

**Methods of Deposition  
of  
Hydrogenated Amorphous Silicon  
for  
Device Applications**

*Wilfried G.J.H.M. van Sark*

Debye Institute, Utrecht University,

P.O. Box 80000, NL-3508 TA Utrecht, The Netherlands

Chapter 1 in Thin Films and Nanostructures, Vol. 30,  
'Advances in Plasma-Grown Hydrogenated Films',  
Edited by M. H. Francombe, Academic Press, San Diego, 2002, pp. 1-215.

and

Chapter 1 in Handbook of Thin Film Materials, Vol. 1,  
'Deposition and Processing of Thin Films',  
Edited by H. S. Nalwa, Academic Press, San Diego, 2002, pp. 1-102.

## Contents

<b>I</b>	<b>Introduction</b>	<b>5</b>
	A Historical overview . . . . .	6
	B Material aspects of hydrogenated amorphous silicon . . . . .	7
	1 Atomic structure . . . . .	7
	2 Microstructure and hydrogen content . . . . .	8
	3 Electronic structure and transport . . . . .	9
	4 Optical properties . . . . .	11
	5 Metastability . . . . .	12
	6 Alloys . . . . .	13
	7 Device quality characteristics . . . . .	16
<b>II</b>	<b>Research and industrial equipment</b>	<b>17</b>
	A General aspects . . . . .	17
	B Reactor configurations . . . . .	19
	C Scale-up to systems of industrial size . . . . .	20
	D ASTER, a research system . . . . .	22
<b>III</b>	<b>Physics and chemistry of PECVD</b>	<b>29</b>
	A General introduction . . . . .	29
	B Plasma physics . . . . .	29
	1 Plasma sheath . . . . .	29
	2 Scaling laws . . . . .	31
	3 Effective discharge power . . . . .	34
	4 Paschen law . . . . .	34
	C Plasma chemistry . . . . .	35
	1 Species considered . . . . .	36
	2 Electron impact collisions . . . . .	36
	3 Chemical reactions . . . . .	39
	4 Plasma-wall interaction . . . . .	39
	5 Transport coefficients . . . . .	40
<b>IV</b>	<b>Plasma modeling</b>	<b>41</b>
	A 1D fluid discharge model . . . . .	41
	1 Model description . . . . .	41
	2 Particle balances . . . . .	43
	3 Particle fluxes . . . . .	44
	4 Poisson equation . . . . .	44
	5 Energy balances . . . . .	44
	6 Boundary conditions . . . . .	45
	7 Electron energy distribution function . . . . .	47
	8 Calculation scheme . . . . .	48
	9 Simulation results . . . . .	49
	10 Comparison with experiments . . . . .	50
	B 2D fluid discharge model . . . . .	57
	1 Model description . . . . .	57
	2 Comparison with experiments, small reactor . . . . .	58

	3 Comparison with experiments, large reactor . . . . .	62
C	Particle-In-Cell discharge models . . . . .	65
	1 Particle-in-Cell model principles . . . . .	65
	2 Hybrid PIC/MC-fluid model . . . . .	66
	3 Simulation of RF discharges in silane/hydrogen with the hybrid model . . . . .	68
	4 Simulation of RF discharges in silane/hydrogen with a full PIC/MC model . . . . .	71
<b>V</b>	<b>Plasma analysis</b>	<b>77</b>
A	Optical emission . . . . .	77
B	Electrostatic probes . . . . .	78
C	Mass spectrometry . . . . .	82
	1 Analysis of neutrals . . . . .	82
	2 Analysis of radicals . . . . .	86
	3 Analysis of ions . . . . .	88
D	Ellipsometry . . . . .	99
<b>VI</b>	<b>Relation between plasma parameters and material properties</b>	<b>103</b>
A	External parameters . . . . .	103
	1 Plasma power . . . . .	103
	2 Total pressure . . . . .	104
	3 Gas flow . . . . .	104
	4 Hydrogen dilution . . . . .	104
	5 Substrate temperature . . . . .	105
	6 $\alpha$ - $\gamma'$ transition . . . . .	105
	7 Other deposition parameters . . . . .	108
B	Internal parameters . . . . .	108
	1 The role of ions . . . . .	108
	2 The ion flux . . . . .	111
	3 Relation between ion flux and material quality . . . . .	112
	4 Ion-surface interactions . . . . .	120
<b>VII</b>	<b>Deposition models</b>	<b>123</b>
A	Surface adsorption . . . . .	124
B	Solubility of hydrogen in silicon . . . . .	127
C	Elimination of hydrogen from <i>a</i> -Si:H . . . . .	127
	1 Thermal reactions . . . . .	128
	2 Athermal reactions . . . . .	130
D	Dangling bond and weak bond density . . . . .	131
<b>VIII</b>	<b>Modifications of PECVD</b>	<b>132</b>
A	VHF . . . . .	132
	1 General . . . . .	132
	2 Optimization of deposition conditions . . . . .	132
	3 Discharge analysis . . . . .	140
B	Chemical annealing . . . . .	142
C	RF modulation . . . . .	144

<b>IX</b>	<b>Hot Wire Chemical Vapour Deposition</b>	<b>150</b>
	A General description . . . . .	150
	B Experimental set-up . . . . .	151
	C Material properties and deposition conditions . . . . .	152
	D Deposition model . . . . .	154
<b>X</b>	<b>Expanding Thermal Plasma Chemical Vapour Deposition</b>	<b>156</b>
	A General description . . . . .	156
	B Apparatus, setup . . . . .	156
	C Material properties and deposition conditions . . . . .	158
	D Deposition model . . . . .	160
<b>XI</b>	<b>Applications</b>	<b>161</b>
	A Solar Cells . . . . .	161
	1 Operation principle . . . . .	161
	2 Single junction solar cells . . . . .	162
	3 Multiple junction solar cells . . . . .	164
	4 Stability . . . . .	166
	B Thin Film Transistors . . . . .	168
	1 Operation principle . . . . .	168
	2 Stability . . . . .	168
	3 Fabrication . . . . .	169
	4 Application in active matrix displays . . . . .	170
	C Light sensors . . . . .	170
	1 Linear arrays . . . . .	170
	2 Photoreceptors . . . . .	171
	3 Position sensor . . . . .	171
	4 Colour sensors . . . . .	172
	5 IR sensor . . . . .	172
	6 X-Ray sensor . . . . .	172
	D Chemical sensors . . . . .	173
	1 pH-ISFET . . . . .	173
	2 Hydrogen sensor . . . . .	173
	E Other applications . . . . .	173
	1 Loudspeakers . . . . .	173
	2 Erbium in <i>a</i> -Si:H . . . . .	175
	3 Miscellaneous . . . . .	178
<b>XII</b>	<b>Conclusion</b>	<b>179</b>
<b>XIII</b>	<b>Acknowledgements</b>	<b>180</b>
<b>XIV</b>	<b>References</b>	<b>181</b>

## I. INTRODUCTION

This chapter describes the deposition of hydrogenated amorphous silicon ( $a$ -Si:H) and related materials by employing a low-temperature, low-density plasma. The method basically is a special form of Chemical Vapour Deposition (CVD), which is known as Plasma Enhanced Chemical Vapour Deposition (PECVD) or Plasma-CVD. Essentially, silane gas ( $\text{SiH}_4$ ) is excited by a radio-frequent (RF, 13.56 MHz) plasma, which causes silane molecules to dissociate. Subsequently, dissociation products are deposited on heated substrates and form a layer. Most research and industrial reactor systems consist of two parallel electrodes in a stainless steel chamber. Because of the relative ease to deposit  $a$ -Si:H uniformly over large areas, the original parallel plate geometry with RF excitation frequency is commonly used in industry, and has not changed much over the past 2-3 decades [1–13]. Material and device optimization is mostly done empirically, and so-called "device quality"  $a$ -Si:H layers are made by PECVD having an excellent uniformity.

Nevertheless, several modifications have evolved since the first demonstration of deposition of  $a$ -Si:H, e.g. the use of higher excitation frequencies (from VHF (50–100 MHz) up to the GHz-range), the use of remote excitation of the plasma, plasma beams, and modulation in time or frequency of the plasma. Even methods without the assistance of a plasma have evolved, i.e., the Hot-Wire CVD (HWCVD) method.

The material properties of layers deposited in a PECVD reactor strongly depend on the interaction between the growth flux and the film surface. Therefore, a central theme in this chapter is the relation between material properties and deposition parameters. Considering the plasma as a reservoir of species, we can distinguish neutrals, radicals and ions, which can be either positive or negative. In a typical RF discharge, which is weakly ionized, the neutral species are the most abundant, having a concentration of about  $10^{16} \text{ cm}^{-3}$ , while the concentration of radicals and ions is only about  $10^{14} \text{ cm}^{-3}$  and  $10^{10} \text{ cm}^{-3}$ , respectively. The energy of these species may differ considerably: the neutrals, radicals and ions *within* the plasma are not energetic at all, below 0.1 eV. The ions that reach substrates and reactor walls are much more energetic, e.g. 1–100 eV. This can have enormous consequences for the impact of species on a growing film. For example, the amount of energy that is present in the plasma amounts to  $10^{13} \text{ eV/cm}^3$  for radicals and  $10^{12} \text{ eV/cm}^3$  for ions. This shows that although ions are much less present in the plasma, their impact might be comparable to that of radicals.

In this chapter we will treat the common RF-PECVD method for  $a$ -Si:H deposition, with emphasis on intrinsic material. First, a short introduction on the material properties of hydrogenated amorphous silicon is given. Subsequently, details are given on experimental and industrial deposition systems, with special emphasis on the UHV multichamber deposition system ASTER (Amorphous Semiconductor Thin film Experimental Reactor) at Utrecht University [14,15]. This is not only done because many experimental results presented in this chapter are obtained in this system, but to our opinion this system can also be seen as a generic multichamber deposition system. Then, a thorough description of the physics and the chemistry of the discharge is presented, followed by plasma modeling and plasma analysis results. Subsequently, relations will be formulated between discharge parameters and material properties and models for the deposition of  $a$ -Si:H are presented.

In further sections extensions or adaptations of the PECVD method will be pre-

sented, such as VHF-PECVD [16], the chemical annealing or layer-by-layer technique [17], and modulation of the RF excitation frequency [18]. The HWCVD method [19] (the plasma-less method) will be described and compared to the PECVD methods. The last deposition method that is treated is Expanding Thermal Plasma CVD (ETP-CVD) [20,21]. Other methods of deposition, such as remote-plasma CVD, and in particular Electron Cyclotron Resonance CVD (ECR-CVD), are not treated here, as to date these methods are difficult to scale-up for industrial purposes. Details of these methods can be found in, e.g., Luft and Tsuo [6].

As all these methods are used in research for improving material properties with specific applications in mind, a summary of important applications for which *a*-Si:H is indispensable, is given in the last section.

It will be clear throughout this chapter that it is biased to research performed at or in collaboration with the research group at the Debye Institute at Utrecht University. However, numerous references to other work are presented in order to put this research in a much broader perspective.

For further reading one may find excellent material in the (edited) books by (in chronological order) Pankove [1], Tanaka [2], Street [3], Kanicki [4,5], Luft and Tsuo [6], Bunshah [7], Bruno, Capezzuto, and Madan [8], Machlin [9,10], Schropp and Zeman [11], Searle [12], and Street [13].

### I.A. Historical overview

The research on amorphous semiconductors in the 1950s and 1960s was focused on the chalcogenides, i.e. materials containing group VI elements (sulfur, selenium and tellurium), e.g.  $\text{As}_2\text{Se}_3$ . These glasses are formed by cooling from the melt, their structure being similar to oxide glasses. Of particular interest was the relation between disorder of the structure and the electronic properties. This still is a question not fully answered, see e.g. Overhof in his Mott Memorial Lecture [22].

Amorphous silicon (and germanium) was prepared in those days by thermal evaporation or sputtering, see e.g [23]. This unhydrogenated material was highly defective, which inhibited its use as a useful semiconductor. Research on incorporating hydrogen as a passivating element was pursued by introducing hydrogen in the sputtering system, which indeed improved the electrical properties [24–26].

In 1965 it was discovered that deposition of amorphous silicon employing glow discharge as the deposition technique yielded a material with much more useful electronic properties [27,28]. The deposition occurs on a moderately heated substrate (200–300 °C) through reactions of gas radicals with the substrate. At that time the infrared absorption bands of silicon-hydrogen bonds were observed, but they were not recognized as such. Some years later, Fritsche and co-workers in Chicago confirmed that *a*-Si produced from a glow discharge of  $\text{SiH}_4$  contains hydrogen [29,30]. Recently, a personal account of the early years in amorphous silicon research shows the struggle to reach this conclusion [31]. Spear and co-workers in Dundee succeeded in improving the electrical properties, and finally, in 1975, a boost in research activities occurred when it was shown that *a*-Si:H could be *n*- or *p*-type doped by introducing phosphine ( $\text{PH}_3$ ) or diborane ( $\text{B}_2\text{H}_6$ ) in the plasma [32]: a variation of resistivity of more than 10 orders of magnitude could be reached by adding small amounts of these dopants.

