Mass Spectra of Metal Tropolonates

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The mass spectra of the complexes MT_n (TH = tropolone; M = Co, Ni, Cu, Zn, Pd; n = 2. M = Al, Ga, In, Sc, Cr, Fe; n = 3) have been investigated. The main decomposition pathways involve loss of T-radicals and CO molecules. For complexes derived from transition metals loss of CO groups leads to cyclopentadienyl metal ions reflecting the ability of such metals to form complexes with unsaturated hydrocarbon ligands. In complexes of type MT_3 successive loss of T-radicals occurs only when M = Ga, In, Cr or Fe. This behaviour is rationalized in terms of the ability of these metals to change oxidation state. The spectrum of the copper complex exhibits several hydrogen-migration reactions.

Introduction

The mass spectrometric study of metal chelates, especially of those derived from β -diketones, has received considerable attention in recent years. On the basis of these studies it has been established that the metal has a pronounced effect on the fragmentation behaviour of metal chelates. Following our studies of various types of metal complexes we have now investigated several metal tropolonates of formula (I):

$$n = 2; M = Co, Ni, Cu, Zn, Pd$$

 $n = 3; M = Al, Ga, In, Sc, Cr, Fe$

TABLE I. Ion Abundances^a for MT₂.

Ion ^b	M in MT ₂									
	Co m/e		Ni m/e ^c		Cu m/e		Zn m/e	F 200.	Pd m/e	
$[MT_2]^+$	301	34(3) ^d	300	40(4)	305	21(1)	306	50(5)	346	35(5)
$[MT_2-H_2O]^+$	_	` `	_	. ,	287		-		_	
[MT ₂ -CO] ⁺	273	4	272	5	277	1	278	11	318	1
[MT ₂ -H ₂ O-CO] ⁺	_		_		259	1	_		_	
$[MT_2-2CO]^+$	245	1	244	1	249	1	250	1	_	
$[MT_2-C_5H_4O]^+$	_		_		_		226	1	_	
$[MT_2-2CO-C_2H_2]^+$	219	1	218	1	223	1	_		_	
[MT ₂ -3CO] ⁺	_		216	1	221	1	222	1	262	2
MT ₂ -4COj ⁺	189	1	188	1	_		_		_	
[MT+H] ⁺			_		185	11	_		_	
[MT] ⁺	180	30	179	18	184	22	185	29	225	2
[MT_H] ⁺	_		178	1	183	8	_		_	
[MT-CO] ⁺	152	1	151	2	156	9	157	1	197	6
[MT_HCO] ⁺	_		150	1	155	1	_		_	
MTH-CO-H ₂ O] ⁺	_				139	1	_		_	
[MT-2CO] ⁺	124	10	123	11	128	4	_		169	48
$[MC_5H_4]^+$	123		122		127	3	_		_	`
$[MC_3H_3]^+$	98	3	97	4	102	1	_		_	
$[MC_3H_2]^+$	_		96	1	101	1	-		_	
[M] ⁺	59	6	58	4	63	23	64	1	104	6

^a Abundances are expressed as percentage of the total ion current due to metal-containing ions. All species are corrected for isotopic abundance. ^b Only metal-containing ions are recorded. ^c m/e values are given for the ions containing ⁵⁸Ni, ⁶³Cu, ⁶⁴Zn and ¹⁰⁴Pd. ^d Figures in brackets indicate abundance of doubly charged species.

TABLE II. Metastable Ions in MT₂.

