Metallic nanostructures as electronic billiards for nonlinear terahertz photonics

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The optical properties of metallic nanoparticles are most often considered in terms of plasmons, the coupled states of light and quasifree electrons. Confinement of electrons inside the nanostructure leads to another, very different type of resonances. We demonstrate that these confinement-induced resonances typically join into a single composite "super-resonance," located at significantly lower frequencies than the plasmonic resonance. This super-resonance influences the optical properties in the low-frequency range, in particular, producing giant nonlinearities. We show that such nonlinearities can be used for efficient down-conversion from optical to terahertz and midinfrared frequencies on the submicrometer propagation distances in nanocomposites. We discuss the interaction of the quantum-confinement-induced super-resonance with the conventional plasmonic ones, as well as the unusual quantum level statistics, adapting here the paradigms of the quantum billiard theory and showing the possibility to control the resonance position and width using the geometry of the nanostructures.

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I. INTRODUCTION

Light propagating in the vicinity of metallic surfaces or metallic nanostructures is strongly coupled to the "electronic fluid" formed by quasifree electrons in the metal, resulting in joint electron-photon excitations which are called surface or particle plasmons, depending on the geometry. Plasmons and the corresponding plasmonic resonance (PR) are at the very heart of optics of nanostructures.

PRs appear by matching of the incoming light to the intraparticle fields, leading to strong surface charges and resonance peaks of linear and nonlinear response of metallic nanoparticles at certain frequencies [1–3]. That is, PRs are defined via the matching condition of the fields, rather than electrons themselves, and have no direct relation to the electron confinement inside the nanostructure. Plasmonic resonances are located at quite high frequencies, commonly in the visible range.

As soon as we consider very small metallic nanoparticles, quantum confinement of electrons in the finite volume of a nanoparticle comes into play. Possible confined-based resonances have rarely attracted attention *per se*, separately from the properties of plasmons. On the other hand, any calculation of the properties of small nanoparticles does in principle include quantum mechanical confinement of electrons as an ingredient, noticeably influencing the position and the width of PR [2,4–7]. In the recent years, huge progress was made in both calculations and measurements [2,7–14] of properties of small nanoparticles. Theoretical approaches are used, starting from relatively simple analytical techniques [4,15–17], through the single electron in a box [15,18], jellium model [19], hydrodynamiclike equations [20,21], and quantum hydrodynamic theory [22] towards direct *ab initio* density functional theory methods fully taking into account the ionic core structure [9,14,23–25]; for a review see [13]. Whereas analytical and hydrodynamic approaches can address nonlinear properties [16,20,21], complex *ab initio* methods focus solely on the linear susceptibility, unless very small nanoclusters are considered [26,27].

For very small nanoclusters and nanoparticles (a few hundred atoms and below), the response is molecularlike [9,14], typically including a cacophony of resonances replaced by a single PR [14,28,29]. At lower frequencies, a prominent molecularlike resonance is the highest occupied molecular orbital–lowest occupied molecular orbital (HOMO-LUMO) transition [14,30–32]. It describes an excitation of a single localized electron, and depends heavily on the molecular structure [30].

As we shift to larger nanoparticles, the HOMO-LUMO transition effectively fades out [14] because its decay rate grows exponentially with the size of the nanostructure [31,32]. However, the PR is not the only one which remains. Signatures of a single resonance well below the plasmonic one but above the HOMO-LUMO transition were observed experimentally [33,34] and later confirmed theoretically [8,35] involving *ab initio* simulations [35]. This resonance is however mostly overlooked since then, and there is no consensus in explanations of its nature. Whereas in [8,34] the confinement-based argument were put forward, [33,35] tries to explain it without leaving the plasmonic framework by introducing "restoring force on the electrons."

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FIG. 1. A spherical gold nanostructure (left) of 2.75-nm diameter and the band level structure (right) appearing due to the electron confinement (quantum billiard resonances). Fermi energy \mathcal{E}_F and work function \mathcal{E}_W as well as exemplary virtual transitions contributing to linear (orange arrows) and nonlinear (red arrows) properties are indicated. One of the wave functions $\psi(x, y, z)$ is visualized inside the nanostructure (a sector of the sphere is cut for visualization purposes). Even for the simplest nanoparticles, energy level structure is quite complex.

Here we show that confinement-based resonances under normal conditions (at least for nanostructures with regular enough geometry) merge together and form a single broadband "super-resonance" located commonly in the terahertz and midinfrared (MIR) frequency range, with both the width and position widely controllable with the nanostructure geometry. Such super-resonance must be considered as one of the universal signatures of metallic nanostructures, yet fundamentally different from the PR.

We study the statistical features of confinement-based resonances by adapting and modifying the paradigm of neighboring-level statistics from the field of quantum billiards [36]. Electrons confined in a nanostructure certainly represent a type of quantum billiard. Yet, up to now, with only very few exceptions [37,38], nonmetallic billiards such as semiconductor quantum dots [39–45] were considered. As we show, the metallic nature of our billiard provides a unique opportunity to observe certain features of the level statistics directly in the optical properties.

Furthermore, the super-resonance provides a broadband nonlinear response, leading to giant nonlinearities. We demonstrate how these nonlinearities can be used for an efficient optical rectification and difference-frequency mixing in the nanocomposites, enabling broadband conversion from optical to MIR or terahertz ranges by submicrometer devices.

II. THE MODEL

We used a simple analytical approach of a single particle in a box [6,8,16,34] (see also more details in Appendixes A and B) with electrons fully confined inside a nanoparticle. For calculation of optical properties, we consider electron energy structure characterized by Fermi energy \mathcal{E}_F and work function of \mathcal{E}_W , as illustrated in Fig. 1 right. This approximation works well for metals with a simple Fermi surface such as alkali metals (Li, Na, Ca, Rb); it is an acceptable simplification for metals with somewhat more complicated Fermi surfaces such as Cs, Cu, Ag, or Au; and it is barely applicable at all for other metals. The linear ($\chi^{(1)}$) and nonlinear ($\chi^{(3)}$) susceptibilities were calculated using a version of the standard perturbative iterative approach [46] which takes into account selection rules following from the Fermi-Dirac statistics (see Appendixes D, F, and G). The linear and nonlinear optical properties can be described via a sum of contributions of virtual transitions from inside the Fermi sea to outside and back [see orange (linear) and red (nonlinear) arrows Fig. 1]. We also assumed fast population decay time $T_1 = 50$ fs and dephasing time $T_2 = 5$ fs [16].

To be more specific, in our numerical simulations we consider gold since it is a very widespread material, and is suitable for composites due to its low imaginary part of susceptibility. Yet we note that conclusions we draw below are basically metal independent (taking into account precautions mentioned above). In gold, a simple ideal-metal picture discussed above neglects several linear and nonlinear effects, such as interband transitions, influence of finite temperature, and hot-electron nonlinearities. However, these effects are negligible for lowfrequency response in the terahertz or MIR range driven by femtosecond pulses, and our model remains adequate in this regime (see Appendix F for justifications and detailed estimates).

III. QUANTUM BILLIARD (CONFINEMENT-BASED) RESONANCES

The typical level structure obtained by the above model for an exemplary spherical gold nanoparticle of the diameter d = 2.75 nm is shown in Fig. 1 (see examples for other diameters in Appendix E and Fig. 5). These levels originate from electron confinement in the nanostructure. Even in the presented case of a very simple particle, the levels look quite irregular. This is a familiar picture in the framework of quantum billiard theory [36], where the statistical properties of the level distribution play one of the central roles. For instance, one can consider the neighboring level statistics (NLS), which allows one to distinguish between integrable (regular) and nonintegrable (chaotic) billiards. For the regular billiards, such as spheres, the probability density $P(\omega)$ of neighboringlevel distance ω obeys Poissonian statistics: $\ln P(\omega) \propto -\omega$. This is also true in our case: NLS corresponding to Fig. 1 is plotted in Fig. 2(a) by yellow bars and coincides well with the Poissonian distribution (black dashed line).

