Mechanistic study of 1,1-dimethylhydrazine transformation over Pt/SiO₂ catalyst†

Andrei V. Smirnov, Pavel A. Kots, Maksim A. Panteleyev and Irina I. Ivanova

Catalytic oxidation of 1,1-dimethylhydrazine (UDMH) with molecular oxygen over Pt/SiO₂ was studied by in situ FTIR spectroscopy coupled with online MS monitoring of the gas phase. An unusual two-step oxidation process was detected in experiments with the pulse UDMH feeding: initial UDMH oxidation over a fresh platinum surface quickly terminates due to the blockage of active sites; a time-separated second oxidation step corresponds to combustion of the surface residue. This residue consists of \( \text{C} = \text{N} \) nitrile groups formed via decomposition of the products of non-oxidative UDMH conversion, such as dimethylamine. The two-step oxidation picture is observed over a broad range of reaction temperatures and oxygen to UDMH ratios.

Moreover, the concentration of \( \text{CO}_2 \) as a product of complete oxidation is sharply raised in the temperature range 280–300 °C from almost zero to the maximum level, while for non-platinum catalysts \( \text{CO}_2 \) concentration at the outlet increased monotonically.\(^6,7\) The reason for these effects is definitely connected with peculiarities of UDMH transformation over the Pt surface but it has not been discussed in detail. However, mechanistic insights are important for understanding the oxidation processes and selectivity control.

The surface chemistry of UDMH in non-oxidative transformation on Pt in a temperature-programmed regime was studied in ref. 8. UDMH decomposition was described in terms of \( \text{N} = \text{N} \) bond breaking and formation of amino and dimethyl amino groups. Hydrogenation of amino groups led to ammonia desorption; other products detected in the gas phase were \( \text{HCN} \), \( \text{C}_2\text{N}_2 \), \( \text{N}_2 \) and surface C. Since reaction products were detected under ultrahigh vacuum conditions, these results cannot be directly transferred to real “working” conditions.

FTIR spectroscopy is a useful tool to investigate compounds formed on the catalyst surface and follow successive transformation of different reaction products.\(^{5,11}\) In situ FTIR spectroscopy was successfully used for the study of photocatalytic oxidation of UDMH in a batch reactor over \( \text{TiO}_2 \) as the photocatalyst.\(^{12}\) Adsorbed \( \text{N}_2\text{O} \) species were detected as the main surface intermediate. It was concluded that accumulation of surface nitrates was responsible for catalyst deactivation in the long-term process.

Adsorption and oxidation of UDMH on the surface of the supported spinel (20%) \( \text{Cu}_9\text{Mg}_{17} \cdot \text{Cr}_2\text{O}_4/\text{Al}_2\text{O}_3 \) was studied by FTIR spectroscopy in ref. 13. An increase in the adsorption temperature from 200 to 300 °C resulted in the appearance of the bands above 2000 cm\(^{-1}\) attributed to \( \text{C} = \text{N} \) bonds pointing to some surface reactions. In the presence of oxygen at 200 °C...
these bands along with the band characteristic for adsorbed dimethylamine (DMA) were also observed. While at 300 °C only adsorbed water as a product of complete oxidation was present on the surface. Thus, catalytic oxidation of UDMH is accompanied by formation of some intermediate products which can be detected in an adsorbed state by FTIR spectroscopy. These species may serve as spectators or catalyst poisons due to irreversible binding to metal sites, or on the contrary participate in oxidation as true reaction intermediates. The exact role of those species in oxidation catalysis and their impact on the state of the Pt sites under working conditions are not well understood yet.

In the present work in situ FTIR spectroscopy coupled with MS detection of reaction products was used to study non-oxidative and oxidative UDMH conversion on Pt/SiO2 catalyst via dynamic analysis of the residue formation, transformation and degradation.

Experimental

The Pt/SiO2 sample with a platinum content of 3 wt% was prepared by incipient wetness impregnation of 3 g of silica (0.25–0.5 mm fraction, specific surface area of 110 m2 g−1) with 3 ml of 0.15 M [Pt(NH3)4]Cl2 solution followed by drying at 110 °C for 24 h and calcination at 500 °C for 3 h.

Interaction of UDMH with the catalyst surface was studied by in situ FTIR-spectroscopy. The setup for spectroscopic measurements included a Thermo IS-10 FTIR-spectrometer and heated flow cell (dead volume 1 cm3) described in ref. 14. The cell was equipped with IR transparent CaF2 windows and was used simultaneously as a reactor and IR cell. Gases at the outlet of the cell were analyzed by mass-spectrometer MS7-200.

