



High Efficiency c-Silicon Solar Cells Based on Micro-nanoscale Structure

by Fred Semendy, Priyalal Wijewarnasuriya, and Nibir K. Dhar

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14. ABSTRACT This report describes our aim to develop high-efficiency crystalline silicon (c-Si) solar cells with increased power conversion efficiency (>30%) in order to reduce solar array mass, stowed volume, and cost. Our approach is based upon increasing the electrical junction area per unit volume using microblock design and fabrication. Current thin-film and c-Si solar cells have a limited conversion efficiency of 10–20% and cost \$3–\$5/W-peak and state-of-the-art crystalline multijunction solar cells have a ~30% efficiency and cost \$30–\$40/W-peak. Increasing the conversion efficiency to >30% enables a reduction in cost to <\$1/W-peak, making the cells viable as power platforms supporting mobile wireless, laptops, residential, and commercial applications. To do this, we are moving from a two-dimensional standard flat cell to three-dimensional blocks of cells. Incorporating nanoscaled blocks in solar cell structures enhances the performances by (1) increasing the surface area-to-volume ratio; (2) bringing the junction closer to the carrier generation region, which eliminates carrier recombination; (3) increasing absorption of all incident photon flux; and (4) broadening the absorption spectrum. We have achieved a >20 times greater short-circuit current than conventional structures. We have also modeled the proposed structure to simulate key performance parameters so that they can be incorporated into the design and fabrication of the device.					
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1. Introduction

In the current energy crisis, the search for a viable alternative to hydrocarbons has taken many paths: nuclear, wind, solar, etc. Solar cells provide an attractive form of limitless alternative energy. The placement of solar cells can be unobtrusive and provide not only a source of thermal energy, but electricity. However, the development and implementation of effective photovoltaic (PV) cells is hindered by two primary components: cost and efficiency. Research into cheaper and more efficient solar cells has been underway for several decades, from the development of thin-film solar cells with efficiencies greater than 10% in the 1970s to the most recent developments in new PV materials achieving greater than 24% efficiency, as shown in figure 1.

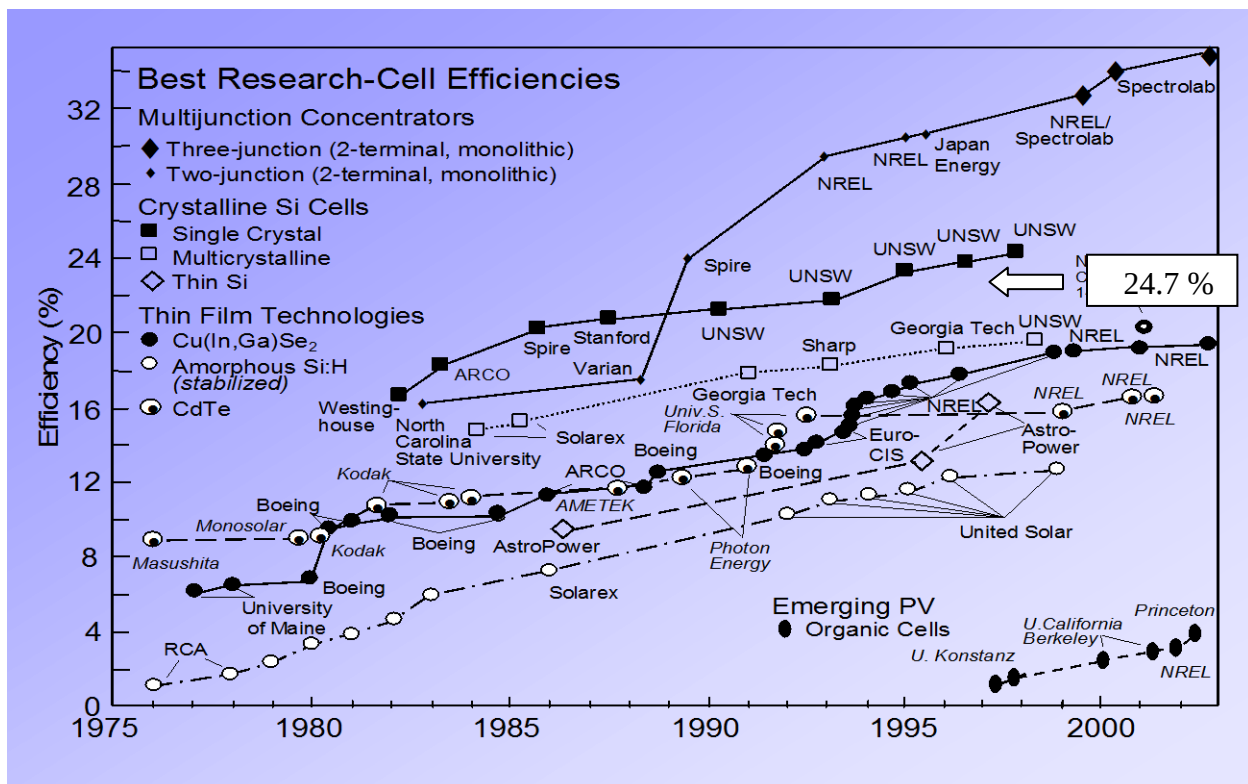


Figure 1. PV efficiency solar cells of different materials.

In general solar energy and its many applications are not widely accepted compared to conventional electrical energy solutions. The main reason for the low acceptance of solar energy is the high manufacturing costs involved. Especially in Japan and Germany, many efforts have been made to reduce manufacturing costs and expand the usability of solar energy in numerous industrial, residential, and commercial applications. In the United States, research and development (R&D) and applications are far behind other developed countries like Japan and Germany. In order to cope with the current energy crisis, it is highly desirable to build solar

energy infrastructures inside the United States alongside other energy solutions so that the United States can leapfrog other countries in the field of solar energy (1, 2).

