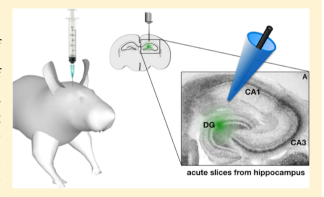


Graphene Oxide Flakes Tune Excitatory Neurotransmission in Vivo by Targeting Hippocampal Synapses

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ABSTRACT: Synapses compute and transmit information to connect neural circuits and are at the basis of brain operations. Alterations in their function contribute to a vast range of neuropsychiatric and neurodegenerative disorders and synapse-based therapeutic intervention, such as selective inhibition of synaptic transmission, may significantly help against serious pathologies. Graphene is a two-dimensional nanomaterial largely exploited in multiple domains of science and technology, including biomedical applications. In hippocampal neurons in culture, small graphene oxide nanosheets (s-GO) selectively depress glutamatergic activity without altering cell viability. Glutamate is the main excitatory neurotransmitter in the central nervous system and growing evidence suggests its involvement in neuropsychiatric disorders. Here we demonstrate that s-GO directly targets the



release of presynaptic vesicle. We propose that s-GO flakes reduce the availability of transmitter, via promoting its fast release and subsequent depletion, leading to a decline ofglutamatergic neurotransmission. We injected s-GO in the hippocampus in vivo, and 48 h after surgery ex vivo patch-clamp recordings from brain slices show a significant reduction in glutamatergic synaptic activity in respect to saline injections.

KEYWORDS: Graphene, synapses, hippocampal network, glutamate, quantum dots

raphene is a two-dimensional (2D) material made of sp²-hybridized carbon atoms organized in a hexagonal lattice and characterized by excellent physical features, including outstanding electron mobility and mechanical flexibility. 1-3 Because of its properties, 4-6 graphene is considered a rising star in a growing number of technological developments, including biomedical ones. 2,4,5,7 In neurology, graphene-based neuronal implants or biodevices may overcome current technical limitations in treating pathologies that range from neurooncology to neuroregeneration. 8,9 We reported recently the ability of small, thin graphene oxide sheets (s-GO) to alter specifically neuronal synapses with no impact on cell viability. In particular, in cultured hippocampal networks upon chronic long-term exposure to s-GO glutamatergic release sites were sized down. 10 It is well-known that glutamate is the main

excitatory neurotransmitter in the mammalian central nervous system (CNS) and mediates neuronal development, migration, synaptic maintenance, and transmission. An uncontrolled release of glutamate in the extracellular space may lead to excitotoxicity, neurodegeneration, and neurological disorders, including pain. Localized targeting and fine-tuning of the glutamatergic system are attractive objectives in neuroscience. To achieve a deep understanding of the interactions between s-GO and the machinery governing nerve cell functions is mandatory to translate these findings into potential therapeutic applications. In particular, graphene translocation or adhesion

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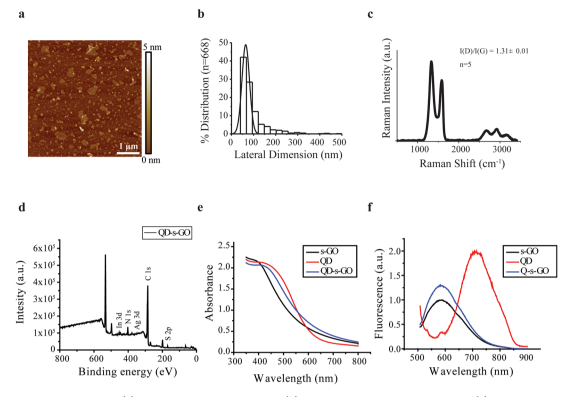


Figure 1. Characterization of s-GO. (a) AFM measure of s-GO sheets. (b) s-GO lateral dimension distribution. (c) S-GO Raman spectrum. (d) XPS survey of QD-s-GO. UV-vis (e) and fluorescence (f) spectra of s-GO, QD, and QD-s-GO.

to cell membranes 15,16 may potentially interfere with activities such as the exocytic and endocytic trafficking systems, essential to physiological synaptic transmission. 15,17 Here, we describe by single cell electrophysiology how s-GO nanosheets acutely tune synaptic release in excitatory synapses of hippocampal cultured neurons and acute slices by interfering with the probability of vesicle release. We propose that such interference leads to transmitter depletion and subsequent depression of the glutamatergic activity. We next address whether such material similarly affects glutamatergic transmission in vivo, by injecting s-GO in the dentate gyrus of the hippocampus of juvenile rats. We patch-clamped single neurons from ex vivo hippocampal slices 48 and 72 h after s-GO microinjections. We demonstrate that s-GO targets and down-regulates glutamatergic synapses in vivo and further illustrates the potential of s-GO flakes to be engineered as specific synaptic transmission modulators.

GO Functionalization and Characterization. The produced s-GO dispersion was visually homogeneous and of a brownish-translucent appearance. The dispersions did not show any evidence of sedimentation or any other observable changes for over 6 months, indicating their physical stability. The characterization of the s-GO nanosheets is presented in Figure 1 and in Supplementary Figure S1. The morphological features of the s-GO nanosheets were examined using AFM (Figure 1a) and TEM (Supplementary Figure S1a). Both analytical methods showed that the lateral dimension of the s-GO nanosheets was predominantly between 100-300 nm with very few larger sheets into the micrometer range (Figure 1b). Moreover, atomic force microscopy (AFM) revealed that the material is composed of sheets from single to few-layer thickness (Supplementary Figure S1b). The material structural features were studied by Raman spectroscopy which evidenced the presence of the characteristic G and D scatter bands at

1595 and 1330 cm⁻¹, respectively (Figure 1c). The D scatter band was markedly higher than the G band. The intensity ratio of these two peaks, known as the I(D)/I(G), was calculated to be 1.31 \pm 0.01, indicating that the material hexagonal lattice was defected. XPS analysis corroborated to the presence of functional groups (Supplementary Figure S1c) and further indicated that the defects correspond to oxygen-containing functionalities. The C/O ratio was found to be 2.1 and the material chemical purity was 99.8%. The surface functionalization was further supported by laser Doppler electrophoresis to indicate that the dispersed sheets had a surface charge of -55.9 ± 1.4 mV. Aiming to track the s-GO flakes within neuronal tissue, we performed covalent labeling of s-GO with quantum dot (QD) luminescent nanoparticles. For this purpose, we first synthesized the AgInS2/ZnS-doped QDs capped with cysteine as described in the literature (see TEM images in Supplementary Figure S1d). 19,20 Subsequently, the coupling with s-GO was achieved via epoxy ring opening with the amino groups of the cysteine-capping agent. Transmission electron microscopy (TEM) shows the presence of small dark dots on the s-GO sheets associated with the presence of QDs on the surface (Supplementary Figure S1e, indicated by the arrows) as confirmed by X-ray photoelectron spectroscopy (XPS) survey analysis (Figure 1d). The UV-vis spectrum (Figure 1e) of the functionalized material showed a broadening of the absorption band between 300 and 600 nm due to the presence of the nanocrystals onto GO. Fluorescence characterization is reported in Figure 1f; s-GO showed an emission centered at 585 nm attributed to the electronic transitions from the bottom of conductive band and the nearby localized states to the valence band.²¹ QDs have an emission centered at 706 nm due to transition between the conductive band and the defected carbon lattice.²² Interestingly, when QDs were coupled to graphene oxide only s-GO luminescence

