Thermal analysis of solvatomorphic decakis (dimethylammonium) dihydrogendodecatungstate hydrates

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Abstract
This study aims to describe the thermal decomposition of two solvatomorphs of decakis(dimethylammonium) dihydrogendodecatungstate ((Me2NH2)10H2W12O42·10H2O and 11 H2O) under inert and oxidizing atmospheres. Thermal studies have been done by TG-MS, TG-DSC-MS, XRD and IR methods in both synthetic air and helium atmospheres. The general characteristics of thermal decomposition are similar for both solvatomorphs. Minor differences could be observed in the resolution and shifting of the decomposition peak temperatures depending on the heating rate or atmosphere used. The first step of decomposition is endothermic in both atmospheres and involves 2 and 5 water molecule elimination with ~150 and ~120 °C peak temperatures for the decahydrate and undecahydrate, respectively. The elimination of further water and dimethylamine was observed with increasing the temperature, as well as the disruption of the lattice of compounds. Until 300 °C, these processes are endothermic in both atmospheres, and the further decomposition processes at higher temperatures are left endothermic in helium, but become exothermic in synthetic air atmosphere. In helium atmosphere, above 350 °C, a solid-phase quasi-intramolecular redox reaction takes place when the dimethylamine degradation products react with the W=O bonds with formation of oxidative coupling products of the organic fragments and reduced tungsten oxide with WO~2.93 composition. In synthetic air, above 350 °C, burning of organic fragments takes place, there are no oxidative coupling products and reduced tungsten oxide formation, and the end product of decomposition is monoclinic WO3.

Keywords Thermal studies · Quasi-intramolecular redox reaction · Dehydration · DSC · TG-MS · Paratungstate B

Introduction
Polyoxometalate compounds (POMs), especially polytungstates (POTs), represent an extensively studied area of material chemistry. There is an emerging interest about thermal decomposition of POMs having ammonium cations for the synthesis of metal oxides to be used as catalysts [1–4]. Alkylamino polytungstates [4–7], containing reducing alkylammonium cations with oxidizing polytungstate anions, are particularly interesting because of the possibility of self-oxidation, and because the cation may be volatilized. Recently, some polymorphic and solvatomorphic dimethylammonium dihydrogendodecatungstates have been synthesized [4, 8, 9]. The structure of these modifications is remarkably different, their water content is also different, and the cation–anion interactions [8] involve numerous bifurcated hydrogen bonds. As a result, their thermal properties are rather interesting, and their decomposition involves concomitant evolution of amine and water. The possibility of redox reactions between the oxometalate anionic cage and reducing N–H functional group of the cation offers further curiosities in their thermal behavior, which has not been established yet.
In the present work, continuing our work on dimethylammonium dihydrogenodecatungstate solvatomorphs, [8, 9] the results of combined thermal studies (TG, TG-MS, DSC) on (Me₂NH₂)₁₀[H₂W₁₂O₄₂]·10H₂O (compound 1a) and (Me₂NH₂)₁₀[H₂W₁₂O₄₂]·11H₂O (compound 1b) are presented.

**Experimental**

Chemical-grade ammonium paratungstate, 40% aq. dimethylamine solution and other analytical reagents were obtained from Deuton-X Ltd., Hungary.

WO₃ as precursor was prepared by the method of Fait et al. [10] with heating of ammonium paratungstate ((NH₄)₁₀H₂W₁₂O₄₂) at 600 °C for 10 min. The freshly prepared monoclinic WO₃ was reacted with 40% aq. dimethylamine solution at 50 °C for 2 h and the compound 1a was precipitated out with addition of ethanol. The white powdery material was recrystallized from water to obtain compound 1b. Both compounds were identified by their IR and XRD traces [8, 9].

X-ray powder diffraction measurements were performed on a Philips PW-1050 system with a Bragg–Brentano parafocusing goniometer. The apparatus was equipped with a copper tube (40 kV, 35 mA tube current), a secondary beam graphite monochromator and a proportional counter. Scans were recorded in step mode. The diffraction patterns were evaluated by full profile fitting techniques.

FTIR measurements were recorded on a Jasco FT/IR-4600 system at room temperature. The FTIR instrument was equipped with a Jasco ATR Pro One single reflection ATR accessory (incident angle 45°), furthermore with a DLATGS detector (4000–400 cm⁻¹ region, resolution of 4 cm⁻¹ and co-addition of 64 individual spectra). An ATR correction (Jasco Spectra Manager version 2, Spectra analysis module version 2.15.11) was performed on the raw spectra.

