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Catalyst Interface Engineering for Improved 2D Film Lift-Off and Transfer

Ruizhi Wang,‡ Patrick R. Whelan,§ Philipp Braeuninger-Weimer,† Stefan Tappertzhofen,‡ Jack A. Alexander-Webber,‡ Zenas A. Van Veldhoven,§,¶ Piran R. Kidambi,¶ Bjarke S. Jessen,¶ Timothy Booth,‡ Peter Bøggild,‡ and Stephan Hofmann¶,*

†Department of Engineering, University of Cambridge, Cambridge CB3 0FA, United Kingdom
‡Center for Nanostructured Graphene (CNG), DTU Nanotech, Technical University of Denmark, DK-2800, Kongens Lyngby, Denmark
§Cambridge Graphene Centre, University of Cambridge, 9 JJ Thomson Avenue, Cambridge CB3 0FA, United Kingdom
¶Department of Mechanical Engineering, Massachusetts Institute of Technology, Cambridge, Massachusetts 02139, United States

ABSTRACT: The mechanisms by which chemical vapor deposited (CVD) graphene and hexagonal boron nitride (h-BN) films can be released from a growth catalyst, such as widely used copper (Cu) foil, are systematically explored as a basis for an improved lift-off transfer. We show how intercalation processes allow the local Cu oxidation at the interface followed by selective oxide dissolution, which gently releases the 2D material (2DM) film. Interfacial composition change and selective dissolution can thereby be achieved in a single step or split into two individual process steps. We demonstrate that this method is not only highly versatile but also yields graphene and h-BN films of high quality regarding surface contamination, layer coherence, defects, and electronic properties, without requiring additional post-transfer annealing. We highlight how such transfers rely on targeted corrosion at the catalyst interface and discuss this in context of the wider CVD growth and 2DM transfer literature, thereby fostering an improved general understanding of widely used transfer processes, which is essential to numerous other applications.

KEYWORDS: 2D materials, CVD, transfer, catalyst, graphene, h-BN

INTRODUCTION

Chemical vapor deposition (CVD) has emerged in recent years as the most promising method for the controlled and scalable synthesis of high quality films of 2D materials (2DM) such as graphene and hexagonal boron nitride (h-BN). While much recent attention has focused on 2DM growth mechanisms on the typically used catalyst materials, many applications require transfer of the 2DM films away from the growth substrate, which has become a serious bottleneck. A widely used method for transfer is to release the 2DM by etching away the complete catalyst foil or film, generally referred to as wet transfer. However, this method can lead to considerable contamination of the 2DM, be it by the polymer, process chemicals, or metal. In addition, it results in increased cost and sustainability concerns for industrial manufacturing, as the catalyst material becomes a waste product.

Approaches to overcome the adhesion between the 2DM film and the CVD catalyst have been developed to preserve the catalyst and to allow transfer from materials that cannot be easily dissolved. A seemingly simple method, as a basis of a so-called dry transfer, is to attach an adhesive layer to the 2DM and to delaminate it by applying mechanical force. A key concern is to sufficiently weaken the 2DM/catalyst interaction to permit the transfer of large areas without tearing and introduction of holes/defects into the atomically thin films. In order to aid their release for a wide range of catalyst materials, several electrochemical methods have been introduced. In the case of bubbling transfer, the enhanced 2DM film delamination is mainly attributed to the generation of hydrogen at its catalyst interface. While this method can be relatively fast, it can significantly damage 2DM films due to the generation of gas bubbles at the interface. Another electrochemical method that has been introduced recently for the transfer of graphene grown on Cu is oxidative delamination transfer (ODT). It enables transfer through the oxidation and following reduction of the copper surface. In addition to the above-mentioned methods, a range of iterations on the chemical modification of the 2DM/catalyst interface have been reported to achieve a better combination of speed and quality of the transfer process. These include for instance water/air exposure in the case of graphene/...
Cu$_{24}^{2+}$–$^{26}$ and the use of a high pH solution to release graphene from platinum (Pt).$^{27}$

