Graphene Plasmonic Fractal Metamaterials for Broadband Photodetectors

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(Dated: May 15, 2020)

Metamaterials have recently established a new paradigm for enhanced light absorption in stateof-the-art photodetectors. Here, we demonstrate broadband, highly efficient, polarizationinsensitive, and gate-tunable photodetection at room temperature in a novel metadevice based on gold/graphene Sierpinski carpet plasmonic fractals. We observed an unprecedented internal quantum efficiency up to 100% from the nearinfrared to the visible range with an upper bound of optical detectivity of 10^{11} Jones and a gain up to 10^6 , which is a fingerprint of multiple hot carriers photogenerated in graphene. Also, we show a 100-fold enhanced photodetection due to highly focused (up to a record factor of $|E/E_0| \approx 20$ for graphene) electromagnetic fields induced by electrically tunable multimodal plasmons, spatially localized in self-similar fashion on the metasur-Our findings give direct insight into the face. physical processes governing graphene plasmonic fractal metamaterials. The proposed structure represents a promising route for the realization of a broadband, compact, and active platform for future optoelectronic devices including multiband bio/chemical and light sensors.

Nowadays, the conversion of light into electrical signals is at the heart of several technologies. Notwithstanding the high level of maturity, the large scale, and diversity of application areas, the need for a compact photodetection platform with higher performance in terms of speed, efficiency, spectral range, as well as flexibility, semitransparency, and compatibility with complementary metal-oxide-semiconductor (CMOS), is becoming more eminent¹.

Such features can be greatly fulfilled by combining functional systems with metamaterials² (i.e. structured materials on the subwavelength scale with engineered electromagnetic properties). This leads to the concept of metadevices², a logical extension of the metamaterial paradigm, where interactions are nonlinear and the response is dynamic, using systems with spatially variable elements. Generally, such metadevices can operate at low voltages, which is a competitive advantage over conventional technology exploiting bulk and expensive electrooptical crystals, often non-integrated in silicon photonics. However, the bandwidth of such metamaterials is generally narrow due to their resonant nature and most of them are working in the infrared band. Mastering control in the optical regime is technologically a more challenging task, due to the expensive and demanding fabrication of nanostructures, resulting from the need of highresolution electron- or ion-beam techniques. Also, devices working in a wide spectral range are important for multiband photodetection³, on-chip sensing of multiple analytes⁴, as well as multiplexed fluorescence and Raman detection⁵. Hybridization of low-dimensional carbon allotropes with plasmonic metamaterials $^{6-16}$ is known to provide broadband and strong light-matter interactions, improvement of the nonlinear response due to the electromagnetic field enhancement given by the metamaterial, and compatibility with well-established silicon photonic and CMOS technologies^{17,18}, therefore realizing a platform for high-speed¹⁹ light modulation and detection on the same chip.

In this regard, graphene — a two-dimensional system composed of carbon atoms arranged in a hexagonal lattice — has rapidly established itself as intriguing building block for optoelectronic applications²⁰, with a strong focus on various photodetection frameworks¹. The versatility of this exceptional material enables its application in a number of areas including ultrafast and electrically tunable detection of light over a wide spectral range from UV to $\text{THz}^{3,9,10,13,14,17-19,21-30}$. However, single-layer graphene absorbs 2.3% of the incident light in a large spectral range³¹, which is remarkably high for an atomically thin material, but it leads to a low photocarrier absorption efficiency that constitute a limiting factor for highly sensitive state-of-the-art photodetectors. By coupling plasmonic metamaterials with graphene, localized surface $plasmons^{32}$ (LSP) — collective charge oscillations at the interface between a subwavelength metal object and a dielectric — can be excited in photodetectors to improve graphene absorption by the resulting field enhancement, therefore increasing the device responsivity. However, plasmonic structures with periodic geometries can be used only for filtering light polarization and selective absorption at designed frequencies 3,9,10,14,33 , restricting their application field. Recently, the first graphene plasmonic fractals^{6,7} have been realized, reporting a polarization-insensitive, gate-tunable, and enhanced photodetection. So far, such metamaterials are still narrow-band and have low photodetection efficiency with poor electrical modulation.

