The first study of the thermal and storage stability of arenediazonium triflates comparing to 4-nitrobenzenediazonium tosylate and tetrafluoroborate by calorimetric methods

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**KEYWORDS**

arenediazonium salts; stability; isothermal flow calorimetry; quantum chemical calculation; thermolysis

**ABSTRACT**

Herein, for the first time, using isothermal flow calorimetry and DSC/TGA, we have determined the thermal decomposition energies for the number of solid arenediazonium triflates comparing to 4-nitrobenzene tosylate and 4-nitrobenzentetrafluoroborate. The kinetics of thermal decomposition, activation energies, and half-lives of the studied diazonium salts were found. Using GC-MS and LC-MS, we have elucidated the products formed during thermolysis of the investigated diazonium salts. By DFT quantum chemical calculations at B3LYP/aug-cc-pVDZ level of theory we simulated the thermodynamics of decomposition reactions proceeding via substitution of diazonium group by corresponding nucleophiles. The method applied predicted the decomposition energies of all the studied compounds fairly precise, except for 2-nitrobenzene diazonium triflate. It has been found that 4-nitrobenzene diazonium triflate has increased storage stability under normal conditions comparing to the corresponding tosylate and tetrafluoroborate. The experimental and theoretical results demonstrated that comparing to DSC/TGA, isothermal flow calorimetry more adequately reflects the energetics of the thermal decomposition of diazonium salts and their storage stability under normal conditions.

**INTRODUCTION**

Aromatic diazonium salts (DSs) are versatile synthetic blocks widely used in fine organic synthesis and industry.1 Beyond that, DSs have been increasingly applied in development of macro- and nanoscale composite materials due to their reactivity towards metal and non-metal surfaces.2a

Nevertheless, such disadvantages as a poor storage stability in solid state and a propensity to explosive decomposition upon heating, photo-irradiation or mechanical stress limit the preparation and usage of DSs, especially, on an industrial scale.1a, 2b-d Several approaches to the stabilization of hazardous DSs, allowing for the minimization of risks associated with the processes of their production and utilization, are available. These include, for instance, preparation of polymer-supported DSs,2e freezing of benzenediazonium chlorides up to -84 °C,2f and application of flow-chemistry techniques for DS chemistry.2g Besides, to diminish the hazard level of DSs, twelve rules of handling these compounds were formulated.2b

Recently, we have synthesized arenediazonium tosylates ArN2+TsO- (ADTs)3a and trifluoromethane sulfonates ArN2+TfO- (ADTfs)3b that showed major advantages over traditional diazonium salts. While being surprisingly stable in solid state, they retained high reactivity in a vast array of diazonium chemistry reactions including the formation of aromatic iodides and bromides,3a,3b,4a-f azides 3b,4g and boronic acids,3b and Pd-catalyzed C-C-cross-compling.3b,4h-j Additionally, it has been shown that ADTs can be successfully used for introducing 19F isotope into the aromatic ring,5a carrying out azo-coupling with ethyl-methyl acetoacetate,5b substituting the diazonium group with the triethoxysilyl moiety,5c and covalently grafting aromatic groups to carbonized metal nanoparticles 5d and graphene.5e

The thermal decomposition energies of some ADTs and ADTfs determined by DSC/TGA, in most cases were found to be below 800 J/g. 3a,b, 4g Therefore, according to the safety criteria of the United Nations Economic Commission for Europe (UNECE), they can be referred as compounds that can be transported safely.6 However, the products of their thermal decomposition have not been studied and remain unknown. Moreover, DSC/TGA provides information about thermal decomposition energy at increased temperatures and does not reflect decomposition processes taking place under normal conditions. To comprehensively assess the possibility of safe use of ADTs and ADTfs in the laboratory and on an industrial scale, it is necessary to determine reliable quantitative characteristics of their storage stability in a solid state and thermal decomposition energies. Such evaluations should be done using various methods and comparing to other types of DSs.

Noteworthy, the DS decomposition in solution has been extensively investigated for a long time (see, for example,1a,g), whereas not many studies cover stability and safety of DSs in the solid state. The paucity of quantitative data, describing the decomposition of solid DSs, reflects the lack of reliable generally accepted procedures for measuring the above-mentioned properties.

A comparative study of the stability of the solid DSs (chlorides, tetrachlorozincates and tetrafluoroborates) was reported in 7a,c, however the decomposition products were not given. The investigation of the thermal decomposition of 14N- and 15N-substituted arenediazonium chlorides and tetrafluoroborates has shown that isotope effect is insensitive to the nature and position of substituents in the aromatic ring and the nature of the counterion.7b The stability test of arenediazonium chlorides has demonstrated that the sensitivity to detonation decreases from *ortho-*, through *meta-*, to *para-* substitution.7a It was shown that the nature of the substituent in the aromatic nucleus has a pronounced effect on DS stability as the nitro derivatives were significantly more sensitive to impact than the chlorine derivatives. The decrease in detonation sensitivity with an increase in molecular weight was observed, which was associated with a decrease in the specific value of the energy released per unit mass. The authors noted that the detonation sensitivity of DSs depends on many factors such as the size and shape of the crystals, as well as the presence of impurities. No correlation between detonation sensitivity and thermal stability was found.7a The values of the thermal decomposition energies of some ADTs and ADTfs, determined by DSC/TGA, do not correlate with the structural features of the diazonium cation or the nature of counterion. 3a,b, 4g The commonly occurred term “the storage stability of DS” has not yet been described quantitatively. To address this issue, in most cases the ability of DSs to be stored without changes for a certain time is indicated. Besides, to the best of our knowledge, up to the present, there are no theoretical methods for predicting the energies of thermal decomposition of DSs based on their chemical structure.

Our work aims to comprehensively address the challenges associated with the thermal and storage stability assessment of DSs. To achieve that, we for the first time have studied the kinetics and thermodynamics of thermal decomposition of arenediazonium triflates **1a-d** comparing to 4-nitrobenzenediazonium tosylate **2** and tetrafluoroborate **3** by DSC/TGA and isothermal flow calorimetry. Additionally, we have endeavored to develop the criteria for the evaluation of the storage stability of an array of diazonium salts with various counterions and substituents. The electron-withdrawing nitro group and the electron-donating methoxy group were chosen as substituents in the aromatic core of ADTfs **1a-d** for the following reasons. First, according to 7a, diazonium salts with NO2- moieties are the most explosive, therefore they represent the highest threshold of these properties. Secondly, DSs with NO2- and MeO- substituents in the aromatic ring differ sharply in their properties,3a,b i.e. these two examples should cover the widest range of properties studied.

We also aimed to determine the possibility to apply DFT quantum chemical calculations for the theoretical evaluation of the DS thermal decomposition and clarification of its mechanism. To the best of our knowledge, DFT methods have not been previously used for these purposes. To investigate the mechanism, a GC-MS and LC-MS study of the decomposition products of DSs **1a-d, 2, 3** was carried out. The obtained results are valuable for both applied and theoretical field of diazonium chemistry. On the one hand, they allow to assess the stability, capabilities and limitations of DSs for industrial use. On the other hand, they provide the better understanding of the mechanisms of DS thermal decomposition and allow to establish the structure-stability relationship.

**EXPERIMENTAL SECTION**

Arenediazonium triflates **1a-d** and 4-nitrobenzenediazonium tosylate **2** were synthesized according to the procedure described previously.3a,b 4-Nitrobenzenediazonium tetrafluoroborate **3** was purchased from Aldrich (CAS № 456-27-9). All samples used in calorimetric studies were dried in vacuum for 48 h. This procedure is recommended in the study of the decomposition of arendiazonium chlorides 7a.

The DSC/TGA runs were made in argon atmosphere using open sample pans on Q600 SDT instrument (TA Instruments), a heating rate of 5 °C·min-1, and a temperature range of 20-600 °C. A typical sample size was 10 mg.

Heat flow was measured under isothermal conditions in nitrogen atmosphere using TAM III microcalorimeter (TA Instruments). The experiments were conducted according to the conventional approach that is used for calorimetric studies of the safety of high-energy materials. 8 Sample was put in a glass beaker placed in a standard calorimeter ampoule made from *Hastelloy* with a volume of 1 mL. The ampoule was evacuated, then purged with nitrogen, argon or air depending on experimental conditions and sealed. The heat flow was measured at three different temperatures: 75, 80, and 85 °C. We conducted all flow calorimetry experiments at temperatures significantly lower than the melting points in order to adequately approximate kinetic curves to normal conditions. The isothermal test was carried out until the heat flux decreased below 2 μW. This value is less than 1% of the maximum heat flux and corresponds to the conversion degree α> 0.99. Time was varied from 1 to 50 days depending on sample and temperature. The acquired experimental curves were approximated using the model of an autocatalytic process and the Arrhenius equation. The experimental data were processed using TAM Assistant Software v1.3.0.153. Gnuplot 4.5 9a and R Statistics v3.3.3 9b were used for mathematical processing, statistical analysis, and dependencies building.

The DS decomposition products were studied by GC-MS on an Agilent 7890A\5975C instrument. The typical sample size was 50 mg. Samples were heated in a thermostat at 85 ° C for 14 days. Then an aqueous solution of KI was added to the sample to convert the undecomposed diazonium salts into the corresponding volatile aryl iodides,3a,b the products were extracted with ethyl acetate and organic layer was filtered through a silica pad. The obtained ethyl acetate extracts were then analyzed by GC-MS.

All LC-MS experiments were carried out on a high-resolution time-of-flight mass spectrometer Agilent LC-1260 MS QTOF 6530 equipped with electrospray ionization source (ESI) and atmospheric pressure chemical ionization source (APCI). A chromatographic method was developed using a Zorbax Eclipse Plus column (C18, 2.1x50 mm, 1.8 micron). The following gradient elution with water as “A” and acetonitrile as “B” was used at a flow rate of 0.25 mL/min: 0-40 min, 0% B  100% B followed by isocratic elution with B for 20 min. The operating parameters of the ESI-QTOF-MS were: gas flow rate, 9 L/min (N2); drying gas temperature, 350 °C; nebulizer, 35 psi; sheath gas flow, 11 L/min; sheath gas temperature, 300 °C, capillary, 3500 V; skimmer, 65 V; octopole RF voltage, 750 V; fragmentor 150 V; energy of collision 20 eV. The operating parameters of the APCI-TOF-MS were: gas flow rate, 6 L/min (N2); drying gas temperature, 300 °C; nebulizer, 35 psi; vaporizer temperature, 400 °C; capillary, 3500 V; skimmer, 65 V; octopole RF voltage, 750 V; fragmentor, 150 V; energy of collision 20 eV. For the LS-MS experiments, the DS decomposition products were dissolved in a water : acetonitrile mixture (1:1 v/v) at a concentration of 1 mg/mL. The volume of sample injected was 5 µL. Acquired LS-MS spectra were processed using the OpenMS 2.0 software package.10

For the theoretical study of suggested DS decomposition routes the quantum-chemical calculations were performed using Kohn−Sham density functional theory (DFT), global-hybrid GGA functional B3LYP and aug-cc-pVDZ basis set in Gaussian 09 software package.11 At the first step, the geometry of all molecules participating in the reactions was optimized. To prove the nature of the stationary points, the harmonic frequency calculations were done. Thereafter, the vibrational frequencies and thermodynamic corrections were calculated at normal conditions (25 ° C, 1 atm) and at temperatures used for isothermal decomposition experiment (75, 80 and 85 ° C).

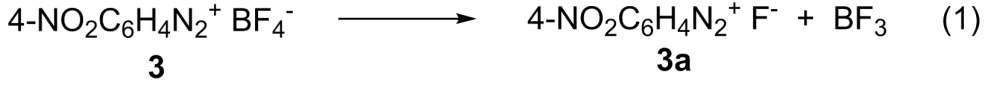
**RESULTS AND DISCUSSION**

**DSC/TGA results**

Our study of the thermal and storage stability of arenediazonium triflates **1a-d**,4-nitrobenzenediazonium tosylate **2** and tetrafluoroborate **3** has begun from collecting DCS/TGA data. Fig. 1-6 show DSC/TGA curves of thermal decomposition of DSs **1-3**. It can be seen that upon heating all the investigated DSs decompose with energy and weight loss. The temperatures and decomposition energies are summarized in Table 1.

