

Origin of passivation in hole-selective transition metal oxides for crystalline silicon heterojunction solar cells

Luis G. Gerling^{a,b,*}, Cristobal Voz^a, Ramón Alcubilla^{a,b}, Joaquim Puigdollers^{a,b}

^a Electronic Engineering Department, Universitat Politècnica de Catalunya, Jordi Girona 1-3, 08034, Barcelona, Spain

^b Centre de Recerca en Nanoenginyeria, Pascual i Vila 15, 08028, Barcelona, Spain

* *Corresponding author. Tel.: +34 93 401 1002; fax: +34 93 401 6756.*

E-mail address: guillermo.gerling@upc.edu

ABSTRACT

Transition metal oxides (TMOs) have recently attracted interest as an alternative to boron/phosphorous doped layers in crystalline silicon heterojunction solar cells. In this work, the interface between n-type c-Si (n-Si) and three thermally evaporated TMOs (MoO₃, WO₃ and V₂O₅) was investigated by transmission electron microscopy and secondary ion-mass/x-ray photoelectron spectroscopy. For the oxides studied, chemical passivation of n-Si was attributed to an ultra-thin (1.9 – 2.8 nm) SiO_{x-1.5} interlayer formed by chemical reaction, leaving oxygen-deficient species (MoO, WO₂ and VO₂) as byproducts. Field-effect passivation was also inferred from the inversion (hole-rich) layer induced on the n-Si surface, a result of Fermi level alignment between two materials with dissimilar electrochemical potentials (work function delta $\Delta\phi \geq 1$ eV). Therefore, the hole-selective and passivating functionality of these TMOs, in addition to their ambient temperature processing, could prove an effective means to lower cost and simplify solar cell processing.

I. INTRODUCTION

As today's photovoltaic market continues to be dominated by crystalline silicon (c-Si) technology, two cost-reduction efforts are being contemplated: thinner silicon wafers (< 100 μm) and the use of novel materials deposited at low temperatures with high throughput processes. Maximization of the solar cell open-circuit voltage (V_{OC}) to record values of 750 mV [1] has been possible by combining two principles of solar cell design [2]: (1) chemical passivation of surface dangling bonds with hydrogenated amorphous silicon (a-Si:H) and other dielectrics (SiO_2 , Al_2O_3 , Si_3N_4 , SiC); and (2) field-effect passivation by inducing a carrier-concentration gradient via electron- and hole-selective contacts, traditionally achieved by phosphorous/boron doping. Eventually, the ideal and simplest strategy would require a *passivating/selective contact*, dual-functionality materials that effectively passivate the c-Si surface while selectively conduct a specific charge carrier [3,4].

In parallel, thin-film photovoltaics including organic and perovskite solar cells, have introduced a wide variety of electron- and hole-selective materials whose optoelectronic properties are comparable or superior to the standard p- and n-doped layers used in c-Si. Despite being quite different from silicon's properties and technology, such materials have been successfully applied to c-Si heterojunction devices by low temperature ($T < 200^\circ\text{C}$) and solution-based processes, including organics [5], alkali salts [6,7] and transition metal oxides (TMOs) [8-18]. TMOs offer themselves as excellent candidates to substitute traditional c-Si dopants given their wide range of work function values (3 – 7 eV) and marked p- or n-type semiconductivity [19]. Additionally, their semi-insulating properties allow for passivation of the c-Si surface without compromising carrier-conductivity, although their passivation potential in abrupt TMO/c-Si heterojunctions is limited to experimental V_{OC} values of ~680 mV [11,20]. This has prompted the use of a-Si:H

passivating interlayers in order to surpass the 700 mV barrier, as has been demonstrated by a 725 mV V_{OC} and a 22.5% conversion efficiency device with MoO_3 hole contacts [21]. Therefore, a detailed understanding of the surface passivation mechanisms occurring at the TMO/c-Si interface (either by chemical bonding, charge selectivity or both) is still needed to potentiate the use of these passivating/carrier-selective materials.

In this work we will explain the origin of passivation in three TMO/n-type c-Si (n-Si) heterojunctions, molybdenum trioxide (MoO_3), tungsten trioxide (WO_3) and vanadium pentoxide (V_2O_5), deposited by vacuum thermal evaporation. The heterojunction interface will be characterized in detail by microscopy (TEM) and spectroscopy (ToF-SIMS, XPS) techniques, establishing two contributions to c-Si surface passivation: 1) through chemical reaction between c-Si and TMOs; and 2) via carrier-concentration gradients established during Fermi level alignment under equilibrium.

II. EXPERIMENTAL

Sample structures were fabricated on polished float-zone Si wafers (100 orientation) with a thickness of 280 μm and 2.5 $\Omega\text{-cm}$ resistivity. After RCA standard cleaning, wafers were dipped in 1% HF during 1 min to etch SiO_2 away. In this step, the surface Si atoms become bonded to hydrogen to form a continuous H-terminated surface. Unfortunately, this configuration is unstable and will be substituted by an ultra-thin native SiO_2 layer after only a few minutes [22]. Therefore, special care was taken to avoid a long air exposure (<30 seconds) by transferring the HF-cleaned wafers into a N_2 glovebox before TMO deposition. The three transition metal oxides, MoO_3 , WO_3 and V_2O_5 (powdered 99.99%, Sigma Aldrich) were thermally evaporated under vacuum ($<8 \times 10^{-6}$ mbar) from a tantalum boat at a rate ~ 0.2 $\text{\AA}/\text{s}$. Films thicknesses were measured by quartz crystal micro-balance at 70, 25 and 5 nm for MoO_3 , WO_3 and V_2O_5 respectively. During the evaporation process, heating

of the Si substrate was negligible and remained at ambient temperature. After a brief air exposure, samples were transferred to a RF-magnetron sputtering chamber for deposition of an 80 nm indium-tin oxide (ITO) film, under an argon partial pressure of 3.5×10^{-3} mbar and 50 W power. This ITO layer, which works as a transparent conducting layer in a TMO/n-Si solar cell, protects the layers beneath from degradation and work function changes [23].

