Electronic and compositional properties of the rear-side of stoichiometric CuInSe2 absorbers

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Abstract
In-depth understanding and subsequent optimization of the contact layers in thin film solar cells are of high importance in order to reduce the amount of nonradiative recombination and thereby improve device performance. In this work, the buried MoSe2/CuInSe2 interface of stoichiometric absorbers is investigated with scanning tunneling spectroscopy and Kelvin probe force microscopy combined with compositional measurements acquired via photo-electron spectroscopy after a mechanical lift-off process. We find that the local density of states, as measured with scanning tunneling spectroscopy, is similar to the front-side of ultra-high vacuum annealed CISe absorbers. The grain boundaries exhibit a weak upward band bending, opposite to Cu-poor CuGaSe2, and we measure an increased Cu accumulation at the rear CISe surface compared to the bulk composition and a non-zero concentration of Cu on the Mo-side. Grazing incidence X-ray diffraction measurements corroborate that a small amount of a Cu$_x$Se secondary phase is present at the MoSe2/CuInSe2 interface in contrast to reports on Cu-poor material. Our findings shed new light into the complex interface formation of CuInSe2-based thin film solar cells grown under Cu-rich conditions.

KEYWORDS
CuInSe2, Kelvin probe force microscopy, scanning tunneling microscopy

1 INTRODUCTION
Cu(In,Ga)Se2 (CIGSe) thin film solar cells offer high power conversion efficiencies (PCE),1 low energy payback time, and long-term stability.2 Several major breakthroughs in the last decades allowed this material system to surpass the 20% efficiency benchmark on rigid and flexible substrates3,4 and PCEs as high as 23.35% were reported.5 One of the important steps that allowed for high PCE was the introduction of a Gallium back-gradient, which effectively reduced the recombination at the Mo/MoSe2 back-contact.6,7 The optimization of this interface is still an area of intensive research due to the following reasons: The Molybdenum back-contact interface has a poor optical reflectivity and the recombination velocity is high,8 which means that effective passivation strategies are indispensable. This is especially true for ultra-thin devices9 where bandgap grading is not feasible.

Pure CuInSe2 (CISe) solar cells have several advantages compared to CIGSe such as an easier manufacturing process and a lower bandgap, which makes this material more attractive for tandem applications. However, the PCE was stuck for a long time at 15%8 without post deposition treatment. Recently, KF treatment improved this number to 16%,9 which is still far away from the record CIGSe absorbers. It was shown that the likely reason for the lower CISe solar cell performance...
was the absence of the Ga back-contact grading. The minority carrier lifetime improved after exfoliation of the absorber layer from the back-contact, which showed that recombination at the back is a major issue in nongraded CISe absorbers. Meanwhile, some laboratories showed that a Ga back-gradation could be introduced in CIGSe in a way that the bandgap was still close to the pure CISe, which resolves the disadvantage for tandem applications, albeit increasing the complexity of the process. In agreement with the previous discussion, the PCE increased, and meanwhile, above 19-% efficient solar cells were reported with a bandgap close to 1.0 eV.

There are already quite a few important reports that focused on the buried MoSe2/Cl(G)Se interface, which mostly relied on the fact that the absorber could be peeled off mechanically from the back-contact. This made it possible to use surface sensitive techniques such as X-ray and UV photo-electron spectroscopy (XPS/UPS) to study compositional and electronic variations at that interface. It was shown that there is a Ga accumulation at the back-contact and the CuInS2 (CIS) back-surface was Cu-depleted despite Cu-rich growth conditions. Transmission electron microscopy measurements (TEM) showed that there was some in-diffusion of Cu into the Mo and atom probe tomography (APT) measurements showed an increased Cu and Ga concentration in the first 50 nm of the CISe absorber (measured from the back-surface). Most of these measurements were performed on Cu-poor CISe (Cu/In<1). For the case of CuGaSe2, grown under Cu-excess, it was shown via TEM measurements that CuSe precipitations could also exist in the bulk of the absorber layer. Already at that time, it was speculated that this may be the reason for the low efficiencies of Cu-rich absorber layers. However, until today, the general consensus in the community is that Cu-rich absorbers exhibit a secondary phase free bulk with CuSe on top, which can be etched away via a potassium cyanide etching step.

