Observed Enhancement in LIBS Signals from Nano vs. Bulk ZnO Targets: Comparative Study of Plasma Parameters

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ABSTRACT

In this article, we will report an experimental evidence of enhanced LIBS emission upon replacing a Bulk-Based ZnO target by the corresponding Nano-Based target. The plasma was initiated via interaction of a Nd:YAG laser at the fundamental wavelength with both targets in open air under the same experimental conditions. The measurements show an enhanced emission from the Zn I-lines at the wavelengths of 328.26, 330.29, 334.55, 468.06, 472.2, 481.01, 636.38 nm. The measurements were repeated at different delay times in the range from 1 to 5 μs at constant irradiation level and fixed gate time of 1 μs. The average enhancement over the different Zn I-lines was found increases exponentially up to 8-fold with delay time. The electron density to each plasma was measured utilizing the Hα-line appeared in the emitted spectra from each plasma and was found to give similar values. The electron temperatures were measured via Boltzmann plot method utilizing the relative intensities of the Zn I-lines and were found to give very close values. Moreover, the relative population density of the ground state of the zinc atoms (relative concentration) was measured spectroscopically utilizing the Boltzmann plot method and was found to increase in a very similar trend to that of enhancement. The results of the spectroscopic analysis conclude that these signal enhancements can be attributed to the higher concentration of neutral atoms in the Nano-Based material plasma with respect to the corresponding Bulk-based ZnO material.

Keywords: LIBS; Enhancement; ZnO Nonmaterial; Hα-Line; Zn I-Lines; Spectroscopy

1. Introduction

LIBS (acronym standing for laser induced breakdown spectroscopy) is one of the potential growing fields of analytical applications. It was successfully used for elemental analysis [1], quality of metal industry [2], cleaning [3], studying of old archeology [4], characterization of soils [5] and in jewelry industry [6]. This passive spectroscopic technique is based on utilizing the light emitted from plasma, assuming that the emitted radiation is sufficiently influenced by the plasma properties [7,8].

However, the relatively small signal to background ratio presents one major problem which imposes a limitation on the use of this technique, hence the poor the ability of the LIBS technique to detect the very small concentration of the different matrix elements [9,10]. Different techniques were devised to overcome this difficulty, such as the double pulse technique [11,12] in which the target material is irradiated by consequent double laser pulses separated by a certain delay time. Moreover, the introduction of the femtosecond laser source provides one basic advantage, which is the very small continuum component appeared under the emitted lines from plasma, hence the better the signal to background ratio [13,14].

On the other hand, the nanomaterials are categorized as those which have structured components with at least one dimension less than 100 nm [15]. Two principal factors cause the properties of nanomaterials to differ significantly from bulk materials namely; the tremendous increase in the relative surface area and the quantum effects. These factors can substantially change and/or enhance the well known bulk properties, such as chemical reactivity [16], mechanical strength [17], electrical and...
magnetic [18] and optical characteristics [19]. As a particle size decreases, a greater proportion of atoms are found at the surface compared to those inside [15]. The quantum effects can begin to dominate the properties of matter as its size is reduced to the nano-scale. Therefore, nanoparticles are of interest because of its inherent new properties when compared with larger particles of the same materials. For example, titanium dioxide and zinc oxide become transparent at the nano-scale however they are able to absorb and reflect UV light [20].

The accelerating progress in the field of nanomaterial science may provide a new facility to reach signal enhancement. To the best of our knowledge no work has been done in a systematic manner to examine the optical signal emitted from the laser produced plasmas utilizing nanomaterials base targets instead of bulky ones.

In this work we will demonstrate the ability of combining the LIBS technique with Nano-Based materials (used as a target) to get strong enhanced optical signals.

