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Calibration-free quantitative analysis of thin-film oxide layers in semiconductors using Laser Induced Breakdown Spectroscopy (LIBS)

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Abstract:

Current largest market share and continually growing industry of the semiconductor manufacturing sector in the US demands rapid and cost-effective quality control and characterizations of thin film semiconducting materials. To this end, we demonstrate Laser Induced Breakdown Spectroscopy (LIBS) as a facile and effective analytical tool for rapid process-line characterization of metal-oxide-semiconductors (MOS) transistors and capacitors. Specifically, we carry out quantitative LIBS analysis on silicon oxide (SiO₂) thin-films of various thicknesses grown out of high-temperature moisture-free oxidation on industrial-grade Si wafers. The stoichiometric ratios of oxygen to silicon ([O]/[Si]) in various SiO₂ films are measured from LIBS analyses using an internal calibration technique. The results are verified against the benchmark analyses based on oxide layer thicknesses and laser-induced crater profile topographies from ellipsometry, scanning electron microscopy (SEM), atomic force microscopy (AFM), and profilometry measurements. The stoichiometric ratios of [O]/[Si] calculated from thickness and profilometry measurements are used to compare with our direct LIBS measurements. Our results indicate good agreement between the LIBS and profilom teric calculations results demonstrating the future capability of LIBS for thin film characterizations during their industrial processing.

Keywords: Laser Induced Breakdown Spectroscopy (LIBS); metal oxide semiconductors (MOS); thin-film SiO₂ layer; calibration-free; quantitative analysis.

1. Introduction:

The global energy demands in recent decades have led to significant manufacturing of semiconducting materials for solar cell applications. The efficiency and performance of solar cells strongly depend on the quality and characteristic properties of cost-effective semiconducting materials. The estimated global market share of US in the semiconductor sales is approximated at~ 50% with the most well-known materials of interest being Silicon (Si), and heterojunction materials such as Gallium arsenide (GaAs) and copper indium gallium diselenide (CIGS) [2]. Specifically, for multi-junction solar cells, the primary parameter that affects the efficiency of thin film semiconductors such as CIGS [CuIn1-_xGa_xSe₂] is the chemical composition of the absorber layer that determines the charge density and transport properties [3-5]. Variations in stoichiometric ratios of the main constituents change both the structural and electronic properties of these materials [3-5]. Typically, in the photovoltaic grade, Si impurities such as boron and oxygen change the electrical and mechanical properties significantly [6, 7]. While oxygen contents determine the defects that decrease the hardness of the Si wafers, it is essential for internal gettering [8, 9]. On the other hand, the efficiency of metal-oxide-semiconductors field-effect transistors (MOSFET) is largely driven by the property and reliability of the SiO_2 layer [10-12] wherein the main gate current leakage gets affected by the SiO_2 thickness [13].

To this end, various techniques such as secondary ion mass spectrometry (SIMS) [14], x-ray fluorescence (XRF) [15], inductively coupled plasma optical emission spectroscopy (ICP-OES) [16] have been used for the elemental analysis of thin-film semiconductor materials. Both SIMS and ICP-OES require extensive sample preparations and standards with ICP-OES specifically requiring the use of strong and harmful acids such as hydrofluoric acid. Such requirements render them as destructive methods [17] that cannot be employed for in-situ characterizations during most industrial manufacturing processes. But these methods are largely relied upon for post-processing off-line quality control in the industrial semiconductor sectors due to their ability to provide accurate measurements. On the other hand, only XRF can be operated in atmospheric conditions with minimal sample preparations [18]. However, XRF is not accurate for light elements measurements [19]. To this end, Fourier transform infrared (FTIR) spectrometry is commonly adopted as the industrial standard technique for oxygen concentration

