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One-Dimensional TiO₂ Nanostructured Photoanodes: From Dye-Sensitised Solar Cells to Perovskite Solar Cells

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Abstract: This review presents one dimensional (1D) TiO_2 nanostructured photoanodes for next generation solar cells such as dye-sensitised solar cells (DSCs) and perovskite solar cells (PSCs). Due to the unique morphological properties, 1D TiO_2 nanostructures can act as express electron channels as well as light scattering layer, leading to improved charge transport properties, such as charge separation, electron injection, and electron lifetime, and light harvesting efficiency. As 1D TiO_2 nanostructures are applied to solar cells, 1D TiO_2 nanostructures should be further modified to overcome some drawbacks. In this review, we have described some solutions by introducing various 1D TiO_2 synthetic methods and device fabrication processes for solar cell applications, where we have described some important surface engineering and hierarchical device design strategies that facilitate charge transport and light utilisation in 1D TiO_2 nanostructured photoanode system.

Keywords: one-dimensional (1D) TiO₂ nanostructure; photoanode; charge transport; light harvesting efficiency; dye-sensitised solar cells (DSCs); perovskite solar cells (PSCs)

1. Introduction

To address the global concerns about the energy crisis and environmental issues, one of the most promising solutions is the utilisation of solar energy. Solar energy is an abundant and free energy source, which can offer an amount of energy equivalent to that consumed by the whole world's population in an entire year through just 90 min-irradiation on the surfaces of the Earth [1]. Solar cells would be one of the commercially available approaches to make a good use of solar energy. In addition, due to the sustainability and cleanliness of solar cells, it has attracted great attention as the key green technology to replace fossil fuel energy-based technologies. Basically, the energy conversion from the sunlight to the electricity on solar cells is driven by electron-hole charge separation and transfer processes, occurring at semiconducting materials with suitable bandgap energy under the sunlight. The key technology of solar cells is closely related with the development of semiconducting photovoltaic materials, and thus current studies on solar cells are focusing on high efficiency solar cells using low-cost semiconducting photovoltaic materials as an alternative to Si-based solar cells requiring expensive processes and materials. In particular, due to the use of low-cost materials and simple fabrication processes, relatively high efficiency, and the possibility of making flexible/wearable devices, dye-sensitised solar cells (DSCs) and perovskite solar cells (PSCs) are of great interest as the alternative next generation solar cells. In 1980, Matsumura et al. [2] first reported the basic idea of dye

sensitization and surface semiconductor electrodes where the power conversion efficiency (PCE) of the reported DSCs with Rose Bengal dye was 2.5%. The current design of DSCs was proposed by O'Regan and Gratzel in 1991 [3], which provided a starting platform in further developing high efficiency DSCs, and this has led to the development of PSCs, based on the architecture of solid-state DSCs. The development progress of PSCs shows an amazing breakthrough in a short period of time since the original work on PSCs was first reported by Kojima et al. in 2009 [4]. Up to now, the certified best *PCE* of PSCs is 22.1%, which is comparable to those of high efficiency thin film solar cells such as Si-, CuInGaSe₂- and CdTe-based solar cells [5]. Interestingly, the working mechanism and device structure of the recently developed PSCs are very similar to those of DSCs, and it is particularly important to note that the use of TiO_2 in both solar cells is critical to achieve high performance [6]. In fact, TiO_2 is a versatile semiconducting material that has widely been used in photocatalytic applications associated with energy and environmental research fields [7–11]. In DSCs and PSCs, thin and compact TiO₂ blocking layers added onto the conductive substrates act as an electron transport layer (ETL) by efficiently separating and transferring charges because the conduction and valence band positions of TiO_2 , interfacing with the active semiconducting layers, are ideal to selectively transfer electrons and block holes, generated from the active layers including dyes or perovskites as a sensitiser. In addition to TiO₂ blocking layers, mesoporous TiO₂ active layers provide a scaffold to adsorb the large amount of sensitisers so that overall light harvesting efficiency of solar cells can be improved, followed by increasing photocurrent density. Therefore optimising TiO₂ layers as a photoanode material in DSCs and PSCs has been extensively studied to achieve high efficiency solar cells, and therein controlling crystallinity, thickness, and morphology of TiO_2 layers has been mainly considered [6,12].

In the past few years, one-dimensional (1D) TiO₂ nanostructured materials have continuously been investigated as the photoanode materials for DSCs. The unique morphology and novel properties of 1D TiO₂ nanostructured materials lead to improved photovoltaic performance in DSC systems by the excellent charge separation, fast charge injection, and efficient light utilisation, whereas trap sites existing in the randomly oriented mesoporous TiO₂ structure result in charge recombination, which is one of the main reasons for poor photovoltaic performances [13,14]. Like 1D TiO₂ nanostructures, 1D ZnO nanostructures also provide good charge transport properties with high crystallinity under low temperature processes, but their intrinsic photocorrosion properties are a critical drawback in terms of stability when applied to solar cell applications [15]. Therefore, in solar cell applications, the anti-photocorrosion effect of 1D TiO₂ nanostructures would be a distinctive advantage compared to 1D ZnO nanostructures [16].

In the meantime, although the utilisation of 1D TiO₂ nanostructured materials in PSC systems has been less reported than the case of DSC systems, it has received great attention as an approach in advancing PSC systems as the advantages of 1D TiO₂ nanostructured materials have already been demonstrated in DSC systems. In this review, diverse material properties of 1D TiO₂ nanostructured materials will be investigated first, and then their synthetic methodologies will be summarised along with the growth mechanism. As 1D TiO₂ nanostructured materials are applied as the photoanode of solar cell applications, the strategical design of 1D TiO₂ photoanodes will be investigated by considering various device structures and fabrication processes. Finally, the current progress and status of 1D TiO₂-applied solar cell applications from DSCs to PSCs will be comprehensively reviewed with the recent information.

2. Novel Charge Transport and Optical Properties of 1D TiO2 Nanostructured Materials

In general, 1D nanostructures are prepared with a high aspect ratio, and at least one of the dimensions should be in the range of 1–100 nm. Based on the geometrical features, 1D nanostructures are typically of four different types: nanotubes (NTs), nanorods (NRs), nanowires (NWs), and nanobelts (NBs) [17]. NTs in particular possesses a hollow channel, whereas other three different types of nanostructures have a solid structure with no hollow channel. To clarify the differences between the NR, NW, and NB nanostructures, NR usually indicates a NW with a small aspect ratio, that is to say,

a short NW [17,18]. NBs show a special NW structure with a rectangular cross-section, comprising single crystals with well-defined side crystal facets. Such geometrically unique 1D nanostructures offer novel material properties that differ from those of conventional nanoparticulate (NP) systems [17]. Obviously, the differences between 1D nanostructures and NP systems become apparent when they are applied to solar cell applications. Figure 1 shows schematic diagrams of the electron pathways in the randomly oriented mesoporous TiO_2 NP- and well-oriented 1D TiO_2 NT-based DSCs [19].