Much less is known about the lateral variations of the electrical properties on the nanometer scale of the back-surface. A detailed combined electron beam induced current (EBIC) and cathodoluminescence (CL) study showed that, for a number of absorbers with different compositions, the grain boundary properties varied and there was no direct correlation between the CL signal and the collection efficiency, as measured via EBIC. For low symmetry grain boundaries, CL measurements showed a reduced signal, which pointed towards an increased number of defects at the grain boundaries. Furthermore, there are some Kelvin probe force microscopy (KPFM) measurements available on CuGaSe2, which showed that most of the grain boundaries exhibited downward band bending in the order of approximately 50 meV compared to the grain surfaces. Furthermore, Ga remnants were found on the Mo-side. So far, no scanning probe microscopy measurements are available for CISe.

It is well known in the Cl(G)Se community that the Cu-content is very important. High performance CISe devices are grown under Cu-poor conditions (Cu/In<1), which improves VOC and quasi-Fermi level splitting compared to Cu/In>1. For Ga-free CISe absorbers, the difference in quasi-Fermi level splitting between Cu-poor and Cu-rich absorbers is small. However, in agreement with the Ga-containing absorbers, the PCE is higher for Cu-poor than for Cu-rich devices. In the case of Cu-rich absorbers, the excess Cu condensates as Cu2Se on top of the CISe absorbers. A subsequent potassium cyanide (KCN) etching removes the Cu2Se (or CuS in the case of CIS), thereby creating a stoichiometric absorber with less Cu-vacancies in the bulk. The KCN etching, however, also has some adverse effects on the composition and defects in the near surface region. The investigation of the rear-surface thereby offers a very nice opportunity to investigate stoichiometric absorbers (grown Cu-rich) without KCN etching.

In this contribution, stoichiometric CISe absorbers (grown Cu-rich and subsequently KCN etched) were mechanically peeled off from their substrate and systematically scrutinized by means of scanning probe microscopy (SPM) techniques such as scanning tunneling microscopy (STM) and Spectroscopy (STS) and KPFM. The local density of states was evaluated by STS and compared to measurements performed at the top surface whereas variations in workfunction of the films were probed by KPFM in order to compare the results to already published data on CuGaSe2. Additionally, XPS and energy dispersive X-ray analysis (EDX) were used to explore the surface and bulk compositions and the impact of the peeling process on quasi-Fermi level splitting was investigated with photoluminescence (PL) imaging. Finally, X-ray diffraction measurements were used to investigate the occurrence of secondary phases in the CISe absorber layers.

## 2 EXPERIMENTAL DETAILS

SPM measurements were performed in a variable temperature ultra-high vacuum system with a base pressure in the 1011 mbar range. The STM and STS measurements were acquired using chemically etched tungsten tips, cleaned in situ by voltage pulses, on a clean gold substrate, anterior to the measurements of the CuInSe2 films. The local density of states (LDOS) of the absorbers were acquired via current imaging tunneling spectroscopy (CITS) maps at several regions of the absorbers in order get representative data. The analyzed areas range from 1×1 μm2 to 4×4 μm2. For each CITS map, a grid of 120×120 pixels was defined at which a current-voltage (I-U) curve was acquired while the feedback loop was switched off. In the present study, the voltage was swept from −2V to +2V. Thereafter, the LDOS, which is proportional to dI/dU, was derived by numerical differentiation of the current I with respect to the applied voltage U.

Work function maps were measured using frequency modulation Kelvin probe force microscopy (FM-KPFM), which compensates the electrostatic force gradient between a sharp conductive tip and the sample, providing a credible and accurate contact potential difference (CPD) value, combined with an excellent lateral resolution. The measurements were performed with the bias applied to the sample and Pt-Ir coated Si-cantilevers with the following nominal characteristics were used: resonance frequency 75 kHz, tip radius 20 nm, length 215–235 μm, width 20–35 μm, set-point for topography measurements −5 Hz.

