chambers and the ionic current at varying voltage (-200 mVto 200 mV) was recorded to evaluate the channel conductance. The nanopore was electrically conditioned at a constant 100 mV bias to obtain a stable baseline. The analytes were then added to the trans chamber and pulled electrophoretically to the cis chamber through the nanopore under a constant bias of 200 mV. Single channel blockade recordings obtained from translocating nucleotides and DNA were then filtered (20 kHz 8-pole Bessel filter), amplified (Axon MultiClamp 700B) and finally digitized (Axon Digidata 1550B) for further analysis. ESI Fig. S1b† presents schematic of the sensing assembly.

A customized MATLAB algorithm was used to identify translocation peaks. The raw traces were first loaded as an .abf file. The peaks were identified as change points and the blockade current was calculated by subtracting the mean blockade current from the mean baseline current. The mean was calculated for the 100 samples before and after the peak to account for shift in baseline during the translocation. The dwell time was calculated from the difference of the initial and final changepoint indices multiplied by the sampling frequency. The mean and median of the dwell time were then compared to test whether the distribution is normal or not. We then assumed a null hypothesis that the dwell times for ML and BL MoS₂ nanopore is the same and then tested the hypothesis by conducting T-test at 95% confidence on the dwell time data. All statistical analysis was performed for a population of 300 nucleotides. For the mixed DNA, a MATLAB algorithm was built to first truncate the data at each

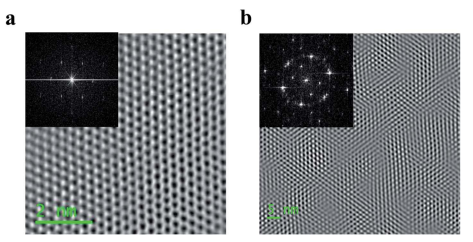


Fig. 1 (a) HRTEM image of a typical exfoliated free-standing ML MoS_2 membrane on SiN_x support with the inset diffraction pattern showing a single hexagon corresponding to monolayer and (b) HRTEM image of AA' stacked BL MoS_2 membrane with the inset diffraction pattern showing two hexagonal domains twisted by 60° (image was filtered to remove noise).

molecular entry and exit. Then the dwell time and blockade current at each individual current drop steps within a molecular translocation event were obtained. The individual nucleotides were then identified by comparing the blockades with that obtained from single nucleotide translocation performed earlier through the same nanopore.

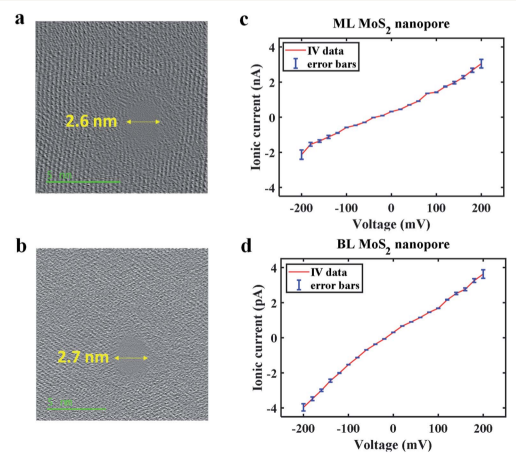


Fig. 2 (a) HRTEM image of 2.6 nm STEM fabricated ML MoS_2 nanopore, (b) HRTEM image of a 2.7 nm STEM fabricated BL MoS_2 nanopore, (c) IV plot for ML MoS_2 nanopore showing an ionic conductance 10.5 nS and (d) IV plot for BL MoS_2 nanopore showing an ionic conductance 17.2 nS. The ionic conductance is determined from the average of the ratio of ionic current to bias value at each voltage. The error bars shown in IV plots correspond to the deviation obtained from 4 pore conductance measurements.

