Implications of Permeation through Intrinsic Defects in Graphene on the Design of Defect-Tolerant Membranes for Gas Separation

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Implications of Permeation through Intrinsic Defects in Graphene on the Design of Defect-Tolerant Membranes for Gas Separation

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Abstract

Gas transport through intrinsic defects and tears is a critical yet poorly understood phenomenon in graphene membranes for gas separation. We report that independent stacking of graphene layers on a porous support exponentially decreases flow through defects. Based on experimental results, we develop a gas transport model that elucidates the separate contributions of tears and intrinsic defects on gas leakage through these membranes. The model shows that the pore size of the porous support and its permeance critically affect the separation behavior, and reveals the parameter space where gas separation can be achieved regardless of the presence of non-selective defects, even for single-layer membranes. The results provide a framework for understanding gas transport in graphene membranes and guide the design of practical, selectively permeable graphene membranes for gas separation.

Keywords: graphene membranes, gas separation, multilayer graphene, nanofiltration, nanofluidics

Separating the components of a gaseous mixture is a critical step in several important industrial processes including natural gas purification, hydrogen production, carbon dioxide sequestration, and oxy-combustion.¹⁴ Membrane separation systems offer definite benefits compared to conventional cryogenic and sorption-based methods, especially for small-to-medium scale applications.¹ Membrane separations employ no moving parts, require no exotic chemicals, and typically exhibit low energy requirements and flexibility in configuration.⁴ However, the inherent trade-off between the permeability and selectivity of conventional membrane materials has largely limited gas separation membranes to systems requiring relatively low production rates or to mixtures with high impurity concentrations in the feed gas.⁵
In contrast, graphene gas separation membranes have the potential to significantly surpass the permeance and selectivity limits of conventional membranes. With this single atom thick material, it is possible to create sub-nanometer-scale pores that allow smaller gas molecules to pass through but produce a physical barrier severely limiting the passage of larger molecules. Theoretical studies employing various computational and analytical tools have predicted permeance and selectivity values for various graphene pore geometries that are orders of magnitude higher than have been achieved with existing membrane technology. The feasibility of constructing such selective pores was recently demonstrated by Koenig et al. using micrometer-scale graphene membranes in which pores were created by UV-induced oxidative etching. They were able to produce two graphene membranes with selectivities of approximately 15,000, one being strongly permeable to gas molecules with kinetic diameters smaller than 3.4 Å and the other permeable to gases with kinetic diameters smaller than 4.9 Å.

However, the imperfect quality of large areas of graphene presents a significant challenge in scaling such single-layer graphene membranes to macroscopic sizes. Small areas of pristine graphene are impermeable even to helium, but due to the difficulty in isolating large areas of pristine graphene, macroscopic graphene membranes inherently have a non-zero permeance due to the presence of defects in the graphene. These defects include intrinsic nanometer-scale holes that develop during the graphene growth process, and tears or other micrometer-scale gaps in the graphene that form during the membrane manufacturing process. Gas leakage through micrometer-scale tears and nanometer-scale intrinsic defects in macroscopic graphene membranes can severely limit their selectivity and make separation impossible. Indeed, to date gas separation has been demonstrated only in multi-layered membranes prepared from graphene oxide, where permeance is 3-4 orders of magnitude lower than that predicted by simulations across single-layer graphene membranes. Even in these multi-layer membranes, gas transport through defects can play a role in addition to interlayer transport. It is therefore critical to understand transport through intrinsic defects and tears in graphene and to develop strategies to mitigate non-selective leakage if the full potential of graphene membranes for gas separation is to be realized.

In this paper, we quantify the effects of tears and intrinsic defects on the gas permeance and separation performance of graphene composite membranes (GCM) comprising graphene situated on a porous support membrane. We show that independent stacking of two to five layers of graphene is a promising method for reducing the effects of defects while maintaining the nanometer-scale thickness of the graphene layer. The inherent permeance of these multi-layer macroscopic graphene membranes is measured and a gas transport model is developed that can accurately explain the experimentally observed flow rates. From this model, the separate contributions of tears and intrinsic defects to the inherent permeance of macroscopic graphene membranes are estimated. This model is further used to identify design constraints for realizing selectively permeable graphene membranes for gas separation.
Results and Discussion

