Laser-Induced Breakdown Spectroscopy Applied on Liquid Films: Effects of the Sample Thickness and the Laser Energy on the Signal Intensity and Stability

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Droplets of organic liquids on aluminum substrate were probed by an Nd:YAG laser, both in a steady state and during rotation at speeds 18–150 rpm. Rotation transforms the droplet into film, which estimated thickness at high speeds was below 3 μm and 20 μm for diesel and peanut oil, respectively. Line intensities from the liquid (C I) and the support (Al I) material were tracked as a function of the film thickness and the laser energy. By film thinning, the line intensities from liquid sample were enhanced up to a factor 100x; simultaneously, the LIBS signal fluctuations were reduced 5–10 times with respect to the steady droplet. In certain experimental conditions, the line intensities from the support material become very weak with respect to the C I line, indicating an efficient screening of the substrate by highly excited plasma from the liquid layer. At a fixed rotation speed, there is a laser energy threshold, dependent on the liquid thickness, above which the LIBS signal becomes stable. Here, we discuss the relative processes and optimization of the experimental conditions for the LIBS measurements from one laser shot to another.

1. Introduction

Laser-induced breakdown spectroscopy (LIBS) provides information about elemental composition of the sample volume involved in the plasma [1–3]. LIBS could be applied for analysis of materials in solid, liquid, and gaseous state, as well as of aerosols [4–6] and of particles suspended in liquid media [7, 8].

Rapid chemical analysis of liquids by LIBS [9] has many potential applications; among them, there are monitoring of waters [10–13], of quality in pharmaceutical or food industry [14–16], medical diagnostics [17, 18], and control of industrial processes or mechanical equipment [19, 20]. Laser-induced plasma on or inside liquids generates intense shockwaves, bubbling, and expulsion of droplets, which together with the laser energy losses due to liquid vaporization reduce the available energy for the plasma excitation. LIBS measurements on liquids usually have low analytical sensitivity, which could be improved by proper experimental approach and sample preparation, as reviewed in [9, 21] and updated in [22].

Excluding some complex methods for preparation of liquid samples that eliminate advantages of LIBS in terms of simplicity, rapidity, and possibility for in-field measurements, on evaporating liquids (like waters and alcohols), an intense LIBS signal could be obtained by probing the residues left on a solid substrate [13, 14, 18, 23]. The sample drying should be performed close to room temperature in order to prevent losses of volatile components. This process is time-consuming, it leaves not uniform residue distribution, and it could not be applied on liquids like oils.

Recently, another simple approach for LIBS measurements of liquids has been reported and it regards the signal generation on a thin liquid film placed over an absorbing substrate [22, 24]. Here, the breakdown initiates on the substrate and the plasma atomizes and ionizes the liquid layer. In this way, the laser-induced splashes, the plasma formation threshold, and the matrix effect (very strong in case of a direct plasma formation on liquids [9]) are importantly reduced. The LIBS signal strongly gains in intensity and so in analytical sensitivity.
Xiu et al. [24] prepared a thin oil film of arbitrary thickness on aluminium substrate by spreading manually the liquid. The plasma distribution in terms of intensity, temperature, and electron density was studied comparatively on dry aluminium and the same was covered with the oil film. It was noticed that on bare aluminium, the plasma remains close to the target surface while in presence of a thin liquid film, the maximum plasma emissivity is shifted away from the surface for a 1.5–2 mm. Vapor confinement by the initial liquid layer, later transformed to the plume, produces the plasma temperature and the electron density higher than in the case of the uncovered substrate.

Well controlled and reproducible oil film thickness could be obtained by rotating substrate at adjustable speed or by electrowetting on dielectric with controlled voltage in case of water solution [22]; the latter is impossible to thin by rotation due to high cohesive forces. This is the only work that reports changes of the LIBS signal with thickness of a liquid film. It was found that progressive thinning of a liquid film first eliminates the laser-induced splashes than the aerosol expulsion, and up to a certain thickness, it increases the emission lines from elements present in the liquid. However, the systematic studies of influence of liquid film height over substrate on the LIBS signal have not been yet performed although such sampling method, simple and applicable also for small sample volumes, could lead to a huge increase in the detection sensitivity [22].