Crystalline silicon (c-Si) and thin films (with multijunction approaches) are two major pathways to achieving solar energy goals. Each of these consists of multiple options (e.g., c-Si approaches include Si wafer and several types of sheet Si). This multiple-path characteristic makes solar cell technology robust and offers high potential for future advances in both performance improvement and cost reduction. For more demanding applications such as residential, industrial, and even, commercial solar generation systems, either polycrystalline or single-crystalline Si is the best choice due to stringent requirements for better reliability and higher efficiency than applications in consumer electronics. c-Si has a well-established technology base and the c-Si industry supplies nearly 90% of solar cell demands. c-Si will continue to dominate the market for at least five years. The technical progress will evolve, but advances will be integrated quickly into the marketplace. This will help to build the infrastructure required for continued rapid growth of solar cell technology. Cell conversion efficiencies for current c-Si approaches vary from 12% to 17% (3). Module efficiencies tend to be 0.5% to 2% lower, based on total area. However, due to the high cost of solar energy solutions (\$3 to \$5/W-peak) compared to other conventional electrical energy solutions, this Si-based solar cell is not yet widely accepted as an alternative to energy solution. As many concerns associated with a steep increase in the amount of the worldwide energy consumption are raised, further developments in c-Si solar cell technology for industrial systems applications are required and have been a primary focus.

Other leading candidates for very low-cost solar cell solutions are the following thin-film materials: (1) amorphous Si (a-Si) (4), cadmium telluride (CdTe) (5), and copper indium diselenide (CIS) (6), which are the most mature thin-film technologies; and (2) thin-film Si, which may encompass both c-Si and/or a-Si. The key metrics of thin-film progress are the same as those for any PV module technology—efficiency at the commercial module level, manufacturing cost, and outdoor module reliability. Many challenges must be overcome before these factors can be optimized. For thin films, the monolithic process blurs the distinction between cells and modules. However, commercial module efficiencies vary between 5% and 11%. The cost of these thin-film solar cells (\$3 to \$5/W-peak) is almost same as that of c-Si. However, due to the toxic natures of these materials and concerns for the environment, thin-film-based solar cells are not widely accepted.

To achieve a higher conversion efficiency (20% and higher), conventional multijunction combinations of III-V semiconductors are usually used (7, 8). The materials and manufacturing cost of the multijunction solar cell costs more than 10 times that of Si solar cells. It is highly desirable to have an alternative solution of a solar cell made of Si, which could offer a higher efficiency (20% and higher) and have a comparable or lower cost than today's Si solar cell.

We propose a solar cell structure that would be environmentally friendly, highly efficient, and cost effective. In this report, we provide an overview of a high-efficiency Si solar cell based on

micro-nanostructures, which can be fabricated using standard Si integrated circuit (IC) technology and is amenable to mass-scale production. Details of simulation and fabrication technology are also provided. We believe the proposed high-efficiency Si solar cell could eliminate the energy crisis and greatly benefit the United States in many ways.

2. Solar Cell Technology

Before providing the technical approach for significantly increasing the conversion efficiency ($>20\%$) of c-Si solar cells using less Si material (0.6 g/W, final target), we present an overview of current solar cell solutions and their limitations with regard to making high-efficiency solar cells.

2.1 Current Solar Cell Technology

Solar cell structures have been studied extensively over the last decade for various applications. Figure 2 is a conventional solar cell comprising a thick p-type semiconductor layer and a thin n-type semiconductor layer formed on an electrically conductive substrate.

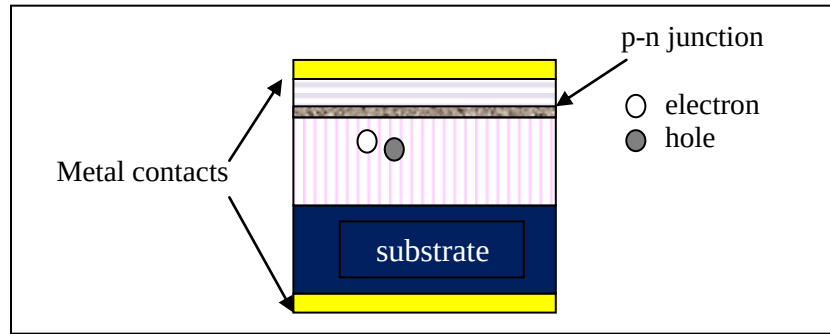


Figure 2. Conventional solar cell.

A p-n-junction is formed at the interface between the p- and n-type semiconductor layers. Incident sunlight entering in the cell generates electron-hole pairs after being absorbed by the p- and n-type semiconductor layers. The incident light generates electrons and holes in the region near the p-n-junction and far from the p-n junction. The photogenerated electrons diffusing toward the p-n junction and entering the p-n junction region contribute to the PV effect. The two key factors that substantially impact the conversion efficiency of solar cell are photocarrier generation efficiency (PCGE) and photocarrier collection efficiency (PCCE). For monochromatic light, a PCGE of $\sim 100\%$ can be achieved by simply making the p-type layer thicker. However, electrons generated at the region far from the p-n junction cannot be collected efficiently due to many adverse recombination processes. These processes prevent photogenerated carriers from diffusing into the p-n junction, thus the basic structure of current PV cells has its own limitation on increasing the conversion efficiency.

The light intensity p at certain depth x can be expressed by $p(x)=Po\exp(-\alpha x)$, where Po is the peak intensity at the surface and α is the absorption co-efficient of the semiconductor into which light enters. The light intensity behavior inside a bulk semiconductor is approximately exponential in behavior. Carriers (not shown in figure 2) generated due to light flux absorbed by the p-n junction are only drifted by the junction field and can be collected efficiently.

