Theoretical study of the formation mechanism of molecules CaF for fluoride determination using electrothermal molecular absorption spectrometry

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In order to select the temperature-time heating program, the calibration method, the prediction of matrix interference and to increase the sensitivity and accuracy of the halogens determination using the electrothermal molecular absorption spectrometry it is essential to know the formation mechanism of the measured molecules. Thermochemical processes of gaseous molecules CaF formation, which are used in the electrothermal molecular absorption determination of fluorine, were studied using the method of the thermodynamic simulation. All calculations were performed using the HSC 6.1 software package with its own database of thermodynamic data. For calculations, the non-equilibrium thermodynamic system, which was realized in the graphite furnace, was divided into the thermodynamic subsystems that correspond to the main stages of time-temperature program: drying of the sample, pyrolysis, vaporization and formation of gaseous diatomic molecules. The composition of the condensed sample residues after drying and pyrolysis stages, the composition of the gas phase in the analytical zone of the graphite furnace during vaporization and the formation of molecules stages were determined by the calculations. This study theoretically describes thermochemical processes of the formation of gaseous CaF molecules using thermodynamic simulation for electrothermal molecular absorption determination of fluorine. The calculations make it possible to establish that gaseous molecules CaF\(_g\) are formed during the course of the thermal decomposition of the condensed fluoride (CaF\(_2\)). The correct execution of thermodynamic simulation was confirmed by the comparison of the theoretical temperature curves of pyrolysis and the formation of molecules CaF\(_g\) with the experimental curves. The developed procedure of the thermodynamic simulation can be applied to study the formation of other dimeric molecules used for molecular absorption analysis with electrothermal atomization.

Keywords: high-resolution continuum source molecular absorption spectrometry, thermodynamic simulation, fluorine, calcium mono-fluoride, temperature of drying, pyrolysis and vaporization.
Introduction

The occurrence of high resolution continuum source atomic absorption spectrophotometers (HR CS AAS) made possible the determination of halogens, sulfur, and phosphorus by the absorption spectra of their diatomic molecules [1]. Simultaneously, there was a question about the formation mechanism of these molecules in commonly used atomizers [2]. Knowledge of this mechanism and the possibility of its control can significantly affect the achievable detection limits and range of determined elements concentrations, calibration method, matrix interference, accuracy, etc.

The experimental method for studying the formation mechanism of SrF\text{g} molecules in the graphite furnace using mixed and separate injection of molecule forming element and the analyte in graphite furnace was proposed in [2]. However, at the present time this experimental method has limited possibilities for investigations. Therefore, more flexible and informative theoretical method and study procedure of SrF molecule formation using the method of equilibrium thermodynamic simulation was proposed in [3]. Control of the developed simulation procedure was carried out based on the experimental results [2, 4] and own experimental investigation. The aim of this work was to use the developed simulation procedure to study the formation mechanism of diatomic CaF\text{g} molecules for electrothermal atomic absorption determination of fluorine.

Calculation procedure of thermodynamic simulation

The method of the equilibrium thermodynamic simulation consists of the behavior analysis of high-temperature heterogeneous multicomponent system based on the calculation of its full equilibrium chemical composition [5, 6]. The calculation is performed at the extremum of the thermodynamic potential (entropy maximization or free-energy (Gibbs energy) minimization) taking into account most of the possible components in the equilibrium. The ability to apply the equilibrium thermodynamic simulation to the study of thermochemical processes of elements atomization in the graphite furnace, which is a non-equilibrium opened thermodynamic system, has been well founded and confirmed in [4, 7]. For the calculations, it was suggested to use the approach based on the division of a non-equilibrium system into the successive quasi-equilibrium subsystems corresponding to the sequence of thermal transformations of the initial sample: drying, pyrolysis subsystems corresponding to the sequence of thermal transformations of the initial sample: drying, pyrolysis.

