Investigation of on-site inter-orbital single electron hoppings in general multi-orbital systems

Qingguo Feng, and P. M. Oppeneer

Department of Physics and Astronomy, Uppsala University, Box 516, S-75120 Uppsala, Sweden

(Dated: February 18, 2022)

PACS numbers: 71.10.-w, 71.10.Fd, 71.27.+a, 71.30.+h

Strongly correlated electron systems have drawn great interest, as they exhibit many exotic phenomena, such as metal-insulator transitions, unusual forms of magnetism, superconductivity, and heavy-fermion behavior. These systems are therefore investigated theoretically in both model and \textit{ab initio} calculations. One of the most essential tasks is to model the system appropriately, i.e., such that a realistic physical picture can be attained. In condensed matter theory the Hubbard model, which was originally proposed in the early sixties, is one of the simplest, yet also the most important and most frequently studied lattice model to investigate strongly correlated electron systems. It sets up a competition between an inter-site quantum mechanical hopping term and an on-site Coulomb interaction term. As a consequence the model can describe various non-trivial phenomena. Due to its simplicity and because the model has captured the essence of strongly correlated electron systems, the Hubbard model has been widely used.

In a realistic situation an atom in a correlated system will usually have several partially filled orbitals and should therefore be described with a multi-orbital (MO) Hubbard model. With the inclusion of orbital degrees of freedom, inter-orbital interactions have to be included for such a system. The inter-site hoppings should now be a sum of the inter-site intra-orbital hoppings and the inter-site inter-orbital hoppings, where the two kinds of hopping are defined according to the orbital indices of the start and the end orbitals for hopping electron are identical or not, respectively. Moreover, besides the remain competition of the inter-site hopping and on-site Coulomb interactions, one new kind of on-site interactions will join into this competition, i.e., the on-site inter-orbital hoppings. Acting as the inter-site hoppings, the on-site inter-orbital hoppings are also associated to the kinetic energy of electrons. All the inter-site hoppings, on-site inter-orbital hoppings and on-site Coulomb interactions (which also divides as on-site intra-orbital Coulomb interactions and on-site inter-orbital Coulomb interactions) constitute a competition for various interactions existing in a realistic multi-orbital system. Therefore, for an arbitrary multi-orbital system, we propose a generalized MO-Hubbard model as follows,

\[
\mathcal{H} = - \sum_{i\neq j} \sum_{lm} t_{ijlm} f_{il\sigma} f_{jml\sigma} + \sum_{il} U_{il} \hat{n}_{il\uparrow} \hat{n}_{il\downarrow}
+ \sum_{il\sigma, l < m} U_{ilm\sigma} \hat{n}_{ilm\sigma} \hat{n}_{ilm\sigma'}
+ \sum_{ilm\sigma, l < m} (J_{ilmf_{il\sigma} f_{jml\sigma}} + J_{ilmf_{ilm\sigma} f_{ilm\sigma'}}),
\]

where \(i, j\) are site indices, \(l, m\) are orbital indices, and \(\sigma, \sigma'\) are spin indices. The \(f_{ilm\sigma} f_{ilm\sigma'}\) are the creation and annihilation operator, respectively, for spin \(\sigma\) in \(l\)-th orbital on site \(i\). The first term is the inter-site hopping term, where \(t_{ijlm}\) is the hopping amplitude for spin \(\sigma\) hopping from \(m\)-th orbital on site \(j\) to \(l\)-th orbital on site \(i\). The second term is the intra-orbital Coulomb interaction and the third term is the inter-orbital Coulomb interaction, where \(U_{ilm\sigma} = U_{ilm\uparrow\uparrow} + J\) and \(J\) is the Hund’s coupling constant. The last two terms in the Hamiltonian are the on-site inter-orbital single electron direct hopping terms, where the \(J_{ilmf_{ilm\sigma} f_{ilm\sigma'}}\) and \(J_{ilmf_{ilm\sigma} f_{ilm\sigma'}}\) are the hopping amplitudes, as \(t_{ijlm}\), in the first term. Note that this part is written in a form of one term and its conjugate, so that one can easily observe that the system is in a statistical equilibrium state. For the half filled case, \(J_{ilm\sigma} = \hat{n}_{ilm\uparrow} = \hat{n}_{ilm\downarrow}\), the model automatically obeys spin rotational invariance. Here the on-site inter-orbital single electron hopping is to be considered as one basic interaction along with the inter-site hopping and the on-site Coulomb interaction. One may argue that with specially chosen basis the on-site inter-orbital hopping will disappear. However, we are here discussing a general multi-orbital system with arbitrary basis. It is convenient to investigate such a system because in some experiments the natural chosen basis in a material may not be orthogonal. Thus studying a system theoretically with a same basis set as in the experimental material can easily gives the results comparative to the physical observables obtained in experiments. Moreover, we find...
that the inclusion of on-site inter-orbital single hoppings has introduced some new and interesting theory.