Here, we show that plasmonic fractal metamaterials can be designed to significantly improve broadband and polarization-independent light absorption in graphene photodetectors, thus enabling multiband photodetection. A novel graphene-based electrically active metadevice for efficient control and detection of radiation in the visible (VIS)-mid-infrared (MIR) range was realized. The metamaterial consists of an Au Sierpinski carpet (SC) fractal³⁴ fabricated on a graphene/SiO₂/Si substrate. The gold/graphene (Au/G) fractal absorbs light 25 times stronger than an Au SC. As a result of the strong electromagnetic field enhancement (up to $|E/E_0| \approx 20$) within the interstructure gaps, the graphene-light interaction is greatly enhanced (up to an unprecedented factor of $|E/E_0|^4 \approx 10^5$) at designed frequencies. This provides a multimodal plasmonic response that covers a broad range of the electromagnetic spectrum (VIS-MIR). Owing to the continuous nature of the Au/G fractal metamaterial, both its absorption intensity and bandwidth can be controlled electrically by varying the charge carrier concentration in graphene, through a gate voltage. This approach allows modulation of the device optical reflectance up to 23% in intensity and 60% in photon energy, much higher than that reported by Aygar et al.⁶. We observed in the optical range up to 100% internal quantum efficiency, remarkably higher than that reported by Fang et al.⁷, with a gain up to 10^6 and an upper bound of optical detectivity of 2×10^{11} cm Hz^{1/2}W⁻¹. Moreover, we present up to 100-fold enhancement of the responsivity, with respect to an unpatterned graphene photodetector.

RESULTS AND DISCUSSION

Deterministic fractals³⁵ are self-similar objects generated by geometrical rules, having a non-integer (Hausdorff-Besicovitch) dimension. Sierpinski carpets can be generated by a recursive geometrical algorithm employing a Lindenmayer system implementation³⁴. We designed Au/G SCs starting from a graphene unit cell of side $L_0 = 10 \ \mu \text{m}$ that is divided into a 3×3 array of sub-cells of lateral size $L_1 = L_0 3^{-1}$, with an Au square placed in the central sub-cell. By iteratively applying the same rule to the generated sub-cells of size $L_t = L_0 3^{-t} = L_0 \mathcal{L}_t$, it is possible to obtain fractals for higher orders t of complexity³⁴. In this way, we indirectly patterned graphene, thereby realizing an Au/G fractal metamaterial that can be thought as a discrete Au SC and a complementary continuous graphene SC. At each iteration the side of the sub-cells is reduced by a factor $\mathcal{L} = 3^{-1}$. Since the number of graphene sub-cells in the SC increases by a factor $\mathcal{N} = 8$ at each iteration, the fractal dimension is $d_H = \log \mathcal{N} / \log \mathcal{L}^{-1} \approx 1.89^{34}$.

By employing electron-beam lithography, reactive-ion etching, metal evaporation, and lift-off techniques (see Methods), we fabricated the first five orders of an Au/G SC on a Si/SiO₂ substrate (see Supplementary Information 1). The nominal thickness of the Au squares is 35 ± 3 nm, while their lateral sizes are $L_1 = 3.33 \pm 0.05 \ \mu$ m, $L_2 = 1.11 \pm 0.01 \ \mu$ m, $L_3 = 370 \pm 17 \ nm$, $L_4 = 123 \pm 7 \ nm$, and $L_5 = 41 \pm 3 \ nm$. A representative optical micrograph of the sample for t = 5 order is shown in Figure 1a along with a scanning electron microscopy (SEM) micrograph in Figure 1b, which illustrates the excellent homogeneity and uniformity of the fractal structure. In Figure 1c, a scheme of an Au/G fractal photodetector is depicted. In order to understand the physics governing the metadevice, before presenting the device characterization we discuss the optical properties of the Au/G fractal metamaterial in the following.

As we have previously reported in Ref. 34, the Au SC can be regarded as a metallo-dielectric photonic quasicrystal. In Figure 1d the singular-continuous³⁶ reciprocal lattice of the SC for t = 5 order is shown, by computing the fast Fourier transform of its SEM micrograph. Such a reciprocal lattice is self-similar as its direct counterpart³⁴. It follows that the diffraction patterns and the related optical spectra of such fractals are themselves self-similar^{34,36}. Also, the SC has a larger number of points in the reciprocal space than a periodic array, since its fractal reciprocal lattice is a superposition of five periodic lattices with different constants $a_t = 3L_t$, under the periodic approximation^{34,36,37}.

