

**In-situ monitoring and characterization
of airborne solid particles in hostile industrial environments
using stand-off LIBS**

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Abstract.

The analytical possibilities of laser induced breakdown spectroscopy (LIBS) to carry out in-situ and real-time detection and compositional characterization of solid aerosols in the atmosphere of a steelmaking factory, have been evaluated. To this aim, a compact and versatile dual-pulse LIBS analyzer, able to sample at distances up to eight meters, has been design to work under these hostile industrial environments. Due to the discrete nature of the particulate matter, the particle sampling rate was less than 2.5% and 6% for single- and double-pulse regimes, respectively. An efficient statistical procedure, based on the calculation of standard deviations, is used to qualitatively characterize the elemental composition of the aerosol. Then, a conditional analysis based on the limit of detection, is employed to assess the elemental sampling rate. This experimental methodology has been used to evaluate the influence on the aerosol formed of the oxycutting process in a continuous casting machine producing steel slabs, revealing a strong presence of elements derived from the casting powder used in the production. Moreover, chromium, present in the steel cast, is detected in suspension in the steel shop. An increase in the concentration of particulate matter when oxycutting is working is suggested. Single-pulse and dual-pulse excitation modes are also evaluated.

Keywords.

Laser-induced breakdown spectroscopy, stand-off analysis, solid aerosol detection, steelshop atmosphere, continuous casting line, sampling rate.

1. Introduction

Air pollution is an old and widespread problem, as well as a today issue. Originated from both natural and anthropogenic sources, contaminants are directly introduced in the atmosphere as gases and particulate matter. In addition, these particles can also come from photochemical processes induced by solar radiation. These airborne particles or atmospheric aerosols are ubiquitous in the environment, and some of them may be hazardous for the environment and health, and affect the economy. Especially adverse effects on human health are due to heavy metal particles that may accumulate in human beings via inhalation and respiratory deposition [1]. So, the analysis of aerosols in ambient air has been a concern of the scientific community in recent years because of those potential adverse effects on health and visibility, and because they play a role in the regulation of climate processes [2-5]. In this context, the acquisition of analytical information of particulate matter present in a specific atmosphere is of a crucial interest.

One of the most important anthropogenic sources of emissions in today's societies is the steelmaking industry. Main processes may include the use of coke as fuel and as reducing agent in smelting iron ore in a blast furnace [6]. Coke ovens are the major source of polyaromatic hydrocarbons (PAHs) emission in the iron and steel facilities [7]. Furthermore, throughout the steelmaking process, the cutting of the produced steel bars, namely slabs, blooms and billets, is performed after the casting in an oxy-fuel unit. In this stage, a torch is used to heat metal to its kindling temperature. Then, a stream of oxygen is guided to the metal, burning it into a metal oxide that flows out of the kerf as slag. Both the

combustion of fuel and the produced metal are involved through a highly energetic reaction, with the inherent generation of particles and combustion gases. Most studies carried out in these industrial environments have focused in the assessment of physical and chemical characteristics of particulate aerosols from important dust sources identified in integrated iron and steel processes, more specifically on sintering, blast furnace, steelmaking and desulfurization slag processing. Fe, Ca, Al, Na, S and Mg have been the major elements usually found [8]. Moreover, the composition of the suspended particulate matter measured around a steel plant is compared with that of the urban environment [1].

Such chemical diagnoses and the specific analytical techniques used to this aim are becoming more and more relevant. The current methodology consists of the use of particle filtration samplers, combining different size-selective inlets, filter media, and flow movers/controllers. However, this approach is time-consuming and uses large sampling air volumes [9,10]. Also, as in many common techniques, treatment processes are required for analysis [11].

Surveillance of solid aerosols demands the rapid, remote and *in situ* evaluation of chemical composition and concentration of particulate matter in air. Relevant analytical challenges emerge because of the significant heterogeneity exhibited by aerosol properties (i.e. particle density, size, morphology and composition) [12]. Thereby, the use of a compact, robust, and versatile analytical system, as well as the capability of working *in situ* and in real time, should be highly desirable. For years, LIBS has been used to obtain information that allows a better understanding of the dynamics of these polluting agents in the form of solid

aerosols. In this context, several research works involving LIBS detection of solid aerosols in ambient air as well as in waste stream effluents for real time monitoring have been reported [13-17]. Thus, LIBS application for aerosols analysis offers many advantages that overcome some of the inherent limitations that the current technologies used for this purpose present. In the last years, this technique has been developed as a novel system for the direct quantitative measurement of particle size [18], extent of oxidation or coating thickness of nanoparticles [19], and separate particulate composition [20]. However, many fundamental issues related to the laser induced plasma interactions with the aerosol particles are still outstanding [12].

Stand-off LIBS approach allows the analysis of physically inaccessible targets or samples which may be located in hazardous environments. Generally, stand-off LIBS systems use nano- or femtosecond pulse lasers for the analysis of both solid and liquid samples located at several meters from the instrument. This approach has been used in numerous and different applications (environmental monitoring, nuclear and metallurgical industries, planetary exploration, cultural heritage, detection and identification of explosives...) [21-24]. Moreover, the possibility to monitor at a safe distance for those cases in which the samples present a risk for both operators and equipment is a clear advantage of this technology [25].

