Analytical Methods

MINIREVIEW

Check for updates

Cite this: Anal. Methods, 2025, 17, 635

Received 1st October 2024 Accepted 25th November 2024

DOI: 10.1039/d4ay01793e

rsc.li/methods

Introduction to laser-induced graphene

Graphene is a unique material which is composed of a single layer of carbon atoms within a hexagonal honeycomb lattice. It exhibits advantageous properties of improved thermal, mechanical, optical, and electrical transport, resulting in it being explored in a range of applications.¹⁻³ There are many ways to produce graphene, which include micromechanical and liquid-phase exfoliation, chemical vapour, and pulsed laser deposition to name just a few.^{4,5} One notable approach is the use of a CO₂ infrared laser cutter system to form laserinduced graphene on polyimide films, where sp³-carbon atoms are photothermally converted to sp²-carbon atoms via pulsed laser irradiation, which has revolutionized how laserinduced graphene can be readily formed.6 This approach results in rapid temperature increases that breaks C-O, C=O, and N-C bonds, whilst releasing CO and C₂H₂, resulting in the instant creation of graphene, at a significantly low cost. Fig. 1A shows a schematic of the process for the creation of laserinduced graphene from polyimide, where various geometries

Electroanalytical overview: the use of laser-induced graphene sensors

Robert D. Crapnell, ^D^a Elena Bernalte, ^D^a Rodrigo A. A. Muñoz ^D^b and Craig E. Banks ^b*^a

Laser-induced graphene, which was first reported in 2014, involves the creation of graphene by using a laser to modify a polyimide surface. Since then, laser-induced graphene has been extensively studied for application in different scientific fields. One beneficial approach is the use of laser-induced graphene coupled with electrochemistry, where there is a growing need for disposable, conductive, reproducible, flexible, biocompatible, sustainable, and economical electrodes. In this mini overview, we explore the use of laser-induced graphene as the basis of electroanalytical sensors. We first introduce laser-induced graphene, before moving to the use of laser-induced graphene electrodes highlighting the various approaches and different laser parameters used to produce different graphene micro and macro structures, whilst describing how these structures are characterised and benchmarked for those working in the field of laser-induced graphene electrodes as the basis of electrochemical sensing platforms towards key analytes and its use in the development of biosensors. We provide a critical overview of the use of laserinduced graphene sensors compared to screen-printed and additive manufactured electrodes, providing future suggestions for the field.

> can be realised. Furthermore, Fig. 1B-D shows scanning electron microscopy (SEM) micrographs that show the orderedporous morphology obtained. The Raman spectrum shows the three peaks that are observed (Fig. 1E), assigned to the D, G and 2D bands, where analysis shows that the 2D band is the same as that of single-layer graphene but it has a larger full width at half maximum of $\sim 60 \text{ cm}^{-1}$ suggesting randomly stacked graphene layers, and the D/G intensity ratio indicates a high degree of order within the graphene formation in the laser-induced graphene.6 Note that the D band is present, which suggests that it is not pristine graphene (which has no D peak) but contains defects.8 X-ray diffraction (XRD) of the laserinduced graphene is shown in Fig. 1E, where an intense peak centred at $2\theta = 25.9^{\circ}$ is seen, giving an interlayer spacing (I_c) of \sim 3.4 Å between the (002) planes in the laser-induced graphene, indicating a high degree of graphitization.6 The authors also explored different laser powers from 2.4 W to 5.4 W, where they monitored the change in the Raman spectra and noted that the increase in power degrades the quality of laser-induced graphene.6 This unique approach has started the trend of how to rapidly make laser-induced graphene, which has now been explored toward medical diagnostics, energy storage and conversion devices, water purification, electronic devices, microfluidic systems, humidity, electrochemical and piezoresistive sensors, health monitoring and wearable sensors, to name just a few.9-11 The main reason why laser-induced graphene has attracted attention is that graphene can be readily



View Article Online

View Journal | View Issue

^aFaculty of Science and Engineering, Manchester Metropolitan University, Dalton Building, Chester Street, Manchester, M1 5GD, UK. E-mail: c.banks@mmu.ac.uk; Tel: +44-(0)-161-247-1196

^bInstitute of Chemistry, Federal University of Uberlândia, Uberlândia, 38400-902, Minas Gerais, Brazil

Open Access Article. Published on 09 2024. Downloaded on 2025/6/1 20:59:50.



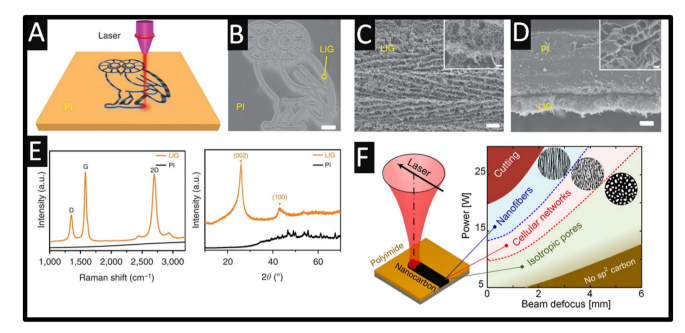


Fig. 1 (A) Schematic of the synthesis process of laser-induced graphene from polyimide. (B) SEM image of laser-induced graphene patterned into an owl shape; scale bar, 1 mm. The bright contrast corresponds to laser-induced graphene surrounded by the darker-coloured insulating polyimide substrates. (C) SEM image of the laser-induced graphene film circled in (B); scale bar, 10 μ m. Inset shows the corresponding higher magnification SEM image; scale bar, 1 μ m. (D) Cross-sectional SEM image of the laser-induced graphene film on the polyimide substrate; scale bar, 20 μ m. Inset shows the SEM image showing the porous morphology of laser-induced graphene; scale bar, 1 μ m. (E) Representative Raman spectrum of a laser-induced graphene film and the starting polyimide film and XRD pattern of powdered laser-induced graphene scraped from the polyimide film. Figure reproduced from ref. 6. Copyright 2014 Nature. (F) Morphology diagram mapping the ranges of laser parameters (power and degree of beam defocusing) for creating different types of LINC (isotropic porous morphology, anisotropic cellular networks, and aligned woolly nanofibers) for a CW CO₂ laser (wavelength $\lambda = 10.5$ mm) scanning at $\nu = 500$ mm s⁻¹. Figure reproduced from ref. 7. Copyright 2021 American Chemical Society.

made without the need for expensive cleanroom equipment, solvents, wet chemical procedures and subsequent treatments.

Abdulhafez and co-workers7 studied morphological transitions in laser-induced graphene on polyimide substrates, where they explored the power as a function of the beam defocus. As shown in Fig. 1F, a useful zone diagram is presented which shows the morphology which ranges from various fluence values (F) corresponding to cutting ($F = 25 \text{ J cm}^{-2}$), nanofibers $(F = 17 \text{ J cm}^{-2})$, anisotropic cellular networks $(F = 12 \text{ J cm}^{-2})$ and isotropic pores ($F = 5 \text{ J cm}^{-2}$). The physicochemical characterisation of laser-induced graphene is dependent upon the material/substrate, laser type and associated parameters, e.g., wavelength, power and scanning speed, and environmental conditions also affect the hydrophilicity/hydrophobicity, nanoand micro-scale morphology, i.e., crystallinity, porosity, and chemical composition of the graphene produced.9,10 For example, various materials have been used to produce laserinduced graphene, which can be classed into thermoplastics (polyimide, poly(vinyl chloride) and polyphenylene sulfide), thermoset materials (polystyrene and phenolic resin), natural polymer materials (lignin and cellulose) and non-polymeric materials (charcoal and activated carbon).9,12 Furthermore, as shown in Fig. 2A, laser-induced graphene can be extended to bread, wood, cotton paper, cardboard and muslin cloth, which allows the approach to be utilised on biodegradable substrates through tuning the lasing parameters,¹² noting that this is the Rice University (USA) Owl Mascot. That approach has used

a CO_2 laser with a wavelength of 10.6 µm, but this can be achieved on various substrates/surfaces using continuous wave, pulsed and ultrashort pulse lasers, as well as visible and infrared laser sources, where the power source and raw materials for the generation of different laser-induced graphene outputs have distinct advantages to form graphene.¹⁵

One particular use of laser-induced graphene is within the realm of electrochemistry, where these materials have found application in batteries, water splitting devices, health and gas monitoring, fuel cells, and supercapacitors and are applied in various wearable and telemedicine platforms.^{12,16–20} In this mini-review, we introduce laser-induced graphene focusing on electrochemical-based sensors which can provide on-site analysis with a highly selective and sensitive output, whilst remaining cost-effective. We therefore summarise the use of laser-induced graphene as the basis of electroanalytical sensors and compare these critically against screen-printed and additive manufactured electrodes.

Laser-induced graphene electrodes

Laser-induced graphene electrodes have been utilised as the basis of sensors due to their rapid and easy approach of fabrication, high conductivity, flexibility, and unique micro- and macro-features. Most importantly, the fact that these are multilayer graphene, it is the presence of edge plane sites/defects which has been extensively shown to be the origin of fast

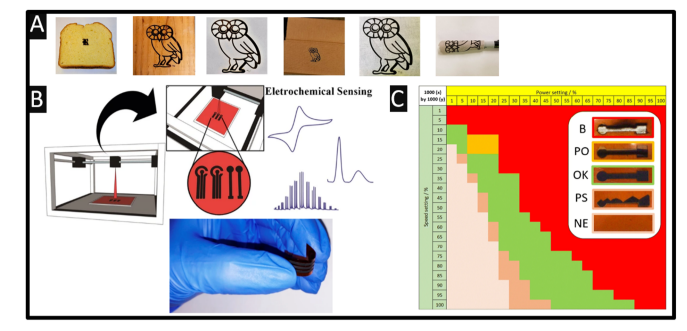


Fig. 2 (A) Laser-induced graphene on (left to right) bread, fire-retardant treated pine wood, cotton paper, cardboard, grey muslin cloth and a muslin cloth wrapped around a marker pen. All owls depicted are 60 mm in height. Figure reproduced from ref. 12. Copyright 2018 American Chemical Society. (B) Illustration of the visible laser-printer machine showing the production of laser-induced graphene where a three-electrode configuration (1.5 cm \times 4 cm) and single working electrode (1 cm \times 4 cm) are shown. Also shown is the flexibility of laser-induced graphene electrodes. Figure reproduced from ref. 13 Copyright 2022 Springer. (C) Heatmap of the electrode outcome vs. power and speed settings at a pulse density of 1000 \times 1000 (x by y) (colour code as indicated in the inset: green = ok, darker brown = partial scribing (PS), lighter brown = no effect (NE), orange = laser-induced graphene peeled off from the substrate (PO), and red = laser burned through the substrate (B)). Figure reproduced from ref. 14. Copyright 2021 Springer. Creative Commons Attribution 4.0 International License.

electron transfer.^{21,22} Another benefit is that production of laserinduced graphene does not need a template, as is the case when using screen-printed electrodes where a stencil design is required. For example, Fig. 2B shows the use of a visible laserprinter machine and the production of laser-induced graphene, where a three-electrode configuration (1.5 cm × 4 cm) and single working electrode (1 cm × 4 cm) can be realised. The real images of the produced laser-induced graphene electrode which highlights their flexibility are also shown.¹³ A useful approach has been reported by Clark and co-workers where the use of a stencil allowed minimal achievable resolution of the laser-induced graphene from 120 μ m to 45 μ m allowing a microarray electrode to be realised.²³

As mentioned above, laser-induced graphene can show variability between different research groups and batch-tobatch variations,²⁴ where the graphene is dependent upon the material/substrate, laser type and associated parameters, *e.g.*, wavelength, power and speed, and also environmental conditions. In each case, where researchers are producing laserinduced graphene, they need to physicochemically characterise their graphene using SEM, Raman, X-ray photoelectron spectroscopy (XPS) and electrochemistry. That said, this is often overlooked.