The achievement of Spear and LeComber [32] immediately paved the way for practical amorphous silicon devices. In fact, it is argued that the research field became "polluted by the applications" [33]. Carlson and Wronski at RCA Laboratories started in 1976 with the development of photovoltaic devices [34]. The first *p-i-n* junction type solar cell was reported by the group of Hamakawa [35,36]. In 1980, Sanyo was the first to market devices: solar cells for hand-held calculators [37]. Also considerable research effort was directed towards amorphous silicon photoconductors for application in photocopying machines and laser printers [38,39]. The *a*-Si:H photoconductor is, amongst other materials, used as light-sensitive component in the electrophotographic process.

The first field effect transistors were reported [40–42] also at about this time. These thin film devices take advantage of the capability to deposit and process *a*-Si:H over large areas. It took only a few years before these thin film transistors (TFTs) were utilized in Active Matrix Liquid Crystal Displays (AMLCD) by various companies. Active matrix addressing can also be used in printer heads. Combining the photoconductive properties and the switching capabilities of *a*-Si:H has yielded many applications in the field of linear sensor arrays, e.g. 2D image sensors, position-sensitive detectors of charged particles, X-rays, gamma rays, and neutrons [43–47].

## I.B. Material aspects of hydrogenated amorphous silicon

### I.B.1. Atomic structure

Hydrogenated amorphous silicon is a disordered semiconductor whose optoelectronic properties are governed by the large number of defects present in its atomic structure. The covalent bonds between the silicon atoms in *a*-Si:H are similar to the bonds in crystalline silicon. The silicon atoms have the same number of neighbours and *on average* the same bond lengths and bond angles. One can represent the disorder by the atom pair distribution function, which is the probability of finding an atom at a distance  $r$  from another atom. A perfect crystal is completely ordered to large pair distances, while an amorphous material only shows short-range order. Because of the short-range order material properties of amorphous semiconductors are similar to their crystalline counterparts.

Amorphous silicon is often viewed as a continuous random network (CRN) [48,49]. In the ideal CRN model for amorphous silicon, each atom is fourfold coordinated, with bond lengths similar (within 1 % [50]) to that in the crystal. In this respect, the short-range order ( $< 2$  nm) of the amorphous phase is similar to that of the crystalline phase. Amorphous silicon lacks long-range order because the bond-angles deviate from the tetrahedral value ( $109.5^\circ$ ). The average bond-angle variation  $\Delta\Theta$  reflects the degree of structural disorder in the random network. Raman spectroscopy is used to determine the vibrational density of states. The transverse optical (TO) peak region has been used to determine  $\Delta\Theta$ . Beeman *et al.* [51] have related the half-width at half-maximum  $\Gamma/2$  of the TO peak to  $\Delta\Theta$ . Values for  $\Gamma/2$  range from  $33$ – $50$   $\text{cm}^{-1}$ , which translates into values of  $8$ – $13^\circ$  for the average bond-angle variation.

The continuous random network may contain defects, but the crystalline concepts of interstitials or vacancies are not valid here. Instead, in the CRN one uses the coordination defect, when an atom has too few or too many bonds. In *a*-Si:H dangling bonds

arise when a silicon atom has too few bonds to satisfy its outer  $sp^3$  orbital. It is the common view that the dominant defect in amorphous silicon is a threefold-coordinated silicon atom. This structural defect has an unpaired electron in a non-bonding orbital, called a dangling bond. Pure amorphous silicon has a high defect density,  $10^{20} \text{ cm}^{-3}$  (one dangling bond for every  $\sim 500$  Si atoms), which prevents photoconductance and doping. The special role of hydrogen with regard to amorphous silicon is its ability to passivate defects. Hydrogenation to a level of  $\sim 10$  at.% reduces the defect density by four to five orders of magnitude.

The incorporation of phosphorus yields fourfold coordinated P atoms, which are positively charged, as phosphorus normally is threefold coordinated. This substitutional doping mechanism was described by Street [52], thereby solving the apparent discrepancy with the so-called  $8 - N$  rule, with  $N$  the number of valence electrons, as originally proposed by Mott [53]. In addition, the incorporation mechanism, due to the charge neutrality to be preserved, leads to the formation of deep defects (dangling bonds). This increase in defect density as a results of doping explains the fact that  $a$ -Si:H photovoltaic devices are not simple  $p$ - $n$  diodes (as in crystalline materials): an intrinsic layer, with low defect density, must be introduced between the  $p$ - and  $n$ -doped layers.

### *I.B.2. Microstructure and hydrogen content*

The structure on a scale from the atomic level to about 10 nm is denoted by the term microstructure. Many structural aspects are represented in here, such as hydrogen bonding configurations (SiH, SiH<sub>2</sub>, SiH<sub>3</sub>, and polysilane ((SiH<sub>2</sub>) <sub>$n$</sub> ) groups), multi-vacancies, internal surfaces associated with microvoids, density fluctuations, unbonded hydrogen distribution (isolated and bulk molecular hydrogen) and bonded hydrogen distribution (clustered and dispersed). Also columnar structure, void size and volume fraction are included in this concept of microstructure, although these are of much larger scale.

The hydrogen content  $C_H$  greatly influences structure and consequently electronic and optoelectronic properties. An accurate measurement of  $C_H$  can be done with several ion beam-based methods, see e.g. Arnold Bik *et al.* [54]. A much easier accessible method is Fourier-transform Infrared Transmittance (FTIR) spectroscopy. The absorption of IR-radiation is different for different silicon-hydrogen bonding configurations. The observed absorption peaks have been indentified [55–57] (for an overview, see Luft and Tsuo [6]). The hydrogen content can be determined from the absorption peak at  $630 \text{ cm}^{-1}$ , which includes the rocking modes of all possible silicon-hydrogen bonding configurations. The hydrogen content is now defined as:

$$C_H = A_{630} \int_{-\infty}^{+\infty} \frac{\alpha_{630}(\omega)}{\omega} d\omega \quad (1)$$

with  $A_{630} = 2.1 \times 10^{19} \text{ cm}^{-2}$  the proportionality constant determined by Langford *et al.* [58]. Alternatively, one can determine the hydrogen content by integration of the absorption peaks of the stretching modes ( $2000$ - $2090 \text{ cm}^{-1}$ ), albeit by using the proportionality constant  $A_{2000-2090}$  determined by Beyer and Abo Ghazala [59],  $A_{2000-2090} = 1.1 \times 10^{20} \text{ cm}^{-2}$ . Typical hydrogen contents are 9-11 %.

In order to distinguish between isolated silicon-hydrogen bonds in a dense network and other bonding configurations, such as clustered monohydride and dihydride bonds, bonds on internal void surfaces, and isolated dihydride bonds, Mahan *et al.* [60] have defined the microstructure factor  $R_*$  as:

$$R^* = \frac{I_{2070-2100}}{I_{2070-2100} + I_{2000}} \quad (2)$$

$I_{2000}$  is the integrated absorption band due to isolated Si-H bonds. For all other bonds the IR absorption shifts to 2070-2100  $\text{cm}^{-1}$ . Isolated bonds are preferred, so  $R^*$  equals about zero for ideal material.

The refractive index of amorphous silicon is, within certain limits, a good measure for the density of the material. If we may consider the material to consist of a tightly bonded structure containing voids, the density of the material follows from the void fraction. This fraction  $f$  can be computed from the relative dielectric constant  $\epsilon$ . Assuming that the voids have a spherical shape,  $f$  is given by Bruggeman [61]:

$$f = \frac{(\epsilon_d - \epsilon)(2\epsilon_d + 1)}{(\epsilon_d - 1)(2\epsilon_d + \epsilon)}, \quad (3)$$

with the effective refractive index  $n = \sqrt{\epsilon}$ , and  $n_d = \sqrt{\epsilon_d}$  the refractive index of the dense fraction.

The presence of a dense material with a varying void fraction results in compressive stress, with typical values of 500 MPa. Compressive stress can be determined conveniently by comparing the curvature of a crystalline silicon wafer before and after deposition of an *a*-Si:H film.

Hydrogenated amorphous silicon is not a homogeneous material. Its structure is thought to consist of voids embedded in an amorphous matrix [62,63]. The size and number density of the voids depend on the deposition conditions. Poor-quality material can have a void fraction around 20 %, while device quality *a*-Si:H has been shown to contain fewer voids,  $\sim 1$  %, with a diameter of  $\sim 10$  Å [64–66]. The surfaces of the voids may partly be covered with hydrogen [62,67], or are also thought to be filled with  $\text{H}_2$  [68–72]. The influence of voids on the defect density in *a*-Si:H is the subject of much debate.

### *I.B.3. Electronic structure and transport*

The preservation of the short-range order results in a similar electronic structure of the amorphous material compared to the crystalline one: bands of extended mobile states are formed (defined by the conduction and valence band edges,  $E_C$  and  $E_V$ ) separated by the energy gap,  $E_g$ , more appropriately termed "mobility gap". Figure 1 shows schematically the density of states distribution. The long-range atomic disorder broadens the densities of energy states, resulting in band tails of localized states that may extend deep into the band gap. Coordination defects (dangling bonds) result in electronic states deep in the band gap, around mid-gap. As electronic transport mostly occurs at the band edges, the band tails greatly determine the electronic transport properties. The deep defect states determine electronic properties by controlling trapping and recombination.

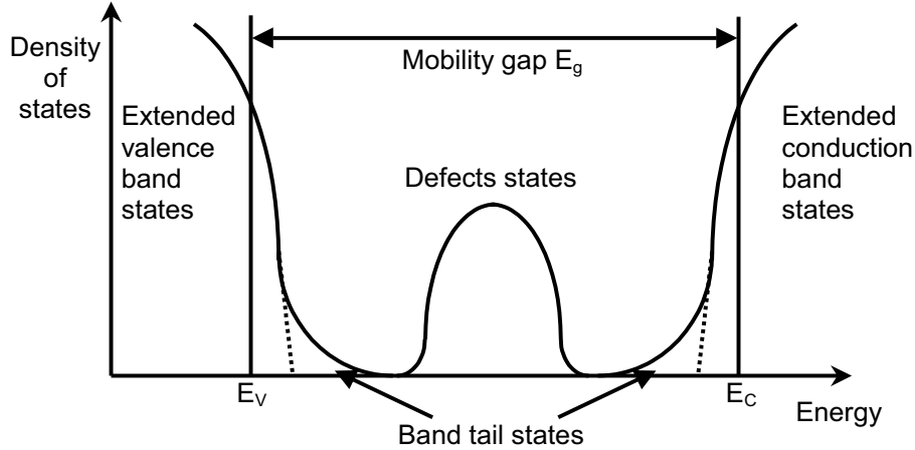


FIG. 1. Schematic density of states distribution. Bands of (mobile) extended states exist due to short range order. Long range disorder causes tails of localized states, whereas dangling bonds show up around mid-gap. The dashed curves denote the equivalent states in a crystal.

Normally the defect density is low, and electronic transport is considered to occur predominantly at the mobility edges. For electron transport one can write the following expression for the conductivity  $\sigma$ :

$$\sigma(T) = \sigma_0 \exp\left(-\frac{E_C - E_F}{k_B T}\right) \quad (4)$$

where  $E_F$  denotes the Fermi energy,  $\sigma_0$  the prefactor, and  $k_B$  Boltzmann's constant, and  $T$  the temperature. Of particular interest for devices is the conductivity in the dark,  $\sigma_d$ , and the photoconductivity  $\sigma_{ph}$ . For typical undoped material  $E_F$  lies at midgap and  $\sigma_d$  equals about  $10^{-10} \Omega^{-1}\text{cm}^{-1}$  or lower.

The activation energy  $E_A$ , defined as  $E_C - E_F$  for the conduction band (and analogous for the valence band), can be used to assess the presence of impurities. Due to their presence, either intentional (B or P dopant atoms), or unintentional (O or N), the Fermi level shifts over several tenths of eV towards the conduction or valence band. The activation energy is determined from plots of  $\log \sigma(T)$  versus  $1/T$ , with  $50 < T < 160$  °C. For undoped material  $E_A$  is about 0.8 eV. The Fermi level is at midgap position, as typically  $E_g$  is around 1.6 eV.

The photoconductivity usually is determined while illuminating the material with light of known spectral content (AM1.5, 100 mW/cm<sup>2</sup>). It has been shown [73] that the photoconductivity is expressed as:

$$\sigma_{ph} = e\mu\Delta n = \frac{e\eta_g\mu\tau(1 - R_f)F_0}{d} [1 - \exp(-\alpha(\lambda)d)] \quad (5)$$

with  $e$  the elementary charge,  $\mu$  the mobility of photogenerated majority carriers, i.e. electrons,  $\Delta n$  their density,  $\eta_g$  the quantum efficiency for generation of carriers,  $\tau$  their lifetime,  $R_f$  and  $d$  the reflectance and thickness of the material under study,  $F_0$  the illumination intensity (photons/cm<sup>2</sup>/s), and  $\alpha$  the wavelength-dependent absorption coefficient. For good intrinsic *a*-Si:H one usually finds  $\sigma_{ph} = 10^{-5} \Omega^{-1}\text{cm}^{-1}$  or higher, which is 5-6 orders of magnitude higher than the dark conductivity. One uses the photoresponse (or photosensitivity)  $\sigma_{ph}/\sigma_d$  as a measure of optoelectronic quality.