These ITO/TMO/n-Si structures were first characterized by High Resolution Transmission Electron Microscopy (HR-TEM), using lamellas prepared by Focused Ion Beam (FIB) milling with Ga^+ ions (Carl Zeiss NEON40). Cross-sectional HR-TEM imaging (Jeol 2010F) was done with a field-emission electron source at 200 KV and lattice resolution. Secondly, Time of Flight Secondary Ion-Mass Spectroscopy (ToF-SIMS) depth profiling (IONTOF Model IV) was performed by sputtering a sampling area of $300 \times 300 \mu\text{m}^2$ with Cs^+ ions (1 keV) at high vacuum (3.8×10^{-8} mbar). The analysis of the sputtered secondary ions (negative) was done with a Bi^{+1} source (25 keV) and a time resolution of 200 ps. Charge compensation (<20 eV) was applied via electron flooding, given the semi-insulating character of TMOs.

For the X-ray Photoelectron Spectroscopy (XPS) scans, separate samples were prepared *ex-situ* by thermally evaporating V_2O_5 at different thicknesses (1 – 8 nm) on n-Si substrates (same conditioning as before). After oxide evaporation, each sample was transferred from the glovebox into the XPS chamber after a brief air exposure. The XPS analysis (SPECS, Phoibos 150 detector) was performed at 3×10^{-9} mbar using a non-monochromatic Al-K α source (1486.6 eV) and a 90° incidence angle guaranteeing a probing depth of at least 10 nm. Deconvolution of the core level spectra was done by Gaussian-Lorentzian fittings after Shirley background subtraction, while percentage atomic concentrations were calculated from the integration of the fitted peak areas. When

quantifying the oxygen content, correction for adventitious carbon C–OH and C–O contamination (<15% atomic C content) was performed. Unless stated otherwise, all binding energies were referenced to the Si2p^{3/2} peak (99.5 eV) of the clean Si sample.

III. RESULTS AND DISCUSSION

A. Interlayer in TMO/c-Si interfaces

As a first approach, detailed imaging of the fabricated ITO/TMO/c-Si structures was performed by HR-TEM, revealing the presence of an unknown 1 – 2 nm thick interfacial layer between the TMOs and the HF-cleaned (and therefore SiO₂ free) c-Si surface. This interlayer (IL), amorphous in nature, is clearly distinguished at the lowest magnifications [Fig. 1(a)] because of its high contrast to the adjacent materials. Its thickness varies among the three TMOs [Fig. 1(b) – (d)] but is relatively uniform and without pinholes. Moreover, it shows an abrupt interface with c-Si and a somewhat diffused one with the oxides. The maximum IL thicknesses are ~2.8 nm (WO₃), ~2.5 nm (MoO₃) and ~2.2 nm (V₂O₅), all of which are greater than the maximum threshold for tunneling (~1.2 nm) [2], suggesting another charge conduction mechanism must be present.

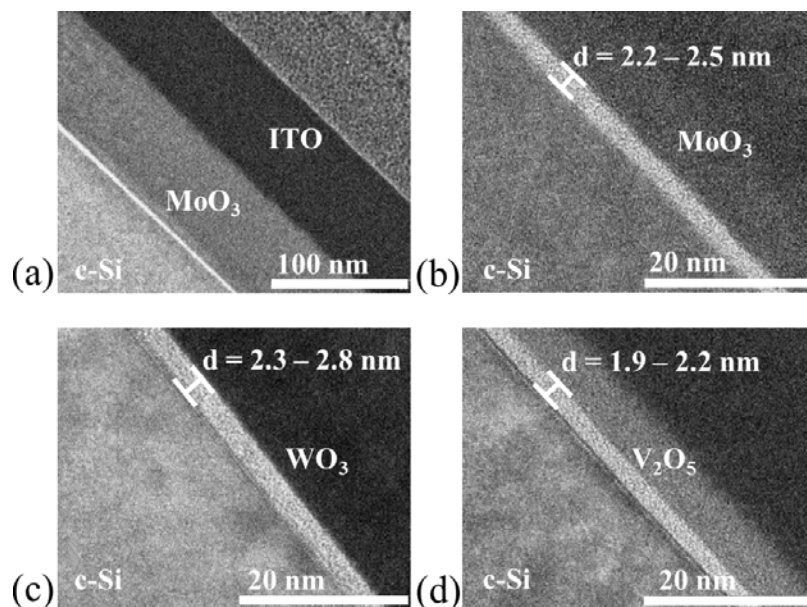


FIG. 1. HR-TEM images of the ITO/TMO/c-Si heterostructures showing an interlayer between c-Si and MoO₃ (a–b), WO₃ (c) and V₂O₅ (d).

