Infrared analysis of the bulk silicon-hydrogen bonds as an optimization tool for high-rate deposition of microcrystalline solar cells

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In general, device grade microcrystalline silicon (μc-Si:H) properties are obtained in a narrow deposition parameter window close to the conditions at which the growth leading to the amorphous silicon (a-Si:H) phase transfers to the μc-Si:H phase.1,2 Device grade μc-Si:H can be classified as dense μc-Si:H without any significant post-deposition oxidation, as bulk oxidation is linked to a reduction in the red response of the p-i-n device.3-5 The most characteristic properties of μc-Si:H, such as microstructure, crystallinity, grain size, defects, and conductivity, have been extensively studied by Raman spectroscopy,6,7 x-ray diffraction,8 transmission electron microscopy,9 electron spin resonance,9 and optoelectronic characterization techniques.9,10,11 However, these easy-to-use analyze techniques are not able to unambiguously determine whether a deposited film is in the good narrow parameter window. The only reliable qualification of “device grade” material is the time-consuming procedure of the integration of an intrinsic film in a p-i-n device. Furthermore, the parameter window for device grade μc-Si:H becomes narrower for the industrially interesting high deposition rate (>2 nm/s) conditions in the high pressure depletion (HPD) (4–25 Torr) regime.3-5 As a result, easy optimization strategies of μc-Si:H material properties, without the necessity to integrate every film in to p-i-n devices, are highly desirable to reduce the time consuming optimization process. Here, we will demonstrate that the unwelcome incorporation of crystalline grain boundaries can easily be detected using infrared (IR) spectroscopy and that this observation can be used as a simple and fast optimization for the properties of high-rate deposited μc-Si:H. We demonstrate this on μc-Si:H deposited by two different high-deposition-rate setups in the very high frequency (VHF)-HPD regime, the first setup using a conventional showerhead electrode3 (SE) and the second setup using a multi-hole-cathode (MHC) electrode.12

The solar cells deposited by SE-VHF and its corresponding SE-VHF conditions A and B (see Table I) have been reported earlier in detail in Ref. 3. The μc-Si:H deposition conditions (C up to G) for the MHC-VHF setup correspond to a pressure of 10 Torr, VHF (80 MHz) power density of 1.3 W/cm², electrode gap of 7 mm, substrate temperature of 180 °C, silane flow of 12 SCCM (SCCM denotes cubic centimeter per minute at STP) under various hydrogen dilutions. Silane profiling was used during the initial 40 s of the deposition for conditions C up to G.31 The films are integrated in a solar cell structure of glass/ZnO/p-i-n/ZnO/Ag with an active area of 0.25 cm². Films with a thickness of 1.8 μm have been deposited on IR transparent c-Si samples for the IR analyses in transmission mode (Perkin Elmer, FT-IR Spectrum 2000). Note that for the depositions using the MHC-VHF (conditions C–G) compared to deposition using the SE-VHF, the texture of the ZnO, the intrinsic film thickness (~1.8 μm compared to 2.2–2.4 μm) and the p-i interface is not optimized and the solar cell structure is exposed to two vacuum breaks, one before and one after the i-layer deposition. Consequently, the solar cell performances of MHC-VHF conditions C up to G are lower than for the SE-VHF condition B (see Table I).

In Fig. 1, an IR spectrum, focused on the range of the hydride (Si-H₃) stretching modes (SMs), for μc-Si:H films deposited on c-Si, is presented and exhibits all modes which have been observed in undoped μc-Si:H solids so far. It is impossible to uniquely resolve all SMs using only one IR spectrum. Nevertheless, by using a large set of samples with

