Ion implantation for silicon solar cells

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ABSTRACT
This paper presents the background and technology development of the use of ion implantation technology in today’s crystalline silicon solar cell manufacturing lines. The recent history of ion implantation development and commercialization is summarized, and an explanation is given for the cell efficiency improvements realized using the technique on p-type mono-crystalline cells. The potential economic impact on the factory is also discussed.

Introduction
There are continuing pressures on PV manufacturers to lower manufacturing costs further while increasing efficiency. The efficiency of the module impacts all downstream area-related costs in the installed PV system, such as wiring and bracing structures. Manufacturers have already brought costs down to very low levels through better yield and economies of scale, and are now evaluating various approaches to improving cell efficiency. While standard screen-printed p-type cell efficiency can still be improved, most roadmaps will transition to advanced cell designs to meet the increased efficiency requirements. Thus, any investment in near-term manufacturing solutions has to also offer extendibility paths towards advanced cells and materials. There are numerous high-efficiency device architectures and various technology roadmaps regarding material type, cell architecture, process flow and process equipment. Cell efficiencies are generally predicted to improve as shown in Fig. 1, with the implication that new designs and processes will be adopted by industry to achieve the higher efficiencies.

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Ion implantation has a unique characteristic in that it is both beneficial to current cell designs and extendible to future cell architectures. In the near term, ion implantation provides higher cell efficiency for P emitters, narrower efficiency distribution and a lower overall cost. Equally important, the manufacturing of many advanced device designs can be significantly simplified by ion implantation. Recent efforts to commercialize high-productivity ion implanters have placed ion implantation technology centre stage in the silicon PV roadmap.

Synopsis of implantation used in solar cells
Throughout the history of implanted solar cells, various implanter tool designs have been utilized, the two main ones being mass-analyzed and non-mass-analyzed. Mass analysis is performed with a large magnet which bends the ion beam through an aperture. Only species with the proper mass and energy pass through the aperture. The filtered ion beam current is typically significantly less than the initial ion beam current. Non-mass-analyzed implanters, however, feature higher beam currents and lower capital costs, although they exhibit a broad range of ion energies and may co-implant other precursor species such as H+ and F+.

Mass-analyzed implantation
Mass-analyzed ion implantation has a unique characteristic in that it is both beneficial to current cell designs and extendible to future cell architectures. Throughout the history of implanted solar cells, various implanter tool designs have been utilized, the two main ones being mass-analyzed and non-mass-analyzed. Mass analysis is performed with a large magnet which bends the ion beam through an aperture. Only species with the proper mass and energy pass through the aperture. The filtered ion beam current is typically significantly less than the initial ion beam current. Non-mass-analyzed implanters, however, feature higher beam currents and lower capital costs, although they exhibit a broad range of ion energies and may co-implant other precursor species such as H+ and F+.

Ion sources have also changed over the decades and include glow discharge, microwave and inductively coupled RF plasmas. One of the earliest references to using ion implantation for solar cells was in 1964 by King and Burrill [2], who used a Van de Graaff electrostatic accelerator to accelerate boron or phosphorus ions generated by a microwave ion source.

Glow discharge plasma source designs were common in the 1970s and 1980s [3–6]. A variety of dopant gases – including BF3, B2H6, PH3 and PF5 – were utilized as the precursors. One of the most advanced and ambitious implementations of implantation for solar cells was that of the Hoxan Corporation in Japan in 1982 [7]. Using 4” round wafers, Hoxan built an integrated, in-line, computer-controlled, 9MW manufacturing line with a non-mass-analyzed implanter using BF3 (40keV) to form the rear boron back-surface field (B-BSF), and a second non-mass-analyzed implanter utilizing vapours from a heated solid P source (25keV) to form the emitter.

Annealing was performed with a halogen lamp and followed by TiO2 ARC and screen printing/firing stages. The entire automated manufacturing line also included tabbing and stringing and then module assembly and lamination. Each process was synchronized to five seconds per wafer for a throughput of 720 wafers per hour (wph).