This study addresses to demonstrate the stand-off LIBS capability for in-situ analysis of variable-sized heterogeneous solid aerosols at Brownian motion, generated in a real industrial scenario. To the knowledge of the authors, it is the first time that a LIBS system

is deployed in a steel factory to detect particulate matter in suspension. The work focuses to the detection of solid aerosols as well as the determination of their major elemental composition. In order to gather preliminary information, measurements were performed by remote and direct sampling of the environment inside the steel making factory, where diverse metallic elements were expected to be detected. Furthermore, in order to assess the efficiency of the methodology, the intensity variability and the sampling rates were estimated.

2. Experimental

2.1 Stand-off LIBS system

A versatile and compact LIBS portable system was built for industrial tests capable to work in stand-off mode with non-invasive technology. The prototype is shown in Figure S1a (see Supplementary Material) and has been previously described [26,27].

Two 1064-nm Q-switched Nd:YAG lasers with 5 ns pulse width, and combined in an axial configuration, were used. The combined maximum energy available was 900 mJ at a maximum repetition rate of 10 Hz. The two laser beams were properly synchronized by means of their own internal electronic device which triggered them with a controlled delay of 2 μ s, and then guided, expanded by means of a Galilean telescope, and focused in order to obtain the level of irradiance required to produce aerosol optical breakdown at typical

distances of 5 m. Selected optical elements allowed an adequate guiding and focusing of the laser beam and an optimal signal collection.

Thus, collection of the plasma light was carried out by a commercial Maksutov-Cassegrain telescope (f 1/15) and focused with a planoconvex quartz lens (FL 50 mm) onto the input aperture of an optical fiber (D 400 μm). This fiber is connected to the entrance slit of a dual-channel asymmetrical crossed Czerny-Turner spectrometer that detects the dispersed radiation with an integrated linear CCD array of 2048 pixels. The CCD aperture is triggered just 1 μs after the firing of the second laser pulse, and the integration time was fixed at 1.1 ms (the minimum integration width allowed). The grating installed inside each spectrometer has 1200 grooves per mm, and the slit width was 10 μm , providing a spectral resolution of 0.1 nm (FWHM) in the selected spectral region. The spectral range for the combination of both spectrometers covers from 340 to 810 nm, which allowed the detection of the spectral lines of the elements of interest. The system was protected with an enclosed box away from the hot and dusty environment in the factory. A detailed description of the system can be found elsewhere [26].

The LIBS analyzer and the auxiliary equipment were then deployed next to the slab continuous casting line, in the proximity of the oxycutting unit. Figure S1b shows a scheme of the location of the LIBS analyzer. The final distance from the LIBS system to the slabs in the continuous casting line was 7.6 m, and the laser output was tilted and pointed just behind the area where oxycutting process of slab took place.

The laser beams were focused onto a region at around 5.6 m of distance of the LIBS system, approximately at a distance of 2 m from the top surface of the slab. This ensures that the depth of focus (DOF) was separated from the slab surface at all times. The beam guiding components play a crucial role since they have to focus the laser energy in a sufficient small spot in order to provide the necessary irradiance to produce the emission plasma of airborne particles. The minimum achievable spot diameter in such analysis conditions was estimated to be 0.5 mm, taking into account the beam quality parameter ($M^2= 2.95$) and the laser beam diameter at focusing optics ($D = 40$ mm). According to these experimental conditions, the DOF of the laser beam was estimated in 125 mm.

LIBS measurements have been conducted using two laser excitation approaches, namely single- and double-pulse. In both cases, two situations were compared: experiments carried out while the oxycutting system is running or not. In addition, measurements were taken separately for the two spectral ranges (from 340 to 595 nm, and from 595 to 810 nm) to obtain a better signal to noise ratio. When double-pulse regime was employed, laser energies of 250 mJ/350 mJ for first and second pulses, respectively, were chosen with a repetition rate set to 5 Hz, whereas for single-pulse, the laser energy was set to 250 mJ. In both cases, the energies involved are enough to generate the breakdown of the sampled aerosol. A value not lower than 2.5×10^{10} W/cm² is obtained for these experimental settings when using single-laser pulse mode.

In the search of differences among both laser excitation modes, aerosol sampling rates, defined as the percentage of laser pulses expected for sampling at least one particle within

the plasma volume [12], were determined. For this estimation, a conditional analysis was performed based on the IUPAC criterion $3s$ (threshold value for the limit of detection), which usually corresponds to a confidence level of about 90% [28,29].

2.2 Standard deviation and conditional analysis methodologies

Because of the discrete, particulate nature of aerosols, the utilization of LIBS is considered in part as a statistical sampling problem involving the finite laser-induced plasma volume, as well as the concentration and size distribution of the target metal species [30]. Single particle hits originate strongly variable spectral signals for specific species. Nevertheless the high intensity found for some elemental spectral features, as a large percentage of laser shots originate particle signature free spectra (i.e. only background and lines unrelated to particles), the ensemble-averaged intensities are strongly reduced.

On the contrary, standard deviation (sd) values for the measured emission intensities become extremely high. Thus, this statistical parameter (sd metric) can be used instead of the absolute LIBS emission intensities to characterize the spectral response of the aerosol particles in low concentration, and, consequently, with a negligible loss of significant information from the analyte [31].

