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The fox and the hound: in-depth and in-grain Na doping and Ga grading in Cu(In,Ga)Se₂ solar cells†

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Cu(In,Ga)(S,Se)₂ (CIGS) thin film solar cells require appropriate depth and lateral distributions of alkali metal dopants and gallium to attain world record photovoltaic energy conversion. The two requirements are interdependent because sodium is known to hamper In/Ga interdiffusion in polycrystalline films. However, such a fact is challenged by recent findings where sodium appears to enhance In/Ga interdiffusion in monocrystalline films. This contribution reviews closely the two cases to the benefits of grain boundary engineering in CIGS. A computational model reveals why Na induces In accumulation at CIGS grain boundaries, confining Ga to grain interiors. The positive technological implications for wider gap chalcopyrites are stressed.

Introduction

Solar power is central to the sustainable growth of humankind. With 23.35% efficiency, Cu(In,Ga)(S,Se)₂ (CIGS) solar cells display the highest ever reported light to electricity conversion of all commercially available thin film photovoltaic technologies.² Excitingly, there is still room for improvement at both cell and module levels.³

Thanks to the possibility to alloy CuInSe₂ (CIS) and CuGaSe₂ (CGS), the CIGS band-gap can be tuned from *ca.* 1.0 eV (CIS) to *ca.* 1.7 eV (CGS), allowing for both single and double junction device fabrication.⁴

Commercial CIGS solar cells are currently based on a highly engineered single junction configuration that is the result of cumulative knowledge gathered during 30+ years of R&D. Fig. 1 shows the historical progress of record power conversion efficiency for laboratory CIGS cells since the inception.⁵ Throughout this (sometimes serendipitous) journey three strategies have played a crucial role for technological success:

(a) The controlled gradient of Ga concentration relative to In through the depth of the film,⁶ Fig. 1a.

(b) The deliberate doping with Na,⁷ Fig. 1b.

(c) The post deposition treatment (PDT) with heavier alkali metal fluorides (KF,^{8–10} RbF and CsF¹¹), Fig. 1c.

Soon after Na was found to be beneficial for cell efficiency, it was also shown to impede the interdiffusion of In and Ga. Later, K was also observed to hinder In/Ga interdiffusion.¹² This effect has been exploited to tailor specifically the Ga grading in films produced by reactive annealing of sputtered metallic precursors.¹³ Therefore, the two above strategies are closely entangled, although they have been historically pursued in parallel without attempting to purposely engineer one another.

The beneficial effect of heavy alkali PDT was first connected to the changes in the absorber surface chemistry and morphology.^{14,15} A phase containing K–In–Se with a wider band gap was observed which was connected to the reduced absorber–buffer interface recombination and increased open-circuit voltage.^{15,16} Moreover, the improved surface morphology enables a thinner buffer layer resulting in reduced optical losses.¹⁴ Most recently, based on the observation of similar open-circuit voltage improvements, instead of different surface conditions it was concluded that the improvements in the bulk absorber play the most important role.¹⁷ Namely, heavy alkali metal atoms can diffuse along grain boundaries accumulating on them and also substitute the proximate Na atoms.^{14,16} This leads to the passivation of charged defects at the grain boundaries which reduces the concentration of tail states inside the grains. The result is reduced recombination and increased open-circuit voltage.¹⁷

This contribution focuses on the influence of Na and K on the interdiffusion of In and Ga. Generally, it is well known that grain boundaries in polycrystalline materials have a strong influence on mass transport properties¹⁸ and the same is true for Na along CIGS grain boundaries.¹⁹ Therefore, it is important to understand if the hindering effect of Na and K on the interdiffusion of In and Ga in CIGS is related to transport at the grain boundaries or is intrinsically related to bulk CIGS properties.

In such cases, a constructionist approach can reduce the complex interdependency between variables, rendering the