Recently we have numerically studied the first three terms in Eq. (1) within the dynamical mean field theory (DMFT). In this work we formulate theoretically and evaluate numerically the influence of the on-site inter-orbital single electron hopping. Using the DMFT the proposed generalized Hubbard model can be mapped to a generalized single impurity Anderson model (SIAM) along with a self-consistency condition. The Hamiltonian of this generalized SIAM is

\[ H_{imp} = \sum_{kl\sigma} \varepsilon_{kl\sigma} c_{l\sigma}^\dagger c_{k\sigma} + \sum_{l\sigma} \varepsilon_{f\sigma} f_{l\sigma}^\dagger f_{l\sigma} + \sum_{l} U_l \hat{n}_{l\uparrow} \hat{n}_{l\downarrow} \]

\[ + \sum_{lm\sigma\sigma',l<m} U_{lm\sigma\sigma'} \hat{n}_{l\sigma} \hat{n}_{m\sigma'} + \sum_{lk\sigma} (V_{lk\sigma} c_{l\sigma}^\dagger f_{k\sigma} + V_{lk\sigma} f_{l\sigma}^\dagger c_{k\sigma}) \]

\[ + \sum_{l=1}^{m-1} (I_{lm\sigma} f_{m\sigma}^\dagger f_{l\sigma} + I_{lm\sigma} f_{l\sigma}^\dagger f_{m\sigma}), \]

where the first term is the energy of the conduction electrons (bath), \( c_{l\sigma}^\dagger \) and \( c_{l\sigma} \) are correspondingly the creation and annihilation operators of conduction electrons. The second term is the energy of the localized electrons. The third (fourth) term represents the intra- or inter-orbital Coulomb interactions. The fifth summation is the hybridization term that gives the interaction between the bath and the localized electrons. The sixth summation represents just the on-site inter-orbital single electron direct hoppings on the impurity site, where for convenience we have dropped the site indices in the symbols. These inter-orbital single electron hoppings are on-site interactions which do not change in the mapping. We can hence study these terms in Eq. (1) by equivalently studying the mapped SIAM.

We use the equation of motion (EOM) method to solve the DMFT impurity problem. In this method we compute the equations of motion according to the equation

\[ \omega \ll A; B \gg = \langle [A, B]_+ \rangle + \langle [A, H_{imp}] ; B \rangle, \]  

where we have used the Fourier transform of the double time temperature-dependent retarded Green's function (GF) \( \langle A(t'); B(t) \rangle \), i.e., \( \langle A; B \rangle \) is defined in the space [23], \( [\cdot]_+ \) means the anti-commutator and \([\cdot]\) the commutator. The first term on the right hand side (RHS) labels the interaction associated to \( \langle A; B \rangle \), the second term describes the involvement of the higher-order interactions, where the higher-order GFs will appear. Calculating the EOM of these higher-order GFs, GFs of even higher-order will appear in the newly derived EOMs. Repeating this procedure, more higher and even higher order GFs will appear. Each GF is associated with an order and one physical interaction. The order of the GF approximately labels the weight of its associated interaction. Approximately, the GF of lower order asso-

\[ \langle \omega + \mu - \varepsilon_{f\sigma} \rangle \ll f_{\sigma} ; f_{\sigma}^\dagger \gg = \]

