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Graphene-based interfaces do not alter target nerve cells

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ABSTRACT

Neural-interfaces rely on the ability of electrodes to transduce stimuli into electrical patterns delivered to the brain. In addition to sensitivity to the stimuli, stability in the operating conditions and efficient charge transfer to neurons, the electrodes should not alter the physiological properties of the target tissue. Graphene is emerging as a promising material for neuro-interfacing applications, given its outstanding physical-chemical properties. Here we use graphene-based substrates (GBSs) to interface neuronal growth. We test our GBSs on brain cell cultures by

measuring functional and synaptic integrity of the emerging neuronal networks. We show that GBSs are permissive interfaces, even when uncoated by cell adhesion layers, retaining unaltered neuronal signalling properties, thus being suitable for carbon-based neural prosthetic devices.

Coupling (nano)materials to organic tissues is crucial for developing prosthetic applications, where the interfacing surfaces should provide minimal undesired disturbance to the target tissue.¹ Ultimately, the (nano)material of choice has to be biocompatible,^{1,2} promoting cellular growth and adhesion with minimal cytotoxicity or dis-regulation of, *e.g.*, cellular activity.²

In neurology, relevant examples in the area of prosthetic devices are deep-brain intracranial electrodes,³ used to control motor disorders, or brain interfaces, such as those used to recover sensory functions⁴ or to control robotic arms for amputated patients.⁵ In all these cases, the inorganic material constituting the interfaced electrode has to preserve unaltered tissue functionality to avoid uncontrolled side effects.⁵ A charge transfer taking place from electrodes to neurons, flexibility and ease of molding into complex shapes are also key requirements.² Current approaches involve the use of tungsten microwire electrodes,⁶ or silicon based electrode arrays.⁶ The clinical relevance of these approaches has been demonstrated.⁴ However, drawbacks are still limiting their long-term performance when implanted.¹ The most common is the formation of an insulating layer around the electrodes, the so-called “glial scar”,⁷ as a consequence of insertion-related brain trauma and long-term inflammation. This can halve the level of the desired signal (electrical stimulus delivered/recorded by the electrode) respect to the level of background noise, namely the signal-to-noise ratio,^{1,7} leading to electrode failure.⁶ Another failure mechanism stems from the electrodes’ stiffness, usually larger than the surrounding tissue, resulting in tissue detachment.¹ Thus, there is a need to develop flexible electrodes, consisting of biocompatible, cell-adhesion-promoting and conductive materials.

Carbon-based nanomaterials, such as carbon nanotubes (CNTs), have been extensively used as neural electrodes.^{1, 8-15} Interfacing neurons with CNTs was shown to increase neuronal activity, at

least *in vitro*, in various experimental models.^{8, 16-20} This can be exploited in neural prostheses/devices to bypass non-functional neuronal tissue (*e.g.* glial scar following a lesion).²¹ Coating extracellular electrodes with CNTs enhances both recording and electrical stimulation of neurons, both in-culture and *in vivo*, in rats and monkeys, by decreasing the electrode impedance and increasing charge transfer.⁹ CNTs can also alter the neuronal behaviour in terms of spontaneous synaptic activity¹⁶ and action potential firing frequencies.¹⁶ Neuroprosthetics applications require low neuronal tissue perturbation:¹ implanted electrodes must excite the neuronal cells, without depressing (or boosting) the surrounding neuronal network.¹

Due to its excellent electrical properties,²² graphene is promising for the development of neural interfaces.^{23,24} A number of studies to date have addressed the issue of graphene toxicity.^{24,25} However, less attention was paid to graphene bio-interfaces, in particular those exploiting non-chemically modified graphene,^{26,27} and even fewer reports have addressed the issue of biocompatibility with neuronal cells.^{28,30} Polylysine-covered graphene was shown²⁸ to be a neuro-favourable laminar material, sustaining viability and improving the growth of specialised neuronal compartments (the neuritis) in dissociated hippocampal cultures.²⁸ Laminin-coated graphene favours the differentiation of neural stem cells into neurons.³¹ However, peptide-based (*e.g.* polylysine or polyornithine) coatings might increase the electrical resistance of the neuron/interface electrical contacts, thus affecting the charge transfer properties.^{28, 32-3} The direct contact and exposure of neurons to GBSs is crucial to promote tight adhesion between cell membranes and interfacing electrodes, a key requirement to detect small (tens of μV) signals during extracellular recordings, and to reduce voltage drops during tissue stimulation, thus improving charge transfer.^{28, 32-34} Ref. 28 reported the biocompatibility of uncoated graphene surfaces with neuronal cells in terms of neuronal survival and morphology. Yet, to the best of our knowledge, thus far no study addressed how uncoated graphene may impact the neuronal electrophysiological behaviour.

Micromechanical exfoliation can be used to produce graphene flakes with outstanding structural and electronic properties.³⁵⁻³⁹ However, its limited yield makes it impractical for large-scale

applications.³⁶ Graphene films can be produced by carbon segregation from metal substrates^{40,41} or SiC^{42,43} or by chemical vapour deposition [44-45] followed by transfer (wet³⁶ or dry⁴⁵) to a target substrate.³⁶ However, such processes require high temperatures (>1000 °C)^{36, 45-47}, costly substrates, besides the additional transfer.³⁶ Solution processing is emerging as a most promising technique to produce single- (SLG) and few-layer (FLG) graphene flakes on large scale,³⁶ both starting from oxidized⁴⁸⁻⁵¹ and pristine graphite.^{36, 52-58} Graphene oxide (GO), produced by exfoliation of graphite oxide, can be mass-produced at room temperature.^{48,49} However, it is insulating,^{49,50} with defects^{49,50} and gap states,^{50, 51} and may not offer the optimal charge transfer between substrate and neurons.³⁶ Liquid phase exfoliation (LPE) of graphite⁵² can be performed without the potentially hazardous chemical treatments involved in GO production,⁴⁸⁻⁵¹ being at the same time scalable, room temperature and high yield.³⁶ LPE dispersions can also be easily deposited on target substrates, by drop casting,⁵³ filtration⁵² or printing.⁵⁴ Another approach to graphite exfoliation is ball milling (BM) with the help of melamine, which forms large H-bond domains and intercalates graphite^{55, 56} and, unlike LPE, can be performed in solid.⁵⁶

Here, we use LPE and BM of graphite to fabricate GBSs. Electrophysiological measurements show the bio-compatibility *in vitro* of both samples with dissociated hippocampal neuronal cultures. Our GBSs allow neuronal adhesion and growth when mammalian, differentiated, post-mitotic neurons are explanted and cultured on them. We also investigate the impact on neuronal, synaptic and network electrophysiological properties, to address the ability of our GBSs to interface and transform neuronal signalling.⁵⁹ We find that our GBSs favour nerve-cell adhesion and survival without altering the cell differentiation, biophysics passive properties, synaptogenesis, spontaneous synaptic activity and plasticity, when compared to control growth-substrates. Our GBSs also retain neuronal signalling properties, thus paving the way to the development of carbon-based neural interfaces able to preserve the neuronal activity.

