Matrix effect on emission/current correlated analysis in laser-induced breakdown spectroscopy of liquid droplets

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Abstract

We have investigated influence of matrix salts on the liquid droplets by laser-induced breakdown spectroscopy (LIBS). An electrospray ionization technique coupled with LIBS is employed to generate the microdroplets of the Na sample solution with various matrix salts added. A sequence of single-shot time-resolved LIB emission signals is detected. The LIB signal intensity integrated within a gate linearly correlates with the plasma-induced current response obtained simultaneously on a single-shot basis. The slopes thus obtained increase with the sample concentration, but appear to be irrespective of different matrix salts, added up to a 2000 mg/l concentration. The matrix salts involved have the same K⁺ cation but different anions. Given a laser radiation emitting at 355 nm with the energy fixed at 23 ± 1 mJ, a limit of detection (LOD) of 1.0 mg/l may be achieved for the Na analysis. The current normalization might have probably taken into account the ablated amount of the sample and the plasma temperature. Accordingly, the LIB/current correlated analysis becomes efficient to suppress the signal fluctuation, improve the LOD determination, and concurrently correct the matrix effect.

Keywords: Laser-induced breakdown spectroscopy; Matrix effect; Electrospray ionization; Correlation method

1. Introduction

Laser-induced breakdown spectroscopy (LIBS) has grown to be a popular analytical technique for rapid analysis of a large variety of materials present in solids, gases, and liquids [1–11]. This technique provides the advantages of multi-elemental detection, non-invasive analysis, suitability to remote measurement, and minimal sample preparation. Nevertheless, it suffers from the emission signal fluctuation, which may restrict the LIBS sensitivity to some extent. However, the matrix composition contained in the analyzed materials has substantial impact on the accuracy of the elemental analysis. How to improve the signal quality and correct the matrix effect have been important issues in this field.

The signal fluctuation problem in the LIBS analysis are caused by complicated sources such as the laser pulse fluctuation, inconsistency of the sampled amount of medium, instability of the plasma position, and the inverse bremsstrahlung radiation. Averaging the spectral signals over a number of laser shots is a way usually used to overcome this problem. Nevertheless, the improvement of the signal-to-noise ratio is offset by the non-linear optical ablation effect [7,8]. Internal standard method is an alternative solution via the measurement of the intensity ratio of analyte and reference element [6,9–11].

Another interference caused by the matrix effect has also been widely investigated to achieve matrix-corrected LIB emission signals [11–16]. For instance, Eppler et al. reported that in the sand and soil samples the LIB emission signals of Pb and Ba depend on the chemical speciation and the matrix composition [14]. They related the influence on the LIB measurement to plasma excitation conditions, the physical properties of the sample compound, matrix absorptivity, electron density, and plasma temperature. Such complicated factors may cause the accuracy of LIB measurement to be degraded if the calibration is not compound and matrix-specific. Chal-
eard et al. quantized the LIB emission signals in air at atmospheric pressure by taking intensities of the emission lines as a function of the vaporized mass and the plasma excitation temperature [15]. Normalization of the LIB emission signals by these two parameters allowed for an efficient correction of matrix effects when the sample composition was changed. Hakkanen et al. studied the matrix effects in analyzing inhomogeneous paper coatings and found linear correlations between the LIB emission signals and the coat weight and the binder contents of the coatings [16]. Winefordner and coworkers investigated the influence of matrix effect on LIB signal of Mg in a powdered sample [17]. They suggested a method for matrix-free calibration based on normalization of the emission intensities by the surface densities of the analyzed materials. The surface density normalization method may account for the surface densities of the analyzed materials. The laser-interacted spot was approximately 100-μm diameter, but not measured precisely. The pulse-to-pulse fluctuation remained at ±5%.