\[ 1 + U_{nm\sigma} \ll \hat{n}_{m\sigma} f_{\sigma}^\dagger f_{m\sigma} \gg + \sum_{l,l \neq m} (U_{lm\sigma} \ll \hat{n}_{l\sigma} f_{\sigma} ; f_{l\sigma}^\dagger \gg + U_{lm\sigma} \ll \hat{n}_{l\sigma} f_{\sigma} ; f_{m\sigma}^\dagger \gg) + \sum_{k} V_{mk\sigma} \ll c_{mk\sigma} ; f_{l\sigma}^\dagger \gg - \sum_{l,l \neq m} I_{lm\sigma} \ll f_{l\sigma} ; f_{m\sigma}^\dagger \gg, \]

where the last term is generated by the on-site inter-orbital single electron hopping terms, and \( \mu \) is the chemical potential. Next, we calculate the second-order EOMs of those newly appeared higher-order GFs on the RHS of Eq. (4), e.g., the EOM of the GF \( \ll f_{l\sigma} ; f_{m\sigma}^\dagger \gg \) is

\[ \langle \omega + \mu - \varepsilon_{f\sigma} \rangle \ll f_{l\sigma} ; f_{m\sigma}^\dagger \gg = \]

\[ \langle [f_{l\sigma} , f_{m\sigma}^\dagger]_+ \rangle + U_{ll} \ll \hat{n}_{l\sigma} f_{\sigma}^\dagger f_{m\sigma} \gg + \sum_{l',l \neq l} (U_{l'l\sigma} \ll \hat{n}_{l'\sigma} f_{l\sigma} ; f_{m\sigma}^\dagger \gg + U_{l'l\sigma} \ll \hat{n}_{l'\sigma} f_{l\sigma} ; f_{m\sigma}^\dagger \gg) + \sum_{k} V_{lk\sigma} \ll c_{lk\sigma} ; f_{m\sigma}^\dagger \gg - \sum_{l',l \neq l} I_{ll'\sigma} \ll f_{l'\sigma} ; f_{m\sigma}^\dagger \gg, \]
n_{m\sigma} f_{l \sigma} f_{l' \sigma} f_{m\sigma} \gg \) as an illustration we obtain

\begin{align*}
(\omega + \mu - \varepsilon_{f \sigma}) & \ll n_{m\sigma} f_{l \sigma} f_{l' \sigma} f_{m\sigma} \Rightarrow \\
&= (n_{m\sigma} f_{l \sigma} f_{l' \sigma} f_{m\sigma}) + U_{m\sigma} \ll n_{m\sigma} f_{l \sigma} f_{m\sigma} \\
&+ U_{l\sigma} \ll n_{l\sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma} \\
&+ U_{l\sigma} n_{l\sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma} \\
&+ \sum_{l', l'' \neq (l, m)} (U_{l' \sigma} \ll n_{l' \sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma}) \\
&+ U_{l' \sigma} n_{l' \sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma}) \\
&+ \sum_{k} (-V_{mk\sigma} \ll c_{m\sigma}^{\dagger} f_{m\sigma} f_{l}\sigma f_{m\sigma}) \\
&+ V_{lk\sigma} n_{l\sigma} c_{k\sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma} ) \\
- I_{m} & \ll f_{l' \sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma} \\
- & \sum_{l', l'' \neq (l, m)} I_{m} \ll f_{l' \sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma} \\
+ I_{m} & \ll f_{l' \sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma} \\
&+ \sum_{l', l'' \neq (l, m)} I_{m} \ll n_{l'' \sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma} \\
&+ \sum_{l', l'' \neq (l, m)} I_{m} \ll n_{l'' \sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma}, \quad (6)
\end{align*}

where \( l' \neq (l, m) \) means \( l' \neq l \) and \( l' \neq m \). The last six terms on the RHS are generated by the on-site inter-orbital single electron hoppings.

Next, noting that

\begin{align*}
[f_{m\sigma}, f_{l' \sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma}] &= -f_{l' \sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma}, \quad (7) \\
[f_{m\sigma}, f_{l' \sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma}] &= -f_{l' \sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma}, \quad (8)
\end{align*}

