Charge and spin stripe in La$_{2-x}$Sr$_x$NiO$_4$ ($x = \frac{1}{3}, \frac{1}{2}$)

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Electronic structure of stripe ordered La$_{2-x}$Sr$_x$NiO$_4$ is investigated. The system with $x = \frac{1}{3}$ is insulator, in LSDA+U calculations, and shows charge and spin stripe, consistent with the experimental results. Highly correlated system of $x = \frac{1}{2}$ is studied by using exact diagonalization of multi-orbital many body Hamiltonian derived from LDA calculations and including on-site and inter-site Coulomb interactions. The fluctuation of the residual spin on Ni$^{3+}$ (hole) site couples with the charge fluctuation between Ni$^{3+}$ and Ni$^{2+}$ states and this correlation lowers the total energy. The resultant ground state is insulator with charge and spin stripe of the energy gap 0.9 eV, consistent with observed one. The on-site Coulomb interaction stabilizes integral valency of each Ni ion (Ni$^{3+}$ and Ni$^{2+}$), but does not induce the charge order. Two quantities, inter-site Coulomb interaction and anisotropy of hopping integrals, play an important role to form the charge and spin stripe order in a system of $x = \frac{1}{2}$.

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I. INTRODUCTION

The stripe order of charge and spin has been found in several layered perovskites and organic conductors. Both of them are of pseudo two-dimensional (2D) electron system and have strong Coulomb interactions compared to hopping integrals. A single band system can be a model of organic conductors, while, in layered perovskites, a number of relevant orbitals depends on filling, crystal field, exchange splitting etc. The inter-site Coulomb interaction is essential to the charge stripe in organic conductors and, then, extended Hubbard model is adopted to explain the mechanism of stripe order there. In layered perovskites, however, an origin of the stripe order is still controversial particularly in the perovskites other than cuprate, while the order in cuprate is attracting much attention in conjunction with marked suppression of $T_c$ with hole doping and rich physics.

Nickel compound La$_{2-x}$Sr$_x$NiO$_4$ (LSNO) is a typical system of static stripe order of charge and spin. It is an insulator with the total spin $S = 0$ in wide range of Sr doping $0 \leq x \leq 0.9$. This stability of insulating phase is quite different from the variety of the phases in cuprate case: cuprate changes to metal, insulator and superconductor depending on the hole concentration. LSNO at $x = \frac{1}{3}$ shows the highest spin order temperature $T_{SO} \sim 200$K, because stripe order of charge and spin is commensurate with lattice periodicity. The periods of charge and spin stripe are not generally commensurate to the lattice. Incommensurability $\epsilon$ increases with increasing $x$ and satisfies $\epsilon \gtrsim x$ in the region $\frac{1}{3} > x > 0$, $\epsilon \lesssim x$ in the region $x > \frac{1}{3}$. Here, incommensurability $\epsilon$ is defined as displacement of peak positions of super structure from reciprocal lattice points. Increase of $\epsilon$ saturates in the region of $\frac{1}{2} > x > \frac{1}{3}$ with the value $\epsilon \sim 0.44$. In the region $x \gtrsim \frac{1}{3}$, there exists commensurate charge ordered phase without magnetic order, called a checkerboard type charge order, between $T_{SO} = 80$K and $T_{CO} = 480$K. Another experimental fact in LSNO is the dependence of the ratio between two lattice constants along with $c$- and $a$-axis ($c/a$-ratio) on $x$. The observed $c/a$-ratio has a maximum value at $x = \frac{1}{2}$, because holes are first doped into the $x^2 - y^2$-orbital in a region $x < \frac{1}{2}$ and, then, additional holes are doped into the $3z^2 - 1$-orbital when $x > \frac{1}{2}$.

We present two issues on LSNO in this paper by using both LSDA+U method and the exact diagonalization of many body Hamiltonian. The first issue is that the inter-site Coulomb interaction is essential to static charge order in doped layered perovskites. Not only Hartree energy but also correlation energy due to the inter-site Coulomb interaction (beyond Hartree-Fock approximation) is important. Because electron configurations fluctuate between hole (Ni$^{3+}$) and non-hole (Ni$^{2+}$) states by hopping, the correlation energy is maximized at $x = \frac{1}{2}$. In this situation, we need to diagonalize the many-body Hamiltonian to know the true ground state at $x = \frac{1}{2}$. We also need to consider JT distortion coupling between hole site and non-hole site in the multi-orbital system with fractional occupation, which is a essential difference from the case of single-orbital systems.

The stripe order in layered perovskites is, in some cases, attributed to the Jahn-Teller (JT) distortion. In the present system, however, this is not the case, because the JT distortion is not consistent with the local symmetry of the observed stripe order of charge and spin in $x = \frac{1}{2}$ LSNO. In the $e_g$ type JT distortion coupling with the $e_g$ orbitals, two oxygen atoms at opposite po-

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positions, centering Ni site, displace in opposite directions with each other. Consequently, the periodicity should be doubled along with -Ni-O- · · · O-Ni- line. It contradicts to the observed tripled structure. Distortion with ungerade mode can be consistent with the observed order but increases the total energy. The spin structure at \( x = \frac{1}{2} \) is inconsistent with the JT distortion, too.

There is another candidate for the origin of stripe order in layered perovskites. That is long-ranged Coulomb interaction. Also it is well known that the inter-site Coulomb interaction stabilizes the charge order in quarter filled (single-orbital) extended Hubbard model. Thus, the inter-site Coulomb interaction can stabilize the electronic structure of insulator with static stripe order. In the electronic structure of LSNO, degeneracy of 3d \( e_g \)-orbitals causes two important parameters: splitting between \( 3z^2-r^2 \) - 1-orbital and \( x^2-y^2 \)-orbital and the hopping integrals. It is reported that different values of these variables bring the system different order. We should get reliable values of these parameters from the first principles electronic structure calculations.

The present paper is organized as follows. The charge and spin stripe of LSNO (\( x = \frac{1}{2} \)) is discussed in Sec. II based on LSDA+U calculation. We show in Sec. III the problem that charge ordered solution is not found in LSDA+U calculations. Then we explain that the LSNO of \( x = \frac{1}{2} \) is highly correlated system and the correlation energy in LSDA+U method is not enough to stabilize the charge ordered solution as a ground state. In later Sections, exact diagonalization of many body Hamiltonian is employed to investigate the electronic structure of LSNO (\( x = \frac{1}{2} \)). The Hamiltonian derived from LDA calculation is explained in Sec. IV. Section V is devoted to discussion of the charge stripe order and charge correlation functions in the system of \( x = \frac{1}{2} \). The excitation spectra and the energy gap of the system with \( x = \frac{1}{2} \) is discussed in Sec. VI. The spin stripe order of LSNO (\( x = \frac{1}{2} \)) is discussed in Sec. VII. Finally Sec. VIII is the conclusion.

II. CHARGE AND SPIN STRIPE OF LSNO

\( x = \frac{1}{2} \) BY USING LSDA+U METHOD

LSDA+U method in conjunction with the linear muffin-tin orbital method with the atomic sphere approximation includes on-site Coulomb and exchange interaction, \( U, J \) with rotational invariant form. This on-site Coulomb term in LSDA+U Hamiltonian is called a “Hubbard correction term”. LSDA+U also includes the inter-site Coulomb interaction by means of Hartree energy. Therefore, LSDA+U method can explain the physics of any charge ordered system, if Hartree energy is enough to describe them.