A clear evidence of this fact is reflected in Figure S2, where a comparison of the ensemble-averaged emission intensities and sd metric is presented. As observed, both amplitude and signal-to-noise ratio (SNR) corresponding to sd metric proved to be noticeably higher than the averaged emission intensities. All detected spectral lines appearing in the ensemble-averaged spectra (Figure S2a) are also present in the sd plot (Figure S2b), thus confirming

that sd method is appropriate for data examination of non-homogeneously dispersed particles. Nevertheless, some assigned elemental lines clearly visible in the sd plot are not present at all in the ensemble-averaged spectra, what proves the interest of this kind of plot [31]. Although the absolute spectral intensity values are lost, the use of the standard deviation method becomes a very powerful tool for further processing of spectral signals with high variability.

On the other hand, the classification of spectra into two groups, one of them with significant spectral response, and the other one with no information, can be used to assess the collected data. This evaluation allows achieving more accurate values for the sampling yield, related to particles concentration levels and aerosol composition. This conditional analysis, which increases the signal-to-noise ratio of the spectral data, has been previously described in the study of non-homogenous analyte systems [32].

In this manuscript, the calculated sd values are used to accomplish the spectral lines to be considered in the conditional analysis. The selection of an appropriate threshold for these spectral lines is the essential task to quantify a given spectrum as corresponding to a particle hit or particle miss. When the value of the emission intensity exceeds this threshold, the corresponding spectrum is classified as representative of a particle hit [33]. The criterion adopted here was that signal-to-noise ratio (SNR) value resulted greater than 3 (according to IUPAC recommendations). Additionally, spectra with SNR values greater than 10 were also taken into account for assessing variability differences between signals belonging to the different sampled volume components, namely particle matter and air

matrix constituents. For the present analysis, SNR is defined as the ratio of the atomic emission line net intensity to the average of the adjacent, featureless, continuum (background).

After hits spectra counting, sampling rate values were then computed both in a global mode (any spectral line) and for selective detection of specific lines (i.e. selective analytes).

3. Results and discussion

3.1 Spectral analysis of aerosols

The examination of the attained spectra from individual breakdown events lets to identify the relevant elements present in the aerosol particles or air susceptible of rupture; the corresponding spectral lines are summarized in Table 1. In general, the most of breakdown hits correspond to air rupture emission, thus confirming that the experimental irradiance level of the LIBS system far exceeds the threshold for clean air breakdown (around 10^{11} W/cm²). Also, the presence of aerosol particles in the focal volume will lower the threshold for gas breakdown by several orders of magnitude (typically to around 10^7 W cm⁻²) [34]. On the other hand, a spectral analysis determines that most of the sampled particles are based on both alkali (Li, Na, K) and alkali earth (Ca) compounds. Figure S3 shows a pair of typical spectra obtained for each of the active channels of the spectrometer (while the oxycutting unit is working) and using double-laser pulse excitation. As observed, the spectrum in Figure S3a is characterized by the presence of Ca II (393.37 and 396.85 nm)

peaks, and the well-known doublet Na I (589.00 and 589.59 nm). On the other hand, the strong emission features corresponding to air rupture are depicted in Figure S3b, in which O I (triplet not resolved at 777.19, 777.42 and 777.54 nm) LIBS signal governs the spectrum.

Figure 1 presents three examples of individual laser events exhibiting emission signal of further metallic species. The LIBS spectra correspond to Ti-, Al- and Cr-containing particles, respectively. Figure 1a illustrates a representative spectrum for the ablation of a Ti-containing particle, in which lines at 453.48 nm and 499.95 nm become the most intense signals for this element. Figure 1b shows a detail of a LIBS spectrum where the Al resonant lines at 394.40 nm and at 396.15 nm can be observed. Figure 1c exhibits a single-shot spectrum with the presence of several Cr emission lines. It should be noted that these elements were detected in a lower number of laser events, compared to Ca, Na and air constituents. This fact will be extensively discussed below (in the section on sampling rates). On the other hand, surprisingly, in spite of being measuring inside a steel shop, Fe lines were not identified in any of the laser event measurements.

In addition, CaF and CaO molecular bands have also been detected in some of the emission plasma events, as shown in Figure 2 [35]. Moreover, in this spectral range, the presence of lithium is revealed by its signature emission line at 670.79 nm.

3.2 Sources of the particulate matter. Steel slab and casting powder

In the steel manufacturing process, once the steel is cast, the resulting slabs/billets have to be cut. The process of cutting with an oxygen flux -or oxycutting- is the most common in

steel industry. This aggressive process is performed with an iron powder supply and an oxygen stream, releasing large amounts of energy on a previously heated metal (near the ignition point of the material). Furthermore, in order to facilitate the cutting of the stainless steel slab, iron powder is provided using compressed air or nitrogen. As the powder burns, a great deal of heat is generated resulting in the formation of a viscous slag due to the combination of combustion phenomena with the melt action of the flame. This slag consists mainly of chromium oxides that opposes the action of the oxygen flame [36]. In order to help the heat dissipation, the oxycutting process is water cooled. It is reasonable to think that this device could be one of the main sources of aerosols generation in this industrial ambient. For this reason, field trials were carried out nearby the steel oxycutting unit.