we recognize that \( \ll f_{l' \sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma} \ll f_{m\sigma} f_{l' \sigma} f_{m\sigma} f_{l}\sigma \) in Eq. (6) is actually associated with the pair-hopping term. While \( \ll f_{l' \sigma} f_{m\sigma} f_{l}\sigma f_{m\sigma} \ll f_{m\sigma} f_{l' \sigma} f_{m\sigma} f_{l}\sigma \) corresponds to the so-called spin-flip exchange term.\(^{13,10}\) One can note that both the pair-hopping and spin-flip exchange terms can be reproduced by on-site inter-orbital single hopping terms. The pair hopping is thus a special type of double hopping in which both the two electrons in the \( l \)-th orbital hop to the \( m \)-th orbital simultaneously. Similarly, the spin-flip exchange term is a certain kind of double hopping, in which one particle with spin \( \sigma \) hops from the \( l \)-th orbital to the \( m \)-th orbital as well as one particle with spin \( \sigma' \) hops from \( m \)-th orbital to \( l \)-th orbital at the same time, where \( \sigma \neq \sigma' \). Thus we have given them a more general definition for arbitrary multi-orbital systems. Notably, in a multi-orbital system the possible double hoppings are not only the pair-hopping and the spin-flip exchange term. The other four terms in the last six terms on the RHS of Eq. (6) also correspond to certain forms of on-site inter-orbital double hoppings at an equal time limit.

If we continue the procedure to calculate higher-order EOMs, more and higher-order GFs will appear which are associated with higher-order interactions. In view of the above, the on-site inter-orbital single hoppings \( f_{m'\sigma} f_{l'\sigma} f_{m\sigma} f_{l\sigma} \) are the most basic physical interactions in Hubbard-like strongly correlated systems. These will furnish the higher-order double-time double-hopping interactions, \( f_{l'\sigma} f_{m'\sigma} f_{l\sigma} f_{m\sigma} \), that is, one single hopping occurs at time \( t_{1} \) and another hopping occurs at time \( t_{2} \). In the same manner higher-order hoppings, as three-time three-hopping interactions and four-time four-hopping interactions \( etc. \) will appear. These multi-hopping interactions are embedded in our formulation and appear in higher-order EOMs.

Next, we investigate in more detail the double-time double hopping at equal times, i.e., the two hoppings occur simultaneously. The general form for the double hoppings is \( f_{l'\sigma} f_{m'\sigma} f_{l\sigma} f_{m\sigma} \), where \( l' \neq l \) and \( m' \neq m \). This equation gives all the double hoppings that possibly exist in a multi-orbital system. When \( l' = m' \), \( l = m \), \( \sigma' \neq \sigma \), it is the pair-hopping between orbitals.\(^{13,10}\) When \( l' = m \), \( l = m' \), \( \sigma' \neq \sigma \), it is the spin-flip exchange term. The whole set of on-site double hoppings include several kinds of interactions. If separated according to initial and final orbital of the hopping, we can schematically represent them as shown in Fig. 1 Writing

\begin{align*}
&f_{l'\sigma} f_{m'\sigma} f_{l\sigma} f_{m\sigma} = \sum_{m \sigma, m' \neq l, \sigma' \neq \sigma} f_{m\sigma} f_{l'\sigma} f_{l\sigma} f_{m\sigma} \\
&+ \sum_{m \neq l', \sigma', m' \neq l, \sigma' \neq \sigma} f_{m\sigma} f_{l'\sigma} f_{m\sigma} f_{l\sigma} \\
&+ \sum_{m' \neq l', l' \neq m', \sigma' \neq \sigma} f_{m'\sigma} f_{m'\sigma} f_{l'\sigma} f_{l\sigma} + f_{l'\sigma} f_{l'\sigma} f_{m\sigma} f_{m\sigma} \\
&+ \sum_{(l' m' \neq) \text{ all different}} f_{l'\sigma} f_{l'\sigma} f_{l\sigma} f_{m\sigma}, \quad (9)
\end{align*}

FIG. 1: Illustration of possible simultaneous double hoppings in a multi-orbital system. The straight arrows label the spin up or down of an electron and the bend arrow gives the hopping direction. (a) the spin-flip exchange term; (b) the pair-hopping term; (c) two electrons in different orbitals with different spins hop to the same orbital; (d) two electrons in one orbital with different spins hop to different orbitals; (e) two electrons in different orbitals hop to different orbitals.
where the first term is just the spin-flip term and the second term is the pair-hopping term. Note that the fifth term can split into three sub-terms according to the two spins present in different orbitals. And, Fig. 1 only gives an illustration of the double hoppings, the terms should sum over all orbitals. If there are less orbitals, the number of terms will be reduced accordingly.