For low doping systems (\( x = 0, \frac{1}{2} \)) where the correlation energy induced by the inter-site Coulomb interaction is small, the results of LSDA+U calculations can explain well experimental results. Details of the calculations are as follows. The values of \( U \) are chosen to be 7.5 eV for Ni\(^{2+}\) ions and to be 7.0 eV for Ni\(^{3+}\) ions. The value of \( J \) are chosen to be 0.88 eV. These values are consistent with those of constrained LDA.

In the present system, NiO\(_6\) octahedra are elongated with the direction of \( c \)-axis and tilted. The elongation splits the two \( e_g \)-orbitals by \( \Delta = 0.97 \) eV in LDA calculation, where we denote the energy splitting between \( 3z^2-r^2 \) - 1-orbital and \( x^2-y^2 \)-orbital as \( \Delta \). On the other hand, the tilt of NiO\(_6\) octahedra does not change the electronic structure in both LDA and LSDA+U calculations; the localized magnetic moments on the Ni ion are 1.56 \( \mu_B \) (with tilt) and 1.54 \( \mu_B \) (without tilt), and the energy gap 3.66 eV (with tilt) and 3.73 eV (without tilt). Therefore, the tilt of NiO\(_6\) octahedra is neglected hereafter. We, then, fix the total volume of unit cell and set the \( c/a \)-ratio to be the observed value 3.26.

Crystallographic coordinates of atoms are fixed at those of \( x = 0 \) and, with changing \( x \), positions of all atoms are scaled. The magnetic unit cell at \( x = 0 \) is a \( \sqrt{2} \times \sqrt{2} \times 1 \) supercell, and a \( m \sqrt{2} \times \sqrt{2} \times 1 \) supercell at \( x = \frac{1}{2} \).

At \( x = 0 \), calculated values of the band gap (3.73 eV) and local magnetic moment (1.54 \( \mu_B \)) agree well with the observed values of 4 eV and 1.68 \( \mu_B \) respectively. Figure I shows the projected density of states of LSNO at \( x = \frac{1}{2} \). The position of this \( x^2-y^2 \)-orbital on the Ni\(^{3+}\) site shifts to energy region just above the Fermi energy. The calculated energy gap is 0.10 eV. The resultant spin structure is such that antiferromagnetic domains are separated by hole stripe, and the magnetic moment localized on each site is \(-1.46, -0.94, 1.51, -1.51, 0.93, 1.46 \mu_B\).

FIG. 1: (color online) Projected density of states for each spin (up and down indicated by arrows) of electron in La\(_2\)Sr\(_2\)NiO\(_4\) by LSDA+U method for (a) Ni\(^{2+}\) and (b) Ni\(^{3+}\). Energy zeroth is fixed at the top of the occupied bands. Inset shows the location of each site. Local magnetic moment of each site is \(-1.46, -0.94, 1.51, -1.51, 0.93, 1.46 \mu_B\), respectively from the left to the right.
respectively, from the left to the right. The center of the hole stripe is located on the Ni$^{3+}$ ions and no neighboring Ni$^{3+}$ ion exists. The spins on two Ni$^{2+}$ ions sharing the same neighboring Ni$^{3+}$ ions are anti-parallel to each other (inset of Fig. 1(a)). Introduction of the multiple Slater determinant decreases the spin moments on Ni$^{3+}$ ions, because the spin configuration with the opposite spin direction on Ni$^{3+}$ ions gives the same energy with the present spin configuration. This order is consistent with the experimentally observed stripe and should be assigned to the real ground state. However, the calculated lowest energy state is of no charge order and the energy is lower by 0.5 eV/cell than that of the real ground state in Fig. 1. The spin ordered alignment of this calculated lowest energy state is different from that of the charge ordered state shown in the inset of Fig. 1(a). The local magnetic moment of the calculated lowest energy state is, on each Ni site in Fig. 1(a), 1.48, 1.33, 1.48, $-1.48$, $-1.33$, $-1.48\mu_B$, from the left to the right. There are two problems responsible for the energy increase of the real ground state with the charge order. One is the absence of correlation energy arising from the charge fluctuation between Ni$^{2+}$ and Ni$^{3+}$ configuration. The other is the fact that LSDA+U method often underestimates the correlation energy of antiferromagnetic bonds.

III. ELECTRONIC STRUCTURE OF LSNO ($x = \frac{1}{3}$, 1) BY USING LSDA+U METHOD

LaSrNiO$_4$ ($x = 1$) is observed to be paramagnetic metal, the occupation of the Ni ion distributes homogeneously and each ion is Ni$^{3+}$. On the contrary, calculated ground state is antiferromagnetic metal and the projected density of states is shown in Fig. 2. This contradiction between the observation and the calculation is due to unrealistic stabilization energy of the magnetically ordered state in LSDA+U calculation of large on-site Coulomb interaction $U$, $J$, against the paramagnetic state. If we assume that the ground state is represented by multiple Slater determinants, paramagnetic state would be represented as a linear combination of random spin configurations. Then spin polarization within atomic plane lowers the on-site Coulomb and exchange interaction energy and correlation energy stabilize the paramagnetic metal phase. However, in the LSDA+U method, the ground state wavefunction is represented by a single Slater determinant. Consequently, the paramagnetic state is only possible in the state where all ions have no spin polarization. This increases the total energy much, coupling with large $U$, $J$. A dominant component of the real ground state at $x = 1$ should be a linear combination of Slater determinants which have a single electron on each site and spins are not ordered.

