Combined approach of density functional theory and quantum Monte Carlo method to electron correlation in dilute magnetic semiconductors

Jun-ichiro Ohe$^1$, Yoshihiro Tomoda$^1$, Nejat Bulut$^{1,2}$, Ryotaro Arita$^3$, Kazuma Nakamura$^3$, and Sadamichi Maekawa$^{1,2}$

$^1$Institute for Materials Research, Tohoku University, Sendai 980-8577, Japan
$^2$CREST, Japan Science and Technology Agency (JST), Kawaguchi, Saitama 332-0012, Japan
$^3$Department of Applied Physics, University of Tokyo, Bunkyo-ku, Tokyo 113-8656, Japan

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We present a realistic study for electronic and magnetic properties in dilute magnetic semiconductor (Ga,Mn)As. A multi-orbital Haldane-Anderson model parameterized by density-functional calculations is presented and solved with the Hirsch-Fye quantum Monte Carlo algorithm. Results well reproduce experimental results in the dilute limit. When the chemical potential is located between the top of the valence band and an impurity bound state, a long-range ferromagnetic correlations between the impurities, mediated by antiferromagnetic impurity-host couplings, are drastically developed. We observe an anisotropic character in local density of states at the impurity-bound-state energy, which is consistent with the STM measurements. The presented combined approach thus offers a firm starting point for realistic calculations of the various family of dilute magnetic semiconductors.

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The discovery of ferromagnetism in dilute magnetic semiconductor (DMS) materials, represented by a (Ga,Mn)As system, has created much activity in the field of spintronics [1, 2, 3, 4]. In the low-doping regime ($\ll 1\%$), (Ga,Mn)As is insulating with a clear experimental evidence for the presence of an Mn-induced impurity band located at 110 meV above the valence band [3]. The position of the impurity level and the impurity-induced carriers are key quantities in generating the ferromagnetism. Various model studies [5, 6, 7, 8, 9, 10] have clarified that this ferromagnetic correlations between the impurities are based on the polarization mechanism of the impurity-induced carriers.

The challenge for predicting new functional properties of the DMS materials and their realistic designs clearly requires many tasks such as nonempirical electronic-band structure of the host compound, reliable estimations for impurity-host hybridizations, and rigorous treatments for local electron correlations at the impurity site. A combined approach of density functional theory (DFT) [11] and quantum Monte Carlo (QMC) technique [12] is clearly useful for this purpose. The hybridization or mixing parameters between the host and impurity orbitals is a crucial parameter characterizing material differences and therefore this parameter should be estimated on the basis of $ab$ $initio$ density-functional calculations. On the other hand, density-functional calculations often fail to describe local electron correlations, so this problem must be treated with more accurate solvers such as the QMC technique. Critical comparisons with the experimental data are needed for a check whether this approach is really reliable in describing the details of the "low-energy" electronic and magnetic properties and/or whether it has an extended applicability for many DMS materials.

In this letter, we present a comprehensive study for the generation mechanism of the ferromagnetism of a dilute (Ga,Mn)As system. A multi-orbital Haldane-Anderson model [13] is employed to describe electronic/magnetic properties of this system. First principles band-structure calculations are performed to evaluate the mixing matrix elements in this model and the resulting model is solved with the help of the QMC technique to examine the inter-impurity and impurity-host magnetic correlations. The hybridization between host and impurity electrons generates the impurity bound state [14]. When the chemical potential is located between the impurity bound state and the valence-band top, a notable long-range ferromagnetic correlation between the impurities and its dramatic enhancement as the temperature decreases are observed. We will show that this ferromagnetic correlation is mediated by antiferromagnetic impurity-host couplings. In addition, our calculated host local density of states is highly anisotropic around the impurity sites, which is

![FIG. 1: (Color online) Square of the impurity magnetic moment $\langle M^2 \rangle$ vs the chemical potential $\mu$ at various temperatures in the single-impurity case. These results are for the single-impurity Haldane-Anderson model. The vertical line ($\mu = 0$) denotes the top of the valence band.](image-url)
follows [13]:

These quantitative agreements with the experiments indicate that the presented combination idea is a useful approach for aiming at realistic calculations and searching new functional high-$T_c$ DMS materials.

