Simultaneous determination of 11 elements in fly ash by oaTOF-ICP-MS after

closed-vessel microwave-assisted extraction with ammonium fluoride

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Abstract

A method for simultaneous multi-element analysis of fly ash samples by inductively coupled

plasma orthogonal acceleration time-of-flight mass spectrometry (oaTOF-ICP-MS) after closed-

vessel microwave extraction with ammonium fluoride was introduced here. Corrosive and and/or

toxic acids like HF, HCl or HClO₄, as well as HNO₃, which are commonly used during sample

preparation of the fly ash samples are avoided in this method. The spectral effects due to the

formation of different Cl, Na, K, Ca, Mg, containing polyatomic species interfering the

determination of number of elements like As, Se or Ni during the oaTOF-ICP-MS analysis are

negligible. Under the optimum experimental extraction conditions evaluated using a fractional

factorial design being 10 mg of the sample extracted with 5 mL of 140 g L⁻¹ NH₄F for 10 min at

200 °C, subsequent analysis of the resulting supernatant enabled under optimum instrumental

conditions and with Rh as an internal standard precise and accurate simultaneous determination

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of 11 elements (Li, Be, Ni, As, Se, Rb, Sb, Cs, W, Tl and U) at trace and ultra-trace levels. The accuracy was assessed by analysing two certified reference materials, namely Fine Fly Ash CTA-FFA-1 and Constituent Elements in Coal Fly Ash Standard Reference Material[®] 1633b. The precision of the reported method was better than 10 %.

Keywords: oaTOF-ICP-MS; Trace elements; Fly ash; Microwave-assisted extraction; Fractional factorial design; Ammonium fluoride

Introduction

Fly ash as a by-product of coal combustion is a significant source of potential toxic elements and its analysis is important for evaluation of the associated and environmental risk as well as for its further use in the industry [1, 2]. From an environmental perspective, trace elements have attracted more attention than the major and minor constituents [3] and many analytical methods like inductively coupled optical emission spectrometry (ICP-OES) [4-17], graphite furnace atomic absorption spectroscopy (GFAAS) [14, 18, 19], hydride generation atomic absorption spectrometry (HGAAS) [14, 19], atomic fluorescence spectrometry (AFS) [20], inductively coupled plasma mass spectrometry (ICP-MS) [3-5, 7, 8, 12, 15, 16, 19, 21-28], instrumental neutron activation analysis (INAA) [6, 19], etc. aim at measuring the elements associated with coal and coal by-products at concentrations down to the trace level.

Inductively coupled plasma mass spectrometry has arisen as one of the preferred methods due to its unique combination of low detection limits, capabilities for rapid multi-elemental and isotopic determination and wide dynamic range [23]. Despite the significant improvements on solid sample introduction techniques such as laser ablation sampling [29] or slurry sampling

electrothermal vaporization [30-34], solution nebulization is still the most preferred method for many ICP-MS applications [35].

Coal fly ash however represents a complex and resistant material mainly composed with oxidized compounds of Si, Al, Fe and Ca [36], due to this fact the quantitative liberation of toxicologically important elements is quite difficult and has been for many researchers an analytical challenge for a long time. The methods involved in sample preparation of fly ash samples employed conventional alkali fusion, wet digestion, sequential extraction [7, 8, 12], ultrasound assisted digestion or extraction [10] and microwave digestion and extraction [20, 37]. Employment of the alkali fusion which is traditionally used for preparation of coal and coal ash samples [38, 39] results however in a high solid environment of the fused materials. The necessity of vast amounts of concentrated reagents then represents a risk of contamination and may be responsible for matrix interferences [39, 40]. Conventional wet digestion methods are time consuming, susceptible to loss of volatile analytes and contamination, requiring large amount of reagents [36]. Chemical sequential extraction for element fractionation in coal fly ashes [7, 8, 12] has been proved an effective tool to investigate various binding forms of different elements. These methods are however relatively complicated, time-consuming and expensive and so should be only used when appropriate [41]. Ultrasound and microwave-assisted extraction have been investigated in an attempt to improve the yield of the extracted metal and to reduce the process time, especially with the increasing demand for more environmentally friendly processes [20, 41-43]. Both methods are increasingly finding applications in analytical chemistry and are useful for fast screening of elements in the environment [41]. They cannot however ensure a quantitative extraction of wide range of analytes compare to closed vessel microwave-assisted digestion being currently the most widely used preparation technique of coal and coal fly ash samples to overcome the main drawbacks of the conventional digestion methods [36]. In contrast

to this fact, ultrasound and microwave-assisted extraction (MAE) are known as one of the best green technologies with advantages like high extraction efficiency and sample throughput, good reproducibility, low consumption of solvents and time, and low carbon dioxide output [44].

