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Cite as: Appl. Phys. Lett. 116, 152402 (2020); https://doi.org/10.1063/1.5143311
Submitted: 21 December 2019 . Accepted: 27 March 2020 . Published Online: 16 April 2020

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ABSTRACT

We find that in the ferromagnetic semiconductor EuS, near its Curie temperature, a single band edge photon generates a spin polaron (SP), whose magnetic moment approaches 20 000 Bohr magnetons. This is much larger than the supergiant photoinduced SPs in antiferromagnetic europium chalcogenides, reported previously. The larger SP in ferromagnetic EuS, and still larger expected for EuO, is explained by a larger Bohr radius of the photoexcited electron state, which encircles and polarizes a greater number of lattice spins. However, because the wave function of the photoexcited electron spreads over a greater volume, the photoexcited electron’s exchange interaction with individual lattice spins weakens, which makes the SP more easily quenched thermally.

Uncovering new efficient techniques by which light magnetizes materials is an interesting scientific modern topic, and paves the way for novel optomagnetic devices. Recently, an ultrastart magnetization mechanism based on the photoinduction of spin polarons for intrinsic antiferromagnetic europium chalcogenides EuTe and EuSe was reported. The magnetization process is described as follows. A photon resonant with the bandgap generates a randomly oriented spin polaron (SP). The magnetic moment of the photoinduced SP is of the order of 600 Bohr magnetons in EuTe and 6000 in EuSe. These photoinduced SPs are orders of magnitude larger than those seen in diluted magnetic semiconductors or detected by muon investigations, which are of a few tens of Bohr magnetons only. Due to the large magnetic moment of the SPs in EuTe and EuSe, a very small magnetic field is sufficient to align their spin with the field, magnetizing the sample. The remarkable characteristic of this process is that the absorption of a single photon leads to the spin alignment of thousands of electrons, thus opening the perspective of controlling magnetism with very low intensity light.

In this paper, we demonstrate that in the ferromagnetic members of the europium chalcogenide series, a single photon generates an even larger SP than in the antiferromagnetic ones, EuO and EuS, than in the antiferromagnetic ones, EuSe and EuTe, is not entirely surprising. The SP is in essence an exciton, well known in diamagnetic semiconductors, but with the difference that in a magnetic semiconductor, an exchange interaction between carriers and lattice spins operates. This interaction favors ferromagnetic arrangement of the lattice spins, encompassed by the exciton wave function. In EuX (X=O, S, Se, Te), the exciton’s hole belongs to the strongly localized 4f shell of an europium atom. In contrast to the hole, the electron in the exciton extends over many lattice parameters, and is described by a Bohr envelope wave function, with an effective Bohr radius

\[ a_B \approx \frac{\varepsilon R^2}{m^* k_e^2}, \]

where \( \varepsilon \) is the dielectric constant of the host material, \( m^* \) is the effective mass of the electron in the conduction band, \( k = 9.0 \times 10^9 \text{ N} \cdot \text{m}^2/\text{C}^2 \) is Coulomb’s constant, and \( e = 1.60 \times 10^{-19} \text{ C} \) is the elementary charge. The Coulomb attraction by the hole is crucial to stabilize the SP. However, the strong intra-atomic localization of the hole suppresses its exchange interaction with the surrounding lattice spins, and the hole does not participate in the lattice spin polarization. The strong localization of the hole also makes the SP immobile.

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We can roughly estimate the potential size of photoinduced SPs in EuX by assuming full ferromagnetic alignment within the exciton’s Bohr sphere, which gives a magnetic moment of

$$\mu_{\text{pol}} \sim \frac{4}{3} \pi a_0^3 N \mu_B,$$

where $\mu_{\text{pol}} = 7 \mu_B$ is the magnetic moment of an Eu atom, $\mu_B = 9.27 \times 10^{-24}$ A m$^2$ is Bohr’s magneton, and $N = \frac{\pi}{a}$ is the density of Eu atoms in the EuX face-centered cubic lattice of parameter $a$.