Prior to in situ FTIR experiments the sample of the catalyst was pelletized into a 2 cm2 self-supported wafer ca. 10 mg cm−2 density. The wafer was placed in the IR cell and purged in He or O2/He flows 20 ml min−1 at 260 °C. UDMH was pulsed by syringe into the gas flow at the IR cell inlet. IR spectra were registered in the 1000–4000 cm−1 range with 4 cm−1 resolution, in situ FTIR spectra were measured with a 3 s acquisition time.

Catalytic experiments with variation of reaction conditions, such as reaction temperature, oxygen concentration and amount of UDMH introduced in the pulséd mode, were performed in the tubular quartz reactor with 30 mg catalyst loading. Gas flow rate was maintained at 10 ml min−1, molar oxygen content in O2/He mixtures was varied from 2 to 20%, the volume of UDMH pulses was 0.05–2.0 μl, the influence of temperature was studied in the range of 250–450 °C. Reaction products were analyzed by an online MS detector; ion current of m/z for individual ions was continuously monitored, overlapping signals were corrected.

Results and discussion

Catalytic oxidation of UDMH on Pt/SiO2

Oxidative conversion of UDMH over Pt/SiO2 catalyst was performed at 260 °C in a flow of 3% O2 in He. UDMH was fed into the reactor in the pulse regime, which allows the provision of more detailed information about the initial stages of the reaction and intermediate products in comparison with the flow mode.

This list of reaction products correlates well with results of numerous studies.1,6,15 MS detection (Fig. 1) revealed the oxygen uptake (m/z = 32) with appearance of total oxidation products as CO2 (m/z = 44) and water (m/z = 18, 17) along with nitrogen (m/z = 28) and hydrogen (m/z = 2). Then other reaction products were also detected, namely dimethylamine (DMA) (CH3)2NH (m/z = 44, 45), ammonia NH3 (m/z = 16, 17), methanol CH3OH (m/z = 31, 32, 29), formaldehyde CH2O (m/z = 28, 29, 30), nitrogen oxides NOx (m/z = 30), formaldehyde dimethylhydrazine (CH3)2NNHCH2 (m/z = 72), dimethylidiazene (CH3)2NN (m/z = 43, 58) and hydrocyanic acid HCN (m/z = 27, 26). Conversion of UDMH was close to 100% (m/z = 60).

Identification of reaction products was based on evaluation of the ion current intensity ratios for different ions. So, the first and third peaks on the curve with m/z = 44 are attributed to CO2, while the second peak is associated with DMA as it totally correlates with the peak on the m/z = 45 curve having intensity ratio characteristics for DMA. Negative peaks on the curve m/z = 32 correspond to oxygen consumed for oxidation, while the appearance of a positive peak indicates the presence of methanol at the outlet as evidenced by similar changes in the concentration of the ion with m/z = 31. Ammonia was identified on m/z = 16, 17 ions taking into account that the m/z = 17 ion also corresponds to water and the m/z = 16 ion presents in the spectra of various compounds, for example, methane or oxygen.

Fig. 1 UDMH conversion over Pt/SiO2 catalyst. O2 (3%) He flow, single UDMH pulse, 260 °C, mass-spectrometric detection of reaction products.
Analysis of the main features of UDMH conversion revealed a very unusual effect of a two-step oxidation process, as was mentioned in our previous work. After UDMH pulsing the peak of oxygen consumption was observed, and then oxygen concentration returned to the initial level. However, after a rather extended period the second stage of oxygen consumption occurs, accompanied by the release of molecular nitrogen, NO and CO₂.

This two-peak shape oxidation indicates that complex processes occur on the surface of the catalyst. Without an understanding of these processes it is impossible to draw up a reliable scheme of UDMH oxidation on Pt-containing catalysts. More detailed information about the intermediate products can be obtained by in situ FTIR spectroscopy performed at the same reaction conditions. The role of the catalyst support as well as peculiarities of the UDMH conversion on Pt/SiO₂ catalyst in non-oxidative and oxidative conditions were successively studied using this technique.

Formation of heavy residues during UDMH conversion on Pt/SiO₂ under non-oxidative conditions

UDMH adsorption on silica. UDMH pulsing followed by FTIR spectroscopy was firstly performed over pure silica in a He flow at 260 °C to check the possible influence of the catalyst carrier on the transformation of UDMH. Fig. 2 shows the main features appearing in the spectra during UDMH adsorption over a silica pellet.