In recent years, both the digestion and extraction steps have been carried out using very strong acidic and oxidizing reagents like HNO₃, HCl, H₂O₂ and their mixtures as the best way to achieve satisfactory results [20]; highly toxic HF and explosive HClO₄ are usually employed to quantitatively determine trace elements in coal fly ash and coal [36]. The employment of HF may however result in a damage of glass components of the instruments and may lead to formation of several volatile fluorides (many analytes such as As, B, Se, Sb, Hg and Cr may volatilize) [24]. The formation of such volatile compounds presents a serious risk of analyte loss after opening the vessel, even if the decomposition of the sample has taken place in a closed system [45, 46]. However, this problematics is much more complex and the final reaction products, which will be created, are strongly dependent on the conditions of the experiment [47, 48]. The addition of H₃BO₃ [24, 36] may solve some drawbacks being attributed to the use of HF, but the addition of H₃BO₃ increases the salinity of the sample, the price of the analysis and disables the determination of boron in the sample [49].

In the ICP-MS analysis of coal and coal ashes, several elements like Si, P, S, K, Ca, Fe and Cl origination from sample matrix or the employment of acids being used in sample dissolution process are problematic as they serve as precursors for the formation of polyatomic ions [15, 16, 24, 26, 36, 50] which can be problematic to be handled with especially with commonly used quadrupole mass spectrometers with low-resolution [7, 21, 26, 27, 37]. The collision/reaction cell ICP-MS is an effective correction technique to minimize the interferences on the analyte of interest for conventional quadrupole ICP-MS [7, 21, 26, 27]. While a decrease of the multi-element capabilities could be observed for older Q-ICP-MS systems, this problem

has been dealt with for modern quadrupole ICP-MS instruments. Samples with a complex matrix can be dealt with a sector-field ICP-MS which can handle polyatomic interferences originating from major elements thanks to the high mass resolution of the spectrometer. The price for the higher resolution is however the loss of sensitivity. On the other hand, the multi-element capability is kept. Nevertheless, the high cost of the instrumentation may make these methods less accessible to routine laboratories.

ICP time-of-flight mass spectrometers are more reachable for routine analytical practice when compared with the HR-ICP-MS instrumentation. These instruments' advantage is the simultaneous mass determination and high speed without any compromise in mass range, sensitivity or precision [51]. The higher resolution of the oaTOF mass analyzer compare to Q-ICP-MS could be an advantage while determining several analytes such as Sc in the presence of large quantities of Si [52], however, it is not adequate to completely deal with the interference signals caused by polyatomic species observed during the analysis of samples with a complex matrix [53-55]. Several approaches such as the mathematical correction equations [53, 55] without [53] or with the use of microwave digestion [55] or methods employing matrix modification [54, 56] were successfully used previously during the analysis of different sample matrices in order to manage the spectral effects during the TOF-ICP-MS analysis.

Several works can be found describing the analysis of fly ash using quadrupole ICP-MS [7, 21, 26, 27, 37] which feature collision and/or reaction cells or mathematical corrections for attenuating the interferences and high resolution ICP-MS instruments [15, 23]. However, no method is currently available for this purpose using the TOF-ICP-MS. Modified sample preparation process is introduced in this work using ammonium fluoride for extraction of a several elements from coal fly ashes prior to TOF-ICP-MS analysis. In combination with a closed vessel microwave-assisted extraction (MAE), with the flexibility of vessel options, choices of

sample size, temperature, amounts of solvent, and number of samples [57], simple, rapid, interference-free and high-throughput method was developed. Using the described method, the use of some other reagents like mineral acids, being usually necessary for complete sample digestion is not needed which is in agreement with green chemistry trends [58].

Experimental

Instrumentation

The detailed description of the Optimass8000 oaTOF-ICP-MS instrument used throughout this study (GBC Scientific Equipment Pty. Ltd., Australia) is given in ref. [51]. The details of the typical operating and instrument conditions can be found in Ref. [53-56]. The resolution (full width at half maximum intensity) of 1800 and sensitivity of 15000 counts s⁻¹ per μ g L⁻¹ (mass integrated peak) was achieved for ¹³⁹La⁺ under these conditions. The same parameters were 2100 and 17000 counts s⁻¹ per μ g L⁻¹, respectively for ²³⁸U⁺.

The microwave-assisted extraction of samples was carried out in SpeedwaveTM MWS-3⁺ (Berghof, Germany) microwave system with the maximum total output of the microwave generator 1450 W. The sample throughput of the MWS-3⁺ system was increased by using the Multitube System (MT) employed with the DAC-100S digestion vessels [49]. This arrangement allows simultaneous extraction/digestion of three samples in one DAC-100S PTFE vessel by placing three MT PFA tubes into each of the vessels.