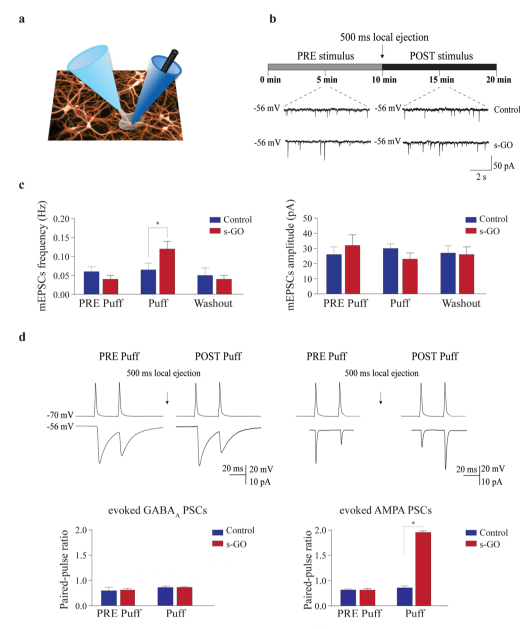


Figure 2. s-GO affects presynaptic glutamate release in hippocampal cultures. (a) Sketch of the experimental setting for simultaneous s-GO pressure-release (puff) and single-cell recording from cultured neurons. (b) Top: diagram of the experimental protocol. Bottom: representative tracings of the spontaneous synaptic activity detected prior and after puff applications of control saline (Control, top) or s-GO (bottom). Recordings of mEPSCs are performed in the presence of TTX. In (c) bar plots of pooled data summarize the average mEPSCs frequency (left) and amplitude (right) before (PRE puff) and after (Washout) saline (Control) or s-GO (100 μ g/mL final concentration) pressure ejections (*P < 0.05 Student's t test). Note the reversible increase in miniatures frequency due to s-GO. In (d) simultaneous pair recordings are shown: top traces represent presynaptic pairs (30 Hz) of action potentials and bottom ones represent the corresponding evoked monosynaptic PSCs (GABA_A-receptor mediated on the left and glutamate AMPA-receptor mediated on the right) prior and after s-GO puffs. The paired-pulse ratios (PPR) measured prior and after saline solution (Control) and s-GO puffs are summarized in the histograms; note that s-GO reduced the first evoked AMPA-receptor mediated PSCs and the PPR of glutamatergic synapses (*P < 0.05 Student's t test) supporting the notion of s-GO affecting presynaptic release.

was detected. The quenching of the QD emission may be attributed to an interfacial electron transferbetween the QDs and the s-GO surface due to their close proximity. Surprisingly, the emission band centered at 585 nm attributed to the GO photoluminescence, appeared stronger in the case of QD-s-GO than in nonmodified s-GO. Most probably, the energy transfer process causes the decrease of donor emission (QD quenching) and increases the s-GO acceptor emission²⁴ allowing us to visualize the s-GO-QD in the biological environment (vide infra).

s-GO Targets Synaptic Vesicle Release at Glutamatergic Synapses in Cultured Hippocampal Neurons. To unravel the mechanisms by which thin s-GO sheets affect neurotransmission, we patch-clamped cultured hippocampal neurons while a second pipette for the local delivery of standard saline solution of s-GO (100 μ g/mL; see Methods) was positioned at 200 μ m distance (by microscopic guidance) from the recorded neuron (sketched in Figure 2a). We estimated that at this distance the application of a brief (500 ms) pulse of pressure should result in a local (i.e., on the

patched cell) and transient delivery of s-GO at a concentration of at least 10% of that contained in the pipette (see Methods). Spontaneous synaptic activity was recorded in the presence of Tetrodotoxin, (TTX; 1 µM). In TTX, synaptic events, termed miniature post synaptic currents (mPSCs), reflect the presynaptic, stochastic release of vesicles at individual synaptic terminals impinging on the recorded neuron. mPSCs frequency reflects the presynaptic release probability and on the number of synaptic contacts, while mPSCs amplitude is dictated by postsynaptic receptor sensitivity. 25 Baseline mPSCs were sampled before and after the local ejection of saline or s-GO (Figure 2b). In cultured neurons, virtually all mPSCs were made up by excitatory (AMPA glutamate receptor-mediated) events, identified by their fast kinetics (decay time constant $\tau =$ 5 ± 0.5 ms; ²⁶) and were thus named excitatory mPSCs (mEPSCs). Figure 2b shows that representative control (top) and s-GO (bottom) current tracings prior and after saline or s-GO solution, respectively, were pressure ejected. In control neurons, mEPSCs frequency did not change (from 0.06 ± 0.03 Hz to 0.065 ± 0.04 Hz after saline-ejection, n = 14; bar plot in Figure 2c, left). On the contrary, acute s-GO ejection significantly increased (* P < 0.05 Student's t test) the mEPSCs frequency (from 0.04 \pm 0.01 to 0.12 \pm 0.02, Hz n=13; bar plot in Figure 2c, left). The increase in mEPSCs appeared with 8-10 s delay from the local s-GO ejection and completely reversed to baseline values (0.04 \pm 0.01 Hz) of 8-9 min following the acute application (bar plot in Figure 2c, left). In all treatments, the mEPSCs amplitude was not affected (bar plot in Figure 2c, right). These transient changes in the frequency of mEPSCs suggest a direct interference of s-GO with the presynaptic release machinery^{27,28} and are consistent with the hypothesized targeting by s-GOs of endoexocytotic mechanisms. This hypothesis is also validated by the colocalization of bassoon (presynaptic terminal marker²⁹) and s-GO detected by confocal microscopy in a different set of experiments, where s-GO was incubated (20 μ g/mL; 30 min), before fixation of the cultures (see Methods; Supplementary Figure S2a,b, controls and s-GO, respectively). We further address the dependency of these effects on the flakes' size. We adopted the same protocol to press-eject GO flakes (same concentration as s-GO) characterized by different lateral dimensions: large GO (l-GO, $\approx 2 \mu m$) or ultrasmall GO (us-GO, ≈ 40 nm).¹⁸ Supplementary Figure S2 c shows representative control (top) and l-GO (bottom) current tracings sampled before and after the local ejection of saline or l-GO solutions. Opposite to s-GO, l-GO did not change mEPSCs frequency (from 0.06 ± 0.01 to 0.07 ± 0.02 Hz after saline ejection, n = 5; from 0.05 ± 0.01 to 0.07 ± 0.02 Hz after 1-GO ejection, n = 5). Similarly (Supplementary Figure S2d), us-GO did not modulate mEPSCs frequency (from 0.05 ± 0.01 to 0.07 \pm 0.02 Hz, after saline, n = 8, and from 0.05 \pm 0.01 to 0.07 \pm 0.02 Hz, after us-GO, n = 8).

