



Article Laser Induced Breakdown Spectroscopy of Er II for Transition Probability Measurements

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Abstract: We present a laser induced breakdown spectrum of Er II in the near ultraviolet region. To use the spectrum for the evaluation of the transition probabilities, an alloy target with a low content of Er was used to suppress the self-absorption. From the linearity of the Boltzmann plot obtained by using the sensitivity corrected experimental intensity and existing transition probability data, the local thermal equilibrium condition of the plasma and the reliability of the transition probability data are confirmed. The linear function obtained in the Boltzmann plot is used for the determination of a previously unreported transition probability for the line at 393.863 nm.

Keywords: laser induced breakdown spectroscopy; transition probabilities; lanthanides; erbium; neutron star merger

1. Introduction

Spectroscopic data of rare-earth elements are important for several fields, especially in astrophysics. For example, data of transition wavelength and probability are essential to estimate the amount of these elements in the stellar atmosphere [1,2]. In addition, such atomic data sets are needed for the accurate radiative transfer simulation for kilonovae produced by neutron star mergers [3,4]. Er is one of the most important among rare-earth elements.

In the NIST atomic spectra database (ASD) [5], 285 lines of Er II are registered on the basis of the data compilation by Meggers et al. [6] in 1975. Among them, only 11 lines are listed with energy level classifications and transition probabilities. Musioł and Łabuz [7] measured relative transition probabilities of 101 lines by arc plasma spectroscopy. By normalizing the relative data to the experimental radiative lifetime by Bentzen and Nielsen [8], absolute values were determined. Xu et al. [9] measured radiative lifetimes of 30 excited levels by time-resolved laser-induced fluorescence (LIF). By using theoretical branching fractions, transition probabilities of 32 lines were obtained.



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Copyright: © 2022 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). Significant progress was made by Lawler et al. [10] (this paper is referred to as Lawler2008 hereafter), who determined the transition probabilities of 418 lines from their branching fraction measurements with a combination of radiative lifetimes measured by Stockett et al. [11]. Some of the transition probability data in Lawler2008 were evaluated in a recent experimental study by Yu et al. [12] using a time resolved laser induced fluorescence method. Good agreement in the *gA* values found between their data implies the reliability of the transition probabilities in Lawler2008. Wyart and Lawler [13] extended the study with the aid of theoretical analysis, using the Cowan code [14], and 130 levels of even parity and 230 levels of odd parity were summarized with configuration mixing coefficients. Ankush and Deo [15] estimated the isotope shift between ¹⁶⁶Er and ¹⁷⁰Er, using the mixing coefficients derived by Wyart and Lawler, and found that the estimated shifts were consistent with the experimentally measured shifts.

In this paper, we present an emission spectrum of Er II, obtained by laser induced breakdown spectroscopy (LIBS), where the emission from plasma produced by an intense laser pulse is observed. LIBS has been mainly developed for element analysis in industrial manufacturing [16], environmental monitoring [17,18], geological studies [19] including planetary exploration [20], and many other applications. On the other hand, LIBS is one of the most powerful methods for the determination of fundamental atomic properties, such as transition probabilities (see the review paper [21] in this Special Issue and references therein). In order to determine transition probabilities properly, a local thermal equilibrium (LTE) condition should be realized. Although the temperature of the plasma varies with time, instantaneous observation of the plasma with a time-resolved intensified camera can ensure the LTE condition. Since the emission intensity I in the LTE condition is proportional to the product of the transition probability A and the Maxwell-Boltzmann population determined by the temperature T and the upper level energy E, the logarithm of I/gA, where g is the statistical weight, should have a linear dependence on E (see Section 2). The linearity can be used to confirm the reliability of the A values used. Once the linearity is confirmed, unknown A values can be obtained by applying the experimental intensity to the linear function obtained. LIBS is a simple method for evaluating or accumulating transition probability data, compared with other methods such as LIF. It also has the advantage that many transitions can be evaluated at the same time and that metal targets can be used as is, without vaporization. From the LIBS spectrum, we confirm the linearity by using the A values in Lawler2008 and a previously unreported transition probability is derived from the obtained linearity.

2. Experiment

Figure 1 shows a schematic of the experimental setup. A Q-switched Nd:YAG laser (Spectra Physics GCR-150) was used in the fundamental mode (1064 nm) for producing plasma. Laser pulses with a width of 8–9 ns and an energy of about 15 mJ were focused with a 150 mm focal length lens and introduced through a fused silica window. The pulse repetition frequency was 10 Hz.