Besides  $\sigma_{ph}$  and the photoresponse the quantum efficiency-mobility-lifetime product,  $\eta_g\mu\tau$ , is used as a figure of merit. Usually this product is measured at a wavelength of 600 nm, and typical values of  $(\eta_g\mu\tau)_{600}$  are  $10^{-7}$  cm<sup>2</sup>/V or higher.

#### I.B.4. Optical properties

The optical properties of *a*-Si:H are of considerable importance, especially for solar-cell applications. Because of the absence of long-range order, the wavenumber  $\vec{k}$  is not a proper quantum number and it is not conserved in electronic transitions. Therefore, in contrast to crystalline silicon, *a*-Si:H behaves as though it has a direct band gap. Its absorption coefficient for visible light is about an order of magnitude higher than that of *c*-Si [74]. Consequently, the typical thickness (sub- $\mu$ m) of an *a*-Si:H solar cell is only a fraction of that of a *c*-Si cell.

In general one can divide the absorption behaviour in three ranges:

1. For the absorption coefficient  $\alpha$  larger than  $10^3$  cm<sup>-1</sup>, absorption takes place between extended states and is described by [75]:

$$(\alpha(E)n(E)E)^{1/1+p+q} = B(\hbar\omega - E_g) \quad (6)$$

with  $\alpha(E)$  the energy ( $E$ ) dependent absorption coefficient,  $n(E)$  the refractive index,  $p$  and  $q$  constants related to the shape of the band edges,  $B$  a proportionality constant,  $\hbar$  Planck's constant, and  $\omega$  the frequency. Tauc has argued that the density of states near the band edges has a square-root dependence on energy, as is the case for crystalline semiconductors [75]. This results in  $p = q = 1/2$ . Thus, extrapolating  $(\alpha(E)n(E)E)^{1/2}$  versus the photon energy to  $\alpha(E) = 0$  for  $\alpha(E) \geq 10^3$  cm<sup>-1</sup> yields the Tauc gap. Klazes *et al.* [76] have proposed the density of states near the band edges to be linear, i.e.  $p = q = 1$ . Evaluating  $(\alpha(E)n(E)E)^{1/3}$  versus  $E$  yields a so-called cubic gap, which is about 0.1-0.2 eV lower than the Tauc gap. The cubic plot is linear in a larger energy range than the Tauc gap, and fitting yields more accurate values. Nevertheless, the Tauc gap is still widely used. For intrinsic *a*-Si:H the Tauc gap typically is 1.7 eV or lower. The cubic gap is 1.6 eV or lower. The refractive index at 600 nm is around 4.3, the absorption coefficient  $4 \times 10^4$  cm<sup>-1</sup>.

2. In the range of  $1 - 10 < \alpha < 10^3$  cm<sup>-1</sup> absorption takes place between subbandgap states. An exponential dependence of  $\alpha$  exists [23]:

$$\alpha = \alpha_0 \exp\left(\frac{E}{E_0}\right) \quad (7)$$

with  $\alpha_0$  a prefactor, and  $E_0$  the so-called Urbach energy. This reciprocal logarithmic slope of the edge,  $E_0$ , depends both on temperature and the disorder in the material. The slope of the absorption edge mainly is determined by the slope of the valence band tail. Typically,  $E_0$  amounts to 50 meV. The slope of the conduction bandtail is about half this value [77].

3. In the low-energy range  $\alpha$  depends on defect density, doping level, and details of the preparation process. Sensitive subbandgap spectroscopy is used to measure  $\alpha$  and relate it to the defect density in the material [78,79].

The optical properties of *a*-Si:H are influenced both by the hydrogen concentration and bonding in the film, and by the disorder in the *a*-Si:H network [80]. A linear relation between band gap and hydrogen concentration is reported to be [81]:  $E_g = 1.56 + 1.27C_H$ . However, Meiling *et al.* [63,82] have identified three regions in the relation between  $E_g$  and  $C_H$ : the band gap first increases linearly as a function of hydrogen content, up to a value of about 0.1. A further increase up to  $C_H \sim 0.22$  does not lead to a change in the band gap. For higher values of  $C_H$  the band gap further increases linearly, but with a smaller slope than in the first region. This is explained as follows: in the first region ( $C_H < 0.1$ ) hydrogen is present only in monohydride bonds. The microstructure parameter is low, hence there are no voids in the material [62,83]. In the middle region ( $0.1 < C_H < 0.22$ ) the increasing amount of hydrogen is no longer incorporated as isolated SiH bonds. The microstructure parameter increases from 0 to 0.6, along with a reduction in silicon density. The material in this region consists of two phases, one which contains only SiH bonds, and one with SiH<sub>2</sub> bonds (chains) and voids. In the third region ( $C_H > 0.22$ ) the material mainly consists of chains of SiH<sub>2</sub> bonds, and the material density is much lowered.

Berntsen *et al.* [84,85] have separated the effect of hydrogen content and bond-angle variation. The structural disorder causes broadening of the valence and conduction bands and a decrease of the band gap by 0.46 eV. Hydrogenation to 11 at.% results in an independent increase of the band gap with 0.22 eV.

For undoped *a*-Si:H the (Tauc) energy gap is around 1.6-1.7 eV, and the density of states at the Fermi level is typically  $10^{15} \text{ eV}^{-1}\text{cm}^{-3}$ , less than one dangling bond defect per  $10^7$  Si atoms. The Fermi level in *n*-type doped *a*-Si:H moves from midgap to approximately 0.15 eV from the conduction band edge, and to approximately 0.3 eV from the valence band edge for *p*-type material [32,86].

### I.B.5. Metastability

An important drawback of *a*-Si:H is its intrinsic metastability: the electronic properties degrade upon light exposure. This was discovered by Staebler and Wronski [87,88], and is therefore known as the Staebler-Wronski effect (SWE). This effect manifests itself by an increase in the density of neutral dangling bonds,  $N_{db}$ , upon illumination, according to  $N_{db}(t) \sim G^{0.6}t^{1/3}$ , where  $G$  is the generation rate and  $t$  the illumination time [89]. The excess defects are metastable as they can be removed by annealing the material at temperatures above  $\sim 150$  °C for some hours. The presence of these excess defects to levels upto  $10^{17} \text{ cm}^{-3}$  leads to a reduction of free carrier lifetime, and hence to a lower conversion efficiency for solar cells.

The SWE is an intrinsic material property as the SWE is also observed in very pure *a*-Si:H, with an oxygen concentration as low as  $2 \times 10^{15} \text{ cm}^{-3}$  [90]. At impurity concentrations above  $10^{18} \text{ cm}^{-3}$  a correlation between SWE and impurity level has been established [91]. The hydrogen concentration, the bonding structure of hydrogen to the silicon network, and the disorder in the silicon network together affect the SWE. It has been long assumed that the origin of the SWE was local. Stutzmann *et al.* [89] have

proposed a model in which photogenerated charge carriers non-radiatively recombine at weak silicon-silicon bonds (e.g., strained bonds). The release of the recombination energy can be used to break the bond, and the thus formed two dangling bonds are prevented to recombine by the passivation of one of the bonds by a back-bonded hydrogen atom. Recent experimental results contradict the local nature of this model. Electron Spin Resonance (ESR) experiments have revealed that there is no close spatial correlation between the presence of hydrogen and light induced dangling bonds [92]. Further, it has been found that the rate of dangling bond creation is independent of temperature, in the range of 4 to 300 K [93]. As hydrogen is immobile at 4 K, diffusion can be ruled out.

The SWE has been found to depend on the hydrogen microstructure. The amount of hydrogen bound to silicon in the dilute phase, i.e., monohydride bonds, determines the saturated density of metastable defects [94]. In addition, only in regions of low hydrogen density light-induced defects have been found in ESR experiments [92], where it may well be that in these regions only monohydride bonds up to the solubility limit of 2 - 4 % of hydrogen in silicon [69,95] are present. The presence of clustered hydrogen (high microstructure parameter  $R^*$ ) leads to faster defect creation kinetics [96,97]. These and other observations have led to the realization that the SWE extends over large regions, in as much that configurational defects are transported over long distances. Branz proposed the hydrogen collision model [98]: the defect-creating recombination takes place at a Si-H bond, and the recombination energy is used to lift the hydrogen to a mobile energy level. The mobile hydrogen atom diffuses interstitially through the material. This mobile hydrogen atom is represented by a mobile complex of a Si-H bond and a dangling bond [99]. As the complex diffuses through the silicon network it breaks one Si-Si bond after another, but each broken bond is reformed after the complex has passed it. If two mobile hydrogen complexes collide, an immobile metastable two-hydrogen complex is formed. The net result is that two dangling bonds are created that are not spatially correlated to hydrogen atoms.

### I.B.6. Alloys

The glow-discharge technique is especially suitable for controlling various material properties by introducing other precursor gases in the plasma. As was first demonstrated by Anderson and Spear, incorporation of carbon or nitrogen in *a*-Si:H results in material with a high band gap [100]. A linear relation between band gap and C carbon fraction ( $x_C$ ) in *a*-SiC:H has been reported [101,102]:  $E_g = 1.77 + 2.45x_C$ . On the other hand, by diluting the plasma with germane ( $\text{GeH}_4$ ), material is obtained with low band gaps [103]: from 1.7 eV (*a*-Si:H) to 1.0 eV (*a*-Ge:H). Here, also a linear relation between band gap and Ge fraction ( $x_{Ge}$ ) has been reported [104], in which in addition the hydrogen fraction ( $x_H$ ) is included:  $E_g = 1.6 + x_H - 0.7x_{Ge}$ .

The incorporation of elements as carbon, nitrogen or germanium however, leads to material with a low mobility and lifetime of charge carriers. This would limit the application of these alloys, and a large research effort has been undertaken to find ways around this problem.

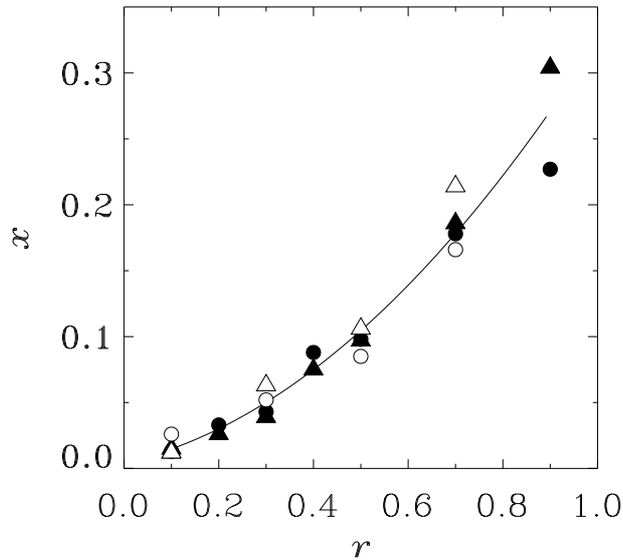


FIG. 2. The dependence of the carbon fraction  $x = [C]/([Si] + [C])$  on the gas-flow ratio  $r = [CH_4]/([SiH_4] + [CH_4])$  for films deposited in the ASTER system, AST1 (filled circles) and AST2 (filled triangles), and for films deposited in a similar system (ATLAS), ATL1 (open circles) and ATL2 (open triangles). (From R. A. C. M. M. van Swaaij, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1994).

The addition of an extra feedstock gas ( $GeH_4$  or  $CH_4$ ) in relatively large quantities (certainly compared to dopant gases) adds an extra degree of freedom to the already complex chemistry of the discharge. It is therefore even more complicated to relate material properties to deposition conditions [105].

As an example,  $GeH_4$  is less stable than  $SiH_4$  in the glow discharge, and Ge is preferentially deposited from  $SiH_4/GeH_4$  mixtures: a low  $GeH_4$  fraction in the gas already results in a high Ge fraction in the solid. Moreover, optimum conditions for  $a$ -Si:H deposition (low power density) cannot simply be translated to  $a$ -Ge:H deposition, in which case High power density is required to obtain good quality material [106]. The deposition of  $a$ -SiGe:H requires elaborate fine-tuning of deposition parameters. Other possibilities have been pursued to improve the properties of  $a$ -SiGe alloys, such as the use of strong dilution of  $SiH_4/GeH_4$  mixtures with hydrogen [107], and the use of fluorinated reactants such as  $SiF_4$  [108]. Modulation of the discharge (see also Sec. VIII C) has been demonstrated to increase the amount of Ge in the alloy with a factor of 2-10, depending on discharge conditions, if compared to a continuous discharge [109].

