Chloride and pseudohalide hydride-carbonyl ruthenium(II) complexes with 4-pyrrolidinopyridine as co-ligand

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Received: 27 September 2012 / Accepted: 31 October 2012 / Published online: 20 November 2012
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Abstract Chloride and pseudohalide (N$_3^-$, NCS$^-$) hydride-carbonyl ruthenium(II) complexes with 4-pyrrolidinopyridine as co-ligand were synthesized and characterized by IR, $^1$H, and $^{31}$P NMR, electronic absorption and emission spectroscopy and X-ray crystallography. The electronic structures of the complexes were calculated by density functional theory (DFT) on their crystal structures. The spin-allowed singlet–singlet electronic transitions of the complexes were calculated by time-dependent DFT, and the UV–Vis spectra have been discussed on these basis. The emission properties of the complexes were also studied.

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Transition Met Chem (2013) 38:133–142 DOI 10.1007/s11243-012-9670-8

Introduction

Pyridine ligands have energetically low-lying $\pi$-antibonding orbitals, which can accept electrons from the occupied $d$ orbitals of metal atoms. Metal complexes with pyridine ligands can exhibit charge transfer bands with interesting spectroscopic properties in the visible region [1]. Hence, ligands containing pyridine rings have been widely studied, and their $\sigma$-donor and $\pi$-acceptor properties are often interesting. Their combination with other donor atoms should in principle afford complexes with tunable spectroscopic properties [2]. 4-Pyrrolidinopyridine (py-4P) is a stronger electron donor N-heteroaromatic ligand compared with pyridine (py-4P pKa = 18.33; pyridine pKa = 12.53) [3]. Hence, 4-pyrrolidinopyridine should be interesting as a ligand, but reports on this topic are rather scarce. The py-4-P ligand has been found to stabilize Zn–Zn bonded complexes [4, 5], five-coordinate zirconium(IV) and titanium(IV) complexes [6, 7] and iridium, hafnium, neodymium, and iron complexes with 4-pyrrolidinopyridine [8–11] have also been reported. Moreover, ruthenium and osmium catalysts containing 4-pyrrolidinopyridine have been claimed in several patents [12–14].

On the other hand, it is known that thiocyanate ligands tune the $t_{2g}$ ruthenium orbitals by distributing the $4d_{Ru}$ energy levels over a wide energy range, due to mixing with orbitals centered on the NCS ligand ($2p_N$, $2p_C$ and $3p_S$) [15]. The calculated density of states showed that both inter- and intramolecular interactions are important and can significantly influence the orbital composition in the frontier electronic structure. The N$_3^-$ ligand, which is similar in properties to thiocyanate, should exhibit comparable characteristics. Thus, studies of the electronic structures of these complexes are an important area of chemistry.

The complexes reported in this paper combine our interest in ruthenium coordination compounds and complexes containing pyridine derivative ligands [16–20]. We describe an experimental and quantum chemical study of ruthenium hydride-carbonyl chloride, isothiocyanate, and azide complexes with 4-pyrrolidinopyridine as co-ligand. As well as the syntheses and spectroscopic ($^1$H, $^{31}$P NMR, IR) characterizations, the X-ray crystal structures and photophysical properties of the complexes are presented. The quantum chemical study included a characterization of the molecular and electronic structures of the complexes by analysis of the optimized molecular geometries and electronic populations using the natural bond orbitals scheme. The latter was also used to identify the nature of the interactions between the ligands and the metal. Finally, time-dependent density functional theory (TD-DFT) was used to calculate and interpret the electronic absorption spectra.
Experimental

All reagents used for the syntheses of the complexes were commercially obtained and were used without further purification. The starting complex [RuHCl(CO)(PPh3)3] was synthesized according to the literature method [21].

Synthesis of [RuHX(CO)(py-4-P)(PPh3)2] (X = Cl; N3; NCS)

The complexes were synthesized by reaction of [RuHCl(CO)(PPh3)3] (0.1 g, 1 × 10⁻⁴ mol), 4-pyrrolidino-pyridine (0.015 g, 1 × 10⁻⁴ mol; py-4-P) (1), and sodium azide (0.007 g, 1 × 10⁻⁴ mol) (2) or ammonium thiocyanate (0.008 g, 1 × 10⁻⁴ mol) (3) in methanol solution (100 cm³). In each case, the mixture was refluxed in methanol for 4 h, then cooled and filtered. Crystals suitable for X-ray crystal analysis were obtained by slow evaporation of the filtrates.

