Fluid-like sliding graphenes but with solid-like out-of-plane compressive rigidity offer unique opportunities for achieving unusual physical and chemical properties for next-generation interfacial technologies. Of particular interest in the present study are graphenes with specific chemical functionalization that can predictably promote adhesion and wetting to substrate and ultralow frictional sliding structures. Lubricity between stainless steel (SS) and diamond-like carbon (DLC) is experimentally demonstrated with densely functionalized graphenes displaying dynamic intersheet bonds that mechanically transform into stable tribolayers. The macroscopic lubricity evolves through the formation of a thin film of an interconnected graphene matrix that provides a coefficient of friction (COF) of 0.01. Mechanical sliding generates complex folded graphene structures wherein equilibrated covalent chemical linkages impart rigidity and stability to the films examined in macroscopic friction tests. This new approach to frictional reduction has broad implications for manufacturing, transportation, and aerospace.

The development of advanced superlubricant materials, defined by a coefficient of friction (COF) near or lower than 0.01, is essential for the sustainability and achieving higher performance from mechanical devices.[1–3] Although pristine graphene has a desirable incommensurability in its lattice plane relative to other materials and can suppress mechanical wear,[4–9] the creation of incommensurate stable contacts at macroscopic interfaces still poses fundamental challenges. In particular, superior lubricant materials will have both the properties of a liquid and a solid.[10–14] At one extreme, a thin layer of a liquid (water) can behave as a lubricant, and hydroplaning behavior reduces frictional contact between tires and a road.[15] Inspired by hydroplaning and graphite, we have designed a fluid-like reconfigurable graphene matrix that shows quasi-hydrodynamic and ultralow frictional behavior. We hypothesized that the graphene matrix with chemically dynamic (reversible) connections can transform into molecular bearings that simultaneously have a large resistance to further compression and a stable top surface with sustained lubricity with ultralow friction.

To create a scalable ultralow frictional material, we designed a coating wherein basal-plane-functionalized graphenes, produced by electrochemical activation, are covalently attached to structurally rigid triptycene molecular cores. Specifically, the interconnected graphene matrix is produced by the reactive formation of Meisenheimer complexes between graphene functionalized with 3,5-dinitrophenyl groups and triaminotriptycene (Triamino-T).[16,17] Triamino-T was designed as a highly stable rigid 3D interlocking group that reversibly covalently links reactive functionalized graphenes into an interconnected matrix that is dynamically configured by physical sliding motions. The COF of the resultant triaminotriptycene-functionalized graphene (FG-T) reaches the ultralow regime (0.01), whereas graphene functionalized with 3,5-dinitrophenyl groups, but lacking Triamino-T modification, shows a higher and erratic friction as a result of chemical interactions with the interacting surfaces. The 3,5-dinitrophenyl-functionalized graphene is reacted with Triamino-T to give a material (FG-T+) that evolves into stable self-leveling tribolayers during sliding. The structural rearrangement of the graphene matrix provides a mechanism to achieve robust superlubricity at both the micro- and macroscales.

The FG-T+ composition (Figure 1a; Figure S1, Supporting Information) undergoes reversible Meisenheimer complexation reactions that provide a reconfigurable interlocking component. Here the NH2 groups of the Triamino-T can be bonded to up to three different 3,5-dinitrophenyl groups, and the stoichiometry of FG-T+ ensures residual reactive primary amines (Figure S2, Supporting Information). The FG-T+ film structure
equilibrates through transient chemical configurations when mechanical shearing (sliding motion) is applied to the material. The reversible reconfiguration of the films under sliding against a diamond-like carbon (DLC)–coated ball further enables self-leveling (recoating) of damaged regions (Figure S3, Supporting Information), which demonstrates strong adhesion and “complete wetting” between FG-T+ and steel substrate. The final matrix contains a highly crosslinked FG-T network with 3D rigid triptycene interlocking units between graphene sheets bond in Meisenheimer complexes (Figure 1a).[18] We postulate that the FG-T+ film is segmented into viscous films on the top surfaces, and a core connection to the substrate through the elastic graphene matrix. These layers help to induce viscous flow on top-surface and to guard against wear, thereby producing quasi-hydrodynamic behavior with reduced friction.[19]