Conversely, carriers generated due to absorption of light flux by the semiconductor region are diffused in all directions. Only those which are generated closer (a separation distance equal to or less than the diffusion-length of the semiconductor) to the p-n junction can be collected. Carriers that are generated far away (a separation distance longer than the diffusion-length of the semiconductor) from the p-n junction are recombined and lost. The light flux is usually lost either by escaping or being absorbed by the substrate. For these reasons, it is difficult to increase the conversion efficiency of the cell if the standard structure is used. Apart from these, one has to consider the bulk recombination process, which reduces the minority carrier lifetime. To alleviate this, we need defect-free materials. In addition, the nature of the surface plays an important role. High reflection at the illuminated surface can be avoided by antireflection coating and other light-trapping effects. Because of the bandgap limitations, long wavelength light escapes in Si, which can be avoided by having multiple passes, optimizing the reflection from front to back and front surfaces, and optimizing the substrate thickness.

The solar spectrum is shown in figure 3. The spectrum, as seen from a satellite, is referred to as the AM0 spectrum (where AM stands for air mass) and closely fits the spectrum of a black body at 5800 K. The total power density is 1353 W/m^2 .

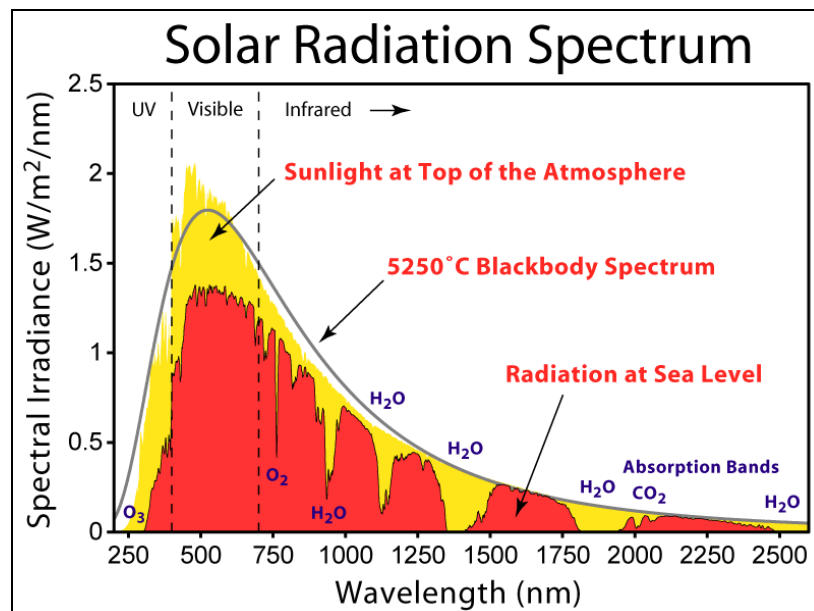


Figure 3. Solar radiation spectrum.

The solar spectrum, as observed on Earth, is modified due to absorption in the atmosphere. For AM1 (normal incidence), the power density is reduced to 925 W/cm^2 ; whereas, for AM1.5 (45° above the horizon), the power density is 844 W/m^2 . The irregularities in the spectrum are due to absorption at specific photon energies. Figure 4 gives the current-voltage (I-V) graph for a solar cell.

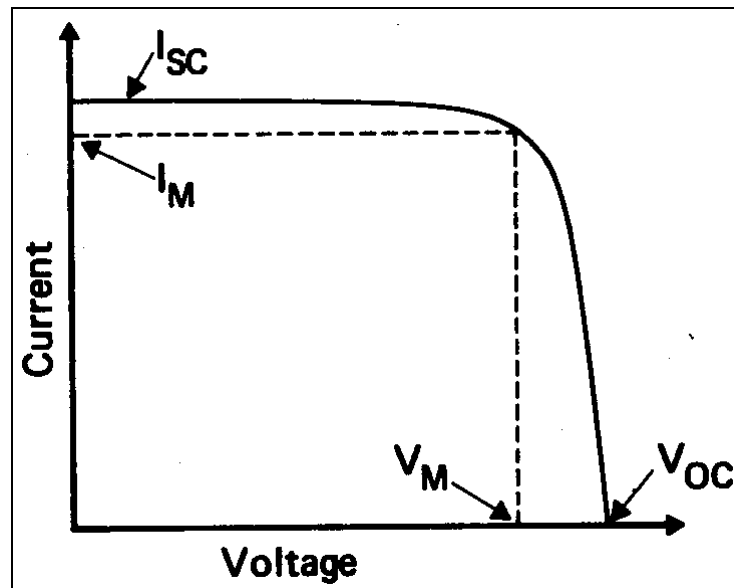


Figure 4. I-V characteristics of a solar cell.

The important parameters for calculating power conversion are the following:

- The short-circuit current (I_{SC}) is the point at which the I-V curve crosses the x-axis at 0 V. When a solar cell is operated at short circuit (that is, when a low-resistance connection is established by accident or intention between two points in an electric circuit so the current tends to flow through the area of low resistance bypassing the rest of the circuit), $V = 0$ and the current (I) through the terminals is defined as the short-circuit current.
- The open-circuit voltage (V_{OC}) is the voltage at which there is zero current flow. When a cell is operated at open circuit (that is, an incomplete electrical circuit in which no current flows, $I = 0$), the voltage across the output terminals is defined as the open-circuit voltage.
- The maximum power output (P_{MAX}) is the voltage and current points where the cell is generating its maximum power. The P_{MAX} point on an I-V curve is often referred to as the maximum power point (MPP).
- The current at maximum power (I_m) is the cell's current level at P_{MAX} .
- The voltage at maximum power (V_m) is the cell's voltage level at P_{MAX} .
- Fill factor (FF) is P_{MAX} divided by V_{OC} multiplied by I_{SC} . FF is a popular measurement because it indicates the cell's efficiency under a specific spectrum and intensity of light. In

