

19.2% Efficient InP Heterojunction Solar Cell with Electron-Selective TiO₂ Contact

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Supporting Information

ABSTRACT: We demonstrate an InP heterojunction solar cell employing an ultrathin layer (~10 nm) of amorphous TiO_2 deposited at 120 °C by atomic layer deposition as the transparent electron-selective contact. The TiO_2 film selectively extracts minority electrons from the conduction band of p-type InP while blocking the majority holes due to the large valence band offset, enabling a high maximum open-circuit voltage of 785 mV. A hydrogen plasma treatment of the InP surface drastically improves the long-wavelength response of the device, resulting in a high short-circuit current density of 30.5 mA/cm² and a high power conversion efficiency of 19.2%.



KEYWORDS: InP photovoltaics, titanium dioxide, heterojunctions, selective contact

S elective contacts that transmit photogenerated minority carriers and block majority carriers are key to highperformance solar cells.¹ Traditionally inorganic semiconductors use doping to achieve carrier selectivity at the contacts. For organic solar cells, in contrast, transition metal oxides have proven to be efficient selective contacts.^{2–8} In particular TiO₂ is applied widely as an electron contact in dye-sensitized solar cells,^{9,10} polymer solar cells,^{11,12} and high-performance organometallic perovskite solar cells,^{13–16} while substoichiometric MoO_x (x < 3) has emerged as an excellent hole contact.⁴ Recently we demonstrated that MoO_x can also serve as an efficient hole contact for n-type silicon solar cells, enabling a high open-circuit voltage of 711 mV and an efficiency of 18.8%.^{17,18} Analogously, TiO₂ has been used as an electronselective contact to p-type silicon.¹⁹

In this work, we show that transition metal oxides can also be applied as selective contacts for III–V solar cells. We focus in particular on InP, whose band gap of 1.35 eV lies close to the optimum for solar energy conversion.²⁰ The most efficient InP solar cells use a homojunction configuration fabricated by metal–organic vapor deposition (MOCVD), reaching efficiencies of up to 22.1%.²¹ Alternatively, a shallow buried homojunction can be formed in InP by sputtering indium tin oxide (ITO) on the front surface. This approach yields efficiencies of up to 18.9%.²² In another approach, a CdS window layer is used with InP to form a heterojunction solar cell with an efficiency of 17.4%.^{23,24} Here, by employing a TiO₂ electron contact deposited by atomic layer deposition (ALD) at

a low temperature of 120 °C, we demonstrate a TiO_2/InP heterojunction solar cell with a high open-circuit voltage of 785 mV and an excellent efficiency of 19.2%.

The process flow for fabricating TiO₂/InP solar cells is shown in Figure 1a. Zn-doped p-type (100)-oriented InP wafers with a thickness of 350 μ m and a carrier density of 2 \times 10^{17} cm⁻³ are etched in HCl (6%) for 30 s to remove the native oxide. Twenty nanometers of Zn followed by 100 nm of Au are then sputtered onto the back side of the InP wafers, followed by annealing in forming gas (5% H₂, 95% N₂) at 420 °C for 40 min to form an ohmic back contact. TiO₂ is subsequently deposited onto the front side of the InP wafers by ALD using titanium isopropoxide $(Ti[OCH(CH_3)_2]_4)$ and H_2O as precursors. Finally, a transparent conductive oxide (ITO = In_2O_3 :SnO = 90%:10%) electrode is sputtered on top of the TiO₂ layer at room temperature in Ar ambient at a pressure of 6-7 mTorr. For all devices, the thickness of the ITO layer was adjusted according to the thickness of the TiO₂ layer to maintain a total oxide thickness of \sim 70 nm, such that the reflectance minimum falls in the spectral range where the solar spectrum is strongest (around a wavelength of 600 nm) in order to maximize light in-coupling into the InP absorber.¹⁷ The optimum TiO₂ thickness (10 nm) maximizes the opencircuit voltage (see discussion below) and minimizes resistive losses. Photolithography followed by electron beam evapo-

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Figure 1. (a) Process schematics of the TiO_2/InP heterojunction solar cell fabrication, depicting Au:Zn back-contact formation, ALD of TiO_2 electron-selective contact, ITO sputtering, and Ag finger grid evaporation. (b) Photograph of a fully fabricated cell. The area of the cell is defined by the outer Ag frame. A black plastic tape is applied on the edges of the chip (around the outer Ag frame) to ensure no collection is obtained from outside the marked cell area. (c) False-colored, cross-sectional SEM image of a fabricated cell.

ration and lift-off were then used to define a Ag finger grid with finger width and pitch of 11 and 490 μ m, respectively, to reduce series resistance. The final patterned cells have an aperture area of 5 × 5 mm² defined by the outer Ag frame (Figure 1b). Figure 1c shows a false-colored cross section scanning electron microscopy (SEM) image of the final TiO₂/*p*-InP heterojunction solar cell.

