

Spectroscopic (IR, Raman) and thermogravimetric studies of 3d-metal cinchomeronates and dinicotinates

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Abstract In this work, the thermal and spectroscopic properties of 3d-metal (Co, Cu, Zn, Mn, Ni, Fe) and sodium salts with cinchomeronic acid (3,4-pyridinedicarboxylic acid) (3,4pda) and dinicotinic acid (3,5-pyridinedicarboxylic acid) (3,5pda) were studied. The IR and Raman spectra were registered and analyzed in the range of 400–4000 cm⁻¹. The thermogravimetric study of synthesized compounds was done. The degree of hydration and the products of thermal decomposition were estimated. Moreover, it was done the comparison between the effect of 3d-transition metal cations and sodium cation on the degree of perturbation or stabilization of the electronic system of pyridine molecule depending on the position of carboxylic group in the ring.

Keywords Nicotinic acid and derivatives · Cinchomeronic acid · Dinicotinic acid · Metal complexes · Spectroscopy (IR Raman)

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Introduction

Pyridinecarboxylic acids (picolinic, nicotinic and isonicotinic) are biologically important ligands incorporated into some enzymes, and they are active agents in some drugs as well [1–18]. In our previous work, we examined the effect of different metals (alkali [19–21], alkaline earth [22], 3dmetals [23]) on the electronic system of pyridinecarboxylic acids. Research shows that alkali and heavy toxic metals with low ionic potential disturb the aromatic system, while the 3d-transition metals stabilize the aromatic system pyridinecarboxylic acids.

This work is a continuation of our investigations on the metals effect on the electronic structure of pyridinecarboxylic acid. The study of metal complexes with nicotinic acid and cinchomeronic acid has shown that these complexes can have different types of metal-ligand coordination [24–26]. Whitfield et al. [24] studied the complexes of cobalt and nickel with 3,5-pyridinedicarboxylic acid. The complex of cobalt (CoL·2H₂O) is monodentate binding. However, in the nickel complex (NiL·H₂O), there are two types of bonds: monodentate and bidentate chelating. Łyszczek and Mazur studied the structures and thermal properties of lanthanide dinicotinates [25]. Studies have shown that the lanthanides form the types of coordination, such as monodentate, bidentate bridging and chelating bidentate. In the structure of calcium dinicotinate, there are two types of bonds-monodentate and bidentate chelating Starosta et al. [26]. In the structure of manganese cinchomeronate, there are two types of bonds-monodentate and bidentate bridging Tong et al. [27].

The specific objectives of the present study (along with the purpose of implementing the above-mentioned broader theme) are to:

- 1. determine the composition studied complexes on the basis of elemental analysis and thermogravimetric data
- propose the structure of the compounds on the basis of FT-IR and Raman spectral data
- 3. compare the change in electronic charge distribution as a result of complex formation.

In the present work, the impact of 3d-metals on the electron system of pyridinedicarboxylic acids (dinicotinic and cinchomeronic acids) (Fig. 1) was examined and thermal properties of metal complexes of these acids were determined.

Experimental

Sample preparations

3,4-pyridinedicarboxylates (3,4-pdc) and 3,5-pyridinedicarboxylates (3,5-pdc) of 3-d metal ions were prepared in water solution by adding disodium salts of each ligand to appropriate amount of 3-d metal chlorides at 80 °C with continuous stirring for 2 h (molar ratio of sodium salts of ligands to the metal chlorides was 1:1 in case of divalent metal ions and 1.5:1 in case of trivalent metal). The mixtures were left for 24 h at room temperature. The precipitated solids were filtered off and washed with water to eliminate chloride ions. Disodium salts were obtained as microcrystals by adding the of sodium hydroxide solution (0.1 M) to the stoichiometric amount of acid (molar ratio of ligand to the sodium hydroxide: 1:2). Salts were slowly evaporated at room temperature. After precipitation, the salt microcrystals were filtered and washed with a small amount of deionized water. Purity of the salts and complexes was confirmed by elemental analysis and IR spectra analysis. Substances were dried at 30 °C under atmospheric pressure.

Methods

The FT-IR spectra were recorded with an Alfa (Bruker) spectrometer within the range of $400-4000 \text{ cm}^{-1}$. The samples in the solid state were measured in KBr matrix pellets. FT-Raman spectra of solid samples were recorded

Fig. 1 Structures of studied ligands

in the range of 400–4000 cm⁻¹ with a MultiRam (Bruker) spectrometer. Thermogravimetric analyses were conducted on the Setsys 16/18 analyzer in dynamic air atmosphere. Therefore, 6.0–8.0-mg samples were heated in the range of 30–800 °C in the ceramic crucibles using the heating rate of 10 °C min⁻¹. The TG, DTG and DSC curves were registered. The elementary analysis was carried out using the CHN 2400 PerkinElmer Analyzer. The water content was determined from the thermogravimetric curves.