Next, the interlayer composition was qualitatively determined by ToF-SIMS. Figure 2(a) shows the global depth profile of the ITO/MoO₃/n-Si sample, where the three distinct materials are clearly differentiated by their respective ions InO₂⁻/SnO₂⁻, MoO₃⁻ and Si⁻. Extraneous elements include carbon and fluoride (C⁻, F⁻), found mainly at the material interfaces and related to organic contamination during air exposure and to the HF dips respectively. By increasing the resolution of the time scale [Fig. 2(b)], a signal with an atomic mass unit (amu) of 59.97 and a peak intensity of 3x10³ counts is observed in the MoO₃/n-Si interface, corresponding to SiO₂⁻ ions. Since the HF-cleaned silicon surface is free of any native oxide (as will be confirmed later by XPS), the presence of SiO₂⁻ ions suggests that a thin silicon oxide IL has grown during thermal evaporation of MoO₃, proving that chemical passivation of dangling bonds by oxygen actually occurs. Within the same time window, a reduced Mo⁺² oxide (MoO⁻) signal is detected, indicating it is a by-product of the incomplete oxidation of the Si surface by MoO₃ (Si + MoO₃ → SiO₂ + MoO). Since remnants of the HF etch are also observed, it can be confirmed that such interlayer is restricted to the TMO/n-Si interface. A compound SiO₂-MoO₃ signal was also detected but at quite low intensities (<10² counts), more than two orders of magnitude lower than the SiO₂ signal and following the same trend. Therefore, the probability of a having a silicate-like Mo_xSi_yO₃ composition in the IL is low although it cannot be fully discarded. Finally, a SiOH⁻ ion signal is also detected but 10 times smaller than SiO₂, suggesting it is also related to adventitious contamination.

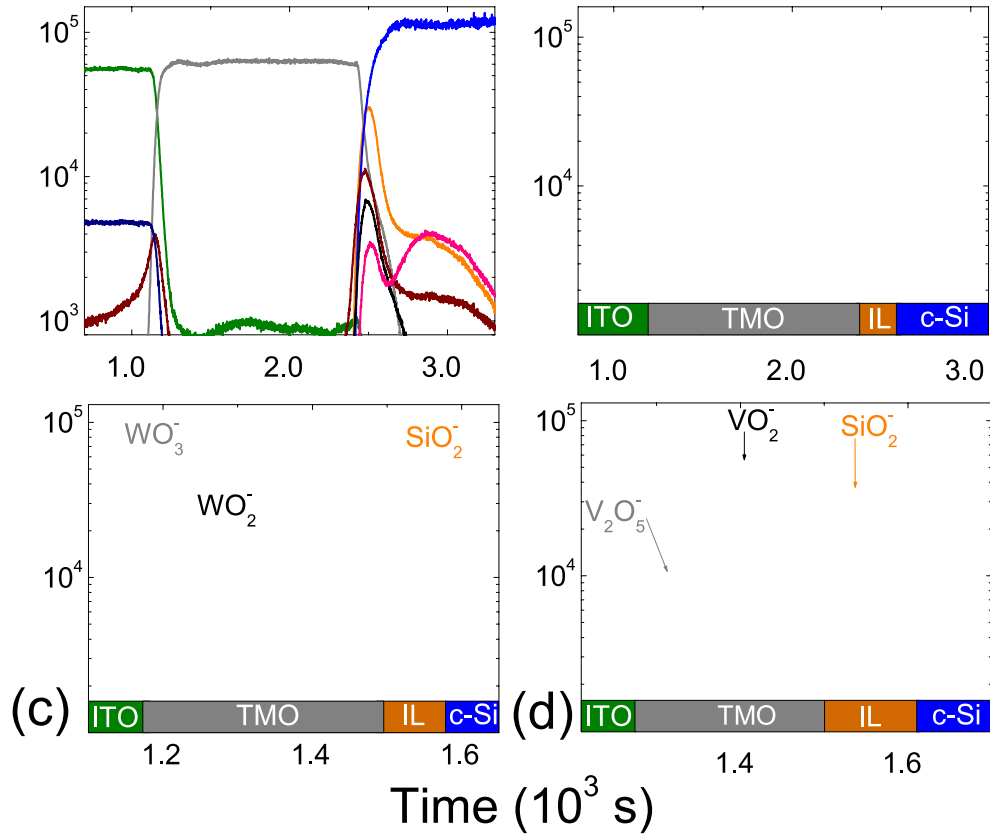


FIG. 2. ToF-SIMS depth profile for the ITO/TMO/c-Si heterostructures showing a SiO_2^- signal at the interface between c-Si and MoO_3 (a–b), WO_3 (c) and V_2O_5 (d). For each TMO, related reduced species (MoO^- , WO_2^- , VO_2^-) are also detected near the interface. Also shown in (a) are the F^- , C^- and SiOH^- signals, attributed to surface contamination of the HF-treated c-Si substrate after air-exposure. (Color Online).

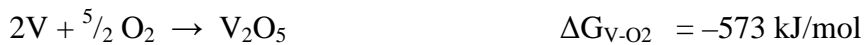
Similarly, the silicon oxide peak is also detected in the $\text{WO}_3/\text{n-Si}$ and $\text{V}_2\text{O}_5/\text{n-Si}$ interfaces in Fig. 2(c-d), at similar intensities and associated with reduced TMO species located adjacent to the interface (WO_2 , V_2O_4). In fact, since the vanadium oxide layer is the thinnest, the extent of the reduction reaction is greater and VO_2^- species surpass V_2O_5 in intensity. Moreover, since the SiO_2^- peak intensities are about the same order of magnitude as their respective TMO signals ($3.0\text{--}6.0 \times 10^4$ counts), it means that the SiO_2 layer is conformal and uniform across the analyzed area, in agreement with the TEM images.

Finally, SiOH⁻ and silicate signals were also detected in the WO₃ and V₂O₅ samples but at the low intensities mentioned before, leading to the same conclusions.

With respect to the abundant presence of the F⁻ ions (1.0 x10⁴ counts) and its effect on the interface, a separate V₂O₅ sample was washed with abundant deionized water after the HF dip, decreasing the F⁻ intensity by a factor of 2 (not shown) while the SiO₂⁻ remained unchanged. Since Si-F terminations are not kinetically viable during HF etching, it is probable these F⁻ are only physisorbed [22]. However, preliminary tests on finished solar cell devices prepared with and without the deionized H₂O quench indicate that F⁻ is not detrimental to the passivation or transport properties of these structures.

