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Optical signatures of the $J_{\text{eff}} = \frac{1}{2}$ state in Ir⁴⁺ halides

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We study the vibrational and electronic excitations in cubic iridium halides (NH₄)₂IrCl₆, K₂IrCl₆, and K₂IrBr₆ by infrared reflectivity and transmission measurements of single crystals. All three compounds show optical excitations in the energy range of 0.5–0.8 eV, which can be ascribed to $j_{\text{eff}} = \frac{3}{2}$ to $j_{\text{eff}} = \frac{1}{2}$ transitions (spin-orbit excitons), and t_{2g} -to- e_g excitations above \sim 1 eV. We observe at least two peaks due to spin-orbit excitons in the bromide and four peaks in both chlorides, suggesting local deviations from the cubic symmetry already at room temperature. We further show that the e_g states lie at lower energies in the bromide compared to the chlorides, in agreement with density-functional band-structure calculations.

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I. INTRODUCTION

Iridium oxides containing Ir⁴⁺ ions with a $5d^5$ electronic configuration have been studied intensively in the past years as relativistic Mott insulators hosting highly anisotropic exchange interactions that may lead, among others, to the Kitaev spin liquid ground state [1–4]. The energy-level scheme of iridates is illustrated in Fig. 1. The octahedral crystal field Δ_{cubic} splits Ir 5d states into the t_{2g} and e_g manifolds. The t_{2g} states are further separated into $j_{\text{eff}} = \frac{1}{2}$ and $j_{\text{eff}} = \frac{3}{2}$ states due to the strong spin-orbit (SO) coupling [5]. The rather narrow $j_{\text{eff}} = \frac{1}{2}$ band could, furthermore, be split into lower and upper Hubbard bands due to the on-site Coulomb repulsion, hence, leading to a Mott insulating state. Additionally, distortions of the IrO₆ octahedra can cause a noncubic crystal field that leads to a splitting of the $j_{\text{eff}} = \frac{3}{2}$ states and to a departure from the pure $j_{\text{eff}} = \frac{1}{2}$ state for the local magnetic moments.

Among the prominent examples of this physics, the honeycomb iridates Na₂IrO₃ and α -Li₂IrO₃ have raised special attention and extensive research investigations since their Ir⁴⁺ magnetic moments are close to $j_{\rm eff} = 1/2$ as witnessed by optical spectroscopy and resonant inelastic x-ray scattering (RIXS) [6–8]. However, magnetic interactions clearly deviate from the pure Kitaev model. Consequently, antiferromagnetic order is observed at low temperatures at ambient pressure. Moreover, the honeycomb iridates undergo a dimerization of the crystal structure under high pressure [9,10].

In this context, the iridium halides $A_2 Ir X_6$ with an antifluorite crystal structure (see Fig. 2), where A can be an alkali metal or NH₄ and X are halogen atoms, have caused attention because of their face-centered cubic symmetry that imparts cubic symmetry also to the isolated $Ir X_6$ octahedra and may lead to the ideal $j_{\rm eff} = \frac{1}{2}$ state of Ir^{4+} [11–16]. Indeed, RIXS studies on several $A_2 Ir X_6$ compounds found evidence for a proximity to the $j_{\rm eff} = \frac{1}{2}$ state, albeit with a small splitting of the excited $j_{\rm eff} = \frac{3}{2}$ quartet into two doublets due to residual structural distortions driven by soft phonon modes

in the crystal structure [13,14]. In this paper, we investigate the electronic ground state of the $A_2 Ir X_6$ materials further by infrared spectroscopy measurements, which enables the study of vibrational and electronic excitations with a better energy resolution as compared to RIXS. The obtained optical conductivity and absorbance spectra are compared to the theoretical density of states for $K_2 Ir Cl_6$ and $K_2 Ir Br_6$ as obtained by density-functional-theory (DFT) calculations.

II. METHODS

Single crystals of K₂IrCl₆, (NH₄)₂IrCl₆, and K₂IrBr₆ were grown from water solutions as reported elsewhere [14,17]. The crystal quality was checked by magnetization measurements and by x-ray powder diffraction performed on finely ground crystals selected from the same batch.

