PROPERTIES OF Cd and Zn PARTIAL ELECTROLYTE TREATED CIGS SOLAR CELLS

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ABSTRACT

We study the influence of Cd partial baths on the photovoltaic properties of CuInGaSe₂ (CIGS) and CuInGaSSe₂ (CIGSS) thin film absorbers. We find that efficient solar cells can be fabricated by this treatment, and we compare their properties with those containing CdS window layers grown by chemical bath deposition. The results suggest that Cd plays a dominant role in establishing efficient photovoltaic junctions in CuInSe₂ alloys. Micron scale photoluminescence scans show non-uniformity along the length probed. Cd treatment quenches one of the luminescence transitions, which indicates a change in shallow acceptor level density. We present a model that helps to explain the evolution of photovoltaic action.

INTRODUCTION

CuInSe₂ (CIS) thin films and their alloys with CuGaSe₂ and CuInS₂ have been the subject of study in the photovoltaic community because of their suitability for low-cost power generation. Excellent cell efficiencies are demonstrated routinely in laboratory scale devices [1], and large area modules are entering the marketplace [2]. A majority of the laboratory scale devices are fabricated from CIGS alloys deposited by co-evaporation of the elements. Large area modules are made from CIGSS absorbers derived from sputtered metal precursors and subsequent reaction with the chalcogens. The two examples provide a sharp contrast in terms of the preparation of the absorber materials. In any further analysis of the physics of the devices, one must take into account the differences in the preparation methods and the resulting absorber properties. A reasonable starting point is that the absorbers are single-phase, chalcopyrite Cu(In,Ga)(SSe)₂ alloy thin films in which the Ga and/or S mole fractions are carefully controlled to be within tolerance limits, and the Cu composition is also controlled to avoid Cu chalcogenide impurities. In all of these materials, the junction formation is achieved by the growth of a thin CdS window layer by chemical bath deposition (CBD). CBD CdS processing produces the most efficient solar cells. However, the mechanism that forms the junction remains unclear. Historically, the CdS/CIGS interface has been treated as a “non-interacting,” or abrupt junction. There are a few reports [3,4] suggesting that the interface is intermixed, or may contain distinct compounds such as CdSe or CdIn₃Se₄. The temperature of CBD growth is deceptively low, but the chemical driving forces are powerful enough to cause dissolution of species and cause atomic movements [5]. This is also illustrated in the case of aqueous treatment of CIS in In-S solutions in which a thin CuInS₂ layer is grown on the surface [6]. Compositional changes such as those mentioned above have not been adequately treated in further analysis of the junction. A more serious omission is the potential role of electrically active impurities, such as Cd or Zn, which are known to produce efficient n-type conductivity [7] in the materials of interest. We have shown previously that the CBD reaction mixture containing only Cd, or a similar bath constituted with Zn, can be used to produce efficient photovoltaic cells in CuInSe₂ thin films [5]. In this report, we extend this work to high-efficiency CIGS absorbers fabricated by the three-stage process [8], and CIGSS absorbers provided by Shell Solar Industries (SSI).

EXPERIMENTAL

CIGS thin films were grown by the three-stage process on Mo/soda lime glass substrates. The resulting absorber material was 2.5 µm thick, and it was p-type with a hole density of 1-3×10¹⁶ cm⁻³. The nominal composition was CuInₓGa₁₋ₓSe₂. The Cu concentration in the near-surface region was carefully controlled so that the overall Cu/(In+Ga) ratio is less than unity. Absorbers provided by SSI were graded CIGSS materials. CdS thin films were deposited from an aqueous solution containing 0.0015M CdSO₄, 1.5M NH₄OH, and 0.0075M thiourea. The temperature of the bath was raised from room temperature to 60°C, and the samples were held in the bath for 15 min, during which a 40 to 60 nm CdS film was deposited. Partial electrolyte treatments, designated as Cd PE and Zn PE, were performed by immersing the samples in a bath similar to the CdS deposition, but excluding thiourea. The maximum temperature during the Cd PE treatment was 80°C, and the total immersion time was 22 min. Solar cells were fabricated by depositing a two-layer ZnO film. The first layer of 80 to 100 nm was deposited from a pure ZnO target, and the second layer of 200 nm was deposited from an Al₂O₃ doped target. Ni/Al contacts were evaporated on the ZnO. Current-voltage characteristics of the solar cells were measured in dark and under light simulated to correspond to AM1.5 Global spectrum. Photoluminescence (PL) was excited by a He-Ne laser and collected by a InGaAs diode array detector. The samples
were held at 4 K in a liquid-He cold finger cryostat, and the excitation level was about 120 nW.