Judging from such statistics, one might expect a conglomerate of resonances near zero frequency, but this is not the case. An optical response $\chi^{(1)}$ resulting from the electron confinement for few exemplary nanostructures is shown in Fig. 2. Note that, in addition to the confinement-based impact shown in Fig. 2, the full linear response includes also the so-called Drude part, representing the action of quasifree electrons (see Appendix F for more details).

The clearly observed feature of the confinement-based linear response is the presence of a single resonancelike peak in the MIR/terahertz range at a nonzero resonance frequency ω_{conf} which quickly decreases with increasing particle size. Such peak was observed experimentally [33,34] and theoretically [8,35]. It is easy to see that this resonance co-



FIG. 2. Billiard resonances and weighted level statistics for a golden sphere (a) of a few exemplary diameters *d* and a cylinder (b) of the diameter *d* and height *h*. The blue and red curves in (a) and (b) show $\text{Im}(\chi^{(1)})$ as the function of frequency for different sizes (see legend). Bars of the same colors show the corresponding weighted level statistics $P(\omega)$ normalized to some P_0 (P_0 is not the same for different curves). Green bars show the statistics for all allowed transitions attached, for visibility, to the upper *x* axis (that is, technically, $1 - P(\omega)/P_0$ is shown; see also larger frequency scales in Appendix G, Fig. 8). Black lines show $\text{Im}(\chi^{(1)})$ for T_2 increased 100 times. Yellow bars show NLS and the dashed black line indicates the Poissonian statistics.

incides well with the minimally possible allowed transitions close to the Fermi energy which can be, for a spherical nanoparticle, analytically estimated as (see Appendixes C and E for details; see also [34])

$$\omega_{\rm conf} \approx \frac{\pi}{r} \sqrt{\frac{\mathcal{E}_F}{2m_e}},$$
 (1)

where m_e is the electron mass. The frequency in Eq. (1) decreases with increasing diameter and, for the diameters above 5 nm, is located in the midinfrared and terahertz region. This makes it fundamentally different from the PR, which is located at a significantly higher frequency. Yet, why does only a single resonance arise? Can we influence its width and position? These and related questions will be addressed in the following paragraphs.

Closer consideration allows us to establish that, starting already from quite small nanosphere diameters $d \approx 2.5$ nm, the resonance near ω_{conf} is composed of many transitions with nearby frequencies [see Fig. 2(a)]. Existance of multiple resonances is especially well observable if we consider much larger T_2 (which would correspond to low temperatures [47]). In this case, many separated resonances are indeed visible in the optical response, as shown in Figs. 2(a) and 2(b) by black curves. The particular structure of the transitions depends significantly on the geometry [see Fig. 2(b) for a cylinder]. At room temperature, these transitions merge into one super-resonance because of the broad line widths. For instance, the position and width of the super-resonance for a cylinder are shifted in comparison to a sphere with the same volume and diameter by the noticeable amount of 35 and 23%, correspondingly. Nevertheless, Eq. (1) remains a valid, yet rough estimation of the position of the resonance.

Both the position of the super-resonance ω_{conf} and its structure can be analyzed using a level-distance statistics similar to NLS. A naïve approach would be to calculate such statistics using all dipole-allowed transitions between the confinementbased levels, shown in downward green rectangles in Fig. 2(a), and covering an extremely broad range around 10 eV [see green bars in Fig. 2(a) and also Appendix G and Fig. 8]. However, it must be modified to include only transitions from below to above \mathcal{E}_F , obviously corresponding to the Pauli principle and absence of population above the Fermi level. In addition, in the statistics we weight the transitions by the square of the corresponding dipole momentum, thus taking into account the known tendency of the transition dipole momenta to rapidly decrease, on average, with the energy difference. The resulting modified statistics is shown by the red and blue bars in Fig. 2 and agrees nicely both with the position of the super-resonance and with its width (for small T_2). Namely, in the limit of small T_2 the super-resonance has a certain "natural" width; for the case of nanospheres it can be estimated as $\omega_{conf}/4$ (see Appendix G). Analysis of the position of the super-resonance for nanospheres (see Appendix G) indicates that the energy of the participating states is located mostly in radial (rather than angular) motion.

IV. NONLINEARITIES

The above described low-frequency resonance is expected to lead also to strong nonlinearities; in our case, $\chi^{(3)} \neq 0$ as calculated using the approach described above. We note that such approach to calculate Kerr nonlinearity was already utilized in [16]; however, instead of the discrete spectrum, approximation of continuous density of states was used. Nevertheless, we checked that our calculations are in quantitative agreement with [16]; they are also in agreement with experimental measurements for short pulses (see [11] and references therein). An example of $\chi^{(3)}(\omega; \omega, \omega, -\omega)$ for the four-wavemixing (FWM) process $\omega + \omega - \omega = \omega$ (corresponding to the Kerr nonlinearity) is shown in Fig. 3(a) for several diameters. The low-frequency resonance we observed in $\chi^{(1)}$ is also well visible here. Whereas in the linear response the Drude part dominates (see Appendix F and Fig. 5), in the nonlinear response it is fully absent.

We now try to exploit this low-frequency resonance. Motivated by detection and spectroscopic applications of terahertz and MIR radiation, we focus on the FWM providing a signal in the terahertz and MIR range, generated from a sub-100-fs pump pulse. Nonlinear susceptibilities $\chi^{(3)}(\delta; \omega, \omega, -2\omega + \delta)$ and $\chi^{(3)}(\delta; \omega, \omega + \delta, -2\omega)$, leading to generation of low-frequency signal at frequency δ as a result of a FWM



FIG. 3. Nonlinear susceptibility for the Kerr nonlinearity (a) and for different FWM processes leading to the optical rectification (b, c) in dependence on the signal δ (c) and pump ω (b) frequency, shown for the nanostructures of different diameters [as indicated by different marker shapes, see legend in (a)].

process in a two-color pump at frequencies ω (fundamental) and 2ω (its second harmonics), are presented in Figs. 3(b) and 3(c). Both Kerr rectification nonlinearities presented in Fig. 3 are several orders of magnitude higher than the Kerr nonlinearity of the fused silica $\approx 2 \times 10^{-22} \text{m}^2/\text{V}^2$. In Fig. 3(a), where pump-frequency dependencies are shown, the billiard super-resonance described above is very clearly visible. This is not a unique property of metallic nanostructures. Giant nonlinearities in semiconductor nanostructures due to billiard resonances were recently predicted in [48].