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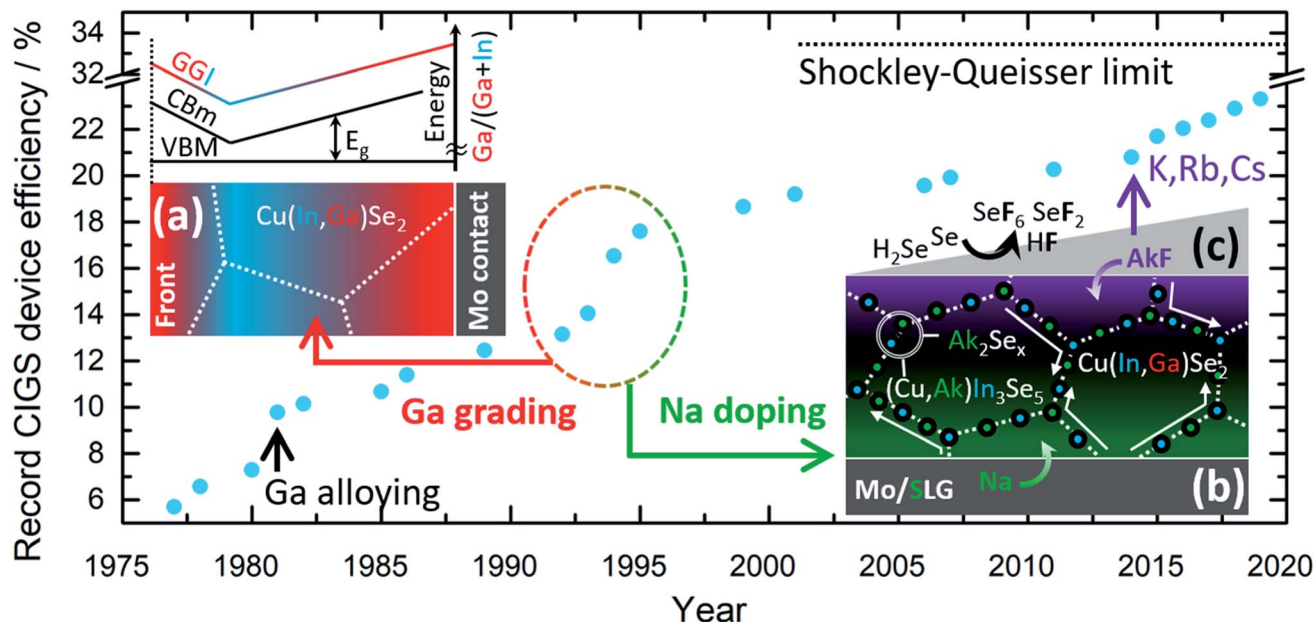


Fig. 1 Historical progress of CIGS solar cell efficiency at laboratory scale with emphasis on the major technological breakthroughs, shown as schematic insets with illustrative purposes: (a) Ga grading, (b) Na doping and (c) the more recent doping by heavier alkali metal post deposition treatments (PDT). Acronyms are as follows; GGI: atomic concentration ratio of Ga over the sum of Ga and In, VBM: valence band maximum, CBm: conduction band minimum, E_g : band-gap energy, AkF: generic alkali metal fluoride, SLG: soda-lime glass.

system more manageable, as demonstrated for macroscopic²⁰ and mesoscopic²¹ crystals.

Through a parallel approach, a recent study has shown that Na can enhance the interdiffusion of In and Ga in monocrystalline CIGS films, contrary to the polycrystalline case.¹ Precisely, Na appears to either enhance or impede the diffusion of In and Ga depending on the relative concentration of Na and Cu.²² This paper reviews Na doping and Ga grading in CIGS with a comparison between the historical and recently discovered interdependences.

Learning from CIGS' past breakthroughs

The fox: sodium doping

Today's CIGS technology relies on Na as an essential extrinsic dopant to ensure the best optoelectronic properties.^{8,23,24} The first encounter between Na and CIGS dates to 1993, when Hedström *et al.* observed that CIGS grown on soda-lime glass substrates yielded cells with substantially higher open circuit voltage and fill factor compared to CIGS grown on Na-free substrates.⁷ It was later established by Granata *et al.* that the Na concentration for optimum performance ranges between 0.05 at% and 0.5 at%.²⁵

The device's improvement induced by Na doping is attributed to increased p-type conductivity and defect passivation,²⁶ but the detailed underlying mechanism is still debated. It has been suggested that Na catalyzes the oxygenation of Se vacancies,²⁷ forms $\text{Na}_{(\text{In,Ga})}$ acceptors,²⁸ eliminates $(\text{In,Ga})_{\text{Cu}}$ donors²⁹ or increases the concentration of Cu vacancies (V_{Cu}) due to differential solubility of Na_{Cu} during CIGS growth and cool

down.³⁰ All these proposed mechanisms have strengths and weaknesses, but none of them can be claimed to unify all the experimental evidence and stand against theoretical scrutiny.

Indeed, the effects of Na doping in CIGS are not limited to electronic properties. Besides impeding atomic diffusion,^{7,12,31–39} Na also affects grain growth and texture.^{7,32,40–44}

These unexplained “symptoms” remain rather understudied, but, conceivably, represent the other side of the coin.⁴⁵ Studying them may add one crucial link to understand CIGS from the material viewpoint and help unlock the full potential of the technology towards the Shockley–Queisser limit.