Figure 1. Schematic diagrams of electron transport pathway in (**a**) TiO₂ nanoparticulate (NP)-based photoanodes; and (**b**) TiO₂ nanotube (NT)-based photoanodes. Reprinted with permission from [19]. Copyright 2009 Royal Chemical Society.

Mesoporous TiO₂ NP layers have usually been regarded as the conventional photoanodes of DSCs due to their large surface area for effectively adsorbing dye sensitisers. However, mesoporous TiO₂ NP layers in DSC structures are not ideal in the aspect of electron transport due to the following two reasons. (1) Electron drift, which is an essential electron transport mechanism in most photovoltaic cells, is prevented in DSCs by ions in the electrolyte that screen macroscopic electric fields and couple strongly with the moving electrons. (2) By the presence of numerous grain boundaries in TiO₂ NP layers, electrons can easily be trapped, resulting in reduced electron lifetimes with the short electron diffusion length [20]. In the meantime, Zhu and co-workers [21] have reported a comprehensive investigation on charge collection efficiency in DSCs using vertically-oriented 1D TiO₂ NT arrays. When comparing the charge transport time and recombination time of TiO₂ NP- and TiO₂ NT-based DSCs with a comparable thickness of TiO₂ layers, both charge transport times were comparable, while the recombination time of TiO₂ NT layers was 10 times slower than that of TiO₂ NP layers, indicating the TiO₂ NT-based DSCs (Figure 2).



Figure 2. Comparison of (**a**) charge transport time; and (**b**) recombination time constants for NP- and NT-based dye-sensitised solar cells (DSCs) as a function of the incident photon flux for 680 nm laser illumination. Reprinted with permission from [21]. Copyright 2007 American Chemical Society.

Likewise, TiO_2 NR-based DSCs also showed longer electron lifetimes than TiO_2 NP-based DSCs with a comparable electron diffusion coefficient where the lower lifetime of the NP layers might be due to the increased charge recombination rate caused by the presence of intraband gap states and electron loss between the grain boundaries [22]. Therefore, considering the higher charge collection

efficiency with longer lifetime of 1D TiO₂ nanostructures, the thickness of 1D TiO₂ nanostructured layers in DSCs can become thicker than that of NP layers for a given recombination loss. In addition to the improvement of charge transport properties, light harvesting efficiency should also be considered for high efficiency solar cells. As seen in the following formula for a short-circuit current density (J_{sc}) of the photovoltaic performance, J_{sc} is determined by light harvesting efficiency as well as charge transport properties such as charge collection and injection efficiencies:

$$J_{\rm sc} = q \times \eta_{\rm lh} \times \eta_{\rm cc} \times \eta_{\rm inj} \times I_0 \tag{1}$$

where *q* is the elementary charge, η_{lh} is the light harvesting efficiency, η_{cc} is the charge collection efficiency, η_{inj} is the charge injection efficiency, and I_0 is the incident photon flux (light intensity) [21]. Therefore, it is necessary to utilise a suitable light management strategy in enhancing the light harvesting efficiency, and one of these is the solar cell architecture. It is well-known that the optical properties of TiO₂ nanostructures are closely related to their atomic arrangement and light injecting direction, so 1D TiO₂ nanostructures with unique morphological properties can provide an excellent light scattering effect. A work by Nakayama and co-workers [23] compared TiO₂ NP- and TiO₂ NT-based DSCs with the same layer thickness of 10 μ m, and their power conversion efficiencies were 5.27% with J_{sc} of 10.1 mA·cm⁻² and 3.41% with J_{sc} of 7.4 mA·cm⁻², respectively. Despite the lower amount of adsorbed dyes of the TiO2 NT-based, Jsc of the TiO2 NT-based was higher than that of the TiO₂ NP-based, indicating a higher performance of the TiO₂ NT-based that could be attributed to their higher light harvesting efficiency by the internal light scattering of TiO₂ NT arrays. Meanwhile, Yun and co-workers [24] reported an investigation on optical modelling of DSCs with different thicknesses of TiO₂ NT layers of 3.3 µm, 11.5 µm, and 20.6 µm via the generalised transfer matrix (GTMM) method. Based on the optical modelling, short 3.3 µm thick TiO₂ NT-based DSCs presented relatively low light absorption with fluctuating light fraction intensity, while thicker TiO₂ NTs with 11.5 μ m and 20.6 μ m showed significantly decreased reflectance results with the increased light absorption, followed by the higher charge generation rate with the increase in thickness of TiO₂ NT (Figure 3) [24].



Figure 3. (**a**–**c**) Calculated absorption, reflectance and (**d**–**f**) charge generation rate of the DSCs with different TiO₂ NT arrays using generalised transfer matrix method (GTMM). (**a**,**d**) 3.3 μ m thick TiO₂ NT arrays; (**b**,**e**) 11.5 μ m thick TiO₂ NT arrays; and (**c**,**f**) 20.6 μ m thick TiO₂ NT arrays. Reprinted with permission from [24]. Copyright 2014 Beilstein-Institut.

Besides vertically-oriented TiO₂ NTs, even randomly-oriented TiO₂ NWs led to intensive light scattering by the bundles of TiO₂ NWs, thereby improving the optical absorption of the photoanodes

(Figure 4). For a 2.5 μ m thick TiO₂ NW layers, 20% to 40% of visible light was observed to be scattered by transmission and refection [25].



Figure 4. Fibrous network structure photoanodes comprising TiO₂ NTs. (**a**) Scanning electron microscope (SEM) image; and (**b**) transmittance and reflectance spectra of the photoanodes. Reprinted with permission from [25]. Copyright 2010 American Chemical Society.

3. Synthesis of 1D TiO₂ Nanostructured Materials

1D TiO₂ nanostructures can be prepared by a variety of synthetic methods, and depending on the method used, the morphological types and physicochemical properties of the 1D TiO₂ nanostructures can be changed, thereby influencing the charge transport and optical properties in solar cell photoanodes. Another key point in synthesising 1D TiO₂ nanostructures is the orientation status of 1D TiO₂ nanostructured layers grown onto the substrates, which can be employed as the form of photoanodes. Thus, both randomly oriented and self-oriented cases of 1D TiO₂ nanostructures are discussed with different synthetic methods.