The room-temperature infrared spectroscopy measurements were carried out in the frequency range from 120 to 20000 cm⁻¹ (15 meV to 2.5 eV) with a Bruker Vertex 80v FTIR spectrometer coupled to a Bruker Hyperion IRmicroscope. The reflection measurements were performed on as-grown samples and for the transmission measurements the samples were polished to a thickness between 37 and 25 µm. In the case of K₂IrBr₆, a crystal was fixed on beeswax for polishing to a smaller thickness of \sim 20 μ m and for mechanical support during the transmission measurements. A commercial unprotected aluminum mirror served as reference for the reflectivity measurements. The real part of the optical conductivity $\sigma_1(v)$ was obtained by Kramers-Kronig analysis. To this end, the reflectivity spectra were extrapolated to low energies by using the Lorentz model. The extrapolation above 2.5 eV was calculated by x-ray atomic scattering functions [18]. The absorbance spectra $A(\nu)$ were calculated from the transmission spectra $T(\nu)$ according to $A(\nu) = -\log T(\nu)$. For the transmission measurements the empty beam path served as reference. For all measurements the spectral energy resolution $\Delta \nu$ amounts to ~ 2 cm⁻¹ (≈ 0.25 meV).

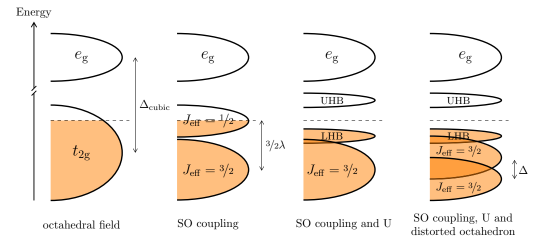


FIG. 1. Energy level scheme of the $5d^5$ orbitals in Ir^{4+} , taking into account the cubic crystal field $\Delta_{\rm cubic}$, spin-orbit coupling (SO) with strength λ , on-site Coulomb repulsion U, and noncubic crystal field Δ . Please note the interrupted energy scale because $\Delta_{\rm cubic} \gg \lambda$, and that the splittings $3/2\lambda$, U, and Δ are not in scale.

Relativistic DFT band-structure calculations were performed using the FPLO [19] and WIEN2K [20,21] codes with the Perdew-Burke-Ernzerhof exchange-correlation potential [22]. Correlation effects in the Ir 5d shell were taken into account on the mean-field DFT+U+SO level. The parameters of the correlated orbital were set to $U_d = 2.2$ eV (K₂IrCl₆) and $U_d = 1.8$ eV (K₂IrBr₆) for the on-site Coulomb repulsion as well as $J_d = 0.3$ eV for Hund's coupling in both compounds. These optimized parameters were previously shown to reproduce magnetic properties of the Ir⁴⁺ hexahalides [12,14]. Optical conductivity was calculated with the internal routines of WIEN2K [23] on the dense $24 \times 24 \times 24$ mesh.

III. RESULTS AND DISCUSSION

The absorbance A and optical conductivity σ_1 spectra of $(NH_4)_2IrCl_6$, K_2IrCl_6 , and K_2IrBr_6 are depicted in Figs. 3(a) and 3(b) over a broad frequency range. The optical data reveal the insulating character of all studied compounds. The profiles of the spectra are characterized by peaks centered at 0.6–0.7 eV due to the excitations of electrons from the

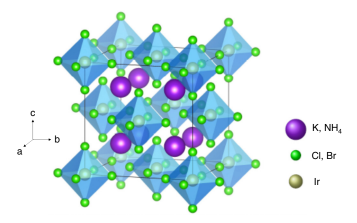


FIG. 2. Crystal structure of iridium halides $A_2 Ir X_6$, consisting of isolated $Ir X_6$ octahedra.

 $j_{\rm eff} = \frac{3}{2}$ to the $j_{\rm eff} = \frac{1}{2}$ states (see level scheme in Fig. 1), denoted as spin-orbit excitons in the literature [24], followed by the transitions from the t_{2g} to e_g states at higher energies. The t_{2g} -to- e_g transitions have a large spectral weight, which causes a steep absorption edge at 1.07 eV for K₂IrBr₆ and at 1.64 eV for (NH₄)₂IrCl₆ and K₂IrCl₆, with a saturation of the absorbance above 1.2 and 1.7 eV, respectively [see Fig. 3(a)]. Below \sim 0.5 eV the stretching and bending modes of the Ir X_6 octahedra can be observed. They are located at frequencies of 191 and 325 cm⁻¹ for (NH₄)₂IrCl₆, at 182 and 331 cm⁻¹ for K₂IrCl₆, and at 124 and 233 cm⁻¹ for K₂IrBr₆. The lower phonon mode frequencies for K₂IrBr₆ as compared to the other two compounds can be qualitatively explained by a simple harmonic-oscillator model with eigenfrequencies $\omega_j = \sqrt{\frac{k_j}{k_j}}$, where m_j is the reduced mass of the ions involved in the silbation, and k_j is the force constant of the model.