RESULTS AND DISCUSSION

The GBSs are produced following two different protocols, LPE and BM, in order to unveil possible effects of materials production, processing, deposition and structure, on neuronal activity.

The LPE protocol is as follows: 120mg of graphite flakes (Sigma Aldrich) are dispersed in 10 ml deionised water (DIW) with 90 mg of sodium deoxycholate (SDC), then placed in an ultra-sonic bath for 9 hours and subsequently ultracentrifuged exploiting sedimentation-based separation (SBS)³⁶ using a TH-641 swinging bucket rotor in a Sorvall WX-100 ultracentrifuge at 10 krpm ($\sim 17,000g$) for 1 hour. After ultracentrifugation, the top 70% of the dispersion is extracted by pipetting and deposited on glass coverslips by vacuum filtration and film transfer. The dispersion is characterized by optical absorption spectroscopy (OAS) and Raman spectroscopy. OAS of the dispersions, diluted to 10% to avoid scattering losses at higher concentrations, is acquired in the range 200–1300 nm with a Perkin-Elmer Lambda 950 spectrophotometer. The concentration of graphitic flakes is determined from the optical absorption coefficient at 660 nm, using $A = \alpha lc$ where l [m] is the light path length, c [gL^{-1}] is the concentration of dispersed graphitic material, and α [$Lg^{-1}m^{-1}$] is the absorption coefficient, with $\alpha \sim 1390 Lg^{-1}m^{-1}$ at 660 nm.^{52, 54} For Raman spectroscopy the dispersions are drop-cast onto a Si wafer with 300 nm thermally grown SiO_2 (LDB Technologies Ltd.), dried on a hot plate and rinsed in a solution of DIW/ethanol (50:50). Raman measurements on both the graphene dispersions and GBSs are collected using a Renishaw InVia spectrometer at 457, 514.5, and 633 nm with a 100x objective and an incident power < 1 mW. The G peak dispersion is defined as $Disp(G) = \Delta Pos(G) / \Delta \lambda_L$, where $Pos(G)$ is the position of the G peak and λ_L is the laser excitation wavelength and Δ indicates the rate of change of $Pos(G)$ as a function of varying λ_L .

LPE-GBSs dispersions are then vacuum filtered *via* 100 nm pore-size filters (Millipore nitrocellulose filter membranes). This blocks flakes, while allowing water to pass through. To remove the residual surfactant, the GBSs deposited on the filters are rinsed by vacuum filtration of 20 mL DIW. The film transfer on glass coverslips is done by applying pressure and heat (~ 90 °C, to

improve adhesion) overnight (~10 hours), followed by dissolution of the filter in acetone. The GBSs are then rinsed in isopropyl alcohol and in DIW baths, and finally dried in oven (~90 °C) for 1 hour. The BM-GBSs are produced by exfoliation of graphite (from Bay Carbon) with melamine,⁵⁵ and then dispersed in dimethylformamide (DMF), followed by drop casting on glass coverslips placed on a hotplate at 100 °C. The substrates are thermally treated for 20 minutes in an oven at 350 °C under nitrogen atmosphere to remove traces of solvent or impurities.

Fig. 1a plots the absorbance of the LPE dispersion. The peak at ~266 nm is a signature of the van Hove singularity in the graphene density of states.⁶⁰ Fig 1b plots a typical Raman spectrum (black curve) measured at 514.5nm, of representative flakes of the LPE dispersion on Si/SiO₂. The G peak corresponds to the E_{2g} phonon at the Brillouin zone center.⁶⁰ The D peak is due to the breathing modes of sp² rings and requires a defect for its activation by double resonance (DR).⁶²⁻⁶⁴ The 2D peak is the second order of the D peak.⁶³ This is a single peak in SLG, whereas it splits in FLG, reflecting the evolution of the band structure.⁶² The 2D peak is always seen, even when no D peak is present, since no defects are required for the activation of two phonons with the same momentum, one backscattered from the other.⁶¹ DR can also happen as intra-valley process, *i.e.* connecting two points belonging to the same cone around K or K'.⁶² This process gives rise to the D' peak. The 2D' is the second order of the D'. Statistical analysis (black dashes plots), based on 30 measurements for each excitation wavelength (457, 514.5, 633nm), gives an average position of the 2D peak, Pos(2D), ~2698cm⁻¹ (Fig. 1c). FWHM(2D) varies from 45 to 72 cm⁻¹ (Fig. 1d) with a peak at ~63cm⁻¹. Pos(G), Fig. 1e, and FWHM(G), Fig. 1f, are 1583 and 24 cm⁻¹. I(2D)/I(G) ranges from 0.4 to 1.1 (Fig. 1g). This is consistent with the samples being a combination of SLG and FLG flakes. The Raman spectra show significant D and D' peaks intensity, with I(D)/I(G) ranging from 0.5 to 2.5 (Fig. 1h). This is attributed to the edges of our sub-micrometer flakes⁶⁵ rather than to the presence of a large amount of structural defects within the flakes. This observation is supported by the low Disp(G) <0.05 cm⁻¹/nm, much lower than what expected for disordered carbon.^{63, 64} Combining I(D)/I(G) with Disp(G) allows us to discriminate between disorder localized at the

edges and disorder in the bulk. In the latter case, a higher $I(D)/I(G)$ would correspond to higher $\text{Disp}(G)$ (see Fig. 1i). The lack of a clear correlation between $I(D)/I(G)$ and $\text{FWHM}(G)$ (see Fig. 1i) is an indication that the major contribution to the D peak comes from the sample edges.