RESULTS AND DISCUSSION

Fig. 1 shows the light current-voltage (J-V) characteristics of three types of solar cells fabricated from NREL CIGS absorbers. Sputter deposition of ZnO directly on the CIGS (open circles) does not produce a well-behaved diode. The Cd PE treatment prior to the ZnO deposition (solid circles) results in a marked improvement, and an efficient solar cell is achieved. A comparison with the solar cell with a CdS window layer (solid line) demonstrates the beneficial effect of Cd in creating a strong electric field within the device.

Fig. 2 is an example of these effects in CIGSS graded absorbers prepared by SSI in a manner that has no similarity to the CIGS case. The trends are similar to the CIGS case, and demonstrate that Cd is equally effective in "real-world" materials.

Table 1. Light J-V Parameters of Solar Cells Made Using Cd PE and Zn PE. Device Structure is MgF2/ZnO/CIGS.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>$V_{oc}$ (V)</th>
<th>$J_{sc}$ (mA/cm$^2$)</th>
<th>FF</th>
<th>Efficiency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cd PE</td>
<td>0.636</td>
<td>34.6</td>
<td>0.72</td>
<td>15.7</td>
</tr>
<tr>
<td>Zn PE</td>
<td>0.558</td>
<td>36.3</td>
<td>0.70</td>
<td>14.2</td>
</tr>
</tbody>
</table>

Internal quantum efficiency of Cd PE and CdS cells fabricated on the same absorber are shown in Fig. 3. The Cd PE cell exhibits a superior collection in the short wavelengths due to the absence of absorption and collection losses related to the CdS. The long wavelength response is maximized as well. The differences in the long wavelength edge are due to the differences in the Ga content. Capacitance-voltage characteristics of the CdS and Cd PE cells were studied as a function of post-heating after the deposition of CdS or Cd PE treatment. The carrier density profiles are changed by short anneals in air at 200°C. In the Cd PE cells, we have observed a lower carrier density and, hence, a larger depletion width. This is likely the result of higher process temperature and a consequent compensation of holes by Cd donors. The red response is enhanced as a result of field assisted collection. This aspect is discussed in another paper at this conference [9].

In general, the efficiency of the Cd PE cells are lower than the CdS cells fabricated on the same absorbers. $V_{oc}$'s and fill factors are lower, whereas the current density is higher. To understand the potential causes for the dif-
ferences, we measured the J-V characteristics and these are shown in Fig. 4. An analysis of the dark and light J-V characteristics was performed, and a summary is shown in Table 2. The reverse saturation current density, $J_0$, of the Cd PE cell in the dark is two orders of magnitude higher than that of the CdS cell, and this difference is maintained under illumination as well. The diode factor for the Cd PE cells is also higher. These can be ascribed to the higher interface recombination or to changes in the current transport mechanisms. The CdS deposition affords the growth of a lattice matched window layer, compositional grading at the interface, and passivation of defect states near the interface. The absence of these beneficial effects and the inclusion of a sputter-deposited contact close to the high field region can contribute to a higher $J_0$ in the Cd PE cells, and hence to a reduction in the $V_{oc}$ and fill factor. However, the difference in performance between the two is much smaller compared to the case when no Cd treatment is given. The demonstration of ~16% devices by using Cd alone clearly establishes the dominant role of Cd.

