Gas flow through atomic-scale apertures

Jothi Priyanka Thiruraman¹,²*, Sidra Abbas Dar³,⁴,⁵*, Paul Masih Das¹, Nasim Hassani⁶, Mehdi Neek-Amal⁷,⁸, Ashok Keerthi⁴,⁸†, Marija Drndić¹†, Boya Radha³,⁴†

Gas flows are often analyzed with the theoretical descriptions formulated over a century ago and constantly challenged by the emerging architectures of narrow channels, slits, and apertures. Here, we report atomic-scale defects in two-dimensional (2D) materials as apertures for gas flows at the ultimate quasi-0D atomic limit. We establish that pristine monolayer tungsten disulfide (WS₂) membranes act as atomically thin barriers to gas transport. Atomic vacancies from missing tungsten (W) sites are made in freestanding (WS₂) monolayers by focused ion beam irradiation and characterized using aberration-corrected transmission electron microscopy. WS₂ monolayers with atomic apertures are mechanically sturdy and showed fast helium flow. We propose a simple yet robust method for confirming the formation of atomic apertures over large areas using gas flows, an essential step for pursuing their prospective applications in various domains including molecular separation, single quantum emitters, sensing and monitoring of gases at ultralow concentrations.

INTRODUCTION
Understanding confined gas flows in angstrom-scale tight spaces plays a major role not only in the design of gas extraction techniques but also for gas separation and production (1). In extremely narrow pores, the mean free path of a gas is much larger than that of the dimensions of the pore itself, which leads to gas dynamics dominated by molecular collisions with walls of the pore rather than the intermolecular collisions (2). This is known as the free molecular regime, and the gas flow through these pores was comprehensively described using the Knudsen equation (2), which has been modified and adapted to explain the flows through various confined systems (3). From a theory standpoint, a pore or an aperture is a simple model system through which gas transmission is proportional to the impingement of gas molecules, i.e., likelihood of a gas molecule encountering a pore, and the activation barrier, if any, to cross the pore. In the cases where the membrane surface can absorb gases, the flow is a combination of direct transmission through the pore and diffusion along the membrane surface (4). Despite the emergence of many nanoscale gas flow conduits such as nanopores (5–9), nanotubes (10–12), nanochannels (11, 13–16), nanolaminates (17, 18), etc., ultimately narrow quasi-zero-dimensional (0D) apertures with atomic-scale dimensions in both the transmembrane and lateral directions, which challenge the applicability of the Knudsen equation for gas flows, have been limited (5, 8, 9, 19). Although atomic vacancies in monolayer 2D materials have been ideal candidates for theoretical simulations and modeling of gas flows (4, 20–22), they have not been studied extensively in experiments (9, 14). Here, we investigate an inert gas, i.e., helium flow through atomic vacancies in freestanding monolayer tungsten disulfide (WS₂) membranes, to validate the Knudsen description in the ultimate atomic aperture limit.

RESULTS
Several studies in the past have explored various sources for atomic defect creation in 2D materials; among those, the popular techniques are oxygen plasma (23, 24), thermal annealing (25), ion and electron beam irradiation (26, 27), acid etching (23, 28), and ultraviolet-induced oxidation etching (5). In particular, ion irradiation offers a precise method of creating atomic vacancies with a controlled localization of defect sites at comparatively high densities (>10¹¹ cm⁻²). Recently, we have illustrated how to create highly controlled single-atom defects by focused ion beam (FIB) irradiation on a monolayer transition metal dichalcogenide (TMD) flake (29). In particular, TMDs provide better imaging contrast in aberration-corrected scanning transmission electron microscopy (AC-STEM) imaging and appear less prone to contamination, enabling easier characterization of the defects. We compared the effects of FIB irradiation on various suspended monolayer TMDs and established that defects with areas down to 8 Å² (single transition metal vacancy) can be produced in WS₂ with a specific low irradiation dose (increasing dose produces larger vacancies in the membrane). Therefore, we choose monolayer WS₂ as an optimal base support for hosting single-atom defects. Apart from being mechanically stable with a Young’s modulus of 270 GPa and a thickness of ~0.7 nm (30), our detailed investigation of the controlled ion irradiation mechanism minimizes the possibility of undesired damage or tears. The FIB irradiation mechanism used here allows for exclusively hosting single atomic apertures on a freestanding monolayer WS₂ membrane, enabling a high density of uniform defects in the range of 9 × 10¹¹ ± 3.5 × 10¹⁰ cm⁻² (see sections S1 to S3). These defect densities are comparable to that achieved in graphene membranes (9).