B. Thermodynamic feasibility

The formation of a SiO₂ layer on a H-terminated Si surface can only be explained by chemical reaction with TMOs during deposition. The growth of SiO_x interlayers on c-Si has been previously reported for dielectric oxides Al₂O₃, HfO₂, Ta₂O₅, LaAlO₃ and SrTiO₃, with interlayers varying from 0.7 to 9 nm in thickness and diverse stoichiometries (SiO_x, Hf_{1-x}Si_xO₂) [24-26]. This is partially explained by the high reactivity of H-terminated Si [22], but also from purely thermodynamic considerations. Parting from the oxidation of metallic Si and V by molecular O₂, the feasibility of these reactions is given by their negative Gibbs formation energies (ΔG):



For the case where Si is oxidized by V₂O₅, the vanadium oxidation reaction should be subtracted from the silicon one, leading to:

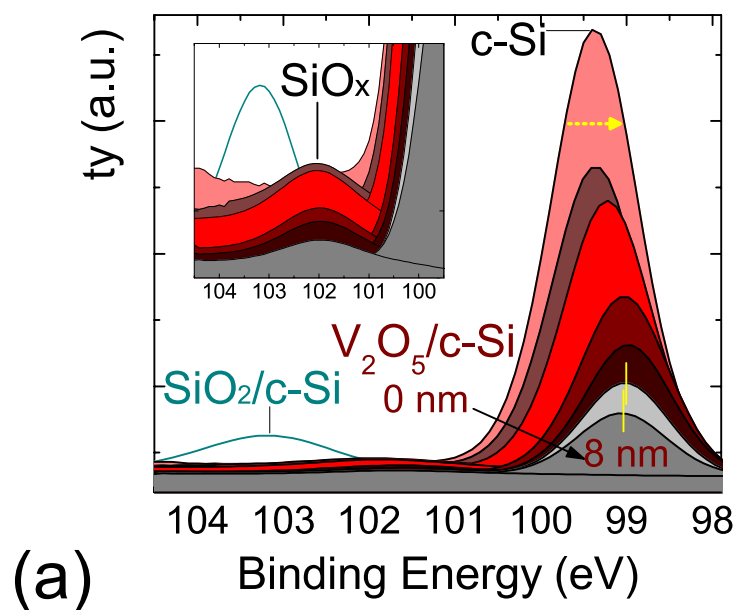


Hence, the oxidation of Si by a deposited metal oxide is thermodynamically feasible (spontaneous) if its Gibbs formation energy is less negative than the Gibbs formation energy of SiO₂. The same condition is met for the rest of the TMOs under study, with $\Delta G_{\text{Si-WO}_3} = -348$ and $\Delta G_{\text{Si-MoO}_3} = -406$ kJ/mol, using the standard Gibbs formation energies from [27] at 300 K. An exception to this rule is observed for HfO₂ and Al₂O₃ (ΔG positive), indicating that the spontaneity of these reactions is also affected by the initial equilibrium conditions, kinetic rates and the reactivity of the interface. Note also that any oxidation reaction essentially takes place by electron transfer, meaning that an electron loss in metallic Si yields an oxidized species ($\text{Si}^0 - 2\bar{e} \rightarrow \text{Si}^{+2}$). This interpretation will be important when explaining Fermi level alignment between TMOs and n-Si (see Section E). Regarding the relative ΔG s magnitude ($\text{V}_2\text{O}_5 > \text{WO}_3 > \text{MoO}_3$), no correlation was found with IL thickness (Fig. 1) or the work function of air-exposed TMOs ($\phi_{\text{V}_2\text{O}_5} \sim 5.3$, $\phi_{\text{WO}_3} \sim 5.0$, $\phi_{\text{MoO}_3} \sim 5.4$ eV)[16].

C. XPS of TMO/interlayer/c-Si region

As suggested before, the degree of completion of TMO-Si reactions could be less than unity, so that Si could partially oxidize ($\text{Si} + \text{V}_2\text{O}_5 \rightarrow \text{SiO}_x + \text{V}_2\text{O}_{5-x}$) and a mixture of sub-stoichiometric species would coexist in the interlayer. Therefore, a detailed chemical composition of the TMO/IL/n-Si interfaces was obtained from XPS analyses of V₂O₅ films deposited on n-Si at incremental thicknesses (1 – 8 nm), as a way to monitor film growth and the chemical/electronic changes taking place in the interface. Figure 3(a) shows the Si2p core level spectra as a function of binding energy, deconvoluted for the Si2p^{3/2} doublet. For the reference case where no V₂O₅ is present, only one peak centered at 99.5 eV

binding energy can be observed, characteristic of the metallic Si^0 substrate [28]. As soon as 1 nm V_2O_5 is deposited, an additional chemical state is detected at a higher binding energy of 102.0 ± 0.1 eV [Fig. 3(a) inset] and is discernible for thicker V_2O_5 films. This peak is attributed to sub-stoichiometric $\text{SiO}_{x-1.5}$ species (Si^{+3}) [28,29] and is identified as one of the components of the interlayer revealed by HR-TEM and ToF-SIMIS. As a reference, a stoichiometric SiO_2 oxide grown by wet etching of c-Si with HNO_3 is shown with its Si^{+4} peak at 103.2 eV. The sub-stoichiometry of the SiO_x interlayer is quite relevant because it explains why its thickness (as measured by HR-TEM) does not represent a barrier for the transversal conduction of carriers. Instead, the conduction mechanism is via defects (oxygen vacancies) and is not limited by the maximum tunneling thickness of SiO_2 passivation layers. The atomic Si^{3+} concentration in the IL was 0.9 – 2.0% and fairly constant for every sample, suggesting SiO_x formation takes place during the first deposition stages until it saturates, remaining constant afterwards.