The current density–voltage (I-V) characteristics of a TiO₂/ InP heterojunction solar cell measured under simulated 1-sun illumination (1000 W/m², global air mass 1.5 spectrum, 25 °C, Solar Light 16S 300W) are shown in Figure 2a. A sharp probe tip was placed on the outer Ag frame to contact the front side of the cell. Comparative tests with a four-point probe configuration gave identical results to the two-point probe configuration used here, as the Ag frame provides a contact point with negligible contact resistance to the probe tip(s). The simulated solar radiance received by the cells was calibrated to a NIST traceable silicon photodiode with similar aperture area as the InP cells and known external quantum efficiency and further cross-checked via an ISO-classified pyranometer (Solar Light PMA 2144). The error in our cell efficiency measurement is estimated to be at least 1%. A more detailed error calculation including spectral mismatch beyond class "A" would require a full spectral analysis of our solar simulator. Also shown in Figure 2a is a reference device with no TiO₂ blocking layer, reaching an open-circuit voltage $(V_{\rm oc})$ of 545 mV and an efficiency of 9.9%. Insertion of the TiO₂ layer leads to a dramatic increase of the $V_{\rm oc}$ to 785 mV. The short-circuit current density (J_{sc}) also slightly improves from 24.6 mA/cm² to 25.8 mA/cm², resulting in an improved conversion efficiency of 15.4%. The higher J_{sc} is not due to a change in cell reflectance, but is attributed to improved carrier collection due to reduced recombination at the InP front interface (see Figure S1 for details). Cell performance parameters are summarized in Table 1.

To optimize device performance, we investigate the dependence of V_{oc} on the TiO₂ thickness and deposition temperature (see Figure 2b and c, respectively). For each condition, at least three devices were fabricated and measured, and the average V_{oc}



Figure 2. (a) Representative J-V characteristics for InP solar cells, with and without TiO₂ electron-selective contact under 1-sun illumination. (b) Dependence of V_{oc} on TiO₂ thickness (at a fixed deposition temperature of 250 °C) and TiO₂ deposition temperature (at a fixed thickness of 10 nm).

Table 1. Summary of InP Solar Cell Performance Parameters

device	$V_{\rm oc}$ [mV]	$J_{\rm sc} \begin{bmatrix} mA/\\ cm^2 \end{bmatrix}$	FF [%]	eff [%]
ITO/InP	545	24.6	73.6	9.9
ITO/TiO ₂ /InP	785	25.8	72.8	15.4
$ITO/TiO_2/InP$ with H_2 plasma	785	30.5	80.1	19.2

values are reported here. When deposited at a fixed temperature of 250 °C, the $V_{\rm oc}$ increases rapidly with increasing TiO₂ thickness and reaches a saturation value of 740 mV for a thickness of 10 nm, beyond which the $V_{\rm oc}$ does not change significantly (Figure 2b). Lowering the deposition temperature to 120 °C at a fixed thickness of 10 nm increases the $V_{\rm oc}$ further to 780 mV (Figure 2c).

In order to better understand the V_{oc} improvement due to the insertion of the TiO₂ layer, we characterized the structural and electronic properties of ALD TiO₂. Raman spectroscopy (Figure S2a) indicates that TiO₂ deposited at 120 °C, for which we observed the highest V_{oc} , is in an amorphous phase in agreement with literature.²⁵ Atomic force microscopy (Figure S2b and c) reveals a smooth, relatively featureless, film morphology consistent with an amorphous film structure (root-mean-square (rms) roughness 0.3 nm). In contrast, TiO₂ films deposited at 250 °C exhibit a weak Raman signal at 140 cm⁻¹ corresponding to a nanocrystalline anastase phase and a rougher surface morphology consistent with the formation of small crystallites (rms roughness 1.2 nm).