The solid 3,5-dinitrophenyl-functionalized graphene (FG) film shows transient shear thinning behavior (Figure 1b). Conversely, dynamic linkages of FG-T+ lead to Newtonian fluid-like behavior, perhaps promoted by the equilibrating Meisenheimer complexation and the associated changes in graphene structure (Figure 1b). The addition of excess Triamino-T lowers the viscosity from the beginning shear rate of 5.0 s⁻¹, and this behavior for the FG-T+ films is consistent with energy dissipation as a result of dynamic rearrangement of the Meisenheimer complexes (Figure 1b).[20,21]

To further probe viscoelastic properties of these reversible and reconfigurable complexes, we have determined the storage modulus (G’) and loss modulus (G”) at different frequencies of oscillating strain. FG and FG-T show a predominant elastic response over the entire frequency range (G’ ≫ G”), whereas the viscoelastic behavior of FG-T+ shows continuous changes over the frequency spectrum investigated (Figure 1c). In case of FG, we note that the decrease in the loss modulus at 7 rad s⁻¹ is consistent with chemical degradation, wherein continuous shearing induces mecanochemical reactions on graphene. Subsequent mecanochemical damage or physical removal of material results in further change the loss modulus contribution during shearing (vide infra). The gradual increases in the viscoelastic response of the FG-T+ film is consistent with our proposed chemical/structural evolution that occurs without chemical degradation. In total, the rheological results are self-consistent with reversible Meisenheimer complex formation between the triptycenes and graphene to give a fluid-like matrix for reducing frictional behavior. These mechanical dynamics offer insights into the kinetics of the reconfigurable bonding in functional graphenes.

Friction tests were conducted by coating stainless steel (SS) with the different graphenes and sliding a DLC-coated SS ball, using a custom-built linear macro-tribometer (Figure 2a; Figure S4, Supporting Information). Initially, the COF of unfunctionalized exfoliated low defect graphene (UEG) is ≈0.022
(Figure 2b), a value comparable to the previously reported graphenes and graphene oxides (GOs), thereby confirming the system’s performance (Figure S5, Supporting Information).[22–24] The UEG is produced from a dimethylformamide (DMF) dispersion of a Hyper-3-Stage-1 graphite intercalation compound (GIC),[16] and although it forms a low-frictional tribolayer, the stability of the tribolayer on the SS substrate was not satisfactory, as it easily peels off or balls up on the substrate, showing poor adhesion and incomplete wetting. To create a stable layer, the coating materials need to remain strongly associated with the surface during mechanical movement (Figure S6, Supporting Information). Friction experiments were repeated three or more times for each sample (one cycle = two linear-sliding motions). c) The average COF of the uncoated SS, FG, FG-T, FG-T+, and Triamino-T films.

d) The proposed dynamic model of reversible Meisenheimer complexation of graphene with Triamino-Txs to create reconfigurable networks “bearing” behavior. First, unrestricted Triamino-T shows unstable sliding motion with temporary low COF in our system. Second, the incompatibility of UEG-T restricts the formation of a stable matrix during sliding. Third, the molecular bearing motion of partially restricted Triamino-T is effectively stabilized in a dynamic FG-T/FG-T+ matrix, maintaining low COF. e) The FG-T+ friction with increasing sliding speed tests from 16 to 40 mm s$^{-1}$ with >0.01 above a 26 mm s$^{-1}$ sliding speed in the quasi-hydrodynamic lubrication regime. f) Load dependency of the COF of FG-T+.

