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Soft X-ray Spectroscopy of a Complex Heterojunction in High-Efficiency Thin-Film Photovoltaics: Intermixing and Zn Speciation at the Zn(O,S)/Cu(In,Ga)Se2 Interface

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ABSTRACT: The chemical structure of the Zn(O,S)/Cu(In,Ga)Se2 interface in high-efficiency photovoltaic devices is investigated using X-ray photoelectron and Auger electron spectroscopy, as well as soft X-ray emission spectroscopy. We find that the Ga/(Ga+In) ratio at the absorber surface does not change with the formation of the Zn(O,S)/Cu(In,Ga)Se2 interface. Furthermore, we find evidence for Zn in multiple bonding environments, including ZnS, ZnO, Zn(OH)2, and ZnSe. We also observe dehydrogenation of the Zn(O,S) buffer layer after Ar+ ion treatment. Similar to high-efficiency CdS/Cu(In,Ga)Se2 devices, intermixing occurs at the interface, with diffusion of Se into the buffer, and the formation of S—In and/or S—Ga bonds at or close to the interface.

KEYWORDS: chalcopyrite thin-film solar cell, chemical structure, alternative buffer layers, Zn(O,S), X-ray photoelectron spectroscopy, X-ray emission spectroscopy

INTRODUCTION

Cu(In,Ga)Se2 (CIGSe) thin-film photovoltaic devices have recently achieved a world-record efficiency of 22.6% on a laboratory scale utilizing a CdS buffer layer.1 While CdS-based CIGSe devices have traditionally dominated the record efficiencies for this materials class,2 other groups also report high conversion efficiencies (up to 22.0%) utilizing an alternative buffer layer based on Zn(O,S).3–8 An understanding of the interface formation is crucial for optimizing these buffer layers and the buffer/absorber interface. While several studies have been published showing theoretically and experimentally derived interfacial band alignments,9–14 there is a need to better understand the chemical interactions at the buffer/absorber interface, as this information can aid in deliberately tailoring the electronic band alignment. We note that current state-of-the-art Zn(O,S)-based devices feature a flat conduction band alignment.14

Previous studies have revealed a S/Se intermixing at the CdS/CIG(S)Se interface of high-efficiency thin-film devices with a chemical bath-deposited buffer layer.15–18 Only few studies report on intermixing at the heterojunction between Zn(O,S) and chalcopyrites.19,20 To gain better insights into the formation of the Zn(O,S)/CIGSe buffer layer and the chemical interactions at the interface in current state-of-the-art devices, we employ X-ray photoelectron spectroscopy (XPS), X-ray-excited Auger electron spectroscopy (XAES), and synchrotron-based soft X-ray emission spectroscopy (XES) on Zn(O,S)/CIGSe samples with varying buffer layer thickness.

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Due to the presence of this adsorbate layer, all adsorbate species, as well as most of the Na and a linear background. For quantification, XPS peaks were analyzed by fitting with Voigt functions, fixed spin–orbit splitting, coupled Gaussian and Lorentzian widths, and a linear background. For quantification, invariant elastic mean free paths (IMFPs) were determined by the QUASES software.

**EXPERIMENTAL SECTION**

A full description of the samples can be found in ref 14. The sample set consists of a CIGSe three-stage bare absorber (nominal Ga/(Ga + In) ratio of 0.3) and three Zn(O,S)/CIGSe interfacial samples produced with varying Zn(O,S) chemical bath deposition (CBD) times (5, 10, and 22.5 min).

XPS and XAES were conducted at UNLV, while XES was performed at the Advanced Light Source (ALS), Lawrence Berkeley National Laboratory. XPS measurements were taken with a SPECS PHIIBOS 150 MCD electron analyzer using Mg and Al Kα radiation and calibrated according to ref 23. The XES spectra were taken with the SALSA endstation on beamline 8.0.1 at the ALS, utilizing a variable line-space grating (VLS) spectrometer. The spectra were calibrated using the prominent emission features of CdS.25 The base pressure for the XPS and XES measurements was below 5 × 10⁻¹⁰ and 1 × 10⁻⁹ mbar, respectively.