In Figure 2a, a comparison between the reflectance rof an Au and Au/G SC divided by the reflectance r_{sub} of the SiO_2/Si substrate is shown. The reflectance of the substrate is dominated by a minimum around 0.7eV due to thin-film interference effects between SiO₂ and Si. The spectra of Au and Au/G SCs exhibit multiple features in the VIS–MIR range, which for the Au SC appear as maxima ω_n' due to the radiative scattering in air, while for the Au/G SC result in shifted minima ω_n owing to the thin-film interference effects between Au and graphene. In Ref. 34, we have reported that the optical spectrum of an Au SC at a given fractal order tmainly manifests t diffraction-mediated LSP resonances $\omega_n \approx 2\pi c/3L_0\mathcal{L}_n$, where $n = 1, 2, \ldots t$, for small LSP wavevectors k_p . For large k_p , $\omega_n \equiv \omega_{sp}/\sqrt{3} = 2.40 \text{ eV}$ with $\omega_{sp} = 4.17 \text{ eV}$ the Au surface plasmon³² (see Supplementary Information 2). These LSP resonances are self-similar as their photon energy has the same scaleinvariance law $(\mathcal{L}_t = 8^{-t/d_H})$ as the SC lattice constant a_t , depending on the fractal dimension d_H (see Supplementary Information 2). On the other hand, graphene surface plasmons polaritons are expected at lower photon energy³⁸. Furthermore, the relative reflectance change $(R_{Au} - R_{Au/G})/R_{Au}$ between the two samples is up to 25% at ω_3 , which means a remarkable relative absorption variation $(R_{Au/G} - R_{Au})/(1 - R_{Au})$ of 2500%. Therefore, graphene can be exploited as an ultrathin, broadband, anti-reflective coating. From the comparison with the reflectance spectra of Au/G periodic arrays in Figure 2b, it is possible to observe that the energy of the Au/G SC resonances scales with the periodicity a_t of the



FIG. 1. Au/G plasmonic fractals. **a**, False color optical microscopy micrograph of a 35 ± 3 nm thick Au SC for t = 5 order (in yellow) deposited on a CVD-grown, single-layer, single-crystal graphene (in blue) transferred on a Si/SiO₂ substrate (in gray). Four leads of the photodetector device are also shown (in yellow). **b**, Scanning electron microscopy micrograph of the Au/G SC. **c**, Scheme of the Au/G photodetector realized by Blender software, www.blender.org. **d**, Fast Fourier transform of the SEM micrograph in **b**. The $\Delta = \overline{\Gamma X}$ and $\Sigma = \overline{\Gamma M}$ directions in the fractal reciprocal lattice are marked along with the first pseudo-Brillouin zones $[-\pi/a_t, \pi/a_t]^2$ of the different fractal orders. Starting from the origin, each point of the SC reciprocal lattice is defined by a reciprocal lattice vector $G_{ij}^{(t)} = 2\pi\sqrt{i^2 + j^2}/a_t$.

Au structures. Also, the SC resonances in Figure 2a are blue-shifted with respect to those of the periodic arrays in Figure 2b, due to a far-field hybridization of LSPs belonging to the different fractal orders³⁴. In particular, the ω_3 mode overlaps with the SiO₂ minimum. From data in Figure 2b, we can draw the dispersion relation of the hybrid Au/G LSP modes in Figure 2c. From the electromagnetic calculations (see Supplementary Information 3), $k_p = 2\pi/L_t = 6\pi/a_t$ is the self-similar LSP wavevector of the Au SC. From the plot we observe that the out-of-plane coupling between the Au SC LSP modes and graphene is two times stronger than the in-plane coupling between the LSP modes of the Au SC squares (see Supplementary Information 2).

Moreover, the Au SC interaction with graphene allows to tune the Au/G LSP modes by applying a gate voltage to the device. The intensity of the relative reflectance to the charge neutrality point (CNP), $(R - R_{CNP})/R_{CNP}$, for the five LSPs can be enhanced up to 23% (2d). Also, we noticed that only the position in energy of the $2E_F$ graphene interband threshold (Figure 2d inset) can be modulated up to 60%, following the law $2E_F = 2\hbar v_F \sqrt{\pi n}$, being $n \approx C (V_G - V_{CNP})$ the charge carrier density, $C = \varepsilon_{SiO_2} \varepsilon_0 / et_{ox} = 7.2 \times 10^{10} \text{ V}^{-1} \text{ cm}^{-2}$ the SiO₂ gate capacitance per area, V_G the gate voltage, $V_{CNP} = 30$ V the voltage at the charge neutrality point, and $v_F = 10^8$ cm/s the graphene Fermi velocity. We did not observe any shift in energy for the the other LSP modes for such values of applied gate voltage. The dispersion of ω_3 is shown in Figure 2d, for instance.