Also in the 1980s, several researchers utilized commercial mass-analyzed beamline implanters for solar cells. Mass-analyzed ion implantation was becoming more important for integrated-circuit applications, in which the energy precision was crucial. Spitzer et al. [8–10] investigated various implantation conditions and anneals for both phosphorus and boron emitters using mass-analyzed implantation; for phosphorus emitters, 18% efficiencies were obtained.

Figure 1. ITRPV Roadmap, March 2012 update: stabilized cell efficiency trend curves [1].

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Later, in 1987, using a glow discharge source, Wood et al. [11] demonstrated 19.5% efficiency (AM1.5) using a non-mass-analyzed B$_2$H$_6$ (5% in H$_2$) implant for the emitter (n-type wafer), and non-mass-analyzed PH$_3$ (1% in H$_2$) implant for the phosphorus back-surface field (P-BSF). Using oxide passivation and photolithography to pattern evaporated metal contacts, a $V_{oc}$ greater than 660mV was demonstrated. The high $V_{oc}$ obtained demonstrates the exceptional quality of the implanted emitter and BSF, and indicates that the co-implanted hydrogen was not deleterious, despite the dilution of the dopant gas in H$_2$.

More recently, numerous groups have shown high cell efficiencies on industrial phosphorus emitter cells using modern commercial mass-analyzed implanters. For example, Suniva Inc. has demonstrated high cell efficiencies of ~19% on an industrial scale using a mass-analyzed ion implanter [12,13]. Non-mass-analyzed implanters have also been targeting solar cell applications. For instance, Intevac has demonstrated a continuous flux non-mass-analyzed implant tool – ENERGi™ – specifically designed for the solar industry, featuring a 2400wph throughput, sheet resistance uniformities of 1% ($\sigma$/mean) wafer to wafer, and a cost of ownership (COO) that is competitive with the standard POCl$_3$ diffusion process. In collaboration with China Sunergy, Intevac has achieved high efficiencies of 19% in an industrial line [14] using the ENERGi implanter.

**Advantages of ion implantation**

Ion implantation in general provides numerous process advantages. Specifically, the advantages of implantation include:

- High-precision dopant doses and profiles
- High uniformity and repeatability
- Single-sided doping capability
- Boron-doping capability
- Patterned doping

For today’s standard screen-printed p-type cells, ion implantation yields improved efficiencies through precision phosphorus profiles and high uniformity. The single-sided doping capability simplifies the process flow by eliminating the need for phosphosilicate glass (PSG) removal and edge isolation. Additionally, in situ thin passivating oxides can be easily incorporated during the subsequent anneal. Single-sided ion implantation can benefit thinner wafers by substituting a high-stress alloyed Al-BSF with B-BSF.

Tools and technologies adopted for today’s processes must also be extendible and compatible with the future cell designs throughout a manufacturer’s development roadmap. With the advent of industrial processes to passivate p-type surfaces, there is increasing benefit in transitioning away from standard cell structures with an aluminum-alloyed BSF. The single-sided doping capability of implantation facilitates this transition by eliminating multiple diffusion-barrier formation and removal steps otherwise required by furnace diffusions to form a B-BSF or P-BSF.

Patterned doping for selective emitters can be accomplished through inserting a shadow mask into the ion beam [15,16]. The ENERGi ion implantation can also pattern dopants for selective emitters by using shadow masks so that both the blanket and SE dopants are implanted in the same single-wafer pass through the ion beam.

Because boron diffusion has been difficult to implement cost-effectively in industry, implanting both phosphorus and boron, followed by a single co-anneal, greatly simplifies many designs, such as passivated rear cell designs [13,17,18] and n-type cells with boron emitters [19–22]. For very high efficiencies, ion implantation holds the promise of significant process simplification for interdigitated back-contact (IBC) cells [23–25]. Patterned, single-sided P and B doping with a co-anneal has the potential to make IBC manufacturing far simpler than using repeated diffusion masks and furnace diffusions, as shown in Fig. 2.

This comparison of a furnace-based IBC process flow and an implant-based process flow...
flow illustrates the potential process simplification using implantation.