The effective mass was taken to be $m^* = 0.3$, in units of the free electron mass, $m_0 = 9.1 \times 10^{-31}$ kg, for all members of the EuX family.\textsuperscript{16,17} Table I shows the remaining parameters entering Eqs. (1) and (2), and the estimated photoinduced SP produced by Eq. (2). Also shown in Table I is the measured maximum size of photoinduced SPs. Table I shows that, for ferromagnetic EuO, Eq. (2) predicts an SP approaching 100 000 Bohr magnetons. This hypergiant size is a consequence of the increasing ratio between the effective Bohr radius and the lattice constant, the greater the ratio, the greater the number of Eu spins within the Bohr sphere that can be potentially polarized to form an SP.

To test our predictions, photoinduced SPs were investigated in an epitaxial EuS sample, grown on a BaF$_2$ (111) substrate. The BaF$_2$ substrate was cleaved of from a bar along the (111) plane in air, and cleaned by dry nitrogen gas before being loaded into an UHV chamber. The substrate was annealed at 800°C for 30 min to clean the surface, and then kept at 600°C. EuS (99.9%) was deposited at 600°C from Ref. 13 and (2), and the estimated photoinduced SP produced by Eq. (2). Also shown in Table I is the measured maximum size of photoinduced SPs. Table I shows that, for ferromagnetic EuO, Eq. (2) predicts an SP approaching 100 000 Bohr magnetons. This hypergiant size is a consequence of the increasing ratio between the effective Bohr radius and the lattice constant, the greater the ratio, the greater the number of Eu spins within the Bohr sphere that can be potentially polarized to form an SP.

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Because of the lattice mismatch of less than 4%, (lattice constant $a = 5.968$ Å and $a = 6.196$ Å for EuS and BaF$_2$, respectively), the epitaxial layers are expected to have a high structural quality. Our \textit{in situ} Reflective High-Energy Electron Diffraction (RHEED) confirmed the epitaxial growth of EuS on the BaF$_2$ (111) substrate (see Fig. 1). The sharp streaky RHEED pattern for the 5 nm thick EuS film, shown in Fig. 1(b), indicates a 2D growth. After 10 nm, due to the relaxation of strain, growth changes to a 3D mode, as revealed by the spotty RHEED pattern shown in Fig. 1(c). A 60° rotation symmetry confirmed by RHEED indicates EuS film growth along the (111) direction.

Next, we address the fundamental optical and magnetic properties of the EuX series. All members of the EuX series have a common band edge electronic energy level structure,\textsuperscript{13,18} independent of their natural magnetic arrangement. The primary effect of a change in magnetic order on the band edge energy level structure is a rigid shift of conduction band energy levels relative to the top valence state,\textsuperscript{19,20} which is a consequence of the ferromagnetic electron-lattice exchange interaction. Thus, for example, on cooling through the Curie temperature,\textsuperscript{16} the bandgap of ferromagnetic EuO shrinks due to the onset of the ferromagnetic order. Similarly, when a magnetic field imposes a ferromagnetic order, the bandgap of antiferromagnetic EuX shrinks.\textsuperscript{19,21,22}

Circular dichroism in EuX reflects the magnetic moment of the sample in the direction of light travel.\textsuperscript{23} In zero magnetic field, ferromagnetic EuX naturally forms a multiple microdomain structure, and no net magnetic moment exists. In antiferromagnetic EuX, in zero magnetic field, the magnetic moment is always zero. Therefore, circular dichroism in zero field is absent for all EuX members. However, a strong enough magnetic field turns EuX into a single ferromagnetic domain, and a huge magnetic circular dichroism develops.\textsuperscript{19,20,22,24}

To verify if our EuS sample displayed the expected circular dichroism, we measured its band edge polarized optical absorption at $T = 5$ K, and the result is shown in Fig. 2. Since $T < T_C$ for EuS (see below), the sample is in the ferromagnetic phase. Despite ferromagnetic order, circular dichroism is absent for $B = 0$, because the photon crosses many magnetic domains, oriented randomly. However, when a magnetic field imposes a single domain oriented along the light wave vector, the huge circular dichroism inherent to EuX emerges, demonstrating the excellent quality of the sample. At saturation, the absorption onset shows a $\sigma^+$ peak at 1.84 eV and a $\sigma^-$ peak at 2.13 eV. The size of the $\sigma^+ - \sigma^-$ splitting is nearly the same as reported for other EuX members.\textsuperscript{19,24} From the spectrum at $B = 0$, the EuS bandgap is estimated to be 1.65 eV (see the inset of Fig. 2). For europium chalcogenides, the bandgap is found by extrapolation of the linear absorption region to zero. The EuX bandgap determined in this