![Graph with absorption spectra](https://example.com/graph.png)
a wide variety of hydrogenated silicon (Si:H) phases, ranging from amorphous up to highly crystalline porous material, we were able to assign a consistent set of SMs capable of fitting the wide variety of spectra measured. The SM frequency position of a hydride in the bulk depends on the unscreened eigen frequency of the hydride, local hydride density, bulk screening, and possible mutual dipole interactions of the hydrogen incorporation configuration. The low SM (LSM) (1980–2010 cm−1) and the high SM (HSM) (2070–2100 cm−1) originate from the a-Si:H tissue in the bulk. The LSM ranges in μc-Si:H, broadens by two additional modes ~2120 and 2150 cm−1, due to significant contribution of di- and trihydrides to the macroscopic amorphous surfaces in the bulk. Furthermore, three narrow HSMs (NHSM) (2083, 2103, and 2137 cm−1) are observed, reflecting mono-, di-, and trihydrides on crystalline surfaces, assigned to crystalline grain boundaries in the bulk. The assignment of the extreme LSMs (ELSM) (~1895, ~1929, and ~1950 cm−1) is still under discussion. However, to explain its rather large frequency shift with respect to the frequency of unscreened mono-(2099 cm−1) and dihydrides (2124 cm−1), these hydrogen incorporation configurations have to correspond to extreme high local hydride densities combined with mutual hydride dipole-dipole interactions.

First, we consider the NHSMs reflecting hydrogenated crystalline surfaces, which are dominantly present in less dense μc-Si:H with a high crystallinity. The IR spectra of such μc-Si:H film, deposited under high hydrogen dilution and power conditions, are depicted in Fig. 2 as deposited, 10 days, and 10 months after deposition. Figure 2(a) shows the dihydride bending modes at 840–890 cm−1 and the Si–O–Si SMs at 950–1200 cm−1 and Fig. 2(b) shows the hydride SM absorption. The increase in the Si–O–Si SMs show the μc-Si:H film significantly oxidizes in the bulk due to the exposure to ambient air. Simultaneously, the intensity of the NHSMs reduces under air exposure and completely disappears within a few months, while a mode at 2250 cm−1 shows up. This latter mode corresponds to the hydride OSi−Hx vibration with oxygen atoms back bonded to the silicon atom. These trends reflect that the bulk oxidation occurs at least at the crystalline grain boundaries by most probably water and that the crystalline Si−H surfaces are fully transferred to OSi−Hx surfaces. This suggests that all crystalline grain boundaries have to be surfaces in an interconnected pore and crack network which ends up at the top surface of the μc-Si:H film.

In Table I, the performances of the solar cells deposited under conditions A up to G are presented. Condition A results in p-i-n efficiency of 4.5% at 2.0 nm/s, while a slight modification of the deposition parameters in to condition B results in good cell efficiency of 9.1% at 2.3 nm/s. The difference in material properties between conditions A and B is also reflected in the SMs, as depicted in Figs. 2(c)–2(f), respectively. The as-deposited film of condition A still reflects a small signature of NHSMs which disappears in 8 days, accompanied with the appearance of the OSi−Hx mode at 2250 cm−1. The presence of a significant postdeposition oxidation is also reflected by the increase in the Si–O–Si SMs. In contrast, in the as-deposited high quality film of condition B, the NHSMs are absent and no postdeposition oxidation is observed, reflecting a denser bulk matrix and the absence of hydrogenated crystalline grain boundaries in the pore network. Furthermore, for the Si:H phase in which the NHSMs are just absent, the total integrated area under the ELSM, LSM, and MSM has its maximum. In our interpretation, this

![Graph showing absorption spectra](image)

**Table I.** Illuminated J-V parameters of the p-i-n devices with the layer deposited under conditions A up to G. The crystalline fraction Xc is determined from Raman measurements of μc-Si:H deposited on Corning glass under conditions C up to G.