The resulting calculated condensed equilibrium chemical composition of the previous subsystem is the starting compound for the subsequent subsystem. This approach takes into account the interaction between all of the components of the system (solvent, sample matrix, analyte, chemical agent - modifier, surface material of the atomizer, protective or active gas with its impurity composition), distillation of volatile components, allows for the determination of the complete chemical composition and the optimal temperature for each stage of atomization, etc.

Procedure of the thermochemical processes simulation for molecular absorption analysis in the graphite furnace and corresponding basic stages of the time-temperature program of the electrothermal atomizer includes the following sequence [3].

The stage of sample drying. Initial chemical composition of the thermodynamic subsystem corresponds to experimental or suggested data: diluent and salt composition of the pipetted solution. The amount of argon in the subsystem is set based on the molar ratio of \textit{Ar}: \textit{solvent} = 1: 1. The interaction of the sample solution with carbon atomizer is excluded. Calculations take into account the gaseous and condensed phase,
considering the individual condensed and water compounds in the latter. The calculation is performed in the 30-150 °C temperature range with the step of 10 °C, taking into account losses of the gaseous products in an open system of the atomizer at each step.

**The stage of pyrolysis.** Initial chemical composition of the thermodynamic subsystem for this stage is determined by the calculated composition of the condensed product of the previous drying stage under experimental or selected temperature of this step. In calculations a “thick” layer of the sample is divided into two zones: in contact and not in contact with the carbon atomizer. All sample components in these zones are uniformly distributed and may form, by reaction between each other, condensed individual substances and condensed dilute solutions (usually - oxides) based on the sample matrix and chemical agent - modifier. The molar ratio of carbon : sample matrix is assumed to be equal to 10, $\text{Ar : condensed sample} = 10^2-10^4$. The calculation is carried out in the range of temperature of the drying stage up to 2000 °C with the steps of 50 °C, taking into account at each step the loss of gaseous products in an open atomizer system.

**The stage of vaporization and formation molecules.** This step, which is realized experimentally at the maximum heating rate of the graphite furnace and the gas stop mode (“gas-stop”) for calculations, is subdivided into two successive stages.

The first stage – vaporization. Initial chemical composition of the condensed products on the surface of the atomizer for this phase is determined by summing up the calculated compositions of condensed products of the two zones of the sample during the pyrolysis stage at experimental or theoretical temperature of this step (corresponding to the loss of analyte in the gas phase of the graphite furnace). Further calculations are carried out only for a “thin” layer of the sample undivided into zones and reacting with the carbon of the atomizer. This is because most of the volatile components of the sample during the pyrolysis stage have been removed from the space of the graphite furnace by argon flow. The amount of argon for the vaporization stage is set to be equal to the device placed in graphite furnace at this temperature, by its impurities ($\text{O}_2$, $\text{N}_2$, $\text{H}_2\text{O}$) and partial pressure of carbon vapor in the gas phase of the furnace. This calculation is performed completely analogous to the calculation of the atomization element stage [7] and carried out starting from the temperature of the pyrolysis step up to 2700 °C with step of 50 °C.

When applying the components of the sample to different portions of the evaporator surface for the drying and pyrolysis stage, as well as for the vaporization step, calculations are performed individually for each of the initial chemical composition. When calculating the chemical composition of the gas phase of the graphite furnace only the possibility of partial mixing and interaction of the vaporized components is taken into account.

**Results and discussion**

The procedure of thermodynamic simulation, which we developed in [3], was used for studying of the formation mechanism of CaF molecules in the graphite furnace. The applicability verification of this procedure was carried out using experimental results from [8-10], where the possibility of fluorine determination from the molecular absorption CaF using electrothermal vaporization was studied.

For the thermodynamic simulation the initial compositions of thermodynamic systems that were used are completely similar to the experimental data of [8] where fluorine was introduced as NaF and Ca(NO$_3$)$_2$ was used as chemical agent (10 ng of fluorine, 40 µg of calcium, 20 µL H$_2$O), [9] (3 ng of fluorine, 15 µg of calcium, 20 µL H$_2$O) and [10] (16 ng of fluorine, 315 µg of calcium, 20 µL H$_2$O). Calculations were carried out in full compliance with the procedure mentioned in paragraph above. Considered in the calculations of the thermodynamic equilibrium individual substances are given Table. All calculations were performed using the HSC 6.1 software package (minimization of Gibbs energy in thermodynamic system) [11] with its own database.