For comparison, when analyzing water solutions in form of thin liquid sheet in air, it was found that the LIBS line intensities increase up to the sheet thickness of 20 μm [25], and this was explained by the largest liquid-laser interaction volume. For major sheet thicknesses, the LIBS signal decreases due to larger energy losses into mechanical effects and liquid evaporation. For a fixed liquid jet thickness, there is an optimal laser energy for the LIBS signal strength and the further energy growth reduces both the line intensities and their shot-to-shot stabilities [26].

On the other hand, it is well known that the efficiency of wet laser ablation (LA), particularly important for medicine and nanoparticle production, depends on a liquid type and its height above the solid target. In [27], by using KrF laser, the highest ablation rate of silica covered with water was obtained for the liquid layer of 1.1 mm, corresponding to the irradiance 2–21 GW/cm². The plasma emission was collected at angle of about 60° from the target plane by two quartz lenses (focal lengths 100 mm and 70 mm) and brought to a spectrometer (Jobin-Yvon 550), used here with grating 1200 gr/mm, by a fiber bundle arranged at the exit into 0.1 mm × 12.2 mm array. At the spectrometer’s output, an ICCD (Andor iStar DH734) was used for the LIBS signal detection. The acquisition gate and delay from the laser pulse were adjusted by a delay generator (Quantum Composer 9600+) to values of 2 μs and 1.2 μs, respectively. The measurements were performed at the laser repetition rate limited to 2 Hz.

Images of the plasma and of the liquid droplet/film were taken by a commercial photographic camera. The contact area between the liquid and the support material was determined from dimensionally calibrated photographs with help of ImageJ software (free source). The average liquid thickness on the substrate was calculated from the measured area occupied by the droplet and the known, deposited sample volume.

2. Experimental

2.1. Instruments and Layout. Figure 1 shows the experimental layout, where the plasma was generated by an Nd:YAG laser (Quantel, CER Ultra) emitting 6.5 ns long pulses at 1064 nm. The laser beam, without or after passing a beam splitter that transmits about 34% of the incident energy, was focused perpendicularly to the sample by means of a plano-convex quartz lens with focal length f = 100 mm. The spot diameter on the sample, placed slightly above the focal plane, was of about 0.42 mm, and it was determined on silica wafer. The LIBS measurements were performed with the incident laser energy between 18 mJ and 190 mJ, corresponding to the irradiance 2–21 GW/cm².

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2.2. Samples and Preparation. Liquids used in the experiment are a commercial diesel fuel and a refined edible peanut oil. These two liquids were chosen due to very different kinematic viscosities (see Table 1), which produce large differences in the liquid film thicknesses on the substrate.

As a substrate, we used an aluminum disk with diameter of 25 mm, matching the spin coater. If not specified differently in the text, aluminium surface was polished by a sandpaper 400 grits/mm but for some measurements, we also tested the surface polished with 800 grits/mm. Before placing a liquid
droplet on the substrate, the same was washed in a water flux, dried, and then washed by a high-purity methanol.

Liquid sample placed on the substrate had precisely controlled volume (±1%), delivered by an autoclavable pipette (Labgene Scientific). A droplet placed at the disc center was spread into a film by rotating the substrate on a spin coater (Laurrel Technologies, KL-SCI-20), which speed can be adjusted between 18 and 150 rpm. After turning on the spin coater and before starting the LIBS measurements or stopping the rotation to photograph the film surface, we waited at least 30 s to stabilize first the rotation speed and successively the liquid film thickness.