V. EFFICIENT FREQUENCY DIFFERENCE GENERATION

As an interesting application we consider the process of optical rectification and difference frequency generation, governed by three nonlinearities $\chi^{(3)}(\delta; \omega_0, \omega_0, -2\omega_0 +$ $\delta)$, $\chi^{(3)}(\delta; \omega_0 + \delta, \omega_0, -2\omega_0)$, and $\chi^{(3)}(\delta; \omega_0, \omega_0 + \delta, -2\omega_0)$, with a two-color optical pump at around ω_0 and $2\omega_0$ and signal $\delta \ll \omega_0$ in terahertz and MIR. We solve the propagation equations, assuming slowly varying envelope approximation and taking into account dispersion relations, but neglecting nonlinear effects for the pump waves because of very small propagation distance (see Appendix I for details). Both $\chi_{eff}^{(3)}$ and the linear susceptibility $\chi_{eff}^{(1)}$ are calculated from given linear and nonlinear properties of the nanoparticles ($\chi_{NP}^{(1)}, \chi_{NP}^{(3)}$) and host ($\chi_h^{(1)}, \chi_h^{(3)}$) using the effective medium approach [49] (see Appendix H). By calculation of the linear properties the full linear susceptibility containing both confinement-based and Drude parts is included. As a host material, we take fused silica which possesses strong losses in the range between 30 and 40 THz (see Fig. 4), but otherwise is quite transparent [50]. We consider the filling factor of f = 0.01 and neglect the nonlinearity of the host. Resulting effective linear quantities are shown in Fig. 4 and demonstrate the usual PR resonance at around 2.4 eV with the width of around 30 THz. The shortest pulses still supported by this resonance are around 30 fs in duration.

Assuming an exemplary pulse durations of around 30 fs, we must consider two regions for the pump where conversion works significantly different. For the signal in the terahertz range ($\delta/2\pi \leq 30$ THz), the frequencies $j\omega_0$ and $j\omega_0 + \delta$ (j = 1, 2) are both located within the spectrum of the pump. In contrast, for the signal in MIR range $\delta/2\pi > 30$ THz, the components $\omega_0 + \delta$ and $2\omega_0 + \delta$ are not within the pump spectrum anymore. This leads to different treatment of these two frequency ranges for the selected pulse duration (see Appendix I).

The resulting field amplitude at zeroth harmonic A_0 is given in Fig. 4(b) for different parameters and for the pump amplitudes $A_1 = A_2 = 10^{10}$ V/m. This pump for 30-fs pulses corresponds to a fluence around 0.3 J/cm², which is yet below the damage threshold of gold (around 0.5 J/cm² [51]) and of fused silica (around 1 J/cm² [52]). One can see that in the terahertz range the signal amplitude reaches 5×10^8 V/m corresponding to efficiency of around 10^{-5} . In the MIR range, the amplitude can exceed 10^9 V/m, delivering efficiencies above the percent level. Moreover, the maximal efficiency is achieved at 100-nm propagation distance for the terahertz signal and 1 μ m for the MIR signal. From Fig. 4(b) one can also see that the most efficient conversion is achieved for the pump frequency ω_0 centered at the PR (solid lines in Fig. 4). In this case, the coupling of the pump to the signal is most efficient.

VI. DISCUSSION AND CONCLUSIONS

We showed that confinement-based energy levels in metallic nanostructures, representing integrable (or close to integrable) quantum billiards, typically join together into a single super-resonance, the position and width of which can be controlled by the geometry of the nanostructure. Whereas we focused here on (almost) integrable quantum metallic billiards, we anticipate richer resonance structure and control possibilities if truly chaotic billiards are considered. We analyzed the super-resonance, using the level statistics extended in comparison to that typically used in quantum chaos theory. In the linear regime the ballistic super-resonance is "hidden" behind the much stronger Drude response, yet it manifests itself strongly in a giant nonlinearity. This nonlinearity can be in addition enhanced by interaction with plasmons and effectively used to down-convert light to the terahertz and MIR range with high



FIG. 4. Efficient generation of terahertz and MIR light in a composite of gold spheres with two-color pump. (a) Effective refractive index n_{eff} and effective losses α_{eff} in dependence on frequency ω for a gold nanostructure with r = 10 nm immersed into fused silica (f = 0.01). Vertical lines show two variants of the two-color pump at $\omega_0 = 1.55$ eV ($\lambda_0 = 530$ nm, dashed line) and $\omega_0 = 2.4$ eV ($\lambda_0 = 530$ nm, solid line). The horizontal lines connect the spectral components of the two-color pump. (b) The generated field amplitude for different propagation distances *L* and nanostructure diameter *d* (see legend) and the pump as described in text. The solid vertical line in (b) separates the terahertz from the MIR band.

efficiency already after 100-nm distances, despite huge linear and nonlinear losses. Our confinement-based framework might also be helpful in a deeper understanding of the recent experimental work on efficient terahertz generation in nanostructures [53,54] and paves a way to extend newly proposed electronic metadevices [55] into the nonlinear regime. lence Strategy within the Cluster of Excellence PhoenixD (EXC 2122, Project No. 390833453). A.H. acknowledges support from European Union Project No. H2020-MSCA-RISE-2018-823897 "Atlantic."

APPENDIX A: SIMPLIFIED HAMILTONIAN AND WAVE FUNCTIONS FOR THE CASE OF SPHERICAL PARTICLES

To approach the problem analytically, we consider a spherical metallic particle of the radius a (diameter d = 2a). Since we are interested in low frequencies, we neglect the interband



FIG. 5. (a, b) Linear susceptibility due to confinement-based resonances only, in dependence on frequency for spherical nanoparticles of different diameters. (c, d) The Drude contribution as well as the full linear susceptibility (Drude plus confinement) for a few selected diameters.

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transitions. We neglect the temperature effects assuming that N electrons in the conduction band occupy all levels below Fermi energy \mathcal{E}_F (see Fig. 1). The energy structure is calculated assuming one-electron approximation and corresponds to that of a single free electron in an infinite-strength spherical potential of the radius a. To take into account finiteness of the potential, the levels above the work function \mathcal{E}_W are disregarded. The validity of these approximations is justified below.

The corresponding single-particle eigenproblem can be then formulated as $\hat{H}_0\psi = \mathcal{E}\psi$, with $H_0 = -\hbar^2/2m_e\Delta +$ V(r), V(r) = 0 for $r \leq a$ and $V(r) = \infty$ for r > a (m_e is the electron mass). With the approximations above, we neglect various effects of electron-electron and electron-ion interaction such as interband transitions, electron heating, the change of the eigenstates due to the finite height of the potential, and other effects, which play only a minor role at low frequencies and ultrashort sub-100-fs pulse durations. The validity of this approximation is discussed in Appendix F below. As it will be shown there, our simple model is rather adequate for the parameters we consider, despite its simplicity. The advantage of this approach is the possibility to determine the energy structure analytically. The corresponding eigenfunctions are combinations of spherical harmonics. The energies are defined as

$$\mathcal{E}_{nl} = \mathcal{E}_0 \alpha_{nl}^2 \tag{A1}$$

where n and l are quantum numbers,

$$\mathcal{E}_0 = \frac{\hbar^2}{2m_e a^2},\tag{A2}$$

and α_{nl} is the *n*th zero of the Bessel function of order *l*. In contrast to the Coulomb potential, there is no degeneracy in *l*.

The eigenfunctions of the problem described in the main paper are

$$\psi_{nlm}(r,\theta,\phi) = R_{nl}(r)Y_l^m(\theta,\phi), \qquad (A3)$$

where n, m, and l are quantum numbers, $R_{nl}(r) = \frac{\sqrt{2}j_l(\alpha_{nl}r/a)}{\sqrt{a^3}j_{l+1}(\alpha_{nl})}$, Y_l^m are spherical functions $(-l \le m \le l)$, j_l is the spherical Bessel function of order l, and α_{nl} is its lth

Zero.

The radial part of the matrix element is, for the allowed transitions $l' - l = \pm 1$,

$$\mu_{nl,n'l'} = \frac{4ae\mathcal{E}_0\sqrt{\mathcal{E}_{nl}\mathcal{E}_{n'l'}}}{(\mathcal{E}_{n'l'} - \mathcal{E}_{nl})^2},\tag{A4}$$

and zero in other cases (here e is the electron charge).