While a diverse body of literature on improved 2DM transfer is emerging, there is currently very little fundamental understanding of the underlying mechanisms of transfer. Much progress has been made in understanding the role of the different catalysts and the 2DM interface during the growth process.$^{28,29}$ Noticeable advances have also been achieved regarding the understanding of postgrowth passivation and corrosion applications.$^{30–32}$ However, this knowledge has not yet been used in the context of 2DM transfer. Neither 2DM/catalyst material specific mechanisms nor the categorization into general cases such as weakly and strongly interacting catalysts have been applied to transfer in any detailed level. The development of holistic approaches connecting growth and transfer is critical for future 2DM manufacturing and device integration.

Here, we demonstrate how we can draft improved methods of transfer based on the understanding of the properties of the 2DM/catalyst interface. The goal of this study thereby is not to target record electrical mobility or other 2DM properties. Instead, we seek to understand the mechanisms involved in 2DM transfer and to find a general approach to design and improve transfer methods in particular regarding control and reproducibility while minimizing 2DM contamination. We focus on 2DM grown on Cu, as this is currently the most widely used catalyst. The starting point of our rationale is the weak interaction between 2DMs, such as graphene and h-BN, and Cu, which we previously explored in terms of CVD growth.$^{10,33}$ Thus, gases$^{30}$ and also liquids$^{34,35}$ can intercalate into the Cu/2DM interface. This effect is highly undesired when it comes to surface passivation and corrosion protection.$^{30,32}$ In the context of transfer, however, it offers a new pathway toward delamination. Through intercalation, it is possible to access the 2DM/catalyst interface, to change its chemical composition, and to remove for instance interfacial oxide layers. In this way, delamination is achieved by targeted corrosion at the interface. This mechanism allows for a method of transfer, which we refer to here as lift-off transfer (LOT). In general, there are two pathways for LOT, either involving both interface oxidation and selective removal in a single (LOT-I) or two different process steps (LOT-II). This approach allows us...
to demonstrate the transfer of graphene and h-BN single layer films from Cu. The resulting layers are of high quality regarding surface residues, layer coherence, and electronic properties. Furthermore, the catalyst is retained. We discuss our results in the context of the latest literature on CVD growth, corrosion/passivation, and 2DM transfer. We thereby foster an improved general understanding of transfer processes, which is of crucial importance to many applications.

**RESULTS**

Figure 1a outlines the LOT-I transfer process, both schematically and by corresponding optical photographs of a graphene/Cu sample throughout transfer. As a model system, we use a simple Cu-catalyzed CVD process that gives a continuous single layer graphene film with an average graphene domain size on the order of 10–20 μm (see Experimental Methods section). After the growth period, the samples are stored under atmospheric conditions for up to a month before transfer. It is well-known that storing graphene on Cu for extended periods can lead to local Cu oxidation and contamination with adsorbents. Our process here is designed to minimize the effects of uncontrolled factors. The transfer process starts by coating the samples with a polymer support layer [in the given case poly(methyl methacrylate) (PMMA)] and then floating it on top of a 1 M solution of sodium hydroxide (NaOH). It is important to note that removal of backside graphene is not necessary for the given process. As shown in the photographs in Figure 1a, the Cu foil changes its color at the edge of the sample after a short time on NaOH, which we relate to Cu oxidation. Upon extended exposure, the oxidation front moves inward toward the center of the sample. At room temperature (RT), a graphene film of 1 cm × 1 cm will detach completely from the Cu growth substrate after roughly 12 h. When the temperature is raised to 60 °C, delamination occurs after only 2–3 h. We relate the decrease in process time to an acceleration of the same underlying mechanism as the progression of the oxidation front and overall appearance of the foil remain the same. After rinsing in water, the layer is transferred onto the destination substrate, dried, and then dipped into acetone and isopropanol (IPA) for polymer removal (see Experimental Methods section for details).