All bands observed in the spectra can be assigned to well-known UDMH vibrations. Bands at 3339 and 1588 cm⁻¹ correspond to asymmetric stretching and deformation of the NH₂ group of UDMH, respectively. The group of unresolved bands in the 3050–2700 cm⁻¹ region are coupled with CH₃ asymmetric and symmetric stretching (ν₅ at 2961, ν₇ at 2816 and ν₈ at 2775) and overlapping combination band at 2869 cm⁻¹. Deformation vibration of CH₃ groups also appears at 1458 cm⁻¹. The broad band at 3209 cm⁻¹ could be explained through formation of dimeric species with hydrogen bonding (N–H⋯N)¹⁷ or by the (N–H⋯OSi) interaction between NH₂ groups and surface silanols. A negative band at 3742 cm⁻¹ due to perturbed Si–OH groups of the silica is clearly seen. The intensity of the 3742 cm⁻¹ band changes with time in accordance with UDMH bands, indicating that Si–OH groups participate in UDMH adsorption most probably via hydrogen bonding. For the first 3–11 seconds after the pulse all UDMH bands increase in intensity due to UDMH accumulation over the silica pellet. Minor fluctuations in the CO₂ vibration intensity at ca. 2350 cm⁻¹ in this time interval lie within noise level. Then for 17–44 seconds the amount of UDMH adsorbed gradually decreases because of relatively weak interaction with the surface. Prolongation of the flushing with the helium flow for 71 second leads to complete disappearance of all bands corresponding to UDMH. This reveals no residue formation and confirms that the surface of silica remains intact after contact with UDMH in non-oxidative conditions.

Thus, there are no visible signs of UDMH transformation over silica indicating that the carrier is completely inert. In correlation with this observation, MS analysis of the gas flow at the outlet of the IR-cell revealed the presence of pure UDMH. So we assume all further UDMH transformation products over Pt/SiO₂ catalyst originate through interaction with Pt sites.

UDMH interaction with Pt/SiO₂. Significant differences were observed in IR spectra after UDMH pulsing through the IR cell

![Image of IR spectra](image-url)
with Pt/SiO₂ pellet in pure helium flow. The surface species formed have been followed with in situ FTIR as shown in Fig. 3. The first few spectra in Fig. 3 show that part of UDMH adsorbs molecularly on the Pt surface and silica carrier with asymmetrical stretching and scissoring vibrations at ca. 3339 and 1588 cm⁻¹ typical for the NH₂ group. Also other bands corresponding to unreacted UDMH located at 3209, 2869, 2816, 2775 and 1458 cm⁻¹ appear in the spectra 3 and 6 seconds after the pulse. Bands corresponding to UDMH grow to a maximum intensity after 6 seconds and then drop to nearly zero intensity after 35 seconds.

Also, during the first 3–6 seconds after the UDMH pulse, the emission of CO₂ was detected at ca. 2340 cm⁻¹ (Fig. 3). As the Pt/SiO₂ sample was used without preliminary reduction it could be proposed that some adsorbed oxygen can participate in UDMH oxidation as was previously observed for other molecules, in particular, HCN adsorbed onto fresh Pt-containing catalysts.¹⁸ In contrast to the experiment with pure silica, new bands due to products of UDMH transformation on Pt/SiO₂ appear in the spectra already after 3 seconds. A broad and asymmetric band at 1653 cm⁻¹ is coupled with C≡N double bonds in imine-like surface species formed via dehydrogenation of UDMH.¹⁸²⁶ This band remains in the spectra even after the disappearance of the bands corresponding to molecularly adsorbed UDMH. A group of bands in the 2050–2250 cm⁻¹ region have been previously observed and assigned to C≡N groups.¹³ So, bands at 2192, 2148 and 2097 cm⁻¹ could be corresponding to different nitrile species overlaying the Pt surface.¹³²⁷ The variation of the position of these bands could be explained by the diverse nature of the nitrile surface deposits. Some of them may form chains with unsaturated C≡N and C≡C fragments.

Nitrile and some isomeric species were observed over transition metal surfaces with similar C≡N triple bond wavenumbers. The strong interaction of the CN group with metal sites causes a decrease in nitrile vibration from the typical 2245 cm⁻¹ to ca. 2100 cm⁻¹ for surface Au nitriles, observed by Bamwenda et al.²² For nitrile metal-complexes of platinum in liquid phase the typical band position lies also at relatively low wavenumbers at ca. 2192–2165 cm⁻¹.²³ It also should be noted that C≡N vibration in cyanogen (C₂N₂) at 2168 cm⁻¹ (ref. 23) is close to bands observed in this study.