Complex (1) ([RuHCl(CO)(py-4-P)(PPh3)2]): Yield 63 %. IR (KBr; cm⁻¹): 2,058 (w) \(v\) (Ru-H), 1,915 (s) \(v\) (CO); 1,611, 1,528 (m) \(v\) (C=N; C=C). UV–Vis (methanol; log e; nm): 341 (1.58), 276 (3.74), 208 (4.89). ¹H NMR (400 MHz, CDCl3) \(\delta\) (ppm) 8.32 (d, \(J = 7.2\) Hz, py), 8.16 (d, \(J = 7.1\) Hz, py), 7.95–7.15 (m, PPh3), 6.25 (d, \(J = 6.9\) Hz, py), 5.92 (d, \(J = 6.3\) Hz, py), 5.74 (d, \(J = 7.2\) Hz, py), 3.28 (s, pyrrolidine), 3.22 (dd, \(J = 32.4, 5.6\) Hz, pyrrolidine), 2.19 (s, pyrrolidine), −4.45 (t, \(J = 19.4\) Hz, 1H). ³¹P NMR (162 MHz, CDCl3) \(\delta\) (ppm) 39.12 (s, PPh3).

Complex (2) ([RuH(N₃)(CO)(py-4-P)(PPh3)2]): Yield 65 %. IR (KBr; cm⁻¹): 2,052 (w) \(v\) (N₃), 1,939, 1,919 (w, s) \(v\) (Ru-H)/\(v\) (CO); 1,617, 1,572 (m) \(v\) (C=N; C=C); 701 (s) \(\delta\) (N₃). UV–Vis (methanol; log e; nm): 339 (1.92), 274 (3.47), 208 (4.96). ¹H NMR (400 MHz, CDCl3) \(\delta\) (ppm) 7.83–6.85 (m, PPh3, py), 3.11 (d, \(J = 37.4\) Hz, pyrrolidine), 2.19 (s, pyrrolidine), 1.62 (s, pyrrolidine), 1.34 (s, pyrrolidine), −7.17 (dt, \(J = 104.4, 24.8\) Hz, H₉). ³¹P NMR (162 MHz, CDCl3) \(\delta\) (ppm) 40.10 (d, \(J = 15.7\) Hz, PPh3).

Complex (3) ([RuH(NCS)(CO)(py-4-P)(PPh3)2]CH₂OH): Yield 68 %. IR (KBr; cm⁻¹): 2,104 (s) \(v\) (C=O), 1,694 (w, m) \(v\) (Ru-H)/\(v\) (CO); 1,615, 1,585 (m) \(v\) (C=N; C=C); 742, 694 (m) \(v\) (SC from SCN) 519 (m) \(\delta\) (NCS). UV–Vis (methanol; log e; nm): 329 (1.99), 2,589 (3.62), 211 (4.92). ¹H NMR (400 MHz, CDCl3) \(\delta\) (ppm) 7.84–7.00 (m, py, PPh3), 6.96 (t, \(J = 8.4\) Hz, py), 3.04 (d, \(J = 6.7\) Hz, pyrrolidine), 2.01 (d, \(J = 13.3\) Hz, pyrrolidine), 1.63 (d, \(J = 12.1\) Hz, pyrrolidine), 1.26 (d, \(J = 11.1\) Hz, pyrrolidine), −7.18 (dt, \(J = 100.0, 24.4\) Hz, H₉). ³¹P NMR (162 MHz, CDCl3) \(\delta\) (ppm) 39.40 (d, \(J = 15.4\) Hz).

Physical measurements

Infrared spectra were recorded on a Perkin–Elmer spectrophotometer in the range 4,000–450 cm⁻¹ using KBr pellets. Electronic spectra were measured on a Lab Alliance UV–VIS 8500 spectrophotometer in the range of 600–180 nm in methanol solution. The ¹H and ³¹P NMR spectra were obtained at room temperature in CDCl₃ using a Bruker 400 MHz spectrometer. Luminescence measurements were taken in methanol solutions on an F-2500 FL spectrophotometer at room temperature.

Computational methods

The calculations were made using the Gaussian 09 [22] program. Molecular geometries of the singlet ground state of complexes (1), (2), and (3) were fully optimized in the gas phase at the B3LYP level of theory. [23, 24] For each complex, a frequency calculation was made, verifying that the optimized molecular structure corresponded to an energy minimum; thus, only positive frequencies were found. The DZVP basis set [25] with f functions with exponents 1.94722036 and 0.748930908 was used to describe the ruthenium atom, and the basis set used for the lighter atoms (C, N, O, P, H) was 6-31G with a set of d and p polarization functions. The TD-DFT method [26] was employed to calculate the electronic absorption spectra of the complexes using the solvent polarizable continuum model (PCM). In this work, 100 singlet excited states were calculated as vertical transitions for each complex. A natural bond orbital (NBO) analysis was also made for each of the complexes, using the NBO 5.0 package [27] included in Gaussian 09. Natural bond orbitals are orbitals localized on one or two atomic centers that describe molecular bonding in a manner similar to a Lewis electron pair structure, and they correspond to an orthonormal set of localized orbitals of maximum occupancy. NBO analysis provides the contribution of the atomic orbitals (s, p, d) to the NBO σ and π hybrid orbitals for bonded atom pairs. In this scheme, three NBO hybrid orbitals are defined, namely bonding orbital (BD), lone pair (LP), and core (CR), which were analyzed for the atoms directly bonded to, or presenting some kind of interaction with, the ruthenium atom. The contribution of a group (ligands, metal center) to a molecular orbital was calculated using Mulliken population analysis. GaussSum 2.2 [28] was used to calculate group contributions to the molecular orbitals and to prepare the partial density of states (DOS) spectra. The DOS spectra were created by convoluting the molecular orbital information with Gaussian curves of unit height and FWHM (full width at half maximum) of 0.3 eV.