![Image](https://example-image-url.com/)

**FIG. 1.** \textit{In situ} RHEED pattern of (a) BaF$_2$ (111), (b) 5 nm EuS, and (c) 90 nm EuS along $\Gamma - K$. (d) Line profile of (a)–(c).
way is very consistent, and circumvents the shortcomings of Tauc plots or other approaches, as discussed in detail in Ref. 25. The changes in the shape of the optical spectrum with the magnetic field is a consequence of the changes in the oscillator strengths of band edge electronic transitions, due to the alignment of spin domains with the light wave vector.19,24 Nevertheless, in the case of EuS at $T = 5$ K, the energy bandgap itself does not change with $B$, because at this temperature the arrangement is always ferromagnetic.

In europium chalcogenides, in any magnetic phase, the magnetization of the sample is directly proportional to the bulk Faraday rotation angle, $\theta_F$, of light crossing the sample, if the wavelength is within the bandgap.25 The proportionality constant is almost independent of temperature; therefore, the magnetization dependence on $T$ can be inferred from $\theta_F$ measurements. Figure 3 shows the $\theta_F$ vs $T$ trace obtained for $B = 9$ mT, using a photon energy of 0.821 eV, which is well within the bandgap of 1.65 eV (see Fig. 1). Measurements started at room temperature and the sample was slowly cooled. The magnetization of the sample increased as the sample went through the Curie temperature and approached saturation, as expected for a ferromagnet, due to the appearance of spontaneous magnetization. The weak magnetic field of 9 mT was required to inhibit multiple domains that would preclude a macroscopic measurement. The Curie temperature for our sample was taken to be equal to the temperature at which the derivative $d\theta_F/dT$ is minimum, giving $T_C = 15.2 \pm 0.3$ K. The $T_C$ value for our thin epitaxial layer is slightly less than the known $T_C = 16.3$ K for bulk EuS.26

We investigated photoinduced spin polarons in the EuS sample, using a two-color pump-probe Faraday rotation experiment. Electron-hole pairs were generated by photons above the bandgap (2.33 eV), and the photoinduced Faraday rotation (PFR) was measured for photons within the bandgap (1.46 eV). For $T > T_C$, the lattice spins are uncorrelated, hence the Weiss field is absent. In this case, photoinduced SPs rotate freely. We used low pump intensities, so that the average distance between adjacent SPs is much larger than their radius, making them non-interacting. Therefore, the SP ensemble forms a paramagnetic gas, whose magnetization is described by a Langevin function.5,6

Inevitably, the pump pulse causes heating of the illuminated region, which will change the magnetization, hence it will show as PFR. Reference 6 reports a detailed investigation of the effect of pump heating on the PFR signal: the heating adds a PFR signal whose absolute value increases linearly with $B$. Such a linear component is easily distinguished from the step-like photoinduced magnetization associated with photoinduced SPs.6 The dots in Fig. 4 represent the net PFR signal, associated with photoinduced SPs at 16.5 K, as a function of the internal magnetic field. The solid line in Fig. 4 shows a fit of the net PFR signal with a Langevin function. The excellent fit yields $\mu_{\text{pol}} = 4600 \mu_B$. The Langevin fit also yields a PFR angle at a saturation of 1.64 $\mu$ rad, as shown in Fig. 4.

As discussed above, the Faraday rotation angle is proportional to the magnetization.25 On the other hand, the magnetization of the polaron ensemble, at saturation, is proportional to the magnetic moment of an individual SP. Therefore, from Fig. 4, the factor required...
to convert saturation PFR into SP magnetic moment is \(4600/1.64 = 2800 \mu_B/\mu\) rad. This factor can be used to convert the measured saturation PFR into the SP magnetic moment at all temperatures, provided that all measurements be taken for the same pump intensity. This includes \(T < T_C\), when SP orientation may be constrained by the onset of the ferromagnetic phase. Below \(T_C\), a spontaneous magnetization arises, which gives rise to a Weiss field, which may constrain the rotation freedom of the SPs, hindering Langevin magnetism.