Although the LIBS measurements could be performed on a single droplet of a few microliters, here, we used larger sample volumes, sufficient to cover completely the aluminium disk. At low rotation speed (<25 rpm), the excess of the liquid accumulates at the support edges; otherwise, it escapes from the surface. The sampling was performed during the rotation, by registering separately the spectra produced by 10 laser pulses. Between the two LIBS measurements, the whole holder was shifted horizontally for 2 mm in order to avoid development of the craters on support material, and another holder was shifted horizontally for 2 mm in order to avoid pulses. Between the two LIBS measurements, the whole disk was in order of a few μm below which the laser energy of 23 mJ and 70 mJ. By using the lower pulse energy, the line intensities from both C I and Al I steadily grow in range 25–90 rpm, and then the saturation occurs. At this point, the liquid film is probably fully evaporated locally and efficiently atomized by the laser pulse; thus, its further thinning does not contribute to the signal intensity. RSD of the characteristic lines rapidly decays with disappearance of the splashes, that is, passing from a thick droplet to a liquid film. However, RSD of C I line keeps at high values, around 0.5, up to the speed of 90 rpm where the LIBS signal saturates. For the faster rotation, the RSD oscillates around value of 0.2. At the higher laser energy (70 mJ), the LIBS signal growth is much slower than in the previous case, and C I line intensity becomes almost saturated starting from about 50 rpm. For the same speed, the RSD of the C I line drops down and then oscillates around value of 0.1. This value of RSD was obtained also for Al I lines from the rotating support uncovered by a liquid, and we hypothesize that the vibrations present during rotation keep the LIBS signal fluctuations at this limit. From the obtained results here, we might conclude that there exists a threshold thickness of liquid film below which the laser pulse produces a full local sample evaporation and a more stable LIBS signal. This threshold thickness and the LIBS signal stability relative to the liquid increase with the laser energy. From Figures 3(a) and 3(b), we might note that the ratio of C I and Al I line intensities is higher at higher pulse energy. From Figures 3(a) and 3(b), we might note that the ratio of C I and Al I line intensities is higher at higher pulse energy, and this will be discussed later.

In analyzing behavior of the Al I at 256.8 nm, some cautions must be taken as this line is resonant, although much less intense than the nearby line at 257.5 nm, and a self-absorption might occur. The experimentally measured ratio of these two lines for the data shown in Figure 3 was in range 0.58–0.59 while their predicted ratio from the NIST database should be about 0.36 according to the formula valid for the plasma in local thermal equilibrium:

\[
\frac{I_{k_i}}{I_{k_j}} = a \cdot \frac{\bar{g}_k A_{k_i} e^{-(E_h/kT)}}{U(T)},
\]

where \(a\) is constant depending on experimental conditions; \(\bar{g}_k\) and \(A_{k_i}\) are the level degeneracy and the tabulated transition probability, respectively; \(E_h\) is energy of the upper...
transition level, $U(T)$ is the temperature dependent partition function of the considered species with density $N$ in the plasma. The low measured ratio of the two Al I (doublet) means that the more intense transition (257.5) is self-absorbed. We do not know if and how much the less intense line is self-absorbed but the stable line intensity ratio from the doublet in different experimental conditions indicates that behavior of the chosen Al I line (256.8 nm) in Figure 3 is not caused by the line saturation.

We performed the analog measurements on peanut oil, which a droplet leaves much thicker liquid film with respect to diesel (Table 1), by applying the laser energy of 70 mJ. Also in this case, the intensity of C I line becomes more stable close to the rotation speed of 50 rpm (Figure 4) but the corresponding RSD is about three times higher than in the case of diesel (see Figure 3(d)). For the steady droplet, the RSD is much higher than the same one measured on diesel, and it decreases below 1.0 only at speed of about 35 rpm.

