Covalent functionalization of graphene by azobenzene with molecular hydrogen bonds for long-term solar thermal storage

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Reduced graphene oxide-azobenzene (RGO-AZO) hybrids were prepared via covalent functionalization for long-term solar thermal storage. Thermal barrier (ΔEᵦ) of cis to tran reversion and thermal storage (ΔH) were improved by molecular hydrogen bonds (H-bonds) through ortho- or para-substitution of AZO. Intramolecular H-bonds thermally stabilized cis-ortho-AZO on RGO with a long-term half-life of 5400 h (ΔEᵦ = 1.2 eV), which was much longer than that of RGO-para-AZO (116 h). RGO-para-AZO with one intermolecular H-bond showed a high density of thermal storage up to 269.8 kJ kg⁻¹ compared with RGO-ortho-AZO (149.6 kJ kg⁻¹) with multiple intra- and intermolecular H-bonds of AZO according to relaxed stable structures. Thermal storage in experiment was the same order magnitude to theoretical data based on ΔH calculated by density functional theory and packing density. Photoactive RGO-AZO hybrid can be developed for high-performance solar thermal storage by optimizing molecular H-bonds.

Utilization of the sun as a renewable energy source based on the development of new materials and efficient devices is one of great challenges due to an increasing demand for energy impacts. Generally, an efficient and regenerative cycle is basically important for the use of solar energy including three steps – the capture of a great amount of photons, the conversion to useful form of energy, and reversible storage¹. Thus, numerous strategies to utilize sun’s energy are presented such as photovoltaic, photo-catalysis, photo-synthesis and photothermalization, but few progresses on the conversion and storage of solar thermal energy have been made so far²–⁴. Recently, photo-responsive metastable compounds are studied for close-cycle thermal storage because of their high potential capability of storage and reversible response. Such photo-responsive material stores a certain amount of solar energy in the chemical bonds through isomerization irradiated by solar light. By applying an external stimulus, the metastable moiety surmounts thermal barrier (ΔEᵦ for per molecule) and “releases” thermal energy (ΔH for per molecule). The material can be subsequently reused by the irradiation of light to induce isomerization and reversion for multiple cycles. This renewable material, which can be transportable in a liquid or powder without the emission of chemicals to environment, is an ideal candidate for solar thermal storage.

Many photo-responsive molecules capable of undergoing photoisomerization are designed and prepared⁵–⁶, but none achieves long-term and high-density storage. Azobenzene derivatives (AZO) show the great potential for solar thermal storage due to good absorption at 350–450 nm, reversible isomerization and thermal reversion controlled by functional groups and steric structures. However, the utilization of AZO is restricted by low density of thermal storage (ΔH for per molecule) during cis (Z) → trans (E) reversion, short half-lives (τ₁/₂, the time required for a quantity of Z-isomers to fall to half its value by Z → E reversion) of metastable states caused by low ΔEᵦ and low work temperatures⁷,⁸,⁹. An important strategy to overcome these problems is to tune transition bands and the formation of molecular hydrogen bonds (H-bonds) in AZO by the incorporation of electron donating or withdrawing groups in the ortho position⁷–¹⁴. Metastable Z-ortho-AZO is thermally stabilized by changing orbital energy or the formation of intramolecular H-bonds. Bléger et al. reported that Z-isomers of ortho-fluoro-AZO showed a long half-life of thermal reversion up to 714 days due to the separation of n → π* transition bands¹¹, which provided a great promise for a long-term thermal storage. Recently, a carbon-based nanostructure attached to photo-responsive AZO molecules has been established for solar thermal storage¹²,¹³. The calculation based on density functional theory (DFT) predicted that ΔEᵦ and ΔH for per AZO molecule on carbon nanotube (CNT) were controlled by molecular H-bonds. Compared with pure AZO, an increase in ΔH by more than two-times and high ΔEᵦ up to 1.25 eV (τ₁/₂ is longer than a year) were obtained for 2-2’ hydroxyl-AZO with six H-bonds (four intramolecular...
and two intermolecular H-bonds) in E-isomer and four H-bonds (two intramolecular and two intermolecular H-bonds) in Z-isomer\(^6\). On the basis of the interaction between AZO and carbon nanostructures, many studies reported on the control of isomerization and reversion of AZO bounded on graphene controlled by tuning the substitution of organic groups\(^7\). Isomerization of AZO tethered to graphene gives rise to the interaction between AZO and graphene\(^8\), which was also confirmed by the calculation based on DFT\(^9\). In spite of progresses on the calculation, the remarkable increase in both \(\Delta H\) and \(\Delta E\) for per AZO on graphene by optimizing inter and intramolecular H-bonds is still a great challenge because methods to improve the former often result in the decrease in the latter.