Defect engineering synopsis

Ion implantation damage and recrystallization has been studied for decades. Consequently, very sophisticated models of damage formation and annealing exist, which are beyond the scope of this paper, but a brief synopsis of relevant damage and annealing processes will be given. For phosphorus implantation, the strategy is to form an amorphized surface layer which recrystallizes epitaxially with very few defects during a subsequent higher temperature annealing step. This process is called solid-phase epitaxial regrowth (SPER). The key to minimizing defects through the SPER process is to form a fully amorphized layer with a smooth a/c interface and a minimum of damage beyond the a/c interface. While phosphorus implantation does amorphize the silicon well, boron implantation is more complicated because it does not typically amorphize the top layer of silicon for the doses desired for solar cells.

Except for the lightest elements such as H, as the implanting ion enters the silicon, it transfers most of its energy to the target lattice through collisions. These collisions create local heat and, if more than 15–20eV is transferred to the lattice silicon atoms, one or more Frenkel pairs (Si vacancy + Si interstitial) are created [26]. The damage along the path of the implanting ion, i.e. the cascade collapse, is highest near the end of the range where the ion’s velocity is sufficiently slow so that nearly every atomic interaction results in a displacement event. Some of the created defects have a very short lifetime, and, with the localized heating assisting the defect mobility, the defects can begin to annihilate (i.e. anneal) on a nanosecond timescale. This dynamic annealing is undesirable because, as mentioned earlier, a well-formed amorphous layer requires significant damage accumulation. Partially amorphized layers or regions with sub-amorphous damage do not typically result in low defects, even after higher temperature annealing. Dynamic annealing can be prevented by cooling the sample to cryogenic temperatures. Implantation studies at such low temperatures illustrate the strong effects of dynamic annealing at higher temperatures. For example, the minimum dose required to amorphize at ~150°C is less than $1 \times 10^{15}$ B/cm². In contrast, at room temperature the required implantation dose is $2 \times 10^{17}$ B/cm² for amorphization.

Other factors contributing to better amorphization include ion species, continuous flux and dose rate [27]. Heavier ions, such as phosphorus, create more damage per ion and therefore require lower doses for full amorphization. Viewing the amorphization process as a dynamic competition between damage accumulation and dynamic annealing, one can understand that a higher dose rate [28] and continuous implant will accumulate damage faster and help form fully amorphized layers. Numerous ion implantation tools raster or pulse the ion beam, thus allowing time between the beam sweeps for dynamic annealing to occur. On the other hand, continuous fluxes and higher dose rates are expected to result in deeper amorphous layers, with less sub-amorphous damage beyond the a/c interface.

During a subsequent annealing step, the amorphous layers will recrystallize epitaxially, with the a/c interface moving towards the surface at a few hundred nm/s at 700°C [29]. Typically, the recrystallization is fully complete by the time the wafer boat is loaded into a furnace at 700°C. Dopants within this regrown epitaxial layer have high levels of activation, even if the dopant concentration somewhat exceeds the solubility during the SPER [30]. However, during the anneal (typically >800°C), equilibrium concentrations will be restored and supersaturated dopants will precipitate or cluster [31]. Thus proper implant and anneal optimization is important for obtaining emitters of the highest efficiency. Additionally, during the anneal the minor sub-amorphization damage, which lies deeper than the initial a/c interface, can be minimized or completely annealed as the phosphorus diffuses deeper into the silicon.

The ENERGi implantation tool was designed with both the needs of the solar market and defect-free emitters in mind. The ENERGi tool cools the wafer, which undergoes a single pass through a continuous and high-flux phosphorus ion beam. The difference between a rastered and a continuous flux implantation is schematically illustrated in Fig. 3 for the same total dose (area of each coloured region). At any given point on a silicon wafer, the rastered ion beam allows periods of time for dynamic annealing to occur and may result in poorer amorphized layers.

Figure 2. Comparison of process flows for IBC cells by diffusion and by patterned implantation.