LOT-I is based on two main processes. The first is the intercalation of the base into the interface between the 2DM and Cu, which leads to local Cu oxidation. The following selective dissolution of the copper oxide gently releases the 2DM film, which remains floating on the surface of the liquid. The proposed mechanism of LOT-I will be presented in greater detail.
In order to quantify the cleanliness and coverage of the graphene layer after transfer, software-aided optical and Raman analysis was performed. The results are given in Figure 2. The details of the optical mapping method, which has been used previously to assess graphene layers after transfer, can be found in the Experimental Methods section. In short, a number of optical images covering the complete graphene layer are taken. Custom-made software, which is precalibrated for graphene transferred onto 90 nm SiO₂ due to optimal interference contrast, analyzes every pixel of each image. On the basis of the contrast, it discerns whether a pixel represents graphene, SiO₂, or other particles, which are then categorized as residues. This provides a platform to quantify the coverage and cleanliness of graphene after transfer. Figure 2a shows an image consisting of multiple smaller pictures. It was taken by the optical microscope and stitched together during post processing. For the sake of comparison, the image after software analysis is also shown. We analyzed three samples of each of the following methods to transfer continuously grown layers of graphene: LOT-I at RT, LOT-I at 60 °C, wet transfer, and bubbling transfer (see Experimental Methods section for details). The latter two are widely used standard techniques and serve as references. Figure 2b and c show the quantitative results based on the analysis of around 50–100 images across the whole area of about 1 cm² of each respective transferred layer. The fraction of surface covered by pristine single layer graphene after LOT-I at RT is about 99.5%. Residues are found on less than 0.25% of the surface. The graphene cleanliness and coherence slightly deteriorates when the temperature of the transfer is increased. Nonetheless, 98.9% of the surface area consists of single layer graphene (SLG), and only 0.8% is composed of other residues. We suggest the graphene quality decrease for LOT-I at elevated temperatures is due to the impact of a hot alkaline solution on the polymer support layer. On the basis of a similar argument, we point out that reducing the reactivity of the etchant and, as proposed previously,22 improving, though slower, transfer for wet transfer and bubbling, respectively. Nonetheless, LOT outperforms wet transfer (97.8% SLG and 1.4% other residues) and bubbling (96% SLG and 1.4% residues). The spread of residues is also of noticeable difference for the latter two methods. Bubbling results in large local variations of residues, i.e., regions without vs regions with accumulation of residues, as evidenced by the large standard deviation. We suggest that this is linked to the local damage that the film sustains during the violent transfer process,23 which also results in a lower SLG coverage due to holes.

Recent literature highlights that the full width at half-maximum (fwhm) of the 2D peak is an indicator for nanometer-scale strain variations, which negatively impact the electronic properties of graphene.24 Figure 2d shows a comparative Raman analysis of the graphene transferred by LOT-I at RT and by standard wet transfer. Notably for the LOT-I process the fwhm of the 2D peak is only 28 cm⁻¹. This
is well below the average fwhm of samples obtained through wet transfer in our own experiments (41 cm\(^{-1}\)) as well as compared to previous studies, which report a value of around 35 cm\(^{-1}\).\(^{16,43}\) In fact, only dry transfer by directly picking up the graphene flake with a layer of h-BN and placing this heterostructure onto SiO\(_2\) leads to a similar fwhm of about 25 cm\(^{-1}\).\(^{42}\) In order to assess the graphene quality over a larger area, Raman maps of graphene transferred by LOT-I, including the D/G and 2D/G ratios, are presented in Figure 2e and f. For reference, the same measurements were performed on samples obtained through wet transfer. The resulting maps are given in Figure S1. The average D/G peak intensity ratio for LOT-I transfer is below 0.03 (standard deviation <0.01), and the 2D peak is well below the average fwhm of samples obtained through wet transfer. The results are consistent with the reported minimum fwhm of 41 cm\(^{-1}\) for standard wet transfer in our own experiments (41 cm\(^{-1}\)).\(^{42}\) This result is significantly higher than for the wet-transferred samples, which only show 1570 cm\(^{-1}\) s\(^{-1}\) for large grain size and 461 cm\(^{-1}\) s\(^{-1}\) for standard grain size graphene. In the case of wet transfer, we suggest that acid induced polymer cross-linking\(^{16,43}\) and preferential residue accumulation along the grain boundaries\(^{15,45-46}\) are the main contributors to the diminished performance. In particular, the case of cross-linking is avoided in the LOT-I process, as the samples are only in contact with a NaOH solution. In addition to the FET measurement, it is possible to derive the contact resistance by comparing devices with different channel lengths due to the high uniformity of devices. In Figure S3, the channel resistance at 0 V gate voltage is plotted against the channel length. We find a contact resistance of 4400 \(\Omega\)\(\mu\)m, which corresponds to the reported maximum for optical lithography.\(^{47}\)