Considering the origin of several types of surface nitriles with corresponding band positions at 2097, 2148 and 2192 cm⁻¹ band maxima in our case, one can assume the following two reasons: (i) differences in Pt surface atoms coordination (edge, corner atoms or various crystallographic facets) give rise to a distribution of nitrile ligands attached to those sites; (ii) different degree of conjugation within CₙNₙ polymeric chains, so the C≡N wavenumber is decreased in longer and more conjugated species.²⁴ Interestingly, redistribution in intensity within a group of bands in the 2050–2250 cm⁻¹ region was observed. The band at ca. 2148 cm⁻¹ is a minor shoulder after 3 seconds, but became a dominant feature at 140 seconds after the pulse. Another band at 2097 cm⁻¹ remains constant during the whole time of the experiment, while the band at 2192 cm⁻¹ decreases in intensity. These findings may represent dynamic transformations within the pool of nitrile species on the Pt surface, which could convert from less stable to more stable ones. The total intensity of this group of bands remains unchanged after 140 seconds. Thus, the group of bands at 2050–2250 (C≡N) and 1653 cm⁻¹ (C≡N) were tentatively assigned as characteristic of deposits.

For the spectrum after 140 seconds, analysis of C–H and N–H stretching in the 3200–2700 cm⁻¹ region and deformation vibrations of NH₂ and CH₃ groups at 1588 and 1458 cm⁻¹, respectively, indicates nearly total absence of such vibrations in
residues. Definitely, strong dehydrogenating properties of Pt eliminate all hydrogen atoms from both CH₃ and NH₂ groups initially present in the UDMH molecule.

**CO probing of Pt accessibility.** To verify the accessibility of Pt before and after UDMH treatment *in situ* CO probing was performed. Carbon monoxide is known to form stable carbonyls over platinum on fully coordinated and uncoordinated metal and cationic sites. In the flow of He at 260 °C the principal carbonyl band was found in *ex situ* experiment at 2088 cm⁻¹ (Fig. 4a) simultaneously with gas-phase CO (centered at 2142 cm⁻¹) in line with numerous previous reports. This carbonyl band over Pt surface is generally assigned to CO linearly bonded to Pt⁰. No CO in bridged configuration or coupled with cationic Pt⁺ was found, suggesting a relatively small size for the platinum particles and a lack of platinum oxide. Characteristic vibration at ca. 2335 cm⁻¹ revealed formation of gaseous CO₂ probably due to oxidation of carbonyl species on Pt with adsorbed oxygen as was already observed for UDMH.

In order to probe the accessibility of Pt sites a *in situ* UDMH treatment, the UDMH pulse was followed by CO injection with ca. 30 seconds interval. UDMH pulsing caused formation of surface debris with characteristic bands maxima at 2148 and 2097 cm⁻¹ (Fig. 4b). Only gaseous CO molecules at 2142 cm⁻¹ were visible in the spectra over the UDMH pre-covered surface. Thus, carbonyls could not be formed after UDMH contact with the Pt/SiO₂ catalyst confirming the total platinum blockage with strongly adsorbed deposits.

Thereby, *in situ* experiments in He flow reveal that heavy residues are formed over Pt/SiO₂ in the early stage of UDMH conversion. The concentration of these species on the Pt surface remains constant over 140 s, indicating that these deposits adsorb strongly and are unreactive in non-oxidizing media. FTIR spectroscopic investigation shows that residues contain mostly C≡N conjugated bonds and C≡N nitrile triple bonds.

**In situ FTIR study of UDMH oxidation on Pt/SiO₂**

*In situ* FTIR experiments on UDMH oxidation were performed in helium flow with 3% oxygen. FTIR spectra of the products found on the surface of Pt/SiO₂ during UDMH transformation are presented in Fig. 5. The whole process can be divided into three stages: quick oxidation of UDMH in parallel with formation of the residue (0–14 s), elimination of reaction products adsorbed on the silica surface and evolution of residue (17–143 s) and, finally, oxidation of residue (146–150 s). This behavior totally correlates with the two-step oxygen consumption picture observed previously.

Extensive formation of CO₂ (band centered at 2348 cm⁻¹) and N₂O (vibration at 2222 cm⁻¹), with typical for gaseous species peak shapes, was observed during the first 3 seconds after the UDMH pulse (Fig. 5). Then generation of carbon and nitrogen oxides declines and is completely stopped. It is necessary to note that from nitrogen oxides only nitrous oxide is well resolved while mono- and dioxide are hardly detected, but their formation is also possible. Water as one of the possible products of oxidation is not visible in the spectra as its adsorption at the reaction temperature is rather weak.