The absorbers analyzed in this study were synthesized via coevaporation on molybdenum coated soda lime glass (SLG) substrates at a nominal temperature of 580°C. A detailed description of the growth process can be found in Deprédurand et al. Typical device performances from Cu-rich absorbers are in the range of 8%-10%.
In order to examine the back-side of the absorbers, they were peeled off from their original substrates following the procedure schematized in Figure 1. The samples were glued to clean fluorine doped tin oxide coated glass substrates using a ultra-high vacuum (UHV) compatible Ag epoxy glue. Curing was performed in air on a hot plate at approximately 100°C for 45 min. Afterwards the samples were transferred into a nitrogen filled glove box (H2O < 1 ppm; O2 < 1 ppm) where they were exfoliated and introduced into the SPM analysis chamber without exposing them to air. This was accomplished by means of a vacuum suitcase. A similar procedure was followed for the XPS analysis. PL and EDX investigations were accomplished on the same absorbers (produced during the same growth process). For these measurements, a short air exposure could not be circumvented. For the PL measurements, the time between cleaving and measuring was of the order of 10 minutes.

For the XPS measurements, the samples were glued in Luxembourg and then shipped to Münster in a N2-filled transport box. The samples were again introduced into a N2 filled glovebox, which was connected to the UHV system. The samples were cleaved in this box and directly introduced into the XPS chamber. Thereby, the back-surface was only exposed to N2 for a few minutes. Because only one sample could be loaded at a time, the Mo-side was kept in N2 and introduced into UHV the day after. The XPS setup was calibrated on clean single-crystal metal surfaces prior to the measurements of the CISe back-side and of the Mo-side. The base pressure of the XPS system was in the 10−11 mbar range and a monochromatic Al Kα X-ray source was used for the XPS measurements.

In order to compare the near-surface composition to the bulk composition, additional EDX measurements were performed on both the front-side and back-side of the films using 20, 15, 7, and 5 kV acceleration voltage. Elemental compositions were deduced via the well-known ZAF method38 without standardization. Grazing incidence X-ray diffraction measurements were carried out at an incident angle of 1° in a parallel beam configuration. The measurements were carried on samples, which were first KCN etched in order to remove a Cu2Se secondary phase at the front surface and subsequently cleaved, as discussed in the previous paragraphs.

The impact of the cleaving and of the Ag epoxy glue on optoelectronic properties was analyzed with PL imaging performed in a custom built system. A 532-nm pulsed laser illuminated an area of approximately 2x2 cm². The generated PL signal was collected with an InGaAs camera, with a high quantum efficiency in the range of 800 to 1600 nm. The system was calibrated to absolute photon numbers in order to estimate the PL quantum efficiency and the quasi-Fermi level splitting of the absorber before and after the peeling process.

3 | RESULTS

3.1 | The MoSe2/CISe interface

Figure 2A shows a representative STM image of the rear-surface of a Cu-rich CuInSe2 absorber layer. The surface is very flat with an RMS roughness of 8 nm. The locations of the grain boundaries are well visible in the image as they exhibit a slightly lower height than the grain interior. The large grains are covered with tiny little grains, which we observed in all of the images we analyzed. Furthermore, some deep holes (black contrast) were observed, which were likely to be caused by pinholes in the absorber layer prior or during the peeling process. In Figure 2B, the corresponding STM image of the Mo-side is presented, where no indications of grain boundaries were found. The surface was covered with small precipitates, which had an extension of approximately 70 nm.

In order to corroborate that the absorber peeled at the MoSe2/CISe interface, XPS measurements were carried out on both samples (back-side and Mo-side) and the scans are presented in Figure 2C. On the back-side, we did observe all the peaks of the CISe matrix, that is, Cu, In, and Se. Contrary, the Mo-side did only show peaks that are related to Mo and Se in the survey scan. At a later stage of the manuscript, we will discuss high resolution scans in more detail. In contrast to the survey scan presented in Figure 2C, we found a very small amount of Cu on the Mo side but no traces of In. The XPS measurements corroborated that the absorber cleaved exactly at the MoSe2/CISe interface in accordance with previous reports6,11,20 due to the formation of a MoSe2 layer during synthesis, which is known to be a van der Waals solid where exfoliation is feasible.

