Photoemission and x-ray absorption study of MgC$_{1-x}$Ni$_3$

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We investigated electronic structure of MgC$_{1-x}$Ni$_3$ with photoemission and x-ray absorption spectroscopy. Both results show that overall band structure is in reasonable agreement with band structure calculations including the existence of von Hove singularity (vHs) near $E_F$. However, we find that the sharp vHs peak theoretically predicted near the $E_F$ is substantially suppressed. As for the Ni core level and absorption spectrum, there exist the satellites of Ni 2$p$ which have a little larger energy separation and reduced intensity compared to the case of Ni-metal. These facts indicate that correlation effects among Ni 3$d$ electrons may be important to understand various physical properties.

I. INTRODUCTION

Interplay between superconductivity and magnetism is a source of rich solid state physics. In particular, much attention is being paid to the superconductors in close vicinity with ferromagnetism (FM). In Sr$_2$RuO$_4$, the p-wave pairing symmetry is thought to be intimately related with the FM fluctuation of Ru ions. In UGe$_2$ and ZrZn$_2$, the two phases were found to coexist. While it was generally believed that FM and SC are mutually exclusive, these observations suggest that the current understanding needs to be modified.

Recently, following the discovery of superconductivity in MgB$_2$, intermetallic compound MgC$_{1-x}$Ni$_3$ ($x = 0.04 \sim 0.1$) was reported to be superconducting at $T_c \sim 7K$. MgCNi$_3$ has a cubic antiperovskite structure where C is surrounded by six Ni ions to form an octahedral cage. Although the material is paramagnetic at $T > T_c$, Ni ions exhibit ferromagnetic (FM) spin fluctuation as shown by $^{13}$C NMR experiment. At $T < T_c$, tunneling data show the zero conductance peak (ZCP) which may suggest a non-s-wave superconducting gap. These facts seem to indicate that the SC is closely linked with FM in this system, as also discussed theoretically by Rosner et al. On the other hand, the result from specific heat experiment is consistent with conventional BCS behavior. Also, Shim et al. showed that the observed $T_c$ is reasonably explained within the BCS electron-phonon interaction mechanism. Thus, the origin of SC in MgCNi$_3$ is still controversial.

Studying electronic structure of this material, particularly near the Fermi energy, is essential to understand the SC. Band calculations show that the Ni-3$d$ state is hybridized with C 2$p$, which dominate density of state (DOS) at $E_F$. In particular, it is predicted that a van Hove singularity (vHs) peak exist very close to the $E_F$. The vHs peaks give rise to a large DOS at $E_F$ which can be directly related with the superconductive coupling constant. It is thus important to experimentally probe the vHs peak in detail. Photoemission spectroscopy (PES) is a powerful tool to investigate energy-band structures and electron correlation effects. In this work, we performed PES and x-ray absorption spectroscopy (XAS) measurements of MgCNi$_3$. We find that overall electronic structure is in general agreement with band calculation results. The vHs peak is also identified at about 100 meV below $E_F$. However, its intensity is much weaker than the theoretical predictions. We discuss several possibilities of the suppression and its implication on the SC.

II. EXPERIMENT

Polycrystalline sample used in this experiment was synthesized by the powder metallurgical technique. Powders were mixed in nominal composition of Mg$_{1.25}$C$_{1.45}$Ni$_3$. Here, excess Mg and C were added to maximize carbon incorporation in MgC$_{1-x}$Ni$_3$, similar to the previous report. The powders were pelletized, wrapped in a Ta foil, and then quartz-sealed under vacuum. The sample was reacted for about two hour at 900 °C and quenched. X-ray diffraction (Rigaku RINT d-max) showed that the sample is in a single phase. Small amount of MgO impurity and some unreacted carbon were also identified. Magnetic susceptibility was measured with a dc SQUID magnetometer (Quantum Design). As shown in Figure 1, the superconducting onset temperature of this as-grown sample was 6.8 K and the transition width measured from 10 \sim 90 % transition was about 0.3 K. These results were close to the previous reports of MgC$_{1-x}$Ni$_3$. Afterward we put the sample in a high pressure cell and annealed under 3GPa at 900 °C for 30 minutes to make densified sample for PES and XAS measurements. After this treatment, the pellet density increased significantly and was almost identical to its theoretical value. Figure 1 shows that $T_c$ also increased by about 0.7K and the transition became sharper.
PES experiments were performed using both He I \((h\nu = 21.2\text{ eV})\) source at home laboratory and 120 eV photon at Pohang Light Source (PLS) in Korea. Mg \(K_{\alpha}\) line \((h\nu = 1253.6\text{ eV})\) was used in x-ray photoemission spectroscopy (XPS). The resolution for He I, 120 eV and Mg \(K_{\alpha}\) are 40 meV, 100 meV and 1 eV, respectively. Also, we carried out Ni \(L_3\)-edge XAS at PLS. Samples were in situ fractured to obtain clean surface. Base pressure was \(2 \times 10^{-10}\text{ torr}\). For the energy calibration, a bulk palladium was measured at the same time.