The WS₂ membranes incorporating atomic vacancies are supported on silicon chips (SiNₓ/Si) with a size of 11 mm by 11 mm (Fig. 1A and fig. S1). In the center of each chip, one or more submicrometer holes were drilled in a freestanding silicon nitride (SiNₓ) membrane (Fig. 1B). A monolayer WS₂ flake was suspended on submicrometer holes present on these silicon chips (see fig. S1). The suspended
Atomic apertures in the WS₂ membrane act as the only connecting paths between the two chambers where gas molecules can flow. As control samples, three replicas were done for each of the pristine silicon nitride membrane with and without holes and nonirradiated WS₂ membrane suspended over the silicon nitride hole. The control samples used to establish baseline flow are different set of membranes than those used for gas transport; however, they all come from the same fabrication procedures and chemical vapor deposition (CVD) growth. A bare hole without WS₂ layer exhibits large gas flow, as expected, and has been used as a standard leak for validation of the experimental setup (Fig. 2). A freestanding pristine WS₂ monolayer covering nine holes each with diameters of ~250 nm, without any irradiated defects, exhibited negligible helium flow below our detection limit (~$10^{-18}$ mol s$^{-1}$ mbar$^{-1}$). It is remarkable that the WS₂ monolayer grown by CVD methods (see fig. S2) has such low intrinsic defect density that it is practically impermeable over a suspended area of a few square micrometers. Let us recall the ultralow permeability, ranging from $10^{-22}$ mol s$^{-1}$ mbar$^{-1}$ to only few gas molecules per hour (31, 32), of intrinsic defects in 2D materials with high crystal quality such as mechanically exfoliated graphene, studied extensively with specialized device architectures and atomic force microscopy measurements done over several days.

Apart from the high crystal quality of our WS₂ monolayer, the impermeability emphasizes the excellent sealing of the WS₂ layer on the SiN, membrane in our devices, which is achieved by repeated annealing of the samples in H2/Ar atmosphere at 350°C, both right after the WS₂ monolayer transfer and also before the gas flow measurements (see section S3).

Next, we did helium (He) gas flow measurements on FIB-irradiated samples containing atomic vacancy defects in suspended WS₂ membranes. Three typical irradiated samples along with controls are shown in Fig. 2A, in which sample no. 1 has WS₂ suspended on nine holes each with a diameter of ~200 nm. From our fabrication method, mainly three types of defects are possible such as 1W (~69%), 2W (~22%), and 3W (~9%) defects, respectively. Here, the pore configuration is focused on W atoms since experimentally, the AC-STEM contrast from S atoms is weak, and hence, their locations are not identified with certainty. Unlike the geometric area given in Fig. 1D, to get the effective area (accessible pore area for He atom), the van der Waals (vdW) diameter of each atom on the aperture edge is subtracted (see section S2) (14). The total estimated tungsten defects is given as $N = N_1 + N_2 + N_3$, where $N_1$, $N_2$, and $N_3$ are estimated numbers of 1W, 2W, and 3W vacancies, respectively, from AC-STEM image analysis. In our samples, $N \sim 2070 \pm 830$ for sample no. 1; the sample no. 2 was WS₂ on a single hole with a diameter of 250 nm to yield total defects $N \sim 440 \pm 170$; the sample no. 3 was WS₂ on a 200-nm-diameter hole to give estimated defects, $N \sim 300 \pm 120$. He flow through atomic apertures increases linearly with increasing pressure and also with increasing number of defects in a sample (Fig. 2A). The gas flux is not exactly linearly correlated with the increase in $N$. The additional sources of error include the variation in the substrate hole area leading to an overall WS₂ suspended membrane area variation and errors in the ion irradiation dose, in total amounting to about ~40% error. At the experimental working pressure $P$ ranges from few to 200 mbar, the mean free path of helium is >0.5 μm, and the Knudsen number for these atomic apertures in our WS₂ monolayer is $>10^3$. Here, the defects are not circular and have a well-defined atomic structure (see inset in Figs. 1D and 3), meaning that a diameter (typically used for larger circular pores) is...
Fig. 2. Gas flow through atomic apertures. (A) Helium gas permeation measured through atomic apertures in WS$_2$ made with same irradiation dose ($5.1 \times 10^{13}$ ions/cm$^2$) but with different WS$_2$ areas leading to difference in the number of defects in a given sample. The proportion of the 1W, 2W, and 3W vacancies in each sample is 69:22:9%, sample no. 1, $N \sim 2070 \pm 830$; sample no. 2, $N \sim 440 \pm 170$; sample no. 3, $N \sim 300 \pm 120$. Pristine WS$_2$ membranes without any irradiation and large bare hole not covered with WS$_2$ (shown in red color) are shown alongside as controls. Solid lines running through symbols indicate best fit to the data. Inset shows a schematic of our experimental setup. (B) Comparison of normalized permeance per unit pressure of irradiated samples and controls, with the Knudsen estimates. The colors within the gray bars representing Knudsen estimates arise from the flow contributions due to 2W (red) and 3W (dark blue) vacancies. Horizontal light blue color bar indicates the detection limit. Error bars on the helium flow of samples are from the SD of flow values recorded over a series of helium flow measurements (measured three times) on the same sample repeated after annealing three times. For the Knudsen theory estimates, the error bars arise from the error values associated with the number of defects and with the total area of the apertures $A$ in each sample.

Fig. 3. Atomic apertures and electron density isosurfaces. Typical aperture sizes with the irradiation dose ($5.1 \times 10^{13}$ ions/cm$^2$) are shown. The top panel shows the density functional theory (DFT) results for electron density isosurfaces of atomic apertures in a WS$_2$ membrane (isovalue of 0.2 \text{e/Å$^3$}). The S and W atoms are shown in yellow and blue, respectively. The blue and red arrows inside the images are the accessible dimensions in width $a$ and length $b$ across the pores (1W+6S, $a \sim 3.2$ Å; 2W+6S, $a \sim 2.1$ Å, $b \sim 6.2$ Å; 3W+12S, $a \sim 5.8$ Å, $b \sim 6.2$ Å). The accessible aperture areas for helium molecules are $\sim 0.08$, $\sim 0.13$, and $\sim 0.23$ nm$^2$, respectively. The bottom row includes the AC-STEM images of the defect, while the top row illustrates the observed atomic configuration.