Figure 2. Illustration of COF and results for graphene films. a) Schematic illustration showing the geometry of a linear tribometer equipped with a DLC-coated stainless steel ball. The friction tests were performed with graphenes and/or Triamino-T to demonstrate the synergetic coupling effect of reversible Meisenheimer complexes and molecular bearings. b) The COF for a DLC ball sliding at 16 mm s$^{-1}$ and a contact pressure of 0.3–0.4 GPa, against UEG (COF = 0.022 ± 0.003), UEG-T (COF: unstable), FG (COF = 0.036 ± 0.005), FG-T (COF = 0.021 ± 0.002), FG-T+ (COF = 0.011 ± 0.001), and Triamino-T (COF: unstable) films on an SS substrate in N$_2$ environment (relative humidity ≈ 3%). Friction experiments were repeated three or more times for each sample (one cycle = two linear-sliding motions). c) The average COF of the uncoated SS, FG, FG-T, FG-T+, and Triamino-T films.
motion give rise to a natural gradual loss of graphene coverage from metallic substrates in macroscopic friction tests. As a result, graphene tribolayers can undergo detrimental transformations in very short periods. These adhesion challenges are the reason that typical multilayer graphene systems, which are nominally expected to have the low frictional properties, fail to provide stable superlubricity.[7]

FGs, in general, have enhanced interactions with most substrate materials and hence have utility for increasing the stability of a tribofilm (Figure 2b). However, the high COF observed for the FG used in this study appears to produce adhesive interactions between contacting materials, but without special additives, this material degrades and does not produce stable tribofilms in the friction tests. These results are consistent with previous atomic force microscopy (AFM) friction measurements on fluorinated graphene,[25,26] wherein functionalized graphene monolayers exhibited higher friction forces relative to pristine graphene. In short, the friction originates from basal plane functional groups that increase van der Waals interactions. To overcome these seemingly intrinsic limitations associated with functionalized graphene, we targeted the generation of dynamic nanostructures to realize a stable ultralow frictional state.

The FG-T matrix relaxes under applied stress through reversible chemical bonding events with Triamino-T. We observed the further reduction in COFs with FG-T+, which produces an ultralow frictional state (Figure 2b,c) comparable
to UEG-nanodiamond combinations (Figure S8, Supporting Information). We attributed this result to the reconfiguration of the stable graphene matrix, wherein the molecular cross linker connects graphene surfaces and stabilizes bilayers and/or folded structures to create dynamic “molecular bearings” with mechanical stiffness and stability (Figure 2d; Figure S9, Supporting Information). This process gives a dense stable tribolayer with liquid-like properties, which allows for self-leveling of the lubricant layer, to counteract the continuous mechanical removal caused by the sliding and pressure gradient. In contrast to FG-T/FG-T+, the mixture of UEG and Triamino-T (UEG-T) exhibits unstable sliding behavior due to the lack of the chemical function for stabilizing bearing molecules (Figure 2b,d). The friction coefficient of FG-T stabilizes at 0.011 at 16 mm s$^{-1}$ and drops to 0.01 at 26 mm s$^{-1}$ (Figure 2e). The FG-T+ also shows low frictional behavior (COF < 0.015) at loads varying from 3 to 1.1 N (Figure 2f). After long-term cycles, we observed the lower wear rate of DLC-coated SS ball against an FG-T+ (0.94 × 10^{-9} mm$^3$ N$^{-1}$ m$^{-1}$) on SS as compared to uncoated SS (9.94 × 10^{-9} mm$^3$ N$^{-1}$ m$^{-1}$) (Figure S10, Supporting Information). FG-T+ also outperformed an uncoated SS in ambient air and initially showed reduced friction for an SS/SS pair (Figure S11, Supporting Information).

