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Magneto-Optical Evidence of Double Exchange in a Percolating Lattice

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Substituting Eu by Ca in ferromagnetic EuB₆ leads to a percolation limited magnetic ordering. We present and discuss magneto-optical data of the $Eu_{1-x}Ca_xB_6$ series, based on measurements of the reflectivity $R(\omega)$ from the far infrared up to the ultraviolet, as a function of temperature and magnetic field. Via the Kramers-Kronig transformation of $R(\omega)$ we extract the complete absorption spectra of samples with different values of x. The change of the spectral weight in the Drude component by increasing the magnetic field agrees with a scenario based on the double-exchange model, and suggests a crossover from a ferromagnetic metal to a ferromagnetic Anderson insulator upon increasing Ca content at low temperatures.

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Because of the potential technological applications, materials exhibiting colossal magnetoresistive (CMR) effects are of high current interest in solid state physics. Europium hexaboride (EuB₆) and the well-known manganites, for which the onset of ferromagnetism is accompanied by a dramatic reduction of the electrical resistivity, are primary examples that have intensively been studied. Although a lot of experimental data on the magnetic and electronic properties are available, a complete understanding of the underlying physical mechanisms is still lacking.

We concentrate on the series of cubic $Eu_{1-x}Ca_xB_6$, which displays interesting correlations between magnetic, transport, and optical properties. The high temperature electronic transport of EuB₆, a ferromagnet (FM) below $T_C \sim 12$ K, relies on a small effective electron density [1]. The magnetic properties are dominated by the half-filled 4f shell of divalent Eu, which accounts for the measured magnetic moment of $7\mu_B$ per formula unit [2]. The strong coupling of transport properties to the magnetization was revealed by measurements of the magneto-optical properties, which unveiled a substantial blueshift of the plasma edge in the optical reflectivity with decreasing temperature and increasing magnetic field [3]. The remarkable results correlate with the precipitous drop in the dc resistivity just below T_C , and the large negative magnetoresistance observed near T_C [4–6].

A variety of models was invoked to address those properties. Full potential spin-polarized self-consistent electronic band structure calculations suggest a semimetallic or degenerate semiconductor state of EuB_6 [7]. This view does not agree with results from angular-resolved photoemission spectroscopy (ARPES) that reveal a considerable gap between valence and conduction bands [8]. Our magneto-optical Kerr effect data set [9] implies that the inclusion of a Drude contribution is essential in interpreting the experimental optical data for EuB₆. Lin and Millis were able to reproduce T_C , the enhancement of the plasma frequency in the ordered phase [1,3], and the specific heat [1] with a calculation based on the dynamical mean-field approximation [10]. The major finding of this latter approach establishes that the splitting of the Eu derived conduction band, due to the coupling to core electrons at each Eu site, causes the phase transition. Another model, based on the formation of magnetic polarons [11], suggests that a Hund-type coupling of holes to local spins and magnetic ordering via the Ruderman-Kittel-Kasuya-Yoshida (RKKY) interaction are responsible for the ferromagnetic phase transition and CMR. Wigger et al. [6], on the other hand, reproduced the negative magnetoresistance in the paramagnetic regime by a model in which the spin disorder scattering of the itinerant electrons is reduced by the applied magnetic field, i.e., by the increasing magnetization, and also accounts for the results of de Haas-van Alphen, optical reflectivity, and ARPES measurements.

The intimate relation between magnetization and electronic conductivity also emerged from experiments on the $Eu_{1-x}Ca_xB_6$ series [12–15]. The Ca substitution leads to significant changes of the magnetic and electronic properties. The FM transition temperature decreases with increasing Ca content and stoichiometric CaB₆ exhibits no magnetic order [14,15]. Evidence was also established for a spin-filter effect in the charge transport and dynamics for x = 0.4 [12,13]. The electrical transport properties seem to be governed by percolation-type phenomena across the $Eu_{1-x}Ca_xB_6$ series [14,15]. At about x = 0.7, which coincides with the site-percolation limit in a simple cubic lattice, the long-range order disappears and for 0.7 < x < 0.9 a spin-glass type ground state is adopted [15].

A recently published approach to explain the behavior of EuB₆, as well as offering specific predictions for the electronic properties of the Eu_{1-x}Ca_xB₆ series, is based on a double-exchange scenario [16]. This scenario may be regarded as an effective theory for the Kondo lattice problem in the limit of a very small number of carriers. The reduced itinerant carrier concentration places the Fermi level near a magnetization dependent mobility edge, which emerges in the spectral density because of the disordered spin background and/or Ca doping. A FM metal to insulator crossover is expected as a function of the position of the Fermi level with respect to the mobility edge, which can be tuned by the Ca content [16]. The model also addresses [17] the region of stability of magnetic polarons in the paramagnetic phase near T_C [18].