Figure 5 shows the measured magnetic moment of the photoinduced SP in EuS, as a function of temperature. Above \(T_C\), the magnetic moment of the SP was determined from a Langevin fit to the PFR, whereas for \(T < T_C\), the SP magnetic moment was obtained from the saturation PFR, using the conversion factor of \(2800 \mu_B/\mu\) rad described above.

The photogenerated SP size shows a sharp peak near the Curie temperature. This contrasts to antiferromagnetic EuTe, where the SP maximum size is two orders of magnitude smaller, and its size varies very slowly with temperature. This enormous difference in behavior is explained by the greater effective Bohr radius in EuS than in EuTe (Table I), as follows. The phenomenon that fathers SPs is the exchange interaction between the photoexcited electron and the surrounding Eu spins. This interaction is described by an effective magnetic field, \(B_{eq}(r)\), acting on the lattice spins

\[
B_{eq}(r) = \frac{J_S S}{N \mu_B} |\psi(r)|^2,
\]

where \(J_S\) is the band-lattice exchange interaction constant, \(S\) is the Bohr envelope wave function describing the photoexcited electron, and \(r\) is the distance to the center of the SP. In EuTe, the average effective magnetic field within the SP is about 1 T, which is approximately equal to \(B_{eq}(a_B)\) (see the last column in Table I). When \(a_B\) increases, on one hand, the electron wave function covers an increasing number of spins, producing a larger SP, but on the other hand, the average effective magnetic field acting on an individual lattice spin drops, which reduces the polarization power of the photoexcited electron. As Table I shows, in EuS, the typical exchange magnetic field inside the SP is only about 0.2 T. Nevertheless, the reduced average exchange magnetic field is still enough to polarize the lattice spins in close vicinity to \(T_C\), when the magnetic susceptibility is very high. On either side of the Curie temperature, the magnetic susceptibility decreases, and the magnetic moment of an SP decreases in size. Well below \(T_C\), the spontaneous magnetization saturates, which completely quenches SP formation.

To provide support for our interpretations, we performed Monte Carlo calculations. A cube of 18 EuS lattice parameters, centered on an Eu atom, was analyzed. A randomly oriented \(S = 7/2\) spin was associated with all Eu sites in the cube. The exchange interaction between lattice spins was incorporated via the first and second neighbor exchange constants, \(J_1\) and \(J_2\), respectively. To exclude surface effects, Born-Kármán periodic boundary conditions were imposed on the cube. The exchange interaction between Eu spins and the photoexcited electron was incorporated via the effective magnetic field acting on lattice spins, \(B_{eq}(r)\), given by Eq. (3). Then every lattice spin in the cube was reoriented individually, using the Monte Carlo stochastic method; this process was repeated 1600 times at each temperature. The magnetic moment of the SP was taken to be the difference between two magnetic moment calculations of a sphere centered on the cube, one including and the other excluding the exchange field produced by the photoexcited electron (see the inset in Fig. 5). The radius of the sphere was determined from the condition \(B_{eq}(r) > B_0\). Here, \(B_0\) is the minimum value of the exchange field, produced by the photoexcited electron, required to overpower other competing interactions. One obvious competing field is the applied one. As Fig. 4 shows, measurements of PFR require the application of fields equivalent to an internal field of a few tens of milliteslas. In the region where the exchange field is much less than the applied field, the photoexcited electron loses the competition and does not polarize the lattice spins. We assumed \(B_0 = 0.005\) T. The Monte Carlo result is depicted in Fig. 5 by the full curve, using \(J_1 = 0.2\) K, \(J_2 = -0.09\) K, and \(J_S S \approx 0.3\) eV. The calculated SP magnetic moment curve shows a peak with both the maximum value and position in excellent agreement with the experiment. The parameters \(J_1, J_2\) are \(\sim 10\%\) less than values reported in the literature for bulk EuS; the small difference could be associated with residual strain in our epitaxial sample.