Thus, only s-GO transiently increased the frequency of mEPSCs. This apparent discrepancy with our previous results, where prolonged exposure to s-GO decreased glutamatergic activity, may be explained by the emergence of glutamate depletion due to forced glutamate release. The latter leading to a transient facilitation was followed, when s-GO is applied longer than the duration of the facilitatory effects, by a depression of vesicle release and thus a down-regulation of glutamate transmission. 10

To investigate the s-GO interference with presynaptic release and whether this was truly selective for excitatory

synapses, we tested the local delivery of s-GO nanosheets on the occurrence of evoked PSCs (ePSCs), by simultaneous whole-cell recordings from two monosynaptically connected neurons.²⁶ Action potentials were induced in the presynaptic neuron and the evoked postsynaptic unitary PSCs (delay 2 ms) were examined. In our in vitro system, monosynaptically coupled pairs of neurons typically display either GABA_A or glutamate AMPA receptor-mediated evoked currents. ^{26,30,31} We identified the different populations of ePSCs on the basis of their kinetic properties and pharmacology. 26,32 In fact, GABAergic ePSCs were characterized by a slow decay time constant ($\tau = 23 \pm 7$ ms, n = 15 for each condition, control and s-GO; Figure 2d, left) and were fully abolished by administration of 5 μ M Gabazine (n = 3). Glutamatergic AMPA receptor-mediated ePSCs displayed fast decay ($\tau = 7 \pm$ 1.2 ms, n = 7 for each condition; Figure 2d, right) and were further blocked by application of 10 μ M CNQX (n = 3). To investigate the presynaptic properties, we adopted paired-pulse stimulation protocols. 33,34 In paired-pulse stimulation the second response can be either facilitated or depressed. Usually, at a specific synapse an increased probability of neurotransmitter release will favor paired-pulse depression, whereas a decrease in the release probability favors facilitation. 33,35,36 Thus, differences in postsynaptic responses to paired-pulse stimulation indicate variations in presynaptic transmitter release. 33,36–38 To probe the changes in efficacy of unitary ePSCs, paired-pulse protocols were performed with short interstimulus interval (50 ms). Figure 2d shows representative presynaptic pairs of action potentials (top) and the corresponding monosynaptic GABAergic (left tracings) or glutamatergic AMPA receptor (right) evoked currents (bottom) before and after s-GO local pressure ejection. We indirectly assessed the GABA and glutamate release probability before and after saline (control) or s-GO ejection by measuring the paired-pulse ratio (PPR, calculating the ratio between the mean peak amplitude of the second and the first PSC^{37,38}). In control GABAergic and glutamatergic ePSCs, the resulting PPR indicated the presence of paired pulse depression and did not change upon saline solution applications (for GABA_A receptor-mediated pairs, 0.5 ± 0.2 before and 0.6 ± 0.2 after saline; for AMPA receptor-mediated pairs, 0.5 ± 0.1 before and 0.6 ± 0.2 after saline, plot in Figure 2d). When investigating the impact of s-GO ejection, we detected a reduction (on average -32%) in the amplitude of the first glutamatergic ePSC and a significant difference (* P < 0.05 Student's t test) in PPR, indicative of paired-pulse facilitation, while the PPR did not change in GABAA ePSCs (for AMPA mediated pairs, 0.5 ± 0.1 before and 2 ± 0.9 after s-GO; for GABA_A mediated pairs, 0.5 ± 0.2 before and $0.65 \pm$ 0.2 after s-GO; summarized in the bar plots in Figure 2d, bottom). Altogether these experiments strongly support a direct interference of s-GO flakes with synaptic vesicle release, with an initial high rate of release followed by a decline when s-GOs are applied longer 10 or when acting synergistically to the action potential-evoked activation of the exocytotic apparatus, ultimately depleting evoked release, typically reflected by changes in ePSC amplitude. 39,40 Notably, only glutamatergic synapses were targets of the s-GO.

s-GO Exposure Specifically Affects Glutamatergic Synapses in Acute Hippocampal Slices. Since cultured networks are simplified 2D models of immature brain circuits, we explored the ability of s-GO to regulate glutamate synaptic activity in acute hippocampal slices, thus scaling up the

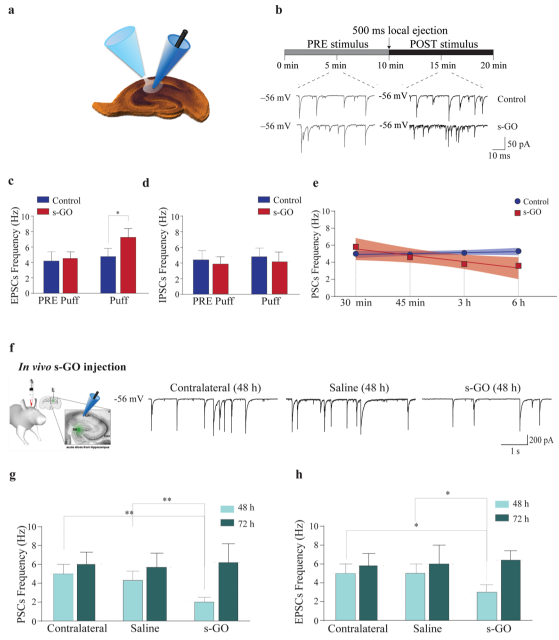


Figure 3. EPSCs frequency modulation by s-GO in hippocampal slices in vitro and in vivo. (a) Sketch of the experimental setting for simultaneous s-GO release and recording from hippocampal pyramidal cells. In (b) top: diagram of the experimental protocol. Bottom: representative current tracings recorded prior and after saline (Control; top) and s-GO (bottom) local pressure ejections. Glutamate AMPA-receptor mediated PSCs or GABAA-receptor mediated ones (EPSCs and IPSCs, respectively) were pharmacologically isolated and bar plots in (c) summarize the mean values of EPSCs and in (d) of IPSCs frequency before and after saline (Control) or s-GO puffs (*P < 0.05 Student's t test). Note that also in hippocampal slice explants only glutamatergic activity was transiently affected by brief local injection of s-GO. In (e), plots of pooled data represent the average PSCs frequency upon 30 min, 45 min, 3 h, and 6 h s-GO incubation (50 μ g/mL final concentration; Control, blue circles; s-GO, red squares). Note that prolonged incubation in s-GO depresses spontaneous synaptic activity. Linear regression analysis of the two time progressions is depicted as blue and red fitting lines (y = 4.91 + 0.11x for Control and y = 5.56 - 0.74x for s-GO, respectively) together with their corresponding confidence interval in pale blue and pale red, respectively. Regardless, the significance of the difference between each two conditions at a specific time point (at 30 m, P = 0.91; at 45 m, P = 0.60; at 3 h, P = 0.07; and at 6 h, P = 0.06), multiple regression statistical analysis revealed that the zero slope hypothesis is accepted for Controls but not for s-GO. The equal slope hypothesis between the two trends was instead rejected by a TU.K.ey test on the two slopes. (f) In vivo intrahippocampal s-GO delivery reversibly reduces glutamatergic synaptic activity in adult rats: sketch of the experimental settings (left) and (right) spontaneous synaptic activity recorded from ex vivo hippocampal slices isolated from juvenile rats after 48 h from the surgery. Recordings were taken from the contralateral, control (saline), and s-GO (50 µg/mL final concentration) injected hemisphere after 48 h from surgery. In (g) bar plots summarize the PSCs and in (h) the EPSCs frequency in control and s-GO treated animals after 48 h and after 72 h from surgery (**P < 0.001 two-way ANOVA; *P < 0.05 two-way ANOVA). Note that the specific reduction in EPSC frequency at 48 h that was entirely reversed at 72 h.