Thus far, our study has focused on the LOT-I process. As mentioned earlier, based on the rationale of transforming the catalyst interface composition and selectively removing an as-formed interface layer, another approach is possible. The method which we refer to here as LOT-II is depicted schematically in Figure 4a. LOT-II essentially splits the interface oxidation of the catalyst and the dissolution of the oxide layer into two separate processes. There are many ways to oxidize Cu at the interface. As oxygen can easily intercalate into the interface between Cu and graphene,\(^{31}\) a simple method is to leave the sample under ambient conditions for an extended period.\(^{35,30}\) However, in order to speed up the process, we generate Cu oxide by immersing samples in water at 50 °C for 12 h.\(^{54}\) The samples are then spin-coated with PMMA and floated on hydrochloric acid, which was chosen because it attacks solely the Cu oxide but not Cu itself. Within a few minutes, the graphene/PMMA stack detaches itself from the Cu foil, which then sinks to the bottom of the beaker. The film can then be processed identically to LOT-I, i.e., transferring to the destination substrate, drying, and removing the support layer.

Figure 4b shows an optical image of the graphene sample after transfer on SiO\(_2\). Our preliminary results indicate that LOT-II results in a higher degree of surface contamination, as shown by large particles on the sample that are completely absent for LOT-I. We relate this result to the impact of acids on polymers, which is known to be one of the main contributors to contamination in wet transfer.\(^{37}\) Nonetheless, the graphene is
still of good quality as shown by the absence of a significant D peak in the Raman spectrum given in Figure 4c. The 2D peak is well fitted by a single Lorentzian with a fwhm of only 27 cm\(^{-1}\), which is similar to samples transferred by LOT-I. We propose that the generation of a sufficiently thick interfacial Cu oxide, as given for LOT-I and LOT-II, decreases the nanometer-scale strain variations in the graphene layer on the substrate, which is a state that is retained after transfer. This finding suggests that preoxidizing the Cu interface prior to wet transfer could potentially also lead to a reduction of nanometer-scale variation post-transfer and thus improve the overall quality of the 2DM layer.

LOT transfer processes can also be applied to other 2D materials including h-BN, which, while being isostructural to graphene, is a wide band gap semiconductor and has otherwise distinct properties.\(^3\)\(^5\) This allows us to establish that, despite different 2DM properties, the main mechanisms of the LOT transfer are in fact quite similar. Here, we focus on continuous h-BN films with grain sizes of 1−5 μm, which are catalytically grown by CVD directly on Cu.\(^1\)

Following again the LOT-I process outlined in Figure 1a, the h-BN single layer on Cu is spin coated with PMMA, floated on 1 M NaOH, rinsed, and then transferred to the desired substrate. Figure 5a shows optical images of LOT-I and wet transfer samples for comparison. Similar to graphene transfer, LOT-I results in a clean interface with no large visible particles. Figure 5b presents Raman spectra of an as-transferred h-BN sample on SiO\(_2\) by LOT-I and wet transfer. In both cases, the Raman peak position at 1371 cm\(^{-1}\) indicates that the h-BN is single-layered.\(^4\)\(^8\)\(^9\) Raman spectroscopy can be used to assess the contamination by organic residues of an h-BN film. Previous studies have shown that organic contaminants can lead to a significant broad luminescent background.\(^49\) In the spectra given in Figure 5b, it is apparent that wet transfer leads to an increased background in the region between about 2000 and 3500 cm\(^{-1}\), which can be related to residues. This result strongly suggests that LOT-I produces samples of improved cleanliness.