Additionally, in Ref. 34 we demonstrated that having a centrosymmetric geometry, plasmonic SCs are insensitive to the linear polarization of the incident light (i.e., the SC response does not change if light polarization is parallel to the edge of the SC or to its diagonal). This is an important feature in order to realize

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FIG. 2. Au/G fractal optical properties. **a**, Experimental reflectance $R = r/r_{sub}$ spectra of 35 ± 3 nm thick Au (blue solid line) and Au/G (red solid line) SCs for t = 5 order. Localized surface plasmon modes $\omega'_1 = 0.17 \text{ eV}$, $\omega'_2 = 0.50 \text{ eV}$, $\omega'_3 = 1.11 \text{ eV}$, $\omega'_4 = 2.04 \text{ eV}$, and $\omega'_5 = 2.40 \text{ eV}$ for the Au and $\omega_1 = 0.16 \text{ eV}$, $\omega_2 = 0.28 \text{ eV}$, $\omega_3 = 0.63 \text{ eV}$, $\omega_4 = 1.67 \text{ eV}$, and $\omega_5 = 2.30 \text{ eV}$ for the Au/G SCs are marked. **b**, Experimental normalized reflectance $R = r/r_{sub}$ of 35 ± 3 nm thick Au/G periodic arrays with lattice constants a_t and LSP modes $\omega_1 = 0.16 \text{ eV}$, $\omega_2 = 0.28 \text{ eV}$, $\omega_3 = 0.59 \text{ eV}$, $\omega_4 = 1.59 \text{ eV}$, and $\omega_5 = 2.20 \text{ eV}$. Curves are offset by 1. **c**, Dispersion relations of the LSP modes as a function of the plasmon wavevector for Au (blue squares), Au/G periodic arrays (red dots). Black curve is the result of electromagnetic calculations. **d**, Experimental relative reflectance $(R - R_{CNP})/R_{CNP}$ of the Au/G SC at t = 5 order for different gate voltages. The LSP mode ω_3 (red diamonds) and the interband absorption threshold $2E_F$ (blue diamonds) are marked. **d**, **inset**, Dispersion relations of the interband absorption threshold $2E_F$ (blue diamonds) and the Au/G SC ω_3 mode (red diamonds) as a function of the graphene Fermi level. The blue solid line is the law $\hbar \omega = 2E_F$.

polarization-independent devices⁷, since other aperiodic metamaterial^{6,12,15,16} and periodic metamaterials^{3,9,10,14} usually depend strongly on light polarization.

Plasmonic fractals provide a hierarchical spatial distribution of near-field electromagnetic enhancement, due to LSP modes supported on the surface of the self-similar resonators forming the whole structure^{34,36}. When a mode of the Au fractal is excited, a considerable field enhancement not only near the squares but also across the graphene layer is observed. In fact, graphene represents a

unique model system where nonlinear applications, such as surface enhanced Raman spectroscopy (SERS), can be studied in detail^{39–43}. In Figure 3a, average Raman spectra for the t = 5 order of the Au/G SC and for an unpatterned graphene sample (t = 0) are plotted. A SERS enhancement factor on the Au/G SC can be evaluated in Figure 3b as a function of the SC lattice constant a_t for the graphene G and 2D phonon bands. As for Au SCs, the SERS enhancement factor follows a power law³⁴, but for Au/G SCs is 10-fold higher for the smallest feature,

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FIG. 3. Au/G fractal surface electromagnetic enhancement. **a**, Averaged Raman spectra at $\lambda_{ex} = 514$ nm of the Au/G SC for t = 5 order and for a reference unpatterned graphene. Curves are offset by 50. The graphene D, G, and 2D vibrational bands are shown at 1343-1349 cm⁻¹, 1584-1589 cm⁻¹, and 2674-2689 cm⁻¹, respectively. **b**, SERS enhancement of the graphene G (black squares) and 2D (red dots) bands at $\lambda_{ex} = 514$ nm as a function of the SC lattice constant a_t , in log-log scale. Experimental electric field enhancement $|E|^2/|E_0|^2$ maps of the BCB vibrational band $\omega^* = 1655$ cm⁻¹ at $\lambda_{ex} = 532$ nm for the t = 5 order SC (**c**) and its electromagnetic simulation obtained by CST Microwave Studio, www.cst.com (**d**). Each map is normalized to its maximum value. Note that since the laser spot diameter is 1 μ m, the instrument averages out over L_4 and L_5 structures.

achieving a value of 10^5 . Additionally, as shown in Figure 3a, depositing metal structures on graphene represents an elegant way to pattern it without directly etching it, thus without altering the graphene crystalline structure (see Supplementary Information 4). Furthermore, by depositing on the t = 5 order Au/G SC a thin layer of Brilliant Cresyl Blue (BCB) dye as a probe (see Supplementary Information 5), it is possible to map the SERS enhancement all over the fractal surface (Figure 3c). The contour plot presents a maximum of contrast among the squares with size L_4 and L_5 in excellent agreement with

the electromagnetic simulation illustrated in Figure 3d (see Supplementary Information 3). At the probed excitation wavelength the dye is not resonant³⁴, therefore the observed signal is entirely related to a surface-enhanced electromagnetic effect.