Table 2. Diode Quality Factor, $A$, and Reverse Saturation Current Density, $J_0$ (A/cm$^2$), of CdS and Cd PE cells.

<table>
<thead>
<tr>
<th>Type</th>
<th>$A$ (dark)</th>
<th>$J_0$ (dark)</th>
<th>$A$ (light)</th>
<th>$J_0$ (light)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CdS</td>
<td>1.7</td>
<td>$10^{-9}$</td>
<td>1.7</td>
<td>$10^{-8}$</td>
</tr>
<tr>
<td>Cd PE</td>
<td>1.9</td>
<td>$10^{-7}$</td>
<td>2.0</td>
<td>$10^{-7}$</td>
</tr>
</tbody>
</table>

Fig. 4. Dark J-V characteristics of Cd PE and CdS solar cells.

Fig. 5. PL spectra of CIGS thin film treated in NH$_4$OH.

Fig. 6. PL spectra of CIGS thin film treated in Cd PE.

The PL intensities exhibit variation along the length probed, which is attributed to the local variations in composition between the grains and intergrain regions. The spectral dependence shows two broad peaks centered around 1150 nm and 1250 nm for the ammonium hydroxide case. It is well known that the two transitions are donor-to-acceptor pair transitions between shallow energy levels [10]. In the Cd PE sample, the higher energy emission is quenched by the action of Cd. This is likely due to the neutralization of a shallow acceptor-like defect. The filling of Cu vacancies by Cd, or the out-diffusion of Se are the most probable chemical changes that are likely to occur. We can see that changes in the ionization states of shallow levels can lead to a change in the compensation ratio and the conductivity type of the region close to the interface. Similar observations were made in CuInSe$_2$ single crystals in which the electrical activity of Cd and Zn dopants was closely linked to changes in luminescence, and was associated with a downshift of the peak emission wavelength [11].
In the above, we have shown a series of examples that clearly demonstrate the strong effect of Cd in creating efficient solar cells in CIS alloy thin films prepared by vastly different methods (i.e., co-evaporation and selenization). We attribute this to the ability of Cd and Zn to produce n-type doping or a strong inversion of the surface region. This is most logical explanation for the observed effects. Experimental evidence for this was provided in the studies of Cd and Zn diffusion in CulnSe2 single crystals [7,11]. An appreciation of the differences between the single crystal devices and the present day thin film solar cells is essential. However, one must not dismiss the obvious connection. The results shown here suggest that the ability to produce n-type doping could well be at the heart of the CIS alloy thin film solar cells. The evolution of the solar cell can be viewed as a combination of the following three components:

1. Formation of Cd doped, n-type region in the absorber surface region.
2. Compositional mixing and interface grading. Band discontinuities or spikes can be softened by grading. This can minimize the effect of defect states in the interface region and reduce the interface recombination velocity. Shifting the peak of the electric field away from the metallurgical interface can also mitigate the effect of interface recombination. Sulfur can play a useful role in passivating dangling bonds.
3. Wide band gap window layer (CdS) protects the electrical junction from further processing, and facilitates shaping of the electric field within the n-region of the device.

It is clear that the optimal combination of all three components would give rise to an efficient solar cell. Indeed, the same picture can be used to explain the behavior of solar cells made using Zn compounds in CBD, CVD, or ALE environments. The ability to dope CIS alloys n-type, and the inability to do the same in CuGaSe2 might offer clues to the differences in their performance.

CONCLUSIONS

The use of Cd and Zn PE treatments are shown to produce efficient solar cells in CIGS and CIGSS thin film materials. The properties of these cells are compared with those with CdS window layers, and the causes for the differences are identified. We show that the high efficiencies achieved in CBD CdS/CIGS thin film solar cells arise from a set of key interactions that could produce n-type doping or inversion, compositional grading, and interface passivation. The results are shown to be applicable to laboratory samples and industrial scale samples from which modules are fabricated. We present a model for the operation of CdS/CIGS devices.

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REFERENCES