This LIBS experiment was performed during continuous casting of a ferritic steel grade (AISI 430), whose elemental composition is showed in Table 2.

In order to gain evidence on the origin of the different elemental species observed during aerosol monitoring, two different spectra were measured and compared. Figure 3a shows the real-time LIBS spectrum focusing the laser beam precisely onto the slab surface (at around 900°C), at a distance of 7 m from the laser, while the steel was being cast at a speed of ca. 1 m/min. The spectrum shown is the average of 100 laser shots at different fresh positions of the slab surface. In this spectrum, the main emission features of ferritic steel are observed: Fe spectral lines, accompanied by some lines of chromium and manganese as major alloy elements. However, strong emission peaks (mainly of Ca-containing species), due to elements not present in the chemical composition of the stainless steel, are also

detected [27]. These unexpected spectral emissions have been demonstrated to come from the scale layer formed on the top surface of the steel, contaminated with rests of the casting powder (used to ensure the reliability of the continuous casting process). These casting powders or mould fluxes are synthetic slags constituted by a complex mix of oxides, with fluorite and carbonaceous materials in their composition. The chemical content of the mould flux employed during the experimental measurements is detailed in Table 3. Figure 3b shows a LIBS spectrum of a mould powder pellet, acquired in the laboratory at a high temperature, simulating the experimental conditions of the casting line, and averaging 20 laser shots on the same spatial position of the sample. A comparison of Figure 3a and 3b proves the presence of remains of mould powder spread on the slab surface examined at the steel shop. Species as Ca, Na, Li, Al, Ti and CaO were identified in both spectra. Ti emission, for instance, is well observed despite its low concentration ($\text{TiO}_2 < 0.7\%$). However, it should be noted that, although iron is also present in the chemical composition of the mould powder in low concentrations, in the form Fe_2O_3 ($<1\%$), Fe lines are not detectable.

Elements as Ca, Na and K are present in many of the aerosol breakdown events (see Figures 1, 2 and S3), since they take part of the main oxides existing in the mould powder composition. Moreover, these atomic species are easily ionizable and their ablation and excitation thresholds are low.

The detection of Li in some of the collected spectra from aerosols could be attributed to its presence -in very low levels ($<0.5\%$)- in the mould powder composition, as it is also

observed in the LIBS spectrum of the mould powder pellet. However, its presence in the aerosol could also be associated to lithium soap. This grease, used for the lubrication of rollers and bearings of continuous casting lines, as well as the hot-rolling mill, are Li-based compounds. This lithium soap could eventually be evaporated due to the high temperatures generated in the oxycutting process. Notwithstanding, there is no evidence of organic molecular bands (CN, C₂, CH, NH, OH...) in the collected spectra that could confirm such an origin; this source may not be excluded as secondary source since the emission intensity achieved is often very significant in the aerosol, as observed in spectrum depicted in Figure 3b. The observation of CaO and CaF molecular bands, characteristic of the particulate matter spectra, is an additional fact supporting that casting powder is the main source of the measured aerosols.

Furthermore, the only metallic species detected in the aerosol analysis, but not present in the casting powder, is chromium, the main alloying element of the steel grade cast, as observed in Figure 1c. In this case, its origin can be associated unequivocally to the scale layer formed on the surface of the slab. This layer is enriched in chromium oxides due to the diffusion of Cr atoms from the bulk to the external regions of the slab, favored by high temperatures in the casting process [37,38]. The lower density of the scale layer compared to that of the steel (7.8 g/cm³) along with the weak joining to the slab surface, allows the pulverization of the oxidized layer in the oxycutting process, thus generating Cr-containing particles in the aerosol. Volatility of chromium oxides has been widely studied [39,40] as well as the oxidation of steel in different atmospheres and temperature ranges [37,41,42].

3.3 Signal variability

Once the suspended material has been spectrally identified in the steel shop atmosphere, and its most probable origins discussed, the next step to be carried out is a statistical study of the data. Due to the nature of the particulate matter, a large signal variability is expected.

The uncertainty of measurements is assessed by means of the ensemble-average of the intensities and SNR values. These values were calculated after applying a conditional analysis to the spectra, as discussed in the Experimental Section, and are summarized in Table 4. RSD of the ensemble-average values of net intensities and SNR, for the sets of spectra with SNR greater than 3 and greater than 10, respectively, were computed. These results, for both spectral lines corresponding to particle matter and air matrix constituents, were then compared. The most intense spectral lines measured for this purpose were: Ca II lines at 393.37 and 396.85 nm in the first spectral range, N I lines at 744.23 and 746.83 nm, and the O I triplet, not spectrally resolved, at 777.19, 777.42 and 777.54 nm in the second one. This statistical study has been performed using single- and double-laser pulse regimes.

As it can be observed in Table 4, uncertainties computed for SNR values (in terms of the averaged relative standard deviation -RSD-) are systematically larger than those calculated for the own spectral intensities, since the variability of the background is lower than that of the signals. When the averaged RSD values of all spectral data with $\text{SNR} > 3$ are compared with only those with $\text{SNR} > 10$, differences are found between the behavior of particles constituent element (calcium), and air major elements constituents (nitrogen and oxygen).