Recently, the quantum properties of electrons in fractals have been investigated both theoretically⁴⁴ and experimentally⁴⁵, however their classical electronic behavior is has not been experimentally reported so far. As shown in Figure 1a, c, the phototransistor channel in the photodetector devices consists of a $40 \times 40 \ \mu m$ CVD-

FIG. 4. Au/G fractal electrical properties. **a**, I_{SD} - V_{SD} characteristics at $V_G = -100$ V of the Au/G SC for t = 5 order. **a**, **inset**, Scheme of the conduction band E_c spatial profile of the Au/G devices between two Au electrodes, under a positive bias V_{SD} . E_F is the junction Fermi level at equilibrium and $\Delta \Phi$ the contact potential. **b**, Resistivity at $V_{SD} = 0.1$ V of the Au/G SC for t = 5 order and a reference unpatterned graphene as a function of the carrier density and the gate voltage shift with respect to the unpatterned graphene CNP. **b**, **inset**, Fermi level shift as a function of the Au/G fractal fill factor. The red solid line is a linear fit of the data. **c**, Carrier mobility at $V_{SD} = 0.1$ V of the Au/G SC for t = 5 order and a reference unpatterned graphene as a function of the unpatterned graphene CNP. **b**, **inset**, Fermi level shift as a function of the au/G fractal fill factor. The red solid line is a linear fit of the data. **c**, Carrier mobility at $V_{SD} = 0.1$ V of the Au/G SC for t = 5 order and a reference unpatterned graphene as a function of the carrier density and the relative gate voltage to the unpatterned graphene CNP. **d**, Normalized mobility at a carrier density $n = 6 \cdot 10^{12}$ cm⁻² of Au/G fractals for t = 0.5 to the value for t = 0 as a function of the relative Fermi level shift to the value for t = 0. Error bars are due to averages over multiple resistivity measurements on the same sample.

grown, single-crystal single-layer graphene flake sensitized with an Au SC fractal (see Methods). Graphene is the active material of the device and the Au fractal acts as a plasmonic sensitizer to improve the overall light sensing. In the inset of Figure 4a, a scheme of the electronic band profile of the Au/G photodetector device is depicted. Upon contact, graphene forms an ohmic junction with the Au electrodes (Figure 4a). Therefore, an electrostatic contact potential $\Delta \Phi = 0.3$ eV is built, due to the difference between the graphene and Au work functions $\Phi_G = 4.5 \text{ eV}^{20}$ and $\Phi_{Au} = 4.8 \text{ eV}^{46}$, respectively. For the same reason, the Au fractal also provides an hole charge-transfer to graphene (Figure 4b), resulting in a p-type doping. As illustrated in the inset of Figure 4b, such a doping is linear with the Au/G fill factor, i.e. the area fraction of graphene uncovered by Au (see Supplementary Information 1). This effect leads to an increase of the hole carrier density n in graphene, therefore to a

FIG. 5. Au/G t = 5 order fractal photodetectors. **a**, Photocurrent at $V_G = -50$ V and $V_{SD} = 0.1$ V as a function of the power of 1.46 eV (blue dots), 1.94 eV (red dots), and 2.38 eV (green dots) lasers. Solid lines are power law fits $J \propto P^{\alpha}$, with $0 < \alpha < 1$, for P > 6 mW/cm². The fits return $\alpha = 0.2$. **b**, Photocurrent at $V_G = -50$ V as a function of the V_{SD} bias for a 1.94 eV laser of power 10 mW/cm².

decrease of the carrier mobility $\mu_e = \sigma/ne$, as a function of the fractal order or fill factor (Figure 4c, d).

Since all the leads are made of the same material, a symmetric potential that traps the photogenerated electrons occurs in the device (Figure 4a, inset). By applying a bias voltage V_{SD} , the photogenerated holes can be extracted, provided that the graphene Fermi level E_F compensates the trapping potential by applying for instance a gate voltage V_G (see Supplementary Information 6).

In Figure 5a, it is possible to observe that for a t = 5 order device, the photocurrent divided by the illuminated

active area $J = I_{ph}/(4L_0)^2$ exhibits a nonlinear dependence on the incident power of light P, due to electrostatic and thermal effects, trapping, and recombination occurring at high values of power^{7,47}. The photocurrent response results linear up to 6 mW/cm² for $V_G = -50$ V at a photon energy of 1.94 eV. Furthermore, by photogating effect, the device produces a DC photovoltage that is dependent on the incident photon energy and power (see Supplementary Information 6). Moreover, the Au/G SC can linearly detect light down to 2 nA for $V_G = -50$ V at 1.94 eV (Figure 5b).