In this paper, two AZO molecules were covalently bound on reduced graphene oxides (RGO-AZO), and molecular H-bonds were controlled by the substitution (ortho or para) and close-packing arrays of AZO on RGO. Chemical structures, packing densities (the weight percentage of AZO for RGO-AZO) and the formation of H-bonds of AZO on RGO were studied by fourier transform infrared spectroscopy (FT-IR), proton nuclear magnetic resonance spectroscopy (NMR), X-ray photoelectron spectroscopy (XPS), and thermo gravimetric analysis (TGA). The effect of molecular H-bonds of AZO on \(\Delta H\) and \(\Delta E\) of RGO-AZO hybrids was investigated by isomerization and reversion using ultraviolet-visible (UV-vis) absorption spectra. The densities of thermal storage of RGO-AZO hybrids were experimentally measured by a microcalorimetric sensor and also calculated by DFT based on the relaxed stable structures.

**Results**

RGO was prepared by partial reduction of graphene oxide (GO) using NaBH\(_4\) controlled by reaction time (1 h) and temperature (80°C). Two AZO chromophores, 4-sulfo-4′-amino-AZO (para-AZO) and 2-sulfo-2′-hydroxy-4′-amino-AZO (ortho-AZO, Figure S1) were covalently functionalized with RGO by the diazotization using diazonium salts\(^{10,11}\). Figure 1 shows the synthesis route and chemical structures of RGO-para-AZO and RGO-ortho-AZO hybrids. Morphologies of RGO-AZO hybrids and RGO were observed by transmission electron microscopy (TEM). GO resembles crumpled silk veil with the clear boundary (Figure S2). RGO shows wrinkled nanosheets with slightly rough surfaces, and crystal lattices of RGO retains its hexagonal pattern after partial reduction according to corresponding fast fourier transforms (FTTIs, inset of Figure 2b). After the functionalization, the rough surface of dispersed nanosheets in Figure 2c and 2d appears to be covered by the addends, which form randomly distributed domains of immobilized AZO molecules\(^{12}\). This microstructure is different from that of noncovalently functionalized graphene with organic molecules\(^{13}\). The domains affect the crystal lattices of the flakes in FTTIs pattern (the inset of Figure 2d). Topological morphologies (Figure S3) are also found in RGO-AZO hybrid films. The compact structure favors the formation of molecular H-bonds among AZO molecules on the surface of nanosheets.