3.1. Hydrothermal Method

The hydrothermal method is a suitable synthetic approach to prepare single crystalline 1D TiO_2 nanostructured materials which are dependent on the solubility of the precursors in water under high temperature and pressure conditions. Usually, the hydrothermal process is performed in an autoclave with a Teflon liner under controlled conditions of temperature, pressure, pH and additives. With such experimental conditions, different morphologies and crystal phases such as anatase, rutile and brookite can be obtained. As TiO_2 nanoparticle-based suspension solution in alkaline condition are used as precursor for the hydrothermal reaction, suspended TiO₂ nanoparticles are hydrothermally transformed into TiO_2 NTs (Figure 5a) [26]. The formed TiO_2 NTs can be selectively formed as anatase and rutile phases by using different precursor ratios, hydrothermal conditions, and salt incorporation [27,28]. Besides, it was reported that single crystalline anatase TiO₂ NRs were transformed from NTs by the local shrinkage of the NT walls and the subsequent oriented attachment of crystallites when the pH of NT suspension in 10 M NaOH was controlled by dilute HCl solution in the ion-exchange process [29]. 1D TiO₂ nanostructures can also be directly grown onto Ti metal substratse or conductive glass substrates through the hydrothermal process, and this would be beneficial to fabricate the 1D TiO₂-based photoanodes for solar cells. Although the synthesis of the randomly oriented 1D TiO₂ nanostructures can produce large amounts of 1D TiO₂ materials with a large surface area, the randomly oriented 1D TiO_2 -based photoanodes should be prepared through the additional fabrication steps such as TiO₂ paste preparation and doctor-blade printing. Therefore, in-situ growth of 1D TiO₂ nanostructures onto the substrates would offer a simple and easy process and improved interfacial properties between 1D TiO_2 layers and the surfaces of the substrates. Recently, many studies have reported the fabrication of vertically-oriented TiO₂ NRs and NWs onto various substrates using hydrothermal methods as seen in Figure 5b,c. Without any substrates or seeding layers, the as-prepared 1D TiO₂ materials were random and existed in a powder form, while vertically-oriented 1D TiO₂ layers

can be achieved in the presence of substrates with thin TiO_2 seeding layers. The seeding layer could be prepared by dip- or spin-coating TiO_2 precursor solution, followed by calcination of the layers at 450 °C for 1 h. Typically, titanium (IV) butoxide in HCl solution was applied as the precursor for the hydrothermal reaction, performed at 170 °C for 6 h. Then, the concentrated HCl constraint on the hydrolysis of the TiO₂ precursor results in 1D TiO₂ NRs. The growth of oriented 1D TiO₂ NRs requires slow hydrolysis of the precursor in a strong acidic aqueous medium, while reduction in acidity of the precursor solution might have prompted hydrolysis of titanium (IV) butoxide. Through the optimised hydrothermal process with tuned hydrolysis condition, well-oriented single crystalline rutile TiO₂ NRs with an excellent uniformity are obtained [30].



Figure 5. SEM images of 1D TiO_2 nanostructures using a hydrothermal method. (a) Randomly oriented TiO₂ NTs. Reprinted with permission from [31]. Copyright 2006 American Chemical Society; (b) Well-aligned TiO₂ NRs. Reprinted with permission from [30]. Copyright 2011 American Chemical Society; (c) Well-aligned TiO₂ NWs. Reprinted with permission from [32]. Copyright 2013 Nature Publishing Group.

3.2. Template-Assisted Method

Template-assisted synthesis offers a reproducible synthetic approach for precise control over the size, shape and configuration, and growth direction of 1D nanostructured materials. In a typical template-assisted synthesis synthetic process, an anodised aluminium oxide membrane (AAM) is used as a template that possesses a self-oriented porous structure with tightly controlled pore size, density, and intervals. Thus, as Ti-precursors penetrate into the porous membrane structure through various deposition processes such as electrochemical deposition (ECD), atomic layer deposition (ALD), chemical vapour deposition (CVD), physical vapour deposition (PVD), sol-gel deposition, reactive ion etching (RIE), and vapor-liquid-solid (VLS) growth etc., well-oriented 1D TiO₂ nanostructured materials with tunable dimensions such as lengths and diameter can be achieved. Figure 6 shows the general processes for the fabrication of well-oriented nanostructured arrays using an AAM template.



Figure 6. Schematic diagrams on the fabrication of well-oriented nanostructured arrays of: (a) nanoholes; (b) nanodots; (c) NRs; and (d) NWs using an AAM template. Reprinted with permission from [33]. Copyright 2014 American Chemical Society.

In Figure 6a,b, nanoholes and nanodots are a kind of intermediate stage prior to fabricating 1D nanostructures, and they can be prepared by RIE and PVD methods, respectively. In particular, nanodots can simply be achieved by depositing the precursors of the desired materials through an AAM template where the array pattern with controlled size and interdistance of the nanodots is defined by the template, and further well-oriented 1D nanostructured arrays are grown along the nanodot pattern.

Hoyer [34] first reported 1D TiO₂ NT arrays using the template-assisted method via replication and electrodeposition process. Figure 7a shows fabrication of TiO₂ NT arrays by electrodeposition of TiO₂ onto the surfaces of polymethylmethacrylate (PMMA) rods as a template with gold film substrates replicated from porous AMM template. After dissolution of PMMA template in acetone at 40 °C, well-oriented TiO₂ NT arrays were fabricated (Figure 7b).



Figure 7. (a) Schematic diagram of the template-assisted synthetic process via replication and electrodeposition; and (b) SEM image of top and cross-section views for TiO₂ NTs obtained after 20 min-electrodeposition. Reprinted with permission from [34]. Copyright 1996 American Chemical Society.

Similar to the AAM template-assisted method, ZnO NR arrays can be employed as the template for 1D TiO₂ NT arrays. Yodyingyong and co-workers [35] reported the formation of well-oriented nanoporous TiO₂ NT arrays on transparent conductive oxide (TCO) substrates using ZnO NR arrays as the template. As seen in Figure 8, the ZnO NR template is prepared by the hydrothermal method, and then TiO₂ NTs are formed onto the surfaces of ZnO NR template by immersing the template substrate in Ti-precursor solution at room temperature for 1.5 h, and further the ZnO templates are dissolved by TiO₂ deposited ZnO template substrate in acidic solution for 1 h. More importantly, this ZnO NR template-assisted method offers in-situ assembly of well-oriented 1D TiO₂ NT arrays with porous tube walls onto the TCO substrates. Although well-oriented 1D TiO₂ nanostructures can be achieved through the AAM template-assisted method, adhesion issues near the interfaces between the 1D nanostructures and the surfaces of substrates occur. In particular, wet-etching the AAM template and transplanting 1D TiO₂ arrays to the TCO substrates are necessary to fabricate the photoanodes for solar cells, but those processes may cause breakdown of unstable parts of 1D TiO₂ nanostructures and poor interface properties due to adhesion issues, while ZnO NR templated onto TCO substrates may offer a proper solution due to the preferential etching characteristic along the (0001) orientation, facilitating the mild template removal with a top-down etching process. Additionally, the formation of

porous walls of 1D TiO₂ NT arrays by etching processes using TiCl₄ contributes to a high surface area of 119 m²·g⁻¹, expected to increase the amount of dye adsorption in DSCs.