Transition	M in MT ₂	m/e Values for	m*	Neutral		
		Transition	Found ^a	Calcd.	Fragment	
$[MT_2]^+ \rightarrow [MT_2 - H_2O]^+$	Cu	305→287	270	270.06	H ₂ O	
$[MT_2]^+ \rightarrow [MT_2-CO]^+$	Co	$301 \rightarrow 273$	247.5	247.60	20	
	Ni	$300 \rightarrow 272$	246.8	246.61	00	
	Cu	$305 \rightarrow 277$	251.5	251.57	CO	
	Zn	$306 \rightarrow 278$	252.5	252.56		
$[MT_2]^+ \rightarrow [MT_2-2CO]^+$	Co	$301 \rightarrow 245$	199.5	199.41 \	260	
	Zn	$306 \rightarrow 250$	204	204.24 ∫	$2 \times CO$	
$[MT_2]^+ \rightarrow [MT+H]^+$	Cu	$305 \rightarrow 185$	112.1	112.21	[T–H]	
$[MT_2]^+ \rightarrow [MT]^+$	Co	$301 \rightarrow 180$	107.6	107.64)	. ,	
	Cu	$305 \rightarrow 184$	111.0	111.00 }	T	
	Zn	$306 \rightarrow 185$	112	111.85		
$[MT_2]^+ \rightarrow [MT-H]^+$	Cu	$305 \rightarrow 183$	109.7	109.80	[T + H]	
$[MT_2-CO]^+ \rightarrow [MT_2-2CO]^+$	Ni	$272 \rightarrow 244$	219.0	218.88	2×CO	
$[MT_2-CO]^+ \rightarrow [MT]^+$	Co	$273 \rightarrow 180$	118.5	118.68)		
	Ni	$272 \rightarrow 179$	117.6	117.80 }	C_6H_5O	
	Zn	278→185	123	123.11 J	0 3	
$[MT_2-2CO]^+ \rightarrow [MT]^+$	Co	$245 \rightarrow 180$	132.2	132.24	0.11	
	Zn	$250 \rightarrow 185$	136.7	136.90	C_5H_5	
$[MT_2-4CO]^+ \rightarrow [MT-2CO]^+$	Co	$189 \rightarrow 124$	81.2	81.35)		
	Ni	$188 \rightarrow 123$	80.5	80.47 }	C_5H_5	
$[MT]^+ \rightarrow [MT-CO]^+$	Co	$180 \rightarrow 152$	128.3	128.36)		
	Ni	$179 \rightarrow 151$	127.4	127.37 }	CO	
	Cu	$184 \rightarrow 156$	132.3	132.26		
[MT] ⁺ → [MT–2CO] ⁺	Co	$180 \rightarrow 124$	85.5	85.42)	2. 60	
	Ni	$179 \rightarrow 123$	84.5	84.52	$2 \times CO$	
$[MT]^+ \rightarrow [M]^+$	Co	180→ 59	19.3	19.34	T	
[MT-CO] ⁺ → [MT-2CO] ⁺	Co	$152 \rightarrow 124$	101.2	101.16)	200	
-	Ni	$151 \rightarrow 123$	100.2	100.19	$2 \times CO$	
$[MT-2CO]^+ \rightarrow [MT-2CO-C_2H_2]^+$	Co	$124 \rightarrow 98$	77.9	77.45	0.11	
	Ni	123→ 97	76.5	76.50	C_2H_2	
$[MT-2CO]^+ \rightarrow [M]^+$	Co	124→ 59	28.0	28.07	CH	
	Ni	123→ 58	27.3	27.35	C_5H_5	

^a For the compounds MT_2 (M = Cu, Ni, Zn) only transitions involving ⁶³Cu, ⁵⁸Ni, ⁶⁴Zn are given.

Experimental

All the compounds except ScT₃ (TH = tropolone) have been prepared as described in the literature. The complex ScT₃ has been prepared by heating under reflux a mixture of scandium trichloride (0.1 g, 1 mol equiv.) and tropolone (3 mol equiv.) in ethanol. The crude product was purified by sublimation at 250° C /0.1 mm (Found: C, 61.6; H, 3.6. $C_{21}H_{15}O_6Sc$ requires: C, 61.8; H, 3.7%).

