Entanglement spectrum in cluster dynamical mean-field theory

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Received 23 June 2014
Accepted for publication 10 December 2014
Published 9 January 2015

Online at stacks.iop.org/JSTAT/2015/P01016
doi:10.1088/1742-5468/2015/01/P01016

Abstract. We study the entanglement spectrum of the Hubbard model at half filling on a kagome lattice. The entanglement spectrum is defined by the set of eigenvalues of a reduced thermal density matrix, which is naturally obtained in the framework of the dynamical mean-field theory. Adopting the cluster dynamical mean-field theory combined with continuous-time auxiliary-field Monte Carlo method, we calculate the entanglement spectrum for a three-site triangular cluster in the kagome Hubbard model. We find that the results at the three-particle sector well capture the qualitative nature of the system. In particular, the eigenvalue of the reduced density matrix, corresponding to the chiral degrees of freedom, exhibits a characteristic temperature scale \( T_{\text{chiral}} \), below which a metallic state with large quasiparticle mass is stabilized. The entanglement spectra at different particle number sectors also exhibit characteristic changes around \( T_{\text{chiral}} \), implying the development of intertriangular ferromagnetic correlations in the correlated metallic regime.

Keywords: Hubbard model (theory), entanglement in extended quantum systems (theory)
1. Introduction

It is a central issue of condensed matter physics to describe the nature of strongly correlated electron systems both qualitatively and quantitatively. Dynamical mean-field theory (DMFT) [1] provides a promising theoretical tool for this purpose. DMFT bridges the gap between two solvable cases, the atomic limit and the free fermion limit, and successfully describes the properties of correlated electron systems. In particular, correlation-driven metal-insulator transition (MIT) is successfully explained by DMFT. In addition to the success in describing various thermodynamic quantities in a quantitative way, DMFT also provides a qualitative understanding of MIT in terms of insulating Hubbard bands and a renormalized conduction band. These aspects of MIT can be qualitatively captured by focusing on a simple observable quantity, a doublon density, i.e. the fraction of sites where spin-up and spin-down states are simultaneously occupied. The temperature dependence of this quantity is easily computable in the scheme of DMFT and it, indeed, gives crucial insights into the nature of MIT [2].

While DMFT is quite useful to understand MIT in a simple single-orbital correlated model, however, realistic compounds have complex degrees of freedom composed of spin and orbitals and their spatial correlations, which lead to rich behavior in MIT. Correspondingly, the cluster extension of DMFT (CDMFT) [3] has been used to explore this fertile field, by taking into account spin, orbital, and intersite degrees of freedom, in
a quantitative way. However, the complexity of the system inevitably makes it difficult to achieve qualitative understanding. It is then, desirable to set up a measure to characterize these systems in a qualitative way, i.e. the counterpart of doublon density in these complex systems. For this purpose, in this article, we propose the entanglement spectrum as a measure to qualitatively describe the systems composed of multiple degrees of freedom.

This quantity can be computed with a small numerical cost, and gives an account of the qualitative nature of the system. In contrast to the usual terminology of entanglement spectrum, it is defined from the reduced thermal density matrix. Accordingly, standard properties required for the entanglement spectrum, e.g. the area law of entanglement entropy, will not be satisfied. Nonetheless, we show that this quantity will be quite useful to disentangle the complexity of correlated systems with multiple degrees of freedom.

This article is organized as follows. In section 2, we introduce CDMFT, and especially the algorithm to compute the entanglement spectrum. In section 3, we show the results of the entanglement spectrum, and discuss how this quantity is associated with the actual behavior of the systems, by taking the Hubbard model on the kagome lattice as an example. Section 4 is devoted to the discussion and summary.

2. Model and method

2.1. Model

As a prototypical model of correlated electron systems composed of multiple degrees of freedom, we consider the single-orbital Hubbard model on the kagome lattice. The Hamiltonian is given in the standard notation as

\[ H = -t \sum_{\langle ij \rangle, \sigma} (a_{i\sigma}^\dagger a_{j\sigma} + \text{h.c.}) + U \sum_i n_{i\uparrow} n_{i\downarrow}, \tag{1} \]

where the sum of \( \langle ij \rangle \) is taken over the nearest-neighbor sites on the kagome lattice (figure 1). Hereafter we set \( t = 1 \) and the Boltzmann constant \( k_B = 1 \), and fix the electron density at half filling. This model has been extensively studied as a minimal model with electron correlation and geometrical frustration [5–11].