essence, it calculates the percentage of performance of the real cell versus an ideal cell with no internal losses. Thus, by using the diode equation and V_{OC} , and manipulating a number of equations, one can arrive at the power conversion efficiency,

$$\eta = \frac{V_m I_m}{P_{in}}, \quad (1)$$

which can be further modified by substituting for I_m

$$\eta = \frac{V_m q A_a \int \phi(\lambda) QE(\lambda) d\lambda}{A_t \int \phi(\lambda) \frac{hc}{\lambda} d\lambda}, \quad (2)$$

where A_a is the active area and A_t is the total area. Thus, by increasing the junction area, one can maximize the power conversion efficiency. Therefore, our goal is to increase the volume-to-surface area ratio using microblock design masks. Using this technique, large junction areas can be made using microstructural fabrication, in which the junction is closer to the carrier generation region, thus reducing carrier recombination. The design is easy to implement and is independent of the material system.

2.2 Our Approach and Goal

The proposed solar cell structure consists of a number of micro-nanoscaled blocks (micro-nanorods/wires) around and on which the p-n junction is formed, thereby increasing surface (junction) area, as shown in figure 5. The p-n junction is formed very close to the region where the carriers are generated; thus, all photogenerated carriers can be collected before recombination. Furthermore, incorporation of these nanoscaled blocks into the solar cell structure can create quantum confinement and intermediate states within the bandgap to harvest photons with energy less than that of the bandgap of the host material. In this way, nanoscaled block structures in the proposed solar cells can be optimized to absorb a large portion of the solar spectrum. The front-side contact is transparent so that light can pass through the semiconductor, and the back-side contact is formed on the substrate. Differences in the proposed solar cell over conventional solar cells include a larger junction area due to use of nano (or micro) structures and the creation of a junction closer to the carrier generation region, which eliminates carrier recombination. In addition, most of the incident photons flux is absorbed due to the angular nature of the microstructure. Furthermore, there is less usage of Si, which, in turn, makes the device more cost effective, enabling it to be used as the universal structure for significantly high-efficiency solar cells. Precise control of the nano (or micro) cylindrical blocks' (a.k.a., pillars) length, density per unit area, reliability, and mechanical robustness will be ensured. The proposed unique and novel structure approaches will open doors to the incorporation of nano/microscaled cylindrical blocks for designing significantly high-conversion-efficiency solar cells.

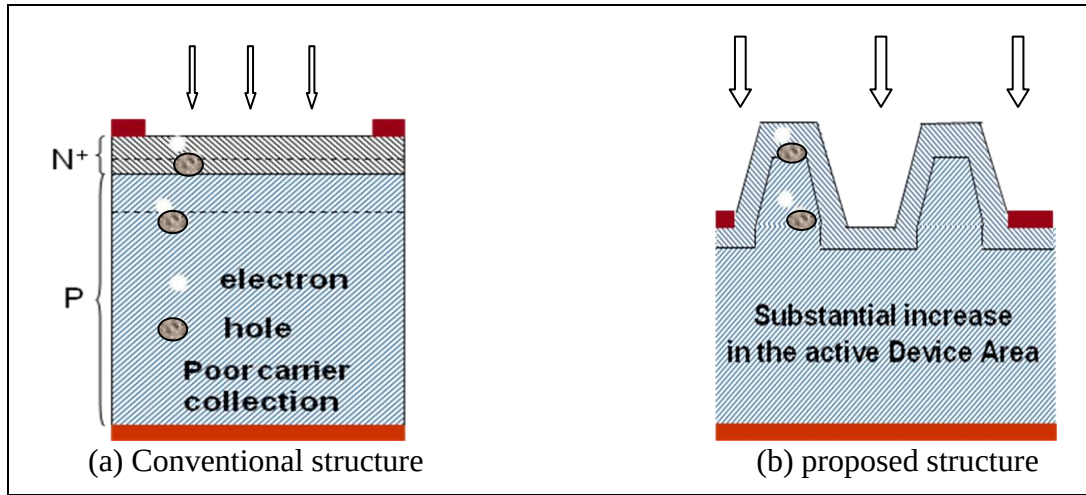


Figure 5. (a) A conventional solar cell and (b) the proposed solar cell in this study.

The proposed solar cell can be made from any kind of system material. For example, the substrate material can be any semiconductor, such as Si, gallium arsenide (GaAs), indium phosphide (InP), etc., glass, metal, or polymer. If the substrate is a semiconductor such as Si, micro-nanopillars can be formed using the etching approach, and if the substrate base material is InP, similar micro-nanopillars can also be made on the substrate. The proposed structure can also be fashioned from polymer to make the solar cells flexible. In this case, nanorods can be incorporated into the polymer materials to make a flexible solar cell using the proposed structure. We propose to fabricate the solar cell based on micro-nanocylindrical blocks, having a high conversion efficiency and high Si intensity.

We have carried out the simulation and also fabricated test structures on the proposed solar cell incorporating cylindrical structures. Initial experimental results demonstrated that the proposed solar cell showed significant conversion efficiency (>20 times greater compared to conventional cells). Theoretical estimated results showed that the conversion efficiency can be increased to >30% for a c-Si solar cell. Simulation and experimental results are explained in sections 3 and 4.

The proposed structure was calculated for a Si-based solar cell to see the benefits of the structure, varying the side tilting angle at fixed height of 5 μm . Noted here that the thickness of the c-Si wafer ($5 \times 10^{17} \text{cm}^{-3}$) considered in the simulation was 350 μm , which is frequently used in solar cell fabrication. A shallow p-n junction of 0.1 μm was considered in the simulation. A vertical trapezoidal structure was assumed to be formed using the Si wafer. In simulation, it was also assumed that the carrier movement was mainly dominated by diffusion, rather than drift, as the estimation of the depletion region due to built-in-potential was 120 nm. We used standard solar irradiance and the absorption coefficient for Si in simulation.