Graphene composite membranes

Graphene composite membranes were fabricated by transferring graphene grown on copper foil
by chemical vapor deposition (CVD) to a polycarbonate track-etched membrane (PCTEM)
support using a direct transfer method developed previously (Fig. 1a). PCTEMs were chosen as
the porous support because they contain a high density of straight, well-defined pores with
uniform sizes that can be tuned from 10 nm to over 10 µm. The porous support allows the
graphene layer to be handled without damage. The series of parallel pore channels in the
PCTEM also isolate small areas of graphene, such that the graphene composite membrane is
effectively formed from numerous smaller graphene membranes arrayed in parallel. Flow
through any defect in the graphene is restricted by the narrow channels in the PCTEM, which
can potentially prevent leakage from dominating over the flow through intentionally created
selective pores.

In addition, we hypothesized that the effects of leakage may be further mitigated by
independently stacking multiple layers of graphene on the porous support, so that defects in one
layer are covered by another layer. To achieve this goal, the transfer process was repeated to
create GCMs with multiple independently stacked layers of graphene (Fig. 1a). In a multi-layer
graphene composite membrane (Fig. 1b), individual layers of graphene can be distinguished by
the successively darker areas over the white PCTEM. Scanning electron microscope (SEM)
images of GCMs with a single layer of graphene (Fig. 1c) clearly show several polycarbonate
pores with micrometer-scale tears in their graphene coverage. In contrast, SEM images of a five
layer graphene membrane (Fig. 1d) show a much smaller fraction of tears as compared to the
single layer membrane. The images clearly show that independently stacked graphene layers
improve graphene coverage on the PCTEM.

Independent stacking of multiple layers of graphene reduces leakage

Gas transport through the GCMs was measured by supplying the desired gas on the graphene
side of the membrane and monitoring the pressure rise in a downstream reservoir (Fig. 2a, inset).
Transferring a single layer of graphene onto a PCTEM reduced the flow rate of helium through
the PCTEM by approximately 60%, as seen in Fig. 2a. Since no selective pores were created in
graphene, we expect that the residual 40% leakage represents helium flow through intrinsic
defects and tears in the graphene. Next, we examined helium flow rates through GCMs with one
to five layers of graphene, and observed an exponential reduction in the helium flow rate as the
number of graphene layers was increased (Fig. 2). By stacking multiple layers of graphene on a
PCTEM, it was possible to produce a macroscopic graphene membrane with a 99% smaller
permeance (i.e., only 1% leakage) compared to the bare support membrane. Furthermore, the
apparent exponential decay of helium flow rates with increasing number of layers of graphene
suggests that approximately the same fraction of polycarbonate pores are covered by graphene in
each layer, and that the locations of the micrometer-scale tears in each layer are independent of those in the other layers.

Modeling transport through graphene composite membranes

We developed a model to quantitatively understand the effects of defects and independent stacking of graphene layers on the gas transport behavior of the membranes. Since pristine graphene is impermeable to gases, we assume that the net permeance is a result of flow through micrometer-scale tears over the PCTEM pores (see Fig. 1c,d), intrinsic nanometer-scale defects dispersed across areas of continuous graphene, and any intentionally created selective nanopores (Fig. 3).

We account for any micrometer-scale tears by assuming that each layer of graphene covers a fraction \( \gamma \) of the pores in the PCTEM support, with subsequent layers covering the same fraction of pores independent of the other layers (Fig. 3a,b). As a result, a two-layer graphene membrane has polycarbonate pores with zero, one, or two layers of graphene (Fig. 3c). For a general \( N \)-layer graphene composite membrane, the fraction of polycarbonate pores with \( n \) layers of graphene coverage, \( \gamma_n \), is given by,

\[
\gamma_n = \frac{N!}{n!(N-n)!} (1 - \gamma)^{N-n} \gamma^n
\]  

(1)

When multiple layers of graphene are stacked, we assume that intrinsic defects in one layer are covered by graphene in another layer, except where intrinsic defects in all layers randomly align.