The kagome lattice has three sites in the unit cell, which constitute the internal degrees of freedom. The interplay between the intra-cell degrees of freedom and electron correlation leads to nontrivial heavy-fermion behavior in the metallic region, as we will show later.

2.2. Cluster DMFT

The model (1) is not easily tractable with a standard unbiased numerical method, e.g. exact diagonalization or quantum Monte Carlo simulation, due to the exponentially large numerical cost or notorious negative sign problem. Here we adopt the cluster dynamical mean-field theory (CDMFT) [1, 3] as an approximate method to take into account the electron correlation. In CDMFT, the Hubbard model (1) on the original infinite lattice is mapped to the effective Anderson model with interacting impurity sites on an \( N_c \)-site cluster embedded in the effective medium. In this study, we mainly focus on the case of \( N_c = 3 \), namely, we adopt the triangle unit cell as an impurity cluster, as shown in figure 1(a). In order to solve the effective Anderson model, we adopt the
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Figure 1. (a) Schematic picture of the mapping of the kagome lattice to three-site cluster in the CDMFT calculations. The three-site density matrix is calculated for the numbered three sites. (b) Schematic pictures of the doublon, spin-polarized, and chiral states.

2.3. Entanglement spectrum

The doublon density, defined by $D \equiv \langle n_{i\uparrow}n_{i\downarrow} \rangle$, is a standard measure to characterize the effect of electron correlation. $D$ corresponds to the fraction of sites where the up-spin and down-spin states are simultaneously occupied. It shows a characteristic temperature change, accompanying a discontinuity between metallic and Mott insulating states. This quantity is, however, not enough to characterize the complicated energy scales in the correlated electron systems with multiple degrees of freedom.

To generalize $D$ for the systems with multiple degrees of freedom, we focus on the aspect of doublon density as an eigenvalue of the reduced density matrix. In DMFT, the whole system is divided into the single target site and the environment. By tracing out the environment degrees of freedom, the reduced density matrix $\hat{\rho}$ is constructed for the target site, as

$$\hat{\rho} = \rho_{\uparrow\downarrow}\langle \uparrow\downarrow | + \rho_{\uparrow\uparrow}\langle \uparrow\uparrow | + \rho_{\downarrow\downarrow}\langle \downarrow\downarrow | + \rho_0|0\rangle\langle 0 |.$$ (2)

Here, $|\uparrow\downarrow \rangle = a_{\uparrow}^\dagger a_{\downarrow}^\dagger |0\rangle$, $|s\rangle = a_s^\dagger |0\rangle$, and $|0\rangle$ are a doubly-occupied state, a singly-occupied state with spin $s$ electron, and the empty state, respectively. In equation (2), $\hat{\rho}$ takes a diagonalized form by choosing the particle number $N_p$ and spin quantum number to label the bases, in the systems with conserved particle number and spin. In this representation, the doublon density $D$ is nothing but the coefficient $\rho_{\uparrow\downarrow}$, i.e. the eigenvalue of $\hat{\rho}$ in the $N_p = 2$ sector, in equation (2).

2.4. CDMFT treatment

Now, let us consider the CDMFT calculations of the entanglement spectrum for the systems composed of multiple degrees of freedom. The cluster impurity mapping of
CDMFT divides the system into the (A) cluster sites and (B) environment, which gives a natural definition of reduced density matrix, \( \hat{\rho}_A \equiv \text{Tr}_B \hat{\rho}_\text{thermal} \). Here \( \hat{\rho}_\text{thermal} \equiv \exp(-\beta H) / \text{Tr}_A \exp(-\beta H) \) is the equilibrium thermal density matrix, and the trace \( \text{Tr}_A(B) \) is taken over the cluster (environment) degrees of freedom. The reduced density matrix \( \hat{\rho}_A \) can be expanded with the \( 4^N_c \) basis states \( \{ \psi_p \} \) within the cluster sites, as \( \hat{\rho}_A = \rho_{A,pq} |\psi_p\rangle \langle \psi_q| \). \( \hat{\rho}_A \) can be diagonalized to give \( 4^N_c \) eigenvalues, \( \{ \rho_{A,pq} \} \), which constitute the entanglement spectrum.