In Figure 6a, b, the broadband and electrically tunable responsivity $\mathcal{R} = J/P$ of a t = 5 order Au/G SC 3×3 array can be observed as a function of the applied gate voltage. An enhancement factor $\mathcal{R}_{Au/G}/\mathcal{R}_G$ up to 100-fold of the Au/G SC device over an unpatterned graphene device (see Supplementary Information 6) is presented in the inset of Figure 6a. The array was introduced in order to increase the photocurrent signal-to-noise ratio. In addition, the device total absorption A = 1 - r, and the external $EQE = \mathcal{R}\hbar\omega/e$ and internal IQE = EQE/A quantum efficiencies are evaluated in Figure 6c. It is worth noting that in the spectral range investigated, the Au/G SC photodetector IQE is up to 100% in correspondence of the ω_5 LSP resonance.

From the application perspective, the relevant figure of merit to characterize a photodetector is the optical noise-equivalent-power $NEP = \sum_i N_i / \mathcal{R}$, which corresponds to the lowest detectable power in 1 Hz output bandwidth⁴⁷. Usually, the noise level of photodetectors in current-operating mode is due to the shot noise⁴⁷ $N_1 = \sqrt{2eI_{SD}}$ and to the thermal Johnson-Nyquist contribution⁴⁷ $N_2 = \sqrt{4k_B T \sigma}$, which ultimately limits the NEP. In Figure 6d NEP is shown as a function of the incident photon energy for $V_G = -50$ V. At room-temperature, we observed a minimum NEP of $2 \times 10^{-14} \text{ W} \text{Hz}^{-1/2}$ for a 2.38 eV photon energy, while $D^* \equiv L/NEP = 2 \times 10^{11} \text{ cm Hz}^{1/2} \text{W}^{-1}$ or Jones represents the device maximum specific detectivity at the same photon energy. Such a high value of detectivity is compatible with state-of-the-art two-dimensional $photodetectors^{1,48}$.

Generally, five main phenomena compete in the operation of a graphene-based photodetector: the photovoltaic, the bolometric, the photothermoelectric, the photogating, and the plasma-wave-assisted effect¹. In our case of study, we can reasonably exclude the latter mechanism, as it is likely to occur for THz radiation²⁸ $(\omega \tau \ll 1, \text{ with } \tau \approx 0.1 \text{ ps the graphene carrier scattering})$ time in our samples). In a metal-graphene-metal configuration, in which graphene is contacted with metal electrodes as source and drain, a p-n junction occurs close to the contacts because of the difference in the work functions between the metal and graphene⁴⁹. Consequently, a charge transfer with a shift of the graphene Fermi level in the region below the metal pads occurs. The junction results in an internal electric field able to separate the light-induced electron-hole pairs (photovoltaic effect).

FIG. 6. Au/G t = 5 order fractal array photodetector efficiencies. **a**, Responsivity of an Au/G SC array and an unpatterned graphene photodetector at $V_{SD} = 0.1$ V as a function of the gate voltage. **a**, **inset**, Responsivity enhancement factor of an Au/G SC array with respect to an unpatterned graphene photodetector (t = 0) at $V_G = -50$ V and $V_{SD} = 0.1$ V. **b**, Responsivity of an Au/G SC photodetector as a function of the photon energy of a panchromatic source at $V_G = -50$ V. **b**, **inset** False color optical microscopy micrograph of the 3×3 array on the device channel. **c**, Au/G SC device total absorption (A = 1 - r), internal and external quantum efficiencies at $V_G = -50$ V and $V_{SD} = 0.1$ V. **d**, Total NEP (red solid curve) and only thermal contribution (solid black curve) at $V_{SD} = 0.1$ V as a function of a panchromatic source at $V_G = -50$ V in linear-log scale.

Unless the contacts are made of different materials, the photocurrent produced at both the contacts will be of opposite polarity for symmetry reasons, resulting in zero net current at small gate voltages close to the CNP^{19} . Usually, in such devices a light-induced photogating and photothermoelectric effect are also present at gate voltages close to the CNP, while at larger gate voltages the bolometric effect is typically dominant^{1,26,49}. We observed the photovoltaic and photogating effects in the unpatterned graphene photodetectors (see Supplementary Information 6). In the Au/G SC devices, at gate