The LSDA+U calculation fails to present the paramagnetic metal phase as the ground state at $x = 1$, but the correlation effects cause correct change of the splitting of two $e_g$ bands. Figure 2 shows that the $3z^2-1$ and $x^2-y^2$ bands locate above and below the Fermi energy, respectively. Comparing Figs. 1 and 2, one can see the change of the hole characters from the case of $x = \frac{1}{3}$ to that of $x = \frac{1}{2}$. In the $x = \frac{1}{3}$ system, the hole is doped in $x^2-y^2$-orbital, while it is doped in $3z^2-1$-orbital in the $x = 1$ system. This change of the hole character agrees with the experimental results described in Sec. I. The origins of this change are the strong on-site Coulomb interaction $U$ between two orbitals and the difference of the dispersion of two orbitals/bands. In the LDA calculation of the system of $x = 0$, two $e_g$ bands cross the Fermi energy. The $3z^2-1$-band is narrow and lying on the energy region $-0.62$ eV$\sim-0.69$ eV, and the $x^2-y^2$-band is wide and lying on the energy region $-0.86$ eV$\sim-2.29$ eV, where the energy zeroth is fixed at the Fermi energy. When small amount of hole is introduced, the hole is doped into $x^2-y^2$-band, because $x^2-y^2$-band is averaged located higher than $3z^2-1$-band. At the hole concentration $x = 1$, orbital polarization is maximized if only one band is occupied and the other is push up over the Fermi energy. Because the lowest energy levels of $e_g$-band are mainly consist of $x^2-y^2$-orbitals, the $x^2-y^2$ orbital is preferable to be occupied. Thus, the hole character changes from $x^2-y^2$-orbital to $3z^2-1$-orbital at $x = 1$. In LSDA+U calculation, however the spin and orbital polarization could not be completely maximized and the system becomes metal. The potential correction in Table I shows this relation between the hole character and the strong on-site Coulomb interaction $U$. Potential corrections are the derivatives of Hubbard correction term of LSDA+U Hamiltonian with respect to each element of local occupation matrix $\left< c_{\alpha}^\dagger c_{\beta} \right>$ on each atom, where $\alpha$ and $\beta$ denote atomic orbitals and $\sigma$ spin. In the present system, potential corrections are diagonal matrices due to symmetry and only diagonal elements are listed in Table I. In such case, each diagonal elements shows the energy level shift of respective orbital induced by on-site Coulomb interaction. Potential correction of
3z^2 − 1 ↑ (x = 1/2) nearly equals to that of x^2 − y^2 ↑ (x = 1) and that of x^2 − y^2 ↑ (x = 1/3) nearly equals to that of 3z^2 − 1 ↑ (x = 1). Therefore, the on-site Coulomb interaction applied to the narrow 3z^2 − 1- and wide x^2 − y^2-band induces the difference of the hole character between the low concentration (x = 1/3) case and the high concentration (x = 1) case.

In high-doped x = 1/2 case, the real system has an incommensurate stripe type charge order, as is described in Sec. I. The ground state of LSDA+U calculation shows, however, no charge order. Consequently, the electronic structure of the calculated system is metallic as shown in Fig. 3.

The charge order of the system of x = 1/2 would be expected to be a commensurate checker-board type order, because of following three reasons. The first is that the periodic boundary condition forces the charge order to be commensurate to the lattice, and once we assume the incommensurability ε to be a rational value 1/2, the resultant charge order is checker-board type. The second is that in the temperature range T_{SO} < T < T_{CO}, the checker-board type charge order is observed in the real system with x = 1/2. The third reason is that the positive value of inter-site Coulomb interaction stabilizes the checker-board type charge order. We must notice that the inter-site Coulomb interaction is included in LSDA+U Hamiltonian by means of Hartree energy.

We could not find a converged charge ordered solution, even as a metastable state, by using LSDA+U calculation. Localized magnetic moment of the four Ni ions in the unit cell equal to 1.29, 1.32, −1.29, −1.32μB, respectively. Therefore, all the Ni ions are of the same type, allowing the difference of local magnetic moment by 3%. Figure 3 shows the partial density of state of each orbitals of Ni ions.

The Hartree energy may be insufficient to stabilizes the charge order. In the real system, the electron localized on the Ni^{2+} site has extending tail on Ni^{3+} site and the center of localized electron can be fluctuating among Ni^{2+} and surrounding Ni^{3+} sites, in order to lower the correlation energy. Because the ground state wavefunction is represented by a single Slater determinant in LSDA+U calculation, the fluctuation of the center of localized electron is neglected, then the correlation energy is underestimated. Moreover, in the Hartree energy, the contribution of the tail to the charge on Ni^{3+} site and the contribution of the head to the charge on Ni^{2+} site are treated as if they were separate electrons. Such treatment gives an additional self-interaction and causes the increase of the estimated energy. Because the number of nearest neighbor (n.n.) pairs of Ni^{2+} and Ni^{3+} ions are maximized in the checker-board type charge ordered system, above two reasons become serious for the increase of estimated energy.

Now the prescription for resolving these problems are follows. We need to preserve the antisymmetry for exchanging of any pair of single electron wavefunctions in order to remove self-interaction and need to employ as many configurations of Slater determinants as possible in order to correctly estimate the correlation energy. Thus we employ the exact diagonalization of many body Hamiltonian in following Secs.

| 3z^2 − 1 ↑ x^2 − y^2 ↑ 3z^2 − 1 ↓ x^2 − y^2 ↓ |
|-------------------------------|
| Ni^{2+} (x = 1/2) | -2.07 | -2.06 | 3.80 | 3.29 |
| Ni^{3+} (x = 1/3) | -1.61 | -0.18 | 2.92 | 2.10 |
| Ni^{3+} (x = 1) | -0.01 | -1.67 | 2.56 | 2.92 |

TABLE I: Diagonal elements of potential correction (in unit of eV). Potential correction is a derivative of Hubbard correction term in LSDA+U Hamiltonian with respect to respective elements of occupation matrix. That gives orbital dependent potential corresponding to fluctuations of orbital occupation.

IV. DOUBLE ORBITAL EXTENDED HUBBARD HAMILTONIAN FOR LSNO (x = 1/2) BASED ON LDA CALCULATION

To understand the ground state at x = 1/2, we adopt the extended Hubbard Hamiltonian ̃H of e_g electrons on two-dimensional (2D) simple square lattice derived from results of LSDA+U calculations, and diagonalize this many body Hamiltonian exactly by using the Lanczos method with the inverse-iteration method;

\[ ̃H = \sum_{i,j,\alpha,\beta,\sigma} t_{i\alpha j\beta} \hat{c}^\dagger_{i\alpha \sigma} \hat{c}_{j\beta \sigma} + \sum_{i,\alpha,\sigma} \varepsilon_{i\alpha} \hat{c}^\dagger_{i\alpha \sigma} \hat{c}_{i\alpha \sigma} + \frac{1}{2} \sum_{i,\alpha,\beta,\gamma,\delta,\sigma,\sigma'} U_{\alpha\beta\gamma\delta} \hat{c}^\dagger_{i\alpha \sigma} \hat{c}^\dagger_{i\beta \delta} \hat{c}_{i\beta \sigma} \hat{c}_{i\gamma \sigma'} + \frac{V}{2} \sum_{(i,j),\alpha,\beta,\sigma,\sigma'} \hat{c}^\dagger_{i\alpha \sigma} \hat{c}_{i\alpha \sigma} \hat{c}^\dagger_{j\beta \sigma} \hat{c}_{j\beta \sigma'}, \]  (1)

where the braces (⋯) denotes the summation over n.n. pairs and the symbol t Slater-Koster type hopping pa-
parameters. The on-site energy $\varepsilon_{\alpha\alpha}$'s are determined by LDA calculation at $x = 0$. $t_{\alpha\beta\sigma}$, $t_{dd\sigma}$, $t_{dd\delta}$ are determined by LDA calculation at $x = 0$. The one-n.n. pairs and $t'_{dx\sigma} + \frac{3}{4}t'_{dx\delta} = -0.018$ eV, $t'_{dx\delta} = -0.023$ eV for the second n.n. pairs and $\Delta = \varepsilon_{x\sigma+1} - \varepsilon_{x\sigma-1} = 0.97$ eV. The energy zero-th is set at the midst of $\varepsilon_{x\sigma-1}$ and $\varepsilon_{x\sigma+1}$. The matrix elements of the intra-atomic Coulomb interactions $U_{\alpha\beta\delta}$ are represented as functions of averaged Coulomb and exchange interaction $U$, $J$, by using the same expressions as in LSDA+U method. The values of $U$, $J$ are chosen to be $7.5$, $0.88$ eV, respectively, on all sites. The value of the inter-site Coulomb interaction $V$ is chosen as $0.5$ eV, except explicit indication of the value of $V$. Later we will explain the reason of this choice, $V = 0.5$ eV, in detail.