not an optimal measure of size. We quote their geometric area, i.e., mean area $<A>$, and the characteristic sizes in table S1. As an example, the 1W (+6S) defect has a pore with characteristic size, $\sim 3.15$ Å, and an area of 0.08 nm$^2$, whereas the 3W (+6S) defect has a pore size of $\sim 5.25$ Å and an area of 0.216 nm$^2$. To represent the size of the He, we use the kinetic diameter ($\sim 2.6$ Å), which is a semiclassical notion; however, for monoatomic spherical molecules like He, this is quite close to the quantum-mechanical size of the electron cloud around the nuclei (33). As the aperture size is much smaller than the mean free path, the mass flow of the gas $Q$ (moles per second) through the aperture is simply the impingement rate upon the area of the pore (see section S6), as described by Knudsen (22, 34).

$$Q = \frac{PA}{2\pi MR^2}$$

where $P$ is the inlet pressure, $A$ is the total area of the conducting apertures, $M$ is the atomic mass of the gas being transported ($M = 4$ g mol$^{-1}$ for helium), $R = 8.314$ J mol$^{-1}$ K$^{-1}$ is the gas constant, and $T$ is the temperature ($T = 295$ K in our experimental setup). In our case, $A$ is the sum of all individual atomic aperture areas in the WS$_2$ membrane, which is, on average, $N_1 \times A_1 + N_2 \times A_2 + N_3 \times A_3$ (in our samples, $N$ can be varied from 300 to 2000 for an individual sample by increasing the membrane area; $A_1$, $A_2$, and $A_3$ are the accessible aperture areas for 1W, 2W, and 3W vacancies, i.e., $\sim 0.08$, $\sim 0.13$, and $\sim 0.23$ nm$^2$, respectively, given in table S1. In the case of control devices, i.e., large bare holes, $A$ is the sum of individual hole areas (with diameters in the range of $\sim 200$ to $\sim 300$ nm). Taking into consideration the proportion of the various defects observed in AC-STEM images, we calculated the Knudsen estimates from all the vacancies such that 3W vacancies contribute to $\sim 9\%$ of the flux, whereas $\sim 91\%$ of the flux is contributed by 1W (+6S) and 2W vacancies together (Fig. 2B). If the pores are much narrower than
the size of the molecule, then there can be a finite energy barrier, which is expressed as an exponential term in the Eq. 1, \( \exp(-E/RT) \), where \( E \) is the energy barrier that substantially reduces the transmission coefficient of the gas even for small \( E \) \( (14, 22) \).

The 2D membranes with atomic apertures show measurable helium flux \( \sim 10^{-16} \) to \( 10^{-15} \) mol s\(^{-1}\) mbar\(^{-1}\), over two to three orders higher than the detection limit and significantly lower than the leakage rate for submicrometer holes (Fig. 2B). This corresponds to a flow of about \( \sim 10^2 \) to \( 10^8 \) He molecules per second per millibar pressure difference across the membrane. Theoretical simulations with such small pores in graphene have estimated a pass through frequency of \( 10^9 \) He molecules per second per defect (close to our flow values per millibar for a sample with \( \sim 2000 \) defects; Fig. 2), which can be likely ascribed to a low diffusion barrier for He resulting from its noninteracting nature \( (14, 20) \). As a check for the reproducibility of devices with atomic apertures, we would like to point out that in a given batch of irradiated samples, about 40 to 50% samples show the measurable flow commensurate with the number of defects estimated by the irradiation dose \( (29) \). Other devices either did not conduct gas or showed much higher gas flow. From the He flow, we back-calculated the number of defects contributing to the flow using Eq. 1, and \( N \) matches close to the defect numbers from AC-STEM image analysis for the sample nos. 1 to 3. For high leaking samples, substituting experimental \( Q \) in Eq. 1 gave large contributing defect area \( A \), which does not correlate with the defect density statistics attained from AC-STEM, hinting that these samples might have tears or cracks in the membrane. Examination of these samples by scanning electron microscopy (SEM) on large areas indeed indicates that they are damaged samples (see tears in suspended WS\textsubscript{2}; fig. S6).

The measured He flow values are within an order of magnitude of the values predicted by the Knudsen estimates, despite the uncertainties in our defect densities and the distribution of the type of defects. Equation 1 is based on the kinetic theory and has been applied to describe the flows through apertures using effusion mechanism in both theoretical \( (4) \) and experimental literature \( (5, 6) \).

To more precisely validate the agreement to Eq. 1 and/or evaluate its limitations in our experiments, we would need to further decrease our experimental sources of error, including the calibration of all defect areas, which is a challenging experimental task. To verify whether there is any energy barrier, we measured gas flows by varying the temperature between \( \sim 25^\circ \) to \( \sim 100^\circ \)C, by using a heating tape wound around the chamber, interfaced with a temperature controller. We did not see noticeable variation in the gas flow with temperature, hinting an absence of an energy barrier (an exponential dependence is expected if an energy barrier is present). As per Eq. 1, a \( T^{-1/2} \) dependence would lead to variation of flux by \(<10\% \) within the temperature range tested, and this is beyond our experimental measurement error.