A multi-orbital Haldane-Anderson model is given as follows [13]:

$$
\mathcal{H} = \sum_{k,\alpha,\sigma} (\epsilon_{k\alpha} - \mu) c_{k\alpha}^\dagger c_{k\alpha} + \sum_{i,\xi,\sigma} (E_{i\xi} - \mu) d_{i\xi}^\dagger d_{i\xi}^\sigma + \sum_{k,\alpha,\xi,\sigma} (V_{k\alpha,i\xi} c_{k\alpha}^\dagger d_{i\xi}^\sigma + \text{h.c.}) + U \sum_{i,\xi} d_{i\xi}^\dagger d_{i\xi}^\dagger d_{i\xi}^\sigma d_{i\xi}^\sigma.
$$

Here, $c_{k\alpha}^\dagger$ ($c_{k\alpha}$) creates (annihilates) a host electron with wavevector $k$ and spin $\sigma$ in the valence or conduction bands denoted by $\alpha$, and $d_{i\xi}^\dagger$ ($d_{i\xi}^\sigma$) is a creation (annihilation) operator for a localized electron at impurity $d$ orbital $\xi$ located at transition-metal site $i$. The first term in Eq. (1) represents a kinetic energy $\epsilon_{k\alpha}$ of a host $k\alpha$ electron, and the second term describes a bare onsite energy $E_{i\xi}$ of the impurity and $\mu$ is a chemical potential. The third term specifies impurity-host hybridization $V_{k\alpha,i\xi}$ and the last term represents the onsite repulsion $U$ at the impurity site. In the present study, we keep only diagonal terms in the onsite interaction by neglecting the off diagonal terms in the interaction (i.e., Hund’s rule coupling).

When a Mn atom is substituted in place of a Ga site, the five $3d$ orbitals of the Mn ion are split into the three-fold degenerate $t_{2g}$ orbitals and the two-fold degenerate $e_g$ orbitals by the tetrahedral crystal field. In the present Mn$^{2+}$ case with a $3d^5$ configuration, the $e_g$ orbitals are fully occupied and therefore are inactive for the magnetic properties. In contrast, the three-fold $t_{2g}$ orbitals being a partially-filled state are active for the low-energy magnetic excitations.

The onsite repulsion was set to $U=4.0$ eV following a photoemission spectrum [10] and this value is very close to theoretical values 4-5 eV for metallic Mn [17]. We note that both $U$ and $E_{i\xi}$ can shift the energy of the impurity bound state. Experimentally, it is widely agreed that an impurity band exists in the dilute limit of (Ga,Mn)As at 110 meV above the valence band [5]. To realize this situation, we found that $E_{i\xi}$ should be set to $-2.0$ eV.

Estimations for the hybridization parameters $V_{k\alpha,i\xi}$ in Eq. (1) need careful treatments. Empirical estimations with the Slater-Koster table fail to describe the dispersion of the conduction bands, thus leading to a serious failure in the hybridization-parameter estimation. We estimate the hybridization parameter $V_{k\alpha,i\xi}$ in Eq. (1) from ab initio density-functional calculations [18]. The matrix element is given by

$$
V_{k\alpha,i\xi} = \langle \psi_{k\alpha} | \mathcal{H}_0 | \psi_{i\xi} \rangle,
$$

where $\mathcal{H}_0$ is the one-body part of Eq. (1), $|\psi_{k\alpha}\rangle = c_{k\alpha}^\dagger |0\rangle$, and $|\psi_{i\xi}\rangle = d_{i\xi}^\dagger |0\rangle$. $|\psi_{k\alpha}\rangle$ can be expressed with a linear combination of Wannier functions as

$$
|\psi_{k\alpha}\rangle = \frac{1}{\sqrt{N}} \sum_{\mu} \sum_{\mathbf{R}} a_{\alpha\mu}(\mathbf{R}) e^{i\mathbf{kR}} |w_{\mathbf{R}\mu}\rangle,
$$

where $|w_{\mathbf{R}\mu}\rangle$ is the Wannier orbital at the lattice $\mathbf{R}$ and the label $\mu$ specifies the type of the orbitals. In the present GaAs case, the $\mu$ specifies the four types of $sp^3$ orbitals centered at a Ga or As site. Expansion coefficients ($a_{\alpha\mu}(\mathbf{k})$) are obtained from solving the equation $\mathcal{H}_0 |\psi_{k\alpha}\rangle = \epsilon_{k\alpha} |\psi_{k\alpha}\rangle$. By substituting Eq. (3) into Eq. (2), we obtain the form of

$$
V_{k\alpha,i\xi} = \frac{1}{\sqrt{N}} \sum_{\mathbf{R}} a_{\alpha\mu}(\mathbf{k}) e^{-i\mathbf{kR}} \langle w_{\mathbf{R}\mu} | \mathcal{H}_0 | \phi_{i\xi} \rangle.
$$