For the DSs **1a, 1b, 1d, 3** additional low-temperature endothermic effects are observed in the temperature range close to their melting point, whereas DSs **1c, 2** decomposed only exothermically (Fig. 1-6, Table 1). Note that DS **2** decompose giving two exothermic peaks (Fig. 5). The first one at 69.37 °C is characterized by a small amount of heat released (24.47 J/g) and can be caused by the baseline issue, or crystallization from amorphous phase. While for the second one, located at 146.6 °C, the heat release of 323.0 J/g is observed. In the case of DSs **1a**, **1b**, **1d** the endothermic peaks are not accompanied by a weight loss and are probably associated with the rearrangement of the crystal lattice.

The weight loss upon heating DS **3** in the endothermic process, starting at 58.7 °C is 28.5%, which corresponds exactly to the elimination of volatile BF3 (28.6%) according to reaction (1):

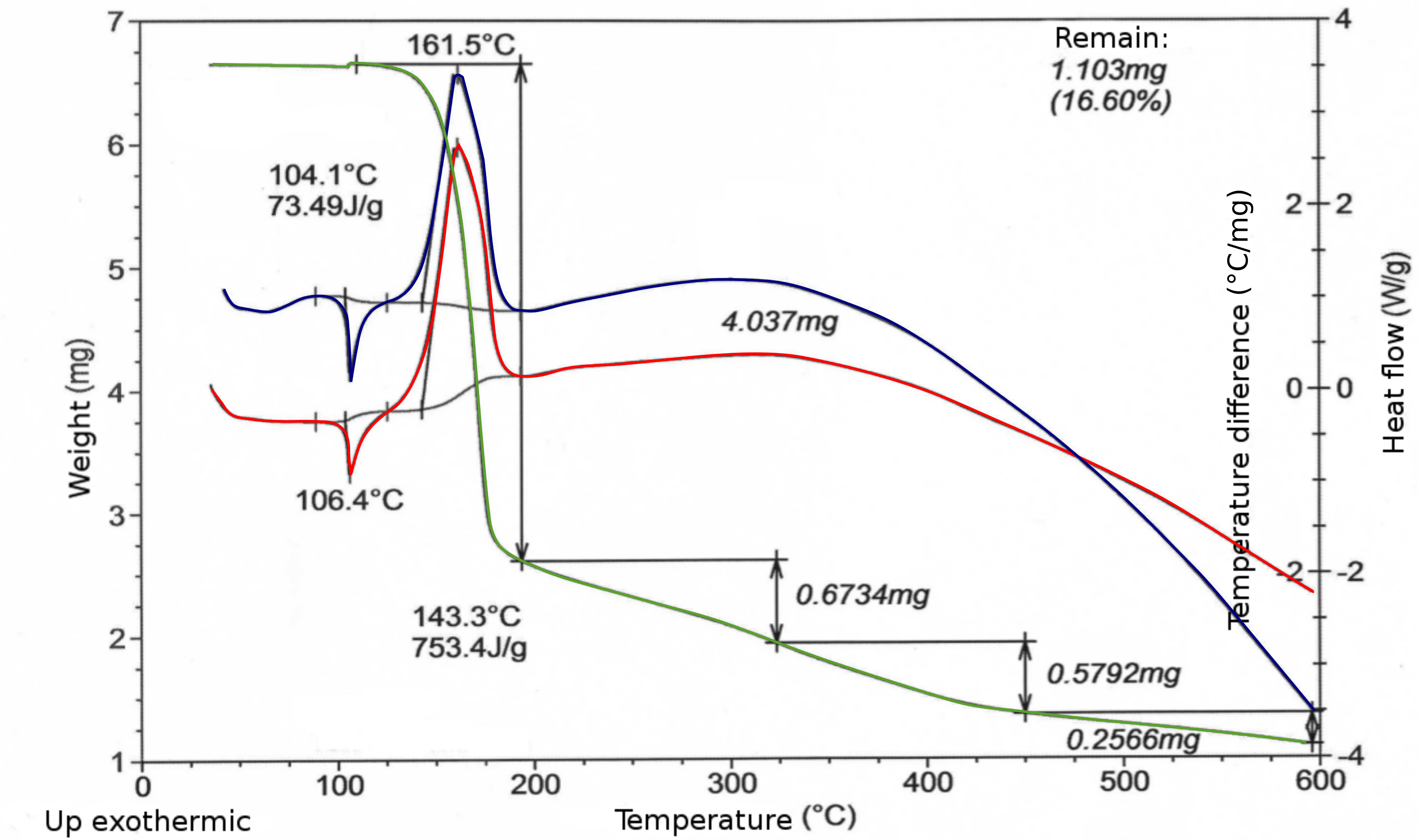


It might seem that the loss of BF3 at 58.7 °C indicates that intermediate 3a undergoes decomposition at 146.5 °С (equation 1). However, upon thermal decomposition of PhN2+BF4-, a small endothermic peak at 63 °C was also observed on the DSC/TGA curves. The appearance of this peak was explained by the removal of water bound to DS via hydrogen bonds. 12 It cannot be applied to DS **3** thought, as the weight loss of DS **3** at 58.7 °C corresponds to 5.2 mol of water per 1 mol of DS **3**. Should it be such a large amount of water, DS **3** would have to be partially dissolved, but not crystalline. Hence, there are contradictions in explaining the cause of the endothermic peak appearance during the heating of DS **3** and PhN2+BF4-. Obviously, this issue requires further special studies. Therefore, at present, we can propose equation 1 only as a hypothesis, partially consistent with the results of quantum-chemical modeling of DS decomposition reactions.

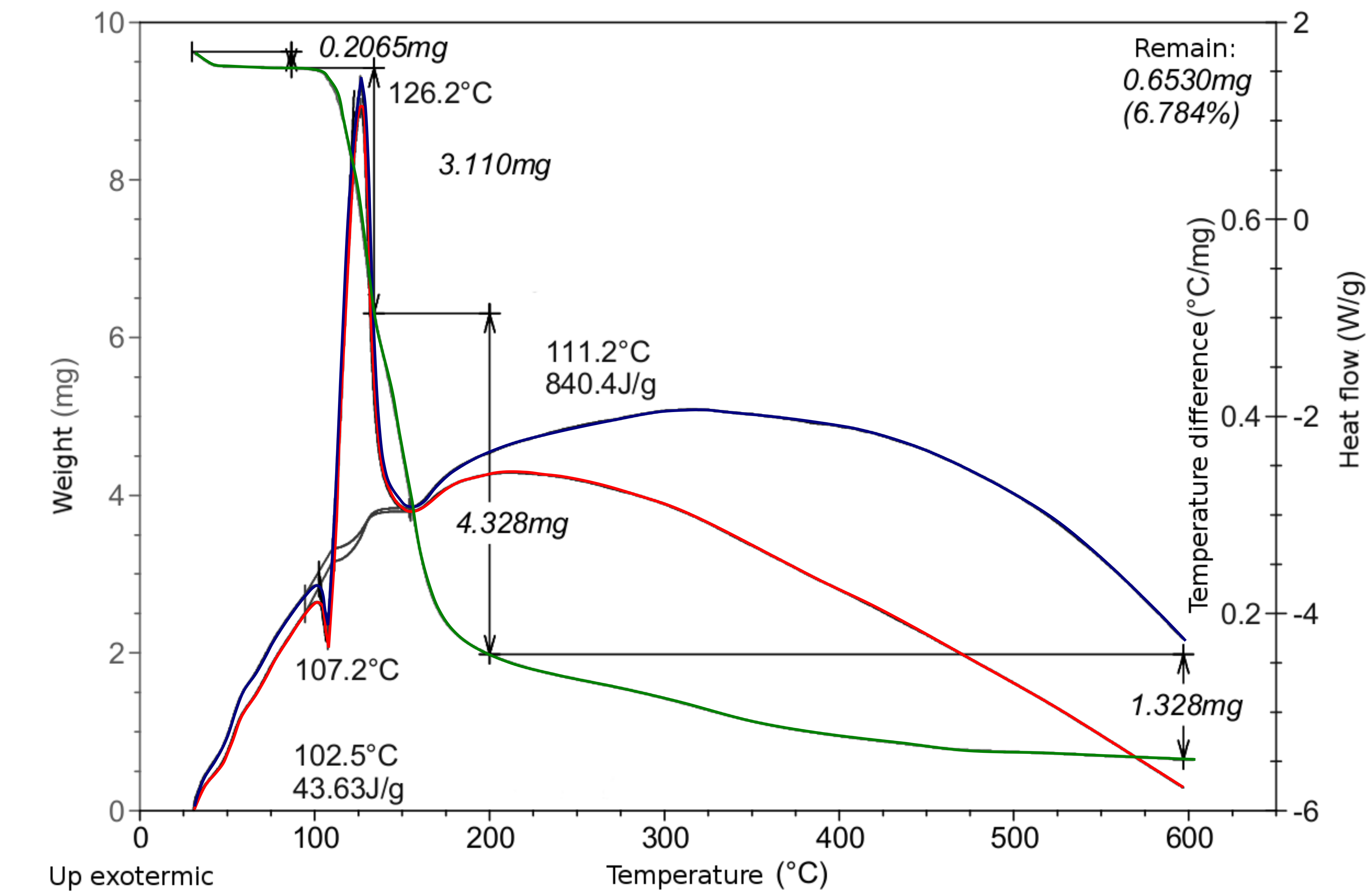
The major weight loss upon heating of DSs **1-3** occurs in exothermic processes, which is clearly associated with the formation of volatile decomposition products (Fig. 1-6).