Similarly to LPE, Fig 1b plots a typical Raman spectrum (red curve) measured at 514.5nm of representative BM flakes on Si/SiO₂. A statistical analysis based on 30 measurements for each excitation wavelength (457, 514.5, 633nm) gives an average $\text{Pos}(2D) \sim 2708 \text{ cm}^{-1}$ (Fig. 1c) and $\text{FWHM}(2D)$ peaked at 75 cm^{-1} (Fig. 1d). $\text{Pos}(G)$, Fig. 1e, and $\text{FWHM}(G)$, Fig. 1f, are 1583 and 22 cm^{-1} . $I(2D)/I(G)$ peaks at ~ 0.5 (Fig. 1g). The Raman spectra show lower $I(D)/I(G)$ and $I(D')/I(G)$ than the LPE samples, with $I(D)/I(G)$ having a bimodal distribution peaked at 0.2 and 1.0 (Fig. 1h). However, the correlation between $I(D)/I(G)$ and $\text{FWHM}(G)$, Fig 1i, indicates that defects are present inside the samples, and that the D peak does not only come from the edges. The comparison between the Raman data on LPE and BM samples indicates that the latter mostly comprise defective graphite flakes a few layer thick, while the former mostly consist of SLG, or FLG, but electronically decoupled, and with fewer defects.

Before cell deposition, both LPE and BM-GBSs are thermally annealed for 20 minutes in oven at $350 \text{ }^\circ\text{C}$ in nitrogen in order to remove residual solvent. Electrical measurements show that for LPE-GBSs, the sheet resistance, R_s , is $\sim 3.4 \text{ k}\Omega/\square$, which corresponds, using $R_s = 1/\sigma \cdot t$ [with $t \sim 150 \text{ nm}$], to an average value of conductivity (σ) $2600 \pm 400 \text{ S/m}$, comparable with other films produced by LPE graphene³⁶ and reduced GO ($\sigma \sim 10^3 \text{ S/m}$).⁶⁶ The BM-GBSs have a lower $\sigma = 1010.1 \pm 90.0 \text{ S/m}$ with respect to the LPE-GBSs.

Fig. 2a compares 514.5nm representative Raman spectra for LPE and BM dispersions with the spectra of the resulting LPE-GBS and BM-GBS after the annealing process. Figs. 2b-e compare $\text{Pos}(2D)$ and $\text{FWHM}(2D)$ distributions. The LPE-GBS has a narrower distribution of both $\text{Pos}(2D)$ (peak at $\sim 2698 \text{ cm}^{-1}$) and $\text{FWHM}(2D)$ (peak at $\sim 72 \text{ cm}^{-1}$), with respect to the LPE dispersion. However, the 2D peak still shows a Lorentzian lineshape distinctly different from that of graphite. The BM-GBS has a similar distribution of both $\text{Pos}(2D)$ (ranging from ~ 2698 to 2718 cm^{-1}) and

FWHM(2D) (peak at $\sim 85\text{ cm}^{-1}$), to those of the BM dispersion and the 2D peak lineshape indicates the predominance of thick flakes.

AFM measurements are also used to characterize the GBSs. They are carried out with a Bruker Dimension Icon equipped with NanoScope V Controller in ScanAsyst PeakForce Tapping Mode using ScanAsyst-Air Silicon Nitride probes with a nominal tip radius of 2 nm. Topographic images are taken on surface areas of $100\ \mu\text{m}^2$ to measure roughness, while $80\ \mu\text{m}$ profiles across the film edges are used to estimate the thickness.⁶⁷ Fig. 3a shows an average root means square (RMS) roughness of $14.1\pm 2.9\text{ nm}$ and roughness average (Ra) of $9.0\pm 1.4\text{ nm}$, with an average film thickness (t) of $131\pm 44\text{ nm}$. The BM-GBS shows a sub-micrometric morphology, with the lateral dimensions of the flakes $< 200\text{ nm}$ (Fig. 3b).

Following the structural characterization, neuronal cultures are placed on the GBSs and electrophysiology studies are performed as detailed in Methods. Both peptide-free and polyornithine-covered glass coverslips are used as control substrates. We design the first set of experiments (reported in details in the Supplementary Information, S.I.) to separately compare the ability of the different GBS substrates, prepared from LPE and BM, to interface and allow neuronal network formation *in vitro*. Thus, we preliminarily match (S.I. and Fig. S1 and S2) sample cultures grown on the LPE or on the BM-GBSs with their respective control substrates. To examine neuronal cells when interfaced to the various substrates, we compare their membrane passive properties, *i.e.* the input resistance and cell capacitance (Fig. S1), which are also indicative of neuronal health conditions.⁵⁹ In individual data-set neurons grown on LPE or on BM are measured against their control sister cultures (*i.e.* within the same culture series) and these results show that both tested GBSs conditions allow neuronal growth without apparent differences with controls in the measured parameters (Fig. S1 and S2). The detected membrane passive properties are in agreement with previous control assessments for hippocampal neurons in culture.^{8, 17, 18} Thus, data on both GBSs and on pure glass or polyornithine-covered glass are pooled and collectively named GBSs or control, respectively.

Neurons are investigated by electron microscopy and immunofluorescence. They show normal morphology characterized by well-defined round soma and extended neurite arborisation (Fig. 3c) and a cell density similar to control substrates. The density of neuronal cells, quantified by staining neuronal nuclei with the specific marker NeuN, is 91 ± 13 neurons/mm² on the control substrate and 104 ± 11 cells/mm² on GBSs (immunofluorescence shown for control (Fig.3d) and LPE-GBS (Fig.3e)). Fig. 3f plots the neuronal density for GBSs pooled data (n=12 fields from 4 cultures). Similarly, the growth of glial cells, identified by their marker glial fibrillary acidic protein, GFAP, is unaffected by both GBSs. The glial cells density is 84 ± 5 cells/mm² on the control substrate and 100 ± 8 cells/mm² on GBSs (immunofluorescence shown for control (Fig. 3g) and LPE-GBS (Fig.3h)). Fig. 3i shows the glial cell density in the GBSs culture condition pooled data (n=12 fields from 4 cultures).

It is important to test whether graphene affects the network synaptic behaviour, since this is predictive of information processing,¹⁷ in view of developing biocompatible GBSs able to fully preserve neuronal functionality. To this aim, we analyze both spontaneous network activity and neuronal synaptic connectivity in cultures developed on GBSs or on control substrates. Fig. 4 (a,b) plots typical heterogeneous spontaneous post-synaptic currents (PSCs) recorded from a voltage-clamped control neuron (Fig 4a) and from neurons grown onto the GBSs surface (Fig 4b). Graphene interfacing does not affect the frequency of spontaneous PSC (4.0 ± 0.9 Hz, n=25 and 3.6 ± 0.8 Hz, n=21, for control and GBSs cultures, respectively; Fig. 4c) and their amplitude (51 ± 7 pA for controls and 54 ± 6 pA for GF, respectively; Fig. 4d). The network sizes of graphene-interfaced and control cultures are similar, as indicated by the comparable neuronal density, consequently they contribute equally to PSC frequency. GBS interfaces do not alter the spontaneous PSC frequency. This is further supported by the results shown in Fig. S2 where we measure PSCs in sampled cultures individually (LPE or BM). The observation that GBSs are inert growth substrates to synaptic networks is also strengthened by an additional set of experiments where we interface neurons to multi-wall carbon nanotubes (MWNTs). In the same cultures series we compare the

frequency of PSCs amongst MWNTs, GBSs and control growth substrates (S.I. and Fig. S3). We measure a selective increase in activity only in MWNT neurons (Fig. S3), confirming the reported^{8, 16-20} ability of MWNTs to boost synaptic activity. This differs from what reported in Ref. [30] where graphene induced enhancement of electrical signalling in neural network. However, we believe that the results in Ref. 30 are, at least in part, biased by the presence of different network sizes in the two culturing conditions.