The average distance between intrinsic defects in one-layer graphene, \( L_{ip}^{(1)} \), was obtained from the defect size distribution data in Ref. 13 (where we reported characterization of graphene from the same source), combined with the intrinsic porosity of graphene, \( \eta \), i.e., the fraction of graphene area occupied by intrinsic defects. The average spacing between aligned intrinsic defects in \( n \) stacked layers of graphene, \( L_{ip}^{(n)} \), was computed by Monte Carlo simulations (see Supporting Information Section 1 for details). Given the spacing between aligned intrinsic defects, the fraction of polycarbonate pores (of diameter \( D_{PC} \)) with \( n \) layers of graphene that have \( k \) aligned intrinsic defects can then be calculated from the Poisson distribution,

\[
f_{n,k} = \frac{1}{k!} \left[ \frac{\pi}{4} \left( \frac{D_{PC}}{L_{ip}^{(n)}} \right)^2 \right]^k \exp \left[ -\frac{\pi}{4} \left( \frac{D_{PC}}{L_{ip}^{(n)}} \right)^2 \right]
\]  

(2)

To limit complexity, we assumed identical permeance for all intrinsic defects regardless of the size, an assumption that was independently validated (see Supporting Information Section 1.3). The average permeance due to intrinsic defects in \( n \)-layer graphene (\( P_{IP,n}^A \)) was estimated as the equilibrium ideal gas flux through the fraction of graphene area occupied by aligned intrinsic defects, assuming that the intrinsic pores are much larger than the gas molecules. Finally, any selective nanopores intentionally introduced into the graphene will provide a flow path in
parallel to the intrinsic defects. We assume that their density is sufficiently high to represent flow through these nanopores by an average selective pore permeance, \( P_S^A \), defined as the molar flow rate per unit graphene area per unit pressure difference.

Gas transport through the graphene membrane was computed using an equivalent resistance network\(^{18}\) (Fig. 3d). Each branch of this network represents polycarbonate pores with a certain number of layers of graphene coverage and a certain number of intrinsic defects. The graphene composite membrane permeance for arbitrary gas \( A \) is then computed as,

\[
P^A = y_0 P_{PC}^A + \sum_{n=1}^{N} \sum_{k=0}^{\infty} y_n f_{n,k} \left[ \frac{1}{P_{PC}^A} + \left( \frac{4k}{\pi} P_{IP,n}^A \left( \frac{r_{IP}^{(n)}}{\delta_{PC}} \right)^2 + P_S^A \right)^{-1} \right]^{-1}
\]  

(3)

The polycarbonate permeance (\( P_{PC}^A \)) for each gas was measured on a bare PCTEM without any graphene (see Supplementary Information Section 2 and Fig. S9). This leaves three unspecified parameters for this model: (1) the fraction of polycarbonate pores covered by a single layer of graphene, \( \gamma \), (2) the fraction of graphene area occupied by intrinsic defects, \( \eta \), and if applicable, (3) the intentionally created selective nanopore permeance, \( P_S^A \).

Model predicts transport characteristics of independently stacked graphene membranes

The number density and distance between intrinsic defects is an important parameter governing the membrane performance. Power-law fits to the Monte Carlo simulation results for a range of intrinsic porosities (Fig. 4a) show a scaling \( \left( L_{IP}^{(n)} \right)^{-2} \sim \eta^n \), i.e., the average number density of aligned intrinsic defects in independently stacked layers of graphene decreases as the \( n^{\text{th}} \) power of the intrinsic porosity of graphene. This scaling behavior is explained by the fact that when a layer of graphene with porosity \( \eta \) is placed on a previous layer, only a fraction \( \eta \) of the existing defects align with defects in the new layer. Therefore, each successive layer reduces the number of defects by a fraction \( \eta \). The exact coefficients of the scaling relations depend on the intrinsic defect size distribution.

Next, we compared model predictions (Supporting Information Section 1.3) of gas transport through the multi-layer membranes with measured flow rates of gases with a range of molecule sizes (He, N\(_2\), and SF\(_6\), with kinetic diameters of 2.6 Å, 3.64 Å, and 5.5 Å, respectively\(^{19}\)), without any intentionally created selective nanopores (\( P_S^A = 0 \), Fig. 4b,c). Using least-squares fitting, the graphene composite membrane characteristics were well described by \( y = 0.698 \) and \( \eta = 6.87 \times 10^{-3} \). The model results accurately matched the exponential decrease of measured flow rates as the number of layers of graphene was increased from one to four (Fig. 4b). The model also captures the gradual rise in flow rate ratios of the gases with increasing number of graphene layers. Since the intrinsic defects were measured to be larger than the kinetic diameter of the gases, the intrinsic defect selectivity is approximately \( P_{IP,1}^A/P_{IP,1}^B = \sqrt{M_B/M_A} \) from the kinetic theory of gases, which is somewhat higher than that for flow through the polycarbonate.

