Graphene overcoats for ultra-high storage density magnetic media

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Hard disk drives (HDDs) are used as secondary storage in digital electronic devices owing to low cost and large data storage capacity. Due to the exponentially increasing amount of data, there is a need to increase areal storage densities beyond ~1 Tb/in². This requires the thickness of carbon overcoats (COCs) to be <2 nm. However, friction, wear, corrosion, and thermal stability are critical concerns below 2 nm, limiting current technology, and restricting COC integration with heat assisted magnetic recording technology (HAMR). Here we show that graphene-based overcoats can overcome all these limitations, and achieve two-fold reduction in friction and provide better corrosion and wear resistance than state-of-the-art COCs, while withstanding HAMR conditions. Thus, we expect that graphene overcoats may enable the development of 4–10 Tb/in² areal density HDDs when employing suitable recording technologies, such as HAMR and HAMR+bit patterned media.
there has been an incessant increase in data generation over the past few decades. The annual data creation rate was ~16.1 zettabytes/year (ZB, 1ZB = trillion gigabytes (GB)) in 2016, and is expected to increase to ~163 ZB/year by 2025. While various devices are used to store digital information, e.g., tape and flash drives, hard disk drives (HDDs) remain the primary choice as a secondary storage device, due to their low cost <0.1$/GB at 2016 prices and large storage capacity >10 TB with 3.5 inch HDDs. HDDs will rule storage technologies at least for the next 5–10 years in terms of capacity, price, production, and shipment. Supplementary Note 1 provides the key background concepts on HDD technology and the main technological challenges.

Solid state drives (SSDs) are the main competing technology, posing a threat to future HDD viability. However, SSDs are not yet used as HDD overcoats due to the presence of macro-particles.

Cost, oxidation, and corrosion protection are seen as anti-friction, wear resistance, Young’s Modulus, surface smoothness, and corrosion protection, with corrosion current densities < 5 nA/cm². Reduced graphene oxide was suggested to reduce the steel COF ~ 0.15 for 1LG, indicating high wear resistance and corrosion protection, with corrosion current densities < 5 nA/cm². Graphene is an emerging material for lubrication as well as oxidation and corrosion protection. Reference 37 reported that single-layer graphene (1LG) reduced the steel COF from 0.9 to 0.3 with a coating lifetime up to 6500 cycles, and decreased the wear rate by two orders of magnitude. Multilayer graphene (MLG) (3–4 layers) showed excellent tribological performance with a COF < 0.2, a decrease of wear rate by three orders of magnitude, and sliding lifetime up to 47000 cycles on steel. Reference 35 reported that 1LG exhibits superlubricity and reduced the COF by 2–3 times and wear rate by two orders of magnitude for Au-based electrical contacts. 1LG decreases the oxidation and corrosion of various metals, such as Ni, Co, Fe, Pt, Cu, Ag. Reduced graphene oxide was also used as a barrier coating. Reference 43 reported that suspended 1LG has good thermal stability up to 2600 K, with a thermal conductivity up to 2000 W/mK. In ref. 47, we showed that it is possible to achieve high mobilities ~30,000 cm²/Vs at room temperature in wet transferred, polycrystalline 1LG. Thus, scalable processes, such as wet transfer, can be used for integration and packaging. All these characteristics make 1LG promising as a protective overcoat for both existing and HAMR-based technologies.

Here we use 1–4 layers of chemical vapor deposition (CVD) grown graphene (1–4LG) transferred on Co-alloy (current technology) and FePt-based (HAMR technology) HDDs, and test friction, wear, corrosion, thermal stability, and lube compatibility. We demonstrate that 1LG shows better performances than current 2.7 nm COCs on Co-alloy-based HDD, and good tribological properties with a COF < 0.2 for >10,000 cycles. We achieve very low COF ~ 0.15 for 1LG, indicating high wear resistance and corrosion protection, with corrosion current densities < 5 nA/cm² for 2–3LG. Thermal stability tests confirm that 1LG on FePt can withstand HAMR-like conditions, without degradation. Graphene’s superior performance and its thinness can enable the development of ultra-high-density magnetic data storage technologies, based either on current technology, as well as on HAMR, or HAMR combined with bit patterned media (BPM), where the magnetic storage layer is patterned into an array of pillars, each representing a single bit. The combination of 1LG + HAMR + BPM may increase AD > 10 Tb/in².

**Results**

**Magnetic media substrates.** We use CoCrPtOxide-based bare HDM (BM) from Hitachi Global Storage Technologies (now Western Digital). These comprise multiple layers: seed bottom layer, soft-magnetic under layers (SUL), antiferromagnetic layer (A-FM), intermediate layers, and CoCrPtOxide storage layer (Fig. 1a, b). We also use other HDD from Western Digital with ~2.7 nm commercial COC (CMC, Fig. 1c), and ~2.7 nm commercial COC + ~1 nm commercial PFPE lube (CMCL, Fig. 1d) as reference.

The details and nomenclature of the COCs are in Table 1. The structural properties of the FCVA and sputter-deposited COCs are compared in Table 2.
can be found in refs. 50,51. HAMR-compatible FePt-based HDMs are shown in Fig. 1f. See Supplementary Notes 2 and 3 for further details.

