Magnetic circular dichroism in Ga$_x$Mn$_{1-x}$As: Theoretical evidence for and against an impurity band

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(Received 17 August 2009; published 13 October 2009)

Magneto-optical properties of the ferromagnetic semiconductor GaMnAs are studied in a material-specific multiband tight-binding approach. Two realistic models are compared: one has no impurity band while the other shows an impurity band for low Mn concentrations. The calculated magnetic circular dichroism (MCD) is positive for both models proving that, unlike previously asserted, the observed positive MCD signal is inconclusive as to the presence or absence of an impurity band in GaMnAs. The positive MCD is due to the antiferromagnetic $p$-$d$ coupling and the transitions into the conduction band.

DOI: 10.1103/PhysRevB.80.161201 PACS number(s): 75.50.Pp, 71.55.Eq, 75.50.Dd, 78.20.Ls

The ferromagnetic semiconductor GaMnAs (Refs. 1–3) is a fascinating material not only because of its importance for technological applications$^4$ but also because its understanding poses great challenge. Indeed, understanding the electronic structure of GaMnAs requires resolving complexities due to disorder, correlations, and magnetism. Over the past years, an intense debate has sparked about the nature of the electronic structure of GaMnAs$^{8,11–15}$ giving important insights into such a view applying to the antiferromagnetic $p$-$d$ coupling, in contrast to what is expected for the presence or absence of an impurity band.

The apparent picture coming from experiments$^8,13,15,24$ is that the signal results from the spin-resolved electronic level ordering in GaMnAs which is present despite the strong disorder. The Fermi-level shift due to doping by Mn acceptors is strong enough to make the MCD signal positive, regardless whether or not the Fermi level lies in the valence band (Moss-Burstein shift) or in the impurity band, while holding to the antiferromagnetic $p$-$d$ coupling in both cases. The dominant transitions are those involving the conduction band (CB), not the impurity one. We thus find most previous conclusions drawn from the positive MCD signal unfounded.

Our argument is based on large-scale tight-binding calculations of MCD in disordered GaMnAs using two models. One model reproduces the valence-band picture, the other shows an impurity band. Both have antiferromagnetic $p$-$d$ coupling, and both show a positive MCD around the fundamental absorption edge, in agreement with experiment.

Our simulations use a material-specific microscopic tight-binding approach. The models are based on $1.5\text{ eV}$ valence and conduction bands of GaAs, which are approximated throughout the entire Brillouin zone to fit the experimentally determined band structure.$^{25,26}$ We use two different parameter sets for the inclusion of Mn impurities into the models. One of the models$^{27}$ is characterized by an almost identical change in the band structure, and the Fermi energy at zero temperature and various Mn concentrations. Within this framework we can treat the disorder effects nonperturbatively. While this approxima-
tion seems justified for large concentrations of Mn impurities, an explicit inclusion of the carrier-carrier interactions (computationally infeasible) may give qualitative corrections of our findings.

As already stated, we investigate two different sets of tight-binding parameterizations to study in detail how an impurity band affects the MCD results. These two models were thoroughly analyzed in Ref. 29 concerning the density of states, the size of the band gap, the position of the Fermi energy, localization properties of holes, optical effective masses, and mean-free paths. The first model, which we call model A, was suggested by Mašek.27 This model, derived from a first-principles approach, describes GaMnAs qualitatively very similar to what is expected from a p-doped GaAs.29 Its main characteristic is the inclusion of additional energy levels of the host material rather strongly.29 Applying this model to disordered systems with many Mn impurities leads to the formation of an isolated impurity band for Mn concentrations \(x \approx 1\%\). At higher concentrations the impurity band starts to merge with the host valence band. Hence, the concentrations leads to the formation of an isolated impurity band for Mn.