III. RESULT AND DISCUSSION

Inset in Figure 2 shows PES data taken with 120 eV photon source. The peak centered at 1 eV below \(E_F\) corresponds to the Ni 3d derived conduction band. After the high pressure (HP) treatment, the spectrum below 3 eV decreased substantially. In general, photoemission intensity below the conduction band arises from non-intrinsic sources such as grain boundary or surface contamination. Decrease of it, together with the \(T_c\) enhancement, suggests that the sample quality is improved by the HP sintering. Note, however, that the band structure above 3 eV is almost independent of the sintering. Afterwards, we show the data of the HP sintered sample. Even the high-pressure sintered sample shows structures around \(\sim 6\text{ eV}\) binding energy which we presume mostly come from remnant MgO precipitates or grain boundaries. It may also be related with the structural inhomogeneity and the nanoprecipitates which were recently reported in this system.

To see the spectra near \(E_F\) more closely, we took spectrum using He I source (Figure 2, solid circles). For comparison, we also show predicted spectra from the three band calculations by Shim et al. [dotted], by Rosner et al. [dash dot] and by Dugdale et al. [short dot]. To get these lines, the theoretical densities of states were first convoluted by the Fermi-Dirac distribution, followed by broadening procedures: the energy-dependent Lorentzian broadening with \(\sim \alpha |E - E_F|\) \((\alpha = 0.3)\) and the Gaussian instrumental broadening of 40 meV linewidth. The curves were normalized to give the same integrated spectral weight as the experimental data. The short dash-dotted line shows the background based on the Shirley method.

In the data, four features are observed at \(\sim 2.7\text{ eV}, 1.2\text{ eV}, 0.7\text{ eV}\) and \(0.1\text{ eV}\). The first three features agree roughly with the theoretical lines, particularly with that by Dugdale. In the curve by Shim, \(1.2\text{ eV}\) peak position is deeper than the data. The curve by Rosner doesn’t properly predict the structure centered at \(\sim 1.2\text{ eV}\). According to band calculations, the three features are from non-bonding Ni 3d states.

The peak located very close to \(E_F\) \((0.1\text{ eV})\) corresponds to the vHs as predicted in the theoretical lines. Note, however, that the peak height is much smaller than the predictions. Rough estimation of the peak intensity shows about \(1/2 \sim 1/4\) of the theoretical value. The vHs arise from the saddle point near \(\Gamma\) in the band structure. The peak strength is then determined by the band curvature or the effective mass at this point. Our observation suggests that the calculations are overestimating the peak intensity or equivalently understimating the curvature. Also note that the height is different significantly among the three theory results, which shows that band structure near the saddle point depend sensitively on the calculation methods. Thus, correct estimation of the peak strength seems a non-trivial work.

In spite of the uncertainty in the calculation, the observed peak is smaller than any of the three curves. This may suggest that the peak is suppressed for reasons not accounted for in the band theory. For example, when electron-phonon interaction or electron-electron interaction are present, part of spectral weight will shift to higher energy. \(YNi_2B_2C\), a superconductor which bears some similarity with \(MgCNi_3\) where Ni-B and Ni-C bondings are important, is another system that has vHs close to \(E_F\). There, the observed peak is also suppressed compared with theoretical prediction and Kobayashi et al. interpreted it in terms of electron-electron or electron-phonon interaction.

It is also possible that the spectral weight suppression may be due to matrix element effects. In this case, the peak intensity will depend on the photon energy. We thus performed PES using various photon energy from 40 eV to 150 eV, but the spectra didn’t change. Surface effect to which UPS is somewhat sensitive is another possible source. The vHs is a bulk property which arises from the saddle point \((\Gamma)\) of the Fermi surface. As one approaches the surface, the band structure will change and the vHs feature may possibly be smeared.