Graphene growth and transfer. Graphene is grown by chemical vapor deposition (CVD) and placed on the HMD by wet transfer52,53, as described in the "Methods" section. The quality and uniformity of the samples are assessed by Raman spectroscopy54. Unpolarized Raman spectra are recorded at 514.5, as well as at 785 nm, close to the 850 nm used for HAMR writing tests25,55, with a Renishaw InVia spectrometer equipped with a Leica DM LM microscope and a ×100 objective with a numerical aperture 0.85. A 514.5 nm Raman spectrum of 1LG on Cu before the transfer is shown in Fig. 2a. The photoluminescence (PL) background due to the Cu foil is removed using baseline subtraction56. The D to G intensity ratio $I(D)/I(G)$ is <0.1, indicating a defect concentration $n_d < 2.4 \times 10^{10}$ cm$^{-2}$ 54,57–59 (Fig. 2b). The 2D peak position can be fitted with a single Lorentzian with Pos(2D) ~ 2705 cm$^{-1}$ and full-width-half-maximum,
Table 1 List of samples.

| Sample Code | Description                                                                 |
|-------------|-----------------------------------------------------------------------------|
| BM          | CoCrPt-oxide-based BM without COC and lube                                  |
| BML         | CoCrPt-oxide-based BM without COC, with PFPE lube (1.2 ± 0.2 nm)             |
| CMC         | CoCrPt-oxide-based commercial HDM+CO (2.7 nm) without lube                   |
| CMCL        | CoCrPt-oxide-based commercial HDM+CO (2.7 nm)+PFPE lube (~1 nm)              |
| 1LG         | 1LG on CoCrPt-oxide-based commercial HDM                                   |
| 2LG         | 2LG on CoCrPt-oxide-based commercial HDM                                   |
| 3LG         | 3LG on CoCrPt-oxide-based commercial HDM                                   |
| 4LG         | 4LG on CoCrPt-oxide-based commercial HDM                                   |
| 1LGL        | PFPE lube (1.2 ± 0.2) nm-coated 1LG on CoCrPt-oxide-based commercial HDM    |
| 2LGL        | PFPE lube (1.2 ± 0.2) nm-coated 2LG on CoCrPt-oxide-based commercial HDM    |
| 3LGL        | PFPE lube (1.2 ± 0.2) nm-coated 3LG on CoCrPt-oxide-based commercial HDM    |
| 4LGL        | PFPE lube (1.2 ± 0.2) nm-coated 4LG on CoCrPt-oxide-based commercial HDM    |
| 3CF         | FCVA-deposited ta-C (~0.3 nm) on CoCrPt-oxide-based commercial HDM           |
| 6CF         | FCVA-deposited ta-C (~0.6 nm) on CoCrPt-oxide-based commercial HDM           |
| 12CF        | FCVA-deposited ta-C (~1.2 nm) on CoCrPt-oxide-based commercial HDM           |
| 18CF        | FCVA-deposited ta-C (~1.8 nm) on CoCrPt-oxide-based commercial HDM           |
| 8CS         | Pulsed DC sputtered sp² rich carbon (~0.8 nm) on CoCrPt-oxide commercial HDM|
| 12CS        | Pulsed DC sputtered sp² rich carbon (~1.2 nm) on CoCrPt-oxide commercial HDM|
| FePt         | FePt/MgO/CrRu/glass samples grown by magnetron sputtering. CrRu and MgO seed layers are grown at 400 °C and 1.5 mTorr; FePt is grown at 600 °C and 3.5 mTorr as discussed in ref. 22. |

Bare media, commercial media with an overcoat, graphene-coated media, FCVA, and sputtered carbon overcoats with and without lubricant (indicated by L), with details on sample structure and corresponding acronyms.

Fig. 2 Raman and irradiation measurements. a) 514.5 nm Raman spectra of as-grown 1LG on Cu (before transfer) and 1-4LG transferred on CoCrPt:Oxide after CoCrPt:Oxide background subtraction. b) 514.5 nm Raman spectra of CoCrPt:Oxide substrates and 1-4LG transferred onto CoCrPt:Oxide. c, d) 785 nm Raman spectra of 1LG on (c) Si/SiO₂ and (d) FePt-based HAMR-HDM, for different laser power densities.
deposited by plasma-based processes. The thickness and high diation stability. To test whether 1LG can withstand HAMR and on Si/SiO₂. Reference achieved a specific value. The thickness of 4LG on HDM is ~2.1 nm (Supplementary Fig. 3), FWHM(2D) ~ 33 cm⁻¹, a signature of 1LG. We first consider Si/SiO₂/1LG. We report spectra at different power densities for ~4 min to achieve a good signal-to-noise ratio. Figure 2c shows that 1LG on Si/SiO₂ has no D peak even for the highest power density. Pos(G) downshifts from ~1588 to 1583 cm⁻¹, indicating an increase in T to ~312.5 K with respect to room temperature, by taking ~0.016 cm⁻¹/K as a shift of Pos(G) with T. Figure 2d plots the data for 1LG on FePt HDD, with I(G) normalized to that at 1.3 mW/μm². In this case, the FePt substrate is rotated at 4100 rpm on a circular track with a diameter ~4 mm to simulate rotating HDD conditions, with a 20 min acquisition time, much larger than the total laser irradiation time expected for a 5-years life of HAMR-based HHDs. No D peak is seen for all power densities, thus confirming the stability of 1LG. The Raman data show that no visible changes occur during the irradiation test measurements.

