Effect of sodium diffusion on the properties of CIGS solar absorbers prepared using elemental Se in a two-step process

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The influence of Na diffusion from various glass substrates during a high-temperature selenization process on the microstructure and morphology of two-step formed CIGS absorber layers is investigated. In order to minimise the CIGS absorber formation time, elemental Se vapour is used to prepare the CIGS absorber. The grain sizes of the CIGS films are found to increase with increasing sodium in the glass substrates (extra clear glass, soda-lime glass, borosilicate glass). TiN and SiN thin films are used as diffusion barrier layers inserted between the glass substrate and the Mo rear contact to tune the Na diffusion from the soda-lime glass. The interdiffusion between the In-rich CuInSe₂ surface layer and the Ga-rich CuGaSe₂ layer is promoted by the barrier layer, leading to larger CIGS grains. Efforts are also taken to understand the differences in Na diffusion (from the glass substrates) and their effects on the MoSe₂ intermediate layer formation during the high-temperature CIGS absorber formation processes. We find that excess amounts of Na and Se are essential for the MoSe₂ growth. The excessive Na in the form of Na₂Se at the CIGS/Mo interface works as a Se source and catalyses the MoSe₂ formation. The Se flow in the two-step CIGS formation process must be sufficiently high to obtain high-efficiency CIGS solar cells.

The copper indium gallium di-selenide (CIGS) solar cell is one of the most efficient solar cells for standard 1-sun application, with efficiencies of up to 22.9% at the lab scale using a very thin absorber layer (< 3 μm)¹. Molybdenum (Mo) thin films deposited by magnetron sputtering is the most widely used rear contact for CIGS photovoltaic, as it can meet most of the back contact requirements and ensures high-efficiency CIGS solar cells²–⁶. Mo films deposited by magnetron sputtering show low electrical resistivity (sheet resistance < 0.5 Ω/sq), high thermal stability, and good corrosion resistance during the CIGS absorber formation process which normally takes place at a high temperature of 500–600 °C²–¹¹. In addition, Mo film also offers a low resistive ohmic contact between the Mo back contact and the CIGS absorber by forming a thin MoSe₂ layer at the Mo/CIGS interface during the high temperature absorber formation step¹²–¹⁴.

During the two-step CIGS formation process, Na diffusion from the substrate plays an important role in forming a uniform CIGS film: the Na content can postpone the inter-diffusion of In and Ga atoms to form a uniform CIGS layer¹⁵–¹⁹. However, the Na diffusion is related to the properties of the rear contact and the selenization process. The rear contact properties have a strong influence on the formation of the intermediate MoSe₂ layer and the properties of the formed CIGS absorber layer¹⁶,¹⁷,¹⁹. The MoSe₂ layer plays a significant role in ensuring a low contact resistance and good adhesion of the CIGS absorber to the rear contact. The CIGS layer tends to delaminate from the Mo coated substrate after CdS buffer layer deposition if its adhesion to the substrate is not promoted by the MoSe₂ layer, which is normally formed at the Mo and CIGS interface during high temperature CIGS formation process. The existence of a thin MoSe₂ layer can decrease the apparent Schottky barrier height and thereby provide a better ohmic contact to the CIGS absorber. Simultaneously, MoSe₂ is able to enhance the mechanical peel strength of CIGS to Mo back contact. However, an excessive thickness of MoSe₂ layer formed

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respectively. Solar cells with these CGI and GGI ratios are reported to achieve high efficiencies of above 20%.