We then indirectly measure the impact of GBSs on the *in vitro* formation of functional contacts (*i.e.*, synaptogenesis) by simultaneously patch clamping randomly selected neuron pairs,¹⁸ see Fig. 5a. In each cell pair, an action potential is elicited in the presynaptic neuron, and the presence of a monosynaptic connection between the two neurons is assessed by the PSC response in the postsynaptic neuron. The probability of finding mono-synaptically coupled neurons pairs (expressed as %) is a measure of functional synaptic connections formed in the *in vitro* network.¹⁸ We observe a % pairing on GBSs similar to that found on control substrates ($29 \pm 10\%$ and $39 \pm 11\%$, $n=5$ culture series; total $n=30$ pairs on GBSs and $n=28$ pairs on control substrate), thus indicating that GBSs interfacing does not alter synaptogenesis, see Fig. 5b. No differences in the amplitude of induced presynaptic action potentials are found between neurons in control and GBSs (91 ± 9 mV $n=10$ cells in control and 93 ± 4 mV $n=11$ cells on GBS, $P=0.7$).

We then investigate short-term synaptic plasticity, by eliciting a pair of action potentials in the presynaptic neuron (at 20 Hz), measuring the amplitude of the evoked PSCs in the postsynaptic neuron (Fig. 5c,d), and evaluating the ratio between the amplitudes of the second and first PSCs in the pair (paired-pulse ratio, Fig.5e). We find a moderately depressing response (paired-pulse ratio 0.89 ± 0.07 , $n=9$ pairs) in the control, very similar to GBSs (paired-pulse ratio 0.86 ± 0.10 , $n=7$ pairs; amplitude of the first PSC 36 ± 6 pA, $n=10$ and 44 ± 9 , $n=8$ for control and GBS substrates, respectively).

The observation that GBSs support neuronal functional development in the absence of any perturbation of neuronal network synaptic performance is intriguing. Other electrical conductive

carbon-based nanomaterials, such as CNTs, are known^{16, 19} for their ability to reshape spontaneous network activity, synaptogenesis, and for their short-term synaptic plasticity and single-cell electrogenic properties^{16, 19} in the same *in vitro* models. The precise mechanisms determining the different impact on neuronal activity of these two carbon based substrates requires further investigation. Our data using CNTs control surfaces strengthens the hypothesis that the tight and intimate interactions amongst CNTs and neurons, combined with the conductivity,¹⁷ are responsible for their ability to interfere with synaptic networks.^{18, 19} The lower roughness of GBSs (~14 or ~9 nm in GBSs against ~30 nm in MWNT¹⁸), might explain their inert nature in spite of their high electrical conductivity.

CONCLUSIONS

Our data show that GBSs, produced by liquid phase exfoliation or ball milling of graphite, are inert neuron-interfacing materials, able to preserve the basal physiological level of neuronal activity. The ability to interface neuronal circuit regrowth without altering cell and synapse behaviour may enable the fabrication of graphene-based devices for medical applications,²⁵ such as biosensors²⁵ and neuroprosthetics,²³ whereby GBSs at the tissue interface result in efficient interaction with cells. In the future design of electrically-functional implants, the use of novel materials characterized by the ability to integrate with tissue and, at the same time, flexible and not affecting excitable tissue behaviour, is highly relevant. Our data indicate that both GBSs are promising for next generation bio-electronic systems, to be used as brain-interfaces. In this framework, it is important to note the uncommon ability of our GBSs to support neuronal development (in terms of neuronal passive properties, spontaneous synaptic activity, synaptogenesis, and short-term synaptic plasticity) without pre-coating with adhesion-promoting peptides (*e.g.* polylysine or polyornithine). Previous works demonstrated the full biocompatibility of peptide-coated chemical vapour deposited graphene interfaces with hippocampal neurons (polylysine-coated graphene²⁸) or neural stem cells (laminin-coated graphene^{30, 31}). However, peptide coating might weaken neuron/interface electrical contacts and electrical signal transmission, resulting in non-optimal charge transfer.^{28, 32-34}

METHODS

Materials preparationMulti wall carbon nanotubes preparation

Multi wall carbon nanotubes-coated substrates are prepared similarly to our previous Refs. 17-19 (optimized MWNT dispersion concentration 0.1 mg/mL; final MWNT film density 7×10^{-5} mg/mm²). Briefly, MWNTs (Nanostructured & Amorphous Materials, Inc., Stock#: 1237YJS, outer diameter 20÷30 nm), used as received, are functionalized using 1,3-dipolar cycloaddition. Ethyl acetate solution of functionalized MWNTs (0.1 mg/mL) is sprayed on glass coverslips placed on a hot plate at 200 °C, then the substrates are heated at 350 °C under nitrogen atmosphere to restore the pristine structure of MWNTs.

Electrical Characterization

A Jandel station with 4-Probe head, 100 µm titanium tips arranged in a straight line 1mm apart, combined with a Keithley 2100 digital multimeter is used to measure R_s pre and post thermal annealing, avoiding sample edges, in order to verify the approximation of the four probe method.⁶⁸ The measurement accuracy is verified against a $12.93 \Omega/\square$ ITO on glass reference (Jandel Engineering Ltd., tested against a NIST traceable sample).

Scanning electron microscopy

GBs and cellular adaption to substrates are qualitatively assessed through scanning electron microscopy (SEM). Images are acquired collecting secondary electrons on a Gemini SUPRA 40 SEM (Carl Zeiss NTS GmbH, Oberkochen, Germany) working at 5 keV. Cellular samples are washed with 0.1 M cacodylate buffer (pH = 7.2) and fixed with a solution containing 2 % glutaraldehyde (Fluka, Italy) in 0.1 M cacodylate buffer for 1 h at RT. Cultures are then washed in a cacodylate buffer and dehydrated by dipping in water/ethanol solutions at progressively higher alcohol concentrations (50%, 75%, 95% and 100% ethanol for 3 minutes each). Samples are dried at room temperature in a N₂ chamber overnight. Prior to SEM imaging samples are gold metalized in a metal sputter coater (Polaron SC7620).