As  $SiH_4$  is less stable than  $CH_4$ , a low C fraction in the solid is obtained for a high  $CH_4$  fraction in the gas. This is illustrated in Fig. 2, which shows the dependence of the carbon fraction  $x = [C]/([Si] + [C])$  on the gas-flow ratio  $r = [CH_4]/([SiH_4] + [CH_4])$  for films deposited in the ASTER system and a system with a similar reactor (ATLAS) [101,102,110]. These films were Deposited in the so-called ‘low-power’ deposition regime [111,112], which is defined as the regime in which the applied power density is lower than the threshold power density demanded for the decomposition of methane [112]. In this regime the deposition of films is dominated by the decomposition of silane and is not dependent on the methane concentration [111,112]. Incorporation of carbon results from chemical reactions between methane molecules and silane species which are created by the plasma. The deposition rate in this regime is almost

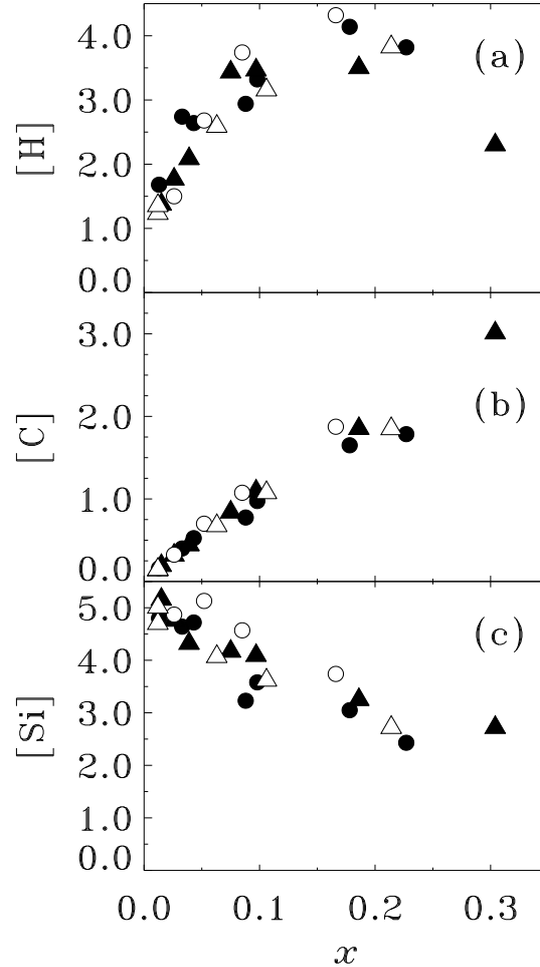


FIG. 3. Absolute atomic concentrations in units of  $10^{22}$  at./ $\text{cm}^3$ , determined by ERD, RBS and optical reflection and transmission spectroscopy, of (a) hydrogen, (b) carbon and (c) silicon as a function of the carbon fraction  $x$ . Results are presented for the series AST1 (filled circles), AST2 (filled triangles), ATL1 (open circles) and ATL2 (open triangles). (From R. A. C. M. M. van Swaij, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1994).

independent of the gas-flow ratio  $r$ , provided the gas in the reactor is not depleted in the discharge [101,102,113].

The dependence of the absolute atomic concentration of hydrogen, silicon, and carbon on the carbon fraction is shown in Fig. 3 [101,102]. The atomic concentrations were determined by using Elastic Recoil Detection (ERD) and Rutherford Backscattering Spectrometry (RBS) [114]. The most striking observation is the rapid increase of the hydrogen concentration upon carbon alloying (Fig. 3a), which can be ascribed to the incorporation of  $\text{CH}_n$  groups in the material during deposition [112,115]. It can be inferred from the data that up to a carbon fraction of about 0.1, three hydrogen atoms are incorporated per carbon atom. Above  $x = 0.15$  the rate of increase of the hydrogen concentration becomes smaller. For higher  $x$  values the hydrogen concentration even tends to decrease.

In Fig. 3b and 3c the absolute atomic concentrations of carbon and silicon, respectively, are shown as a function of the carbon fraction. As expected, the carbon con-

centration increases upon alloying. In contrast, the silicon content decreases rapidly, which implies that the material becomes less dense. As it was reported that the Si-Si bond length does not change upon carbon alloying [116] it thus can be inferred that the  $a$ -SiC:H material contains microscopic voids, which is in agreement with small-angle X-ray scattering (SAXS) results [62].

Much more can be said about Ge- and/or C-alloying, but this is not within the scope of this chapter. We refer to the books by Kanicki [5], and Luft and Tsuo [6] for further reading.

### I.B.7. Device quality characteristics

Over the past decades the term *device quality* has evolved to denote intrinsic PECVD hydrogenated amorphous silicon that has optimum properties for application in a certain device. Of course, depending on the type of device different optimum values are required, nevertheless the properties as listed in Table 1 are generally accepted, e.g. [6,11]. Many of these properties are interrelated, which has to borne in mind, when attempting to optimize only one of them.

Optimum properties for  $p$ - and  $n$ -type doped  $a$ -Si:H have been identified. For  $p$ -type doping boron is used as dopant element. Due to alloying with silicon, the band gap is reduced. This can be compensated by adding carbon. Typically silane ( $\text{SiH}_4$ ), methane ( $\text{CH}_4$ ), and diborane ( $\text{B}_2\text{H}_6$ ) are used, with silane and carbon in about equal amounts and diborane a factor of thousand lower. This yields  $p$ -type material with  $E_g = 2.0$  eV,  $E_A = 0.5$  eV,  $\sigma_d = 10^{-5} \Omega^{-1}\text{cm}^{-1}$ , and  $\alpha_{600} = 10^4 \text{ cm}^{-1}$ . Adding phosphine ( $\text{PH}_3$ ) to silane in a ratio 0.025:1 yields  $n$ -type material of good quality:  $E_g = 1.8$  eV,  $E_A = 0.3$  eV,  $\sigma_d = 10^{-3} \Omega^{-1}\text{cm}^{-1}$ , and  $\alpha_{600} = 4 \times 10^4 \text{ cm}^{-1}$ .

TABLE 1. Selected properties of *device quality* hydrogenated amorphous silicon films.

property	symbol	value	unit
optical bandgap (Tauc)	$E_g$	1.8	eV
optical bandgap (cubic)	$E_g$	1.6	eV
refractive index	$n$	4.3	
absorption coefficient (600 nm)	$\alpha_{600}$	$4 \times 10^4$	$\text{cm}^{-1}$
Urbach energy	$E_0$	50	meV
dark conductivity	$\sigma_d$	$10^{-10}$	$\Omega^{-1}\text{cm}^{-1}$
photo conductivity (AM1.5)	$\sigma_{ph}$	$10^{-5}$	$\Omega^{-1}\text{cm}^{-1}$
activation energy	$E_A$	0.8	eV
mobility-lifetime product	$\mu\tau_{600}$	$10^{-7}$	$\text{cm}^2\text{V}^{-1}$
hydrogen content	$C_H$	8-12	%
microstructure factor	$R^*$	0-0.1	
HWHM of TO Raman peak	$\Gamma/2$	33	$\text{cm}^{-1}$
intrinsic stress	$\sigma_i$	400-500	MPa
defect density	$N_s$	$10^{15}$	$\text{cm}^{-3}\text{eV}^{-1}$

## II. RESEARCH AND INDUSTRIAL EQUIPMENT

### II.A. General aspects

A plasma deposition system usually consists of several subsystems, each providing different functions [117]. The gas handling system includes process gas storage in high-pressure cylinders, mass flow controllers to measure and control the different gases to the reactor, and tubing. The vacuum system comprises pumps and pressure controllers. The plasma reactor is operated between 1 mTorr and 10 Torr\*. A much lower background pressure in the Ultra High Vacuum (UHV) range ( $10^{-9}$  mbar) is often required to ensure cleanliness of the process. High vacuum rotation pumps are used in combination with turbomolecular pumps [118].

The deposition set-up as shown in Fig. 4a is the central part of the most commonly used planar diode deposition system. The power to the reactor system is delivered by means of a power supply connected to the reactor via appropriate DC or RF circuitry (matchboxes). Power supplies can consist of generator and amplifier combined in one apparatus, with a fixed RF-frequency. More flexible is to have an RF-generator coupled to a broadband amplifier [119,120].

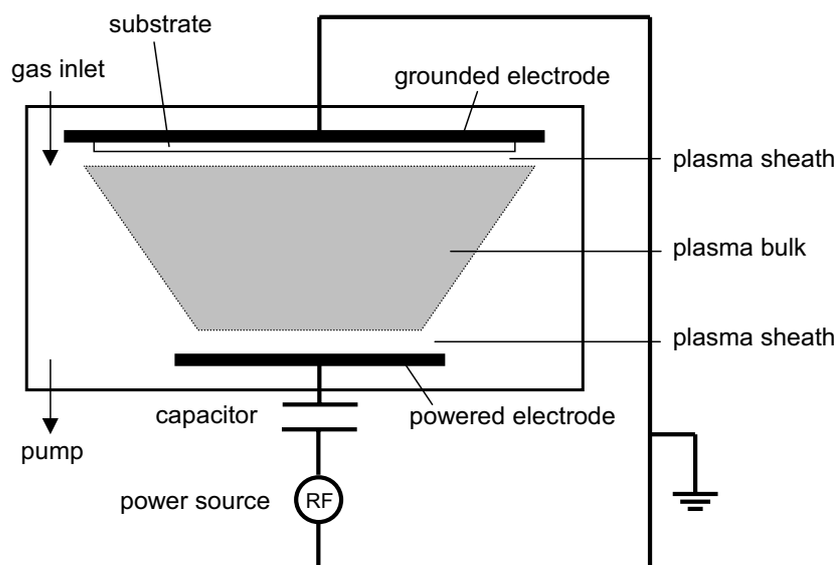
Using the planar diode geometry *p-i-n* devices have been made in single-chamber reaction systems, either with [121] or without loadlock [122]. In such single-chamber systems cross-contamination occurs. After deposition small amounts of (dopant) gases will remain in the reactor due to adsorption on the walls. If no load-lock is used, water vapour and oxygen are also adsorbed upon sample loading. These residual gases will desorb during subsequent depositions and may be built in as impurities in the amorphous layer [123,124]. Additionally, the desorption of gases after a deposition may cause contaminated interfaces, e.g., a graded boron concentration profile may be present in the first monolayers of the *i*-layer of a  $p^+i$  interface [125,126]. Contaminated interfaces combined with (too) high impurity levels in the layers have a negative influence on the electrical properties of the layer [127,128], which results in a low efficiency of solar cells.

By using a multichamber system [129] exchange of residual gases between successive depositions will strongly be decreased and very sharp interfaces can be made. Furthermore, the use of a load-lock system ensures the quality of the background vacuum, and thus the low levels of contaminants in the bulk layers. Multichamber reactor systems have been used for the fabrication of solar cells and considerable improvements in energy conversion have been achieved [130,131].

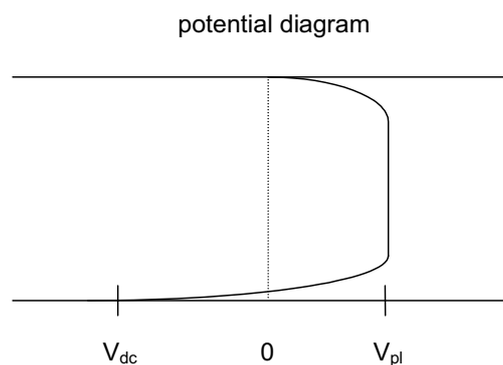
Most of the gases used are hazardous, i.e., they can be corrosive, flammable, explosive, and/or highly toxic. Silane is pyrophoric, the dopant gases diborane, phosphine, and arsine are extremely toxic, with TLV values in the low-ppm range. Therefore extreme care has to be taken in dealing with these gases. One generally installs a.o. flow limiters (between the valve of the supply cylinder and pressure regulator to prevent excessive gas flow in case of breakdown of the regulator), cross-purge assemblies (in order

---

\*Throughout this chapter units of pressure Torr, mbar, and Pa are used, without converting them into the S.I. unit Pa. Rather the original data are used. Note that  $1 \text{ Pa} = 0.01 \text{ mbar} = 0.0076 \text{ Torr} = 7.6 \text{ mTorr}$ .



(a)



(b)

FIG. 4. Schematic representation (a) of a parallel-plate, capacitively coupled RF-discharge reactor, with unequally sized electrodes. The potential distribution (b) shows the positive plasma potential  $V_{pl}$ , and the negative DC selfbias voltage  $V_{dc}$ .

to purge the regulators and prevent release of the gas when exchanging (empty) gas cylinders, scrubbers or diluters to avoid above-TLV levels at the exhaust, and detectors mounted at several critical locations. More on safety issues can be found in, e.g., the Proceedings of the 1988 Photovoltaic Safety Conference [132].

In the design of the reactor chamber one needs to address issues such as electrode geometry, gas flow patterns, heater design, reactor volume (discharge and discharge free regions). Hot walls will influence the gas temperature, and gas phase chemistry. Electrode size and electrode spacing as well as dark-space shields directly influence the discharge properties. Asymmetry introduces a bias, which may can be controlled

externally. Also the driving frequency is an important parameter. Over the years many reactor designs have been developed. Studies on the effects of process parameters on material quality not always take geometric causes into account. This has prompted the development of a standard type of reactor, i.e., the GEC reactor cell [133]. Nevertheless generalizations of the effect of process parameters on material quality have been made. For an overview see, e.g., Luft and Tsuo [6].