SLG/SiN/Mo/CIGS samples are clearly observed in Fig. 3(c,g), while white Na2Se precipitates are found on the surface of SLG/Mo/CIGS samples. The different glass substrates and rear contact designs utilized in this study for CIGS solar cell fabrication are shown in Fig. 1. The optimized deposition conditions of the individual layers used to form various back contact stack designs are described in our previous publication. Bilayer Mo films comprising of a high pressure (HP, 6.0 × 10⁻⁵ mbar) deposited bottom Mo layer and a low pressure (LP, 1.5 × 10⁻³ mbar) deposited top Mo layer are deposited on three different glass substrates (size: 30 cm × 40 cm) with different Na concentrations: extra clear glass (solar grade opti-white diamond glass, Dia, high Na concentration), soda-lime glass (SLG, intermediate Na concentration), borosilicate glass (Boro, negligible Na concentration), see Fig. 1(a–c). On Na-free borosilicate glass substrate, for comparative purposes, an additional 200 nm thick Mo:Na (3 wt%) film is deposited on top of the bilayer Mo stack, which serves as an extrinsic Na source (see Fig. 1(d)). In addition, TiN and SiN thin films are utilised as sodium diffusion barrier and adhesion enhancement layer for the LP-deposited 500-nm Mo film onto SLG substrates (see Fig. 1(e,g)). Finally, a 200 nm thick Mo:Na layer (see Fig. 1(f,h)), used as extrinsic Na source, is deposited onto these rear contacts with a thin TiN or SiN sodium diffusion barrier layer.

The Mo films using different back contact designs were cut into 10 cm × 10 cm substrates for CuGa/In precursor deposition. A four-layer CuGa/In/CuGa/In stack design was used to prepare the Cu-In-Ga precursors by direct current (DC) magnetron sputtering. The selenisation of the precursors was performed in nitrogen atmosphere at ambient pressure condition. The heater temperatures of the RTP (rapid thermal process) were 580 °C and the duration of the high temperature process was about 12 minutes. Thermally evaporated (420 °C) Se vapour was used as the Se source to form CIGS absorber layers. The average GGI ([Ga]/([Ga] + [In])) and CGI ([Cu]/([Ga] + [In])) ratios of the CIGS absorber measured by X-ray Fluorescence (XRF) were about 0.30 and 0.93, respectively. Solar cells with these CGI and GGI ratios are reported to achieve high efficiencies of above 20%.

Results

In this study, CIGS absorbers with different Se contents (50 and 57 at.%) were fabricated via changing the Se crucible temperature in order to investigate the effect of the Se content on the formation of the MoSe₂ layer. Figure 2 shows Scanning electron microscopy with energy dispersive X-ray spectroscopy (SEM-EDX) images of Se elemental mapping (at.%) across the absorber layer thickness. It can clearly be seen that the Se content in Fig. 2(a) decreases gradually from the top to the bottom of the absorber. At the rear surface of the absorber, i.e. at the Mo/CIGS interface, an approximately 200 nm thick CIGS layer was found to contain about 33% to 42% of Se, indicating a Se-poor absorber. However, for the sample shown in Fig. 2(b), the Se is distributed uniformly at about 50 at.% showing a Se-rich absorber. The effects of the different rear contacts on the properties of the Se-poor and Se-rich CIGS absorber regions are investigated and discussed in the following sections.

Se-poor CIGS absorber. Surface morphology. The surface morphology of the CIGS films formed on different rear contact structures was studied by SEM, see Fig. 3. The surface grain boundaries of Boro/Mo/CIGS and SLG/SiN/Mo/CIGS samples are clearly observed in Fig. 3(c,g), while white Na₂Se precipitates are found on the surface of SLG/TiN/Mo/CIGS samples and samples having a Mo:Na layer at the rear contact as well.