**Table 1. Temperatures and decomposition energies of diazonium salts 1-3 according to DSC\TGA experiments data.**

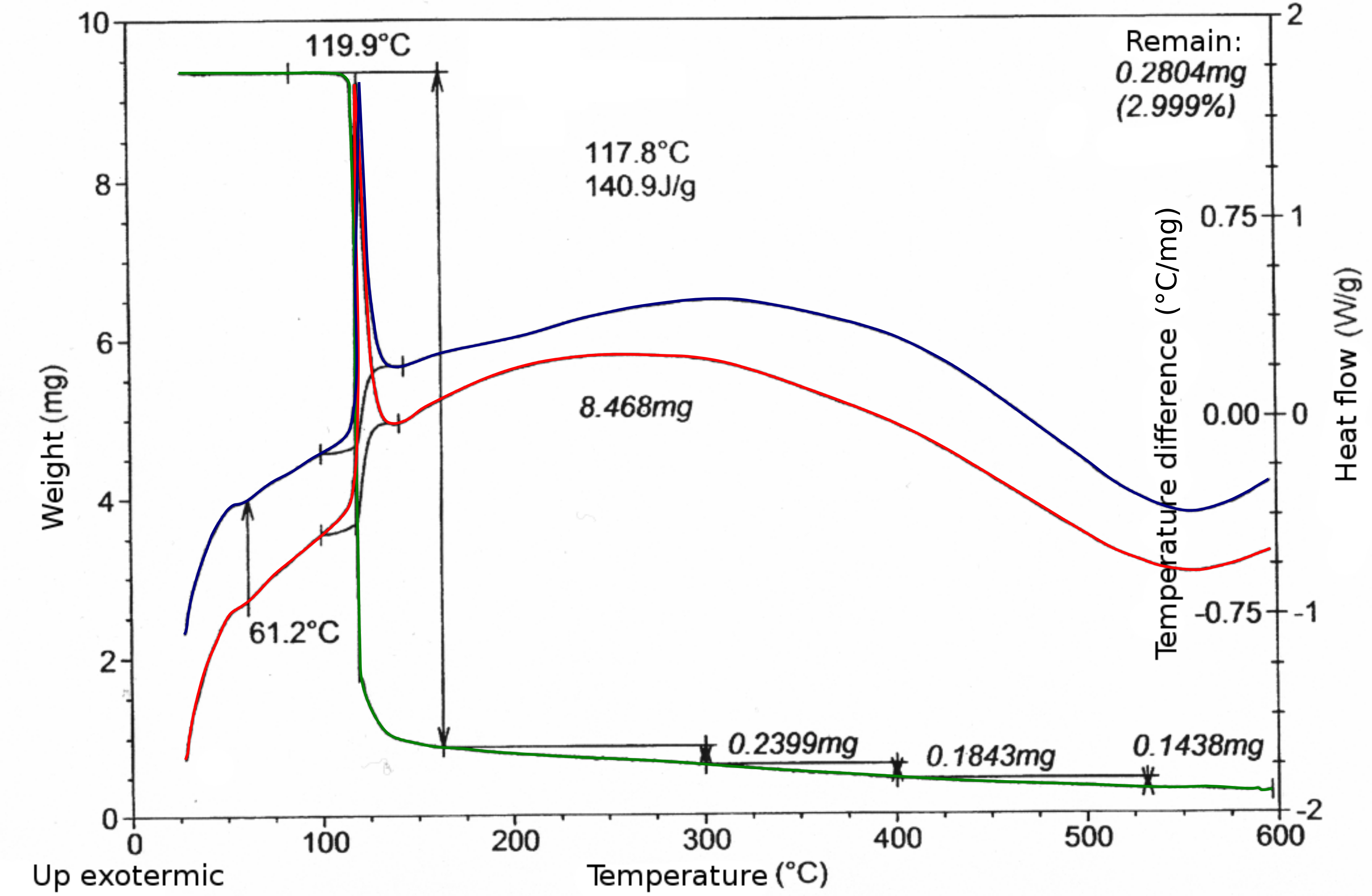
| Diazonium salts | Endothermic process | | Exothermic process | | Mp, (° C) |
| --- | --- | --- | --- | --- | --- |
| Т, ° C | Н, J/g (kJ/mol) | Т, ° C | Н, J/g (kJ/mol) |  |
| 2-NO2С6H4N2+TfO-  **1a** | 104.1 | 73.49 (21.98) | 143.3 | -753.4 (-225.3) | 110 |
| 3-NO2С6H4N2+TfO- **1b** | 102.5 | 43.6 (10.04) | 111.2 | -840.4 (-251.3) | 108-109 |
| 4-NO2С6H4N2+TfO- **1c** | - | - | 116.4 | -219.9 (-65.7) | 104 |
| 4-MeOС6H4N2+TfO- **1d** | 88.4 | 102.2 (29.05) | 136.6 | -328.9 (-93.5) | 94-97 |
| 4-NO2С6H4N2+TsO- **2** | - | - | 69.37  146.6 | -24.47 (-7.8)  -323.0 (-103.7) | 132 |
| 4-NO2С6H4N2+ BF4- **3** | 58.7 | - | 146.5 | -229.2 (-54.3) | 144 |



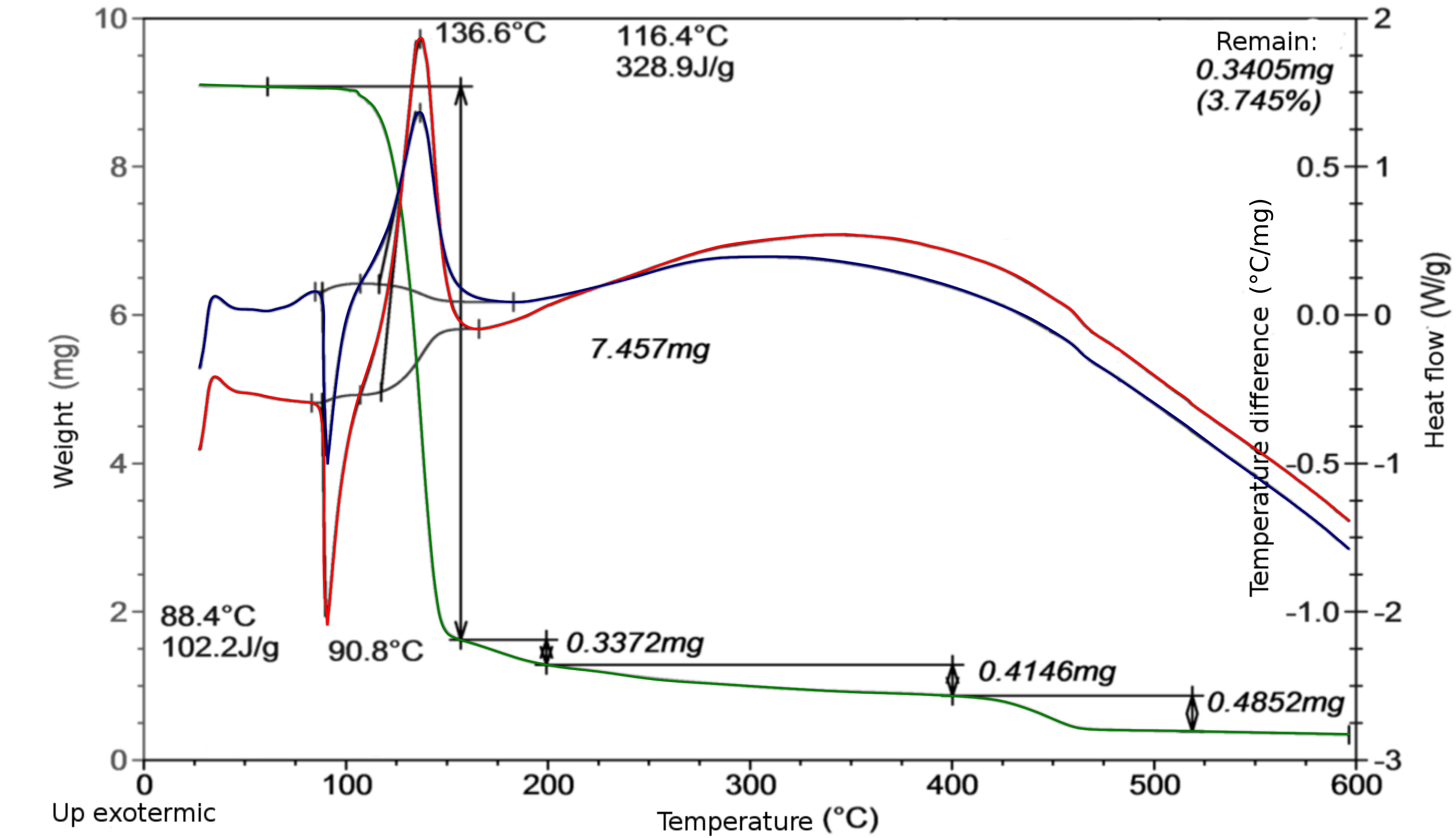
**Figure 1.** DSC/TGA thermograms of thermal decomposition of 2-NO2С6H4N2+TfO- **1a**. The temperature difference is shown in red; the sample weight is shown in green; the heat flow is shown in blue.



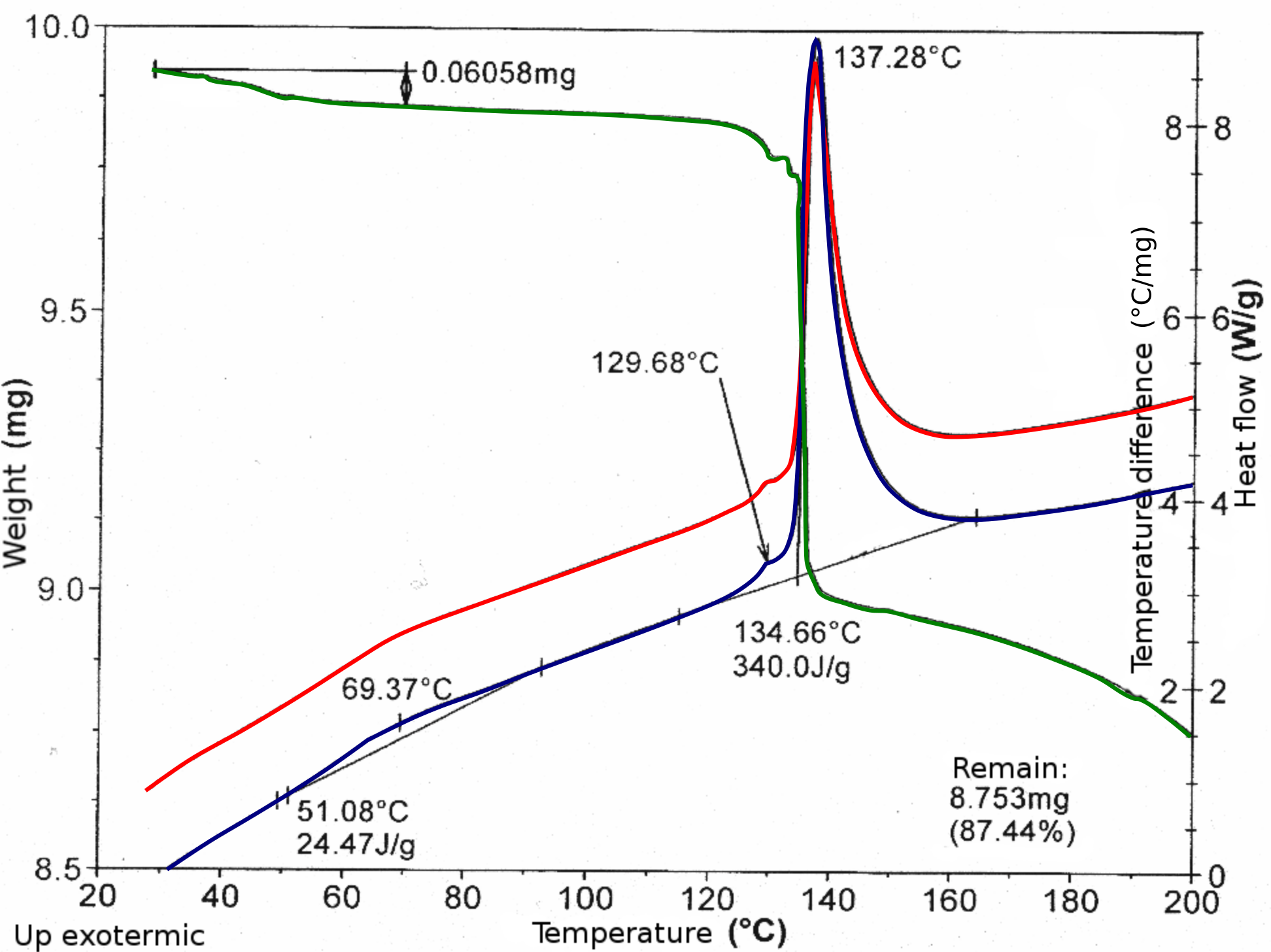
**Figure 2.** DSC/TGA thermograms of thermal decomposition of 3-NO2С6H4N2+TfO- **1b**. The temperature difference is shown in red; the sample weight is shown in green; the heat flow is shown in blue.