Figures 6 and 7 show the simulation results of conversion efficiencies at various tilting angles. Pillar structure 1 shows less angle and structure 3 shows a 86° angle, nanopillar (conventional), without the pillar case. Decreasing the tilting angle offers higher conversion efficiencies over 35%, which are thought to be due to (1) increasing the junction area, (2) increasing the

absorption spectra, and (3) increasing the light intensity due to the self-concentrating effect of the cylinder (pillar). More simulation and optimization are necessary to accurately access the conversion efficiency for the proposed structure.

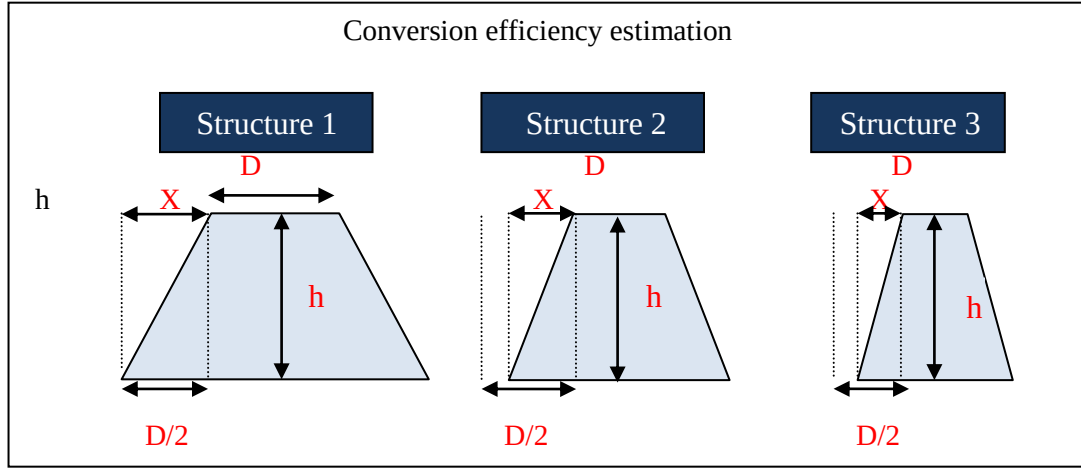


Figure 6. Schematics showing the various pillar structures used in simulation.

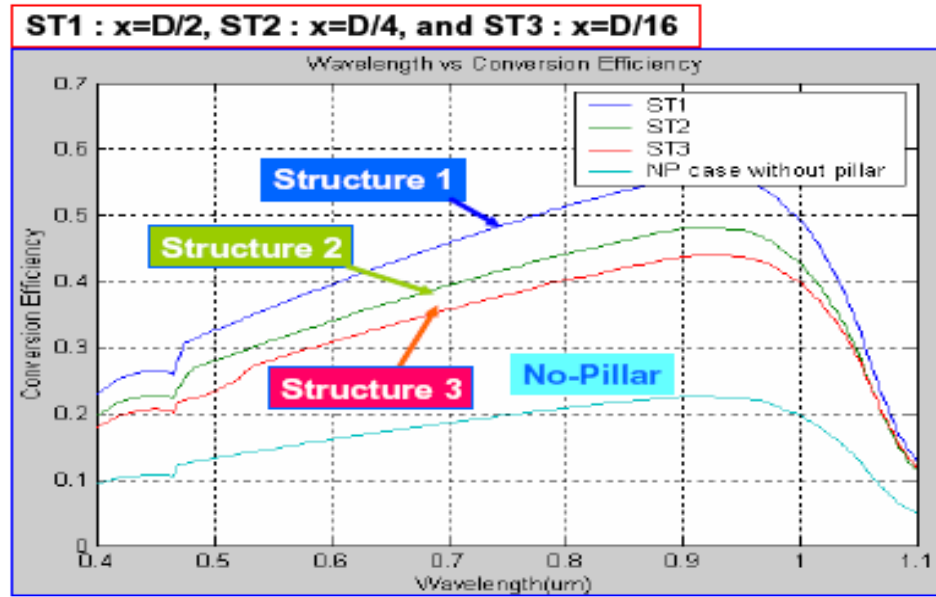


Figure 7. Conversion efficiency comparison as parameter of pillar inclinations with structure 1: $x=D/2$, structure 2: $x=D/4$, and structure 3: $x=D/16$.

3. Experimental

The proposed solar cell structure based on micro-nanopillars was fabricated using a c-Si solar cell. Figure 8 shows the mask design part of the work with (a) the pillar size in consideration and (b) for a 4-in Si wafer. Figure 9 shows (a) a 1-cm die and (b) the micro-pillar etching area showing the size of the pillar ($2\ \mu\text{m}$).