To gain insight on the tribochemical reactions and how FG-T and FG-T+ promote stable ultralow frictional states, we have conducted Raman and X-ray photoelectron spectroscopy (XPS) analyses of the as deposited materials and the mechanically produced tribolayers (Figure 3; Figure S12, Supporting Information). In general, graphenes slip by each other and then evolve into graphene tribolayers with the reduced COF against the DLC-coated ball. The final state of graphene on an SS substrate after sliding can produce changes in the Raman 2D- and D-bands, which are quantified by the D-band to the G-band ratio, $I_D/I_G$. An increasing $I_D/I_G$ indicates the introduction of defects in the graphene layers (Figure 3a), and this is readily apparent for UEG. In addition, the G-band (1591 cm$^{-1}$) for the lubricated graphene shifts after sliding, compared with that of pristine graphene (1580 cm$^{-1}$), which also indicates changes in chemical or physical state. This is expected as the G-band...
is sensitive to damage within the graphene sheets. Additionally, XPS analysis reveals increases in the relative ratios of sp² C−C (≈285.5 eV) to sp³ C−C peaks (≈284.5 eV), which is also an indication of mechanically induced chemistry (Figure 3a).

FG initially has a high density of pendant groups (Raman \( I_D / I_G \) ratio of ≈0.93 in Figure 3b) with a stronger interaction with the DLC surface and a higher frictional resistance toward sliding. The nitrogen-containing functional groups are removed with sliding (Figure 3b), and the XPS N 1s signals associated with functional groups on the surface of graphene are not detectable after sliding. However, the XPS signals associated with sp³ carbons (≈285.5 eV) and the Raman \( I_D / I_G \) remains high (Figure 3b; Figure S9, Supporting Information). The fate of removed nitrogen-containing functional groups and the nature of the sp³ Cs are not known; however, mecanochemical reactions are expected under the applied load.\[26,31–33\]

The addition of the triptycene stabilizes the 3,5-dinitrophenyl groups on the graphene, and in contrast to FG the XPS N 1s signals associated with the 3,5-dinitrophenyl functional groups of FG-T and FG-T+ are retained after sliding (Figure 3c,d). Although it is a challenge to experimentally probe the precise physical and chemical interior states of the graphene matrix, it would appear that FG-T+ is stabilized by the dynamic covalent chemically interlocking reactions within the material (Figure S9, Supporting Information).\[34\]

Tribolayers are produced after sliding tests as shown in Figure 4. Scanning electron microscopy (SEM) images (Figure 4a; Figure S6, Supporting Information) reveal that with frictional testing the graphene is partially removed and reconfigured. The formation of a stable surface-bonded tribolayer at a contacting surface reduces wear by minimizing contact of material pairs.\[39\] Our lubricant with its quasi-hydrodynamic nature is initially in a form that can be characterized as crumpled graphene nanoballs. The mechanical sliding under load then leads to a coalescence of particles to produce continuous solid lubricant film that strongly adheres to and completely wets the substrate. The very rigid framework within these films results from the interlinking of graphenes by Meisenheimer complexation.

To observe the structural progression of our graphene lubricant, we performed microstructure and surface analyses of tribolayers after sliding using transmission electron microscopy (TEM) and AFM (Figure 4; Figure S13, Supporting Information).\[36–38\] As shown in Figure 4b, that graphenes initially appear as a continuous interconnected material (Figure 4b). During sliding at a stress of 0.4 GPa, the matrix undergoes compression and structurally evolves from a spherical configuration to compressed interconnected layers that have a spacing of 11.6 Å, which is consistent with the interlayer organic linkages (Figure 4c). The tribolayer of FG-T+ produced by the sliding is reasonably flat with ±5 nm surface features (Figure 4d; Figure S14, Supporting Information), which is also consistent with a compressed layered structure. This planar tribolayer allows DLC to slide and only encounter relatively small energetic barriers. The high out-of-plane rigidity of the FG-T+ tribolayer is also critical and compares favorably to the bare SS (grade 316). Nanoindentation reveals a hardness with ≈5.3 GPa for the SS 316 substrate and ≈7.3 GPa for the tribolayer of FG-T+ (Figure 4e). The higher value of hardness is also consistent with a strongly textured FG-T+ tribolayer.