The higher oxygen signal at the Mo-side was likely to be caused by the longer storage time of this sample compared to the back-side as discussed in the experimental part.

We also found an Ag peak in the XPS spectra, which we related to glue that was present at the edges of the exfoliated absorber due to the small size of these samples (see Figure 2F). Complementary EDX measurements showed that the Ag concentration in the bulk is lower than 1 at%, which is within the experimental error of the machine, considering the close distance of the In and Ag L-lines.
In order to check if the absorber layer was still intact after the peeling process, we carried out PL imaging measurements before the gluing and after the peeling process and the results are presented in Figure 2D–F. The scale bar is given in units of PL quantum yield (PLQY), which is defined as the quotient of the PL yield (photons/cm² · s) emitted from the sample $Y_{out}$ to the impinging photon density $Y_{in}$ generated by the laser. This quantity can then be used to extract the quasi Fermi-level splitting $\Delta E_F$ of the absorbers via Equation (1):\[\Delta E_F = \Delta E_F^{rad} + k_B T \ln \left( \frac{Y_{out}}{Y_{in}} \right). \tag{1}\]

$\Delta E_F^{rad}$ refers to the maximum achievable $V_{OC}$ in the absence of non-radiative recombination. In our case, the bandgap of the absorber is 1.0 eV, which translated to $\Delta E_F^{rad}=749$ meV. The remaining constants in Equation (1) are the temperature $T$ at which the sample was measured and the Boltzmann constant $k_B$. From the measurement right after KCN etching, we deduced a $\Delta E_F=471$ meV (measured by illuminating the front side of the CISe), which is a typical value for Cu-rich CuInSe$_2$ absorbers.\cite{23,24} After the gluing process and the subsequent liftoff, the non-cleaved part of the absorber did not change significantly and $\Delta E_F=465$ meV was very close to the fresh case (still measuring the front-side of the CISe absorber). The result of the back-surface, which is presented in Figure 2F shows a somewhat reduced $\Delta E_F$ of only 426 meV. This showed that the gluing and exfoliation procedure had a negative impact on the opto-electronic properties of the absorber layer. However, it is important to note here that the absorber was still intact and $\Delta E_F=426$ meV was only 39 meV lower, which ruled out extensive metallic in-diffusion of Ag that would have increased nonradiative recombination massively. We also note that the roughness of the exposed back-contact was extremely low, in contrast to the front-surface, which may also have altered the reflection coefficient of the impinging laser light and thereby the reabsorption probability of the emitted photons, which also has an impact on PLQY.\cite{39} From this analysis, we concluded that the peeling process of the back-surface was successful and that this procedure did...
Figure 3 (A) STM topography image of the back-surface of the CISe absorber layer. The black lines highlight the position of the grain boundaries. (B–D) $dI/dU$-CITS maps at different applied voltages of the region shown in Figure 2A. The measurements were performed at a tunneling set-point of 1 nA and the voltage was changed from $-2V$ to $+2V$. The blue solid, red dotted, and black dashed curves displayed in (E) represent the average density of states extracted from the back-side, Mo-side, and the UHV annealed front-surface [Colour figure can be viewed at wileyonlinelibrary.com].

not deteriorate the absorber much and an in-depth analysis of the back-surface with scanning probe techniques was well justified.

In Figure 3A, the visible grain boundaries of Figure 2A are highlighted together with the $dI/dU$-CITS maps at different applied voltages (Figure 3B–D). In all three cases, ($U = -1V$, $U = 0V$, and $U = 1V$) the maps are relatively featureless, which is in contrast to previously reported top-view measurements on potassium cyanide etched or air exposed CISe absorber layers.\textsuperscript{27,28,30} Especially, no distinct grain boundary contrast at $U = 0V$ could be observed, which indicated that the density of states at the grain boundaries was similar to the grain surfaces. The measurements showed some similarity to absorbers after UHV annealing, where a passivation of the defect states at the Fermi-level ($U = 0V$) was observed.\textsuperscript{27,28,30} Because the CITS maps at all specific voltages were featureless (no lateral variations), we extracted an average $dI/dU$ curve from the scanned region, shown in Figure 3A. The curve is shown in Figure 3E as a solid line together with a measurement on the Mo-side (see Figure 2B) and one that was measured after heat induced passivation on the front surface of a CISe absorber synthesized with the same process in the same physical vapor deposition system.\textsuperscript{27}