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measurements in single crystalline Si wafers [20, 21]. However, it suffers from interference from free carrier absorption present in heavily doped substrates [22]. To overcome such issues, the semiconductor industry currently resorts to gas fusion analysis (GFA) to measure oxygen concentrations in heavily doped epitaxial substrate. But, GFA is a high destructive technique that has increasingly proven to be energy expensive and time consuming for the semiconductor manufacturing sector. In turn, such techniques fail to get implemented as rapid and on-line quantitative analyses tool for quality control purposes. Specifically, for MOS devices, the challenge rests on the accurate monitoring and tailoring of thin oxide layers on silicon or other semiconducting materials to tune the inversion layer charge accumulation, mobility and transport [23, 24]. Typical commercial-grade MOSFETS have been classically fabricated and extensively analyzed for gate oxide layers ranging between ~3 nm and 20-50 nm thicknesses approximately [23, 25]. To this end, the effect of gate oxide layer thicknesses on the charge mobility and capacitance in Si/SiO_2 systems become a critical parameter in the design of MOSFETS and have been investigated in great details over the years [26]. In light of the aforementioned challenges in the physical and chemical characterizations of thin films for semiconductor manufacturing sectors, there exits an acute need for a robust and efficient analytical technique that can accurately monitor and measure gate oxide film characteristics during the manufacturing and processing of metal/metal oxide semiconductor materials in order to tailor their wafer electronic properties for specific device fabrication requirements.

Laser Induced Breakdown Spectroscopy (LIBS) is a relatively non-destructive spectrochemical characterization technique that can address the aforementioned issues in a facile, rapid and yet effective manner. Being a non-contact technique that is not limited by the type or physical state of the analyte materials, LIBS provides advantages that are specifically attractive for large-scale manufacturing sectors. Furthermore, it is cost-effective and capable of carrying out rapid multi elemental analysis in one shot without compromising on the precision that makes it well-suited for in-situ implementation in industrial environments. In the past, LIBS has been employed for chemical analysis in highly diverse applications ranging from combustion [27-29], environmental/bio-hazard analysis [30-33], and intermetallic nanocatalysts [34, 35] to forensics [36], explosives

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detection/energetic studies [37-40], pharmaceutical [41], and biological/biomedical studies [42-44].

In recent years, there has been a growing interest in employing LIBS for material characterizations in the semiconductor industry – specifically, in regards to its choice as the future on-line quantitative analysis tool. To this end, Owens et al. [45] showed that LIBS can effectively measure heavy metal dopants in thin films. Seok et al. [46] had carried out semi-quantitative LIBS analysis to obtain the elemental ratios of constituents in thinfilm CIGS absorption layer. Added to this, Darwiche et al. [17] had extended the technique to quantify the extent of boron dopants in solar grade Si wafers using standard calibration curves. More recent efforts by Axente et al. [47] and Popescu et al. [48] have demonstrated the application of LIBS for the characterization of indium zinc oxide thin films, while Banerjee et al. [49] had employed the technique for multi-layered profiling of organic photovoltaic coatings. Specifically, the work by Axente et al. had compared recorded spectra with spectra computed based on local thermodynamics equilibrium to obtain atomic fraction of indium quantitatively. On the other hand, the work by Banerjee et al. had qualitatively shown that LIBS can be employed as a feedback control system during realtime laser scribing processes. It needs to be mentioned here that the calibration-free LIBS techniques proposed by Popescu et al. and Axente et al. resorted to computational models in deducing the quantitative elemental fractions, as compared to a methodology based on raw spectral data collected from analyte samples as developed by our previous works and implemented in the current study. Additionally, while the average thin films used in these studies were about 350 nm (Popescu et al. [48]) and 500 nm (Axente et al. [47]) thick, they did not attempt to report the lowest film thickness analytically detectable by LIBS.

In our efforts to investigate the feasibility of LIBS as the future analytical tool for rapid quantitative characterization of industrial grade thin film MOS, this study presents a calibration-free LIBS technique for quantifying thin-film oxide layers grown atop industrial-grade Si wafers. Commercial Si wafers, obtained were treated at different temperatures in order to synthesize and control the oxide layer thicknesses systematically. The ratio of ablated oxygen (O) to Si is measured based on the O and Si atomic emission signals by using the LIBS internal calibration technique developed in our earlier works [32, 34, 40]. The oxide layer thicknesses along with the crater sizes and profiles are measured

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using ellipsometry, scanning electron microscope (SEM), atomic force microscopy (AFM), and optical profilometer. Based on the profilometric measurements, the ratios of O to Si in the oxide layers and their film thicknesses on the Si wafers are estimated, and compared to the LIBS results.

1. Experimental details:

1.1 LIBS set-up:

The experimental set-up is illustrated in Fig. 1. The laser-induced plasma is generated on the sample target surface using a Q-switched Nd-YAG laser of nominal wavelength 1064 nm operating at 10 Hz repetition rates, 200 mJ/pulse energy, and a pulse width of 8 ns (Make: Insight Model: 122551-R). The laser is focused with a 25 mm diameter fused silica lens (focal length = 35mm) at the focal point on the target surface (Fig. 1). The spot size on the sample is set to 75 μ m diameter, which creates a plasma volume of ~1-2 mm³ approximately. Since the oxide layer is thin, only one laser shot per spot is collected. To improve the statistical average of the signal intensity for each analyte of interest, spectral data over approximately 100–150 spots are collected.