The mass spectra were obtained on a Hitachi RMS-4 spectrometer operating at 80eV, and by use of a direct insertion probe at *ca.* 250°C. Exact mass measurements and enhanced metastable spectra were obtained on an AEI MS-9 instrument at the Physico-Chemical Measurements Unit, Harwell.

Scheme 1

Results and Discussion

The spectra are given in Tables I–VI. Only the metal-containing ions are recorded. The spectra also showed the presence of non-metal species the most abundant of which had m/e values 122, 121, 94, 66, 65 and 39. In all cases, the sum of the intensities of these ions was less than 15% of the total ion-current.

MT₂ Complexes

The spectrum of each MT₂ complex is characterised by an intense molecular ion and exhibits a fairly intense doubly charged molecular ion. In general the molecular ions [MT₂]⁺ fragment by loss of a ligand radical to give the even-electron ions [MT]⁺ and by reactions involving elimination of CO groups (Scheme 1*).

The subsequent fragmentation of the ions $[MT]^+$ proceeds by similar routes except for the ion $[ZnT]^+$ which shows little tendency to decompose. The spectra of all compounds show fairly intense ions arising by loss of one CO group from the molecular ion and, except when M = Pd, weak ions due to loss of two such groups. The reactions leading to the ions $[MT_2-nCO]^+$ (n = 1 or 2) are strongly metastable supported in most cases.

TABLE III. Accurate Mass Measurements on Selected Peaks for MT_2 .

Com- pound	Nominal Mass	Deter- mined Mass	Ion Assignment	Calculated Mass
CoT ₂	∫ 189	189.0710	C ₁₀ H ₁₀ Co	189.0114
C01 ₂	123ª	122.9644	C ₅ H ₄ Co	122.9645
	(300	299.9937	$C_{14}H_{10}O_4^{58}N_1$	299.9932
NiT_2	{ 244	244.0001	$C_{12}H_{10}O_2^{58}N_1$	244.0034
	122 ^b	121.9640	C ₅ H ₄ ⁵⁷ Ni	121.9666
	(287	286.9761	C ₁₄ H ₈ O ₃ ⁶³ Cu	286.9768
	277	277.9928	$C_{13}H_{10}O_3^{63}Cu$	276.9926
	259	258.9820	$C_{13}H_8O_2^{63}Cu$	258.9820
	186	185,9557	C ₇ H ₅ O ₂ ⁶⁵ Cu	185.9567
	185°	184.9642	$C_7H_6O_2^{63}Cu$	184.9664
CuT ₂	185°	184.9487	$C_7H_4O_2^{65}Cu$	184.9489
-	184	183.9578	C ₇ H ₅ O ₂ ⁶³ Cu	183,9585
	183	182,9495	$C_7H_4O_2^{63}Cu$	182,9507
	156	155.9635	$C_6H_5O^{63}Cu$	155.9636
	139	138.9611	C ₆ H ₄ ⁶³ Cu	138.9609
	127	126.9608	C ₅ H ₄ ⁶³ Cu	126.9608
	(250	249.9972	$C_{12}H_{10}O_2^{64}Zn$	249.9972
ZnT ₂	226	225.9605	$C_9H_6O_3^{64}Zn$	225,9608
•	157	156.9630	$C_6H_5O^{64}Zn$	156,9631
PdT_2	347	346.9626	$C_{14}H_{10}O_4^{105}Pd$	346.9630

 $[^]a$ Peak overlaps with $[{C_6}^{13}{\rm CH_6O_2}]^+.$ b Peak overlaps with $[{C_7}{\rm H_6O_2}]^+.$ c Doublet intensity ratio $[{C_7}{\rm H_6O_2}^{63}{\rm Cu}]^+/$ $[{C_7}{\rm H_4O_2}^{65}{\rm Cu}]^+=2.$

TABLE IV. Ion Abundances^a for MT₃.