Crystal structure determination and refinement

The crystals of [RuHCl(CO)(py-4-P)(PPh3)2] (1), [RuH(N₃)-(CO)(py-4-P)(PPh3)2] (2), and [RuH(NCS)(CO)(py-4-P)(PPh3)2] (3) were mounted in turn on an Xcalibur, Atlas, Gemini Ultra Oxford Diffraction automatic diffractometer.
eqipped with a CCD detector, and used for data collection. X-ray intensity data were collected with graphite monochro-
mated MoKα radiation (λ = 0.71073 Å) at a temperature of
295.0(2) K, with ω scan mode. Ewald sphere reflections were
collected up to 2θ 50.10. The unit cell parameters were
determined from least-squares refinement of the setting angles
of 6,947, 15,306, and 10,064 strongest reflections for com-
plexes (1), (2), and (3), respectively. Details concerning
crystal data and refinement are gathered in Table 1. Lorentz,
polarization, and empirical absorption corrections using
spherical harmonics implemented in the SCALET3 ABSPACK
scaling algorithm [29] were applied. The structures were
solved by the Patterson method and subsequently completed
by difference Fourier recycling. All the non-hydrogen atoms
were refined anisotropically using full-matrix, least-squares
techniques. Bearing in mind the limits of Fourier synthesis and
the problems in recognizing artifacts in the immediate
neighborhood of heavy atoms, it is doubtful if a reliable
position for the hydrogen atom bound to the Ru atom can be
found in the difference Fourier map while avoiding the danger
of mistaking the effects of the series termination errors for a
true atomic position. In the studied complexes, the Ru–H bond
length of 1.50 Å is normal. The Olex2 [30] and SHELXS97,
SHELXL97 [31] programs were used for all the calculations.
Atomic scattering factors were incorporated in the computer
programs.

Results and discussion

Spectroscopic characterization

In the 1H NMR spectra of the complexes, as well as signals
corresponding to the PPh3 and 4-pyridilidopyridine
ligands, there are signals at high field indicating the presence
of the hydride ligands. The chemical shifts of these signals
are due to the shielding effect of the metal and to the partial
charge of the hydrogen atom. The Ru–H signals are observed
as a triplet in (1) and doublet of triplets in (2) and (3) with JHP
~ 100 and 20 Hz. Even if (2) and (3) were asymmetric with
inequivalent phosphines, a doublet of doublets would have
been expected, but the asymmetry could be partially
removed on the NMR timescale. The signals are observed at
−4.45, −7.17, and −7.18 ppm for complexes (1), (2), and
(3), respectively, and the differences are connected with the
increasing π-acceptor properties in the chloride, nitride,
and isothiocyanate ligands. The 31P NMR spectra of all three
complexes show signals close to 40 ppm. The signals are
doubled in the case of complexes (2) and (3) which suggests
two triphenylphosphine groups, not in perfect trans
positions. In the 31P spectrum of complex (1), the observed singlet
may be caused by electronic interactions (π–π stacking)
between PPh3 phenyl and pyridine rings.

The IR spectra of the complexes show strong bands at
1,934–2,004 and 1,915–1,944 cm−1, assigned to the Ru–H
and C≡O stretching bands (see the selected IR frequencies
given in Experimental section). The differences in the
maxima of these bands are connected with the different
(pseudo)halides present in the coordination sphere. The
electron–donor hydride ligand delivers electron density via
backbonding to the antibonding orbitals of the CO,
resulting in a decrease in the frequency of the CO vibra-
tion. However, the acceptor properties of the (pseudo)halide
ligands vary as Cl− < N3− < NCS− and the positions of the νRu–H and νCO bands in the azide complex (2)
suggest that this pseudohalide anion exerts a much weaker
effect than isothiocyanate in these complexes. This is
supported by the theoretically determined charge values
which indicate charges on the Ru(II) centers of −0.905,
−0.844, and −0.855 in (1), (2) and (3), respectively. The
charges on the hydride ligands are close to zero, being
0.074, 0.036 and 0.021 in complexes (1), (2), and (3),
respectively. The charges on the CO ligands, calculated by
summing the individual charges on the carbon and oxygen
atoms, are 0.213 (1), 0.198 (2), and 0.229 (3). Hence, in the
complex (2), the charges on ruthenium and the carbonyl
ligand are the smallest, and this is in accordance with the
largest decrease in CO vibration frequency for this com-
plex. The natural charges on the chloride, azide, and iso-
thiocyanate ligands of −0.553, −0.602, and −0.663,
respectively, are in accordance with their acceptor
properties.

