Pressure-induced spin transition and site-selective metallization in CoCl₂

Jose A. Barreda-Argüeso¹, Lucie Nataf², Fernando Aguado¹, Ignacio Hernández¹, Jesús González³, Alberto Otero-de-la-Roza³, Victor Luaña³, Yating Jia⁴, Changqing Jin⁴, Bongjae Kim⁵,⁶, Kyoo Kim⁷, Byung I. Min⁷, Wilhem Heribert⁸, Andrew P. Jephcoat⁹ & Fernando Rodríguez⁵,⁶

The interplay between spin states and metallization in compressed CoCl₂ is investigated by combining diffraction, resistivity and spectroscopy techniques under high-pressure conditions and ab-initio calculations. A pressure-induced metallization along with a Co²⁺ high-spin (S = 3/2) to low-spin (S = 1/2) crossover transition is observed at high pressure near 70 GPa. This metallization process, which is associated with the p-d charge-transfer band gap closure, maintains the localization of 3d electrons around Co²⁺, demonstrating that metallization and localized Co²⁺ - 3d low-spin magnetism can coexist prior to the full 3d-electron delocalization (Mott-Hubbard d-d breakdown) at pressures greater than 180 GPa.

Pressure-induced structural phenomena have received considerable attention in transition-metal (M) dihalides and oxides MX₂ (X: Cl, Br, F, O) due to their ample and subtle polymorphism¹⁻¹⁰ and their intriguing electronic properties associated with changes of metal coordination, spin state, and/or insulator-metal transition. Most MX₂ are antiferromagnetic Mott insulators¹¹ that present strong d-d electron-electron correlation¹². The breakdown of the Mott-Hubbard d-d or charge-transfer d-p electron correlation leading to a metallization, concurrent with a collapse of magnetism via electron delocalization, is the typical pressure-induced electronic/magnetic behavior. These phenomena involve external pressures that induce a large crystal-field strength at M, causing the high-spin to low-spin (HS-LS) transition¹³⁻¹⁵. In fact, spin crossover (SCO) is often portrayed as a trigger for metallization either by volume collapse¹⁶,¹⁷, or as a result of the ground-state change from HS to LS¹⁸⁻²⁰. Nevertheless, the coupling between structure, spin state, and electron delocalization (Mott-Hubbard metal-insulator transition) governing the electronic properties in MX₂ requires clarification. The access to both electronic ground and excited states via optical spectroscopy at high pressure, combined with the modelling of the electronic properties through ab initio calculations and precise crystal structure determination and resistivity measurements, can provide a definitive description of electronic phenomena in compressed MX₂ systems.