Because two diagonal directions are inequivalent in the real LSNO ($x = \frac{3}{4}$), we introduce a parameter $\delta$ showing the anisotropy of the second n.n. (diagonal) hopping parameter $t'$. We add to or subtract from $t'_{\alpha\beta+1,\alpha\beta-1,\alpha\beta+2}$ as $t'_{\alpha\beta+1,\alpha\beta-1,\alpha\beta+2} \pm \delta$, depending on the direction, as depicted in Fig. 4(c). We investigate the $\delta$ dependence in the range of $0 \leq \delta \leq 0.02$ so that the sign of the second n.n. hopping parameter does not change. This anisotropy $\delta$ reserves translational symmetry. As a result, finite value of $\delta$ does not induce charge order, at least in the present range of $\delta$. For example, the difference between the charge-charge correlation of the ground state at $V = 0$ eV under $\delta = 0.02$ and $\delta = 0$ is at most 3%. However, once charge order is induced by other quantities (in the present case, by inter-site Coulomb interaction $V$), finite value of $\delta$ changes essentially the spin order of the system, as is shown in Sec. VII. Finite value of $\delta$ neither changes essentially excitation spectra and energy gap. The width of energy gap reduced by a few percent at $V = 0.5$ eV. Therefore, we fix the value of $\delta$ at $\delta = 0$ in Secs. IV, V and VI. We will change the value of $\delta$ in Sec. VII where the change of spin order induced by $\delta$ is discussed.

We diagonalize the many-body Hamiltonian of a system of twelve electrons on the planar $\sqrt{8} \times \sqrt{8}$ supercell, where each site has two $e_g$-orbitals. A periodic boundary condition is imposed, avoiding the bunching of electron at corners, and causes commensurate checker-board type charge order (SDCCO in short), where closed and open circle denote non-hole and hole Ni site respectively. Another type of distribution is obtained by exchanging the hole site and non-hole site. (c) The occupation distribution of SDCCO in (b). A number in the open circle shows the occupation number of respective site.

FIG. 5: (color online) $V$ dependence of total energies and charge-charge correlations at $x = \frac{3}{4}$ and $\delta = 0$, where $V$ is inter-site Coulomb interaction. The state labeled “S” is singly degenerate while the other state labeled “D” is doubly degenerate. Energy-zero-th is set at the ground state energy. State S is a ground state on the region $0 \leq V < 0.41$ eV, State D is a ground state on the region $0.41$ eV $< V < 1$ eV. Suffix 1, 2, 3, 7 denotes following sites: origin, nearest neighbor, second nearest neighbor and third nearest neighbor, respectively. These site indexes are depicted in Fig. 5(a).

FIG. 6: (color online) (a) Site index of the present system at $x = \frac{3}{4}$ in a $\sqrt{8} \times \sqrt{8}$ cell. (b) A type of hole distribution of the Slater determinants with the complete checker-board type charge order (SDCCO in short), where closed and open circle denote non-hole and hole Ni site respectively. Another type of distribution is obtained by exchanging the hole site and non-hole site. (c) The occupation distribution of SDCCO in (b). A number in the open circle shows the occupation number of respective site.

V. CHARGE ORDER OF LSNO ($x = \frac{3}{4}$)

At first, we discuss the ground state of LSNO at $x = \frac{3}{4}$. With increasing $V$, the ground state changes from singly degenerate state (State S) in the region $0 \leq V < 0.41$ eV to doubly degenerate (State D) $0.41$ eV $< V < 1$ eV. Smoothness of the connection among respective series of States S and D with respect to $V$ is discussed in Appendix A.

Figure 5 shows the $V$ dependence of the total energies and the correlation functions of charge fluctuation $\langle \delta n_i \delta n_j \rangle$, where $\delta n_i = n_i - \langle n_i \rangle$ and $\delta$ is fixed at 0. At $V = 0.5$ eV, charge correlations $\langle \delta n_i \delta n_j \rangle$ of State D equal to $-0.135$, $0.093$, $0.096$ for n.n. $(i,j) = (1,2)$, second
FIG. 7: (color online) $V$ dependence of the weight of Slater determinants with complete checker-board type charge order (SDCCO in short), where $V$ is inter-site Coulomb interaction. The state labeled “S” and “D” are the same states as in Fig. 5. n.n. $(i, j) = (1, 3)$ and third n.n. $(i, j) = (1, 7)$ pairs respectively, and this corresponds to checker-board type charge order shown in Fig. 5(b). The checker-board type charge order in State D exists even at $V = 0$ eV, though the amplitude is very small. The charge correlations for n.n., second n.n. and third n.n. pairs at $V = 0$ eV equal to $-0.074, 0.007, 0.019$, respectively. These values satisfy the checker-board type charge order: The positive correlation between second n.n. pair nearly equals to that between third n.n pair, and the correlation between the first n.n. pair is negative. The charge correlations in State S are homogeneous in comparison with those in State D. The charge correlation in State S at $V = 0$ eV of first, second and third n.n. pairs equal to $-0.046, -0.013, -0.056$, respectively. Therefore, State S at $V = 0$ eV does not have any charge order. The absence of charge order in State S in $0 \text{ eV} < V < 0.4$ eV is attributed to a slow increase of charge correlations with increasing $V$ in this region of $V$.

Now we discuss three features in Fig. 5, two are about correlation functions of State S and one about the energy difference between States S and D: (i) Rapid increase of the charge correlation of State S in the region $0.4 \text{ eV} < V < 0.6$ eV, (ii) merging charge correlation of State S into that of State D, in the region $V > 0.6$ eV and (iii) the constant energy difference between States S and D in the region $V > 0.6$ eV. All these features are related to respective wavefunctions of States S and D.

Let us define the ideal complete checker-board type charge order whose Slater determinant wavefunction (SDCCO) should satisfy the following two conditions, (a) and (b), then we specify the characteristics of real wavefunctions of States S and D by using the overlap with SDCCO’s: (a) every site has occupation one or two, and (b) all neighboring sites of a singly occupied site are doubly occupied sites and vice versa. A type of their distribution of occupation is shown in Fig. 5(c). Assuming both transfer integral and on-site exchange parameter $J$ equal to zero, SDCCO’s would be the ground states, where all the spin configurations are degenerate, because the numbers of neighboring hole $(\text{Ni}^{3+})$ and non-hole $(\text{Ni}^{2+})$ pairs are maximized. In an actual calculation, transfer integrals are finite and the SDCCO’s are not eigenstate. The value of $V$ determines how much SDCCO’s are hybridized into respective eigenstates. The more $V$ increase, the more hybridization of SDCCO’s are preferable energetically.