Ion ^b	M in MT ₃											
	Al		Ga		In		Sc		Cr		Fe	
	m/e		m/e	2	m/e		m/e		m/e		m/e	
$[MT_3]^+$	390	8.0(0.4)d	432	2.3	478	3.6(0.5)	408	20.0	415	16.6(1.7)	419	8.7(0.4)
$[MT_2]^+$	269	75.0	311	90.7	357	59.0	287	74.0	294	57.6(0.6)	298	66.0(0.6)
$[MT_2-C_2H_2]^+$	243	0.6		_		_	261	w		_		
$[MT_2-CO]^+$	241	0.6(0.3)	283	w	329	w	259	1.0	266	0.1	270	1.0
[MT ₂ -HCO] ⁺	240	0.3		_			258	w		~		_
$[MT_2-2CO]^+$		_		_		_		_	238	0.2		_
$[MT_2-2CO-C_2H_2]^+$		_		_		_			214	0.5	218	0.1
$[MT_2-3CO]^+$		_		-		_		-	212	0.5	216	0.2
$[MT_2-C_5H_4O]^+$	189	3.4	231	2.2		_	207	2.0				_
$[MT_2-C_5H_4O_2]^+$	173	1.8		_		_	191	1.0				_
[MTOH] ⁺	165	3.5	207	1.1		_	183	0.6		_		_
[MT] ⁺	148	1.7	190	2.1	236	15.4	166	0.4	173	20.0	177	21.0
$[MT-C_2H_2]^+$		_		_		_		-	147	0.6		_
[MT-CO] ⁺		_	162	0.5	208	0.2		-	145	0.4	149	0.3
[MT_HCO]+		_		_	207	0.2	137	1.0		-		
[MT-2CO] ⁺		_		_				_	117	0.2	121	0.1
[M] ⁺		_	69	3.9	115	18.2		_	52	2.5	56	1.7

^a Abundances are expressed as percentage of the total ion current due to metal-containing ions. All species are corrected for isotopic abundance. ^b Only metal-containing ions are recorded. ^c m/e values are given for the ions containing ⁶⁹Ga, ¹¹⁵In, ⁵²Cr and ⁵⁶Fe. ^d Figures in brackets indicate abundance of doubly charged species.

^{*} In the fragmentation schemes solid arrows represent processes confirmed by presence of metastable peaks whilst the dotted arrows represent unconfirmed but possible processes.

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TABLE V. Metastable Ions in MT₃.