The covalent linkage between AZO and RGO was studied by FT-IR, NMR, and XPS spectra. As shown in Figure 3, C-N bands (brown circle) corresponding to (Ph)\(\mathrm{C-NH}_2\) at 1310 cm\(^{-1}\) of para-AZO and at 1330 cm\(^{-1}\) of ortho-AZO are disappeared in the spectra of RGO-para-AZO and RGO-ortho-AZO hybrids after the attachment. This feature is confirmed by N 1s XPS spectra (Figure S4), in which the N 1s regions of RGO-para-AZO and RGO-ortho-AZO hybrids show only one band of (Ph)\(\mathrm{N}=\mathrm{N}\) (Ph) at 400.1 eV\(^{22}\), and no peaks at 399.1 eV for (Ph)\(\mathrm{NH}_2\) are observed on the shoulder. Furthermore, the chemical shift (\(\delta\)) at 5.39 corresponding to –\(\mathrm{NH}_2\) group in ortho-AZO is also not observed in RGO-ortho-AZO hybrids (not shown). Results indicate that AZO molecules are covalently attached on RGO rather than \(\pi\)-stacking. Intermolecular H-bonds of \(=\mathrm{S-OH---OH-}\) between two adjacent AZO molecules on RGO were confirmed by red-shifted peaks of -\(\mathrm{SO}_3\)H group on RGO-ortho-AZO and RGO-para-AZO hybrid in FT-IR spectra\(^{23}\). The shifts of three characteristic peaks are clearly distinguished by dash lines in Figure 3. Red shifts by dozens of wavenumbers in sulfonic groups are observed in both RGO-para-AZO and RGO-ortho-AZO hybrids. RGO-ortho-AZO shows large red-shifts of two peaks of S-O vibration at 1180 and 1116 cm\(^{-1}\) and one band of S-phenyl vibration at 1008 cm\(^{-1}\) compared with ortho-AZO (1211, 1136 and 1018 cm\(^{-1}\)). Besides, red-shifted peaks are also observed in RGO-para-AZO (1164, 1110 and 1024 cm\(^{-1}\)) by comparison of para-AZO (1215, 1122 and 1037 cm\(^{-1}\)).

High packing density of AZO molecules on RGO is crucial for the control of closed arrays favoring the formation of intermolecular H-bonds between two adjacent molecules. The amounts of AZO functionalized on RGO were estimated by TGA (Figure 4a and 4b) and XPS spectra (Figure S5). RGO exhibits a good thermal stability and its weight loss is the elimination of remaining oxygen groups. The residual weight is 71.39%, 69.74%, and 67.63% at 700, 750, and

![Figure 1](image-url)

**Figure 1 |** The synthesis route and chemical structures of RGO-para-AZO and RGO-ortho-AZO hybrids by the diazotization.
Para-AZO and ortho-AZO molecules are thermally stable from 30 to 220°C and subsequently show a sharp weight loss between 220 and 370°C with a slow loss from 370 to 800°C. AZO molecules are almost completely combusted at 800°C. In Figure 4a and 4b, both two RGO-AZO hybrids show weight loss properties because of the elimination of oxygen groups of RGO and the decomposition of AZO molecules with two stages. Thermal decomposition temperatures of two AZO molecules on RGO are slightly increased in comparison of AZO molecules.

On the basis of the distinctive weight loss of RGO and AZO, the packing density of AZO on RGO can be determined by residual weight percentages of RGO, AZO and RGO-AZO hybrids at three temperatures. As shown in Table S1, the weight percentages of AZO molecules on RGO-para-AZO are 59.5%, 60.4%, and 59.1% based on residue weights at 700, 750, and 800°C. Similarly, RGO-ortho-AZO hybrid shows the functionalization degree of AZO by 55.4%, 56.8%, and 58.2% at 700, 750, and 800°C. In addition, the degree was also estimated by element composition data in XPS. Based on the relative atomic percentage of sulphur and nitrogen in Table S2, weight percentages of para-AZO and ortho-AZO in hybrids are 61.14% and 63.78%, respectively. The average quantitative calculation based on TGA and XPS indicates that one AZO molecule is approximately covalently supported by per 11–12 carbon atoms of RGO by the diazotization. The densities confirmed by TGA and XPS pattern are much higher than that of AZO/graphene hybrid reported recently. The dense attachment favors the formation of intermolecular H-bonds of AZO on RGO.