**Figure 1.** Different back contact designs used for CIGS solar cell fabrication: (a) Dia/Mo, (b) SLG/Mo, (c) Boro/Mo, (d) Boro/Mo:Na, (e) SLG/TiN/Mo, (f) SLG/TiN/Mo:Na, (g) SLG/SiN/Mo, and (h) SLG/SiN/Mo:Na.
The grain size of the CIGS films formed on bilayer Mo-coated diamond glass and soda-lime glass was much larger than that of the CIGS film formed on borosilicate glass, which was attributed to Na diffusion from the glass substrate through the bilayer Mo rear contact into the absorber. Thus, Na can diffuse through the voids between the columnar structures of bilayer Mo films. The addition of a 200 nm thick Mo:Na layer on top of the bilayer Mo rear contact slightly increased the grain size of the CIGS film. For the modified rear contacts deposited on SLG substrates, the grain size of the CIGS films formed on SLG/TiN/Mo rear contacts was much larger than that of CIGS films formed on SLG/SiN/Mo rear contacts (see Fig. 3(g)). In addition, the grain sizes of the Mo films deposited on SLG/SiN/Mo and on Boro/Mo substrates were quite similar (in the 0.5–1 μm range). These two findings indicate that a SiN barrier layer is more efficient in terms of blocking the Na diffusion from the soda-lime glass substrate than a TiN barrier layer. Using a Mo:Na film as extrinsic Na source, the grain size of the CIGS films formed on SLG substrates with a TiN or SiN barrier layer were in the same range (about 1–3 μm) as those without a barrier layer in the rear contacts.

The grain sizes of the CIGS absorber were measured via surface SEM images. Histogram plots of the CIGS absorber grain size formed on different glass substrates and rear contacts are depicted in Fig. 4. The calculated average grain sizes are also shown in Fig. 4. The SEM studies clearly reveal that the Na incorporation either from the glass substrate or from the Mo:Na layer can enhance the CIGS grain size. It should also be noted that the CIGS film deposited on the SLG/TiN/Mo substrate showed the best lateral uniformity.

**Chemical composition.** The XRF measurement results for the CIGS absorber with 50 at.% Se formed on different rear contacts are shown in Table 1. The SeC and SeGI are \( \frac{[\text{Se}]}{[\text{Cu}]} \) and \( \frac{[\text{Se}]}{([\text{Ga}]+[\text{In}])} \) ratios. The average CGI and GGI ratios of these CIGS absorbers are about 0.93 and 0.30. The calculated Se/M \( ([\text{Se}]/([\text{Cu}]+[\text{In}]+[\text{Ga}]) \) ratio was slightly higher than 2.0. It is reported in the literature that the excess Se in CIGS solar cells exists in the form of a MoSe\(_2\) layer\(^{20}\). The thickness of the MoSe\(_2\) \( (t_{\text{MoSe}_2}) \) layer can be estimated from the following equation\(^{20}\):

\[
t_{\text{MoSe}_2} = t_{\text{CIGS}} \left( \frac{C_{\text{Se}}^{\exp} - C_{\text{MoSe}_2}^{\text{nom}}}{C_{\text{Se}}^{\text{nom}}} \right) \frac{n_{\text{CIGS}}}{n_{\text{MoSe}_2}}
\]

where \( C_{\text{Se}}^{\exp} \) is the atomic percentage of Se in the absorber as measured by XRF, \( C_{\text{Se}}^{\text{nom}} \) is the nominal atomic percentage of Se in the absorber, \( C_{\text{MoSe}_2}^{\text{nom}} \) is the atomic percentage of Se in the MoSe\(_2\) (i.e. 0.67), \( n_{\text{CIGS}} \) is the atomic density of CIGS (i.e. \( 4.23 \times 10^{22} \text{cm}^{-3} \)), and \( n_{\text{MoSe}_2} \) is the atomic density of MoSe\(_2\) (i.e. \( 5.15 \times 10^{22} \text{cm}^{-3} \)). The nominal atomic percentage of Se in the absorber \( C_{\text{Se}}^{\text{nom}} \) is calculated by:

\[
C_{\text{Se}}^{\text{nom}} = \frac{1 + 3/x}{3 + 5/x}
\]

where \( x \) is the GGI ratio \( ([\text{Ga}] / ([\text{Ga}]+[\text{In}])) \) in the absorber.