 $\omega_j = \sqrt{\frac{k_j}{m_j}}$, where m_j is the reduced mass of the ions involved in the vibration, and k_j is the force constant of the mode [25,26]. In the case of (NH₄)₂IrCl₆ we observe an additional mode at 130 cm⁻¹, which we assign to NH₄ modes, following the interpretation for K₂IrCl₆ in Ref. [16]. The corresponding modes of the K atoms in K₂IrCl₆ and K₂IrBr₆ are below 120 cm⁻¹ [16] and, hence, out of our measurement range. Besides, we observe NH₄⁺ molecular vibrations in (NH₄)₂IrCl₆ and in the case of K₂IrBr₆ the absorbance spectrum contains additional vibrations that are reminiscent of the OH groups and may indicate partial hydrolysis of the hexahalide ions during crystal growth [see Figs. 3(c) and 3(d)] [27]. We believe that [IrBr₆]²⁻ is more prone to the hydrolisis than [IrCl₆]²⁻, hence, these OH lines are observed in the bromide only. The NH₄⁺ and OH⁻-excitations are better seen in Fig. 5, which will be discussed in more detail later.

Since all materials are opaque above the energy of the absorption edge, the transition energies of the t_{2g} -to- e_g excitations cannot be extracted from the absorbance spectra. Instead, they can be obtained from the high-energy optical conductivity spectra where several well-defined peaks are observed. In order to determine the t_{2g} -to- e_g transition energies, the σ_1 spectra were fitted with the Lorentz model (see Fig. 4), and the so-obtained energies of the most pronounced and well-separated peaks are given in Table I.

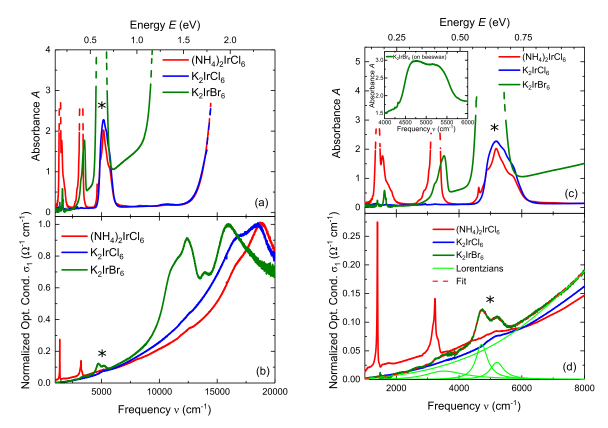


FIG. 3. Absorbance A and optical conductivity σ_1 spectra of three iridium halides: (a) absorbance spectra A in the energy range from 1000 to 20 000 cm⁻¹ (0.12 to 2.48 eV) and (b) corresponding optical conductivity σ_1 , normalized to the maximum value for comparison reasons, respectively. (c) and (d) are zoom-ins of (a) and (b) on the frequency range from 1000 to 8000 cm⁻¹ (0.12 to 0.99 eV). The asterisks mark the excitations from the $j_{\text{eff}} = \frac{3}{2}$ to the $j_{\text{eff}} = \frac{1}{2}$ energy levels. The inset in (c) shows the nonsaturated absorbance spectrum A of a 25- μ m thick K₂IrBr₆ sample for frequencies around the $j_{\text{eff}} = \frac{3}{2}$ -to- $j_{\text{eff}} = \frac{1}{2}$ excitations.