The first thing to note is that the UHV annealed sample and the measurement from the back-surface were very similar. The parts at positive voltages were almost identical whereas the curves at higher negative applied voltages were different. We relate this difference to a different densities of states of the tips, which were certainly different in the two measurements. Important however, was the excellent agreement of the valence band and conduction band onsets. The Fermi-level, which is located at $U = 0V$ was very close to mid-gap, in agreement with the UHV annealed case.\textsuperscript{27,29} However, we also note that the valence band and conduction band edges were not sharp and there was substantial tailing into the bandgap region. A possible reason for this will be discussed later in the manuscript.

The measurements on the Mo-side, which mainly consisted of MoSe\textsubscript{2} were different and we did see a finite conductance at $E_F$. Furthermore, the Fermi-level was closer to the conduction band, which points towards n-type doping in agreement with UPS measurements.\textsuperscript{14} The finite conductance at $E_F$ was an indication that the Mo-side had a lot of defects and explained why we did not observe a well-defined semiconducting gap as we would expect for MoSe\textsubscript{2}.

The absence of a clear grain boundary contrast in STS on the back-side is somewhat surprising because it was shown via a combined electron beam induced current (EBIC) and cathodoluminescence (CL) study that carrier collection efficiency and the CL yield were different at distinct grain boundaries. Our STS measurements do not support this interpretation. However, we need to keep in mind that the information depth was very different for EBIC/CL (several hundred nanometers) and STS (<1 nm). Furthermore, the oxygen content may have been very different. In the present study, the cleaving was carried out in a glovebox preventing air exposure. This is usually not done for EBIC/CL measurements, which are much less surface sensitive. Another question to be discussed is the sensitivity limits of the STS measurements to detect changes in work function. This can conveniently be done using the fact that the tunneling conductance can be approximated by the following formula:\textsuperscript{40}

$$\frac{dI}{dU} \propto \rho_T(eV)T(d, V, eV) \equiv LDOS(eV).$$

This equation assumes that the density of states of the tip $\rho_T$ is constant for all applied voltages and that the voltage dependence of the tunneling coefficient $T$ is small and can be neglected. The density of states of the sample is denoted with $\rho_S$. The tunneling coefficient
KELVIN PROBE FORCE MICROSCOPY

Figure 4 shows a technique that is more sensitive to changes in the work function than STS.33 The results are presented in Figure 5 where a topography image (Figure 5A) and a CPD map (Figure 5B) are depicted. The CPD map is related to the work function via Equation (5).

\[
V_{\text{CPD}} = \frac{\phi_S - \phi_T}{e}.
\] (5)

Consequently, the brighter regions in the CPD map corresponded to regions with higher work functions, assuming a constant work function of the tip. In the topography image, we again observed the grain boundaries as slightly lower regions, in agreement with the STS measurements. Furthermore, we also observed very small dots on the rear-surface. The contact potential difference map showed, in contrast to STS a slightly higher contrast at the grain boundaries. A line profile along such a grain boundary is shown in Figure 5C. The shaded area in the graph highlights the grain boundary region. We did observe a decrease of the topography by approximately 10 nm, together with an increase in the work function of approximately 40 meV. This anti-correlation was observed on all grain boundaries in the image. Comparing this result to the STS measurements (Figure 3) and the calculations (Figure 4) it is clear that values of grain boundary band bending of approximately 40 meV were too small to be detected in STS.