The thickness of the MoSe\(_2\) layer estimated using Eq. 1 is listed in Table 1. The estimated thicknesses of the MoSe\(_2\) layers of all the Se-poor samples are quite similar (\( \sim 54 \) nm). Therefore, it is not possible to derive a correlation between the MoSe\(_2\) layer thickness and the Na content, as the thickness of the MoSe\(_2\) intermediate layer is the same regardless of the Na content in the glass substrates. Also, it is not possible to visibly observe the MoSe\(_2\) layer at the Mo/CIGS interface in the cross-sectional SEM measurements\(^{18}\), which indicates that the estimated MoSe\(_2\) layer thickness (54 nm) from Eq. 1 is reliable. Thus, the Na concentration is not the only factor that affects the formation of the MoSe\(_2\) layer during the high-temperature selenisation process.

**Figure 2.** Cross-sectional SEM-EDX Se elemental mapping of CIGS absorbers formed on SLG/Mo substrates with different Se contents: (a) Se-poor (50 at.% measured by XRF) and (b) Se-rich (57 at.% measured by XRF). The different colours at the bottom denote the corresponding atomic percent values.
However, it should be noted that the Se/M ratio in the CIGS absorber is higher for SLG/Mo and extra clear (diamond glass)/Mo rear contacts as compared to those with Boro/Mo and SLG/TiN/Mo (or SLG/SiN/Mo) rear contacts. For example, the Se/M ratio of the absorbers formed on Dia/Mo substrates was the highest (1.25) of the three investigated glass substrate types. Thus, Na can assist the incorporation of Se into the absorber by the formation of Na$_2$Se$_x$. It should also be noted that the thickness of the CIGS layer increased with increasing Na content in the glass substrate, which confirms that a certain percentage of Na$_2$Se is always present at the CIGS grain boundaries, and also at the Mo/CIGS interface, which was confirmed by the SIMS measurements.
Microstructure. The XRD patterns and the CIGS (112) peak profiles of the Se-poor CIGS absorbers deposited on glass substrates containing different sodium content and rear contacts are shown in Figs 5 and 6. The peak position, peak intensity and other important parameters derived from XRD measurements are summarised in Table 2. As shown in Figs 5 and 6, all CIGS films show a polycrystalline structure, indicating the formation of stoichiometric CIGS. Moreover, there is no diffraction peak at 2θ = 26.2° indicating that Cu$_2$−xSe secondary phase was not formed on the surface of the CIGS absorber. The high-intensity peak of the Mo (110) orientation was found at 2θ = 40.5°. The intensity ratio I(220/204)/I(112) shown in Table 2 was calculated to evaluate the crystal growth of the CIGS films. There was little variation in the I(220/204)/I(112) ratio between the samples prepared on different glass substrates and rear contacts. A slight reduction in the (220/204) peak intensity was observed in samples deposited on SLG/TiN/MoNa and SLG/SiN/MoNa substrates compared to those without Mo:Na layer, which was attributed to the extrinsic Na incorporation from the Mo:Na layer.

As reported in our previous publication, the absorbers formed by the two-step sputtering-selenisation process are consisted of a bilayer structure: i.e., a Ga-rich CGS layer near the back contact and an In-rich CIS layer near the CdS buffer. The final Cu(In$_{1-x}$Ga$_x$)Se$_2$ film was formed through the interdiffusion between the CGS and CIS layers. However, the diffusion rate of Ga is much lower than that of In in the absorber. Hence, CIS and CGS phases are observed from XRD measurement. As the lattice constant increases linearly with reducing Ga content (x), the XRD diffraction peaks of the CIS phase will shift to lower angles compared to the CGS phase. The dashed lines in Figs 5 and 6 at 2θ = 26.65° and 27.65° are the (112) reflection position of the single CIS and CGS layers, respectively. The x value in Cu(In$_{1-x}$Ga$_x$)Se$_2$ is derived from the 2θ value of the CIS (112) reflection in the XRD pattern, using the following equation:

$$x = \frac{(y - 26.65)}{0.24} \times 0.3 = (y - 26.65) \times 1.25$$

where y is the 2θ value of the CIGS (112) peak. The calculated x values are listed in Table 2. It is evident from Table 2 that the CIGS absorbers formed on borosilicate glass substrates show the highest x values, which are closer to the GGI ratio in the films measured by XRF. In contrast, CIGS films formed directly on the SLG substrate without any diffusion barrier layer show the lowest x value. The diffusion of Na from the SLG substrate into the CIGS
Figure 5. XRD patterns and the CIGS (112) peaks of CIGS absorbers deposited on different glass substrates: (a) Diamond/Mo, (b) SLG/Mo, (c) Boro/Mo, and (d) Boro/MoNa. Attributions of each XRD peak are indicated in the plot as well.

Figure 6. XRD patterns and the CIGS (112) peaks of CIGS absorbers deposited on different modified rear contacts: (a) SLG/TiN/Mo, (b) SLG/TiN/MoNa, (c) SLG/SiN/Mo, and (d) SLG/SiN/MoNa.
absorber hinders the In and Ga inter-diffusion, and thus reduces the Ga content within the front region of the CIGS absorber. Both TiN and SiN barrier layers can hinder the Na diffusion, and thus promote Ga diffusion to the front. The lower $x$ values of the samples deposited on Na-containing glass substrates indicate lower Ga contents near the front surface of the CIGS films.

Another set of Se-rich (≥57 at.%) CIGS absorbers was prepared using a CuGa and In multilayer precursor with a thickness of about 525 nm. The CGI and GGI ratios in all samples were maintained at around 0.93 and 0.30, respectively. Properties of this Se rich absorber formed on various type of back electrodes are discussed below.

**Se-rich CIGS absorber.** Another set of Se-rich (≥57 at.%) CIGS absorbers was prepared using a CuGa and In multilayer precursor with a thickness of about 525 nm. The CGI and GGI ratios in all samples were maintained at around 0.93 and 0.30, respectively. Properties of this Se rich absorber formed on various type of back electrodes are discussed below.

**Morphology.** SEM images of the Se-rich CIGS absorber formed on SLG/Mo and Boro/Mo substrates are shown in Fig. 7. In the SEM surface images (Fig. 7(a)) white sodium selenide precipitates are seen on the SLG/Mo samples (which have a high Na content in the CIGS absorber due to Na diffusion from the substrate into the absorber). In contrast, the samples deposited on Na-free Boro/Mo substrate showed compact and well faceted grains with clear grain boundaries (Fig. 7(b)). SEM measurements also reveals a thick MoSe$_2$ layer formation.
between the Mo and the CIGS layer in the sample deposited on SLG/Mo substrate. The measured MoSe₂ thickness (~200 nm) is comparable to the estimated value using Eq. 3 (see Table 3).

**Chemical composition.** Table 3 summarises the XRF measurement results of Se-rich CIGS absorbers formed on various rear contact structures. It should be noted that the measured Se/M ratio, MoSe₂ layer thickness, and the thickness of the Se-rich CIGS absorber are quite different from the values of samples with Se-poor CIGS absorber. The results can be classified into two groups, based on the Na concentration: (i) absorbers with low or negligible Na concentration, for example SLG/SiN/Mo and Boro/Mo and also with optional Mo:Na capping layer) rear contacts, and (ii) absorbers with high Na concentration, for example SLG/Mo (500 and 900 nm) and SLG/TiN/Mo(Mo:Na) samples. From the XRF measurements, the values of Se/M ratio, MoSe₂ thickness and absorber thickness of the low-Na samples were observed to be lower than those of high-Na samples. However, for the Se-poor samples, as discussed in section 3.1, no significant difference is found in the thickness of both the CIGS layer and the intermediate MoSe₂ layer, regardless of the rear contact structure. Thus, a strong correlation between the Se and Na contents and their influences on the growth of the CIGS absorber and the MoSe₂ thickness is observed. This observation suggests that presence of excess Na in the CIGS absorber can enhance the Se/M ratio in the CIGS absorber, which further promotes the MoSe₂ growth, as well as increases the thickness of the CIGS absorber.