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pores (Fig. S9b). As a result, increasing the contribution of intrinsic defects compared to tears upon adding more layers of graphene causes a slight increase in the flow rate ratio.

Quantitative agreement between the experimentally measured and theoretical flow rate and flow rate ratio with increasing number of graphene layers provides validation for the model developed here. The model was further verified by comparing its predictions to those of a more detailed model, discussed in Supporting Information Section 1. We considered the possibility that the deviation of the model trends from the measured flow rates for five layer graphene composite membranes (Fig. 4b) is in-plane gas transport between layers of graphene, which scales weakly (inversely) with the number of layers and will eventually govern transport as the number of layers increases. If the deviation for the five-layer membrane is attributed entirely to in-plane transport, we can estimate a conservative lower bound on in-plane transport resistance of $1.2 \times 10^4$ Pa·m$^2$/mol for helium, corresponding to an interlayer diffusivity of $1.2 \times 10^{-15}$ m$^2$/s (see Supporting Information Section 3). However, such a low interlayer transport resistance would cause significant deviation between measured flow rates and model predictions for two, three, and four layer membranes, which we do not observe. Therefore, it is concluded that interlayer transport is not responsible for the deviation between model and measured flow rates for the five layer membranes in Fig. 4b, and the computed in-plane transport resistance represents a very conservative lower bound. The discrepancy between the measurements and model predictions for the five layer membrane in Fig. 4b are thus likely due to damage induced during the repeated mechanical pressing steps, which could become a significant factor at low flow rates.

A recent report in which transport across stacked graphene layers was measured speculated that the increase in flow rate ratio is due to interlayer transport. Our experiments and modeling results indicate that the permeability for interlayer transport in graphene is low and that intrinsic defects alone can account for the increased selectivity observed from stacking successive layers. This result is consistent with measurements of in-plane gas diffusion along a graphene-copper interface, which show significantly lower transport than has been observed between layers of graphene oxide.

**Feasibility of selective transport in the presence of non-selective defects**

We now use the gas transport model to explore whether gas separation may be accomplished in macroscopic, single or few-layer graphene membranes regardless of the presence of defects for two model gases, He and SF$_6$. To do this, we estimated intentionally created selective nanopore permeances, and used these in the model to predict overall membrane selectivities in the presence of intentionally created selective nanopores. Based on molecular dynamics simulations and gas flux measurements by Koenig et al. (assuming that the reported flux corresponds to one measured pore), the achievable resistance of a single selective nanopore to He is estimated to be in the range of $9 \times 10^{19}$ to $2 \times 10^{22}$ Pa·s/mol. We further assume a selective pore density of $10^{12}$
cm² (one nanopore per 10 × 10 nm² area) to calculate the permeance due to selective pores, \( P_{S}^{He} \). Based on the measurements of Koenig et al.,⁸ the selective pore selectivity is set to \( P_{S}^{He}/P_{S}^{SF_6} = 15,000 \).

The available graphene quality (specified by \( \eta = 6.87 \times 10^{-3} \) and \( L_{ip}^{(n)} \) in Fig. 4a) and the graphene permeance due to selective pores are limited by the graphene synthesis and selective pore creation procedures. However, it is possible to precisely control the support membrane pore size and permeance, and to enhance the graphene coverage through the graphene transfer process and the number of layers used. The model predicts that for the 1 µm diameter polycarbonate membranes with ~70% graphene coverage fabricated in this study, it is unlikely that any selectivity will be obtained even with five layers of graphene for the expected range of selective pore permeance and for different polycarbonate pore resistances (top left plot). This lack of selectivity results primarily from the high permeance of polycarbonate pores, causing leakage through micrometer-scale tears to dominate. By increasing the polycarbonate pore resistance to mitigate the effect of leaks through tears, it is possible to achieve modest selectivity with five layers of graphene (top row of plots). Further increase in polycarbonate pore resistance reduces selectivity because flow is then limited by the non-selective polycarbonate pores.