## II.B. Reactor configurations

Diode reactors can be powered by an RF or DC electric field. In case of RF excitation the deposition typically is on the grounded electrode. As the combined area of grounded electrode and reactor walls usually is larger than the area of the powered electrode, a DC self bias is developed, as shown in Fig. 4b. The potential drop at the powered electrode is much larger than at the grounded electrode. The powered electrode is more negative with respect to ground, and therefore is often called the cathode. The grounded electrode then is the anode. For a DC glow discharge, the potential distribution is similar to the one shown for the RF discharge in Fig. 4b. In both cases ion bombardment at the cathode is larger than at the anode. Due to this, deposition of films on the cathode or anode leads to different microstructural properties. Deposition rates at the cathode are usually higher than at the anode. Films deposited at the cathode are dense, but also stressed. Anodic films are more porous. One sometimes uses a mesh in DC discharges positioned above the cathode, which is polarized at the same potential as the cathode [134]. Ions are thus slowed down by gas-phase collisions in the region between mesh and cathode, and much better material is obtained [135]. Moreover, the mesh acts as a screen for reactive radicals. The  $\text{SiH}_2$  radical has a large sticking probability, and it will stick to the mesh easily. As a consequence, the  $\text{SiH}_2$  radical will be filtered out, and  $\text{SiH}_3$  will dominate the deposition. The resulting material quality has been shown in RF triode discharges to have been improved [136–139].

Planar triode RF discharges used in research generally are asymmetric; the area of the powered electrode is much smaller than the area of all grounded parts taken together (the grounded electrode may only be just a small part of the grounded area). The DC self bias therefore is large. One can reduce the asymmetry by confining the discharge with a grounded mesh [140,141] or wall [142]. Such confinement also allows for a higher power density in the discharge, which leads to enhanced deposition rates.

An external magnetic field has also been used to confine the plasma [143]. An arrangement where electromagnets are located under the cathode is known as the controlled plasma magnetron method [144]. The diffusion of electrons to the walls is prohibited by the magnetic field between cathode and anode. This results in an increase in electron density, and consequently in a faster decomposition of silane and a higher deposition rate. At a deposition rate of 1 nm/s device quality material is obtained [144]. In addition, a mesh is located near the anode and the anode can be biased externally, both in order to confine the plasma and control ion bombardment.

Hot reactor walls are sometimes used as a means to increase the density of the films that are deposited on the walls. This reduces the amount of adsorbed contaminants on the walls, and leads to lower outgassing rates. A hot wall is particularly of interest for single-chamber systems without a load-lock chamber. Material quality is similar to the

quality obtained with a cold reactor wall [145].

Other configurations that are used are an concentric electrode set-up in a tubular reactor, where the discharge still is capacitively coupled. Also an inductive coupling has been used, with a coil surrounding the tubular reactor [146,147].

### II.C. Scale-up to systems of industrial size

A plasma process that has been demonstrated to yield good quality materials in the laboratory one day will need to be scaled up to a technology that can produce the particular material in larger sizes and larger quantities. Such a transfer is not straightforward, and many technological difficulties will have to be overcome before a scaled-up process is commercially viable.

A plasma process is characterised by many parameters and their interrelations are very complex. It is of paramount importance to understand, at least to a first approximation, how the plasma parameters have to be adjusted when the geometrical dimensions of the plasma system are enlarged. Especially of use in scaling up systems are scaling laws, as formulated by Goedheer *et al.* [148,149] (see also Sec. III B 2).

In general, the substrate temperature will remain unchanged, while pressure, power, and gas flow rates have to be adjusted such that the plasma chemistry is not affected significantly. Grill [117] conceptualizes plasma processing in two subsequently occurring processes, i.e., the formation of reactive species, and the mass transport of these species to surfaces to be processed. If the dissociation of precursor molecules can be described by a single electron collision process, the electron impact reaction rates depend only on the electric field over pressure ratio,  $E/p$ , because the electron temperature is determined mainly by this ratio. The mass transport is pressure dependent, because the product of diffusion constant and pressure,  $Dp$ , is constant. Hence, preservation of both plasma chemistry and mass transport is obtained by keeping both  $E$  and  $p$  constant during scale-up. As the area of the scaled-up reactor is larger (typically about  $1 \text{ m}^2$ ) than its laboratory equivalent ( $1\text{-}100 \text{ cm}^2$ ), the total power or current supplied to the discharge must be enlarged in order to keep  $E$  constant. For a parallel plate reactor the power simply scales linearly with the area increase, while for other configurations this will not be a simple relation, because the plasma is not confined between the electrodes. Note, that changing the area ratio of grounded to powered electrode will affect the DC self bias voltage, and ion bombardment. The gas flow rate is the only external parameter that needs adjustment. It should be scaled up such that the average flow velocity is identical for laboratory- and industrial scale reactors. Alternatively, one can keep the average residence time constant, which is defined as  $\tau_r = pV/Q$ , with  $V$  the volume of the reaction zone and  $Q$  the total mass flow rate. The residence time is a measure of the distance over which the reactive species diffuse in the reaction zone. With the pressure being constant, the requirement of constant residence time scales the gas flow rate proportional to the reaction volume.

The scale-up from a small to a large plasma reactor system requires only linear extrapolations of power and gas flow rates. However, in practice, the change in reactor geometry may result in effects on plasma chemistry or physics that were unexpected, due to a lack of precise knowledge of the process. Fine-tuning, or worse, coarse readjustment is needed, which is mostly done empirically.

A critical issue in scaling-up of a process is the uniformity in deposition rate and material quality. In general, once the deposition rate is constant within 5 % over the whole substrate area the material properties also do not vary much. After fine-tuning the power and gas flow rates operators still may face non-homogeneity issues. These can be caused by local changes in temperature, RF voltage, and gas composition, due to various causes. As an example, it has been reported that improper attachment of the substrate to the grounded electrode results in a local decrease of the deposition rate [150,151].

Low contamination levels are more or less intrinsic in laboratory scale UHV systems. Very high costs inhibits the use of UHV in industrial scale systems, however another "local-UHV" approach has been proposed, i.e., the plasma box reactor [152]. The substrate is mounted in a box, that is surrounded by a shell, which is pumped to a low pressure. The process pressure in the box is maintained by a throttle valve. As the pressure in the box is larger than the pressure in the surrounding shell contaminants diffuse outwards and the incorporation of contaminants in the deposited layer is low.

Schropp and Zeman [11] have classified current production systems for amorphous silicon solar cells. They argue that cost-effective production of solar cells on a large scale requires that the product of deposition time needed per square meter and the depreciation and maintenance costs of the system is to be small. Low deposition rates must be accompanied by low costs. The costs requirement is the main drive for ongoing investigations on the question how to increase the deposition rates while maintaining device quality material. Current production system configurations can be divided into three classes: 1) single chamber systems, 2) multichamber systems, and 3) roll-to-roll systems [11].

In single chamber systems there is no transport of substrates under vacuum conditions, which makes the system simple, but contamination levels may be relatively high. The addition of a load-lock lowers contamination levels, and requires only a linear transport mechanism. During the deposition process temperature and geometry cannot be changed easily, while changes would be required for different layers in the solar cell. Substrates are loaded consecutively, and for every substrate a complete pump-down and heating is needed. The actual deposition time is much smaller than the total processing time. A higher throughput can be achieved by loading substrates in cassettes, as is general practice in semiconductor industry. The investment is considered low, as well as the flexibility of the system.

Industrial scale multichamber systems offer much of the advantages that are characteristic of laboratory scale systems. Typically, the deposition of each layer in a solar cells is performed in a separate chamber, and process parameter optimizations can be done for each layer individually. However, this high flexibility comes with high investment costs. Two types of multichamber systems can be distinguished, a cluster configuration, and an in-line configuration. The cluster configuration more or less is the enlarged copy of a laboratory scale system, and offers the highest flexibility. Transport and isolation chambers can be shared by many deposition chambers, and, depending on the actual configuration, production can be continued even while some reactors are down. A cassette of substrates (batch) can be processed completely automatically, which increases throughput of the production system.

The in-line configuration consists of deposition chambers that are separated by isolation chambers [153]. The layer sequence of a solar cell structure prescribes the

actual sequence of deposition chambers. Flexibility is much less compared to a cluster configuration, and costs are generally much higher, but the throughput can also be much larger. In an in-line system the substrates can move while deposition takes place, which leads to very uniformly deposited layers, as only uniform deposition in one dimension (perpendicular to the moving direction) is required.

In the roll-to-roll configuration the substrate consists of a continuous roll, e.g. stainless steel [154], SnO<sub>2</sub>:F coated aluminium [155], or plastic [156–158], which is pulled through a sequence of deposition chambers.

#### II.D. ASTER, a research system

As a large part of experimental and simulation results as shown in this chapter are obtained with or specifically calculated for the UHV multichamber deposition system ASTER (Amorphous Semiconductor Thin film Experimental Reactor) at Utrecht University [14,15], it is described in this section. It also is a more or less generic parallel plate multichamber research system. Similar commercially available multichamber equipment has been developed and described by Madan *et al.* [159]. It has also been installed at Utrecht University and is named Process equipment for Amorphous Silicon Thin-film Applications (PASTA).

The ASTER system is intended for use in research and consists of three identical plasma reactors, a spare chamber and a load lock, all having the same outer dimensions. These chambers are connected radially via gate valves to a central transport chamber, in which a transport system or robot arm is located. All chambers are made of stainless steel, and all seals with valves, windows, gas-supply lines, and measuring devices are Conflat connections, ensuring the vacuum to be of UHV quality. The ASTER system was completely designed and partly built in-house, the chambers and transport arm were manufactured by Leybold companies. A schematic drawing is shown in Fig. 5, which illustrates the situation around 1990 [14,15]. The spare chamber was designed first to be used as an evaporation chamber, with which it was possible to evaporate contacts without exposing the sample to air. This was stopped for practical reasons. It was used then as a chamber where samples could be parked, to enhance productivity. As research shifted in the direction of plasma analysis, this fourth chamber was retrofitted with a quadrupole mass spectrometer [160]. After having built a separate system for plasma analysis in which the mass spectrometer was integrated [161], the fourth chamber now is in use for deposition of silicon nitride.

The central transport chamber is a 80 cm diameter stainless steel vessel, and is pumped by a 1000 l/s turbomolecular pump, which is backed by a small (50 l/s) turbomolecular pump to increase the compression ratio for hydrogen, and by a 16 m<sup>3</sup>/h rotating-vane pump. Ultra-high vacuum is obtained after a bake-out at temperatures above 100 °C (measured with thermocouples at the outside surface) of the whole system for about a week. A pressure in the low 10<sup>-11</sup> mbar range then is obtained. With a residual gas analyzer (quadrupole mass spectrometer, QMS) the partial pressures of various gases can be measured. During use of the system, the pressure in the central chamber is in the low 10<sup>-10</sup> mbar range due to loading of samples. Water vapor then is the most abundant species in the chamber.

The robot arm can transport a substrate holder for substrate sizes up to 10 × 10 cm<sup>2</sup>.