Figure 3a presents the X-ray photoelectron spectrum (XPS) of the Ti 2p core level of the ALD TiO₂ layer deposited at 120 °C measured with monochromated Al K α photons with an



Figure 3. X-ray photoelectron spectra for (a) Ti 2p core level (vertical arrows indicate the energy positions of the Ti⁴⁺ and Ti³⁺ valence states), (b) the valence band region with the Fermi level (E_F) at zero binding energy, and (c) the secondary electron cutoff of 10 nm TiO₂ on a p-InP wafer, deposited at 120 °C. (d) Absorption coefficient as a function of photon energy for an ALD TiO₂ film on glass. (e) Schematic band diagram of the TiO₂/InP heterojunction solar cell simulated by PC1D.³⁶ The horizontal axis represents the distance away from the top surface, and the vertical axis shows the energy with respect to the vacuum level. The extended space charge region in InP after the H₂ plasma treatment is indicated by light green lines. A work function of 4.3 eV and electron affinity of 4.4 eV were assumed for ITO.³⁷

energy of 1486.7 eV. The core level is spin split into the $2p_{3/2}$ and $2p_{1/2}$ doublet centered at 458.7 and 464.4 eV, respectively, corresponding to the Ti⁴⁺ valence state of the Ti atoms.²⁶ No appreciable signal from the Ti³⁺ components, typically located at 462.0 and 456.8 eV (see arrows in Figure 3a), is detected. Also a satisfactory fit obtained by adding only two Voigtian components for the Ti⁴⁺ doublet on top of a Shirley background indicates that our films are close to stoichiometric TiO₂ (see SI S5 for technical details). A fit including four Voigtian components for the Ti⁴⁺ and Ti³⁺ doublets results in the Ti³⁺ peaks vanishing during convergence, indicating no detectable evidence for oxygen vacancies with Ti³⁺ configuration.

Figure 3b shows the valence band spectrum of the TiO₂ film, which is dominated by O 2p states. Consistent with the conclusion from the Ti 2p core level, we do not observe evidence for an oxygen-vacancy-derived defect band in the band gap of TiO₂ (compare for example to MoO_{x} , where such a defect band is clearly observed¹⁷). A linear extrapolation (red line) of the valence band edge fixes the valence band maximum at 2.9 eV below the Fermi level located at zero binding energy (dashed black line). Figure 2c shows the secondary electron cutoff of the photoelectron spectrum (the spectra are corrected for an externally applied bias of -9 V on the sample into the detector), from which we extract the workfunction by subtracting the cutoff energy of 1482.2 eV determined again by linear extrapolation (red line) from the photoel energy of

1486.7 eV. Thus, the work function of TiO_2 is determined to be 4.5 eV.

To determine the optical band gap of our TiO_2 , we measured the absorption coefficient of a TiO_2 film deposited on a glass substrate (Figure 3d). Typical for an amorphous material, we observe a tail in the absorption spectrum that rises exponentially toward higher energy. The Tauc band gap determined to be the energy where the absorption coefficient starts to deviate from the exponential (red line in log scale) is 3.4 eV, which is slightly higher than the value of 3.2 eV reported for crystalline anastase TiO_2 .²⁷

From the position of the Fermi level with respect to the vacuum level given by the workfunction and the valence band maximum determined with respect to the Fermi level, we conclude that the valence band maximum of TiO_2 is located at 7.4 eV below the vacuum level. Together with the information on the optical band gap, the conduction band minimum consequently falls at 4.0 eV below the vacuum level.

A similar analysis for the InP surface (Figure S3) places the valence band of InP at 5.4 eV below the vacuum level. With a band gap of 1.35 eV, the conduction band maximum falls to 4.05 eV. These values are roughly consistent with values determined by the Mott–Schottky technique, which confirms that the conduction band of TiO₂ (4.3 eV) is energetically well matched to that of InP (4.2 eV) although slightly further away from vacuum.²⁸ More detailed investigations are necessary to compare and unify band positions extracted from different experimental techniques.²⁹

Table 2.	Comparison	of State-of-the-Art	InP Sola	r Cell 1	Performance	Parameters	(Measured	under AM	1.5, for	: Cell Areas	s ^c)
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device type	description	$V_{\rm oc} [{\rm mV}]$	$J_{\rm sc} \left[{\rm mA/cm^2} \right]$	FF [%]	eff [%]	ref
MOCVD	homojunction	878	29.5	85.4	22.1	21, 30
TiO ₂ /InP	heterojunction	785	30.5	80.1	19.2	this work
ITO/InP	buried homojunction	813	$28.0 (28.2^{a})$	82.9	18.9	22
CdS/InP	heterojunction	750	$32.3 (28.6^b)$	72.0	17.4	24

^aAfter applying recalibration due to replacement of the IEC 60904-3 Ed. One (1989) standard by the IEC 60904-3 Ed. Two (2008) standard.³¹ ^bExtracted from EQE using IEC 60904-3 Ed. Two (2008) standard. ^cThe total cell area of the current InP world record cell is specified as 4.02 cm² in ref 30. No further details are given. The ITO/InP cell in ref 22 is reported to have a mesa area of 0.108 cm², of which 5% is shaded by the Au finger grid. The TiO₂/InP cell described in this work has an area of 0.25 cm² masked by the outer Ag frame (Figure 1b), of which 2% is shaded by the Ag finger grid. Ref 25 specifies a diameter of 1.5 mm for the CdS/InP cells, but cell area/shape and grid shading are not specified.