Figure 3. The ion beam flux observed for any given point on a wafer for a rastered ion beam and for a continuous flux ENERGi ion beam for the same dose (area of each coloured region).
At Intevac, fully amorphized layers with doses as low as $1 \times 10^{15}$ P/cm$^2$ have been demonstrated. Examples of amorphization at different energies using the ENERGIi continuous-flux implantation tool are shown in Fig. 4, as well as the defect-free recrystallization of the 20kV implanted sample.

Boron does not amorphize silicon at doses of interest for solar cells. Instead of SPER, the sub-amorphous implant damage needs to be annealed at higher temperatures. Good emitters have been demonstrated by annealing at high temperatures such as 1000°C. Some problems with $V_{oc}$ or internal quantum efficiency (IQE) response have been reported for B+ implanted emitters despite low $J_{sc}$ values having been achieved [32–34]. However, other studies have shown excellent results with similar B+ implanted emitters [21,22]. To be successful, boron implantation for solar cells will depend on the right tool, implant and anneal.

**Efficiency improvements for standard P emitters by ion implantation**

The efficiency gain from ion implantation in today’s phosphorus emitters is based on precisely controlling the near-surface phosphorus dose. Any phosphorus emitter optimization entails maximizing the sheet resistance while minimizing carrier recombination and contact resistance to the front metallization. While the sheet resistance is a function of the entire profile shape, the recombination and the contact resistance are very sensitive to the portions of the emitter containing the highest concentration of phosphorus, i.e. the first ~50nm of the emitter. Fig. 5 shows a PC1D simulation, illustrating the general importance of the near-surface region. The black curve indicates the phosphorus profile, with concentration values on the left axis; the red curve is an approximation of the power loss (as a function of depth) caused by carrier recombination. (Fig. 5 is intended only to illustrate the importance of the near-surface region – various assumptions and simplifications were made that are outside the scope of this paper.)

By lowering the near-surface phosphorus dose, improved emitters and higher efficiency cells can be fabricated. An improvement in $V_{oc}$ and $J_{sc}$ with decreasing dose is shown in the data plotted in Fig. 6: as the total phosphorus dose is decreased, recombination in the emitter is reduced, which results in improved $V_{oc}$ and $J_{sc}$.

However, below a threshold phosphorus dose the contact resistance to the front metallization may increase significantly. The industrial Ag-based screen-printed metallization requires high (degenerate) doping levels at the Ag-Si interface in order to make low-resistance tunnelling junctions [35]. This effect places a lower limit on the reduction of the near-surface phosphorus dose. In Fig. 6, the fill factor (FF) is also plotted as a function of implanted phosphorus dose. For a fixed anneal, the near-surface dose will be proportional to the total implanted dose. Below a threshold dose, the paste and firing conditions result in higher contact resistance and degraded FF. While new pastes are able to contact lower surface concentrations, a strong drop in FF will still occur if the concentration is too low. Thus, for industrial high-efficiency emitters the near-surface phosphorus dose...
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must be precisely optimized and controlled across the wafer and from wafer to wafer.

Such near-surface phosphorus dose control is more difficult with POCl₃ diffusion. In this case the phosphorus dose and phosphorus drive-in are increasing during the anneal. The near-surface dose is very sensitive to temperature and ambient gas variations in the furnace. This process trajectory (increase in dose and \( J_{0E} \) with reduction in sheet resistance) is illustrated in Fig. 7 by the red arrow. More complex furnace recipes involving multiple anneal plateaus and ambient gases [9] can, to a limited extent, alter the near-surface phosphorus concentration and improve \( J_{0E} \). The POCl₃ process is therefore depicted as a wider range by the grey area in Fig. 7; for illustration purposes, the emitter etch-back [36] process trajectory is also included and is indicated by the green arrow. During the etch-back, the surface and peak concentrations are reduced, resulting in a lower \( J_{0E} \) emitter and higher sheet resistance.

In contrast, the ion implantation process separates the dose step and the drive-in step in order to precisely control and tailor the final phosphorus profile. During the anneal, the as-implanted phosphorus dopant starts to diffuse into the silicon. This redistribution of the implanted P results in a reduction in both \( J_{0E} \) and sheet resistance as indicated by the blue lines in Fig. 7, where each blue line represents a different phosphorus dose. This process trajectory is somewhat orthogonal to the typical POCl₃ diffusion.