The plasma emission is collected with a fiber optic port carrying a pair of collimationfocusing lens and mounted at 45 degree collection angle (Fig. 1), which yields the optimum intensity from the plasma volume. The focused light is transmitted to a Czerny-Turner spectrometer (Make: Andor Technology; Model: Shamrock - SR-303i-A) with 1200 grooves/mm grating (resolution of 0.1 nm at 500 nm) and a nominal dispersion of 2.58 nm/mm. The slit width is fixed at 95µm for all experiments carried out here in order to have the optimum spectral line intensity and resolution. A time-gated intensified chargecoupled device (ICCD) detector array (1024x1024 CCD) (Make: Andor Technology; Model: DH334T-18U-E3) detects the spectral lines at the spectrometer exit focal plane. The time gating is synchronized with the laser Q-switch through an in-built timing generator in the ICCD camera set-up.

1.2 SiO₂ film synthesis:

Commercial silicon wafers, purchased from (UniversityWafer, Inc. ID#2444), were cut to approximately 1x1 cm² pieces. The wafers underwent dry thermal oxidation in laboratory furnaces at various temperatures in order to create and control the oxide layer thickness on

top of the wafer. Specifically, the wafers were baked for 20 minutes at oven temperatures ranging from 900^{0} C- 1000^{0} C with a 25^{0} C step, and with one final sample being baked at 1050^{0} C. The samples were tested using LIBS in an in-house built reactor cell while both helium (6 lpm) and argon (2 lpm) were purged into the cell as the buffer gas. The choice of flow rates is discussed in the results and discussion (2.1) To account for any unwanted background trace oxygen that might exist in the buffer gas or the control Si wafer samples, the background emission for O I is subtracted from the emissions from the baked Si wafer samples. Here, the background is considered as the emissions from the Si wafers without any oxide layer. To accomplish that, Si wafers are dipped into 10% hydrofluoric (HF) acid for 10 minutes.

1.3 LIBS methodology:

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In this study, atomic emission lines were selected from the NIST Atomic Energy Levels Data Center [1] according to their relative lines strength, and transition probabilities. To gain maximum accuracy in the results, the optimal gate delays are determined based on the variations of signal-to-noise ratios (SNR) as a function of gate delays for the respective elemental species. The SNR is calculated by measuring the peak signal value at the specific wavelength of interest for a spectral line divided by the noise of the spectra. The noise is defined as the root mean square over the baseline (~ over 40 pixel) adjacent to the analyte peak. The optimum SNR is used as the effective emission (I_{em}) for the population density calculations of the respective atomic species based on the Maxwell-Boltzmann distribution:

$$I_{em} = hc \frac{A_{ki}}{\lambda_{ki}} N_i \frac{g_k}{g_i} exp(-\frac{\Delta E_{ki}}{k_B T_{exc}})$$
(1)

where N_i, A_{ki}, λ_{ki} , g_k , g_i are atomic number densities at the lower energy state, Einstein's transition probability, emitted wavelength, and statistical weights for the higher (k) and lower (i) energy states respectively for the specific atomic transition of interest. ΔE_{ki} is the energy difference between the k and i states, and T_{exc} is the plasma excitation temperature at the optimal gate delays. Boltzmann and Planck's constant are indicated by k_B (1.38064×10⁻²³ m² kg/ s² K) and h (6.62607×10⁻³⁴ m²kg/s) respectively, and c (3x10⁸ m/s) is the speed of light in vacuum.