The Fermi level of InP as determined by XPS is located 0.6 eV above the valence band maximum (see Figure S3a). It is important to note that due to the extreme surface sensitivity of XPS, this value represents the surface Fermi level and may be heavily affected by surface defects, surface oxidation, or charge transfer from surface adsorbates (Figure S3c). The position of the Fermi level in the InP bulk can be calculated from the hole concentration $(2 \times 10^{17} \text{ cm}^{-3})$ and is located 0.1 eV above the valence band maximum. Consequently the bare InP surface already exhibits some band bending at the surface. In fact, it is well known that intentional surface defect formation via ion bombardment of the InP surface can lead to type conversion resulting in buried homojunctions.³⁰ However, here the bare InP surface is still slightly p-type, as the position of the intrinsic Fermi level in InP lies 700 mV above the valence maximum (slightly above midgap due to the different effective masses between electrons and holes).

On the basis of these results, we draw the band diagram for the TiO₂/p-InP heterostructure shown in Figure 3e. At equilibrium, the Fermi level of TiO₂ aligns with the Fermi level of InP, inducing an inversion region in p-type InP close to the TiO₂/InP interface. Minority electrons generated by the absorbed photons in InP can easily diffuse toward the front contact and do not experience a barrier at the TiO₂ interface. Majority holes in contrast are prohibited from flowing into the front contact due to the combined effect of the band bending in the space charge region and the very large band offset (~2 eV) between the valence band of InP and the valence band of TiO₂. Therefore, TiO₂ acts as an efficient electron-selective contact for InP.

While the $V_{\rm oc}$ obtained with TiO₂ compares well with the best values reported for ITO/InP buried homojunction and CdS/InP heterojunction cells (Table 2), the J_{sc} turns out to be much lower. External quantum efficiency (EQE) measurements shown in Figure 4a reveal that the low current can be attributed to a poor long-wavelength response of the TiO₂/InP solar cell (black line). The reduction in EQE toward the infrared is only partially due to reflectance losses. As the TiO₂ and ITO thicknesses were optimized to minimize reflectance in the spectral range around 600 nm, the reflectance (shown as 1-R in Figure 4a) increases continuously toward longer wavelengths. This is expected for an antireflection coating that relies on destructive interference between incoming and reflected light to cancel reflection. However, the EQE drops much faster toward the band edge of InP than the 1-R curve. Thus, the poor infrared response cannot be explained solely by reflectance losses, but must also be attributed to collection losses.

We recently showed using electron-beam-induced current (EBIC) measurements that the effective minority carrier collection length in InP can be increased dramatically from



Figure 4. (a) EQE, 1-R, and (b) J-V characteristics measured under 1-sun for TiO₂/InP solar cells, with and without H₂ plasma treatment prior to ALD of TiO₂.

0.5 μ m to 2.6 μ m by reactive ion etching in a H₂:CH₄:Ar ambient, resulting in a much enhanced near-band-gap response in the EQE.³² In addition, it is well known that hydrogen effectively neutralizes shallow acceptors in InP, resulting in a decrease in effective hole concentration and an extension of the effective minority carrier collection depth (see Figure 3e).^{33,34} However, these buried homojunction-type cells suffered from a strongly suppressed response at short wavelengths due to the formation of a defective surface layer with strong nonradiative recombination limiting cell efficiencies to 14.4%.

In order to boost the long-wavelength response, while maintaining a strong short-wavelength response, we subjected the front side of the InP wafers to a H₂ plasma (pressure of ~85 mTorr, power density of 0.5 W/cm², and bias voltage of -450 V) before TiO₂ deposition. The H₂ plasma causes the loss of phosphorus and formation of indium droplets at the InP surface, as shown in Figure S4, which are detrimental for device performance. To eliminate these indium residues, the InP surface was wet etched in piranha solution (H₂SO₄/H₂O₂ =

10:1) for 30 s and rinsed in deionized water, which leaves a flat specular surface similar to InP before H_2 plasma treatment.