The interlayer effect and the crystallization of RGO-AZO hybrid are analyzed by XRD spectra (Figure 4c). The characteristic 2θ peak of GO at 11.5° corresponds to (0 0 1) interplanar spacing of 1.16 nm consistent with the thickness of GO in atomic force microscopy (AFM) images (Figure S2). RGO shows a broad band of (0 0 2) peak of graphite at 2θ = 23.2° corresponding to small d-spacing of 0.38 nm because of the aggregation of partially reduced sheets. The aggregation might arise from van der Waals interaction among...
nanosheets because the reduction weakens electrostatic repulsion. Therefore, the ultrasonication is necessary to disturb the stacking of nanosheets for well-dispersion of RGO before the functionalization with AZO molecules\(^{27}\). The poor crystallization is found in RGO-AZO hybrids due to the thick organic layer attached on RGO. The weak peaks (0 0 2) of RGO-\(_{\text{para}}\)-AZO and RGO-\(_{\text{ortho}}\)-AZO with remarkably low intensity shifted to \(2\theta = 24.8^\circ\) and 25.3\(^\circ\). This feature arises from stacked layers of sheets coated by non-crystalline AZO through the radical addition.

**Discussion**

Isomerization and thermal reversion of AZO are investigated by UV-vis absorption spectra. Time-evolved absorption spectra of RGO-\(_{\text{para}}\)-AZO and RGO-\(_{\text{ortho}}\)-AZO under the irradiation and after the irradiation in darkness were shown in Figure 5. Generally, one of strategies to optimize solar thermal storage is to maximize the absorption of sunlight. Compared with RGO-\(_{\text{para}}\)-AZO at 344 nm, the maximum peak of E-\(_{\text{ortho}}\)-AZO on RGO is 424 nm with a broad band with the edge up to 550 nm ascribed to \(\pi-\pi^*\) transition, indicating the control of isomerization by visible light\(^{8,10,11,28}\). Photoisomerization of RGO-\(_{\text{para}}\)-AZO (Figure 5a) and RGO-\(_{\text{ortho}}\)-AZO (Figure 5c) hybrids is indicated by continuously decreased bands of \(_{\text{para}}\)-AZO at 344 nm and \(_{\text{ortho}}\)-AZO at 424 nm owing to the transformation from E- to Z-isomers. Electron transition in visible region and long-time isomerization (25 h until a photostationary equilibrium) indicate efficient absorption of high-energy solar power. This absorbed solar energy is basically important because it is required to be two-fold or three-fold higher than activation barrier \(\Delta E_i\) (energy required for isomerization of per AZO molecule)\(^2\). According to previous studies, \(\Delta H\) (energy stored by per AZO molecule) for per metastable Z-isomer must be less than \(\Delta E_i\) in terms of equation (1), and the storage efficiency (the ratio of stored energy to absorbed solar energy) usually was <30% according to a review paper\(^2\).

\[\Delta H = \Delta E_i - \Delta E_a\]  

(1)

The kinetics and thermal barrier (\(\Delta E_a\) for per Z-AZO molecule) were also studied. Thermal reversion of AZO with \(_{\text{ortho}}\)- and \(_{\text{para}}\)-substitution is observed in Figure S6. Compared with rapid reversion of \(_{\text{para}}\)-AZO, Z-\(_{\text{ortho}}\)-AZO is thermally stable with a long \(t_{1/2}\) (the time required for a quantity of Z-AZO to fall to half its value by \(Z \rightarrow E\) reversion) of 116 hours at room temperature due to the formation of intramolecular H-bonds in Z-isomer (Figure S7). Thermal reversion of RGO-\(_{\text{para}}\)-AZO (Figure 5b) and RGO-\(_{\text{ortho}}\)-AZO (Figure 5d) in darkness is further studied by absorption spectra. After the functionalization, both two RGO-AZO hybrids show an appreciably slower thermal reversion of \(Z \rightarrow E\) isomerization with first-order rate constant several orders of magnitudes lower than that of AZO molecules (Figure 5e). An interesting result is found that RGO-\(_{\text{ortho}}\)-AZO hybrid exhibits remarkable long-term thermal reversion of Z-isomers with a \(t_{1/2}\) of 5400 hours (225 days), which is much longer than that (116 h) of RGO-\(_{\text{para}}\)-AZO. The long \(t_{1/2}\) indicates high \(\Delta E_a\) for per metastable Z-AZO on RGO\(^{13}\). As shown in Table 1, \(\Delta E_a\) for Z-\(_{\text{ortho}}\)-AZO on RGO up to 1.2 eV calculated by equation (2) substantially outperforms other AZO derivatives\(^{5,7-10,29}\) and noticeably conquers the limitation of short reversion of Z-AZO.
The activation energy, $\Delta E_a$, is given by:

$$\Delta E_a = -RT\ln\frac{h\ln 2}{\tau_{1/2}k_BT}$$

where $T$ is the storage temperature and $\tau_{1/2}$ is the half-life. $k_B$, $R$, and $h$ are the Boltzmann, gas and Planck constants, respectively.

An important strategy to tune thermal reversion of AZO is the incorporation of electron-withdrawing or donating groups in the ortho position with molecular interaction. Intramolecular H-bonds in RGO-AZO hybrid are indicated by long-term reversion and large chemical shifts in $^1H$ NMR. According to previous calculation, the di-ortho-substitution (-OH) of Z-AZO on CNT resulted in a large increase in $\Delta E_a$ in Table 1 based on the stabilization of Z-isomer by intramolecular H-bonds. Thus, compared with RGO-para-AZO showing no intramolecular H-bonds, high $\Delta E_a$ indicates the formation of intramolecular (=N---H-O) in ortho-AZO molecules on RGO. This analysis is supported by large chemical shifts (-O1H) of RGO-ortho-AZO at $\delta = 13.85$ in $^1H$ NMR, which is consistent with the chemical shift at (-O1H) $\delta = 14.19$ of ortho-AZO. Furthermore, RGO-AZO hybrids also show a good cycling stability of isomerization and thermal reversion, which follows first-order kinetics (equation S1). As shown in Figure 5, first-order rate constants for reversion ($k_{rev}$) of RGO-para-AZO and RGO-para-AZO only fluctuate in the range of $3.52-3.63 \times 10^{-8} s^{-1}$ and $1.64-1.68 \times 10^{-8} s^{-1}$ respectively during five cycles. The measure-
ment of \( \kappa_{\text{rev}} \) of RGO-ortho-AZO is based on the kinetics of thermal reversion before \( \tau_{1/2} \) (5408 h). This feature indicates a good photostability of ortho-AZO and para-AZO on RGO. Good cycling stability of reversible isomerization-reversion is important for solar thermal storage.

The capacity of solar thermal storage is another key performance parameter for the utilization. The theoretical capacity of thermal energy of RGO-AZO hybrids is calculated using the PAW formalism of spin polarized DFT based on \( \Delta H \) for per AZO molecule and packing density. The relaxed steric structures of Z- and E-isomers of RGO-para-AZO and RGO-ortho-AZO hybrids were shown in the inset of Figure 6, in which green and brown dash line indicate inter- and intramolecular H-bonds, respectively. The model is that one AZO molecule is supported by 16 carbon atoms of RGO. The \( \Delta H \), packing density (the weight percentage of AZO for RGO-AZO according to experimental measurement and calculation) and energy density (thermal energy stored by a quantity of RGO-AZO) are shown in Table 2. Compared with pure AZO molecules (para-AZO: 0.558 eV, ortho-AZO: 0.77 eV), one-fold increase in \( \Delta H \) for per AZO molecule is obtained due to the increased energy level between Z- and E-isomers by the formation of intermolecular H-bonds between two adjacent AZO molecules on RGO according to stable configurations (dash line Figure S8).