voltages close to the CNP and for a symmetrical electronic band profile, the photovoltaic and bolometric effects are expected to play a minor role in comparison to the photothermoelectric and photogating effects²⁶, due to the plasmonic Au SC sensitizer (see Supplementary Information 6). On the other hand, at sufficiently larger gate voltages ($V_G < -40$ V), where the photothermoelectric and photogating effects become smaller, the bolometric effect should be dominant. Therefore, we propose the following explanation for the photogeneration mechanism in Au/G SC photodetectors. When light is shone

on the metadevice, it is absorbed by Au fractal LSPs generating electron-hole pairs in the sensitizer (gating laver), which are transferred to graphene (channel laver). This results in a photocurrent and a photovoltage due to the change in electronic temperature between the Au SC and graphene regions, thus a variation of the carrier density $\Delta n = C \Delta V_G$ (see Supplementary Information 6). Since the device electronic bands are symmetric, the photogenerated electrons are trapped in graphene for $\tau_{trap} \equiv \Delta n \hbar \omega / I Q E P \approx 27 \text{ ms}$ (see Supplementary Information 6). Therefore, only holes at relatively high gate voltages contribute to the photocurrent. For the t = 5 order Au/G SC array at $V_G = -50$ V the hole carrier transit time is $\tau_{trans} \equiv L^2/\mu_e V_{SD} \approx 20$ ns, with FET channel length $L = 40 \ \mu m$ and carrier mobility $\mu_e \approx 8 \times 10^3$ $cm^2V^{-1}s^{-1}$. Therefore, the lower bound (i.e. by neglecting the circuit RC constant) of the device bandwidth is $\Delta f_{-3dB} \approx 3.5/2\pi \tau_{trans} = 30$ MHz, while the photocon-ductive gain is up to $G \equiv \tau_{trap}/\tau_{trans} = 10^6$, providing evidence of a multiple carrier generation^{29,50} in graphene (see Supplementary Information 6). Evidently, such a high gain, produced by prolonged carrier lifetime due to trapping, limits the photodetector response speed and bandwidth.

In summary, we presented a novel graphene-based, plasmonic metamaterial having a multiresonant optical response hierarchically distributed on the fractal structure, which can be employed to realize a highly sensitive and polarization-independent, broadband photodetector. The metadevice is compact and can operate at room temperature under small voltages, with low power consumption. The detectivity could be further improved by depositing the Au fractal on high-mobility graphenebased van der Waals heterostructures, such as G/hBN⁵¹. Moreover, the deterministic chemical vapor deposition (CVD) growth of large area graphene⁵² would grant integration over wafer scale at a low cost of graphene-based photodetectors. Finally, the wide-range and high-gain spectral response of the Au/G fractal and its ability to hierarchically localize strong electromagnetic fields on a sub-wavelength scale could pave the way to a new generation of materials for integrated optics and optoelectronics, such as multiband photosensor pixel arrays for imaging cameras and optical communication devices.

METHODS

Device fabrication

Single-layer, single-crystal graphene was grown on copper foil by CVD using an Aixtron BM cold-wall reactor at a temperature of 1060 °C and a pressure of 25 mbar. A semi-dry transfer approach was used to transfer the graphene from Cu to the target substrate⁵². The foil was spin-coated by a PMMA membrane and the graphene was electrochemically delaminated from the growth substrate in 1M NaOH solution. Using a frame, the graphene/PMMA membrane was removed from the electrolyte, rinsed in deionized water and dried in ambient conditions. The membrane was deposited on the target Si/SiO₂ substrates at 120 °C and the PMMA carrier polymer was removed in acetone. Graphene and Au SCs were patterned by EBL (Raith 150-Two) on Si wafers (1-5 m Ω cm) with a $t_{ox} = 285$ nm tick layer of thermal oxide. A layer of approximately 160 nm of 950K polymethyl-methacrylate (PMMA A2 1:1, 1% in anisole) was deposited by spincoating on the substrates, which were then post-baked for 7 min on a hot plate at 180° C. The electron beam was operated at 20 kV with a current of 35 pA. After exposure, graphene was etched by O_2 (5 sccm) reactive ion etching (Sentech instruments ICP-RIE 500) for 30 s at a RF power of 35 W and temperature of 15° C in an Ar inert atmosphere (80 sccm). The SC patterns were exposed with a varying dose inversely proportional to the size of the squares of the fractal. Contacts pads and probes were exposed for the electrical measurements. The resist was developed for 30 s in a cold (8° C) 1:3 mixture of MIBK:IPA. A 35/5 nm thick Au/Ti film was deposited on the substrates at a rate of 0.2 Å/s by electron-beam evaporation (Kenosistec) at an operating pressure of 10^{-6} mbar. Lift-off was then performed by hot (80° C) acetone. A wire bonder (F&S Bontec Series 56) was used to wire the contact pads to an external circuit board. For SERS measurements, SCs were dipcoated for 1 h in 1 mM BCB $[(C_{17}H_{20}ClN_3O)_2ZnCl_2]$ aqueous solution, then rinsed in deionized water to wash the excess molecules in order to form a thin layer 12 ± 2 nm thick, and finally dried in nitrogen flow.

