The present invention relates to a photovoltaic cell comprising of first and second electrodes and a plurality of aligned semiconducting nanorods disposed between the electrodes, each nanorod being electrically connected to the first electrode and electrically insulated from the second electrode. A plurality of photoresponsive nanocrystals surround and are bound to the nanorods. A semiconductor polymer surrounds the nanorods and is bound to the nanocrystals and to at least the second electrode. The nanocrystals act as heterojunctions channeling a first charge carrier into the nanorods and a second charge carrier into the polymer.
HYBRID PHOTOVOLTAIC CELLS AND RELATED METHODS

The present invention relates to solar cells and their fabrication, and in particular to nanorod-nanocrystal-polymer hybrid solar cells.

To create useful electrical current from electromagnetic radiation, photovoltaic (PV) cells must absorb incident radiation such that an electron is promoted from the valence band to the conduction band (leaving a hole in the valence band), and must be able to separate the electron and hole and deliver these charge carriers to their respective electrodes before they recombine.

Many different strategies based on diverse materials have been employed, with varying degrees of success, to realize these basic behaviors with commercially satisfactory efficiency. Representative devices include crystalline inorganic solar cells (e.g., silicon, germanium, GaAs), nanocrystalline dye-sensitized solar cells, semiconductor-polymer solar cells, nanoparticle solar cells, and more recently, composite solar cells that incorporate and combine the aforementioned components from other strategies.

Silicon is by far the most commonly used material for fabricating inorganic photovoltaics. These cells rely on the ability of silicon to absorb light and, consequently, to generate an excited electron-hole pair that is then separated at a p-n junction. The electric field set up by the p-n junction facilitates this separation because of the way electrons and holes move through materials: electrons move to lower energy levels while holes move to higher energy levels.

Creation of p-n junctions generally involves high-temperature processing in inert atmospheres to form very pure, crystalline silicon wafers, which are inflexible and expensive. Because silicon is an indirect semiconductor, a relatively thick layer is typically needed to achieve a good level of absorption, which increases material costs further. Efficiencies for the most pure (and expensive) silicon photovoltaics are on the order of 20%; efficiencies for the cheaper amorphous silicon cells are approximately 5-10%.

Today's commercial PV systems can convert from 5% to 15% of sunlight energy into electricity. These systems are highly reliable and generally last 20 years or longer. The possibility of fabricating solar cells by less expensive, lower-temperature techniques is very
attractive. Accordingly, nanocrystalline dye-sensitized solar cells (DSSCs), semiconductor-polymer solar cells and nanoparticle solar cells have enjoyed widespread interest.

Semiconducting polymers can be used to make organic photovoltaics. The properties of these polymers can be tuned by functionalization of the constituent monomers. As such, a wide range of polymers with suitable bandgaps, absorption characteristics and physical properties is available. In order to achieve separation of the electron-hole pair, organic photovoltaics rely on donor-acceptor heterojunctions. In polymers, the excited-state electron and hole are bound together, and travel together, as a quasi-particle called an exciton. They remain together until they encounter a heterojunction, which separates them. Unfortunately, excitons are very short-lived and can only travel about 10 nm before recombining. Hence, any photon absorbed more than this diffusion length away from a heterojunction will be wasted. Charge mobilities for polymers are typically low (0.5-0.1 cm² V⁻¹ s⁻¹) compared to silicon, which is much higher (1500 cm² V⁻¹ s⁻¹). Current state-of-the-art polymer photovoltaic cells have efficiencies of 1-2%. Although such efficiencies are low, these materials hold promise for low-cost, flexible solar cells.

Inorganic nanoparticles (or nanocrystals) have been used to prepare colloidal, thin-film PV cells that show some of the advantages of polymer photovoltaics while maintaining many of the advantages of inorganic photovoltaics. For example, such cells can contain a bi-layer structure comprising a layer of donor and a layer of acceptor nanoparticles, wherein the two layers exhibit little intermixing, and both contribute to the measured photocurrent. The strong photoconductive effect exhibited by these devices suggests that these materials have a large number of trapped carriers and are better described by a donor-acceptor molecular model than by a p-n band model. Increased bandgap energy compared to that of the bulk semiconductors minimizes the number of carriers available, and spatial separation of the donor and acceptor particles in different phases traps the excitons so that they must split at the donor-acceptor heterojunction. There is no band-bending, so splitting of the exciton is more difficult.

It should be stressed that simply blending the donor and acceptor nanoparticles together will not create a film that produces a photovoltage. The lack of selectivity at the electrode towards one particle or another means that the electrodes can make contact with both the donor and acceptor species. These species may take the form of nanorods rather than nanospheres because nanorods with high aspect ratios help to disperse the carriers. Quick transfer of the
exciton along the length of the nanorods improves the chance of splitting the exciton at the donor-acceptor heterojunction.

Solution processing of, for example, CdSe rods can achieve a size distribution of 5% in diameter and 10% in length with an aspect ratio of 20 and a length of 100 nm. The substantial control available through solution processing allows for optimization of the cell by variation of nanorod length and bandgap energy.

The combination of nanomaterials and polymer films has been shown to give good power conversion efficiencies while affording low-temperature solution processes for fabrication. In one approach, nanomaterials are used to conduct charges while the polymer is used as the absorbing material, or alternatively, the nanomaterial serves as a chromophore, i.e., the light absorber, and the semiconductor polymer is employed as a hole conductor. In the former case, a wide-bandgap semiconductor (e.g., TiO₂) receives the excited electron from the conduction band of the chromophoric polymer semiconductor; and in the latter case, light-absorbing semiconductor nanocrystals absorb photons and transfer the resulting negative charge to the transparent primary electrode, while the semiconducting polymer transfers the holes to the counter electrode. In both types of cell, a heterojunction between the nanocrystal and the polymer separates the exciton created in the nanocrystal or polymer. The electron is transferred to the conduction band of the nanocrystal and the hole stays in the valence band of the polymer, or the electron stays in the conduction band of the nanocrystal, and the hole is transferred to the valence band of the polymer.