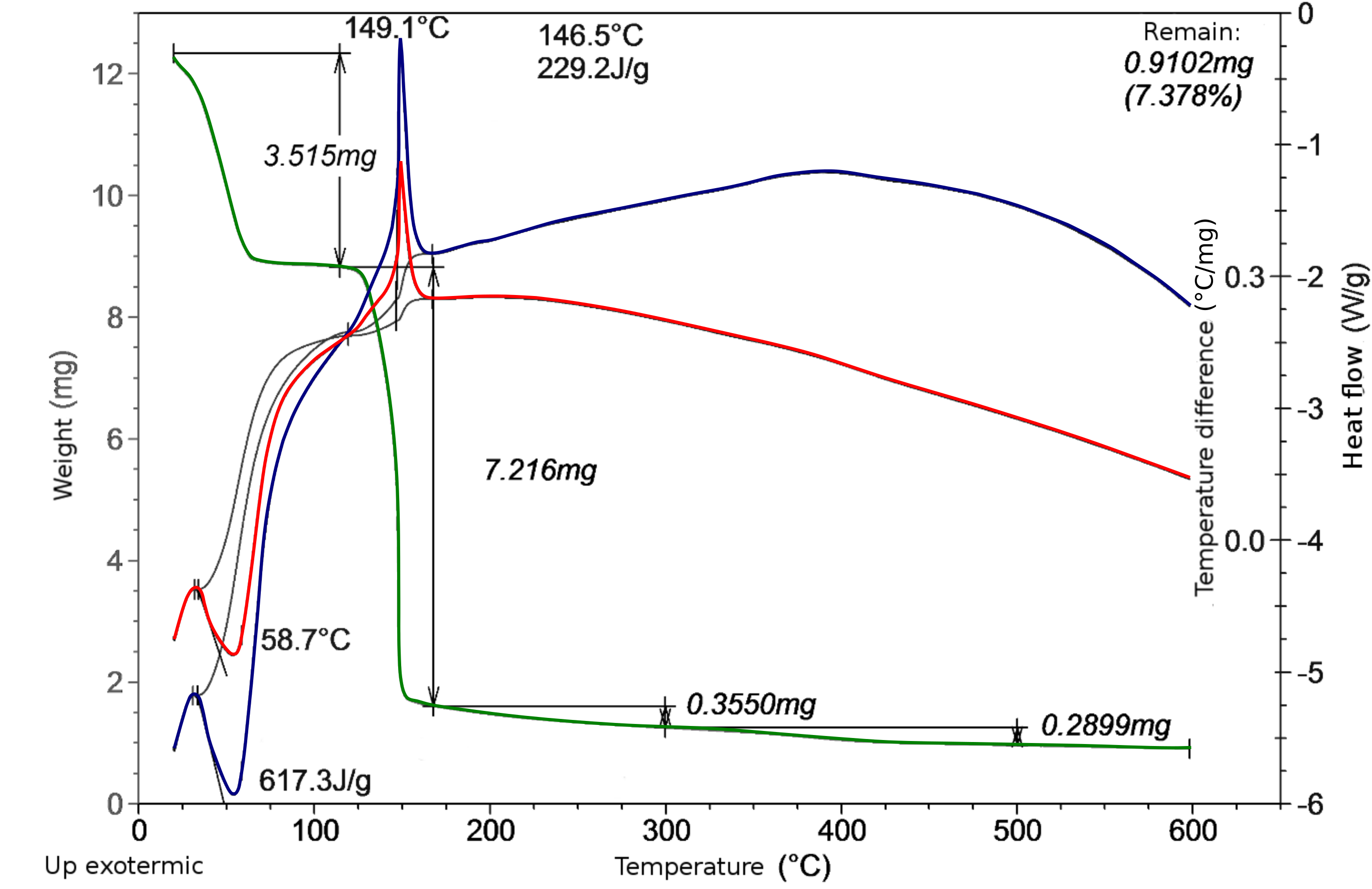
**Figure 3.** DSC/TGA thermograms of thermal decomposition of 4-NO2С6H4N2+TfO- **1c**.The temperature difference is shown in red; the sample weight is shown in green; the heat flow is shown in blue.



**Figure 4.** DSC/TGA thermograms of thermal decomposition of 4-MeOС6H4N2+TfO- **1d**. The temperature difference is shown in red; the sample weight is shown in green; the heat flow is shown in blue.



**Figure 5.** DSC/TGA thermograms of thermal decomposition of 4-NO2С6H4N2+TsO- **2**. The temperature difference is shown in red; the sample weight is shown in green; the heat flow is shown in blue.



**Figure 6.** DSC/TGA thermograms of thermal decomposition of 4-NO2С6H4N2+BF4- **3**. The temperature difference is shown in red; the sample weight is shown in green; the heat flow is shown in blue.

The exothermic effects Hexoterm of DSs **1-3** thermal decomposition are the most important for the DS safety evaluation. As can be seen from Table 1, these effects strongly depend on the nature and position of the substituent in the benzene ring and partly on the nature of counterions. Among salts **1a-d** with a triflate counterion, the energy release during decomposition is the highest for 3-nitro-derivative **1b** and decreases noticeably in the row **1b**>**1a**>**1d**>**1c**. Among DSs having the same 4-nitrobenzene diazonium cation and different counterions, the energy release decreases in the row **2c**>**1c**≈**3**. Therefore, the highest heat release is observed for DS with a TsO- counterion. However, a change in the counterion nature affects exothermic decomposition process substantially less than the position and type of substituents in the benzene ring.

**Isothermal flow calorimetry results**

With DSC/TGA data in our hands we moved on to isothermal flow calorimetry analysis of DSs studied. As the stability of DS could be affected by the presence of impurities, which are often cannot be controlled by conventional analytical methods,1a,7a we first decided to evaluate the reproducibility of thermal decomposition parameters measured. To do that we took samples of ADTS **2** from three synthetic batches obtained under identical conditions (samples 1-3) and samples of ADTS **2** purified by single and double precipitation of sample 3 from acetic acid/ether solutions (repricipitation 1 and 2). During the isothermal decomposition the following parameters have been determined: the integral enthalpy, initial and maximum heat flows, decomposition rate constants, and initial product concentrations approximated by equation 2. Additionally, we have varied the purged gas and studied sample decomposition in air, argon and nitrogen atmosphere. The results show that the collected data are consistent, and the results do not depend statistically significantly on the degree of sample purification. Even though there is a slight tendency to the reduction of the initial heat flow depending on the number of reprecipitations, the differences between the first and second reprecipitation are minimal. No qualitative and statistically significant differences were observed when decomposition was conducted in air, argon and nitrogen atmosphere. The average statistical deviation (RMD) for the rate constants (according to equation 2) and the enthalpy was 5%. Therefore, the further isothermal flow calorimetry experiments were carried out in a nitrogen atmosphere and after a single reprecipitation of the initial DS.

**Table 2. The reproducibility of the DS 2 thermal decomposition parameters measured at 85 ° C depending on synthetic batch and a number of reprecipitations.**

|  |  |  |  |
| --- | --- | --- | --- |
| Sample | k·103, s-1 | H, kJ·mol-1 | P0, mW/g |
|  | | | |
| Sample 1 Nitrogen (reprecipitation 1) | 0.2723 ± 0.0364 | 234.2 ± 3.0 | 31.2 ± 8.1 |
| Sample 2 Nitrogen, (source, reprecipitation 1) | 0.2568 ± 0.0371 | 243.6 ± 20.7 | 12.7 ± 3.5 |
| Sample 3 Nitrogen, (source, reprecipitation 1) | 0.2639 ± 0.0283 | 235.5 ± 7.0 | 10.1 ± 3.5 |
|  | | | |
| Sample 3 Air, reprecipitation 2 | 0.2683 ± 0.0333 | 229.0 ± 28.9 | 8.7 ± 8.5 |
| Sample 3 Argon, reprecipitation 2 | 0.3119 ± 0.0352 | 242.3 ± 8.0 | 12.8 ± 2.4 |
| Sample 2, 3 Nitrogen,  (source, reprecipitation 1, 2) | 0.2608 ± 0.0184 | 238.9 ± 7.8 | 11.2 ± 2.3 |
|  | | | |
| Sample 4 Nitrogen, source | 0.2589 ± 0.0340 | 241.7 ± 16.2 | 12.0 ± 2.0 |
| Sample 4 Nitrogen, reprecipitation 1 | 0.2608 ± 0.0492 | 235.1 ± 17.9 | 10.2 ± 6.3 |
| Sample 4 Nitrogen, reprecipitation 2 | 0.2898 ± 0.0221 | 238.1 ± 6.8 | 11.3 ± 2.3 |
|  | | | |
| General statistics | 0.2745 ± 0.0140 | 237.7 ± 4.6 | 13.8 ± 3.3 |

k – rate constant, H - integral enthalpy, P0 – initial heat flow

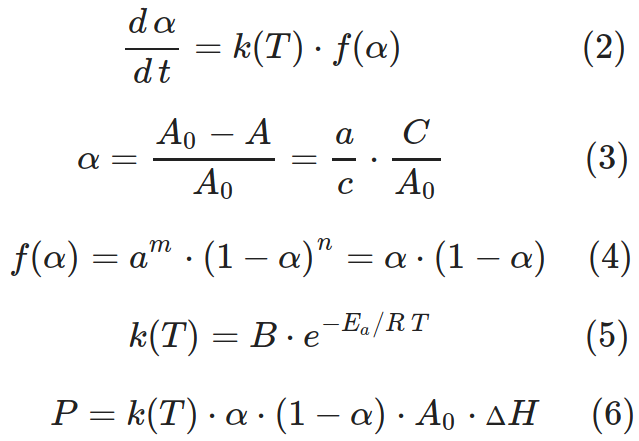
Table 3 and Fig. 7-10 show the results of isothermal decomposition of DSs **1-3** at75 оС, 80 оС, and 85 оС. At 75 оС, the maximum heat flow values obtained for the 4-nitrobenzene diazonium salts **1c, 2, 3** almost do not depend on the counterion nature. However, it can be noted that the heat flow curve acquired during DS **1c** decomposition is much steeper. Additionally, the Pmax values obtained for this DS at 80 and 85 оС are significantly higher than those for DS **2** and **3**. The Pmax values found for 4-methoxybenzenediazonium triflate **1d** are noticeably smaller comparing to 4-nitrobenzenediazonium triflate **1c.** Among the nitrobenzenediazonium triflates **1a-c**, the maximum heat flow decreases from *para-***1c**, through *meta-***1b**, to *ortho-***1a** substitution. The values of Pmax are important not only for mathematical modeling of the reaction kinetics, but also for quantitative description of the compound safety that is essential for practical application.

**Table 3.** Integral enthalpy and maximum heat flow values found during isothermal decomposition of diazonium salts **1-3**.

| Diazonium salt | ∆H, kJ/mol | | | Pmax, mW/g | | |
| --- | --- | --- | --- | --- | --- | --- |
| 75 oC | 80 oC | 85 oC | 75 oC | 80 oC | 85 oC |
| 2-NO2С6H4N2+TfO- (**1a**) | 414.0 | 386.0 | 396.0 | 0.705 | 1.49 | 2.33 |
| 3-NO2С6H4N2+TfO- (**1b**) | 227.9 | 230.0 | 225.4 | 6.57 | 14.15 | 28.8 |
| 4-NO2С6H4N2+TfO- (**1c**) | 200.0 | 235.1 | 250.0 | 20.12 | 47.89 | 119.5 |
| 4-MeOС6H4N2+TfO- (**1d**) | 183.1 | 183.2 | 106.0 | 1.34 | 2.97 | 5.37 |
| 4-NO2С6H4N2+TsO- (**2**) | 253.0 | 232.4 | 231.0 | 21.10 | 34.82 | 64.60 |
| 4-NO2С6H4N2+ BF4- (**3**) | 173.0 | 156.0 | 147.0 | 20.02 | 31.63 | 66.87 |