The second model, which we call model B, was suggested by Tang and Flatté.28 Within this model the Mn impurities are described by a modified on-site potential and a spin-dependent potential at the four nearest As neighbors. The two relevant parameters are adjusted such that the experimental binding energy of 0.11 eV of the Mn impurity is recovered. Qualitatively, this model not only shifts the Fermi energy due to the additional holes but it also affects the positions of the energy levels of the host material rather strongly.29 Applying this model to disordered systems with many Mn impurities leads to the formation of an isolated impurity band for Mn concentrations \(x \approx 1\%\). At higher concentrations the impurity band starts to merge with the host valence band. Hence, the comparison of these two different parameterizations, models A and B, allows us to draw conclusions on how the existence of an impurity band influences the MCD signals that are seen in the experiments.

We analyze the MCD signal by evaluating the diagonal and off-diagonal elements of the optical-absorption matrix. The matrix elements \(\sigma_{\alpha\beta}\) can be written in terms of the eigenstates \(|n\rangle\) and eigenenergies \(E_n\) as

\[
\sigma_{\alpha\beta}(\Delta, E_F) = \frac{\hbar^2}{2m^2 \Omega} \sum_{n n'} f_n f_{n'} \Delta \Delta_{n'n} \left( \Delta + i \delta - \Delta_{n'n} \right),
\]

with the system volume \(\Omega\), the Fermi function \(f_n = f_{E_n}(E_F)\), energy difference \(\Delta = E_n - E_{n'}\), and momentum matrix elements \(P_{\alpha\beta} = (\langle n'|\hat{p}\cdot\hat{r}|n\rangle)\). The absorption \(\sigma_{\alpha\beta}\) for right (left) polarized light can be obtained by replacing \(p - p^\pm = p^\pm \mp i\delta\) in Eq. (1). The MCD signal, for a constant index of refraction (or poor conductors), is

\[
\text{MCD} = (\sigma_+ - \sigma_-)/(\sigma_+ + \sigma_-).
\]

In order to numerically evaluate conductivity (1) we calculate the eigenenergies and the matrix elements of the momentum operator using a multiband tight-binding approach.29 This approach is applied to periodically repeated finite-size supercells containing up to 2000 atoms. The resulting conductivity is furthermore averaged over several disorder configurations. Therefore, our approach goes beyond the single-impurity calculation presented on model B in

![FIG. 1. (Color online) Left panel: model A; right panel, model B. Upper panel: schematic level ordering deduced from absorption data. Lower panel: absorption of right (solid line) and left circularly polarized (dashed line) light. The system size is 2000 atoms with one Ga atom replaced by Mn. The absorption rates are evaluated at the \(E_p\) point in the superlattice Brillouin zone. Levels are labeled according to their carrier character in pure GaAs: conduction-band (CB), heavy-hole (HH) and light-hole bands, and the spin-orbit split-off (SO) band. Magnetic quantum numbers are indicated. Since substitutional Mn is an acceptor, the highest valence-band level ("HH-3/2") is not occupied so that the transitions "a" are suppressed.](161201-2)

Ref. 23 as we explicitly include disorder averaging effects. In our calculations, for numerical reasons, we use a smearing temperature corresponding to 1 meV in the Fermi functions and a peak width \(\delta = 5\) meV. The integration over the superlattice Brillouin zone is performed by summation over up to 2048 different \(\vec{k}\) vectors.29 As the systems under consideration are disordered one cannot restrict this summation to the irreducible part of the Brillouin zone. Furthermore, the finiteness of the systems limits the Mn concentrations to \(x \approx 1\%\) in our present simulations.

For a better understanding of the numerical MCD results we first analyze the effect of a single Mn impurity in a supercell of 2000 atoms with periodic boundary conditions. The absorption was evaluated at the \(E_p\) point, i.e., \(\vec{k} = 0\) in the superlattice Brillouin zone. Due to the magnetic moment of the Mn impurity the spin degeneracies of the conduction \(s\) and the valence \(p\)-bands are lifted. The order in which the levels appear can be concluded from the absorption peaks of \(\sigma_{\pm}\) at \(E_p\approx 1.5\) eV. We show the numerical results in Fig. 1. The Fermi energy was chosen to be (artificially) above the highest impurity level, to show all relevant optical transitions. The absorption peaks then correspond to the six possible transitions between the six heavy-hole, light-hole, and spin-orbit split-off bands to the two conduction bands.17 Other transitions are suppressed because of the selection rules for the momentum matrix elements. Among the allowed transitions, "a" and "c" are the strongest ones due to their larger momentum matrix elements. From the position of each \(\sigma_{\pm}\) peak one can uniquely identify the involved bands. The extracted ordering of the levels is shown schematically in the upper panel of Fig. 1.
As a Mn impurity also acts as an acceptor, the highest impurity level, that is the levels labeled “HH-3/2” in Fig. 1, is in fact unoccupied. This means that the actual Fermi energy lies just below this highest impurity level. Therefore, all transitions starting from this level are suppressed. For the absorption this means that the peak labeled “a” is not observable at zero temperature.