In the region of $0 \text{ eV} < V < 0.4$ eV, the ground state is homogeneously extending State S since the kinetic energy is a source of gain of the total energy. On the other hand, in the region of $0.4 \text{ eV} < V$, the ground state changes to the charge ordered State D since the source of energy gain is the correlation energy due to the inter-site Coulomb interaction $V$. The dependence of the total weight of SDCCO’s is shown in Fig. 7. Comparing Figs. 5 and 7, the above features (i), (ii) and (iii) appear as follows. Since the wavefunction of the ground state State S is homogeneous, the overlap with SDCCO is small in the region of $0 \text{ eV} < V < 0.4$ eV. We discuss more detail about the weight of SDCCO’s of State S at $V = 0$ eV in Appendix B Then, in the range $0.4 \text{ eV} < V < 0.6$ eV, the weight of SDCCO’s rapidly increases. Because both States S and D are eigenstates of $\hat{H}$ and orthogonal with each other, the coefficients of SDCCO’s in State S are different from those in State D. In the range $0.4 \text{ eV} < V < 1$ eV, States S and D have the same charge order but the spin order is quite different, which is discussed in Sec. VI Then, the difference of the total energies between States S and D is attributed to the difference of spin configurations. As is discussed in Sec. III $T_{SO} < T_{CO}$ in the real system with $x = \frac{1}{2}$ and, therefore, the energy scale of spin order is smaller than that of charge order. In fact, we can see a large energy difference between States S and D in the region of $0 \text{ eV} < V < 0.4$ eV, since the charge order different between States S and D here. Then, the characteristics of the charge order are the same in States S and D in the range $0.6 \text{eV} < V < 1$ eV (Feature (ii)), as seen in the charge correlation functions and the same weights of SDCCO’s in States S and D, and the constant energy difference between States S and D (Feature (iii)). The observed high temperature state ($T > T_{SO}$) in LSNO of $x = \frac{1}{2}$ might be a mixture of charge ordered eigenstates with significant weight of SDCCO’s, including the doubly degenerate States D and the singly degenerate State S.

As mentioned in Sec. III, the real ground state of LSNO ($x = \frac{1}{2}$) shows charge stripe order, incommensurate to the lattice. And as mentioned in Sec. III, the periodic boundary condition forces the charge order to be commensurate charge order of checker-board type. Therefore, the ground state of the Hamiltonian should have checker-board type charge order, and we choose $V = 0.5$ eV from now on. The value $V = 0.5$ eV is not very unrealistic because a recent report shows $V = 0.34$ eV in GW approximation of LaMnO$_3$.

VI. EXCITATION SPECTRA AND ENERGY GAP OF LSNO ($x = \frac{1}{2}$)

We show calculated single-particle spectral functions of the system with $x = \frac{1}{2}$ in Fig. 6 in order to see whether the spectra have energy gap and the system is insulator as the observed case. Inter-site Coulomb interaction is fixed at $V = 0.5$ eV as discussed in Sec. VI. Three $k$-points are chosen: $k = (0, 0.5), (0.25, 0.5)$ and $(0.25, 0.0)$, where $a$ is the nearest Ni-Ni distance, the positions of...
The ionization level (HIL) is located at 9 eV and the affinity level (LAL) at 10 eV. Three panels are corresponding to respective k-points, \((\frac{\pi}{a}, 0)\), \((\frac{\pi}{a}, \frac{\pi}{a})\), and \((0, 0)\), from the top to the bottom. Each value of k corresponds to different boundary condition. Two arrows near 10 eV shows the position of the highest ionization level 9.4 eV and the lowest affinity level 10.3 eV in case of \(k = (0, 0)\). Spectral functions are approximated by the polynomial of 800 degree in \(H\), where \(\hat{H}\) denotes the Hamiltonian. A small imaginary part (0.01 eV) is added to the energy \(\omega\) for smearing \(\delta\)-function peaks.

Ni ions are \((n, m)a\), and the two translation vectors are \((-2, 2)a\) and \((2, 2)a\). Each k-point corresponds to periodic or antiperiodic boundary condition along respective translation vector. In order to obtain continuous spectra, we introduce an imaginary energy \(\eta\) of 0.01 eV and the value of \(\eta\) determines the resolution of the spectra.

The value of \(\eta\) must satisfies \(\eta \gtrsim \frac{\text{total width of spectra}}{m}\), where \(m\) denotes the the highest power of \(H\), because \(m\) is the upper limit of the number of \(\delta\)-functions in the spectra. In the Lanczos method, the truncation error easily breaks down the orthogonality of Lanczos vectors. Because calculating spectral function may need the large \(m\), we employ the shifted-COCG method which is a variant of the conjugate gradient (CG) method. It must be noticed that the shifted-COCG method is numerically stable. Because shifted-COCG method requires an reference energy in the energy region of a peak of spectral weight, we need to know rough profile of spectral weight. Therefore, the shifted-COCG method should be used only when we need very fine profile of spectral function. More details will be explained elsewhere.

In the present case, the “total width of spectra” is roughly estimated as \(E(Ni^{2+}) - E(Ni^{3+}) \sim U = 7.5\text{eV}\) for ionization levels, and affinity levels, separately. Then we choose \(m = 800\). We calculate the spectral function of \(m = 160\) by Lanczos method, and then, calculate that of \(m = 800\) by the shifted-COCG method. Actually, the width of spectra is wider than \(U\) due to the mixing of higher/lower occupation configurations. The spiky structure of spectra is an artifact due to the choice of smaller \(\eta = 0.01\text{eV}\). Our choice of the value of \(\eta\) is intending to show the gap structure between ionization levels and affinity levels, at every k-point. We fix the value of hopping anisotropy \(\delta = 0\). The hopping anisotropy \(\delta\) does not change the shape of the spectral function in the range of \(0 \leq \delta < 0.02\). The width of the gap at \(k = (0, 0)\) in Fig. 8 is smallest among three k-points, where the highest ionization level (HIL) is located at 9.4 eV and the lowest affinity level (LAL) 10.3 eV. Therefore, we conclude that this system is insulator with an energy gap of 0.9 eV.

It is important to know the symmetry of the single electron wavefunctions of HIL and LAL, in order to show that the “gap” is not an artifact of discretized k-point.

| \(E_b, E_t\) (eV) | Peak area | The largest eigenvalue and its degeneracy | Symmetry of eigenvectors \(k(\frac{\pi}{a})\) |
|-----------------|-----------|--------------------------------------|----------------------------------|
| \(HIL\) at \(k = (0, 0)\) | 9.15 9.55 | 0.37 0.09 \(\times 4\) | \((\frac{\pi}{a}, \frac{\pi}{a})\) |
| \(LAL\) at \(k = (0, 0)\) | 10.05 11.14 | 2.87 0.41 \(\times 4\) | \((\frac{\pi}{a}, \frac{\pi}{a})\) |
| \(HIL\) at \(k = (\frac{\pi}{a}, \frac{\pi}{a})\) | 8.18 8.98 | 1.72 0.63 \(\times 2\) | \((\frac{\pi}{a}, \frac{\pi}{a})\) |
| \(LAL\) at \(k = (\frac{\pi}{a}, \frac{\pi}{a})\) | 10.02 11.02 | 2.99 0.44 \(\times 4\) | \((\frac{\pi}{a}, \frac{\pi}{a})\) |
| \(HIL\) at \(k = (\frac{\pi}{a}, \frac{\pi}{a})\) | 8.18 9.18 | 1.88 0.38 \(\times 4\) | \((\frac{\pi}{a}, 0)\) |
| \(LAL\) at \(k = (\frac{\pi}{a}, \frac{\pi}{a})\) | 10.12 11.12 | 2.90 0.42 \(\times 4\) | \((\frac{\pi}{a}, \pi)\) |

TABLE II: Bottom and top of the energy of peak-like structure, area of the structure, the largest eigenvalue (and its degeneracy after “x” symbol) related to “single electron wavefunction” (see text) and the representative k which shows the symmetry of the single electron wavefunction. In the right column, single \(\hat{k}\) is written and the other equivalent \(\hat{k}\) are omitted. HIL and LAL stands for highest ionization level (HIL) and lowest affinity level (LAL) respectively. Each k-points corresponding to the boundary conditions are written in the leading lines. \(\delta = 0\).