2.2. Electrospray ionization

A homemade electrospray ionization device was employed to generate a stream of microdroplets, as described previously [22]. In brief, the analyte solution was syringe-pumped into a 200-μm i.d. stainless steel spray needle. The needle served as the anode, biased at 3 kV, through which the analyte solution was sprayed toward the other metal base as the cathode. As carrier solvent evaporated, the analyte solution was then broken into droplets [20, 21]. The laser pulse interacted with the sprayed droplets at ~2 mm downstream from the needle tip. The irradiated spot was close to the tip of Taylor cone where a higher density of microdroplets may be obtained.

2.3. LIB and current detection

Upon laser-pulse interaction with the liquid droplets, the produced luminous plume was imaged with a set of 15- and 20-cm focal length lenses onto a 0.35 m Czerny-Turner monochromator (Model 270, McPherson) behind which a photomultiplier tube (PMT, R955, Hamamatsu) was mounted. The LIBS emission was collected in perpendicular direction relative to the impinging laser beam. The grating in a monochromator, with a reciprocal linear dispersion of 2 nm/mm, was centered at 589 nm for the detection of the Na emission in the 3p2P®3s2S transition. Since the shape of Taylor cone in the electrospray ionization process was easily disturbed under ambient conditions, the irradiated spot size varied from shot to shot. To exempt from influence of such plume instability, the slits were open to 500-μm wide, allowing transmission of the total plasma plume, which was magnified by a factor of 1.3 as imaged onto the entrance slit. Meanwhile, the slit width was narrow enough to substantially reduce the transmitted continuum background emission at the wavelength 589 nm.

The obtained time-resolved LIB emission was fed into either a transient digitizer (Model 9450A, LeCroy) for single-shot profile recording or a boxcar integrator (Model SR250, Stanford Research System) for signal processing. The time-resolved spectra of the Na analyte
were integrated within a gate on a single-shot basis. In this work, the gate was opened in 900 ns delay relative to the onset of the continuum background emission. The LIB emission could then be the least interfered with by the intense continuum background. The gate width was optimized to 12 \( \mu \)s. The adjustment of the gate width and position may help suppress the LIB signal fluctuation to improve the signal-to-noise ratio.

The current was collected from the cathode via a high-pass filter into a second channel of either a transient digitizer or a boxcar integrator. It was monitored concurrently with the LIB emission. The current profile oscillated rapidly in a nanosecond scale, as displayed in the previous work [22]. Its first maximum peak was then integrated within a 20 ns gate throughout all the experiments. In fact, an appropriate selection of RC time constant may slow down the oscillation and extend the life time of each peak. To acquire a better LIB emission/current (LIB/C) correlation linearity, it must rely on an appropriate adjustment of the current gate width and position.

### 2.4. Reagents

The Na element as the analyte was made from its chloride solution. A 1000 mg/l NaCl (Reagent grade, Merck) solution was first prepared with water/methanol \((v/v = 1:1)\) (HPLC grade, Baker) as solvent and then diluted to the desired concentrations. For gaining a better electrospray efficiency, we used a water/methanol \((v/v = 1:1)\) mixed solvent throughout this work. To study the matrix effect, the KCl, KNO\(_2\), and KH\(_2\)PO\(_4\) (Reagent grade, Merck), prepared at concentration from 100 to 2000 mg/l, were added individually to the NaCl solution as the matrix salt.

### 3. Results and discussion

#### 3.1. Matrix effect on LIB emission/current correlation

Three diatomic and polyatomic salts, KCl, KNO\(_2\), and KH\(_2\)PO\(_4\), were added individually as the matrix to the NaCl solution, and the generated microdroplets were irradiated by a laser pulse at 355 nm. The resultant LIB emission signal of Na and the induced current were acquired concurrently up to 200 shots. Fig. 1 shows an example of single-shot time-dependent LIB emission spectrum of Na detected from the sample microdroplets. The spectrum is then integrated within a 12 \( \mu \)s gate in 900 ns delay relative to the onset of the continuum background emission. The LIB emission intensity thus obtained can be linearly correlated with the plasma-related current intensity. The correlation linearity relies on adjustment of the current gate width and position, as mentioned in Section 2. Fig. 2 shows comparison for the linearity acquired under different current gate conditions. The slope in the LIB/C correlation plot increases with the concentration of the analyte solution, as shown in Fig. 3. A possible interpretation for the sources leading to the LIB signal fluctuation is referred to our previous work [22]. Given the NaCl solution with Na concentration fixed at 200 mg/l, Fig. 4 shows the slopes of LIB/C correlation plots as a function of 100, 500, and 1000 mg/l KCl, KNO\(_2\), and KH\(_2\)PO\(_4\) solutions added individually as matrix. The LIB/C slopes in the figure remain almost constant without influence of the matrix added. The figure also shows the results of blank solutions, which contain the same matrix solutions but without the sample added. Under otherwise identical