Using polycarbonate membranes with the smallest commercially available pore size of 10 nm shows the potential for much better selectivity, using the experimentally measured resistance of those polycarbonate membranes (center row of plots in Fig. 5). Since the support membrane pore diameter is much smaller than the average distance between intrinsic defects (see Fig. 4a) it tends to isolate the defects to only a small fraction of graphene-covered polycarbonate pores, leaving the majority of the pores covered with defect-free graphene. Three layers of graphene and a factor of 1000 increase of the polycarbonate pore resistance further mitigates leaks, yielding selectivities on the order of 100. With a modest increase in the fractional area coverage of graphene from 70% to 85%, selectivities on the order of 1000 are feasible (bottom row of plots in Fig. 5). These results suggest the feasibility of gas separation using graphene with typical quality obtained by chemical vapor deposition, provided that an appropriate porous support is chosen.

**Dimensionless parameters defining regimes of selective transport**

We now specify dimensionless parameters to define gas permeation regimes through graphene membranes for the purpose of guiding their design and optimization. For simplicity, we focus on membranes with a single layer of graphene and omit notations specifying the number of layers. Given two gases \( A \) and \( B \), under the assumptions that intrinsic defects are non-selective and that the polycarbonate pore resistances for both gases are equal to those under rarefied gas flow, we can express the membrane selectivity \( P_A/P_B \) in terms of dimensionless parameters as follows (see Supporting Information Section 1.3):
\[
\frac{P^A}{P^B} = f\left(\frac{L_{IP}}{D_{PC}}, \frac{P_{PC}}{P^A_S}, \frac{P_{IP}}{P^A_S}, \frac{P_{PB}}{P^B_S}\right)
\]  

The number of intrinsic defects per polycarbonate pore is proportional to \((D_{PC}/L_{IP})^2\). \(P_{PC}/P^A_S\) and \(P_{IP}/P^A_S\) define the permeance of the support membrane and intrinsic defects, respectively, relative to that due to selective pores, while the selectivity of the selective pores is defined by \(P^A_S/P^B_S\) (assumed equal to 15,000 here, see Supporting Information Section 1).

Analytical expressions for the GCM selectivity \((P^A/P^B)\) function in Eq. 4 and corresponding GCM permeance \((P^A/P^A_S)\) are derived in Supporting Information Section 1.3. Given a certain intrinsic defect permeance \((P_{IP}/P^A_S)\), both the GCM selectivity \((P^A/P^B)\) and permeance \((P^A/P^A_S)\) are defined by the choice of support permeance \((P_{PC}/P^A_S)\) and intrinsic defect spacing relative to the support pore size \((L_{IP}/D_{PC})\). This results in a trade-off between the selectivity \((P^A/P^B)\) and permeance \((P^A/P^A_S)\), illustrated for \(P_{IP}/P^A_S = 0.01\) and \(P_{IP}/P^A_S = 1\) for \(\gamma = 0.99\) (Fig. 6a,b). When the permeance of intrinsic defects is small (Fig. 6a), high GCM selectivity can be achieved for any value of \(L_{IP}/D_{PC}\). However, for \(P_{IP}/P^A_S = 1\) (Fig. 6b) high selectivity is only achieved when the spacing between intrinsic defects is large compared to the support pore diameter \((L_{IP}/D_{PC} \gtrsim 1)\). This illustrates that, when the selective pore permeance dominates over the intrinsic defect permeance, intrinsic defects in the graphene over the support membrane pore will not compromise selectivity. However, when the intrinsic defect permeance is significant, it is important that the support membrane pore diameter be sufficiently smaller than the intrinsic defect spacing, so that the intrinsic defects will be isolated to a small fraction of the support pores, leaving a large number of support pores with high selectivity. In addition to the dependence on \(L_{IP}/D_{PC}\), the membrane is selective only for a certain range of permeance \(P^A/P^A_S\). Non-selective transport through tears result in poor selectivity at high \(P^A/P^A_S\), corresponding to high \(P_{PC}/P^A_S\), whereas the dominant non-selective backing membrane resistance prevents selective transport at low \(P^A/P^A_S\), corresponding to low \(P_{PC}/P^A_S\). Selectivity is observed at intermediate \(P^A/P^A_S\) values, when the support resistance approaches the selective pore permeance \(P_{PC}/P^A_S \approx 1\). Further decrease in the support permeance decreases the total permeance by several orders of magnitude for only modest gain in selectivity.