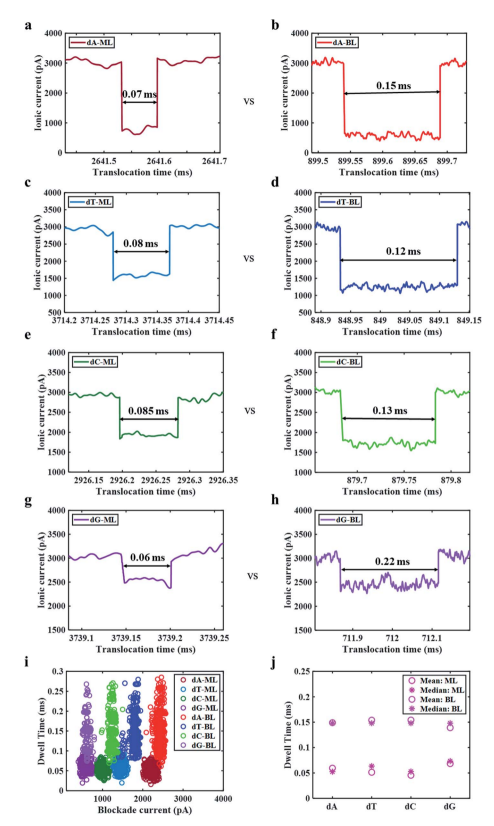


Fig. 3 Truncated typical single nucleotide peaks for (a and b) dA, (c and d) dT, (e and f) dC and (g and h) dG translocations through ML and BL MoS_2 nanopores respectively at 200 mV bias showing characteristic dwell time and blockade current values, (i) Scatter plots of 3000 translocation events showing four different levels for each nucleotide for both ML and BL MoS_2 nanopores and distinctly higher dwell times for BL MoS_2 compared to ML MoS_2 nanopores and (j) plots showing mean and median dwell time values to be similar for each of the four nucleotides through ML and BL MoS_2 nanopores, suggesting a normal distribution.

Results

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The free-standing ML and BL MoS2 membranes were first characterized by High Resolution TEM (HRTEM) microscopy to confirm number of layers. Fig. 1a shows STEM image of molecular arrangement of a ML MoS₂ membrane. Fig. 1b shows STEM image of an AA' stacked BL MoS2 membrane showing Moiré pattern of two overlapped domains rotated by 60°. The TEM structure obtained for ML MoS2 and Moiré patterns observed for BL MoS₂ are in agreement with the literature.³³⁻³⁵ All the HRTEM images are low pass filtered using Gatan Microscope software to remove noise for prominent visualization of the molecular structure. No additional artifacts were produced from the simple filtering. ESI Fig. S2a and b† shows the profilometric studies of ML and BL MoS₂ on SiN_r support showing the corresponding membrane thicknesses. ESI Fig. S2c and d† shows additional STEM images of the MoS2 membranes.

The ionic conductance of the fabricated ML and BL MoS₂ nanopores were measured following the procedure described in the methods section to ensure properly cleaned and conducting nanopores. Fig. 2a and b shows MoS₂ nanopores with diameters of 2.6 nm for ML and 2.7 nm for BL respectively. Ionic current through each nanopore was measured by applying a trans bias in the range of -200 mV to 200 mV in 300 mM KCl. 4 consecutive ionic conductance measurements were conducted through the nanopore along with intermediate cleaning steps. Here, we have reported the average conductance for each pore. Fig. 2c and d shows the IV plot for both the nanopores. The mean of ionic current recorded at each bias is displayed as scatter plot, with the error bars showing the current variation between the 4 measurements at each voltage. The mean nanopore conductance (calculated as the average of the ratio of ionic current and voltage at each bias value) was found to be 10.5 \pm 0.016 nS for 2.6 nm ML MoS $_2$ nanopore and 17.2 \pm 0.023 nS for 2.7 nm BL MoS₂ nanopore.