The stretching vibrations for N3− and S=C≡N− are
observed at 2,052 and 2,104 cm−1, respectively. The coor-
dination mode of thiocyanate in complex (3) cannot be
determined from the IR spectrum. For N-bonded complexes,
generally the C–N stretching band is in a lower region,
around 2,050 cm−1, compared with 2,100 cm−1 for S-bonded
complexes. However, the frequencies of the bands are sen-
sitive to other factors such as co-ligands; hence, the struc-
tures of these complexes were determined by X-ray analysis.
While the M–S–C angles of S-bonded thiocyanato ligands in
such complexes are bent at around 110°, the M–N–C angles
of N-bonded isothiocyanato ligands are the smallest, and this is in accordance with the
largest decrease in CO vibration frequency for this com-
plex. The natural charges on the chloride, azide, and iso-

thiocyanate ligands of −0.553, −0.602, and −0.663,
respectively, are in accordance with their acceptor
properties.

Molecular structures

Crystals of the complexes suitable for single crystal X-ray
analyses were obtained by slow evaporation of the reaction
mixtures. Complexes (1) and (3) crystallize in monoclinic
P21/c and P21/n space groups and (2) in triclinic P-1. The
azide complex (2) has two independent molecules in the
asymmetric unit, while complex (3) crystallizes as a sol-
vate with one methanol molecule. The molecular structures
of the complexes are displayed as ORTEP representations in Fig. 1, and selected bond distances and angles are collected in Table 2. The Ru(1)–N(1) bond lengths in the complexes are normal and comparable with other ruthenium hydride complexes with pyridine derivative ligands [17, 18].

The structures of all three complexes can be considered as distorted octahedral, with the largest deviation from the expected 90° bond angles for N(1)–Ru(1)–H(1), equal to 82.0(9)° in (1) and 84.5(10)° (average value) in (2), and P(1)–Ru(1)–H(1) (83.7(9)°) in complex (3). The angles between carbonyl C(1) and the chloride or pseudohalide (N₅⁻, NCS⁻) ligands differ by about 7° from a right angle. The P–Ru–P angles are lower than 180°, being in the 168.23(2)°–171.16(2)° range. As shown in Fig. 2, the CO groups are trans to the 4-pyrollidinopyridine ligands, and the halide and hydride ligands are mutually trans disposed. In the parent complexes with general formula [RuHX(CO)(PPh₃)₃] where X = Cl⁻, N₅⁻, NCS⁻, the halide ligands are trans to the carbonyl, and the hydride and one PPh₃ ligand are also mutually trans disposed [32, 33]. In the complexes with 4-pyrollidinopyridine, the trans position to the X ligand is occupied by hydride, whilst the carbonyl is located opposite to the py-4-P ligand. The Ru–X bonds lengths in the complexes are longer by about 0.04, 0.097, 0.08 Å, and the Ru–CO bonds shorter by 0.015, 0.031, 0.014 Å in complexes (1), (2), and (3), respectively, compared with the [RuHX(CO)(PPh₃)₃] complexes. Moreover, the C≡O distances in the carbonyl ligands are longer than in the corresponding [RuHX(CO)(PPh₃)₃] complexes; in the chloride complex (1), the C≡O bond length is longer by 0.015 Å, in (2) by 0.07 Å and in (3) by 0.06 Å. Hence, the CO bond...
length is minimally elongated in the case of chloride complex (1), which is connected with the weak acceptor property of Cl\(^-\) compared with pseudohalide ligands. The ruthenium–py-4-P bond distances are similar in the chloride and azide complexes (1) and (2), being close to 2.18 Å and in isothiocyanate complex (3) the distance is shorter by about 0.02 Å. Similarly, the Ru–N(3) distance in complex (3) is shorter by 0.04 Å than in complex (2). The Ru(1)–C(1) distances are similar, and the differences are more visible in the C\(\equiv\)O distances. The shortest carbonyl bond length is in the azide complex (2) (Table 2). In the molecular structures of the complexes several inter- and intra-molecular hydrogen bonds [34] exist and are collected in Table 3. Additionally, some \(\pi\)--\(\pi\) stacking between the PPh\(_3\) phenyl and pyridine rings is also visible. The plane-to-plane distances between the phosphine phenyl centroids, determined by C(11)–C(16) in (1), C(41)–C(46) and C(75)–C(80) in (2), and C(41)–C(46) in (3), and the pyridine rings are 3.595, 3.62 Å.
(average value) and 3.764 Å, indicating weak π–π stacking interactions. Moreover, in the structure of complex (2), a T-shaped C–H···π stacking interaction is visible between the two molecules in the asymmetric unit, involving the C(17)–C(22) and C(81)–C(86) phenyl rings with a distance of 2.829 Å.