The Avanta P double beam atomic absorption spectrometer (GBC Scientific Equipment Pty. Ltd., Australia) was used for determination of Na, K, Ca, Fe, Si and Mg in the flame emission and absorption mode, respectively. The operating and instrument condition are in detail summarized in Ref. [49, 56].

The determination of C was performed by an independent accredited analytical laboratory using the Eltra CW-800M analyzer (Eltra GmbH, Germany).

Reagents and standards

The UltraClear (SG, Germany) pure water system was used to purify water to $0.05~\mu S$ cm $^{-1}$, which was further employed for preparation of all solutions.

Nitric acid (65%, w/w) of Selectipur quality (Lach-Ner, Neratovice, Czech Republic) was further purified by sub-boiled distillation. Stock solution containing 1 g L⁻¹ Rh obtained from SCP Science (Canada) was used to prepare internal standard solution. Multi-element stock solution "A" comprising of 10 mg L⁻¹ of Ag, Al, As, B, Ba, Be, Bi, Cd, Co, Cr, Cs, Cu, Ga, Ge, Hf, Hg, Li, Mn, Mo, Ni, Pb, Pd, Pt, Rb, Re, Ru, Sb, Se, Sn, Sr, Ta, Te, Th, Ti, Tl, U, V, W and Zr was prepared from single element standards of 1 g L⁻¹ obtained from SCP Science or Analytika Ltd. (Czech Republic). Stock solution "B" containing 100 mg L⁻¹ of Ce, La, Nd, Pr, U ("B1") and 20 mg L⁻¹ of Dy, Eu, Er, Gd, Ho, Lu, Sc, Sm, Tb, Tm, Y, and Yb("B2") was prepared using a multi-element solution purchased from Analytika Ltd. and 10 g L⁻¹ of U. Ammonium fluoride was of pro-analysis grade (p.a.) and obtained from Sigma-Aldrich (USA). The stock solutions of Ca, Fe, K, Mg, and Na (1 g L⁻¹) were purchased from Analytika Ltd. (Czech Republic).

Quality assurance and quality control

The commercially supplied reference materials Constituent Elements in Coal Fly Ash Standard Reference Material® 1633b (National Institute of Standards & Technology, USA) and certified reference material Fine Fly Ash CTA-FFA-1 (Institute of Nuclear Chemistry and

Technology, Warszawa, Poland) intended primarily for use in calibration and verification of apparatus and methods respectively for determining components in fly ashes were analyzed.

Microwave-assisted extraction

The certified reference material Fine Fly Ash CTA-FFA-1 was employed for optimization of the MAE procedure. The portion around 250 mg of this sample was at first ground in the vibration mill Wig-L-Bug 30 (Crescent Dental Mfg. Co.) for reassurance of the sample homogeneity. Then, 10 mg of the CTA-FFA-1 were weighed into the MT-tubes and NH₄F was added at constant volume 5 mL with different concentration of NH₄F being 60, 100 and 140 g L⁻¹, respectively, which equals to the absolute amount of 300, 500 and 700 mg of NH₄F in the MT-tubes. These MT-tubes were placed into the outer digestion vessel, where 20 mL of the appropriate concentration of NH₄F were added. The level of NH₄F is higher in the outer vessel than in the PFA tubes. The evaporation of the solution from the PFA tubes is thus prevented by compensation of the vapour pressures [49].

The extractions were performed at the temperature of 160, 180 or 200 °C held for 5, 10 or 15 minutes (with a ramp set at 5 min at 50 % power of the MWS-3⁺). The resulting solutions were diluted to 10 mL with deionised water. Before analysis all solutions were filtrated through a 0.45 µm Nylon syringe filters (Whatman Autovial).

Sample analysis

Quantification of the concentration of the trace element by oTOF-ICP-MS was done by constructing calibration curves by linear regression. The differences in the calibration slopes of fly ash extracts and aqueous standards were insignificant at significance level 0.05 as will be

discussed later thus the employment of simple calibration standards was possible. Calibration solutions: blank, 1, 2, 5, 10, 20 and 50 μ g L⁻¹ of Ag, Al, As, B, Ba, Be, Bi, Cd, Co, Cr, Cs, Cu, Ga, Ge, Hf, Hg, Li, Mn, Mo, Ni, Pb, Pd, Pt, Rb, Re, Ru, Sb, Se, Sn, Sr, Ta, Te, Th, Ti, Tl, U, V, W and Zr and 0.1, 0.2, 0.5, 1, 2 and 5 μ g L⁻¹ Ce, La, Nd, Pr, U and 0.02, 0.04, 0.1, 0.4 and 1 μ g L⁻¹ of Dy, Eu, Er, Gd, Ho, Lu, Sc, Sm, Tb, Tm, Y, and Yb were prepared by dilution of multielement stock solutions "A" (500 μ g L⁻¹) and "B" (50 + 10 μ g L⁻¹) in 50 mL volumetric flasks on a daily basis. Any possible instrumental drift and non-spectral effects were compensated by addition of 1.0 μ g L⁻¹ of Rh to all samples and solutions. Regression coefficients greater than 0.999 were obtained regularly. Five second data acquisition time, ten replicates, and peak area mode were used for the measurement. The solutions were analyzed along with a blank containing the same amount of ammonium fluoride and prepared in the same way as the samples. Before the analysis, the samples were diluted with deionized water in 1:4 ratios.