**The stage of sample drying**

The calculation results of the equilibrium composition of the system Ca-F-Na-O-H-Ar at the drying stage using the experimental data in [8] are shown in Fig. 1. It is observed that the exchange reactions during the formation of CaF$_2$ are already possible at the heating of the solutions on the graphite platform during the stage of sample drying. With the low-temper-
nature decomposition of crystalline calcium nitrate (Fig. 1, B) due to the hydrolysis, the formation of Ca(OH)$_2$ is thermodynamically possible.

Calculated functions according to the experimental data of [9, 10] showed a similar composition of the final products for the drying stage at temperatures above 100 °C.

**The stage of pyrolysis**

Input amounts of condensed substances of this subsystem are defined by the results of calculations at the drying step at temperatures of 120 °C [8], 140 °C [9] and 110 °C [10], which correspond to the experimental temperatures of the drying stage. Amount of argon for each zone of “thick” sample was set according to the molar ratio of $Ar: matrix = 2.3 \times 10^3$. The results of calculation of the equilibrium composition for the subsystem [8] are shown in Fig. 2.

According to calculations (Fig. 2, B) in the zone of sample surface and already at the initial stage of pyrolysis, the decomposition of Ca(OH)$_2$ to a stable over a wide temperature range CaO will occur. Fluoride is present in the subsystem after the drying stage mostly in CaF$_2$ and partly in NaF. During the pyrolysis stage it can be redistributed between these individual substances (Fig. 2, A). However, such small amounts of calcium fluoride and sodium fluoride remain in dilute condensed solution based on CaO. Small losses of fluoride in the gas phase in the form of NaF at the temperature range 800-1000 °C are possible. Significant losses of fluorine

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**Table**

<table>
<thead>
<tr>
<th>Gas (upper index – g)</th>
<th>Condensed (upper index – c)</th>
<th>Aqueous (upper index – a)</th>
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<tr>
<td>F, F$_2$, FOO, FNO, FNO$_2$, FNO$_3$, F$_2$NO, CF, CF$_2$, CF$_3$, F$_2$C, C$_F$, F$_2$C, F$_3$C, F$_4$C, F$_5$C, F$_6$C</td>
<td>Ca(NO)$_3$,$_2$H$_2$O, Ca(NO)$_3$,$_3$H$_2$O, Ca(NO)$_3$,$_4$H$_2$O, CaO, Ca$_2$O, Ca(OH)$_2$</td>
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<td>HNO$_3$, H$_2$N$_2$, H$_3$N, H$_4$N, N$_2$O, N$_2$O$_2$, N$_2$O$_3$, N$_2$O$_4$, N$_2$O$_5$</td>
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<tr>
<td>O, O$_2$, O$_3$</td>
<td>-</td>
<td>O, O$_2$, O$_3$</td>
</tr>
</tbody>
</table>

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Fig. 1. Calculated functions of logarithms for the amounts of individual compounds containing fluorine MeF (A), calcium CaX (B) and sodium NaX (C) from the temperature of the drying sample $T$ using the experimental data of [8].
should be observed at temperatures above 1300 °C in the form of CaF$_3$ and CaF$_2$.

In the zone of sample contact with the surface of the atomizer (Fig. 2, D) already at the initial temperatures of the pyrolysis stage the interaction of calcium compounds with the carbon surface atomizer to form CaCO$_3$, which decomposes to CaO at 500-550 °C, is thermodynamically possible. This is confirmed by experimental data [12]. Fluoride is present in the subsystem at the pyrolysis stage predominantly as CaF$_2$ and partly as NaF, which remains in the diluted solution based on CaO. Significant losses of fluoride in the gas phase of the graphite furnace should occur above 1200 °C in the form of CaF$_2$.