However, one important aspect still needed to be discussed in more detail. A careful inspection of the density of the small dots that cover the complete back-surface showed that, at the grain boundaries the density seemed to be slightly lower (see Figures 2A and 5A). We therefore needed to analyze if the changes of the dot density at the grain boundaries were responsible for the observed work function changes. At least the dots seemed to have an impact on the work function because we also measured variations in CPD on the grain surfaces. The line profiles in Figure 5C showed that the variations were approximately 20–30 mV in the area outside of the shaded region. In Figure 5C, we highlighted all the prominent minima in the topography with green dashed lines whereas the maxima are highlighted with black dotted lines. For both situations (maxima or minima) in the topography, we also measured CPD variations in both directions, that is, no direct correlation between the maxima and minima in topography and CPD. We concluded that the dots are not the main reason for the systematic changes in the work function that we observed at the grain boundaries. However, we realized from this analysis that the dots do have an impact on CPD albeit not always in the same direction.

In the following, we would like to discuss the EDX and XPS results in more detail in order to link them to the SPM measurements.

5 | ELEMENTAL COMPOSITION AT THE REAR-SURFACE

Table 1 summarizes the elemental compositions we measured with EDX and XPS. The EDX measurements were acquired at 20, 15, 7, and 5 kV in order to gain some insights into the compositional changes as a function of depth. In addition to these measurements of the back-side, we also included a measurement carried out at 20 kV on the front-side of the absorber after KCN etching. This measurement can be considered as a reference because an etched stoichiometric CISe absorber layer should exhibit a Cu/In ratio of 1 in the bulk in agreement with our elemental composition at the rear-surface measurements.
measurements. The systematic error for the elemental concentration is at least 1 at% for an EDX measurement and consequently the small deviations from the expected values were well within the error of the measurement technique. Although the systematic errors in EDX are large, changes in the composition at varying acceleration voltages are much less susceptible to large errors if the quantification is done with the same characteristic X-ray lines. Consequently, the measurements performed at different acceleration voltages suggested that the Cu-content is slightly higher at the back-contact region. However, EDX measurements are not surface sensitive enough to analyze compositional changes in the near surface region.

XPS measurements are much more surface sensitive (≈ 6–8 nm) than EDX measurements, which are more bulk sensitive (information depth ≈ 0.1–1 μm at 5–20 kV acceleration voltage). Interestingly, the Cu concentration, as deduced via XPS was much higher close to the interface with a Cu/In ratio of 1.7. This was a clear indication that excess Cu was present at the back-contact and further strengthens the EDX results. At the Mo-side we measured a small amount of Cu and no In, which corroborated that there was no CISe remnants on the Mo-side. The Mo/(Se+O) ratio is 0.6, which points towards a MoSe₂ layer, partially oxidized due to the rather long storage time of this substrate in the glovebox before introduction into the XPS system. The oxygen concentration was more than doubled compared to the back-side absorber, which was introduced directly after the exfoliation. However, residual oxidation of the substrate prior to the growth process cannot be excluded and may also have contributed to the higher oxygen content.

We will now discuss the high resolution scans of the individual elemental transitions acquired via XPS. In Figure 6A, the Cu₂p line of the back-side and of the Mo-side are shown. The binding energy of 932.4 eV was in excellent agreement with the reported values for CuInSe₂. The Auger parameter, which is only sensitive to the chemical environment and not to changes in binding energy as a result of different band bending, was 1849.7 eV, which was also in good agreement with CISe. The Cu concentration at the Mo-side was only 1 % (see Table 1). In Figure 6D, the Cu 2p spectra were normalized to the peak maxima in order to improve the visibility. Within the error of the measurement, we could not observe a significant shift in the binding energy and the Auger lines were too weak to estimate an Auger parameter. The In 3d lines are presented in Figure 6B. The binding energy was in accordance with the reported values, and we could not observe Indium at the Mo-side. This showed that the detected Cu that we measured on the Mo side is not bound to CISe. The Se 3d lines are depicted in Figure 6C. A clear shift in the binding energy was visible when comparing the two substrates. On the backside the Se 3d½ was located at 54.2 eV whereas on the Mo-side the value...
TABLE 1  EDX and XPS compositions

|                  | Cu | In | Se | Cu/In |
|------------------|----|----|----|-------|
| **Back-side (20 kV)** | 24 (24) | 25 (25) | 50 (50) | 0.9 (0.9) |
| **Back-side (15 kV)** | 28 (28) | 24 (24) | 48 (48) | 1.2 (1.2) |
| **Back-side (7 kV)** | 30 (30) | 24 (24) | 46 (47) | 1.3 (1.3) |
| **Back-side (5 kV)** | 32 (30) | 23 (23) | 45 (47) | 1.4 (1.3) |
| **Front-side (20kV)** | 25 | 24 | 51 | 1.0 |