For three-hopping, four-hopping, and even higher-order multi-hopping interactions, a schematic representation similar to Fig. 1 can be made.

From the above discussions, it can be recognized that the usually studied spin-flip and pair-hopping terms are higher-order consequences of the on-site inter-orbital single electron hoppings and Coulomb interactions. As the on-site inter-orbital single hopping is of lower order it will have a larger weight (or probability) than the spin-flip and pair-hopping terms. We note that, from the Eqs. (4) and (5), the off-diagonal GFs \( \langle f_{i\sigma} f_{j\sigma}^\dagger \rangle \) relate to the on-site inter-orbital single hopping. If one neglects the on-site inter-orbital single hopping but includes the pair-hopping and spin-flip terms in a model Hamiltonian, the obtained off-diagonal GFs will be inexact. Furthermore, in any self-consistent theory where the number of orbitals is larger than two, more inter-orbital double hopping terms have to be included, as shown by Eq. (9). Such terms can be well treated in our theory, because they will be automatically generated from the on-site inter-orbital single hoppings.

The hopping amplitudes \( t_{lm\sigma} \) have the physical meaning of the hopping amplitudes. They are to be determined for each studied system, but not fixed. For example, if in a certain system the \( \sigma \) spin channel is fully occupied for one orbital all the incoming single hoppings of spin \( \sigma \) into this orbital will be suppressed. In a paramagnetic system one can consider, for simplicity, that each orbital has the same incoming and outgoing inter-orbital single hoppings. Importantly, all studies of higher-order hopping terms must be made on top of having the on-site inter-orbital single hoppings included.

As an example we have numerically studied the two-orbital Hubbard model in paramagnetic case to show the influence of the on-site inter-orbital hopping. This interaction was explored in Ref. 15 but not in a self-consistent manner. Theoretically, in the situation where the neighboring sites are identical to the impurity site, the influence of the on-site inter-orbital hopping is similar to that of the inter-site inter-orbital hopping, as the orbitals on the neighboring sites are identical to the orbitals on the impurity site. In order to cleanly distinguish the contribution of the on-site inter-orbital hopping from that of the inter-site inter-orbital hopping, we set the inter-site inter-orbital hopping and on-site inter-orbital fluctuations to zero. We used the MO-EOM impurity solver of Ref. 10 in combination with genetic algorithm techniques 12,19.

The computed quasiparticle densities of states (DOS) are shown in Figs. 2 and 3 for the cases that the two orbitals have identical or different band widths, respectively. Line 1) is calculated without on-site inter-orbital hopping, line 2) with the same inter-site intra-orbital hopping and a nonzero on-site inter-orbital hopping, and line 3) with both inter-site intra-orbital and on-site inter-orbital hopping but the total hopping amplitude equals that for line 1) 20. The narrow orbital’s DOS for lines 2) and 3) with \( U=3 \) in (c) have been shifted vertically for visibility by 0.1 and 0.05, respectively.

FIG. 2: Quasiparticle DOS for the paramagnetic half-filled two-orbital Hubbard model on the Bethe lattice, computed for the case of identical band widths. The parameters used are \( D_2=D_1=1 \), temperature \( T=0.01 \), and the Coulomb \( U \) as specified in the panels. Line 1) is obtained for the inter-site intra-orbital hopping parameters \( t_2=t_1=D_1/2 \) and no on-site inter-orbital hopping. Line 2) uses the same parameters as 1) but the on-site inter-orbital hopping \( t_{12}=t_1/2 \). Line 3) uses the parameter \( t_{12}^0=t_1+t_{12}=D_1/2 \) and \( t_1=t_{12} \).

FIG. 3: As Fig. 2 but for the case of different band widths. The left panels depict the DOS for the narrow orbital and the right panels that for the wide orbital. The parameters used are \( D_2=2D_1=2 \) and \( T=0.01 \). The lines 1) - 3) are obtained as explained in Fig. 2 only \( t_2 \) is modified correspondingly with \( D_2 \). The narrow orbital’s DOS for lines 2) and 3) with \( U=3 \) in (c) have been shifted vertically for visibility by 0.1 and 0.05, respectively.
teraction the Hubbard bands are somewhat broadened. This happens because the on-site inter-orbital hopping effectively increases the total hopping amplitude so that the electrons gain some itineracy. In Fig. 3 line 1) shows that in the absence of on-site inter-orbital hoppings there is an orbital selective Mott transition (OSMT). However, from both lines 2) and 3) one observes that the narrow and wide orbitals simultaneously change from metallic states to insulating states along with the increase of $U$. Therefore, the OSMT shown with line 1) is suppressed with the inclusion of the on-site inter-orbital hopping.