APPENDIX B: CYLINDRICAL GEOMETRY

In this section we determine the eigenvalues for the cylindrical geometry. We assume here that the main axis of the cylinder oriented along the z direction, and the light is assumed to be linearly polarized also in the z direction. Note the difference in the denotations with the part where propagation is considered: there, the z direction is the direction of the light propagation.

The eigenfunctions in
$$(r, \theta, z)$$
 coordinates are [56]

$$\psi_{nlm}(r,\theta,z) = C_{nlm} J_m(r\tilde{\alpha}_{lm}/a) \cos(m\theta) \cos(\pi nz/h), \quad (B1)$$

where C_{nlm} is the normalization factor, *h* is the height of the cylinder, and $\tilde{\alpha}_{lm}$ is the *l*th zero of the Bessel function $J_m(x)$ of order *m*. These eigenfunctions are described by three integer quantum numbers: *n* describes the localization in the *z* direction, and *m* and *l* describe that along the orthogonal directions.

The energies of these eigenstates are given by the expression

$$\mathcal{E}_{nlm} = \frac{\hbar^2}{2m_e} \left(\frac{\tilde{\alpha}_{lm}^2}{a^2} + \frac{4\pi^2 n^2}{h^2} \right). \tag{B2}$$

Because the light is assumed to be linearly polarized in the z direction, only z components of the dipole moments

$$\mu_{nlm,n'l'm'} = \langle \psi_{nlm} | z | \psi_{n'l'm'} \rangle \tag{B3}$$

play a role, and they can be calculated as

$$\mu_{nlm,n'l'm'} \propto \frac{2nn'[(-1)^{n+n'}-1]}{(n-n')^2(n+n')^2} \delta_{ll'} \delta_{mm'}, \qquad (B4)$$

where we omitted for simplicity a prefactor which comes from the normalization of the wave functions.

APPENDIX C: DERIVATION OF EQ. (1)

As we see in the main paper (see also below), the main role in the super-resonance is played by the transitions close to the Fermi energy \mathcal{E}_F . For not very small nanostructures, this corresponds to relatively large *l* and *n*. For large *n* and *l* an analytical estimation

$$\alpha_{nl} \approx (2n+l)\pi/2 \tag{C1}$$

is possible. Based on this, the energy difference between the allowed transitions $\Delta l = \pm 1$ near the energy \mathcal{E} is

$$\Delta \mathcal{E} = \pi \sqrt{\mathcal{E} \mathcal{E}_0}.\tag{C2}$$

Near the Fermi energy $\mathcal{E} \approx \mathcal{E}_F$, substituting Eq. (A2), we obtain Eq. (1), with $\hbar \omega_{\text{conf}}$ identical to $\Delta \mathcal{E}$.

We note that Eq. (C1) is valid for $n \gg l$. In contrast, for $n \approx 1$ we have $\alpha_{nl} \approx (2n + l)$, that is, the positions of the resonances would be shifted in this latter case by the factor $\approx \pi/2$ to the lower frequencies. The resonances shown in Fig. 2 and Fig. 5 in Appendixes coincide well with Eq. (C1), indicating that large *n* are involved (see also Appendix G).

In addition to the derivation of the position of the superresonance, we are able to approximately deduce its natural shape and width. For this, see Appendix G below.

APPENDIX D: LINEAR AND NONLINEAR SUSCEPTIBILITIES OF A SINGLE NANOSTRUCTURE

The corresponding expressions for $\chi^{(1)}$ and $\chi^{(3)}$ are obtained by the method of iterations [46]: The evolution of the density matrix ρ in the presence of damping can be, under suitable approximations, written as

$$\dot{\rho} = -\frac{i}{\hbar}[\hat{H}, \rho] - \Gamma(\rho - \rho^{(\text{eq})}), \qquad (D1)$$

where \hat{H} is the full Hamiltonian, $\rho^{(eq)}$ is the equilibrium value for ρ , and Γ describes the decay. The Hamiltonian consists of the action of the potential well H_0 (see the main paper) as well as the action of the field $V = e\mathbf{r}\mathbf{E}$ (in dipole approximation). In terms of the eigenfrequencies of \hat{H}_0 , $\omega_{mn} = (\mathcal{E}_m - \mathcal{E}_n)/\hbar$, Eq. (D1) can be rewritten in terms of perturbations as

$$\dot{\rho}_{mn} = -i\omega_{mn}\rho_{mn} - \frac{\iota}{\hbar}[V,\rho] - \gamma_{mn}(\rho_{mn} - \rho_{mn}^{(eq)}), \quad (D2)$$

where $\rho_{mn} = \langle m | \rho | n \rangle$, $\rho_{mn}^{(eq)} = \langle m | \rho^{(eq)} | n \rangle$, $\gamma_{mn} = \langle m | \Gamma | n \rangle$, and $|m\rangle$ and $|n\rangle$ are eigenstates of \hat{H}_0 corresponding to eigenvalues \mathcal{E}_m and \mathcal{E}_n (note that here, in contrast to previous sections, we denote by *n* and *m* the "multi-indices," fully describing the eigenstate). One can obtain ρ iteratively, in the form of $\rho = \rho^{(0)} + \theta \rho^{(1)} + \theta^2 \rho^{(2)} + \ldots$, where θ is a formal small parameter, assuming thereby that V is a small perturbation of order θ . As an initial approximation we obtain

$$\rho^{(0)} = \rho^{(\text{eq})},\tag{D3}$$

and $\rho^{(n)}$ is related to $\rho^{(n-1)}$ as

$$\dot{\rho}_{mn}^{(n)} = -(i\omega_{mn} + \gamma_{mn})\rho_{mn}^{(n)} - \frac{i}{\hbar}[V, \rho^{(n-1)}].$$
(D4)

The nonlinear polarization of *n*th order is defined via $P_i^{(n)} = \epsilon_0 \sum_{j,k,\dots} \chi_{i;jk\dots}^{(n)} E_j E_k \dots$, where $P_i^{(n)}$ and E_i are the components of the vectors $\mathbf{P}^{(n)}$ and \mathbf{E} , respectively. $\mathbf{P}^{(n)}$ is given in terms of $\rho^{(n)}$ as $\mathbf{P}^{(n)} = -eN \operatorname{tr}(\rho^{(n)}\mathbf{r})$, where *N* is concentration of the particles. The expression for $\chi^{(n)}$ is obtained by comparing the two expressions for $\mathbf{P}^{(n)}$ above. For the first-order susceptibility $\chi_{ij}^{(1)}(\omega_p)$ we thus obtain

$$\chi_{ij}^{(1)}(\omega_p) = \chi_D^{(1)} + \frac{N}{\epsilon_0 \hbar} \sum_n \left[\frac{\mu_{an}^i \mu_{na}^j}{(\omega_{na} - \omega_p) - i\gamma_{na}} + \frac{\mu_{an}^i \mu_{na}^j}{(\omega_{na} + \omega_p) + i\gamma_{na}} \right].$$
(D5)

We note once more that *a* and *n* are multi-indices, that is, every one of them describes a particular set of quantum numbers *n*, *l*, and *m* fully characterizing the system; $\omega_{mn} = \mathcal{E}_{mn}/\hbar$, $\gamma_{mn} = \delta_{mn}\gamma$ (δ_{mn} is a Kronecker symbol), $\gamma = 1/T_2$; T_2 and T_1 are given in the main paper. For the isotropic case we consider here, the indices *i* and *j* in Eq. (D5) are disregarded. According to Eq. (D3), $\rho_{ll}^{(0)}$ describes the unperturbed populations (see more details below). The first term in Eq. (D5) describes the Drude dispersion. It must be introduced into Eq. (D5) as an additional phenomenological term since its proper first-principle treatment is possible only if electron-phonon interactions [57] are taken into account, which is not the case in our approach:

$$\chi_D^{(1)} = \chi_\infty - \frac{\omega_{pl}^2}{\omega(\omega + i\gamma_D)},\tag{D6}$$

with $\omega_{pl} = Ne^2/(\epsilon_0 m_e)$ and the effective phenomenological quantities are taken as $\chi_{\infty} = 8.84$, $\gamma_D = 0.067 \text{ eV}$, and $N = 5.9 \times 10^{28} \text{ m}^{-3}$ as given in [58].