The hound: gallium grading

Thanks to the nearly invariant CIS and CGS valence band levels, typical v-shaped gradients of $\text{Ga}/(\text{Ga} + \text{In})$ (GGI) through the depth of CIGS films correspond to a v-shaped conduction band level. Such a configuration has been beneficial because it acts as a back surface field, and is thought to ease the collection of photogenerated electrons while reducing interface recombination^{39,46} (Fig. 1a).

GGI depth gradients were first deliberately engineered at NREL by Contreras *et al.* in 1993 with the introduction of the 3-stage co-evaporation process.^{6,47–49} However, it soon became apparent that (steep) gradients tend to form spontaneously also during sequential processing (chalcogenization of metal precursor films) if Na is present during the growth.^{50–52} As such, the phenomenon concerns the two major industrial routes of commercial CIGS modules.

Simulations by Witte *et al.*³⁹ show that cell efficiency depends critically on the exact position of the notch point, with a trade-



off between open circuit voltage and short circuit current. In fact though, the minority carrier diffusion length in modern devices is sufficient to ensure good current collection without the need for v-shaped gradients.⁵³ Hence, the community will soon have to adjust the film growth conditions to allow for a modified outcome of In/Ga interdiffusion.

Besides the deliberate *depth* grading, accidental GGI gradients can also occur *laterally*, leading to so-called band-gap fluctuations.⁵⁴ These inhomogeneities are a source of voltage losses in record cells below 20% efficiency.^{55,56}

When going from lab-scale to large industrial areas the range of fluctuations increases from sub-micrometer to micrometer length scale. This is because absolute control of In and Ga lateral concentration in CIGS modules can be problematic.^{3,57} Such fluctuations can contribute to performance losses beyond the effect of parallel electrical connection,⁵⁸ as the module's minority carrier diffusion length is progressively increased.⁵⁶

The elusive friendship between the fox and the hound

The hindering effect of Na on In/Ga interdiffusion in polycrystalline CIGS films is supported by evidence gathered over 20 years by independent research groups using different experimental approaches.^{12,31–33,35–39,50,59} If Na is present during the growth, its actual source does not appear to be very important. Decreased In/Ga interdiffusion was observed regardless of the Cu/III ratio^{33,38} and of the Na origin such as: NaF pre-deposited on either Mo-coated soda-lime glass (SLG)/Al₂O₃ (ref. 31) or polyimide foils,³⁶ or supplied during the growth^{31,37} or during the selenization⁵⁹ or resulting from diffusion respectively out of soda-lime glass^{31–33,38,50} or enamel coated steel substrates,¹² when the films are compared to respective “Na-free” reference samples. Retrospectively, this is consistent with the high Na mobility in the CIGS lattice revealed both by theory and experiment.^{19,60} In other words, Na will diffuse and yield similar effects over its large diffusion length scale, regardless of the source location (front/back), if present during the growth.

However, larger-scale inhomogeneous distributions of alkali dopants during film growth may still translate into undesired GGI distributions in the final device, given the influence of the dopants on In/Ga diffusion. Since these fluctuations are

detrimental to performance, controlling the uniformity of sodium and potassium doping is crucial.^{26,61,62}

Understanding the interdependence between alkali metal doping and GGI gradients would then allow to take better control over the intermixing behaviour of In and Ga in CIGS (both laterally and in-depth) with positive repercussions on the technology. A deeper diffusion mechanistic knowledge will also prove useful when tackling changes in cell geometry from the conventional configuration towards thinner⁶³ and micro-pattern^{64,65} CIGS devices.

Rodriguez-Alvarez³⁸ and Witte *et al.*³⁹ offer an exhaustive account of the interdependence between Na doping and Ga grading in polycrystalline CIGS. Their studies reveal a clear relationship between Na and Cu, confirming that:

(a) Diffusion of In and Ga occurs through the Cu sublattice, as previously suggested by Schroeder *et al.*⁶⁶ (Fig. 2a);

(b) Na is likely to interfere with Cu vacancies, as inferred by Lundberg *et al.*³³ (Fig. 2b).

It is then commonly agreed that the hindering effect of Na at high temperature should be ascribed to a decreased concentration of Cu vacancies resulting from the capture of Na and formation of Na_{Cu} substitutional defects,³⁰ as originally proposed by Rudmann *et al.*³² An analogous argument was recently proposed by Cai *et al.* for potassium, with the formation of K_{Cu} substitutionals.¹³

In and Ga atoms occupy the same CIGS lattice sites and thus diffuse interdependently. Repins *et al.* were able to simulate GGI depth profiles in CIGS films obtained by the 3-stage process through a modification of the diffusion equations taking into account In/Ga atomic site sharing.⁶⁷ However, engineering GGI gradients by means of tuning the Na or K concentration gradients in the films requires knowledge of the migration paths also *via* the grain boundaries.