To study the effect of ML and BL MoS₂ on single nucleotide translocation and sensing, 10 pM concentration of dA, dT, dC and dG were electrophoretically pulled through the abovementioned ML and BL MoS₂ nanopores. For these nucleotide translocations 200 mV bias was applied and the translocation traces were recorded. The ESI Fig. S3a-h† shows the recorded data for single nucleotide (for each of dA, dT, dC and dG) translocation for 5 s and 1.5 s through ML MoS2 and BL MoS2 nanopores respectively. We also observe a detection rate of 10-15 nucleotides per s and 50-60 nucleotides per s for ML and BL MoS₂ nanopores respectively (see Fig. S3a-h†). The five-fold higher

detection rate obtained for BL nanopores may be due to faster capture of the translocating nucleotides at the nanopore due to stronger electrophoretic pull towards the layered BL MoS₂.

Fig. 3a-h presents truncated single nucleotide peaks obtained for ML and BL MoS₂ nanopores for a direct comparison of dwell times. It is observed that the dwell times are higher for BL as compared for the ML MoS₂ nanopores for all the different nucleotides. Blockade current is plotted as a function of dwell time for 3000 single nucleotide transport events in Fig. 3i. We observe four distinct blockade current regions for the different nucleotides. Thus, we can conclude that both ML and BL MoS₂ nanopores are capable of detecting single nucleotides. The blockade current for the nucleotide translocation is plotted as histograms to observe their distribution (ESI Fig. S4†). We observe normal distribution for all the nucleotides for both ML and BL nanopores. Thus, the mean blockade current values along with their standard deviations can be obtained.

Table 1 lists the dwell time and blockade current values for all single nucleotides sensed using ML and BL MoS₂ nanopores. The blockade current magnitudes measured are found to be the dA, dT, dC and dC in decreasing order which agrees with previous study.13 The mean blockade current obtained through BL MoS₂ is found to be slightly higher than or comparable to ML MoS₂, which may be due to better charge interaction at the van der Waals separated nanopore interface. Based on these numbers we can conclude that the current resolution obtained for ML is preserved in BL MoS₂ nanopores. On the other hand, it was observed that the dwell time values of the translocations for BL nanopore are spread over a larger range as compared to the ML nanopore. The events detected through ML nanopores show mean dwell times in range of 0.053-0.061 ms with a 17-20% of deviation. For BL nanopore detection the mean dwell time range is 0.141-0.143 ms with a deviation of 28-30% (see Table 1 and ESI Fig. S5†). The mean and median dwell time values were then evaluated for ML and BL nanopores (see Fig. 3j) and the values were found to be pretty close to each other, supporting an assumption of a normal distribution. Therefore, T-tests were done on the values for a statistical comparison. T-Tests thus performed, indicate that for all the nucleotides, the dwell times for ML and BL nanopores are different at a 95% confidence level (shown in ESI Table T1†). The sample mean of the dwell time for the BL nanopore is approximately twice that of the ML nanopore. A higher dwell time indicates an improvement in sensing.

To demonstrate the single nucleotide detection capability of ML and BL MoS₂ nanopores mixed nucleotide translocations through ML and BL MoS₂ nanopores were conducted. Fig. S6a

Table 1 Blockade current and dwell time for the single nucleotide translocation for both ML and BL MoS₂ nanopores along with the deviation

Analyte	ML MoS ₂ nanopore		BL MoS ₂ nanopore	
	Blockade current	Dwell time	Blockade current	Dwell time
dG	$0.595 \pm 0.057 \text{ nA}$	$0.0529 \pm 0.0092 \; ms$	$0.608 \pm 0.036 \; \text{nA}$	$0.1409 \pm 0.0424 \text{ ms}$
dC	$0.987 \pm 0.066 \text{ nA}$	$0.0577 \pm 0.0102 \text{ ms}$	$1.256 \pm 0.048 \; \mathrm{nA}$	$0.1430 \pm 0.0424 \text{ ms}$
dT	$1.454 \pm 0.076 \; \mathrm{nA}$	$0.0610 \pm 0.0110 \text{ ms}$	$1.809\pm0.07~\mathrm{nA}$	$0.1422 \pm 0.0405 \text{ ms}$
dA	$2.245 \pm 0.079 \; \mathrm{nA}$	$0.0615 \pm 0.0122 \text{ ms}$	$2.434\pm0.06~\mathrm{nA}$	$0.1412 \pm 0.0401 \; ms$

and b† show translocation traces for 5 s and 1.5 s through ML and BL MoS_2 nanopores respectively. The nucleotide distinction is made based on the current blockades obtained for individual nucleotides sensed separately. The dwell time range for all nucleotides (as observed in the scatter plot in Fig. S6c†) is found to be the same as that when detected separately through ML and BL MoS_2 nanopores.