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The plasma excitation temperatures (T_{exc} , K) are determined from the slope of the fit to the linear Boltzmann plot of ln($I_{em}\lambda_{ki}/g_kA_{ki}$) as a function of normalized upper energy levels (E_k/k_B) (derived from Eq. 1) for multiple strong lines with sufficiently widespread upper energy levels. It is worth mentioning that the Boltzmann distribution (Eq. 1) is valid under the assumption of local thermodynamics equilibrium (LTE) conditions for the laser-induced plasma. The condition is established when the plasma quenching processes are predominantly collisional rather than radiative. Typically, such conditions are observed within longer times of plasma evolution (> 1 μ s) [34, 40, 50]. Thus, all our LIBS emission lines are collected at gate delays beyond 1 μ s. Finally, the population densities of the analyte species of interest and bulk matrix or background species are calculated at similar plasma conditions. As discussed in details in our earlier works [34, 40], any artifacts arising from the optical/experimental set-up or time dependent plasma characteristics is nullified through the use of the internal calibration technique. Based on this technique, the stoichiometric ratios between any two species X and Y of interest is estimated as:

$$R = \frac{[X]}{[Y]} = \frac{[N_i^{XI}/N_i^{BI}]_{@GD=t_1}}{[N_i^{YI}/N_i^{BI}]_{@GD=t_2}}$$
(2)

where, N_i^X , and N_i^B represents the lower energy state population densities of the analyte X, and the bulk background (matrix) species respectively. The aforementioned normalization of population densities of the desired analyte species are carried out at the optimal gate delays (GD) for the LIBS emissions of the respective species.

1.4 Crater characterizations:

Prior to sample ablations for the LIBS measurements, the oxide layer thicknesses on the respective heat-treated Si wafer samples are measured using an ellipsometer (Horiba Sci.; Model: UVISEL,). Subsequent to the LIBS measurements, the sample crater sizes are also measured using a profilometer (Veeco Wyko Non-contact Profilometer NT3300). Finally, we use atomic-force microscopy (AFM; Make: NT-MDT; Model: NTEGRA Spectra) in the tapping mode with silicon cantilever tips (Model: AppNano Model: ACTA) and scanning electron microscopy (SEM) in order to verify the crater sizes and profiles further.

2. Results and discussion:

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2.1 Optimizing measurements and plasma parameters:

In this study, atomic transition lines were chosen from the NIST Atomic Energy Levels Data Center [1]. For detailed quantitative LIBS analyses, the atomic emission lines of O I (777.19 nm) and Si I (288.16 nm) were selected. The specific choice of these lines was based on their robust transition probabilities and line strengths. To this end, Table 1 reports the detailed spectral data for the aforementioned emission lines of choice. Fig. 2. a & b show spectral emissions of O I (777.19 nm) and Si I (288.16 nm) lines respectively. It is observed in Fig. 2a that in the presence of 6 lpm helium mixed with 2 lpm argon, O I (777.19 nm) signal enhances significantly. Such observations were also previously reported by Mukherjee et al.'s work [40] where it was discussed that the addition of helium creates a less dense plasma that effectively reduces the background continuum noise. However, reducing plasma density eventually deteriorates the emission signals too. Therefore, based on the specific sample of study, an optimal value for the ratio of helium to argon flow rates provides the ideal signal enhancements. To this end, the O I (777.19 nm) SNR is plotted as a function of He/Ar flow rates ratio as the background buffer gas (Fig. 3). It is clearly observed that an optimal ratio of ~ 3 for He/Ar ratio in the background buffer gas produces the maximum O I SNR of ~18. For lower ratios, the plasma is relatively thick that subdues the signal enhancement while for greater ratios, the plasma becomes too thin and O I (777.19 nm) emission signal is decreased. Hence, for the rest of experiments, 2 lpm of argon and 6 lpm of helium was used as the buffer gas. Furthermore, in order to gain the maximum accuracy for the population density calculations from the LIBS signals, the optimal gate delays for each of the Si I and O I emission lines are determined by plotting their respective SNRs as a function gate delays. Fig. 4.a & b shows the temporal evolution of O I (777.19 nm) and Si I (288.16 nm) emission lines respectively at the gate widths of 2 µs. From these plots, the optimal gate delays are estimated as 3.5µs and 5µs for the respective O I and Si I emission lines. Based on the linear parts of Fig. 4c and 4d the gate width 14µs is used for both O I and Si I.