∆H – integral enthalpy, Pmax – maximum heat flow value

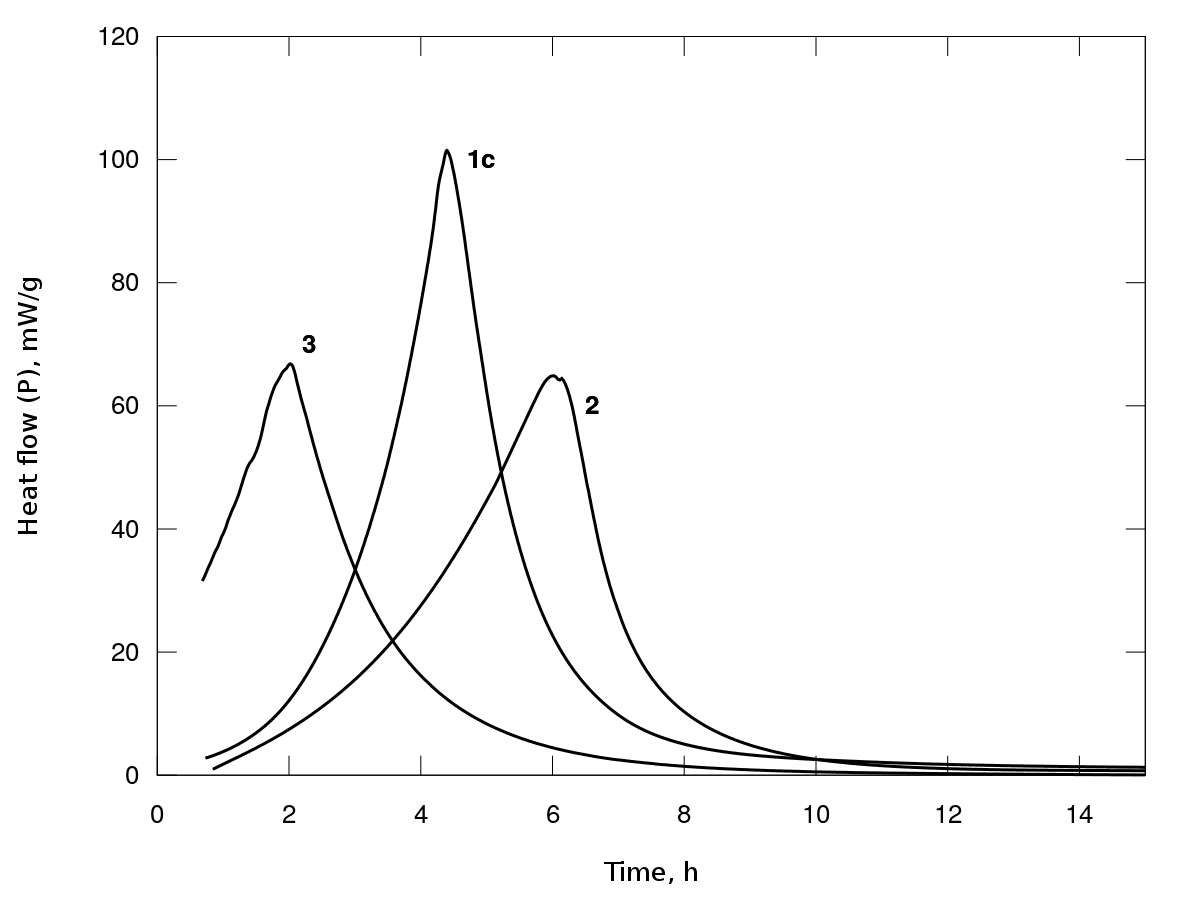
To investigate the kinetics of isothermal decomposition (Fig. 7-10), we approximated the experimental heat flow curves with kinetic equation for autocatalytic reactions (2) that qualitatively describes the heat flow over time dependency. The following main parameters were determined: *k* - rate constant, and P0, Pmax - values of the initial and maximum heat flows in the autocatalytic reaction equation 2. Table 4 shows calculated kinetic parameters found as result of approximation of the experimental heat flow curves of DSs **1a-1d, 2, 3**. The following kinetic equation describing autocatalytic reaction was applied: ***a A → c C***

,

where α – conversion degree; *k(T)* – rate constant, [s-1]; *f(α)* – the kinetic model in differential form; *A0*, *A* – initial and current concentration of DS, which for solid-phase reactions is measured in mol/g units; *С0*, *С* – initial and current concentrations of products, [mol/g]; *a*, *c* – stoichiometric coefficients, for the processes studied *a/c*=1; *P*, *P0* – current and initial heat flow; *ΔH* – reaction enthalpy; *B* - is the pre-exponential factor, a constant for each chemical reaction; *Ea* - an experimentally determined parameter that indicates the sensitivity of the reaction rate to temperature; *R* - the universal gas constant (8.315 J⋅K−1⋅mol−1); *T* - the absolute temperature (in K).

The kinetic equation 2 is used to model solid phase reactions 13, and can be expressed in terms of the conversion degree α (equation 3). The kinetic curves for the reactions studied are best described by equation 4 at *m*=1 and *n*=1. The temperature dependence of the reaction rate is expressed by the Arrhenius equation 5. The reaction enthalpy *ΔH* is calculated by integrating a curve approximating the experimental heat flux (equation 6), in the time interval (0, ∞). The half-life corresponds to the conversion degree of 0.5. Due to the symmetry of the approximating kinetic curve 4, the half-life coincides with the maximum heat flux (in the case of more complex kinetics, this rule is not fulfilled).

Figure 7 shows the experimental heat flow curves describing the isothermal decomposition of DSs **1c**, **2** and **3** at 85 ° C. As can be seen, the half-life of DSs depends on the counterion: for tetrafluoroborate **3** it is 2h, for triflate **1c** – 4h, and for tosylate **2** – 6h.



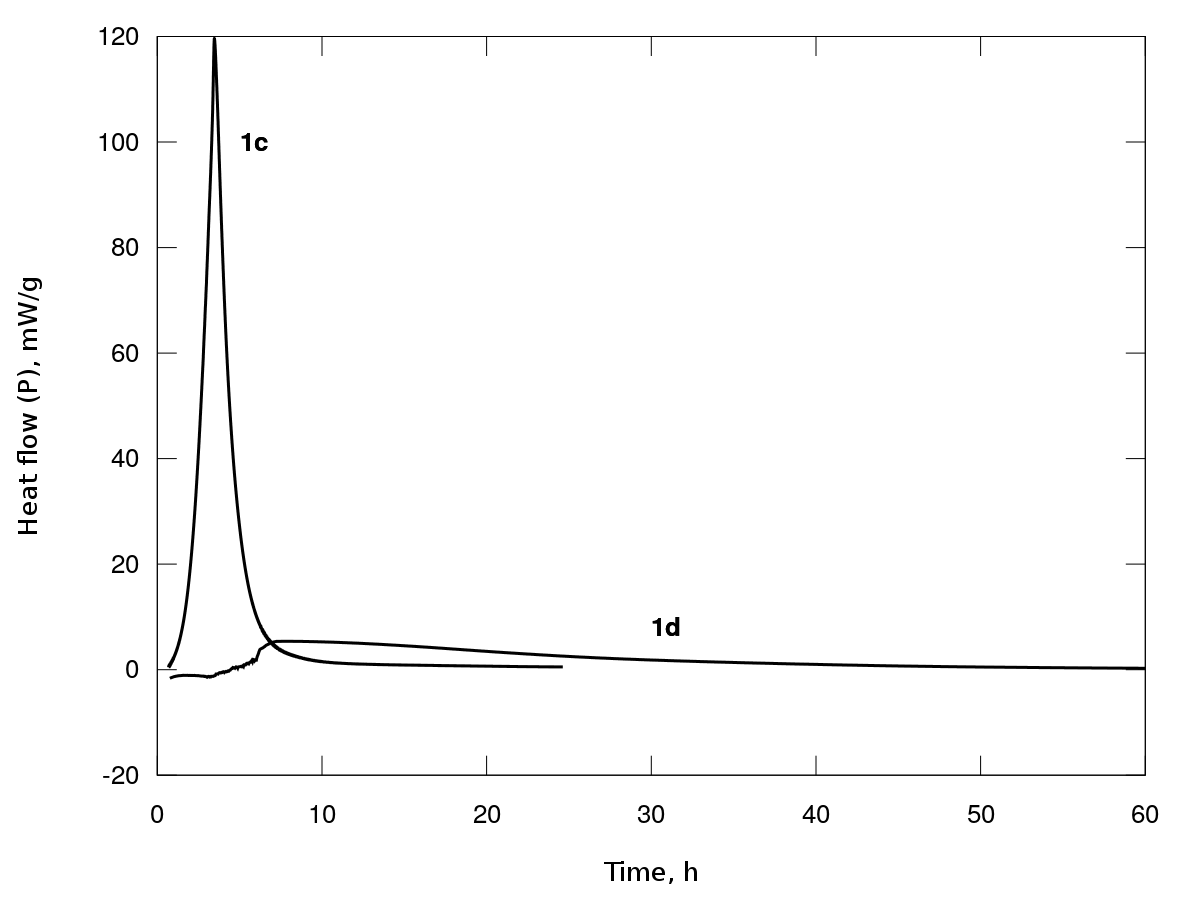
**Figure 7.** The heat flow (P) during isothermal decomposition of nitrobenzenediazonium salts: tetrafluoroborate **3**,tosylate **2**, and triflate **1c** at 85 ° C.

The experimental heat flow curves describing the isothermal decomposition of salt **1a** are more complex and, therefore, differ from the curve characteristic for the autocatalytic process. At the initial stage, a decrease in the heat flow is observed, which is associated with a higher rate of endothermic process. Then, the heat flow is increased indicating the predominance of the exothermic process. The presence of the endothermic process is consistent with the DSC/TGA data (Table 1). To describe the kinetics of this reactions, we used the model of two sequential and two parallel autocatalytic processes. The deconvolution results are given in detail in Supporting Information 1s. Table 4 shows the parameters of the heat flow kinetics calculated when isothermal decomposition reaction is approximated by a single autocatalytic process corresponding to the main exothermic stage. This stage is the most important as it determines the main characteristics of safety and stability of DSs.

**Table 4.** Kinetic parameters of the isothermal decomposition of DSs **1-3 (**P0 – initial heat flow value, *k* – rate constant calculated according to the equation 2).

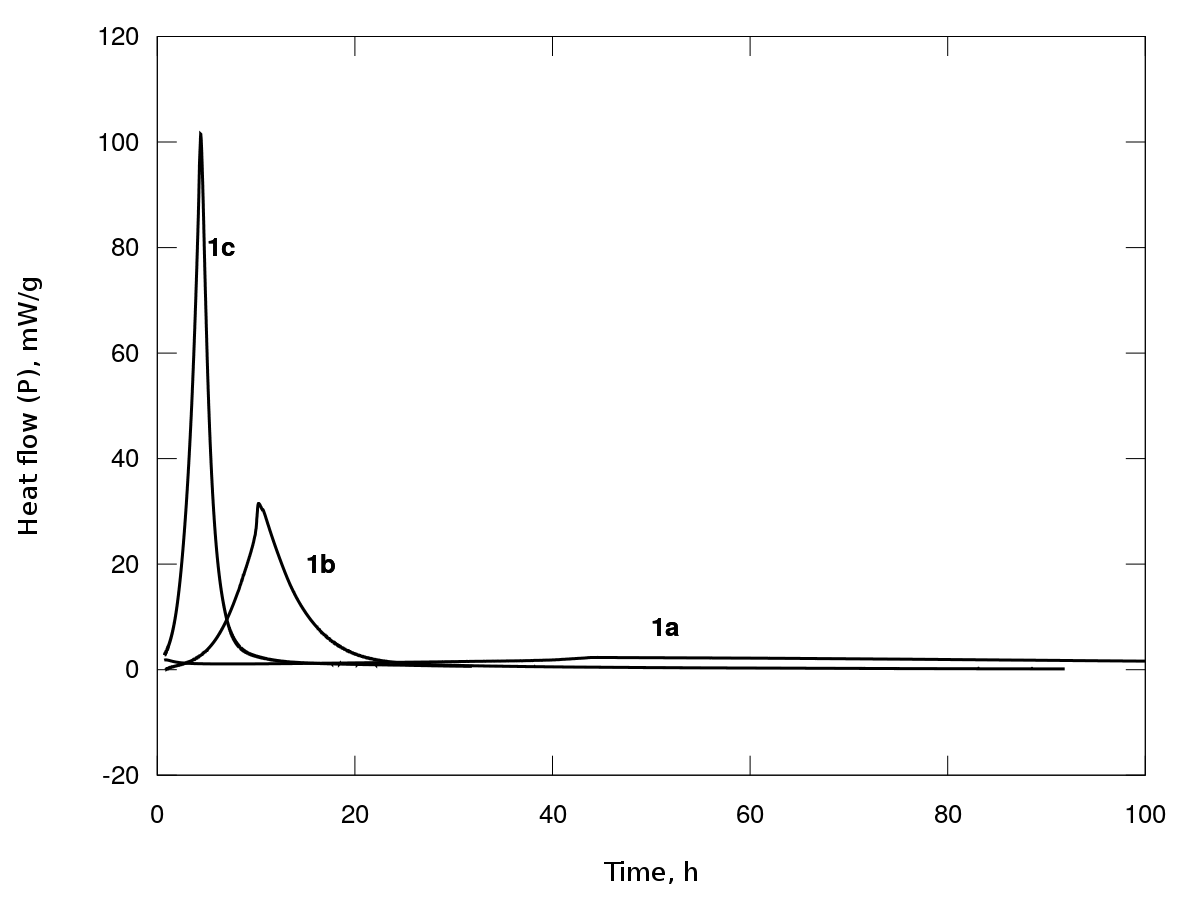
| Diazonium salt | P0, mW/g | | | k·103, s-1 | | |
| --- | --- | --- | --- | --- | --- | --- |
| 75o | 80o | 85o | 75o | 80o | 85o |
| 2-NO2С6H4N2+TfO- **1a** | 40.6 | 88.2 | 147.0 | 0.00087 | 0.00184 | 0.00254 |
| 3-NO2С6H4N2+TfO- **1b** | 0.94 | 11.0 | 44.6 | 0.03048 | 0.06545 | 0.14339 |
| 4-NO2С6H4N2+TfO- **1c** | 9.0 | 13.1 | 48.4 | 0.10562 | 0.21057 | 0.53446 |
| 4-MeOС6H4N2+TfO- **1d** | 14.78 | 32.00 | 113.8 | 0.00739 | 0.01548 | 0.04504 |
| 4-NO2С6H4N2+TsO- **2** | 19.57 | 27.5 | 41.5 | 0.08995 | 0.16775 | 0.32057 |
| 4-NO2С6H4N2+ BF4- **3** | 392.4 | 896.0 | 1804.2 | 0.07851 | 0.14520 | 0.30432 |