2. Experimental Setup

The experimental setup is shown in Figure 1(a). The irradiation system comprises a Nd:YAG laser able to deliver an energy of 650 mJ/pulse at the fundamental wavelength 1064 nm, at constant duration of 5 ns. The detection system consists of an SE-200 echelle type spectrograph equipped with time control iStar®-ICCD camera. The emitted light from plasma is spatially integrated and collected at the entrance hole of spectrograph via a 25 μm quartz fiber, positioned with the help of precise xyz-translational stage holder at 15 mm from the laser-plasma axis. The relative spectral response of the camera-MCP (micro channel plate) and spectrograph including optical fiber over the entire wavelength window from 200 to 1000 nm was measured using a Deuterium-Halogen lamp (type, DH-2000-CAL) with the result as shown in Figure 2(a). The processing of the experimental data was carried out using home-made routine built under the MATLAB7®package [21]. Each data
point was taken as the average over three different single laser shots on fresh target condition which enable us to account for any fluctuations in measurements. A monitor to the incident laser power on the target was made via a beam splitter which reflects 4%, absorb 4% and with the help of absolutely calibrated power-meter (type Ophier). Unless specified, the ZnO (MK\textit{NANO}-ZnO-020) material (as shown in Figure 1(b)) with almost spherical shaped powder samples was purchased from “MK\textit{NANO}” and was used without further purification.

The nano and bulk powder targets were compressed to a disk shape (Pressure ~ 5 ton/cm\(^2\)), under the same ambient conditions for a time period of 5 min. Both tables were positioned side by side perpendicular to laser axis on a xy-translational stage. It is worth noting that, no further chemical or heat treatment was carried out and to the best of our knowledge, both the nano and bulk materials were synthesized using the same base material.

### 3. Plasma Parameters

The plasma state can be described by two measurable parameters namely; the electron density and temperature \([7,8]\). At relatively large electron density \(\sim 10^{19} \text{ cm}^{-3}\), the plasma state is said to be in complete thermodynamical equilibrium (CTE). In this situation the emitted light from the plasma displays a continuous spectrum and the plasma is characterized by single temperature. This condition is rarely verified in the laser induced plasma experiments \([7]\).

In LIBS experiments, it was verified that the electron density is in the range from \(10^{16} \text{ to } 10^{18} \text{ cm}^{-3}\) and the radiation field is dominated by a line rather than continuous spectrum. This state is called local thermodynamical equilibrium (LTE), at which the particle species are characterized by unique temperature which is different from the temperature of the radiation field.

At a rather lower electron density regimes \((10^{16} > \text{ne} > 10^{8} \text{ cm}^{-3})\), the electron gas in plasma tends to divide the energy levels of the atoms into two main categories. We call this state as partial local thermodynamic equilibrium (PLTE) \([7,8]\).

#### 3.1. Measurement of Plasma Electron Density

Spectroscopically, the electron number density can be measured by different methods namely; measurement of the optical refractivity of the plasma \([7]\), calculation of the principal quantum number at the series limit \([7,8]\), measurement of to Stark profile of certain optically thin emitted spectral lines \([22]\), the measurement of the absolute emission coefficient (spectral intensity) of spectral line \([7]\) and finally from the measurement of the absolute emissivity of the continuum emission \([8]\).

Among the different proposed methods, the Stark broadening of emitted lines has been the most widely used method \([8]\). This method is based on the assumption that the Stark effect is the dominant broadening over the Doppler broadening and the other pressure broadening mechanisms resulted from collisions with neutral atoms (\textit{i.e.}, resonance and Van der Waals broadenings) \([22]\). The theoretical calculation of Stark broadening parameters of hydrogenic lines is described in detail by Griem \([8]\).

For the H\(_{\text{a}}\)-line, the following expression can be used to calculate the electron density \([22]\):

\[
 n_e (H_{\text{a}}) = 8.02 \times 10^{12} \left( \frac{\Delta \lambda_1 (H_{\text{a}})}{\alpha_{\text{e}/2}} \right)^{3/2} \cdot \text{cm}^{-3}
\]

(1)

In this expression \(\alpha_{\text{e}/2}\) is the half width of the reduced Stark profiles in Å, it is a weak function of electron density and temperature through the ion-ion correlation and Debye-shielding correction and the velocity dependence of the impact broadening.