To summarize, we introduced a general MO-Hubbard model that can readily be employed to study a broad range of multi-orbital systems. We have shown analytically and numerically the influence of introducing the on-site inter-orbital single hoppings. When these exist in a correlated electron system they will greatly change its properties. Higher-order effects, such as spin-flip exchange and double hoppings, have to be studied on top of the on-site inter-orbital single hoppings, which, as outlined, can be done in a well-defined way for arbitrary correlated systems. The developed theory is expected to be beneficial for studies of unsolved correlation-related phenomena and to trigger inspiring theoretical studies and discoveries.

We thank the Swedish Research Council (VR) and SKB for financial support. Computer time from the Swedish National Infrastructure for Computing (SNIC) is acknowledged.

---

1 N. Grewe and F. Steglich, Handbook on the Physics and Chemistry of Rare Earths, ed. K.A. Gschneidner, Jr. and L. Eyring (Elsevier, Amsterdam, 1991), Vol. 14, p. 343.
2 M. Imada, A. Fujimori, and Y. Tokura, Rev. Mod. Phys. 70, 1039 (1998).
3 J. Hubbard, Proc. Roy. Soc. A 276, 238 (1963).
4 M. C. Gutzwiller, Phys. Rev. Lett. 10, 59 (1963).
5 J. Kanamori, Prog. Theor. Phys. 30, 275 (1963).
6 J. Quintanilla and C. Hooley, Physics World 22, 32 (2009).
7 T. Maier, M. Jarrell, Th. Pruschke, and J. Keller, Phys. Rev. Lett. 85, 1524 (2000).
8 D. J. Scalapino, J. Supercond. Nov. Magn. 19, 195 (2006).
9 Q. Feng and P. M. Oppeneer, J. Phys.: Condens. Matter 23, 425601 (2011).
10 W. Metzner and D. Vollhardt, Phys. Rev. Lett. 62, 324 (1989).
11 Q. Feng and P. M. Oppeneer, J. Phys.: Condens. Matter 24, 055603 (2012).
12 A. Georges and G. Kotliar, Phys. Rev. B 45, 6479 (1992).
13 D. N. Zubarev, Sov. Phys. Usp. 3, 320 (1960).
14 A. Koga, N. Kawakami, T. M. Rice, and M. Sigrist, Phys. Rev. B 72, 045128 (2005).
15 P. Werner, E. Gull, and A. J. Millis, Phys. Rev. B 79, 115119 (2009).
16 X. Dai, G. Kotliar, and Z. Fang, arXiv:cond-mat/0611075.
17 Q. Feng, Y.-Z. Zhang, and H. O. Jeschke, Phys. Rev. B 79, 235112 (2009).
18 Q. Feng, Ph.D. Thesis, University Frankfurt (2009).
19 Specifically, the parameters used in Fig. 2 are: line 1) $t_1=t_2=D_1/2$, $t_{12}=t_{21}=0$; line 2) $t_1=t_2=D_1/2$, $t_{12}=t_{21}=D_1/4$; line 3) $t_1=t_2=D_1/4$, $t_{12}=t_{21}=D_1/4$. The parameters used in Fig. 3 are: line 1) $t_1=D_1/2$, $t_2=D_2/2$, $t_{12}=t_{21}=0$; line 2) $t_1=D_1/2$, $t_2=D_2/2$, $t_{12}=t_{21}=D_1/4$; line 3) $t_1=D_1/4$, $t_2=D_2/2-D_1/4$, $t_{12}=t_{21}=D_1/4$. The parameters used in Fig. 4 are: line 1) $t_1=D_1/2$, $t_2=D_2/2$, $t_{12}=t_{21}=0$; line 2) $t_1=D_1/2$, $t_2=D_2/2$, $t_{12}=t_{21}=D_1/4$; line 3) $t_1=D_1/4$, $t_2=D_2/2-D_1/4$, $t_{12}=t_{21}=D_1/4$.