The analysis by flame atomic absorption/emission spectrometry for the determination of Na, K, Ca, Fe, Si and Mg, respectively was performed using the same way as described previously [49, 56].

Experimental design and statistical data treatment

Three-level 3(k-p) fractional factorial design (where k is the number of factors investigated, and p describes the size of the fraction of the full factorial used), with three factors (3(3-1)) and with three replicates (n = 3), was used to find the optimal conditions for the extraction and to evaluate the significance of the factors (variables), as well as the interactions between them [59, 60]. The fractional factorial experimental design 3^{3-1} (n = 3) was used because of its reduced number of experiments (27 against 81 experiments using the full design) but also because no loss

of significant information occurs. The studied factors were: mass of the extraction agent (A), extraction temperature (B) and extraction time (C). All factors were evaluated at three levels, low (denoted as -1), middle (denoted as 0) and high (denoted as +1). Table 1 summarizes the levels for the experimental design.

"INSERT TABLE 1 HERE"

The responses for each experiment were calculated as the average mean of concentrations obtained for Li, Be, Ni, As, Se, Rb, Sb, Cs, W, Tl and U in CTA-FFA-1 Fine Fly Ash. The yield of the extraction data (%) for the Li, Ni, As, Rb, Sb, Cs, W, U, i.e. elements whose concentration was certified and resulted in a quantitative extraction, are summarized in Table S1.

The Statistica 12 computer program (StatSoft, Inc., USA) and QC ExpertTM 2.5, TriloByte Statistical Software, (Pardubice, Czech Republic) were used for data processing.

Results and discussion

Selection of extraction agent

We have reported previously [49, 61] that ammonium fluoride is able to impact effectively both silicate and alumina matrix. If ammonium fluoride or ammonium hydrogen fluoride are the reagents, different reaction products such as oxides, hydroxides, salts, etc. are created for many analytes and their compounds compare to the reaction with HF [62]. For ammonium fluorides, both in the anhydrous state or in the aqueous solution, the formation of ammonium fluorometallates or oxofluorometallates was reported [62]. The resulting products are in contrast to those obtained with HF more thermally stable [49, 62], NH₄F is also significantly

less toxic than HF, which is a requirement for the replacement of the hydrofluoric acid with all their handicaps and the utilization of NH₄F in a sample preparation step. When the experimental conditions employing a closed vessel microwave-assisted extraction with the use of NH₄F as an extractant are used, Si and B can be quantitatively determined in various environmental samples [49]. When the experimental conditions employing a closed vessel microwave-assisted extraction with the use of NH₄F as an extractant are used, Si and B can be quantitatively determined in various environmental samples [50]. Quantitative extraction of larger number of analytes with the use of only NH₄F would be favourable in this point of view. Relating the ICP-MS analysis all the above mentioned facts may be important in several aspects. Increasing number of reaction agents may influence the formation of different polyatomic species arising from ions from both of the reagents and from those matrix components which have been by this agent released to the analysed solution. In addition, higher number of reagents increases the risk of contamination. Due to this fact NH₄F was investigated for extraction of different analytes prior to ICP-MS measurement.

Optimization of the experimental conditions for microwave extraction

Three parameters were examined on the extraction results and efficiency, respectively, for individual analytes, which include the mass of the extraction agent, temperature and extraction time (see Table 1). The effects and significance of the variables in the extraction process were evaluated using Pareto's charts, including a vertical line that corresponds to the 95% limit which indicates the statistical significance. Pareto's charts were constructed for total of 11 elements for which accurate concentration was evaluated via the two certified reference materials of fly-ashes as will be discussed later. Figure 1 shows that the factors influencing the extraction efficiency are

strongly dependent on the type of the element. For the extraction of Li, Be, Se, Rb, Sb, Cs, Tl and U the mass of the extractant has the most positive effect on the extraction efficiency. While for Li and Be, the analytical response will increases linearly, respectively quadratically with the amount of extractant, for other of the above mentioned elements both linear and quadratic effects are important relating the factor of the mass of the extractant. What can be further seen from the Fig. 1 is that the extraction temperature significantly influences the extraction efficiency of Li, Rb, Cs, As and U. While for As and Cs the analytical response is improved linearly by increasing the extraction temperature, for Li, Rb, and U the effect is opposite and the extraction efficiency drops quadratically with increasing temperature. However, for Li the values obtained via the experiment are within the acceptable interval given by the producer of the CRM and analytical recoveries fall within the range of 75–110 %. The time of the extraction has influenced the extraction of Be, Ni, W, Rb and Tl. While for Be, Ni and W positive effect on the extraction efficiency was observed, for Rb and Tl the situation was opposite. Analytical recoveries for Rb were however within 53–104 %, while for Tl it could not be calculated as in the CRM used for optimization the Tl level was not certified. Important interactions between the mass of the extractant and temperature were observed for the extraction of Li, Be, Se, Rb, U, Sb, W and Tl. The interactions observed influence in all cases the analytical results positively.