Pressure-induced transformations in MX₂ involve a large variety of energetically-equivalent dense structures, which are characterized either by an increase of the M coordination number¹¹,¹², or by stacking up dense MX₂ layers keeping the metal coordination¹³,¹⁴, depending on the metal/ligand ionic radii ratio. Due to their simple structure, phase diagrams of crystals in the MX₂ family show common features regarding the coordination polyhedra and the stacking sequence. This distinct structural behavior has important implications in their electronic properties, which are substantially modified upon compression. Besides, these studies of MX₂ under pressure are of importance in geophysics to understand polymorphism in the Earth's interior, where SiO₂ plays an important role in the Earth's interior, where SiO₂ plays an important role in the Earth's interior, where SiO₂ plays an important role in the Earth's interior, where SiO₂ plays an important role in the Earth's interior, where SiO₂ plays an important role in the Earth's interior, where SiO₂ plays an important role in the Earth's interior, where SiO₂ plays an important role in the Earth's interior, where SiO₂ plays an important role in the Earth's interior, where SiO₂ plays an important role in the Earth's interior, where SiO₂ plays an important role in the Earth's interior, where SiO₂ plays an important role in the Earth's interior, where SiO₂ plays an important role in the Earth's interior, where SiO₂ plays an important role in the Earth's interior, where SiO₂ plays an important role in the Earth's interior, where SiO₂ plays an important 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¹Universidad de Cantabria, MALTA Consolider Team - DCITIMAC, Santander, 39005, Spain. ²Synchrotron SOLEIL, L’Orme des Merisiers, St Aubin BP48, 91192, Gif-sur-Yvette cedex, France. ³Universidad de Oviedo, Departamento de Química Física y Analítica, Oviedo, 33006, Spain. ⁴Institute of Physics, Beijing National Laboratory for Condensed Matter Physics, Chinese Academy of Sciences, Beijing, 100190, China. ⁵Pohang University of Science and Technology, Department of Physics, PCTP, Pohang, 37673, Korea. ⁶Kunsan National University, Department of Physics, Gunsan, 54150, Korea. ⁷Pohang University of Science and Technology, Max Planck POSTECH/Hsinchu Center for Complex Phase Materials, Pohang, 37673, Korea. ⁸Diamond Light Source Ltd, Chilton, Didcot, Oxfordshire, OX11 0DE, United Kingdom. ⁹Okayama University, Institute for Planetary Materials, Yamada 827, Misasa, Tofuku, Tottori, 682-0193, Japan. Correspondence and requests for materials should be addressed to F.R. (email: rodriguf@unican.es)
CoCl₂ is an attractive system for studying combined structural and electronic effects because the octahedral coordination of Co²⁺ (3d⁶) is thought to be stable in a wide pressure range from 0 to 60 GPa, which enables us to exclude the physics coming from the coordination number changes. Besides, this system has a relatively small energy difference between its HS and LS phases, and the SCO of the system can be observed in a very accessible pressure regime. Furthermore, it is the member of the MCl₆ series (M: Cr, Mn, Fe, Co) where SCO is expected to take place at the lowest pressure. In addition, SCO phenomena involving transition-metal ions with 3d⁶ electronic configurations are very scarce. Thus, this system can be a good model system to explore the various origins of SCO and to give answer to questions such as whether the pressure-induced LS states (3d⁶ configuration) originate from large crystal-field effects enhanced by the Jahn-Teller effect associated with LS configuration, or from the metallization, which can be induced either by the p-d charge-transfer gap closure or breakdown of d-d Mott-Hubbard correlation.

Here we report on the stability of the Co²⁺ coordination under compression in CoCl₂, and the relationship between SCO, crystal structure, and metallization. In order to achieve these goals we perform a combined experimental and theoretical study using optical absorption, Raman spectroscopy, X-ray diffraction (XRD) and resistivity under pressure, and first-principles Density Functional Theory (DFT) calculations.

Results

The Co²⁺ (3d⁶) Tanabe-Sugano diagram describing the electronic states energy in terms of the octahedral crystal field (in Racah parameter B units, Fig. 1) shows that, for CoCl₂ (B ≈ 80 meV at 50 GPa), the HS-LS (4T₁₁/₂ → 2E, S = 3/2 → 1/2) transition should occur at ΔSCO = 1.7 eV. Importantly, the LS state may be affected by a strong Jahn-Teller effect, providing an additional lattice relaxation energy which, in principle, could reduce the SCO crystal-field strength triggering metallization at unexpectedly low pressure. Alternatively, high pressure could, in turn, suppress the Jahn-Teller distortion causing the HS-LS to occur at higher pressures than expected, or even disappear if CoCl₂ transforms into a fluorite-type structure (3d⁶ configuration). XRD shows that CoCl₂ exhibits nearly-degenerate layered structures at ambient conditions. The CoCl₂ pressure-induced phase transition sequence as determined experimentally with support of DFT calculations is indicated in Fig. 1. The three represented layered structures are more stable than the rutile, cotunnite, and fluorite phases at all pressures up to 100 GPa. This result, which is confirmed by both single-crystal and powder XRD experiments at high pressure, demonstrates the stability of the hexagonal layered structure of CoCl₂ and thus the Co²⁺ sixfold coordination in a wide pressure range (0–60 GPa), in a way similar to FeCl₂ and MgCl₂, and contrary to CoF₂.