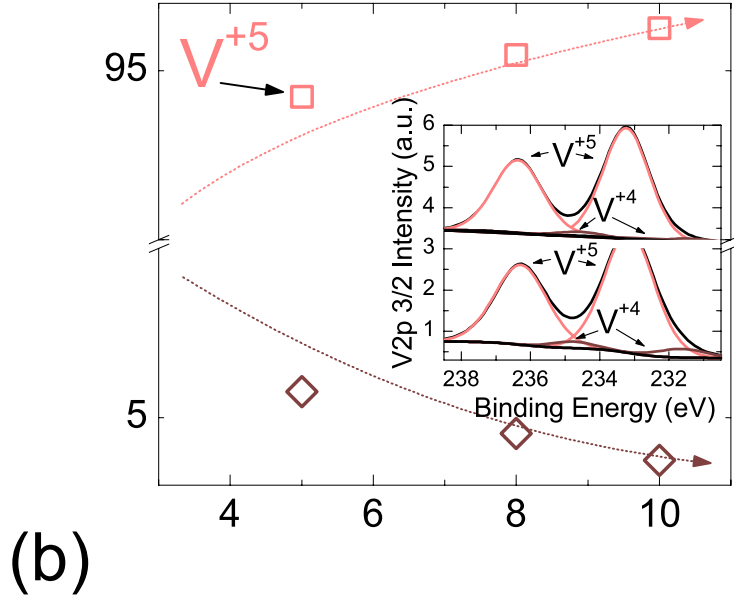


FIG. 3. XPS analysis of the V_2O_5 /interlayer/c-Si region at increasing V_2O_5 thicknesses (1 – 8 nm). (a) $Si2p^{3/2}$ doublet spectra showing the characteristic features of the silicon substrate (Si^0 , 99.5 eV), an stoichiometric SiO_2 /c-Si reference (Si^{+4} , 103.2 eV) and a sub-stoichiometric $SiO_{x-1.5}$ (Si^{+3} , ~102.0 eV) attributed to the TMO/c-Si interlayer. Inset shows a detailed view of these features. The peak maxima, marked with a vertical line, shifts to lower energies as thickness increases. The $Si2p^{1/2}$ doublet, shifted 0.63 eV to higher binding energies and smaller by a ratio 1:2, is omitted for clarity. (b) Evolution of the vanadium cation species V^{+4} and V^{+5} close to the c-Si interface. Inset shows the $V2p^{3/2}$ doublet spectra showing the characteristic features of V^{+4} (516.5 eV) and V^{+5} (518.1 eV) states. The $V2p^{1/2}$ doublet is shifted 7.6 eV to higher binding energies. (Color Online).

Since the $Si \rightarrow SiO_x$ partial oxidation reaction implies a complementary $V_2O_5 \rightarrow V_2O_{5-x}$ partial reduction, the XPS analysis of the vanadium $V2p$ core level spectra should show the vanadium oxidation states involved. This is depicted in Fig. 3(b) inset where the

$V2p^{3/2}$ doublet is deconvoluted into two peaks centered at 518.1 and 516.5 ± 0.1 eV binding energies, corresponding to the V^{+5} and V^{+4} cation species [30]. The area of each cation provides the relative V^{+4} and V^{+5} atomic concentrations as a function of V_2O_5 film thickness [Fig. 3(b)]. The observed tendency is the following: V^{+4} species, associated to oxygen-deficient films ($V_2O_{x<5}$) and generated by reduction ($V^{+5} \rightarrow V^{+4}$), are primarily found in the vicinity of the SiO_x IL and tend to decrease as the V_2O_5 film thickens; inversely, V^{+5} species are lowest (but still predominant) close to the IL and tend to increase until reaching a relatively fixed concentration in the bulk (away from the interface and its electronic influence). Similar trends have been observed previously for MoO_3 /metal [31] and MoO_3 /Si [32] systems, where reduced cation states (Mo^{+5}) exist near the oxidized interface. The presence of V^{+4} cations was already expected from the ToF-SIMS depth profiles, where VO_2^- ions were detected adjacent to the SiO_x interlayer. Therefore, although no detailed XPS analyses were performed with incremental WO_3 and MoO_3 thicknesses, the similarity between the ToF-SIMS/HR-TEM results indicate that analogue interlayers with $SiO_{x \leq 2}$ composition and reduced cation transitions ($Mo^{+6} \rightarrow Mo^{+5}$, $W^{+6} \rightarrow W^{+5}$) can be expected as well, as has been recently confirmed by [33,34].

D. Film stoichiometry and growth mode

Compared to the qualitative composition obtained from ToF-SIMS, XPS allows the quantification of film stoichiometries as oxygen to metal ratios (O/M). However, the specific distribution of oxygen species (SiO_x , V_2O_x) from the total oxygen content is not straightforward. To clarify this issue, we focus on the oxygen O1s XPS spectra shown in Figure 4 for the V_2O_5 /IL/n-Si region. From the deconvolution of the spectra, two components representing different chemical states are recognized. The first (larger)

component is centered at 530.9 ± 0.3 eV, attributed to O in vanadium oxide [30]. The second (smaller) component is centered at a higher binding energy (532.3 ± 0.3 eV) and is detected for every V_2O_5 thickness (1 – 8 nm), comprising ~25% of the total O1s area. This secondary-O, which has been previously described as a *defective oxide* inherent to many other TMOs [30], has its source on surface hydroxides (V–OH) and adventitious carbon (C–O) physisorbed during exposure to humid air [31,35] when transferred out of the glovebox into the XPS chamber. Additionally, the sub-stoichiometric SiO_x of the underlying interlayer also contributes to this secondary-O, as it lies very close to the SiO_2 reference (centered at 532.8 eV).