In fact, the accepted paradigm is challenged by newer findings on monocrystalline CIGS films.¹ These show that diffusion of Ga from GaAs substrates into epitaxial CIS films (*i.e.* free from high-angle grain boundaries) is substantially enhanced by the presence of Na (introduced even from the front surface, Fig. 2c). Given that these findings *seemingly* dispute an established body of literature, the observation was supported by several techniques: secondary ion mass

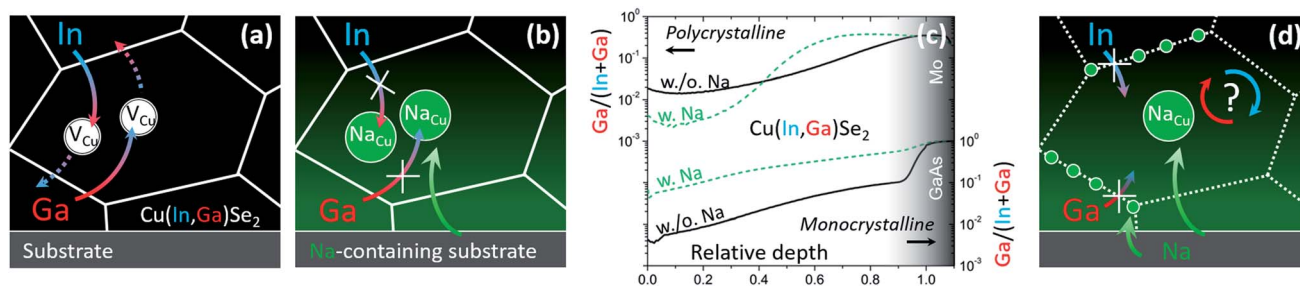


Fig. 2 (a) Schematics of In/Ga interdiffusion mediated by Cu vacancies in Na-free CIGS and (b) impeded interdiffusion according to the commonly accepted rationale. (c) Ga/(In + Ga) secondary ion mass spectrometry depth profiles of Na-free (solid black) and Na-containing (dashed green) polycrystalline (top)⁵⁹ and monocrystalline (bottom)¹ CIGS films. (d) Phenomenological interpretation following the findings in monocrystalline CIGS.¹



spectrometry, energy dispersive X-ray spectroscopy from scanning and transmission electron microscopes, nano-Auger electron spectroscopy, atom probe tomography, X-ray diffraction analysis, spectrophotometry, photoluminescence spectroscopy and Raman spectroscopy.

Sodium-induced atomic diffusion is not limited to CIGS matrix elements. Heavier alkali dopants also happen to be displaced to and from surfaces and grain boundaries, depending on temperature and relative concentration.^{8,10,68} This phenomenon is not clear yet, but it has obvious implications on the more recent strategy for efficiency enhancement with heavier alkali metals²⁴ (cf. Fig. 1c).

These newer findings do not contradict the established literature on polycrystalline films, they merely suggest that different diffusion mechanisms are likely to take place at CIGS grain interior and grain boundaries (Fig. 2d).²²

Defect migration dynamics in CIGS has been reported previously.^{13,30,39,60,69,70} However, due to the computational demand associated to sufficiently large unit cells, these studies are limited to the migration of individual species in bulk CIGS. Therefore, simulating the effects of grain boundaries and migration interdependence is not trivial.

Challenging the polycrystalline paradigm

Two hypotheses

Na is well known to influence CIGS grain growth. On the one hand, it is reported to *promote* the growth when used in rapid thermal processing⁴⁰ or when it is finely dispersed among the matrix elements.⁴² On the other, Na *impedes* the growth both during the (relatively high GGI stage) co-evaporation and during sequential processing.^{26,41,43,71}

Since grain growth is often a synonym of mass diffusion,⁷² it is useful to relate the Na-induced *crystallization* effects to the Na-induced *diffusion* effects.

Due to the chemical affinity between Na and Se, several sodium (poly)selenides are reported to form during CIGS growth.⁷³ These compounds occur as segregations at the grain boundaries, but during the growth they modulate selenium and copper activities,⁷⁴ acting as fluxing agents. Sodium effectively increases the copper activity in the intermediate Cu_xSe phases, by lowering the selenium availability. The enhanced copper diffusivity facilitates CIS crystallization by reaction between Cu_xSe and In_2Se_3 . Conversely, the hindered grain growth under relatively high GGI is explained by the sluggish reaction between Cu_xSe and Ga_2Se_3 , resulting in the common bilayer microstructure with strong GGI gradient.⁷⁴ It is then generally agreed that the resulting similarity of Na and Ga distributions is purely consequential.