The stability of the given CoCl₂ structures, which involve the different packing sequence of layers of face-sharing CoCl₆ octahedra, is a consequence of the subtle competition of inter-layer van der Waals interactions. The CoCl₂ equation-of-state can be phenomenologically described by two Murnaghan’s equations:

\[ \Delta \approx \frac{21}{2} \] for the crystal-field region (a). Besides the gap energy, these spectra allow us to determine the excited-state electronic structure in the transparency window of CoCl₂ (≈50 GPa). At ambient conditions, the main sub-gap absorption peaks within the D₂h (nearly O₃) CoCl₄⁻ octahedron correspond to crystal-field transitions ⁴T₁(F) → ⁴T₂(G), ⁴A₂(F), and ⁴T₁(P) and are located at 0.79, 1.66 and 2.10 eV, respectively. In terms of the Tanabe-Sugano diagram for d⁶ ions (Fig. 1b), the transition energies at ambient pressure correspond to Δ = 0.87 eV and B = 97 meV with Δ/B = 9.0 (see Supplementary Table S1). According to this diagram the crystal-field strength required to induce the SCO is (Δ/B)SCO = 21. Interestingly, the SCO also involves crossing of the ⁴T₁(F) and ⁴T₂(G) excited states, hence these states, which are well observed by optical absorption, can be used to efficiently probe the HS-LS transition.

Figure 1. (a) Structural phase-transition sequence in CoCl₂ under high-pressure conditions determined by density-functional-theory calculations and confirmed experimentally by x-ray diffraction. (b) Simplified Tanabe-Sugano diagram for octahedral Co²⁺ (3d⁶) showing the effect of pressure (i.e. Δ/B) on the excited state energies, and HS-LS crossover (Δ/B = 21). See also Supplementary Figs S1–S9.
The variation of the absorption spectra with pressure shows that the band gap energy decreases linearly with pressure at a rate of $-43 \text{meV/GPa}$ (Fig. 3). Such a large shift is responsible for the intense piezochromism exhibited by CoCl$_2$ (Fig. 2a). The pressure-induced redshift of the bandgap follows a quadratic dependence with the crystal volume yielding gap closure at $V = 17.5 \AA^3$/Co ($V/V_0 = 0.56$) – i.e. 80 GPa – (see Supplementary Fig. S5(c)). This redshift is produced by the hybridization enhancement of the Cl $-p$ and Co$^{2+}$ $-d$ orbitals with pressure which in turn causes a broadening of the mainly 3$p$- and 3$d$-orbital valence band and an energy decrease of the mainly 3$d$-orbital intermediate band both reducing the $p$-$d$ charge-transfer bandgap. DFT reproduces the decrease in the band gap of HS state ($P < 67 \text{GPa}$) reasonably well (see Supplementary Fig. 6(c)). The plots of the electron band and density of states certainly show a clear energy delocalization of the $d$-orbital manifolds with pressure yielding band broadening (see Supplementary Fig. S7(c,d)). Concurrently, the increasing crystal-field energy as obtained from the optical spectra, and the reduction of $B$ from 97 to 82 meV in the 0–50 GPa range yield a $\Delta/B$ variation from 9.0 to 18.5, which implies an almost doubled splitting between $e$ and $t_2$ orbitals, $\Delta$, from 0.87 to 1.52 eV (see Supplementary Table S1 and Fig. S5).

As Figs 2 and 3 show, the variation of the absorption spectrum and its associated peak energies with pressure reveal that Co$^{3+}$ has a HS state in the crystal transparency range. However, extrapolating the linear dependence of the transition energies with pressure we obtain a HS to LS $[4T_1(F) \leftrightarrow 2E(G)]$ transition at 67 GPa. It is worth noting that the SCO is observed in the DFT + $U$ results using a Coulomb correlation energy $U = 3 \text{eV}$ (Figs 3 and 4). As indicated in the Methods section, this method cannot capture the evolution of the Coulomb correlation parameter upon volume changes and could underestimate the SCO pressure for high pressure regime. However, the essence of the electronic and magnetic properties before and after the transition is valid. The spectroscopic determination of the 3$d$-electron structure together with the DFT estimates make CoCl$_2$ a reference system to validate theories dealing with SCO phenomena and metallization processes in transition-metal systems.