complexity of the tissue to the third dimension and testing more mature synapses. Single neuron patch-clamp recordings were obtained from visually identified pyramidal cells in the CA1 hippocampal region. A second pipette was again

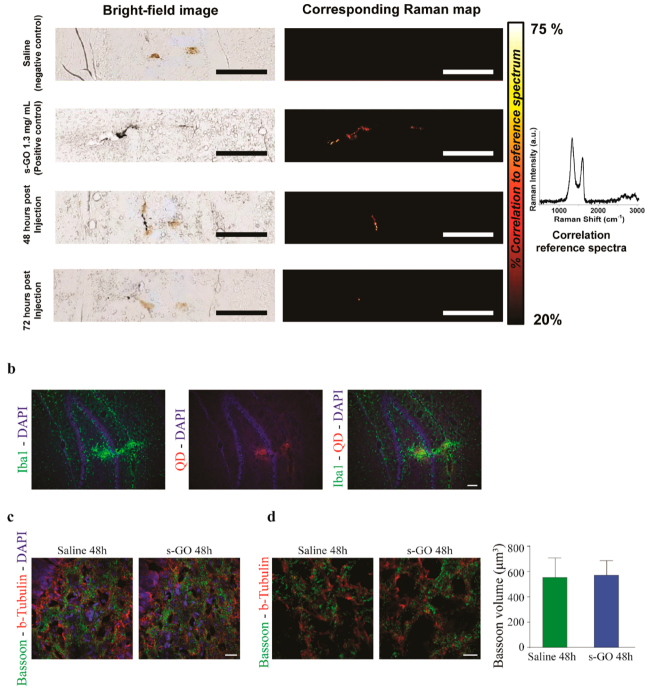


Figure 4. In vivo delivery of s-GO is localized and does not alter excitatory synaptic density. In (a), confocal Raman maps were acquired to establish the location of s-GO within cryosectioned dendate gyrus' of s-GO (1.3 mg/mL) treated animals over time (48 and 72 h). Scale bars = 500 nm. Maps were generated based on the acquired spectra's correlation to a s-GO reference spectra, shown on the right. The dentate gyrus of rats injected with saline (Controls) and rats treated with a higher concentration (1.3 mg/mL) (positive control) were also examined for comparison. The acquisition parameters were as follows λ = 633 nm, laser power = 1 mW, frequency = 25 Hz, and a pixel size of 1.6 μm. In (b), ex vivo fluorescence imaging of hippocampal slices processed for Iba1-positive microglia (in green) and QD linked to s-GO (in red) at the injection site after 48 h. DAPI for nuclei is in blue. Note the precise localization of s-GO within the target area. (c) Ex vivo confocal images of hippocampal synapses at the injection site, excitatory presynaptic terminals were identified by the marker bassoon (in green), in neurons colabeled with β-tubulin III (in red) and results are shown for saline (control) and s-GO injections after 48 h. DAPI for nuclei is in blue. Analysis has been performed at the higher magnification on 70 × 70 μm² ROIs shown in (d) and results are summarized by the bar plots. No differences in bassoon quantification were detected between saline and s-GO injection after 48 h. Scale bars: 100 μm in (b), 25 μm in (c), and 10 μm in (d).

positioned at a distance of 200 μ m from the recorded cell (sketched in Figure 3a) and filled with standard saline solution or with s-GO (100 μ g/mL). Baseline PSCs were recorded

before and after the local saline or s-GO ejection. Figure 3b shows representative current tracings of controls (top) and s-GO (bottom) before and after saline or s-GO solutions,

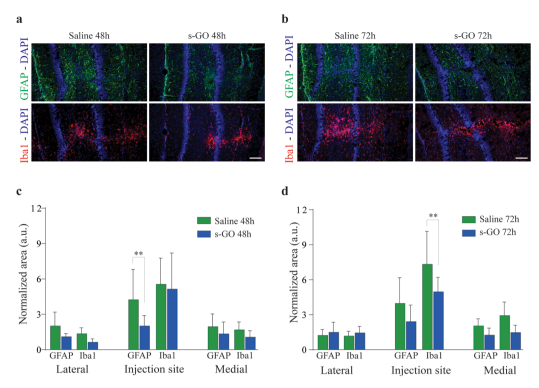


Figure 5. Brain tissue reactivity to surgery and s-GO injections after 48 and 72 h. Ex vivo hippocampal slices from saline (control) and s-GO injected brains were labeled for GFAP-positive astrocytes (in green, top row) and Iba1-positive microglia (in red, bottom row) and the injection site (left dentate gyrus, saline vs s-GO 50 μ g/mL) are shown after 48 h (a) and 72 h (a). DAPI for nuclei is in blue. Scale bar: 100 μ m. Bar plots in (c,d) quantify the glial reaction 48 and 72 h post-surgery. Comparable values of GFAP and Iba1 immunoreactivity between saline and s-GO were observed in the hippocampus at 300 μ m distance from the injection site, either lateral or medial in both 48 and 72 h postsurgery. Notably, at the injection site s-GO induced lower GFAP immunoreactivity at 48 h and lower Iba1 immunoreactivity at 72 h when compared to controls. (** P < 0.01 two-way ANOVA).

respectively, were pressure ejected. In neurons exposed to saline solution, spontaneous PSCs frequency did not change (6 \pm 2 Hz before the pipette saline-ejection and 5 \pm 1 Hz after the pipette saline-ejection, n = 14). On the contrary, acute s-GO ejection significantly increased (* P < 0.05 Student's ttest) the PSCs frequency (from a baseline of 5 ± 2 Hz to a post ejection frequency of 8 ± 2 Hz, n = 13). The increase in PSCs after the local s-GO ejection was reversible. In fact, PSC frequency fully returned to baseline values 7-8 min following the acute application. In all treatments, the PSC amplitude was not affected. We further dissected the nature of PSCs by the use of CNQX or Gabazine, isolating GABAA or AMPA receptor-mediated IPSCs or EPSCs, respectively. When EPSCs were measured after s-GO ejection, we detected a strong increase (* P < 0.05 Student's t test) in their frequency when compared to the saline solution pressure application (4 \pm 1 before and 4.7 \pm 1 Hz after the pipette saline-ejection; 4.2 \pm 0.9 before and 7.3 \pm 1 Hz after the s-GO ejection, Figure 3c). On the contrary, when we measured IPSCs, their frequency was not affected both by saline (4.2 ± 1) before and 4.8 ± 1 Hz after, Figure 3d) and s-GO (3.8 \pm 1 before and 4.2 \pm 1 Hz after, Figure 3d) pressure applications. Such results support the notion of s-GO ability to specifically target excitatory synapses, even in tissue explants. In cultured neurons as well as in acute hippocampal slices, the brief pressure ejection of s-GO transiently increased the excitatory activity, apparently affecting glutamate release machinery at the presynaptic site.