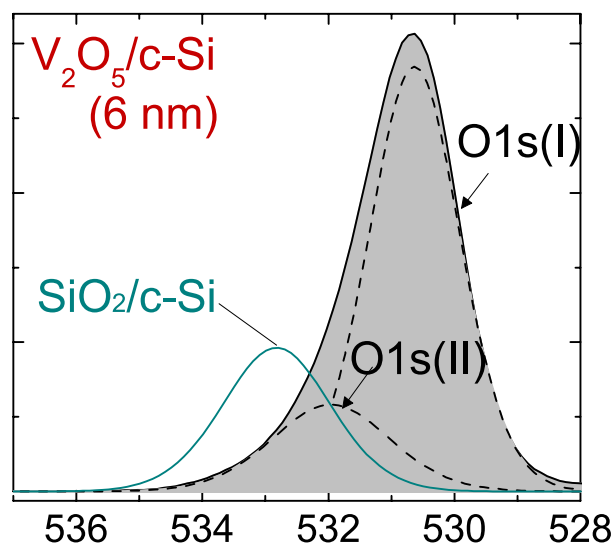


FIG. 4. O1s spectra of the V_2O_5 (6 nm)/interlayer/c-Si region. The larger feature (labeled I, 530.6 eV) belongs to oxygen in vanadium oxide, while the smaller one (labeled II, 532.0 eV) is attributed to oxygen in the $SiO_{x-1.5}$ interlayer and to a *defective-oxide* associated to adventitious oxygen physisorbed during air exposure. Also shown is a reference spectra for $SiO_2/c-Si$ (532.8 eV). (Color Online).

Under these assumptions, the approximate stoichiometry of the V_2O_5 films can be calculated from the oxygen to metal (O/V) ratios, where O is the total oxygen minus the surface hydroxides, adventitious carbon and interlayer contributions. Figure 5 (left axis) shows the variation of O/V ratios with film thickness, observing over-stoichiometric ratios $O/V > 2.5$ for the thinnest films ($V_2O_{x-6.3}$). Considering that other oxygen sources have already been neglected, an excess of oxygen near the interface seems correct but in conflict with the presence of V^{+4} cations attributed to oxygen deficient films [Fig. 3(b)]. Similar ratios have been reported for Al_2O_3/n -Si interfaces where excess oxygen is explained as the source of a negative fixed charge in Al_2O_3 [25,36]. Finally, it is worth noting that oxygen-deficient stoichiometries have been reported as inherent to the bulk of these TMOs (>8 nm), with oxygen vacancies, or defect states lying deep within the bandgap, being responsible for their conductivity [31].

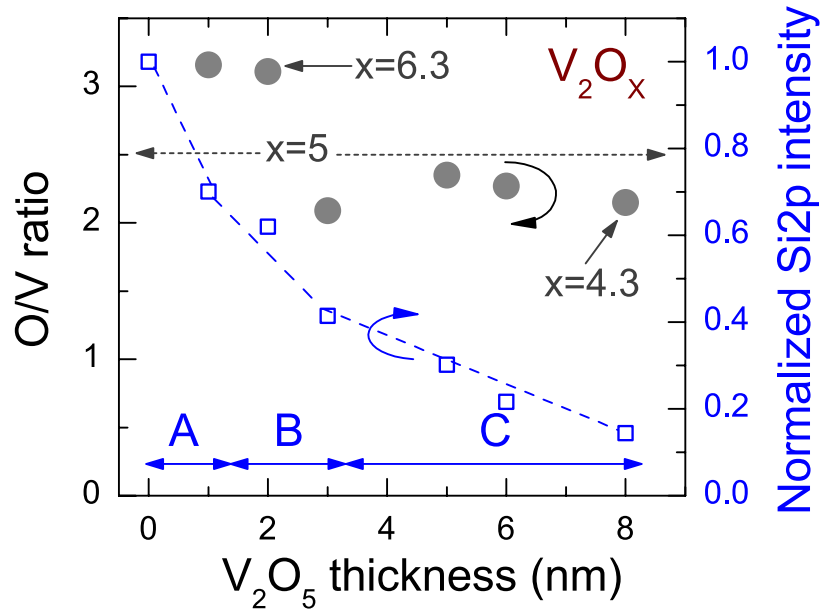


FIG. 5. (Left axis) Evolution of O/V ratios as a function of V_2O_5 film thickness, indicating a high stoichiometry ($V_2O_{x>5}$) close to the interface. (Right axis) Normalized area of the substrate $Si2p^{3/2}$ peak for each V_2O_5 thickness, identifying three growth regimes: (A) SiO_x

interlayer formation, (B) layered growth of V_2O_5 and (C) nucleation of V_2O_5 islands. (Color Online).

A qualitative measure of silicon substrate coverage by V_2O_5 can be obtained from plotting the normalized Si2p peak areas as a function film thickness, as shown in Figure 5 (right axis). At a first glance, three growth regimes are observed. During the first nanometers, a rapid linear increase of oxide coverage is observed until a critical thickness of ~ 3 nm is reached, followed by a less steep linear growth. Given that the IL thicknesses in Fig. 1 are quite uniform, we propose that the first regime (labeled A) is limited by the reaction rate between Si and the TMO, where SiO_x is formed until it saturates at a fixed thickness (and atomic concentration). Next (regime B), V_2O_5 continues to deposit on a layer-by-layer basis for several mono-layers until growth occurs by island nucleation (regime C), in what is called a Stranski–Krastanov growth mode [26,37]. Diffusion of species is also possible [26] but improbable at the low substrate temperatures ($\sim 25^\circ\text{C}$) at which thermal evaporation takes place.