Sample characterization

Samples were characterized as in Ref. 34. Scanning electron microscopy micrographs of Au/G SC samples deposited on Si/SiO₂ were acquired by FEI Helios NanoLab DualBeam 650. Atomic force microscopy height profiles of the samples deposited on Si/SiO₂ were measured in tapping mode by Bruker Innova in combination with v-type cantilever and SiN tips. Optical spectroscopy in reflection mode was performed with unpolarized light on SCs deposited on CaF_2 substrates, by Thermo Fisher FTIR spectrometer and Thermo Fisher Nicolet Continu μ m microscope equipped with NO₂cooled MCT and Si detectors, KBr and quartz beamsplitters, and a $15 \times (0.58 \text{ N.A.})$ Cassegrain objective. Surface enhanced Raman spectroscopy was carried out by Renishaw inVia micro-Raman microscope equipped with a $100 \times (0.85 \text{ N.A.})$ and $150 \times (0.95 \text{ N.A.})$ objectives, and 514 nm and 532 nm lasers at power 1.2 W/cm² and 0.1 W/cm^2 , respectively, and integration time 1 s. All spectra were calibrated with respect to the first-order silicon LO phonon peak at 520 cm^{-1} and recorded in backscattering geometry at room temperature. Raman measurements were performed on Au/G SCs and as a reference on bare graphene deposited on the same Si/SiO₂ sub-

strate. Raman maps of graphene G and 2D, and BCB $\omega^{\star} = 1655 \text{ cm}^{-1}$ vibrational bands were scanned at 0.3 μm steps in both the directions in the plane of the sample. Renishaw WiRE 3.0 software was used to analyze the collected spectra, whose baseline was corrected to the third-order polynomial. Two- and four-probe electrical measurements in van der Pauw configuration were performed in vacuum (10^{-6} mbar) by Keithley 2612A digital source-measure unit and Agilent 34410A digital voltmeter. In order to estimate the value of the residual carrier density $n_0 \approx 10^{11} \text{ cm}^{-2}$, two-probe resistance curves were fitted by the formula $R_{tot} = 2R_c + R_{sh}$, with $R_{sh} = L/We\mu_e \left(n^2 + n_0^2\right)^{1/2}$ the sheet resistance and L, W the FET channel length and width, respectively. For four-probe resistance curves $R_c = 0$ and $R_{sh} = 4.53 R_{tot}$, due to the van der Pauw geometrical correction. Carrier mobility was then evaluated by $\mu_e = \sigma/ne$. Photocurrent was measured using 0.80 eV, 1.46 eV, 1.94 eV, and 2.38 eV laser sources at different powers. The output

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power density was calibrated by a power meter. Responsivity was measured by using the following custom optical setup. An ASB-XE-175 source emits panchromatic light passing through a SP CM110 monochromator controlled by PC. Emerging monochromatic light is coupled to an optical fiber, modulated at 173 Hz by a mechanical chopper, and focused by a lens first on the samples, then on Si and Ge photodiodes. The AC current from the sample is measured through a current pre-amplifier (DL 1211) with a amplification factor $\mathcal{C} = 10^{-6} \text{ A/V}$, whose output is read by a Signal Recovery 7265 lockin amplifier. The photocurrent is given by the relation $J = 2\pi\sqrt{2}V_{lock-in} * \mathcal{C}/4$, where the factor 2 is due to the peak-to-peak magnitude, the factor $\sqrt{2}$ originates from the lock-in amplifier rms amplitude, $\pi/4$ is the fundamental sine wave Fourier component of the square wave produced by the chopper, and $V_{lock-in}$ the amplitude of the signal read by the lock-in.

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ACKNOWLEDGMENTS

The authors acknowledge that this work has received funding from the European Union's Horizon 2020 research and innovation programme under grant agreement No. 696656 - GrapheneCore1.

AUTHOR CONTRIBUTION

F.D.N., A.T., M.P., R.K., and V.P. conceived the experiments. V.M. and C.C. grew and transferred the graphene. N.P.P. fabricated the devices. F.D.N. performed the sample optical characterization, Raman spectroscopy, simulations, and data analysis. F.D.N., D.S., and N.P.P carried out the electrical measurements. All the authors discussed the experimental implementation, the results, and contributed in writing the paper.

The authors declare no competing financial interests.