In summary, we have demonstrated that FG-T+ forms a fluid-like graphene matrix that produces stable tribolayers and a low frictional force against the DLC (COF ≈ 0.01). The dynamic reactivity and flexibility of functional graphene and the rigid frame of triptycene combine to produce adherent and shearing structures under compressive sliding deformation. This approach to an ultralow frictional state is potentially generalizable and is an important step to understanding how to control friction at the macroscale.

Supporting Information
Supporting Information is available from the Wiley Online Library or from the author.

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I.J. and G.P. contributed equally to this work. T.M.S., J.L., and I.W.H. directed the project. I.J. synthesized graphene materials and G.P. conducted the friction measurement apparatus. I.J. and G.P. carried out experiments and the data analysis. P.W. synthesized triamimotriptycene. All the authors discussed the results and contributed to the writing of the manuscript. This work was supported by the US Army Research Office through the use of facilities at the Institute for Soldier Nanotechnologies under Contract Number W911NF-13-D-0001 and National Science Foundation, DMR-1809740. The authors appreciate Maggie He for synthesizing the 3,5-dinitrobenzenediazonium salt.

Conflict of Interest
A patent has been filed on this work.

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coefficient of friction of 0.01, fluid-like graphene, molecular bearings, triptycene, ultralow friction

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Dynamic Fluid-Like Graphene with Ultralow Frictional Molecular Bearing

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Dynamic Fluid-Like Graphene with Ultralow Frictional Molecular Bearing

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Materials and Methods

Synthesis of 3,5-dinitrophenyl functionalized graphene (FG): Highly oriented pyrolytic graphite (HOPG) was used as the cathode in a three-electrodes electrochemical cell, containing 1.5 M tetrabutylammonium perchlorate (TBAP), acetonitrile (MeCN, 2 mL) and dimethylformamide (DMF, 2 mL) (Ag/Ag+ as reference electrode and graphite as counter electrode). The application of continuous voltage ramp (−3.05 to −3.25 V as −2 μV s⁻¹) induced the intercalation of tetrabutylammonium cations (TBA⁺) into graphene interlayers, resulting in Hyper-3-stage-1 graphite intercalation compound (GIC). Graphene and functionalized graphene dispersions were obtained by the following methods: (i) The Hyperstage-1 GIC was sonicated in DMF for exfoliation and then removal of the intercalated TBA⁺ was achieved by washing with MeCN and DMF. (ii) For covalent functionalization, the Hyper-3-Stage-1 GIC was added carefully an electrochemical cell, containing 0.1 M 3,5-dinitrobenzenediazonium tetrafluoroborate, 1 M TBAP and 4 mL MeCN solution. Again, continuous voltage ramp (−1.40 to −1.80 V as −2 μV s⁻¹) was passed through the Hyper-3-Stage-1 GIC. FGs were spontaneously exfoliated from the Hyper-3-Stage-1 GIC. Then the functionalized graphene sheets were collected through vacuum filtration and thoroughly washed with MeCN and DMF.

Synthesis of Trinitrotriptycenes a and b (Figure S1): Triptycene (2.0 g, 8 mmol) and concentrated HNO₃ (80 mL) were added in a round flask. The mixture was heated to 75 °C for 24 h. After cooling down to room temperature, the solution was poured into ice water. The precipitate was filtered, washed with ice water and dried. The product was further purified by column chromatography to afford a and b as white solids (a/b = 2.9/1, 1.3 g, 42% yield). Compound a (major): ¹H NMR (600 MHz, CDCl₃): δ 8.35-8.32 (m, 3H), 8.07-8.04 (m, 3H), 7.64-7.63 (m, 3H), 5.83 (s, 1H), 5.81 (s, 1H).