For the determination of transition probabilities, it is important to realize a thin plasma condition [21] as well as the LTE condition. If the plasma is so thick that it can absorb the emission occurring within the plasma (self-absorption effect), the observed intensity does not reflect the transition probability. An Al alloy with a low content ($5 \text{ wt}\% \sim 0.8 \text{ at}\%$) of Er was thus used as a target in the present experiment for producing thin Er plasma. A rod-shaped target with a 6 mm diameter was installed in the chamber. The target rod was rotated while being moved up and down with a motion feedthrough system similar to that used in [22]. The target motion enabled each laser pulse to hit the unirradiated surface of the target. The target chamber was filled with 200 Pa of Ar after evacuating the chamber with a scroll pump.



Figure 1. Schematic diagram of the experimental setup.

The emission from the plasma was observed through a fused silica window placed at 90° with respect to the laser direction. A set of synthetic silica plano-convex lenses was used to focus the emission onto the entrance of the fiber connected to an echelle spectrometer (LTB ARYELLE 200). A fast-gated image-intensified sCMOS (Andor iStar-sCMOS-18F-03) was used as a detector. The spectral resolution (specification value) is 9200 at 253 nm and 8000 at 404 nm. The data acquisition period was 1 µs after a delay of 5 µs from each laser pulse. The timing was controlled by a digital delay generator installed in the sCMOS detector. The spectrometer response function was obtained using a calibration light source (Ocean Optics DH-3*plus*-BAL-CAL).

The emission line intensity I (photon number emitted in a unit of time) is determined by the product of the population of the upper level and the transition probability A. It is thus represented as the following equation for the lines emitted from the local thermal equilibrium (LTE) plasma with a temperature T:

$$I = gA \cdot C \exp\left(-\frac{E}{kT}\right),\tag{1}$$

where *g*, *E*, and *k* represent the statistical weight, the energy of the upper level, and the Boltzmann constant, respectively, and *C* is a constant depending on the ion species. By taking the natural logarithm, we obtain

$$\ln\left(\frac{I}{gA}\right) = -\frac{E}{kT} + C',$$
(2)

where C' is a new constant. Thus the plot of the left side value as a function of E (Boltzmann plot) should indicate linearity when the LTE condition is established.

3. Results and Discussion

Figure 2a shows the LIBS spectrum for the whole spectrometer region (200–1080 nm) obtained by about 1000 shot accumulations. Close-up views of the 300–400 nm, 330–340 nm and 390–400 nm regions are also shown in Figure 2b–d, respectively. In addition to the emission lines of Er, lines from Ar (atmospheric gas) and Al (the main content of the target alloy) were also confirmed, including strong lines of Al I [5] at around 310 nm and 395 nm. Among the observed Er lines, 30 Er II lines in Lawler2008 were found to have enough intensity to analyze. All of them, which are listed in Table 1, were concentrated in the 300 nm range shown in Figure 2b. For example, in the 330–340 nm range shown in Figure 2c, eight lines from Er II were observed, as indicated by the red arrows. They are well represented by a Gaussian line shape, as shown in the inset for the 337.271 nm line as an example. The peak area of each observed line was thus obtained by fitting a Gaussian function, and the relative intensity *I* was obtained, as listed in Table 1, by correcting the peak area with the response function obtained with the calibration light source.



Figure 2. LIBS spectrum obtained for an Al-5 wt% Er alloy target in an Ar atmosphere for the (**a**) whole spectrometer (200–1080 nm), (**b**) 300–400 nm, (**c**) 330–340 nm, and (**d**) 390–400 nm regions. It is noted that the sensitivity correction is not applied. It is also noted that the undulated baseline of the spectrum reflects the response of each diffraction order of the Echelle grating. The lines indicated by the red arrows represent those given in Lawler2008 [10]. The blue arrow indicates the 393.863 nm line listed in the NIST database without a transition probability.