To ascertain whether prolonged interference of s-GO with excitatory synapses might indeed reduce the activity of synapses capable of releasing glutamate, as observed in

dissociated cultures, 10 we incubated acute slices with s-GO (50 μ g/mL) and we monitored PSCs frequency after 30 min (n = 5), 45 min (n = 5), 3 h (n = 5), and 6 h (n = 8). Under these experimental conditions, s-GO will be repeatedly presented at synapses in the absence of the fast clearance brought about by saline flow rate in the previous experiments (see Methods). The plots in Figure 3e compare the frequencies of PSCs in Control and s-GO treated samples against four different exposure time points. A progressive reduction in PSCs frequency was observed from 30 min to 6 h (from $5.8 \pm$ 1 to 3.6 \pm 0.8 Hz), and such changes were not detected in control (from 5.0 ± 1 to 5.3 ± 1 Hz). In Figure 3e, by linear regression analysis of the two time progressions (Control and s-GO) combined to multiple regression statistical analysis, we show that the zero slope hypothesis is accepted for Controls but not for s-GO, indicative of a significant progressive decrease in PSCs frequencies due to s-GO prolonged

In Vivo Intrahippocampal s-GO Delivery Reversibly Reduces Glutamatergic Synaptic Activity in Juvenile Rats with Minimal Tissue Reaction. To gain more insights into the synapse specificity, tissue reactivity and kinetics of s-GO in vivo, we injected in juvenile rats (P15) 1 μ L of s-GO (50 μ g/mL in saline solution; Figure 3f) in the dentate gyrus of the hippocampus and we patch-clamped single neurons to measure glutamatergic synaptic activity from ex vivo hippocampal slices isolated after 48 and 72 h after the brain surgeries. As control, we injected 1 μ L of saline solution in the same anatomical region. Figure 3f shows representative current tracings of the recorded electrical activity in acute slices

isolated from the contralateral (not subjected to the injection), the control saline- and s-GO-injected hemispheres, after 48 h from injection. After this time period hippocampal slices isolated from s-GO treated animals showed a clear and significant (** P < 0.01 two-way ANOVA) reduction in PSCs frequency $(2 \pm 0.5 \text{ Hz}, n = 7)$, when compared with slices from the contralateral untreated hemisphere (5 \pm 1 Hz, n = 9) or with saline treated ones $(4.3 \pm 1 \text{ Hz}, n = 8)$; see plot in Figure 3g). Remarkably, such effects were reversible: upon 72 h recovery post injections the reduction in synaptic PSCs frequency in s-GO treated slices is absent $(6 \pm 1.3 \text{ Hz}, n = 8)$ for contralateral slices; 5.7 ± 1.5 Hz, n = 7 for saline-injected slices; 6.2 ± 1.8 Hz, n = 7 for s-GO-injected slices; see plot in Figure 3g). In all treatments, the PSCs amplitude was not affected. When pharmacologically discriminating GABA, and AMPA receptor-mediated PSCs, we specifically detected after 48 h of s-GO a significant (* P < 0.05 two-way ANOVA) reduction in EPSCs frequency (2.9 \pm 0.8 Hz, n = 7), when compared with slices from the contralateral untreated hemisphere $(4.8 \pm 1 \text{ Hz}, n = 9)$ or with saline treated ones (5 ± 1.3) Hz, n = 8; see right plot in Figure 3h). GABA_A receptormediated PSCs were not affected by any treatment (from $3.5 \pm$ 1 to 3.9 \pm 0.8 Hz after saline-ejection, n = 8, and from 3 \pm 0.5 to 3.4 \pm 0.5 Hz after s-GO-ejection; n = 7). To prove the presence of s-GO and gain more insight regarding its fate within the hippocampus in vivo, we used bright-field microscopy with correlative Raman based mapping (Figure 4a). Forty-eight hours following intrahippocampal delivery (50 μ g/mL final concentration; 1 μ L injected volume), the presence of s-GO could be positively identified within the 20 μ m sections of injected hippocampi, specifically within the confines of the dentate gyrus. However, 72 h post injection, the material presence is shown to decrease. We also tested the hippocampi of rats that were injected with a saline control and a higher concentration of s-GO (1.3 mg/mL; 1 μ L), which served as negative and positive controls, respectively, to verify our data. The localization of s-GO by using QD-s-GO (50 μ g/ mL; 1 μ L) was performed next. Figure 4b shows the immunofluorescence labeling of slices isolated from the treated hippocampus where the area of injection is highlighted by the typical microglia reaction (Iba1 positive cells in green) due to the surgery per se. 41,42 QD-s-GO was typically localized in the area of injection after 24 h (red staining). Next we investigated whether s-GO injection was affecting the number of synapses in the injected brain area. We used bassoon marker for presynaptic terminals present in both glutamatergic and GABAergic synapses.²⁹ We quantified the colocalization of bassoon with neurons (labeled with β -tubulin III) and we did not detect any difference between saline and s-GO treated (48 h) animals in terms of bassoon volume at the injection site (saline 552.68 \pm 155.06 μ m³ and s-GO 570.40 \pm 115.74 μ m³; number of animals = 2 for each experimental group; Figure 4c,d). To investigate tissue reactivity, in particular neuroglia responses, to s-GO following 48 and 72 h, we performed immunohistochemistry experiments on treated animals to identify GFAP-positive astrocyte and Iba1-positive microglia (number of animals = 3 for each experimental group; Figure 5a,b). We measured astrocytes and microglia located 300 μ m apart to the injection site in the medial and lateral directions, at such a distance we detected only a low tissue response in all groups (Figure 5c,d). Conversely, at the injection site the tissue reactivity was higher, as expected, 41,42 yet comparable between saline and s-GO. To note, astrocyte recruitment was decreased in s-GO treated animals, particularly after 48 h; while microglia reactivity was similar in saline and s-GO groups after 48 h but it was significantly lower in s-GO treated animals after 72 h (Figure 5; ** P < 0.01; two-way ANOVA).