Synthesis of Triaminotriptycenes (Triamino-T) c and d (Figure S1): Trinitrotriptycenes a, b (1 g, 2.5 mmol) and MeOH (40 mL) were added in a flame-dried Schlenk flask. Palladium on activated charcoal was added to the solution, bubbling with hydrogen for 10 h. The mixture was then filtered through a pad of celite. The solvent was evaporated to give Triamino-T as yellow solids (c/d = 3.0/1, 0.73 g, 95% yield). Compound c (major): ¹H NMR (600 MHz, CDCl₃): δ 7.07-7.04 (m, 3H), 6.73-6.71 (m,3H), 6.27-6.22 (m, 3H), 5.04 (s, 1H), 5.02 (s, 1H), 3.43 (br.s, 6H).

3,5-dinitrophenyl-triaminotriptycene functionalized graphene (FG-T): Figure S1 illustrates the synthesis of the functionalized graphene with 3,5-dinitrophenyl groups anchoring triaminotriptycene groups. Meisenheimer complexes form by the addition of Triamino-T to the FG (50 mg : 4 mg weight ratio) in DMF. And then FG-T was collected through a polytetrafluoroethylene (PTFE) filter membrane with 0.1 μm pore size by vacuum filtration. FG-T+: 4 mg of Triamino-T was added to the above FG-T. For a stable sliding condition, the optimal weight ratio of FG/T is from 0.017 to 1.

Rheometric measurements of graphenes: Graphene dispersions were deposited on a substrate and cut them to the appropriate size for rheometric tests. Graphenes were measured on a TA AR2000 rheometer in a 25-mm-diameter plate-plate geometry at 25 °C. Frequency sweeps were performed by applying a sinusoidal strain, γ = γ₀sin(ωt), with γ₀ = 0.5%, within the linear viscoelastic range.

Graphene assembly: Graphene solutions were dispersed into DMF with the aid of sonication to generate a uniform and stable graphene solution. The FG and FG-T+ solutions were
dispersed in DMF. These graphenes assembled into uniform films on the surface of water when several drops of ethyl acetate were added to the dispersion in dimethylformamide (DMF) that had been added to water (Figure S1). In this process the graphene moves to the water-air surface by Rayleigh-Bénard convection and then simultaneously assemble into an interconnected films driven by Marangoni forces. The film is then transferred onto a stainless steel (SS) substrate (Figure S1). After transfer, solvent is removed by vacuum evaporation at room temperature for 30 min. The SS plate samples were cleaned by sonication in acetone and isopropanol alcohol to remove any organic contaminants and then kept in a dry chamber. This densely covered surface was required to prevent the DLC from directly contacting the SS (grade 316) substrate.

Friction characterization: The friction measurement system, shown in Figure S4, consists of a linear flexure bearing stage driven linearly back and forth by a custom built Lorenz force motor, two load cells connected in series and perpendicularly to each other, a stepper motor driving the sensor assembly up and down, and an environmental chamber in which the measurement system is enclosed. A reconfigurable input-output (cRIO) system (cRIO-9024, National Instruments) was used to control the velocity of the bearing stage and the normal force applied on the sample. To quantify COF ($\mu = F_f/F_n$) the DLC ball slides back and forth (linear velocity: 16 to 52 mm s$^{-1}$) repeatedly along a horizontal line on a film, while simultaneously measuring fictional force ($F_f$) and normal force ($F_n$). The resolutions of each load cell are 5 mN (normal) and 0.5 mN (lateral), and forces are sampled at 1 kHz. 3 N of normal force was applied to the SS plate by moving the DLC-coated (a-C:H, Richter precision inc.) SS ball (grade 440C, diameter of 9.5 mm), attached at the tip of the sensor assembly, down to the sample on the moving bearing stage. The sample on the bearing stage slides 4 mm backward and forward against the DLC-coated SS ball and the linear velocity was maintained constant as 16 mm s$^{-1}$ during the measurement. All measurements were conducted at room temperature. The COF was calculated by dividing the measured frictional force by normal force and averaged over each backward and forward cycle to generate one data point on the COF plot. The first and the last 5% of the measured COFs in each cycle were omitted as they reflect the static friction of the materials. Friction experiments were repeated more than three times for each sample in order to ensure the reliability. After the friction test, the tribo-layer on the wear track were collected to investigate the structure of films. To validate the accuracy of our custom built friction measurement setup, the friction coefficient of stainless steel ball (grade 440C, diameter of 9.5 mm) against a Teflon sheet (thickness of 0.9 mm) is measured and compared to data presented in other literature$^{[1,2]}$ (see Figure S5). The sliding speed of the experiments is 2.5 mm s$^{-1}$ and normal force of 1.5 N, 3 N and 6 N are used to vary the normal pressure of the contact. The contact areas are approximated based on the optical and SEM images of the Teflon sheet after the experiments. As is shown, the measured COFs are ranging in between 0.018 and 0.026, comparable to the data reported by Thompson et al.$^{[1]}$ and Mokha et al.$^{[2]}$. However, in our result, the trend of lower COFs with higher normal pressures are not observed. This can be attributed to the difference in calculating the normal pressure of the contact: the contact area is assumed to be held constant with different normal forces in the literature whereas it is observed to change as the Teflon sheet at the contact deforms.