E. Band bending in c-Si and interface dipole

Figure 6 depicts the proposed energy band diagram for TMO/n-Si before contact (a) and after the formation of the SiO_x IL has taken place (b). In the previous sections, passivation of the surface dangling bonds by SiO_x was proposed as the origin of chemical passivation. Additionally, the TMOs under study tend to induce an inversion (hole-rich) layer on n-Si as a result of their particularly large work functions ($\phi_{\text{TMO}} > 5.5$ eV), forming the contact needed for photogenerated holes extraction [16,38,39]. This transition from n-doping in the Si bulk to p-doping near the surface manifests as a band bending whose

magnitude is the built-in potential (V_{bi}) of the heterojunction. Thus, passivation through field-effect is established by driving photogenerated electrons away from the interface.

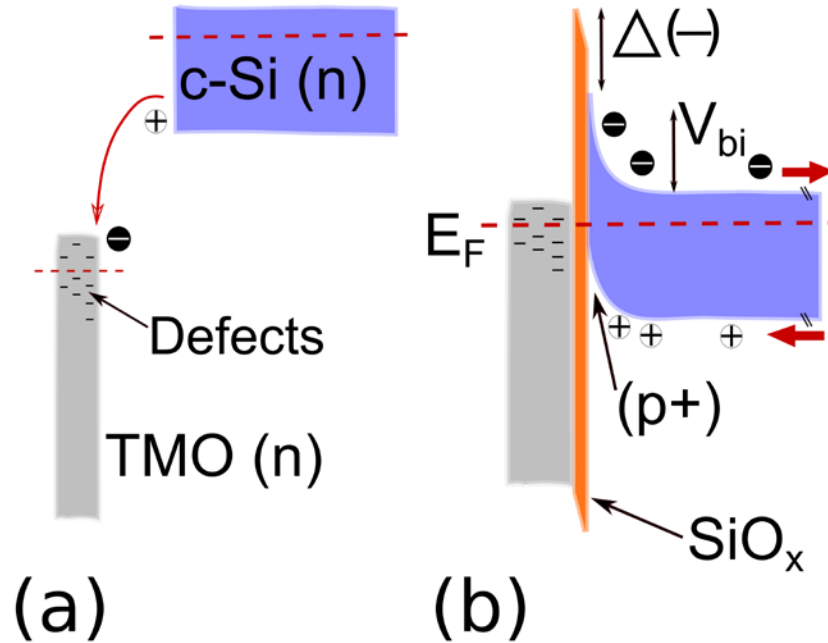


FIG. 6. Proposed band diagram for the TMO/n-Si heterojunction. (a) Before contact, depicting the transfer of electrons from n-type c-Si (oxidation) towards the higher work function TMO (reduction). (b) After contact, following the formation of a passivating SiO_x interlayer. Electron transfer also induces a p^+ inversion layer on the c-Si surface and possibly a negative dipole $\Delta(-)$ until Fermi levels reach equilibrium. (Color Online).

Evidence of silicon band bending resulting from TMO deposition can be found in the $\text{Si}2p$ core level spectra in Figure 2(a). A shift of the $\text{Si}2p^{3/2}$ peak maximum to lower binding energies is observed as the V_2O_5 film thickness increases. This shift implies the ‘distance’ between the Fermi level (E_{Fermi}) and the valence band (E_{VB}) is decreasing, or in other words, up-bending occurs as V_2O_5 is deposited [40]. A maximum shift of 0.36 eV is obtained for the 5 nm film, although these values are only approximate given the sources of

error in these measurements, mainly 1) the mismatch between the photoelectron sampling depth (<10 nm) and the inversion layer full width (> 100 nm), and 2) the generation of a surface photovoltage from x-ray illumination [40].

The formation of an inversion layer in n-Si by high work function TMOs can also be explained as a consequence of band alignment across the interface [41,42]. Before both materials are contacted, a large electrochemical potential difference exists between them, as defined by the difference in their work functions $\Delta\phi = \phi_{\text{TMO}} - \phi_{\text{nSi}} = 5.4 - 4.2 \text{ eV}$. Such energy mismatch is the driving force that initiates E_{Fermi} alignment in a series of steps. First, electron transfer occurs from the valence band of n-Si ($E_{\text{VB}} \sim 5.2 \text{ eV}$) into the defect states lying deep within the TMO bandgap ($E_{\text{states}} > 6 \text{ eV}$) (see Fig. 6). As explained in Section B, an electron loss in Si results in its oxidation to SiO_x , while the TMO suffers a reduction to lower valence states due to electron gain. Then, as the Si surface is depleted of electrons, the valence band of silicon approaches E_{Fermi} and a p^+ layer is induced. This inversion-effect might be reinforced by negative charges fixed in the IL or in the first $\text{V}_2\text{O}_{x>5}$ layers, as evidenced from the high O/V ratios. Since band bending cannot occur indefinitely, we presume the n-Si surface reaches a strong inversion condition with built-in voltage values close to $V_{bi} \sim 0.7 \text{ eV}$. At this point, the large $\Delta\phi$ has only been distributed partially in the Si surface and the remaining energy difference needed for equilibrium is allocated by a negative dipole $\Delta(-)$, so that $\Delta\phi = V_{bi} - \Delta$. Similar mechanisms have been previously described for TMO/organic semiconductor systems where electron transfer into the TMO results in the formation of interfacial dipoles and p-doping of the organic Highest Occupied Molecular Orbital (HOMO) levels [23,43]. Regarding the origin of the dipole, it is most probably located in the $\text{SiO}_x/\text{c-Si}$ interface as surface states, although it could also lie in the

TMO in the form of uncoordinated (dangling) oxygen atoms, an inherent characteristic of TMOs with large stoichiometries (Mo_2O_6 , W_2O_6 and V_2O_5) and multi-layered bulk structures [44,45].