Results and discussion

Elemental analysis

The composition of the synthesized complexes was determined by elemental analysis. The data are gathered in Tables 1 and 2. All the complexes prepared in accordance with the procedure presented in experimental section were hydrated. It was confirmed by thermal analysis. Complexes of manganese and zinc as well as sodium salts were white similarly as the ligands from which they were synthesized. Other complexes were colorful. The obtained complexes were poorly soluble in water. Synthesis was carried out several times to give a reaction yield of 60–80 %. The reaction yields for the different complexes are given in Tables 1 and 2.

Thermogravimetric analysis

The complexes of 3,4-pyridinedicarboxylic and 3,5pyridinedicarboxylic acids with transition metal ions have the general formula: $M[(C_5NH_3(COO)_2] \cdot nH_2O$ where M = Mn, Co, Ni, Cu, Zn(II) ions, n = 1-3 and Fe₂[(C₅. NH₃(COO)₂]₃·nH₂O where n = 4 or 6. The complexes of both series are obtained as hydrated, crystalline well-defined compounds. The simultaneous TG/DTG/DSC curves in dynamic dry air atmosphere of all compounds are shown in Figs. 2–9. The analytical and thermoanalytical data for the synthesized compounds are given in Tables 1–4, respectively (Figs. 2–9). These results indicate the stoichiometry of two series of compounds, which are consistent with their general formulae.



Table 1 Elemental analysis, analytical and physical data for 3,4-pyridinedicarboxylates

Compound	Color	Yield/%	Content C/%		Content H/%		Content	N/%	Content M/%	
			Exp.	Calc.	Exp.	Calc.	Exp.	Calc.	Exp.	Calc.
Na ₂ L·H ₂ O	White	70–75	36.18	36.66	2.14	2.18	6.02	6.11	19.78	20.08
MnL· H ₂ O	White	70	35.16	35.29	2.21	2.10	5.69	5.88	22.33	23.08
Fe ₂ L ₃ ·6H ₂ O	Brown	65-70	35.10	35.15	2.07	2.09	5.97	6.11	16.50	16.89
CoL·1.5H ₂ O	Violet	65-70	33.78	33.46	2.28	2.39	5.64	5.58	23.72	23.47
NiL·3H ₂ O	Green	60–65	32.06	32.33	2.60	2.69	5.48	5.39	22.97	22.59
CuL·H ₂ O	Blue	65-70	33.95	34.05	1.98	2.03	5.61	5.68	25.44	25.76
ZnL·H ₂ O	White	75–80	33.49	33.80	2.13	2.01	5.47	5.63	25.56	26.31

Table 2 Elemental analysis, analytical and physical data for 3,5-pyridinedicarboxylates

Compound	Color	Yield/%	Content C/%		Content H/%		Content N/%		Content M/%	
			Exp.	Calc.	Exp.	Calc.	Exp.	Calc.	Exp.	Calc.
Na ₂ L·H ₂ O	White	70–75	36.44	36.66	1.93	2.18	5.88	6.11	19.32	20.08
MnL·2H ₂ O	White	65-70	32.41	32.80	2.61	2.73	5.44	5.47	21.34	21.46
Fe ₂ L ₃ ·4H ₂ O	Brown	60-70	33.70	33.88	2.41	2.42	5.44	5.65	20.92	21.73
CoL·2H ₂ O	Pink	70–75	32.71	32.30	2.57	2.69	5.37	5.38	22.62	22.66
NiL·3H ₂ O	Green	60–65	35.29	35.34	1.29	1.31	6.01	6.12	20.73	21.11
CuL·0.5H ₂ O	Blue	60–70	33.20	32.62	2.06	2.33	5.48	5.44	26.94	26.74
ZnL·1.5H ₂ O	White	75-80	36.44	36.66	1.93	2.18	5.88	6.11	24.43	25.39



Fig. 2 TG/DTG/DSC curves of $Mn(3,4-pdc) \cdot H_2O(a)$ and $Mn(3,5-pdc)(H_2O)_2(b)$

Generally, the complexes of both series decompose in a similar way when heated in the two ground stages: the first stage is dehydration of the complexes with formation of more or less stable anhydrous compounds, and the second stage is decomposition of anhydrous compounds connected with oxidation of organic ligands and formation of metal oxides as residues. The first stage the endothermic processes of dehydration occur for the majority of complexes in only one step at low temperatures with the onset temperatures of water evaporation of the hydrates in the range of 333–343 K. Their value points to outer-sphere character of water molecules in the structure of complexes. However, higher temperature of dehydration at 423–503 K as it takes place in the case of