The goal of this Letter is to present and analyze our magneto-optical data of the $Eu_{1-x}Ca_xB_6$ series. Replacing Eu by Ca has direct consequences on the electrodynamic response. It influences the distribution of the spectral weight between the metallic (Drude) component and excitations at nonzero energy in the absorption spectrum for different Ca contents at different temperatures and magnetic fields. Our analysis provides support for the phase diagram that emerges from the double-exchange model predictions [16]. The boundary between the metallic and the insulating ferromagnetic state at low temperatures is found to be close to x = 0.5.

 $Eu_{1-x}Ca_xB_6$ single crystals of high structural quality were prepared by solution growth from Al flux, using the necessary high-purity elements as starting materials [14,15]. The optical reflectivity $R(\omega)$ was measured in a broad spectral range from the far infrared (FIR) to the ultraviolet, and as a function of both temperature (1.6-300 K) and magnetic field (0-7 T) [1,3]. Because of the broad coverage of spectral range, $R(\omega)$ spectra allowed for a reliable Kramers-Kronig (KK) transformation providing the complete absorption spectrum represented by the real part $\sigma_1(\omega)$ of the optical conductivity [19,20]. At low frequencies, i.e., below our low frequency experimental limit of about 30 cm⁻¹, $R(\omega)$ was extended using the Hagen-Rubens extrapolation, and inserting the $\sigma_{\rm dc}$ values obtained with dc transport measurements [6,12,14]. Above the highest measurable frequency, $R(\omega)$ was extended into the electronic continuum with the standard extrapolations $R(\omega) \sim \omega^{-s}$ (2 < s < 4) [19,20].

The temperature and magnetic field dependence of $R(\omega)$ for EuB₆ was already presented and discussed in Refs. [1,3]. Here, we complement those results with data for x = 0.3, 0.4 [Ref. [13]], 0.55, and 0.8. For the presentation in Fig. 1, we chose spectra that were recorded at 10 K (i.e., $T > T_C$) for all compounds, and in magnetic fields of 0 T and 7 T. $R(\omega)$ is progressively enhanced with increasing magnetic field for x = 0.3 and 0.4, while for x = 0.55 and 0.8, basically no field dependence was registered. Although $R(\omega)$ exhibits metallic character for all these Ca contents, the onset of the plasma edge in $R(\omega)$ is quite broad in all compounds. This is distinctly different



FIG. 1 (color online). Reflectivity spectra $R(\omega)$ of $\operatorname{Eu}_{1-x}\operatorname{Ca}_x\operatorname{B}_6$ in the infrared spectral range at T = 10 K ($T > T_C$), for magnetic fields of 0 T and 7 T. Note that for x = 0.8 the 0 and 7 T spectra are identical. The thin lines represent the Lorentz-Drude fit (see text).

from the previously observed rapid increase of $R(\omega)$ with decreasing ω for EuB₆ [Refs. [1,3]].

In order to analyze the spectra, we apply the phenomenological Lorentz-Drude approach [19,20]. The fit procedure consists in considering a Drude term for the zerofrequency mode excitation due to the itinerant charge carriers, and an appropriate number of Lorentz harmonic oscillators (HO) for the absorptions at nonzero energies. The latter components represent phonon modes, electronic interband transitions, and localized-state excitations. Although of phenomenological character, this approach allows for the evaluation of the spectral-weight distribution among the various components of the absorption spectrum. For all Ca substitutions, we consistently reproduce each spectrum for any combination of temperature and magnetic field with the same fit procedure and the same number of fit components. As an example, we show $\sigma_1(\omega)$ at 10 K and 0 T for x = 0.8 in Fig. 2. The individual components to the fit may readily be identified. Both Figs. 1 and 2 demonstrate that in this way the raw experimental data for $R(\omega)$ and the resulting $\sigma_1(\omega)$ can be reproduced very well. We can thus disentangle the Drude spectral weight corresponding to the squared plasma frequency from the spectral weight associated with the excitations at nonzero energies. The total spectral weight encountered in the excitations at higher energies is proportional to the sum of the squared mode strengths [19,20].