**Microstructure.** The XRD patterns of Se rich CIGS absorbers deposited on different rear contacts are shown in Fig. 8. All of these CIGS films show a polycrystalline structure. The intensity ratio I(220)/I(110) of the samples on SLG substrates is higher than that of the samples on borosilicate glass substrates (see Fig. 8). Hence it can be concluded that, in the case of Se-rich CIGS films, Na diffusion from the glass substrate enhances the growth of CIGS grains in the (220) orientations. As shown in Table 4, the estimated Ga content (x value in Eq. 3) of the samples on borosilicate glass is higher than that of the samples on SLG substrates, as the CIS and CGS interdiffusion is delayed by the presence of Na. However, the Ga content near the CIGS front surface is reduced by a TiN barrier layer for the samples deposited on SLG substrates, which can be ascribed to the suppressed Na diffusion by the TiN barrier from the glass substrate to the CIGS absorber.

From Fig. 8(e), it can be seen that for the samples formed on borosilicate glass substrates with an additional Mo:Na layer as extrinsic Na source, no MoSe₂ diffraction signal was detected from the XRD measurement, which is due to the low amount of MoSe₂ at the Mo/CIGS interface. The low growth rate of the MoSe₂ layer can be explained by insufficient Na₂Se intermediate compound formation at the rear surface. As discussed in our previous publication, for the CIGS films with higher Na diffusion from the substrate, it is observed to have higher Na and Se content in the form of Na₂Se compound at the Mo/CIGS surface. Additionally, from XRF measurements (Table 4), the observed MoSe₂ thickness was significantly higher for Na-rich glass substrates than for Na-deficient ones. Thus, it is reasonable to conclude that the growth of the MoSe₂ layer can be accelerated by Na₂Se. In other words, Na₂Se can be considered as a catalyst to form a thick MoSe₂ layer.

If all the Na from the 200 nm thick Mo:Na capping layer is assumed to be uniformly distributed in the completed CIGS solar cell, the Na content in the CIGS absorber would be about 0.5 at.%, which is higher than the observed Na content in CIGS solar cells formed on SLG/Mo substrates. Thus, it can be concluded that most of the Na atoms diffusing from the Mo:Na layer are trapped within the Mo grains and cannot easily diffuse into the CIGS film.

From the XRD measurements, the value of the MoSe₂ peak intensity ratio I(100)/I(110) was calculated, see Table 4. Since this value is higher than 1.0, this indicates a preferred grain orientation of the MoSe₂ layer in the <110> direction. It has been reported that a MoSe₂ grain orientation in the <110> direction reduces the contact resistance between the Mo and the CIGS layers, and enhances the adhesion of the CIGS layer to the rear contact. 

| Sample number | CGI | GGI | SeC | SeGI | Se/M | Mo | MoSe₂ | CIGS |
|---------------|-----|-----|-----|------|------|----|-------|------|
| Boro/Mo       | 0.94 | 0.31 | 2.44 | 2.29 | 1.18 | 573 | 81 | 1629 |
| Boro/MoNa     | 0.91 | 0.30 | 2.48 | 2.26 | 1.18 | 745 | 75 | 1599 |
| SLG/Mo(500 nm)| 0.93 | 0.31 | 2.72 | 2.53 | 1.31 | 548 | 140 | 1680 |
| SLG/Mo(900 nm)| 0.91 | 0.30 | 2.74 | 2.49 | 1.30 | 848 | 133 | 1703 |
| SLG/TiN       | 0.91 | 0.30 | 2.67 | 2.44 | 1.28 | 513 | 122 | 1673 |
| SLG/TiN/MoNa  | 0.93 | 0.31 | 2.73 | 2.53 | 1.31 | 775 | 140 | 1681 |
| SLG/SiN/Mo    | 0.93 | 0.29 | 2.32 | 2.16 | 1.12 | 511 | 46 | 1492 |
| SLG/SiN/MoNa  | 0.93 | 0.31 | 2.37 | 2.20 | 1.14 | 765 | 57 | 1514 |