The samples were briefly air-exposed (less than 5 min) at NREL before being packaged in a vacuum-sealed container. At UNLV, the samples were immediately introduced into an inert gas-filled glovebox, mounted on a sample holder, and transferred into the ultrahigh vacuum system. The samples were measured “as-received” and also after a low-energy (50 eV) Ar⁺ ion treatment (two subsequent 60 min treatments for CIGSe, and three subsequent 20 min treatments for each of the 5 and 22.5 min Zn(O,S)/CIGSe samples) at low incidence angle.26,27 XPS peaks were analyzed by fitting with Voigt functions, fixed spin–orbit splitting, coupled Gaussian and Lorentzian widths, and a linear background.26 For quantification, inelastic mean free paths (IMFPs) were determined by the QUASES software.

**RESULTS AND DISCUSSION**

XPS survey spectra of the bare CIGSe absorber and the 5 and 22.5 min Zn(O,S)/CIGSe samples series are shown in Figure 1. The respective “as-received” spectra are shown in black, while the “ion-treated” spectra are shown in red. The pertinent CIGSe and ZnO spectra show Auger lines of Cu, In, Ga, Se, Zn, and S, as well as the Na lines, and are labeled, in addition to the peak associated with C. For the as-received CIGSe absorber, the O 1s peak is particularly pronounced, especially when the photoionization cross section is taken into account.30 Due to the presence of this adsorbate layer, all high-binding (i.e., low-kinetic) energy peaks in the survey spectrum (e.g., Ga 2p, Cu 2p, In MN, and Zn 2p) are suppressed in intensity. The low-energy ion treatment removes the majority of the adsorbate species, as well as most of the Na surface species, and thus the surface-sensitive high-binding energy peaks gain in intensity. In Figure 1 and the subsequent analysis, data after the first 20 min treatment is shown/used in order to minimize ion beam influences. In fact, as will be discussed later, the two subsequent ion treatments of the Zn(O,S)/CIGSe samples induced signs of surface alteration (dehydrogenation of the buffer). This indicates that Zn(O,S)—or more precisely the buffer hydroxide component (see below)—is more susceptible to changes due to low-energy ion treatments (and other irradiation) than the CIGSe absorber, CdS, or ZnO.26,27,32,33 As the Zn(O,S) thickness is increased, Figure 1 shows that all peaks from the CIGSe absorber surface are attenuated, as expected. Small CIGSe-related core-level peaks are detected in the spectra of the thin Zn(O,S)/CIGSe sample (e.g., the In 3d and Se 3d peaks in Figure 1, center). In contrast, the 22.5 min Zn(O,S) sample shows no evidence of absorber-related peaks, suggesting that it is a continuous layer.

The Na peak intensity for the CIGSe bare absorber decreases with ion treatment, in parallel to a reduction of the oxygen and carbon signals [we note that the carbon signal for the Zn(O,S) layers is quite sizable, which we assign to an incorporation during the CBD process].20 To gain further insight into the chemical state of Na on the CIGSe surface and the cause of this intensity decrease, the modified Auger parameters α′ of Na were calculated. Figure 2 shows the Wagner plot for the CIGSe absorber surface (red, ion-treatment times as indicated), along with relevant references (black).35 The modified Auger parameter of α′ of Na is calculated by summing the binding energy of the Na 1s core level and the kinetic energy of the KL2,3L2,3 Auger peak, and information about the chemical state is gained by comparing with reference data along three axes: the Na 1s binding energy (abscissa), the Na KLL kinetic energy (ordinate), and α′ (diagonal and right ordinate). The location of the CIGSe data on the Wagner plot is indicative of Na in an oxidized chemical environment. It is clearly different from metallic Na, but close to a large variety of O and/or C containing Na compounds [for example, we find evidence for a carbon species at ~289.7 eV, which would be in agreement with the presence of a (sodium) carbonate]. Thus, upon ion treatment, Na is likely to be desorbed along with the surface adsorbates. Based on earlier studies, this is not unexpected for air-exposed CIGSe surfaces and is also assumed to happen in the chemical bath solution.

![Figure 1](image-url)

Figure 1. XPS survey spectra of the untreated (black) and 50 eV Ar⁺-ion-treated (red) Zn(O,S)/CIGSe sample series: CIGSe bare absorber (bottom), 5 min Zn(O,S) (center), and 22.5 min Zn(O,S) (top).
Core-level peaks from the CIGSe absorber can be detected in the spectra of the thin Zn(O,S)/CIGSe sample, indicating that the 5 min CBD of Zn(O,S) results in a layer that is inhomogeneous and/or thinner than the region from which XPS information can be derived (note that XPS signals are governed by an exponential attenuation function, not a “box” function associated with a specific information depth). The absorber peaks detected for Cu, In, and Ga are of low intensity, while, in comparison, the Se 3d peak is larger, indicating the possibility of Se diffusion into the Zn(O,S) layer, as will be discussed now.