As shown in Figure 4a, the H₂ plasma treatment leads to a dramatic increase of the EQE (shown in red) over a wide spectral range, reaching from 500 nm all the way up to the band gap at 920 nm. As expected, the reflectance of the H₂ plasma treated cell remains identical to the cell without H₂ plasma treatment. The EQE now follows the 1-R curves closely, indicating much reduced collection losses. Remarkably the short-wavelength response remains almost unaffected and is maintained above 60% at 400 nm. Figure 4b compares the 1sun J-V curves for cells with and without H₂ plasma treatment. The V_{oc} remains unchanged, reaching 785 mV for the best cell. The enhanced EQE due to the H₂ plasma treatment accounts for a massive gain in J_{sc} of 4.7 mA/cm², resulting in an excellent photocurrent of 30.5 mA/cm². This current could be further improved by applying an additional MgF₂ antireflection coating. Combined with a fill factor of 80.1%, a high efficiency of 19.2% is reached.

As for the role of the H_2 plasma treatment versus TiO_2 , the former enhances band bending, which helps in collecting minority carriers generated by long-wavelength photons and consequently improves the red response. On the other hand, TiO₂ provides a large barrier for majority holes (associated with the large valence band offset with respect to InP) but allows minority electrons to pass through (due to the small conduction band offset with respect to InP). This reduces the dark current and therefore improves $V_{\rm oc}$. From experimental data, band bending induced by H_2 plasma treatment alone is not sufficient to reach high V_{oc} . Table 2 compares the performance parameters of our TiO2/InP heterojunction solar cell with the values reported in the literature for alternative device configurations. Remarkably, our heterojunction solar cell reaches the highest efficiency after the world record InP homojunction solar cell fabricated by expensive MOCVD. In particular, our cell fabrication process is compatible with InP thin films grown directly on cheap metal foils using our recently developed thin film vapor liquid solid growth technique, which circumvents the main cost drivers for III-V photovoltaics: the epitaxial wafer substrate, low precursor utilization yield, and long growth times associated with MOCVD. Our cell configuration therefore provides a promising route toward lowcost large-area III-V photovoltaics with high efficiency.

In conclusion, we have demonstrated a simple TiO_2/InP heterojunction solar cell achieving an efficiency of 19.2% using an electron-selective TiO_2 contact prepared by ALD. A H_2 plasma treatment substantially improves the spectral response of the cell. Importantly our process is fully compatible with InP thin films, rendering InP interesting not only for space applications but also for terrestrial deployment.

ASSOCIATED CONTENT

S Supporting Information

Additional materials and device characterization data. This material is available free of charge via the Internet at http:// pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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19.2% efficient InP heterojunction solar cell with electron selective TiO₂ contact

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Supporting information

Figure S1



Fig. S1: External quantum efficiency (EQE) and reflectance R (shown as 1-R) for ITO/InP and ITO/TiO₂/InP solar cells. The TiO₂ layer yields an improved short wavelength response in the EQE at almost identical values of R, indicative of reduced surface recombination. The thickness of the ITO front electrode was adapted to maintain a total oxide thickness of ~70 nm.

Figure S2



Fig. S2: (a) Raman spectra of bare InP wafer and \sim 30 nm thick TiO₂ deposited on InP by ALD showing that TiO₂ deposited at 120°C is amorphous, whereas TiO₂ deposited at 250°C exhibits a peak at 140 cm⁻¹ consistent with a nanocrystalline anastase phase. Atomic force microscopy (AFM) images of \sim 30 nm thick TiO₂ deposited on InP at (b) 120°C and (c) 250°C.



Fig. S3: X-ray photoelectron spectra (XPS) for (a) the valence band region with the Fermi level (E_F) at zero binding energy, (b) secondary electron cutoff, and (c) In 3d core level of a bare *p*-type InP wafer, with and without HCl etching right before the XPS measurement.

Figure S4



Fig. S4: Scanning electron microscopy (SEM) image of an InP wafer right after H_2 plasma treatment showing the resulting In droplets.

S5: Explanation to analysis of core level photoemission spectra:

Voigtian line profiles, named after Woldemar Voigt, are often used in spectroscopy to account for the combined effect of Lorentzian and Gaussian peak broadening mechanisms. In x-ray photoelectron emission the natural lineshape of a core level resonance is Lorentzian. The broadening is related the lifetime of the photohole generated during the photoemission process. The Gaussian profile takes into account the finite resolution of the electron analyzer.

The Shirley background, named after David Shirley, is the most commonly applied background correction in x-ray photoelectron spectroscopy. It takes into account the photoelectrons which are inelastically scattered out of a core level resulting in an increased background to the high binding energy/low kinetic energy side of the elastic core level peak. More details on the definition and application of the Shirley background can be found in J. Végh, J. Electron Spectropscopy and Related Phenomena, 51, 159 (2006).