Figure 8. Schematic diagram of formation of TiO₂ NT arrays by ZnO NR template-assisted method. Reprinted with permission from [35]. Copyright 2010 American Chemical Society.

3.3. Electrochemical Anodisation Method

As discussed earlier, the template-assisted method seem to be more beneficial compared to the hydrothermal method in terms of reproducible preparation of well-oriented 1D TiO₂ nanostructures with precise control of the geometric features such as lengths and diameters. However, the template-assisted methods require an additional dissolution process to remove undesired templates such as AAM, PMMA mold, and ZnO NR arrays. Furthermore, the dissolution process may also damage the desired 1D TiO₂ nanostructures. Thus, the electrochemical anodisation approach for 1D TiO₂ nanostructures has been of great interest in fabricating stable vertically-oriented TiO₂ NT arrays because it is a template-free process with no additional post-treatment. The geometry of the anodised TiO₂ NT array can readily be controlled by modulating the anodising conditions such as anodising time, electrolyte condition, and applied potential. In contrast, the template-assisted method is limited by the geometry of the applied template [14].

Typically, the anodic TiO_2 NT arrays are grown by anodising pure Ti foil in an electrolyte including fluoride ions. During anodisation, the growth of TiO_2 NT arrays is driven by the competition between anodic oxidation and chemical/field-assisted dissolution (equations in Figure 9). Figure 9 shows the comprehensive TiO₂ NT growth mechanism by anodisation. When starting the anodization, dense and thin TiO_2 layers are quickly formed onto the Ti foil substrates (Figure 9a), and subsequently undergo localised dissolution to form small pores (Figure 9b). The localised dissolution makes the barrier layer thinner, leading to an increasing electric field intensity across the barrier layer to further grow pores (Figure 9c). In fact, the barrier layer acts as a resistance to the flow of ions such as Ti⁴⁺ and O²⁻, which need to transport through the anodic oxide layer to maintain the oxidation active. In the meantime, higher resistance caused by thicker barrier layers can be overcome by increasing the applied potentials for a higher electric field. The presence of fluoride ions in the electrolyte allows the creation of channels in which current flows to keep the oxidation process active [36]. The thickness of the barrier layers underneath the pores/tubes is continuously reduced by dissolution, and simultaneously regenerated by oxidation. In Figure 9d, the electric field distribution at the bottom surface of the pore causes widening and deepening of pores. As the pores get deeper, the electric field in the exposed metallic regions increases, thus enhancing the field-assisted oxide growth and oxide dissolution. At the same time, the inter-pore voids start to be formed. Thereafter, both voids and pores grow in equilibrium. The TNT length increases until the oxidation rate at the metal/oxide interface equals the chemical dissolution rate of the top surface of the tubes (at the oxide/electrolyte interface). Afterwards, the TNT length is independent of anodizing time, as a result of a given electrolyte concentration and applied potential [37,38].



Figure 9. Schematic diagram of the TiO_2 NT growth mechanism. (a) Growth of dense and thin oxidation layer; (b) formation of pores onto the layer; (c) growth of the pores and repassivation of the pore tips; (d) growth of voids in the metallic part between pores under oxidation and chemical dissolution; and (e) the growth of fully developed TiO_2 NT arrays. Reprinted with permission from [38]. Copyright 2012 The University of New South Wales.

As the anodisation process is properly modified, defect-free anodic 1D TiO_2 NT arrays can be fabricated. Wang and co-workers [39] demonstrated a novel protocol to prepare defect-free 1D TiO_2 NT arrays using a two-step anodisation process. In the first anodisation step, the grown NT arrays are peeled off thereby the ordered footprint of NTs are left on the surface of the Ti substrate. After the second anodisation step, nanoporous-layer-covered well-aligned NT arrays are obtained. The defect-free condition of 1D TiO_2 nanostructures as the photoanodes for solar cells would suppress charge recombination by reducing charge trapping sites.

The morphology of well-oriented TiO₂ NT arrays can be tuned by modulating anodising conditions, thereby length, pore diameter, and wall thickness, and wall surface roughness are controllable. Yun and co-workers [14,40–42] comprehensively investigated the geometrical properties of TiO₂ NT arrays with different experimental conditions. Under the optimised anodisation conditions with the applied potential of 60 V in the electrolyte containing 0.5 wt % NaF and 5 wt % water, geometry of TiO₂ NT arrays was significantly changed with anodising time. As seen in Figure 10a,b, 3 h-, 10 h-, and 15 h-anodised TiO₂ NT arrays present 3.3 μ m, 11.5 μ m, and 20.6 μ m in length, respectively, and the pore diameter also increases from 50 nm to ~90 nm with the increase in anodising time.



Figure 10. Geometry of TiO_2 NT arrays: (a) length; and (b) pore diameter and wall thickness as a function of anodising time; (c) cross-sectional SEM images of TiO_2 NT arrays for 3 h, 10 h, and 15 h under 60 V applied potential in the electrolyte containing 0.5 wt % NaF and 5 wt % water. Reprinted with permission from [14]. Copyright 2011 American Chemical Society.

The proportional increase in NT lengths can be attributed to the continuous oxidation of Ti foil, whereas the decrease in tube wall thickness is due to chemical dissolution of the oxide. In Figure 10c, well-oriented TiO_2 NT arrays are observed over the large area of Ti foil substrates with the uniform growth of lengths and pore diameters, and those TiO_2 NT arrays can directly be employed as the photoanodes of solar cells without further fabrication processes [14,24].

4. 1D TiO₂ Nanostructure-Based Solar Cell Applications: DSCs and PSCs

In light absorber-sensitised solar cell systems such as DSCs and PSCs, the photovoltaic performance is driven by the charge transport properties and light harvesting efficiency, and therefore strategic approaches to enhance electron transport and lifetime and light scattering should be considered. In this regard, 1D TiO₂ nanostructured materials would be an excellent candidate to meet such requirements in the cases of both DSCs and PSCs owing to the novel charge transport and optical features of unique 1D TiO₂ nanostructures. In this review, thus, as 1D TiO₂ nanostructures are applied as the photoanodes, the desirable strategical architectures in DSCs and PSCs will be discussed with various ideas suggested by the current research progress.

4.1. 1D TiO₂ Nanostructure-Based DSCs

In 1991, O'Regan and Gratzel [3] reported low-cost DSCs fabricated using mesoporous TiO₂ layers. The DSCs are basically a thin-layer solar cell fabricated by sandwiching two TCO electrodes (typically fluorine-doped tin oxide or indium-doped tin oxide). The photoanoode is composed of a mesoporous TiO₂ layer adsorbed with a dye as a photosensitiser deposited onto the TCO substrate. The counter electrode is composed of finely distributed Pt films deposited on another TCO substrate. The inner space between a photoanode and a counter electrode is filled with a liquid electrolyte containing I^-/I_3^- redox couple in acetonitrile. Figure 11 presents a schematic diagram on the principles of DSC operation. Under sunlight illumination, excited electrons generated from the dyes are injected into the conduction band of TiO₂, and then the injected electrons are extracted to the external load.