The self-absorption effect can be estimated from the line intensity ratio between the intense transitions with the same upper level [23,24]. When the thin condition is valid, the intensity ratio of such a transition pair should be equivalent to the branching ratio determined only by the transition probability ratio. On the other hand, when the thin condition is not valid, the intensity ratio of such a pair tends to deviate from the branching ratio, as the self-absorption effect decreases the observed intensity more significantly for the transition with a larger transition probability. In the present spectrum, the ratio between the 389.623 nm and the 383.048 nm lines, which are relatively intense and have the same upper level with an energy of 3.236 eV, is a suitable probe for the self-absorption effect. The sensitivity corrected experimental intensity ratio for those lines was 1.2 ± 0.2 , which is in good agreement with the branching ratio 1.23 ± 0.09 [10] within the uncertainties. We also measured this ratio using a target with twice the content (10 wt%~1.8 at%) of Er. Although consistency with the branching ratio was also confirmed for the higher content target, we used the lower content (5 wt%~0.8 at%) target to ensure the thin plasma condition more certainly.

Table 1. Transitions used in the present analysis. Wavelength (λ), parity, 2*J* value, level energy (*E*), and transition probability (*A*) are taken from Lawler2008 [10] except for the 393.863 nm line. For the 393.863 nm line, wavelength, parity, 2*J* value, and level energy taken from the NIST database [5] and the transition probability obtained in the present measurement are listed. The sensitivity-corrected relative intensities *I* in the present LIBS spectrum, normalized to the 390.631 nm line, are listed for all the lines.

		Lower			Upper		A	
λ (nm)	Parity	2 <i>J</i>	<i>E</i> (eV)	Parity	2 <i>J</i>	<i>E</i> (eV)	(10^6 s^{-1})	Ι
290.447	odd	15	0.846164	even	17	5.113654	93 ± 5	0.05 (1)
291.036	odd	15	0.846164	even	15	5.105013	204 ± 11	0.12 (1)
296.452	odd	15	0.846164	even	17	5.027212	159 ± 8	0.14 (2)
300.241	odd	15	0.846164	even	13	4.974455	130 ± 7	0.14 (2)
323.058	even	11	0.054607	odd	11	3.891327	92 ± 5	0.42 (5)
323.798	even	9	0.636362	odd	9	4.464322	19.1 ± 1	0.05 (1)
326.478	even	13	0.000000	odd	13	3.796533	69 ± 3	0.44 (5)
330.557	even	9	0.892110	odd	9	4.641802	32.1 ± 1.7	0.03 (1)
331.243	even	11	0.054607	odd	13	3.796533	40.9 ± 2.1	0.31 (3)
332.319	even	9	0.636362	odd	11	4.366164	34.3 ± 1.9	0.11 (1)
334.603	even	11	0.054607	odd	9	3.758950	25.6 ± 1.3	0.15 (2)
336.408	even	11	0.054607	odd	11	3.739083	18.5 ± 0.9	0.17 (2)
336.802	even	11	0.054607	odd	11	3.734768	18.8 ± 0.9	0.16 (2)
337.275	even	13	0.000000	odd	15	3.674999	145 ± 7	0.89 (10)
339.199	even	13	0.000000	odd	15	3.654160	28.3 ± 1.4	0.29 (3)
349.910	even	11	0.054607	odd	9	3.596907	105 ± 5	0.62 (7)
359.950	odd	15	1.402160	even	13	4.845660	52.2 ± 2.7	0.20 (2)
360.490	odd	19	1.588866	even	17	5.027212	42 ± 3	0.07 (1)
361.657	even	13	0.000000	odd	11	3.427253	21 ± 1.1	0.37 (4)
369.265	even	11	0.054607	odd	13	3.411246	67 ± 3	0.95 (10)
370.764	odd	15	1.770584	even	17	5.113654	46 ± 4	0.06 (1)
373.127	even	9	0.892110	odd	11	4.214012	19.5 ± 1	0.08 (1)
373.816	odd	13	1.619322	even	15	4.935095	46.8 ± 2.5	0.06 (1)
374.264	even	9	0.636362	odd	9	3.948168	20.6 ± 1.1	0.11 (1)
383.048	even	13	0.000000	odd	13	3.235860	19.4 ± 1	0.49 (6)
385.839	even	9	0.892110	odd	11	4.104563	18.5 ± 0.9	0.08 (1)
388.061	even	9	0.636362	odd	7	3.830423	31.3 ± 1.6	0.15 (2)
388.289	even	11	0.886441	odd	9	4.078630	31.6 ± 1.6	0.12 (2)
389.623	even	11	0.054607	odd	13	3.235860	23.9 ± 1.2	0.59 (6)
390.631	even	13	0.000000	odd	11	3.173046	48.2 ± 2.4	1.00 (10)
393.863	even	13	0.000000	odd	11	3.147014	12 ± 2	0.26 (3)