The active layer in a polymer-nanocrystal cell has two components: a light absorber and a nanoparticulate electron carrier. Typically, the light absorber is a p-type polymeric conductor, e.g., poly(phenylene vinylene) or poly(3-hexylthiophene), and the nanoparticulate electron carrier is a wide-bandgap semiconductor such as ZnO or TiO₂. In this configuration, the polymer serves to absorb light, to transfer electrons to the electron acceptor/carrier, and to carry holes to the primary electrode. The electron acceptor accepts electrons and transfers the electrons to the metal back contact.

The morphology of the phase separation is crucial. For example, a bi-layer structure in which each layer has only one component results in a cell with poor performance. The reason is that the lifetime of the excited state of the light-absorbing polymer is generally shorter than the
transfer rate of the exciton to the interface, and, consequently, the majority of the excitons formed in the bulk of the polymer never reach the interface separating electrons and holes, resulting in loss of photocurrent. Morphologies in which a bulk heterojunction is formed tend to show greater efficiencies. If the absorber and electron acceptor are in intimate contact throughout the entire active layer, the shorter exciton path length will result in increased electron transfer and higher efficiencies. The best efficiencies obtained from cells of this configuration are around 2%.

This technology shows promise, but there are obstacles to overcome. One problem is incomplete absorption of the incident radiation. The polymer — which absorbs light very strongly and is referred to as a polymeric dye — has a large extinction coefficient (\(>100,000 \text{M}^{-1} \text{cm}^{-1}\)), but due to low exciton migration rates, the films must generally be thinner than 100 nm, which contributes significantly to incomplete absorption. This effect can be combated by means of an interdigitated array structure of donor and acceptor species.

A problem associated with the light-absorbing polymer strategy is underutilization of available solar energy due to the narrow absorption bandwidth of typical polymers. Approximately 40% of the light (from about 600 nm out into the near IR) can be wasted. An alternative configuration is to utilize nanocrystals as light absorbers and electron carriers, and employ the polymer as a light absorber and a hole carrier. CdSe nanorod and tetrapod/polymer systems have demonstrated power-conversion efficiencies of up to 1.7%. These systems have the advantage that the absorption of the nanocrystal can be tuned via the size of the nanocrystal, and systems that absorb essentially all of the incoming radiation can therefore be fabricated.

Unfortunately, it is difficult to disperse inorganic nanocrystals into a solution of monomers. The two phases tend to agglomerate and minimize the electrical contact essential to form the heterojunction which enables charge separation. Dispersion of nanocrystals in polymer phases is an area of great interest.

Typically, the strategy employed for dispersing the nanocrystals is to functionalize the nanocrystal with a capping agent that has an organic tail, which enhances solubility in the solvent in which the polymerization is carried out. Capping agents for this purpose typically have a head-group with a strong affinity for the nanocrystal; amine, carboxylate, phosphine, thiol, phosphine oxide and phosphonic acid, for example, all bind strongly. The organic tail of the
capping agent should be compatible with solvents in which the polymer is soluble. Long hydrocarbon chains typically provide high solubility but are non-conducting; accordingly, it is necessary to balance optimum solubility against conductivity.

The most popular polymers used for composite studies are PDFC, P3Ht and MEH-PPV (where PDFC refers to \{9,9-dihexylfluorenyl-2,7-diyl)-alt-co-(9-ethyl-3,6-carbazole)\}, P3Ht refers to poly(3-hexylthiophene), and MEH-PPV refers to poly(2-methoxy-5-(2'-ethylhexyloxy)-1,4-phenylene vinylene)). Each of these polymers has sites for functionalization, allowing the manipulation of the valence/conduction band energies to achieve optimal conditions for charge transfer to and from the nanocrystals. It has been suggested that the capping agent may also serve as the organic acceptor phase; for example, P3HT functionalized with phosphonic acid groups has been shown to isolate CdSe nanocrystals.

DSSCs incorporate a substrate which has been coated with a transparent conducting oxide (which serves as the primary electrode). The counter electrode may also be coated with a transparent conducting oxide, but may also be a non-corrosive metal, such as titanium coated with a very thin layer of platinum. A porous layer of a wide-bandgap semiconductor (such as TiO$_2$) is deposited on the conductive surface of the primary electrode. This porous layer is then coated with a dye having a strong absorption in the visible region of the spectrum. To be optimally effective, the dye concentration should be limited to a monolayer of dye molecules. Because of this, a huge surface area is necessary to accommodate enough dye to absorb all of the incoming light. Therefore, nanocrystals (e.g., TiO$_2$) are used to make the highly porous films. Electrolyte containing a redox couple (typically $\Gamma/\Gamma^-$) is absorbed into the titania layer. To complete the cell, the substrate bearing the primary electrode and the sensitized titania layer is brought into face-to-face contact with the counter electrode.

Typical dyes are inorganic-ruthenium-based, although organic dyes are receiving increased interest. The dye absorbs visible light, and the excited state injects an electron into the TiO$_2$ conduction band. Before back electron transfer can occur, the oxidized dye is reduced by a redox active species in solution (typically $\text{FV}_{13}^-$), regenerating the dye. The oxidized redox active species diffuses to the counter electrode, where it is reduced, finishing the cycle and completing the circuit. Work can be done by passing the injected electron through an external load before allowing it to reduce the oxidized redox active species at the counter electrode.
Inexpensive DSSC devices, which exhibit up to 10% energy conversion efficiency, can be fabricated. There are many issues to be addressed with this technology to improve performance and stability, including replacing the best performing liquid electrolytes with solid-state or higher-boiling electrolytes; improving spectral overlap; using a redox mediator with a lower redox potential; and lowering recombination losses due to poor electron conduction through the nanoparticle TiO₂ layer.