Table 3. Summary of XRF measurements of Se-rich CIGS absorbers prepared using different rear contacts and glass substrates. The same method as in Table 1 was used to calculate the MoSe₂ thickness and the CIGS thickness.
Discussion and Conclusion

Dependence of morphology and microstructure of the CIGS absorber on the Mo-coated glass substrates. The sodium diffusion either from the glass substrate or from the Mo rear contact plays a large role in the grain growth of the CIGS absorber layer, as the sodium can promote the grain growth of the CIGS absorber layer. Na diffusion to the front surface of the CIGS absorber is beneficial for forming large crystallites near the front surface. The CuInSe2 on the CIGS surface can be formed through the reaction of either CuSe2 or CuSe with InSe. However, the grain size of Cu2Se is larger than that of CuSe34,35. The sodium selenide formed at the front surface of the CIGS can act as a Se source to promote the formation of larger CuSe2 grains, resulting in large CIS grains near the surface. At the same time, the diffused Na suppresses the CuInSe2 formation via reaction of smaller grain CuSe with InSe.

The CIGS absorber formed on the TiN modified Mo rear contact shows the best uniformity in grain size, indicating the good control of Na diffusion from the SLG glass substrate by utilizing the TiN diffusion barrier layer.

All the CIGS films show a bilayer microstructure verified by XRD measurement, i.e., a Ga-rich CGS layer near the back contact and an In-rich CIS layer near the CdS buffer. The growth of the CuInSe2 and CuGaSe2 grains across the film thickness requires different formation energy, time and temperature during the selenisation process. The final Cu(In1−xGa)xSe2 chalcogenide film is formed via inter-diffusion between the CGS and CIS layers. However, the Na diffusion from the substrate affects the inter-diffusion between the CGS and CIG layers.

Table 4. XRD measurement results of Se-rich CIGS absorbers formed on different glass substrates and different back contacts. The intensity ratio of CIGS (220/204) peak to CIGS (112) peak of CIGS absorber was also derived for comparison. x is the GGI value derived from Eq. 2. The intensity ratio of (100) to (110) peak of the MoSe2 layer are also calculated.

| Sample        | CIGS (112) | (220/204) | I(220/204)/I(112) | x   | MoSe2 (100)/I(110) |
|---------------|------------|-----------|------------------|-----|-------------------|
| SLG/Mo        | 26.80      | 44.39     | 0.32             | 0.20| 1.41              |
| SLG/TiN/Mo    | 26.78      | 44.39     | 0.32             | 0.16| 2.28              |
| SLG/TiN/MoNa  | 26.75      | 44.38     | 0.29             | 0.13| 2.03              |
| Boro/Mo       | 26.93      | 44.54     | 0.25             | 0.35| —                 |
| Boro/MoNa     | 26.93      | 44.57     | 0.25             | 0.35| —                 |

Figure 8. Full range XRD patterns (10–70°) and CIGS (112) peak profiles of Se-rich CIGS absorbers deposited on different rear contacts: (a) SLG/Mo, (b) SLG/TiN/Mo, (c) SLG/TiN/MoNa, (d) Boro/Mo and (e) Boro/MoNa. The two MoSe2 peaks are highlighted by red dashed lines.
CIGS absorbers deposited on borosilicate glass substrates show the highest x values, which are closer to the GGI ratio in the films measured by XRF. In contrast, CIGS films formed directly on the SLG substrate without any barrier layer show the lowest x value. The Na diffusion from the SLG substrate into the absorber reduces the In and Ga inter-diffusion, and thus reduces the Ga content at the front surface of the CIGS absorber. The TiN and SiN barrier layers can hinder the Na diffusion\(^9\), and thus promoted Ga diffusion to the front surface. The lower x values of the samples deposited on Na-containing glass substrates indicate a lower Ga content at the front surface of the CIGS absorber.