In order to obtain each coefficient \( \rho_{A,pq} \) in the CDMFT formalism, it is convenient to adopt the second quantized representation. We choose the basis to diagonalize the particle number and the \( z \) component of spin:

\[
|\phi\{m_{is}\}\rangle = (a_{1\uparrow}^\dagger)^{m_{1\uparrow}} (a_{1\downarrow}^\dagger)^{m_{1\downarrow}} \cdots (a_{N_c\uparrow}^\dagger)^{m_{N_c\uparrow}} (a_{N_c\downarrow}^\dagger)^{m_{N_c\downarrow}} |0\rangle.
\]

Here, \( m_{is} = 0 \) or \( 1 \) is the number of electrons at site \( i \) and spin \( s \). The projection operator \( |\psi_p\rangle \langle \psi_p| \) can be simply expressed as

\[
|\phi\{m_{is}\}\rangle \langle \phi\{m_{is}\}| = \hat{f}_{1\uparrow} [m_{1\uparrow}] \hat{f}_{1\downarrow} [m_{1\downarrow}] \cdots \hat{f}_{N_c\uparrow} [m_{N_c\uparrow}] \hat{f}_{N_c\downarrow} [m_{N_c\downarrow}],
\]

where \( \hat{f}_{is}[m] = 1 - n_{is} \) \( (n_{is}) \) for \( m = 0 \) \( (1) \). Accordingly, the diagonal components of \( \hat{\rho}_A \), \( \rho_{A,pp'} \), can be obtained as the expectation value of the corresponding operator \( |\phi\{m_{is}\}\rangle \langle \phi\{m_{is}\}| \).

The calculation of the off-diagonal components of \( \hat{\rho}_A \) is slightly more involved. To depict this, suppose the three-site cluster \( (N_c = 3) \) and consider an off-diagonal component, \( |\uparrow, 0, 0\rangle \langle 0, \uparrow, 0| \). Here, we denote \( |\uparrow\uparrow, \uparrow, 0\rangle \equiv a_{1\uparrow}^\dagger a_{2\uparrow}^\dagger a_{2\downarrow}^\dagger |0\rangle \), and so on. Using the expressions, we can write down the creation operator as

\[
a_{i\uparrow}^\dagger = \sum_{k_2,k_3} (|\uparrow, k_2, k_3\rangle \langle 0, k_2, k_3| + |\downarrow, k_2, k_3\rangle \langle 1, k_2, k_3|).
\]

Here, the summation of \( k_i \) is taken over the four possible states at site \( i \): \( |0\rangle, |\uparrow\rangle, |\downarrow\rangle \) and \( |\uparrow\downarrow\rangle \). Then, we obtain

\[
|\uparrow, 0, 0\rangle \langle 0, \uparrow, 0| = a_{1\uparrow}^\dagger a_{2\uparrow}^\dagger a_{2\downarrow}^\dagger |0, \uparrow, 0\rangle \langle 0, \uparrow, 0| = a_{1\uparrow}^\dagger a_{2\uparrow}^\dagger (1 - n_{1\uparrow}) (1 - n_{1\downarrow}) n_{2\uparrow} (1 - n_{2\downarrow}) (1 - n_{3\uparrow}) (1 - n_{3\downarrow}) = a_{1\uparrow}^\dagger a_{2\uparrow}^\dagger (1 - n_{1\uparrow}) a_{2\downarrow}^\dagger (1 - n_{2\downarrow}) (1 - n_{3\downarrow}).
\]

The expectation value of this product can be simplified with the help of Wick’s theorem,\( \rho_{\uparrow\downarrow, 0, 0} |0, \uparrow, 0\rangle \langle 0, \uparrow, 0| = \langle a_{1\uparrow}^\dagger a_{2\uparrow}^\dagger \rangle \langle (1 - n_{1\uparrow}) \rangle \langle (1 - n_{2\uparrow}) \rangle \langle (1 - n_{3\uparrow}) \rangle \langle (1 - n_{3\downarrow}) \rangle \rangle + \cdots \)

where each correlator \( \langle \cdots \cdots \rangle \) can be evaluated by Monte Carlo sampling. All the other components of the reduced density matrix, \( \{ \rho_{A,pq} \} \), can be expressed as the product of fermion creation and annihilation operators, and are evaluated in a similar way.