Neuronal cultures and electrophysiology

Dissociated hippocampal neurons are obtained from postnatal (P2÷P3) rats, as previously described.¹⁶⁻¹⁸ For each culture series, sister cultures are prepared by plating cells on control substrates or on GBS-covered coverslips, and the same number of cells (~30,000) is plated on each coverslip. Cultures are used for experiments over 8–10 days *in vitro*. Neuronal and glial cell densities are quantified by immunofluorescence stainings and microscopy analysis.^{16, 18, 20} Cultures are fixed with 4% paraformaldehyde in phosphate buffer solution (PBS), incubated in blocking solution and stained by incubation with primary antibodies (against NeuN for neuronal staining, mouse monoclonal, 1:100, Millipore; or against glial fibrillary acidic protein -GFAP- for glial cells staining, mouse monoclonal, 1:400, Sigma). Upon washout in PBS, cultures are incubated with the secondary goat antibody (Alexa Fluor-488-conjugated; 1:500; Invitrogen) and with 4',6-diamidino-2-phenylindole (DAPI) for nuclei staining. Samples are mounted in a Vectashield medium and images are acquired using a conventional epifluorescence microscope (Leica DM6000). Whole-cell patch clamp recordings are performed with pipettes filled with: 120 mM K gluconate, 20 mM KCl, 10 mM HEPES, 2 mM MgCl₂, 4 mM MgATP, 0.3 mM GTP (pH adjusted to 7.35 with KOH). The external solution contains: 150 mM NaCl, 4 mM KCl, 1 mM MgCl₂, 2 mM CaCl₂, 10 mM HEPES, 10 mM glucose (pH adjusted to 7.4 with NaOH). Experiments are performed at room temperature. Recordings are taken with a Multiclamp 700B amplifier (Molecular Devices). Current and voltage clamp signals are digitized at 10 kHz by a Digidata 1440A equipped with the pCLAMP 10 software (Molecular Devices) and stored for analysis. Neuronal passive membrane properties are evaluated by applying a 10 mV hyperpolarizing step (250 ms). Dual recordings are taken by eliciting action potentials in the presynaptic cell (in current-clamp mode) by injecting short square current pulses (1 nA, 2 ms) at -60 mV resting membrane potential, and recording response postsynaptic currents from the postsynaptic cell voltage-clamped at -56 mV holding potential (potential values not corrected for 14 mV liquid junction potential). Paired-pulse ratio (PPR) analysis is performed on mono-synaptically connected (latency between the peak of the elicited presynaptic action potential

and the onset of the postsynaptic current response less than 3 ms) neuron pairs.¹⁸ A pair of spikes at 20 Hz is induced in the presynaptic neuron (10–20 times repetitions at 10 s intervals) and currents evoked in the postsynaptic neuron are recorded (in our experimental setting the majority of postsynaptic currents observed in dual recordings are GABA-mediated, as in Ref. 18; GABAergic synapses are therefore analysed). Recordings showing ongoing spontaneous activity occasionally superimposed to the evoked postsynaptic current are excluded from the analysis.

Statistics

Statistically significant differences between datasets are assessed by the Student's test (after validation of variances homogeneity by Levene's test) for parametric data and by Mann-Whitney for non-parametric ones.⁶⁹ For synaptic pairing, comparisons are performed by paired t-test between datasets obtained from sister cultures from the different culture series. A P value <0.05 is taken as indicative of statistically significant difference. n is the number of neurons, if not otherwise indicated. Data are shown as mean ± standard error of the mean.

Ethical statement

The work on animals (neonatal rats) was performed according to the EU guidelines (2010/63/UE) and Italian law (decree 26/14). The use of animals was approved by the Italian Ministry of Health, in agreement with the EU Recommendation 2007/526/CE. Animals were hosted by the University of Trieste Animal Facility (Life Sciences Department, Italy, authorized by the Italian Ministry of Health), and breeding conditions and procedures complied with the 2010/63/UE EU guidelines and Italian law (26/14). Neonatal rats were sacrificed by rapid decapitation and the tissue of interest (hippocampus) harvested, all efforts were made to minimize suffering. The work was performed on explanted tissue and did not require ethical approval. The entire procedure employed in the present study is in accordance with the regulations of the Italian Animal Welfare Act, with the relevant EU legislation and guidelines on the ethical use of animals and is approved by the local Authority Veterinary Service. The procedures were also in accordance with the ethical assessment established in the EU FP7-ICT-2013-FET-F GRAPHENE Flagship project (No. 604391).

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FIGURE CAPTIONS

Figure 1: a) Optical absorption of LPE dispersion in water/SDC diluted 1:12 with pure water/SDC.
 b) Raman spectrum at 514.5nm for representative LPE (black curve) and BM (red curve) flakes.

Distribution of c) Pos(2D), d) FWHM(2D), e) Pos(G), f) FWHM(G), g) I(2D)/I(G), h) I(D)/I(G), and distribution of I(D)/I(G) as a function of i) FWHM(G) and l) Disp(G) for LPE (black dashed histograms and dots) and BM (red dashed histograms and dots) flakes, respectively.

Figure 2: a) Representative Raman spectra of flakes, compared with films. LPE (black curve) and BM (red curve). Pos(2D) and FWHM(2D) for (b, c) flakes in dispersion and (d, e) deposited on the glass substrate for LPE (black dashed histograms) and BM (red dashed histograms).

Figure 3. AFM images of a) LPE-GBS and b) BM-GBS. c), SEM image of dissociated hippocampal neurons grown onto LPE-GBS. Immunofluorescence staining of cultures developed on (d) control and (e) LPE-GBS substrates, marked for neurons (NeuN, green) and nuclei (DAPI, blue). (f) plot summaries of neuronal density in the two GBSs culturing conditions. Immunofluorescence staining for glial cells (marked for GFAP, green; DAPI in blue) on (g) control and (h) LPE-GBS. (i) Plot summaries of glial cells density in the two GBSs culturing conditions. Histograms are mean \pm standard error.

Figure 4. Representative PSC current tracings recorded from voltage clamped neurons grown on (a) control substrates or on (b) LPE-GBS. Frequency and amplitude of spontaneous PSCs are almost identical in neurons grown on (c) control and on (d) GBSs. Each dot represents values from a single neuron. Histograms are mean \pm standard error.