Hybrid cells combine dye-sensitized titania, coated and sintered onto a transparent semiconducting oxide, with a p-type polymer that carries electrons to the oxidized dye. Since just one polymer replaces the multi-component electrolyte, these cells can be fabricated conveniently and reproducibly. Ruthenium dye-sensitized, nanorod-based DSSCs tend to exhibit low efficiency, however, because the lower surface area does not accommodate enough dye to absorb all of the incident light. The most efficient dyes found so far only have extinction coefficients on the order of -20,000 M⁻¹cm⁻¹, and therefore a large surface area is needed to bind enough dye to get maximal absorbance.

An object of the present invention is to obviate or mitigate one or more of the disadvantages associated with existing photovoltaic cells and/or the fabrication of components of photovoltaic cells.

Aspects of the present invention provide a photovoltaic (PV) cell comprising a semiconducting nanorod-nanocrystal-polymer hybrid layer, as well as methods for fabricating the same. In PV cells according to this invention, the nanocrystals serve both as the light absorbing material and as the heterojunctions at which the excited electron-hole pairs (i.e., excitons) split. The nanorods function as electron carriers and are electrically connected to the anode of the cell, and the polymer acts as the hole carrier and is electrically connected to the cathode of the cell.

One of the advantages of the invention lies in the use of small particles, the nanocrystals, as both light absorber and heterojunction. The resulting spatio-temporal proximity of excitor generation and splitting entails a significant reduction in recombination losses, compared, for example, with those of conventional polymer PV cells, and consequently in higher conversion efficiencies of photons into electricity. Embodiments of the invention offer the additional advantages of mechanical flexibility and low cost manufacturing processes.
Accordingly, in a first aspect, the invention provides a photovoltaic cell containing two electrodes and, in between these electrodes, a plurality of aligned semiconductor nanorods surrounded by and bound to a plurality of photoresponsive nanocrystals, and a semiconductor polymer surrounding the nanorods and bound to the nanocrystals. The nanocrystals act as heterojunctions channeling electrons into the nanorods and holes into the polymer, or vice versa. The nanorods are electrically connected to the first electrode, and electrically insulated from the second electrode by a thin layer of polymer bound to the second electrode. In various embodiments, the polymer is a hole-transfer polymer, and consequently, the nanocrystals channel holes into the polymer and electrons into the nanorods. In various embodiments, the nanocrystals are bound to the nanorods by a bifunctional capping agent, which can, for example, be mercaptoacetic acid. For example, the nanorods may be grown on the first electrode, and the other electrode can later be deposited on the nanorod-nanocrystal-polymer layer in a manner ensuring insulation of the nanorods from the second electrode.

Advantageous nanorods have aspect ratios (i.e., ratios of the longest dimension to the shortest dimension of the particle) of at least 3, and their shortest dimension is not greater than 100 nm. Preferred nanorods are single-crystalline. Suitable nanorod materials according to the invention include, but are not limited to, wide bandgap semiconductors such as, for example, ZnO, SnO, and TiO₂, whereby ZnO is the preferred material.

Suitable nanocrystals according to the invention include semiconducting, monocry stalline or polycrystalline nanoparticles of diameter not greater than 20 nm, which may (but need not) be generally spherical in shape. Suitable nanocrystal materials include, but are not limited to CuInSe₂, CuInS₂, CuInₙGaₓSe₂ (where 0 ≤ x ≤ 1), GaAs, InAs, InP, PbS, PbSe, PbTe, GaSb, InSb, CdTe and CdSe. Nanocrystals with extinction coefficients of at least 100,000 M⁻¹ cm⁻¹ are preferred. In various embodiments, the largest spatial dimension of the nanocrystals is no greater than the average diffusion distance of the excitons created in the nanocrystal upon absorption of light.

Suitable polymer materials include, but are not limited to, poly(3-hexylthiophene), polyphenylenevinylene (PPV) and its derivatives, and polyfluorene (PFO) and its derivatives. In various embodiments, the polymer is bound to the nanocrystals but not to the nanorods.
In a second aspect, the invention provides a method of fabricating a semiconductor structure with heterojunctions; the structure can be used in a photovoltaic cell. Embodiments of the method involve providing a plurality of nanorods and a plurality of photoresponsive nanocrystals capped with a first capping agent; exposing the nanorods or the nanocrystals to a second, bifunctional capping agent; then combining the nanocrystals with the nanorods so that the nanocrystals bind to the nanorods via the bifunctional capping agent; combining the bound nanorods and nanocrystals with a functionalized monomer which has a binding group with (i) stronger affinity for the nanocrystals than the first capping agent and (ii) weaker affinity for the nanorods than the bifunctional capping agent, so that the monomer preferentially displaces the first capping agent and binds to the nanocrystals; and polymerizing the monomer. The bifunctional capping agent can first bind to the nanorods, and then bind to the nanocrystals, replacing some of the first capping agent. Alternatively, the bifunctional capping agent can first bind to the nanocrystals (replacing some of the first capping agent), and then bind with its free ends to the nanorods. In various embodiments, the first capping agent contains a thiol, selenol, amine, phosphine, phosphine oxide, and/or aromatic heterocycle functionality. A non-limiting example of a suitable capping agent is octanethiol.

The foregoing discussion will be understood more readily from the following detailed description of the invention when taken in conjunction with the accompanying drawings.

FIG. 1A schematically depicts an embodiment of a nanorod-nanocrystal-polymer hybrid solar cell according to the invention.

FIG. 1B is an enlarged schematic view of the three major components of the hybrid semiconductor layer of FIG. 1A, and of their interconnections.

FIG. 2A is a flow diagram detailing a method of fabricating the structure depicted in FIG. 1A according to one embodiment.

FIG. 2B is a flow diagram detailing a method of fabricating the structure depicted in FIG. 1A according to an alternative embodiment.