**DISCUSSION**

On the basis of our observations, the following paragraphs provide a detailed discussion on the proposed mechanisms of LOT-I and -II. The transfer process consists of two distinct steps. The first is the intercalation of the solution into the interface and its oxidation. We mentioned previously that during LOT-I the PMMA/2DM/Cu stack is floated on an alkaline solution instead of being immersed. Thus, we assume intercalation only occurs from the edge of the sample. The second step is the dissolution of the interfacial oxide. Due to the presence of Cu oxide after transfer, we suggest that the rate of dissolution is smaller than the rate of oxidation.

The interface oxidation during LOT-I depends critically on how easily atmospheric oxygen can access the 2DM/Cu interface. It has been shown that in the case of 2DMs on Cu, gases and liquids will permeate into the interface.\(^50\)\(^53\) However, this is less due to direct permeation through the 2DM, but rather a result of intercalation through defects or the edges of the sample.\(^34\)\(^35\) Thus, we suggest that, while not being perfect, 2DMs act as a permeation barrier for atmospheric oxygen. In contrast, PMMA does not contribute significantly to the barrier function as it is highly permeable to oxygen.\(^50\)

Whether or not the 2DM film is continuous has an important impact on the overall process. We will first discuss the case in which a continuous 2DM film acts as a permeation barrier for atmospheric oxygen, which we refer to here as reaction path A. Assuming complete dissociation of NaOH, a 1 M solution has a pH of 14. The ongoing chemical reaction can be predicted using the Pourbaix diagram for Cu.\(^53\) Oxidation occurs through the half-reactions given below:

\[
\begin{align*}
2\text{Cu} + 2\text{OH}^- & \rightarrow \text{Cu}_2\text{O} + \text{H}_2\text{O} + 2\text{e}^- \quad \text{(anodic reaction)} \\
\text{O}_2 + 2\text{H}_2\text{O} + 4\text{e}^- & \rightarrow 4\text{OH}^- \quad \text{(cathodic reaction)}
\end{align*}
\]

In highly alkaline solutions, cuprous oxide (Cu\(_2\)O) is not stable. It will corrode by dissolution as cuprite ions (CuO\(_2^{2-}\)) via the following half-reactions:

\[
\begin{align*}
\text{Cu}_2\text{O} + 6\text{OH}^- & \rightarrow 2\text{CuO}_2^{2-} + 3\text{H}_2\text{O} + 2\text{e}^- \quad \text{(anodic reaction)} \\
\text{O}_2 + 2\text{H}_2\text{O} + 4\text{e}^- & \rightarrow 4\text{OH}^- \quad \text{(cathodic reaction)}
\end{align*}
\]

As mentioned in the Results section, when the process temperature is raised to 60 °C, delamination of a 1 cm × 1 cm
sample will result after only 2–3 h as opposed to more than 12 h for the same process at room temperature. We suggest that the reaction path remains identical but that the process time decreases purely due to thermally increased chemical reaction rates. The process of LOT-I is driven by intercalation, oxidation, and Cu oxide dissolution. An increase of temperature accelerates the rate limiting step and thus leads to the improved process speed.

A detailed schematic of the 2DM/Cu interface is presented in Figure 6a. When the sample is first placed onto the NaOH solution, the solution will start to intercalate into the interface. It is important to emphasize that intercalation is crucial to the transfer process. If LOT were based on oxidation and dissolution of the oxide only, the result would be equivalent to wet transfer; i.e., the Cu foil would be completely dissolved. Once the liquid has penetrated the interface, Cu is oxidized to form cuprous ions (Cu+), which then react with oxygen to form CuO. For reaction path A, i.e., atmospheric oxygen is not present, the oxidation will occur uniformly from the edge of the sample toward the center. Previous studies on the oxidation of graphene-covered Cu have claimed that graphene serves as the cathode for the half-reaction leading to oxidation in aqueous media. In our experiments, we have shown that LOT-I leads to identical results for h-BN or graphene. This demonstrates that the presence of graphene is not imperative for the cathodic half-reaction; i.e., the corrosion reaction can take place without a conductive 2DM. Instead, we suggest that just as for bare metal surface corrosion in aqueous media, certain regions on the surface act as local anodes and certain others as local cathodes, thus forming local cells for the overall reaction.