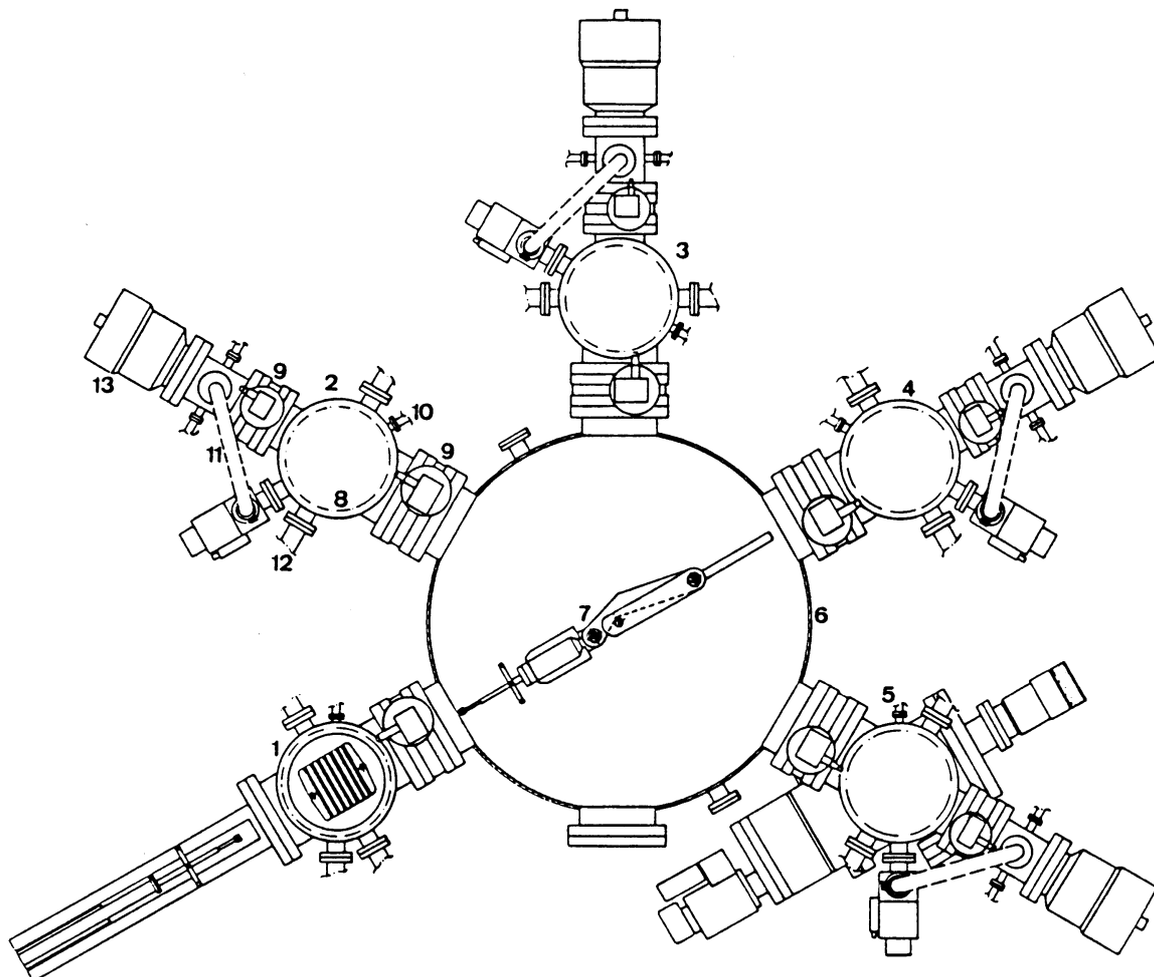


FIG. 5. Schematic representation of the ASTER deposition system. Indicated are: (1) loadlock, (2) plasma reactor for intrinsic layers, (3) plasma reactor for *p*-type layers, (4) plasma reactor for *n*-type layers, (5) metal-evaporation chamber (see text), (6) central transport chamber, (7) robot arm, (8) reaction chamber, (9) gate valve, (10) gas supply, (11) by-pass, (12) measuring devices, and (13) turbomolecular pump.

The arm can both rotate and translate, and is driven mechanically by two external DC motors, one for each movement. The translation mechanism was specially designed for fast transport of the samples. The mechanism is based on an eccentrically moving hinge-joint, much like a stretching human arm, and it combines a high speed with a shock-free start and stop of the movement. It takes only 4 seconds to translate the substrate holder from the transport arm to the outermost position in the reaction chamber. A gear transmission inside the transport chamber is used to decrease the torque to be applied to the feed-throughs.  $WSe_2$  is used as lubricant for the gear

wheels. The arm position is measured by potentiometers. The position of the arm in angular direction is reproducible to within 0.03 degrees of the preset position.

A clear advantage of the central transport chamber is that a sample can be transported to the reaction chambers in any arbitrary sequence, without breaking the vacuum in the reaction chambers. Due to the very low pressure in the central chamber, the vacuum in the reaction chambers in fact is improved, when a gate-valve is opened. After a complete stretch of the robot arm the substrate holder can be taken off from the arm by a lift mechanism and is clamped to the upper (grounded) electrode. Experienced operators are manually able to remove a substrate holder from one chamber and deliver it to any other within 30 seconds, including the time needed to open and close the gate valves. Automatic control by means of a programmable logic controller, coupled to a personal computer, only slightly shortens this transfer time. Because of the large heat capacity of the 12 mm thick titanium substrate holder and the short transfer time the cooling of the sample is limited to a few degrees. Fast transport and a low background pressure in the central chamber ensure that clean interfaces are maintained. Transport at  $10^{-10}$  mbar for 30 seconds will give rise to a surface contamination of only about 0.01 monolayer of oxygen.

New substrates are mounted on the substrate holder which then is loaded in the load-lock chamber, which subsequently is evacuated down to  $10^{-6}$ - $10^{-7}$  mbar with a 150 l/s turbomolecular pump in combination with a 25 m<sup>3</sup>/h rotating vane pump. In this chamber the substrates are pre-heated to the desired temperature for the deposition process, and at the same time they can be cleaned by DC argon sputtering. Usually the heating time (about 1 hour) is much longer than the pumpdown time (less than 10 minutes). The substrate is transferred to the central chamber by a sequence of events: opening of the gate valve, translating the arm, lowering the substrate holder with the lift mechanism, retracting the arm (now with substrate holder), closing the gate valve. The pumping capacity of the central chamber is such that the  $10^{-10}$  mbar background pressure is re-established within 10 seconds after closing the gate valve.

Each plasma reactor consist of a reaction chamber, an individual pumping unit, pressure gauges, and a dedicated gas-supply system with up to four different gases, i.e., according to the layer type to be deposited. Residual gas exchange is prevented due to the separated pumping units, as well as due to the presence of the central chamber. The background pressure in the plasma reactors is lower than  $10^{-9}$  mbar, with partial pressures of water vapour and oxygen lower than  $10^{-10}$  mbar, as a result of the capacity of the pumping unit, i.e., a 360 l/s turbomolecular pump backed with a 40 m<sup>3</sup>/h rotating-vane pump. The process pressure typically is between 0.1 and 0.5 mbar. This can be accomplished by closing the gate valve between reaction chamber and pumping unit (9 in Fig. 5). The chamber then is pumped only via the bypass (11 in Fig. 5), in which a butterfly valve regulates the process pressure.

A vertical cross-section of the reaction chamber is shown in Fig. 6. The inside diameter of the chamber amounts to 200 mm. The diameters of the grounded and powered electrodes are 180 mm and 148 mm, respectively, with a fixed interelectrode distance of 36.5 mm. The interelectrode distance has also been changed to 27 mm, and recently a modified powered electrode assembly has been retrofitted, with which it is possible to vary the interelectrode distance from 10 to 40 mm, from the outside, i.e., without breaking the vacuum [162]. With process pressures in the range of 0.1-0.6 mbar the product of pressure and interelectrode distance  $pL$  may range from about 0.1

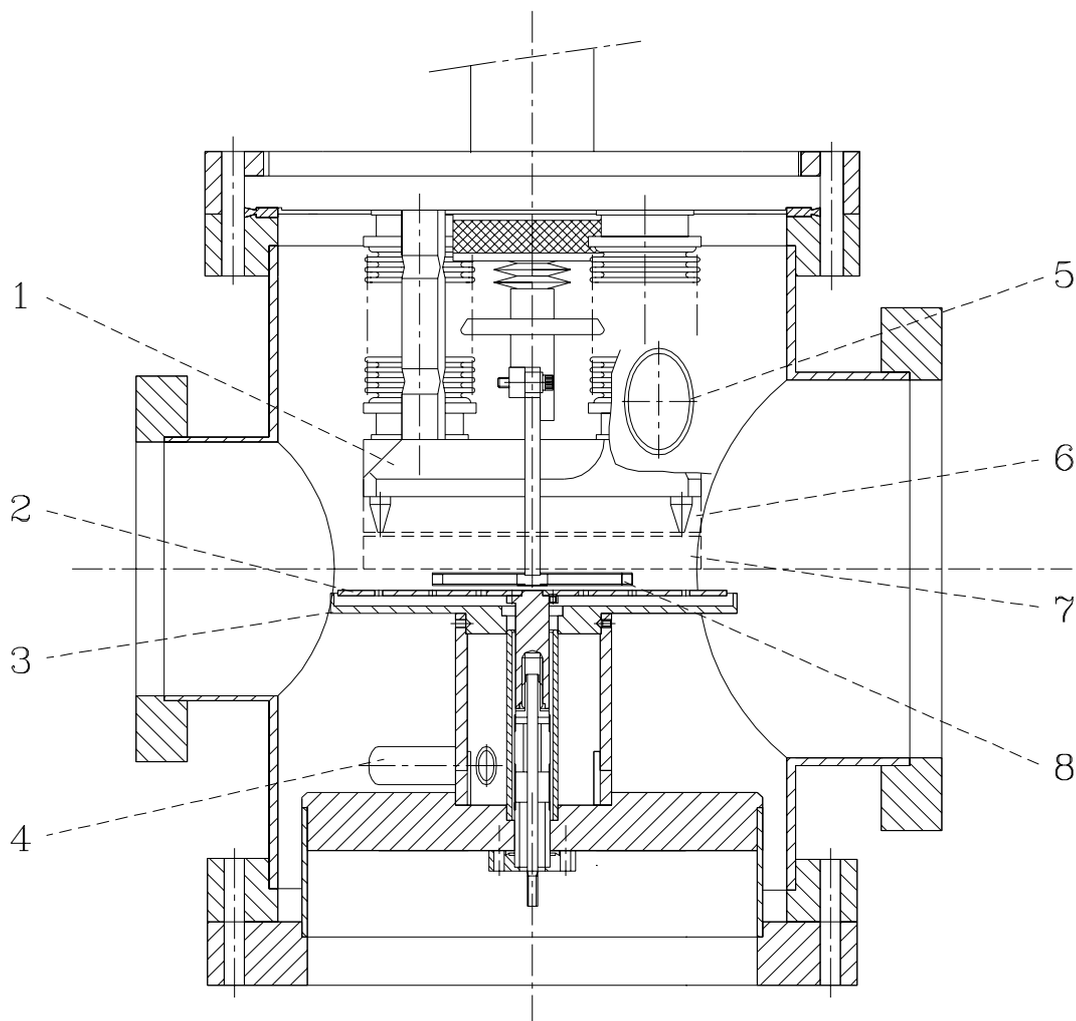


FIG. 6. Vertical cross-section of the reaction chamber. Indicated are: (1) the grounded electrode, (2) the RF electrode, (3) dark space shield, (4) the gas supply, (5) the gas exhaust, (6) position of the sample holder during deposition, (7) position of the sample holder when loaded, and (8) the lift mechanism.

to 6 mbar·cm. In practice,  $pL$  values are between 0.4 and 0.5 mbar·cm, i.e., around the Paschen law minimum (see Sec. III B 4).

The lower electrode is coupled via a  $\pi$ -type matching network to a 13.56 MHz generator. This network provides a power matching between the RF power cable (50  $\Omega$ ) and the plasma. Power levels are between 1 and 100 Watt, or between 6 and 600 mW/cm<sup>2</sup>, using the area of the powered electrode.

The substrate holder is positioned face down (6 in Fig. 6) to the upper electrode, thus deposition is upwards, which prevents dust formed in the plasma to fall onto the substrate. The plasma is confined between the powered and grounded electrode, and the reactor walls, which in addition can be water-cooled. The substrate temperature can be varied up to about 500 °C by means of fire-rods in the grounded electrode. The temperature is regulated by a PID-type temperature controller, which measures the temperature with a thermocouple, also mounted in the upper electrode. The actual

substrate temperature deviates from the temperature set by the controller, due to heat losses in the system. Therefore the actual substrate temperature has been measured by thermocouples on the substrate over a wide range of controller temperatures, and a calibration graph is used. Throughout this chapter, the actual substrate temperature is given.

The appropriate gas mixture can be supplied to the center of the reactor (4 in Fig. 6) via holes in the lower electrode (5 in Fig. 6), and is pumped out through the space between substrate electrode and the reactor wall to the exhaust (4 in Fig. 6). Alternatively, the gas mixture can be supplied horizontally, parallel to the electrodes, through a flange in the reactor wall, positioned between the electrodes (perpendicular to the plane of the cross-section in Fig. 6, not shown). In this case, the gas is pumped out at the opposite side of the supply.

The volume of the reactor is about 10 l. At a typical process pressure of 0.2 mbar and a total gas flow rate of 60 sccm, the average residence time of molecules in the reactor amounts to about 1.3 s.