Quantum calculations

The ground states geometries of the complexes were optimized in singlet states using the DFT method with the B3LYP functional. The calculations were made for gas phase molecules [without the solvent molecule of complex (3)], and in general, the predicted bond lengths and angles are over-estimated by about 0.1 Å and 5°, respectively. Nevertheless, the general trends observed in the experimental data are reproduced in the calculations, as can be seen from the data collected in Table 2. The calculated IR frequencies of the complexes show good agreement with the experimental spectra; the differences can be explained by the neglect of intermolecular interactions for the gas phase. From the data collected in Table 2, the major

| Bond lengths (Å) | (1) | (2) | (3) |
|------------------|-----|-----|-----|
|                  | Exp | Calc| Exp | Calc | Exp | Calc |
| Ru(1)–C(1)      | 1.822(3) | 1.86 | 1.826(3) | 1.86 | 1.815(3) | 1.86 |
| Ru(1)–N(1)      | 2.1797(19) | 2.25 | 2.176(2) | 2.186(2) | 2.25 | 2.156(2) | 2.24 |
| Ru(1)–N(3)      | 2.231(2) | 2.219(2) | 2.23 | 2.177(2) | 2.18 |
| Ru(1)–P(1)      | 2.346(6) | 2.43 | 2.358(7) | 2.3699(7) | 2.43 | 2.346(7) | 2.44 |
| Ru(1)–P(2)      | 2.363(6) | 2.43 | 2.351(7) | 2.343(7) | 2.43 | 2.368(7) | 2.44 |
| Ru(1)–Cl(1)     | 2.541(6) | 2.62 | 1.482(2) | 1.60 | 1.51(3) | 1.58(2) | 1.62 | 1.56(2) | 1.63 |
| C(1)–O(1)       | 1.156(3) | 1.16 | 1.150(3) | 1.155(3) | 1.16 | 1.161(3) | 1.16 |
| C(1)–Ru(1)–N(1) | 173.35(9) | 171.9 | 173.37(10) | 171.13(10) | 173.7 | 174.76(11) | 176.3 |
| C(1)–Ru(1)–N(3) | 97.64(11) | 103.94(10) | 98.6 | 96.95(12) | 96.5 |
| C(1)–Ru(1)–P(1) | 89.92(8) | 88.4 | 89.01(8) | 88.59(9) | 88.1 | 89.84(9) | 88.9 |
| N(1)–Ru(1)–P(1) | 91.04(5) | 90.9 | 89.44(5) | 89.56(6) | 91.1 | 90.90(6) | 90.6 |
| N(3)–Ru(1)–P(1) | 91.76(7) | 90.4 | 90.26(8) | 88.18(9) | 88.3 | 87.13(9) | 88.3 |
| C(1)–Ru(1)–P(2) | 89.45(8) | 89.4 | 89.02(8) | 85.1(9) | 87.3 | 91.98(6) | 91.6 |
| N(1)–Ru(1)–P(2) | 88.26(5) | 90.4 | 90.28(5) | 92.10(6) | 91.9 | 91.98(6) | 91.6 |
| N(3)–Ru(1)–P(2) | 97.07(7) | 90.4 | 97.04(7) | 94.51(7) | 93.8 | 91.31(6) | 91.8 |
| P(1)–Ru(1)–P(2) | 168.23(2) | 173.5 | 171.16(2) | 169.41(2) | 174.5 | 170.44(2) | 171.5 |
| C(1)–Ru(1)–Cl(1) | 95.37(8) | 98.5 | 96.94(9) | 95.9(10) | 98.3 | 96.8(9) | 97.9 |
| N(1)–Ru(1)–Cl(1) | 90.09(5) | 89.6 | 90.04(5) | 88.18(9) | 88.3 | 87.1(9) | 88.3 |
| P(1)–Ru(1)–Cl(1) | 95.58(2) | 93.9 | 95.78(2) | 94.5 | 95.78(2) | 94.5 |
| P(2)–Ru(1)–Cl(1) | 96.17(2) | 94.5 | 96.17(2) | 94.5 | 96.17(2) | 94.5 |
| C(1)–Ru(1)–H(1) | 91.6(9) | 86.7 | 89.0(10) | 86.6(9) | 88.3 | 86.8(9) | 89.7 |
| N(1)–Ru(1)–H(1) | 82.0(9) | 85.2 | 84.4(10) | 84.6(9) | 85.4 | 88.19(9) | 86.6 |
| N(3)–Ru(1)–H(1) | 173.0(10) | 169.4(9) | 172.8 | 176.19(9) | 173.8 |
| P(1)–Ru(1)–H(1) | 82.8(9) | 85.6 | 86.2(9) | 85.1(9) | 87.1 | 83.7(9) | 84.3 |
| P(2)–Ru(1)–H(1) | 85.5(9) | 88.1 | 85.0(9) | 84.6(9) | 88.5 | 87.19(9) | 87.6 |
| Cl(1)–Ru(1)–H(1) | 171.8(9) | 174.8 | 171.86(9) | 174.8 | 171.86(9) | 174.8 |
| N(1)–Ru(1)–N(3) | 88.86(9) | 84.88(8) | 87.6 | 88.10(9) | 87.2 |
| Ru(1)–C(1)–O(1) | 176.7(2) | 175.9 | 177.3(3) | 175.1(2) | 171.1 | 177.5(3) | 179.6 |
| Ru(1)–N(3)–N(4) | 128.3(2) | 130.7(2) | 124.7 | 128.3(2) | 130.7(2) | 124.7 |
| Ru(1)–N(3)–C(47) | 174.7(2) | 176.7 | 174.7(2) | 176.7 | 174.7(2) | 176.7 |
| N(3)–N(4)–N(5) | 173.7(4) | 177.3(3) | 177.5 | 173.7(4) | 177.3(3) | 177.5 |
| N(3)–C(47)–S(1) | 179.7(3) | 179.4 | 179.7(3) | 179.4 | 179.7(3) | 179.4 |
distances (0.084 and 0.094 Å, respectively) in complex (1). Differences between the experimental and calculated geometries are found in the Ru(1)–N(1) and Ru(1)–P(1) distances (0.084 and 0.094 Å, respectively) in complex (3). In the case of chloride complex (1), the experimental and calculated Ru(1)–H(1) distances differ by about 0.12 Å. Based on the optimized geometries of the complexes, NBO analyses were performed in order to reveal the nature of the coordination between ruthenium and the donor atoms of the ligands. These analyses showed that the bonding between the py-4-P ligand and ruthenium is largely non-covalent; the Coulomb-type interaction between the ruthenium center and py-4-P ligand is clearly visible in the calculated Wiberg bond index, which is considerably lower than one and close to 0.4 (similar in all these complexes). The Ru–P bond orders are also smaller than 1 (0.7).