However, the presence of graphene can change the spatial distribution of local cells due to its function as a permeation barrier, as will be discussed below.

The quantity and location of Cu2O that is formed depends critically on the permeability of the PMMA/2DM stack since PMMA is permeable to oxygen. In reaction path A, the continuous 2DM film acts as a permeation barrier. When the 2DM film is not continuous, atmospheric oxygen can access those regions that are not covered by the 2DM. The reaction changes to what will be called reaction path B. The process in place is schematically depicted in Figure 6b. The given situation is similar to oxygen mediated localized oxide formation in the case of metal corrosion in aqueous media. A well-known example is the iron oxide formation at the edge of a water droplet on an iron surface. In our case, the Cu region without graphene coverage becomes the cathode of the half-reaction due to the local abundance of oxygen. Cu oxide is now primarily formed at these locations, whereas very little is formed underneath the graphene layer. Indeed, we observe that, during LOT-I transfer, discontinuous graphene layers do not delaminate—not even after numerous days. However, when discontinuous samples are immersed completely in NaOH instead of floating at the surface (like LOT-I), the duration of transfer is identical to that of continuous layers in LOT-I. We relate this to the fact that the influence of atmospheric oxygen is alleviated when the sample is completely immersed. In order to achieve an even better understanding of the influence of oxygen, future studies could involve LOT-I using degassed electrolytes in an oxygen-free environment or less permeable polymers.

Previous studies have shown that while water-assisted oxidation does facilitate transfer, ultimately delamination does not occur. Therefore, both oxidation and dissolution are of critical importance for LOT-I transfers. In fact, dissolution can be indirectly observed during LOT-I. When a large piece of graphene on Cu is floated on a small quantity of 1 M NaOH (e.g., four pieces of 1 cm × 1 cm in 100 mL of 1 M NaOH solution), one can observe that the solution will turn blue after an extended period (about 48h). This can be related to the presence of CuO2-, which has a blue color. The solubility of CuO2- is only 10⁻⁶ to 10⁻⁴ M between pH 13 and 14 at room temperature. We observed that if the quantity of NaOH solution is not sufficient, as in the example given above, delamination of the 2DM does not occur, despite oxidation of the Cu foil.
Besides temperature, the concentration of the solution and thus the pH also have a major impact on the transfer characteristics and mechanism. The maximum pH of 15.4 as dictated by the solubility of NaOH, which is 25 M, gives an upper limit to the parameter space. In order to find the dependency of LOT-I on the concentration of the solution, experiments were performed at pH 15, i.e., using a 10 M solution. The first consequence of raising the pH of the solution is the increase in solubility of CuO$_2$.$^{32}$ However, at extremely high pH, e.g., pH 15, the reaction mechanism itself changes. Cu will directly dissolve, without the intermediate step of oxidation. The reaction then follows:$$\text{Cu} + 4\text{OH}^- \rightarrow \text{CuO}_2^{2-} + 2\text{H}_2\text{O} + 2e^-$$ (anodic reaction)$$\text{O}_2 + 2\text{H}_2\text{O} + 4e^- \rightarrow 4\text{OH}^- \quad \text{(cathodic reaction)}$$

Using 10 M NaOH, we successfully transferred discontinuous graphene layers. We suggest that since dissolution can take place without prior oxidation, localized oxidation plays a smaller role in the overall process.