"INSERT FIGURE 1 HERE"

The results of the 3³⁻¹ design are presented in Table S1 from which corresponding response surfaces were constructed as illustrated in Fig. 2. Each plot shows the effects of two variables within their studied ranges and visualizes the tendency of each factor to influence the extraction efficiency of investigated element. In order to ensure the most robust conditions for the

utilization of the method to as many analytes as possible, experimental conditions were finally selected as follows: 700 mg of the extractant, 200 °C extraction temperature held for 10 min.

Quantitative extraction was achieved for all of the elements mentioned above (see Fig. 2) under these extraction conditions.

"INSERT FIGURE 2 HERE"

Evaluation of matrix effects

Concentration of matrix elements in analysed samples significantly influences the occurrence of both spectral and non-spectral matrix effects during the oaTOF-ICP-MS multielement analysis [53-56]. The quantitative values of apparent element concentration as measured on the Optimass 8000 in the presence of 0–1000 mg L⁻¹ of the elements (C, Ca, Cl, K, Mg, Na, P, and S) suspected to generate polyatomic ions interfering the determination of important elements such as As, Co, Cr, Cu, Ga, Mn, Ni, Se, Ti, V or Zn, etc. were previously published [53, 55]. Table 2 summarizes the concentration data of important matrix elements determined in the extract of CTA-FFA-1 and NIST SRM 1633b as well as the concentrations declared by the producers. It is obvious from the data that the concentration of major elements like Ca and Mg resulted in very low concentration values which can be attributed to the formation of insoluble fluorides. The complete dissolution of Si from the materials was indicated by the obtained values, which were in accordance to the certified values. Relating the concentration of Na, the quantitative extraction was achieved only for the NIST SRM 1633b. For CTA-FFA-1 whose Na concentration was one order higher than for NIST SRM 1633b, the

recoveries were of about 32 %. Similarly, the recoveries of K in CTA-FFA-1 and NIST SRM 1633b were only 60 and 61 %, respectively. Although KF and NaF are easily soluble in the water, different reaction products may be created with ammonium fluoride like oxofluorometallates, etc. for which different solubility will be observed.

The presence of major matrix elements whose quantitative extraction is not achieved will significantly less influence the determination of problematic elements in contrast to procedures where total sample decomposition is used. Relating the concentration of Ca, Mg, Na and K whose concentration in supernatants is lower than 25 mg L⁻¹ these values will not significantly interfere during the oaTOF-ICP-MS analysis [53, 55]. Similarly, the interference from polyatomic ions containing P or S is not expected to be significant as the concentration of the analytes in original samples is about one order lower in contrast to the above mentioned elements. According to the certificates the concentration of P and S in NIST SRM 1633b is 2.3 g kg⁻¹ and 2.075 ± 0.011 g kg⁻¹, respectively and 725 ± 74 mg kg⁻¹ for P in CTA-FFA-1.

"INSERT TABLE 2 HERE"

As the ammonium fluoride matrix can have a considerable effect on the progress of the analysis and the quality of the obtained results a calibration curve using aqueous standard samples and one containing ammonium fluoride in the same concentrations as used for sample preparation was constructed for all elements determination. Table 3 shows the comparison of the slope values of matrix matched calibration with the slopes of aqueous calibration. The presented slopes of the calibration lines are expressed as relative sensitivities. Relative intensity was analytical response defined by counts per second (cps) of the peak area of individual analytical

signal calculated and averaged from the ten measurements at each concentration and divided by the response of the internal standard element Rh. The paired *t*-test (at significance level 0.05) revealed that the slope values of matrix matched calibration and that of aqueous calibration are not significantly different as its *p*-value was 0.185. The presence of ammonium fluoride thus does not influence the sensitivity and is not responsible for inducing non-spectral effects during the oaTOF-ICP-MS analysis. Thus, for quantification a direct aqueous calibration method using Rh as internal standard element, which is less time consuming than standard additions or the matrix matched method thus may be used.