Figure 5. (a) Schematic representation of the experimental setting of dual recordings: an action potential is elicited in the presynaptic neuron (current-clamped; top), while recording response postsynaptic current from the postsynaptic neuron (voltage-clamped, bottom). (b) The fraction of synaptically coupled neuron pairs is similar in the two culturing conditions. Each dot represents the % pairing from neurons recorded from the same culture series. Dual recordings from neurons interfaced to (c) control or (d) BM-GBS substrates, in which a pair of action potentials at 20 Hz is induced in the presynaptic neuron (top traces) and the response postsynaptic current recorded in the postsynaptic neuron (bottom traces), to investigate short-term synaptic plasticity (paired pulse). The action potentials pair decrease the amplitude of the second postsynaptic current with respect to the

first one (paired pulse depression) in both control and GBSs-interfaced neurons. (e) Histograms quantifying synaptic plasticity as the ratio between the amplitudes of the second and first postsynaptic currents (paired-pulse ratio). Each dot represents the value from one neuron pair.

Histograms are mean \pm standard error

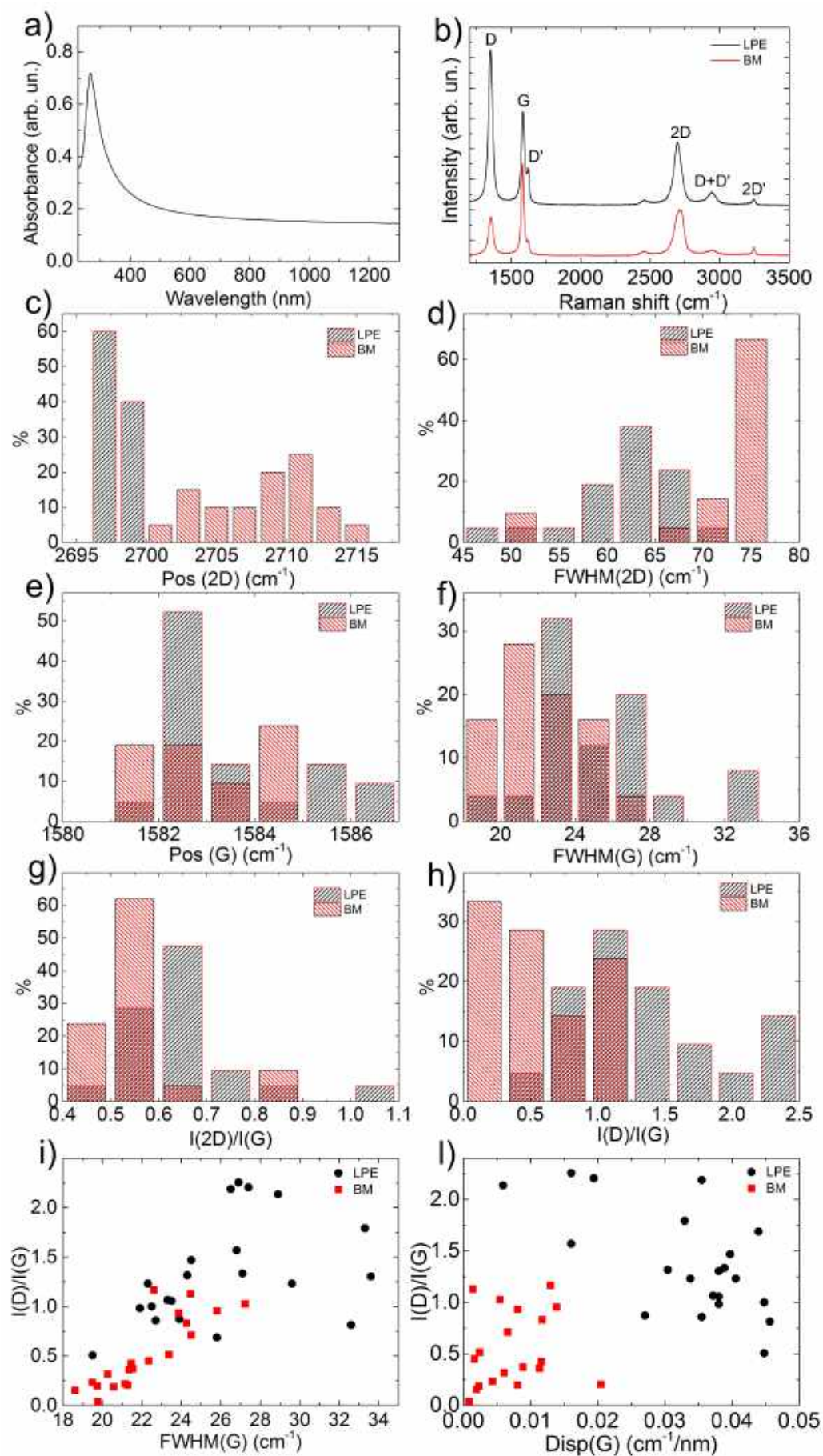


FIGURE 1

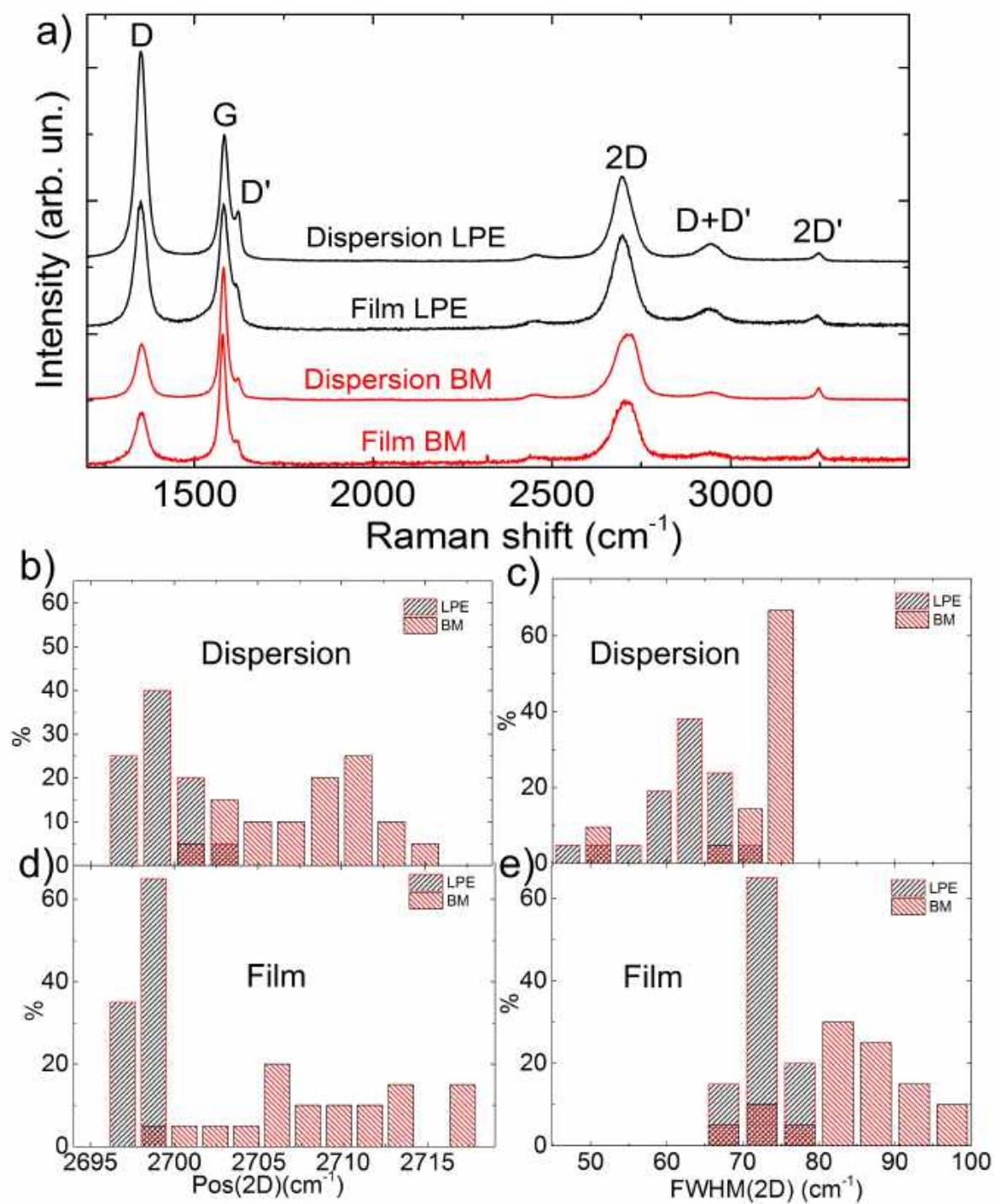


FIGURE 2

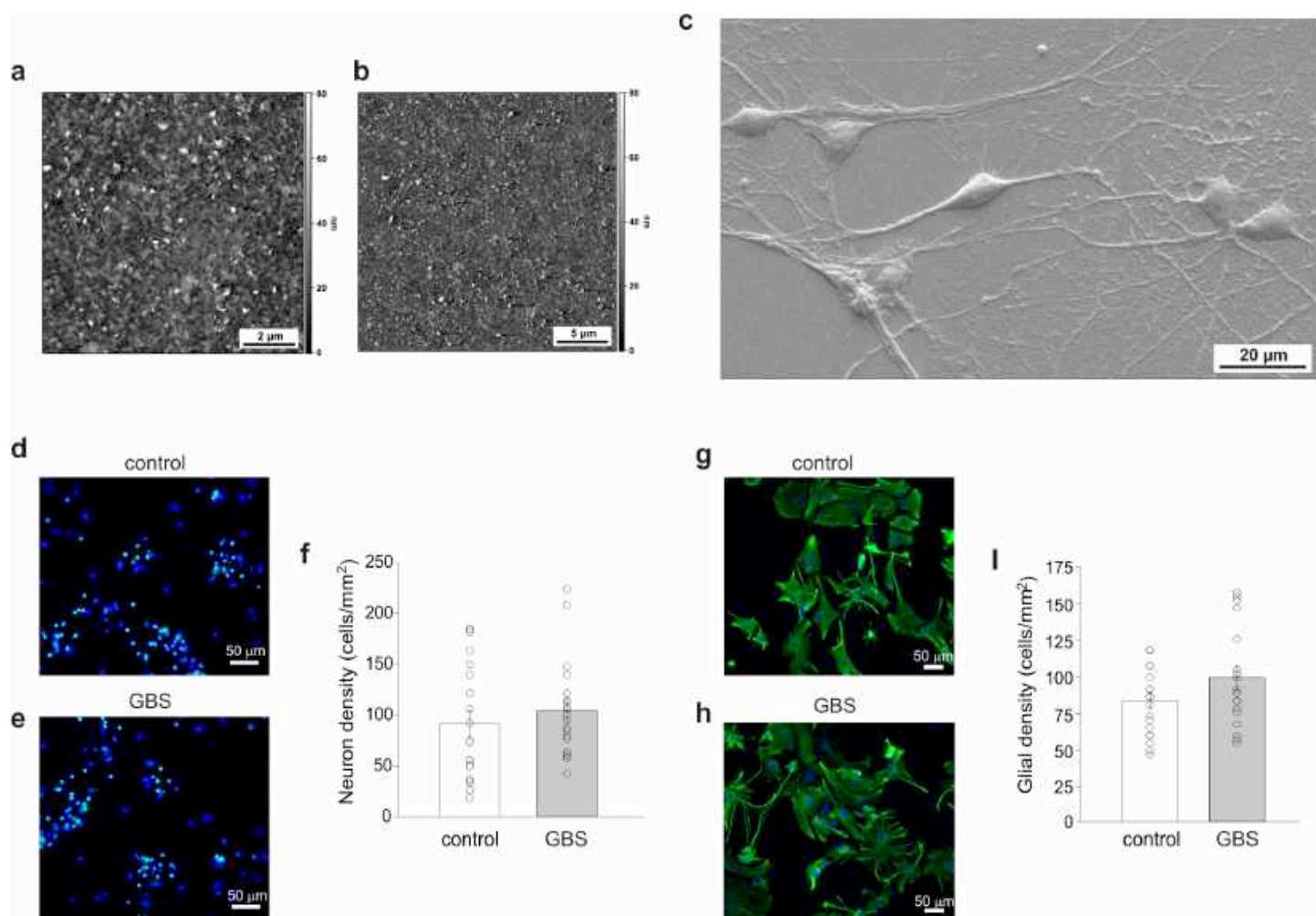


FIGURE 3

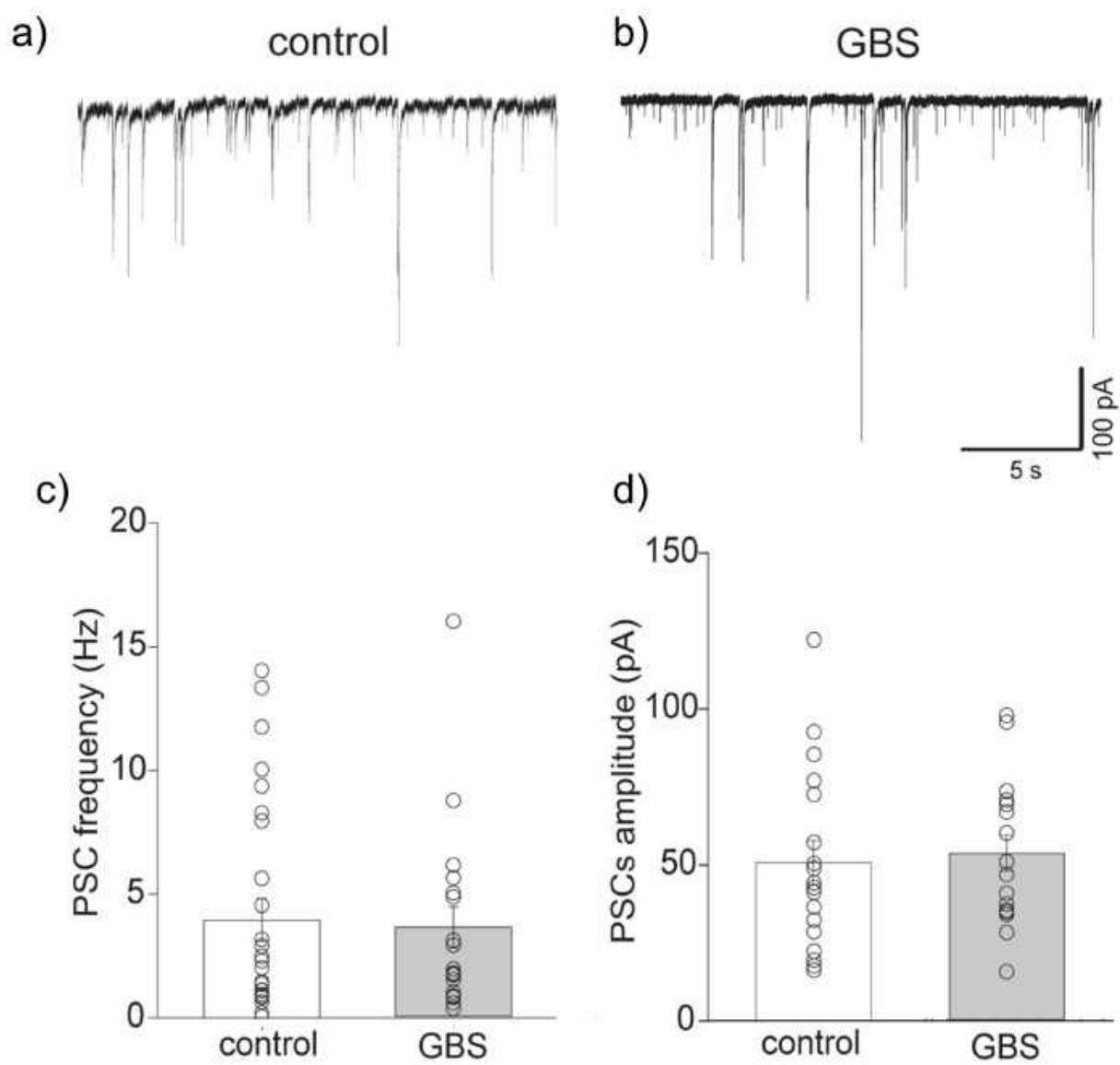


FIGURE 4

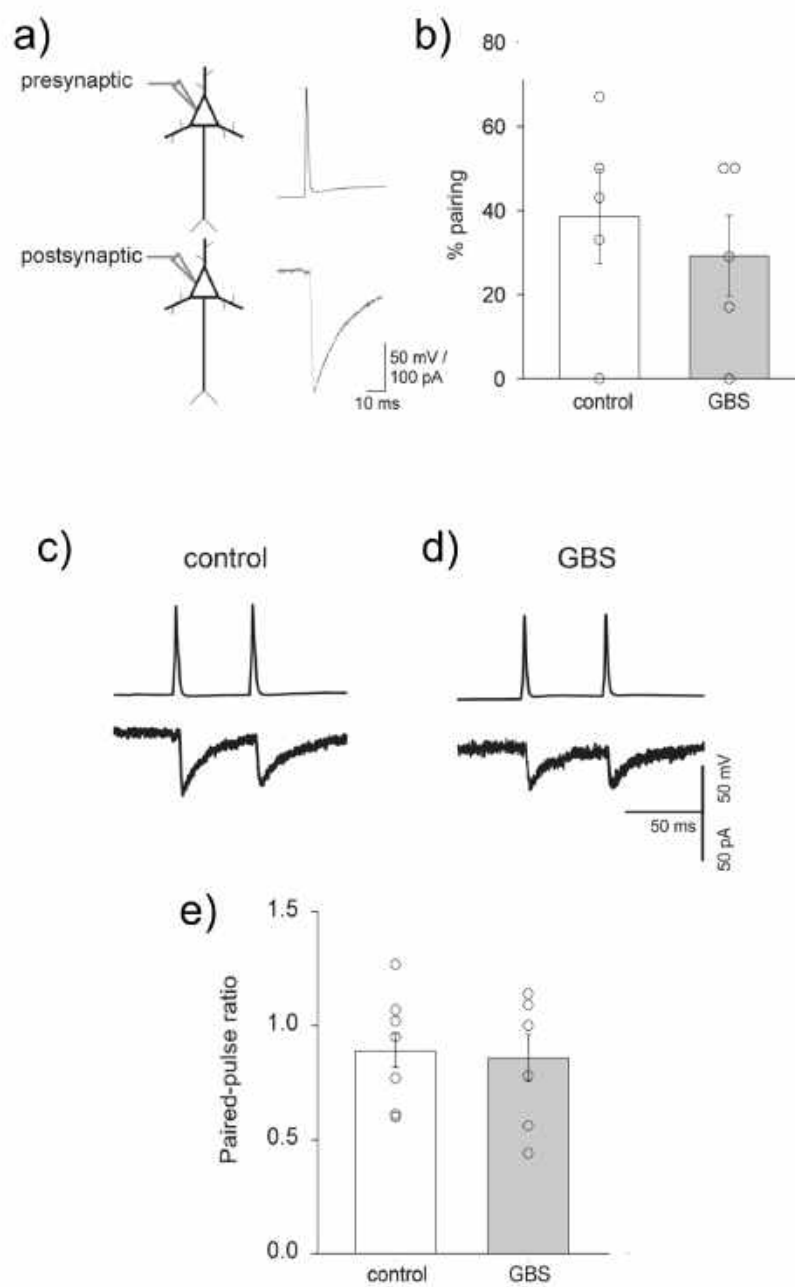


FIGURE 5