Figure 11. Schematic diagram on working mechanism of DSCs. Reprinted with permission from [38]. Copyright 2012 The University of New South Wales.

Meanwhile, the resulting oxidised dyes are reduced rapidly to the ground state by I⁻ in the electrolyte and the electrons reaching the Pt counter electrode through the external load reduce back the formed I₃⁻ to I⁻, thereby the DSCs have a cyclic electron transfer process [3,43,44]. The current best *PCE* obtained from the DSCs is 12.3% using porphyrin sensitisers and cobalt (II/III) redox electrolyte [45]. In the development of 1D TiO₂ nanostructure-applied DSCs, 1D TiO₂ NTs prepared by a hydrothermal method were first applied to DSCs. Compared to mesoporous TiO₂ NP systems using commercial P25 TiO₂ nanoparticles, TiO₂ NT-DSCs showed higher *PCE* of 6.4% than TiO₂ NP-DSCs with *PCE* of 5.5%, attributed to the improved electron transfer along the NT structure. After TiCl₄ post-treatment, the *PCE* of 1D NT-applied DSCs further increased to 7.1%, which is

attributed to the facilitation of the electron transport by the improved interparticle connectivity [46]. Karthik and co-workers [47] reported backside-illuminated DSCs with vertically oriented TiO₂ NT arrays with long lengths of 20 μ m, achieving a *PCE* of 6.9%. However, 1D TiO₂-DSCs have still been struggling to reach the high photovoltaic performance of the mesoporous TiO_2 -based DSCs. The lower efficiency of 1D TiO₂-DSCs is mainly due to the relatively low light harvesting efficiency that can be attributed to the small amount of dye adsorption by their small surface area and low transmittance by backside-illumination. Recently, various strategies to overcome those drawbacks by surface engineering and electrode architecture have been reported. So and co-workers [48] achieved high PCE of 8.09% with conical-shaped TiO₂ NT arrays as the photoanodes under backside-illumination conditions, wherein the conical-shaped TiO₂ NT arrays were prepared by two step anodisation processes using an electrolyte containing 0.2 M NH₄F, 50 vol.% lactic acid and 50 vol.% triethyleneglycol (Figure 12a). In spite of light absorption loss by the backside-illumination, the significant improvement of 1D TiO₂ NT-DSCs was mainly due to the efficient light utilisation management by the conical shaped tips of the tubes. Although the modification of the 1D nanostructures can contribute to improving the light utilisation in solar cells, the backside-illumination condition still limits the overall light harvesting efficiency so that the frontside-illumination condition is more preferable to enhance the overall photovoltaic performances of the cells. As shown in Figure 12b, Lv and co-workers [49] applied porous rutile TiO₂ NR arrays onto TCO substrates for enhancing the performance of DSCs, where $\sim 30 \ \mu m$ thick TiO₂ NR arrays were prepared by a hydrothermal method, and the completed NR arrays were further etched by HCl. The etched TiO₂ NR arrays showed much higher dye loading compared to their counterpart mesoporous TiO_2 , leading to an improved PCE of 7.91% with J_{sc} of 20.49 mA·cm⁻². Figure 12c shows the anodised 1D TiO₂ NT arrays are directly grown onto the TCO substrates. In this case, instead of Ti foil substrates, Ti metal deposited TCO was used as the source material for the anodisation, and thus TiO_2 NT arrays can be strongly grown onto the TCO substrates [50]. Hierarchical designs integrating more than two heterostructured nanomaterials provide multiple advantages in optimising the solar cell architectures. As seen in Figure 12d,e, 0D and 3D nanostructured materials can be incorporated with 1D nanostructured materials in various arrangements, leading to efficient light scattering and electron transport properties and large dye loading, followed by improving overall photovoltaic performances. Similar to Figure 12c, Figure 12d also presents frontside-illuminated 1D TiO2 NT-DSCs using the anodised TiO2 NT arrays. However, in this work, the fabrication procedure of the anodised 1D TiO_2 NT arrays onto the TCO substrates is different from that of Figure 12b. Instead of using Ti metal deposited TCO substrates, the anodised TiO_2 NT arrays were transplanted onto the TCO substrates by imprinting the peeled off the TiO_2 NT arrays from Ti foil substrates. When applying the peeled off TiO₂ NT arrays as the photoanodes, randomly oriented TiO₂ NPs are incorporated onto the bottom or the top of the TiO₂ NT arrays. In the junction structure of "FTO-double TiO₂ NP layers-17 µm thick TiO₂ NT arrays", the photoelectron lifetime was extended by 237.5% and the PCE was 8.8% with J_{sc} of 18.89 mA·cm⁻² [51]. In Figure 12e, the photoanodes are composed of branched 0D-3D TiO₂ composites on TiO₂ NT arrays at the bottom, where 0D TiO₂ nanoparticles are on the top of the 3D TiO₂ microspheres. Particularly, the branched 1D TiO₂ NT arrays and 0D-3D TiO₂ nanocomposite layers provide rough surfaces, contributing to improving the charge transport capacity and light harvesting efficiency, resulting in a PCE of 9.1% with $J_{\rm sc}$ of 17.9 mA·cm⁻² [52].



Figure 12. Schematic diagrams on the photoanode architectures in the 1D TiO₂ nanostructure-applied DSSCs. (a) Cylindrical- and conical-shaped TiO₂ NT arrays; (b) porous rutile TiO₂ NR arrays by the post-etching process. Reprinted with permission from [53]. Copyright 2012 American Chemical Society; (c) Frontside-illuminated TiO₂ NT-DSCs using Ti metal-deposited TCO substrates. Reprinted with permission from [50]. Copyright 2006 American Chemical Society; (d) Hierarchical layer-by-layer photoanode incorporating TiO₂ NT array and TiO₂ NP layer. Reprinted with permission from [51]. Copyright 2011 American Chemical Society; (e) Hierarchical structured photoanode incorporating branched TiO₂ NT arrays and TiO₂ hollow microspheres. Reprinted with permission from [52]. Copyright 2014 American Chemical Society.