The symmetry of respective single electron wavefunction can be labeled by crystal momentum \(k\). If the “gap” is an artifact and the ground state is metal, HIL and LAL are labeled by different \(k\). Actually many levels overlap and form a single peak-like structure because of complicated interactions, especially in the energy range near LAL. Even the definition of the “single electron wavefunction” are not clear in such a circumstance. Therefore, we define a “single electron wavefunction” as follows, here. First we choose the bottom \(E_b\) and top \(E_t\) of a single peak-like structure. Next we integrate the Green function of matrix form in the energy range \(E_b < E < E_t\):

\[
-\frac{1}{\pi} \int_{E_b}^{E_t} \text{Im}G_{\text{vac}}^{R}(\omega) d\omega.
\]

and then we diagonalize it. This matrix corresponds to energetically partitioned \(\langle \hat{c}^\dagger_{i\alpha} \hat{c}_{j\beta}\rangle\) \((E_b < E_t < E_F)\) or \(\langle \hat{c}^\dagger_{i\alpha} \hat{c}_{j\beta}\rangle\) \((E_F < E_b < E_t)\). Therefore, the eigenvector of above matrix corresponds to the single electron wavefunction related to the peak-like structure and the eigenvalue corresponds to the occupation number of the single electron wavefunction. We summarize in Table II the \(E_b, E_t\), eigenvalues and the \(k\) representing the symmetry of eigenvectors for the peak-like structure near HIL and LAL. At \(k = (0, 0)\), HIL and LAL single electron wavefunctions share the same symmetry. This can not occur in the metallic system. Therefore, we can conclude that the split between HIL and LAL is not an artifact and the system is an insulator.

Now we show that the inter-site Coulomb interaction \(V\) induces the energy gap between occupied- and
unoccupied-states, not on-site Coulomb interaction $U$. Figure 9 (a), (b) shows the “spectral function” with respect to State D with $V = 0.5$, 0.2 and 0 eV. Note that, since State D is not the ground state at $V = 0.2$ eV and $V = 0$ eV, they (for $V = 0.2$ eV and $V = 0.5$ eV cases) are not satisfying the definition of spectral function. These spectral functions are approximated by rational function of the degree of $\hat{H}^2$ by using Lanczos method. The top panel of Fig. 9 (a) and the bottom panel of Fig. 9 are the same, except that the latter uses higher degree of $\hat{H}^2$ and the shifted-COCG. They are sharing characteristic peaks of HIL and LAL. Therefore, 160 degree is enough to discuss the characteristics of spectra. At $V = 0.5$ eV (top panel), HIL and LAL are $(x^2 - y^2)$- and $(3z^2 - 1)$-orbitals, respectively, and this agrees well with experimental observation and the spectrum in Fig. 9. At $V = 0.2$ eV (middle panel), HIL lowers to 7.9 eV and the energy gap becomes smaller than that at $V = 0.5$ eV. At $V = 0$ eV (bottom panel), HIL lowers to 6.9 eV and two excitation peaks overlap with each other in HIL and LAL, i.e., the energy gap vanishes. Thus we can conclude that the system with $x = \frac{1}{2}$ becomes insulator due to $V$, not $U$, and the inter-site Coulomb interaction $V$ causes the energy stabilization and the opening the gap of the ground state State D. The gap width in the bulk limit is discussed in Appendix C.

VII. SPIN ORDER OF LSN0 ($x = \frac{1}{2}$)

We discuss the spin stripe order induced by the anisotropy of hopping integral between the n.n. sites. Introduction of the anisotropy does not contradict to the local symmetry of observed charge and spin stripe. This anisotropy with $\delta$ may couple with orthorhombic lattice distortion of $T_{2g}$ symmetry. Coupling to the distortion reminds us of JT mechanism. However, the $T_{2g}$ distortion is not a JT mode for the $e_g$ electron, and therefore, this mechanism is not of JT mechanism.

The spin correlation function $\langle S_i \cdot S_j \rangle$ at $\delta = 0$ equals to $-0.37$, 0.12, −0.07 respectively when $V = 0.5$ eV and shows a lack of long range order. Equivalence of two diagonal directions is an origin of the two-fold degeneracy, as is shown in the result of $k = (\frac{2\pi}{a}, \frac{2\pi}{b})$ in Sec. VI.

Because large amount of holes are doped in $x^2 - y^2$-orbital, the electron in $x^2 - y^2$-orbital on Ni$^{3+}$ can have a long tail extending to Ni$^{4+}$ site. The hopping part of the total energy per electron is $-0.30$ eV and its absolute value is as twice large as that in the parent material ($x = 0$, $N = 16$), $-0.15$ eV. Following the localized electron picture, increase of hopping energy indicates larger hybridization of $x^2 - y^2$-orbital between Ni$^{2+}$ and Ni$^{3+}$. Then, spin on Ni$^{2+}$ site has strong correlation with that on Ni$^{3+}$ site. Thus the spin correlation in $x = \frac{1}{2}$ system is more complicated than that in $x = 0$ system where the localized spin picture is applicable.

The present parameter set in $\hat{H}$ is another origin of the complexity of spin structure. If $J \gg \Delta$, the majority spin is preferable in order to lower the exchange energy when the electron visited the Ni$^{3+}$ site. In contrast, if $\Delta \gg J$, the minority spin is preferable in order to lower on-site energy. The present values of $J$ and $\Delta$ satisfy neither of two conditions. In addition to that, transfer integral and the inter-site Coulomb interaction $V$ has the same energy scale. This causes the competition between the extended state and the localized single electron wavefunction. All these competitions may cause the instability against the perturbation.

In the case of $\delta \neq 0$, the ground state energy decreases with increasing $\delta$ and can be expressed as $(36.75 - 104.35\delta^2)$ eV. The spin correlation function is depicted in Fig. 10. The value of the correlation function between nearest neighbors $\langle S_1 \cdot S_2 \rangle = -0.37 \sim -0.38$, does not change. The local spin moment is $\langle S_i \rangle = 1.31$. If two electrons localized on one site in a triplet state, four ele-
trons are localized on the four n.n. sites, and the total spin $S$ of this totally six electron system equals to 1, then $\langle S_1 \cdot S_2 \rangle = -0.375$, which is almost the same as the calculated value of $\langle S_1 \cdot S_2 \rangle$. In the range of $\delta \gtrsim 0.01$ eV, correlation functions of the two types of second neighbor pairs have opposite signs and the structure of the calculated spin stripe order is consistent with the observed one. This critical value $\delta = 0.01$ eV is less than 2% of the absolute value of the n.n. hopping $t_{\text{dd}}$, and thus small anisotropy changes the spin structure drastically.