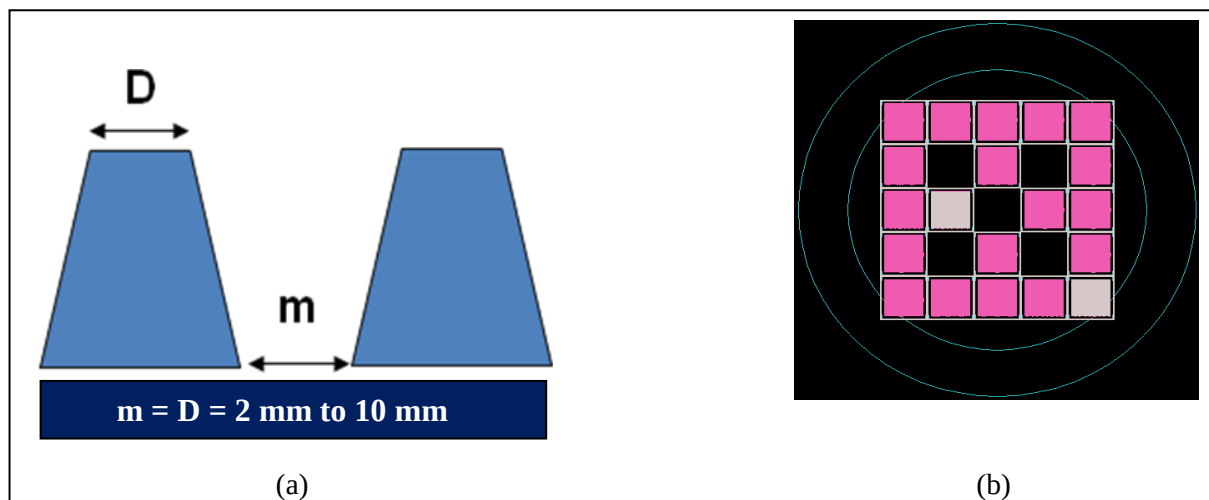


Figure 8. Mask design: (a) pillar size and (b) for a 4-in Si wafer.

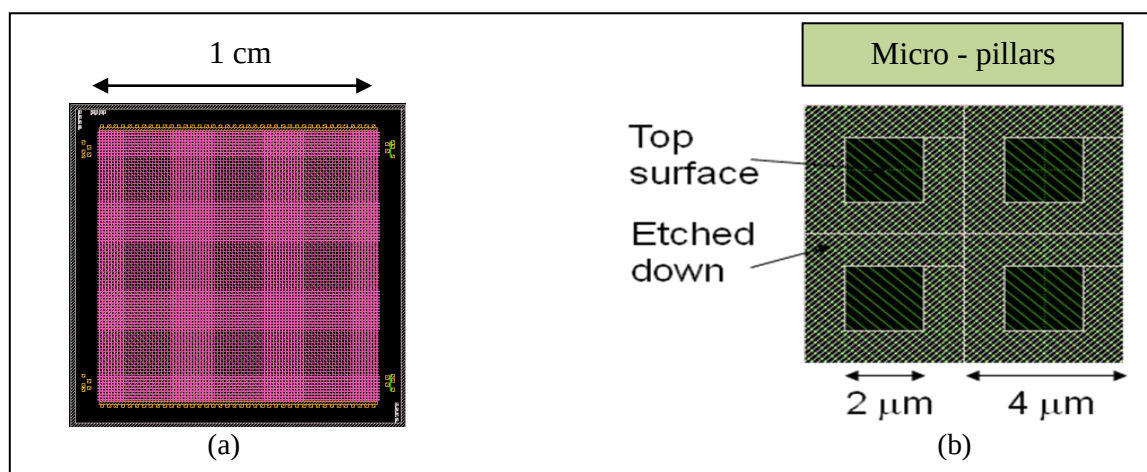


Figure 9. Mask design: (a) 1-cm die and (b) micro-pillars.

Boron-doped, 4-in crystalline Si wafers with a conductivity of $10\ \Omega\text{cm}$ and a thickness of $500\ \mu\text{m}$ were used for the fabrication. The wafer was p-type doped at $\sim 10^{15}\ \text{cm}^{-3}$. For the junction formation, phosphorus was ion implanted at an energy level of $100\ \text{keV}$ with a dosage of $3 \times 10^{13}\ \text{cm}^{-2}$ to enhance the layer to n^+ . Secondary ion mass spectrometry (SIMS) analysis was performed to verify the doping level and the results are shown in figure 10.

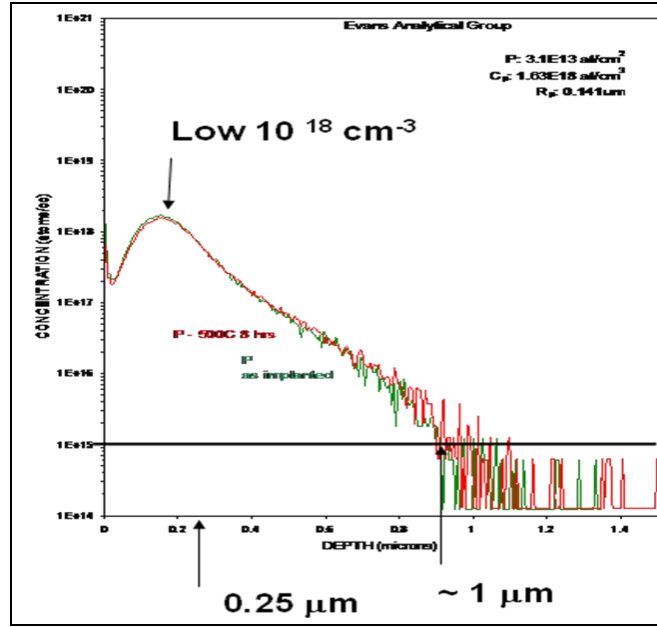


Figure 10. SIMS analysis of the n^+ Si after phosphorus implantation.

The fabrication process flow for the c-Si solar cell with micro-nanopillars is shown in figure 11. To form vertically arranged and well-defined microblocks in the c-Si ($<100>$) substrate, the standard dry-etching technique was used. After cleaning the wafer using the standard cleaning procedures, photolithographic techniques were used to define the micro-pillar structure. This was followed by etching using the inductively coupled plasma (ICP) deep reactive ion etching (DRIE) technique.