From the electron density isosurface (EDI) of defects shown in Fig. 3 (and fig. S4) using first-principles density functional theory (DFT) calculations, accessible pore areas are obtained (table S1). The accessible pore areas from EDI closely match with those estimated using \( \text{vdW} \) radii (section S2). The surface transport contributions can be negligible as He does not adsorb sufficiently on the WS\textsubscript{2} basal plane, similar to that observed on graphene (typical adsorption layer thickness, \( \sim 4 \) to \( 6 \) Å) \( (35) \).

Ideal atomic apertures with only 1W missing and with no missing sulfurs on the pore edge would likely, theoretically, be nonconducting. On the other hand, 1W defects with six sulfur vacancies with an accessible pore diameter \( \sim 3.2 \) Å, which is larger than the kinetic diameter of helium \( (2.6 \) Å), would be amenable to gas flows. As shown in Fig. 1D, the minimum area of the defects obtained from AC-STEM imaging was about 0.06 to 0.2 \( \text{nm}^2 \), which corresponds to the defects with 1W and 6S missing \( (\sim 0.08 \text{nm}^2) \), whereas the 1W defects with no sulfur missing would be of significantly lower area \( (\sim 0.002 \text{nm}^2) \). As the characteristic pore size decreases, the electron overlap between He and membrane significantly increases, thereby increasing the energy barriers for the flow. However, the theoretical calculation of energy barriers might not necessarily capture all the favorable electronic interactions in real systems as there can be rearrangements in the vacancies. These discrepancies between the computed energy barriers and the experimental flows have been noted before by Wang et al \( (8) \). Specifically, our results do not imply a significant energy barrier compared to theoretical \( (4, 8, 14, 22) \), and we also note that different levels of DFT calculations (and supercell size, or equivalently, the ratio between the pore area and the simulated defect area in membrane) can give different energy barriers; thus, a separate systematic DFT study is required for carefully probing the energy barriers.

We stored our samples with atomic vacancy defects under ambient conditions. When we tested the samples after a few weeks, it was observed that they tend to get clogged with time, leading to reduced/no He flow. As is well known, all surfaces are likely to be covered by hydrocarbons, which might cause the clogging \( (15) \) of atomic apertures in our case. However, upon reannealing at 350°C, He flow was regained to the previously observed value within a factor of 2. Storage of samples in activated charcoal helped extend the life span of the atomic apertures to be open for few months. Annealing with a combination of storage in charcoal has enabled us to keep the atomic apertures open for about a year. The atomic vacancy defects, once created, are quite stable and show similar helium flow with repeated annealing, indicating that the atomic apertures do not expand or propagate rapidly. Those samples that have been irradiated but did not conduct gas remained nonconductive even after repetitive annealing. We speculate that these samples could have contained a majority of the dominant 1W defect type with one “W” atom missing but with partial or all S atoms at that site, thereby increasing the energy barrier and leading to no He flux. While the ion irradiation dose is optimized for single-atom (W) defect pores, the removal of sulfur atoms from ion irradiation and the subsequent AC-STEM imaging to precisely know the composition of sulfur vacancies at a defect site have remained a challenge. However, in the extreme limit, when there were large holes, tears, or cracks in the membrane, the flow was much higher, and the membranes themselves were not stable. In addition, we performed ion flow measurements in KCl solution on our atomic apertures as a cross-check. Those samples that showed inconsistent and unexpectedly high gas flows showed high ionic conductance \( (\sim 400 \text{ nS in } 1 \text{ M KCl}; \text{see fig. S7}) \), again indicating the presence of tears (visible in SEM; see fig. S6). The samples that showed gas flows commensurate with the Knudsen equation exhibited only a small ionic conductance, \(<1 \text{ nS in } 0.1 \text{ M KCl for total membrane area } \sim 0.4 \) to \( 0.6 \text{ m}^2 \), as shown in fig. S7. This is expected as the atomic apertures with sizes \(<6 \text{ Å} \) exclude ions \( (27) \), but the presence of tears in the membranes leads to bulk ionic flow (see fig. S6). These observations further emphasize that the sealing of WS\textsubscript{2} monolayers to the SiN\textsubscript{3} membranes is leak proof, leaving no gaps, and hence, only irradiated defects contribute to the observed gas flow.
DISCUSSION
Let us analyze the gas flow in nano- and angstrom-scale pores from the literature, in comparison to our atomic vacancy defects presented here. It has been shown in the literature that when the pores are less than the size of the gas molecule passing through (8), activated transport is observed and highly sensitive energy barriers play a major role in the transport, and the barrier is usually estimated using combined theoretical and experimental efforts. In this work, since the atomic vacancy apertures are only slightly larger than the gas molecule (i.e., less than twice the size of helium molecule, 2.6 Å), the flow is governed by simple effusion (14). In general, effusive flow is seen as detrimental and the cause of reduction for selectivity between gases in size exclusion. Because of this, an accurate comparison of the observed permeability and value estimated from the Knudsen description for the known size of the aperture is often overlooked. Such a comparison made for nanometer-sized holes showed the validity of the Knudsen description in graphene pores down to ~7 nm in size, where a N₂ permeance of ~0.05 mol m⁻² s⁻¹ Pa⁻¹ was observed (6). Angstrom-size defects (size, ~0.38 to ~0.43 nm) made by ozone-induced etching in a CVD-grown graphene layer were shown to have a H₂ permeance of about 10⁻⁷ mol m⁻² s⁻¹ Pa⁻¹ (9), which is one order higher than the permeability of graphene membrane, 10⁻⁸ mol m⁻² s⁻¹ Pa⁻¹ hosting the defects (36). However, the observed gas flow values for atomic defects are still much lower than that estimated from the Knudsen description for given defect densities, which might be due to the overestimation of the conductive apertures. In our WS₂ atomic aperture samples, the He gas flow obtained matches the Knudsen estimates within a factor of 2 to 3, and the gas flow normalized by the total area of defects translates to ~0.05 to 0.1 mol m⁻² s⁻¹ Pa⁻¹. From theoretical studies on graphene pores with sizes 3.6 to 4.8 Å, for defect densities up to ~10¹⁴ cm⁻², large permeance of ~0.1 mol m⁻² s⁻¹ Pa⁻¹ has been predicted (14, 20), which closely matches with our experimental values. These permeance values are higher than those typically observed in silica (37), zeolite, and metal-organic framework membranes, graphene oxide membranes (18), which are in the range of ~10⁻⁶ to 10⁻⁷ mol m⁻² s⁻¹ Pa⁻¹. Large permeance values are obtained through our WS₂ apertures, even when normalized with the area of the membrane, i.e., ~10⁻⁷ to 10⁻⁸ mol m⁻² s⁻¹ Pa⁻¹, which indicates that most of the atomic apertures are in the predicted size range and hence conducive to gases.