Fransition	M in MT ₃	m/e Values for Transition	m*	Neutral Fragment	
		Tunismon	Founda	Calcd.	
$[MT_3]^+ \rightarrow [MT_2]^+$	Al	390→269	185.5	185.54	
	Ga	$432 \rightarrow 311$	224	223.89	
	In	$478 \rightarrow 357$	267	266.66	T
	Sc	$408 \rightarrow 287$	202	201.88 (1
	Cr	$415 \rightarrow 294$	208	208.27	
	Fe	$419 \rightarrow 298$	212	211.94 ^J	
$[MT_2]^+ \rightarrow [MT_2-CO]^+$	Al	$269 \rightarrow 241$	216	215.90	
	Ga	$311 \rightarrow 283$	257.5	257.51	
	Sc	$287 \rightarrow 259$	234	233.73 }	CO
	Cr	$294 \rightarrow 266$	241	240.66	
	Fe	$298 \rightarrow 270$	245	244.63 J	
$[MT_2]^+ \rightarrow [MT_2 - HCO]^+$	Al	$269 \rightarrow 240$	214	214.13	HCO
MT_2 ⁺ $\rightarrow [MT_2-2CO]$ ⁺	Ga	$294 \rightarrow 238$	193	192.66	$2 \times CO$
MT_2] ⁺ $\rightarrow [MT_2 - C_5H_4O]^+$	Al	$269 \to 189$	132.8	132.79)	
	Ga	$311 \rightarrow 231$	171.6	171.58	C_5H_4O
	Sc	$287 \rightarrow 207$	149.3	149.30	5 .
$[MT_2]^+ \rightarrow [MTOH]^+$	Al	$269 \to 165$	101.0	101.20	CHO
-, ,	Sc	$287 \rightarrow 183$	116.5	116.68	C_7H_4O
$[MT_2]^+ \rightarrow [MT]^+$	Ga	$311 \rightarrow 190$	116	116.08)	
	In	$357 \rightarrow 236$	156	156.00	OTP.
	Cr	$294 \rightarrow 173$	101.8	101.79	T
	Fe	298→177	105.0	105.13	
MT_2 - CO] ⁺ \rightarrow [MTOH] ⁺	Al	$241 \to 165$	113	112.96	0.17
	Sc	$259 \to 183$	129	129.30	C ₆ H₄
$[MT_2-CO]^+ \rightarrow [MT]^+$	Cr	$266 \rightarrow 173$	112.5	112.51	CHO
J t J	Fe	270→177	116.0	116.03	C_6H_5O
$MT_2-2CO]^+ \rightarrow [MT]^+$	Cr	238→173	125.5	125.75	C_5H_5
$MT_2-C_5H_4O]^+ \rightarrow [MT]^+$	Ga	$231 \rightarrow 190$	156.3	156.27	C ₂ HO
MTOH] ⁺ → [MT] ⁺	Al	$165 \rightarrow 148$	132.8	132.75	_
	Sc	183→137	102.6	102.56	ОН
$[MT]^+ \rightarrow [MT-CO]^+$	Ga	$190 \rightarrow 162$	138	138.12)	
	Cr	173→145	121.5	121.53	CO
	Fe	177 → 149	125.5	125.42	
$[MT]^+ \rightarrow [MT-2CO]^+$	Fe	177→121	82.7	82.71	
MT] ⁺ \rightarrow [M] ⁺	Ga	190→ 69	25.1	25.06	T
, (<u>-</u>)	In	236→115	56	56.03	T
$MT-CO]^+ \rightarrow [MT-2CO]^+$	Fe	$149 \rightarrow 121$	98.2	98.26	CO
$MT-2CO]^+ \rightarrow [M]^+$	Fe	121→ 56	25.9	25.91	C_5H_5

^a For the compounds MT₃ (M = Cr, Fe, Ga, In) only transitions involving ⁵²Cr, ⁵⁶Fe, ⁶⁹Ga, ¹¹⁵In are given.

When M=Co or Ni loss of all four CO groups from $[MT_2]^+$ leads to the biscyclopentadienylmetal ion $[M(C_5H_5)_2]^+$ ($[MT_2-4CO]^+$) which decomposes to the ion $[MC_5H_5]^+$ ($[MT-2CO]^+$) by the metastable-supported elimination of a cyclopentadienyl radical. The ions $[MC_5H_5]^+$, which are present in high abundance, are also formed by elimination of two CO groups from $[MT]^+$. This process is metastable-supported, occurs when M=Cu or Pd also but is not observed for the zinc compound. The formulation of the ions $[MT_2-4CO]^+$ and $[MT-2CO]^+$ as cyclopentadienyl metal ions is supported by their fragmentation behaviour (Scheme 2) which involves processes, such as the metastable-

$$[M(C_5H_5)_2]^+ \to [M(C_5H_5)]^+ \longrightarrow [M]^+$$

$$([MT_2-4CO]^+) \quad ([MT-2CO]^+)$$

Scheme 2

supported elimination of $C_5H_5^{\cdot}$ and C_2H_2 species, analogous to those observed in the spectra of metal cyclopentadienyl compounds.⁵ The formation of these ions in all cases except when M=Zn is in accord with the ability of transition metals to form complexes with unsaturated hydrocarbon ligands. However, the zinc

TABLE VI. Accurate Mass Measurements on Selected Peaks for MT_a .