A useful approach has been reported by Behrent and coworkers¹⁴ who, as shown in Fig. 2C, have produced a heatmap of laser-induced graphene electrodes showing how power *versus* speed can either make a useful laser-induced graphene electrode, or not. They have identified these as follows: green is ok,

darker brown is a result of partial scribing (see inset of Fig. 2C: PS), lighter brown indicates no effect (NE), orange resulting from laser-induced graphene which peeled off from the substrate (PO), while red is a result of the laser burning through the substrate (B); more authors need to follow the valuable approach by Behrent and co-workers.14 A useful review paper has been published by Muzyka and Xu25 which covers laserinduced graphene characterisation using Raman spectroscopy, sheet resistance, electrochemical active surface area, wettability (contract angles), and the heterogeneous electron transfer rate constant and compares different laser wavelengths with heterogeneous electron transfer rate constants, which can show great variations.25 We next summarise the physicochemical characterisation of their laser-induced graphene electrodes. In Table 1, one can see the different SEM images of laserinduced graphene electrochemical surfaces that are a result of different laser sources, where the Raman spectrum, the electrochemical area (A_{real}) and the heterogeneous rate constant (k^0) reported for $[Ru(NH_3)_6]^{3+/2+}$ and $[Fe(CN)_6]^{4-/3-}$ redox probes are also shown. It is expected that the laser-induced graphene sensors will be dependent upon the surface and chemical morphology, indicating that researchers need to conduct physicochemical characterisation of their graphene surfaces.

SEM (scanning electron microscopy)

SEM is critical to observe the structure of the formed laserinduced graphene structures, where one can see the micro and macro structures (see Table 1).

Table 1 An overview of the laser-induced graphene electrodes with the SEM, Raman spectroscopy, electrochemical area (A_{real}) and the heterogeneous rate constant (k^0) reported for outer-sphere [Ru(NH₃)₆]^{3+/2+} and inner-sphere [Fe(CN)₆]^{4-/3-} redox probes 13 $k^0/[\text{Fe}(\text{CN})_6]^{4-/3-}$ $0.0056~{\rm cm~s^{-1}}$ This article is licensed under a Creative Commons Attribution 3.0 Unported Licence. $k^0/[Ru(NH_3)_6]^{3+/2+}$ $0.20~\mathrm{cm}^2$ A_{real} 4000 20, 20, 300 20 Raman shift / cm "Q+Q 2000 G 000 Raman spectra Normalized intensity / a.u. (cc) SEM

IR CO₂ laser (10.6 μ m)

11

 $0.0044 \ {\rm cm \ s^{-1}}$

 1.3 cm^2

2

۵

Intensity / a.u.

3000

20

0

Raman shift / cm⁻¹

C

1.4 1.2 1.0 0.8 0.6 0.4

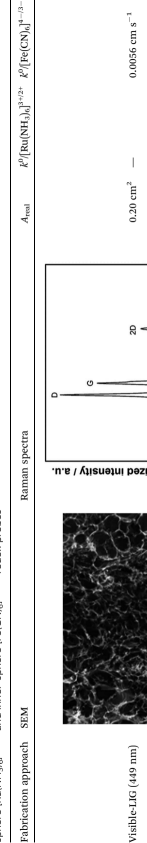
This journal is © The Royal Society of Chemistry 2025

IR CO₂ laser (10.6 μ m)

Analytical Methods

References

Open Access Article. Published on 09 2024. Downloaded on 2025/6/1 20:59:50.



638 | Anal. Methods, 2025, 17, 635-651

26

3500

3000

2000 2500 Raman shift, cm⁻¹

1500

1000

UV LIG

0.2 0.0

 26

 $0.282~{\rm cm~s^{-1}}$

 $0.0231~{\rm cm~s^{-1}}$

2D' 0+0

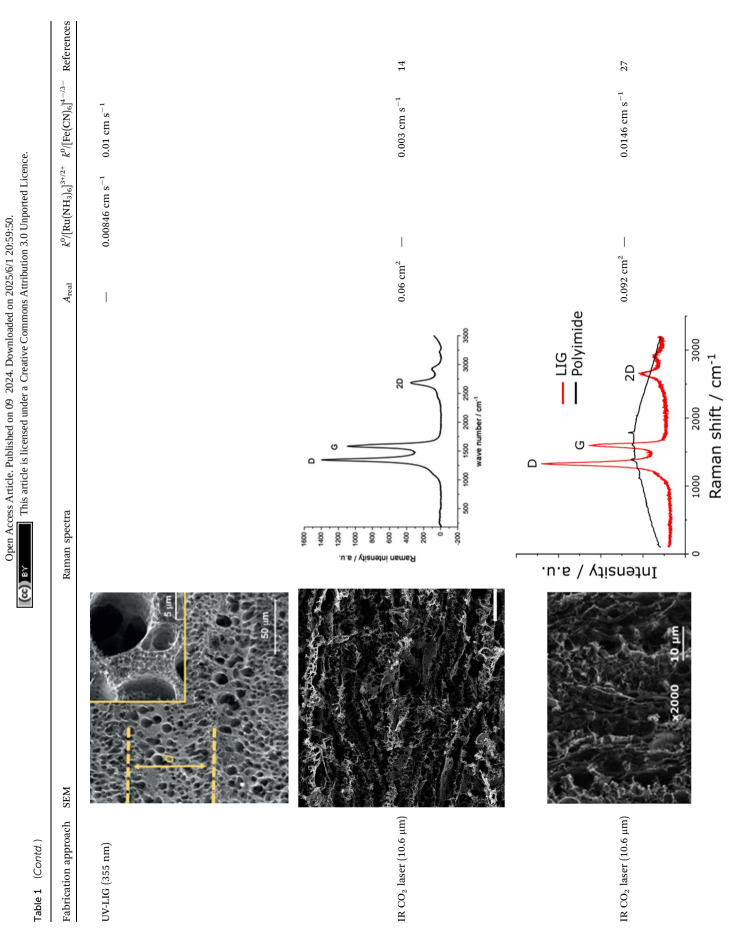
"0+D

RIG

'n

Normalized intensity, arb. units

20



Goniometry

Goniometry is used to determine the hydrophobicity, where solutions are carefully applied onto the laser-induced graphene surface where the average contact angle is reported. For example, an average contact angle using deionised water is reported to be 56° but it has been shown that the contact angle varied by more than 30% across four batch-to-batch variations.²⁴

Raman spectroscopy

Raman spectroscopy is critical to identify the presence of micro and macro structures. As shown in Table 1, a typical Raman spectrum for laser-induced graphene is shown (first image) where the main peaks are identified, which are D, G and 2D bands and also D + D'' and 2D' bands. If we elaborate on this figure, ¹³ we can observe the D (1343 cm^{-1}), G (1591 cm^{-1}), and 2D (2691 cm^{-1}) vibrational bands. These arise from the Stokes phonon energy shift, where the G band is a primary in-plane vibrational mode, and the 2D band is a second-order overtone of a different in-plane vibration, the D band. Note that for graphene, one should only observe G and 2D bands, and the presence of the D band indicates defects. The G band represents the in-plane stretching vibrations of the sp² bonded carbon atoms. Its position is affected by the number of layers present, where the peak shifts to lower wavenumbers as the number of graphene layers increases. The number of layers can be calculated from the peak position using the following equation where $\omega_{\rm G}$ is the band position, and *n* the number of layers present:²⁸

$$\omega_{\rm G} = 1581.6 + 11(1 + n^{1.6}) \tag{1}$$

Of further note is that the intensity of the G band decreases as the layers increase from mono- to tri-layer. The 2D band is also indicative of the number of layers present in the sample, which originates from a double resonance enhanced twophonon lateral vibrational process. The 2D band changes in shape and position as the layers of graphene increase. The Dband is the peak which highlights defects within the sample, where the more intense this band is, the higher the level of disorder within the sample. This band is a result of ring breathing mode of sp²-carbon rings and must be adjacent to a defect to be Raman active. The D-band is a 1-phonon lattice vibrational process and is a double resonance band. The ratio of the intensities of I_{2D} and I_G bands, I_{2D}/I_G will be equal to 2 for high quality single layer graphene. The ratio changes to 1, 0.8 and 0.5 indicating that in each case, these correspond to double layer, few layer and multi-layer graphene. See ref. 29 for a calculator to help with the deduction of graphene layers. Lastly, one can observe a minor band D + D' (2936 cm⁻¹) which indicates the disorder structure of graphene with oxygencontaining groups where combined overtones of D + D'' and 2D' are also identified which indicate the formation of a graphite-like carbon structure.

XPS (X-ray photoelectron spectroscopy)

XPS can be used to analyse the elemental composition and chemical state of the material's surface. XPS can be used to

measure the elemental composition and the chemical and electronic state of atoms. This approach is useful as it can be used to determine the carbon-oxygen content and identify them. Using a CO₂ pulsed laser working at 10.6 μm using a scan speed of 160 mm s^{-1} , manufacturing laser-induced graphene has shown, via XPS, an % atomic C of 76.6, % atomic O of 16.6, while % atomic nitrogen and silicon are 6 and 0.8 respectively, which are ascribed to C-O-C, C-N and C=O functional groups.30 The use of functional groups enhances surface wettability assisting with the adsorption and desorption of ions and enhancing interaction with the electrolyte. For example, it has been shown that oxygenated species decrease the electron transfer kinetics of ferrocyanide using basal plane and edge plane pyrolytic graphite electrodes.31 On the other hand, others report that oxygenation of the single-walled carbon nanotube ends is known to speed up the electron transfer kinetics³² understanding which oxygenated species are present is critical and needs to be reported.