For the third-order susceptibility we have

$$\begin{split} \chi_{kjlh}^{(3)}(\omega_{p} + \omega_{q} + \omega_{r}; \omega_{r}, \omega_{q}, \omega_{p}) &= \frac{N}{\epsilon_{0}\hbar^{3}} \mathcal{P}_{I} \sum_{vnnl} \rho_{ll}^{(0)} \\ &\times \left\{ \frac{\mu_{lv}^{k} \mu_{v}^{l} \mu_{m}^{l} \mu_{ml}^{l}}{\left[(\omega_{vl} - \omega_{p} - \omega_{q} - \omega_{r}) - i\gamma_{vl} \right] \left[(\omega_{nl} - \omega_{p} - \omega_{q}) - i\gamma_{nl} \right] \right] (\omega_{nl} - \omega_{p}) - i\gamma_{nl} \right] \\ &+ \frac{\mu_{lv}^{h} \mu_{m}^{k} \mu_{ml}^{l} \mu_{ml}^{l}}{\left[(\omega_{nv} - \omega_{p} - \omega_{q} - \omega_{r}) - i\gamma_{nv} \right] \left[(\omega_{nv} - \omega_{p} - \omega_{q}) - i\gamma_{mv} \right] \left[(\omega_{vl} + \omega_{p}) + i\gamma_{vl} \right] \right] \\ &+ \frac{\mu_{lv}^{l} \mu_{v}^{k} \mu_{mn}^{l} \mu_{ml}^{l}}{\left[(\omega_{nv} - \omega_{p} - \omega_{q} - \omega_{r}) - i\gamma_{nv} \right] \left[(\omega_{vn} + \omega_{p} + \omega_{q}) + i\gamma_{vm} \right] \left[(\omega_{nl} - \omega_{p}) - i\gamma_{ml} \right] \right] \\ &+ \frac{\mu_{lv}^{l} \mu_{v}^{l} \mu_{v}^{k} \mu_{mn}^{l} \mu_{ml}^{l}}{\left[(\omega_{mn} - \omega_{p} - \omega_{q} - \omega_{r}) - i\gamma_{mn} \right] \left[(\omega_{vn} + \omega_{p} + \omega_{q}) + i\gamma_{vl} \right] \left[(\omega_{vl} + \omega_{p}) + i\gamma_{vl} \right] \right] \\ &+ \frac{\mu_{lv}^{l} \mu_{v}^{l} \mu_{v}^{k} \mu_{mn}^{l} \mu_{ml}^{l}}{\left[(\omega_{vn} - \omega_{p} - \omega_{q} - \omega_{r}) - i\gamma_{mn} \right] \left[(\omega_{nl} - \omega_{p} - \omega_{q}) - i\gamma_{ml} \right] \left[(\omega_{vl} + \omega_{p}) + i\gamma_{vl} \right] \right] \\ &+ \frac{\mu_{lv}^{l} \mu_{v}^{l} \mu_{vm}^{k} \mu_{mm}^{l} \mu_{ml}^{l}}{\left[(\omega_{vn} + \omega_{p} + \omega_{q} + \omega_{r}) + i\gamma_{vm} \right] \left[(\omega_{ml} - \omega_{p} - \omega_{q}) - i\gamma_{mn} \right] \left[(\omega_{vl} + \omega_{p}) + i\gamma_{vl} \right] \right] \\ &+ \frac{\mu_{lv}^{l} \mu_{v}^{l} \mu_{vm}^{l} \mu_{mm}^{l} \mu_{ml}^{l}}{\left[(\omega_{mn} + \omega_{p} + \omega_{q} + \omega_{r}) + i\gamma_{mm} \right] \left[(\omega_{vn} + \omega_{p} - \omega_{q}) - i\gamma_{mn} \right] \left[(\omega_{vl} - \omega_{p}) - i\gamma_{ml} \right] \right] \\ &+ \frac{\mu_{lv}^{l} \mu_{v}^{l} \mu_{vm}^{l} \mu_{mm}^{l} \mu_{ml}^{l}}{\left[(\omega_{mn} + \omega_{p} + \omega_{q} + \omega_{r}) + i\gamma_{mm} \right] \left[(\omega_{vm} + \omega_{p} + \omega_{q}) + i\gamma_{vm} \right] \left[(\omega_{vl} - \omega_{p}) - i\gamma_{ml} \right] \right] } \\ &+ \frac{\mu_{lv}^{l} \mu_{v}^{l} \mu_{vm}^{l} \mu_{mm}^{l} \mu_{ml}^{l}}{\left[(\omega_{mn} + \omega_{p} + \omega_{q} + \omega_{r}) + i\gamma_{mm} \right] \left[(\omega_{vm} + \omega_{p} + \omega_{q}) + i\gamma_{vm} \right] \left[(\omega_{vl} + \omega_{p}) + i\gamma_{vl} \right] \right] } \right\}, \quad (D7)$$

where \mathcal{P}_I denotes permutations of the frequencies ω_p , ω_q , and ω_r with the Cartesian indices h, i, and k permuted simultaneously.

As it was mentioned in the main paper, every term in the expressions for $\chi^{(1)}$ and $\chi^{(3)}$ can be seen as a sum over all transitions through the intermediate virtual states [46], with the initial and final state being the same. Since we do not consider effects of finite temperatures here, the initial populations are taken in the form

$$\rho_{ll}^{(0)} = \begin{cases} 1, & \text{if } \mathcal{E} \leq \mathcal{E}_F; \\ 0, & \text{if } \mathcal{E} > \mathcal{E}_F. \end{cases}$$
(D8)

Moreover, in Eq. (D7), due to the Pauli principle, we keep only the transitions over the intermediate virtual levels which are outside of the "Fermi sea," that is, with $\mathcal{E} > \mathcal{E}_F + \mathcal{E}_0$, where \mathcal{E}_0 is the ground state. In Eq. (D5), in contrast to Eq. (D7), this preselection happens automatically. To take into account the finite depth of our potential, we also do not consider levels with $\mathcal{E} > \mathcal{E}_W + \mathcal{E}_F + \mathcal{E}_0$, where \mathcal{E}_W is the work function. For this paper, we have taken $\mathcal{E}_W = 5.1 \text{ eV}$, $\mathcal{E}_F = 5.53 \text{ eV}$.

For practical computations, in Eqs. (D5) and (D7) we use the radial parts of the dipole moments given by Eq.(A4), and average over angular parts [16]. This is possible if assuming that only the transitions with $l \gg 1$ are relevant, which is indeed the case even for smallest diameters we consider, as one can see in Fig. 1. In this situation, averaging over the angular dependencies for $-l \leq m \leq l$ gives [16] the constant factor $\mathcal{A} = 1/3$ for the linear susceptibility $\chi^{(1)}$ and $\mathcal{A}' = 2/15$ for third-order susceptibility $\chi^{(3)}$. Furthermore, when calculating $\chi^{(3)}$, we take into account a population-induced correction factor T_1/T_2 (see [16]). As a result, the susceptibilities obtained in Eqs. (D5) and (D7) are corrected as

$$\chi^{(1)} \to \mathcal{A}\chi^{(1)}, \ \chi^{(3)} \to \frac{T_1}{T_2}\mathcal{A}'\chi^{(3)}.$$
 (D9)

APPENDIX E: LINEAR SUSCEPTIBILITY FOR DIFFERENT DIAMETERS

Whereas in Fig. 1(a) only two particular examples of the linear susceptibility for spherical nanoparticles are shown, in Figs. 5(a) and 5(b) more examples are given, to illustrate further the dependence of the super-resonance position on the particle size.