For the carbonyl ligands, three natural bond orbitals were detected for the C–O bond, and one for the Ru–C bond. The Ru–C bond orbitals are polarized toward the carbon atom, and the C–O bond orbitals are polarized toward oxygen. The oxygen atom of the carbonyl ligand has one lone pair (LP) orbital. The occupancies and hybridization of the Ru–H, Ru–C, and CO bonds are gathered in Table 4 (antibonding NBOs are given in round brackets). The Wiberg indexes of the CO bonds in the complexes are reduced (by about 0.2) with respect to free CO (W_{CO} = 2.23). The maximum reduction of Wiberg index is calculated for complex (2) which is consistent with the lowest charge on the carbonyl ligand as well as the relatively short C=O bond in the azide complex.

Analysis of the frontier molecular orbitals is useful for understanding the spectroscopic properties such as electronic absorption and emission spectra. The electronic structures of the complexes are similar because of their similar compositions. The density of states (DOS) in terms of Mulliken population analysis were calculated using the GaussSum program, and Fig. 2 presents the composition of the fragment orbitals contributing to the molecular orbitals for the complexes along with the N_3^- and NCS^- participations in the HOMOs. The HOMO of complex (1) is localized mainly on the ruthenium atom (54 %), with a contribution from the chloride ligand (36 % Cl). In complexes (2) and (3), the HOMOs are shifted to higher energy and composed of pseudohalide π orbitals (~80 %) with antibonding participation of ruthenium d orbitals (~17 %). This change in ordering of the molecular orbitals influences the luminescent properties of the complexes. The LUMOs are composed of π* orbitals on the PPh_3 ligands (~80 %) with a contribution from the ruthenium d^2 orbital (~16 %). The py-4-P ligand plays a role in higher virtual orbitals (LUMO+3/ +4 ~70 %). The HOMO-2 and HOMO-4 show antibonding interactions involving the π orbitals of 4-pyrrolidino-pyridine and carbonyl ligands with ruthenium d orbitals.