Complex	Stage	TG	DTG(DSC)	Peak nature/	Mass loss/%		Loss	Final	
		$T_{\rm range/K}$	T _{max. peaks/K}	kJ/mol H ₂ O	Calc. Found			residue	
Mn(3,4-pdc)·H ₂ O	Ι	343-473	453 (448)	endo (39.3)	8.75	8.05	H ₂ O	MnO	
	II	613–688	648 (673)	exo	70.21	69.38	CO ₂ ; CO; py		
							(pyridine rings, fragmentsof aromatic rings) [30]		
$Fe_2(3,4-pdc)_3 \cdot (H_2O)_6$	Ι	333-473	373 (373)	endo (22.7)	15,10	15,60	H ₂ O	Fe ₂ O ₃	
	II	573-653	603 (633)	exo	74,80	77,60	CO ₂ ; CO; py		
	III		753 (753)	exo			CO		
Co(3,4-pdc)·(H ₂ O) _{1,5}	Ι	333-473	358 (359)	endo (27.2)	10.75	10.43	H ₂ O	CoO	
	II	563-653	613 (643)	exo	70.15	68.87	CO ₂ ; CO; py		
$[Ni(3,4-pdc)(H_2O)_2] \cdot H_2O$	Ι	343-398	363 (373)	endo (29,3)	10.76	9.95	1H ₂ O	NiO	
	Π	398-453	423 (433)	endo	19.43	20.77	$2H_2O$		
	III	603-733	643 (673)	exo	73.12	73.35	CO ₂ ; CO; py		
	IV		693 (693)	exo			CO ₂ ; CO		
Cu(3,4-pdc)·H ₂ O	Ι	343-473	423 (348)	sł.endo (49,7)	7.29	7.00	H ₂ O	CuO	
	Π	513-533	528 (533)	exo	~ 40	67.77	CO_2		
	III	533-633	623 (623)	exo	67.75		CO ₂ ; CO; py		
Zn(3,4-pdc)·H ₂ O	Ι	343-413	358 (353)	endo (78,5)	7.24	7.08	H ₂ O	ZnO	
	Π	573–763	668 (643)	exo	67.30	68.08	CO ₂ ; py		
	III		728 (733)	exo			CO_2 ; CO		
	IV		873 (873)	exo			CO_2		
$Na_2(3,4-pdc) \cdot H_2O$	Ι	423–493	463 (468)	endo (34,4)	7.85	8.69	H ₂ O	Na ₂ CO ₃	
	II	703–783	733 (743)	exo	-	-	CO ₂ ; CO;py		

Table 3 Thermoanalytical results (TG,DTG,DSC) for complexes of 3,4-pyridinedicarboxylates



Fig. 3 TG/DTG/DSC curves of $Fe_2(3,4\text{-pdc})_3(H_2O)_6$ (a) and $Fe_2(3,5\text{-pdc})_3(H_2O)_4$ (b)

 $Mn(3,5-pdc)(H_2O)_2$ complex points to the inner-sphere water molecules [28–30].

There are some complexes for which the dehydration is of double-stage character as for $Co(3,5-pdc)(H_2O)_2$; $Ni(3,4-pdc)(H_2O)_3$; and $Ni(3,5-pdc)(H_2O)_3$ (Figs. 4b, 5a, b). For the $Co(3,5-pdc)(H_2O)_2$ complex, there are two onset temperatures of dehydration at 343 and 403 K, two peaks on the DTG and two endothermic peaks on DSC curves. This fact as well as loss of mass allows to determine the formula of the complex as $[Co(3,5-pdc)(H_2O)] \cdot (H_2O)$ which points to on one microcrystal- and one coordination water molecule. Two stages of dehydration are more visible on the TG, DTG and DSC curves of Ni(II) complexes.

For Ni(3,4-pdc)(H₂O)₃ onsets of dehydration are at 343 and 398 K, for Ni(3,5-pdc)(H₂O)₃ at 343 and 493 K. Dehydration at lower temperature (343 K) in both cases is



Fig. 4 TG/DTG/DSC curves of $Co(3,4-pdc)(H_2O)_{1,5}$ (a) and $[Co(3,5-pdc)(H_2O)] \cdot H_2O$ (b)



Fig. 5 TG/DTG/DSC curves of [Ni(3,5-pdc)(H₂O)₂]·H₂O (a) and [Ni(3,5-pdc)(H₂O)]·2H₂O (b)

connected with a loss of outer-sphere water, but dehydration at higher temperatures (398 and 493 K) is connected with a loss of coordination, inner-sphere water in the complex structures. Similarly to the Co(II) complex, two peaks on the DTG and DSC curves as well as calculated and found loss of mass allow to determine the formulae of complexes as $[Ni(3,4-pdc)(H_2O)_2]$ (H₂O) and $[Ni(3,5-pdc)(H_2O)]\cdot 2H_2O$.

Analysis of the TG/DTG curves course indicates that after dehydration the formed anhydrous complexes are not generally stable and the TG curves show a slight but systematic loss in mass. The complex of Mn(II) is the most stable in the series of complexes with 3,4pda ligand (Fig. 2a), while the complexes of Mn(II) and Zn(II) are the most stable in the series with 3,5-pda ligand (Figs. 2b and 7b). The complexes of 3,5-pda ligand seem to be more stable, and the onset temperatures for the decomposition of the anhydrous compounds, except for Fe(III) and Cu(II), are higher than those for 3,4pda ligand about 50–90 K (Tables 3 and 4). It can be due to more symmetric charge distribution in the complexes with 3,5-pdca ligand.