Calculations based on experimental data [9, 10] have shown a similar chemical composition and possible thermochemical processes for the zone of the sample surface and for the zone of the contact sample with the carbon atomizer.

Normalized calculated curves of the pyrolysis were derived using Fig. 2 calculated chemical compositions curves. These normalized curves correspond to the grade of fluorine loss in the gaseous form with an increase of the atomizer temperature (Fig. 3), and are close enough to confirm the experimental temperatures of the pyrolysis stage (700 °C [8], 900 °C [9] and 725 °C [10]). There is some difference in the course of the experimental and theoretical curves of the pyrolysis because the experimental curves of these three authors [8-10], obtained in similar operational conditions, have significant differences among each other. Nonetheless, all experimental and theoretical data coincide with the temperature of the pyrolysis stage.

**The stage of vaporization and formation molecules**

In the case of molecules CaF$_2$ formation, input amounts of condensed substances of this subsystem for the vaporization stage were set based on the results of calculations of the pyrolysis stage (Fig. 3) at a temperature of 827 °C. The calculation results of the subsystems’ chemical composition based on the experimental data of different authors for a “thin” layer of the sample interacting with carbon atomizer (tenfold molar excess) are shown at Fig. 4. Based on this data, the composition of the gas phase of the furnace, defined with the transition

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Fig. 2. Calculated functions of logarithms of amounts of individual substances containing fluorine MeF (A, C) and calcium CaX (B, D), from the temperature of pyrolysis stage T for the zone of sample surface (A, B) and the zone of contact sample with the carbon atomizer (C, D). Calculations were carried out according to the experimental data [8].

Fig. 3 Comparison of normalized experimental and theoretical curves of the pyrolysis stage: A – experimental data [9] (2) and [8] (3), theoretical calculation based on these data (1), B – experimental data [10] (2) and the theoretical calculation based on these data (1); ○ – calculated points, —— – approximation.
of sample components during vaporization ($C_g$, $CO_g$, $Cag$, $CaF_g$, $CaF_2_g$, $Na_g$, $NaF_g$), impurity composition of argon and carbon vapors at appropriate temperature was determined. On this basis, we calculated curves of efficiency of molecules $CaF$ formation and compared them with the experimental data (Fig. 5). There is a close agreement between the experimental and calculated curves. High temperature reduction of values $\eta(CaF)$ is due to a loss of vapor from the analytical zone of the graphite furnace as a result of diffusion and convection, which is not included in the thermodynamic calculations.

Fig. 4 shows that formation of molecules $CaF$, used for the molecular absorption determination of fluorine, occurs by the thermal decomposition of $CaF_2$ on the surface of the graphite heater.

Using the model of a “thin” layer that interacts with the carbon atomizer, a theoretical calculation of the calibration dependence on experimental data [8] with the introduction of various amounts of fluorine as NaF and 40 mg of calcium as calcium nitrate (Fig. 6) was carried out. There is a very good agreement between the experimental and theoretical normalized calibration curves.

**Conclusions**

By method of thermodynamic simulation the mechanism of gaseous molecules formation using a previously developed algorithm for $SrF$ was defined for $CaF$. It was shown that the formation of gaseous molecules $CaF$ is due to the decomposition of condensed $CaF_2$ during the vaporization. The mechanisms of formation $SrF$ and $CaF$ in the graphite furnace for electrothermal molecular absorption determination fluorine are similar.

The proposed algorithm can be used to study the formation mechanisms of diatomic molecules of halogen, phosphorus and sulfur for the purpose of optimizing the determination of these elements by electrothermal molecular absorption spectrometry.

**Acknowledgements**

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**References**

4. Pupyshev A.A. [Acting mechanism of inorganic chemicals modifiers in electrothermal atomic absorption spectrometry].
Литература

3. Зайцева П.В., Пупышев А.А., Курмачев Ю.А. О механизме образования молекул SrF при электротермическом определении фтора // Аналитика и контроль. 2014. Т. 18. № 3. С. 287-301.