### 2.5. Entanglement spectrum of the kagome Hubbard model

Now we focus on the CDMFT analysis of the kagome Hubbard model with \( N_c = 3 \). In this case, the total \( 4^N_c = 64 \) eigenvalues of \( \hat{\rho}_A \) can be classified into several irreducible representations under the \( U(1) \otimes SU(2) \otimes C_3 \) symmetry of the kagome Hubbard model. Each state can be labeled by three quantum numbers: particle number \( N_p = 0 \) \( \pm 6 \), the \( z \) component of total spin, \( s_z \), and the helicity \( h = 0, \pm 1 \). The helicity is defined from the

\doiref{10.1088/1742-5468/2015/01/P01016}{J. Stat. Mech. (2015) P01016}
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response of the state to the 120 degrees rotation $\hat{L}$, with respect to the axis perpendicular to the kagome plane at the center of a triangle, as $\hat{L}|\psi\rangle = \omega^h|\psi\rangle$, with $\omega = e^{\frac{i\pi}{3}}$.

In the $N_p = 3$ sector, the following states are of particular interest: the doublon state, $|D\rangle = \frac{1}{\sqrt{3}}(|\uparrow \downarrow, \uparrow, 0\rangle + |0, \uparrow \downarrow, \uparrow\rangle + |\uparrow, 0, \uparrow \downarrow\rangle)$, the spin-polarized state, $|S\rangle = |\uparrow, \uparrow, \uparrow\rangle$, and the chiral state, $|K\rangle = \frac{1}{\sqrt{3}}(|\downarrow, \uparrow, \uparrow\rangle + \omega|\uparrow, \downarrow, \uparrow\rangle + \bar{\omega}|\uparrow, \uparrow, \downarrow\rangle)$ ($\bar{\omega} = \omega^2$). See figure 1(b). We denote the eigenvalues corresponding to these states as $\rho_D$, $\rho_S$, and $\rho_K$, respectively. Among these states, the chiral states are of special interest. $|K\rangle$ and its time-reversal and reflection conjugate states constitute fourfold degeneracy, labeled by $s_z = \pm 1/2$ and $h = \pm 1$.

3. Results

In this section, we will introduce our results on the entanglement spectrum. In section 3.1, we will introduce our previous results on the heavy-fermion behavior, and associate it with the temperature dependence of the entanglement spectrum at the $N_p = 3$ sector. In section 3.2, we will show the entanglement spectrum at $N_p \neq 3$ sectors, and discuss how the change of electron coherency is reflected in these spectra.

3.1. Heavy-fermion behavior

First, we summarize our results on the entanglement spectrum at $N_p = 3$ sector, together with the thermodynamic behavior of this system, which were previously discussed in [5]. In figure 2(a), we plot the temperature dependence of $\rho_D$, $\rho_S$, and $\rho_K$ at $U = 6$, which is slightly smaller than the critical value of $U$ for MIT, $U_c \sim 8.3$ [8]. In this correlated metallic region, the negative sign problem is not severe: for instance, the average of the sign is larger than 0.99 at $U = 6$ and $T = 0.1$.

At high temperature $T \sim 100$, all the eigenvalues take almost the same value $\sim 1/64$ and all the 64 states are equally probable within the three-site cluster in the high-temperature limit. With decreasing temperature, $\rho_D$ decreases at $T = T_{\text{charge}} \simeq 2.0$, while the other states continue to show a gradual increase. As the temperature is further lowered, $\rho_S$ turns to decrease at $T_{\text{spin}} \simeq 0.37$. Meanwhile, $\rho_K$ continues to show enhancement even below $T_{\text{spin}}$, until another temperature scale $T_{\text{chiral}} \simeq 0.18$ is reached. These three energy scales are defined from the inflection points of the corresponding eigenvalues.

The two energy scales $T_{\text{charge}}$ and $T_{\text{spin}}$ can be basically understood from the energetics of an isolated triangle. $T_{\text{charge}}$ is associated with the on-site repulsion $U$, and $T_{\text{spin}}$ is related to the energy scale of spin sector, $J = 4t^2/U$ [5]. These states are suppressed, if temperature is lowered below $T_{\text{charge}}$ or $T_{\text{spin}}$. In other words, below $T_{\text{spin}}$, the chiral states are stabilized at each triangle, and loosely interact with each other, storing a large amount of entropy $\sim \log 4$ per triangle, arising from the fourfold degeneracy of the chiral state.