Nanoindentation: Instrumented nanoindentation experiments were performed using a Triboindenter (Hysitron 950) at room temperature, calibrated on fused quartz. The FG-T+ tribo-layer was used after sliding test. During indentation process, the displacements versus the applied loads are recorded. Those records are then used to calculate the indentation hardness of the FG-T+ tribo-layer. Each point is separated by a distance of 2 µm.
Figure S1. (a) Synthesis of Triamnio-T. (b) FG-T from Hyper-3-Stage-1 graphite intercalation compound (GIC). The Hyper-3-Stage-1 GIC represents the expanded gallery with intercalated tetrabutylammonium ion (TBA⁺) domains between the basal planes of graphenes (TBAP = tetrabutylammonium perchlorate. (c) The size distribution of FG. (d) The optical micrograph of a graphene film on water surface. The graphene assembly on water surface is driven by Rayleigh–Bénard convection and Marangoni force.
Figure S2. (a) Attenuated total reflectance Fourier transform infrared spectroscopy (ATR-FTIR) data of Triamino-T, FG, FG-T, and FG-T+. (b) Nuclear magnetic resonance spectroscopy (NMR) spectra of FG-T, and Triamino-T. FG-T (major): $^1$H NMR (400 MHz, DMF-d7): δ 6.96-6.90 (m, 3H), 6.72-6.67 (m, 3H), 6.19-6.14 (m, 3H), 4.91 (s, 1H), 4.93 (s, 1H), 4.76 (br.s, 6H).
**Figure S3.** Top: Mechanical sliding-induced reconfiguration of a scratched FG-T+ film: The reversible Meisenheimer complexes across the cracked film (intentionally scratched by a razor blade) was cured under mechanical shear force. Raman signature of FG-T+ from the recovered region. Bottom: Schematic illustration showing the formation of a tribo-layer against a DLC-coated SS ball.
Figure S4. (a) Experimental demonstration and coefficient of friction (COF) results of graphene films. An image of the linear tribometer equipped with a DLC-coated SS ball. Tests were performed at room temperature (RT) under 3 N load (contact pressure: 0.3 ~ 0.4 GPa). Each sample was tested more than three times in order to ensure the reliability. The tapping-mode atomic force microscopy (AFM) surface scans and the Raman spectra of (b) a SS substrate, and (c) a DLC-coated SS ball.
**Figure S5.** Comparison of the COF between stainless steel balls and Teflon sheets measured by the custom-built linear macro tribometer and the data from literature.\textsuperscript{[1,2]} The ball diameter of 9.5 mm is used to slide at the linear speed of 2.5 mm s\textsuperscript{-1} on the Teflon sheet of 0.9 mm thickness. The contact pressures for the normal force of 1.5 N, 3 N and 6 N are estimated to be 12 N mm\textsuperscript{-2}, 13 N mm\textsuperscript{-2}, and 11 N mm\textsuperscript{-2}, respectively.
**Figure S6.** (a) Scanning electron microscope (SEM) images with the corresponding energy dispersive X-ray spectroscopy (EDS) carbon mapping of UEG, (b) FG, and (c) FG-T+ after sliding.