Because of the specific model prediction [16] of a FM metal to insulator crossover, our primary aim is to evaluate the change of spectral weight (SW) of the metallic component of $\sigma_1(\omega)$ (red area in Fig. 2) at $\tilde{T} > T_C$ between 0 and 7 T. This difference is calculated on the basis of our Lorentz-Drude results and is defined as [21]: $\Delta SW^{Drude} = SW^{Drude}(\tilde{T}, 7 \text{ T})-SW^{Drude}(\tilde{T}, 0 \text{ T})$. The magnetic field of 7 T is high enough to drive the system into magnetic saturation for all values of *x*, such that $\sigma_1(\omega)$ at 7 T reflects



FIG. 2 (color online). (a) Real part, $\sigma_1(\omega)$, of the optical conductivity of Eu_{0.2}Ca_{0.8}B₆ at 10 K and 0 T. The total Lorentz-Drude fit as well as the fit components are displayed, and one can appreciate the very satisfying fit quality. The colored red area identifies the Drude spectral weight, while the sum of the red and yellow areas represents the total spectral weight encountered in $\sigma_1(\omega)$.

the maximum metallicity reached by increasing the magnetic field. \tilde{T} (20 K for x = 0, 10 K for x > 0) was chosen to be close enough to, but above, T_C for each compound [22]. In order to allow for a comparison for different Ca contents, we renormalize the change of the Drude spectral weight ΔSW^{Drude} by the total spectral weight (red plus yellow areas in Fig. 2) encountered in $\sigma_1(\omega)$ up to about $\omega_c = 1$ eV, at either 0 or 7 T, i.e., SW^{TOT} (\tilde{T} , 0, or 7 T) [Ref. [21]]. The cutoff frequency ω_c coincides approximately with the onset of the high-frequency electronic interband transitions, where both the temperature and the magnetic field dependence in $\sigma_1(\omega)$ vanish. Figure 3 displays the variation of the normalized Drude spectral weight (i.e., $\Delta SW^{Drude}/SW^{TOT}$) as a function of x, in comparison with the variation of T_C for the Eu_{1-x}Ca_xB₆ series [14,15]. The ranges on the vertical axes were chosen such that $T_C(x=0)$ coincides with the renormalized changes of the Drude spectral weight, inserting SW^{TOT} for either 0 or 7 T. For both values of SW^{TOT}, we obtain the same type of variation with x. ΔSW^{Drude} decreases sharply between x = 0 and 0.3, reaching zero at approximately 50% Ca content.

The *x* dependence of ΔSW^{Drude} reveals the reduction of the maximum of itinerant charge carriers with increasing *x* in Eu_{1-x}Ca_xB₆, previously indicated by results of resistivity [5,12], Hall effect [14], and optical response [1,3,13] measurements. The field-induced enhancement of ΔSW^{Drude} as a function of *x* and its correlation with the evolution of $T_C(x)$ is fully consistent with the microscopic mechanisms that are considered in the low-density doubleexchange model applied to a percolating lattice [16]. The crucial detail of this description of the interaction between the conduction electrons and the *classical* (S = 7/2) lo-



FIG. 3 (color online). Dependence of the Curie temperature, T_C , on Ca doping [14,15], compared to $\Delta SW^{Drude}/SW^{TOT}$, the change of the spectral weight in the Drude component, normalized by the total spectral weight (at 0 or 7 T) encountered in $\sigma_1(\omega)$ up to ω_c . The dashed lines for $T_C(x)$ above $x \sim 0.7$ schematically define the interval of T and Ca content, where a cluster spin-glass phase was established [14,15].

calized magnetic moments is not the usual strong Hund coupling condition $(J_H \gg t)$, but rather the extremely reduced density of itinerant carriers that permits the highenergy electronic states to be projected out. Part of the experimental evidence [4-6,12-15] for the strong interplay between transport and magnetization in the hexaborides is then the consequence of Anderson localization effects. In this sense, the magneto-transport in $Eu_{1-r}Ca_rB_6$ is dominated by a magnetization dependent source of disorder in the random magnetic background in which the electrons itinerate. This is quantified by the position of the mobility edge, $E_C(\mathcal{M})$, relative to the Fermi energy, $E_F(\mathcal{M})$ [23]. The reduced electronic densities, characteristic of the $Eu_{1-x}Ca_xB_6$ series, place these two energies very close to each other. Direct experimental consequences of this are the marked enhancement of the itinerant carrier density upon driving the system into a ferromagnetic regime by either reducing T to below T_C or by applying an external magnetic field. With $E_C(\mathcal{M})$ shifting towards the band edge upon increasing \mathcal{M} , more extended (metallic) states are occupied in the system.