FIG. 3 illustrates at a microscopic level some of the steps of the method shown in FIG. 2 and the resulting products.
1. Nanorod-nanocrystal-polymer hybrid structure

In polymer-based photovoltaics, excitons travel on average of the order of 10 nm before recombining; accordingly, there is a need to separate the excitons, i.e., to have them encounter a heterojunction as soon as possible. This need is met in embodiments of the present invention, in which nanocrystals (quantum dots) serve as a bridge between a hole-transfer polymer and a wide-bandgap semiconductor electron acceptor, thus constituting the heterojunction, and serve simultaneously as the light absorber, i.e., the place where the excitons are created. The diameter of a nanocrystal according to the invention is approximately equal to, or smaller than, the diffusion distance of an exciton. As a result, an exciton generated in the nanocrystal will generally encounter the interface of the nanocrystal with the electron acceptor or the hole-transfer polymer within its average diffusion distance, regardless of the direction in which it migrates. Consequently, the exciton splits very efficiently, and recombination within the nanocrystal occurs infrequently. The electron enters into the wide-bandgap semiconductor, and the hole enters into the polymer.

The structure of a PV cell 100 according to the invention is illustrated in FIG. IA. In between two electrodes, an anode 101 and a cathode 103, a plurality of aligned wide-bandgap semiconductor nanorods 106, which constitute the electron acceptor, is arranged. As shown in the detail of FIG. IA, the nanorods 106 are each surrounded by photosensitive nanocrystals 109. The sensitized nanorods, in turn, are surrounded by the hole-transfer polymer 112, which fills the remaining space between the electrodes 101, 103. The polymer 112 also forms a thin layer underneath the cathode 103, which electrically isolates the cathode 103 from the sensitized nanorods 106.

FIG. IB shows how these three components are interconnected in preferred embodiments of the invention. The nanocrystals 109 are bound to the nanorods 106 by means of a bifunctional binding molecule 115. In various embodiments, the bifunctional capping agent 115 has thiol and carboxylate moieties. The thiol groups bind preferentially to the nanocrystals 109, and the carboxylate groups bind preferentially to the (metal oxide) nanorods 106. The intervening chain should be short enough so that charge transfer from nanocrystal 109 to nanorod 106 is not impeded. A representative bifunctional capping agent 115 is mercaptoacetic acid. The hole-
transfer polymer 112 is directly bound to the nanocrystals 109, but preferably not to the nanorods 106.

A representative, non-limiting example of a system of nanorods, bifunctional molecules, nanocrystals, and polymers comprises ZnO nanorods capped with mercaptoacetic acid, CuInSe$_2$ quantum dots, and poly(3-hexylthiophene).

1.1 Nanocrystals 109

The semiconductor material used for nanocrystals in a particular application depends on the suitability of valence and conduction band energy levels. The conduction band should be of sufficient energy to be able to inject electrons efficiently into the nanorods, while the valence band should be of sufficiently low energy to inject holes into the polymer valence band. The latter constraint is generally straightforward to satisfy, as suitable polymers having a higher-energy valence band than the nanocrystal can readily be identified. Subject to the above constraints, the bandgap of the nanocrystal should be small enough to allow for a large portion of the solar spectrum to be absorbed. Suitable nanocrystal materials include materials based on copper-indium-diselenide and variants thereof, for example, CuInS$_2$, CuInSe$_2$, or CuIn$_x$Ga$_{1-x}$Se$_2$ (wherein $0 \leq x \leq 1$), as well as CdSe, GaAs, InAs, and InP.

Nanocrystals can be synthesized using techniques described, for example, in U.S. Patent No. 6,379,635 and co-pending U.S. Patent Application Serial Nos. 11/579,050 and 11/588,880.

Methods for preparing CdSe, InS, GaS, ZnS, CdS, ZnAs, CdAs and further related nanocrystalline materials are described in Sections 1. to 10. of the experimental section of US Patent No. 6,379,635. This describes how nanocrystal precursors can be provided in a single precursor complex (such as, but not limited to an alkyl carbonate complex of the required metal ions), which is then thermally decomposed under suitable conditions (e.g. temperatures of around 200 to 300° C) to yield the final desired nanocrystalline material.

By way of example, nanocrystalline CdSe can be formed as follows, 1.2 MeCddsc (0.5 mmol) was placed in 10 ml of TOP (98%, Aldrich) and the mixture formed was filtered after which was injected in 30 g of TOPO at 200° C. The temperature of the solution was then raised to 250° C and heated for half an hour. The deep red solution that formed was allowed to cool
down to 75°C after which a large excess of dry CH₃OH (BDH) was added. A flocculate precipitate formed and was isolated by centrifugation and redispersed in toluene, any insoluble material was then discarded. The toluene was pumped off under vacuum (10⁻² Torr) to give a deep red material which was washed with CH₃OH. The solid was redispersed in toluene to give solutions with a Port wine red colour which remained optically clear for weeks. Size selective precipitation was performed by adding CH₃OH to this solution until turbidity was observed followed by centrifugation the solid. This procedure was successively applied to the supernatant solutions obtained during the fractionation process until no optical absorption was detected.

Similar methods can be employed to produce III-VI nanocrystalline materials (e.g. InS, GaS) by replacing MeCddsc with a complex of the general formula, (AlK₁)₂M¹₁E₂CN(AlK₂)₂, where AlK₁ and AlK₂ are individually alkyl groups, such as Me,Et,Np etc, each AlK₁ and AlK₂ being the same or different, M¹ is a group III metal ion such as In, Ga etc, and E is a group VI ion, such as S,Se etc; and to produce II-VI or H-V nanocrystalline materials (e.g. ZnS, CdS, ZnAs and CdAs) by replacing MeCddsc with a complex of the general formula [M²[E₂CN(AlK₃)₂]₂], where each AlK₃ is an alkyl group, such as Me,Et,Pr etc, each AlK₃ being the same or different, M² is a group II metal ion, such as Zn, Cd etc, and E is a group V or VI ion, such as As or S,Se etc.