Figure S7. (a) A SEM image of the surface of diamond-like-carbon (DLC)-coated ball. The sliding motions with the rough surface at contact edges highly cause the damages of coated materials. (b) The COF for DLC-coated ball sliding against UEG. As a result of the unstable tribo-layer formation on the SS substrate, the COF increases at the beginning of sliding and many large COF spikes appear. Similarly, graphene films on SiO$_2$ surfaces display a substantial amount of variance in the COF in sliding experiments against a DLC ball.$^{[3]}$ This erratic nature of COF of pristine graphene coatings at macroscale levels will limit the operational lifespan of a mechanical components and/or require frequent reapplication of the lubricant.
Figure S8. The COF of UEG-nanodiamond in our system. The average of COF is 0.011.
Figure S9. Optical images and high resolution X-ray photoelectron spectroscopy (XPS) C 1s spectra of (a) UEG (b) FG (c) FG-T, and (d) FG-T+ after sliding cycles. (e) (Top) The optical profile of the tribo-layer of FG-T+ on a SS substrate after sliding cycles. After sliding, the tribo-layer of FG-T+ was formed. (Bottom) Schematic illustration of reconfigurable networks of a FG-T+ matrix during sliding against a DLC-coated SS ball. The bottom of a FG-T+ matrix adheres to a SS substrate and the rest of the matrix provides reversible slip by Meisenheimer complexes.
Figure S10. (a) The optical profile of a DLC-coated SS ball before a friction test. The optical profile and the COF of an intact DLC-coated SS ball against (b) a FG-T+-coated SS substrate and (c) an uncoated SS substrate in N₂ environment. (b) Over 20000 cycles, the wear rate of DLC-coated SS ball against FG-T+ on a SS substrate is $0.94 \times 10^{-9}$ mm³ N⁻¹ m⁻¹. (c) The wear rate of DLC-coated SS ball against an uncoated SS substrate is $9.94 \times 10^{-9}$ mm³ N⁻¹ m⁻¹. (d) An AFM surface scan of the uncoated SS substrate after sliding against a DLC-coated SS ball. We observed that the tribo-chemically generated particles due to the wear of a DLC/SS pair. The morphology is different from the FG-T+ in Figure 4d in main text. (e) Raman spectra of a SS substrate before and after sliding against DLC.
Figure S11. The COF of a DLC-coated ball against (a) a FG-T+coated SS substrate, and (b) a SS substrate in ambient environment. The COF of an uncoated SS ball against (c) a FG-T+coated SS substrate and (f) a SS substrate in N₂ environment. The COF of an uncoated SS ball against (e) a FG-T+coated SS substrate, and (f) a SS substrate in ambient environment.
Figure S12. XPS spectra of (a) uncoated SS, (b) Triamino-T, and (c) UEG-T before/after sliding against DLC. During sliding, the Triamino-T and UEG-T also undergo tribo-chemically damage that causes unstable frictional sliding. (d) Raman spectra of the tribo-layer of uncoated SS, UEG, FG, UEG-T, UEG-T, FG-T, and FG-T+ after sliding against DLC. The tribo-layers of graphenes clearly show different chemical signatures after sliding against DLC, compared with the tribo-layer of uncoated SS. For example, the G_{FG-T+} band (1606 cm$^{-1}$) of the tribo-layer of FG-T+ is distinguishable from the G_{DLC}-band (1578 cm$^{-1}$) of DLC debris.
Figure S13. AFM surface scans and TEM data of (a) UEG, and (b) FG tribo-layers.
**Figure S14.** AFM surface scan of the initial FG-T+ coating on a SS substrate before a friction test.

Reference