#### 3.2. Measurement of Plasma Temperature

The second important plasma parameter is the electron temperature, which determines the strength of the different distribution functions \([7]\). In laser produced plasmas, a combination of the line and continuum spectra is appeared in the emission spectra. In this condition, the LTE is almost being fulfilled and one should employ the optical emission spectroscopy technique to measure the temperature \([7,8]\).

The most direct method to estimate the temperature is via measurement of the relative spectral intensity of two or more lines emerging from the same element and ionization stage with small wavelength separation and large separation in upper state excitation energy. Moreover, these lines should be optically thin \([7,8]\). The following expression is recommended:

\[
 \ln \left( \frac{I_{1\lambda} A_1 C_{1\lambda}}{A_{2\lambda} C_{2\lambda}} \right) = \ln \left( \frac{h c N_\lambda}{4 \pi U_\lambda} \right) - \left( \frac{\Delta E}{k T_e} \right)
\]

(2)

In this expression, \(I, \lambda, A, g\) are the spectral intensity, wavelength, transition probability and statistical weight of the upper state respectively. The subscript numbers indicate different lines. \(N\) and \(U\) are the population density and the parathion function of the atom at temperature \(T_e\). The constants \(h\) and \(c\) are the Planck constant and speed of light, respectively.

The lines intensities should be corrected by the relative response factors \(C_{1\lambda,2}\) at the different emitted wavelengths; these factors are saved from the absolute calibration curve, shown in Figure 2(a). Mathematically, the plot of the LHS of Equation (2) with excitation energy (RHS) yields a straight line of negative slope which determines the temperature.
3.3. Measurement of Relative Concentration

In order to measure the relative concentration of the zinc atoms in both plasmas created from the nanomaterial and the corresponding bulk material targets, we have developed the following simple method. From Equation (2), the term \((hcN_o/4\pi U_o)\) is a constant defined by the intersection of the backward extrapolation of the Boltzmann line with the vertical axis. This method is not new, since it was suggested before [23,24] in what is called multi-element Saha-Boltzmann plot in which the relative concentration of the different matrix elements in the target material can be estimated.

After the construction of the two Boltzmann plots, one for the plasma emerging from the Nano-Based ZnO material target and the other for the plasma created from the Bulk-Based ZnO target, we should get two straight lines as shown in Figure 2(b).

The upper line proves a higher concentration than the lower one. The two points of intersection with the vertical axis should give two similar quantities \((hcN_o/4\pi U_o)_{Nano}\) for the Nano-Based target and \((hcN_o/4\pi U_o)_{Bulk}\) for the Bulk-Based target.

This means that, if the lower bulk line is multiplied by factor \(\eta\) we can make the two straight lines coincide (of course, with little orientation defined by the different temperatures of the two plasmas). This factor represents the relative concentration of the atoms (stoichiometry) in the plasmas created from the Nano vs. Bulk targets i.e. \(\eta = (N_{Nano}/N_{Bulk}) = (C_N/C_B)\).

4. Results and Discussion

Figure 3(a) shows an example of one emitted optical signal at the Zn I-lines from the plasmas created from the Nano-Based target material (red colored) in comparison to that from the Bulk-Based material (black color), showing the emission from the Zn I-lines as depicted in figure; (b) Demonstration to enhanced spectral radiance from the Zn I-lines at different wavelength regions from the Nano-Based target (red colored) in comparison to that from the Bulk-Based target at different three arbitrary delay times of 1, 3, 5 \(\mu\)s, are shown in subsequent rows.