Methods for preparing large amounts of nanocrystalline materials are described in US Patent Application Serial Nos. 11/579,050 and 11/588,880. These applications describe for the first time how molecular clusters can be used to seed controlled growth of any desirable nanocrystalline material, which can then be provided with one or more outer shells of a different material to the material forming the nanocrystal core grown on the molecular cluster seed.

By way of example, US Patent Application Serial No. 11/579,050 describes in Examples 1 to 9, the preparation of the clusters [HNEt₃MCd₄(SPh)I₀] and [HNEt₃]₄[CdIOSe₄(SPh)i₆] which can then be used to produce CdSe nanocrystals by controlled addition of Cd ions and Se ions (from, e.g. TOPSe and Cd(CH₃CO₂)₂) over a suitable time period, in combination with suitably controlled increases in the temperature of the reaction mixture. For example, Example 1 describes the preparation of CdSe nanoparticles from [Et₂NH]₄[CdIOSe₄(SPh)i₆] / TOPSe / Cd(CH₃CO₂)₂ in HDA. HDA (300g) was placed in a three-neck flask and dried/degassed by heating to 120°C under a dynamic vacuum for 1 hour. The solution was then cooled to 70°C. To this was added 1.Og of [Et₃NH]₄[CdIOSe₄(SPh)i₆] (0.31 lmmol), TOPSe (20ml, 40.00mmol)
[previously prepared from dissolving selenium powder in TOP] and Cd(CH\textsubscript{3}CO\textsubscript{2})\textsubscript{2} (10.66g 40.00mmol) the temperature of reaction mixture was gradually increased from 70°C to 180°C over an 8 hour period. The progressive formation/growth of the nanoparticles was monitored by their emission wavelength by taking aliquots from the reaction mixture and measuring their UV-vis and PL spectra. The reaction was stopped when the emission spectra had reached 572nm by cooling the reaction to 60°C followed by addition of 200ml of dry "warm" ethanol which gave a precipitation of nanoparticles. The resulting CdSe were dried before re-dissolving in toluene filtering through Celite followed by re-precipitating from warm ethanol to remove any excess HDA and Cd(CH\textsubscript{3}CO\textsubscript{2})\textsubscript{2}. This produced 9.26g of HDA capped CdSe nanoparticles.

By way of further example, US Patent Application Serial No. 11/588,880 describes the preparation of CdSe nanoparticles from [HNEt\textsubscript{3}]\textsubscript{4}Cd\textsubscript{4}Se\textsubscript{4}(SPh)\textsubscript{16}/TOPSe/Me\textsubscript{2}CdTOP in HDA by dropwise addition of Me\textsubscript{2}CdTOP, and the preparation of ZnS nanoparticles from [Et\textsubscript{3}NH]\textsubscript{4}ZnO\textsubscript{4}(SPh)\textsubscript{6} seeds in HDA by dropwise addition of Et\textsubscript{2}Zn and S-octylamine. Nanoparticles formed in this way can then be provided with one or more shell layers to provide CdSe/ZnS-HDA capped nanoparticles, and ZnSe/ZnS nanoparticles. For example ZnS nanoparticles can be formed from [Et\textsubscript{3}NH]\textsubscript{4}ZnO\textsubscript{4}(SPh)\textsubscript{6} seeds in HDA by dropwise addition of Et\textsubscript{2}Zn and S-octylamine as follows. A 200 g portion of hexadecylamine (HDA) was placed in a three-neck round-bottomed flask and dried and degassed by heating to 120 °C under a dynamic vacuum for > 1 hour. The solution was cooled to 60 °C, the reaction flask was filled with nitrogen and the following reagents were loaded into the flask using standard airless techniques: 0.6 g [HNEt\textsubscript{3}]\textsubscript{4}ZnO\textsubscript{4}(SPh)\textsubscript{6} (0.2 mmol), 4 mL of a 0.5 M solution of Et\textsubscript{2}Zn in trioctylphosphine (2 mmol), and 4 mL of a 0.5 M solution of elemental sulphur in octylamine (2 mmol). The temperature was increased to 120 °C and allowed to stir for 2 hours. At this point a programmed temperature ramp from 120 °C to 210 °C at a rate of ~0.2°C/min was initiated.

Simultaneously, 8 mL of 0.5 M Et\textsubscript{2}Zn and 8 mL of 0.5 M S-octylamine were added dropwise at a rate of -0.05 mL/min. The reaction was stopped when the PL emission maximum had reached the required emission (\(\lambda_{\text{max}} = 391\) nm, FWHM = 95 nm) by cooling to 60°C followed by addition to 300 ml of dry ethanol or acetone to precipitate the particles from solution. This precipitate was isolated by filtration. The resulting ZnS particles were further purified (to remove excess HDA, sulphur and zinc) by re-dissolving in toluene, filtering the solution through Celite and re-precipitating from warm ethanol (product yield: 0.9g). Capping or shelling of CdSe
nanoparticles can then be carried out, for example, as follows HDA (800g) was placed in a three
neck round-bottom flask, dried and degassed by heating to 120°C under a dynamic vacuum for > 1 hour. The solution was then cooled to 60°C, to this was added 9.23g of CdSe nanoparticles that have a PL maximum emission of 585nm. The HDA was then heated to 220°C. To this was added by alternate dropwise addition a total of 20ml of 0.5M Me₂Zn*TOP and 0.5M, 20ml of sulfur dissolved in octylamine. Three alternate additions of 3.5, 5.5 and 11.0ml of each were made, whereby initially 3.5ml of sulphur was added dropwise until the intensity of the PL maximum was near zero. Then 3.5ml of Me₂Zn*TOP was added dropwise until the intensity of the PL maximum had reached a maximum. This cycle was repeated with the PL maximum reaching a higher intensity with each cycle. On the last cycle, additional precursor was added once the PL maximum intensity been reached until it was between 5 - 10% below the maximum intensity, and the reaction was allowed to anneal at 150°C for 1 hour. The reaction mixture was then allowed to cool to 60°C whereupon 300ml of dry "warm" ethanol was added which resulted in the precipitation of particles. The resulting CdSe-ZnS particles were dried before re-dissolving in toluene and filtering through Celite followed by re-precipitation from warm ethanol to remove any excess HDA. This produced 12.08g of HDA capped CdSe-ZnS core-shell nanoparticles. Elemental analysis C = 20.27, H -3.37, N = 1.25, Cd = 40.11, Zn = 4.43 %; Max PL 590nm, FWHM 36nm.