Taking all the above 1D TiO₂ photoanode designs for DSCs into account, the use of 1D TiO₂ nanostructures as the photoanode in solar cells leads to a significant change in designing solar cell device structures compared to the conventional mesoporous TiO₂ photanode systems with randomly oriented structures, wherein both advantages and disadvantages of 1D TiO₂ nanostructures in solar cell application can be found. Although the relatively smaller surface area of 1D TiO₂ nanostructures results in low device performance due to the low dye loading amount, such drawbacks can be overcome through strategic approaches to modify 1D TiO_2 nanostructures and the architecture of 1D TiO_2 -DSCs. Specifically, the low light harvesting efficiency of 1D TiO_2 -DSCs is a key issue in improving device performance, and it can be improved by excellent light scattering effect of the modified conical shaped TiO₂ NTs (Figure 12a) and increasing incident light absorption by the front illumination device architecture (Figure 12c). Another approach to enhance light harvest efficiency is by increasing the surface areas of 1D TiO_2 photoanodes to uptake a large amount of dyes. This can be achieved by using porous 1D TiO₂ nanostructures (Figure 12b) and 0D/3D/1D integrated TiO₂ nanostructures (Figure 12d,e). In particular, the integrated 1D TiO_2 photoanodes lead to significantly improved light harvesting efficiency through the synergistic effect of the integrated 0D/3D nanoparticles with large amount of dye loading and enhancing light scattering. Recently, Mahmood and co-workers [54] have reported highly efficient hyperbranched 3D TiO₂ photoanodes consisting of four-layered 1D TiO₂ nanostructured arrays with a PCE of 11.22%, which is comparable to that of high efficiency DSCs using mesoporous TiO₂ photoanodes. The remarkable performance of the unique hyperbranched structured DSCs is due to their significantly enhanced electron lifetime and light harvesting efficiency, demonstrating the great potential of the hyperbranched structures in next generation solar cells.

As the intermediate stage in DSC progress prior to PSCs, solid-state DSCs (SSDSCs) have been extensively studied as the alternative way to solve some of the potential drawbacks of DSCs. Although DSCs have already achieved a high *PCE* of 12.3%, the liquid electrolyte in DSCs may cause potential leakage and corrosion problems. Thus it would be meaningful to replace liquid electrolytes with solid electrolytes to overcome such disadvantages of DSCs. In addition, 1D TiO₂-SSDSCs have also been investigated as one of the promising approaches for high performance. Most importantly,

it is noted that the working mechanism and the device structure of SSDSCs are analogous to those of PSCs. Thus, it would be meaningful to briefly review 1D TiO₂-SSDSCs before moving on to the 1D TiO₂-PSC applications. The main key in SSDSCs would be the use of solid-state hole transport materials (HTMs) to replace the liquid electrolytes found in DSCs. Typically, p-type inorganic materials or organic materials are applied as the HTMs in SSDSCs. spiro-MeOTAD (2,2',7,7'-tetrakis-(N,N-di-p-methoxyphenylamine)-9,9'-spirobifulerene) is used as HTM in SSDSCs [55]. The SSDSCs using spiro-MeOTAD as a HTM have already achieved a PCE of 5.1%, but this PCE is still lower than that of DSCs using liquid electrolytes [56]. Ding and co-workers [57] recognised that the main reason would be a pore filling issue of HTM into the dye-sensitised photoanode internal structure, and it was indicated that a 2.5 μ m thick TiO₂ layer with 60%–65% pore filling prepared by optimising the spin coating conditions was the best solution. Likewise, in the case of 1D TiO_2 -SSDSCs, the optimised pore filling conditions should be investigated by considering the thickness of 1D TiO_2 arrays and spin coating conditions. Mor and co-workers [58] investigated 1D TiO₂ NT-SSDSCs, mimicking a typical bi-layered organic photovoltaic architecture as seen in Figure 13a. In this device structure, 1D TiO₂ NT arrays were fabricated by anodising Ti metal-deposited TCO substrates (refer to Figure 12b), and poly(3-hexylthiophene-2,5-diyl) (P3HT) as a p-type HTM and PEDOT:PSS as an electron blocking layer were applied to SSDSCs. In the charge transfer processes of Figure 13b, excited electrons from dye and P3HT are injected to the conduction band of TiO₂, while holes are efficiently separated and transferred to Au through the PEDO:PSS layers by a p-type P3HT HTM.



Figure 13. Schematic diagram of (**a**) 1D TiO_2 NT-SSDSCs in the frontside-illumination condition; and (**b**) energy level positions and charge transfer processes of the SSDSCs. Reprinted with permission from [58]. Copyright 2009 American Chemical Society.

As pore filling is a key issue in SSDSCs, the deposition of P3HT into the NTs was performed by spin coating after pre-wetting NT arrays with *p*-xylene, a non-polar solvent, acting to penetrate P3HT inside the NT arrays. With 600–700 nm thick and ~35 nm pore diameter, the achieved *PCE* was 3.8%. Roh and co-workers [59] reported high efficiency SSDSCs prepared using hierarchical anatase pine tree-like TiO₂ NT (PTT) arrays in which the hierarchical anatase PTT arrays were fabricated directly onto the TCO substrates by a one-step hydrothermal method. The applied electrolyte was poly((1-(4-ethenylphenyl)methyl)-3-butylimidazolium iodide) (PEBII), which was deposited onto the PTT arrays by two-step drop casting. The SSDSCS assembled with 19 μ m thick PTT arrays showed an excellent *PCE* of 8%, which was twice higher than that of commercially available TiO₂ paste with a *PCE* of 4%, representing one of the highest records achieved from N719 dye-based SSDSCs. In Table 1, we summarise the photovoltaic performance of DSCs and SSDSCs using 1D TiO₂ nanostructured photoanodes.

TiO ₂ Type; Fabrication Method ¹	Architecture ²	η	$J_{ m sc}$	$V_{\rm oc}$	FF	Dye Type; Dye Concentration	Reference				
		%	mA·cm ⁻²	v	%	nmol⋅cm ⁻²					
DSCs with 1D TiO ₂ Nanostructured Photoanodes											
Layered NTs; HT	FI; 10 μm thick NT particulate layer on FTO	7.10	13.3	0.79	0.68	N719; 150	[46]				
Layered and Hyper-branched NRs; HT/ES	FI; 29 μm thick 4 layered Hyperbranched NRs on FTO	11.22	18.60	0.84	0.69	N719; 240	[54]				
Vertical NT arrays; SG + AAO Template	FI; 25 μm thick NT-AAO arrays on ITO	2.70	5.5	0.72	0.69	N3	[60]				
Vertical NW arrays; HT	FI; 47 μm thick NW arrays on FTO	9.40	18.25	0.79	0.65	N719; 160	[<mark>61</mark>]				
Vertical NT arrays; Anodisation	BI; 6 µm thick NT arrays on Ti foil	4.24	8.79	0.84	0.57	N719	[62]				
Vertical and Conical shaped NT arrays; Anodisation	BI; 13 μm thick NT arrays on Ti foil	8.09	15.63	0.79	0.66	N719; 144	[48]				
Vertical NT arrays; Anodisation	FI; 63 µm thick NT arrays on FTO	9.10	18.50	0.77	0.64	N719; 152	[63]				
Vertical and Porous rutile NR arrays; HT	FI; 30 µm thick NR arrays on FTO	7.91	20.49	0.71	0.54	N719; 133	[49]				
Vertical and Hierarchical NT arrays;	FI; 17 μm thick; FTO/double 0D NP/1D NT	8.80	18.89	0.75	0.62		[51]				
Anodisation/SG	FTO/ 1D NT/double 0D NP	6.03	13.07	0.75	0.62	N719					
Vertical and Hierarchical NT arrays; HT/SG	BI; 28 μm thick; 0D NP/3D Microsphere/1D NT/Ti foil	9.10	17.90	0.74	0.69	N719; 148	[52]				
SSDSCs with 1D TiO ₂ Nanostructured Photoanodes											
Vertical NT arrays; Anodisation	FI; FTO/600–700 nm thick NT-dyes/P3HT/PEDOT:PSS/Au	3.2	10.75	0.55	0.55	SQ-1	[58]				
Vertical and Hierarchical NT arrays; HT	FI; FTO/19 μm thick Hierarchical NT arrays-dyes/PEBII/Pt	8.00	17.70	0.74	0.62	N719; 106	[59]				