Experimental and theoretical electronic spectra

The UV–Vis spectra of the complexes are similar and present two bands with maxima in the range 300–250 nm. A third high-energy band close to 210 nm may result from transitions in the PPh_3 ligands and/or from π → π* transitions.
Table 4 The occupancies and hybridization of the calculated R–H, Ru–C and C≡O natural bond orbitals (NBOs) of [RuHCl(CO)(py-4-P)(PPh3)2] (1), [RuH(NCS)(CO)(py-4-P)(PPh3)2] (2) and [RuH(NCS)(CO)(py-4-P)(PPh3)2] (3) complexes

| BD (2-center bond) | Occupancy | Hybridization of NBO | Wiberg bond indices |
|-------------------|-----------|----------------------|---------------------|
| Ru–H              | 1         | 1.858(0.120)         | 0.734(sp^0.60d^2.59)_{Ru} + 0.679(s)_{H} | 0.79 |
|                   | 2         | 1.710(0.061)         | 0.674(sp^0.01d^1.90)_{Ru} + 0.739(s)_{H} | 0.79 |
|                   | 3         | 1.862(0.119)         | 0.713(sp^0.83d^2.64)_{Ru} + 0.701(s)_{H} | 0.79 |
| Ru–C              | 1         | 1.941(0.148)         | 0.578(sp^0.81d^2.47)_{Ru} + 0.816(sp^0.50)_{C} | 1.30 |
|                   | 2         | 1.931(0.134)         | 0.574(sp^0.67d^1.98)_{Ru} + 0.819(sp^0.40)_{C} | 1.32 |
|                   | 3         | 1.943(0.142)         | 0.581(sp^0.86d^2.61)_{Ru} + 0.814(sp^0.45)_{C} | 1.30 |
| C≡O               | 1         | 1.997(0.231)         | 0.489(p)_{C} + 0.872(p)_{O} | 2.04 |
|                   | 2         | 1.996(0.210)         | 0.494(p)_{C} + 0.870(p)_{O} | 2.02 |
|                   | 3         | 1.994(0.010)         | 0.556(sp^0.03)_{C} + 0.832(p^1.13)_{O} | 2.05 |

Table 5 Selected calculated electronic transitions for [RuHX(CO)(py-4-P)(PPh3)2] (X = Cl, NCS, N3) complexes

| (nm)   | f   | Major contributions | Character |
|--------|-----|---------------------|-----------|
| (1)    |     |                     |           |
| 344.0  | 0.0243 | HOMO → LUMO (54 %), HOMO → L + 1 (14 %) | d_{Ru}/π_{C3} → d_{Ru}/π^*_{ppPh3} |
| 319.1  | 0.0310 | H-2 → LUMO (31 %), H-1 → LUMO (33 %) | d_{Ru}/π_{C3}/π_{py-4-P} → d_{Ru}/π^*_{ppPh3} |
| 315.1  | 0.0357 | H-2 → LUMO (31 %), H-1 → LUMO (22 %) | d_{Ru}/π_{C3}/π_{py-4-P} → d_{Ru}/π^*_{ppPh3} |
| 311.9  | 0.0300 | HOMO → L + 2 (15 %), HOMO → L + 3 (11 %), HOMO → L + 10 (15 %), HOMO → L + 13 (10 %) | d_{Ru}/π_{C3}/π_{py-4-P} → d_{Ru}/π^*_{ppPh3}/π^*_{py-4-P} |
| 305.2  | 0.0009 | HOMO → LUMO (16 %), HOMO → L + 1 (67 %), HOMO → L + 2 (10 %) | d_{Ru}/π_{C1} → d_{Ru}/π^*_{ppPh3} |
| 300.2  | 0.0046 | HOMO → L + 2 (63 %), HOMO → L + 3 (16 %) | d_{Ru}/π_{C1} → π^*_{ppPh3}/π^*_{py-4-P} |
| (2)    |     |                     |           |
| 380.07 | 0.0302 | HOMO → LUMO (76 %) | d_{Ru}/π_{N3} → d_{Ru}/π^*_{ppPh3} |
| 343.7  | 0.0157 | H-1 → LUMO (22 %), HOMO → LUMO (13 %), HOMO → L + 1 (25 %), HOMO → L + 3 (10 %) | d_{Ru}/π_{N3} → d_{Ru}/π^*_{ppPh3}/π^*_{py-4-P} |
| 342.1  | 0.0066 | H-1 → LUMO (53 %), HOMO → L + 1 (11 %), HOMO → L + 3 (11 %) | d_{Ru}/π_{N3} → d_{Ru}/π^*_{ppPh3}/π^*_{py-4-P} |
| 332.4  | 0.0011 | HOMO → L + 2 (87 %) | d_{Ru}/π_{N3} → π^*_{ppPh3} |
| 326.6  | 0.0031 | HOMO → L + 1 (48 %), HOMO → L + 3 (41 %) | d_{Ru}/π_{N3} → π^*_{ppPh3}/π^*_{py-4-P} |
| 317.4  | 0.0566 | H-2 → LUMO (63 %) | d_{Ru}/π_{py-4-P} → d_{Ru}/π^*_{ppPh3} |
| 314.4  | 0.0474 | H-2 → LUMO (14 %), H-1 → L + 1 (46 %) | d_{Ru}/π_{py-4-P} → d_{Ru}/π^*_{ppPh3} |
| (3)    |     |                     |           |
| 356.6  | 0.0448 | HOMO → LUMO (76 %) | d_{Ru}/π_{NCS} → d_{Ru}/π^*_{ppPh3} |
| 321.3  | 0.0085 | H-1 → LUMO (78 %) | π_{NCS} → d_{Ru}/π^*_{ppPh3} |
| 318.0  | 0.0046 | H-2 → LUMO (22 %), HOMO → L + 1 (26 %), HOMO → L + 3 (19 %) | d_{Ru}/π_{py-4-P}/π_{NCS} → d_{Ru}/π^*_{ppPh3}/π^*_{py-4-P} |
| 313.9  | 0.0464 | H-2 → LUMO (45 %), HOMO → L + 1 (32 %) | d_{Ru}/π_{ppPh3}/π_{py-4-P} → d_{Ru}/π^*_{ppPh3} |
| 307.1  | 0.0288 | HOMO → L + 1 (34 %), HOMO → L + 3 (43 %) | d_{Ru}/π_{ppPh3}/π_{py-4-P} → d_{Ru}/π^*_{ppPh3}/π^*_{py-4-P} |