**XPS composition [at%]**

|                  | Cu2p | In3d | Se3d | Cu/In | Na1s | Mo3d | O1s |
|------------------|------|------|------|-------|------|------|-----|
| **Back-side**    | 29   | 17   | 42   | 1.7   | 0.5  | /    | 11  |
| **Mo-side**      | 1    | 0    | 36   | /     | 0.3  | 38   | 24  |

Note. EDX and XPS measurements were acquired on large areas (compared to the grain size) in order to get representative averages. EDX values in brackets were measured after KCN etching of the back side. All samples were etched on the front side prior to the peel off process. All measurements performed on the back side were carried out in one session to limit the error bar. The measurements on the front side were done with another SEM machine and different systematic errors may influence the absolute numbers.

FIGURE 6  (A–D) High-resolution XPS measurements of the back-side and the Mo-side (A) Cu 2p, (B) In 3d, (C) Se 3d, (D) normalized Cu 2p curves depicted in (A). (E) Grazing incidence XRD measurement performed on a peeled absorber (the front-side of the absorber was etched whereas the back-side was not). (F) SEM micrograph of the back-side of the absorber prior to KCN etching. (G) Identical absorber after KCN etching. XRD references: CuInSe2:04-007-6152; CuSe2: 04-007-6152, *: Kβ peak of the (112) and (204) reflex [Colour figure can be viewed at wileyonlinelibrary.com]

was located at 54.4 eV. The shift towards higher binding energies was attributed to a change in the chemical environment of the Selenium from a CISe matrix on the back-side to a MoSe2 matrix on the Mo-side. The measured values of the binding energies were in agreement with reported values.42,43 The elemental composition of the MoSe2 (Table 1) was not equal to 2 and the large amount of oxygen was likely to influence the MoSe2 concentration we measured. For both surfaces, the Na concentration was on a similar level with approximately 0.5%.

The XPS and the EDX measurements strongly suggested that there was excess Cu at the back contact of the absorber layer. In order to corroborate the existence of a Cu-Se secondary phase, grazing incidence X-Ray diffraction was used. The results are presented in Figure 6E. It needs to be emphasized that this absorber layer was KCN etched before the peeling process in order to make sure that all the Cu,Se detected in the diffractogram originated from precipitates in the bulk or from the back contact. We observed a peak at 25.6° and additional shoulders around the (312) CISe peak at 52.39°, which were also reported to originate from Cu-Se.19,44 Consequently, the XRD data confirmed that precipitations exist in Cu-rich absorbers, despite the fact that the front surface was etched with KCN prior to the peel off process. In Figure 6F,G, high-resolution SEM measurements are presented, which show the morphology of the CISe back side.
prior to KCN etching and after KCN etching. The small white grains completely disappeared after KCN etching, and furthermore, we did observe substantial changes at the grain boundaries. Complementary AFM measurements (not shown) corroborated the observations made with SEM. The grain boundaries measured in AFM however were much deeper compared to the non KCN etched sample. The brighter contrast in SEM as shown in Figure 6G is not indicative of an accumulation of material but rather due to a higher secondary electron yield at the grain boundaries, i.e. a different electron affinity or workfunction. Currently, we assume that the etching rate at the grain boundaries was enhanced, suggesting that there was a larger amount of Cu$_2$Se present compared to the absorber back-surface. We attribute the small grains in Figure 6F to Cu$_2$Se precipitates, which were removed during the KCN etching process. EDX measurements performed on the same sample showed a reduction of the Cu/In ratio from 1.4 to 1.3 due to the removal of a Cu$_2$Se secondary phase.