In conclusion, we demonstrate the fast He transport through quasi-0D atomic-scale apertures (~W sites). To our knowledge, this is the first experimental observation on He gas transport in such angstrom-scale pores in WS₂/TMD monolayers where we attempt to evaluate the applicability of the Knudsen equation down to the ultimate limit. Our results indicate the necessity of future theoretical models to explore the role of sensitive variation of the energy barriers of various gas molecules and their critical dependence on the size of the aperture at this atomic limit, especially taking the electron density isosurfaces of atoms to account for the enhanced flux. In addition, this work provides a new method for corroborating atomic pore formation and their density over large areas via a simple bulk measurement technique of measuring gas flow through them. This is analogous to using nanoholes (few hundred nanometers in size) as standard leak elements for calibration (38). So far, the only way to inspect and confirm the atomic pore formation in the case of atomic vacancy defects is by AC-STEM, which is limited to relatively small areas. Conventional ionic transport measurements through nanopores are mediated by Hille’s equation (27), which yields a linear relationship between conductance and total aperture area. However, for pores that are smaller in dimensions than many common salts, ionic conductance measurements do not yield a predictable or a measurable signal, leading to the breakdown of Hille’s model for atomic aperture limit (28). With our study, we would like to highlight that once the pore creation is confirmed and calibrated by AC-STEM locally, gas flow measurements can act as a standard to test for the presence of these pores and their density over large areas. Let us also note that the stability of pore/aperture configuration of these atomic scale defects needs further investigation (27, 39) as the pores could change over time, which can be attempted in future experiments by imaging the pores over time. More advanced methods including dynamic scanning tunneling microscopy need to be devised to locate S atoms (40) and also any other atoms such as possible contaminants (C, O, and N) to be certain of pore compositions and sizes. Future work may involve efficient gas separation investigations using a scaled-up version of such membranes.

MATERIALS AND METHODS
Sample fabrication
**CVD growth of monolayer WS₂ flakes**
Monolayer WS₂ flakes were grown using CVD processes similar to previously reported methods (27, 29). Solutions of 2% sodium cholate growth promoter and 15 × 10⁻³ M ammonium heptamolybdate (metatungstate) were spun onto piranha-cleaned silicon (Si) substrates coated with 150 nm of SiO₂, which were then loaded into the center of a tube furnace (Thermo Fisher Scientific Lindberg/Blue M). For WS₂, samples were heated in argon atmosphere [100 standard cubic centimeters per minute (sccm)] at a rate of 65°C min⁻¹ and held at 800°C for 10 min, during which time 15 sccm of H₂ was also added. Approximately 100 mg of sulfur precursor placed 22 cm from the substrates was kept at 180°C during the growth procedures. Both samples were rapidly cooled to room temperature by cracking open the furnace.

**Position-controlled transfer of WS₂ flakes**
With WS₂ flakes grown on Si/SiO₂ wafer pieces, we cut smaller pieces with areas about 3 × 3 mm² that contain good WS₂ flakes. Using a wet transfer technique, the TMD flake side of the SiO₂ pieces is coated with poly(methyl methacrylate) (PMMA) (MicroChem C4, 4000 rpm for 60 s). The PMMA-coated substrates are left to dry at ambient temperature for a few minutes before being floated onto the KOH etchant (8 g per 100 ml of water). Depending on the thickness of the SiO₂ wafer, the time for etching away the SiO₂ layer varies. Last, we observe the PMMA + TMD flake floating on the KOH etchant. These PMMA + TMD flakes are then rinsed three times in clean water baths before being used to transfer them onto a SiNₓ/Si chip with a hole (diameter range, 200 to 300 nm) drilled by FIB. Position transfer is practiced when the flake size and their number density are high, so that in this case, it is possible to place one PMMA + TMD flake onto the SiNₓ, FIB hole. This step of placing the PMMA + TMD flake over the SiNₓ, FIB hole is somewhat challenging since the coverage of the FIB hole depends on the number of the as-grown flakes. However, because in a given batch of CVD growth, the number of grown flakes is large, the transfer is completed within several transfer trials. In addition to this manual transfer, we have also used custom-built manipulators, with needles and tweezers.
to move the flake on the substrate to increase device yield. These manipulations came at a cost, since there was now a higher risk of device failure caused by the usage of additional tools, which can sometimes break the membrane. In this work, all devices were fabricated by manually “fishing” (moving) the PMMA + TMD flake onto the SiN, FIB hole. After we transfer the flake and optically verify the coverage, the sample is then dried in ambient temperature for 30 min. Furthermore, the sample is placed into hot acetone (90°C) to remove the supporting PMMA.