For K_2IrBr_6 the $j_{eff}=\frac{3}{2}$ -to- $j_{eff}=\frac{1}{2}$ transitions in the energy range of 0.6–0.7 eV have a larger spectral weight as compared to the other two studied compounds as can be seen in the optical conductivity spectra [Figs. 3(b) and 3(d)] where these transitions are marked with an asterisk. Since the reflectivity measurements on all samples were carried out on as-grown crystal surfaces, the features in the corresponding optical conductivity are broadened by the scattering effect. From the σ_1 spectrum of K_2IrBr_6 we could resolve two $j_{eff}=\frac{3}{2}$ -to- $j_{eff}=\frac{1}{2}$ transitions. The corresponding absorbance spectra obtained by transmission measurements are less sensitive regarding scattering effects and, therefore, allow a more detailed view on the $j_{eff}=\frac{3}{2}$ -to- $j_{eff}=\frac{1}{2}$ excitations. However, the high spectral weight of these excitations in

TABLE I. Energies of the t_{2g} - e_g transitions as obtained from the Lorentz fittings of the optical conductivity σ_1 spectra [Fig. 4]. All parameters are in eV. The contribution for $(NH_4)_2IrCl_6$ marked with an asterisk leads to a bump in the spectrum with very small spectral weight.

Material	$\hbar\omega_1$	$\hbar\omega_2$	$\hbar\omega_3$	$\hbar\omega_4$	$\hbar\omega_5$
(NH ₄) ₂ IrCl ₆	1.55*	2.09	2.32		
K ₂ IrCl ₆	1.59	2.07	2.32		
K_2IrBr_6	1.38	1.55	1.73	1.97	2.21

 K_2IrBr_6 causes a saturation of the absorbance spectrum in this frequency range [Fig. 3(c)]. Even in a very thin crystal with thickness ~25 μm stabilized by beeswax the absorbance is close to saturation [inset of Fig. 3(c)], and, therefore, it was impossible to resolve all the contributions to the $j_{eff} = \frac{3}{2}$ -to- $j_{eff} = \frac{1}{2}$ transitions. In contrast, for $(NH_4)_2IrCl_6$ and K_2IrCl_6 all contributions can be well resolved in the absorbance spectra since there is no saturation.

In the following, we focus on the transitions from the $j_{\rm eff}=\frac{3}{2}$ to the $j_{\rm eff}=\frac{1}{2}$ energy levels in the energy range of 0.6–0.7 eV as observed in the experimental optical spectra. For a quantitative characterization, we carried out Lorentz fittings of the absorbance spectra [see Figs. 5(a), 5(b) and 5(d)] and the σ_1 spectrum of K₂IrBr₆ [see Fig. 3(d)]. In the case of K₂IrCl₆, we can resolve four contributions according to the fitting of the absorbance spectrum. For (NH₄)₂IrCl₆, the absorbance spectrum shows additional excitations of stretching modes of (NH₄)⁺ and combinations thereof located at energies of 0.15–0.25 eV and 0.30–0.45 eV [28–30]. We ascribe the sharp peak located at 0.57 eV to a combined vibration of (NH₄)⁺.

As already mentioned above, for K_2IrBr_6 it is difficult to reveal all contributions to the $j_{eff} = \frac{3}{2}$ -to- $j_{eff} = \frac{1}{2}$ transitions from the absorbance spectra since the absorbance saturates in the range of 0.6–0.7 eV, even for a very thin crystal stabilized by beeswax [see Figs. 5(c) and 5(d)]. The additional excitations in the spectrum at around 0.35 eV are due to beeswax [31], and the spectra of the bromide compound contain hints

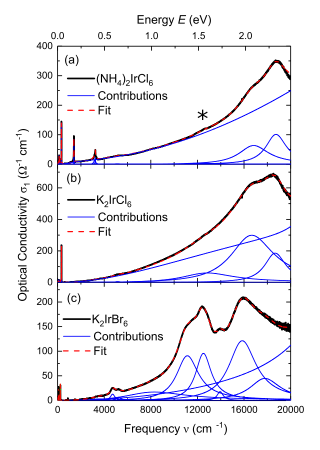


FIG. 4. Fits of the optical conductivity spectra σ_1 of three iridium halides with the Lorentz model together with the Lorentz contributions. The asterisk in panel (a) indicates the small bump, which is also marked with an asterisk in Table I.

for OH excitations. Based on the fit of the σ_1 spectrum [see Fig. 3(d)] we could identify only two contributions to the $j_{\rm eff}=\frac{3}{2}$ -to- $j_{\rm eff}=\frac{1}{2}$ excitations in K₂IrBr₆ since a broad background obscures features with less spectral weight. Hence, for K₂IrBr₆ we could determine the energies of two $j_{\rm eff}=\frac{3}{2}$ to $j_{\rm eff}=\frac{1}{2}$ transitions but cannot rule out additional transitions.