FWMH(2D) ~ 33 cm⁻¹, a signature of 1LG. The G peak at 1583 cm⁻¹ and 16 cm⁻¹, J(2D)/I(G) and A(2D)/I(G) are ~2.3 and 4.9. Representative Raman spectra of 1–4LG transferred on HDM are in Fig. 2a, with the HDM spectrum subtracted. The HDM spectrum and the spectra of 1L–4LG on HDM before background subtraction are in Fig. 2b. By subtracting the reference spectrum, using the N₂ Raman peak from the air for normalization, from the spectrum taken on 1LG-coated HDM, we can reveal the 1LG contribution. After transfer, all peaks are downshifted and broader, with Pos(2D)₁LG ~ 2672 cm⁻¹, Pos(2D)₂LG ~ 2681 cm⁻¹, Pos(2D)₃LG ~ 2687 cm⁻¹, Pos(2D)₄LG ~ 2682 cm⁻¹, FWHM(2D)₁LG ~ 61 cm⁻¹, FWHM(2D)₂LG ~ 57 cm⁻¹, FWHM(2D)₃LG ~ 49 cm⁻¹, FWHM(2D)₄LG ~ 53 cm⁻¹. Assuming 2686 cm⁻¹ as unstrained Pos(2D)₁LG, and a rate of change in Pos(2D) with strain δPos(2D)/ δε ~ −64 cm⁻¹/％, this would give ~0.22% uniaxial strain in 1LG, consistent with the large FWHM(2D). We assume the strain to be mostly uniaxial, because it is unlikely to have biaxial, i.e., perfectly isotropic strain, in all directions, unless this is used for a specific purpose. There is a small increase in I(D)/I(G) to ~0.2 for 2–4LG on HDM.

Current commercial HDD technology uses a ~2.5–3 nm COC deposited by plasma-based processes. The thickness and roughness of 1–4LG are measured by Atomic Force Microscopy (AFM), as discussed in Supplementary Note 5. This shows that the thickness of 4LG on HDM is ~2.1 nm (Supplementary Fig. 3), well below that of the current commercial COC. Graphene is transferred without such an energetic process as that used for commercial COC deposition. Thus, it does not adversely affect the magnetic performance. Even considering that COCs slightly affect the HDM magnetic properties, the influence of graphene will be much less than commercial COCs deposited by energetic processes, where energetic atoms/ions (neutrals also) hit and participate in deposition onto the HDM. Reference 64 reported that 1LG grown by CVD on Co (and Ni) does not affect the hysteresis loop. The coercivity changes from 510 mT (BM) to ~97 mT (~4LG-coated BM), which is unchanged at ~97% of the saturation magnetization when compared to BM. The coercivity and the remanent magnetization of 1–4LG-coated BM are similar to BM+COC, indicating that graphene coating does not have any detrimental effect on the media, see Supplementary Note 6 for details.

Laser irradiation stability. HAMR is not possible without laser heating. References 22,25,65 reported that FePt does not degrade upon laser heating, with lubricant and COC being the main concern when it comes to laser irradiation. Consistent with refs. 25,55, here we use an IR laser to examine the 1–4LG irradiation stability. To test whether 1LG can withstand HAMR conditions we consider 1LG transferred on L₁₄Fe₇Pt-based HDM and on Si/SiO₂. Reference 55 achieved Tₘ ~ 1.3 mW/μm² by optimizing aperture optics. Reference 55 suggested using a laser power density ~10⁷ W/cm² and ~0.35 × 10⁷ W/cm² for FePt-based HDD. We perform Raman measurements at 785 nm, the available Raman wavelength closest to that used in HAMR (~830 nm), with a spot size ~1.24 μm, as determined by the razor blade technique, see Supplementary Note 7 for details. We vary the power density from ~1.3 mW/μm² (0.013 × 10⁷ W/cm²) to ~31.5 mW/μm² (0.315 × 10⁷ W/cm²) in order to examine the laser irradiation-driven evolution of the SLG Raman spectrum under HAMR power densities.

We first consider Si/SiO₂/1LG. We report spectra at different power densities for ~4 min to achieve a good signal-to-noise ratio. Figure 2c shows that 1LG on Si/SiO₂ has no D peak even for the highest power density. Pos(G) downshifts from ~1588 to 1583 cm⁻¹, indicating an increase in T to ~312.5 K with respect to room temperature, by taking ~0.016 cm⁻¹/K as a shift of Pos(G) with T. Figure 2d plots the data for 1LG on FePt HDD, with I(G) normalized to that at 1.3 mW/μm². In this case, the FePt substrate is rotated at 4100 rpm on a circular track with a diameter ~4 mm to simulate rotating HDD conditions, with a 20 min acquisition time, much larger than the total laser irradiation time expected for a 5-years life of HAMR-based HHDs. No D peak is seen for all power densities, thus confirming the stability of 1LG. The Raman data show that no visible changes occur during the irradiation test measurements.