Simultaneous thermogravimetric, differential scanning calorimetric and mass spectrometric evolved gas analysis (TG-DSC-MS) measurements under inert conditions were performed on a Setaram LabsysEvo thermal analyzer, in high-purity helium (6.0) atmosphere, with a flow rate of 90 mL min⁻¹. The measurements were recorded in the 25–700 °C temperature range, with a heating rate of 20 °C min⁻¹, and samples were weighed into 100 µL alumina crucibles. The obtained data were baseline corrected and further evaluated by the thermoanalyzer’s processing software (Calisto Processing, ver. 2.01). Simultaneously with the TG-DSC measurement, the analysis of the evolved gases/decomposition products was carried out on a Pfeiffer Vacuum OmniStar™ quadrupole mass spectrometer coupled to the above-described TGA. The gas splitters and transfer lines to the spectrometer were thermostated to 220 °C. The measurements were taken in SEM Bargraph Cycles acquisition mode, in which the total ion current (TIC), the analog bar graph spectra (for structure determination), and the separate ion current of each scanned individual mass (96 masses) were recorded. The scanned mass interval was 5–100 amu, with a scan speed of 20 ms amu⁻¹, and the spectrometer was operated in electron impact mode. The measurements under synthetic air atmosphere were done with the same conditions and instruments.

**Results and discussion**

The thermal decomposition of compounds 1a and 1b were studied in inert atmosphere (helium) and in synthetic air. The decomposition temperatures, mass loss and the evolved gas fragments together with TG-DSC-MS characteristics are given in both atmospheres for compounds 1a and 1b in Table 1 and 2.

First, we summarize the general features of the heating curves and the mass losses of the two compounds in the two atmospheres. In the graphical representation shown in Figs. 1 and 2 and ESI Figs. 1–5, one can see that the TG curves measured for compound 1a in inert and oxidizing atmospheres are almost identical until about 300 °C. The mass loss in the oxygen-containing atmosphere becomes first (starting at about 330 °C) faster, but above about 380 °C, which is significant slower than in helium. The same holds for compound 1b, except that for the latter the steps are more expressed.

The multiple steps indicate that the leaving constituents (water and dimethylamine) evolve in several steps from both compounds, and their binding energies are different in the crystal lattices of 1a and 1b. The number of steps and the mass losses associated with each step are different for the two compounds. All reaction steps in inert atmosphere are strongly endothermic (Figs. 3, 4 and Table 1), while in synthetic air, the decomposition processes were found to be endothermic below and strongly exothermic above 300 °C (Figs. 3, 4 and Table 2).

Since in our compounds there are strongly reducing cations and oxidizing anions, one can expect a thermally induced redox reaction between them. The occurrence of such kind of redox reactions within one compound having oxidizing and reducing components is well documented in the literature [11–17]. Since in inert atmosphere the decomposition steps are endothermic, one can conclude that the energy required for elimination of water and dimethylamine as well as the disruption of the lattice exceeds the exothermicity of the redox interaction. The behavior of compounds 1a and 1b is similar, in spite of the important difference between their crystal structures [8].

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Now we turn to the detailed analysis of the decomposition curves. In inert atmosphere, the decomposition of compounds 1a starts with elimination of the two most weakly bound water of crystallization molecules/mol (m/z = 18, 17, 16) up to 150 °C (Fig. 5). Compound 1b, in contrast, loses 5 mol H₂O/mol already by 120 °C (Fig. 6). According to the XRD patterns, the lattice of compound 1a does not collapse due to removal of the first two moles of water (ESI Fig. 6), which means that the water is eliminated from the holes of the structure. The intensity of IR bands related to the water of crystallization molecules decreased (ESI Fig. 7) upon heating. The lattice of compound 1b also survives the loss of