The anhydrous compounds of Mn(II), Fe(III), Co(II) and Ni(II) with 3,4pda and 3,4pda ligands above 600 K decompose in one step which correspond to the sharp exothermic peaks at about 700 K which can be also attributed to oxidation of organic matter and the gaseous products that evolve during thermal decomposition. In the case of Ni(3,4-pdc) complex (Fig. 5A), the exothermic peak splits into two. Processes of decomposition of anhydrous Cu(II) complexes of both ligands proceed in two stages (Fig. 6a, b). In both cases, the first step is connected with about 40 % mass loss and the second one leads to CuO formation. The first step is attributed to small, but the second to large exothermic effects. The large effect is attributed to the oxidation of carbonaceous residue, formed in the former step [31].

TG DTG(DSC) Peak nature/ Mass loss/% Final Complex Loss Stage kJ/mol H₂O residue Trange/K Tmax. peaks/K Calc. Found $Mn(3,5-pdc) \cdot (H_2O)_2$ I 423-503 473(483) endo (50.0) 14.06 15.07 H₂O MnO Π 663-748 703(728) 71.84 70.81 CO₂; CO; py exo (pyridine rings, fragmentsof aromatic rings)[30] $Fe_2(3,5-pdc)_3 \cdot (H_2O)_4$ I 343-423 358(363) endo(33.2) 10.60 10.10 H₂O Fe₂O₃ Π 573-748 643(653) exo 76,48 74,85 CO₂; CO; py [Co(3,5-pdc)(H₂O)]·H₂O I 343-393 363(363) endo(32.5) 6.92 7.00 H₂O CoO Π 403-463 12.31 433(433) endo 12.50 H_2O Ш 636-733 70.18 70.69 CO₂; CO; py 668(693) exo NiO [Ni(3,5-pdc)(H₂O)]·2H₂O I 343-443 423(430) endo(24.1) 12.94 13.00 $2H_2O$ Π 493-543 513(523) endo 19.42 19.10 $1H_2O$ Ш 636-713 673(713) 73.14 73.47 CO₂; CO; py exo Cu(3,5-pdc)·(H₂O)_{0.5} I 323-393 343(351) endo(21.0) 3.79 4.02 H₂O CuO Π ~ 45 64.94 CO_2 523-573 563(563) exo Ш 573-748 66.73 693(693) CO₂; CO; py exo Zn(3,5-pdc)·(H₂O)_{1.5} I 343-393 368(353) endo(18.6) 3.50 3.30 H₂O ZnO Π 393-493 10.10 10.48 443(447) endo H₂O Π 663-753 738(743) exo 68.39 68.72 CO₂; CO; py I Na2(3,5-pdc)·H2O 393-443 423(423) endo (38.8) 7.86 8.03 H₂O Na₂CO₃ Π 728-798 773(793) 53.19 53.73 CO₂; CO; py exo

 Table 4
 Thermoanalytical results (TG,DTG,DSC) for complexes of 3,5-pyridinedicarboxylates



Fig. 6 TG/DTG/DSC curves of Cu(3,4-pdc)·H₂O (a) and Cu(3,5-pdc)(H₂O)_{0,5} (b)

Thermal decomposition of both Zn(II) complexes proceeds in a different way. The Zn(3,4-pdc)·H₂O complex at 573 K undergoes decomposition and oxidation with two exothermic effects and next oxidation of carbonaceous residue with the exothermic effect at 873 K. The Zn(3,5pdc)·H₂O complex is more stable, and at the temperature higher by about 100 K decomposition and oxidation take place in one step without intermediates products. In all complexes of both series, the total mass loss above 800 K corresponds to formation of metal oxides (MO; Fe_2O_3) as final residues.

Additionally the simultaneous TG/DTG/DSC curves in dynamic dry air atmosphere of free 3,4-pyridinedicarboxylic and 3,5-pyridinedicarboxylic acids are presented in Fig. 8a, b. In both cases, these curves show total mass loss due to heating but in one step for 3,4- and in two steps for 3,5-pyridinedicarboxylic acid. The beginning of mass loss for 3,4-pdc acid starts at 473 K, and the maximum of mass



Fig. 7 TG/DTG/DSC curves of $Zn(3,4-pdc) \cdot H_2O(a)$ and $Zn(3,5-pdc)(H_2O)_{1,5}(b)$



Fig. 8 TG/DTG/DSC curves of 3,4-pdc (a) and 3,5-pdc (b)

loss is observed at 558 K with the simultaneous large endothermic effect with the maximum at 563 K. At 573 K the process is over. The endothermic peak at 563 K on the DSC curve is attributed to the melting of acid and partial evaporation as well as decarboxylation and decomposition [31, 32]. The 3,5pda is more stable, and the beginning of mass loss is observed at 523 K with the maximum at 608 K and the endothermic effect at 623 K. At 633 K the carbonaceous residue is formed and the exothermic peak at 753 K is attributed to its oxidation.