An additional process advantage of ion-implanted phosphorus emitters is that an in situ passivating oxide can be rapidly grown during the anneal. Bhosle et al. [37] have shown a +0.35% absolute efficiency gain by utilizing an in situ oxide. The high surface concentrations of phosphorus accelerate oxide growth and allow for thicker oxides in short periods of time [38].

The final profile of the cell can be finely tailored by a combination of implant dose and energy, as well as anneal time, temperature and ambient. Fig. 8 summarizes the main parameters of the implant and anneal which control the various portions of the final P profile. Notice that, even after the anneal, the near-surface dose is largely controlled by the initial implant dose and energy. As discussed earlier, it is vital to optimize and control this near-surface P to achieve the best balance of contact resistance to the surface metallization and the highest \( V_{oc} \) and \( J_{sc} \).

Performing such an optimization in an industrial line results in a higher efficiency than typical POCl₃ diffusion is capable of. This has been demonstrated by Rohatgi et al. [13], who showed an increase in absolute efficiency of +0.3% to +0.8%, and by Wang et al. [39], who showed an improvement in absolute efficiency of more than 0.4%.
over POCl$_3$. In cooperation with China Sunergy, Intevac, using their ENERGi ion implantation tool, compared implanted cells with standard POCl$_3$ cells processed on a China Sunergy industrial process line [14]. The normalized histogram shown in Fig. 9 already shows a clear improvement using ion implantation with only preliminary optimization. Intevac recently fabricated cells on an industrial manufacturing line with another partner and demonstrated average cell efficiencies exceeding 19.1% (n>2000 wafers). This result also represented a significant improvement over the cell efficiencies for standard POCl$_3$ diffusion. The basic process steps are shown in Fig. 10.

**Economics**

The COO of the recent commercial high-throughput ion implanters is compelling because not only are higher cell efficiencies obtained, but also the distribution of the cell efficiency is narrower. Owing to its high throughput, the ENERGi ion implantation tool is cost competitive with the standard POCl$_3$ process via the higher final cell efficiency, the throughput and cost reduction by the elimination of PSG etch and edge isolation.

After process optimization, the ENERGi phosphorus implant operating at 2400wph can conservatively improve the cell efficiency by more than 0.5% absolute over the diffusion process. The impact of this on the output of a manufacturing line is an increase of more than 2MWp/year with a value of $1.4 million/year. Narrowing the distribution in cell efficiencies can increase the value even more by virtue of a reduction in scrap, and higher module prices at the higher efficiencies. The investment in the line is also minimized by the fact that adding ion implantation removes the need for wet-etching steps (phosphorus glass etch and edge isolation). This elimination removes approximately $1 million of capital cost from the line and also removes the acid and water running costs (~$0.3 million/year). With these considerations, the payback time of an ion implantation system running at 2400wph is approximately 1.3 years. This is one of the most efficient payback schedules possible in the solar manufacturing business today. The payback time is even shorter for boron implant with higher cell efficiency improvements. As this value is realized, ion implantation systems will become ubiquitous and highly valued by today’s Tier 1 solar cell and module manufacturers. Lastly, the Intevac ENERGi tool has a very small footprint compared with other tools which handle similar throughput of wafers on the fab line (Fig. 11). This makes retrofitting existing lines for phosphorus implantation easier.

**Conclusion**

In the solar cell technology roadmap a transition is taking place in which diffusion processes that have been used for many years are now being replaced by new, cell-efficiency-enhancing ion implantation solutions. Ion implantation offers solar cell manufacturers the capability to extend cell efficiency beyond 19% with a narrower cell efficiency distribution. Additionally, ion implantation offers a path towards the commercialization of new advanced cell architectures, n-type material and IBC cells. A continuous-flux plasma-based ion implantation solution provides the necessary economic solution to achieving this with a very efficient payback schedule.

**References**


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