**Gallium ion irradiation**

Monolayer TMD flakes were irradiated with a Ga⁺-sourced ion beam FEI Strata Dual-Beam instrument. The acceleration voltage of the ion beam was set to 30 kV and incident normal to the surface. The beam spot size was observed to be 100 nm for a flash second at 10-pA current. To produce atomic defects, an area of 250 nm by 250 nm was irradiated with the dwell time (16 μs) and current (10 pA). Pixel resolution (1024 × 884) was kept constant. The exposure was carried out in an imaging mode, which followed a raster pattern where the beam sequentially exposed each pixel in a row. The instrument, FEI FIB Strata DB 235, has an option to “grab frame,” which takes a single scan at a set resolution; this option was used for all the scans. The dose was varied by changing the number of scans. Suspended and substrate-supported samples were exposed to FIB irradiation while sitting on holey carbon TEM grids and Si/SiO₂ substrates, respectively.

**AC-STEM imaging**

AC-STEM images of WS₂ samples were acquired using a Cs-corrected JEOL ARM 200CF STEM operating at 80 kV. Images were obtained using a high-angle annular dark-field (HAADF) detector with a collection angle of 54 to 220 mrad and 10-cm camera length. Pixel current (22 pA), focusing time (<2 s), and electron dose (=6.0 × 10⁻⁶ e⁻ nm⁻²) were kept low to minimize beam-induced knock-on damage.

**First-principles calculations**

The DFT was performed with the generalized gradient approximation (41) form with exchange-correlation potential parametrization of Perdew-Burke-Emzerhof (42). The calculations were carried out by the Quantum ESPRESSO package (43) and have been performed on the basis of the plane-wave basis sets and ultrasoft nonlocal pseudopotentials (44). To take the vdW contributions into account in the total energy, the Tkatchenko-Scheffler (45) method was used. The cutoff of kinetic energy in the plane-wave expansion and the convergence threshold for the self-consistent field calculations were chosen as 280.0 and 10⁻⁶ e⁻ V per atom, respectively. The WS₂ membrane was relaxed both in plane and out of plane.

Calculations were done at 0 K for a rectangular supercell shape with size 15.953 × 16.579 Å² for all considered defects consisting of 29(60), 29(54), 28(58), 28(54), 27(54), and 27(48) atoms of W (S) for 1W, 1W (+6S), 2W (+2S), 2W (+6S), 3W (+6S), and 3W (+12S) with a vacuum layer of 30 Å between periodic images in the vertical direction. We calculated the EDI using the above aforementioned functionals and energy cutoff. For the EDI images (Fig. 3 and fig. S4), the isovalue was set to be 0.2 e/A³.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/6/51/eabc7927/DC1

**REFERENCES AND NOTES**

1. H. B. Park, J. Kamcev, L. M. Robeson, M. Emilevich, B. D. Freeman, Maximizing the right stuff: The trade-off between membrane permeability and selectivity. Science 356, eaab0530 (2017).

2. M. Knudsen, Die Gesetze der Molekulärströmung und der inneren Reibungsströmung der Gase durch Röhren. Ann. Phys. 333, 75–130 (1909).

3. M. V. Smoluchowski, Zur kinetischen Theorie der Transpiration und Diffusion verdünnter Gase. Ann. Phys. 338, 1559–1570 (1910).

4. C. Sun, M. S. H. Boutillier, H. Au, P. Poesio, B. Bai, R. Karnik, N. G. Hadjiconstantinou, Mechanisms of molecular permeation through nanoporous graphene membranes. Langmuir 30, 675–682 (2014).

5. S. P. Koenig, L. Wang, J. Pellegrino, J. S. Bunch, Selective molecular sieving through porous graphene. Nat. Nanotechnol. 7, 728–732 (2012).

6. K. Celebi, J. Buchheim, R. M. Wyss, A. Droudian, P. Gasser, I. Shorubalko, J.-I. Kye, C. Lee, H. G. Park, Ultimate permeation across atomically thin porous graphene. Science 344, 289–292 (2014).

7. Y. Yamada, K. Murota, R. Fujita, J. Kim, A. Watanabe, M. Nakamura, S. Sato, K. Hata, P. Ercius, J. Ciston, C. Y. Song, K. Kim, W. Regan, W. Gannett, A. Zettl, Subnanometer vacancy defects introduced on graphene by oxygen gas. J. Amer. Chem. Soc. 136, 2232–2235 (2014).

8. L. Wang, L. W. Drahushuk, L. Cantley, S. P. Koenig, X. Liu, J. Pellegrino, M. S. Strano, J. Scott Bunch, Molecular valves for controlling gas phase transport made from discrete ångström-sized pores in graphene. Nat. Nanotechnol. 10, 785–790 (2015).