The different numbers of H-bonds between ortho-AZO and para-AZO on RGO are indicated by high \( \Delta E_a \) (1.2 eV) of RGO-ortho-AZO controlled by intramolecular H-bonds in Z-ortho-AZO and high \( \Delta H \) (1.38 eV) of RGO-para-AZO controlled by one intramolecular H-bond in E-para-AZO. This analysis is also confirmed by relaxed steric structures of RGO-para-AZO and RGO-ortho-AZO hybrids. As shown in Table 2, RGO-para-AZO hybrid shows a higher \( \Delta H \) of 1.38 eV than that of RGO-ortho-AZO (1.224 eV). This feature indicates that \( \Delta H \) for per AZO on RGO is controlled by the number of intra- and intermolecular H-bonds formed in both Z- and E-isomers on RGO. Previous calculation demonstrated that maximizing the number of H-bonds in E-AZO and minimizing H-bonds in Z-AZO provided a high \( \Delta H \). As shown in Figure S8, there is one intramolecular H-bond of \(-\text{S-OH}--\text{OH-S}^+\) in E-para-AZO (confirmed by red-shifted bands in Figure 3a) and no H-bonds in Z-para-AZO on RGO, while both Z-ortho-AZO and E-ortho-AZO on RGO show one intramolecular H-bond of \(=\text{N}--\text{HO}^+\) (confirmed by large chemical shift at 13.85) and one intermolecular H-bond of \(=\text{S-OH}--\text{OH-S}^+\) (confirmed by red-shifted bands in Figure 3d). Intramolecular H-bonds of RGO-ortho-AZO are also indicated by high \( \Delta E_a \) E-para-AZO and Z-para-AZO on RGO exhibit a larger difference in number of H-bonds than that of RGO-ortho-AZO. Results suggest that RGO-para-AZO with only one intramolecular H-bond results in a high \( \Delta H \) but a low \( \Delta E_a \). In contrast, intramolecular H-bonds in Z-ortho-AZO on RGO increase \( \Delta E_a \) but multiple (intra- and intermolecular) H-bonds in both Z- and E-ortho-AZO lead to a low \( \Delta H \). Table 2 displays that compared with RGO-para-AZO, the increase in \( \Delta E_a \) by 9% \((\tau_{1/2} up to \approx 5000 h) \) of RGO-ortho-AZO is gained at the sacrifice of the decrease in \( \Delta H \) by 11%. This feature suggests that \( \Delta H \) and \( \Delta E_a \) are determined by different molecular H-bonds of Z- and E-AZO controlled by the substitution and microstructures on RGO. Thus, maximizing both \( \Delta H \) and \( \Delta E_a \) by optimizing intra- and intermolecular H-bonds of AZO bound on graphene still needs more deep studies in the future.

The amount of heat released by RGO-AZO solution during Z \( \rightarrow \) E reversion determined by \( \Delta H \) for per AZO on RGO was measured using a three-dimensional high-precision calorimetric sensor in a sealed insulating vessel. The solution was completely sealed by thick walls to keep the constant temperature at 19–20 °C during the measurement. The densities of heat (related to the capacity of thermal energy stored by a quantity of RGO-AZO) released by (a) RGO-para-AZO and (b) RGO-ortho-AZO hybrids with the time in darkness after the irradiation are shown in Figure 6. Curves follow three-order fitting exponential decay (Table S3).

The density of heat released by RGO-para-AZO hybrids is 269.8 kJ kg\(^{-1}\) when hybrids are kept in a sealed dark vessel after the irradiation for 600 h, which is the time required for almost complete reversion (>95 wt.% Z-para-AZO transforms to E-para-AZO) calculated based on the change in absorption intensity (Figure 5). Meanwhile, compared with RGO-para-AZO, RGO-ortho-AZO shows a relative low density of 149.6 kJ kg\(^{-1}\) when hybrids are kept for 5400 h, which is only \( \tau_{1/2} \) for Z-ortho-AZO on RGO. Thus, this density might be 50% capacity of energy stored by hybrids. It’s definitely difficult to measure the maximum capacity of heat storage for RGO-ortho-AZO hybrid because of a very long time of complete reversion up to 23333 hours (972 days, according to \( \kappa_{\text{rev}} \) in Table 2). The measurement might be disturbed by the environmental temperature. Despite that, the densities of thermal storage of RGO-AZO hybrids are better than that of azo-polymers and derivatives reported recently due to the formation of intra- and intermolecular H-bonds in RGO-AZO. As shown in Table 2, the densities of heat released by RGO-AZO hybrids in experiment are of the same order magnitude to that (RGO-para-AZO: 312.2 kJ kg\(^{-1}\) and RGO-ortho-AZO: 274.9 kJ kg\(^{-1}\)) based on \( \Delta H \) calculated by DFT and