<table>
<thead>
<tr>
<th>Setup</th>
<th>H2 (SCCM)</th>
<th>Rx (nm/s)</th>
<th>Xc (%)</th>
<th>Voc (V)</th>
<th>Jsc (mA cm−2)</th>
<th>FF</th>
<th>η (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>SE+opt. TCO 670</td>
<td>2.0</td>
<td>0.467</td>
<td>15.5</td>
<td>0.62</td>
<td>4.5</td>
<td></td>
</tr>
<tr>
<td>B</td>
<td>SE+opt. TCO 670</td>
<td>2.3</td>
<td>0.528</td>
<td>23.7</td>
<td>0.73</td>
<td>9.1</td>
<td></td>
</tr>
<tr>
<td>C</td>
<td>1200</td>
<td>1.6</td>
<td>80 ± 3</td>
<td>0.435</td>
<td>20.1</td>
<td>0.65</td>
<td>5.7</td>
</tr>
<tr>
<td>D</td>
<td>MHC</td>
<td>1200 → 600</td>
<td>1.7</td>
<td>73 ± 3</td>
<td>0.497</td>
<td>20.6</td>
<td>0.68</td>
</tr>
<tr>
<td>E</td>
<td>MHC</td>
<td>600</td>
<td>1.8</td>
<td>69 ± 5</td>
<td>0.491</td>
<td>21.2</td>
<td>0.66</td>
</tr>
<tr>
<td>F</td>
<td>MHC</td>
<td>400</td>
<td>1.9</td>
<td>70 ± 7</td>
<td>0.502</td>
<td>20.5</td>
<td>0.68</td>
</tr>
<tr>
<td>G</td>
<td>MHC</td>
<td>200</td>
<td>1.9</td>
<td>47 ± 10</td>
<td>0.518</td>
<td>18.9</td>
<td>0.65</td>
</tr>
</tbody>
</table>

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reflected thin hydride-dense $a$-Si:H tissue, which either passivates the crystalline grain boundaries or fills the small pores, to prevent any postdeposition oxidation of grain boundary surfaces. Considering the fact that these postdeposition oxidizing crystalline grain boundaries are linked with the reduction of the red response of the $p$-i-$n$ device, a significant amount of the unwelcome recombination of charge carriers generated in the crystalline grains seems to take place at these surfaces, reflecting the poor surface passivation properties of a native oxide.

An issue to be addressed is the fact that the IR analysis is performed on $\mu$C-Si:H deposited on IR transparent $c$-Si samples and not on glass/TCO/ $\mu$C-Si:H($p$) substrates as used for the solar cells. Different substrate materials could induce different initial $\mu$C-Si:H growths. As the crystalline grain boundaries reflected by the NHSMs are only present in the postinitial growth zone (>200 nm), the relation between the IR spectrum and the solar cell performances is bulk dominated and therefore independent of the nature of the substrate. Consequently, the first optimization step of the $p$-i interface (mainly controlling the $V_{oc}$) has been performed by combined IR absorption and Raman spectroscopy (to guarantee the absence of an $a$-Si:H incubation) on thin films of 50–100 nm deposited on $c$-Si and Corning glass, respectively.

To demonstrate the generality of the approach presented above, $\mu$C-Si:H films deposited using MHC-VHF setup have been optimized by using the IR spectrum corresponding to condition B as a reference. The inclusion of a silane-profiling preparation is significantly larger. Taking condition D as a starting point, conditions C up to G have been obtained by variation of the hydrogen dilution with the purpose to create slightly different SM signatures in the IR. Figure 3 shows that the IR spectrum of condition C exhibits NHSMs, reflecting less dense material accompanied with postdeposition oxidation (not shown). In line with the trend observed for the films deposited using SE-VHF, the solar cell deposited under condition C has a significant reduced performance of $\eta$=5.7% in a $p$-$i$-$n$ device compared to conditions D–G, as a result of a reduced red response (not shown) and lower $V_{oc}$. Going from conditions E up to G, the general trend of increasing $V_{oc}$ and decreasing $J_{sc}$ versus decreasing hydrogen dilution is observed due to the fact that the amorphous fraction of the film becomes larger. This trend is also reflected in a larger contribution of the MSM and LSM to the IR spectrum for condition F. In our experience, the optimum efficiencies are only found for films having an IR signature such as conditions D, E, and F, independent of the deposition rate and the type of TCO substrates used.

In summary, we have demonstrated that the unique signature of the SMs of bulk hydrides in $\mu$C-Si:H can be used as an easy-to-use tool to optimize $\mu$C-Si:H properties at high deposition rates, without the necessity to integrate every film in a solar cell device during the film optimization process.