IV. CONCLUSIONS

The mechanisms behind the passivation of n-type c-Si surfaces by thin layers of transition metal oxides MoO_3 , WO_3 and V_2O_5 were investigated by probing the TMO/c-Si interface by HR-TEM, ToF-SIMS and XPS. Chemical passivation due to silicon bonding with oxygen was identified as a sub-stoichiometric $\text{SiO}_{x-1.5}$ interlayer with thickness averaging 1.9 – 2.8 nm. This silicon sub-oxide is formed during the partial oxidation of H-terminated silicon by the first TMO monolayers, leaving behind reduced and oxygen-deficient species (MoO , WO_2 and VO_2). The redox reaction was determined to be thermodynamically favorable ($-\Delta G_{\text{rxn}}$) and driven by the large difference in the electrochemical potentials (Fermi levels) of both materials. Therefore, Fermi level equilibration across the interface results not only in SiO_x formation but also on electron depletion (inversion) of the n-Si surface. Such inversion layer (band bending) thus acts as the hole-selective contact in this type of heterojunction solar cells.

Additionally, the detailed investigation of the $\text{V}_2\text{O}_5/\text{SiO}_x/\text{c-Si}$ region by XPS determined that 1) film stoichiometry (V_2O_x) varies with thickness, 2) different growth regimes (layer-by-layer, island nucleation) are present, and 3) band bending is observed as a shift of the Si2p core level spectra to lower binding energies, confirming that field-effect passivation (i.e. carrier-concentration gradients) is present as well.

Overall, the described behavior of the TMO/c-Si interface allows for further improvements in its passivating properties, including surface functionalization, H_2 diffusion

to enhance chemical passivation or fine-tuning of TMO stoichiometry by other deposition methods (sputtering, atomic layer deposition). Once a genuine passivating/selective contact is developed, while maintaining (or improving) the conversion efficiency of the device, its eventual implementation could lower processing costs and enhance the competitiveness of c-Si photovoltaics.

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FIGURE LIST

FIG. 1. HR-TEM images of the ITO/TMO/c-Si heterostructures showing an interlayer between c-Si and MoO₃ (a–b), WO₃ (c) and V₂O₅ (d).

FIG. 2. ToF-SIMS depth profile for the ITO/TMO/c-Si heterostructures showing a SiO₂⁻ signal at the interface between c-Si and MoO₃ (a–b), WO₃ (c) and V₂O₅ (d). For each TMO, related reduced species (MoO⁻, WO₂⁻, VO₂⁻) are also detected near the interface. Also shown in (a) are the F⁻, C⁻ and SiOH⁻ signals, attributed to surface contamination of the HF-treated c-Si substrate after air-exposure. (Color Online).

FIG. 3. XPS analysis of the V₂O₅/interlayer/c-Si region at increasing V₂O₅ thicknesses (1 – 8 nm). (a) Si2p^{3/2} doublet spectra showing the characteristic features of the silicon substrate (Si⁰, 99.5 eV), an stoichiometric SiO₂/c-Si reference (Si⁺⁴, 103.2 eV) and a sub-stoichiometric SiO_{x~1.5} (Si⁺³, ~102.0 eV) attributed to the TMO/c-Si interlayer. Inset shows a detailed view of these features. The peak maxima, marked with a vertical line, shifts to

lower energies as thickness increases. The $\text{Si}2p^{1/2}$ doublet, shifted 0.63 eV to higher binding energies and smaller by a ratio 1:2, is omitted for clarity. (b) Evolution of the vanadium cation species V^{+4} and V^{+5} close to the c-Si interface. Inset shows the $\text{V}2p^{3/2}$ doublet spectra showing the characteristic features of V^{+4} (516.5 eV) and V^{+5} (518.1 eV) states. The $\text{V}2p^{1/2}$ doublet is shifted 7.6 eV to higher binding energies. (Color Online).

FIG. 4. $\text{O}1s$ spectra of the V_2O_5 (6 nm)/interlayer/c-Si region. The larger feature (labeled I, 530.6 eV) belongs to oxygen in vanadium oxide, while the smaller one (labeled II, 532.0 eV) is attributed to oxygen in the $\text{SiO}_{x-1.5}$ interlayer and to a *defective-oxide* associated to adventitious oxygen physisorbed during air exposure. Also shown is a reference spectra for $\text{SiO}_2/\text{c-Si}$ (532.8 eV). (Color Online).

FIG. 5. (Left axis) Evolution of O/V ratios as a function of V_2O_5 film thickness, indicating a high stoichiometry ($\text{V}_2\text{O}_{x>5}$) close to the interface. (Right axis) Normalized area of the substrate $\text{Si}2p^{3/2}$ peak for each V_2O_5 thickness, identifying three growth regimes: (A) SiO_x interlayer formation, (B) layered growth of V_2O_5 and (C) nucleation of V_2O_5 islands. (Color Online).

FIG. 6. Proposed band diagram for the TMO/n-Si heterojunction. (a) Before contact, depicting the transfer of electrons from n-type c-Si (oxidation) towards the higher work function TMO (reduction). (b) After contact, following the formation of a passivating SiO_x interlayer. Electron transfer also induces a p^+ inversion layer on the c-Si surface and possibly a negative dipole $\Delta(-)$ until Fermi levels reach equilibrium. (Color Online).