| Step | Temperature range/°C | Δm/ % | DTG peak temperature/°C | DSC peak temperature/°C | ΔH/kJ mol⁻¹ | Main fragments/m/z values |
|------|----------------------|-------|-------------------------|-------------------------|-------------|---------------------------|
| 1a   | 60–155               | 1.04  | 140.5                   | 89.672                  | 18, 17, 16  (H₂O) |
| 2    | 155–269              | 8.73  | 200.6                   | 551.857                 | 18, 17, 16  (H₂O); 45, 44, 43, 42  (Me₂NH) |
| 3    | 269–399              | 6.69  | 361.1                   | 648.645                 | 18, 17, 16  (H₂O); 45, 44, 43, 42  (Me₂NH); 60, 59, 58 (coupling products) |
| 4    | 399–487              | 1.10  | 417.6                   | 134.248                 | 18, 17, 16  (H₂O); 45, 44, 43, 42  (Me₂NH, small); 30, 28 (NO, CO/N₂ as oxidation products, small); 60, 59, 58 (coupling products, small) |
|      | 487–703              | 0.49  | –                       | –                       | 18, 17, 16  (H₂O) |
| 1b   | 35–143               | 4.33  | 117.4                   | 210.534                 | 18, 17, 16  (H₂O) |
| 2    | 143–176              | 1.04  | 154.2                   | 36.904                  | 18, 17, 16  (H₂O); 45, 44, 43, 42  (Me₂NH, small); 30, 28 (NO, CO/N₂, oxidation product, small); |
| 3    | 176–283              | 6.99  | 234.7                   | 332.2                   | 18, 17, 16  (H₂O) |
| 4    | 283–409              | 6.07  | 365.4                   | 330.562                 | 45, 44, 43, 42  (Me₂NH) |
| 5    | 409–490              | 1.12  | 419.7                   | 105.447                 | 18, 17, 16  (H₂O) |
| 6    | 490–702              | 0.51  | –                       | –                       | 30, 28 (NO, CO/N₂, oxidation product, small) |

| Step | Temperature range/°C | Δm/ % | DTG peak temperature/°C | DSC peak temperature/°C | ΔH/kJ mol⁻¹ | Main fragments/m/z values |
|------|----------------------|-------|-------------------------|-------------------------|-------------|---------------------------|
| 1a   | 60–159               | 1.27  | 150.7                   | 151.0                   | 35.00       | 18, 17  (H₂O) |
| 2    | 159–264              | 8.38  | 199.3                   | 204.3                   | 535.22      | 18, 17, 16  (H₂O); 45, 44, 43, 42  (Me₂NH) |
| 3    | 264–344              | 1.73  | 332.2                   | 364.2                   | 3145.03     | 18, 17, 16  (H₂O) |
| 4    | 344–444              | 3.86  | 363.5                   | 464.0                   | 461.0       | 1732.99     | 44  (CO₂, N₂O) |
| 5    | 444–535              | 1.28  | 464.0                   | –                       | –           | –           |
| 6    | 535–698              | 0.40  | –                       | –                       | –           | –           |
| 1b   | 35–178               | 5.35  | 119.9                   | 124.5                   | 307.79      | 18, 17  (H₂O) |
| 2    | 152.8                | 6.77  | 230.4                   | 233.0                   | 427.09      | 18, 17, 16  (H₂O) |
| 3    | 178–262              | 5.35  | 281.3                   | 334.7                   | 378.3       | 1317.24     | 44  (CO₂, N₂O) |
| 4    | 300–347              | 1.32  | 334.7                   | 378.3                   | –           | –           |
| 5    | 347–443              | 3.75  | 365.4                   | 485.5                   | –           | –           |
| 6    | 443–523              | 1.66  | 484.3                   | 485.5                   | –           | –           |
| 7    | 523–698              | 0.15  | –                       | –                       | –           | –           |
the first 5 water molecules. The reduction of the intensities of the IR peaks of water of crystallization is more expressed for 1b than for 1a. It is obvious that compound 1a is thermally more stable than compound 1b and larger amount of water is eliminated from compound 1b before disruption of lattice (Tables 1, 2, ESI Fig. 3 and 4).

The more strongly bound water molecules also leave the crystals in the next endothermic decomposition steps, occurring at ~140/~200 °C and ~120/~160 °C (both in He and in synthetic air) for 1a and 1b, respectively. However, at these temperatures simultaneously with water, dimethylamine also evolves. The decomposition curves are almost the same in both inert and oxidizing atmospheres (Figs. 5, 6 and Tables 1, 2). From this, one can conclude that the aerial oxygen is not involved in these processes directly. The loss of the amine and the additional water molecules, however, disrupts the crystal lattice, and an XRD amorphous intermediate is formed (ESI Fig. 6). On further heating, two additional endothermic processes are observed for both compounds at around 348 and 365 °C in inert atmosphere (ESI Figs. 8–12). Besides the appearance of dimethylamine, its degradation and oxidative coupling products (m/z = 58 and 59), and as a consequence of the oxidation, formation of water can also be observed (Figs. 7, 8) and ESI Fig. 13–16).