We report here the ability of s-GO nanosheets to interact selectively with glutamatergic synapses, affecting the efficacy of neurotransmission, in vitro and in vivo. In particular, in cultured hippocampal neurons, brief exposures to s-GO promote an initial high rate of glutamate quantal release, presumably by modifications at the presynaptic site, as indicated by the increase in frequency of spontaneous mEPSCs^{25,43,44} and by the paired-pulse experiments.^{33,35–38} We hypothesize that this initial high rate of release depletes presynaptic glutamate and in the continuous presence of s-GO inhibits glutamatergic transmission. Indeed, in the same preparation the decline in action potential-evoked monosynaptic EPSCs upon s-GO exposure supports the notion of a subsequent reduction in the probability of release following vesicle depletion⁴⁵ brought about by s-GO. In all tests, the mere pressure ejection of saline solution without s-GO, or GO of different dimensions, did not change spontaneous or evoked synaptic responses. Notably, GABAergic synapses were never affected. The biphasic effects of s-GO, characterized first by a transient increase in neurotransmitter release which upon a potential reduction in the vesicle-pool size is followed by a depression, hint at the ability of s-GO to engage the presynaptic exocytotic machinery, as also supported by the colocalization with presynaptic terminal markers. Neurotransmitter release at the presynaptic site in the CNS is controlled by specific proteins that function in large complexes, displaying multiple roles in synaptic vesicle recycling.⁴⁶ In addition to release-proteins, another potential target of s-GO is represented by intracellular Ca²⁺ levels, known to regulate evoked neurotransmitter release⁴⁵ and recently reported to be modulated by chronic exposure to s-GO.⁴⁷ Although we cannot exclude a role of presynaptic Ca²⁺ influx contributing to acute s-GO effects, the detected increase in spontaneous miniature current frequency, much less dependent on Ca2+ levels, 44,48 and the absence of modulation by s-GO of the GABAergic terminals, usually regulated by presynaptic Ca2+ dynamics, ⁴⁹ are suggestive of a Ca²⁺ independent mechanisms. The responses evoked by pressure ejected s-GO are reminiscent to those induced by hypertonic solutions, 50 however a simple osmotic mechanism is ruled out by the selectivity of the effects (restricted to glutamatergic terminals) observed in all conditions tested and by direct osmotic pressure measures (see Methods). The current data are in agreement with our previous report, where a long-term (days) exposure to s-GO selectively down regulated excitatory neurotransmission leaving inhibitory synapses unchanged. 10 We previously speculated that differences in GABAergic and glutamatergic synaptic cleft ultrastructure, in particular in the cleft size and organization,⁵¹ might explain why the latter terminals became ideal targets to s-GO interactions. To note, larger or smaller GO flakes did not modulate glutamatemediated synaptic transmission. In this framework, we propose a simply mechanistic interpretation of our current experiments: glutamatergic synapses, in virtue of their relatively larger size and less structured organization,⁵¹ allow penetration of s-GO flakes which remain trapped within the cleft and adhere to the plasma membrane at active release sites. GO nanosheets have been suggested to adhere to complex patches of cellular membranes, rather than specific ones. 52 Also in our experiments, the s-GO adhesion to the membrane may be supported by nonspecific interactions (as described in other cell types⁵²) accompanied by variable degrees of membrane deformations, a mechanism further supported by our previous results showing astrocyte vesicle shedding when exposed to s-GO. 10 A deformation of synaptic active zones would interfere with the exocytosis and neurotransmitter vesicle release by a mechanical mechanism reminiscent of, for example, stress-induced ones, 50 not necessarily implying an impairment of membrane integrity. In fact, we never observed any functional sign of membrane damage, and in addition the alterations in vesicle release were reversible. An alternative mechanism due to the physical properties of nanoparticles is related to their surface potential and able to tune neuronal excitability.⁵³ We previously documented that the s-GO surface potential, measured as zeta potential value, 10 is negative (-50 mV), thus the negative charge may favor the s-GO interactions with neuronal membrane influencing the excitability of neurons.⁵³ Although we cannot exclude this mechanism, the short- and long-term regulation and the selectivity for excitatory synapses are not explained by this interpretation. The interface between dispersed s-GO sheets and the cell membrane is currently subject to active investigation due to its potential in modulating cellular mechanosensing for diverse biomedical applications; nevertheless, the nature of such interactions is still elusive. 52,54 Synaptic vesicle recycling machinery represents a feasible therapeutic target, regardless of the direct involvement of presynaptic function in a pathological process. Even subtle alterations in (pre)-synaptic communication hold the potential to compensate for deficits without interfering with postsynaptic signaling. Presynaptically targeted drug development might be challenging due to the sophiticated molecular complexity of the release machinery. The ability of s-GO to specifically hook glutamatergic presynaptic nerve terminals is thus highly promising, however conventional 2D cultures may lack appropriate cell-extracellular matrix interactions, providing an artificially higher access of exogenous agents to synapses. s-GO specificity toward glutamatergic synapses may be virtually restricted to 2D biosystem models. This potential pitfall is excluded by our experiments on acute hippocampal slices, the neuroscientist gold standard to investigate synaptic functions in intact circuitries. The selective effect of s-GO on glutamatergic transmission is preserved in tissue slices, where excitatory EPSCs are reversibly affected by s-GO, with a short-term up-regulation of release, turned into a down regulation upon prolonged exposure. The ultimate potential of any s-GO sheets in the design of therapeutic strategies based on synaptic targeting resides in testing their efficacy in vivo. We demonstrated the delivery of s-GO in vivo by stereotactic injection and we have shown that such an administration of s-GO (but not the surgery per se) in the hippocampus of juvenile rats significantly and selectively sized down glutamatergic activity, in the absence of direct reduction in the number of synapses. We have also shown that local tissue responses to stereotactic injections 41,42 were not increased by the presence of s-GO in terms of patterns of microglia together with astrocyte aggregation at the injection site. These results are supportive, within the concentrations tested, of the in vivo biocompatibility of the s-GO dispersions. In general, GO is characterized by better biocompatibility when compared to other types of graphene (such as pristine graphene or reduced GO) and additional functionalization might even further reduce the risk of inflammation and subsequent tissue toxicity.⁵⁵ Interestingly, our results also suggest a possible anti-inflammatory effect by limiting the aggregation of astrocytes surrounding the stereotactic injection and lessening prolonged microglia reactivity. 41,42 This result, although preliminary, is in accordance with previous observations⁵⁶ and renders further investigation. Exploiting s-GO in presynaptic drug design development certainly requires additional studies, as well as to ascertain a more precise s-GO mechanism of action and clearance, because in our experiments due to diffusion perfusion flow rate in vitro, potential membrane recycling,⁵⁷ and microglial uptake,⁵⁸ we probably had only a "local" tissue clearance. Besides, most of the studies evaluating the clearance in vivo of GO suggested that GO is rapidly cleared but have been performed in nonmammalian organisms. 59,60 In our proof-of-concept in vivo study the coherence between the low detection of residual s-GO at 72 h and the reversibility of the synaptic silencing upon 72 h are supportive of a direct, mechanical interaction at the presynaptic plasma membrane.

Methods. Graphene Oxide Nanosheets Synthesis. GO was manufactured under endotoxin-free conditions through our modified Hummers' method as previously described. 10 Briefly in this procedure, 0.8 g of graphite flakes was added to 0.4 g of sodium nitrate (Sigma-Aldrich, U.K.). This was followed by the slow addition of 18.4 mL of sulfuric acid 99.999% (Sigma-Aldrich, U.K.). After a homogenized mixture was achieved through stirring, 2.4 g of potassium permanganate (Sigma-Aldrich, U.K.) was added and maintained for 30 min. Thereafter, 37 mL of water for injection (Fresenius Kabi, U.K.) was added. This resulted in an exothermic reaction. The temperature was strictly kept at 98 $^{\circ}\text{C}$ for 30 min. The mixture was next diluted with 112 mL of water for injection (Fresenius Kabi, U.K.). Twelve milliliters of 30% hydrogen peroxide (Sigma-Aldrich, U.K.) was then added to reduce the residual KMnO₄, MnO₂, and MnO₇ to soluble manganese sulfate salts. The resulting mixture purified by repeated centrifugation cycles at 9000 rpm for 20 min until an orange/brown gel-like layer of GO began to appear on at the pellet-supernatant interface which occurred at around pH 6-7. This layer was carefully extracted with warm water for injection (Fresenius Kabi, U.K.). This layer contained large GO sheets; the obtained material was diluted in water for injection to yield an aqueous suspension with a concentration of 2 mg/mL. A portion of this obtained material was then lyophilized, reconstituted in water for injection (Fresenius Kabi, U.K.), and then sonicated in a bath sonicator (VWR, 80W) for 5 min. The resulting dispersion was then centrifuged at 13 000 rpm for 5 min at room temperature (RT); the supernatant which contained the desired s-GO nanosheets was separated from the unwanted pellet. A thorough physicochemical characterization of us-GO and l-GO has already been reported. 18 Structural properties such as lateral dimension and thickness of the GO materials were then studied by AFM and TEM. Raman spectroscopy and ζ -potential measurements were used to define the materials surface properties. TGA was also performed to examine the functionalization degree of the s-GO sheets. Moreover, XPS was used to examine the composition of the GO sheets, C/O ratio, and the presence of the different functional groups.