"INSERT TABLE 3 HERE"

Trueness and precision

Coal Fly Ash NIST SRM 1633b and Fine Fly Ash CTA-FFA-1 certified reference materials were analyzed in order to prove the trueness and precision. As seen from Table 2, good agreement between the results obtained using an aqueous calibration for quantification was achieved for 11 elements by the proposed microwave-extraction method and the declared values. The accuracy was for elements whose concentration was not certified assessed using a second approach. This involved the calculation of analytical recoveries, which were determined for samples of extracts in three replicates using standard additions as shown in Table 2. For all elements, the obtained recoveries were between 91 and 106 %, which confirms that the accuracy of the method is within 10 %. Using the described procedure an accurate results were obtained also for the elements like As, Se or Ni for which the use of collision/reaction cell [7, 21, 26] or mathematical corrections [37] are needed when using Q-ICP-MS.

However, the determination of some of the trace elements like Cd, Pb, Ba, Cr, Hg, etc. which can be considered as potentially toxic [63], could not be accurately done by the presented method. Low recovery yields of trace elements may be attributed to the formation of insoluble fluorides or co-precipitation with the fluoride. The insoluble fluoride precipitates formed during NH₄F extraction of fly-ash samples may incorporate trace elements differentially into the structures of the precipitated major element fluorides, which therefore act as carriers for trace elements [35]. In addition, some analytes like Pb, Zn, Cd or Cu exist in carbonate phase [12] of fly ash that influences their leaching behavior.

Intra-day comparison was used to evaluate the precision of the method. It is expressed as relative standard deviation (RSD) of analyses of a sample, which was independently prepared and analysed three times during one day. The value used for the computation was obtained as a mean of an analysis having ten replicates. Table 2 shows that the values of the intra-day RSD were typically lower than 10 %. Thus, the precision of the method was acceptable. This also confirms the sufficient homogeneity of the material as these values of RSD were obtained for sample masses being 10 mg. These observations that the use of such small sample mass does not affect the trueness or the precision of the analysis of coal and coal ash are in accordance to those reported previously by Lachas et al. [3].

Limits of detection and quantification

Table 3 shows the limits of detection (LODs) and limits of quantification (LOQ) for each isotope, which were calculated as a three times, respectively 10 times the standard deviation of the blank response at the given m/z position divided by the slope of the calibration graph. The values are evaluated as the instrument LOD and method limits of detection and quantification.

The time of the integration was set to 5 s and blanks and standards were determined in ten replicates. In order to evaluate the instrument limits of detection the blanks containing deionized water alone or with ammonium fluoride and internal standard in all cases to reflect how the simple presence of NH₄F influences the detection capability were analysed. The problems with contamination were of a little importance except for Ni, Rb, or Sb as is shown in Table 3. This was ensured by using ammonium fluoride available in sufficient purity.

Sample throughput

The analysis of each sample comprised of sample introduction, ten-replicate measurements and rinsing. Under the conditions summarized in experimental section, the simultaneous analysis of all elements took for each sample approximately three minutes, which means that ca 20 samples could be analysed in an hour. Using the full capacity of the 8-position carousel of the MWS-3+ microwave oven and the PFA multitube system [49] three MT PFA tubes were placed into a standard DAC-100S PTFE vessel, thus three samples were extracted simultaneously in one DAC-100S digestion vessel and therefore it was possible to extract up to 24 samples in one microwave run. The total time needed for the microwave procedure is 30 min (including extraction and cooling down of the reaction vessel) for 24 samples. The sample preparation, i.e. weighing, pipetting of the extractant, diluting etc., takes approximately 90 min for 24 samples. With this setup, it is possible to prepare and analyse a minimum of 96 samples in an 8 h working day, which equals to 12 samples h⁻¹.

Conclusions

The presented method using ammonium fluoride for extraction of the elements from fly ash samples offers an efficient and simple way how to accurately quantify 11 elements in this difficult matrix by oaTOF-ICP-MS method. The main benefit of the proposed approach is that the matrix interference is insignificant. Thus, simple calibration method can be performed for precise and accurate determination of elements even such as As, Se or Ni, which are more problematic if ICP-MS instruments with inadequate resolving power are used. Another advantage of this procedure is its speed and simplicity. The determination of all elements is done simultaneously; individual analytes do not require specific setting, which may be another benefit. As the purpose of this proposed method is to eliminate different polyatomic ions causing spectral interferences, the procedure is expected to be beneficial both for Q- and for TOF-ICP-MS. The method is avoiding handling of the very corrosive and toxic HF. The relatively low price of NH₄F compared to HF, no significant waste generation and speed of MAE are other benefits. The method thus contributes to the green chemistry. However, the quality of the used ammonium fluoride limits the LODs. Additionally, some elements of potential interest like Ba, Cd, Hg, Pb, etc. could not be determined accurately using the presented approach due to problems with inclusion and/or co-precipitation in the precipitated matrix. Further, relating the analysis of solutions with high fluoride content, special attention should be paid to the care of ICP-MS cones. Rather than using nickel cones using more chemically resistant Pt-cones is recommended as they can usually be used for longer before they need cleaning. Nevertheless, this procedure is despite the above mentioned drawbacks for oaTOF-ICP-MS instrumentation this procedure is currently the sole solution to reduce the interferences to a manageable level. On the other hand, it is expected that the quantification of elements like Cd, Hg, Pb, etc. would be easily performed using this instrumentation when employing some of the more conventional sample preparation

procedures ensuring quantitative release of desired analytes from the sample matrix as for the elements above 80 amu the spectral effects are significantly less important.