![Fig. 1. Single-shot time dependent LIB spectra for a 200 mg/l Na solution. The spectrum within a gate, set at 12 \( \mu \)s in width and delayed by 900 ns relative to the onset of continuum background emission, is integrated to obtain the intensity of LIB emission.](image1)

![Fig. 2. Comparison of the correlation plots of Na LIB emission intensity vs. corresponding current intensity acquired with different gate conditions. (a) The first maximum peak of the current profile is gated optimally within a 20 ns duration; (b) The gate width remains the same, but the position is slightly shifted by <5 ns away from the onset of the current profile.](image2)
conditions, the blank solutions give rise to a non-zero slope, probably caused by the contribution of the residual continuum background emission.

The calibration curve is represented by the plot of the obtained slopes vs. the corresponding sample concentrations. The subsequent calibration curves, given in Fig. 5, exhibit insignificant difference between the cases without and with the matrix added up to 2000 mg/l. In the LIB/C correlation plots, the obtained regression coefficients for the slopes are larger than 0.95. The error bar represented in the calibration curve denotes the standard deviation of the slope in the correlation plot. Note that the LIB/C slopes are in arbitrary units, but Figs. 4 and 5 are not in the same scale, since both experiments were carried out in different days. The gate conditions have to be re-adjusted to achieve better correlation linearity in the LIB/C plots. Given the slope of the calibration curve and the standard deviation obtained from the slope in correlation plot of the blank solution, the LOD can be determined to be 1.0 mg/l at the laser energy of 23 ± 1 mJ. The resulting LOD of Na sample can be improved with increasing the laser energy [22].

3.2. Interpretation of exemption from matrix interference

In this work, the added matrices, each with the same K⁺ cation but different anions, seem not to cause any marked interference with the LIB/C correlated signals of the analyte solution alone. Such findings of resistance of the matrix interference may be reasonably interpreted in the following. When the pulse energy is large enough to ionize the droplets, the current collected is substantially from the electrons, which move more rapidly than the ions and are in a much shorter distance toward one end of the collector [23,24]. The electrons result mainly from the ionized water/methanol droplets and their fragments, while little from the contained sample salt. Such polyatomic molecules used as the solvent may lead to the electron multiplication as fragmented and ionized [22]. Like the sample salt, the electron contributions from the added matrix may be negligible, as compared to those from the water/methanol droplets. This consequence is demonstrated in Fig. 6, showing that 200 shots of current signals, each acquired from either an analyte plus matrix solution or a water/methanol solvent, are distributed randomly in the same intensity range. The fact is also reflected in the coordinate of current signal in Fig. 3, in which the current
4. Conclusion

We have inspected the matrix influence on the LIBS analysis of the Na sample. The microdroplets of the mixed solution were generated via an electrospray ionization needle. In spite of a smaller amount of sample ablated, the resultant LIB emission signal is well correlated linearly with the corresponding plasma-induced current. In this manner, the obtained calibration curve yields the Na LOD value approximately 1 mg/l, as comparable as those reported previously. In the LIBS analysis, the electron contributions generated by the matrix salts are negligible in an electron-rich atmosphere. The matrix effect studied in this work shows insignificant influence on the accuracy of the elemental analysis by LIBS with the current normalization. The LIB/C normalized signal might have probably taken into account both the ablated weight of the sample and the plasma excitation temperature. This method is capable of reducing signal fluctuation, resisting matrix interference, and achieving a lower LOD.

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