The energies of the $j_{\rm eff} = \frac{3}{2}$ -to- $j_{\rm eff} = \frac{1}{2}$ transitions in all studied compounds are listed in Table II. According to Ref. [32] these transition energies can be used to estimate the value of the spin-orbit coupling constant λ since their aver-

TABLE II. Energies of the transitions from the $j_{\text{eff}} = \frac{3}{2}$ to the $j_{\text{eff}} = \frac{1}{2}$ energy levels as obtained from the Lorentz fittings of the absorbance spectra [Figs. 5(a), 5(b) and 5(d)] and the σ_1 spectrum [Fig. 3(d)], together with the spin-orbit coupling strength λ (see text). All parameters are in meV.

Material	$\hbar\omega_1$	$\hbar\omega_2$	$\hbar\omega_3$	$\hbar\omega_4$	λ
(NH ₄) ₂ IrCl ₆	605	643	671	708	438
K ₂ IrCl ₆	616	641	674	710	440
K ₂ IrBr ₆ (absorbance)	582	654			412
K_2IrBr_6 (σ_1)	585	646			410

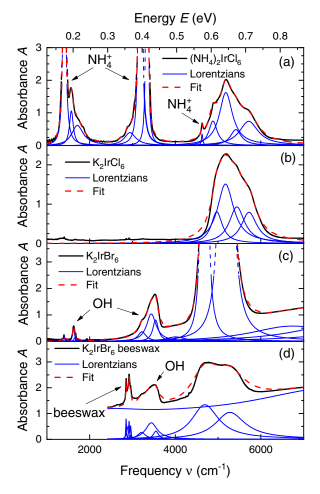


FIG. 5. Absorbance spectrum A with the corresponding Lorentz fit and Lorentz contributions for (a) $(NH_4)_2IrCl_6$, (b) K_2IrCl_6 , (c) K_2IrBr_6 , and (d) a thinner K_2IrBr_6 crystal fixed on beeswax. The vibrational excitations related to NH_4^+ , OH^- , and beeswax are indicated.

age energy corresponds to $\frac{3}{2}\lambda$. For all studied iridium halide compounds, the values of λ lie in the range of 410–439 meV (see Table II), which is in good agreement with the values reported in Refs. [13,14] obtained by RIXS studies with an overall energy resolution of \sim 35 meV. However, in contrast to the RIXS results, we could resolve four contributions for the compounds (NH₄)₂IrCl₆ and K₂IrCl₆ due to the very good energy resolution of infrared spectroscopy (\sim 0.25 meV), and this most probably holds also for K₂IrBr₆.

It has been argued that the observation of two excitations of the intra- t_{2g} levels in the RIXS spectra can be attributed to a splitting of the $j_{\rm eff}=\frac{3}{2}$ quartet state into two doublet states due to noncubic crystal field [13]. However, such a splitting cannot explain the four excitation peaks in the infrared spectra. For an explanation of this finding, we carried out orbital-resolved DFT band-structure calculations, and the calculated density of states for K_2IrCl_6 and K_2IrBr_6 are shown in Fig. 6. Accordingly, the lower Hubbard band of $j_{\rm eff}=\frac{1}{2}$ is strongly mixed with $j_{\rm eff}=\frac{3}{2}$ and, moreover, with the ligand p states that are dominant even in the vicinity of the Fermi level (the inset of Fig. 6). Possible explanations for our findings of

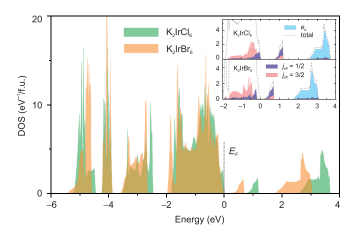


FIG. 6. Comparison of DFT+U+SO density of states (DOS) for cubic K₂IrCl₆ and K₂IrBr₆. The Fermi level is at zero energy. In the case of bromide, the lower value of U_d and the smaller crystal-field splitting both shift Ir e_g states to lower energies. The inset: Orbital-resolved DOS for K₂IrCl₆ and K₂IrBr₆.