**Formation of MoSe\(_2\) intermediate layer.** The precise control of the MoSe\(_2\) thickness and its crystal orientation are the most important factors determining the solar cell efficiency. Based on this study, the formation of the MoSe\(_2\) layer depends on both Na diffusion from the substrate and the Se content, which are related to the sodium content on the substrate, back contact structure and the selenisation process.

All selenisation processes in this study were conducted at a substrate temperature of 580 °C for a duration of 12 minutes. From the experiments, it was observed that there is no difference in the thickness of the MoSe\(_2\) layer in the Se-poor CIGS absorbers, while the thickness of the MoSe\(_2\) layer varies significantly in the Se-rich CIGS absorbers. In particular, for the Se-poor absorber, the Se is fully consumed by the metallic CIG precursor to form the CIGS absorber. Thus, the formation of MoSe\(_2\) is inhibited due to the lack of Se in the absorber forming environment. In this situation, the CIGS/Mo interface is stable even during long-time annealing in a Se-deficient condition, regardless of the amount of Na diffusing from the substrate.

In contrast, in Se-rich absorbers, Na\(_2\)Se forming at both the Mo/CIGS interface and the CIGS grain boundaries tend to increase the MoSe\(_2\) formation probability. In this case, the excessive Se reacts with the Na diffusing from the substrate to form Na\(_2\)Se\(_x\), which is usually located at the CIGS/Mo interface as a Se source and catalyses the MoSe\(_2\) formation. When there is insufficient or no Na available to form Na\(_2\)Se\(_x\), it is less probable to form MoSe\(_2\) by the direct reaction between Se and Mo atoms. Hence, a very thin MoSe\(_2\) layer is found in the Se-rich samples made on rear contacts with low Na concentrations, for example the Na-free substrates and SLG substrates with SiN acting as a good Na diffusion barrier layer. A thick MoSe\(_2\) layer is only observed in samples with high Na diffusion from the substrate, and its thickness increases with increasing Na content in the glass substrate. However, if the MoSe\(_2\) layer gets too thick, the series resistance tends to increase and the solar cell efficiency was found to decrease.

In this study, the influence of the sodium diffusion from the substrate on the properties of two-step process formed CIGS absorber using elemental Se vapour are investigated. Using a TiN layer on SLG glass as a barrier layer to adjust Na diffusion, the CIGS film shows the best uniformity in grain size, indicating the good control of Na diffusion from the SLG glass substrate. The TiN barrier layers can partially reduce the Na diffusion, and thus enhance Ga and In inter-diffusion to form large CIGS grains. We also find that both Na and Se play crucial roles in the formation of the MoSe\(_2\) intermediate layer. A Se-rich atmosphere is required to grow a thick MoSe\(_2\) layer, while a Se-poor CIGS formation process leads to a thin MoSe\(_2\) layer, regardless of the Na content in the substrates. Since the series resistance of the CIGS solar cells increases with increasing MoSe\(_2\) layer thickness, an intermediate MoSe\(_2\) layer thickness will give the best device performance. Thus, the Se flow in the two-step CIGS formation process must be sufficiently high to obtain high-efficiency CIGS solar cells.

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Author Contributions

Dr. Weimin Li prepared the Mo back contact and CIGS precursor, measured the SEM-EDX of the CIGS absorbers, and wrote the main manuscript text; Dr. Xia Yan measured the XRF and XRZ of the CIGS absorbers; Prof. Armin Aberle wrote the introduction of the manuscript and the conclusion of the manuscript; Dr. Selvaraj Venkataraj prepared the CIGS absorber and Figures 1 and 4. He also contributed to prepare the abstract of the manuscript.

Additional Information

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