In the case of calcium, the RSD values calculated for spectral intensities in the ensemble $\text{SNR}>10$ are higher than the values for the subset corresponding to $\text{SNR}>3$. This fact could be attributed to a number of variables, including the presence of a larger number of sampled lower-size particles, the calcium content level, and the higher variability of the sampled major-size particles (the number of major-size particles is lower).

In contrast, in the case of elements present in air, as nitrogen or oxygen, the averaged RSD values behave in the opposite way. For the $\text{SNR}>3$ spectra ensemble, the means of RSD values are higher than the values obtained from the subset with $\text{SNR}>10$. These results confirm the heterogeneity of the particles sampled. However, as the number of LIBS measurements is not similar in all the cases, due to the random nature of the experiment, this kind of conclusions is not straightforward. Moreover, the discrete behavior of an aerosol breakdown event, in terms of particle size, composition, density of particles in the interrogated pathway (focal volume where plasma threshold is attained) and so on, is another source of uncertainty.

Additionally, it should be noted that the results acquired in both excitation modes, namely single-pulse and dual-pulse, are not comparable between them, as the energy delivered in both excitation modes differs. Generally, a comparison of experimental results obtained under different experimental conditions (laser irradiance, laser focusing and plasma light collection distances, environmental conditions, signal acquisition parameters...) is very problematic [21].

3.4 Sampling Rates

As it has been explained in the Experimental section, particles sampling rates (PSR) were obtained from the calculation of percentage of events with plasma breakdown from the total of delivered laser pulses [12]. Again, the studies were carried out (while the oxycutting unit is working) and using single and double-laser excitation.

Table 5 compares the computed values of the overall sampling rates, measured while the oxycutting process is working and while it is off, and under the two different excitation approaches: single and double-pulse. These values have been obtained in two different spectral ranges, when two different spectrometers (spectral ranges 340-595 nm, and 595-810 nm) were used. In the second spectral range, most of the breakdown events correspond apparently to only air-rupture (where main emission lines corresponding to H, N and O species are found, but not emission from rest of elements detected), obscuring the interpretation in terms of particulate matter detection. The air rupture frequency can be inferred from the sampling rate values calculated in the second spectral range, providing values between 19 and 23% of the total of laser events, regardless the operating conditions (and taking into account that species emission from discrete aerosol particles appearing in this spectral range were always concomitant to air emission).

On the other hand, results confirmed a significant impact of the oxycutting process in the averaged sampling rates, at least in the case of double-pulse excitation, and therefore, a higher concentration of discrete particles in the ambient during and immediately after the steel-cutting interval. Moreover, a higher interaction rate in the case of double-pulse operation mode compared to single-pulse regime was observed.

Finally, individual sampling rates corresponding to each detected element were computed and compared in Figure 4. For example, Cr detection exhibits a maximum value of 0.2%. Taking into account that particle sampling rates calculated by LIBS are on the order of 0.1% for many aerosol loading conditions [30,43], this value may be considered as significant and corresponding to a notable concentration. It has been demonstrated that for submicrometer sized metal particle distributions, and for effluent levels of about 10 $\mu\text{g}/\text{acm}$ (typical of many treatment process wastestreams), LIBS particle sampling rates may fall to 1% or less [30].

Several conclusions can be drawn from Figure 4. First, the number of LIBS spectra where Ca lines are detected was calculated in order to know its statistical weight with respect to the total of the breakdown events (in both spectral ranges). In the first spectral range, the amount of these spectra detected while the oxycutting is active represented the 87% of the total of the breakdown events. On the other hand, this emission percentage was reduced to 3% in the second spectral range, under the same experimental conditions, as a result of the presence of air rupture emission. If air emission signals are not taken into account, the number of events including Ca emission represents the 71% for the second spectral channel, value closer to 87% obtained for the first one. Second, the oxycutting process seems to increase the probability of the ablation mechanism, and the further excitation and emission of Ca and metallic elements (Cr, Ti, Al). However, breakdown events corresponding to emission of alkali metals (Li, Na and K) were detected in the ambient

independently of the laser excitation mode as well as the presence of the oxycutting process.

4. Conclusions

In this study, the analytical possibilities of in-situ detection of solid aerosols generated in a steelmaking factory, as well as the determination of their elemental composition, have been evaluated, using a stand-off LIBS analyzer able to work at distances up to 8 meters. Two modes of interaction between laser beam and airborne particles, namely single-pulse and double-pulse laser regimes, were assessed. Experimental results confirmed the main source forming aerosol particles: casting powder probably removed from the slab surface during the oxycutting process of stainless steel slabs. Both elemental (Al, Ti, Li) and molecular (CaO, CaF) emission signals confirmed the presence of particles of this lubricant powder in the analyzed atmosphere. An additional identified source of particulate matter, obviously, was the steel itself being cut, since Cr emission, coming either from the steel bulk or from the oxidized steel surface, was also detected in suspension. Computed values for sampling rates from the analysis of aerosols in a region next to the oxycutting unit indicated an enhancement of the concentration of airborne particles for the case to be running the cutting process of the slabs, compared to periods with the oxycutting unit off. Likewise, double-pulse laser measurements seem to slightly improve the sampling rate values compared to those obtained in single-pulse regime.

Field tests have proved the viability of stand-off LIBS technology for the in-situ and real-time detection of solid aerosols in industrial environments, thus becoming an analytical on-line tool for a rapid monitoring and control of particulate emissions generated in hostile scenarios.