On the other hand, $T_{\text{chiral}}$ cannot be associated with the energy scale of an isolated triangle. This temperature rather indicates the energy scale of electron coherency, i.e. the

$^1$ Although $|D\rangle$ has finite off-diagonal components with the state, $\frac{1}{\sqrt{3}}(|\downarrow, \uparrow, \uparrow\rangle + |\uparrow, \downarrow, \uparrow\rangle + |\uparrow, \uparrow, \downarrow\rangle)$, we ignore it and focus on the diagonal component, to characterize the energy scale associated with the charge degree of freedom.

doi:10.1088/1742-5468/2015/01/P01016
Figure 2. (a) The entanglement spectrum at the $N_p = 3$ sector for the kagome Hubbard model at half filling for $U = 6$. $T_{\text{charge}} \simeq 2.0$, $T_{\text{spin}} \simeq 0.37$ and $T_{\text{chiral}} \simeq 0.18$ are the characteristic temperature scales of charge, spin, and chiral degrees of freedom, which are determined from the inflection points of corresponding eigenvalues. (b) The temperature dependence of specific heat at $U = 0$ and 6. (c) The temperature dependence of entropy per triangular cluster, $S$, normalized by its value in the high-temperature limit, log 64. (d) Imaginary part of the diagonal component of self-energy is plotted for $T = 0.1, 0.2$ and 0.4. The horizontal axis is the Matsubara frequency, $\varepsilon_p = (2p + 1)\pi T$, with integer $p$.

The system gains kinetic energy from electron motion between triangles below $T_{\text{chiral}}$. Enhanced electron coherency leads to the formation of a Fermi liquid state below $T_{\text{chiral}}$. This metallic state shows particularly large quasiparticle mass, since the large entropy associated with the chiral states is now bequeathed to itinerant electrons, which leads to the steep slope of the specific heat as shown in figure 2(b). In fact, the entropy takes $\sim \log 4$ at $T_{\text{chiral}}$ as shown in figure 2(c).

The Fermi liquid formation with large quasiparticle mass can also be confirmed from the enhanced magnetic susceptibility $\chi$ [5]. As $T$ is lowered, $\chi$ approaches a constant Pauli paramagnetic value, which grows with increasing $U$, reflecting the enhanced quasiparticle mass. The Fermi liquid formation also leads to the development of a sharp coherence peak in the one-particle spectrum as discussed in [5]. The crossover to the Fermi liquid state is also clearly shown in the self-energy. In figure 2(d), we plot the imaginary part of the site diagonal component of self-energy, $\text{Im} \Sigma(i\varepsilon_p)$. $\text{Im} \Sigma(i\varepsilon_p)$ shows insulating behavior $\sim -1/i\varepsilon_p$ in the high $T$ regions. Meanwhile, as $T$ is lowered, it approaches the Fermi liquid behavior, i.e. $\text{Im} \Sigma(i\varepsilon_p) \to 0$ as $\varepsilon_p \to 0$.

3.2. Charge sectors

The development of electron coherency at $T \sim T_{\text{chiral}}$ is reflected in the entanglement spectra at various particle number sectors. The electron itinerancy enhances charge fluctuations, i.e. each triangle becomes more likely to have particle numbers different from the average value, $N_p \neq 3$. In figure 3, we plot the entanglement spectra at $N_p = 2$ and 4 sectors. We here show the eigenvalues of the states labeled by the $z$ component of total spin $s_z$, helicity $h$, and doublon number $d$. While the states with different numbers of doublon mix with each other, the mixing can be ignored at low temperatures $T \ll T_{\text{charge}}$. 

doi:10.1088/1742-5468/2015/01/P01016
Figure 3. Temperature dependence of the entanglement spectra at (a) $N_p = 2$ and (b) $N_p = 4$ sectors. $T_{\text{chiral}}$ is shown by the vertical dashed line. The plotted data correspond to the states labeled by the set of quantum numbers: in (a), $(s_z, h, d) = (0, 0, 0), (1, 1, 0), (0, 1, 0), (0, 0, 1), (1, 0, 0), \text{and} (0, 1, 1)$ from top to bottom at $T = 1.6$, while in (b), $(s_z, h, d) = (1, 0, 1), (0, 0, 1), (0, 1, 1), (1, 1, 1), (0, 0, 2), \text{and} (0, 1, 1)$ from top to bottom.