s-GO Functionalization. QD have been prepared according to the literature by controlled decomposition of Ag and In salts. Briefly, 88.4 mg of $InCl_3$ (0.4 mmol, Sigma-Aldrich) and 17 mg of $AgNO_3$ (0.1 mmol, Sigma-Aldrich) were placed in a

100 mL round-bottom flask. Then, 190 μ L of oleic acid (0.6 mmol, Sigma-Aldrich), 720 µL of ododecylthiol (0.6 mmol, Sigma-Aldrich), and 8 mL of 1-octadecene (Sigma-Aldrich) were added under argon. The solution was heated at 60 °C for 15 min, at 90 °C for 15 min, and left at 110 °C stirring until no precipitate was visible (15-30 min). Then, 4 mL of solution S (9.6 mg 0.3 mmol, Sigma-Aldrich) was added and the mixture turned reddish. Finally, 5 mL of ZnCl₂ solution (70.5 mg of 0.5 mmol, Sigma-Aldrich) in oleylamine and 1-octadecene were added and the temperature was raised to 150 °C. After 15 min. the reaction was cooled with an ice bath. The QDs were purified by precipitation with ethanol, resuspended with cyclohexane, washed several times with ethanol/acetone and stored in CH2Cl2. For the water transfer reaction, 2 mL of solution oil QDs (2 mg/mL dispersion) was added to 1 mL of a cysteine (50 mg 0.4 mmol) basic solution in methanol. Immediately, the QD precipitated. After 20 min, 5 mL of distilled water was added and the QD passed throw the aqueous phase. Subsequently, the water-soluble QDs have been precipitated with acetone, washed several times with acetone/ethanol, and stored in distilled water. For GO conjugation, QDs were mixed with GO (1 mg/mL) at 1/10 mass ratio in distilled water. The mixture was left stirring for 3 days and then purified via dialysis against distilled water.

Preparation of Hippocampal Cultures and Acute Hippocampal Slices. Primary hippocampal cultures were prepared from neonatal rats at 2-3 postnatal days (P_2-P_3) as previously reported. All procedures were approved by the local veterinary authorities and performed in accordance with the Italian law (decree 26/14) and the UE guidelines (2007/526/CE and 2010/63/UE). The animal use was approved by the Italian Ministry of Health. All efforts were made to minimize suffering and to reduce the number of animals used. All chemicals were purchased by Sigma-Aldrich unless stated otherwise. Cultures were then used for experiments after 8-12 days in vitro. Hippocampal acute slices were obtained from P_7-P_8 rats and from juvenile P_{15} rats (n=18 animals) using a standard protocol. 63,64

Electrophysiological Recordings. In dissociated hippocampal cultures, single and paired whole-cell recordings were obtained with pipettes (5-7 M Ω) with the following intracellular saline solution (in mM): 120 K gluconate, 20 KCl, 10 HEPES, 10 EGTA, 2 MgCl₂, 2 Na₂ATP, pH 7.3; osmolarity 300 mOsm. The extracellular saline contained (in mM) 150 NaCl, 4 KCl, 1 MgCl₂, 2 CaCl₂, 1 MgCl₂, 10 HEPES, 10 glucose, pH 7.4. Data were recorded by Multiclamp 700B patch amplifier (Axon CNS, Molecular Devices) digitized at 10 kHz by pClamp 10.2 software (Molecular Devices LLC, USA). Basal PSCs were recorded at -56 mV holding potential (liquid junction potential of 14 mV was not corrected for). mPSCs were recorded in the presence of TTX (1 µM) to block fast voltage-dependent sodium channels. In voltage-clamp recordings, PSCs and mPSCs were detected by the use of the AxoGraph X (Axograph Scientific) event detection program and by the Clampfit 10 software (pClamp suite, Axon Instruments) as previously reported.⁶⁴ On average, ≥500 PSCs were analyzed from each cell and from the average of these events we measured the peak amplitude and the decay time constant (expressed as τ) by fitting a monoexponential function. In paired recordings, the presynaptic neuron in current clamp mode was held at −70 mV (by ≤0.02 nA negative current injection), and action potentials were evoked by delivering short (4 ms) square current pulses (1 nA). Monosynaptic connections were identified by their short delay (<2 ms).⁶⁴ To characterize the short-term dynamics of synaptic contacts, we delivered to pairs of connected neurons paired pulse stimulations at 20 Hz (1 pair every 20 s; 10 times that were pooled together and averaged). For acute hippocampal slices, a patch-clamp amplifier (Multiclamp 700B, Axon Instruments, Sunnyvale, CA, U.S.A.) allowed recordings from CA1 pyramidal neurons, identified by visual inspection at an upright microscope (Eclipse FN1; Nikon, Japan) equipped with differential interference contrast optics and digital videocamera (Nikon, Japan). All recorded events were analyzed offline with the AxoGraph 1.4.4 (Axon Instrument) event detection software (Axon CNS, Molecular Devices). s-GO was acutely delivered, 10 both in dissociated cells and in acute hippocampal slices, by an injection of pressurized air (500 ms duration, 8 PSI; by a Picospritzer PDES-02DX; NPI electronic GmbH, Germany). Once the neurons were patch-clamped, a second pipette identical to that used for patch-clamp recording was positioned at a distance of 200 μ m (under microscopy control) from the recorded cell. The pipet was filled with standard saline solution (control; osmolarity 300 mosmol L⁻¹) or with s-GO,l-GO and us-GO (100 μ g/mL in Krebs solution; osmolarity 300 mosmol L⁻¹). The concentration of GO reaching the cell was at least 10% of that contained in the pipet, considering 1 mL of extracellular solution in the recording chamber. Baseline PSCs were sampled before (10 min) and after (10 min) the local ejection. Analyses were performed between 4 and 8 min after the local ejection, sampling 2 min of recordings.

Confocal Microscopy in Hippocampal Cultures. Cultured hippocampal neurons (3 cultures; 6-8 DIV) were incubated for 30 min with s-GO (20 μ g/mL). Cultures were then fixed by 4% formaldehyde (prepared from fresh paraformaldehyde; Sigma) in PBS at RT and blocked and permeabilized in 5% fetal bovine serum (FBS), 0.3% Triton-X 100 in PBS for 30 min at RT. Samples were incubated with primary antibodies (mouse monoclonal antibassoon, 1:400 dilution; rabbit anti- β tubulin III, 1:500 dilution) diluted in PBS with 5% FBS for 45 min. Cultures were finally incubated with secondary antibodies (Alexa 488 goat antimouse, Invitrogen, 1:500 dilution; Alexa 594 goat antirabbit, Invitrogen, 1:500 dilution) and DAPI (Invitrogen, dilution 1:200) to stain the nuclei for 45 min at RT and finally mounted on 1 mm thick glass coverslips using the Fluoromount mounting medium (Sigma-Aldrich). To visualize s-GO, localization was used in the reflection mode of confocal microscopy.⁵⁸ Images were acquired using a Nikon C2 Confocal, equipped with Ar/Kr, He/Ne, and UV lasers. Images were acquired with a 60× (1.4 NA) oil-objective (using oil mounting medium, 1.515 refractive index). Confocal sections were acquired every 0.4 μ m.