Disodium 3,4- and 3,5-pirydinedicarboxylates were necessary as the models for spectroscopic examination of complexes. They were obtained both as monohydrate microcrystalline compounds, but their thermal properties differ significantly. The thermal analysis of Na₂(3,4pdc)·H₂O exhibits three mass loss steps. The lower temperature mass (8.0/7.9 %) at 423-493 K corresponds to the dehydration with a loss of one molecule of water resulting in the endothermic peak. Anhydrous compound is stable to about 693 K. In this range of temperatures, the salts Na₂(3,4-pdc)·H₂O and Na₂(3,5-pdc)·H₂O exhibit the same way of decomposition but the anhydrous salt Na₂(3,5-pdc) is a bit more stable (up to 723 K). Further heating causes gradual decomposition and oxidation of anhydrous salts but in different ways. The anhydrous salt Na₂(3,4-pdc) decomposes in two steps with the mass loss at range of 733 K and about 873 K. The exothermic process with three peaks at 760; 765; and 773 K is attributed to oxidation of the organic matter and gaseous products evolved during the thermal decomposition and oxidation. In the case of $Na_2(3,5-pdc)$, the thermal decomposition occurs at 773 K in one step. The total mass loss (53.73/



Fig. 9 TG/DTG/DSC curves of Na_2(3,4-pdc)·H_2O (a) and Na_2(3,5-pdc)·H_2O (b)

Compound		<i>v</i> _{as} (COO ⁻)	v _{sym} (COO ⁻)	Δv	$ \beta_{sym} \\ (COO^-) $	$_{(COO^{-})}^{\beta_{as}}$	$\Delta\beta$	Proposed type of coordination
3,4-pyridined	icarboxyla	tes						
Na ₂ L·H ₂ O	IR	1602 vs, 1585 vs	1411 vs, 1381 s	191, 204	834 s	433 s	401	
	Raman	1606 s, 1581 m	1424 s, 1382 m	182, 199	840 m	431 vw	409	
MnL·H ₂ O	IR	1571 vs	1406 vs	165	840 m	469 m	381	Bidentate chelation
	Raman	1575 m	1411 vs	164	844 m	456 w	388	
Fe ₂ L ₃ ·3H ₂ O	IR	1576 vs, 1558 vs	1401 vs	175, 157	834 m	483 s	351	Bidentate chelation
CoL·1.5H ₂ O	IR	1607 vs, 1585 vs	1405 vs	202, 180	840 w	477 w	363	Bidentate chelation
NiL·2H ₂ O	IR	1585 vs	1405 vs	180	836 m	456 m	386	Bidentate chelation
CuL·H ₂ O	IR	1620 vs, 1585 vs	1386 vs	234, 199	837 w	499 w	338	Bidentate bridging and monodentate
ZnL·H ₂ O	IR	1610 vs, 1598 vs	1392 vs	228, 206	838 w	482 w	356	Monodentate
	Raman	1609 s, 1588 m	1401 s	208, 187	840 s	487 w	357	
3,5-pyridined	icarboxyla	tes						
Na ₂ L·H ₂ O	IR	1644 vs	1377 vs	267	824 w	425 m	399	
	Raman	1665 m	1376 vw	289	825 m	414 w	411	
MnL·2H ₂ O	IR	1614 vs, 1603 vs	1398 vs	216, 205	848 m	440 m	404	Bidentate chelation
	Raman	1604 m	1393 w	211	852 m	427 w	425	
Fe ₂ L ₃ ·4H ₂ O	IR	1608 vs	1384 vs	224	803 w	472 m	321	Bidentate chelation
CoL·2H ₂ O	IR	1615 vs	1401 s, 1379 s	214, 236	826 w	446 w	380	Bidentate chelation
NiL·3H ₂ O	IR	1623 vs	1377 vs	246	819 w	457 w	362	Bidentate chelation
CuL·0.5H ₂ O	IR	1634 vs	1377 vs	257	823 m	469 m	354	Bidentate chelation
ZnL·1.5H ₂ O	IR	1624 vs	1395 vs	229	837 s	448 m	389	Bidentate chelation
	Raman	1613 w	_	_	840 vs	_	_	

 Table 5
 Spectral parameters from the IR and Raman spectra of 3,4- and 3,5-pyridinedicarboxylates

53.19 %) suggests the formation of sodium carbonate (Fig. 9b; Table 4).