The microscopic model of Ref. [16] considers an optical sum rule that includes only the Drude contribution due to the extended states: $\omega_p^2(\mathcal{M}) \propto \int_{E_C(\mathcal{M})}^{\infty} N(E, \mathcal{M}) f(E) E dE$, with $N(E, \mathcal{M})$ as the magnetization dependent electronic density of states, and f(E) the Fermi-Dirac function. Upon replacing Eu by Ca, the Eu sublattice and consequently the magnetic subsystem are diluted. Given the isovalency of Ca and Eu, from the electronic point of view, the main effect of this is a strong enhancement of the electronic disorder, and thus $E_C(\mathcal{M}, x)$ will also depend on the Ca content. For this reason, with increasing x less extended states are occupied, and once $E_C(\mathcal{M}, x_{\text{MI}}) = E_F(\mathcal{M}, x_{\text{MI}})$, a metal-insulator transition (at T = 0) or a metal-insulator crossover (T > 0) emerges. Figure 3 shows that the spectral weight of the Drude term in $\sigma_1(\omega)$ is enhanced when the system is driven from a paramagnetic metallic state into full polarization up to $x \leq 0.4$. The progressive reduction of this enhancement with x reflects the drift of E_C and consequently less metallic conduction. For higher Ca concentrations, the Drude spectral weight is insensitive to the spin polarization, a signal that the mobility edge $E_C(\mathcal{M} =$ 1) went past the Fermi energy and thus the polarization no longer releases any of the localized states. The tail-like behavior of $\Delta SW^{Drude}/SW^{TOT}$ in Fig. 3 for $x \ge 0.4$ can be understood as originating from the nonzero temperature excitations of carriers across the mobility gap, $E_C - E_F$ (all measurements were made at $T > T_C$). Both this interpretation and the features of the optical response are perfectly consistent with the variation of the number of carriers in the paramagnetic phase, $n_0(x)$, presented in Ref. [14]. It is also significant that the results do not depend on the magnetic field used for the normalization of ΔSW^{Drude} , as evidenced by the two curves in Fig. 3, where ΔSW^{Drude} is normalized by SW^{TOT} at either 0 or 7 T.

With respect to magnetic order, Fig. 3 shows that the Curie temperature decreases with x, as expected if site percolation is important. Unlike the Drude spectral weight, long-range magnetic order survives until the Ca concentration coincides with the threshold of the simple cubic magnetic lattice site percolation. This is in agreement with the double-exchange scenario in which the effective magnetic coupling is the result of the electron itinerancy among sites with localized moments. This magnetic coupling, local in nature, is present even in the localized regime as long as typical localization lengths allow wave functions to spread over nearest neighbors. From the optical point of view, Fig. 3 perfectly reflects the phase diagram predicted in Ref. [16]. Up to $x_{\rm MI} \simeq 0.4$ one has a metallic ferromagnet; for higher Ca concentration the system remains a ferromagnetic Anderson insulator until it reaches the percolation threshold. Near and above the percolation threshold the number of disconnected Eu-rich magnetic clusters becomes significant. Even though the tendency should be towards ferromagnetism, it is not surprising that the regime above $x \ge 0.7$ (beyond percolation, and at very low temperatures) seems to be characterized by glassy magnetism [15], on account of the possible presence of superparamagnetic clusters and competing dipolar interactions at such extreme dilutions of the magnetic moments [24].

Finally the Ca driven metal-insulator transition found in transport measurements, now corroborated with our magneto-optical measurements, is a detail of paramount relevance in the phase diagram of $Eu_{1-x}Ca_xB_6$. Disorder is one of the essential building blocks of the double-exchange interpretation and the current confirmation of the phase diagram foreseen in Ref. [16] certainly underlines the significance of localization effects in the description of these compounds.

In conclusion, our magneto-optical investigations on the $Eu_{1-x}Ca_xB_6$ series reveal a phase diagram in support of a scenario based on the close proximity of the Fermi level

and a magnetization dependent mobility edge. A ferromagnetic metal-insulator crossover occurs upon increasing the Ca content in $\text{Eu}_{1-x}\text{Ca}_x\text{B}_6$ to above x_{MI} . The magneto-optical data suggest a critical Ca content $x_{\text{MI}} \sim 0.5$.

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- [20] M. Dressel and G. Grüner, in *Electrodynamics of Solids* (Cambridge University Press, Cambridge, England, 2002).
- [21] SW^{Drude} ~ ω_p^2 and SW^{TOT} ~ $\omega_p^2 + \sum_j \omega_{pj}^2$, where ω_p is the plasma frequency of the Drude term and ω_{pj} are the mode strength of the Lorentz harmonic oscillators covering the spectral range up to ω_c . An equivalent procedure for calculating SW^{Drude} and SW^{TOT} consists in integrating $\sigma_1(\omega)$ up to about 0.15 eV for the effective Drude spectral weight, and up to about ω_c for the total weight.
- [22] Any choice of temperature $T_C < T < \tilde{T}$ for each Ca content leads to equivalent results.
- [23] The band curvature also depends on the strength of disorder.
- [24] In Ref. [16], the scattering is due to spin disorder and also from the disorder induced by the random position of atoms. The treatment of scattering processes is more involved in the case of the spin-glass phase close to percolation. This requires a detailed understanding of the spin-glass nature, an issue which is beyond the scope of the theory [16].