The enhanced Ga indiffusion induced by Na in monocrystalline films¹ plays havoc with the current understanding of diffusion equilibria in polycrystalline CIGS. Since record cells are historically polycrystalline in nature, the study is easily referred to as irrelevant for the technology. However, besides the fact that monocrystalline CIGS cells have now reached notable efficiencies up to 20%,⁷⁵ the findings must be related to

the broader context if the community aspires to gain a deeper understanding of the fundamental material properties.

A phenomenological integration of the findings into the accepted paradigm passes through the simple schematics in Fig. 2d. There, the hindering effect of Na on In/Ga interdiffusion in polycrystalline films is suggested to originate from accumulation of Na *at the grain boundaries* and local drop of Cu vacancy concentration. As the vacancy concentration drops, so do the diffusivities of In and Ga *across the grain boundaries*, according to the very same rationale invoked by Rudmann *et al.*^{32,76}

Since diffusion along the grain boundaries is way faster than at grain interior,¹⁹ grain boundaries are likely to dominate the overall mass diffusion in polycrystalline CIGS. It follows that the enhanced diffusivity induced by Na at the interior of the grain is largely dwarfed by the hindered diffusion across the grain boundaries.

What remains unknown is why Na appears to enhance interdiffusion at grain interior in the first place.²² Here, it should be stressed that Na concentration within the grains is typically one order of magnitude lower than at grain boundaries.^{1,19,77} Therefore, it is not surprising to expect different diffusion mechanisms at the two locations. The schematics in Fig. 3 aim to reconcile established and new experimental evidence into a coherent hypothetical model, to be tested using the density functional theory (DFT).

The model considers the existence of supercomplex defects,¹ as first suggested by Zhang *et al.* in 1998,⁷⁸ and detected recently by Stokes *et al.*⁷⁹ These complexes are the result of coulombic interaction between copper vacancy acceptors V_{Cu}^- and indium/gallium on copper antisite donors $(\text{In,Ga})_{\text{Cu}}^{2+}$. The concentration of these complexes is ruled by an equilibrium that depends on the composition of the CIGS matrix (*i.e.* on the chemical potential of the matrix elements).

The Na incorporation and ensuing capture by CIGS copper vacancies⁶⁹ may lead to a mass action effect on the chemical equilibria of these complexes. If less vacancies are available for binding, the concentration of the complexes would drop, in accordance to Le Chatelier.

Indeed, Stanbery *et al.* suggested that Na uptake destabilizes $\{[\text{In}_{\text{Cu}}^{2+}]_2[\text{V}_{\text{Cu}}^-]\}$ neutral defects, easing In migration away from CIS bulk.^{80,81} In the same matrix, the stability of the corresponding Ga complex $\{[\text{Ga}_{\text{Cu}}^{2+}]_2[\text{V}_{\text{Cu}}^-]\}$ would clearly differ from that of $\{[\text{In}_{\text{Cu}}^{2+}]_2[\text{V}_{\text{Cu}}^-]\}$, due to the smaller Ga radius,⁸² which should result in different mass transport properties for In and Ga. The binding energy of the analogous Ga complex was reported to be 0.7 eV higher in CuGaSe_2 than that of $\{[\text{In}_{\text{Cu}}^{2+}]_2[\text{V}_{\text{Cu}}^-]\}$ in CIS.⁸³ It follows that $\{[\text{Ga}_{\text{Cu}}^{2+}]_2[\text{V}_{\text{Cu}}^-]\}$ should be less susceptible to Na-induced mass action in Ga-rich CIGS, compared to the In complex. The next section reveals if the hypothesis also applies to In and Ga in CIGS having the same composition.

If Na helps to dissolve In complexes, the following effects observed in *In-rich* CIGS are understood:

- (i) Na-induced increased p-type conductivity;³⁰
- (ii) In enrichment of grain boundaries, *i.e.* the consequential passivating character;⁸⁴
- (iii) Na-induced enhanced grain growth.^{40,42}





Fig. 3 Hypothetical model that attempts to reconcile the findings from polycrystalline and *monocrystalline* CIGS. From left to right. Legend describing the point defect and defect complexes discussed. Hypothetic effect of sodium doping on the chemical equilibria involving the defect complexes: whereas the more labile $\{[In_{Cu}^{2+}]2[V_{Cu}^{-}]\}$ gets dissolved unleashing copper vacancies, the same does not hold for $\{[Ga_{Cu}^{2+}]2[V_{Cu}^{-}]\}$. Induced preferential diffusion of indium to CIGS grain boundaries (with formation of Na-rich ODC phase) and retention of gallium at grain interior. The model is tested by DFT and discussed in the text.