IR and Raman spectra

After metal complexation by 3,4-pyridinedicarboxylic and 3,5-pyridinedicarboxylic acids, some characteristic changes in the IR and Raman spectra of ligand and metal

complexes occurred. It was caused by changes in the electronic charge distribution in ligand and complex molecules. In the (Fig. S1 and S2) spectra of 3d-metal complexes the bands assigned to the stretching vibrations of the carboxylate anion occurred, whereas the bands derived from the C=O and C-O disappeared (these bands were present in the spectra of ligand) (Table 5). In the spectra of 3,4-pyridinedicarboxylates the strong bands

assigned to the asymmetric stretches $v_{as}COO^{-}$ were in the range 1620-1571 cm⁻¹ (IR) and 1609-1575 cm⁻¹ (Raman), whereas symmetric stretches ($v_{sym}COO^{-}$) in the range: 1406–1392 cm⁻¹ (IR) and 1411–1401 cm⁻¹ (Raman). In the spectra of 3,5-pyridinedicarboxylates v_{as} - COO^- occurred within 1634–1603 cm⁻¹ (IR) and 1624–1605 cm⁻¹ (Raman) as well as symmetric stretches $(v_{sym}COO^{-})$ in the range 1401–1377 cm⁻¹ (IR) and at 1393 cm^{-1} (Raman). In the spectra of 3,4-pyridinedicarboxylates bands derived from the deforming in-plane vibrations of the carboxylate anion ($\beta_{sym}COO^{-}$) were in the range 840–834 cm^{-1} (IR) and 844–840 cm^{-1} (Raman), whereas $\beta_{as}COO^{-}$: 499–469 cm⁻¹ (IR) and 487–456 cm⁻¹ (Raman). In the spectra of 3,5-pyridinedicarboxylate $\beta_{sym}COO^-$ were present within 848-803 cm⁻¹ (IR) and 852-840 cm¹ (Raman) as well as β_{as} COO⁻: 472–440 cm⁻¹ (IR) and 427 cm⁻¹ (Raman). On the basis of the differences between the wave numbers of the asymmetric and symmetric stretching vibrations of the carboxylate anion COO⁻, some assumption on the type of metal-ligand coordination might be stated [33, 34]. Especially the parameter $\Delta v = v_{as}COO^{-}-v_{svm}COO^{-}$ might be useful [33]. In the spectra of 3,4- and 3,5-pyridinedicarboxylates, more than one band assigned to the stretching vibrations of the carboxylate anion were present. In some cases, these bands differed in the values of their wave numbers which

was caused by the possibility of different type of metal coordination in one metal complex (Table 5) (Fig. 10). On the basis of the Δv (COO⁻), it was concluded that in the copper 3,4-pyridinedicarboxylate two type of copper ion coordination occurred, i.e., bidentate bridging and monodentate. In the Mn, Fe, Ni and Co 3,4-pyridinedicarboxylates and Mn, Fe, Co, Ni, Cu and Zn 3,5pyridinedicarboxylates, the metal ion was coordinated by the bidentate chelation type of bonding. In the case of Zn 3,4-pyridinedicarboxylate, the monodentate type of coordination existed.

The effect of metal ions on the electronic system of ligands was studied on the basis of the differences between location of bands assigned to the aromatic ring vibrations in the IR and Raman spectra of ligands and their metal complexes. The decrease in the wave numbers of these bands or decrease in their intensities or their disappearance shows destabilization of the aromatic system of ligand by metal ion. In the spectra of 3,5-pyridinedicarboxylate (Table 6), most bands assigned to the ring vibrations were located at higher wave numbers comparing with the appropriate band in ligand (i.e., in the case of 19b, 12, 17b in the IR and Raman spectra; deforming in-plane vibrations of β CH). In the spectra of metal complexes, some bands occurred (they were absent in the spectra of ligand), i.e., 8b (IR) and 8b, 3 and 19b (Raman). Some bands were shifted

Fig. 10 Proposed types of metal–ligand coordination: a 3,5-pyridinedicarboxylic acid with manganese, iron, cobalt, nickel, copper, zinc (bidentate chelation); b 3,4pyridinedicarboxylic acid with manganese, iron, cobalt, nickel (bidentate chelation); c 3,4pyridinedicarboxylic acid with zinc (monodentate); d 3,4pyridinedicarboxylic acid with copper (bidentate bridging and monodentate)



 Table 6
 Wave numbers and intensity of selected wave numbers from the IR and Raman spectra of 3,5-pyridinedicarboxylic acid and its metal complexes