2.2 LIBS measurements:

Fig. 5 shows the characteristic emission spectra of O I (777.19 nm) acquired from the control (devoid of oxide layer) and specific samples baked at different temperatures. It can

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be observed from Fig. 5 that the control sample still shows some background emission at 777.19 nm, which is associated with the ubiquitous oxygen contents in the reactor cell that is almost impossible to eliminate in the absence of extremely high-end vacuum systems. However, the specific heated samples show distinct enhancement of the emissions at O I (777.19 nm) as compared to the background emission which allows for background subtractions. Moreover, the O I (777.19 nm) emissions increase systematically as the oxide layers of the specific samples increase due to their baking at increasing temperatures. This increasing oxide layer thickness is also confirmed by ellipsometry measurements in the next section. The plasma excitation temperatures, T_{exc} required for the population density calculations are estimated from linear Boltzmann plots as explained in the experimental sections earlier. Here, the linear Boltzmann plots are constructed at the two different time delays (3.5µs and 5µs) of the plasma evolution using a series of Ar I lines with known spectral properties as listed in Table 2. Fig. 6a and 6b illustrate the Boltzmann plots acquired at 3.5 μs and 5 μs gate, whose slopes indicate Texc=17,126 K and 15,166 K respectively. Similarly, the Texc values for all the different heat-treated samples are estimated and reported in Table 3. The estimated excitation temperatures are employed to calculate the relative population densities of oxygen $(N_i^{O I})$, silicon $(N_i^{Si I})$ and argon (N_i^{Ar}) ^I) from the respective atomic emission lines of O I (777.19 nm), Si I (288.16 nm) and Ar I (series of lines indicated in Table 2). Using Ar as the bulk species in the plasma for the normalization shown in Eq. 2, the ratio of oxygen to silicon ([O]/[Si]) is calculated as:

$$R = \frac{[O]}{[Si]} = \frac{\left[\frac{N_{i}^{O \ I \ (777.19 \ nm)}}{N_{i}^{Ar \ I}} \right]_{@\ GD = 3.5 \mu s}}{\left[\frac{N_{i}^{Si \ I \ (288.16 \ nm)}}{N_{i}^{Ar \ I}} \right]_{@\ GD = 5 \mu s}}$$
(3)

The results for [O]/[Si] (LIBS) ratios are tabulated in Table 3. In order to compare the results obtained from the quantitative LIBS analyses, the amounts of ablated materials are also estimated by characterizing the oxide layer thicknesses and crater profile using ellipsometry, SEM, AFM and profilometer measurements. Based on the profilometric calculation, discussed in the following sections, the amount of ablated oxygen and silicon are estimated, and used for calculating [O]/[Si] (ablation) ratios in Table 3 from the crater ablation analyses that will be presented in the following sections.

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2.3 Oxide layer thickness measurements:

As the first step toward characterization of the amount of elemental species in the laser ablated crater from the LIBS measurements, the initial oxide layer thicknesses from the samples are measured using ellipsometry. A classical dispersion model based on SiO_2 on top of silicon substrate is used for curve fitting, and observed that model and the acquired data are in excellent agreements as shown in Fig. S1 in the supporting information. For each oxidation temperature, four different pieces and on each piece four different spots are analyzed for the ellipsometry measurements. The oxide layer thicknesses are reported in Table 4. As expected, the oxide layer thickness grows further as the oxidation temperature increases. Moreover, the small standard deviations indicate that the oxide layer is uniform across the samples surface.

2.4 Crater measurements:

Fig. 7 shows the SEM image for the crater obtained from the ablation of a representative sample (Si wafer baked at 950^{0} C). The image indicates that ablation spot diameter is approximately 200µm, while the crater depth can be qualitatively observed to be shallow. In order to obtain the crater depth quantitatively, the samples are measured using a profilometer (Fig. 8a). The profilometer result also indicates that the ablation spot diameter is 200µm, while the crater depth is approximated to be ~2.5µm. For a secondary verification of the crater depth, the ablation spots are also topographically imaged using AFM in tapping mode with slow scan rate. Due to the AFM instrument limitations, only a 100µmx100µm window can be measured. Hence, the measured window was chosen such that it spanned over ~30µm of the spot edge and ~70µm of the crater. As seen from the z-profile of the AFM topography image for the selected window in Fig. 8.b, the crater depth can be confirmed to be ~ 2.5 as also corroborated from our profilometer measurements.