Furthermore, if Na tends to cluster with Ga into vacancy supercomplexes, the following effects observed in *Ga-rich* CIGS are understood:

(i) Higher open circuit voltage deficit,⁸⁵ possibly due to higher compensation and increase of electrostatic potential fluctuations;

(ii) Less frequent Ga enrichment of grain boundaries, *i.e.* their detrimental nature for device efficiency;^{43,86}

(iii) Na-induced hindered grain growth.^{26,41,43}

A second hypothesis relates to the asymmetry between In and Ga diffusivity, as already suggested by Witte *et al.* in Na-free CIGS.³⁹ In this sense, the effect of Na on the migration dynamics of In and Ga has never been considered. Does the presence of Na atoms around the In and Ga diffusing species lower the migration barriers, *e.g.* due to reversible lattice distortions, as proposed in ref. 22? Is the impact equal for In and Ga or does the diffusion asymmetry still hold in the presence of Na dopants?

Testing the defect equilibria hypothesis. The binding energies of the $\{[III_{Cu}^{2+}]2[V_{Cu}^{-}]\}$ vacancy antisite complexes in CIS, as defined in (1), were calculated by DFT.

$$E_b([III_{Cu}^{2+}]2[V_{Cu}^{-}]) = E_f([III_{Cu}^{2+}]2[V_{Cu}^{-}]) - E_f([III_{Cu}^{2+}]) - 2E_f([V_{Cu}^{-}]), \quad (1)$$

where $E_f([In_{Cu}^{2+}]2[V_{Cu}^{-}])$, $E_f([In_{Cu}^{2+}])$, and $E_f([V_{Cu}^{-}])$ are the formation energies of the complex, In antisite, and Cu vacancy, respectively.

In order to assess the reliability of the results, several supercell sizes were considered with both the semilocal PBE⁸⁷ and hybrid HSE06 (ref. 88) (with parameters $\alpha = 0.25$ and $\omega = 0.20 \text{ \AA}^{-1}$) functionals. The calculations were performed by the VASP program package^{89,90} ensuring convergence with respect to the plane wave energy cutoff and the *k*-point sampling. The defect formation energies were obtained by using the equilibrium bulk lattice parameters and eqn (2).

$$E_f = E_{tot}^{defect} - E_{tot}^{bulk} - \sum_i n_i \mu_i + qE_F + E_{corr} \quad (2)$$

where E_{tot}^{defect} is the total energy of the supercell containing the defect, E_{tot}^{bulk} the total energy of the bulk supercell, μ_i is the

chemical potential of the atom of type *i*, n_i is the number of added atoms when creating the defect in the supercell, q is the charge state of the defect, and E_F is the Fermi level measured from the VBM. E_{corr} is an energy correction term accounting for the errors due to the finite size of the supercell containing a charged defect and is calculated with the method of Freysoldt, Neugebauer, and Van de Walle⁹¹. When calculating the complex binding energy, the terms involving the chemical potentials μ_i and Fermi level E_F cancel out. On the other hand, E_{corr} terms do not cancel, since they are always positive for charged defects and zero for neutral complexes. The atomic configuration minimizing the complex formation energy corresponds to the initial state of the migration step shown in the inset of Fig. 4b.

The results for the binding energies of the vacancy antisite complexes (Table 1) reveal similar magnitudes and trends obtained with both functionals. The converged binding energies are on the order of -0.6 eV to -0.7 eV and their uncertainty below 0.2 eV . The defect formation energies have converged within 0.1 eV , but their uncertainties do not cancel out when calculating the binding energies. The magnitudes of the binding energies indicate clearly that the complexes are stable, more so, compared to the HSE06 results by Pohl and Albe⁸³ who obtained a value of -0.29 eV using a different method for finite size correction and also a different value for the HSE06 parameter ω .

Regardless of the absolute magnitude of the complexes binding energies, Table 1 clearly shows that the difference of stability between $\{[In_{Cu}^{2+}]2[V_{Cu}^{-}]\}$ and $\{[Ga_{Cu}^{2+}]2[V_{Cu}^{-}]\}$ calculated by using the two functionals converges consistently towards a value less than 0.1 eV (the complex with Ga being more stable). It follows that the defect equilibria hypothesis cannot be supported on the basis of a large stability difference. The next subsection directs the focus to the differences in migration properties of In and Ga.