The half-life of 4-nitrophenyldiazonium triflate **1c** is significantly less than the half-life of 4-methoxyphenyldiazonium triflate **1d**: the values found are 4 h and 16 h, respectively. At the same time, the heat flow observed during decomposition of 4-nitrophenyldiazonium triflate **1c** is much higher than that of 4-methoxyphenyldiazonium triflate **1d** (Fig. 8): 119.5 mW/g for **1c** versus 5.37 mW/g for **1d (**Fig.8).



**Figure 8.** The heat flow (P) during isothermal decomposition of 4-nitrobenzenediazonium **1c** and 4-methoxybenzenediazonium **1d** triflates at 85 ° C.

The substitution pattern in aromatic ring has a pronounced effect on the DS stability. Among DS with triflate counterion, the longest half-life time at 85° C has *ortho-* derivative **1a** (62 h). The *meta-* derivative **1b** is less stable (11 h) and *para-* derivative **1c** is the least stable (4 h). The maximum heat flow values follow the opposite pattern and decrease in a row **1c**> **1b**> **1a**. The results discussed are presented in Tables 3 and 4 and Fig.9.



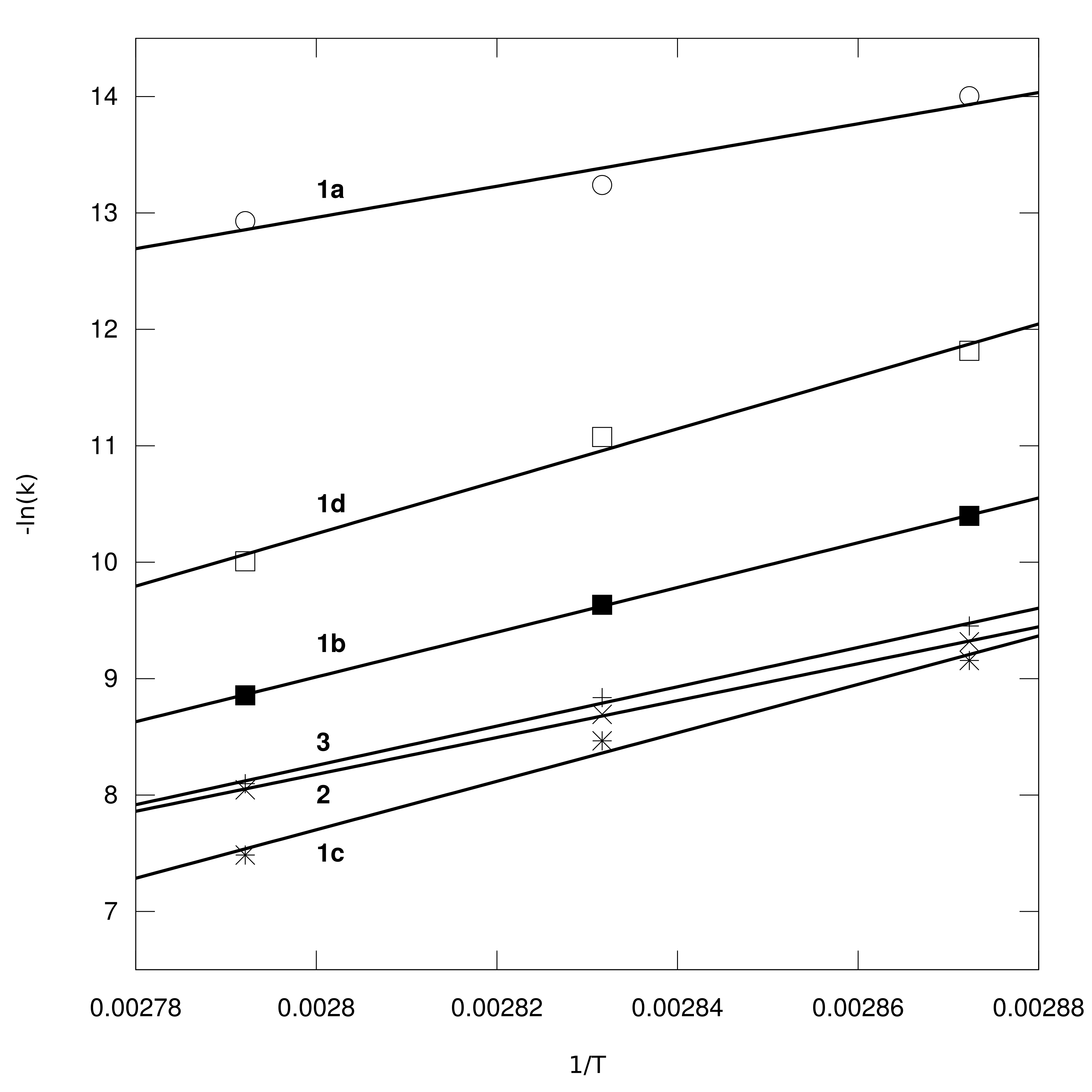
**Figure 9.** The heat flow (P) during isothermal decomposition of 2-, 3- and 4-nitrobenzenediazonium triflates **1a-c** at 85 ° C.

The analysis of the kinetic data obtained at different temperatures followed by approximation with the Arrhenius equation allowed us to find the kinetic parameters of DS decomposition reactions occurring at 25 ° C. The obtained results are presented in Table 5 and Fig. 10. The arenediazonium triflates **1b-d** have higher activation energies comparing to tosylate **2** and tetrafluoroborate **3**. However, 2-nitrobenzenediazonium triflate is out of this line as it has the lowest activation energy among the DS studied.

**Table 5.** The calculated kinetic parameters of DS **1-3** decomposition reactions occurring at 25 ° C.

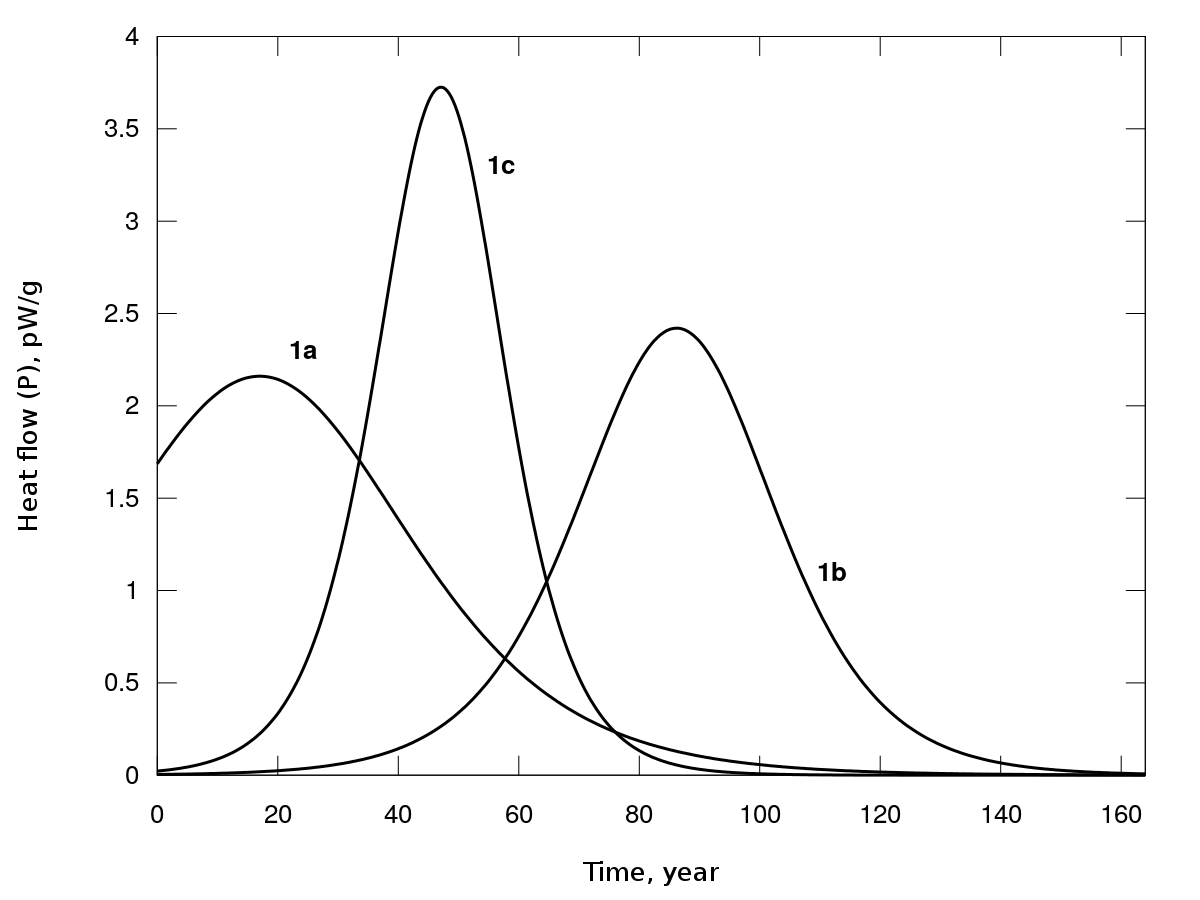
| Diazonium salt | k298 ·109, s-1 | Ea, kJ/mol |
| --- | --- | --- |
| 2-NO2С6H4N2+TfO-**1a** | 1.39 | 111.4 |
| 3-NO2С6H4N2+TfO-**1b** | 2.90 | 159.7 |
| 4-NO2С6H4N2+TfO-**1c** | 4.45 | 173.0 |
| 4-MeOС6H4N2+TfO-**1d** | 0.049 | 187.1 |
| 4-NO2С6H4N2+TsO-**2** | 48.91 | 131.7 |
| 4-NO2С6H4N2+ BF4-**3** | 18.75 | 140.3 |