Electrochemical characterisation

The heterogeneous electrochemical rate constant, k^0 , is calculated as an average from three sets (minimum) of scan rate studies for each electrode using an appropriate redox probe using the quasi-reversible electrochemical reactions through the following Nicholson formula:³³

$$\psi = k^0 [\pi D n \nu F/(RT)]^{-1/2} \tag{2}$$

where ψ is a kinetic parameter, *D* is the diffusion coefficient of the redox probe, *n* is the number of electrons that are taking part in the process, *F* is the Faraday constant, *v* is the scan rate, *R* is the gas constant, and *T* is the temperature in Kelvin. In order to calculate the heterogeneous electrochemical rate constant, we use the peak-to-peak separation (ΔE_p) to deduce ψ , where ΔE_p is obtained at various voltammetric scan rates. The heterogeneous electrochemical rate constant can be calculated *via* the gradient when plotting ψ against $[\pi DnvF/RT]^{-1/2}$. Lavagnini *et al.*³⁴ proposed the following function of $\psi(\Delta E_p)$, which fits Nicholson's data, and for practical usage, rather than producing a working curve, we suggest the following approach:

$$\psi = (-0.6288 + 0.0021X)/(1 - 0.017X) \tag{3}$$

where $X = \Delta E_p$. In cases where ΔE_p is higher than 212 mV, the following equation should be implemented:³⁵

$$k^{0} = 2.18[D\alpha n\nu F/(RT)]^{1/2} \exp[-(\alpha^{2} nF/RT)\Delta E_{\rm p}]$$
(4)

where α is the transfer coefficient (usually assumed to be close to 0.5). If we consider the use of the Randles–Ševćik equation under non-standard conditions for reversible and quasireversible electrochemical processes:^{36–38}

$$I_{\rm p.f}^{\rm Rev} = \pm 0.446 n F A_{\rm real} C \sqrt{\frac{n F D \nu}{R T}}$$
(5)

$$I_{\rm p.f}^{\rm Quasi} = \pm 0.436 n F A_{\rm real} C \sqrt{\frac{n F D \nu}{R T}}$$
(6)

where $I_{p.f}$ is the voltammetric current/analytical signal using the forward peak of the electrochemical process, A_{real} is the electroactive area of the electrode. Eqn (5) and (6) can be used to determine the electroactive area (A_{real}) through a simple cyclic voltammetry experiment. In this approach, typically, a redox probe is used to determine a plot of the forward peak current, $I_{p.f}$, as a function of applied voltammetric scan rate ($v^{1/2}$) where the electrode should be flat and non-porous, where in the case of laser-induced graphene electrodes these are providing an estimate that can help others to benchmark and compare the electrochemical surface area with others. Lastly, it is also good to compare the geometric area (A_{geo}), using the following equation:

$$\% \text{Real} = (A_{\text{real}}/A_{\text{geo}}) \times 100) \tag{7}$$

As can be seen in Table 1, there are many different micro and macro structures produced using different lasers forming laserinduced graphene, which are multi-layer graphene where the heterogeneous electrochemical rate constant and different electrochemical areas can vary. This indicates that laserinduced graphene electrodes are similar to solid carbon electrodes.39-41 As mentioned above, it has been shown that oxygenated species decrease the electron transfer kinetics of ferrocyanide using basal plane and edge plane pyrolytic graphite electrodes³¹ but others report that oxygenation of the single-walled carbon nanotube ends is known to speed up the electron transfer kinetics.32 One must understand the use of the redox probes in terms of their inner-sphere and outer-sphere nature. For example, it is known that $[Ru(NH_3)_6]^{3+/2+}$ is a nearideal outer-sphere redox probe, while $[Fe(CN)_6]^{4-/3-}$ is known as an inner-sphere redox probe which is strongly influenced by the state of the electrode surface in terms of the surface chemistry, microstructure and oxygenated species. This may account for the fast heterogeneous electrochemical rate constant reported using laser-induced graphene, as shown Table 1. Ultimately, it is highly encouraged for researchers reporting the formation and use of laser-induced graphene to characterise and benchmark their material using the list of approaches highlighted above.

Laser-induced graphene sensors

Table 2 provides a snapshot of the use of laser-induced graphene electrochemical sensors, which it is shown in alphabetical order in terms of the analyte being measured alongside. These cases are reported together with the substrate graphene is formed on, the wavelength of the laser source and both the applied power and speed. Moreover, Table 2 summarises the use of laser-induced graphene towards various analytes along with their analytical performance (linear range and limit of detection (LoD)) and real sample application.

One can see that aristolochic acid and roxarsone have been simultaneously measured with a very low LoD of 16.5 and 2.31 nM.⁴⁴ Aristolochic acid has been associated with urinary epithelial cancer, renal tumours, and renal failure related to the intake of herbal medicine derived from *Aristolochia* plants, while roxarsone is an organoarsenic chemical widely used in poultry farming which can be linked to toxic and carcinogenic effects in humans; therefore, there is a need for an electrochemical sensor for monitoring both compounds.⁴⁴

An innovative approach for the measurement of methane has been reported using laser-induced graphene substrates.⁶¹ Using an interdigitated laser-induced graphene design, palladium nanoparticles (50 nm) are decorated on porous polyvinylidene fluoride (PVDF) with 1-ethyl-3-methylimidazolium TFSI (EMImTFSI) in an N-methylpyrrolidone (NMP) layer and then painted onto the electrodes. The interdigitated laserinduced graphene design (see Fig. 3) is composed of 14 anode/cathode fingers, each with a dimension of 13×0.4 mm and a spacing of 0.5 mm between them. Also shown in Fig. 3B is the characterisation of laser-induced graphene electrodes with the Raman spectrum of the laser-induced graphene electrode material, which reports multi-layer graphene. Also shown is a schematic (Fig. 3B) of the dimensions of the laser-induced graphene electrodes and an indication of where the electrolyte is applied. A cross-section SEM image of two adjacent electrodes shows that the electrode structure is not completely embedded within the polyimide, but the porous laser-induced graphene electrodes are $\sim 100 \ \mu m$ above the plane of the unexposed polyimide film. The sensor is applied to the measurement of methane, where the electrochemical procedure transforms methane into carbon dioxide and water. The author reports that this sensor gives a linear range of 10-50 ppm and a LoD of 9.2 ppm, which is 4 orders of magnitude lower than other electrode designs.61

Fig. 3C shows the experimental set-up for the laser-induced technique used in the fabrication of reduced graphene oxide.42 A graphene oxide solution is coated onto an alumina ceramic (Al₂O₃) substrate through drop coating. It is placed in an oven for 2 h at 60 °C, after which, a 1064 nm laser is used with fibre optics via a telecentric lens (specification: f = 125mm, $\lambda = 1064$ nm). The laser is passed over an excitation filter and reflected by a shrinking mirror into a mirror spot and focused by using an objective lens to excite graphene on the ceramic substrate.42 The ceramic plate was placed on a sample holder and fixed with a linear actuator, where the substrate was scanned by moving the stage in the xy-direction.⁴² In essence, the use of the laser forms reduced graphene oxide when used within an open atmosphere. This is confirmed via XRD, Raman spectroscopy and XPS. To show the use of laser-induced reduced graphene oxide, the authors explore the sensing of acetaminophen, one of the most commonly used medications in the world, where they can measure over the range of 0.099-1978 µM, and a LoD of 5.2 nM is reported. They apply their sensor to the detection of acetaminophen in spiked river and urine samples, as well as within a pharmaceutical tablet; this approach provides a low-cost, environmentally friendly method, and offers large-scale manufacturing of reduced graphene oxide. In another example, the use of platinum decorated laserinduced graphene electrodes has been reported for the sensing of carbendazim.⁴⁸ Carbendazim is a fungicide that is regulated in many countries and banned in others. As such, there is a need for the sensing of carbendazim to ensure that high levels are avoided, which can result in potential health risks and

This article is licensed under a Creative Commons Attribution 3.0 Unported Licence. (cc) BY

Open Access Article. Published on 09 2024. Downloaded on 2025/6/1 20:59:50.

Table 2 An overview of the use of laser-induced graphene electrochemical sensors^a