In synthetic air, strongly exothermic processes take place at similar temperatures, indicating the oxidation of dimethylamine by the aerial oxygen. The analysis of the gas evolved from compound 1a in inert atmosphere showed that in inert atmosphere, no redox reaction takes place at 203 °C (Fig. 7) and only water (of crystallization) and dimethylamine peaks appear. Above 300 °C (Fig. 8), the intensities of dimethylamine peaks compared with those of water decrease with simultaneous appearance of oxidative condensation products of dimethylamine (m/z = 58 and 59). Since the water of crystallization is eliminated from polyoxometalates at temperatures below 250 °C, and the dehydroxylation temperatures [10, 14, 18] of the polytungstate cages are
above 400 °C, the source of water at around 300 °C can only be the polytungstate cage via the expected redox reaction between the amine/ammonium and W=O functionalities. Further evidence of the redox reaction is that according to the XRD pattern, the residue formed by heating up to 700 °C in inert atmosphere is reduced tungsten oxides like WO$_{2.93}$. The color of the final decomposition product (which is the same for both 1a and 1b) is green, confirming the presence of blue reduced Mo$^{5+}$ centers in the yellow WO$_3$ matrix. The amount of the products of the oxidation of dimethylamine,
namely CO$_2$ ($m/z$ = 44), CO ($m/z$ = 28), NO ($m/z$ = 30), N$_2$O ($m/z$ = 44) or NO$_2$ ($m/z$ = 46), cannot be determined precisely because their characteristic peaks coincide with those of the main fragments of Me$_2$NH (e.g., $m/z$ = 44 (Me$_2$N) and 28 (CH$_2$N) (ESI Fig. 17). Note, however, that the intensities of the peaks at $m/z$ = 44, 28 and 30 relative to each other and to the Me$_2$NH peak at $m/z$ = 45 differ from those observed for dimethylamine (ESI Fig. 17).

Since the majority of the Me$_2$NH evolves at lower temperatures, it is reasonable to assume that the dimethylammonium ions that react above 300 °C with the polytungstate cage are the most strongly bound ones, and the strong interaction between the N–H and O=W functionalities might initiate a solid-phase quasi-intramolecular redox reaction between the organic cation and the oxometallate anion. It cannot be excluded, however, that the gaseous Me$_2$NH liberated is oxidized by the solid-state polytungstate cages. The
number and binding energy of dimethylammonium ions are different in the structures of compounds \(1\text{a} \) and \(1\text{b} \) [8], and thus it is not surprising that some differences can be seen in the mass losses and the temperatures of lattice disruption and the collapse of the polytungstate cages.

In the final phase of decomposition of compound \(1\text{a} \) that takes place in inert atmosphere at 441 °C, i.e., after the complete removal of organic components, evolution of some gaseous product is observed, which was found to contain only water (Fig. 9). The only source of water in this case can only be the polytungstate cage, namely its inner hydroxyl groups. From this, one can conclude that the polytungstate cage collapse completely at around 450 °C.

The total mass losses of the samples in inert atmosphere were found to be a bit higher than the theoretical value calculated for formation of WO\(_3\) from \((\text{Me}_2\text{NH}_2)_{10}\text{H}_2\text{W}_{12}\text{O}_{42}\cdot n\text{H}_2\text{O} \ (n = 10 \text{ or } 11)\. The reason for this, as mentioned above, is that the final decomposition product in the absence of external oxygen is WO\(_{2.93}\) reduced tungsten oxide.

The TG-MS decomposition curves observed in synthetic air for both \(1\text{a} \) and \(1\text{b} \) at low temperatures show the
same steps as in inert atmosphere, namely evolution of pure
water in the first step, followed by simultaneous evolution
of water and dimethylamine. At higher temperatures, the
gaseous products correspond to the oxidation of dimethyl-
amine (Figs. 10–12) until it is completely consumed. The
final product in this case proved to be pure monoclinic WO3
(ESI Figs. 6 and 7).

The MS-EGA spectrum obtained at 109 ºC (Fig. 10)
shows that the components of synthetic air, N2 (m/z = 28
(N2)) and 14 (N+), O2 (m/z = 32 (O2) and 16 (O+) and water
(m/z = 18 (H2O+), 17 (OH+), 16 (O+)) could be detected. The
intensity of the m/z = 16 peak is higher than the intensity of
m/z = 18 and 17 peaks, although the fragmentation pattern
of OH2+ shows that the peak intensities of the fragment ions
decreases in the OH2+ – OH+ – O+ order [19]. The relatively
more O+ formation can obviously be attributed due to the
fragmentation of the mother ion of aerial oxygen, O2 to yield
O+ also.