APPENDIX F: INFLUENCE OF OTHER MECHANISMS

Although the approach presented in our paper is rather universal and largely material independent, in the main paper we, to be specific, considered the parameters of gold as our basic case since gold has the most practical importance. However, gold is quite a "complicated" metal in the sense that many other mechanisms contribute to nonlinearity. Besides, in all metals, in addition to quantum confinement (billiard part) the Drude part of $\chi^{(1)}$ is contributing. In this section we will clarify in more detail the question of how other mechanisms influence the overall nonlinear and linear response.

1. Drude part of $\chi^{(1)}$

In Fig. 2 we show the linear response without the influence of the Drude part of $\chi^{(1)}$ [the first part in the left side of Eq. (D5) above, also described by Eq. (D6) above]. The influence of the Drude part is much larger than the confinement-based part, especially at small frequencies in the MIR and terahertz range. In particular, Figs. 5(c) and 5(d) show the Drude part alone and together with the quantum confinement part of the linear susceptibility for few particular diameters. One can clearly see that the Drude part absolutely dominates in the linear response at low frequencies. However, this does not mean that the confinement-based super-resonance considered in this paper "disappears" as we take into account Drude. It can be still deconvoluted and separated from the Drude part [34]. Besides, at it was shown in the main text, it is directly visible when considering the nonlinear optical properties such as Kerr effect.

2. Effect of interband transitions on the nonlinear response

In the main paper, we considered a rather simplified band gap consisting only of one band. Whereas such approximation is very well suitable for some materials such as alkali metals, for other materials such as gold it could be claimed to be a rather bad approximation. However, at least in the particular case of gold, as soon as we consider low-frequency response, it can be shown that the nonlinearity due to interband transitions is much smaller than the one due to the billiard resonances.

In the case of gold [59], there is a strong resonance in the optical response around 2.4 eV, responsible for the transitions from the 5d valence band to the 6sp conduction band, as well as a number of less pronounced resonances in the range between around 2 and 10 eV. That is, the effects of the interband transitions might be pronounced for the photon energies above 1 eV. Even at the frequency resonant with the interband transition the impact of confinement-based resonances to the rectification-like FWM processes we consider is one or two orders of magnitude larger than the effect of the interband transitions, as we will see in the next paragraphs.

Let us first consider the Kerr nonlinearity. Experimentally measured Kerr susceptibility for the photon energy around 2.3 eV, that is, close to resonance of the above mentioned transition, is (for short, 100-fs-scale pulses) $\chi^{(3)}(\omega; \omega, \omega, -\omega) \approx$ 10^{-18} m²/V² (see for instance [11]; note that in many other references long, picosecond pulses are considered, demonstrating higher nonlinearity as discussed in the subsection below). These measurements, of course, include all effects simultaneously, in particular intraband transitions and confinement-induced effects. We see that our calculations give the same order of magnitude of susceptibility for this frequency [see Fig. 3(a)] with only the confinement-based nonlinearity included. This means that the interband transitions do not dominate the intraband even at the interband resonance frequency; the impact of confinement-based resonances is of the same order of magnitude or higher.



FIG. 6. Real (a, c, e) and imaginary (b, d, f) part of the nonlinear susceptibility $\chi^{(3)}(\delta; \omega, \omega, -2\omega + \delta)$ for different diameters (see legend), corresponding to Fig. 3(b), effectively taking into account (orange curves) and without taking account (blue lines) the interband transitions as introduced by Eq. (F1). The latter correspond to Fig. 3(b).

On the other hand, as we decrease the frequency from the interband resonance towards terahertz range, the influence of the interband resonance quickly decreases whereas the influence of the confinement-based resonance increases [see Fig. 3(a)]. Therefore, we come to the conclusion that in the terahertz range the Kerr nonlinearity $\chi^{(3)}(\omega; \omega, \omega, -\omega)$ is indeed dominated by the confinement-based (intraband) transitions.

The same is also true for the FWM processes responsible for rectificationlike effects considered in the main paper: the influence of the interband transitions must be significantly smaller than the intraband (confinement-based) ones. In order to make our estimation more quantitative at this point, we use the estimation technique described in [60]. We start from the Kerr process $\chi^{(3)}(\omega; \omega, \omega, -\omega)$ assuming it to be fully resonant to the interband transition ("worst-case scenario," where the interband action is maximal). We consider then the processes $\chi^{(3)}(\delta; \omega + \delta, \omega, -2\omega)$ and $\chi^{(3)}(\delta; \omega, \omega, -2\omega + \delta)$. Because of the missing resonant terms in Eq. (D7) (which gives a factor $\sim \omega^2$ in comparison to the interband resonant case) and also because of smaller population of the virtual levels (factor $\sim T_2^2$), the nonlinear susceptibility is reduced by a factor $(\omega T_2)^2 \approx 10^2$ comparing to the Kerr susceptibility at the interband resonance. Since, as it was established before, even at the interband resonance the confinement-based Kerr nonlinearity is at least of the same order of magnitude as the interband-induced Kerr nonlinearity, we therefore conclude that for the FWM processes the intraband (confinementbased) resonances are at least by the factor of 100 larger than the interband ones. Based on our calculations of the intraband nonlinearities [see Fig. 3(b)], we can estimate the interband effect to the nonlinear susceptibility for the considered FWM processes as 10^{-20} – 10^{-21} m²/V² for ω at the interband resonance (and even lower away from that resonance). This is much less than the confinement-based impact as shown in Fig. 3.

As an alternative and fully independent method to estimate the impact of the interband transitions we introduce a gap directly into our numerical model. That is, we modify our single-band structure as the following:

$$\mathcal{E}_{nl} = \begin{cases} \mathcal{E}_0 \alpha_{nl}^2, & \text{if } |\alpha_{nl}| \leq \frac{2\pi}{\Lambda}, \\ \mathcal{E}_0 \alpha_{nl}^2 + E_g, & \text{if } |\alpha_{nl}| > \frac{2\pi}{\Lambda}, \end{cases}$$
(F1)

where Λ is the lattice constant (for gold $\Lambda \approx 4$ Å), $E_g \approx 2.4$ eV. This modification mimics the band gap which opens near the edges of the Brillouin zone. The comparison of two calculations, that is, using the single band Eq. (A1) model and two band model Eq. (F1), is shown in Fig. 6 for different exemplary diameters. One can see that, whereas the interband transition does provide some limited modification to the confinement-based dynamics for very small nanoparticles $d \leq 3$ nm, this influence quickly decreases and becomes negligible for larger diameters.

3. Thermal, hot-electron, and related effects

Taking into account temperature introduces several effects, which were neglected in the main paper. First of all, it leads to an additional hot-electron contribution into nonlinearly [11], which can overcome, by several orders of magnitude, the



FIG. 7. Real (a, c) and imaginary (b, d) part of the linear (a, b) and nonlinear (Kerr) (b, c) susceptibility for the structure d = 2.75 nm assuming sharp drop of the conducting electron density at the energy $E = \mathcal{E}_F$ (blue lines), corresponding to T = 0 K, and taking into account more smooth Fermi-Dirac distribution for T = 300 K (orange lines).

nonlinearities considered in this paper up to now. The hot electron mechanism involves laser-induced intraband excitation in the conduction band, followed by the energy dissipation of the excited electrons. This process leads to a modification of Fermi-Dirac distribution which depends on the frequency and intensity of the pump, leading thereby to frequency-dependent nonlinearity [61–63]. The key role in the quick thermalization is played by the electron-electron and electron-phonon interactions.