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excitations in the py-4-P type ligands. The electronic spectra of the complexes were calculated with the TD-DFT method with methanol as solvent in the polarizable continuum model (PCM). Table 5 shows the calculated electronic transitions for the complexes; only transitions to 300 nm are included so the character of the first band is presented. The lowest energy bands in the UV–Vis spectra have Metal–Ligand Charge Transfer (MLCT) with admixture of ligand field character. The frontier orbitals HOMO, HOMO-1, and HOMO-2 plus LUMO, LUMO+1 to LUMO+3 are engaged in transitions. These molecular orbitals are constructed from $d$ ruthenium and $\pi$ halide and py-4-P orbitals. The next bands in the vicinity of 260–270 nm have mixed MLCT and LMCT character with admixture of Ligand-to-Ligand Charge Transfer (LLCT) transitions.

The emission characteristics of the complexes have been examined in methanol solutions (concentration of $1 \times 10^{-3}$ mol dm$^{-3}$) at room temperature, as shown in Fig. 3. The solutions of the complexes excited at 327, 333, and 322 nm for complexes (1), (2), and (3), respectively, gave emissions with maxima at 381, 376, and 441 nm, respectively. The solution of isothiocyanate complex (3) when excited at 361 nm also results in emission at 441 nm. The red shifts of the emissions maxima are typical of ruthenium(II) complexes, and the emissions originating from the MLCT states are derived from excitation involving $d_e \rightarrow \pi^*_{\text{ligand}}$ transitions. The assignments are supported by the analysis of the frontier orbitals of the corresponding complexes, which reveal the contributions of the ligands. In Table 5, the transitions near the excitation wavelengths are marked in italics. The fluorescence of complex (1) is connected with $d_{\text{Ru}}/\pi_{\text{Cl}}/\pi_{\text{py-4-P}} \rightarrow d_{\text{Ru}}/\pi^*_{\text{PPh3}}$ transitions that show considerable participation of the triphenylphosphine ligand. For this reason, the fluorescence of this complex has the lowest intensity. In the pseudohalide complexes (2) and (3), the 4-pyrrolidinopyridine ligands play a role in the emission processes which is possible due to the effect of the $\text{N}_3^-$ and $\text{NCS}^-$ ligands on the electronic structure (energies and compositions) of the frontier molecular orbitals.

**Conclusion**

In summary, three new ruthenium(II) complexes with 4-pyrrolidinopyridine ligands were synthesized and characterized by spectroscopy and X-ray crystallography. The crystal structures of the complexes reveal noncovalent interactions between the aromatic rings. The theoretical results obtained from NBO and analysis of the interactions between ruthenium and the pyridine derivative, and carbonyl and hydride ligands were used to explain the differences in bond lengths as well as the differences in the IR band positions of the complexes. Additionally, comparison of the carbonyl band position in the spectrum of complex (1) with similar ruthenium(II) hydride-carbonyl complexes with 4-phenylpyridimide or pyridine ligands [18, 35] confirms the strong $\sigma$-donor property of 4-pyrrolidinopyridine. Electronic structures of the complexes characterized in particular by density of states diagrams have been correlated with their fluorescence properties.

**Supplementary data**

CCDC 879949, CCDC 879950 and CCDC 879951 contain the supplementary crystallographic data for [RuHCl(CO)(py-4-P)(PPh3)$_2$] (1), [RuH(N$_3$)(CO)(py-4-P)(PPh3)$_2$] (2) and [RuH(NCS)(CO)(py-4-P)(PPh3)$_2$]CH$_3$OH (3) complexes respectively. These data can be obtained free of charge from http://www.ccdc.cam.ac.uk/conts/retrieving.html, or from the Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: (+44) 1223-336-033; or e-mail: deposit@ccdc.cam.ac.uk.

**Acknowledgments** Calculations have been made in Wroclaw Centre for Networking and Supercomputing (http://www.wcss.wroc.pl).

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