Friction and wear. The HDD comprises magnetic/metallic layers, including Co-alloys-based magnetic storage layers, with a high COF ~0.6–0.811,15 and wear11,15. Therefore, they can experience mechanical damage whenever intermittent contact occurs with the head11,15, and are susceptible to corrosion11,14 and oxidation11,15, leading to HDD degradation or damage. The current ~2.7 nm commercial COCs have a high COF ~0.3–0.511,15 and wear in a ball-on-disc tribological environment11,15, which can result in damage, hence durability concerns, see Supplementary Note 1 for more details.

Reference 11 used FCVA to deposit ~1.5–2 nm carbon films for protection of Co-alloy-based HDD and reported low COF ~0.25, wear, and corrosion. Here, we use 1–4LG grown by CVD for Co-alloy-based and HAMR HDDs.

Since in HDDs the HDD spins during operation, COF and wear need to be examined in a setup mimicking HDD operation. AFM and other tip-based tools are used to measure COF and wear39. In contrast to setups with rotating geometry, tip-based tools measure the COF based on the movement of the tip in the lateral direction. Ball-on-disc measurements (rotation-based geometry) have a similar assembly as HDDs, with samples rotating while the counterface is in contact with the surface, thus measuring COF. We perform ball-on-disc tests using a nano-tribometer (CSM Instruments) in a cleanroom, to have a controlled environment with T = 23 ± 1°C and a relative humidity ~55 ± 5%. A sapphire ball (Al₂O₃) of diameter ~2.0 ± 0.1 mm and surface roughness ~5.0 ± 0.1 nm is used as the counterface because the hard disk head is made of an Al₂O₃-based composite. During the test, a normal load ~20 mN and a rotational speed ~100 rpm are used, corresponding to a linear speed ~1.05 cm/s for 10,000 cycles.

Since in HDDs the contact occurs occasionally, the 10,000 cycles in our setup are much higher than the HDD operational lifetime. After each test, the wear track and ball images are captured using an optical microscope. To check repeatability, tests are performed 2–7 times. When two surfaces are in contact, and at least one of the surfaces starts to slide with respect to the other, friction and wear occur. As a result, a wear track is formed.

Figure 3 plots representative friction curves for BM, CMC, CMCL, and BML (Fig. 3a) coated with 1–4LG (Fig. 3b), and BM coated by FCVA (Fig. 3c). The average COFs are in Fig. 4, including HDM coated by DC-sputtering. BM has the highest COF ~0.511,15, 0.311,15 and wear11,15. Therefore, they can experience mechanical damage whenever intermittent contact occurs with the head11,15, and are susceptible to corrosion11,14 and oxidation11,15, leading to HDD degradation or damage. The current ~2.7 nm commercial COCs have a high COF ~0.3–0.511,15 and wear in a ball-on-disc tribological environment11,15, which can result in damage, hence durability concerns, see Supplementary Note 1 for more details.

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(without lube) is ~4 times lower than BM and ~2 times lower than CMC, despite a reduction of thicknesses ~7 times (for 1LG) to 2 times (for 4LG), using the theoretical 1LG thickness, with respect to CMC. Figures 5b, d and 6b, d reveal that the wear track width of 1–4LG-coated samples is ~2–4 times lower and debris transferred to the ball are smaller than BM and CMC, indicating higher wear resistance. All FCVA COCs with thicknesses from ~0.3 to ~1.8 nm show ~2–5 times higher COF than 1–4LG-coated samples, with and without lube, apart from 1LG without lube. Pulsed DC sputtered COCs have ~2–3 times higher COF than 1–4LG-coated samples.

To further analyze the wear tracks, Auger Electron Spectroscopy (AES) imaging is performed using a JEOL JAMP Auger Microprobe, see Supplementary Note 4 and Supplementary Fig. 1 for details. Before AES, scanning electron microscope images are taken to select the AES locations. The AES images inside and outside the wear tracks show the carbon-containing sites, and indicate that the amount of carbon on the wear track increases with increasing number of graphene layers, N. The Co and Cr intensities inside the wear tracks are higher for 1LG, and decrease with increasing N, due to the increase in C and the <1–3 nm sampling depth of AES68. The O signal in the wear track appears due to ambient oxygen as the samples are exposed to air before AES, with some contribution from the HDM oxide.