Table 1. Summary of the photovoltaic characteristics for DSCs and SSDSCs using 1D TiO₂ nanostructured photoanodes.

¹ In type and fabrication method, "NTs", "NRs", and "NWs" indicate "nanotubes", "nanorods", and "nanowires", respectively. "HT", "ES", and "SG" indicate "hydrothermal method", "electrospinning method", and "sol-gel method"; ² In Architecture, "FI" and "BI" indicate "frontside illumination" and "backside illumination", respectively.

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4.2. 1D TiO₂ Nanostructure-Based PSCs

Since Kojima and co-workers [4] first reported PSCs comprising perovskite light absorbers such as CH₃NH₃PbI₃ and CH₃NH₃PbBr₃ in a liquid electrolyte, the progress of PSCs has been dramatically fast in a short-term period. Recently, Seok's group [5,64] has reported a world-best *PCE* record of 22.1% achieved by incorporating methylammonium (MA) and formamidinium (FA) ions, which has attracted great attention to the potential of low-cost and simple solution processed solar cells compared to expensive Si-based solar cells. The working mechanism of PSCs is shown in Figure 13. Basically, PSCs are composed of an n-type perovskite-sensitised photoanode, p-type HTM layer, and an Au metal contact. Depending on the presence of TiO₂ active layers, PSCs can be classified to two different device structures: Figure 14a indicates planar structured PSCs without a TiO₂ active layer, while Figure 14b presents mesoscopic PSCs with a mesoporous TiO₂ active layer. Similar to the working principles of DSCs and SSDSCs, electron-hole charge separation by injecting electrons from perovskites to TiO₂ take place in PSCs, and subsequently the injected electrons are extracted to an external load through the TCO (FTO) substrate, while the separated holes from the perovskite are transferred to Au metal contact via a HTM layer. However, due to unique perovskite properties such as charge transport and charge accumulation, the working mechanism should be different from DSCs and SSDSCs.



Figure 14. (**a**) Planar structure without a mesoporous layer; and (**b**) mesoscopic PSCs with mesoporous TiO₂ layer. Reprinted with permission from [65]. Copyright 2014 American Chemical Society.

For instance, $CH_3NH_3PbI_{3-x}Cl_x$ perovskite thin films shows both electron and hole diffusion lengths are over 1 μ m, indicating a dimension 5–10 times greater than the absorption depth. The larger diffusion length leads to a much longer recombination lifetime [66]. Electron and hole mobilities were also found to be as high as $25 \text{ cm}^2 \cdot \text{Vs}^{-1}$, and both mobilities were almost balanced and remained high along with a slow microsecond time scale for recombination [67]. However, the photovoltaic performance may decrease in case the mobility of the injected electrons in the TiO₂ layer is slower than that of the perovskites. Thus, as considering the diffusion length and the mobility of the injected electrons, strategical design of the TiO_2 layers would be important to improve the performance [6]. Since the advantages of 1D TiO₂ nanostructures in DSCs are still a promising option in designing the device structure of PSCs, the use of 1D TiO_2 nanostructures would contribute to improving the performance of PSCs. In comparison with the absorption coefficient of Ru-based N719 dye, the absorption coefficient of CH₃NH₃PbI₃ perovskite was an order of magnitude higher than that of the N719 dye. Furthermore, despite the small amount of perovskite loading onto TiO_2 , the photocurrent density was twice higher than that of the fully loaded N719 dye onto TiO₂ [68]. In this regard, Kim and co-workers [69] first employed TiO₂ NR arrays as the photoanode to PSCs using CH₃NH₃PbI₃ perovskite (Figure 15a). In Figure 15c, the *PCE* was the highest at 9.4% with 560 nm thick NR arrays, and the performance decreased with an increase in NR lengths. This indicates that the shorter NRs more efficiently utilise the short wavelength light than the longer NRs, related with the higher pore filling fraction of the short NRs.



Figure 15. (a) Cross-sectional SEM image of 1D TiO₂ NR-PSCs; (b) *I-V* curves of 1D TiO₂ NR-PSCs with different lengths; and (c) *IPCE* of 1D TiO₂ NR-PSCs with different lengths. Reprinted with permission from [69]. Copyright 2013 American Chemical Society.

Wu and co-workers [70] investigated the dimensional effects of TiO₂ electron transport layers (ETL) in the PSC performance by comparing 0D NP, 1D NW, and 2D nanosheets (NS), where TiO₂ NP, NW, and NS were tuned by the different organic solvents during the solvothermal reaction. These fabricated TiO₂ layers presented enhanced optical transparency, minimising light absorption loss. The photovoltaic performances of PSCs using 1D NW and 2D NS photoanodes were higher than that of the 0D NP counterparts. Basically, efficient infiltration of perovskite sensitisers in 1D NW and 2D NS structures would improve light absorption efficiency. Particularly, 1D NW-based PSCs demonstrated an excellent performance with a *PCE* exceeding 16%, which was due to the bilayered ETL system consisting of a self-assembled compact and thin TiO₂ blocking layer and NW active layer. This bilayered ETL film can lead to blocking the photogenerated holes and improve electron extraction, thereby enhancing PSC performance.

Similar to the DSCs, 3D/1D combined layered hyperbranched TiO₂ NR photoanodes also led to improving PSC performance, indicating a *PCE* of 15.5% with J_{sc} of 20.5 mA·cm⁻², V_{oc} of 1 V, and *FF* of 0.75. 3D hyperbranched structures are suitable for high light harvesting efficiency with large surface area, while 1D NRs included in the 3D hyperbranched structures can efficiently transfer electrons over long distances, suppressing recombination losses.