Actoninoptiend (60) (64) (66) (66) (66) (66) (66) (66) (66) (66) (66) (66) (67) (66) (75) (67)	Analyte	Modification	Laser wavelength	Parameters (substrate/power/ speed)	Linear range	Limit of detection	Sample medium	Comments	References
	Acetaminophen	RGO	1064 nm	GO/30 W	0.099–1978 μM	5.2 nM	River, urine and	I	42
Model Model <t< td=""><td>4-Aminophenol</td><td>MWCNTs- polvaniline</td><td>405 nm</td><td>Kapton/2 W</td><td>0.1–55 µM</td><td>0.006 µM</td><td>Tap water and bharmaceutical</td><td>Ι</td><td>43</td></t<>	4-Aminophenol	MWCNTs- polvaniline	405 nm	Kapton/2 W	0.1–55 µM	0.006 µM	Tap water and bharmaceutical	Ι	43
Ibe add, interaction, bit addition, bit addition	Aristolochic acid and roxarsone	MoS ₂ /MXene	10.6 µm	CO_2 laser/PI/3.2 W/2.8 cm s ⁻¹	0.01-874 µM	16.5 and 2.31 nM	Human blood serum	Simultaneous sensing	44
Bit and utcle PNNs 10.6 µm P12.4-5.4 W/ 10-890, 05-56, 6.1, 0.07, and - Simultaneous ordinat capacity - 10.6 µm $0.7-23 \ln s^{-1}$ $and 1-63 \mu M$ $0.6 \mu M DPH$ Ta Bindrate analysis ordinat capacity - 10.6 µm C0, laser P(1,4) $2-30 \mu M DPH$ $0.6 \mu M DPH$ Ta Bindrate analysis inter analysis - 10.6 µm C0, laser P(1,4) $2-30 \mu M DPH$ $0.6 \mu M DPH$ Ta Bindrate analysis inter analysis - 10.6 µm C0, laser P(1,4) $1-30 \mu M DPH$ $0.6 \mu M DPH$ Ta Bindrate analysis inter analysis - 10.6 µm C0, laser P(1,4) $1-10 \mu M$ $0.6 \mu M DPH$ $0.6 \mu M DPH$ $0.6 \mu M DPH$ $0.6 \mu M DPH$ inter analysis - 10.6 µm C0, laser P(1,4) $1-10 \mu M$ $0.6 \mu M DPH$ $0.6 \mu M DPH$ $0.6 \mu M DPH$ inter analysis - 10.6 µm C0, laser P(1,4) $1-10 \mu M$ $0.6 \mu M DPH$ $0.6 \mu M DPH$ $0.6 \mu M DPH$ $0.6 \mu M DPH$	Ascorbic acid, dopamine and uric	I	10.6 µm	PI/2.4-5.4 W/ 0.7-23 in s ⁻¹	10–680, 0.5–32.5, and 0.5–19.5 µM	7.3, 0.27, and 0.22 μM		Simultaneous sensing	45
didant capacity - 10.6 μ CO, laser P(1,4) 2-900 μ M DPH Tea Indirect analysis inter - 10.6 μ CO, laser P(1,0) 5-35 μ M 1 μ M Beverges (cola Effection inter - 10.6 μ m CO, laser P(1,0) 5-35 μ M 1 μ M Beverges (cola Effection inter - 10.6 μ m CO, laser P(1,4) 10-160 μ M 0.07 μ M Beverges (cola Effection interval - 10.6 μ m P(0.84 W20 mm s^{-1} 1-40 μ M 0.07 μ M Beverges (chink, a 3.D)minterval beter while white, a 3.D)minterval beter white white a 3.D)minterval beter white a 3.D)minterval beter white a 3.D)minterval beter white 3.	actu Ascorbic acid, dopamine and uric acid	Pt NPs	10.6 µm	PI/2.4-5.4 W/ 0.7-23 in s ⁻¹	10–890, 0.5–56, and 1–63 μM	6.1, 0.07, and 0.22 μM	I	Simultaneous sensing	45
Indexime - 10.6 µm CO ₂ laser P(0.9) 5-35 µM 1 µM Berenges (cola Electrode indaxim Pt NPs - $W/20 \text{ mm s}^{-1}$ $1-40 \mu M$ $0.67 \mu M$ Berenges (cola Electrode amphenicol - $10.6 \mu m$ CO_2 laser $P(1/4)$ $1-0 \mu M$ $0.67 \mu M$ Wastewine, 3.37 printed cell amphenicol - $10.6 \mu m$ CO_3 laser $P(1/4)$ $10-160 \mu M$ $0.67 \mu M$ Wastewine, $3.75 \mu M$ $1.75 \mu M$ $1.0 \mu M$ Wastewine, $3.75 \mu M$ $1.75 \mu M$ $1.75 \mu M$ $0.2 \mu M$ $M M M M M$ $M M M M M M$ $M M M M M M$ $M M M M M M M M M M M M M M M M M M M $	Antioxidant capacity	I	10.6 µm	CO_2 laser PI/1.4 W/40 mm s ⁻¹	2-900 µM DPPH	0.6 µM DPPH	Tea	Indirect analysis by DPPH detection	46
Indazim Pt NP - Pt/I.38 W/4 cm s ⁻¹ 1-0 μ M 0.67 μ M Wasteward - amphenicol - 10.6 μ M CO3 laser Pt/1.4 10-160 μ M 0.67 μ M Wasteward - floxacin - 449 μ M Pt/0.38 W/3 cm s^{-1} 1-75 μ M 0.2 μ M Milk Hypercube 3D floxacin - 449 μ M Pt/0.34 W/20 mm s^{-1} 1-75 μ M 0.2 μ M Milk Hypercube 3D floxacin - 600 m s^{-1} 0.001 n g mL ⁻¹ Huma serum - - mine PtANNPs/PDMS - CO2 laser/Pt/3.2 0.001-200 ng 0.001 ng mL ⁻¹ Huma nerum - DNA apamer - CO2 laser/Pt/3.5 5.0 μ M 75 nM Huma nerum - - Rec PEDOT/Au NPs/ 450 nm Pt/5.5 W 5.1 M -2 mM 7.5 nM Attificial sevet Nao Pro-III Sec Chioan/GOx 450 nm Pt/5.5 W 5.1 M -2 mM 0.41 m m - - Sec <td>Atropine</td> <td>I</td> <td>10.6 µm</td> <td>CO₂ laser PI/0.9 W/20 mm s⁻¹</td> <td>5-35 µM</td> <td>1 µM</td> <td>Beverages (cola soft drink, energetic drink, beer, white wine, and whiskv)</td> <td>Electrode assembly within a 3D-printed cell</td> <td>47</td>	Atropine	I	10.6 µm	CO ₂ laser PI/0.9 W/20 mm s ⁻¹	5-35 µM	1 µM	Beverages (cola soft drink, energetic drink, beer, white wine, and whiskv)	Electrode assembly within a 3D-printed cell	47
amphenicol - 10.6 μ m CO ₂ laser PI/1.4 10-160 μ M 1.0 μ M Honcy - floxacin - 449 μ m W/40 $mm s^{-1}$ 1-75 μ M 0.2 μ M Milk Hypercube 3D floxacin - 449 μ m V/40 $mm s^{-1}$ 1-75 μ M 0.2 μ M Milk Hypercube 3D fine Pt-MUPS/PDMS - 0.001 η gmL ⁻¹ Human serum - - mine Pt-MUPS/PDMS - CO ₂ laser, PI/7.6 5-30 μ M 75 n M Human serum - - mine Pt-MUPS/PDMS - CO ₂ laser, PI/7.6 5-30 μ M 75 n M Human urine - - fine Pt-MUPS/PDMS - CO ₂ laser, PI/7.6 5-30 μ M 75 n M Human urine - - for W/2.0 $mm s^{-1}$ mL ⁻¹ 75 m M 75 m M Human urine - - for W/2.0 $mm s^{-1}$ mL ⁻¹ 5 m M 2 μ M Artificial sweat Nano Pro-111	Carbendazim	Pt NPs		$PI/1.38 W/4 cm s^{-1}$	1-40 µM	0.67 µM	Wastewater		48
floxactin - 449 nm $PI0.84 W/20 mm s^{-1}$ $1-75 \mu$ M 0.2μ M Milk Hypercube 3D nine ZnFe_O4 NPs/ 10.6 µm CO_2 laser/PU3.2 0.001-200 ng 0.001 ng mL^{-1} Human serum a visible laser mine PrAuNPs/PDMS - CO_2 laser/PU3.2 0.001-200 ng 0.001 ng mL^{-1} Human serum - wisible laser W/2.8 cm s^{-1} mL^{-1} ML^{-1} Human serum - - see PEDOT/Au NPs/ 450 nm V/2.8 cm s^{-1} mL^{-1} M Artificial sweat Nano Pro-III see Chitosan/GOx 450 nm PU/5.5 W 0-8 mM 0.431 mM - - see GOX 10.6 µm PU/7 W 0-8 mM 0.431 mM - - - see GOX 10.6 µm PU/7 W 0-8 mM 0.431 mM - - - - see GOX 10.6 µm PU/7 W 0-8 mM 0.431 mM - - -	Chloramphenicol		10.6 µm	CO ₂ laser PI/1.4 W/40 mm s ⁻¹	10–160 µM	1.0 µM	Honey		49
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Ciprofloxacin	I	449 nm	$PI/0.84 W/20 mm s^{-1}$	1–75 µM	0.2 µM	Milk	Hypercube 3D printer with a visible laser source	13
nePt-AuNPs/PDMS- $CO_2 laser: Pl/7.6$ $5-30 \ \mu M$ $75 \ nM$ Human urine-PEDOT/Au NPs/450 nm s^{-1} $W/200 \ mm s^{-1}$ $5 \ \mu M - 2 \ mM$ $2 \ \mu M$ Artificial sweatNano Pro-IIIGOX450 nm $Pl/5.5 \ W$ $5 \ \mu M - 2 \ mM$ $2 \ \mu M$ Artificial sweatNano Pro-IIIGOX450 nm $Pl/7 \ W$ $0-8 \ m M$ $0.431 \ m M$ $-$ Laser engravingGOX10.6 μm $CO_2 laser/paper/5 \ W/150 \ m s^{-1}$ $0.431 \ m M$ $-$ Laser engravingPt NPs GOX10.6 μm $CO_2 laser/paper/5 \ W/150 \ m s^{-1}$ $0.003-2.1 \ m M$ $0.13 \ m M$ $ -$ Au NPs/Ni NPs10.6 μm $CO_2 laser/Pl0.03-4.5 \ m M0.023 \ \mu MSweat -Au NPs/Ni NPs10.6 \mu mCO_2 laser/Pl0.30-4.5 \ m M0.023 \ \mu MSweat -Au NPs/Ni NPs10.6 \mu mCO_2 laser/Pl0.30-4.5 \ m M0.023 \ \mu MSweat-$	¢Tn-I	ZnFe ₂ O ₄ NPs/ DNA aptamer	10.6 µm	CO_2 laser/PI/3.2 W/2.8 cm s ⁻¹	0.001–200 ng mL ⁻¹	$0.001~{ m ng~mL^{-1}}$	Human serum	I	50
PEDOT/Au NPs/450 nmPI/5.5 W5 μ M-2 mM2 μ MArtificial sweatNano Pro-IIIGOxGOx450 nmPI/7 W0-8 mM0.431 mM-Iaser printerGOx10.6 μ mCO ₂ laser/paper/5 W/1-15 mM0.13 mMPt NPs GOx10.6 μ mCO ₂ laser/paper/5 W/1-15 mM0.13 mMPt NPs GOx10.6 μ mCO ₂ laser/Pl0.003-2.1 mM300 nMSweatAu NPs/Ni NPs10.6 μ mCO ₂ laser/Pl0.03-4.5 mM0.23 μ MSweatAu NPs/Ni NPs10.6 μ mCO ₂ laser/Pl0-30 mM1.5 μ MSweat and cell-	Dopamine	Pt-AuNPs/PDMS	I	CO_2 laser. PI/7.6 $W/200 \text{ mm s}^{-1}$	5-30 µM	75 nM	Human urine	I	51
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Glucose	PEDOT/Au NPs/ GOx	450 nm	Pl/5.5 W	5 µM-2 mM	2 µМ	Artificial sweat and urine and fetal bovine serum	Nano Pro-III laser printer	52
GOx 10.6 μ m CO ₂ laser/paper/5 W/ 1–15 mM 0.13 mM - </td <td>Glucose</td> <td>Chitosan/GOx</td> <td>450 nm</td> <td>PI/7 W</td> <td>0-8 mM</td> <td>0.431 mM</td> <td>Ι</td> <td>Laser engraving machine</td> <td>53</td>	Glucose	Chitosan/GOx	450 nm	PI/7 W	0-8 mM	0.431 mM	Ι	Laser engraving machine	53
Pt NPs GOx 10.6 μ m Pl/9.6 W/150 mm s ⁻¹ 0.0003-2.1 mM 300 nM Sweat - Cu NPs 10.6 μ m CO ₂ laser/Pl 0.03-4.5 mM 0.023 μ M Saliva - Au NPs/Ni NPs 10.6 μ m CO ₂ laser/Pl 0-30 mM 1.5 μ M Sweat -	Glucose	GOx	10.6 µm	CO ₂ laser/paper/5 W/ 127 cm s ⁻¹	1–15 mM	0.13 mM			54
Cu NPs 10.6 μm CO2 laser/PI 0.03-4.5 mM 0.023 μM Saliva Au NPs/Ni NPs 10.6 μm CO2 laser/PI 0-30 mM 1.5 μM Sweat and cell	Glucose	Pt NPs GOx	10.6 µm	$PI/9.6 W/150 mm s^{-1}$	0.0003-2.1 mM	300 nM	Sweat	I	55
Au NPs/Ni NPs 10.6 μ m CO ₂ laser/PI 0–30 mM 1.5 μ M Sweat and cell –	Glucose	Cu NPs	10.6 µm	CO, laser/PI	0.03-4.5 mM	0.023 µM	Saliva	I	56
	Glucose	Au NPs/Ni NPs	10.6 µm	CO ₂ laser/PI	0-30 mM	1.5 µM	Sweat and cell		57

Access Article. Published on 09 2024. Downloaded on 2025/6/1 20:59:50.	This article is licensed under a Creative Commons Attribution 3.0 Unported Licence.
Open A	(cc)) BY

Table 2 (Contd.)