Figure 3. (a) Demonstration to emission enhancement at the Zn I-lines from the Nano-Based target (red color) in comparison to that from the Bulk-Based material (black color), showing the emission from the Zn I-lines as depicted in figure; (b) Demonstration to enhanced spectral radiance from the Zn I-lines at different wavelength regions from the Nano-Based target (red colored) in comparison to that from the Bulk-Based target at different three arbitrary delay times of 1, 3, 5 \(\mu\)s, are shown in subsequent rows.
to signals at the same wavelengths arise from the corresponding Bulk-Based material target (black colored) at an arbitrary delay time of 4 μs. Qualitatively, one can notice the clear enhancement in the spectral intensity from the nano to bulk plasmas. Moreover, Figure 3(b) displays the same comparison between the two signals at the same wavelengths at regular delay times of 1, 3 and 5 μs, respectively, which demonstrates the temporal increase in the signal enhancement as the delay time increases.

This is in contrast to the behavior of the spectral intensity of the Hβ-line. The reason for the decrease in the spectral intensity is not yet been resolved which grasps a need for more extensive investigations.

Moreover, there is an obvious difference in the enhancement factor at the different Zn I-lines (This will be treated in a separate publication).

An attempt to know the reason which causes this behavior can be achieved via careful examination of the different plasma parameters (electron density and temperature) in both plasmas. The electron density was measured utilizing the optically thin Hβ-line appeared in both spectra under the same condition. Utilizing Equation (1), in conjunction with special software routine written in MATLAB7®. This program was constructed to compare the spectral line shape of the measured Hβ-line to the theoretically constructed Voigt profile [21] and hence to extract the Lorentzian component of the line FWHM used to calculate the electron density as shown in Figure 4 (first two columns).

![Figure 4](image-url)

**Figure 4.** Demonstration of the Hβ-line fitting and the measured electron densities at three different delay times of 1, 3, 5 μs from the Nano-Based plasma target (black color) and the Bulk-Based plasma (blue color). The third column demonstrates the measurement of the plasma temperature at three delay times of 1, 3, 5 μs from the Nano-Based plasma (upper red line) in comparison to Bulk-Based plasma (lower black line) in addition to method of calculation for the relative concentration η.
An example of the fitting of the $H_\alpha$-line at regular delay times of 1, 3, 5 $\mu$s to both plasmas arises from Nano-Based target (first column with red curves) in comparison to the second column in Figure 4, which shows the fitting of the $H_\alpha$-line appeared in the spectra arises from the Bulk-Based plasma (blue curves) together with the estimated electron densities.

On the other hand, in order to calculate the plasma electron temperature, we have constructed the Boltzmann plots utilizing the spectral lines intensities emitted by the Zn I-lines at 330.29, 334.55, 472.2, 481.01, 636.38 nm with the results as shown in Figure 4 (third column).

The atomic parameters of the Zn I-lines used to construct the Boltzmann plot are presented in Table 1.

Also, Figure 4 (third column) reveals that; first, the concentration of the zinc atoms in the plasma created from the Nano-Based target is higher than the corresponding plasma arising from the Bulk-Based material and second, both plasmas show nearly similar temperatures at the different delay times, since both of the Boltzmann lines are almost parallel.

Finally, in order to calculate the relative concentration (stoichiometry) of the neutral zinc in both plasmas (factor $\eta$); we have utilized the predictions of set of Boltzmann plots in Figure 4 in which the factor $\eta$ was calculated at each delay time. The overall result of the measurement of the average enhancement factor over the different emission wavelengths from the neutral Zn I-lines at the wavelengths of 328.26, 330.29, 334.55, 468.06, 472.2, 481.01, 636.38 nm with delay time is shown in Figure 5, (red squares) whereas an obvious increase in an exponential manner to the signal enhancement from the nano vs. bulk based materials is shown.

For a comparison reason, we have plotted both of the measured relative electron density (blue solid squares) and the relative electron temperature (black disks) as well as the measured relative concentration (inverted green triangles) with delays times in the range from 1 to 5 $\mu$s, respectively, in Figure 5.