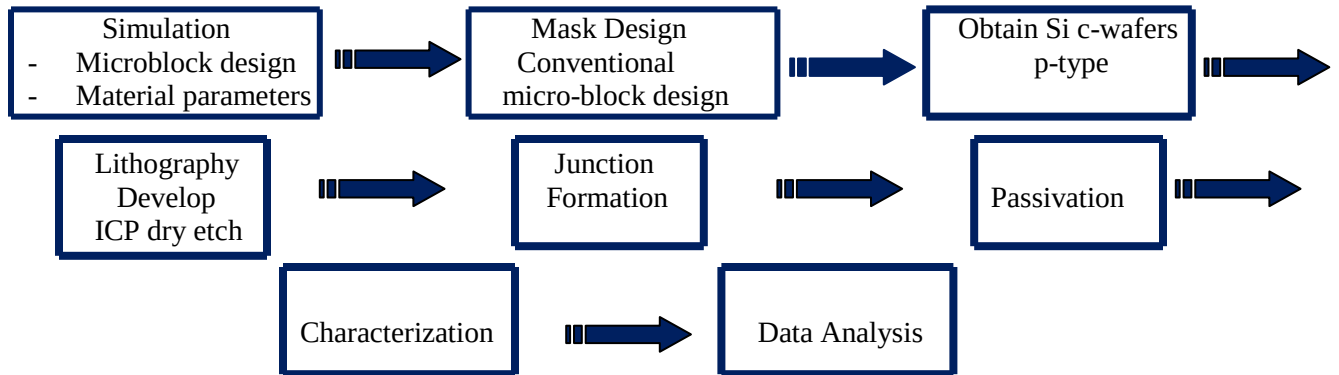


Figure 11. Fabrication process flow.

A mixed gas system of sulfur hexafluoride (SF_6) and oxygen (O_2) was used for the dry etching with varying pressures, gas ratios, times, and temperatures. Cleaned wafers were n^+ ion implanted, followed by passivation using thermal growth of silicon dioxide (SiO_2). The thickness of the thermally grown SiO_2 is calculated as shown in figure 12.

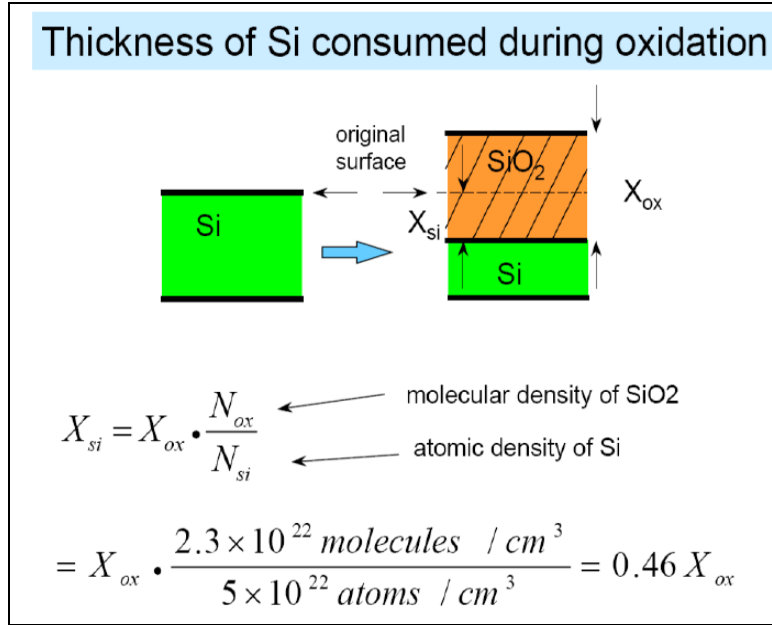


Figure 12. Thickness calculation of thermally grown SiO₂.

After passivation and metallization, devices were characterized and data were analyzed.

4. Results

The dry-etch process to create the micro-pillars was found to be complicated and difficult to optimize. Parameters such as pressure, amount of gas, radio frequency (RF), ICP power, time, and gas ratio had to be varied to obtain the right recipe to etch down and create the micro-pillars. Figure 13 shows some of the micro-pillars obtained by ICP-DRIE process. As shown in the image, the walls are vertical in the etch process. The actual case (as shown in figure 13) demonstrates how not enough O₂, the right amount of O₂, and excess amount of O₂ can provide the best possible etch required by the modeling. In the actual experiment, along with the variations in the amount of O₂ used, changes in other parameters were also made to fine-tune to optimize the etch process.

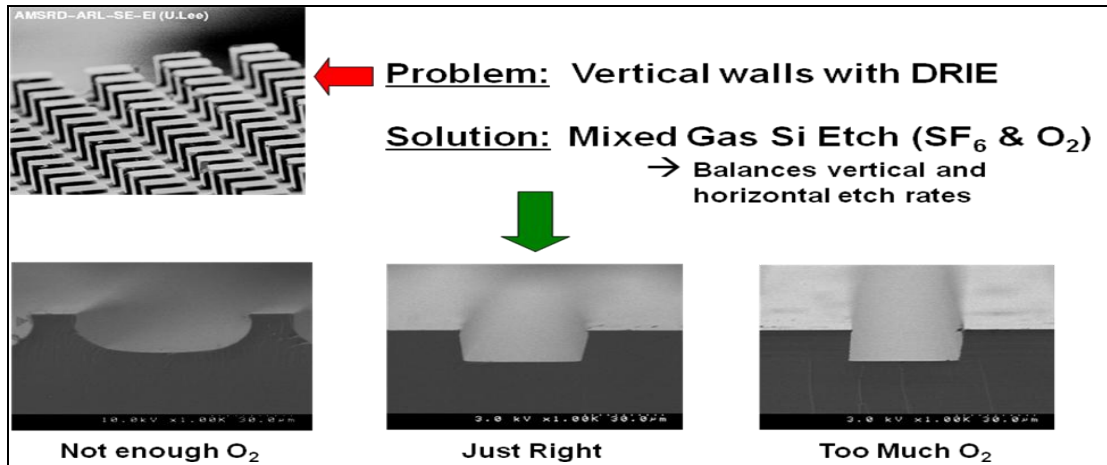


Figure 13. Dry etch of c-Si and variations in etch profile by changing the amount of O_2 .