Compared to LOT-I, the mechanism behind LOT-II is much simpler. It is known that Cu oxidation under a 2DM can be achieved by leaving samples under ambient condition for extended periods of time.$^{30,50}$ However, in order to achieve sufficient oxidation in a well-controlled and fast way, we chose to oxidize the Cu foil in our experiments by immersing the sample in water at 50 °C. The oxidation reaction is the same as for LOT-I without atmospheric oxygen. When the sample is then floated on HCl, the oxide is etched according to the reaction$$\text{Cu}_2\text{O} + 2\text{H}^+ \rightarrow 2\text{Cu}^+ + \text{H}_2\text{O}$$

As Cu is stable to hydrochloric acid, the surface oxide is selectively etched.

We demonstrated lift-off transfer of 2DM via LOT-I and LOT-II, which both make use of the same underlying principle of local interfacial oxidation and oxide dissolution. The focus of our discussion has thereby been on the chemical processes at the interface. However, we emphasize that the chemicals used in the transfer process must also be compatible with both the 2DM and the polymer stack. Previous literature highlights that for wet-transfer, the PMMA contamination can potentially be a consequence of acid induced cross-linking.$^{16,43,54}$

**EXPERIMENTAL METHODS**

Graphene Growth. Graphene growth was conducted on a 25-μm-thick Cu foil with 99.8% purity (Alfa Aesar). All samples were grown in a commercially available Aixtron Black Magic Pro 4-in. PECVD system. The growth process begins by filling the chamber with a mixture of argon and hydrogen at a ratio of 4:1 (200 sccm/50 sccm). The temperature is then increased at a rate of 100 °C/min. The foil is annealed at a growth temperature of 1065 °C for 30 min and for large domain graphene samples, for 120 min. The precursor gas mix, which consists of 250 sccm argon, 26 sccm hydrogen, and 9 sccm methane for standard and 7 sccm for large domain graphene (0.1% diluted in argon), is then injected into the chamber. The growth time depends on the desired graphene grain size. At all stages of the process, the pressure is kept at 50 mbar. Post-growth, the chamber is cooled naturally to room temperature in an argon only atmosphere.

Graphene grain size control is achieved by a combination of foil pretreatment and growth time variation. For standard grain size graphene (10–25 μm), the foil is not pretreated and growth performed over 45 min. Large grain size graphene (200–400 μm) is obtained by using electropolished Cu foils and a growth time of 4 h. The grain size is determined by stopping the respective growth process shortly before the nuclei merge completely to form a complete film and SEM/optical microscope measurement.

h-BN Growth. h-BN growth used processes reported elsewhere.$^{11}$ Two different processes have been used for the samples used in the given studies. For one of these, 25-μm-thick Cu foil with 99.8% purity (Alfa Aesar) was initially cleaned by sonication in ammonium persulfate and rinsed thoroughly in deionized water. The foil was subsequently heated in H$_2$ to 1050 °C and annealed for 60 min in a hot wall furnace. Liquid borazine was used to initiate h-BN growth. However, transfer should be possible for weakly interacting systems such as 2DM on Pt. In fact, a prior study has demonstrated that when graphene/Pt is floated on high pH solutions at high temperature, delamination will occur.$^{27}$ We suggest that the main underlying mechanism is similar to the one in LOT-I, i.e., corrosion at the 2DM/metal interface.

**CONCLUSIONS**

The interaction between a 2DM and its growth catalyst is of crucial importance for all transfer methods that seek to overcome the interfacial adhesion. Weak interaction and weak mechanical adhesion allow for the intercalation of reactive species into the 2DM/catalyst interface. On the basis of this understanding, we have demonstrated that the interplay between the 2DM and the catalyst is not only important for growth but also decisive for transfer processes. We introduced an approach, referred to here as LOT, which relies on the selective removal of the interface layer between the 2DM and its growth catalyst. While 2DM are often discussed in the context of corrosion protection, we have shown how targeted corrosion at the interface can actually enable improved transfer.