After ball-on-disk tests, the wear tracks are analyzed by recording Raman spectra across the wear track at different positions. Reference 58 introduced a three-stage model of amorphization. Stage 1: graphene to nanocrystalline graphene. Stage 2: nanocrystalline graphene to low $sp^3$ amorphous carbon. Stage 3: low $sp^3$ to high $sp^3$ amorphous carbon. For all spectra in Figs. 5 and 6, as the D peak increases, $D'$ and $D+D'$ appear, whereas the $I(2D)$ weakens when approaching the wear track, indicating an increase in disorder according to stage 1. The broad peak between 500 and 1000 cm$^{-1}$ is due to the glass substrate69. At the center of the wear track, all second-order Raman features (i.e., 2D, $2D'$, and $D+D'$) merge, while $I(D)/I(G)$ in Fig. 7a, from at least 5 positions as a function of COF, without and with lubricant (L). Overall, there seems to be no significant trend of $I(D)/I(G)$ with COF. However, a difference is seen without lubricant: $I(D)/I(G)$ is ~0.95 and ~1 for 3–4LG, while for 1–2LG it is ~1.33 and ~1.25. Thus, there are less defects in 3–4LG compared to 1–2LG. The peaks <1000 cm$^{-1}$ in Fig. 7b are due to the HDM: those ~470 cm$^{-1}$ ($E_g$), 513 cm$^{-1}$($F_{2g}$), 606 cm$^{-1}$($F_{2g}$), 673 cm$^{-1}$
(A1g) are from Co3O4, while the A2g mode ~550 cm$^{-1}$ is from Cr2O3.

**Effect of lubricant.** Current commercial HDDs use a layer of perfluoropolyether (PFPE) lubricant (lube) on top of COCs to further reduce friction and wear, and minimize surface energy. Given the lubricating and corrosion protection properties of 1LG itself, lube may not be needed for 1LG. However, exploring the compatibility of 1LG with lube is useful from the fundamental viewpoint.

Friction measurements on lube-coated samples (Fig. 3) show that in certain cases the COF is higher with L, than without. Taking the error bars into account, a significant difference in COF is only observed for 1LG and 1LGL. 1LG has wetting transparency, i.e., it is thin enough that its introduction does not affect the wettability of an underlying substrate. Without overcoat, the L containing HDM, as in BML, shows very high COF ~ 0.55 and wear, due to the metal (medium) L-induced catalysis. 1LG is not enough to avoid the interaction of L with medium, hence higher COF ~ 0.45 and wear is observed in 1LGL. Without L, 1LG shows very low COF ~ 0.15 and wear. A minor difference in case of 2LG versus 2LGL can be seen. However, for 3LG versus 3LGL and 4LG versus 4LGL the difference in average COF is marginal. L on 1LG results in higher and inconsistent COF and wear, as some measurements show lower COF ~ 0.15, some higher ~ 0.78, with COF increasing after few thousands cycles. The COF of 1LG, on an average, is ~0.45, i.e., ~3 times its non-lubricated counterpart. Thus, COFs of 2–4LG are more or less similar to those of the non-lubricated counterparts, suggesting that 2–4LG without L are lubricious enough, and that L does not improve lubricity. The D and G peaks in the Raman measurements in Figs. 5 and 6 on 1–4LG confirm the presence of carbon on the wear tracks, and transfer of debris to the balls, similar to the non-lubricated counterparts.

**Corrosion.** Co-alloys have a great propensity to corrode, mainly due to Co oxidation. This results in the loss of magnetic properties, hence this is one of the major concerns for the long-term functionality and durability of HDDs. To examine the corrosion protection efficiency of 1–4LG and compare their performance with state-of-the-art COCs, the corrosion of different uncoated- and coated-HDM, exposed to an electrolyte solution ~0.1 M NaCl similar to that used in refs. on a ~0.24 cm$^2$ area, is investigated using an electrochemical corrosion method. The measurements are performed with a 3-electrode setup with a Pt wire as counter electrode, Ag/AgCl as reference electrode and HDM as working electrode, to which the potential is applied. Each test consists of anodic and cathodic sweeps, where the potential is varied and the corresponding current measured. The so-called Tafel’s analysis is done by plotting anodic and cathodic curves on a semi-logarithmic scale of potential versus log current. The linear part of the logarithmic anodic and cathodic currents are...
extrapolated, and the intercept of these lines gives the corrosion current, and the corrosion current density \( J_{\text{corr}} \) when divided by the contact area.

The corrosion protection efficiency (CPE) defines how efficient the COC is in protecting against HDM corrosion. This is defined as:

\[
\text{CPE (\%)} = \frac{J_{\text{corr}}^0 - J_{\text{corr}}}{J_{\text{corr}}^0} \times 100,
\]

where \( J_{\text{corr}}^0 \) is the BM corrosion current density and \( J_{\text{corr}} \) that of coated HDM.

When a metal is exposed to a corrosive solution, it releases ions that leave behind electrons, which can be observed in an anodic reaction as:

\[
M \rightarrow M^{n+} + ne^-,
\]

where \( M \) represents the metal and \( n \) the number of electrons released by it. For Co-alloy-based systems, the metal dissolution, which decreases the anode conductivity, can be written as:

\[
\text{Co} \rightarrow \text{Co}^{2+} + 2e^-,
\]

As the corrosion reaction involves the transfer of electrons and ions between metal and solution, the corrosion rate varies with corrosion current, hence \( J_{\text{corr}} \) varies inversely with corrosion resistance, i.e., the COC ability to reduce the HDM corrosion, which could further add galvanic corrosion.