Figure 3 plots the calculated magnetic moment as a function of pressure. According to DFT calculations, the Co$^{3+}$ magnetic moment abruptly decreases from HS, $\mu_{\text{eff}} = 2.6 \mu_B$, to LS, $\mu_{\text{eff}} = 0.9 \mu_B$, at around 67 GPa with a hysteresis of 8 GPa. The experimental SCO pressure and the crystal and electronic structures demonstrate that the Jahn-Teller coupling is not involved in the stabilization of the LS ground state, $2E$. This contrasts with one of the hypotheses given elsewhere, that the high-pressure conditions required for SCO could be relaxed by the strong Jahn-Teller effect. The Jahn-Teller coupling in those measured in the CuCl$_6$ system. The lack of a HS-LS transition at $\approx 35 \text{GPa}$ in CoCl$_2$ indicates that the Jahn-Teller effect is unable to distort the Co$^{3+}$ environment in the severe high-pressure conditions required for SCO.

The pressure dependence of the optical gap, $E_{\text{Gap}}$, allows us to infer that the $p$-$d$ charge-transfer gap closure (metallization) takes place at 80 GPa. This result is confirmed by electrical resistance measurements under pressure (inset of Fig. 3 and Supplementary Fig. S12). Its pressure dependence $R(P)$ unveils two distinct regions corresponding to HS and LS states. The associated SCO pressure, $P_{\text{SCO}} = 70 \text{GPa}$, is close to that derived from optical absorption. Interestingly, the progressive decrease of $R(P)$ in LS shows a change of slope for $P > 80 \text{GPa}$ indicating the metallization onset. Spin density- and DOS simulations indicate that the charges are mainly localized at the Co$^{3+}$ site and small $p$-$d$ hybridized ones can be observed at Cl$^-$ sites for HS. For $P > 67 \text{GPa}$ (LS) a progressive decrease of the localized charges at Co$^{2+}$ occurs due to hybridization increase. However, the hybridized spin density spreads out over the entire crystal for $P > 180 \text{GPa}$ ($E_{\text{Gap}} = 0$) in the Cl$^-$ plane, while it is strongly localized at Co$^{3+}$, indicating that metallization mainly involves Cl$^-$ sublattices rather than the Co$^{2+}$ ones. Furthermore, a full electron delocalization is completed for $P > 180 \text{GPa}$ (Fig. 4). This result is noteworthy since it correlates two
distinct electronic features: (1) the insulator-to-metal transition involves $p$-$d$ charge-transfer states and can be induced under compression in close proximity right after the HS-LS transition; (2) Mott-Hubbard $d$-$d$ electron breakdown should occur for $P > 180$ GPa. Thus, $3d$-electrons still keep their local character at the band-gap closure, albeit pressure-induced progressive delocalization occurs within LS ground state up to approximately 180 GPa, at which delocalization process is completed (Figs 3 and 4).

Conclusions

In summary, with various types of experimental and theoretical approaches, we have thoroughly analysed the physics of the pressure-induced spin-state transition and metallization phenomena in CoCl$_2$. We have shown that the layered structure of CoCl$_2$, and hence the Co$^{2+}$ sixfold coordination, is stable in the 0–200 GPa range, in contrast to CoF$_2$, whose high-pressure phases involve increasing coordination numbers (6 $\rightarrow$ 8 $\rightarrow$ 9). We demonstrate that pressure-induced metallization is associated with $p$-$d$ charge-transfer band gap, closing at about 80 GPa. Although the HS-to-LS transition (67 GPa) can trigger insulator-to-metal transition, DFT calculations also show that after the SCO metallization Co$^{2+}$ preserves the local character of the $3d$-electrons and that Mott-Hubbard-electron breakdown takes place for $P > 180$ GPa (detailed information in Supplementary Figs S4–S7). The inset shows the variation of the electrical resistance of CoCl$_2$ with pressure, $R(P)$, in a semilog plot. It must be noted that $R(P)$ behaves differently below and after 71 GPa indicating SCO transition. The change of slope above 80 GPa unveils the metallization onset. Linear dashed line corresponds to the least-square fit to $R(P)$ taking the linear variation of $p$-$d$ charge-transfer band gap, $E_{GAP}$, obtained by optical absorption, and a thermal activated hoping term, $E_{hop}(P)$ associated to the carrier mobility (see also Supplementary Fig. S12).