Surface engineering of 1D TiO₂ NRs by the atomic layered deposition (ALD) method provides a uniform and ultrathin passivation layer on the surface of NRs, leading to improved photovoltaic performance with high V_{oc} and *FF*. The enhancement of V_{oc} and *FF* is mainly due to the interface properties. The ultrathin passivation layer by ALD offers pinhole-free surfaces over the TCO substrates and TiO₂ NR arrays, thereby improving pore filling and charge generation, and reducing recombination. Particularly, the back-flow of electrons from the TiO₂ conduction band to CH₃NH₃PbI₃ and HTM was minimised by the ALD passivation layer (Figure 16). With the optimised ALD passivation thickness of 4.8 nm, the *PCE* of 1D TiO₂ NR-PSCs was 13.45% with V_{oc} of 0.945 V, J_{sc} of 19.78 mA·cm⁻², and *FF* of 72% [54].





Figure 16. Schematic diagram of the energy level of 1D TiO₂ NR-PSCs. Reprinted with permission from [71]. Copyright 2015 American Chemical Society.

Recently, Lin and co-workers [72] have reported the best efficiency of 1D TiO₂ NR-PSCs by optimising NR diameter, densities, and surfaces through a facile solvothermal method. Surface treatment using UV-ozone on TiO₂ NR array also contributed to increasing the *PCE* from 13.58% to 14.72%. Through the surface treatment, the remaining organic contaminants resulting from the solvothermal process could be successfully removed, and further the adhesion between NR array and the perovskite layer was strengthened by the surface modification, which could promote faster electron injection between the perovskite layer and TiO₂. Therefore, an enhancement of *J*_{sc}, *V*_{oc}, and particularly *FF* was obtained from the PSCs with surface-treated TiO₂ NR arrays, leading to a higher *PCE*. With the optimised perovskite capping layer atop the TiO₂ NR array, an encouraging *PCE* of 18.22% has been achieved with *J*_{sc} of 22.9 mA·cm⁻², which is the highest efficiency reported so far for 1D nanostructure-applied PSCs. Most importantly, the high performance and simple surface modification method of the TiO₂ NR-PSCs are encouraging to compete with the conventional mesoscopic PSCs with mesoporous TiO₂ photoanode [72]. Table 2 summarises the photovoltaic performance of the 1D TiO₂ nanostructured photoanode-employed PSCs.

TiO ₂ Type/Fabrication Method 1	Architecture ²	η (%)	$J_{\rm sc}$ (mA·cm ⁻²)	V _{oc} (V)	FF (%)	References
Vertical and Rutile NR arrays; HT	FTO/NR-MAPbI3/Spiro-MeOTAD/Au					
	0.56 μm thick NR	9.4	15.6	0.96	0.63	[60]
	0.92 μm thick NR	7.3	12.6	0.93	0.62	[69]
	1.58 μm thick NR	5.9	11.2	0.87	0.61	
Vertical NW arrays; ST	FTO/0.15–0.20 μm thick NW-MAPbI ₃ /Spiro-MeOTAD/Au	16.04	22.60	1.01	0.71	
	0.03–0.05 μm thick NP	11.61	21.83	0.90	0.59	[70]
	0.07–0.10 μm thick NS	13.09	21.26	0.95	0.65	
Layered Hyper-branched NR arrays; HT/ES	FTO/1 layered Hyperbranched NR-MAPbI ₃ /Spiro-MeOTAD/Au					
	0.60 μm thick NR	15.50	20.5	1.00	0.75	[54]
	0.85 µm thick NR	6.85	11.0	0.77	0.75	
	1.23 μm thick NR	5.88	9.5	0.75	0.74	
Vertical and Anatase NR arrays; HT/ALD	FTO/1.8 µm thick NR-MAPbI ₃ /Spiro-MeOTAD/Au					
	TiCl ₄ -treated NR arrays	7.98	16.16	0.85	0.58	[71]
	ALD-treated NR arrays (4.8 nm thick)	13.45	19.78	0.94	0.72	
Vertical NR arrays; ST/UV-O ₃	FTO/0.18 μm thick NR-MAPbI ₃ /Spiro-MeOTAD/Au					
	UV-O ₃ treatment	18.22	22.09	1.04	0.76	[72]
	No treatment	13.80	20.10	1.02	0.67	

Table 2. Summary of photovoltaic characteristics for PSCs using 1D TiO₂ nanostructured photoanodes.

¹ In type and fabrication method, "NRs" and "NWs" indicate "nanorods" and "nanowires", respectively. "HT", "ES", "ALD", "ST" indicate "hydrothermal method", "electrospinning method", "atomic layer deposition" and "solvothermal method"; ² In Architecture, "MAPbI₃" indicate "CH₃NH₃PbI₃ perovskite. "NP" and "NS" indicate "nanoparticle" and "nanosheet", respectively.

5. Conclusions

In this review article, the novel properties, synthetic methods, and applications of 1D TiO_2 nanostructures have been reviewed, with a focus on photoanodes for solar cell applications. Basically, the photovoltaic performance of solar cells is driven by the charge transport properties and light harvesting efficiency, which can be improved by optimising the photoanode materials and device architecture. Thus, it is important to note that the unique morphology of 1D TiO₂ nanostructures offers excellent charge transport properties such as charge separation, electron injection, and electron lifetime. Besides, 1D TiO₂ nanostructures lead to efficient light utilisation management by light scattering and channeling effects. In comparison to randomly oriented TiO₂ nanoparticulate systems, the vertically oriented 1D TiO₂ NTs and NWs present significantly improved charge transport properties and light harvesting efficiency in DSCs, followed by the improvement of overall photovoltaic performance of the devices. In particular, the hierarchical architecture incorporating 0D TiO₂ nanoparticles or 3D hollow TiO_2 microspheres to 1D TiO_2 nanostructures can further improve the performance with their large surface area as well as 1D structured light scattering layer and express electron channels. In PSCs, 1D TiO₂ nanostructures can also be a promising candidate for high efficiency solar cells. Especially, the much higher light absorption coefficient and extremely long charge diffusion lengths of the perovskites would be beneficial to remedy a drawback of 1D TiO_2 nanostructures, their relatively low light harvesting efficiency caused by the lower dye loading in DSCs. However, the 1D TiO₂-PSCs have not been extensively studied yet, compared to the case of 1D TiO₂-DSCs, indicating there is still much room to further develop 1D TiO₂-PSCs. Since the key issues of 1D TiO₂-PSCs are pore filling and optimising the device architecture using 1D TiO₂ nanostructured layers, various deposition methods, newly developed perovskites and HTMs, and advanced PSC architectures should be preferentially considered to achieve high performance of 1D TiO₂-PSCs. In addition, studies of 1D TiO₂-applied flexible and large scale devices may provide insights into the commercialisation of DSCs and PSCs in the near future.

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