For the purpose of studying Se diffusion at the interface between Zn(O,S) and CIGSe, we also include the Zn(O,S)/CIGSe sample of intermediate thickness (10 min CBD). Figure 3 shows XPS spectra of the Se 3d peak (left) and the Ga 3d/In 4d peak (right) for the CIGSe bare absorber and the Zn(O,S)/CIGSe samples of increasing thickness. Due to their similar kinetic energies, the Se 3d, Ga 3d, and In 4d peaks are expected to have similar inelastic mean free paths (IMFPs), allowing the attenuation factors of these peaks to be compared as an initial step. The Se peak is strongly attenuated as the Zn(O,S) CBD time increases, but nevertheless it is still detectable even after the standard deposition time of 22.5 min. In contrast, the Ga 3d and In 4d peaks of the 10 min Zn(O,S)/CIGSe and 22.5 min Zn(O,S)/CIGSe samples are extremely small (if present at all), even when magnified 100x (note that the Ga 3d/In 4d peaks lie on the onset of the O 2s peak, at ≈26 eV, giving rise to the steep background observed for the 10 and 22.5 min sample). The detection of a Se signal even after 22.5 min of Zn(O,S) CBD suggests an outdiffusion of Se during the CBD process, similar to the CdS/CIGSe and CdS/CIGSSe interfaces. All other absorber-related core-level and Auger peaks (not shown) were analyzed in a similar fashion, but no absorber-related peaks were visible for the 10 and 22.5 min Zn(O,S) CBD sample (note that these peaks are governed by shorter attenuation lengths due to their lower kinetic energy and will be discussed in greater detail below).

In order to analyze the possibility of Se, Ga, and/or In diffusion more quantitatively, Figure 4 presents an “effective Zn(O,S) layer thickness” as a function of the Zn(O,S) CBD time. The nominal thickness values (determined by Scanning Electron Microscopy images of corresponding cross sections at NREL) are shown as black data points and a best-fit line, while the effective thicknesses derived from the Se 3d (blue), Ga 3d (red, 5 min), In 4d (green, 5 min), Ga 3d/In 4d combined (pink, 10 min), and nominal thickness (black) as a function of Zn(O,S) CBD time. The 10 min nominal thickness and Ga 3d/In 4d data were shifted slightly along the abscissa to differentiate between the data points.
to be below the nominal thickness line in the case of diffusion into the buffer layer. The effective thickness derived for a given peak is calculated using the following:

$$d_{\text{eff}} = \lambda \ln(I_0/I)$$

The IMFP is represented by $\lambda$, $I_0$ is the peak intensity in the bare absorber, and $I$ is the peak intensity in the corresponding Zn(O,S)/CIGSe sample. We find that the Se 3d-based effective layer thickness lies significantly below the nominal thickness line, even when taking the error bars into account. Thus, the attenuation of the Se signal is lower than expected, suggesting that Se is diffusing into the buffer layer during the CBD process. In contrast, for the 5 min sample, the Ga 3d and In 4d effective thicknesses are larger and on (Ga) or very close (In) to the nominal line. For the 10 min sample, we note that the Ga 3d/In 4d region also includes the Mg K$\alpha$ excitation of the O 2s line, which would give rise to a satellite approximately 9 eV lower than the O 2s peak ($\sim 17$ eV), and/or contributions from inelastically scattered Zn 3d electrons (peak at $\sim 10$ eV). Nevertheless, because the peak in the Ga 3d/In 4d region is clearly present for the 10 min sample, we show the effective layer thickness based on the attenuation of the combined Ga 3d/In 4d peak (pink), addressing the possible O 2s or Zn 3d area contributions by including a larger and asymmetric error bar. The effective layer thickness is again close to the nominal thickness line, suggesting that there is no significant diffusion of In or Ga into the buffer. For the 22.5 min sample, we argue that there is no detectable Ga 3d/In 4d peak, noting the absence of the (normally dominant) In 3d$_{5/2}$ peak. The finding of Se outdiffusion into the Zn(O,S) buffer layer is reminiscent of the CdS/CIGSe interface, in which a pronounced S—Se exchange at the interface is found. In the present case of a Zn(O,S) buffer, the diffusion of Se into the buffer layer leads us to speculate that in the proximity of the interface most likely Zn—Se bonds are formed.