four intra- t_{2g} transitions are, therefore, (i) a finite width of both $j_{\rm eff}=\frac{1}{2}$ and $j_{\rm eff}=\frac{3}{2}$ bands, which cause multiple absorption features since optical spectroscopy probes the transitions at all points in the Brillouin zone and (ii) transitions between the lower and the upper Hubbard bands of $j_{\rm eff}=\frac{1}{2}$, which coincide with the $j_{\rm eff}=\frac{3}{2}$ -to- $j_{\rm eff}=\frac{1}{2}$ transitions. Besides, the structural distortion in K_2IrCl_6 and $(NH_4)_2IrCl_6$, whose nature is presently unknown, might lead to the formation of several Ir^{4+} sites. It is important to note that recent Raman spectroscopy studies on K_2IrCl_6 single crystals also observed multiple $j_{\rm eff}=\frac{3}{2}$ -to- $j_{\rm eff}=\frac{1}{2}$ transitions from which $\lambda\approx0.41$ eV was extracted, in good agreement with our results [16]. The multiple excitations were attributed to spin-orbit excitons with a possible coupling to electron-hole excitations of the $j_{\rm eff}=\frac{1}{2}$ orbitals.

Consistent with the above interpretation, the calculated optical conductivity σ_1 spectra of K_2IrBr_6 and K_2IrCl_6 , depicted in Fig. 7, contain several excitation peaks in the energy range 0.4–0.7 eV. These $j_{\rm eff}=\frac{3}{2}$ to $j_{\rm eff}=\frac{1}{2}$ and $j_{\rm eff}=\frac{1}{2}$ to $j_{\rm eff}=\frac{1}{2}$ transitions appear with a rather large spectral weight in the theoretical spectra, which might be a downside of the mean-field treatment of electronic correlations within DFT+U+SO. The steep onset of the optical conductivity at around 2.8 and 3.6 eV is ascribed to t_{2g} -to- e_g transitions for K_2IrBr_6 and K_2IrCl_6 , respectively. The different energetic positions of the onset can be traced back to the calculated DOS where the Ir e_g states are shifted to lower energies in the case of the bromide compound (see Fig. 6). This happens because of the smaller cubic crystal-field splitting in the bromide since the Ir-ligand distances are longer compared to the chlorides.

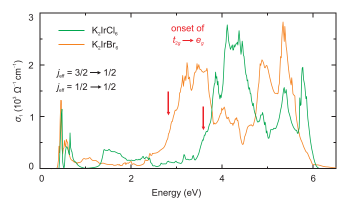


FIG. 7. Comparison of the optical conductivity σ_1 calculated for K_2IrBr_6 and K_2IrCl_6 on the DFT+U+SO level.

Additionally, the lower value of the Hubbard parameter U_d may lead to the smaller splitting between the filled t_{2g} states and empty e_g states in the bromide. This finding is in qualitative agreement with the experimental data [see Fig. 3(b)]; however, theory overestimates the energy of the t_{2g} -to- e_g transitions as compared to the experimental results.

IV. CONCLUSION

The absorbance and optical conductivity spectra of the nominally cubic iridium halides (NH₄)₂IrCl₆, K₂IrCl₆, and K_2IrBr_6 show vibrational modes of the IrX_6 octahedra and NH_4^+ molecular vibrations below ~ 0.5 eV and electronic transitions between $j_{\text{eff}} = \frac{3}{2}$ and $j_{\text{eff}} = \frac{1}{2}$ states and between the lower and the upper Hubbard bands of $j_{\text{eff}} = \frac{1}{2}$ at energies 0.6–0.7 eV, followed by t_{2g} -to- e_g transitions above \sim 1 eV. We confirm that the splitting of the SO exciton peaks occurs already at room temperature. This indicates local deviations from the cubic symmetry and the departure from the ideal $j_{\rm eff} = \frac{1}{2}$ state of Ir⁴⁺ even if macroscopic symmetry probed by x-ray diffraction remains cubic. Additionally, in contrast to the recent RIXS measurements, we could resolve four contributions to the $j_{\text{eff}} = \frac{3}{2}$ -to- $j_{\text{eff}} = \frac{1}{2}$ excitations for $(NH_4)_2$ IrCl₆ and K₂IrCl₆ due to the advanced energy resolution of infrared spectroscopy. The lower energies of the electronic excitations in the experimental optical spectra of K₂IrBr₆ are due to the lower on-site Coulomb repulsion U_d and the smaller crystal-field splitting according to our DFT band structure calculations. The optical data of K₂IrBr₆ and K₂IrCl₆ are in qualitative agreement with the calculated optical conductivity spectra.

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