Analyte	Modification	Laser wavelength	Parameters (substrate/power/ speed)	Linear range	Limit of detection	Sample medium	Comments	References
Hydrogen peroxide	Ag NPs	I	Infra-red laser. 780 nm 5 mW	10-10 000 M	7.9 M	Milk	I	58
Hydrogen peroxide	Prussian blue	10.6 μm	CO_2 laser PI/0.9 W/20 mm s ⁻¹	1-200 µM	0.26 µM	Milk and mouthwash	Combination with batch- injection analysis	59
Ivermectin	1	10.6 µm	CO_2 laser $PI/0.9$ W/20 mm s ⁻¹	10–100 µM	1.6 µM	Tap water, synthetic urine and tablets	, 	60
Methane	Pd NPs/PVDF/ EMImTFSI	I	CO ₂ laser. PI/10% power/25 mm s ⁻¹	10–50 ppm	9.2 ppm			61
m6A-RNA and 5mC- ssDNA	Au NPs		PI/power at 30%, speed at 20%, and a focus height of 25 mm	0.01-10 M	2.81 and 9.53 pM	HeLa cells	Laser engraving	62
Neonicotinoids (elothianidin, imidacloprid, thiamethoxam and dinotefuran)	1	10.6 µm	PI/5.25 W/279 mm s ⁻¹	10-40 µM	823, 384, 338, and 682 nM	River water	Simultaneous sensing	63
Nitrite	COOH- MWCNTs/Au NPs	405 nm	Kapton/50 mm $\rm s^{-1}$	10–140 µM	Mtl 9.0	Tap water	1	64
Picric acid		10.6 µm	CO ₂ laser/cardboard/5 W/10 mm s ⁻¹	0.48–2.0 mM	Ι	I	Proof-of-concept	65
Potassium and ammonium ions	Ionophore membrane	10.6 µm	CO ₂ laser/PI/speed 7%/power 10% ¹	0.3 mM-150 mM	100 and 30 µM	Human urine		66
Uric acid and nitrite	I	449 nm	PI/0.84 W/20 mm s ⁻¹	10–100 and 10– 70 μM	0.07 and 0.27 µM	I	Hypercube 3D printer with a visible laser source	13
Uric acid	Ι	10.6 µm	CO ₂ laser/paper/850 W/30 mm s ⁻¹	10-250 µM	3.97 µM	Human urine	Ι	64
Sulfamethoxazole	2D hexagonal boron nitride	450 nm	$PI/5.2 W/0.5 cm s^{-1}$	0.5-362.5 μM	0.11 μM	Milk and lake water	Compared with HPLC	67
Thrombin	1-Pyrenebutyric acid/Aptamer	10.6 µm	$PI/0.81 W/5.8 cm s^{-1}$	1-100 pM	1 pM buffer, 5 pM in serum	Fetal calf serum	X-660 laser cutter platform	11
Tyrosine	I	10.6 µm	CO_2 laser PI/0.9 W/20 mm s ⁻¹	5-30 µM	1.5 µM	Artificial urine	I	68

Minireview

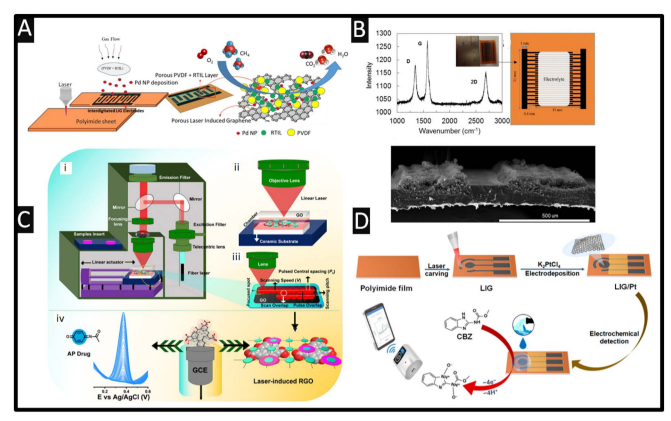


Fig. 3 (A) Schematic of the fabrication process of the electrochemical methane sensor. A CO₂ laser is used to convert the polyimide sheet into a patterned, interdigitated laser-induced graphene design. Palladium nanoparticles are decorated where porous polyvinylidene fluoride (PVDF) with 1-ethyl-3-methylimidazolium TFSI (EMImTFSI) in an *N*-methylpyrrolidone (NMP) layer is painted onto the electrodes. (B) Characterisation of laser-induced graphene electrodes with Raman spectra of the laser-induced graphene electrode material, photograph of electrodes prior to applying the electrolyte, a sketch of the interdigitated electrodes with dimensions and a SEM image of the cross section of the laser-induced graphene electrodes. Figures reproduced from reference ⁶¹. Copyright 2018 American Chemical Society. (C) Schematic setup of the fibre laser system (i), zoomed-in schematic showing the scanning of the laser (ii), laser scanning path (iii) and the electrochemical detection of the anti-pyretic (iv). Figure reproduced from reference ⁴². Copyright 2024 Royal Society of Chemistry. (D) A summary of a platinum modified laser-induced graphene electrochemical sensor and its application for the real-time detection of carbendazim in wastewater samples. Figure reproduced from ref. 48. Open Access.

environmental contamination.⁶⁹ The sensor developed for this application is shown in Fig. 3D, where a polyimide film is exposed to a laser and then modified using electrodeposition of platinum nanoparticles (70 nm diameter) onto the formed graphene. It shows a porous structure and has lattice defects as reported using SEM, Raman spectroscopy and XPS. This sensor was shown to be successful in the measurement of carbendazim with an LoD of 0.67 μ M with a linear range of 1–40 μ M, where good recoveries (88.89-99.50%) are shown in wastewater samples. In another study involving a facile approach to form laser-induced graphene electrodes, Kucherenko and co-workers used this as a solid-state ion-selective electrode for the simultaneous measurement of potassium and ammonium ions using an ionophore membrane.⁶⁶ This sensor provided a linear range of 0.3-150 mM and LoD of 100 µM and 30 µM for potassium and ammonium ions respectively. The authors used their sensor for the measurement of potassium and ammonium ions in urine to help monitoring hydration levels in different patients.

Others have shown that laser-induced graphene can be realised using paper-based electroanalytical devices. As shown in Fig. 4A, one can see that they can readily produce laserinduced graphene upon paperboard.65 This approach allows the researchers to produce laser-induced graphene on paperboard using a CO_2 laser, after which, they painted a reference electrode with conductive silver ink. This was used as a proof-ofconcept approach, which they tested towards the sensing of picric acid. The electrochemical reduction of this compound provides three distinct reduction processes at -0.35, -0.47, and -0.62 V (vs. silver) corresponding to the electroreduction of the nitro groups of picric acid, where the reduction wave at the lowest potential (-0.15 V) corresponds to the formation of a radical anion species.⁶⁵ This approach has been extended to develop a laser-induced graphene electrode for the sensing of uric acid, which gave a linear range of 10-250 µM with a LoD of 3.97 µM which was tested using spiked human urine.64 The use of laser-induced graphene electrodes has been also investigated for the measurement of nitrite, as shown in Fig. 4B.64 In this work, the use of a bare laser-induced graphene electrode has been initially explored using a redox probe of potassium ferricyanide/ferrocyanide with a reported electrochemical area

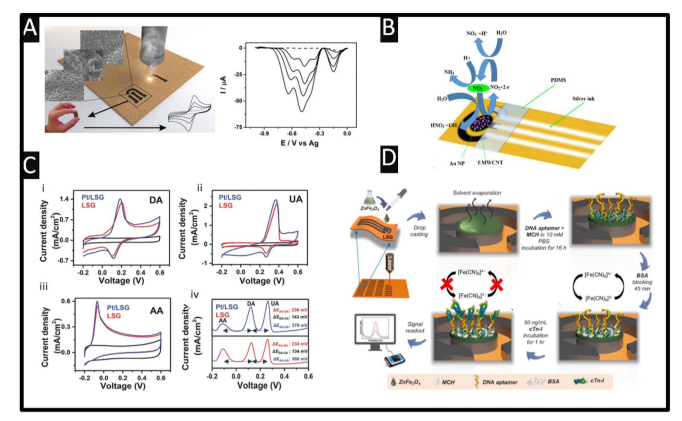


Fig. 4 (A) The use of laser-induced graphene formed upon paperboard. Also shown are the differential pulse voltammograms obtained towards the measurement of picric acid in the presence of 0.48, 0.91, 1.30, and 2.0 mM. Figure reproduced from ref. 65. Copyright 2017 Wiley. (B) An image showing the proposed mechanism of the electro-oxidation of nitrite using a laser-induced graphene electrode modified with COOH-MWCNT and with gold nanoparticles (all drop cast). Figure reproduced from ref. 64. Copyright 2021. (C) Cyclic voltammograms of laser-induced graphene and platinum laser-induced graphene electrodes in a pH 7.0 buffer containing (i) 1 mM dopamine (DA), (ii) 1 mM uric acid (UA) and (iii) 1 mM ascorbic acid (AA). Scan rate: 50 mV s⁻¹. Differential pulse voltammetry curve (iv) of laser-induced graphene and platinum laser-induced from ref. 45 Copyright 2016 Wiley. (D) Schematic illustration of the ZnFe₂O₄ modified laser-induced graphene cTn-I aptasensor. Fabrication of laser-induced graphene electrodes using the laser-scribing method and modification with ZnFe₂O₄ by drop-casting to prepare a laser-induced graphene/ZnFe₂O₄ electrode. A mixture of a thiol-modified DNA aptamer specific to cTn-I and mercaptohexanol (MCH) was immobilized on the electrode surface. MCH helps in the proper folding of the DNA aptamer. Incubation of BSA solution for 45 min to reduce non-specific adsorption. Different concentrations of cTn-I solution were incubated on the cTn-I aptasensor. The electrochemical signal was measured before and after the binding of cTn-I. Due to the attachment of cTn-I, the diffusion of the redox probe to the electrode surface was hindered, resulting in a decrease in the peak current values, directly correlated with the amount of cTn-I molecules attached to the aptasensor surface. Figure reproduced from ref. 50. Copyright 2021 Elsevier.

of 7.62 mm². The use of gold nanoparticles (12.9 nm diameter), MWCNTs and gold nanoparticles/COOH functionalised MWCNTs (drop cast) increase the electrochemical area to 9.27, 9.45 and 13.60 mm², respectively. Furthermore, the authors report that the charge transfer resistance decreased from 2911 Ω using a bare laser-induced graphene electrode to 448 Ω with gold nanoparticles/COOH functionalised MWCNTs. These electrode configurations are explored toward the electrochemical oxidation of nitrite, where the bare induced graphene electrodes gave rise to a peak observed at +0.68 V which moved to +0.75 V using the gold nanoparticles/COOH functionalised MWCNTs, but results in a larger peak height which will provide a useful response for the sensing of nitrite. The COOH-MWCNTs/gold nanoparticle modified laser-induced graphene electrodes are shown to report a linear range of 10-140 µM towards nitrite and an LoD of 0.9 µM. They applied this to

spiked tap water samples reporting recoveries of 91.5–96.1%, the mechanism of the electrochemical oxidation of nitrite is summarised in Fig. 4B.