Different gases experience different conductances towards the pump. The conductance is a combined result of the manually adjustable butterfly valve in the bypass line and the pumping system. This results in gas-dependent pressures for identical gas flows, as the flow  $Q$  is related to pressure  $p$  and conductance  $C_{gas}$  as  $Q = pC_{gas}$ . In Fig. 7 the pressures of pure argon, silane, and hydrogen are shown

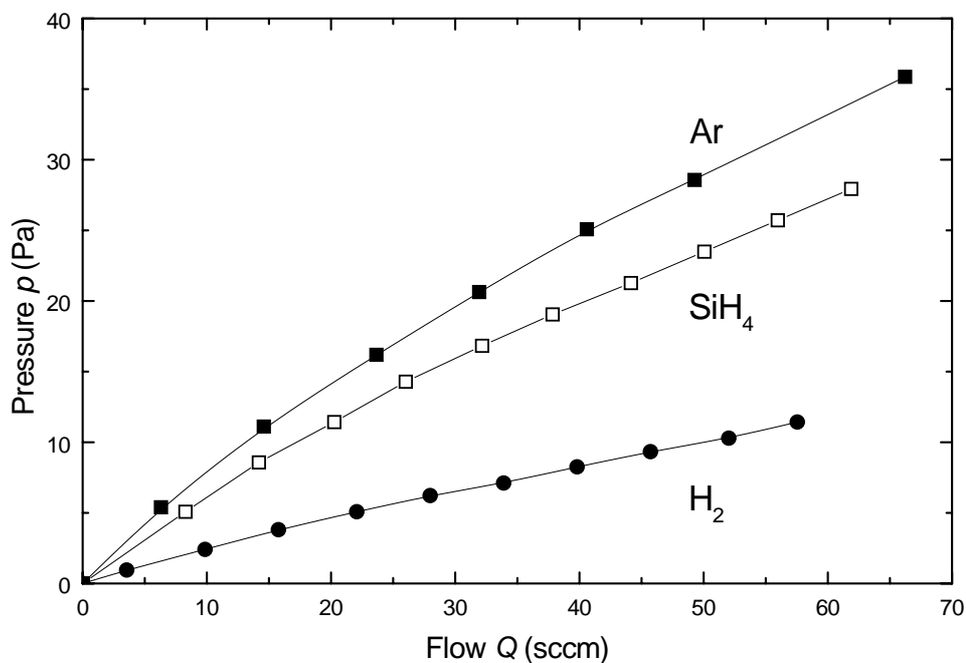


FIG. 7. The pressure  $p$  as a function of flow rate  $Q$  for different gases (Ar, SiH<sub>4</sub>, H<sub>2</sub>) at one and the same setting of the pressure regulating butterfly valve in the bypass (11 in Fig. 5). (From E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

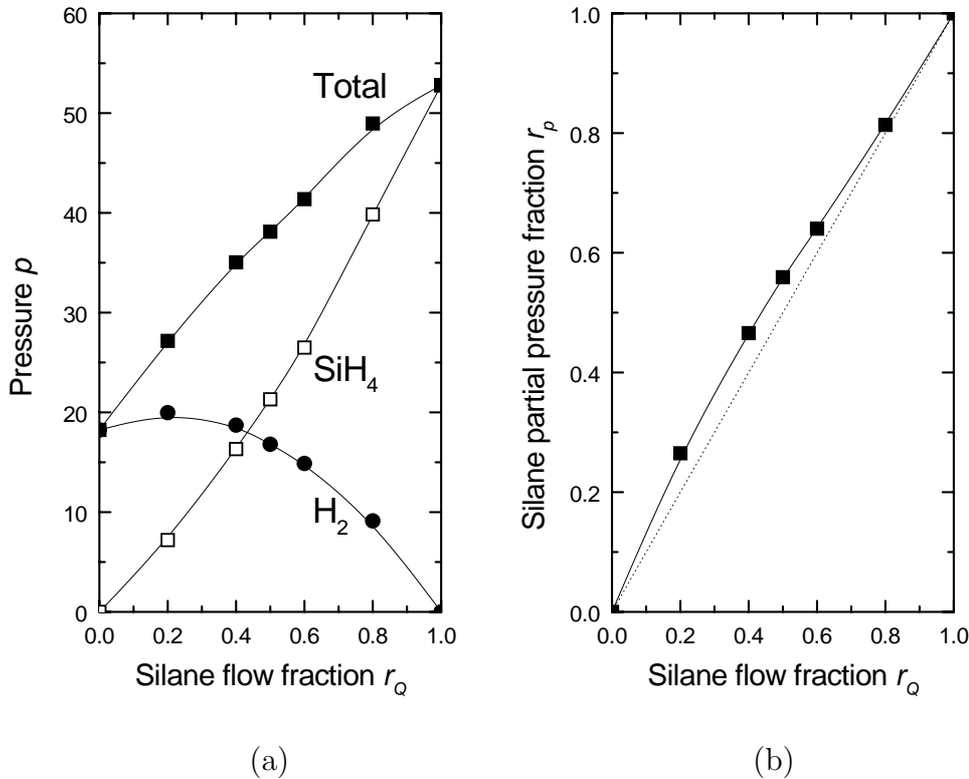


FIG. 8. The total and partial pressures  $p$  (a) the partial pressure ratios  $r_p$  (b) of silane and hydrogen in a silane/hydrogen mixture, at different flow ratios  $r_Q$ . The total flow rate is 30 sccm. (Adapted from E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

as a function of flow rate [163]. The setting of the butterfly valve was the same for all cases. As can be seen the conductance for hydrogen is about a factor of 2-3 larger than the one for argon and silane. From the fact that the rate  $dp/dQ = C^{-1}$  decreases with increasing flow, it is inferred that the flow of the gases leaving the reactor is a mixed laminar/molecular or Knudsen flow [118].

As a consequence of gas-dependent conductances, the composition of a gas mixture may differ from the one expected on the basis of flow ratios. As an example, the partial pressures (measured with QMS) of a range of silane/hydrogen mixtures at a total flow of 30 sccm is shown in Fig. 8a as a function of the flow fraction silane  $r_Q = Q_{\text{SiH}_4}/(Q_{\text{SiH}_4} + Q_{\text{H}_2})$  [163]. A flow of hydrogen yields a lower pressure (about 3 times lower) than the same flow of silane, cf. Fig. 7. The actual partial pressure ratios  $r_p = p_{\text{SiH}_4}/(p_{\text{SiH}_4} + p_{\text{H}_2})$  for these mixtures is shown in Fig. 8b. It is clear that equal flow rates of different gases (in this case hydrogen and silane) do not lead to equal partial pressure ratios. This is important when comparing experimental and simulation data.

Each of the reaction chambers has its own gas-supply system consisting of two channels of 3-5 m in length. It also includes a nitrogen purge system. One channel is connected to a gas manifold for three types of gases, the other channel is connected

to a single gas bottle. With this configuration it is possible to supply the reaction chamber with either a continuous flow of different gases or a mixture of gases, which may rapidly change during deposition (c.f. chemical annealing [164,165]). The gases are connected via pressure regulators to the supply channels. They can be switched to go via a mass flow controller to the reaction chamber or to go to a drain, which is continuously purged with argon and pumped with a 8 m<sup>3</sup>/h rotating-vane pump. When the pressure in the drain or the exhaust lines is too high, a safety exhaust is opened with a much larger diameter than the regular exhaust line from the reaction chamber. Nitrogen is available for pressurizing the system, and also is used to dilute the exhaust lines from the reaction chambers, to ensure below-TLV values on the roof of the building. If one needs to clean a reaction chamber, a separate channel with NF<sub>3</sub> is available. Etching with a NF<sub>3</sub> discharge has been found to reduce the amount of Si<sub>x</sub>H<sub>y</sub> compounds adsorbed on the wall considerably [166].

In the plasma reactor dedicated for intrinsic material deposition (2 in Fig. 5) only hydrogen and silane are used, as well as argon. A mixture of trimethylboron (5 % TMB in H<sub>2</sub>), SiH<sub>4</sub>, and methane (CH<sub>4</sub>) is used in the *p*-plasma reactor (3 in Fig. 5). Also diborane can be used. A mixture of phosphine (PH<sub>3</sub> (1 % in H<sub>2</sub>)) and SiH<sub>4</sub> is used in *n*-plasma reactor (4 in Fig. 5). All gases are of 6.0 quality (99.9999% pure), if available from manufacturers, otherwise as pure as possible.

The low background pressure (10<sup>-9</sup> mbar) together with the purity of the gases used ensures a low concentration of contaminants. Amorphous silicon films made in the intrinsic reactor have been analyzed by using Elastic Recoil Detection (ERD), which is available in our laboratory [114]. The determined oxygen content in these films typically is lower than 3×10<sup>18</sup> cm<sup>-3</sup>, which is somewhat lower than the values required for obtaining device quality films reported by Morimoto *et al.* [167].

It goes without saying that safety is a very important issue in working with such a set-up. A risk analysis has been performed. Safe operation procedures have been formulated. Personnel has been trained to work safely with the set-up, which includes a.o. exchanging gas bottles using pressurized masks. Interlocks are installed, as well as emergency switches. Emergency power is available, in case of power failures. Gases are stored in gas cabinets that were designed and installed with assistance of gas manufacturers. All valves in the gas-supply lines are normally-closed valves: if compressed air (6 bar) is not available, all valves are by definition closed. Valves for nitrogen supply are normally-open valves. Gate valves are double-action valves: compressed air is needed to either open or close the valve.

### III. PHYSICS AND CHEMISTRY OF PECVD

#### III.A. General introduction

A plasma can simply be defined as a partially ionized, quasineutral gas, consisting of about equal numbers of positive and negative charges, and a different number of not ionized neutral molecules. An external source of energy is needed to sustain the plasma for a sufficiently long time. The simplest and most widespread method that is used is the electrical discharge, DC or radiofrequent (RF). High electric fields applied at mbar-pressures yield non-equilibrium plasmas: free electrons are accelerated to 1-10 eV, while ions and neutrals have low energy (0.1 eV). These "hot" electrons initiate chemical reactions through collisions with the "cold" neutrals. Therefore, processing temperatures can be much lower than in thermal CVD, which has tremendous consequences for the applicability of PECVD. Without PECVD modern-day electronic chips would otherwise never have been possible. For general references see, e.g. Bruno [8], Grill [117], Chapman [134], and Ricard [168].

This section treats the plasma physics and plasma chemistry of the typical silane/hydrogen RF discharge, with occasionally some examples that employ a somewhat higher excitation frequency. Electrical characterization of the discharge is followed by an analysis of the silane chemistry. An appropriate set of gas phase species is presented, which are then used in the modeling of the plasma. A comparison is made between modeling results and experimental work in ASTER. Extension to 2D modeling is presented as well.

Plasma analysis is essential in order to compare plasma parameters to simulated or calculated parameters. From the optical emission of the plasma one may infer pathways of chemical reactions in the plasma. Electrical measurements with electrostatic probes are able to verify the electrical properties of the plasma. Further, mass spectrometry on neutrals, radicals, and ions, either present or coming out of the plasma, will elucidate even more of the chemistry involved, and will shed at least some light on the relation between plasma and material properties. Together with ellipsometry experiments, all these plasma analysis techniques provide a basis for the model of deposition.

#### III.B. Plasma physics

##### *III.B.1. Plasma sheath*

A schematic layout of typical parallel-plate RF-discharge system is depicted in Fig. 4a. The RF power is capacitively coupled to the discharge between the two electrodes. Silane is introduced between the electrodes, and reaction products and unreacted gas is pumped away from the reactor. The substrate onto which *a*-Si:H is deposited is mounted on the grounded (top) electrode (anode). The RF-voltage  $V(t) = V_{RF} \sin \omega t$  is applied to the cathode, with  $\omega/2\pi = 13.56$  MHz. This particular frequency has been chosen in the past, as radiating energy at this frequency would not interfere with communications [134]. However, higher harmonics do, and one should be careful in designing proper shielding.

Accelerated electrons in the applied electric field ionize gas molecules, and in these ionization processes extra electrons are created. In the steady-state the loss of charged

particles is balanced by their production. Due to their much lower mass, electrons move much faster than ions. As a result, charge separation creates an electric field, compensating the differences in electron and ion velocities. The potential in the central region, or bulk plasma, is slightly positive with respect to the electrodes, due to the small surplus of positive ions.

The time-averaged potential profile is shown in Fig. 4b. As ions cannot follow the oscillations in the applied electric field, it is this profile that ions experience. The bulk plasma is characterized by a constant potential,  $V_{pl}$ . In both sheaths, the regions between plasma bulk and the electrodes, the ions experience a potential difference, and are accelerated towards the electrodes. This yields energetic ion bombardment onto the electrodes. Electrons are expelled from the sheaths, so all ionization and dissociation processes must occur in the plasma bulk. Plasma light as a result from emission from excited molecules is only present from the plasma bulk; the sheaths are dark.

The electrons follow the oscillations in the electric field, and experience the time-dependent plasma potential. Due to the capacitor by which the RF power is coupled to the electrodes, no DC current flows through the plasma. The ion and electron current towards each of the electrodes balance each other within one RF period.

In most systems the substrate electrodes are larger than the powered electrodes. This asymmetric configuration results in a negative DC self bias voltage  $V_{DC}$  on the powered electrode. The difference in electrode areas would result in a net electron current per RF period. A negative DC shift prohibits this [134,169]. It has been shown that the ratio of the time-averaged potential drops for the sheaths at the grounded ( $V_{sg}$ ) and powered electrode ( $V_{sp}$ ) are inversely proportional to the ratio of the area of both electrodes ( $A_g, A_p$  [134,170–172])

$$\frac{V_{sg}}{V_{sp}} = \left(\frac{A_p}{A_g}\right)^n \quad \text{with } 1 < n < 4 \quad (8)$$

For PECVD conditions one usually find  $n \leq 2$  [173]. The plasma potential  $V_{pl}$  equals the voltage drop at the sheath of the grounded electrode. It has been reported [170] that  $V_{pl} = V_{sg} = \frac{1}{2}(V_{RF} + V_{DC})$ . In the case of an RF frequency of 13.56 MHz, typical values are 10-30 V for the plasma potential, and with  $n = 2$  and with  $1 < A_{sg}/A_{sp} < 2$  typical DC self bias voltages are -50 V for low power (a few W) to -250 V for high power (upto 100 W).