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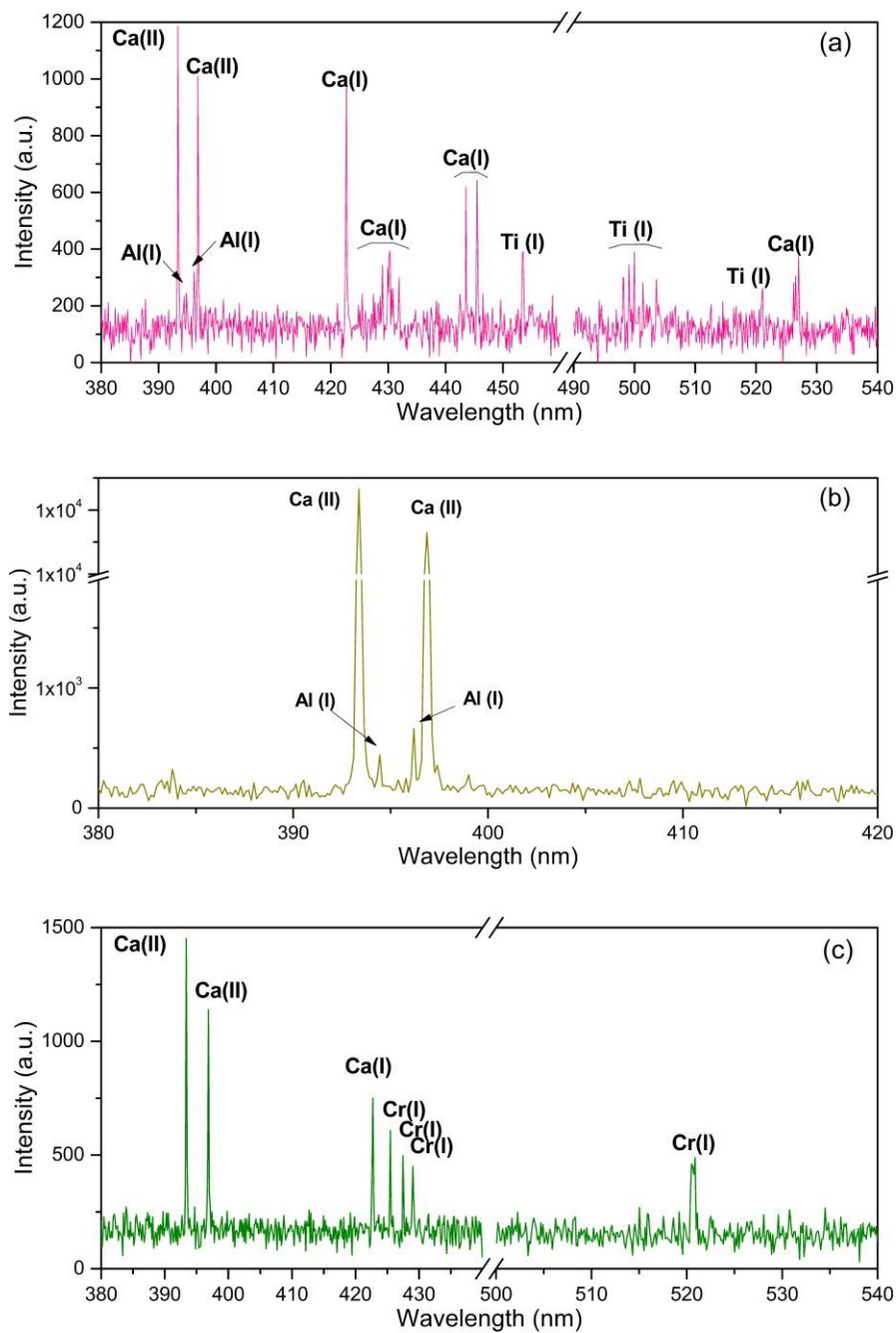


Figure 1. Examples of LIBS spectra from single laser events for (a) Ti-containing particle, (b) Al-containing particle (only a detail of spectral range is presented), (c) Cr-containing particle. Double-pulse mode. Oxycutting system, *on*.