Simultaneously with UDMH oxidation, formation of the residue takes place from the beginning of the UDMH contact with Pt/SiO₂ catalyst. The group of bands characteristic of surface residues in the 2050–2250 cm⁻¹ region (see Fig. 5, inset) have been previously observed over Pt in the case of oxygen-free conditions. For the first seconds after the UDMH pulse the

---

**Fig. 4** CO pulsing over Pt/SiO₂ sample: (a) fresh, (b) pre-covered with residues from UDMH conversion.
dominant band in this region is the 2097 cm\(^{-1}\) band associated with C\(\equiv\)N nitrile groups. Prolonged exposure for 14–17 seconds leads to additional bands at ca. 2148 and 2125 cm\(^{-1}\) assigned to other nitrile species bonded to Pt. The presence of other forms of surface nitriles at wavenumbers higher than 2200 cm\(^{-1}\) could not be totally ruled out due to overlay with gas phase N\(_2\)O bands. Band maxima match to values observed in the case of non-oxidative conditions in pure He flow. Analysis of the FTIR spectra reveals that UDMH oxidation and residue formation compete with each other over Pt sites. As a result, termination of CO\(_2\) and NO\(_x\) formation is connected with the total coverage of the Pt surface by nitrile-based deposits after 14 seconds.

From 17 to 143 seconds the most significant changes in the spectra were recorded in the areas of C–H and N–H vibrations. Already mentioned bands at 2961, 2869, 2816 and 1458 cm\(^{-1}\) due to CH\(_3\) groups and 1583 cm\(^{-1}\) due to NH\(_2\) groups were observed. More accurate examination reveals that the intensity ratio between various bands in the C–H stretching region significantly differs from UDMH observed on the surface of the SiO\(_2\) support. We suppose that those C–H vibrations arise not only from UDMH itself, but mostly due to the products of its conversion, namely, DMA. Separate in situ FTIR experiments with DMA pulsing verified that some bands absent in the spectrum of pure unreacted UDMH indeed correspond to DMA molecules (Fig. S1†).

Some new bands attributed to other products of UDMH transformation also appeared. So, the band at 3069 cm\(^{-1}\) corresponds to C–H stretching in the (\(\equiv\)C–H) fragment, and could be assigned, for instance, to formaldehyde dimethylhydrazine (\(\text{CH}_3\)\(_2\)N\(-\)N\(\equiv\)CH\(_2\)). Also, a new broad asymmetric band at 1709 cm\(^{-1}\) arises due to formation of imine-like surface species which could be coupled with C–N double bonds.

Intensities of C–H and N–H vibration bands decreased with time and almost totally vanished by 143 s. The negative band at 3740 cm\(^{-1}\) of Si–OH groups of silica changes in line with the C–H and N–H bands indicating that adsorption of these products occurs mostly via interaction with silanol groups of the carrier surface. The bands of C\(\equiv\)N groups at 2097, 2125 and 2148 cm\(^{-1}\) remained up to 143 seconds after the pulse, revealing their low reactivity. Some changes in intensity distribution between different C\(\equiv\)N vibration bands were observed with time. The intensities of the bands at 2148 cm\(^{-1}\) and 2125 cm\(^{-1}\) grow in the course of the experiment, while the 2097 cm\(^{-1}\) band intensity slightly decreased.

Dramatic changes in the spectra were observed from 143 to 146 seconds after the UDMH pulse. In this narrow time interval bands at 2148 cm\(^{-1}\) and 2097 cm\(^{-1}\), characteristic of Pt blocking residue, totally disappeared from the spectrum. Decomposition of the residue is accompanied with emission of gaseous CO\(_2\) (2348 cm\(^{-1}\)) and N\(_2\)O (2222 cm\(^{-1}\)). A small residual band at 2125 cm\(^{-1}\) remains in the spectrum, indicating the high stability of this type of nitrile.

After the disappearance of C–H and N–H containing substances (unreacted UDMH and some products of its
conversion) interacting with silanol groups on the silica support, the C,N$_2$ residue removal occurs. This could be traced from the changes in total area of all bands in the C–H stretching (3000–2700 cm$^{-1}$) and C=NN stretching (2200–2000 cm$^{-1}$) vibrations (Fig. 6). It could be proposed that the carrier supplies C–H and N–H containing species towards Pt thus maintaining blockage of the platinum surface by newly formed C=NN species.

**Mechanism of residue formation and oxidation**

Basing on the gas phase products variety detected by MS and adsorbed species registered by in situ FTIR, some consideration about the mechanism of UDMH oxidative transformation on Pt/SiO$_2$ and the nature of the residue could be proposed (Scheme 1).

The first step is UDMH adsorption on Pt (1). Then, two parallel processes can take place on the platinum surface. One is the expected step of direct interaction of UDMH with oxygen atoms adsorbed on Pt (2), resulting in the products of total and partial oxidation. Another way is UDMH dehydrogenation leading to appearance in the gas phase of hydrogen and traces of dimethylazene (3–6). According to in situ FTIR data, the oxidation reactions (2) stopped rather quickly as Pt appeared to be blocked by the residue and becomes unavailable for oxygen activation.