9. J. Zhao, G. He, S. Huang, L. F. Villabobos, M. Dakhchoune, H. Bassas, K. V. Agrawal, Etching gas-sieving nanopores in single-layer graphene with an angstrom precision for high-performance gas mixture separation. Sci. Adv. 5, eaav1851 (2019).

10. B. J. Hinds, N. Chopra, T. Rantell, R. Andrews, V. Gavalas, L. G. Bachas, Aligned multilayered carbon nanotube membranes. Science 303, 62–65 (2004).

11. J. H. Li, Z. Song, X. Zhang, Y. Huang, S. Li, Y. Mao, H. J. Ploehn, Y. Bao, M. Yu, H. Li, Z. Song, X. Zhang, Y. Huang, S. Li, Y. Mao, H. J. Ploehn, Y. Bao, M. Yu, Ultrain, molecular-sieving graphene oxide membranes for selective hydrogen separation. Science 342, 95–98 (2013).

12. M. H. Khan, M. Moradi, M. Dakhchoune, M. Rezaei, S. Huang, J. Zhao, K. V. Agrawal, Hydrogen sieving from intrinsic defects of benzene-derived single-layer graphene. Carbon 153, 485–466 (2019).

13. S. Blankenburg, M. Bieri, R. Fasel, K. Müller, C. A. Pignedoli, D. Passerone, Porous graphene as an atmospheric nanofilter. Small 6, 2266–2271 (2010).

14. D. de Jong, V. R. Cooper, S. Dai, Porous graphene as the ultimate membrane for gas separation. Nano Lett. 9, 4019–4024 (2009).

15. L. W. Drahushuk, M. S. Strano, Mechanisms of gas permeation through single layer graphene membranes. Langmuir 28, 16671–16678 (2012).

16. S. C. O’Hern, M. S. H. Boutillier, J.-C. Idrobo, Y. Song, J. Kong, T. Laoui, M. Atieh, R. Karnik, Selective ionic transport through tunable subnanometer pores in single-layer graphene membranes. Nano Lett. 14, 1234–1241 (2014).

17. J. Jadwiszczak, C. O’Callaghan, Y. Zhou, D. S. Fox, E. Weitz, D. Keane, C. P. Cullen, I. O’Reilly, C. Downing, A. Shmeliov, P. Maguire, J. J. Gough, C. McGuinness, M. S. Ferreira, A. L. Bradley, J. J. Boland, G. S. Duesberg, V. Nicoloisi, H. Zhang, Oxide-mediated recovery of field-effect mobility in plasma-treated MoS₂. Sci. Adv. 4, eaao5031 (2018).

18. J.-A. Ke, S. Garaj, S. Gradešič, Nanopores in 2D MoS₂: Defect-mediated formation and density modulation. ACS Appl. Mater. Interfaces 11, 26228–26234 (2019).

19. Z. Bi, L. Zang, H. Li, L. Liu, Nanopore creation in graphene by ion beam irradiation: Geometry, quality, and efficiency. ACS Appl. Mater. Interfaces 8, 24803–24809 (2016).
30. K. Liu, Q. Yan, M. Chen, W. Fan, Y. Sun, J. Suh, D. Fu, S. Lee, J. Zhou, S. Tongay, J. Ji, et al., "Defect-induced photoluminescence in monolayer semiconducting transition metal dichalcogenides." ACS Nano 9, 1520–1527 (2015).

31. J. S. Bunch, S. S. Verbridge, J. S. Alden, A. M. van der Zande, J. M. Parpia, H. G. Craighead, et al., "Simultaneous measurement of ionic and electronic states in monolayer MoS2 from a polyphenylene network." Nano Lett. 18, 1150–1154 (2014).

32. Y. Tao, Q. Xue, Z. Liu, M. Chen, W. Fan, Y. Sun, J. Suh, D. Fu, S. Lee, J. Zhou, S. Tongay, J. Ji, et al., "Simultaneous identification of low and high atomic number atoms in monolayer 2D structures and formation mechanism of sub-nanometer pores in 2D monolayer MoS2." Nano Lett. 19, 5097–5103 (2019).

33. J. S. Bunch, S. S. Verbridge, J. S. Alden, A. M. van der Zande, J. M. Parpia, H. G. Craighead, et al., "Simultaneous identification of low and high atomic number atoms in monolayer 2D structures and formation mechanism of sub-nanometer pores in 2D monolayer MoS2." Nano Lett. 19, 5097–5103 (2019).

34. S. Wang, C. Fiolhais, A. Züttel, M. S. Strano, K. V. Agrawal, "Single-layer graphene membranes by crack-free transfer for gas mixture separation." Science 279, 1710–1711 (1998).

35. Y. Tao, Q. Xue, Z. Liu, M. Chen, W. Fan, Y. Sun, J. Suh, D. Fu, S. Lee, J. Zhou, S. Tongay, J. Ji, et al., "Simultaneous identification of low and high atomic number atoms in monolayer 2D structures and formation mechanism of sub-nanometer pores in 2D monolayer MoS2." Nano Lett. 19, 5097–5103 (2019).

36. Y. Tao, Q. Xue, Z. Liu, M. Chen, W. Fan, Y. Sun, J. Suh, D. Fu, S. Lee, J. Zhou, S. Tongay, J. Ji, et al., "Simultaneous identification of low and high atomic number atoms in monolayer 2D structures and formation mechanism of sub-nanometer pores in 2D monolayer MoS2." Nano Lett. 19, 5097–5103 (2019).