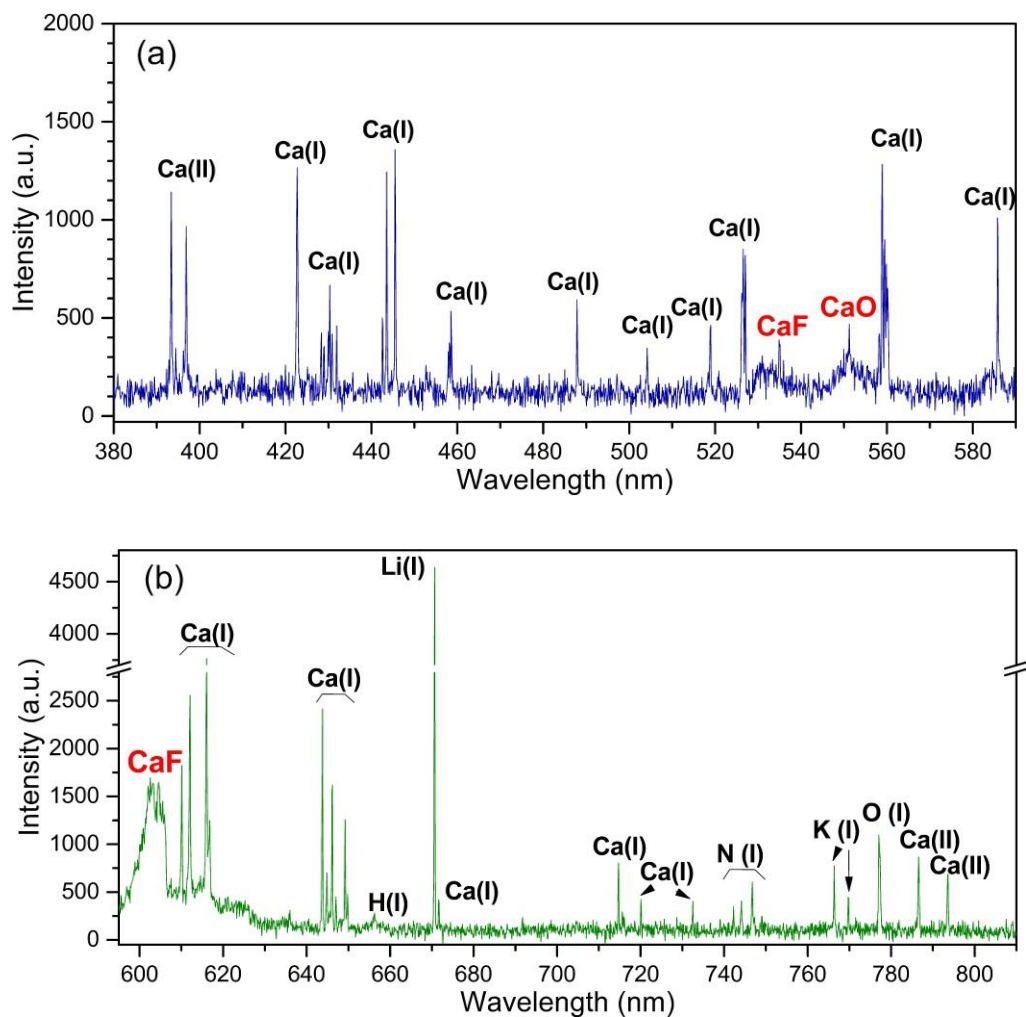


Figure 2. Example of two single shot LIBS spectra for ablation of particulate material containing mould powder particles: (a) without Li emission, (b) with Li emission. Double-pulse mode. Oxycutting system, *on*.

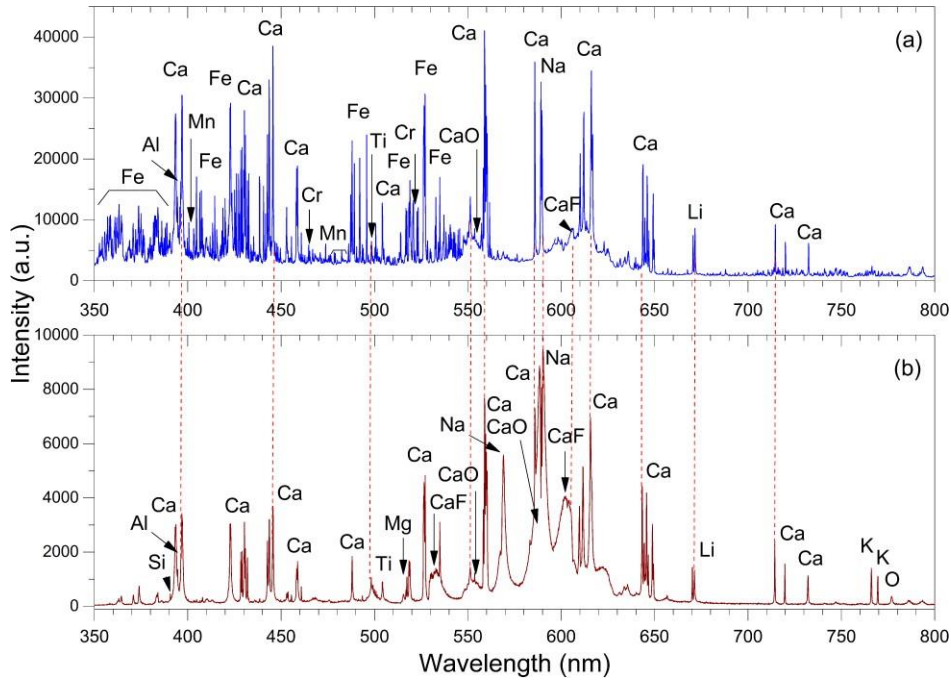


Figure 3. Real-time LIBS spectra corresponding to (a) stainless steel AISI 430 acquired at high temperature during continuous casting; (b) pellet of mould powder acquired in the laboratory at 900°C in similar experimental conditions. Laser operating in double-pulse mode and focusing the laser beam on the sample surface. Two spectrometer channels were connected simultaneously for light collection using a bifurcated-400 μm diameter fiber.