Adsorbed hydrogen species [H] generated by dehydrogenation reactions participate in typical for Pt catalysts hydrogenolysis processes, followed by ammonia (7) and DMA (8–9) formation and their release in the gaseous phase. [(CH$_3$)$_2$N] adsorbed species undergo transformation via dehydrogenation and C–N bond cleavage reactions (10) resulting in [HCN] and [CH$_3$] species with further release to the gas phase as HCN (11) and methane (12). Furthermore, [CH$_3$] can be the source of methanol (13, 14) and formaldehyde (15, 16) appearing in the gas phase. Formaldehyde in the gas phase is detected in a very narrow time interval after the UDMH pulse while methanol release becomes significant in line with decreasing formaldehyde content. Apparently, the deactivated Pt surface could not produce dehydrogenation products. Formation of formaldehyde dimethylhydrazene (CH$_3$)$_2$NNCH$_2$ is described by different mechanisms, other possible ways are also proposed (17–19).

Direct involvement of DMA in residue formation was confirmed by a separate run of DMA conversion over Pt/SiO$_2$ (Fig. S2†). Upon contact with DMA, surface nitrile groups are immediately formed with bands close to typical for UDMH-derived species. Also, residues formed from DMA are stable in oxidizing conditions and the two-step oxidation picture as in the case of UDMH oxidation is observed. So, the DMA-mediated path clearly leads to very similar surface residues according to their reactivity, composition and vibrational spectra.

According to a number of studies, [HCN] formed from adsorbed dimethylamino-groups undergoes dissociation on Pt (20) resulting in [CN] species strongly adsorbed on the Pt surface. It can be considered that these [CN] species could be a building block for oligomerization and formation of cyanogen ([CN]$_2$) and ([CN]$_x$) polymeric species with $x > 2$ (21). Formation of C=NN nitrile groups in a different environment results in variation of corresponding bands in the 2100–2150 cm$^{-1}$ region in the IR spectra (Fig. 4).

Most probably, the residue oxidation (22) occurs similarly to the reactions summarized in ref. 18 for HCN oxidation on P/Al$_2$O$_3$ catalyst. Under this scheme, oxidation by [O] adsorbed proceeds via [NCO] formation followed by its decomposition to [N] and [CO] adsorbed with a further number of surface reactions leading to the evolution of N$_2$, NO, NO$_2$, N$_2$O and CO$_2$ to the gas phase. But the key step is [O] formation, which is definitely limiting the residue oxidation stage.

Dissociation of [O$_2$] requires the presence of free platinum sites to form [O] adsorbed species. In the case where the entire surface of the platinum is totally blocked with deposits, the availability of such sites is complicated. It can be assumed that platinum sites, uncovered due to some rearrangement of overlaying (CN)$_x$ chains on the platinum surface, will be immediately covered with UDMH or the molecule of its transformation, such as DMA, transferred from the silica support. And when the amount of CN-containing molecules adsorbed on the support is significantly reduced, formation of the clean platinum areas becomes possible. This initiates O$_2$ adsorption and dissociation, after that, according to FTIR data, surface residues rapidly disappear via combustion. As soon as fewer residues remain on the surface, more sites for oxygen activation appear leading to oxidation rate increase and avalanche-type combustion. And it is this effect that we observe as the second step of the oxidation process.

**Effect of oxygen and UDMH concentration**

Experimental results showed that both oxygen activation and UDMH transformation to residue proceed on the same platinum centers. So, the oxygen to UDMH ratio and reaction temperature should affect significantly the process of UDMH oxidation, in particular, the time of the presence of deposits on the catalyst surface. Fig. 7 shows oxygen consumption and CO$_2$ evolution vs. time revealed by MS detection. As was mentioned before, peaks 1 and 3 on the m/z = 44 curve are attributed to CO$_2$, while the peak 2 is due to DMA formation; negative peaks on the m/z = 32 curve point to oxygen consumption, while the positive peak is associated with the presence of methanol.

![Fig. 6](#) Total area of all bands in C–H and C=NN stretching regions vs. time after UDMH pulse in 3%O$_2$/He flow.
Two-step O2 consumption and CO2 formation is observed over a broad range of oxygen concentrations. The lowering of oxygen content in the flow (Fig. 7a) leads to a smaller amount of total oxidation products formed in the first step (peak 1). So, in 3% O2 flow or less peak 1 becomes hardly detectable along with the increase in intensity of peak 2 owing to DMA (Fig. 7a). This corresponds to a decrease in surface coverage of \([\text{O}_2]\) and \([\text{O}]\) at low oxygen partial pressures and, as a consequence, initial oxidation upon UDMH dosing is less pronounced. An increase in the oxygen partial pressure also leads to the appearance of methanol in the interval between peak 1 and 2, indicating a higher methyl group conversion via reactions (13, 14) depicted in Scheme 1.