Next, we shall compare the measured SP size at \(T_C\) to the value expected by the molecular field (Weiss field) approximation. In this model, the magnetic moment direction of each atom in the crystal is replaced by an average over the ensemble. Then, the magnetization as a function of temperature and field is given by the transcendental equation

\[
M(B, T) = N_B B_S \left[ \frac{\mu(B + l_M)}{k_B T} \right],
\]

where \(B_S\) represents a Brillouin function of order \(S\), \(\mu\) is the absolute value of the magnetic moment of the atoms in the lattice, and \(k_B = 1.38 \times 10^{-23}\) J/K is the Boltzmann constant. The term \(l_M\) in Eq. (4) represents the molecular field, which describes the ferromagnetic coupling between spins. However, the molecular field model neglects correlated fluctuations in the orientation of different spins. Despite this drastic approximation, the molecular field model
remarkably exhibits all the main features of ferromagnetism, including the temperature dependence of the magnetization on going through a paramagnetic/ferromagnetic phase transition in nickel for example, as discussed in Ref. 30 (p. 433–434). The molecular field model also describes the magnetization dependence on the magnetic field at the Curie temperature in ferromagnets (see, for instance, Ref. 27, p. 90), which is exactly the approach we shall use here.

Expanding the Brillouin function in a power series, $B_S(x) = x_S - B_S x^3$, where $x_S = \frac{2x}{B_S}$ and $B_S = \frac{2x^2}{1 - x^2}$, and substituting $x = \frac{2T_k}{k_B T}$, Eq. (4) can be solved for the magnetization at $T = T_C$, if its basic parameters are known.

$$M(B, T_C) = N_H \left( \frac{z x_B B}{B_S k_B T} \right)^{1/3}.$$  

Substituting $B$ in (5) by $B_X(r)$, as given by Eq. (3), and integrating (5) in a sphere where $B_X(r) > B_0$, we obtain $\mu_{pol}$ at $T = T_C$, in units of the Bohr magneton

$$\mu_{pol} = 27 A \left( \frac{16\pi^2 x_S}{B_S} \right)^{1/3} \left( \frac{a_B}{a} \right)^2 \left( \frac{I_{12} S}{k_B T} \right)^{1/3} \mu_0,$$

where $A = 1 - \left( \frac{1}{2} + x_0 + 1 \right) e^{-x_0}$, and $x_0 = \frac{1}{2} \ln \frac{I_{12} S}{k_B T}$.

To estimate the SP size at $T_C$ for EuS using (6), we substitute $S = 7/2$, $\mu = \mu_{pol}$, $a_B/a = 3.28$ from Table I, the experimental value $T_C = 15.2$ K, $I_{12} S = 0.30$ eV, and $B_0 = 0.005$ T, as used in the Monte Carlo calculations, to obtain $\mu_{pol} = 20 \times 10^6$. This value agrees well with the estimates given in Fig. 5. Of course, because it is based on the exclusion of fluctuations, Eq. (6) may not be accurate, nevertheless it provides a very useful simple practical formula for a first estimate of what kind of SP size can be expected in any concentrated ferromagnetic semiconductor, if its basic parameters are known.

In conclusion, we demonstrated that in the ferromagnetic members of the europium chalcogenide series, the magnetic moment of a photoinduced spin polaron can be much larger than the supergiant spin polaron observed in the antiferromagnetic ones. However, the hypergiant SP photogeneration in ferromagnetic EuX is only efficient in a narrow temperature interval around the Curie temperature. Above $T_C$, thermal fluctuations rapidly destroy the SPs. Below the critical phase transition temperature, in the ferromagnetic members, the onset of spontaneous magnetization quenches SP formation, whereas in the antiferromagnetic ones, SPs are formed most efficiently. For EuO, the exciton effective Bohr radius is larger than in EuS, so potentially an even larger SP is possible in EuO. However, because the Curie temperature for EuO ($T_C = 69$ K) is higher, thermal quenching could be more effective in EuO than in EuS. This remains to be clarified experimentally.

This work was funded by CNPq (Project Nos. 303757/2018-3 and 420531/2018-1), FAPESP (Project Nos. 2019/04207-7 and 2019/12678-8), the National Science Foundation (NSF-DMR No. 1700137), the Office of Naval Research (No. N00014-16-1-2657), and the Center for Integrated Quantum Materials (No. DMR-1231319).