37. Y. Tao, Q. Xue, Z. Liu, M. Chen, W. Fan, Y. Sun, J. Suh, D. Fu, S. Lee, J. Zhou, S. Tongay, J. Ji, et al., "Simultaneous identification of low and high atomic number atoms in monolayer 2D structures and formation mechanism of sub-nanometer pores in 2D monolayer MoS2." Nano Lett. 19, 5097–5103 (2019).

38. Y. Tao, Q. Xue, Z. Liu, M. Chen, W. Fan, Y. Sun, J. Suh, D. Fu, S. Lee, J. Zhou, S. Tongay, J. Ji, et al., "Simultaneous identification of low and high atomic number atoms in monolayer 2D structures and formation mechanism of sub-nanometer pores in 2D monolayer MoS2." Nano Lett. 19, 5097–5103 (2019).

39. Y. Tao, Q. Xue, Z. Liu, M. Chen, W. Fan, Y. Sun, J. Suh, D. Fu, S. Lee, J. Zhou, S. Tongay, J. Ji, et al., "Simultaneous identification of low and high atomic number atoms in monolayer 2D structures and formation mechanism of sub-nanometer pores in 2D monolayer MoS2." Nano Lett. 19, 5097–5103 (2019).

40. Y. Tao, Q. Xue, Z. Liu, M. Chen, W. Fan, Y. Sun, J. Suh, D. Fu, S. Lee, J. Zhou, S. Tongay, J. Ji, et al., "Simultaneous identification of low and high atomic number atoms in monolayer 2D structures and formation mechanism of sub-nanometer pores in 2D monolayer MoS2." Nano Lett. 19, 5097–5103 (2019).

41. Y. Tao, Q. Xue, Z. Liu, M. Chen, W. Fan, Y. Sun, J. Suh, D. Fu, S. Lee, J. Zhou, S. Tongay, J. Ji, et al., "Simultaneous identification of low and high atomic number atoms in monolayer 2D structures and formation mechanism of sub-nanometer pores in 2D monolayer MoS2." Nano Lett. 19, 5097–5103 (2019).

42. Y. Tao, Q. Xue, Z. Liu, M. Chen, W. Fan, Y. Sun, J. Suh, D. Fu, S. Lee, J. Zhou, S. Tongay, J. Ji, et al., "Simultaneous identification of low and high atomic number atoms in monolayer 2D structures and formation mechanism of sub-nanometer pores in 2D monolayer MoS2." Nano Lett. 19, 5097–5103 (2019).

Acknowledgments: We acknowledge assistance with AC-STEM imaging from R. Keyse of Lehigh University and D. Yates of the University of Pennsylvania. We thank J. Ford and M. Brukman of the University of Pennsylvania for help with FIB irradiation and PL/Raman spectroscopy, respectively. We also appreciate G. Danda for initial support and fabrication effort. Funding: B.R. acknowledges the funding from EPSRC grant no. EP/R013063/1. Royal Society fellowship and enhancement award RGF/EA1181000, and the European Union’s H2020 Framework Programme/ERC Starting Grant agreement no. 852674–AngstroCAP. A.K. acknowledges the Ramsay Memorial Fellowship, the Royal Society research grant RGS/ R2/202036 and also the University of Manchester’s research collaboration fund. S.A.D. was funded by a scholarship from the University of Engineering and Technology, Lahore, Pakistan. M.D. acknowledges support from the NSF through the University of Pennsylvania Materials Research Science and Engineering Center (MRSEC) (DMR-1720530) and NSF grant EFRI 2-DARE (EFRI-1542707), NSF DMR 1905045 and NSF CMMI Advanced Manufacturing (AM) grant 2002477. This work was performed, in part, at the University of Pennsylvania’s Singh Center for Nanotechnology, an NNIN member supported by NSF grant ECCS-1542153. J.P.T. acknowledges fellowship support from the Vagelos Institute of Energy Science and Technology (VIEST). N.H. was supported by the ISF under postdoc grant no. 2345. N.H. and M.N.-A. would like to acknowledge high-performance computing support from the Shahid Rajaei Teacher Training University. Author contributions: B.R., M.D., and A.K. designed and directed the project. J.P.T. and P.M.D. grew TMD materials and fabricated and characterized the devices. Specifically, J.P.T. carried out sample fabrication, FIB irradiation, and TEM image analysis. P.M.D. conducted AC-STEM imaging and Raman/PL measurements/analysis. S.A.D., A.K., and B.R. performed the measurements and their analysis. N.H. and M.N.-A. performed the DFT calculations. B.R., A.K., and M.D. wrote the manuscript with the inputs from J.P.T., P.M.D., and M.N.-A. All authors contributed to discussions. Competing interests: The authors declare that they have no competing interests. Data and materials availability: All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Additional data related to this paper may be requested from the authors.

Submitted 14 May 2020 Accepted 2 November 2020 Published 18 December 2020 10.1126/sciadv.abc7927

Citation: J. P. Thiruraman, S. A. Dar, P. Masih Das, N. Hassan, M. Neek-Arnal, A. Keerthi, M. Drndić, B. Radha, Gas flow through atomic-scale apertures. Sci. Adv. 6, eabc7927 (2020).