Fig. 4C shows the use of laser-induced graphene and platinum nanoparticle laser-induced graphene electrodes in the presence of 1 mM dopamine (DA), 1 mM uric acid (UA) and 1 mM ascorbic acid (AA) recorded separately.⁴⁵ Analysis by the authors showed that the electrochemical area corresponds to 9.177 mm² for laser-induced graphene and in comparison a value of 10.383 mm² is reported for the platinum nanoparticle laser-induced graphene electrodes, which increased the area by 13.14%.⁴⁵ There is very little difference between the laserinduced graphene and platinum laser-induced graphene electrodes, which questions the need of carrying out such surface functionalisation with platinum. Interestingly, the separation between dopamine, uric acid and ascorbic acid is feasible where the peaks are well-resolved with and without Pt, as shown in Fig. 4C. This approach is summarised by the authors due to (i) rapid electron transport properties of a porous graphene network, (ii) rich in edge plane like sites/defects, which can provide catalytic activity, and (iii) the 3D hierarchical and porous network providing enough accessibility to biomolecules.⁴⁵

Another notable study is reported by Zeng and co-workers67 who produced a laser-induced graphene surface modified with 2D-hexagonal boron nitride via drop-casting. This sensor has been used in the measurement of sulfamethoxazole, which occurs via an electrochemical oxidation mechanism and undergoes one electron and one proton loss, giving a linear range of 0.5-362.5 µM and an LoD of 0.11 µM. The repeatability has been explored using 20 successive measurements, and the RSD was found to be 4.01%. In terms of the number of electrodes, the authors explored 10 different electrodes, which gave a relative standard deviation (RSD) of 4.65%. The selectivity towards the measurement of sulfamethoxazole is explored where they added in NaCl, KCl, MgCl₂, and CaCl₂ at 100-fold excess concentration and glucose with 10-fold excess concentration to 50 µM sulfamethoxazole. Also, they studied glutamic acid, arginine, and hypoxanthine with 1-fold excess concentration. All these compounds did not affect the measurement of sulfamethoxazole with RSD values less than 5%. This sensor was shown to be successful in the measurement of lake water and milk samples. In this work, while water samples collected from Jiangxi Agricultural University were processed using a 0.22 µm filter to remove the suspended particles, the commercial milk underwent an extraction procedure in a buffer/acetonitrile 50% (v/v) solution before analysis. Subsequent samples spiked with 27 and 55 µM sulfamethoxazole were analysed by HPLC and electrochemistry. Both produced recoveries in the range of 97.5-108.2%, with RSD values below 5% which showed there was no significant difference between the two methods, which also suggests a promising practical application in spiked sample analyses of the sensor.⁶⁷ It is important to highlight that validation of electrochemical sensors using laboratory-based instrumentation is of critical importance when such sensing devices are applied for real sample analysis.

Laser-induced graphene biosensors

Laser-induced graphene biosensors have been extensively reported, where the laser-induced graphene does not contribute to the sensing of the biomolecules under investigation but instead supports the use of nanoparticles which can increase active sites, enhance the number of immobilised biomolecules, and improve the electroanalytical response of the sensor. For example, a laser-induced graphene surface has been utilised as the basis of an aptasensor directed toward the sensing of cardiac troponin-I (cTn-I) proteins that are a clinically validated biomarker for the detection of acute myocardial infarction – as shown in Fig. 4D which shows how they fabricated the sensor.⁵⁰ In this approach, $ZnFe_2O_4$ is synthesised using a wet chemical methodology and drop cast onto the laser-induced graphene surface which is allowed to dry. Next, DNA aptamer (thiol

modified Anti-cTn-I: CGTGCAGTACGCCAACCT TTCTCATGCG CTGCCCCTCTTA) and mercaptohexanol are incubated for 16 h, after which bovine serum albumin is applied for 45 min to reduce non-specific adsorption. This approach uses the redox probe potassium ferricyanide/ferrocyanide, where the cTN-I binds with the surface DNA aptamer and blocks the surface sites causing the signal to decrease. A linear range of 0.001-200 ng m L^{-1} and an LoD of 0.001 ng m L^{-1} are shown to be feasible. Note that cTn-I levels below 0.6 ng mL⁻¹ present in serum are considered normal, while 0.7-1.4 ng mL⁻¹ suggest a minor myocardial injury and over 1.5 ng mL^{-1} indicate myocardial necrotic damage suggesting that this sensor can discriminate between a normal situation and myocardial necrotic damage. This sensor was explored in the presence of potential interference from cholesterol, glucose, myoglobin, cTn-T, and cTn-C and compared to the response for cTn-I. The sensor was explored toward 8.8, 8.2, 7.3, 6.5 and 5.9 fold excess for the same concentration (50 ng mL⁻¹) of each analyte, respectively.⁵⁰ The only interferents were cTn-C and cTn-T, which showed more adsorption onto the sensors, but the authors assure us that their sensor is highly selective for the measurement of cTn-I; potentially a change in blocking solution would benefit this work. This sensor was shown to be used for the measurement of cTn-I in spiked human serum using different amounts of cTn-1 of 0.001, 0.01, 0.1, 1, 10 and 100 ng mL⁻¹. The authors concluded that the performance of their sensor is a result of the use of ZnFe₂O₄ increasing the electroactive surface area and electrocatalytic activity where the laser-induced graphene electrode surface acts as "electrically wiring", where presumably other electrodes could be used; as shown by the authors⁵⁰ there are more sensitive approaches with very low linear ranges \sim pg mL^{-1} levels using other carbon substrates (e.g., screen-printed carbon electrodes).70

Other approaches have developed biosensors for the measurement of glucose using laser-induced graphene electrodes, as shown Table 2. There is still an obvious need for the measurement of glucose that can provide rapid, on-the-spot readings within human blood. The concentration of glucose needs to be in the normal human body range, where a reduction and excess can lead to hypoglycemia and hyperglycemia, respectively. For example, a linear range of 5 µM-2 mM and an LoD of 2 µM using poly(3,4-ethylene dioxythiophene) (PEDOT) and gold nanoparticles (20 nm diameter) with glucose oxidase upon a laser-induced graphene electrode have been reported.52 As shown in Fig. 5A, 3,4-ethylene dioxythiophene (EDOT) monomers are electrochemically polymerized on the laserinduced graphene electrode surface to form a PEDOT film. Next, gold nanoparticles are electrochemically deposited on the laser-induced graphene electrode surface by potentiostatic deposition.52 This is then modified with 11-mercaptoundecanoic acid (MUA) onto which a mixed solution of glucose oxidase (GOx) and glutaraldehyde (GA) is drop cast. Using the Randles-Š evčík equation they deduced the effective electrochemical area to be 4 times greater than the geometric area, which gave rise to the performance of the sensors. This sensor was used to sense glucose in artificial urine, fetal bovine serum and artificial sweat, where the recoveries of glucose with different

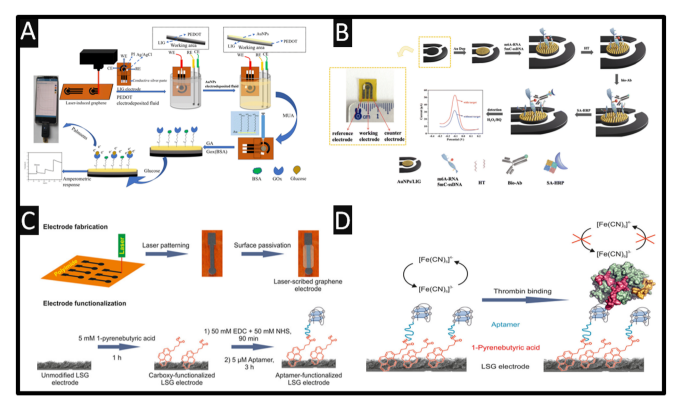


Fig. 5 (A) An overview of the glucose sensing analysis. Figure reproduced from ref. 52. Copyright 2024 Royal Society of Chemistry. (B) Schematic illustration of the biosensor modification process showing how the sensor works. Figure reproduced from ref. 62. Copyright 2024 Royal Society of Chemistry. (C) Electrode fabrication and functionalization process. (D) Schematic of the electrochemical thrombin detection mechanism with increasing concentrations of thrombin, and the diffusion of the redox marker hexacyanoferrate(III) to the electrode surface is hampered. This results in a decrease in the peak currents obtained from voltammetric measurements. Figures reproduced from ref. 11. Copyright 2017 American Chemical Society.

concentrations are 61.34-111.93%, 81.72-159.66% and 72.68-125.00% respectively.52 Non-enzymatic glucose-based laserinduced graphene sensors outperform enzymatic ones when one considers the cost, sensitivity, stability, and of course, operating duration. To this end, laser-induced graphene modified with copper nanoparticles gives rise to the measurement of glucose reporting a sensitivity of 2665 μ A mM⁻¹ cm⁻², a limit of detection of 0.023 µM and a linear range of 0.03-4.5 mM.⁵⁶ Another notable study reported laser-induced porous graphene which has been fabricated and decorated with nickel and gold nanoparticles towards the sensing of glucose, reporting a high sensitivity of 3500 μ A mM⁻¹ cm⁻², a linear range of 0–30 mM, a LoD of 1.5 μ M and a response time of less than one second.⁵⁷ Interestingly, the authors applied their sensor to the measurement of glucose through sweat providing a flexible sensor attached to the arms of a human subject. The authors consider the response of sensing in a flat or bending state reporting excellent mechanical durability using up to 500 cycles of bending and electrochemical stability through immersion in phosphate buffer solution for up to 30 days. Using a commercial sensor, they validated their laser-induced graphene-based sensor within sweat, demonstrating that their non-enzymatic laser-induced graphene glucose sensor could also be integrated with wireless measurement units to provide real-time monitoring capabilities.57

The use of laser-induced graphene biosensors has also been reported for the detection of methylation in DNA and RNA which is essential for the diagnosis and treatment of a wide range of diseases.⁶² As shown in Fig. 5B, a biosensor for the measurement of N⁶-methyladenosine (m6A-RNA) and 5methylcytosine-single strand DNA (5mC-ssDNA) is designed and implement in HeLa cells. This approach utilised electrochemically deposited gold nanoparticles, sulfhydryl-modified nucleic acid chains, biotin-modified antibodies, and streptavidin-modified horseradish peroxidase (SA-HRP), where the peak current/signal is proportional to the concentration of m6A-RNA and 5mC-ssDNA in the hydrogen peroxide-hydroquinone (H₂O₂-HQ) system. This sensor allowed the measurement across the linear range of 0.01-10 nM with an LoD of 2.81 pM for m6A-RNA and 9.53 pM for 5mC-ssDNA. It is reported by the authors that the method utilises gold-sulfur bonding to immobilise the detection target, which improves the conductivity of the laser-induced graphene electrode and introduces an amplified portion of the signal by taking advantage of antigenantibody specific binding.62 The stability of the biosensor was explored when they stored their sensor in a refrigerator (4 °C) for 0, 3 and 5 days and showed that the %RSD decreased by 9.42 for the 5-day detection, suggesting that the biosensor should not be stored for more than 5 days. This sensor was evaluated for the measurement of m6A-RNA and 5mC-ssDNA within

spiked HeLa cells where they report recoveries of 105.4–111.0% and 95.3–98.8% respectively with a low %RSD of 0.8–1.3% and 0.7–1.3% respectively. This approach has potential to be explored further for point-of-care techniques for nucleic acid methylation detection in clinical applications.⁶²