For the fabrication of the devices, we used conventional cells of $1 \times 1 \text{ cm}^2$ from a 4-in silicon wafer. After the fabrication process, each cell was diced out and wire bonded for characterization. A 20-W white lamp light was used for the initial I-V characterization. During the measurement, the lamp was kept at a vertical height of 18 in, directly above the sample. Figure 14 shows the preliminary I-V characteristics of the solar cells and the conventional (flat, without pillar) cell.

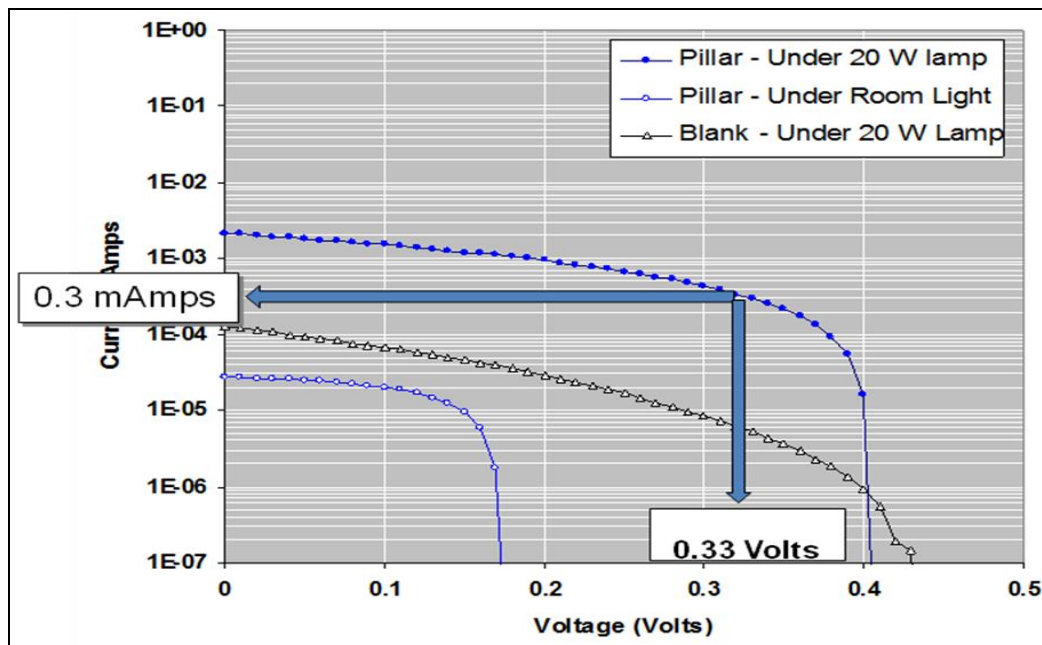


Figure 14. Preliminary I-V results of fabricated solar cells under a 20-W white lamp light and room light illumination. The lamp was at a vertical distance of 18 in away from the target cells.

Conversion power for the micro-pillar and conventional cells was found to be 200 and 20 μW , respectively, showing a more than $\sim 180 \mu\text{W}$ power difference. Further optimization of fabrication process and structural designs are necessary to further enhance the conversion efficiency performance. *Initial experimental results indicate that the proposed solar cell structure promises to improve the conversion efficiency using thinned substrates of any material system.* The proposed solar cell structure is expected to offer higher efficiency for all semiconductor materials as compared with the standard solar cell. Even though the final power conversion efficiency was not as expected, we intend to optimize the process by (1) dry etching specially on smaller scale pillars, (2) optimizing the lithography, (3) improving the P and n^+ contacts, (4) implanting conditions for the P dopant, (5) diffusing P into silicon, (6) solving the passivation issues, and (7) exploring the possibility of boron doping in Si. These optimizing processes can be tried on c-Si as well as in thin solar cell materials.

3. Conclusion

In this report, we present a high-efficiency Si solar cell that consists of micro-nanoblocks with an increased surface area-to-volume ratio and a p-n junction over the 3-D blocks for the top and bottom contacts. Simulation results indicate that (1) the absorption efficiency was increased due to self concentrating or photon trapping and (2) the absorption spectra increased in comparison with the flat device. Simulation from first principles indicates that the short-circuit current increased due to bringing the junction closer to the carrier generation sites. These phenomena increased the conversion efficiency of the Si solar cell. Experimental results of the solar cell fabricated using conventional photolithographic techniques showed that the short-circuit current of the solar cell based on the micro-nanoscale structures was 18 times greater than the standard solar cell fabricated on the same Si substrate.

In order to develop this high-efficiency solar cell, we will need to further optimize the design and iterate the parameters to accurately model the proposed solar cell, and then evaluate and demonstrate our ability to fabricate the optimized solar cell.

We believe that proposed solar cell technology will enable a high-conversion-efficiency, cost-effective solar cell technology that can be mass-produced and customized for commercial and defense systems.

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List of Symbols, Abbreviations, and Acronyms

a-Si	amorphous SI
CdTe	cadmium telluride
c-Si	crystalline silicon
CIS	copper indium diselenide
DRIE	deep reactive ion etching
FF	fill factor
GaAs	gallium arsenide
I	current
IC	integrated circuit
ICP	inductively coupled plasma
InP	indium phosphide
I _{SC}	short-circuit current
I-V	current-voltage
MPP	maximum power point
O ₂	oxygen
P _{MAX}	maximum power output
PCCE	photocarrier collection efficiency
PCGE	photocarrier generation efficiency
PV	photovoltaic
R&D	research and development
RF	radio frequency
SF ₆	sulfur hexafluoride
SiO ₂	silicon dioxide
SIMS	secondary ion mass spectrometry
V _{OC}	open-circuit voltage

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