The spin order is quite different between systems of the single-orbital and of the multi-orbitals. The former on 2D square lattice is the Néel order on $\sqrt{2} \times \sqrt{2}$ cell, and the latter is the charge and spin stripe. Once we assume only $x^2 - y^2$-orbital at each site and put 8 electrons on 16 sites, we get the value of $\langle S_1 \cdot S_2 \rangle$ equal to $-0.03$ which is very small compared to the value $-0.38$ in LSNO. This reduction of $\langle S_1 \cdot S_2 \rangle$ in the single-orbital system is attributed to the absence of localized spin on a hole site and consistent with the Néel order mentioned above. However, a spin remains on a hole site in LSNO ($x = \frac{1}{2}$). Since two neighboring sites of a hole site have spins of opposite direction with each other, the spin state of a hole site is a linear combination of up- and down-spin due to symmetry. Because no term in the Hamiltonian flips a spin on a hole site, two antiparallel-spin electrons on two hole sites must exchange their positions through hopping process, in order to create such a linear combination. This hopping process lowers correlation energy and the ground state must be represented by the multiple Slater determinant. In all hopping processes to create the multiple Slater determinants, that of exchanging a hole and a non-hole site is most probable, due to Large $U$, and the number of these pairs of sites is maximized at $x = \frac{1}{2}$. This causes insufficient description of the ground state by single determinant in LSNO at $x = \frac{1}{2}$.

Finally, we stress that the above stripe spin correlation induced by the anisotropy of the transfer integrals is only seen under the checker-board type charge order. When the anisotropy $\delta = 0.02$ eV is introduced to State S (see Sec. V) in the range $0$ eV $\leq V \leq 0.4$ eV, the resultant state shows the spin order of Néel order type, which is also seen in $\delta = 0$ case in the range $0$ eV $\leq V \leq 0.4$ eV. The Néel order of half-filled $3z^2 - 1$-orbitals appears under the homogeneous charge distribution and the small $\delta$ does not affect the spin order. Under the charge ordered condition, the single electron wavefunction is rather localized, if not site-localized, then the transfer integrals are reduced effectively. Then the small $\delta$ changes the spin order drastically into the spin stripe.

VIII. CONCLUSION

We discussed the charge and spin order of LSNO ($x = \frac{1}{2}$) by using LSDA+U method and those of LSNO ($x = \frac{1}{2}$) by using exact diagonalization of the double orbital extended Hubbard model derived from LDA calculations. In the exact diagonalization, charge and spin order is discussed by using charge-charge or spin-spin correlation functions. Excitation spectra of LSNO ($x = \frac{1}{2}$) are also calculated to show that the system is insulator.

In conclusion, LSNO with $x = \frac{1}{2}$ and $\frac{3}{2}$ are both insulator with charge and spin stripe order. In both systems, diagonal hole stripes are separately located on Ni$^{3+}$ site in order to reduce hole-hole interaction energy induced by inter-site Coulomb interaction $V$. We discussed the important role of multi-orbitals and mixing of multiple Slater determinants especially in high-doped $x = \frac{1}{2}$ system, where the charge order is induced by the correlation energy of the inter-site Coulomb interaction $V$. Charge order and the inter-site Coulomb interaction $V$ are directly related to the energy gap in the excitation spectra of the system with $x = \frac{1}{2}$. Spin stripe occurs only under the condition of the existence of the charge order, with a help of anisotropy $\delta$ in diagonal hopping. Thus the spin stripe is determined by the electronic structure with smaller energy scale than that of the charge stripe. This is consistent with the observation of $T_{\text{CO}} > T_{\text{SO}}$. Though the spin stripe is related to anisotropy, the mechanism is not of Jahn-Teller type unlike the usual ordering in transition metal oxides, because anisotropy in LSNO does not couple to the Jahn-Teller mode.

The stability of LNSO in different hole concentration $x$ depends sensitively upon several physical quantities, e.g. $t_{\alpha \beta} \Delta$, $U$, $J$, $V$ and $\delta$, and we believe that we have successfully shown the new scope of the combination with the first principles electronic structure calculations and the many-electron theory. We have developed a very useful novel tool for the extremely large matrix of extended Hubbard Hamiltonian, the shifted-COCG method, which will be explain more details elsewhere.

Lastly we comment on layered cuprates. The band gap of La$_2$CuO$_4$ (LCO) is 2 eV, narrower than La$_2$NiO$_4$, and hole doping makes systems metallic. Due to these facts, the screened Coulomb interaction in LCO becomes smaller than in LSNO and the energy gain by hole hopping is more important in doped cuprates. This may be one of reasons why the hole stripe in La$_{1.46}$Nd$_0.4$Sr$_{0.12}$CuO$_4$ runs in a direction along the n.n. pair and the hole concentration is one per two Cu sites in the stripe.

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APPENDIX A: METHOD TO GET SMOOTH EIGENVECTORS WITH RESPECT TO A PARAMETER IN HAMILTONIAN

We use Lanczos method and the inverse-iteration method with CG method alternatively in order to obtain the exact eigenvectors. These methods work well for getting ground states, but careful treatment must be taken for solving two smooth eigenvectors with respect to $V$ in Sec. V. Because both methods are the energy minimization process, iterative application of them, with no care, gives the eigenvector of lower eigen energy than that of the aiming eigen energy smoothly connected from the solved eigen energy at adjacent values of $V$, though one chooses the solved eigenvector at adjacent value of $V$ as a starting vector. When the two levels are nearly degenerate, this problem becomes seriously important. In the present case, at least three energy levels cross with changing $V$ from 0 eV to 0.5 eV, because at $V = 0.5$ eV, the lowest three levels are 36.7549, 36.7551, 36.7773 in the unit of eV, (the first one is a level of State S and all the three levels are doubly degenerate). State S locates at the 36.7939 eV at $V = 0.5$ eV, higher than these three levels, while State S is the ground state at $V = 0$ eV.

The first technique to avoid this problem is to use small number of the dimension of the submatrix tridiagonalized by Lanczos method, since Lanczos method has stronger tendency to reduce trial eigenvalue than that of inverse-iteration with CG method. We tried successfully the dimension in the range $0 \sim 80$ depending on the case. The second technique is to control the trial eigenvalue so that the residual is minimized and the trial eigenvalue does not jump, in the whole processes. Here, the residual is defined as $|| (H - (trial\ energy)) / (trial\ vector) ||$. Because increase of the residual is a sign of the transition into lower level, once it occurs, we dispose a new (and possibly lower) trial eigenvalue.