Figure 5 shows fits of the Ga 3d/In 4d peaks of the bare CIGSe absorber and the 5 min Zn(O,S) sample in order to see if the Ga/(Ga+In) ratio at the absorber surface changes with the addition of the buffer layer. The peaks were fit with a linear background and Voigt profiles, using coupled Gaussian widths for all components, and coupled but separate Lorentzian widths for In and Ga, respectively. The ratios of the spin–orbit split peaks were fixed according to their multiplicity, along with their respective peak separation, 0.86 eV for In$^{14,42}$ and 0.46 eV for Ga.$^{33}$ The data points are shown with black dots, the In 4d components in green, the Ga 3d components in blue, and the resulting fit in red. The residual of the fit is shown below each spectrum (purple). We note that these shallow core levels already possess some band character, and thus the quality of the fit is surprisingly high, especially given all the above-mentioned boundary conditions included in the fit. The surface Ga/(Ga+In) ratio for the CIGSe absorber and 5 min Zn(O,S)/CIGSe samples are 0.33 and 0.32 ($\pm 0.10$), respectively, indicating no change in the surface ratio with the addition of the buffer layer. Having thus gained a detailed description of the absorber surface, we can now take a closer look at the overlayer and its contributions to the interface formation.

As mentioned previously, the 5 and 22.5 min Zn(O,S)/CIGSe samples were ion treated in three 20 min increments. Figure 6(left) shows XPS spectra of the 22.5 min Zn(O,S)/CIGSe O 1s peak as a function of ion treatment time. Indeed, we see that with the first 20 min treatment, the O peak is reduced (partial removal of adsorbates) and the main peak position is identical to the untreated surface. However, with each subsequent treatment, the peak shifts toward lower BE and the shape changes as well. This is also true for the Zn and S peaks (not shown): the untreated and 20 min treated surface peak positions are identical, and with each subsequent treatment, the peak shifts to lower BE. The broadness and shape of the O 1s peak suggest that there are multiple chemical species of O in the (untreated) Zn(O,S) film. On the right of Figure 6, a fit analysis of the peak (as a function of ion treatment) shows that at least three species are present at the untreated surface, and at least two O species after each treatment. The peaks were normalized to peak height in order to be consistent.
to easily visualize changes in peak component ratios. The peaks were fit with a linear background, identical Gaussian widths, identical Lorentzian widths, and fixed positions for all species. The residuals shown below the spectra indicate that the quality of the fit is quite high for the ion-treated surfaces. In contrast, the untreated O 1s peak shows a characteristic oscillation, which can be reduced by decoupling the Lorentzian width of this fit from that of the other three spectra. We thus derive that the spectrum of the untreated surface is a convolution of at least three species. Because there is a reduction in the O 1s peak intensity with the first ion treatment step, we believe that such additional species need to be ascribed to surface adsorbates. Based on their binding energies, the three components for the peak prior to ion treatment are assigned to H$_2$O, Zn(OH)$_2$, and ZnO.$^{35,44}$ For the peaks after ion treatment, the components are assigned to Zn(OH)$_2$ and ZnO only. The presence of Zn(OH)$_2$ suggests that the dehydrogenation of the Zn(O,S) layer during deposition is incomplete, indicating that a better description of this layer would be “Zn(O,H,S)”. Also, it cannot be ruled out that Zn(OH)$_2$ might be formed at the surface during the (brief) air exposure. The fits of the O 1s peaks show that, with increasing ion treatment time, the OH/(O+OH) ratio changes—a reduction in Zn(OH)$_2$ and an increase in ZnO suggests that the hydroxide component of the Zn(O,S) buffer surface is more susceptible to the low-energy Ar$^+$-ion treatment than, e.g., the ZnO surface commonly used as a window layer in CIGSe devices. This beam-induced dehydrogenation is not unexpected, as similar results have been found utilizing intense X-rays and electron flood gun irradiation on Zn(OH)$_2$-rich samples.$^{32,33}$ Nevertheless, we use data obtained after the first 20 min ion treatment for further analysis because this surface represents the best compromise between reduced contributions from surface adsorbates and minimal ion beam damage.