When the support membrane resistance matches that of the selective pores \((P_{PC}/P^A_S = 1)\), high permeance occurs for large \(P_{IP}/P^A_S\) and small \(L_{IP}/D_{PC}\) (Fig. 6c), for which all support pores have highly permeable intrinsic defects in the graphene over them resulting in poor selectivity (Fig. 6d). Under these conditions, \(P^A/P^A_S = P_{PC}/P^A_S = 1\), because the resistance of the graphene becomes negligible compared to that of the support membrane. For low intrinsic defect permeance \(P_{IP}/P^A_S\) or large spacing between intrinsic defects \(L_{IP}/D_{PC}\), transport through intrinsic defects becomes negligible and the permeance approaches \(P^A/P^A_S = (1 - \gamma)P_{PC}/P^A_S + \gamma/(1 + P^A_S/P_{PC})\), as shown in Supporting Information Section 1.4. It is immediately seen from Fig. 6d that, even for a properly balanced membrane, selective transport is possible only if either of the two criteria are met: a) the permeance due to intrinsic defects is small compared to the
permeance due to selective pores \((P_{IP}/P_S^A \lesssim 1)\), OR b) the spacing between intrinsic defects is large compared to the support pore diameter \((L_{IP}/D_{PC} \gtrsim 1)\). The first criterion implies insignificant leakage through intrinsic defects. We find that this is practically unlikely for the currently available quality of graphene, based on the estimated \(\eta\) and permeance due to selective pores, which suggest \(2 \lesssim P_{IP}/P_S^A \lesssim 2 \times 10^4\). However, if leakage due to intrinsic defects is negligible with advances in graphene quality and selective pore creation, the selectivity is governed by micrometer-scale tears. For \(P_{PC}/P_S^A = 1\), the selectivity approaches \(P^A/P^B = (1 - 1/\gamma)/(1 - \gamma)\) for infinite selectivity \((P_S^A/P_S^B \to \infty)\), and is only slightly lower for \(P_S^A/P_S^B = 15,000\) (Supporting Information Section 1.4). The second criterion results in isolation of intrinsic defects. Under these conditions, most of the support membrane pores have no intrinsic defects in the graphene suspended over them and flow through intrinsic defects can be restricted by controlling the support membrane pore resistance, enabling high overall membrane selectivity. Leakage is then limited to the remaining pores that have graphene with intrinsic defects, approaching the same selectivity limit, \(P^A/P^B = (1 - 1/\gamma)/(1 - \gamma)\). The non-selective regime occurs when these two criteria are not satisfied, and it is not possible to achieve selectivity through control of the support membrane resistance. In this regime, the support membrane pore diameter is larger than the average distance between intrinsic defects \((L_{IP}/D_{PC} < 1)\), resulting in most of the support membrane pores having intrinsic defects in the graphene over them. Consequently, the gas flow into the support membrane pores is dominated by the lower resistance path through the intrinsic defects, bypassing the selective nanopores and producing no detectable overall membrane selectivity.

We now address the question of how to select an appropriate support membrane to achieve gas separation for graphene of a given quality, as characterized by the distance between intrinsic defects and the permeance due to intrinsic defects. The support membrane permeance that maximizes selectivity and the corresponding maximum possible selectivity depend on the quality of graphene (Fig. 6c,f). As in Fig. 6d, selectivity requires either low \(P_{IP}/P_S^A\) or high \(L_{IP}/D_{PC}\). Under these conditions, the GCM approaches a limiting selectivity of \(P^A/P^B = 1/(1 - \gamma)\) for \(P_S^A/P_S^B \to \infty\) rather than the selectivity of \(P^A/P^B = (1 - 1/\gamma)/(1 - \gamma)\) obtained for the non-optimal value of \(P_{PC}/P_S^A = 1\) shown in Fig. 6d. For finite \(P_S^A/P_S^B\), the GCM selectivity is maximized at low \(P_{IP}/P_S^A\) or high \(L_{IP}/D_{PC}\) to \(P^A/P^B = [\left(1 - 1/\gamma\right) + 1]/\left(\sqrt{P_S^B/P_S^A} + 1\right)^2\) (Fig. 6f, Supporting Information Section 1.4) with a support membrane permeance of \(P_{PC}/P_S^A = \sqrt{P_S^B/P_S^A}/(1 - \gamma)\) (Fig. 6e). However, as seen by comparing the plot in Fig. 6f to Fig. 6d for \(\gamma = 0.99\), selectivity is not significantly compromised with a membrane permeance of \(P_{PC}/P_S^A = 1\) instead of the optimal value of \(P_{PC}/P_S^A = 0.082\), due to the relatively wide peak observed for high \(L_{IP}/D_{PC}\) in Fig. 6a,b. However, choosing a support permeance \(P_{PC}/P_S^A = 0.082\) drastically compromises the overall membrane permeance, which shows that
approximately matching the support permeance to that of selective pores \( P_{PC}/P_{S} \approx 1 \) is practical, rather than optimizing the support permeance for selectivity.