Figure 3. Variations of the Co$^{2+}$ crystal-field state energy of the $^{4}T_1(a)$ $\rightarrow$ $\Gamma_i$ transition and of the charge-transfer band gap energy of CoCl$_2$ ($E_{GAP}$) with pressure. The color of the experimental points ($\Gamma_i$) is the same one employed for representing the energy of $\Gamma_i$ state in the Tanabe-Sugano diagram of Fig. 1. Note that the HS-LS crossover corresponds to the $^{4}T_1(F)$ $\leftrightarrow ^{2}T_1(G)$ crossing point, and the $p$-$d$ band gap closure to $E_{GAP} = 0$ (metallization). The continuous curves in brown color are the calculated Co$^{2+}$ local moment within DFT + U method keeping the $P3m1$ phase in the whole pressure range. The HS-LS transition and $p$-$d$ band gap closure take place at 67 and 80 GPa, respectively, whereas the Mott-transition ($\sigma$-electron delocalization) in LS is found for $P > 180$ GPa (detailed information in Supplementary Figs S4–S7). The inset shows the variation of the electrical resistance of CoCl$_2$ with pressure, $R(P)$, in a semilog plot. It must be noted that $R(P)$ behaves differently below and above 71 GPa indicating SCO transition. The change of slope above 80 GPa unveils the metallization onset. Linear dashed line corresponds to the least-square fit to $R(P)$ taking the linear variation of $p$-$d$ charge-transfer band gap, $E_{GAP}$, obtained by optical absorption, and a thermal activated hoping term, $E_{hop}(P)$ associated to the carrier mobility (see also Supplementary Fig. S12).

Methods

Crystal structure: X-ray diffraction. Both single-crystal plates (100 $\times$ 80 $\times$ 30 $\mu$m$^3$) and powder of CoCl$_2$ (Merck) were used for high-pressure experiments. CoCl$_2$ crystallizes in the trigonal space group $R3m$ at ambient conditions. The evolution of the crystal structure with pressure was studied by x-ray diffraction (XRD) using the I15 beam station at the DIAMOND synchrotron under proposals 832, 1655 and 6078. The pressure was applied by means of Almax-Boehler and MALTA-type Diamond Anvil Cell (DAC). DACs were loaded with several Ruby...
spheres (10 μm diameter) as pressure gauge using helium, silicone oil and paraffin as pressure-transmitting media for powder and single crystal XRD experiments (see Supplementary Figs S1–S3).

Optical absorption and Raman spectroscopy. Optical absorption and Raman experiments were performed on single-crystal plates (100 × 80 × 35 μm³) of CoCl₂. The optical spectroscopy experiments were carried out in membrane and Almax-Boehler DACs. 200-μm-thick Inconel 625 gaskets were preindented to 40 μm. 170-μm-diameter holes were perforated with a BETSA motorized electrical discharge machine. The DAC was loaded with a CoCl₂ single crystal and ruby microspheres (10 μm diameter) as pressure probes using silicone oil as pressure-transmitting medium in an argon atmosphere inside a globe box to avoid sample hydration. Optical absorption under high-pressure conditions was performed on a prototype fiber-optics microscope equipped with two 20× reflecting objectives mounted on two independent x-y-z translational stages for the microfocus beam, and the collector objective and a third independent x-y-z translational stage for the DAC holder (Fig. 5). Optical absorption data and images were obtained simultaneously with the same device. Spectra in the UV-VIS and NIR were recorded with Ocean Optics USB 2000 and NIRQUEST 512 monochromators using Si- and InGaAs-CCD detectors, respectively.

Unpolarized micro-Raman scattering measurements were performed with a triple monochromator Horiba-Jobin-Yvon T64000 spectrometer in subtractive mode backscattering configuration, equipped with a Horiba Symphony liquid-nitrogen-cooled CCD detector. The 514.5-nm and 647-nm lines of a Coherent Innova 70 Ar⁺-Kr⁺ laser were focused on the sample with a 20× objective for micro-Raman, and the collector objective and a third independent x-y-z translational stage for the DAC holder (Fig. 5). Optical absorption data and images were obtained simultaneously with the same device. Spectra in the UV-VIS and NIR were recorded with Ocean Optics USB 2000 and NIRQUEST 512 monochromators using Si- and InGaAs-CCD detectors, respectively.