To demonstrate that the BL MoS₂ improves the dwell time of the translocation and can detect single nucleotides, we used 10 pM single stranded (ss) DNA oligos with customized mixed sequence. The translocations were conducted using ML and BL nanopores. Fig. 4a and b show the translocation trace for a single ssDNA strand through the ML and BL MoS₂ nanopore respectively. For both the nanopores distinct molecular current

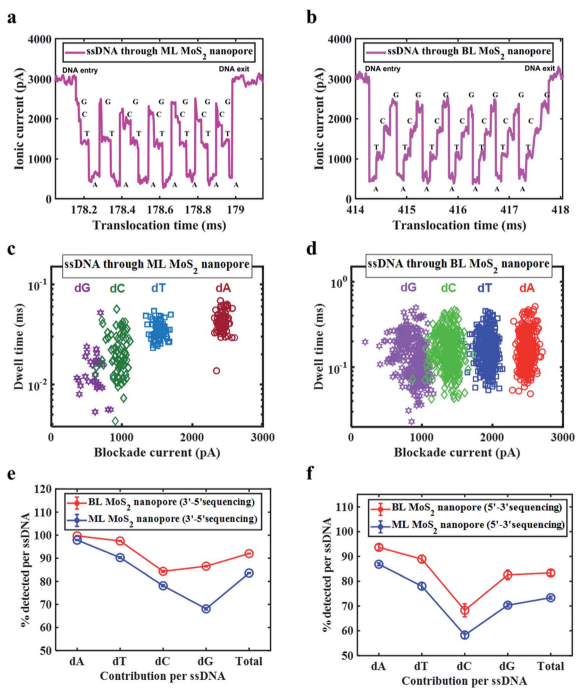


Fig. 4 A typical truncated ssDNA translocation data through (a) ML MoS₂ nanopore and (b) BL MoS₂ nanopore with single nucleotides being detected, dwell time vs. blockade current scatter plot for the single nucleotides detected from ssDNA translocation through (c) ML MoS₂ nanopore and (d) BL MoS₂ nanopore. (e and f) Percentage efficiency of individual nucleotide detection per ssDNA strand and ssDNA sequencing efficiency for 3'-5' and 5'-3' orientations respectively showing that BL MoS₂ nanopore demonstrates better sequencing efficiency than ML MoS₂ nanopore for both sequencing orientations.

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blockades were observed. From careful inspection of each molecular blockade, it can be clearly understood that the discrete blockade steps correspond to the constituent nucleotides. All the translocations for the ML and BL were analysed using the code developed and dwell time as a function of blockade current for 3 s translocations are plotted in Fig. 4c and d. For BL MoS₂ nanopore, we observe four different current levels similar to observed for each nucleotide with dwell times ranging from 0.02-0.6 ms with the highest density obtained around 0.08-0.3 ms which is similar to what we obtained for single nucleotide translocations. On the other hand, for ML MoS₂ nanopore, four different nucleotides are also detected. However, the time resolution obtained for dC and dG are particularly low (0.004-0.04 ms for dG and 0.004-0.06 ms for dG) compared to dA and dT, the latter nucleotides showing dwell times in the range of 0.01–0.07 ms, with the mean \sim 0.05 ms, which is close to that obtained for single nucleotide triphosphates.