A method for producing CIGS nanocrystals of any desirable stoichiometry employing a selenol compound is disclosed in U.S. Provisional Application Serial No. 60/991,510. Embodiments of the method involve dispersing at least a first portion of a nanocrystal precursor composition (comprising sources of at least one of Al, Ga, and/or In, and at least one of Cu, Ag, Zn, and/or Cd) in a solvent (e.g., a long-chain hydrocarbon solvent); heating the solvent to a first temperature for an appropriate length of time; adding a selenol compound to the solvent and heating the solvent; adding a second portion of the nanocrystal precursor composition to the reaction mixture; heating the mixture to a second temperature higher than the first temperature over an appropriate length of time; and maintaining the temperature for up to 10 hours. Once the particles have been formed, the surface atoms of the particles will typically be coordinated to a capping agent, which can comprise the selenol compound employed in the method. If a volatile selenol compound is used, this capping agent can be driven off with heating to yield 'naked'
nanocrystals amenable to capping with other coordinating ligands and further processing. Examples 1 and 2 provide further details regarding the implementation of this method:

**Example 1:** Cu(I) acetate (1 mmol) and In(III) acetate (1 mmol) are added to a clean and dry RB-flask. Octadecene ODE (5 mL) is added the reaction mixture heated at 100 °C under vacuum for 30 mins. The flask is back-filled with nitrogen and the temperature raised to 140 °C. 1-octane selenol is injected and the temperature falls to 120 °C. The resulting orange suspension is heated with stirring and a transparent orange/red solution is obtained when the temperature has reached 140°C. This temperature is maintained for 30 minutes, then 1M tri-octyl-phoshine selenide TOPSe (2mL, 2 mmol) is added dropwise and the solution heated at 160°C. The PL is monitored until it reaches the desired wavelength, after which it is cooled and the resulting oil washed with methanol/acetone (2:1) 4-5 times and finally isolated by precipitation with acetone.

**Example 2 (Large Scale Production):** A stock solution of TOPSe was prepared by dissolving Se powder (10.9, 138 mmol) in TOP (60 mL) under nitrogen. To dry, degassed ODE was added Cu(I) acetate (7.89 g, 64.4 mmol) and In(III) acetate (20.0 g, 68.5 mmol). The reaction vessel was evacuated and heated at 140 °C for 10 min, backfilled with N₂ and cooled to room temp. 1-Octane selenol (200 mL) was added to produce a bright orange suspension. The temperature of the flask was raised to 140 °C and acetic acid distilled from the reaction at 120 °C. On reaching 140 °C the TOPSe solution was added dropwise over the course of 1 hour. After 3 hours the temperature was raised to 160 °C. The progress of the reaction was monitored by taking aliquots from the reaction periodically and measuring the UV/Visible and photoluminescence spectra. After 7 hours the reaction was cooled to room temperature and the resulting black oil washed with methanol. Methanol washing was continued until it was possible to precipitate a fine black material from the oil by addition of acetone. The black precipitate was isolated by centrifugation, washed with acetone and dried under vacuum. Yield: 31.97 g.

For the purpose of optimizing the composition, size, and charge of the nanocrystals, they can be characterized by conventional techniques, including, but not limited to, XRD, UV/Vis/Near-IR spectrometry, SEM, TEM, EDAX, photoluminescence spectrometry, and elemental analysis.

Some embodiments of the invention utilize nanocrystals with extinction coefficients of at least 100,000 M⁻¹·cm⁻¹. At such high absorptivities, fewer nanocrystals are needed to achieve the
same overall absorption. Consequently, embodiments of this invention based on these nanocrystals can benefit from increased absorption without incurring losses in efficiency due to enhanced recombination.

1.2 Nanorods

Nanorods can be produced by direct chemical synthesis, utilizing a suitable combination of ligands such as trioctylphosphine oxide (TOPO) and various phosphonic acids, e.g., octadecylphosphonic acid, for shape control. Moreover, different types of metal oxides can be grown in ordered nanorod arrays, using techniques such as, for example, electrochemical etching of metal foil, or substrate seeding followed by nanorod growth, in a chemical bath, in a direction perpendicular to the substrate. See, e.g., D. C. Olson et al, J. Phys. Chem. C, 2007, 111, 16640-16645; and J. Yang et al, Crystal Growth & Design, 2007, 12/2562.

In preferred embodiments of the invention, the nanorods have high aspect ratios exceeding 3, and are up to 200 nm long. A preferred nanorod material is ZnO. Other materials that might be suitable include SnO, TiO₂, and other metal oxides.

As mentioned previously, the small size of the nanocrystals greatly reduces recombination within the particle. In order to further reduce recombination losses, preferred embodiments of the invention utilize single-crystal nanorods. While in nanoporous particle-based films, such as those employed in DSSC cells, electrons percolate slowly through the film, enabling recombination with the electrolyte to take place, electron transfer through single-crystal nanorods is very fast, which limits the recombination of electrons from the nanorods with holes in the nanocrystals or the polymer.

In preferred embodiments and as discussed in greater detail below, the nanorods are coated with a layer of a bifunctional capping agent, which binds the quantum dots closely to the nanorods, thereby preventing the semiconductor polymer from coming into the proximity of the nanorod, which diminishes nanorod-polymer recombination losses even further.

1.3 Polymer
Polymer 112 should have a valence band energy that allows holes to efficiently transfer from the nanocrystal valence band to the polymer valence band. Suitable polymers include poly(3-hexylthiophene), polyphenylenevinylene (PPV) and its derivatives, and polyfluorene (PFO) and its derivatives. These polymers are efficient hole-transfer polymers due to the high hole mobility in organic materials.