Com- pound	Nom- inal Mass	Deter- mined Mass	Ion Assignment	Calculated Mass
	(390	390.0680	C ₂₁ H ₁₅ O ₆ Al	390.0684
	243	243.0240	$C_{12}H_8O_4Al$	243.0237
	241	241.0442	$C_{13}H_{10}O_3Al$	241.0445
AlT ₃	{ 240	240.0365	$C_{13}H_9O_3Al$	240.0366
ŭ	189	189.0135	C ₉ H ₆ O ₃ Al	189.0183
	173	173.0182	C ₉ H ₆ O ₂ Al	173.0183
	165	165.0128	C ₇ H ₆ O ₃ Al	165.0132
	(432	432.0127	$C_{21}H_{15}O_6^{69}Ga$	432.0125
GaT ₃	{ 231	230.9591	C ₉ H ₆ O ₃ ⁶⁹ Ga	230.9573
J	207	206.9567	C ₇ H ₆ O ₃ ⁶⁹ Ga	206.9573
T 00	478	477.9908	$C_{21}H_{15}O_6^{115}In$	477.9908
InT ₃	1 207	207.9370	$C_{16}H_5O^{115}In$	207.9379
	(408	408.0405	C ₂₁ H ₁₅ O ₆ Sc	408.0428
0. T	259	259.0187	C ₁₃ H ₁₀ O ₃ Sc	259.0189
ScT ₃	207	206.9877	C ₉ H ₆ O ₃ Sc	206.9876
	183	182.9877	C ₇ H ₆ O ₃ Sc	182.9876
CrT ₃	J 415	415.0274	$C_{21}H_{15}O_6^{52}Cr$	415.0274
C1 13	52	51.9408	⁵² Cr	51.9405
	(270	269.9970	$C_{13}H_{10}O_3^{56}Fe$	269.9979
FeT ₃	149ª	148.9960	$C_{14}H_{10}O_4^{56}Fe^b$	297.9928
3	149a	148.9682	C ₆ H ₅ O ⁵⁶ Fe	148.9689

^a Doublet intensity ratio $[C_{14}H_{10}O_4Fe]^{2+}/[C_6H_5OFe]^+ = 2$.

^b Doubly charged ion.

compound exhibits the ion $[ZnT_2-3CO]^+$, which might be represented as $[C_6H_5O\cdot Zn\cdot C_5H_5]^+$, in higher abundance than any of the other compounds.

In addition to loss of a ligand radical and CO elimination the molecular ion of the copper compound exhibits several other reactions which involve hydrogen transfer. These reactions lead to the ions $[CuT_2-H_2O]^+$, $[CuT+H]^+$ and $[CuT-H]^+$ (Scheme 3). The reaction $[CuT_2]^+ \rightarrow [CuT_2-H_2O]^+$

$$[CuT_2-H_2O]^+ \longrightarrow [CuT_2-H_2O-CO]^+$$

$$[CuT_2]^+ \longrightarrow [CuT+H]^+ \longrightarrow [CuTH-CO-H_2O]^+$$

$$[CuT-H]^+ \longrightarrow [CuT-H-CO]^+$$

Scheme 3

is metastable-supported and a grouping of metastable peaks at m/e 109–112 is probably due to the reactions $[CuT_2]^+ \rightarrow [CuT]^+$, $[CuT_2]^+ \rightarrow [CuT+H]^+$ and $[CuT_2]^+ \rightarrow [CuT-H]^+$. Ions analogous to $[CuT_2-H_2O]^+$, $[CuT+H]^+$ and $[CuT-H]^+$ have been observed in the spectra of several copper(II) β -diketonates and ions analogous to $[CuT+H]^+$ have also been observed in the spectra of cobalt(II) and nickel(II) β -diketonates.^{1,6,7} The ions $[CuT_2-H_2O-CO]^+$, $[CuTH-CO-H_2O]^+$ and $[CuT-H-CO]^+$ may originate as indicated in Scheme 3.