For an actual evaluation of the matrix element $\langle w_{\mathbf{R}\mu} | \mathcal{H}_0 | \phi_{i\xi} \rangle$, we utilize a supercell calculation: We first arrange a supercell containing $3\times3\times3$ fcc primitive cells, where a Ga atom located at the origin in the supercell is replaced with an Mn atom. We then make maximally localized Wannier functions [19] for this system [20]. The width of an energy window was set in order to include all the valence-bonding and conduction-anti-bonding states. The total number of the resulting Wannier functions is 217 in which there are 212 GaAs Wannier orbitals $\{|w_{\mu\nu}\rangle\}$ with $\mu$ and $n$ specifying four

![FIG. 2: (Color online) Inter-impurity magnetic correlation function $(M^x_x M^x_y)$ vs. the impurity separation $|\mathbf{R}|/a$ at various temperatures for $\mu$ fixed at 0.05 eV.](image1)

![FIG. 3: (Color online) $(M^x_x M^x_y)$ vs. $|\mathbf{R}|/a$ at $T = 360$ K for various values of $\mu$.](image2)
types of the sp³ orbitals and sites of Ga or As, respectively, and five Mn-3d Wannier orbitals {w_\xi} sitting on the origin site i. Then, we calculate the matrix element ⟨w_\mu|\hat{H}_KS^3|w_\xi⟩ with \hat{H}_KS^3 being the Kohn-Sham Hamiltonian for the supercell. This matrix element is used instead of ⟨w_\mu|\hat{H}_0|\varphi_\xi⟩ in Eq. 1 for R ≠ i. We note that the error due to this replacement is scaled by \frac{1}{M} with M being the total number of the atoms in the supercell. For R = i, ⟨w_\mu|\hat{H}_0|\varphi_\xi⟩ is expected to be negligibly small, since wave-function symmetry of |w_\mu⟩ and |\varphi_\xi⟩ are different. We found that the hybridization parameters are rather short-ranged; these are nearly zero except for the nearest and next-nearest sites; for example, the parameters between the Mn-t₂g orbitals and the nearest As-sp³ orbitals are 0.16-0.74 eV, while the values between the t₂g orbitals and the next-nearest Ga-sp³ orbitals are 0.06-0.14 eV. Notice that, if the empirical Slater-Koster estimation is employed, the latter next-nearest values are zero.

We perform the quantum Monte Carlo simulation for a one- or two-impurity Haldane-Anderson models, in which we use the Hirsch-Fye QMC algorithm [12] for obtaining the electronic correlations in the one- and two-impurity Haldane-Anderson models. In particular, we study with QMC the magnetic correlations and the local density of states around the impurity site. We define the magnetization operator at the ξ orbital of the i’th impurity site as M_i^ξ = n_i^ξ − n_i̅^ξ, and at site r of the host as m^r(r) = n_r^↑(r) − n_r^↓(r). Here, n_i^ξ and n_r are the spin-dependent number operators at the impurity and host sites, respectively. For the single-impurity case, we present data on the average value of the square of the local moment at the impurity site, ⟨(M^2)⟩ = \frac{1}{3} \sum_\xi ⟨(M_i^ξ)^2⟩, where \xi sums over the t₂g orbitals, while for the two-impurity case we show data on the magnetic correlation function defined by ⟨M_i^ζ M_j^\xi⟩ = \frac{1}{3} \sum_\xi ⟨M_i^ζ M_j M_j^\xi⟩ for impurities located at sites R₁ and R₂. In the single-impurity case, we will also present data on the impurity-host magnetic correlation function, ⟨M_i^ζ M_j^\xi⟩ = \frac{1}{3} \sum_\xi ⟨M_i^ζ M_j^\xi⟩, where r is the host site and the impurity is located at the origin.

Figure 1 shows the square of the impurity magnetic moment ⟨(M^2)⟩ as a function of the chemical potential \mu at various temperatures T for the single-impurity Anderson model. With decreasing T, we observe that a step-like discontinuity develops centered at \mu \approx 0.1 eV, which indicates that the impurity bound state is located at this energy [11]. Within the Hartree-Fock picture [10], the impurity-host hybridization creates both the impurity bound state (IBS) and the host split-off state at ω_{IBS}. For \mu < ω_{IBS}, the hybridization with the valence band reduces the impurity magnetic moment. When \mu is increased to above ω_{IBS}, the polarization of the split-off state cancels the polarization of the valence band, which in turn decreases the screening of the impurity moment. This is the reason for why ⟨(M^2)⟩ increases at ω_{IBS}.