2. Method for fabricating a nanorod-nanoparticle-polymer hybrid structure

Hybrid semiconductor structures according to the invention can be fabricated using low-cost deposition technologies, such as printing, dip coating, or chemical bath deposition. An important consideration regarding fabrication is control over where the various pieces bind together. For example, binding of the polymer to the nanorod would most likely result in substantial losses in efficiency due to recombination. In preferred embodiments, the nanocrystals are bound to both the nanorods and to the semiconducting polymer to promote optimal performance as a heterojunction, and the polymer is not directly bound to the nanorods. This structure can be achieved with suitable capping agents in appropriate processing steps.

FIGS. 2A and 2B illustrate representative process sequences 200A and 200B implementing embodiments of the present invention. Some steps of these sequences, and the structures they result in, are further illustrated in FIG. 3 at a microscopic level. In a first step 202, nanorods are grown on an anodic substrate, e.g., by printing seeds on the substrate and then growing the nanorods perpendicularly to the substrate via a chemical bath. In this structure, the nanorods are inherently in electrical contact with the substrate. In subsequent steps, the nanocrystals and monomers are introduced to the resulting film of aligned nanorods.

In step 204, nanocrystals capped with a (first) capping agent which contains functionalities that bind weakly to the nanocrystals are provided. Suitable functionalities include thiol, selenol, amine, phosphine, phosphine oxide, and aromatic heterocycles. Typically, the nanocrystals are dissolved in a non-polar organic solvent. The capping agent serves to control binding of the nanocrystals to the nanorods and the polymer; the bond is reversible and the capping agent can later be exchanged for other ligands. Examples of capping agents suitable for use with CuInSe₂ nanocrystals are octanethiol or pyridine.
In steps 206, 208, the nanorods are coated by the nanocrystals, whereby the bond between nanorods and nanocrystals is established via the bifunctional capping agent 115 (e.g., mercaptoacetic acid), which has strong binding groups for both the nanorods and the nanocrystals. This can be accomplished in different ways. In some embodiments, as illustrated in FIG. 2A and FIG. 3, the nanorods are capped with the bifunctional capping agent (step 206A), for example, by dipping the substrate with the nanorods into a solution of the bifunctional capping agent. For example, the capping agent may be bound to the nanorods via a carboxylate functionality. The capped nanocrystals 302 are then introduced to the film of capped nanorods 300 (step 208A), for example, by dipping the rinsed substrate with nanorods 300 into the nanocrystal solution(s). At this stage, a fraction of the weak capping agent of the nanocrystals is replaced by the stronger binding groups of the bifunctional capping agent, e.g., the thiol functionality of mercaptoacetic acid, which results in nanocrystal-sensitized nanorods 304.

In alternative embodiments, as illustrated in FIG. 2B, a solution of the capped nanocrystals in a non-polar organic solvent is added to a solution of the bifunctional capping agent in a polar organic solvent which is not miscible with the non-polar solvent, and the solution is shaken to ensure good mixing (step 206B). The nanocrystals undergo ligand exchange and transfer from a non-polar organic phase to a polar organic phase. Subsequently, the substrate with the aligned nanorods on the surface is dipped into the nanocrystal solution or otherwise exposed to the nanocrystals (step 208B), whereby the nanorods bind the nanocrystals via a carboxylic acid functionality of the capping agent. These embodiment likewise result in nanocrystal-sensitized nanorods 304.

The monomers are functionalized (step 210) with a binding group that has a stronger affinity for the nanocrystals than the (first) nanocrystal capping agent, but a weaker affinity for the nanorods than the bifunctional capping agent. Moreover, the affinity of the binding group at the monomer for the nanocrystal is preferably weaker than the affinity of the bifunctional capping agent for the nanocrystal. The monomer functionality should not interfere with the polymerization reaction. Binding groups with suitable differential binding affinities are straightforwardly identified by those of skill in the art without undue experimentation based on the identities of the capping agents and their substituents (e.g., whether they are unidentate or multidentate, or on the presence of electron withdrawing groups, etc.) and the size of the nanocrystal. The functionalized monomers are then combined with the nanocrystal-sensitized
nanorods (step 212), where they bind to the nanocrystals (but not the nanorods), preferentially replacing the weak capping agent on the nanocrystal, but leaving the nanorod-nanocrystal bond intact, resulting in structure 306. A subsequent polymerization step 214 results in the nanorod-nanocrystal-polymer semiconductor structure 308.

Finally, a metal cathode (e.g., Al) can be deposited on the structure (step 216), for example, by sputtering or metal evaporation, so that the nanorods form an array of aligned rods deposited between two opposing electrodes. The polymer layer below the cathode should be sufficiently thick to electrically isolate the cathode from the nanorods.

Although the present invention has been described with reference to specific details, it is not intended that such details should be regarded as limitations upon the scope of the invention, except as and to the extent that they are included in the accompanying claims.
CLAIMS

1. A photovoltaic cell comprising:
   a. first and second electrodes;
   b. a plurality of aligned semiconducting nanorods disposed between the electrodes, each
      nanorod being electrically connected to the first electrode and electrically insulated from the
      second;
   c. a plurality of photoresponsive nanocrystals surrounding and bound to the nanorods;
   and
   d. a semiconductor polymer surrounding the nanorods and bound to the nanocrystals and
      to at least the second electrode, whereby the nanocrystals act as heterojunctions channeling a
      first charge carrier into the nanorods and a second charge carrier into the polymer.

2. The cell of claim 1 wherein (i) the polymer is a hole-transfer polymer, (ii) the first charge
   carrier is electrons, and (iii) the second charge carrier is holes.

3. The cell of claim 1 or 2 wherein the polymer is poly(3-hexylthiophene), polyphenylenevinylene
   or a derivative thereof, or polyfluorene or a derivative thereof.