REFERENCES

[1] A. Kirilyuk, A. V. Kimel, and T. Rasing, Rev. Mod. Phys. 82, 2731 (2010).
[2] S. F. Siegert, J. A. Gessner, M. Osliander, C. Denker, Y.-P. Chang, M. C. Schröder, A. Gugenmos, N. Cui, J. Walowski, U. Martens, J. K. Dewhurst, U. Kleineberg, M. Münzenberg, S. Sharma, and M. Schulze, Nature 571, 240 (2019).
[3] A. B. Henriques, G. D. Galgano, E. Abramof, B. Diaz, and P. H. O. Rappl, Appl. Phys. Lett. 99, 091906 (2011).
[4] A. B. Henriques, F. C. D. Moraes, G. D. Galgano, A. J. Meaney, B. M. Christiansen, J. C. Maan, E. Abramof, and P. H. O. Rappl, Phys. Rev. B 90, 165202 (2014).
[5] A. B. Henriques, G. D. Galgano, P. H. O. Rappl, and E. Abramof, Phys. Rev. B 93, 201201(R) (2016).
[6] A. B. Henriques, A. R. Naupa, P. A. Usachev, V. V. Pavlov, P. H. O. Rappl, and E. Abramof, Phys. Rev. B 95, 045205 (2017).
[7] A. B. Henriques and P. A. Usachev, Phys. Rev. B 96, 195210 (2017).
[8] A. B. Henriques, X. Graten, P. A. Usachev, V. A. Chitta, and G. Springholz, Phys. Rev. Lett. 120, 217203 (2018).
[9] A. Wolf, Theory of Bound Magnetic Polarons in Semimagnetic Semiconductors, Semiconductors and Semimetals Vol. 25, edited by J. K. Furdyna and J. Kossut (Academic Press, San Diego, 1988), p. 414.
[10] M. Nawrocki, R. Planel, G. Fishman, and R. Galazka, Phys. Rev. Lett. 46, 735 (1981).
[11] V. G. Stormach, O. E. Parfenov, J. H. Brewer, P. L. R. ans, S. L. Stubbs, R. L. Lichti, G. E. D. Morenzoni, T. G. Aminov, V. P. Zlomanov, A. A. Vinokurov, R. L. Kallacher, and S. von Molnár, Phys. Rev. B 80, 235203 (2009).
[12] A. Mauger and D. L. Mills, Phys. Rev. B 31, 8024 (1985).
[13] A. Mauger and C. Godart, Phys. Rev. B 141, 51 (1986).
[14] D. Ax, Phys. Chem. Sol. 30, 1403 (1969).
[15] G. Güntherodt, J. Phys.: Condens. Matter. 16, 25 (1973).
[16] S. J. Cho, Phys. Rev. B 1, 4589 (1970).
[17] M. Umehara, Phys. Rev. B 60, 445 (1999).
[18] Wachtel, Crit. Rev. Solid State Sci. 3, 189 (1972).
[19] A. B. Henriques, M. A. Manfrini, P. H. O. Rappl, and E. Abramof, Phys. Rev. B 77, 035204 (2008).
[20] A. B. Henriques, P. H. O. Rappl, and E. Abramof, Int. J. Mod. Phys. B 23, 2769 (2009).
[21] K. Hanamoto, A. B. Henriques, N. F. Oliveira, P. Rappel, E. Abramof, and Y. Ueta, J. Phys.: Condens. Matter 16, 5597 (2004).
[22] A. B. Henriques, G. D. Galgano, B. L. Diaz, P. H. O. Rappl, and E. Abramof, J. Phys.: Condens. Matter 19, 406234 (2007).
[23] A. B. Henriques, G. D. Galgano, and E. Abramof, J. Phys.: Condens. Matter 20, 255209 (2008).
[24] A. B. Henriques, A. Wierts, M. A. Manfrini, G. Springholz, P. H. O. Rappel, E. Abramof, and A. Y. Ueta, Phys. Rev. B 72, 155337 (2005).
[25] C. P. van Kooten, P. A. Usachev, X. Graten, A. R. Naupa, V. A. Chitta, G. Springholz, and A. B. Henriques, J. Appl. Phys. 126, 095701 (2019).
[26] G. Güntherodt, J. Schoenes, and P. Wachtler, J. Appl. Phys. 41, 1083 (1970).
[27] S. Blandell, Magnetism in Condensed Matter (Oxford University Press, Oxford, 2001).
[28] H. G. Bohn, W. Zinn, B. Dorner, and A. Kollmar, J. Appl. Phys. 52, 2228 (1981).
[29] N. Goncharenko and I. Mirabeau, Phys. Rev. Lett 80, 1082 (1998).
[30] F. Reif, Fundamentals of Statistical and Thermal Physics (McGraw-Hill, 1965).