An outstanding approach using laser-induced graphene electrodes involves the use of an X-660 laser cutter platform with a wavelength of 10.6 μ m – see Fig. 5C.¹¹ These electrodes are modified with 1-pyrenebutyric acid for 1 h, which forms *via* π -stacking interactions. This then provides COOH-groups upon the electrode surface which allows covalent coupling using EDC/NHS with the antithrombin aptamer (5'-H₂N-C6-GGTTGGTGGTGGGTTGG-3') – see Fig. 5D. This sensor is used for thrombin detection, where increasing concentrations of thrombin stop the diffusion of the redox marker potassium hexacyanoferrate(m) to the electrode surface and produce a decrease in the peak currents. This sensor is shown to report

a linear range of 1–100 pM with a low LoD of 1 pM in buffer and 5 pM in fetal calf serum. The sensor has a reported incubation time of 30 min which is shorter than others reported and the LoD compares favourably.¹¹

Comparison of laser-induced graphene electrodes with screenprinted and additive manufactured electrodes

Screen-printed electrodes are a consolidated technology applied to the development of electrochemical sensors while additive manufacturing has emerged in the last ten years as a potential technology for the same purpose with additional advantages, which is also valid for laser-induced graphene electrodes. Table 3 summaries the advantages and disadvantages of laser-

	Laser-induced graphene electrodes	Additive manufactured electrodes	Screen-printed electrodes
Advantages	• Economical	• Economical	• Economical
	 No need for screens to define the working electrode and associated electrodes 	 No need for screens to define the working electrode and associated electrodes 	• Flexible/bendable electrodes
	• Flexible/bendable electrodes	• Can provide flexible/bendable electrodes based on the filament that is used	• Bulk modified
	• Control of surface morphology	• 3D electrodes can be easily realised	• Abundant edge plane sites/defects
	 Abundant edge plane sites/defects 	• Bulk modified	• Multiple sensing
	• Multiple sensing	• Control of surface morphology	• Different electrodes surface can be easily made (gold, silver, CNTs <i>etc</i>)
Disadvantages	• Highly reproducible	• Abundant edge plane sites/defects	• Comparable to gold standard laboratory approaches
	 Comparable to gold standard laboratory approaches 	• Multiple sensing	• Portable
	• Portable	• Different electrodes surface can be easily made (<i>e.g.</i> , bulk modified metal nanoparticles)	• Comparable to solid carbon electrodes
	 Comparable to solid carbon electrodes They are generally fabricated 	 Comparable to gold standard laboratory approaches Portable 	• Commercially available inks (standardisation is feasible)
	under ambient conditions; however, the inner gas environment can generate new morphologies and insert functional groups	• Comparable to solid carbon electrodes	
	• It cannot be bulk modified, so it needs surface-modification with nanoparticles, electrocatalysts/ mediators, metallic films (<i>e.g.</i> , gold, mercury, and bismuth), which can potentially limit their use	• Filaments need to be made to optimise for every application	• Screens are needed and need to be maintained
	• Difficult standardisation across groups/methods and batch-to-batch variation and power, speed and beam defocus need to be adjusted and depend on the laser source	• 3D printer parameters need to be evaluated for every filament	• Longer time for preparation due to ink curing time; every different material needs to be cured separately (<i>e.g.</i> , silver, carbon and insulating inks)
	 Conductive graphene tracks are easily removed by mechanical contact (electric contact requires extra attention) 	• In some cases, electrochemical surface activation is required	

Table 3 A summary of the advantages and disadvantages of laser-induced graphene electrodes compared with other relevant electrodes

Minireview

induced graphene electrodes in comparison to screen-printed electrodes^{39,71} and additive manufactured electrodes.^{72,73} We take every point in turn, and we comment upon each approach summarising the advantages and disadvantages of laser-induced graphene, screen-printed and additive manufactured electrodes.

Laser-induced graphene, additive manufactured and screenprinted electrodes are shown to be economical, reproducible, comparable to those obtained by standard laboratory approaches, and allow one to control the surface morphology. They are easy to combine with portable electroanalytical setups, and therefore allow one to transfer the laboratory approach into the field. They all have abundant edge plane sites/defects, which promote fast electron transfer and results in sensitive electroanalytical approaches. Furthermore, they can all be fabricated onto flexible materials and they are all electrochemically comparable to solid carbon electrodes. Additionally, new designs, such as microbands, microdiscs, microdisc arrays, etc. using a stencil can be manufactured and also they can be used for multi-analyte sensing. One downside of laser-induced electrodes is that these need to be modified with gold nanoparticles, and with electrocatalysts/mediators as a post-processing approach, which may limit their use, such as on using electrochemical deposition. In comparison, bulk modified additive manufactured and screen-printed electrodes can be readily produced where screen-printed inks are modified with the desired micro/nanomaterial (e.g., macro/nanosized silver, gold, palladium, CNTs, graphene) which reduces the resistance between the material and the electrode surface and improves electron transfer properties.39 Such an approach provides a simple but effective strategy allowing one to create massproduced electrochemical platforms.³⁹ Furthermore, screenprinted metallic electrodes can be realised through changing the ink, and one can readily manufacture different electrode types, for example silver, gold, palladium, etc. The downside with screen-printed electrodes is that one needs to design and use a screen which needs to be maintained,³⁹ while in the case of laser-induced graphene and additive manufactured electrodes, one does not need a screen. One aspect that also limits the use of additive manufactured electrodes, is that in some cases, electrochemical surface activation is required. Lastly, in all cases, these electrode configurations are comparable to solid carbon electrodes ($\sim 10^{-3}$ cm s⁻¹) where the k^0 compares well, suggesting that these fabrication approaches should be routinely used for sensing.

Conclusions

We have overviewed the use of laser-induced graphene electrodes as the basis of sensors and we can summarise as follows:

(1) The rapid and easy fabrication of laser-induced graphene sensors is an interesting approach for use as the basis of electroanalytical sensors. The electrodes are highly conductive, flexible, cost-effective, and have unique micro- and macrofeatures and eco-friendly production and the manufacturing of laser-induced graphene typically uses fewer chemicals and produces less waste than traditional graphene fabrication methods, making laser-induced graphene sensors more environmentally friendly, aligning well with sustainable manufacturing goals. Most importantly, these are multi-layer graphene, which contain using C–O/C=O *etc.* functional groups where edge plane sites/defects are favourable for electroanalysis.

(2) As shown throughout, there are many types of laserinduced graphene and there can be variability between different research groups and batch-to-batch variations.²⁴ The graphene is dependent upon the material/substrate, laser type and associated parameters and in future manufacturing laser maintenance, operational frequency and batch size should be considered.²⁴ In each case, these need to be physicochemically characterised using SEM, Raman spectroscopy, XPS and electrochemistry, helping to benchmark the performance of the laser-induced graphene electrodes. Another area is to vary the length of the electrical connection of laser-induced graphene electrodes, which has shown that a shorter connection give rise to improvement in the electrochemical response using screenprinted electrodes.⁴⁰

(3) The use of laser-induced graphene sensors is comparable with that of screen-printed and additive manufactured electrodes, but they have advantages and disadvantages which need to be considered – see Table 3 (*e.g.*, they cannot be bulk modified with catalysts). Laser-induced formation of catalystmodified graphene occurs through a single laser step; metal nanostructures have been incorporated as a post-treatment protocol by different strategies and steps to improve sensing properties of laser-induced graphene, which increases the time of preparation and decreases reproducibility.

(4) The use of laser-induced graphene sensors has been explored toward a range of useful analytes, but many do not validate their sensors with laboratory-based instrumentation which stops uptake and commercialisation.

(5) One needs to consider how do the different laser-induced graphene sensors compare with each other and how to choose the appropriate ones for a specific target application? The answer lies in the fabrication of laser-induced graphene sensors which is varied, providing different electrochemical based responses (see Table 1) where researchers need to physico-chemically characterise and decide whether their sensor is useful for direct electron transfer processes, or they need to modify the surface using catalytic nanoparticles and enzymes/ proteins, for instance.

Laser-induced graphene sensors have an optimistic future where researchers need to develop their sensors further exploring more challenging analytes in difficult media, *e.g.*, real blood, saliva, sewage, *etc.* New strategies to incorporate different metal-based catalysts within graphene through a single laser irradiation procedure (one-step laser protocol) may enable the formation of next generation electrochemical laser-induced sensing materials with improved selectivity and sensitivity. Polyimide has been the main substrate used to generate laser-induced graphene sensors; however, other substrates can be investigated and paper is likely a promising material considering the generation of sustainable electrochemical sensors. On the other hand, paper as a substrate for

View Article Online

laser-induced graphene sensors brings additional challenges to be overcome, aiming at high-performance benchmarked sensors. Lastly, the use of integration of laser-induced graphene sensors with internet of things (IoT) extends their use in smart infrastructure for real-time data gathering and responsive systems. Overall, we can state that laser-induced graphene sensors are likely to become increasingly versatile and accessible, supporting innovative solutions across health, environmental monitoring and smart infrastructure, to name just a few.

Data availability

Analytical Methods

No new data were generated or analysed as part of this review.

Conflicts of interest

The authors declare no competing interests.

Acknowledgements

RAAM is grateful to Conselho Nacional de Desenvolvimento Científico e Tecnológico (CNPq) for his fellowship (315838/2021-3).

References

- 1 A. R. Urade, I. Lahiri and K. S. Suresh, JOM, 2023, 75, 614-630.
- 2 E. Bernalte, R. D. Crapnell, O. M. Messai and C. E. Banks, ChemElectroChem, 2024, 11, e202300576.
- 3 N. Sharma, R. Dev Gupta, R. Chandmal Sharma, S. Dayal and A. Singh Yadav, Mater. Today: Proc., 2021, 47, 2752-2755.
- 4 R. Ramesh, in Graphene Production and Application, eds A. Sadia, M. S. Akhtar and S. Hyung-Shik, IntechOpen, Rijeka, 2020, Ch. 2, DOI: 10.5772/intechopen.92258.
- 5 J. Phiri, P. Gane and T. C. Maloney, Mater. Sci. Eng., B, 2017, 215, 9-28.
- 6 J. Lin, Z. Peng, Y. Liu, F. Ruiz-Zepeda, R. Ye, E. L. G. Samuel, M. J. Yacaman, B. I. Yakobson and J. M. Tour, Nat. Commun., 2014, 5, 5714.
- 7 M. Abdulhafez, G. N. Tomaraei and M. Bedewy, ACS Appl. Nano Mater., 2021, 4, 2973-2986.
- 8 A. C. Ferrari, J. C. Meyer, V. Scardaci, C. Casiraghi, M. Lazzeri, F. Mauri, S. Piscanec, D. Jiang, K. S. Novoselov, S. Roth and A. K. Geim, Phys. Rev. Lett., 2006, 97, 187401.
- 9 K. Avinash and F. Patolsky, Mater. Today, 2023, 70, 104-136.
- 10 N. Dixit and S. P. Singh, ACS Omega, 2022, 7, 5112-5130.
- 11 C. Fenzl, P. Nayak, T. Hirsch, O. S. Wolfbeis, H. N. Alshareef and A. J. Baeumner, ACS Sens., 2017, 2, 616-620.
- 12 Y. Chyan, R. Ye, Y. Li, S. P. Singh, C. J. Arnusch and J. M. Tour, ACS Nano, 2018, 12, 2176-2183.
- 13 W. R. P. Costa, R. G. Rocha, L. V. de Faria, T. A. Matias, D. L. O. Ramos, A. G. C. Dias, G. L. Fernandes, E. M. Richter and R. A. A. Muñoz, Microchim. Acta, 2022, 189, 185.
- 14 A. Behrent, C. Griesche, P. Sippel and A. J. Baeumner, Microchim. Acta, 2021, 188, 159.