4. The cell of claim 1, 2 or 3 wherein the nanorods are wide-bandgap semiconductors.

5. The cell of claim 1, 2, 3 or 4 wherein the nanorods comprise at least one of ZnO, SnO,
   and/or TiO₂.

6. The cell of any preceding claim wherein the nanorods are single-crystal nanorods.

7. The cell of any preceding claim wherein the nanorods have an aspect ratio of at least 3.

8. The cell of any preceding claim wherein the nanorods are bound to the nanocrystal by a
   bifunctional capping agent.

9. The cell of claim 8 wherein the capping agent is mercaptoacetic acid.
10. The cell of any preceding claim wherein absorption of light by a nanocrystal results in production of an exciton, the nanocrystal having a largest spatial dimension no greater than an average diffusion distance of the exciton.

11. The cell of any preceding claim wherein the nanocrystals comprise at least one of CuInSe$_2$, CuInS$_2$, CuIn$_x$Ga$_{1-x}$Se$_2$, GaAs, InAs, InP, PbS, PbSe, PbTe, GaSb, InSb, CdTe and CdSe, wherein $0 \leq x \leq 1$.

12. The cell of any preceding claim wherein the nanocrystals have extinction coefficients of at least 100,000 M$^{-1}$cm$^{-1}$.

13. The cell of any preceding claim wherein the semiconductor polymer is bound to the nanocrystals but not to the nanorods.

14. A method of fabricating a semiconductor structure comprising heterojunctions and being suitable for use in a photovoltaic cell, the method comprising the steps of:
   a. providing a plurality of nanorods and a plurality of photoresponsive nanocrystals capped with a first capping agent;
   b. exposing the nanorods or the nanocrystals to a second, bifunctional capping agent;
   c. thereafter combining the nanocrystals with the nanorods, whereby the nanocrystals bind to the nanorods via the bifunctional capping agent;
   d. combining the bound nanorods and nanocrystals with a functionalized monomer having a binding group, the binding group (i) exhibiting a stronger affinity for the nanocrystals than the first capping agent and (ii) exhibiting a weaker affinity for the nanorods than the bifunctional capping agent, whereby the monomer preferentially displaces the first capping agent so as to bind to the nanocrystals but not to the nanorods; and
   e. polymerizing the monomer.

15. The method of claim 14 further comprising the step of disposing the nanorods between the first and second electrodes, the nanorods each having one end in electrical contact with the
first electrode, and being electrically insulated from the second electrode through a thin polymer layer at the other end.

16. The method of claim 14 or 15 further comprising, before step (a), the step of growing a plurality of nanorods on a substrate providing the first electrode.

17. The method of claim 14, 15 or 16 further comprising, after step (e), the step of depositing the second electrode, the second electrode being electrically insulated from the nanorods through a thin polymer layer.

18. The method of claim 14, 15, 16 or 17 wherein step (a) involves providing a nanocrystal capped with a first capping agent containing at least one binding functionality selected from the group consisting of thiol, selenol, amine, phosphine, phosphine oxide, or an aromatic heterocycle.

19. The method of any one of claims 14 to 18 wherein the first capping agent is octanethiol.

20. The method of any one of claims 14 to 19 wherein step (b) comprises capping the nanorods with the second capping agent, the first capping agent having a weaker affinity for the nanocrystals than the second capping agent, whereby the second capping agent preferentially displaces the first capping agent so as to bind to the nanocrystal.

21. The method of any one claims 14 to 20 wherein step (b) comprises exposing the nanocrystals to the second capping agent, the second capping agent replacing some but not all of the first capping agent on the nanocrystals, whereby in step (c) the free ends of the second capping agent bind to the nanorods.
GROW NANORODS ON ANODIC SUBSTRATE

CAP NANORODS WITH BIFUNCTIONAL CAPPING AGENT

BIND NANO-CRYSTALS TO NANORODS

ADD FUNCTIONALIZED MONOMERS TO NANOCRYSTAL-SENSITIZED NANORODS

POLYMERIZE

DEPOSIT CATHODE

FIG. 2A
GROW NANORODS ON ANODIC SUBSTRATE

PROVIDE NANO-CRYSTALS CAPPED WITH FIRST CAPPING AGENT

FUNCTIONALIZE MONOMERS

REPLACE SOME OF FIRST CAPPING AGENT WITH BIFUNCTIONAL CAPPING AGENT

BIND NANO-CRYSTALS TO NANORODS

ADD FUNCTIONALIZED MONOMERS TO NANOCRYSTAL-SENSITIZED NANORODS

POLYMERIZE

DEPOSIT CATHODE

FIG. 2B
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT HATTER

INV | H01L51/42

According to International Patent Classification (IPC) and to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

HOIL HOIG

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, INSPEC

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<td>A</td>
<td>paragraph [0066] - paragraph [0103]; figures IA, IF</td>
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<td>Y</td>
<td>EP 1 176 646 A (ECOLE POLYTECH [CH]) 30 January 2002 (2002-01-30) the whole document</td>
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Further documents are listed in the continuation of Box C.

See patent family annex.

* Special categories of cited documents:
*A* document defining the general state of the art which is not considered to be of particular relevance

**E** earlier document but published on or after the international filing date

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**P** document published prior to the international filing date but later than the priority date claimed

**T** later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

**X** document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

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Date of the actual completion of the international search

12 August 2008

Date of mailing of the international search report

21/08/2008

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Authorized officer

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<td>WANG ET AL: &quot;In situ polymerization of amphiphilic acetylene for hole transport in solid-state dye-sensitized solar cells&quot; ORGANIC ELECTRONICS, ELSEVIER, AMSTERDAM, NL, vol. 7, no. 6, 18 November 2006 (2006-11-18), pages 546-550, XP005773063 ISSN: 1566-1199 the whole document</td>
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<td>wo 2007/098378 A (SOLEXANT CORP [US]; REDDY DAMODER [US]) 30 August 2007 (2007-08-30) figures 5-12</td>
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