The BM shows the highest \( J_{\text{corr}} \) indicating the greatest propensity to corrode (Fig. 8). Introduction of 1–4LG reduces HDM corrosion. \( J_{\text{corr}} \) decreases with \( N \), from ~5.3 nA/cm² for 2LG to ~4.4 nA/cm² for 4LG, Fig. 8, and CPE increases. The COC defects and pinholes are the active corrosion sites, and their electrical conductivity could further add galvanic corrosion. The COC should be defect- and pinhole-free, smooth, and with excellent barrier properties to minimize HDM oxidation and corrosion. Reference 43 showed that 1LG can be used as a corrosion barrier for metallic surfaces, and suggested that this should be uniform and defect-free to achieve corrosion protection. In our case, 1LG reduces \( J_{\text{corr}} \) by ~2.5 times with respect to BM, but it is twice that of BM with ~2.7 nm COC. For 2LG, \( J_{\text{corr}} \) is ~3.8 times smaller than BM, and marginally higher than COC. This is remarkable as the 2LG thickness ~0.7 nm, is ~4 times lower than ~2.7-nm-thick COC. Beyond 2LG, \( J_{\text{corr}} \) and CPE remain similar, adding marginal anti-corrosion improvement (Fig. 8). The higher corrosion protection in 2–4LG with respect to 1LG is mainly due to the fact that the increase in \( N \) improves the HDM coverage, leading to the reduction of active corrosion sites. 2–4LG display corrosion protection similar to COC, attributed to better barrier properties. Figure 8 indicates that, for a similar thickness, graphene-based overcoats have lower \( J_{\text{corr}} \) than amorphous COCs, indicating greater protection. Figure 8 shows that the corrosion resistance performance of 1LG is ~2.5 times better than BM, but ~2 times higher than COCs. By adding graphene layers the corrosion resistance is improved. The
The horizontal red dashed line is a reference at 12CS, the yellow region highlights media with graphene based sample in each pair is always slightly thinner than the amorphous COC. Taking into account that the theoretical thickness of 1LG is ~8–9 times lower than state of the art COCs, and that a COC having the same thickness as 1LG would provide no protection and be full of holes19, 1LG comfortably beats any COC of the same thickness (i.e., same storage density).

**Discussion**

Friction is defined as resistance to sliding36,76. At the macroscopic level, Amonton’s law76 states that the frictional force between two bodies varies proportionally to the normal force. Hence:

$$\text{COF} = \frac{F}{W},$$  \hspace{1cm} (4)

where $F$ [mN] is the frictional force and $W$ [mN] is the normal force. This does not take into account the area of contact at the microscopic level76. Reference 76 suggested that the contact between two bodies contains several smaller contacts, called asperities, with the sum of the areas of these asperities being lower than the apparent macroscopic area76. Thus, from ref. 76, $F$ in micro-tribology can be expressed as:

$$F = \tau \sum A_{\text{asp}},$$  \hspace{1cm} (5)

where $\tau$ is the shear strength, i.e., the stress required to shear the contacting interfaces and enable sliding, expressed as shear force/area and $\sum A_{\text{asp}}$ is the sum of the areas of the asperities, also called real contact area76. Hence, friction also depends on $\sum A_{\text{asp}}$. Reference 76 showed that the stresses over regions of contact reach the elastic limit of the material and cause plastic deformation. Thus, the average pressure ($p$) in a contact region is governed by the normal force, and is given by $p = W/A$77. The zone of deformation increases with increase in load, and $p$ on the region of contact tends to a stable value, which eventually causes the deformation to become entirely plastic. The process continues until the area of contact becomes sufficient to support the load $W$. Consequently, the normal force can be expressed as:

$$W = p \sum A_{\text{asp}},$$  \hspace{1cm} (6)

Thus, Eqs. (4)–(6), give:

$$\text{COF} = \frac{F}{W} = \frac{\tau}{p}.$$  \hspace{1cm} (7)

A co-alloy is mechanically softer78 than Al$_2$O$_3$79. When a metallic co-alloy surface slides against Al$_2$O$_3$, it causes the HDM surface to undergo plastic deformation78, generating wear and high COF ~0.8, Figs. 3 and 4, consistent with the friction results for metallic surfaces in ref. 76. This could be due to the formation of an adhesive contact at the HDM-ball interface, due to the high surface energy of Co-alloy-based HDM ~ 42.8 mN/m80, and the presence of contaminants at the interface that may enhance the interaction between the two bodies81. When 1LG is placed on HDM, the COF decreases ~4 times and negligibly changes with $N$.

At the nanoscale, the friction of 1LG on Si/SiO$_2$ against Si tips, measured at a low load ~1 nN and scan speed ~1–10 µm/s, was explained based on out-of-plane deformation in front of a scanning probe tip, the so-called puckering effect39,82. This enhances the contact area, hence friction. The out-of-plane deformation is suppressed with increasing $N$82, resulting in reduced COF. This is not applicable to ball-on-disk tribological tests of 1–4LG, due to the significantly larger dimension (ball radius ~ 2.0 ± 0.1 mm) of the counterface, much higher load ~20 mN and higher speed ~100 rpm or 1.05 cm/s, which do not precisely differentiate the frictional characteristics of 1–4LG. Thus, all coated media show similar friction (Fig. 3b). Since the layers are coupled by van-der-Waals forces, they shear easily, as their interfacial shear strength
is low, with a shear force per unit area \( q \sim 12.8 \cdot 10^{18} \text{ N/m}^2 \) and a resulting shear modulus \( C_{44} \sim 4.3 \text{ GPa} \). The ease of shearing at sliding contacts results in smoother frictional curves (and marginally lower COF) in 2–4LG with respect to 1LG. The improved wear resistance for 1–4LG-coated HDM is attributed to reduced friction, wear, and excellent mechanical properties of 1LG: breaking strength \( \sim 42 \text{ N/m} \), Young’s Modulus \( \sim 1 \text{ TPa} \), flexibility (1LG can be stretched up to \( \sim 20\% \) without breaking).