Figure 3 shows Ni 2p core-level photoemission spectrum of \(MgC_{1-x}Ni_3\). The main peaks, corresponding to \(2p_{3/2}\) and \(2p_{1/2}\), respectively, are accompanied by the weak satellites at higher binding energies. The existence of this satellite structure signals the presence of \(d-d\) electron correlation effect, since such satellite structure originates from the two-hole bound state. For comparison with related compounds, we show Ni 2p spectra of Ni metal and \(YNi_2B_2C\) together with our data in the inset. Note that in \(MgC_{1-x}Ni_3\) the satellite position of Ni \(2p_{3/2}\) is a little large and its intensity is largely reduced compared with Ni metal. In the first order approximation, the relative satellite position and the intensity represent the \(d-d\) correlation energy and the number of \(d\) holes, respectively. Thus our observations imply that the correlation energy is a little farther apart and the \(d\) hole number is smaller in \(MgC_{1-x}Ni_3\) compared with Ni metal.

In \(MgC_{1-x}Ni_3\), Ni is strongly covalent bonded with C and then charge transfer from the Ni atoms to C will occur. This will result in the Ni 2p core level shift to higher binding energy. The observed shift in \(YNi_2B_2C\) is explained in this manner. However, the binding energy of Ni 2p is almost similar to that of Ni metal. This seems
to suggest that there is a large reduction of the binding energy due to, probably, a screening of core hole by free carriers. In fact, the CNi$_3$ cage is fully charged by the two electrons donated by the Mg$^{+2}$ ion and thus Ni hole will be effectively screened.

Figure 4 shows Ni L$_3$-edge XAS spectrum of MgC$_{1−x}$Ni$_3$. It is compared with the calculated Ni PDOS above the $E_F$ in the band calculation result by Shim et al. To get the theoretical curve (dash-dotted line), the Ni partial density of states are broadened by convoluting with the similar way as we did in Figure 2. But in this case we used the Lorenzian broadening with linewidth $\sim \alpha (E - E_F)^2$ ($\alpha = 0.2$) and the Gaussian broadening with 1.6 eV. The two curves are normalized to give the same integrated spectral weight. We see that overall structure is in reasonable agreement with the band calculations. The peak just above $E_F$ is due to the unoccupied part of Ni 3d band. But we also observe the satellite structures around $hv = 863$ eV in the experiments, which represents the correlation effects.

IV. CONCLUSION

In conclusion, we have performed XPS, XAS, and UPS measurements to study the electronic structure of MgC$_{1−x}$Ni$_3$. The satellite structure seen in Ni 2p XPS spectrum suggests that the 3d electron correlation is substantial. L$_3$-edge XAS and UPS data show that the position and width of the Ni 3d derived valence band is in reasonable agreement with theoretical calculation results. The vHs peak is found at $\sim 0.12$ eV below the Fermi energy. Its spectral weight is largely suppressed compared with theoretical predictions and we suggested various possibilities of the suppression including the electron-electron and electron-phonon interaction.

Acknowledgments

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It is possible that these defect structures affect its normal-state and superconducting properties.

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FIG. 1: Normalized magnetic susceptibility from the measured low-field magnetization $M(T, H = 100e)$ of as-grown (solid circles) and sintered at 900°C under 3GPa (open circles).
FIG. 2: He I UPS spectrum (●) of MgC$_{1-x}$Ni$_3$ at room temperature. All lines are theoretical spectra derived from the band-structure calculations. The short dash-dotted line shows the background. Inset: Valence region photoemission spectra with photon energy 120 eV of as-grown (solid circles) and sintered at 900°C under 3GPa (open circles).
FIG. 3: Ni 2p core-level photoemission spectrum of MgC$_{1-x}$Ni$_3$. Inset: Ni 2p spectra of Ni-metal and YNi$_2$B$_2$C are compared with that of MgC$_{1-x}$Ni$_3$. 

hv = 1253.6 eV
FIG. 4: Ni $L_3$-edge XAS spectrum of MgC$_{1-x}$Ni$_3$ (●). The band calculation result (dash-dotted line) by Shim et al. is compared. See text for the convolution.