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Table 1 | The parameter and thermal barrier \( \Delta E_a \) of Z \( \rightarrow \) E thermal reversion

| Parameter          | \( \kappa_{\text{rev}} \) (s\(^{-1}\)) | \( \tau_{1/2} \) (h) | \( \Delta E_a \) (eV) |
|--------------------|----------------------------------------|----------------------|----------------------|
| Para-AZO           | 0.295                                  | 2.35                 | 0.79                 |
| RGO-para-AZO       | 1.66 \( \times \) 10\(^{-6}\)          | 116                  | 1.1                  |
| Ortho-AZO          | 1.60 \( \times \) 10\(^{-6}\)          | 120                  | 1.1                  |
| RGO-ortho-AZO      | 3.56 \( \times \) 10\(^{-8}\)          | 5408                 | 1.2                  |

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Table 2 | \( \Delta H \), packing density and energy density of thermal storage based on experimental measurement and calculation

| Parameter          | \( \Delta H \) (eV) | Packing Density (wt%) | Energy Density (kJ/kg)          |
|--------------------|---------------------|-----------------------|---------------------------------|
| Para-AZO           | 0.558               | /                     | 217.9                           |
| RGO-para-AZO       | 1.38                | 57.9                  | 372.2                           |
| Ortho-AZO          | 0.77                | /                     | 282.4                           |
| RGO-ortho-AZO      | 1.224               | 60.6                  | 274.9                           |

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\( \Delta H \) and \( \Delta E_a \) are determined by different molecular H-bonds of Z- and E-AZO controlled by the substitution and microstructures on RGO. Thus, maximizing both \( \Delta H \) and \( \Delta E_a \) by optimizing intra- and intermolecular H-bonds of AZO bound on graphene still needs more deep studies in the future.

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Figure 6 | The density of heat released by (a) RGO-para-AZO and (b) RGO-ortho-AZO hybrids with the time of storage in darkness (a sealed vessel) after the irradiation and inset shows the stable structures of RGO-AZO hybrids based on the calculated by DFT. White, gray, blue, red and yellow sticks represent H, C, N, O and S atoms, respectively.
packing density. Besides, the volumetric density of thermal storage for RGO-para-AZO and RGO-ortho-AZO hybrid is about 180 and 100 Wh/L.

**Conclusion.** Para-AZO and ortho-AZO bound on RGO with different molecular H-bonds controlled by the substitution and high packing density were presented for solar thermal storage. The covalent attachment showed that per AZO molecule was supported by 11–12 carbon atoms of RGO according to the characterization of chemical structures and thermal stability. Intra- and Intermolecular H-bonds in RGO-para-AZO hybrid were confirmed by large chemical shifts in 1H NMR, red-shifted band in FT-IR, and relaxed structures calculated by DFT, leading to the increase in ΔEh and ΔEH for per AZO on RGO, respectively. Intramolecular H-bonds thermally stabilized Z-ortho-AZO on RGO with a long-term τ1/2 of 5400 h (ΔEg = 1.2 eV), which was much longer than that of RGO-para-AZO (116 h). RGO-para-AZO with one intramolecular H-bond showed high density of thermal storage up to 269.8 kg−1 compared with RGO-ortho-AZO (149.6 kg−1) with multiple intra- and Intermolecular H-bonds in both Z- and E-AZO. Thermal storage in experiment are of the same order magnitude to that (RGO-para-AZO: 312.2 kg−1 and RGO-ortho-AZO: 274.9 kg−1) calculated based on ΔH calculated by DFT and packing density. Results indicated that ΔH and ΔEH are determined by different molecular H-bonds of Z- and E-AZO on RGO. Thus, optimizing intra- and Intermolecular H-bonds of both Z-AZO and E-AZO bound on graphene is critical for maximizing both ΔH and ΔEH. RGO-AZO hybrid with the density of 150 ~ 300 kg−1 and half-lives of 1000–5000 h can be utilized for a long-term and renewable solar thermal storage.