Lastly, the inner product among resultant eigenvectors of respective values of the specific parameter ($V$ in the present case) must be checked. The large value of inner product assures the smoothness of connection to the resultant eigenvectors. In the present case, the inner product between State S at $V = 0$ eV and that at $V = 0.5$ eV equals to 0.54 and the inner product between State S at $V = 0$ eV and that at $V = 0.5$ eV equals to 0.73. These values are large enough to assure the smoothness of these eigenvectors, because no other eigenvectors at the end point $V = 0.5$ eV can have larger inner product than above respective states.

APPENDIX B: WEIGHT OF THE SLATER DETERMINANTS WITH COMPLETE CHARGE ORDERED $V = 0$

One might have a question why State S has a finite weight 0.045 of SDCCO’s at $V = 0$, though the state has no charge order. We conclude that the SDCCO’s are not overweighted by following discussion. Because the energy scale of the charge order is larger than that of spin order in the present system, we neglect the spin configuration in each Slater determinant, and count only the charge degrees of freedom. Strong $U$ inhibits the configurations with site occupation more than two and zero and there exist $sC_4 = 70$ types of the charge configurations, where the site occupations are restricted to be one or two. A number of charge configurations with complete checker-board type order equals to two. Assuming that all configurations have the same weight due to the absence of charge order, resultant weight of SDCCO’s equals to $\frac{2}{10} \sim 0.29$, which is consistent with above 0.045.

APPENDIX C: GAP WIDTH IN THE BULK LIMIT

First we discuss the value of the energy gap 0.9 eV in two way, applying two extreme approximations, and show how inter-site Coulomb interaction opens the energy gap. One of two approximations is such that HIL and LAL are approximated by single particle excitation related to an extended hole/electron, and the other is such that they are approximated by a single particle excitation of a site localized hole/electron. In the actual picture, excitations are described by dressed quasi-particle/hole and not by a single particle/hole, as is discussed later in the present section.

We begin with the case that HIL/LAL is approximated by single particle excitation of an extended state. Due to large splitting $\Delta = 0.97$ eV between $3z^2 - 1$- and $x^2 - y^2$-orbital, hole is doped into $x^2 - y^2$-orbital. Therefore, low energy excitation mostly consists of $x^2 - y^2$-orbital and we neglect $3z^2 - 1$-orbital in the present paragraph. The single electron wave functions of HIL and LAL belong to the same symmetry group labeled $k = (\pm \frac{x}{2}, \pm \frac{y}{2})$ as in Table III. By unitary transformation, four plain waves labeled $k = (\pm \frac{x}{2}, \pm \frac{y}{2})$ change the forms into four checker-board type wavefunctions $\cos(\pi x) \cos(\pi y), \sin(\pi x) \sin(\pi y), \cos(\pi x) \sin(\pi y), \sin(\pi x) \cos(\pi y)$. These four wavefunctions give the two types of charge distribution. One type of them is shown in Fig. 6 (b) and the other is obtained with exchanging hole and non-hole sites. Two wavefunctions $\cos(\pi x) \cos(\pi y)$ and $\sin(\pi x) \sin(\pi y)$ do not interact with on-site Coulomb interaction $U$, because $(\cos(\pi x) \cos^2(\pi y)) (\sin^2(\pi x) \sin^2(\pi y)) = \sin^2(\pi x) \sin^2(\pi y) = 0$ on each lattice point. Instead, the interaction between them with inter-site Coulomb interaction $V$ is roughly estimated as $\frac{1}{n} \times \frac{1}{n} \times 4V \times n = \frac{4V}{n}$, where $n$ is the number of sites whose amplitude is finite $= 4$ in the present case. This is occupied- and unoccupied-splitting. Assuming $\cos(\pi x) \cos(\pi y)$ is occupied, then $\sin(\pi x) \sin(\pi y)$ is unoccupied and vice versa. The same situation occurs for another pair of wavefunctions.
cos(\(\frac{x}{2}\))\sin(\(\frac{y}{2}\)) and \(\sin(\frac{x}{2})\cos(\frac{y}{2})\). Thus, the system becomes insulator with gap roughly estimated as \(\Delta V = 0.5\) eV, except \(V = 0\) eV where all the occupied- and unoccupied- states are degenerate (See Fig.2(b)). Though the correlation induced by inter-site Coulomb interaction causes the splitting between occupied- and unoccupied- states (the energy gap), this mechanism is different from the normal Mott insulator, induced by on-site Coulomb interaction \(U_{\text{site}}\). In addition to that, there is another difference between the current mechanism and that of Mott insulator; the wavefunction of HIL and LAL have checker-board type charge order, not sharing the center of charge distribution in the current mechanism, unlike they shares the center of charge distribution in Mott insulator. However, there is a problem with this estimation of the gap. Following the present estimation, energy gap vanishes in the bulk limit, because the gap is inversely proportional to the system size.

In the next estimation, HIL/LAL excitation is approximated by a single particle excitation of site localized hole/electron. Neglecting hopping integrals, the ground state of the present Hamiltonian consists of four \(\text{Ni}^{2+}\) (spin triplet, \(u^1\) and \(v^1\)) and four doublet \(\text{Ni}^{3+}\) (spin doublet, \(u^1\) located on respective sublattice, where \(u\) and \(v\) denote \(3z^2 - 1\) and \(x^2 - y^2\)-orbital respectively. All these components are assigned to the Slater determinants with complete checker-board type charge order (SDCCO’s) in Sec. \(\Box\). Then the LAL is \(x^2 - y^2\)-orbital of \(\text{Ni}^{3+}\) and the HIL is \(x^2 - y^2\)-orbital of \(\text{Ni}^{2+}\) (both have a majority spin). Consequently, the energy gap is estimated as \(\{E(\text{Ni}^{2+}) + 8V - [E(\text{Ni}^{3+}) + 4V]\} - \{E(\text{Ni}^{2+}) - E(\text{Ni}^{3+})\} = 4V = 0.5\) eV, where \(E(\cdot)\) denote the ground state energy of Ni ion for each ionization state. This is an exact solution even in the bulk limit.

The result of the present calculation is in the intermediate region of above two estimations; the calculated energy gap \(0.9\) eV is greater than \(0.5\) eV in extreme cases by the extended HIL/LAL approximation and less than \(2.0\) eV by the site localized HIL/LAL approximation. Therefore, in the calculated excitations (with respect to State D at \(V = 0.5\) eV in Sec. \(\Box\)) are well described by the wavefunction in intermediate region between site localized and extended states. And the split between occupied- and unoccupied- states occurs due to long-ranged (inter-site) Coulomb interaction, though they are not sharing the center and does not have large overlap.

Finally, we discuss on the bulk limit of the energy gap in the present calculation. Taking the limit such that transfer integrals go to infinity, the system becomes paramagnetic metal. Taking the limit such that transfer integrals go to zero the system becomes antiferromagnetic insulator. Therefore, there exists a critical transfer integral where the bulk limit of the system changes from insulator (metal) to metal (insulator). The high weight of the Slater determinant with complete checker-board type charge order (> 0.4 at \(V = 0.5\) eV, as in Fig.7) strongly suggests that the present choice of transfer integrals makes the bulk limit insulator, and that is consistent with the real LSNO (\(x = \frac{1}{2}\)).

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