These results demonstrate that by choosing a support membrane with appropriate resistance and with a smaller pore size than the average distance between intrinsic defects, it is possible to achieve selective gas transport despite the presence of defects. While we have discussed the design of a porous support with isolated cylindrical pores, the results can also inform the design of membranes where the support comprises a thin isotropic layer with controlled permeance. In this case, the thickness of the layer is the characteristic dimension that replaces the support pore diameter \( D_{PC} \) in the above analysis, i.e., it is important to choose a thickness that is smaller than the spacing between intrinsic defects. The results in Fig. 6 can then be used to identify the appropriate support membrane pore size (or thickness) and permeance once the graphene coverage, intrinsic defect characteristics, and achievable selective pore performance have been estimated.

### Conclusions

Realizing the full potential of graphene for use in gas separation membranes on a macroscopic scale will require understanding and controlling the adverse effects of micrometer-scale tears and nanometer-scale intrinsic defects in the graphene. We quantified these effects by measuring the permeance of macroscopic graphene composite membranes to different gases. It was shown that, by stacking individual layers of graphene, it is possible to exponentially reduce leakage through defects in the membrane. A model for gas permeance through macroscopic graphene membranes accounting for micrometer-scale tears and nanometer-scale intrinsic defects was developed and shown to accurately explain measured flow rates. The contributions of tears and intrinsic defects to the overall permeance of the membrane were quantified by fitting this model to the measured permeance values. The results indicate that inter-layer transport is negligible in graphene membranes, and intrinsic defects can give rise to a modest selectivity. The graphene permeance model was applied to estimate selectivities of graphene composite membranes with intentionally created selective nanopores. Although tears and intrinsic defects have a detrimental effect on the overall membrane selectivity, the model shows that by controlling support membrane pore size and resistance, it is possible to design a selectively permeable graphene composite membrane. Model predictions of the optimal support membrane specifications are provided for a wide range of graphene characteristics to guide in the design of defect-tolerant selectively permeable graphene composite membranes for gas separation.

### Methods

*Membrane fabrication*
Multiple layers of graphene were stacked on a polycarbonate track-etched membrane (PCTEM) support by repeated application of the process for transferring a single layer of graphene onto PCTEMs developed in Ref. 13. This procedure is outlined in Fig. 1a. Graphene on copper foil (ACS Materials) grown by chemical vapor deposition (CVD) was first cut into approximately 5 mm squares and the back sides of these pieces were etched for 5 min in ammonium persulfate (APS-100, Transene) to expose the copper and reduce the foil thickness. After rinsing, the piece of graphene on copper foil was mechanically pressed onto a larger PCTEM with 1 µm pores and without a polyvinylpyrrolidone coating (Sterlitech). The remaining copper foil was then completely removed by floating the stack on an ammonium persulfate solution. The resulting composite membrane, comprised of a single layer graphene on a PCTEM, was then rinsed in ethanol before air drying. Subsequent layers of graphene were transferred one-by-one by mechanically pressing graphene on copper foil onto a PCTEM onto which graphene had already been transferred, etching the copper away, rinsing, and drying.

Membrane characterization

SEM images were acquired with the JEOL 6320FV Field-Emission High-Resolution SEM at the MIT Center for Materials Science and Engineering. All images were obtained at an acceleration voltage of 5 kV in secondary electron imaging mode. Carbon tape was used to create a current path between the graphene on the GCM and the microscope stage to reduce charging of the polycarbonate support material. The CVD graphene was characterized by Raman spectroscopy and scanning transmission electron microscopy with the results reported in Ref. 13.