At 214 ºC (Fig. 11) the disruption of lattice (ESI Fig. 6)
and formation of dimethylamine (m/z = 45), its protonated form (m/z = 46) and fragments could be observed,
together with water evolution. No sign of high amount of
CO\textsubscript{2} (m/z = 44) evolution, the peaks of m/z = 28 may not be evaluated as CO + fragment due to the presence of aerial N\textsubscript{2} (m/z = 28).

At even higher temperature, 523 °C (Fig. 12), after the degradation products of organic part (m/z = 27, 29, 32, 14), water (m/z = 18, 17) and CO\textsubscript{2}/N\textsubscript{2}O (m/z = 44) could be detected. There are no dimethylamine peaks at this temperature, only the fragment ions from the nonvolatile organic products. The compound 1b shows similar behavior in synthetic air (ESI Fig. 18a–e). Only air and water could be detected at 131 °C without any organic fragment. A small amount of dimethylamine was formed between 166 and 236 °C, while the amount of amine evolved together with the degradation products was increased at 373 °C. At 471 °C, mainly degradation products formed with CO\textsubscript{2} as oxidation product without free amine formation. The volatile oxidative coupling products could not be detected at all (ESI Fig. 18a–e), which shows that the aerial oxygen oxidizes them.

When the heating rate is increased, the peak temperatures of thermal decomposition steps are shifted. In the case of compound 1a (ESI Fig. 19–21), the rate of mass loss drastically decreases with increasing the heating rate from 10 to 20 °C min\textsuperscript{-1}. There is no similar change in the case of compound 1b. In the DTG curves (ESI Fig. 20) can be seen that not only the peak temperatures are shifted, but the intensity of each peak is also changed. At 20 °C min\textsuperscript{-1} heating rate, in the case of compound 1a, new peaks appeared at 100 °C and 310 °C.

The DSC curves showed exothermicity for both compounds in the oxidation range (> 300 °C) due to uncontrolled burning processes, but in the low temperature range, only the DSC curves for compound 1a changed with varying the heating rate from 10 to 20 °C min\textsuperscript{-1}. According to the appearance of the new peak in the DTG curve around 100 °C, an endothermic peak appears in the DSC as well as the consequence of water elimination (ESI Fig. 19).

Comparing the type and relative amount oxidation and degradation products in inert atmosphere and aerial conditions for compounds 1a and 1b, it can be concluded that in air, the burning of liberated dimethylamine takes place. The lack of reduced tungsten oxides might be attributed to re-oxidation of the reduced tungsten oxides formed by air at this temperature [20].

**Conclusions**

The general characteristics of thermal decomposition are similar for the two solvatomorphs of decakis(dimethylammonium) dihydrogendodecatungstate ((Me\textsubscript{2}NH\textsubscript{2})\textsubscript{10}H\textsubscript{2}W\textsubscript{12}O\textsubscript{42}·10H\textsubscript{2}O and 11 H\textsubscript{2}O), but some differences could be observed in the amount of water eliminated in the first decomposition steps and in the resolution/shifting of the decomposition peak temperatures depending on the heating rate or atmosphere used.

The first step of decomposition is endothermic in both atmosphere and involving 2 and 5 water molecule elimination with ~150 and ~120 °C peak temperatures for the decahydrate and undecahydrate, respectively. The elimination of further water and dimethylamine was observed with increasing the temperature, and the lattice of compounds started to disrupt above 200 °C. Until 300 °C, these processes are endothermic in both atmosphere, the further decomposition processes at higher temperature are left to be endothermic in helium, but become exothermic in synthetic air atmosphere.

In helium atmosphere, above 350 °C, a solid-phase quasi-intramolecular redox reaction takes place when the dimethylamine degradation products react with the W=O bonds with formation of oxidative coupling products of the organic fragments and reduced tungsten oxide with WO\textsubscript{2.93} composition. In synthetic air, above 350 °C, a burning process of organic fragments takes place, there are no oxidative coupling products and reduced tungsten oxide formation due to oxidation of these by the aerial oxygen. The end products of decomposition are CO\textsubscript{2}, H\textsubscript{2}O and monoclinic WO\textsubscript{3}.

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