Based on the different dwell times and blockade current four different nucleotides can be distinctly identified. The blockade levels are then matched with the single nucleotide data recorded before to obtain and relate the DNA sequence. Fig. S7a and b† shows the sequence of nucleotides detected and left undetected by ML and BL MoS2 nanopore sensing in a color-coded format, thus representing the single nucleotide efficiency of detection in each case. ML MoS2 nanopore is found to demonstrate a single nucleotide detection efficiency of 84.9% ((total - undetected)/total nucleotides) considering 60 DNA molecules translocation. Similarly, the experimental analysis shows that BL MoS₂ nanopore are capable of identifying single nucleotides with about 89.1% efficiency. For both ML and BL nanopores, it is observed that mostly dC and dG (especially at the tailing end or entering end of the DNA) are undetected or detected with poor time resolution. This may be due to the low density of these two nucleotides along with their low blockade current.

In addition, we have extracted the efficiency of ssDNA sequencing for both 3'-5' and 5'-3' orientations. BL MoS₂ nanopores demonstrate 83 \pm 1.07% and 92 \pm 0.14% for 5'-3' and 3'-5'-ssDNA sequencing efficiency respectively. These are 10% better than ML MoS $_2$ for 5′–3′-orientation (73.35 \pm 0.55%) and 8.4% better for 3'-5' sequencing (83.6 \pm 0.16%). Fig. S8 and S9† shows sequencing events for 3'-5' and 5'-3' ssDNA translocations for ML and BL MoS₂ nanopores respectively.

Fig. 4e and f shows the efficiency of detection of individual molecules from ssDNA and ssDNA sequencing as a whole. It is found that the efficiency of BL MoS₂ nanopores is better in both cases. The role of BL MoS₂ is observed to be most prominent in improving the temporal resolution of the signature for dC and dG. It was observed that the 47/60 ssDNA translocations were in 3'-5' orientation for ML MoS₂ nanopore and 51/60 ssDNA translocations were in 3'-5' orientation for BL MoS₂ nanopore.

Discussions

Short dwell time or high translocation speed in a nanopore prevents precise sequencing of nucleotides. Different techniques have been used to slow down the DNA translocation using trans-membrane viscosity gradient, electric field tuning and surface charge modification of the nanopore/electrolyte interface.28-31 DNA molecule sensing has been demonstrated through sub-5 nm nanopores on 25 nm thick SiN_r membranes in 38 pM concentration solution.³⁶ The study demonstrated that unamplified dsDNA molecules (as opposed to single nucleotides) can be sensed by slowing down the translocation using 20-fold salt gradient.36 A ML MoS2 nanopore-based viscosity gradient system has also been used for improved DNA sequencing by slowing down the translocation by two orders of magnitude.13 A simulation study has also been used to demonstrate improved peptide dwell times for BL MoS2 nanopores.22

In this study, we experimentally explore the unaided capacity of BL MoS₂ nanopores in improving real-time sensing resolution, detection rate and efficiency for ssDNA sequencing. We first analysed the blockade current obtained for single nucleotide using ML and BL MoS₂ nanopores. Our ML MoS₂ nanopore current signatures are similar to that obtained in previous study.13 However, we observed geometry defying blockade signatures for guanine. This may be due to the increased hydrogen bonding potential of guanine or increased physical adsorption energy of guanine on MoS2 which may increase the pore conductance, thus producing a lower overall ionic blockade event.37 For ssDNA sequencing, we also observed lower time resolution of guanine base of the ssDNA compared to other bases, which may indicate that the effect of surrounding A and C over nanopore blockade are dominant over G. Few studies have shown exceptions to the geometry-determined blockade when larger A and G have produced lower blockades than C and T, for both single nucleotide and polynucleotide sensing. 13,20,38

We also obtain higher blockade current levels for BL MoS₂ nanopore as compared to ML MoS2 nanopore. In general, for 3D bulk materials like SiNx, the conductance decreases with increase in pore/membrane thickness.39 However, in our study we find that 2D MoS₂ nanopores do not demonstrate this behaviour. Unlike 3D bulk material nanopores, in 2D material nanopores interlayer charge storage, electro-activity and field confinement between the van der Waals coupled monolayers increases the capacitance of the pore.40 COMSOL Multiphysics simulation study indicates that the ionic conductance of MoS₂ nanopores demonstrates the following trend 2-layers > 1-layer > 4-layers > 3-layers > 6-layers > 5-layers and so on above 2 nm diameter nanopore.41 Few studies on 2D materials nanopores like graphene also demonstrate similar behaviour. 18,42,43