Figure 3(a) shows the temperature dependence of entanglement spectra at $N_p = 2$. The dominant contribution comes from the state with $(s_z, h, d) = (0, 0, 0)$. This state keeps the largest eigenvalue of reduced density matrix from high temperature, and starts to develop at the temperature where $\rho_K$ becomes maximum, showing a steep enhancement at $T \sim T_{\text{chiral}}$.

Figure 3(b) gives the behavior of entanglement spectra at $N_p = 4$. In this sector, the states with a finite total spin have the largest eigenvalue, in contrast to $N_p = 2$. Indeed, the states with $s_z = \pm 1$ have the lowest eigenenergy within the states at $N_p = 4$ in the three-site Hubbard model, which explains why they keep dominant. These states, together with several subdominant states, also exhibit a slight enhancement at $T \sim T_{\text{chiral}}$.

These temperature dependences of the entanglement spectra around $T = T_{\text{chiral}}$ give a clue to understanding the nature of metallic states at low temperatures. While the eigenvalue corresponding to the chiral states, $\rho_K$, shows a slight reduction at $T \sim T_{\text{chiral}}$ as shown in figure 2(a), it still keeps dominant among the 64 possible states; locally on each triangle, the chiral state is most likely realized even below $T \sim T_{\text{chiral}}$. Hence, it is reasonable to consider that electron coherency can be achieved by the enhancement of electron hopping in the sea of chiral states.

Suppose each triangle takes one of the fourfold degenerate chiral states. If an electron is moved from one triangle to a neighboring one, then the first triangle is left with two electrons, while the latter ends up with having four electrons. The enhanced eigenvalue of $s_z = 0$ state in the $N_p = 2$ sector suggests that the spin-up (spin-down) electron is more likely to hop from the chiral state with $s_z = 1/2 (-1/2)$. Meanwhile among the states in the $N_p = 4$ sector, $s_z = \pm 1$ states are dominant. This means that the spin-up (spin-down) electron tends to hop to the chiral states with $s_z = 1/2 (-1/2)$ (figure 4). Therefore, for the single-electron hopping process to create $s_z = 0$ state with $N_p = 2$ and $s_z = 1 (-1)$ state with $N_p = 4$ simultaneously, the chiral states have to have a pair of $s_z = 1/2 (-1/2)$ states.
Suppose all the triangles are in the state $|K\rangle$ shown in the top left. If one spin-up electron is transferred from the left bottom site of one triangle to the top site of the neighboring triangle, the first triangle will be left with one up-spin and one down-spin electrons with $s_z = 0$, while the second triangle will acquire total spin $s_z = 1$.

at neighboring triangles. In other words, the entanglement spectra imply that short-range ferromagnetic correlations are developed between neighboring triangles in the correlated metallic phase.

4. Discussion and summary

In this article, we have studied the entanglement spectra defined from the thermal density matrix in the Hubbard model at half filling on the kagome lattice. The cluster dynamical mean-field theory provides a convenient tool to analyze this quantity through the whole temperature range. Indeed, we have clarified that the entanglement spectra for a three-site cluster exhibits nontrivial temperature dependence.

At the $N_p = 3$ sector, the clear separation of energy scales between charge, spin, and chiral degrees of freedom is obtained. In particular, the characteristic temperature scale of chiral degrees of freedom, $T_{\text{chiral}}$, gives a good measure to the enhancement of electron coherency. In fact, the Fermi liquid state with large quasiparticle mass is formed below this temperature, as clearly shown in the analysis of specific heat and entropy.

Moreover, the developed electron coherency affects the entanglement spectra at $N_p = 2$ and 4 sectors. The dominant states at both sectors show enhancement at $T \sim T_{\text{chiral}}$, suggesting enhanced charge fluctuations due to the electron itinerancy. The growth of these states implies short-range inter-triangle ferromagnetic correlations in the low-temperature metallic state. While the complete characterization of the correlated metallic
state is left for a future task, the entanglement spectra will serve as a powerful tool for this purpose.

Acknowledgments

This work was supported by Grants-in-Aid for Scientific Research (No. 24340076, 26400339, and 24740221), the Strategic Programs for Innovative Research (SPIRE), MEXT, and the Computational Materials Science Initiative (CMSI), Japan.

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