Figure 2 shows the impurity-impurity magnetic correlation function ⟨M_i^ζ M_j^\xi⟩ vs |R|/a at various T for the two-impurity Anderson model. Here, |R| = |R_1 − R_2| is the impurity separation and a is the GaAs lattice constant. In addition, \mu is fixed at 0.05 eV so that it is located between the top of the valence band and the impurity bound state. In Fig. 2, we observe that the ferromagnetic correlations become stronger and extended in real space, as T decreases down to 180 K, which is in the physically relevant temperature regime for (Ga,Mn)As [3]. Figure 3 shows ⟨M_i^ζ M_j^\xi⟩ vs |R|/a at 300 K for \mu = −0.15, 0.05, 0.15 eV. For \mu = −0.15 eV, the chemical potential is located in the valence band. In this case, the system is metallic and we observe Ruderman-Kittel-Kasuya-Yosida (RKKY) type oscillations in the magnetic correlations. The ferromagnetic coupling disappears for \mu = 0.15 eV, where the impurity bound state becomes occupied. These results indicate that the chemical potential should be located between the impurity bound state and the valence band for obtaining extended ferromagnetic coupling between the impurity moments. This case corresponds to the insulating state of (Ga,Mn)As found in the dilute limit [3]. In addition, the strong \mu, R and T dependence of ⟨M_i^ζ M_j^\xi⟩ observed in Figs. 3 and 4 makes it difficult to determine phenomenologically the magnetic coupling in DMS materials, instead requiring approaches which treat the band structure and correlation effect on an equal footing and exactly such as the DFT+QMC method used here.

In order to determine the origin of the ferromagnetic correlation between the impurities, we present in Fig. 4 the impurity-host magnetic correlation function ⟨M_i^ζ m^r(r)⟩ vs |r|/a at various T for the single-impurity Haldane-Anderson model. These results are for \mu = 0.05 eV in which case the inter-impurity ferromagnetic correlation is strong. We observe that the impurity-host magnetic coupling is anti-ferromagnetic and it enhances with decreasing temperature. When the impu-
rity bound state is occupied ($\mu = 0.15$ eV), we find that the impurity-host anti-ferromagnetic magnetic coupling becomes much weaker (not shown here) and the inter-impurity ferromagnetic correlations collapse. Hence, the anti-ferromagnetic impurity-host coupling generates the indirect ferromagnetic coupling between the impurities.

Finally, in the inset of Fig. 4 we show local density of states around the Mn site defined by $N(r) = -\frac{d}{dn} \langle n(r) - n(\infty) \rangle_{\mu=0.05 \text{ eV}}$, where $n(r)$ is the electron number operator at site r. The highly anisotropic distribution of local density of states around Mn is consistent with the STM measurements\(^{15} 12\), which again emphasizes the necessity of including both the band structure and correlation effect\(^{23} 24\).

In summary, we have investigated the magnetic properties of the dilute magnetic semiconductors (Ga,Mn)As in the dilute limit within a combined approach of the density functional theory and the quantum Monte Carlo method. For this purpose, we have mapped the electronic state of the system to the multi-orbital Haldane-Andersen model by using the density functional theory with maximally-localized Wannier functions. For calculating the magnetic properties, we have used the Hirsch-Fye quantum Monte Carlo method, which does not exhibit a fermion sign problem. Within this approach, we have correctly reproduced the experimental value for the energy of the impurity band of (Ga,Mn)As in the dilute limit. The numerical results emphasize the role of the impurity bound state in producing the ferromagnetic correlations. We have found that, when the chemical potential is between the top of the valence band and the impurity bound state, the inter-impurity ferromagnetic correlation originating from the anti-ferromagnetic impurity-host coupling is strong and extended in real space. This case corresponds to the insulating state found in the low-doping regime of (Ga,Mn)As. In addition, we have observed a highly anisotropic distribution of local density of states around Mn which is consistent with the STM measurements. The agreements found with the experimental data on the dilute limit of (Ga,Mn)As suggest that the proposed approach is a reliable method for investigating and designing other DMS materials.

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