In the following, we attempted to estimate the average liquid film thickness as a function of the rotation speed. To this aim, we placed on the substrate a droplet with well-controlled volume $V$ ($\mu$L) and then measured the area $A$ (mm$^2$) occupied by the droplet/liquid film after performing dimensional image calibration. The average liquid thickness $d$ ($\mu$m) was then calculated as $d = \frac{V(\mu L)}{A}$ (mm$^2$). In absence of rotation, the error in estimating $A$ is low as the liquid is clearly encircled also when photographing the sample perpendicularly to the support (Figure 5(a)). If the liquid reaches borders of the support (Figure 5(b)), the estimated average film thickness could be higher than the real one as the losses of liquid volume

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**Table 1: Physical properties and the estimated droplet/film thicknesses for diesel and peanut oil.**

<table>
<thead>
<tr>
<th></th>
<th>Diesel</th>
<th>Peanut oil</th>
</tr>
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<tbody>
<tr>
<td>Surface tension at 20°C (mN/m)</td>
<td>25.8 [33]</td>
<td>10.0 [34]</td>
</tr>
<tr>
<td>Kinematic viscosity (mm$^2$/s) at 40°C</td>
<td>2–4$^*$</td>
<td>37 [35]</td>
</tr>
<tr>
<td>Average thickness of 20 $\mu$L droplet ($\mu$m)</td>
<td>141 ± 10</td>
<td>320 ± 20</td>
</tr>
<tr>
<td>Av. thickness of droplet 5 $\mu$L at 18 rpm ($\mu$m)</td>
<td>19 ± 4</td>
<td>102 ± 10</td>
</tr>
<tr>
<td>Av. thickness of liquid film at 100 rpm ($\mu$m)**</td>
<td>≤3.0</td>
<td>≤20</td>
</tr>
</tbody>
</table>

* means specified by the supplier. ** means measured for droplet volumes of 0.2 $\mu$L and 5 $\mu$L for diesel and oil, respectively.

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**Figure 2:** LIBS spectra from diesel, averaged over 10 laser pulses with energy of 23 mJ. Steady droplet (a) and support rotated at 18 rpm (b) and 100 rpm (c).
from the substrate might occur. By applying high rotation speeds, the shape of area occupied by liquid becomes irregular, introducing larger errors in its circle, and it was necessary to take the photographs angularly—this also increases the error due to the perspective (Figure 5(c)).

Initially, we attempted to determine the film thickness of 20 μL oil droplet but the liquid reaches the borders of the support already at the rotation speed of 18 rpm (Figure 5(b)). Further increase of the rotation speed causes losses of the liquid, visibly sputtered away from the support, and to a consequent overestimation of the average film thickness, based on the known deposited liquid volume. For this reason, the measurements of the oil film thickness were performed also for the droplet of 5 μL, on the substrate polished with sandpaper of 400 grits/mm or 800 grits/mm (Figure 6). For the substrate polishing by 400 grits/mm, at the rotation speed of 18 rpm the droplet of 20 μL has the average thickness similar to that of the smaller droplet (5 μL). However, by the applied method here, the estimated film thickness at larger rotation speed differs significantly in the two cases due to the liquid volume loss that affects calculations of the average

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**Figure 3:** Average peak intensity of C I and Al I lines from the diesel droplet, measured over 10 pulses, as a function of the rotation speed for laser energies of 23 mJ (a) and 70 mJ (b); the corresponding RSD is shown in (c) and (d), respectively.

**Figure 4:** RSD of C I and Al I lines from 20 μL droplet of peanut oil as a function of the rotation speed; the laser energy is 70 mJ.
film thickness. On the substrate polished by the finer sandpaper (800 grits/mm), the average liquid height is importantly lower than for the rougher polishing except at high rotation speeds, where the estimated thicknesses converge to about 20 μm. Here, the film height also tends to a constant value at lower rotation speed compared to the rougher substrate, where the exponential decay coefficients are 15.5 rpm\(^{-1}\) and 18 rpm\(^{-1}\), respectively. Comparing the Figures 4 and 6, we might conclude that on peanut oil, the LIBS signal excited by 70 mJ pulses becomes stable for the thickness below 50 μm.