Surgery and s-GO Injection. Four experimental groups were used: standard saline solution (control) and s-GO 50 μ g/mL injection, analyzed at 48 and 72 h. Surgery was performed in P₁₅ Wistar rats anesthetized with ketamine (60 mg/kg i.p.) and xylazine (10 mg/kg i.p.). All animal procedures were conducted in accordance with the National Institutes of Health, international and institutional standards for the care and use of animals in research, and after consulting with a veterinarian. All experiments were performed in accordance with the EU guidelines (2010/63/UE) and Italian law (decree 26/14) and were approved by the local authority veterinary service. All efforts were made to minimize animal suffering and

to reduce the number of animals used. The Italian Ministry of Health, in agreement with the EU Recommendation 2007/526/CE, approved animal use (authorization no. 1135/2015-PR). Animals were fixed in a stereotaxic device (World Precision Instruments, WPI). An incision was made on the top of the head in order to expose the skull and identify bregma and lambda coordinates. The injection of 1 μ L of saline or s-GO solution (10 steps of 0.1 μ L every minute) was performed with a Hamilton syringe (26s gauge; Hamilton). The following coordinates were used to reach the left dentate gyrus: AP, -3.0; ML, -3.0; DV, -3.3; relative to bregma. At the end of the last step, the syringe was left in situ for extra 5 min to optimize the solution permeation. The incision was sutured and animals were constantly monitored and left undisturbed until electrophysiological or histological experiments.

Histology Procedures. After 48 or 72 h, animals were anesthetized and sacrificed by intracardiac perfusion with 0.1 M PBS followed by 4% formaldehyde (prepared from fresh paraformaldehyde; Sigma) in PBS. Brains were promptly removed, postfixed in the same fixative solution for 24 h at 4 °C and cryoprotected in 30% sucrose in PBS at 4 °C for 24-48 h. Finally, brains were embedded in optimal cutting temperature (OCT) compound (Tissue-Tek), frozen at -20 $^{\circ}$ C, and sagittal sections (25 μ m) were obtained using a cryostat (Microm HM 550, Thermo Fisher Scientific) and processed for immunohistochemistry. Tissue-Tek was removed by PBS washing and tissue sections were protein-blocked in 3% BSA, 3% FBS, and 0.3% Triton X-100 in PBS for 45 min at RT. Sections were then incubated overnight at 4 °C with primary antibodies (mouse anti-GFAP, Sigma-Aldrich, 1:400; rabbit anti-Iba1, Wako, 1:500; rabbit anti-β-tubulin III, Sigma-Aldrich, 1:500; mouse anti-Bassoon, Abcam, 1:400) in 5% FBS in PBS. After washing in PBS, sections were incubated in secondary antibodies (goat antirabbit AlexaFluor 594, Thermo Fisher Scientific, 1:400; goat antimouse AlexaFluor 488, Thermo Fisher Scientific, 1:400) in 5% FBS in PBS for 2-4h at RT. Nuclei were labeled with DAPI (Thermo Fisher Scientific, 1:500) in PBS for 20-30 min at RT. Upon final washing (PBS and water), tissue sections were mounted on glass coverslips using Vectashield mounting medium (Vector Laboratories).

Image Acquisition and Analysis. We measured the brain tissue reaction by markers for reactive astrocytes and microglia (GFAP and Iba1, respectively). Fluorescence images were acquired using a Leica DM6000 upright microscope with a 10× dry objective. Identical binning, gains, and exposure times were used for all images of the same marker. Image analysis was performed using Fiji software. For both GFAP and Iba1 intensity measurements, a single region of interest (ROI, 1000 \times 500 μ m²) was selected at the injection site (left dentate gyrus). The background intensity threshold was defined for each section using the labeling intensity measured in the contralateral hemisphere in the same anatomical region (right dentate gyrus). The area within each ROI with intensity above the background threshold was calculated, normalized to the contralateral hemisphere and used for statistics. The ROI for all sections were averaged for each experimental group. We performed this analysis also at 300 μ m medial and lateral to the injection site. We visualized s-GO by linking QD. Fluorescence images were acquired using a Leica DM6000 upright microscope with a 10× dry objective. We further analyzed the amount of synaptic contacts by a specific marker for synapses (bassoon) in two experimental groups: saline and sGO injection at 48 h. Confocal images were acquired using a confocal microscope (Nikon C1) with a 60× oil objective (N.A. 1.4, oil mounting medium refractive index 1.515). Z-stacks were acquired every 350 nm for a total thickness of 7 μ m. Identical binning, gain, and exposure time was used for all images. Nine ROIs (70 × 70 μ m²) for each section were randomly selected at the injection site (left dentate gyrus). Offline analysis was performed using Volocity software (Volocity 3D image analysis software, PerkinElmer, U.S.A.). For each ROI, we used the Z-stack to quantify bassoon signal as 3D objects. From the resulting values, we calculated the volume of only bassoon objects colocalized with the β -tubulin III labeling in order to identify genuine synapses at neuronal level. The ROI for all sections were pooled together and averaged for each experimental group.

Raman Mapping of Brain Sections. Raman mapping of sectioned brain samples was completed using a DXRi Raman Mapping system (Thermo Scientific, U.S.A.) using the following conditions: $\lambda = 633$ nm, 1 mW, pixel size = 1.6 μ m and frequency = 25 Hz. The maps were generated according to the composite spectra's percentage similarity to a correlation GO spectral reference as shown.

Statistical Analysis. All values from samples subjected to the same experimental protocols were pooled together and results are presented as mean \pm S.D., if not otherwise indicated; n = number of neurons, if not otherwise indicated. Statistically significant difference between two data sets was assessed by Student's t test for parametric data. Differences between the logarithmic values of the analyzed variables were assessed using two-way ANOVA and multiple comparisons were adjusted by Bonferroni or Holm-Sidak correction. Statistical significance was determined at P < 0.05, unless otherwise indicated. Significance was graphically indicated as follows: *P < 0.05, **P < 0.01, ***P < 0.001.

ASSOCIATED CONTENT

S Supporting Information

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Additional figures (PDF)

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Author Contributions

R.R. set up the experiments, performed and designed electrophysiology and data analysis, and contributed to the writing of the paper; M.M. performed the surgery, histology, and confocal analysis; L.N. performed Raman spectroscopy and S.V. performed confocal microscopy, and both contributed to the writing of the results and methods; G.R and A.B. performed the synthesis and characterization of QD s-GO and

contributed to the microscopy; M.P. contributed to the experimental design; K.K. contributed with synthesis and characterization of GO sheets and contributed to the experimental design and paper writing; L.B. conceived the idea, the experimental design, and wrote the paper.

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Notes

The authors declare no competing financial interest.

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