The amount of UDMH fed in the pulse (\(V_{\text{UDMH}}\)) was found to affect overall oxidation (Fig. 7b). The increase in \(V_{\text{UDMH}}\) leads to a smaller amount of oxidation products in the first step. In contrast, decreasing \(V_{\text{UDMH}}\) leads to vanishing of the DMA-connected peak 2 and only CO2 evolution is observed (Fig. 7b). This indicates that the UDMH to O2 ratio at the beginning of the pulse is even more important. When a certain ratio is exceeded, excess of UDMH is converted to partial decomposition products, such as DMA with consecutive formation of heavy \([\text{CN}]_x\) residues. These residues burn after a time gap leading to peak 3 in Fig. 7. Peak 3 intensity remains nearly constant at \(V_{\text{UDMH}}\) 0.5–1.3 μl, indicating limited capacity of the surface towards \([\text{CN}]_x\) species. Only peak 2 is growing in this interval, suggesting that any “extra” UDMH converts to partial decomposition products.

The time between first and second oxidation stages is continuously decreasing with increase of oxygen concentration or decrease of UDMH pulse as a result of the shift of competitive adsorption between these molecules and oxygen. Moreover, the higher oxygen content promotes more intensive combustion of the reagent during the first stage of oxidation thus reducing the amount of remaining adsorbed molecules, which could be further transferred to platinum to maintain their blockage. So, the increasing oxygen to UDMH ratio leads to the merger of two oxygen consumption peaks and, for example, after injection of 0.05 μl of UDMH to 3%O2/He flow at 300 °C the one-step oxidation process is observed (Fig. 7b). Hence, under these conditions the blockage of the platinum surface by residue does not occur.

Reaction temperature is also a significant factor for residue oxidation (Fig. 7c). The amount of CO2 released (peaks 1 and 3)
increases with temperature while DMA concentration detected in the gas phase (peak 2) declined and completely vanished above 350 °C. Herewith, the time between two CO₂ peaks diminished significantly when the reaction temperature was raised from 250 to 300 °C, but with further temperature increase almost no change in this parameter was observed. Despite the enhancement in oxidation rate, deposits formation and their separate combustion are still observed at all temperatures studied. The apparent dependence on reaction temperature is affected both by the rate of oxidation and by the rate of surface dehydrogenation. So, the superposition of these two components leads to observed similarity between at 350 and 450 °C.

Thus, experiments with variation of temperature and concentration of oxygen and UDMH led to an important conclusion for UDMH conversion over Pt-catalyst, which will be realized in flow conditions: deep oxidation is possible only when exceeding a certain threshold in the ratio of oxygen to UDMH. Below this level UDMH conversion is low and the reaction products are mainly products of non-oxidative decomposition of UDMH and partial oxidation, such as methanol. After exceeding this level, the UDMH conversion increases sharply, and the products of complete oxidation are mostly recorded in the gas phase. This threshold oxygen to UDMH level depends on reaction temperature, and namely this effect can be the reason for the above-mentioned fast increase in CO₂ concentration during UDMH conversion observed in ref. 6 and 7 at about 300 °C over Pt-catalyst.

Conclusions
An unusual two-step oxidation process was detected in experiments with the pulse 1,1-dimethylhydrazine (UDMH) feeding:
Initially UDMH oxidation takes place on the free platinum surface and terminates quickly due to the blockage of active sites; the second oxidation step is connected with combustion of the surface residue. In situ FTIR study of UDMH oxidation on Pt/SiO₂ catalyst allowed the revealing of peculiarities in the processes occurring on the catalyst surface. It was shown, that, in parallel with oxidation, UDMH undergoes a series of successive transformations leading to formation of deposits strongly adsorbed on the platinum surface. This residue consists mostly of condensed C≡N nitrile groups which totally block the platinum surface. The integrity of the deposits layer is maintained by C≡N species permanently formed via decomposition of UDMH and the products of UDMH conversion, such as dimethylamine, adsorbed on the silica support. The residue layer remains resistant to oxidation as long as the transfer of C–N containing substances from support to platinum proceeds. The two-step oxidation picture is observed over a broad range of reaction temperatures and oxygen to UDMH ratios. The period between the two oxidation stages increases with the amount of UDMH injected and lowering the oxygen concentration in the flow.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors thank the Russian Science Foundation for the financial support (grant no. 14-23-00094). P. A. Kots acknowledges Haldor Topsoe for a PhD fellowship.