Gas permeance measurements

Gas flow rates through the graphene composite membrane were measured using the apparatus sketched in the inset of Fig. 2a. The membrane was sealed between an upstream pressure line and a downstream reservoir, both initially evacuated. During measurements, the upstream line was continuously supplied with a single gas species at an absolute pressure of 1 atm by a regulated gas cylinder. A pressure transducer was used to measure the rate of pressure rise in the downstream reservoir, resulting in time traces such as those presented in Fig. 2b. The slope of the pressure-time line was used in an ideal gas law relation to determine the flow rate through the graphene membrane. (See Supporting Information Section 2 for additional details on the measurement equipment and procedure).

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**Supporting Information Available:** Additional details on gas permeation models, gas flow rate measurement, and estimation of interlayer flux. This material is available free of charge *via* the Internet at [http://pubs.acs.org](http://pubs.acs.org).
Figure Captions

**Figure 1.** Graphene composite membranes. (a) Membrane fabrication process by direct transfer of graphene from copper foil to a polycarbonate track-etched membrane (PCTEM). The copper foil was etched using ammonium persulfate (APS). Multi-layer graphene composite membranes were fabricated by repeated application of the one layer transfer process. (b) Photograph of a four-layer graphene composite membrane. Scale bar 5 µm. (c, d) SEM images of a (c) one-layer and (d) five-layer graphene membrane and tears. Scale bars are 5 µm and 1 µm.

**Figure 2.** Measurements of gas flow rates through graphene composite membranes with multiple layers of graphene. (a) Effect of the number of layers of graphene on helium flow rate, normalized by the flow rate through a bare PCTEM. Data points represent flow rates measured through different GCMs that were manufactured in parallel. Inset shows a sketch of the gas flow measurement setup. (b) Time traces of pressure rise in the downstream reservoir during helium flow rate measurements for the membranes.

**Figure 3.** Transport pathways considered in the gas flow rate model. (a-c) Illustration of the change in tears and intrinsic defects over polycarbonate track-etched pores upon transferring (b) one and (c) two layers of graphene onto the PCTEM. (d) Equivalent resistance network for flow through GCMs. Note that $R_{PC} = 4(\pi D_{PC}^2 P_{PC}^A)^{-1}$, $R_S = 4(\pi D_{PC}^2 P_S^A)^{-1}$, and $R_{IP}^{(n)} = (L_{IP}^2 P_{IP}^A)^{-1}$.

**Figure 4.** Multiple layer graphene composite membrane flow rate measurements and model fit. (a) Computed average distance between intrinsic defects in multi-layer graphene membranes. (b) Flow rate model fit, giving a coverage of 69.8% ($\gamma = 0.698$) and an intrinsic porosity of 0.687% ($\eta = 0.00687$). Flow rates are normalized by the value for a bare PCTEM. (c) Measured flow rate ratios compared to model fit for flow rates.

**Figure 5.** Model predictions of helium (He) to sulfur hexafluoride (SF$_6$) flow rate ratios for various values of graphene fractional area coverage ($\gamma$), support membrane pore diameter ($D_{PC}$), and support membrane pore resistance ($R_{PC}$).

**Figure 6.** Model predictions of the general GCM permeance characteristics. (a,b) Dependence of GCM selectivity ($P^A/P^B$) on GCM permeance ($P^A/P_S^A$) and intrinsic defect spacing ($L_{IP}/D_{PC}$) for 99% coverage ($\gamma = 0.99$) and intrinsic defect permeance of (a) $P_{IP}/P_S^A = 0.01$ and (b) $P_{IP}/P_S^A = 1$. (c,d) Dependence of (c) GCM permeance ($P^A/P_S^A$) and (b) GCM selectivity ($P^A/P^B$) on the intrinsic defect permeance ($P_{IP}/P_S^A$) and intrinsic defect spacing ($L_{IP}/D_{PC}$) for 99% coverage ($\gamma = 0.99$) when the support membrane is matched to that of the selective pores ($P_{PC}/P_S^A = 1$). (e) Support membrane permeance ($P_{PC}/P_S^A$) that maximizes GCM selectivity ($P^A/P^B$) for various graphene area coverage fractions ($\gamma$), intrinsic defect permeances ($P_{IP}/P_S^A$), and intrinsic defect spacings ($L_{IP}/D_{PC}$), and (f) corresponding maximum GCM selectivities ($P^A/P^B$).
References