A steady diesel droplet with volume of 20 μL has the thickness about twice lower than the oil droplet, and this explains the much lower RSD of the C I line (Figure 3(d)). Due to a very low kinematic viscosity, in rotation at the minimum speed (18 rpm), the large diesel droplet already spreads over the whole substrate and partially leaves it, thus compromising the calculations of the average area occupied by the liquid and so, of the film thickness. For the droplet of 5 μL, we estimated the thickness of 19 μm at rotation speed of 18 rpm. To measure the area occupied by diesel droplet at high rotation speeds, it was necessary to reduce its volume to 0.2 μL only; otherwise, the liquid volume was partially lost from the support disc. In this case, the estimated liquid thickness reaches the stable value of about 3 μm at speed of only 35 rpm. Rapid thinning of diesel droplet with the rotation speed explains a drop of the RSD relative to C I line below value of 0.5 already at 18 rpm; on the oil, this value of RSD was achieved at the speeds above 40 rpm. Stability of the C I line intensity from one laser shot to another, much better on diesel than on oil (Figures 3(c) and 3(d) and Figure 4), could be attributed to large differences in the liquid film thickness.

3.2. Influence of Laser Energy on the LIBS Signal. Influence of pulse energy on the LIBS signal was studied at constant rotation speed of 40 rpm, on both diesel and peanut oil; for the latter, we used the support polished with the abrasive papers with 400 or 800 grits/mm. At the chosen rotation speed, the estimated thickness of diesel film was of about 3 μm while peanut oil film was thick of about 40 μm and 27 μm for the polishing by 400 and 800 grits/mm, respectively. For the rougher polishing and the oil sample, by increasing laser energy, the C I line intensity starts to grow from about 50 mJ. At the pulse energy of 190 mJ, the C I peak is about 60 times higher than at the minimum energy used here (Figure 7(a)). Simultaneously, the growth of the Al I emission intensity is much slower compared to the C I line, so the line intensity ratio I(Al)/I(C) rapidly drops from 2.0, reaching values in range 0.1−0.2 only (Figure 7(b)). This means that starting from a certain laser energy, here around 30 mJ, the plasma produced on the liquid film effectively screens the support material from ablation and further energy increase mainly contributes to excitation of the species originating from the liquid. This fact is very important for reducing influence of the substrate’s constituents on the spectra from the analyzed liquid sample. Figure 8 shows the comparative spectra from peanut oil obtained at two very different pulse energies.
On peanut oil, starting from the pulse energy of about 40 mJ, the RSD of C I line steadily decreases from value above 1.1 or 1.3 to 0.2 (Figure 9(a)). On the rougher substrate, the signal stability is initially worse than for the finer polishing, and the stable RSD is reached at higher pulse energy (125 mJ compared to 110 mJ). Further increase of the laser energy corresponds to a stable RSD of C I peak in the spectrum except at the highest energy applied here (190 mJ), where some other instabilities seem to occur. At the same time, the RSD of Al I peak fluctuates at high values, between 0.7 and 1.6 (not shown). On diesel film, about 10 times thinner than the oil, the intensity of C I line is much more stable from one shot to another, and an approximately constant RSD = 0.1 is reached at the laser energy of about 80 mJ. From these results, it is clear that there is a laser energy threshold $E_{st}$ above which the LIBS signal from liquid film becomes stable. Basing on the previously estimated liquid film thickness and the data shown in Figure 9(a), we found a linear relation between the film thickness and the energy threshold $E_{st}$ (Figure 9(b)). We hypothesize that above the threshold energy $E_{st}$, the liquid layer locally detaches from the surface and becomes fully evaporated, resulting in a stable emission intensity from one laser shot to another and independently on a strongly variable support ablation. Further increase of the laser energy mainly contributes to the excitation of the atomized species from the liquid, without affecting importantly the support ablation (see Figure 7).

### 3.3. Optimization of the Experimental Conditions

Previously, we showed that the LIBS signal from an analyzed organic liquid improves in intensity and stability by reducing the liquid film thickness, achieved here by the substrate rotation. Thinning of the liquid film on substrate also reduces the laser energy requirements for obtaining an intense stable signal, and this is very important if a laser source with limited performances is used.