The present investigation shows that Cu-rich Selenium based absorbers exhibit different surface- and back-surface compositions compared to the absorber bulk. The Cu-rich growth conditions trigger the precipitation of a Cu$_2$Se secondary phase, which is not present in Cu-poor material and which segregates at both interfaces. In our STM and AFM measurements, we observed that both sides of the rear-interface were covered with very small grains in the order of several tens of nanometers. We did see a correlation of the work function with the grains although we could not observe one distinct value. It rather seemed like randomly oriented small grains that exhibit slightly different work functions. We assign the small grains and their variations in work function on the grain surfaces (≤20 meV) to Cu$_2$Se. This conclusion was drawn on the basis of our XPS and EDX measurements, which showed a clear increase in Cu at the back-side of the Cu$_2$Se absorbers and the XRD measurements showed that residual Cu$_2$Se was present in the absorber layer. From XRD, we cannot rule out that there was no Cu$_2$Se present in the bulk of the absorber. However, the EDX and SEM measurements confirm that at least a part of the Cu$_2$Se was present at the rear-side.

Another important point that needs to be discussed further are the STS results. Based on our XPS findings, the back-surface of the Cu$_2$Se was partially covered with Cu$_2$Se. In the present study, we focused on large area scans where the dl/dU maps averaged over large areas covered with a large number of small grains. This fact may explain the rather undefined band edges that we observed in our measurements. High resolution STS measurements are necessary to disentangle the signal from the small grains from the one of the underlying substrate. At present this was not possible due to the duration of the CITs scans (approximately 15 hours), which imposed very strict limits for the maximum allowed temperature drift during STS measurements.

We finally would like to discuss the implications for devices. Our measurements corroborate that Cu$_2$Se precipitations do not only form at the front contact but also at the rear-surface under Cu-rich growth conditions in Cu$_2$Se. The interface recombination velocities are likely to be higher than for a pure MoSe$_2$/CuInS$_2$ interface since the MoSe$_2$ is a layered material with pure van der Waals bonding along the growth direction, that is, no dangling bonds. It is likely that the Cu$_2$Se secondary phase makes the interface worse and thereby reduces the PCE of the final device. Furthermore, band bending is different than in Cu-poor CuGaSe$_2$. Although at present we cannot disentangle the influence of the Ga/In from the different Cu content, the results still showed the composition has important consequences for the properties of the grain boundaries as also described in a recent APT study for CuInS$_2$.45
The reported measurements showed an increased concentration of Cu at the grain boundaries in Cu-rich CuInS₂. If we assume a similar grain boundary chemistry in Cu-rich CISe, excess Cu would segregate out at all interfaces (front, back, and grain boundaries) with distinct opto-electronic properties compared to the Cu-poor absorbers. The SEM measurements performed before and after KCN etching of the back side are in line with these observations since we do see a preferential etching of the grain boundaries.

In summary, we showed that for stoichiometric CISe absorbers the rear-surface exhibited an accumulation of Cu, which we assigned to a Cu₂Se secondary phase. This is in contrast to Cu-poor CISe where a Cu and Ga enrichment were measured. We measured a slight upward band bending of approximately 40 meV at the grain boundaries and showed that such low values cannot be measured with STS. The band bending of approximately 40 meV at the grain boundaries and Cu rear-surface exhibited an accumulation of Cu, which we assigned to a preferential etching of the grain boundaries.

Back side are in line with these observations since we do see a SEM measurements performed before and after KCN etching of the out at all interfaces (front, back, and grain boundaries) with distinct grain boundary chemistry in Cu-rich CISe, excess Cu would segregate Cu at the grain boundaries in Cu-rich CuInS₂. If we assume a similar phase, which can be etched off on the front-side is also present at the rear-surface, increasing the back-surface recombination velocity. In the absence of a suitable back-surface field, this may be very detrimental for device performance. Finally, we would like to note that it is likely that similar processes occur in other Selenium based absorbers layers such a Cu₂ZnSnSe₄ where Cu-poor growth conditions that it is likely that similar processes occur in other Selenium based absorbers layers such a Cu₂ZnSnSe₄ where Cu-poor growth conditions are also mandatory to reach the current record power conversion efficiencies.

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