- 15 F. Wang, K. Wang, B. Zheng, X. Dong, X. Mei, J. Lv, W. Duan and W. Wang, Mater. Technol., 2018, 33, 340-356.
- 16 H. Xiao, Y. Li, R. Chen, T. Xie, P. Xu, H. Zhu, J. He, W. Zheng and S. Huang, eScience, 2023, 3, 100134.
- 17 J. Zhang, C. Zhang, J. Sha, H. Fei, Y. Li and J. M. Tour, ACS Appl. Mater. Interfaces, 2017, 9, 26840-26847.
- 18 A. Kothuru and S. Goel, IEEE Trans. Electron Devices, 2022, 69, 1333-1340.
- 19 M. Wang, Y. Yang and W. Gao, Trends Chem., 2021, 3, 969-981.
- 20 Y. Huang, R. Yang and M. G. Li, Adv. Funct. Mater., 2024, 34, 2407503.
- 21 T. J. Davies, C. E. Banks and R. G. Compton, J. Solid State Electrochem., 2005, 9, 797-808.
- 22 D. A. C. Brownson, D. K. Kampouris and C. E. Banks, Chem. Soc. Rev., 2012, 41, 6944-6976.
- 23 K. M. Clark, D. T. Nekoba, K. L. Viernes, J. Zhou and T. R. Ray, Biosens. Bioelectron., 2024, 264, 116649.
- 24 Y. Tang, G. A. Moreira, D. Vanegas, S. P. A. Datta and E. S. McLamore, Micromachines, 2024, 15, 874.
- 25 K. Muzyka and G. Xu, Electroanalysis, 2022, 34, 574-589.
- 26 N. F. Santos, S. O. Pereira, A. Moreira, A. V. Girão, A. F. Carvalho, A. J. S. Fernandes and F. M. Costa, Adv. Mater. Technol., 2021, 6, 2100007.
- 27 R. R. A. Soares, R. G. Hjort, C. C. Pola, K. Parate, E. L. Reis, N. F. F. Soares, E. S. McLamore, J. C. Claussen and C. L. Gomes, ACS Sens., 2020, 5, 1900-1911.
- 28 H. Wang, Y. Wang, X. Cao, M. Feng and G. Lan, J. Raman Spectrosc., 2009, 40, 1791-1796.
- 29 Graphene Number of Layers Calculator From ID/IG and I2D/IG Ratio via Raman Spectroscopy – InstaNANO, https:// instanano.com/all/characterization/raman/graphene-layers/ , accessed June 2.
- 30 A. Lamberti, F. Perrucci, M. Caprioli, M. Serrapede, M. Fontana, S. Bianco, S. Ferrero and E. Tresso, Nanotechnology, 2017, 28, 174002.
- 31 X. Ji, C. E. Banks, A. Crossley and R. G. Compton, ChemPhysChem, 2006, 7, 1337-1344.
- 32 A. Chou, T. Böcking, N. K. Singh and J. J. Gooding, Chem. Commun., 2005, 842-844, DOI: 10.1039/B415051A.
- 33 R. S. Nicholson, Anal. Chem., 1965, 37, 1351-1355.
- 34 I. Lavagnini, R. Antiochia and F. Magno, Electroanalysis, 2004, 16, 505-506.
- 35 R. J. Klingler and J. K. Kochi, J. Phys. Chem., 1981, 85, 1731-1741.
- 36 A. García-Miranda Ferrari, C. W. Foster, P. J. Kelly, D. A. C. Brownson and C. E. Banks, Biosensors, 2018, 8, 53.
- 37 J. E. Randles, Trans. Faraday Soc., 1948, 44, 327-338.
- 38 A. Sevčík, Collect. Czech. Chem. Commun., 1948, 13, 349-377. 39 R. D. Crapnell and C. E. Banks, ChemElectroChem, 2024, 11,
- e202400370.
- 40 M. J. Whittingham, N. J. Hurst, R. D. Crapnell, A. Garcia-Miranda Ferrari, E. Blanco, T. J. Davies and C. E. Banks, Anal. Chem., 2021, 93, 16481-16488.
- 41 H. M. A. Amin, Y. Uchida, E. Kätelhön and R. G. Compton, J. Electroanal. Chem., 2021, 880, 114891.

- 42 K.-Y. Hwa, R. Murugan, S.-F. Tseng, A. Santhan and J.-Y. Lin, *Environ. Sci.: Nano*, 2024, **11**, 951–968.
- 43 S. Nasraoui, S. Ameur, A. Al-Hamry, M. Ben Ali and O. Kanoun, *Sensors*, 2022, 22, 833.
- 44 U. Rajaji, P.-S. Ganesh, S.-Y. Kim, M. Govindasamy, R. A. Alshgari and T.-Y. Liu, *ACS Appl. Nano Mater.*, 2022, 5, 3252–3264.
- 45 P. Nayak, N. Kurra, C. Xia and H. N. Alshareef, *Adv. Electron. Mater.*, 2016, **2**, 1600185.
- 46 N. l. I. Inoque, D. A. Araújo, D. M. de Lima, R. M. F. Sousa, T. R. L. C. Paixão and R. A. Munoz, *ACS Food Sci. Technol.*, 2024, 4(6), 1589.
- 47 T. A. Matias, D. L. Ramos, L. V. Faria, A. de Siervo, E. M. Richter and R. A. Muñoz, *Microchim. Acta*, 2023, **190**, 297.
- 48 L. Wang, M. Li, B. Li, M. Wang, H. Zhao and F. Zhao, *Foods*, 2023, **12**, 2277.
- 49 N. I. Inoque and R. A. A. Munoz, *Anal. Methods*, 2024, **16**, 6793.
- 50 S. Rauf, V. Mani, A. A. Lahcen, S. Yuvaraja, T. Beduk and K. N. Salama, *Electrochim. Acta*, 2021, **386**, 138489.
- 51 X. Hui, X. Xuan, J. Kim and J. Y. Park, *Electrochim. Acta*, 2019, **328**, 135066.
- 52 Z. Zhang, L. Huang, Y. Chen, Z. Qiu, X. Meng and Y. Li, *RSC Adv.*, 2024, **14**, 1034–1050.
- 53 K. Settu, P.-T. Chiu and Y.-M. Huang, *Polymers*, 2021, 13, 2795.
- 54 T. Pinheiro, S. Silvestre, J. Coelho, A. C. Marques, R. Martins,
 M. G. F. Sales and E. Fortunato, *Adv. Mater. Interfaces*, 2021,
 8, 2101502.
- 55 H. Yoon, J. Nah, H. Kim, S. Ko, M. Sharifuzzaman, S. C. Barman, X. Xuan, J. Kim and J. Y. Park, *Sens. Actuators, B*, 2020, **311**, 127866.
- 56 B. E. Nugba, A. A. El-Moneim, N. O. Mousa and A. Osman, *Carbon Lett.*, 2023, **33**, 1767–1780.
- 57 J. Zhu, S. Liu, Z. Hu, X. Zhang, N. Yi, K. Tang, M. G. Dexheimer, X. Lian, Q. Wang, J. Yang, J. Gray and H. Cheng, *Biosens. Bioelectron.*, 2021, **193**, 113606.
- 58 E. Aparicio-Martínez, A. Ibarra, I. A. Estrada-Moreno,
 V. Osuna and R. B. Dominguez, *Sens. Actuators, B*, 2019, 301, 127101.

- 59 T. A. Matias, L. V. de Faria, R. G. Rocha, M. N. Silva, E. Nossol, E. M. Richter and R. A. Muñoz, *Microchim. Acta*, 2022, **189**, 188.
- 60 D. C. Ferreira, N. I. Inoque, A. A. Tanaka, L. M. Dantas, R. A. Muñoz and I. S. da Silva, *Anal. Methods*, 2024, 16, 4136–4142.
- 61 M. Dosi, I. Lau, Y. Zhuang, D. S. A. Simakov, M. W. Fowler and M. A. Pope, *ACS Appl. Mater. Interfaces*, 2019, **11**, 6166– 6173.
- 62 J. Guo, M. Zhao, C. Chen, F. Wang and Z. Chen, *Analyst*, 2024, **149**, 137–147.
- 63 Z. T. Johnson, K. Williams, B. Chen, R. Sheets, N. Jared, J. Li, E. A. Smith and J. C. Claussen, *ACS Sens.*, 2021, **6**, 3063–3071.
- 64 S. Nasraoui, A. Al-Hamry, P. R. Teixeira, S. Ameur, L. G. Paterno, M. Ben Ali and O. Kanoun, *J. Electroanal. Chem.*, 2021, 880, 114893.
- 65 W. R. de Araujo, C. M. R. Frasson, W. A. Ameku, J. R. Silva, L. Angnes and T. R. L. C. Paixão, *Angew. Chem., Int. Ed.*, 2017, **56**, 15113–15117.
- 66 I. S. Kucherenko, D. Sanborn, B. Chen, N. Garland, M. Serhan, E. Forzani, C. Gomes and J. C. Claussen, *Adv. Mater. Technol.*, 2020, 5, 1901037.
- 67 Y. Zeng, Q. Li, W. Wang, Y. Wen, K. Ji, X. Liu, P. He, B. Campos Janegitz and K. Tang, *Microchem. J.*, 2022, 182, 107898.
- 68 T. A. Matias, R. G. Rocha, L. V. Faria, E. M. Richter and R. A. Munoz, *ChemElectroChem*, 2022, **9**, e202200339.
- 69 R. D. Crapnell, P. S. Adarakatti and C. E. Banks, *Anal. Methods*, 2023, **15**, 4811-4826.
- 70 H. Jo, J. Her, H. Lee, Y.-B. Shim and C. Ban, *Talanta*, 2017, 165, 442–448.
- 71 A. García-Miranda Ferrari, S. J. Rowley-Neale and C. E. Banks, *Talanta Open*, 2021, **3**, 100032.
- 72 M. J. Whittingham, R. D. Crapnell, E. J. Rothwell, N. J. Hurst and C. E. Banks, *Talanta Open*, 2021, 4, 100051.
- 73 M. J. Hossain, B. T. Tabatabaei, M. Kiki and J.-W. Choi, *Int. J. Precis. Eng. Manuf. Green Technol.*, 2024, DOI: 10.1007/ s40684-024-00629-5.