3,5pda		3,5-pyridinedicarboxylate											
		Sodium		Mangane	ese	Iron	Cobalt	Nickel	Copper	Zinc			
IR	Raman	IR	Raman	IR	Raman	IR	IR	IR	IR	IR	Raman		
1603 m	1609 m	1605 s	1584 m	1603 vs	1604 s	_	_	-	_	_	1613 s	vCC _{ar}	8a
-	-	1576 s	1575 m	1556 vs	1567 s	1558 s	1559 vs	1558 s	1531 m	-	1586 s	vCC _{ar}	8b
1466 w	1463 m	1445 s	1446 m	1456 s	1463 s	_	_	-	1474 w	_	_	vCC _{ar}	19a
1420 w	-	1416 vs	1415 s	1435 s	_	1437 s	1440 m	1433 m	1445 m	1450 s	1453 s	vCC _{ar}	19b
1304 s	-	1285 m	1284 vw	1299 m	1301 vw	1293 m	1299 m	1290 w	1285 s	1293 m	1293 w	βCH_{ar}	3
1266 m	1266 w	1239 w	1251 vw	1238 m	1247 vw	-	1245 w	1248 w	1266 m	1245 m	1249 w	βCH_{ar}	9a
1219 s	1208 w	_	1189 w	1211 w	-	1204 w	1207 w	1205 vw	1194 m	1205 m	_	vCN _{ar}	
1168 vs	1150 vw	1151 m	1151 w	1150 m	1149 w	-	1150 w	1157 w	1158 m	1156 m	_	βCH_{ar}	
1124 m	1106 vw		1122 w	1124 m	1149 vw	-	1127 w	1134 w	1140 m	1133 m	_	βCH_{ar}	
1110 m	1095 w	1099 m	1089 w	_	1083 vw	-	_	_	1096 m	_	_	βCH_{ar}	18b
1065 s	1035 vs	1031 m	1037 s	1031 m	1033 vs	1030 w	1034 w	1032 w	1031 m	1032 m	1033 vs	βCH_{ar}	18a
942 m	946 w	962 w	965 vw	973 w	988 vw	-	_	_	977 w	977 m	985 m	βCH_{ar}	12
912 m	919 w	925 m	931 vw	935 m	936 vw	929 w	938 w	935 vw	942 w	938 w	938 vw	βCH_{ar}	17b
854 w	869 w	824 m	825 m	848 m	852 m	815 w	826 w	837 w	843 vw	837 m	840 s	γCH_{ar}	
-	801 m	_	_	_	805 w	_	-	-	-	-	807 m	γCH_{ar}	10a
783 m	_	773 s	776 w	769 s	774 w	765 m	769 m	770 m	755 m	770 s	780 w	γCH_{ar}	
697 s	682 w	687 m	699 m	695 m	684 w	693 m	695 w	668 m	677 m	687 m	703 vw	αCCC	6a
642 w	647 w	_	664 w	_	_	_	_	_	_	668 m	_	αCCC	6b
545 m	530 w	564 m	572 vw	_	510 w	535 m	528 vw	531 2	-	545 m	-	αCCC	

band intensity: s strong, m medium, w weak, vw very weak, sh shoulder; type of vibrations: v stretching, β bending in-plane, β ring ring bending in-plane, γ bending out-of-plane, α ring bending, φ ring deformation, sym symmetric, as asymmetric

toward lower wave numbers comparing with ligand (18b, 18a, vCN and γ CH in IR spectra). In the spectra of sodium 3,5-pyridinedicarboxylate, more bands disappeared compared with other metal complexes. In the spectrum of sodium salt, the wave numbers of some bands decreased in their values compared with the spectrum of 3,5-pyridinedicarboxylic acid, whereas in the spectra of 3d-metal complexes wave numbers of some bands increased in comparison with the spectrum of ligand.

In the spectra of 3,4-pyridinedicarboxylates (Table 7), bands no. 8b, 19a, 3, vCN, β CH, 12, γ CH, 6a (IR) as well as 8b, 19a, 3, 12, 17b, vCN and 6a (Raman) were located at higher wave numbers comparing with the appropriate ones in the spectrum of 3,4-pyridinedicarboxylic acid. Whereas bands no. 9a, 18b and γ CH (IR) and 19a, 8a (Raman) were shifted toward lower wave numbers comparing with ligand. Bands no. 19b and 6b (IR) as well as 19b and γ CH (Raman) were present in the spectra of acids and disappeared in the spectra of complexes.

Comparing the IR and Raman spectra of 3,4- and 3,5pyridinedicarboxylic acids and their 3d-metal complexes, it could be concluded that 3d-transitional metal stabilized the electronic system of ligands (similarly as it was stated in the case of 3d-metal complexes of pyridinemonocarboxylic acid, i.e., picolinic acid) [22]. 3d-Metals strongly affected the electronic system of 3,4-pyridinedicarboxylic acid than 3,5-pyridinedicarboxylic acid (on the basis of the changes in the location and intensity of the aromatic ring vibrations in the spectra of ligands and complexes). The carboxylic groups in the 3,5-pyridinedicarboxylic acid are symmetrically located taking into account the location of the nitrogen atom in the aromatic ring. This causes a symmetric distribution of the electronic charge in the molecule of 3,5pyridinedicarboxylic acid. In the case of 3,4-pyridinedicarboxylic acid, the carboxylic groups are unsymmetrically located which translates into unsymmetrically distributed electronic charge in 3,4-pyridinedicarboxylic acid molecule. The differences in the location of the carboxylic groups and the electronic charge distribution in pyridinedicarboxylic acids cause that 3d-transition metal ions in different way affect the molecular structure of studied ligands. This causes different physico-chemical and thermal properties of studied metal complexes. The similar conclusion was described in the papers devoted to the salts