Testing the defect migration hypothesis. A series of DFT calculations has been performed for In and Ga migration barriers in bulk CIS within the Cu vacancy mechanism, *i.e.*, an In_{Cu} or Ga_{Cu} at a Cu site jumping to the neighboring Cu vacancy. Comparing the barriers for the two atoms in bulk CIS to those for breaking the $\{[In_{Cu}^{2+}]2[V_{Cu}^{-}]\}$ and $\{[Ga_{Cu}^{2+}]2[V_{Cu}^{-}]\}$ complexes before



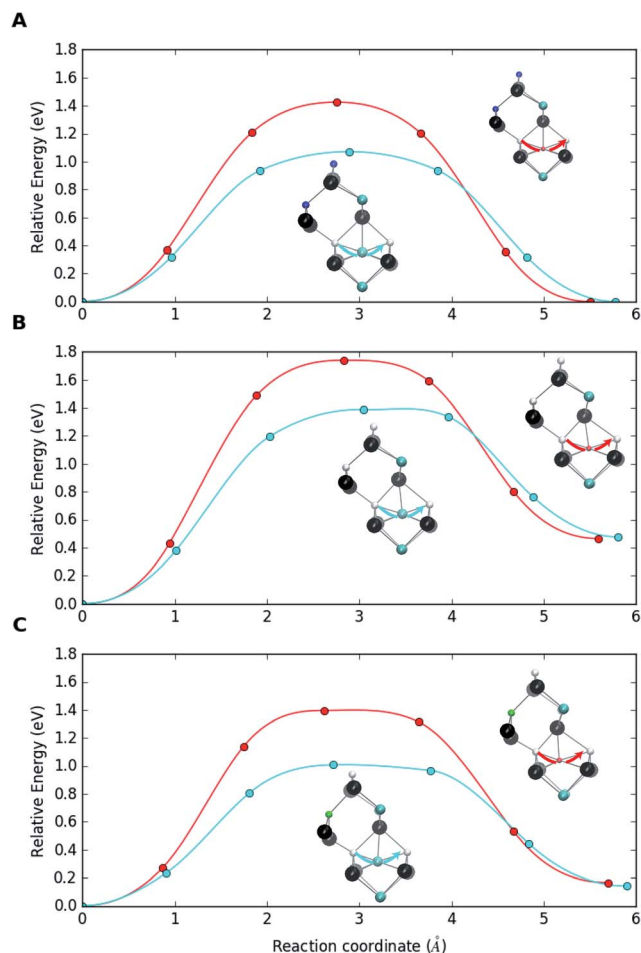


Fig. 4 (a) Migration dynamics of Ga atoms (red curves) and In atoms (cyan curves) in bulk CIS through Cu vacancies. (b) Corresponding migration barriers when In and Ga are part of $\{[(\text{In,Ga})_{\text{Cu}}^{2+}]_2[\text{V}_{\text{Cu}}^-]\}$ defect complexes and (c) in the presence of Na, *i.e.* as $\{[(\text{In,Ga})_{\text{Cu}}^{2+}][\text{V}_{\text{Cu}}^-][\text{Na}_{\text{Cu}}^0]\}$ defect supercomplexes. The structures are shown at the saddle point and truncated for visualization with Cu (blue), In (cyan), Ga (red), Se (black), Na (green) atoms and vacancy sites (white).

and after the complex has trapped a substitutional Na_{Cu} enables us to draw conclusions about the asymmetry of In and Ga behaviors in interaction with Na. In these heavy calculations the PBE functional and the supercell of 64 atoms had to be used.

Table 1 Binding energies (eV) of vacancy antisite complexes in CIS. HSE06 simulation for the 216 atom supercell were beyond the available computer resources

Supercell size (atoms)	Complex	PBE ^a	HSE06 ^b
64	$[\text{In}_{\text{Cu}}^{2+}]_2[\text{V}_{\text{Cu}}^-]$	-0.41	-0.35
	$[\text{Ga}_{\text{Cu}}^{2+}]_2[\text{V}_{\text{Cu}}^-]$	-0.52	-0.47
108	$[\text{In}_{\text{Cu}}^{2+}]_2[\text{V}_{\text{Cu}}^-]$	-0.49	-0.55
	$[\text{Ga}_{\text{Cu}}^{2+}]_2[\text{V}_{\text{Cu}}^-]$	-0.54	-0.62
216	$[\text{In}_{\text{Cu}}^{2+}]_2[\text{V}_{\text{Cu}}^-]$	-0.63	
	$[\text{Ga}_{\text{Cu}}^{2+}]_2[\text{V}_{\text{Cu}}^-]$	-0.67	

^a Lattice dimensions: $a = b = 5.88 \text{ \AA}$, $c = 5.91 \text{ \AA}$. ^b Lattice dimensions: $a = b = 5.82 \text{ \AA}$, $c = 5.85 \text{ \AA}$.