We further observe nearly double the dwell time for BL MoS₂ as compared to ML MoS₂ nanopore, with 95% confidence. To observe the effect of the bias on the blockade current we measured the blockade current for three different voltages. Fig. S11† shows blockade current vs. dwell time scatter plot for single nucleotide translocations through ML and BL MoS₂ nanopores for varying transmembrane bias values (100 mV, 150 mV and 200 mV). However, significant change in dwell times is not observed for either of ML or BL MoS2 nanopore sensing. This further proves that the events considered are indeed translocation events and strengthens the inference Paper

drawn about improved ability of BL MoS₂ nanopores in providing better resolution for single nucleotide sensing.

The effect of the temporal resolution improvement offered by BL MoS₂ nanopore on sequencing single-stranded DNA oligos was investigated next. From these experiments, we found that the capability of BL MoS₂ nanopores can resolve the nucleotide blockade currents and hence can distinguish single distinct nucleotides. Thus, we observed an overall single nucleotide (for 1800 nucleotides in 60 ssDNA strands) detection efficiency of 89% using BL MoS2 nanopore, a 4% improvement compared to that for ML MoS₂ nanopores. We also observed an 8-10% improved ssDNA sequencing efficiency for both 3'-5' and 5'-3' ssDNA orientations. The interlayer potential gradient for BL MoS2 offers improved capture of analytes inside the nanopore even at low (picomolar) concentration leading to $5\times$ better detection rate. Previous studies have shown that improved nanopore/DNA interaction can result in faster capture of DNA within the nanopore and also cause longer residence of DNA at the nanopore.44 We believe that the improved capture rate and nanopore dwell time is a result of the enhanced DNA/ nanopore interaction experienced in BL MoS2 nanopore as compared to ML MoS2 nanopore.

Moreover, the behaviour of even and odd layers of MoS₂ is found to differ under an applied vertical electrical field. ⁴⁵ The electric field confinement within the layers is more for odd-numbered layers than even-numbers, which thus improves the ionic conduction for even layers. Moreover, the BL MoS₂ thickness is close to nucleotide size which helps maintain a good spatial resolution. However, >2 layers can hamper the spatial resolution. Additionally, our simulation results also show a reduction in blockade signal amplitude above two layers. All things considered; we believe >2-layers MoS₂ may not be a better choice for DNA sequencing. The results conclusively suggest that BL MoS₂ nanopores can distinguish four type of nucleotides with significantly higher dwell time. Thus, BL MoS₂ nanopores can be a suitable choice for more efficient and faster DNA sequencing.

Conclusion

In summary, we experimentally demonstrate that BL MoS₂ nanopores are capable of detecting distinct single nucleotides with twice the dwell time as compared to ML MoS₂ nanopores. The van der Waals separated layers of BL MoS2 experience different potentials when a bias is applied across them creating an interlayer potential gradient. This makes each individual molecule experience an increased charge interaction and stronger immobilization at the nanopore, leading to prominently sharp and broad translocation events. The translocation profiles through BL MoS₂ nanopores show distinct peaks for each type of DNA nucleobase of 10 pM concentration. Thus, we have been able to detect nucleotides at 89% efficiency using BL MoS₂ nanopores for 60 detected DNA molecules. Furthermore, we observed an 8-10% improved sequencing efficiency by using BL MoS₂ nanopores. The BL MoS₂ nanopore can resolve singlenucleotide signals temporally, while maintaining a good spatial resolution as well. This high detection rate, efficiency and dwell

time makes BL MoS₂ nanopores promising for high-speed detections in low concentration analyte solutions which is essential for several biosensing applications.

Data availability

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Author contributions

P. S. conducted all the simulations, fabrication, and experiments on the nanopores. P. S. and M. G. analyzed and wrote the manuscript.

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Conflicts of interest

The authors declare that there are no competing interests.

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