k298 – rate constant at 298 оK; Ea — activation energy, kJ/mol

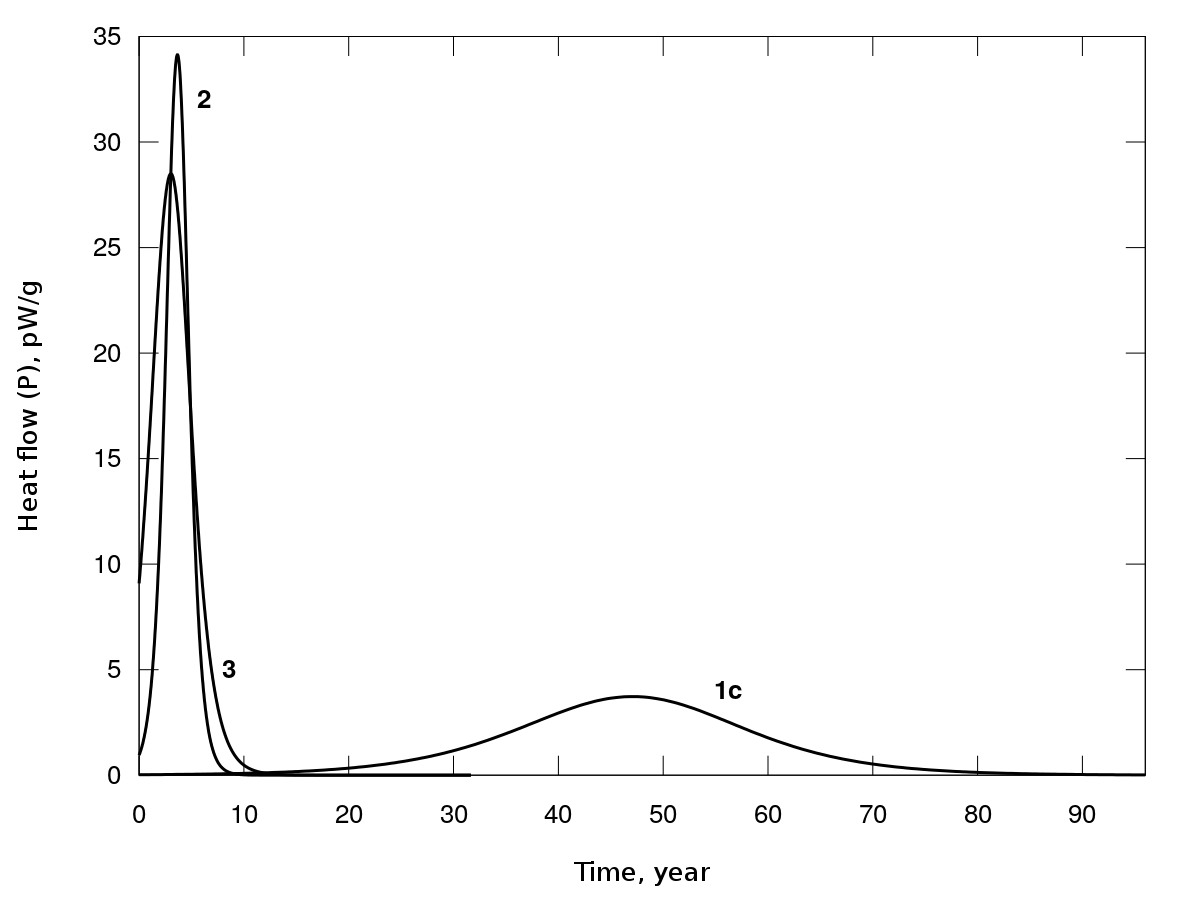


**Figure 10.** Results of the approximation of DS **1-3** decomposition reactions with the Arrhenius equation

Based on the approximation data, we have modeled the process of decomposition of the investigated DS over large time period (Fig. 11-12). According to the results, the stability of arenediazonium triflates depends on the substitution pattern. In particular, at 25 °C 3-nitrobenzenediazonium triflate has the longest half-life time of 83 years, whereas 2-nitrobenzenediazonium triflate has the shortest half-life time of 25 years (Fig. 11). The nature of counterion affects greatly the DS stability (Fig. 12). Thus, 4-nitrobenzenediazonium tosylate **2** and tetrafluoroborate **3** have close half-life times of 4.5 years. Whereas 4-nitrobenzenediazonium triflate **1c** is much more stable with a half-life time of 46 years and a significantly lower maximum heat flow. Note that the effect of counterion on DS stability becomes noticeable only at low temperatures (25 °C), while at elevated temperatures (during DSC/TGA or isothermal flow calorimetry at 75-85 °C) these differences disappear. This fact, as well as the values of activation energies, suggest that the stability of DS under normal storage conditions is largely determined by the strength of the crystal lattice. At higher temperatures, after the destruction of the crystal lattice, the speed and energy of the process are likely to be influenced by both the nature of the Ar-N2+ diazonium cation and the presence and nature of nucleophiles in the immediate environment.



**Figure 11.** The heat flow (P) over the time dependency built upon modelling of the kinetic curves of the decomposition of 2-, 3- and 4-nitrobenzenediazonium triflates **1a-c** at 25 оС.



**Figure 12.** The heat flow (P) over the time dependency built upon modelling of the kinetic curves of the decomposition of 4-nitrobenzenediazonium tosulate **2**, triflate **1c** and tetrafluoroborate **3** at 25 оС.

Thus, considering all the data, we can draw several conclusions. First of all, DSC/TGA alone cannot serve as a reliable criterion for assessing the thermal stability and safety of DSs, since in this case decomposition occurs at higher temperatures and is accompanied by intense evaporation of low-molecular-weight reaction products. In addition, during DSC/TGA analysis DSs decompose at different temperatures, which makes the analysis and comparison of the data complicated. Therefore, the study of DS thermal stability should be complemented by a more detailed investigation of the decomposition kinetics by isothermal flow calorimetry. The values of maximum heat flow, half-lives, and activation energies should be determined and approximation of the kinetics to normal conditions should be done.

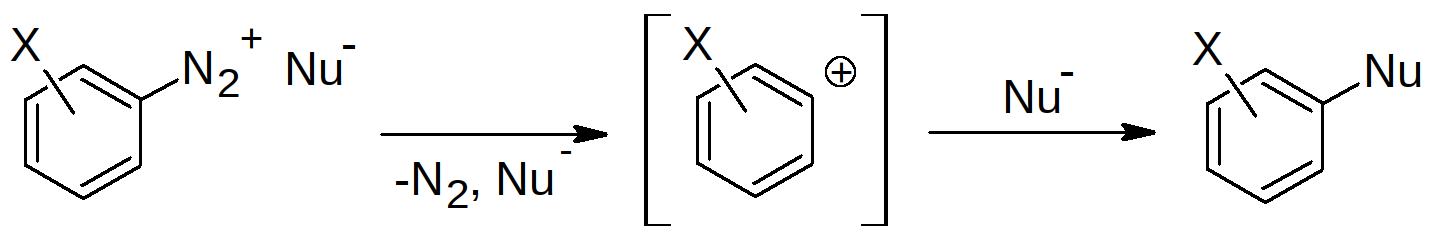
Secondly, the nature of counterion has a pronounced effect on DS stability. The approximation of experimental data with Arrhenius equation showed that arenediazonium triflates are the most stable during the storage, for example, 3-nitrobenzenediazonium triflate has a shelf-life of 83 years. Presumably, changing the counterion effects the probability of the occurrence of various mechanisms during the DS decomposition, which are discussed further and presented in Schemes 1 and 2.

Finally, given the results of isothermal flow calorimetry, the DSs studied decompose with thermal effects close to the threshold value of 800 J/g for safe transportation, according to UNESCO.6 Only in the case of 2-nitrobenzenediazonium triflate, the released energy of 1330 J/g exceeds the permissible value.

**GC-MS and LC-MS study of decomposition products**

While it has long been established that the main products of the thermal decomposition of arenediazonium tetrafluoroborates are the corresponding aryl fluorides (the Balz-Schiemann reaction), the products of the thermolysis of arenediazonium triflates and tosylates remain unknown. We elucidated the structure of the compounds that appear after 14 days of decomposition of DSs **1-3** at 85 оС using GC-MS and LC-MS. To conduct the analysis, the unreacted DS was converted into corresponding aryl iodide by reaction with KI.

According to the GC-MS data (Supporting Information 2s, Fig. 2.s1-11), the main products of the decomposition of DSs **1b**, **1c**, **1d** are the corresponding esters of nitro-phenyl trifluoromethanesulfonates ArOTf. The decomposition of arenediazoniumtosylate **2** resulted in the formation of nitrobenzene and 1-iodo-4-nitrobenzene (GC-MS). LC-MS ESI in negative ionization mode also showed the presence of 4-nitrophenyl 4-methylbenzenesulfonate ester (m/z=292.1) among the decomposition products. In case of arenediazonium tetrafluoroborate **3**, the main decomposition product was the expected 1-fluoro-4-nitrobenzene. It should be noted that during the decomposition of all the DSs studied, significant amounts of resinous products that could not be elucidated by GC-MS were formed. Considering the GC-MS and LC-MS results, the following mechanism can be assumed for thermolysis of DSs **1b**-**d**, **2**, **3** (Scheme 1).



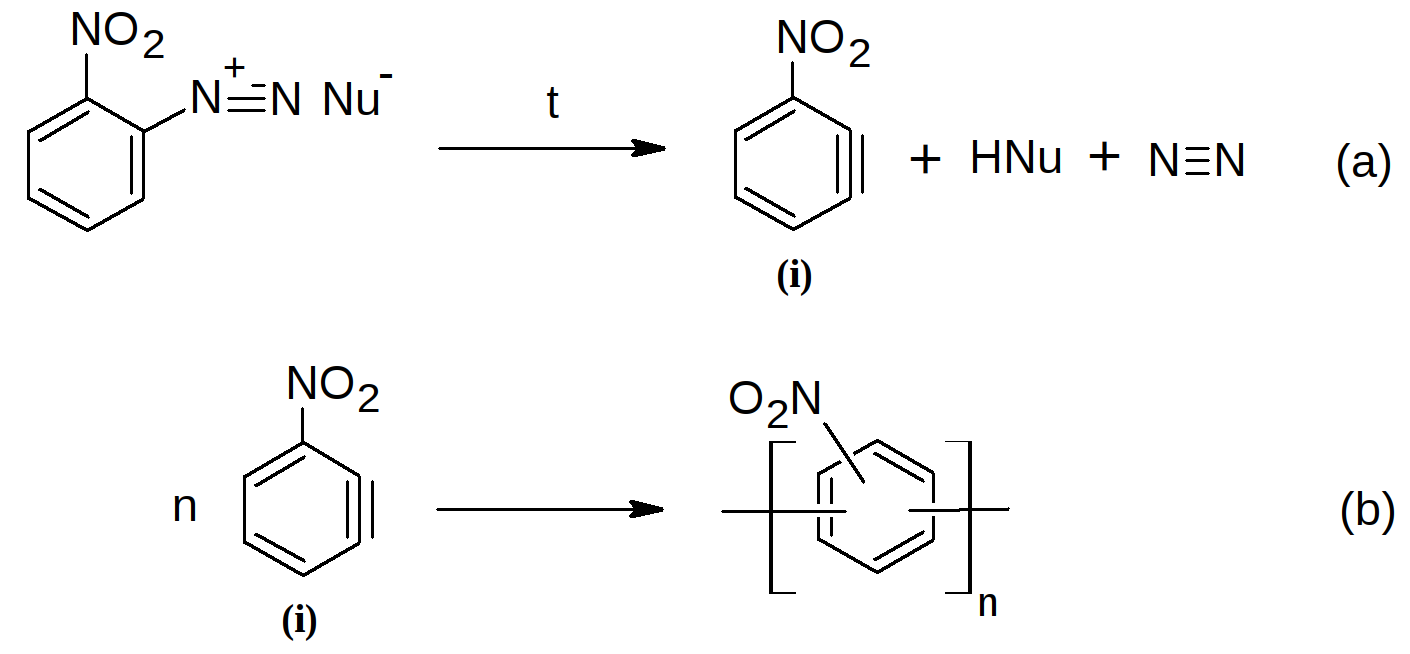
**Scheme 1.** The possible mechanism of thermal decomposition of DSs **1b**-**d**, **2**, **3** (Nu = F, TsO, TfO)

A completely different process takes place when 2*-*nitroarenediazonium triflate **1a** decomposes. In this case, the products of the diazonium group substitution by the triflate anion were not detected. The compounds formed during the DS **1a** decomposition were polymers. We were able to elucidate their structure by LC-MS (Table 6, MS spectra presented in Supporting Information 3s). The polymers found were suggested to be built upon the following repeating units: NO2C6H3 (M/Z=121.0) and [NO2C6H4SCF3] (M/Z=223).