Figure 5 reveals the following conclusions, first, it ensures that the enhanced spectral intensity from the Zn I-lines can’t be attributed to the plasma temperature difference or the difference in the electron densities. Second, from this figure it is clear that the signal enhancement depends only on the relative atomic concentration the plasma created from the Nano-Based material with respect to Bulk-Based plasma. Nevertheless, the relative concentration (inverted green triangles) increases in a very combatable manner with signal enhancement.

This fact can be qualitatively explained in terms of the collisional radiative modeling: The intense emission from the nanomaterial plasma with respect to the bulk one under the same experimental conditions can be related to the higher number of excited atoms to the upper emitting state (labeled $j$). This state can be populated via different atomic processes e.g. collisional excitation from all lower lying states, especially from the ground state and/or collisional de-excitation from the upper states, especially from the ground state of the next ionization stage and/or radiative decay from the upper excited states, especially recombination of both types (radiative or non-radiative three particle recombination).

All the relevant processes can contribute by a certain amount depending on the predominant rate coefficients and the electron density present in the plasma, as well as the electron temperature [7]. In the laser produced plasmas (LIBS), the electron density is in the order of $10^{17}$ cm$^{-3}$, which means that the collisional processes are the dominant ones. For close approximation, the number of collisional processes from the ground state leading to population of the upper emitting state $j$ per unit volume per unit time is in direct proportion to the spectral intensity of the emitted radiation. Hence the relative intensity from either plasma (Enhancement) should amount to;

$$\frac{I_j}{I_{jB}} \propto \frac{N_j}{N_{jB}} \propto \frac{N_{\alpha j}}{N_{\alpha B}} \frac{n_{\alpha j}}{n_{\alpha B}} \frac{R_j}{R_B}$$

Table 1. The atomic parameters of the Zn I-lines.

<table>
<thead>
<tr>
<th>Wavelength (nm)</th>
<th>Transition probability $A$ (s$^{-1}$)</th>
<th>Statistical weight $g$</th>
<th>Energy of upper state $E$ (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>481.01</td>
<td>7.00 $\times$ 10$^7$</td>
<td>3</td>
<td>6.6674</td>
</tr>
<tr>
<td>472.2</td>
<td>4.58 $\times$ 10$^7$</td>
<td>3</td>
<td>6.6673</td>
</tr>
<tr>
<td>330.29</td>
<td>1.07 $\times$ 10$^5$</td>
<td>5</td>
<td>7.7975</td>
</tr>
<tr>
<td>334.55</td>
<td>1.50 $\times$ 10$^4$</td>
<td>7</td>
<td>7.7980</td>
</tr>
<tr>
<td>636.38</td>
<td>4.65 $\times$ 10$^7$</td>
<td>5</td>
<td>7.7585</td>
</tr>
</tbody>
</table>
the population density of this upper state and the population density of the ground state, respectively. \( R, n_e \) are the electron-atom collision excitation rate coefficient and electron density, respectively. The subscripts \( N, B \) indicate nano and the corresponding bulk quantity. Equation (3) indicates that the relative spectral intensity would increase linearly with both of relative concentration \( \eta = \left( N_{eN} / N_{eB} \right) \) and the relative electron density \( \left( n_{eN} / n_{eB} \right) \), and the relative electron temperature through temperature dependant electron-atom collisional excitation rate coefficient [7].

Experimentally, the observed invariance of the relative electron temperature, density and together with the measured relative enhancement compatible relative concentration variation under different conditions leads to conclusion that the higher concentration of atoms in the Nano-Based plasma is decisive in the observed enhancement phenomenon.

5. Conclusion

An enhanced emission optical signal from the Nano-Based material target was observed in comparison to Bulk-Based plasma from ZnO. This signal enhancement, after a careful investigation of the different plasma parameters, was attributed to the larger concentration of the zinc atoms in the plasma created from the Nano-Based material with respect to that created from the Bulk-based one. Further investigation would be appreciated to explore more the effects of the different laser wavelengths and laser fluence as well as the type of the material target.

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This work is completely dedicated to the memory of souls of the Egyptian martyrs during our revolution on Jan. 25, 2011.

REFERENCES