MT₃ Complexes

All the MT_3 complexes show the molecular ion $[MT_3]^+$ which fragments exclusively by loss of a ligand radical to give the even-electron ion $[MT_2]^+$. This ion is present as the most intense ion in all cases. Further fragmentation is dependent on the nature of the metal and reflects primarily its oxidation state characteristics and its ability to form complexes with unsaturated hydrocarbon ligands. When M = Cr, Fe, Ga or In decomposition of $[MT_2]^+$ leads to the ions $[MT]^+$ and $[M]^+$ (Scheme 4).

$$[MT_3]^+ \rightarrow [MT_2]^+ \rightarrow [MT]^+ \rightarrow [M]^+$$
Scheme 4

The presence of these ions in high abundance and the successive loss of odd-electron species from even-electron ions in the spectra of the MT_3 complexes (M=Cr, Fe, Ga or In) can be correlated with the ability of these metals to undergo reduction and the stability of their lower oxidation states. For example the loss of a ligand radical from $[M^{III}T_2]^+$ (M=Ga or In) can be correlated with the stability of oxidation state +I by assuming that this is accompanied by transfer of one electron from the remaining ligand to the metal.

$$\begin{bmatrix} T \\ T \end{bmatrix}^{+} \rightarrow [M^{I}T]^{+} + T$$

The very low abundance of $[MT]^+$ and the absence of $[M]^+$ in the spectra of MT_3 (M = AI or Sc) is in accord with the high stability of aluminium(III) and scandium(III).

When M = Ga, In, Cr or Fe the reaction $[MT_2]^+ \rightarrow [MT]^+$ is supported by the appropriate metastable ions. The detection of metastable ions for this reaction does not necessarily suggest that it occurs in a single step *i.e.* by loss of an intact ligand radical. The transition could also proceed stepwise as indicated in the case of the chromium and iron compounds by the presence of the ions $[MT_2-n CO]^+$ (n = 1 or 2) and the occurrence of the reactions given in Scheme 5.

$$[MT_2]^+ \xrightarrow{[MT_2-2CO]^+ \longrightarrow [MT_2-2CO]^+}$$

$$[MT]^+$$
Scheme 5

For the aluminium and scandium compounds the reaction $[MT_2]^+ \rightarrow [MT]^+$ also occurs stepwise but in these cases the loss of CO from $[MT]^+$ is followed by loss of another even-electron species, C_6H_4 . The ion $[MTOH]^+$ thus produced then gives $[MT]^+$ by loss of an OH radical (Scheme 6).

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$$[MT_2-CO]^+$$

$$[MT_2]^+ \longrightarrow [MTOH]^+ \longrightarrow [MT]^+$$

$$\longrightarrow [MT_2-C_5H_4O]^+ \longrightarrow [MT_2-C_5H_4O_2]^+$$
Scheme 6

All these steps, as well as the step $[MT_2]^+ \rightarrow [MTOH]^+$ are metastable supported. Ions analogous to $[MTOH]^+$ have been observed in the spectra of the acetylacetonate complexes of aluminium and scandium.⁸ Another interesting metastable-supported reaction of $[MT_2]^+$ (M = Al or Sc) involves loss of 80 mass units to give $[MT_2-C_5H_4O]^+$. This ion is also present in the spectra of the compounds GaT_3 and ZnT_2 .

In close similarity to the decomposition of the ions $[MT]^+$ (M = Co, Ni, Cu or Pd) the analogous ironand chromium-containing ions undergo elimination of one or two CO groups to form the ions $[MT-CO]^+$ and $[MT-2CO]^+$ ($[MC_5H_5]^+$). As in the spectra of $M(C_5H_5)_2$ (M = Cr or Fe), further decomposition of the cyclopentadienylmetal ions leads to the bare metal ions. In contrast when M = Ga or In only one CO group is lost from $[MT]^+$. For these metals the reaction $[MT]^+ \rightarrow [MT-CO]^+$ is weak and the main decomposition pathway is $[MT]^+ \rightarrow [M]^+$.

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