3,4pda		3,4-pyridinedicarboxylate											
		Sodium		Mangane	ese	Iron	Cobalt	Nickel	Copper	Zinc			
IR I	Raman	IR	Raman	IR	Raman	IR	IR	IR	IR	IR	Raman		
1607 vs	1613 m	_	1606 w	_	1608 s	_	_	_	_		1609 s	vCC _{ar}	8a
_	-	-	1581 w	_	1575 m	-	_	-	_		1588 m	vCC _{ar}	
1521 m	1534 m	1546 m	1548 w	_	1548 m	_	_	_	_	1558 s	1554 s	vCC _{ar}	8b
1472 m	1492 m	1481 s	1488 w	1491 m	1492 s	1488 s	1493 m	1490 s	1491 m	1491 m	1494 s	vCC _{ar}	19a
1456 m	1453 vw	-	1424 s	-	_	-	_	-	-	-	_	vCC _{ar}	19b
_	1399 s	_	1382 w	_	_	_	_	_	_	_	_	vCHar	14
1276 w	1285 m	1273 w	1287 vw	1285 w	1285 w	1283 w	1286 w	1286 w	1283 w	1286 w	1287 w	βCH_{ar}	3
1260 w	1251 s	1241 w	1239 vw	1229 w	1222 w	_	1231 m	1235 w	_	1234 w	1237 m	βCH_{ar}	9a
1177 w	1170 vs	1197 w	1197 m	1200 w	1203 w	1191 w	1195 w	1194 w	1192 m	1195 w	1201 w	vCNar	
1157 m	_	1160 w	1170 w	1171 w	1170 m	1167 m	1167 w	1167 w	1166 m	1167 w	1168 m	βCH_{ar}	
_	1120 m	1126 vw	1127 vw	1124 w	1125 w	1122 m	1123 w	1124 w	1120 m	1124 w	1127 w	βCH_{ar}	
1078 s	1064 vs	1052 m	1054 w	1058 w	1058 m	1059 m	1064 w	1068 w	1069 m	1072 w	1071 m	βCH_{ar}	18b
973 m	975 w	977 w	_	981 vw	979 vw	_	993 w	_	_	988 w	990 vw	βCH_{ar}	12
_	929 vw	930 vw	_	955 vw	950 vw	_	952 w	_	947 w	950 w	944 vw	βCH_{ar}	17b
844 s	836 vw	853 m	867 vw	862 w	844 m	853 m	840 w	836 m	837 m	838 w	840 s	γCH_{ar}	
_	819 m	_	826 m	_	_	_	_	_	_	_	_	γCH_{ar}	10a
779 m	774 vw	786 m	784 vw	790 m	790 vw	772 m	782 w	782 m	779 m	785 w	_	γCH_{ar}	
731 m	732 vw	708 m	709 vw	708 m	707 vw	707 m	714 m	718 m	722 m	722 w	734 vw	αCCC	
677 w	672 m	678 m	684 w	678 s	684 w	677 s	681 m	683 m	685 m	684 m	686 m	αCCC	6a
660 s	_	646 m	_	_	_	_	_	_	_	650 vw	_	αCCC	6b
587 m	585 vw	572 w	572 vw	595 m	599 vw	582 m	610 w	615 m	609 w	606 w	612 vw	αCCC	

 Table 7
 Wave numbers and intensity of selected wave numbers from the IR and Raman spectra of 3,4-pyridinedicarboxylic acid and its metal complexes

band intensity: s strong, m medium, w weak, vw very weak, sh shoulder; type of vibrations: v stretching, β bending in-plane, β ring ring bending in-plane, γ bending out-of-plane, α ring bending, φ ring deformation, sym symmetric, as asymmetric

and metal complexes of 2, 3- and 4-pyridinecarboxylic acids [19, 20].

Conclusions

The thermal decomposition of the two series of hydrated complexes of Mn(II), Fe(III), Co(II), Ni(II), Cu(II) and Zn(II) with 3,4- and 3,5-pyridinedicarboxylic acids is a multistage process. At first the complexes lose all water molecules in one or two steps which are connected with endothermic effects. After dehydration, in the next stages, the anhydrous complexes decompose exothermally which can be also attributed to oxidation of organic matter and the gaseous products evolved during thermal decomposition. Finally, metal oxides are formed. Larger thermal stability is observed for anhydrous complexes of 3,5-pda ligand which can be a result of more symmetric structure and charge distribution of this ligand contrary to the thermal stability of 3,4-pda series. The differences in the location of the carboxylic groups and the electronic charge distribution in

pyridinedicarboxylic acids cause that 3d-transition metal ions in different way affect the molecular structure of studied ligands. This causes different physico-chemical and thermal properties of studied metal complexes.

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