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Figures

Fig. 1 Pareto charts of the standardized effects in the fractional factorial design 3³⁻¹ for the study of variables (A) amount of NH₄F, (B) extraction temperature, (C) time and their interactions on extraction of selected elements. The L and Q letters indicate the linear and quadratic effect of the factor, respectively.

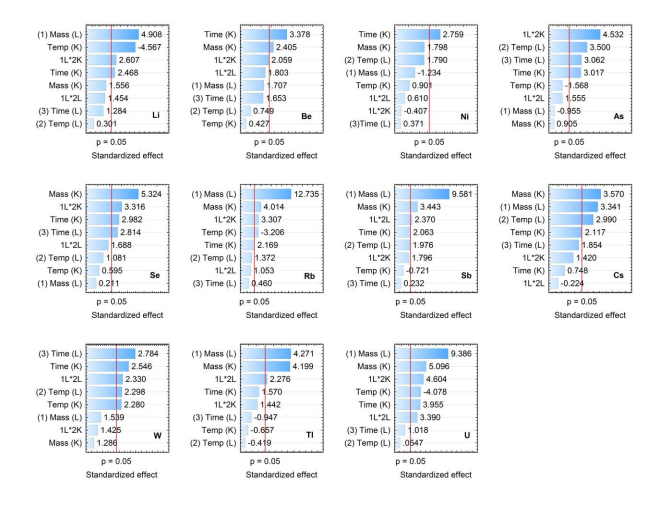
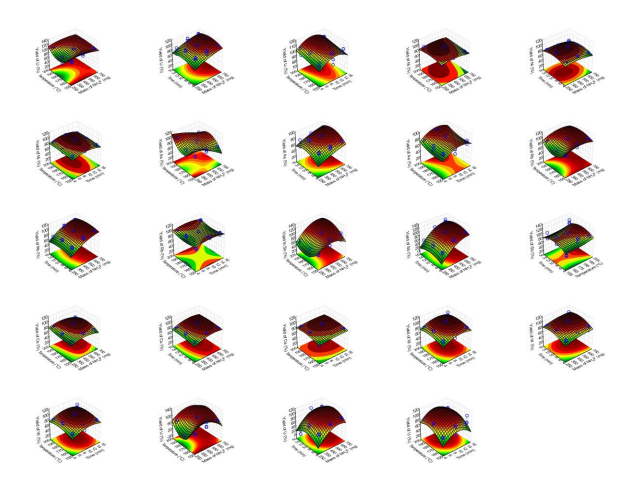


Fig. 2 Response surface plots from 3^{3-1} design for the recovery yield (%) of selected elements in the NH₄F extract as a function of NH₄F amount, extraction temperature and time. Recovery yields (%) were evaluated as (the ratio of the value obtained from an analytical process via a calibration graph/ the reference value) x 100.



TablesTable 1 Variables and levels investigated using the fractional factorial experimental design

Variables		-1	0	+1	
Amount of NH ₄ F (mg)	A	300	500	700	
Extraction temperature (°C)	В	160	180	200	
Extraction time (min)	С	5	10	15	