2.5 LIBS and profile calculations comparison:

Based on the oxide layer thickness, ablation spot diameter and the crater depth, two simple geometric models, namely cylindrical and the frustum of a cone, are constructed to calculate the amount of ablated O and Si, and construct the ablated [O]/[Si] ratio as depicted in Fig. 9.a & b. Based on these geometries for the ablated volume, [O]/[Si] ratio can be determined as:

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$$\frac{[O]}{[Si]} = \frac{2(\frac{\rho_{SiO_2}}{Mw_{SiO_2}})V_{SiO_2}}{(\frac{\rho_{SiO_2}}{Mw_{SiO_2}})V_{SiO_2} + (\frac{\rho_{Si}}{Mw_{Si}})V_{Si}}$$
(4)

where V_{SiO2} and V_{Si} are the ablated volumes of oxide layer and silicon substrate respectively, which can be expressed by the oxide layer thickness (δ_{SiO2}), and crater depth (h). Mw and ρ are molecular weights and densities respectively and are tabulated in Table 5 for Si and SiO₂. Fig. 10 illustrates the results for the ratio of [O]/[Si] as obtained from LIBS plotted as a function of the respective ratios obtained from crater profile or ablation characterizations based on the assumption of a cylindrical and cone frustum geometries. The red and the blue dotted lines represent the linear fits to the calculated values for the cylindrical and the cone-frustum geometries respectively. It can be observed that the LIBS results are relatively in good agreements and indicate a linear correlation with the profilometric calculations. The error bars are calculated based on the error propagation for both methods due to uncertainties in plasma temperature, shot-to-shot variation in laserinduced, and standard deviations in crater measurements. The deviations from the profilometric calculations can be related to the errors in measuring crater profiles and simplistic geometric models, which change the amount of ablated silicon. Analyzing the crater profile of the ablation spot indicates that the amount of ablated silicon based on the cylinder geometry is greater than the actual ablated silicon, while the cone-frustum underestimates the amount of ablated silicon. In fact, the ideal one-to-one linear correlation between the [O]/[Si] ratios from LIBS measurements and the profilometric analyses appears to lie within the two sets of data for the cylindrical and the cone-frustum profile geometry cases (i.e. the red and the blue dotted linear fits). Furthermore, the errors bars on the [O]/[Si] ratios from the LIBS measurements commensurate with the spread in the data for the aforementioned two cases thereby indicating that the deviations are within the experimental margins of error. Finally, we present the LIBS calibration for the quantitative determination of thin-film oxide layer thickness on Si wafer by plotting the O I subtracted signal as a function of oxide layer thicknesses (from ellipsometry measurements) in Fig. 11. Based on the slope of the calibration curve, the minimum distinguishable oxide layer (δ_{\min}) that can be estimated form our current LIBS set-up is found to be:

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$$\delta_{\min} = \frac{3\sigma_{\rm B}}{\rm S} \tag{5}$$

where, σ_B is the standard deviation of the background and S is the slope of the calibration curve. This give the estimated thinnest detectable oxide layer using LIBS as $\delta_{min} \sim 10.51 \pm 1.44$ nm.

3. Conclusion:

We have employed LIBS for calibration-free quantitative analysis of thin-film silicon dioxide layers with various thicknesses synthesized via dry thermal oxidation of Si wafers. To this end, an internal calibration technique was applied that involved effective normalization of the analyte species densities by bulk species densities in the laser-induced plasma. The species densities of O (analyte) and Si (bulk) for the SiO₂ films in this case were estimated from the respective atomic emission lines at identical plasma conditions to generate the ablated oxygen to silicon ([O]/[Si]) ratio. The results from our LIBS analyses were corroborated by the [O]/[Si] ratios directly calculated from the ablated spots by measuring the oxide layer thicknesses and crater profiles using ellipsometry, SEM, AFM and profilometry techniques. The results are in good agreement with each other within the margins of experimental uncertainty. The small deviations of the LIBS results from the profilometric calculations were attributed to the errors in the depth measurements and assumed geometry of the crater profile. Based on our measurements, a calibration curve for the LIBS analysis was established that indicated the detection limit for the thinnest oxide layer thickness to be ~ 10.5 ± 1.4 nm. Results from the current work provide future directions towards the promising potential of LIBS as an analytical tool for rapid screening of oxide layers in MOS transistors/capacitors, and more generally, for surface composition analysis of thin-film materials.

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Tables:

Table 1. Atomic spectral database [1] for Si I and O I emission lines used for the population density calculations during quantitative analysis of ablated spots.