**Table 6.** The main peaks present on the LC-MS chromatograms of the decomposition products of 2*-*nitroarenediazonium triflate **1a**

| Ionization Mode | M/Z | Compound |
| --- | --- | --- |
| Positive ESI | 74.1; 297.1; 520.2; 743.2 | Polymer chain P1 with the mass of repeating unit of 223, possibly, [NO2C6H4SCF3] |
| 432.1; 553.1; 674.1; 795.1; 916.1; 1037.1; 1158.1 | Polymer chain P2 with the mass of repeating unit of 121, [C6H3NO2] |
| 525.1; 646.1; 767.1; 888.1; 1009.1; 1130.1 | Polymer chain P3 with the mass of repeating unit of 121, [C6H3NO2]  *The difference between P2 and P3 is 28, which corresponds to the nitrogen elimination* |
| 588.2; 710.1; 831.2; 952.2; 275.1; 785.8; 915.2; 1087.1 | Polymer chain P4 with the mass of repeating unit of 121, [C6H3NO2] |
| Positive APCI | 394.0; 515.0; 635.9; 756.9; 877.8; 998.8; 1119.7 | Polymer chain P5 with the mass of repeating unit of 121, [C6H3NO2] |
| 597.3; 875.4; 877.4; 879.4 |  |
| Negative ESI | 380.1; 501.1; 622.1; 743.1; 864.1 | Polymer chain P6 with the mass of repeating unit of 121, [C6H3NO2] |
| 528.0; 649.1; 770.1; 891.1 | Polymer chain P7 with the mass of repeating unit of 121, [C6H3NO2] |
| 149.0 | TfO- |
| 276.8; 320.9; 436.7; 563.6; 936.2 |  |

Apparently, the appearance of the above-mentioned resinous compounds is associated with the polymerization processes as proven by the polymeric nature of products formed as a result of DS **1a** decomposition. We can infer that 1-nitrocyclohexa-1,3-dien-5-yne [C6H3NO2] (M/Z=121.0) is a monomer that is derived from DS according to the route (a) presented in Scheme 2.



**Scheme 2.** Possible route of 1-nitrocyclohexa-1,3-dien-5-yne [C6H3NO2] (M/Z=121.0) formation (a) and further polymerization (b)

The mechanism of further chain growth involving C6H3NO2 as monomer is ambiguous and requires special investigation lying beyond the scope of this work. However, based on the available data, we can assume the structure of the polymer formed as a result of the DS **1a** decomposition process (b) presented in Scheme 2. Previously, the appearance of polymer products during the thermal decomposition of *ortho-*carboxybenzenediazonium chloride was also explained by the intermediate formation of didehydrobenzene and naphthalene.2h,2i

Importantly, the unique route found for DS **1a** decomposition including the formation of neutral C6H3NO2 is consistent with the high instability of the intermediate involved in an alternative pathway (Scheme 1). Indeed, the 2-nitrobenzene-1-ylium cation that would be formed in this case is much less stable and energetically unfavorable comparing to aryl cations derived from the other DSs due to fact that electron withdrawing group NO2 is next to carbocationic center.

**Quantum chemical calculations**

Given the identified products of DS **1a-c**, **2**, **3** thermal decomposition, we for the first time predicted the thermodynamics of the occurring processes using DFT calculations at RB3LYP/aug-cc-pVDZ level of theory. We have optimized geometry of DS **1a-c**, **2**, **3** and diazonium group substitution products. The nature of located stationary points was confirmed by the absence of imaginary frequencies in IR spectrum. Cartesian coordinates of all compounds, and calculated thermodynamic parameters are given in Supporting Information 4s. We have explored esters and substituted phenols as the major products of triflates **1a-c** and tosylate **2** decomposition and 4-fluoronitrobenzene as the major product of tetrafluoroborate **3** decomposition. The predicted and experimental thermodynamic parameters of the reactions are presented in Tables 7 and 8. Considering DSC/TGA results that showed that thermolysis of tetrafluoroborate **3** initially causes BF3 detachment with the formation of 4-NO2C6H4N2+F- **3a** (Scheme 1), we calculated the thermodynamics of this reaction along with other possible routes (Table 6, entries 6a, b).

**Table 7.** Predicted thermodynamic parameters of DS **1a‑c**, **2**, **3** decomposition reactions according to quantum chemical calculations at RB3LYP/aug-cc-pVDZ level of theory.

| Entry | Reaction | G298, kJ/mol | H298, kJ/mol | S\*298.15, kJ/mol |
| --- | --- | --- | --- | --- |
| 1 | 2-NO2C6H4N2+-OTf→ 2-NO2C6H4OTf + N2 | -268.6 | -230.1 | 38.5 |
| 2 | 3-NO2C6H4N2+-OTf→ 3-NO2C6H4OTf + N2 | -282.4 | -242.8 | 39.6 |
| 3 | 4-NO2C6H4N2+-OTf→ 4-NO2C6H4OTf + N2 | -287.6 | -248.0 | 39.6 |
| 4 | 4-MeOC6H4N2+-OTf→ 4-MeOC6H4OTf + N2 | -238.1 | -200.5 | 37.7 |
| 5 | 4-NO2C6H4N2+-OTs → 4-NO2C6H4OTs + N2 | -328.4 | -283.9 | 44.6 |
| 6 | 4-NO2C6H4N2+-BF4→ 4-NO2C6H4F + BF3 + N2 | -276.2 | -188.4 | 87.9 |
| 6a | 4-NO2C6H4N2+-BF4→ 4-NO2C6H4N2F + BF3 | 76.2 | 124.6 | 48.5 |
| 6b | 4-NO2C6H4N2F-→ 4-NO2C6H4F + N2 | -352.3 | -312.9 | 39.4 |

**Table 8.** Experimental and predicted enthalpies of decomposition of diazonium salts (RB3LYP/aug-cc-pVDZ).

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| Reaction | Predicted enthalpy | Experimental enthalpy (flow calorimetry) | | | | DSC/TGA |
| ΔH298, (ΔH353) kJ/mol | ΔH348, kJ/mol | ΔH353, kJ/mol | ΔH358, kJ/mol | ΔH, kJ/mol | |
| 2‑NO2C6H4N2+ Tf- → 2‑NO2C6H4OTf + N2 | -230 (-230.4) | -414 | -386 | -396 | -203.4 | |
| 3‑NO2C6H4N2+ TfO- → 3‑NO2‑C6H4OTf + N2 | -243 (-243.2) | -228 | -230 | -225 | -238.5 | |
| 4‑NO2C6H4N2+ TfO- → 4‑NO2‑C6H4OTf + N2 | -248 (-248.4) | -200 | -235 | -250 | -65.8 | |
| 4-MeOC6H4N2+ TfO- → 4‑MeOC6H4OTf + N2 | -201 (-200.8) | -183 | -183 | -106 | -64.5 | |
| 4-NO2C6H4N2+ TfO- → 4‑NO2C6H4OTs + N2 | -284 (-284.0) | -253 | -232 | -231 | -117.3 | |
| 4-NO2C6H4N2+ BF4- → 4‑NO2C6H4F + BF3 + N2 | -188 (-189.5) | -173 | -156 | -147 | -54.3 | |

Overall, for all reactions of DS **1b-d**, **2** and **3** decomposition the calculated enthalpy values are consistent with the experimental ones obtained by isothermal flow calorimetry. Therefore, the suggested reactions make the main contribution to the energy of DS exothermic decomposition (Table 6, 7). The obtained results prove that DFT calculations at RB3LYP/aug-cc-pVDZ level of theory is a convenient and fairly precise method for theoretical estimation of the thermal effects of DS decomposition. A single case where we had significant deviation between the data of isothermal flow calorimetry and calculation results is 2-nitroarenediazonium triflate **1a** (Table 7). However, this is consistent with GC-MS and LC-MS data demonstrating that decomposition of DS **1a** proceeds via different route (Scheme 2) and does not result in 2-NO2C6H4OTf.

In some cases, the enthalpy values of DS **1-3** exothermic decomposition measured by DSC/TGA were found to be significantly less than both predicted ones and values obtained by isothermal flow calorimetry (Table 8). It can be explained by the fact that decomposition of DSs during DSC/TGA analysis occurs at temperatures much higher than 85 ° C (Table 2), resulting in an evaporation process, accompanied by a significant weight loss, which reduces the thermal effect of the reaction.

Importantly, the DFT calculations were carried out assuming reagents and products are isolated molecules, therefore, the electric field of crystal was neglected. However, the very fact that the predicted energies are in good agreement with the experimental values measured by flow calorimetry for the decomposition reactions of five diazonium salts with different counterions and ring substituents **1b**-**d**, **2** and **3** (Table 8) indicates that the major contribution to the reaction thermodynamics is made by the chemical transformations, whereas the effects of changing the crystal lattices are minor. Possibly, it is due to the fact that both reagents and products remain solid (except N2), thus, there is a compensation for the thermal effects of the transformation of the crystal lattices of the starting material and products. From an application prospective, it is also important that the proposed quantum-chemical approach to assessing the thermodynamics of decomposition reactions of DSs is relatively easy to implement. While calculations that take into account unknown transformations of crystal lattice are much more laborious, sophisticated and cannot yet be widely used to predict the thermal effects of chemical reactions, despite the known progress in calculating the crystalline state of benzenediazonium chloride and tetrafluoroborate 2c,d.

**CONCLUSIONS**

In conclusion, for the first time we have determined the thermodynamics and kinetics of thermal decomposition of a series of aromatic diazonium salts ArN2+ X– with various counterions X = TfO, TsO, BF4 by isothermal flow calorimetry and provided the quantitative assessment of the storage stability of solid DSs under normal conditions. Additionally, we have established how the aromatic substitution pattern in diazonium cation and the nature of counterion effect the processes occurring during the thermolysis of DSs studied.

We demonstrated that thermodynamic parameters of DS thermal decomposition reactions calculated by DFT at RB3LYP/aug-cc-pVDZ are consisted with experimental data obtained by isothermal flow calorimetry for all 3- and 4-substituted DSs investigated (**1b-d, 2, 3**). For these DSs the main decomposition route is elimination of nitrogen with the formation of benzene-1-ylium ions that subsequently react with the corresponding anions. A completely different process occurs during the thermolysis of 2-nitrobenzene diazonium triflate **1a**. In this case, the polymeric products are formed, probably through the primary generation of 1-nitrocyclohexa-1,3-dien-5-yne.

**ASSOCIATED CONTENT**

Results of modeling and deconvolution of heat flow experimental kinetic curves acquired in isothermal conditions as well as results of quantum chemical calculations can be found in Supporting Information.

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The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

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**ABBREVIATIONS**

DSs, diazonium salts; ADTs, arenediazonium tosylates; ADTfs, arenediazonium trifluoromethane sulfonates; DSC/TGA, Differential scanning calorimetry and thermal gravimetric analysis; UNECE, United Nations Economic Commission for Europe; GC-MS, Gas chromatography–mass spectrometry; LC-MS, Liquid chromatography–mass spectrometry; ESI, electrospray ionization source; APCI, atmospheric pressure chemical ionization source; DFT, Density functional theory.

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