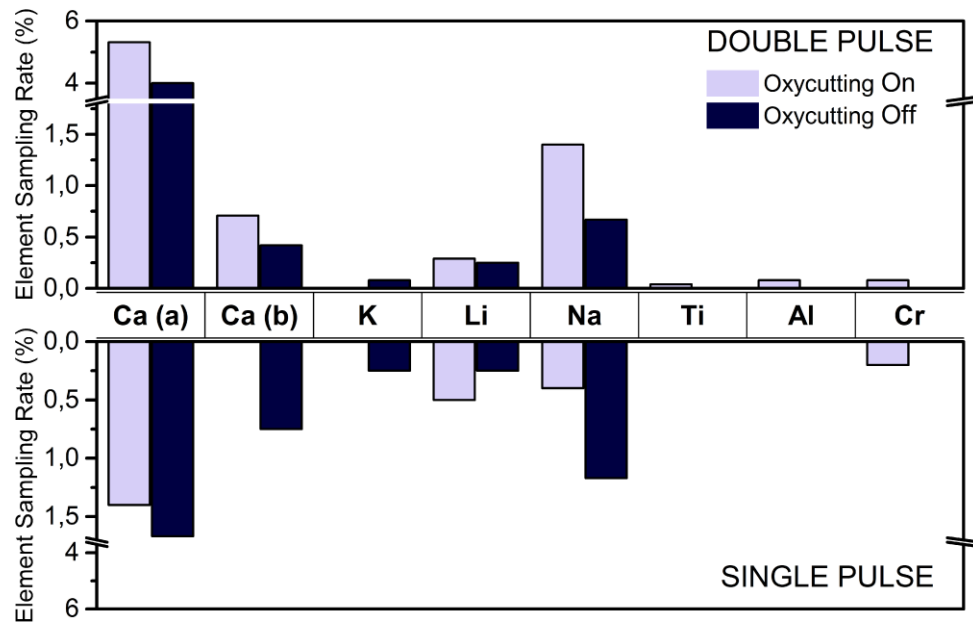


Figure 4. Sampling rate values for individual species, expressed in (%), measured when the oxycutting unit was on (dark bar) and off (light bar). Ca (a) detected in the spectral range 340 – 595 nm. Ca (b) detected in the spectral range 595 – 810 nm.

Table 1. Selected spectral lines used for the calculation of the sampling rates, for both spectral ranges.

Range 340 - 595 nm			Range 595 – 810 nm		
Element	Ion. Stage	Line (nm)	Element	Ion. Stage	Line (nm)
Ca	II	393.37	Ca	I	616.21
Ca	II	396.85	Ca	I	643.90
Ca	I	422.67	H	I	656.28
Al	I	396.15	Li	I	670.78
Ti	I	453.32	N	I	746.83
Cr	I	520.84	K	I	766.49
Na (doublet)	I	588.99/ 589.59	O (triplet)	I	777.19/777.42/777.54

Table 2. Elemental composition of the cast steel grade AISI 430. Samples were analyzed by X-ray fluorescence (XRF) and optical emission spectroscopy (Spark-OES) in Acerinox Europe S.A. labs from lollypops samples extracted from tundish during the casting process.

Concentration (%)										
Steel	EN	Si	Mn	Ni	Cu	Cr	Mo	Nb	Co	V
AISI 430	1.4016	0.42	0.32	0.23	0.05	16.65	0.03	0.01	0.02	0.11

Table 3. Elemental composition of mould powder used in the casting process. Supplied concentrations are averaged values. Synthetic slag density was about 0.64 g/cm³ and the grain-size was lower than 1.0 mm.

		Concentration (%)												
Mould powder	SiO ₂	CaO	Na ₂ O	CO ₂	Al ₂ O ₃	CaF ₂	MgO	C _{free}	TiO ₂	Fe ₂ O ₃	K ₂ O	MnO ₂	Li ₂ O	
MP-1	35.2	30.9	11.2	8.8	5.8	4.0	2.3	1.3	<0.7	<1.0	<0.5	<0.5	<0.5	

Table 4. RSD calculated from the averages of both spectral intensities and SNR values for the set of spectra whose SNR is greater than 3 and 10 respectively, for the examined elemental spectral lines, in experiments carried out using double- and single-pulse excitation.

Ensemble	Peak (nm)	Double Pulse		Single Pulse	
		Int. RSD (%)	SNR RSD %	Int. RSD (%)	SNR RSD %
SNR > 3	Ca II 339,37	39,16	172,30	28,34	100,33
	Ca II 396,86	36,81	180,30	25,80	109,40
	N I 744,23	15,43	46,03	14,62	47,40
	N I 746,83	21,88	61,67	23,02	60,77
	O I 777*	33,02	70,50	27,27	62,24
SNR > 10	Ca II 339,37	42,90	131,85	32,18	71,76
	Ca II 396,86	41,29	138,68	29,33	73,31
	N I 744,23	11,24	14,88	7,03	22,35
	N I 746,83	20,40	26,97	10,44	47,29
	O I 777*	25,24	39,71	24,80	33,77

Table 5. Summary of overall sampling rate values for tested conditions, expressed in (%)

Sampling rate	Double-pulse		Single-pulse	
	Oxyc. on	Oxyc. off	Oxyc. on	Oxyc. off
Range 340 - 595 nm	5.96	4.11	1.60	2.50
Range 595 – 810 nm	23.14	19.83	19.00	23.25
	1.00*	0.58*	0.50*	0.75*

*Spectra excluding events with only-air-rupture features