Species	Wavelength, λ _{ki} (nm)	Transition probability, A _{ki} (10 ⁶ 1/s)	Upper energy level, E _k (eV)	Lower energy level, E _i (eV)	g _k	gi
Si I	288.16	218	5.082	0.781	3	5
ΟΙ	777.19	36.9	10.741	9.146	7	5

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Species	Wavelength, λ _{ki} (nm)	Transition probability, A _{ki} (10 ⁶ 1/s)	Upper energy level, $E_k (eV)$	Lower energy level, E _i (eV)	g _k	gi
Ar I	415.859	1.40	14.528	11.548	5	5
Ar I	420.068	0.967	14.499	11.548	7	5
Ar I	425.936	3.98	14.738	11.828	1	3
Ar I	427.217	0.797	14.525	11.624	3	3
Ar I	433.356	0.568	14.688	11.828	5	3
Ar I	696.543	6.39	13.328	11.548	3	5
Ar I	706.722	3.80	13.302	11.548	5	5

Table 2. Atomic spectral database [1] for different Ar atomic emission lines used for the plasma excitation temperature calculations at $3.5\mu s$, and $5\mu s$ for spectral analysis of ablated spots.

Sample treated	[O]/[Si]	[O]/[Si]	[O]/[Si]	Plasma excitat	tion temp. T _{exc}
temperature	(Ablation-Cylinder)	(LIBS)	(Ablation-Cone frustum)	(K)	
				at 3.5µs	at 5µs
$900^{0}{ m C}$	0.022	0.025	0.046	17,125±3833	15,166±2893
925 ⁰ C	0.027	0.047	0.047	17,803±3706	15,849±3235
$950^{0} \mathrm{C}$	0.032	0.045	0.079	$17,372\pm2898$	16,427±3637
975 ⁰ C	0.051	0.052	0.102	20,802±4162	$17,580 \pm 4550$
$1000^{0} \mathrm{C}$	0.057	0.098	0.133	$21,535\pm4692$	20,311±5826
$1050^{0}{ m C}$	0.078	0.097	0.179	$20,142\pm4620$	$18,255 \pm 4628$

Table 3. [O]/[Si] ratios for various samples and respective plasma excitation temperatures at 3.5µs and 5µs.

Table 4.	The oxide l	ayer thicknesses	for various	oxidation	temperature	obtained by	ellipsomery
measurer	nents.						

Sample treated temperature	900 ⁰ C	925 ⁰ C	950 ⁰ C	975 ⁰ C	$1000^{0} \mathrm{C}$	1050 ⁰ C
Thickness (nm)	50.94±0.25	64.63±0.19	75.62±0.05	122.10±0.01	136.93±0.35	188.82±0.37

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Table 5.	Si and SiO2 properties used for calculating
[O]/[Si] r	atio from profilometric model.

Properties	Molecular weight, Mw (g/mol)	Density, ρ (g/cm ³)	
SiO ₂	60.083	2.65	
Si	28.085	2.33	

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Figures:



Fig. 1. Schematic for LIBS experimental set-up used for characterizing samples.



Fig. 2. Spectral emission signature for (a) O I (777.19 nm) with and without helium in the buffer gas, and (b) Si I (288.16 nm) lines at the respective gate delays of 3.5μ s, and 5μ s.



Fig. 3. Optimizing oxygen SNR with respect to ratio of helium to argon flow rates.

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Fig. 4. Temporal evolution of signal-to-noise ratio for (a) O I (777.19 nm), and (b) Si I (288.16 nm) over gate delyas. The optimal gate delays were determined to be 3.5μ s and 5μ s for the peak signal-to-noise ratio for O I (777.19 nm), and (b) Si I (288.16 nm) respectively. (c) temporal evolution of signal-to-noise ratio for (c) O I (777.19 nm), and (d) Si I (288.16 nm) over gate width.



Fig. 5. Spectral emission signature for O I (777.19 nm) for samples with various oxide layer thickness.



24Fig. 6. The linear Boltzmann plots generated from Ar I lines listed in Table 2, and used for Texc calculations at the $\frac{25}{26}$ respective gate delays of 3.5µs and 5µs.

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Fig. 7. SEM image of the ablated spots used for measuring ablated spot diameter.

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Fig. 8. Depth profile measurement for the ablated spots using (a) optical profilometer, and (b) AFM.



Fig. 9. The simple geometric models (a) cylinder and (b) cone-frustum showing ablation volume used for calculating [O]/[Si] ratio.



Fig. 10. Comparison between [O]/[Si] ratio directly measured by LIBS, and calculated by characterizing ablated spot.



Fig. 11. Calibration curve for O I (777.19 nm) SNR as a function of oxide layer thickness.

Signal (a.u.)
Fig. 1 funct