References

1 Z. R. Ismagilov, M. A. Kerzhentsev, I. Z. Ismagilov, V. A. Sazonov, V. N. Parmon, G. L. Elizarova, O. P. Pestunova, V. A. Shandakov, Y. L. Zuev, V. N. Eryomin, N. V. Pestereva, F. Garin and H. Veringa, Def. Ind. Sci. Technol., A, 1995, 13(3), 1368–1372.
2 A. L. Schwaner, M. Kovar, D. J. Alberas and J. M. White, J. Vac. Sci. Technol., A, 1995, 13(3), 1368–1372.
3 J. P. H. Li, A. A. Adesina, E. M. Kennedy and M. Stockenhuber, Phys. Chem. Chem. Phys., 2017, 19, 26630–26644.
4 J. A. J. Rodrigues, G. M. Cruz, G. Bugli, M. Boudart and M. Odén, Catal. Lett., 2002, 75, 277–285.
5 G. Lunn and E. B. Sansone, Chemosphere, 1994, 29, 1577–1590.
6 Z. R. Ismagilov, Eurasian Chem.-Technol. J., 2001, 3, 241–245.
7 Z. R. Ismagilov, M. A. Kerzhentsev, I. Z. Ismagilov, V. A. Sazonov, V. N. Parmon, G. L. Elizarova, O. P. Pestunova, V. A. Shandakov, Y. L. Zuev, V. N. Eryomin, N. V. Pestereva, F. Garin and H. Veringa, Defense Industries: Science and Technology Related to Security: Impact of Conventional Munitions on Environment and Population, Kluwer Academic Publishers, Netherlands, 2004.
8 A. L. Schwaner, M. Kovar, D. J. Alberas and J. M. White, J. Vac. Sci. Technol., A, 1995, 13(3), 1368–1372.
9 A. Atakan, P. Måki, F. Söderlind, J. Keraudy, E. M. Björk and M. Odén, Phys. Chem. Chem. Phys., 2017, 19, 19139–19149.
10 J. P. H. Li, A. A. Adesina, E. M. Kennedy and M. Stockenhuber, Phys. Chem. Chem. Phys., 2017, 19, 26630–26644.
11 D. Panayotov, E. Ivanova, M. Mihaylov, K. Chakarova, T. Spassov and K. Hadjiivanov, Phys. Chem. Chem. Phys., 2015, 17, 20563–20573.
12 P. A. Kolinko, D. V. Kozlov, A. V. Vorontsov and S. V. Preis, Catal. Today, 2007, 122, 178–185.
13 I. Z. Ismagilov, V. V. Kuznetsov, A. P. Nemudryi and O. Y. Pod’yacheva, Kinet. Catal., 2004, 45(5), 722–729.
14 V. A. Matyshak and O. V. Krylov, Catal. Today, 1995, 25(1), 1.
15 M. A. Mathur and H. H. Sisler, Inorg. Chem., 1981, 20, 426–429.
16 A. V. Smirnov, M. A. Panteleyev, V. V. Krivetskiy and A. M. Gaskov, Russ. J. Appl. Chem., 2016, 89, 1109–1118.
17 J. R. Durig and W. C. Harris, J. Chem. Phys., 1969, 51, 4457–4468.
18 H. Zhao, R. G. Tonkyn, S. E. Barlow, B. E. Koel and C. H. F. Peden, Appl. Catal., B, 2006, 65, 282–290.
19 L. R. Knopke, N. Nemati, A. Kockritz, A. Bruckner and U. Bentrup, ChemCatChem, 2010, 2, 273–280.
20 P. Wu, T. Komatsu and T. Yashima, J. Catal., 1997, 168, 400–411.
21 J. S. Somers and M. E. Bridge, Surf. Sci., 1985, 159, L439–L444.
22 G. R. Bamwenda, A. Obuchi, S. Kushiyama and K. Mizuno, Stud. Surf. Sci. Catal., 2000, 130, 1271–1276.
23 G. Davidson, Spectroscopic Properties of Inorganic and Organometallic Compounds, Royal Society of Chemistry, Cambridge, 1998.
24 R. A. Nyquist, Interpreting Infrared, Raman, and Nuclear Magnetic Resonance Spectra, Academic Press, San Francisco, CA, 2001.
25 C. De La Cruz and N. Sheppard, Spectrochim. Acta, Part A, 1994, 50, 271–285.
26 J. Ruiz-Martinez, A. Sepulveda-Escribano, J. A. Anderson and F. Rodriguez-Reinos, Phys. Chem. Chem. Phys., 2009, 11, 917–920.
27 M. H. Moore and R. K. Khanna, Spectrochim. Acta, Part A, 1991, 47, 255–262.