Figure 1 points to common features and differences between the two models. The order of the first four peaks, “a”–“d,” is the same for both models implying the same ordering of the valence-band states. The spin of the Mn core 3d electrons in our simulation points into the -z direction, that is, antiparallel to the propagation direction of the light. This leads to an acceptor state with spin-down character implying antiferromagnetic p-d exchange coupling for the holes, for both models.

Besides this common feature there are two major differences between model A and model B. First, the magnitude of the level splitting due to the magnetic impurity is very different. For example, model A results in a splitting of the four heavy and light-hole bands of ~4 meV while in model B this splitting is ~0.13 eV. This difference in the splittings is associated with the fact that model B explicitly reproduces the bound-state energies of a single Mn impurity at ~0.11 eV in the host gap by a strong deformation of the host valence band. The second major difference between the two models is the different order of the two conduction bands. In model A the spin-up band has a higher energy in the conduction band while in model B it is the spin-down band.

How are the above effects of a single Mn impurity reflected in the absorption of the left and right circularly polarized light in strongly disordered samples with many impurities? The simulation results for a Mn concentration x=2% are shown in Fig. 2. For this Mn concentration model B still shows an impurity band, which is attached to the host valence band, while model A does not exhibit an impurity band at all as can be seen in the insets of Fig. 2. The Fermi energy in model A lies within the host valence band whereas for model B it lies within the impurity band. For both models we find that the σ− absorption sets in at lower energies Δ=E_gap+Δ_SO ~1.8 eV for model A the Δ peak comes first while for model B it is the σ− peak. This specific in the ordering of the conduction band does not alter the calculated MCD signal qualitatively.

![Graph](image-url)
character are not occupied. Comparing with recent experiments we find very good qualitative agreement for the MCD and IMCD signals.\textsuperscript{12,13} A precise quantitative agreement cannot be expected from a tight-binding model.\textsuperscript{32}

To test the hypothesis put forward in Ref. 15, that the positive MCD is due to the transitions to the impurity band, we have calculated the partial conductivities by excluding the conduction-band states in Eq. (1) for model B; see Fig. 2. These conductivities (and their differences) are much smaller than the complete ones, giving no support to the hypothesis. Instead, the positive MCD around the absorption edge is due to the transitions to the conduction band.

As in Ref. 13, we show the IMCD signal for various Mn concentrations in Fig. 3. With increasing Mn concentration the magnitude of the IMCD signal is decreasing as the total absorption becomes stronger. However, the shape of the signal is rather robust for all investigated Mn concentrations. In particular, we do not observe a change in the sign of the MCD signal within the considered range of concentrations $x$ from $2\%$ to $10\%$.

What would the MCD signal be for a fully compensated GaMnAs? It is particularly interesting to see what the MCD were if the Fermi level were at the top of the impurity band in model B, as this reflects the ordering of the impurity band levels and the deformation of the ordering due to disorder. Our results are shown in Fig. 3 for $x=2\%$ ($E_F=E_0$). The MCD is negative as the $\sigma_\alpha$ transitions are now allowed, dominating the signal in accord with the single-impurity picture in Fig. 1. In summary, we find that the positive MCD signal in GaMnAs (a) is inconclusive as to the presence or absence of an impurity band, (b) implies antiferromagnetic $p$-$d$ coupling also for the valence-band scenario, and (c) originates from the transitions to the conduction band.

We thank D. Weiss and C. Back for helpful discussions. This work was supported by the DFG (Grant No. SFB 689).

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