The left side values of Equation (2) obtained from the experimental intensity *I* and the *gA* values in Lawler2008 are plotted in Figure 3 as a function of the energy *E* of the upper level of the transition. The error bars represent the uncertainty obtained considering four contributions: (i) the fitting error, (ii) the statistical uncertainty of the photon counts estimated from the conversion factor from the digital counts to the photon number, (iii) the uncertainty in the radiometric data of the calibration light source, and (iv) the uncertainty in the *A* values. The linear function corresponding to Equation (2) was obtained by weighted fitting to the data, as shown by the solid line. The 68.3% confidence interval is also shown by the dashed curves. From the gradient of the fitted linear function, the plasma temperature is derived to be 6700 ± 400 K.

As seen in the figure, the Boltzmann plot shows rather a good linearity although some of the data are scattered from the fitted linear function. In general, the linearity in the Boltzmann plot ensures the LTE condition of the plasma and the reliability of the gA values. In other words, the deviation of the data from the linear dependence should be attributed to the deviation from the LTE condition or the error in the gA value (or both). We consider that the scattering is due to an incomplete LTE condition in the present experiment, as the reliability of the data in Lawler2008 was confirmed through comparison with the transition

probability data independently obtained by a different method from another group [12], as described in Introduction. However, the large deviation for the data at E = 4.85 eV may imply an underestimation of the gA value for the corresponding transition (359.950 nm). If the data is applied to the obtained linear function, an A value of $1.5(2) \times 10^8$ s⁻¹ is obtained, whereas it is $5.22(27) \times 10^7$ s⁻¹ in Lawler2008.



Figure 3. Boltzmann plot obtained using the present experimental line intensity *I* and the *gA* values taken from Lawler2008 [10] (red squares). The fitted linear function and the 68.3% confidential interval are shown in the solid and dashed lines, respectively. The blue square represents the linear function value at 3.147 eV for obtaining the *A* value of the 393.863 nm line (see text for details).

In the NIST database, there is one transition at 393.863 nm for which the lower and upper level energies are given but the transition probability is not given. The transition probability of this line is not included either in Lawler2008. As listed in the last line in Table 1, the angular momentum and the parity of the upper level are defined as J = 11/2 and odd in the NIST database, but the electron configuration is not given. According to Wyart and Lawler [13], $4f^{11}5d6s$ is assigned to the corresponding level although $4f^{11}6s^2$, $4f^{12}6p$, and $4f^{11}5d^2$ are mixed with a mixing coefficient of 0.37, 12.55, and 2.71 %, respectively.

This 393.863 nm line was observed in the present LIBS spectrum, as indicated by the blue arrow in Figure 2d, and the intensity was obtained from the peak area of the fitted Gaussian function shown in the inset. By applying this line to the linear function in Figure 3, as plotted by the blue square, the transition probability A is obtained as $1.2(2) \times 10^7 \text{ s}^{-1}$. The uncertainty was estimated from three contributions: (i) uncertainty in the experimental intensity, (ii) uncertainty in radiometric data of the calibration light source, and (iii) the confidence interval of the linear function obtained from the Boltzmann plot, among which the third contribution is dominant.

Our work demonstrated that measurements of the LIBS spectra provide an effective way to confirm the reliability of the existing transition probability data. In addition, we showed that, when there are some existing data of transition probabilities for a certain ion, the LIBS spectra give an estimation of unknown transition probabilities for the lines with identified upper levels. In particular, the latter aspect is useful to systematically give transition probabilities of the lanthanide elements, for which extensive datasets are not available.

4. Summary

In the present laser induced breakdown spectroscopy experiment, emission from Er II excited in the plasma produced by a Q-switched Nd:YAG laser was observed, using an echelle spectrometer with a fast gated intensified detector. The local thermal equilibrium condition of the plasma was confirmed by the linearity in the Boltzmann plot, obtained from the fast-gated spectrum and existing reliable transition probability data. The transition probability of a line at 393.863 nm, which has previously not been reported, was determined by applying its intensity to the linear function obtained in the Boltzmann plot to be $1.2(2) \times 10^7 \text{ s}^{-1}$.

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Abbreviations

The following abbreviations are used in this manuscript:

- LIBS Laser induced breakdown spectroscopy
- NIST National Institute of Standards and Technology
- LIF Laser induced fluorescence
- LTE Local thermal equilibrium
- CMOS Complementary metal oxide semiconductor

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