When compared to BM, the COF reduction of 1–4LG-coated HDM could be attributed to reduced adhesive interaction (lower surface energy)\(^8\),\(^3\)\(^8\)\(^5\), as well as incommensurability of the lattice planes sliding against each other at the tribological interface\(^3\)\(^5\), occurring when the hills of a surface with lattice spacing \( a \) do not match the valleys of another surface of lattice spacing \( b \), such that \( b/a \) is an irrational number\(^3\)\(^5\). This is consistent with what suggested in ref.\(^3\)\(^5\) for 3–4LG on Si/SiO\(_2\) sliding against an a-C:H-coated steel ball, and for 1–3LG on a steel sliding against a steel ball\(^3\)\(^5\),\(^3\)\(^7\) in macroscale ball-on-disk conditions, e.g., with a normal force in the \( z \) direction \( N_z \sim 0.5–3 \text{ N} \), and speeds \( \sim 0.6–25 \text{ cm/s} \).

The AES on the wear tracks and Raman measurements across the wear tracks, after ball-on-disk tribological tests, in Figs. 5 and 6 show that a C signal is still present in all 1–4LG samples, even though disorder-induced peaks appear. Reference\(^8\) also found C on wear tracks after microscale friction and wear tests of graphene on SiC. The counterface also reveals transferred debris, consisting of disordered carbon and underlying substrate atoms. This implies that, when a 20 mN load is applied and the sample starts to rotate at \( \sim 2.1 \text{ cm/s} \), 1LG could be turned into patches, due to the large contact load. Reference\(^8\) reported transformation of continuous layers of 1–2LG into 1–2LG patches during microscale tribology, despite using lower load \( \sim 0.1–1 \text{ mN} \) and speeds \( \sim 30–50 \text{ \mu m/s} \) compared to us. This led to a distribution of 1LG patches along the wear track, and transfer of 1LG-containing debris on the counterface. This also happens for 2–4LG-coated HDM. Raman measurements in Figs. 5 and 6 show disordered carbon on the wear tracks, with disorder lower than for 1LG, as revealed by \( I(D)/I(G) \) (Fig. 7). This implies that 1LG acts as lubricant when in contact with another 1LG, and is not completely removed during tribological tests. AES also reveals a progressive increase in C and C-containing sites on the wear tracks with increasing number of layers. The debris transferred to the ball also contain C. Therefore, the formation of disordered carbon debris on both surfaces facilitates smooth sliding, and contributes to maintaining the low COF. The marginally lower friction in 2–4LG as compared to 1LG can be linked to \( I(D)/I(G) \) (Fig. 7a), with lower disorder corresponding to lower friction, although this does not apply for 1–4LG.

Figure 9 is the proposed mechanism for friction reduction for 1LG and 3LG-coated HDM, where C debris are on both ball and wear track. To validate these assumptions, we perform molecular dynamics (MD) simulations of a Co/Pt–sapphire system with and without 1 and 4LG, as discussed in Supplementary Note 8.

The COF is estimated as the ratio of sliding and normal force, block averaged every 25 ps. Figure 10a–c compares the computed COF and surface disorder of Co/Pt, 1LG, and 4LG. The surface disorder represents the friction-induced damage/wear of the HDM, and its value is derived considering 0% surface disorder before friction measurements. The amount of disorder in HDM is quantified using the centrosymmetry parameter, a measure of the local lattice disorder around an atom. This is 0 for a perfect lattice, whereas when a point defect exists, i.e., when the symmetry is broken, it assumes a larger positive value, see Supplementary Note 8 for details. The simulations indicate that BM develops substrate disorder up to \( \sim 14.5\% \) within 350 ps (Fig. 10b, c).

High wear is also observed for BM in the experiments in Fig. 3a. An additional 250 ps where the forces on atoms are also collected are used to obtain the COF (Supplementary Fig. 5). The COF averaged over these 250 ps is \( -0.82 \pm 0.1 \). 1LG reduces the average simulated COF to \( -0.18 \pm 0.02 \). Figure 10c shows that 1LG maintains its structural integrity with fraction of disorder \( \sim 1\% \) on

![Fig. 9 Friction model](image_url)

Interaction of counterface ball (sapphire) with (a) 1LG and (b) 3LG. The zoomed-in images represent 1LG and 3LG, and underlying substrate atoms (Co, Pt, Cr, etc.) on counterface ball and wear track, where the amount of C debris increases with \( N \). The debris generated on both interacting bodies maintain the lower COF of 1–4LG with respect to BM.
average, as a consequence of the low COF between the blocks, indicating higher resistance to wear as compared to BM. 4LG further improves the tribological behavior. The COF drops slightly to \(-0.14 \pm 0.02\) and substrate disorder is reduced to \(-0.1\%\), much lower than in BM, suggesting that the BM surface remains mostly unaffected. Simulations suggest partial transfer of graphene patches to the sapphire ball. The Raman analysis of wear tracks in Figs. 5 and 6 shows tribo-induced disorder in 1–4LG transfer of C to the ball. Thus, the patches containing debris on both surfaces are responsible for the lower COF in 1–4LG-coated HDM, maintaining higher wear resistance than BM.