The diffusion barriers were calculated by employing the climbing image nudged elastic band (CI-NEB) method.⁹²

Fig. 4a shows the migration barriers for In and Ga in bulk Na-free CIS. The heights of the barriers, 1.07 eV and 1.43 eV, respectively, agree very well with those of Witte *et al.*³⁹ The smaller size of the Ga atom means that it is more deeply bound to the potential minima than the larger In atom, which also explains the higher stability of the complex with Ga. Fig. 4b depicts the situation when an In or Ga antisite of the $\{[(\text{III})_{\text{Cu}}^{2+}]_2[\text{V}_{\text{Cu}}^-]\}$ complex jumps to a neighboring Cu vacancy. The barriers in this breaking process, 1.39 eV and 1.74 eV, respectively, are clearly larger than those for the bulk migration. This is a consequence of the complexes stability, *i.e.* of the Coulomb attraction between negative vacancies and positive antisites. Note that, in the reverse process of antisite returning closer to the two vacancies, the barriers are similar to those in the bulk. Finally, Fig. 4c shows the corresponding barriers when the original complexes have trapped a Na atom to become $\{[(\text{III})_{\text{Cu}}^{2+}][\text{V}_{\text{Cu}}^-][\text{Na}_{\text{Cu}}^0]\}$ supercomplexes. The breaking barriers, 1.01 eV and 1.40 eV for In and Ga, respectively, are even slightly smaller than the migration barriers in bulk CIS. This indicates that trapping a Na atom makes the complex susceptible to breaking *via* the Cu vacancy migration process, which releases $\text{III}_{\text{Cu}}^{2+}$ to diffuse in the bulk CIS. It should be noted that a sufficient amount of energy could be available for breaking the $\{[(\text{III})_{\text{Cu}}^{2+}]_2[\text{V}_{\text{Cu}}^-]\}$ complex, especially for the In case with the lower breaking barrier. Namely, the calculations reveal that trapping an interstitial Na to a Cu vacancy releases *ca.* 1 eV of energy, nearly irrespective to the environment of the vacancy (Ga or In) and whether the substitutional Na lies next to the complex or is separated from it.

These calculations corroborate the mechanism of dopant-induced migration catalysis proposed in ref. 22, wherein sodium enhances atomic diffusivity of other atoms within the metal sublattice due to the Frank–Turnbull dopant migration. The above migration barrier values extend the asymmetry between In and Ga diffusion in the Cu-deficient CIS matrix also to Na-doped material. This asymmetry should reasonably hold also for standard Ga-poor CIGS and could explain the observed accumulation of In and depletion of Ga at CIGS grain boundaries and surfaces.

The model does not exclude the theory of Na-induced CIS crystallization,⁷⁴ but offers an additional insight on the root causes of the hampered growth in CGS. Furthermore, the premise offers a convenient explanation for the detrimental nature of the grain boundaries in Ga-rich CIGS, compared to conventional CIGS.^{43,86}

As vacancy antisite complexes are stable, by lowering the In migration barrier, Na allows the In excess to migrate to its equilibrium location (the grain boundaries). This has long been regarded as a beneficial passivating effect, due to band gap widening at grain boundaries and consequent reduction of charge carrier recombination.⁸⁴

However, even in the presence of Na, the Ga migration barrier remains larger compared to In, favouring the tendency for Ga to stay at grain interior. It follows that grain boundaries remain comparatively poorer from Ga than In. This translates



into Ga-poor grain boundaries in Ga-rich CIGS. The consequently reduced band gap widening at grain boundaries is then unable to prevent parasitic Shockley-Read-Hall recombination, giving a possible explanation for the large deficit of open circuit voltage in Ga-rich CIGS.⁸⁵

Conclusions and outlook

The current CIGS technology relies on accurate Na doping and GGI grading within and across each grain. Na doping hinders In/Ga interdiffusion in polycrystalline films, but can enhance it in monocrystalline films. It appears that Na impedes the diffusion across the grain boundaries, but enhances it within the grains, which is consistent with the micron scale lateral inhomogeneities responsible for band gap fluctuations in early record cells and possibly in current modules.^{39,61,93}

Yet, sodium doping of CIGS is responsible for complex diffusion interdependences with matrix elements and heavier alkali metal dopants that are poorly understood.^{1,8,10,68} This study tackles the diffusivity interdependence between sodium dopants and In/Ga atoms in CIS proposed in ref. 22. It reveals that Cu vacancies cluster with In and Ga antisites but the capture of sodium lowers the corresponding migration barriers of In and Ga. Furthermore, also in Na-doped material, Ga is shown to be a slower diffusing species than In, in line with the evidence of Ga-depleted grain boundaries in CIGS, which is a possible source of SRH recombination in wider-gap CIGS.

More theoretical and experimental efforts devoted to dopant-induced migration catalysis²² will surely benefit not only the CIGS technology, but the chalcogenide materials communities in general, including those working on topological insulators. These communities should find the approach proposed here a useful tool for grain boundary engineering.

Conflicts of interest

There are no conflicts to declare.

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