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Electrical measurements at high pressure. The electrical resistance measurement under pressure up to 96 GPa was performed using diamond anvil cell with solid transmitting medium NaCl (diamond's culet diameter.
of 100 μm). Gasket consists of T301 and the insulate layer is cBN. Pressure was determined by ruby fluorescence method at low pressure and the shift of diamond’s Raman peaks. Figure 6 shows a schematic view of the DAC.

First-principles theoretical calculations. Density functional theory: crystal structure and phase transition. For the description of an accurate phase transition, we have performed total-energy calculations within the framework of dispersion-corrected Density-Functional Theory in the Projector Augmented Wave (PAW) and plane-waves (PW) formulation. The Perdew-Burke-Ernzerhof (PBE) exchange-correlation functional was employed coupled with the exchange-hole dipole moment (XDM) model dispersion correction (damping function parameters $a_1 = 0.0000$ and $a_2 = 3.8036$ Å) as implemented in Quantum ESPRESSO. The calculation parameters are: $6 \times 6 \times 6$ Monkhorst-Pack k-point grid, 60 Ry plane-wave cutoff energy, 600 Ry density-cutoff energy and cold smearing with a smearing parameter of 0.05 Ry. Based on previous studies, we considered the following phases of CoCl$_2$: cotunnite ($Pnma$, orthorhombic, $Z = 4$), CaCl$_2$ (deformed rutile structure, $Pnnm$, orthorhombic, $Z = 2$), fluorite ($Fm\overline{3}m$, cubic, $Z = 1$), CoCl$_2$ ($R\overline{3}m$, rhombohedral, ABCABC stacking), CdI$_2$ ($P\overline{6}$_3 $mc$, hexagonal, ABAB stacking), $\omega$-phase ($P\overline{3}m1$, hexagonal, AAA stacking). All CoCl$_2$ phases were calculated in a range of volumes encompassing the 0–100 GPa range and the internal degrees of freedom (atomic positions and cell shape) were relaxed at each volume.

Density functional theory: spin crossover and metallization. After clarifying structural phase transitions, we identified that the electronic and magnetic transitions occur in $P\overline{3}m1$ phase. Thus, employing the same symmetry, we investigated the SCO behaviors in detail. We further performed electronic-structure calculations within DFT + U
scheme as implemented in Vienna Ab Initio Simulation Package (VASP)\(^\text{37}\). As for layered system, where the van der Waals interactions are important, frequently used generalized gradient approximation (GGA) functionals sometimes fails to predict the correct structural behaviors. We found van der Waals-corrected functions give better description of the ground state volume such that the errors were 1.6% for many-body dispersion and 2.2% for Tkatchenko-Scheffler methods while D3 approach severely underestimates the volume by 8.2%\(^\text{31,34,35}\). From GGA\(^\text{46}\), we found that PBEsol overperforms PBE (3.8% vs. 8.1%) with accuracy similar to van der Waals approach, which enables us to choose PBEsol scheme with safety. Note that in our previous reports, PBEsol successfully explained the spin-state transition behaviors for CoCl\(_2\)\(^\text{31}\). We also carefully tested various \(U\) parameters and found that \(U_{\text{eff}}=3.0\) eV fits best in describing the experimental transition behaviors. To obtain the pressure evolution of the electronic structure and magnetic properties, we fully relaxed the atomic positions until the atomic forces are less than 0.001 eV/Å for each volume point. Once the transition volume is found, we have cross-checked the results employing full potential full relativistic code FPLO\(^\text{69}\), and further analyzed its partial density of states (see Supplementary Figs S6–S9).

Data Availability

All data generated or analysed during this study are included in this published article (and its Supplementary Information files).

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Author Contributions
All authors reviewed the manuscript. F.R. participated in all parts of the project. J.A.B.-A. and I.H. did the optical absorption and J.G. the Raman spectra as a function of pressure. L.N., I.H., W.H. and A.J. conducted the x-ray diffraction measurements and F.A. collected diffraction data and helped with structure refinement. A.O.R. and V.L. did the DFT calculations for structural refinement and phase transitions, and B.K., K.K. and B.I.M. performed the DFT + U electronic structure calculations for spin crossover and metallization processes. Y.J. and C.J. did the electrical measurements at high pressure. All authors participated in the analysis of data and revised the manuscript.

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