The Zn $2p_{3/2}$ and S $2p$/Se $3p$ spectra for the 5 and 22.5 min Zn(O,S)/CIGSe samples are shown in Figure 7, along with the binding energy markers for several references.$^{35,44}$ Because the binding energies of the Zn references overlap (left), and because the observed peak is rather broad, it does not allow for an unambiguous assignment of the different species. This is not necessarily surprising—so far, our analysis has suggested three local bonding partners for Zn, namely O and OH from the O 1s peak fit and Se from the diffusion analysis. Furthermore, bonding to sulfur is also expected for the Zn(O,S) film surfaces. The S 2p/Se 3p spectra (right) show (a) the presence of sulfur in a sulfide environment, as expected, (b) the presence of Se for the 5 min samples (as discussed above; note that the Se 3p peak is much weaker than the Se 3d peak, and hence it is not seen for the 22.5 min sample), and (c) no evidence for sulfates on the surface.

Figure 7. XPS spectra of the Zn $2p_{3/2}$ peak (left) and S $2p$/Se $3p$ peaks (right) for the 5 and 22.5 min Zn(O,S) samples. Reference peak positions$^{35}$ are indicated above both peaks.

Figure 8. S L$_{2,3}$ and Se M$_{2,3}$ emission of the Zn(O,S)/CIGSe interface as a function of CBD time and a ZnS reference. Multiplication factors are shown in parentheses.

The residuals shown below the spectra indicate that the quality of the fit is quite high for the ion-treated surfaces. In contrast, the untreated O 1s peak shows a characteristic oscillation, which can be reduced by decoupling the Lorentzian width of this fit from that of the other three spectra. We thus derive that the spectrum of the untreated surface is a convolution of at least three species. Because there is a reduction in the O 1s peak intensity with the first ion treatment step, we believe that such additional species need to be ascribed to surface adsorbates. Based on their binding energies, the three components for the peak prior to ion treatment are assigned to H$_2$O, Zn(OH)$_2$, and ZnO.$^{35,44}$ For the peaks after ion treatment, the components are assigned to Zn(OH)$_2$ and ZnO only. The presence of Zn(OH)$_2$ suggests that the dehydrogenation of the Zn(O,S) layer during deposition is incomplete, indicating that a better description of this layer would be “Zn(O,H,S)”. Also, it cannot be ruled out that Zn(OH)$_2$ might be formed at the surface during the (brief) air exposure. The fits of the O 1s peaks show that, with increasing ion treatment time, the OH/(O+OH) ratio changes—a reduction in Zn(OH)$_2$ and an increase in ZnO suggests that the hydroxide component of the Zn(O,S) buffer surface is more susceptible to the low-energy Ar$^+$-ion treatment than, e.g., the ZnO surface commonly used as a window layer in CIGSe devices. This beam-induced dehydrogenation is not unexpected, as similar results have been found utilizing intense X-rays and electron flood gun irradiation on Zn(OH)$_2$-rich samples.$^{32,33}$ Nevertheless, we use data obtained after the first 20 min ion treatment for further analysis because this surface represents the best compromise between reduced contributions from surface adsorbates and minimal ion beam damage.

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Figure 8. S L$_{2,3}$ and Se M$_{2,3}$ emission of the Zn(O,S)/CIGSe interface as a function of CBD time and a ZnS reference. Multiplication factors are shown in parentheses.
To gain insights into the component(s) contributing to the S and 10 min spectra, the 22.5 min spectrum [i.e., the S L$_{2,3}$ emission of the Zn(O,S) overlayer] was subtracted after normalizing all spectra to overall peak area. The resulting difference spectra are presented in Figure 9 a). The 5 min intermixing of S at the interface, most likely in an S—In and/or S—Ga environment. We find multiple chemical environments of Zn, best described by (a mixture of) ZnO, Zn(OH)$_2$, ZnS, and ZnSe. There is no evidence for sulfates in the Zn(O,S) layer (photoinduced or otherwise), but we do find evidence for Ar$^{+}$ ion beam-induced dehydrogenation of the Zn(O,S) layer. The resulting chemical interactions during the Zn(O,S)/CIGSe interface formation are found to be similar to those at the CdS/CIGSe interface$^{15,17,18}$ but feature a higher degree of complexity with respect to the local chemical environment of the group II component.

**CONCLUSIONS**

The Zn(O,S)/CIGSe interface has been investigated using XPS, XAES, and XES to investigate the chemical structure. Detailed analysis of the bare CIGSe absorber and Zn(O,S)/CIGSe samples of varying CBD times allowed for a comprehensive analysis of the chemical interactions at this interface. We find evidence for an upward diffusion of Se into the buffer layer, most likely in a Zn—Se environment, and

![Figure 9](image-url)

**REFERENCES**


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**Notes**

The authors declare no competing financial interest.

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