Unfortunately, increase of the rotation speed above a certain value does not bring significant improvements of the LIBS signal as the minimum achievable film thickness depends on kinematic viscosity of liquid, which is a measure of resistance to flow and shear. Kinematic viscosity $\eta$ of oils decreases exponentially with temperature, and it is described by the Arrhenius equation:

$$\eta = A_1 e^{E_a / RT},$$

where $A_1$ and $E_a$ are constants specific for the oil, $R$ is the universal gas constant, and $T$ is the absolute temperature.

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**Figure 7:** (a) Peak emission of C I and Al I lines averaged over 10 pulses as a function of the laser energy. (b) The corresponding line intensity ratio. The sample is peanut oil rotated at speed of 40 rpm.

**Figure 8:** LIBS spectra from peanut oil on aluminum substrate (polishing 400 grits/mm) averaged over 10 laser pulses with energies of 23 mJ (top) and 137 mJ (bottom); the rotation speed is 40 rpm.
Table 2 shows kinematic viscosity at different temperatures for diesel and sunflower oil, published in [36], and the known or extrapolated values by (2) for refined peanut oil [35]. It is evident that close to room conditions, the kinematic viscosity of organic liquids rapidly reduces with raising the temperature. From the practical point of view, this means that a more efficient thinning of a liquid film could be achieved by keeping the sample above room temperature, for example, at 30°C. Anyway, in order to have well-repeatable LIBS measurements, it would be opportune to keep the liquid sample at a stable temperature or to adjust the rotation speed in a way to maintain the wanted liquid film thickness.

If the rotation speed of the motor employed is limited to lower values than here (150 rpm), the film thickness at a certain and not high rotation speed could be further reduced by finer polishing of the substrate. However, due to a simultaneous liquid atomization and the substrate ablation, for detecting the trace elements in liquids, it is necessary to use a support material free of the same, for example, silica wafer [22]. In the case of a metallic substrate, its polishing should not transfer impurities to the material. For example, we compared the spectra from the bare aluminium substrate (not covered by a liquid) after polishing it by a sandpaper or by Al₂O₃ particles. In the first case, by sweeping the monochromator over different wavelength ranges, we detected Cr, Cu, and Zr lines in the spectra. These lines were not present in the LIBS spectra taken on the same substrate polished by Al₂O₃ particles, and this clearly indicates a transfer of impurities from the sandpaper grains to the substrate surface.

4. Conclusions

A droplet of a viscous liquid could be efficiently reduced to a thin film by rotating the support on which it is placed. In this way, the average liquid’s height was reduced here from about 140 μm to 3 μm and from about 320 μm to 20 μm for droplets of diesel and peanut oil, respectively. The minimum film thickness that could be achieved at high rotation speeds is roughly proportional to the kinematic viscosity of liquid, much lower for diesel than for peanut oil.

Thinning of a liquid droplet/film eliminates unwanted laser-induced splashes and spraying during LIBS measurements, thus minimizing losses of the sample volume and a necessity to clean the nearby optical elements. Increase of the support rotation speed enhances the intensities of the spectral lines from liquid up to a factor 100 until the saturation is reached, corresponding to an almost constant liquid film thickness. The saturation speed lowers with the applied laser pulse energy.

Besides an impressive increase in the LIBS signal with thinning of a liquid film, shot-to-shot fluctuations of the line intensities originated from the liquid are reduced many
times. A better signal stability corresponds to a thinner liquid film and to a higher pulse energy.

Raising the laser energy above a certain threshold produces a full local atomization/ionization of the liquid layer and reduces abruptly the LIBS signal fluctuations. Simultaneously, intensities of the lines originating from liquid are largely enhanced while those generated by the support ablation remain relatively weak, and this reduces the substrate interference in the spectra from the liquid sample. The threshold laser energy increases with the liquid thickness, which could be reduced also by a finer substrate polishing.

The proposed simple method for liquid analysis, together with the presented parameter optimizations here (energy, film thickness), allows to obtain a very intense and a relatively stable LIBS signal from small liquid volumes. Further studies should regard the analytical sensibility of the method and the effects of substrate material on the plasma formation.

Conflicts of Interest
The authors declare that there is no conflict of interest regarding the publication of this paper.

References