**Methods**

**Materials.** Graphite was obtained from Qingdao Huarun graphite Co., Ltd., with a particle size of 20 μm. Para-AZO and all other chemicals were bought from Sigma-Aldrich and used as received.

**Synthesis of RGO Single-layer.** GO was prepared through acid oxidation of flake graphite. GO were dispersed and partially reduced in 30 mL of NaBH4 solution (pH = 9) at 80°C for 1 h. RGO was obtained after rinsing with di-water, the filtration and the centrifugation at 500 rpm for 10 min.

**Synthesis of ortho-AZO.** Orthanic acid (18 mmol), NaOH (18 mmol) and NaNO2 (20 mmol) were dissolved in 120 mL of di-water, and the solution was slowly added into 70 mL of HCl (1 mol/L) in an ice bath at 0–5°C for the diazotization. M-Aminophenol (18 mmol) was dissolved in 40 mL of HCl. The diazotized compounds were added into m-amino phenol solution (pH = 5–7) at 0–5°C for 2 hours. The resultant raw materials were re-crystallized by EtOH/H2O (1:1) for several times, and ortho-AZO (14 mmol, Yield: 78%) was collected after washing with di-water, the filtration and dry in vacuum overnight at 70°C. 1H NMR (CD2Cl2, 6): 14.91 (s, 1H, Ar-OH), 5.39 (s, 2H, Ar-NH2), 7.84 (d, J = 8.2 Hz, 1H, Ar-H), 7.49 (dd, J = 7.7 Hz, 1.5 Hz, 1H, Ar-H), 7.45 (d, J = 9.3 Hz, 1H, Ar-H), 7.27 (t, J = 7.7 Hz, 1.5 Hz, 1H, Ar-H), 6.61 (dd, J = 9.3, 1.8 Hz, 1H, Ar-H), 6.21 (d, J = 2Hz, 1H, Ar-H).

**Synthesis of RGO-AZO hybrid.** Azo molecules were covalently functionalized with RGO by the diazotization using diazonium salts. RGO was dispersed in water to form a homogeneous solution by ultrasonication for 2 hours. The aggregate was removed by centrifugation at 1000 rpm for 20 min, yielding a typical concentration of 1 mg/mL RGO solution (150 mg).

**Characterization.** FT-IR spectra were recorded on a Bruker Tensor 27 spectrometer with a disc of KBr. 1H NMR spectra of ortho-AZO and RGO-ortho-AZO were carried out on a Varian INOVA 500 MHz spectrometer with trimethylsilyl as an internal standard. X-ray (1250 eV) source at a base pressure in the 10−7 to 10−8 Torr range. X-ray diffraction (XRD) patterns were taken by a Rigaku D/max 2500 V/pc X-ray diffractometer using Cu Ka radiation (K = 0.154 nm) at a scanning rate of 8/0.01°, using a voltage of 40 kV and a current of 200 mA. Thermal analysis was studied using a Thermogravimetric Analyzer Instrument (NETZSCH STA 449C). The samples were heated from 30°C to 700°C at a rate of 5°C/min in an aluminum crucible under 50 mL/min of nitrogen purging. XPS analyses were performed on a PHI 1600 model surface analysis system with a 2.500 eV MgKa and a 3500 eV AlKA electron source (VG Multilab2000) and scanning electronic microscope (SEM, Hitachi S-4800) were both performed to observe the morphologies. The surface of GO was observed by AFM (Veeco Multimode III).

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