Table 2 Summary of results for the analysis of CTA-FFA-1 and NIST SRM 1633b

	CTA-FFA-1			NIST SRM 1633b				
Trace	Concentratio	n ^{a,b} mg kg ⁻¹	R^d	RSD^f	Concentration ^{a,b} mg kg ⁻¹		R^d	RSD^f
Elements	Declared	Found	%	%	Declared	Found	%	%
⁷ Li ⁺	128 ± 22	133 ± 4	104	1.6	С	187 ± 24	99 ^e	6.3
$^9\mathrm{Be}^+$	27	28 ± 3	104	5.3	с	12.3 ± 1.7	104 ^e	7.0
$^{60}\mathrm{Ni}^{^{+}}$	99 ± 6	95 ± 15	96	8.2	120.6 ± 1.8	121.9 ± 4.2	101	1.7
$^{75}As^+$	53.6 ± 2.7	51.0 ± 2.9	95	2.9	136.2 ± 2.6	133.7 ± 1.4	98	2.3
$^{78}\mathrm{Se}^{^+}$	4.6	4.7 ± 0.7	102	6.9	10.2 ± 0.2	10.2 ± 1.6	100	7.9
$^{85}\text{Rb}^{+}$	185 ± 5	184 ± 11	99	3.0	140	143.1 ± 4.2	102	1.5
$^{121}Sb^{+}$	17.6 ± 2.5	16.8 ± 1.6	95	4.9	6	6.2 ± 0.6	103	4.5
$^{133}\text{Cs}^{+}$	48.2 ± 2.3	45.6 ± 2.3	95	2.5	11	10.0 ± 0.9	91	4.7
$^{182}W^{+}$	10.5 ± 1.1	11.1 ± 0.7	106	2.9	5.6	5.6 ± 0.2	100	2.2
$^{205}\mathrm{Tl}^{+}$	c	7.3 ± 0.2	104 ^e	0.2	5.9	5.5 ± 0.2	93	6.3
$^{238}U^{+}$	15.1 ± 0.8	14.3 ± 1.5	95	5.1	8.8 ± 0.4	8.4 ± 0.2	95	5.8
Major	Concentration ^{a,b} g kg ⁻¹		R^d	RSD^f	Concentration ^{a,b} g kg ⁻¹		R^d	RSD^f
Elements	Declared	Found	%	%	Declared	Found	%	%
Na	21.9 ± 0.8	7.1 ± 0.8	32	5.6	2.01 ± 0.03	2.09 ± 0.16	104	3.8
K	22.0	13.1 ± 0.8	60	3.1	19.5 ± 0.3	11.8 ± 0.8	61	3.4
Ca	22.9	< 0.033	g	g	15.1 ± 0.6	< 0.033	g	g
Mg	15.5	2.3 ± 0.2	15	4.3	4.82 ± 0.08	1.43 ± 0.14	30	4.9
Fe	48.9 ± 1.4	5.6 ± 0.3	12	3.1	77.8 ± 2.3	29.3 ± 3.5	38	6.0
Si	224.8 ± 9.2	216.3 ± 4.1	96	1.0	230.2 ± 0.8	230 ± 18	100	3.9
C a Maan + 2	С	1.15 ± 0.15^{h}	g	6.5	с	$2.7 \pm 0.5^{\rm h}$	g	9.3

^a Mean \pm 2 SD (n = 3).

 $^{^{}b}$ Values were evaluated with the use of calibration with matrix-free standards (n = 5) and Rh as internal standard element.

^c Not available.

^d Recovery (%) expressed as (the ratio of observed value obtained from an analytical process via a calibration graph/ the reference value) x 100.

 $^{^{\}rm e}$ Analytical recovery, R (%) evaluated for samples spiked with 5 $\mu g \; L^{\text{--}1}$ for listed elements.

^f Relative standard deviation (%) of three independent analyses.

^g Not determined.

^h The determination of C was performed in original samples by the Eltra CW-800M analyzer.

Table 3 Comparison of linear calibrations method using a least square regression for matrix-free standards and those containing ammonium fluoride for selected elements determination by oaTOF-ICP-MS method and appropriate LODs and LOQs values

	Relative sensitivity ^a		Instrument LOD (ng L ⁻¹)		Method LOD and LOQ (μg kg ⁻¹)	
	Aqueous ^b	NH ₄ F ^c	Aqueous ^b	NH ₄ F ^c	LOD	LOQ
⁷ Li ⁺	0.1112	0.1298	65	37	37	123
⁹ Be ⁺	0.0437	0.0424	87	98	98	327
$^{60}\mathrm{Ni}^{^+}$	0.1054	0.1207	73	314	314	1047
$^{75}As^+$	0.0609	0.0564	67	150	150	500
$^{78}\mathrm{Se}^{\scriptscriptstyle +}$	0.0106	0.0083	726	476	476	1587
$^{85}\text{Rb}^{+}$	0.4249	0.4663	9.0	209	209	697
$^{121}\mathrm{Sb}^{+}$	0.2226	0.1958	3.4	31	31	103
¹³³ Cs ⁺	0.9086	0.9141	1.3	2.1	2.1	7.0
$^{182}W^{+}$	0.436	0.4367	2.7	17	17	57
$^{205}\text{Tl}^{+}$	1.2432	1.1742	0.37	1.6	1.6	5.3
$^{238}U^{+}$	2.2848	2.1722	0.04	0.09	0.09	0.3

^a Relative sensitivities are simply the computed slopes of the calibration lines where relative intensity was the analytical response defined by counts per second (cps) of the peak area of individual analytical signal calculated and averaged from the ten measurements at each concentration and divided by the response of the internal standard element Rh.

 $^{^{}b}$ Calibration with matrix-free standards (n = 5).

 $^{^{\}text{c}}$ Each calibration solution (n = 5) contains the same amount of NH₄F as the sample.