This nonlinearity has relatively slow, subpicosecond-scale, turn-on time [64], and therefore its influence quickly decreases with decreasing of the pulse duration [11]. For the pulses considered here (10-30 fs) this nonlinearity plays a negligible role. Indeed, the experimentally measured Kerr nonlinearity for gold nanospheres [11,65] for the pulses of 100-fs duration corresponds, by the order of magnitude $(\approx 10^{-18} \text{m}^2/\text{V}^2$, see also discussion in the previous subsection), to our calculations at around the same frequency [see Fig. 3(a)]. Since in our calculations we do not take into account the hot electron nonlinearities, we come to the conclusion that for short pulses such nonlinearities are pretty much negligible in comparison to the confinement-based resonances, or at least do not play a dominant role. This is even more true if we consider lower frequencies towards the terahertz range, since the confinement-based nonlinearity has a resonance at low frequencies, whereas the thermal nonlinearity is not expected to demonstrate a resonant behavior.

A part of the above-described thermalization process is the electron-electron interaction. Fast thermalization, indeed, is the primary consequence of the electron-electron interactions [63,66]. Electron-electron interactions are trackable in the linear properties of the nanostructure (see for instance [61]);

however, the corresponding modification is rather minor and even this small modification starts to be visible at the time scale of a few tens of femtoseconds.

Another thermal effect is the overall nonrectangular shape of the electron distribution near the Fermi zone edge as soon as the temperature is nonzero (in our calculations in the main paper we assumed zero temperature). The effect of nonzero temperature in the vicinity of the Fermi level is shown in Fig. 7 for an exemplary diameter d = 2.75 nm and temperature T =300 K. It is obtained by modifying $\rho_{ll}^{(0)}$ from Eq. (D8) to the Fermi-Dirac distribution for the finite temperature. One can see that this modifies only slightly the linear response. The nonlinear response is also modified quite moderately.

APPENDIX G: LEVEL STATISTICS DETAILS

1. General definitions

In this section we describe different variants of the level statistics, extending the discussion related to Fig. 2. To understand the mechanisms governing the formation of one single super-resonance it is very constructive to consider the level statistics, varying the selection rules included into that statistics. Commonly in the quantum billiard and quantum chaos theory [36] one considers the so-called neighboring level statistics. That is, we consider the difference between the neighboring levels $\delta \omega_i = \omega_{i+1} - \omega_i$, where the eigenfrequencies ω_i are obtained by ordering of the eigenfrequencies in the increasing order, that is, we order them in such a way that $\omega_j \leq \omega_i$ for j < i. In the case of nanospheres the corresponding eigenvalues are $\hbar \omega_{nl} = \mathcal{E}_{nl}$ [see Eq. (A1)], rearranged accordingly. Note that *i* and *j* here are single indices (and not multi-indices as in Appendix D). Then, the



FIG. 8. Different variants of level statistics for the sphere of d = 7 nm diameter [cf. Fig. 1(a)]. (a) $P(\omega_{ij})$ weighted as given by Eq. (G1) with $\Delta_{ij} = 1$, that is not taking into account Fermi sea condition Eq. (G2) (delivering thereby the statistics for dielectrics). Inset to (a): Repetition of the level statistics given by the red bars in Fig. 1(a) Fig. 1 on the larger energy scale, that is, $P(\omega_{ij})$ weighted as given by Eq. (G1) with Δ_{ij} given by Eq. (G2). (b) The same as in (a) but assuming in addition $\mu_{ji} = 1$. (c) The same as in (b) but assuming $w_{ji} = 0$ if $\mu_{ij} = 0$ and $w_{ij} = 1$ otherwise. (d) The same as in (c) but taking into account the Fermi sea restriction Eq. (G2).

probability $P(\omega)d\omega$ that $\delta\omega_i$ is located in the range between ω and $\omega + d\omega$ is calculated. The easiest way to visualize such statistics is to use the histogram technique. For nanospheres this traditional neighboring level statistics is presented in Fig. 2(a) (yellow bars).

The neighboring level statistics do allow us to determine the universality classes of different billiards. It, however, does not take into account the properties of eigenfunctions and of the transition rules, so it seems to be of little use for the optical properties. In order to improve usability for optics, we extend the statistics to take into account all transitions, not only neighboring, and impose additional selection rules. That is, we consider the statistics of energy differences $\omega_{ij} = \omega_i - \omega_j$ for i > j (assuming the ordering of ω_j as discussed above). In addition, when constructing the probability density $P(\omega)$ of ω_{ij} being in the interval $[\omega, \omega + d\omega]$, we take into account the "strength" of the transition by weighting $P(\omega_{ij})$ with a weight

$$w_{ij} = \Delta_{ij} |\mu_{ij}|^2, \tag{G1}$$

where μ_{ij} is the dipole momentum of the corresponding transition $i \rightarrow j$, and Δ_{ij} is defined as

$$\Delta_{ij} = \begin{cases} 1, & \text{if } \hbar\omega_i \leqslant \mathcal{E}_F \text{ and } \hbar\omega_j > \mathcal{E}_F \\ 0, & \text{otherwise,} \end{cases}$$
(G2)

that is, takes into account the Fermi-Dirac distribution Eq. (D8) (we call it below "the Fermi sea condition"). The

resulting statistics is shown in Fig. 2 (red and blue bars) and repeated for convenience as the inset in Fig. 8(a) for the larger frequency range. We note that the traditional statistics discussed in the previous paragraph is a partial case of this more general approach; namely, we obtain the traditional neighboring level statistics assuming $w_{ij} = \delta_{i,j=i+1}$.

2. Natural shape and width of the super-resonance for nanospheres

To find out the natural width of the super-resonance for the case of nanostructures, we use a more precise version of Eq. (C1):

$$\alpha_{nl} = (2n + l + s_{nl})\pi/2,$$
 (G3)

which includes corrections s_{nl} to the values of the roots. All transitions contributing to the super-resonance are characterized by the same value of 2n + l before and after the transition, that is, 2n' + l' = 2n + l + 1. However, values of *n* and *l* can be different. Therefore the energy difference between the eigenfunctions $|\psi_{nl}\rangle$ and $|\psi_{n'l'}\rangle$ will be modified as follows:

$$\Delta \mathcal{E} = \frac{\pi^2 \mathcal{E}_0}{4} [(2n' + l' + s_{n'l'})^2 - (2n + l + s_{nl})^2]$$

$$\simeq \frac{\pi^2 \mathcal{E}_0}{2} (2n + l)(1 + s_{n'l'} - s_{nl}).$$
(G4)

/

For the typical range of n and l actual for our particular situation, we approximate s_{nl} as being distributed in the range [-0.25, 0]. For the probability distribution of the difference $s_{nl} - s_{n'l'}$, given by $P(s_{n'l'} - s_{nl}) \sim \int P(s)P(s + s_{n'l'}) P(s) P(s) P(s) P(s) P(s)$ $s_{n'l'} - s_{nl} ds$, we obtain a triangular symmetric shape with the maximum at zero and full width of 1/2, that is, we have a constant full width at half maximum of 1/4 of the distribution of the difference $s_{nl} - s_{n'l'}$. Using Eq. (C2) we obtain that this translates to the width $\approx \omega_{\rm conf}/4$ in frequency, where ω_{conf} is the position of the super-resonance as defined in Eq. (1). If we consider the dipole-momentum-weighted statistics as described below, this simplified conclusion will be modified by the fact that the dipole momentum depends on the energy level difference. Such dependence will result in a sharper lower-frequency shoulder of the statistics and smoother higher-frequency shoulder. Both such shape and the width of the peak in the statistics are in a surprisingly good agreement with the findings shown in Fig. 2 (for the case of large T_2). Of course, reducing the T_2 to room-temperature values additionally increases the width of the super-resonance beyond the natural width of $\approx \omega_{\rm conf}/4$.