Reference 47 showed that potential corrugations on sliding surfaces can impact friction force and anisotropy of MLG. It reported that higher friction forces result from larger corrugations of the potential energy and depend on \(N\). This is consistent with our experiments in Figs. 3 and 4, and simulations in Fig. 10.

In summary, we analyzed 1–4LG-coated media and found that they can overcome the tribological and corrosion issues of current Co-alloy-based HDM, with laser irradiation stability on FePt-based HDM for HAMR. The overall performance of graphene-coated media exceeds that of thicker commercial COC, as well as other amorphous carbons of comparable/higher thicknesses prepared by FCVA and sputtering. Given the tribological, corrosion, and thermal stability characteristics coupled with an AFM thickness \(~4\) times (for 1LG) to \(~1.3\) times (for 4LG) lower than state-of-the-art 2.7-nm COCs, we expect 1–4LG-based overcoats to meet the requirements for 4–10 Tb/in\(^2\) areal density HDDs, when employing suitable recording technologies, such as HAMR and HAMR+BPM. Graphene-coated media have better corrosion and wear resistance with a COF < 0.2. A single layer of graphene is enough to reduce corrosion \(~2.5\) times and can withstand HAMR conditions. Our results imply that >2LG-based coatings could be

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**Fig. 10 Simulations of friction process.** Light green spheres indicate Co/Pt, red and blue spheres indicate Al\(_2\)O\(_3\). a Simulation setup for evaluating wear between Al\(_2\)O\(_3\) and Co/Pt. Simulations are performed for BM, 1, 4LG at 300 K, with materials sliding against each other at 1 m/s. b Interfacial surface atoms of bare Co/Pt showing disorder. Two representative disorder regions are indicated by circles. c Structure of defected 1LG. d Structure of 4LG. The top two layers are attracted towards the Al\(_2\)O\(_3\) block with the formation of sp\(^3\) cross-links. e Disorder of interfacial Co/Pt surface using the centrosymmetry parameter.
used as a tribological interface for various other materials/devices, such as micro- and nano-electromechanical systems.

Methods

Graphene growth and transfer: 1LG is grown by CVD on a 35 μm Cu foil. The substrate is loaded into a hot wall tube furnace evacuated to ~1 mTorr. The Cu foil is annealed in hydrogen at 1000 °C for 30 min. This reduces the copper oxide surface and increases the Cu grain size. The growth process starts when 5 sccm CH₂ is added to H₂. After 30 min the substrate is cooled down for several hours in vacuum (~1 mTorr) to room temperature and unloaded.

We use a polymethyl methacrylate (PMMA)-based wet transfer. First, ~500 nm PMMA is spin-coated on the sample. The PMMA/1LG/Cu stack is then placed in an aqueous solution of ammonium persulfate to etch Cu. When Cu is fully etched, the graphene/PMMA-stack is placed into a de-ionized (DI) water bath to rinse any acid residuals and subsequently fished out using the HDM substrate. After drying for one day at room temperature, the sample is placed in acetone to remove PMMA, leaving 1LG on the HDM. By repeating the steps described above, several 1LGs are transferred to create an MLG stack. The same procedure is used to place 1LG and MLG onto FePt.

Data availability

All relevant data are available from the authors upon reasonable request. Unique identifiers such as DOI and hyperlinks for any other data sets are also available from the authors upon reasonable request. Source data are provided with this paper.

Code availability

All relevant custom software is available from the authors upon reasonable request.

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Author contributions

A.C.F. and C.S.B. conceived the idea, designed the project and supervised the work. N.D. and T.D. prepared the media substrates. N.D. applied lubricant. A.K.O., C.D., U.S., A.C.F. and C.S.B. conceived the idea, designed the project and supervised the work. N.D. performed tribological measurements. A.K.O. performed Raman spectroscopy and T.D. prepared the media substrates. N.D. applied lubricant. A.K.O., C.D., U.S., A.C.F. and C.S.B. conceived the idea, designed the project and supervised the work. N.D. performed tribological measurements. A.K.O. performed Raman spectroscopy and T.D. prepared the media substrates. N.D. applied lubricant. A.K.O., C.D., U.S., A.C.F. and C.S.B. conceived the idea, designed the project and supervised the work. N.D. performed tribological measurements. A.K.O. performed Raman spectroscopy and T.D. prepared the media substrates. N.D. applied lubricant.

Competing interests

The authors declare no competing interests.

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