Overall, we have demonstrated the feasibility of LOT for both graphene and h-BN. The graphene samples that were obtained were characterized by multiple methods, revealing their high quality over a large area. Using FETs made by LOT, we point out the importance of clean processing, as these clearly outperform devices made using larger grain size graphene, but nonideal transfer methods. We also show that h-BN can be easily transferred and demonstrate the cleanliness of the layer. Further experiments based on our rationale, such as LOT-II, highlight the value and usefulness of the understanding of the interfacial processes. It is shown thereby that LOT can serve as a general process approach for improving and ultimately scaling up 2DM transfer.
for 90 min and Cu foil subsequently cooled to room temperature. For
the other process, 25-μm-thick Cu foil with 99.8% purity (Alfa Aesar)
was directly used for growth. The samples were heated in an Axtron
Black Magic PECVD system in a vacuum to 1000 °C. Liquid borazine
was used as the precursor, and growth took place over 5 min, after
which the samples were cooled down to room temperature.

Lift-Off Transfer I. Graphene or h-BN on Cu as received is spin-
coated with poly(methyl methacrylate) (MicroChem PMMA 950 K
A4) at 3000 rpm for 40 s resulting in layer of 250 nm thickness, which
is then baked for 90 s at 180 °C. The sample is then floated on a 1 M
solution of NaOH (Sigma-Aldrich, product number 306576, 99.999% purity)
dissolved in deionized water. At room temperature, the
PMMA/2DM stack detaches after about 12 h for a sample size of 1 cm
× 1 cm, at 60 °C after only 2–3 h. The sample is then rinsed
repeatedly with deionized water, transferred onto the target substrate,
and dried at 50 °C for 1 h. In the last step, the polymer is removed by
rinsing in acetone and IPA.

Lift-Off Transfer II. Graphene on Cu is immersed in deionized
water and kept at a temperature of 50 °C for 12 h. The sample is then
coated identically to LOT-I and floated on hydrochloric acid (Fisher
Scientific, product code 10251183, 37%) for 5 min, upon which the
sample will detach. After rinsing in water, the sample is transferred
onto the target substrate and dried at 50 °C for 1 h before removing
the polymer by acetone and IPA.

Wet Transfer. The samples are spin-coated identically to LOT.
The backside graphene is removed with oxygen plasma (oxygen partial
pressure of 50 mbar, plasma power of 50 W, duration 10 s), and Cu is
etched using 0.5 M iron chloride over about 2 h. The remaining
transfer process is identical to LOT.

Optical Characterization. Optical maps were made by capturing
and stitching images of the entire sample area at a resolution of ~0.5
μm/pixel. A combination of normalizing all images for intensity
variations and using a high-performance motorized XYZ stage allows
us to generate arbitrarily large and high-resolution optical maps.

To calculate the coverage of graphene and the amount of residues,
we start by calculating the wavelength-dependent contrast of
graphene.33 Using the red, green, and blue (RGB) spectral response
functions of the CCD sensor, we can obtain the numerical values for
pixels of silicon oxide, single layer graphene, as well as bilayer
graphene.34 Pixels that do not fall into any of these three categories
are labeled as residues, and the coverage values presented in this paper
thus represent a lower bound, as pixels containing both graphene and
residues will only count toward the coverage of residues.

Raman Spectroscopy. All measurements were performed with a
commercially available Renishaw Raman system. A 532 nm laser at a
power below 1 mW was used to avoid graphene damage. Spectra were
taken with a 50X objective lens.

Device Fabrication and Characterization. FETs were fabricated
in a two-step photolithography process. The contacts are deposited by
sputtering 70 nm of nickel and subsequent photore sist lift-off. All
samples were measured in air directly after processing without
additional annealing. The drain current $I_D$ is measured while applying a
drain-source voltage $V_{DS}$ of 10 mV and sweeping the gate voltage $V_{GS}$
from 0 V to 60 V. The maximum transconductance $g_m$ is obtained by
fitting the transfer curve. The mobility is calculated using the expression

$$\mu_{FET} = \frac{g_m}{W/C_g}$$

$W$ and $L$ represent the width and length of the channel, respectively.
$C_g$ is the gate capacitance per area, which is 11.6 × 10^{-12} F cm^{-2} for 300
nm SiO₂.