3. Influence of different mechanisms on the statistics

Varying the selection rules incorporated into the weighting Eq. (G1), we can study how these rules influence the statistics. Different variants of statistics are shown in Fig. 8. In particular, in Fig. 8(a) we consider $P(\omega_{ij})$ weighted by Eq. (G1) with $\Delta_{ij} = 1$, that is not taking into account the Fermi sea condition; i.e., all transitions, not only the transitions from below to above the Fermi sea level, are allowed. This situation describes dielectrics rather than metals, and should be contrasted to the "metallic case," that is, the situation where also the Fermi sea condition is satisfied [inset to Fig. 8(a) as well as Fig. 2(a)]. One can see that in the former case the resonance is much more broad and, in addition, noticeably shifted to the lower frequencies.

On the other hand, if we remove all restrictions at all, that is, consider $w_{ii} = 1$ for all i and j, we will see very broad distribution over the scale of many eV [Fig. 8(b)]. The statistics changes not too significantly if we take into account only allowed transitions (but not yet distinguishing between the strengths of the transitions, that is, assuming $w_{ii} = 0$ if $\mu_{ii} = 0$ and $w_{ii} = 1$ otherwise, and also not taking into account the Fermi sea condition). Yet, in this case the peak, corresponding to the low-frequency super-resonance (at around 0.5 eV), does already appear [see Fig. 8(c)]. This peak becomes even sharper if we in addition take into account the Fermi sea condition as shown in Fig. 8(d). But, in addition to this low-frequency peak, in Fig. 8(d) we see also many other peaks at higher frequency. These many peaks disappear as we take into account the "strengths" of the transitions $[\mu^2]$ weighting as given by Eq. (G1); see inset to Fig. 8(a) and Fig. 2(a)].

Therefore, we conclude that there are, in general, many possible confinement-based transitions at low and high frequencies. The key role in the formation of a single superresonance is played by the dipole moments, that is, by the symmetry and composition of the wave functions, and only to a lesser extent by the Fermi sea condition. As we take them into account, we are left with a single bunch of closely spaced resonances, which, taking into account that each of these resonances is broadband, merge into a super-resonance. Whereas the consideration here was focused on the case of nanospheres, we remark that, most probably, this particular situation is quite geometry independent, at least for regular billiards. Indeed, in this case we expect that distantly spaced eigenfunctions have very different numbers of oscillations in every spatial direction, making μ_{ij} small. This, however, is not necessarily the case for the irregular, chaotic billiards which have also rather chaotic eigenfunctions. Finally, as Fig. 2(b) shows, we can tune the positions of the resonances in such a way that the resulting super-resonance is broadened.

APPENDIX H: EFFECTIVE PROPERTIES OF THE NANOCOMPOSITE

The effective linear properties of the nanocomposite for arbitrary frequency ω are calculated using the effective medium approach [49] using the linear properties of the host $\varepsilon_h(\omega)$ (in our case SiO₂), and nanostructure $\varepsilon_{\text{NP}}(\omega)$ [which is given by $\varepsilon_{\text{NP}}(\omega) = 1 + \chi^{(1)}$, where $\chi^{(1)}$ is calculated according to Eq. (D5)], as

$$\varepsilon_{\rm eff}(\omega) = \varepsilon_h(\omega) + f_X(\omega)[\varepsilon_{\rm NP}(\omega) - \varepsilon_h(\omega)] \tag{H1}$$

where *f* is the filling factor and $x(\omega)$ is defined as

$$x(\omega) = \frac{3\varepsilon_h(\omega)}{\varepsilon_{\rm NP}(\omega) + 2\varepsilon_h(\omega)}.$$
 (H2)

We note that at Mie resonance |x| is especially large. Finally, the effective nonlinear susceptibility $\chi_{eff}^{(3)}$ for every process is calculated for given linear and nonlinear properties of the nanostructures and host as follows:

$$\chi_{\rm eff}^{(3)} = f \chi_{\rm NP}^{(3)} x(\omega_0) x_{\rm (}\omega_1)^2 x(\omega_2), \tag{H3}$$

where $\chi_{NP}^{(3)}$ is the nonlinear susceptibility given by Eq. (D5) or Eq. (6), with $\chi^{(3)}$ in those expressions calculated using Eq. (D7).

APPENDIX I: PROPAGATION EQUATIONS

Assuming slowly varying envelope approximation and neglecting nonlinear effects for the pump waves, the governing equations are

$$\frac{\partial A_0}{\partial z} = -\frac{1}{2c\varepsilon_0} \frac{\partial P}{\partial t} = -\frac{i\delta\chi_{\text{eff}}^{(3)}(\delta)}{2c} A_1^2(z) A_2^*(z) - \alpha_0 A_0, \quad (I1)$$
$$\frac{\partial_z A_n = ik_n A_n - \alpha_n A_n, \ n = 1, 2, \quad (I2)$$

where *c* is the speed of light in vacuum, ε_0 is vacuum permittivity, and A_i , $\alpha_i k_i$, i = 0, 1, 2 are correspondingly the slow (complex) amplitudes, linear losses, and wave vectors for the *n*th harmonic (here the signal is assumed to be the zeroth harmonic), and $\chi_{\text{eff}}^{(3)}(\delta)$ is the effective nonlinear susceptibility for the corresponding process. Equations (I1) and (I2) allow an analytical solution, given by

$$A_0 = -iAe^{-\alpha_0 L}(e^{\kappa L} - 1)/\kappa, \qquad (I3)$$

where $A = \delta \chi_{\text{eff}}^{(3)} A_1^2(0) A_2^*(0)/2c$, *L* is the propagation distance, and $\kappa = i(k_0 + 2k_1 - k_2) - 2\alpha_1 - \alpha_2 + \alpha_0$.

Assuming pulse durations of around 30 fs, we must consider two regions for the pump where conversion works significantly differently. For the signal in the terahertz range $(\delta/2\pi \leq 30 \text{ THz})$, the frequencies $j\omega_0$ and $j\omega_0 + \delta$ (j = 1, 2) are both located within the spectrum of the pump. The non-linearity in this case is driven by three contribution types mentioned above, and must be considered as the following sum:

$$\chi_{\rm NP}^{(3)}(\delta) \approx \chi^{(3)}(\delta; \omega_0, \omega_0, -2\omega_0 + \delta) + \chi^{(3)}(\delta; \omega_0 + \delta, \omega_0, -2\omega_0) + \chi^{(3)}(\delta; \omega_0, \omega_0 + \delta, -2\omega_0).$$
(I4)

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In contrast, for the signal in MIR range $\delta/2\pi > 30$ THz, the components $\omega_0 + \delta$ and $2\omega_0 + \delta$ are not within the pump spectrum anymore. In this case, in order to make the conversion efficient, we must, for instance, shift the second harmonic: $2\omega_0 \rightarrow 2\omega_0 + \delta$. The only effective nonlinear process in this case is

$$\chi_{\rm NP}^{(3)}(\delta) = \chi^{(3)}(\delta; \omega_0, \omega_0, -2\omega_0 + \delta).$$
 (I5)

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