The plasma potential is the maximum value with which ions can be accelerated from the edge of the sheath towards the substrate, located at the grounded electrode. This may cause ion bombardment, that may induce ion-surface interactions such as enhancement of adatom diffusion, displacement of surface atoms, trapping or sticking of incident ions, sputtering, and implantation, see Sec. VIB 1.

Meijer and Goedheer [174] have developed a 1D-model with which a.o. the DC self bias and the sheath voltages can be calculated. It is assumed that the potential variation over the *complete* discharge is harmonic:

$$V_{sp}(t) - V_{sg}(t) = V_{RF} \sin \omega t + V_{DC} \quad (9)$$

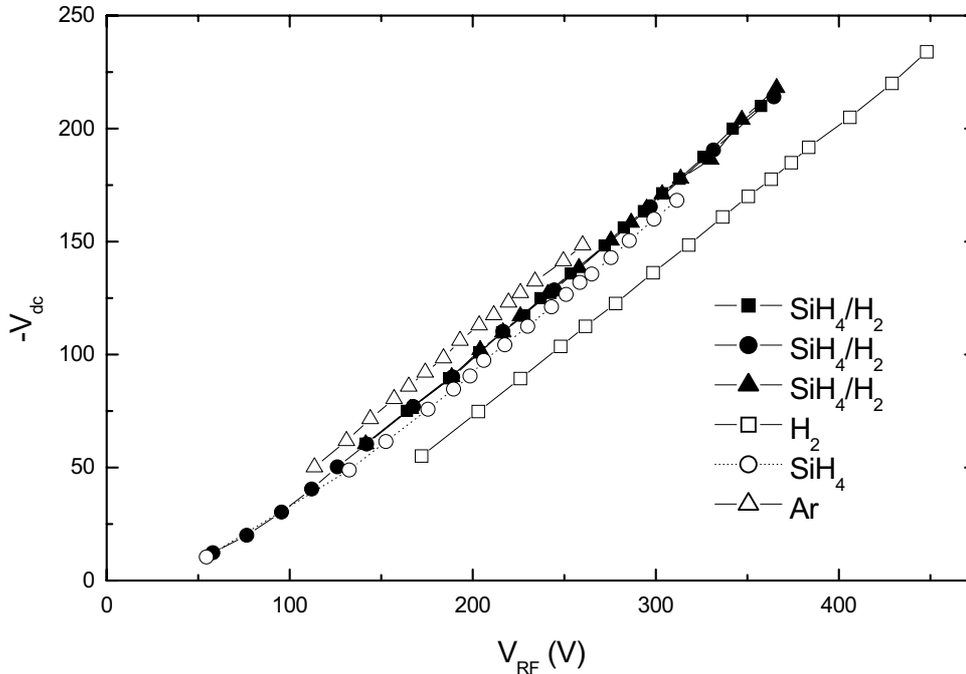


FIG. 9. Relation between DC self bias voltage  $V_{dc}$  and applied RF voltage  $V_{RF}$  for different gases (Ar, H<sub>2</sub>, SiH<sub>4</sub>) and gas mixtures ( $0.1 < \text{SiH}_4/(\text{H}_2 + \text{SiH}_4) < 0.9$ ). Note that the slope is independent of the gas used.

A further assumption is that the time-averaged current is zero, because of the capacitive coupling of the power supply to the electrodes. With this model unequally sized electrode systems can be modeled, by using an area ratio  $\alpha = A_g/A_p$ . The ratio between  $V_{DC}$  and  $V_{RF}$  has been calculated for  $1 < \alpha < 10$ , and is in excellent agreement [174,175] with experimental data [176,177].

Meiling *et al.* [151] have presented measured data on the relation between  $V_{DC}$  and  $V_{RF}$  in the ASTER system with an electrode area ratio  $\alpha = A_g/A_p$  of 2.1 for various silane/hydrogen mixtures and different applied power levels. It was found that  $V_{DC} = aV_{RF} + C$ , with  $a = 0.64$  and  $C = 31$ , independent of power and independent of silane/hydrogen mixture. The values of  $\alpha$  and  $a$  are in excellent agreement with the calculated results from Meijer and Goedheer [174]. Calculations by Goedheer *et al.* [149] in addition show that this relation also is independent of frequency. The slope  $a$  only depends on the geometry, i.e. area ratio, of the reactor system. The intercept  $C$  has a different value for pure argon or pure hydrogen plasmas, as can be inferred from Fig. 9. The intercept has not been predicted in the 1D-model by Meijer and Goedheer [174] and in fact results from this low dimensionality of the model. In a 2D model, the intercept is correctly calculated [178].

### III.B.2. Scaling laws

Goedheer *et al.* [148,149] have attempted to formulate scaling laws. They varied power  $P$ , pressure  $p$ , and frequency  $\omega$  and measured the DC self bias  $V_{DC}$ . From their

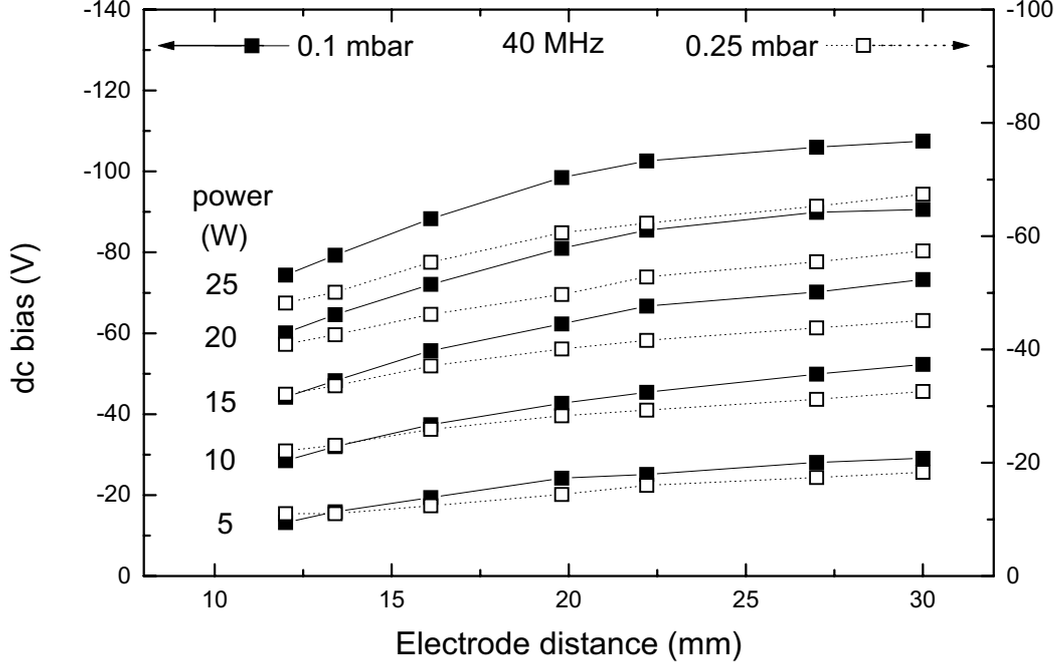


FIG. 10. DC self bias voltage  $V_{dc}$  as a function of electrode distance for two pressures, six power levels, at 40 MHz.

experimental results (frequency range 60-100 MHz, pressure range 20-60 Pa) a scaling law was formulated [148]:

$$P^{exp} \propto \omega^2 p^{2/3} V_{DC} \quad (10)$$

Additional data (frequency range 13.56-100 MHz, pressure range 10-60 Pa) resulted in a slightly changed scaling law [149],  $P^{exp} \propto \omega^2 p^{0.5} V_{DC}$ , which was refined using modeling data to [149,179]

$$P^{model} \propto \omega^{1.5} p^{0.7} V_{DC} \quad (11)$$

In addition a scaling law was derived in which also the electrode distance and radius was introduced:

$$P^{model} \propto \omega^{1.5} p^{0.7} \left(\frac{L}{R}\right)^{-0.4} R^{2.2} V_{DC} \quad (12)$$

In the ASTER system (see Sec. IID) experiments were performed in order to test this scaling law. To this end, a newly designed RF-electrode assembly was retrofitted to a deposition chamber. With this electrode set-up, it was possible to change the electrode distance from the outside, without breaking the vacuum! A large data-set was taken consisting of 420 datapoints [162], i.e. 3 values for the pressure ( $0.1 \leq p \leq 0.45$  mbar), 5 for the RF-power ( $5 \leq P \leq 25$  W), 7 for the electrode distance ( $12 \leq L \leq 30$  mm), and 4 for the RF-frequency ( $13.56 \leq \omega/2\pi \leq 50$  MHz).

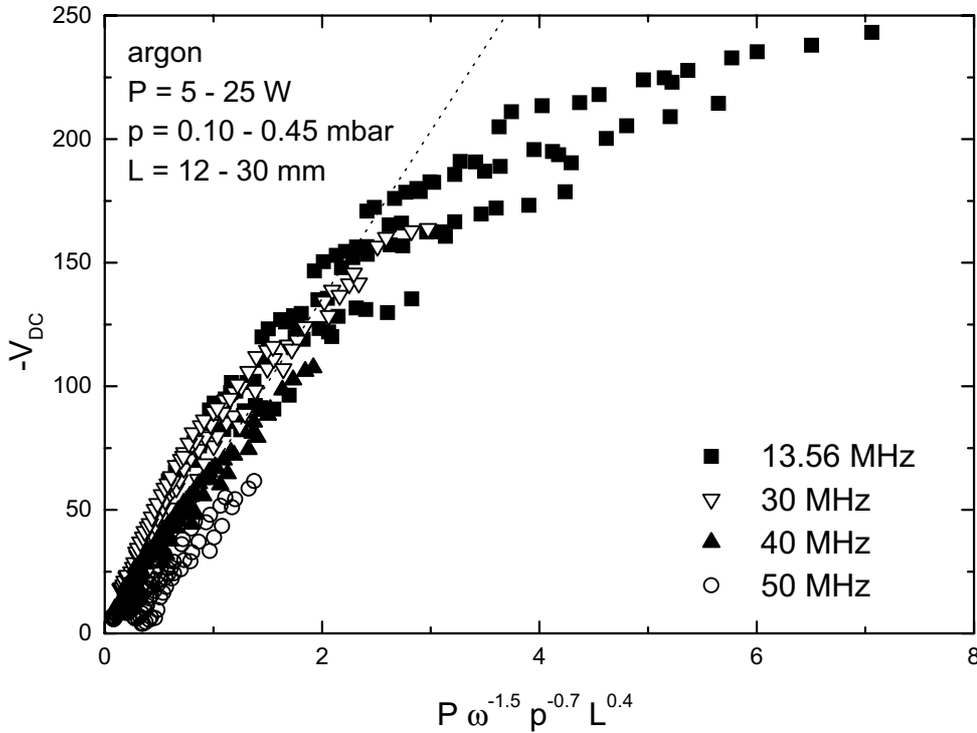


FIG. 11. Scaling law between DC self bias voltage  $V_{dc}$  and  $P \omega^{-1.5} p^{-0.7} L^{0.4}$  for an argon discharge.

A selection of the data-set is shown in Fig. 10. The DC self-bias voltage is shown as a function of electrode distance, for different values of the RF-power, at two pressures and a frequency of 40 MHz. It is clear from the experimental data that the DC self-bias voltage is not strongly dependent on electrode distance. Rauf and Kushner [180] have reported the effect of electrode distance on DC self-bias voltage for the Gaseous Electronics Conference reference cell (GECRC) [133]. They observe a gradual increase of DC self-bias versus electrode distance much like presented here, but only above a certain value of  $L$ . They attribute this to a change in current flows. Bringing the electrodes, which are of equal size, closer together results in a larger part of the current to flow out through the grounded electrode. Hence, the discharge become more symmetric and as a consequence, the DC self-bias decreases.

Comparing published data on the relation between the DC self-bias and RF-voltage between Rauf and Kushner [180] and our group [151] shows that the proportionality constant (slope) in the GECRC is nearly twice the one observed in the ASTER reactor. In other words, the ASTER reactor is less asymmetric than the GECRC. This may be an additional explanation as to why the  $V_{DC}$  data presented here do not show a sudden change as a function of electrode distance.

In Fig. 11 the complete data-set is plotted using the scaling law

$$P^{exp} \propto \omega^{1.5} p^{0.7} L^{-0.4} V_{DC} \quad (13)$$

It is clear that the scaling law, which was deduced from modeling results, also holds

for the experimental results. However, above about a value for the DC self-bias of -150 V, deviations occur. A closer look at the data reveals that the 13.56 MHz-data are responsible for this. Following the arguments discussed by Rauf and Kushner [180] this may well be explained by the fact that two different matching networks have been employed, i.e. a  $\pi$ -network for the 13.56 MHz-data, and an L-network for the other frequencies. These circuits are not included in the model. Another possible explanation is the presence of higher harmonics at high power levels, and large heat dissipation in the network.

### III.B.3. Effective discharge power

It is difficult to determine the true power that is dissipated in the discharge. Usually, the reading of a directional power meter in the 50  $\Omega$  line just before the matching network is reported as the RF power. The effective or true power is not simply the difference between input and reflected power, which is usually also measured with the same power meter. Godyak and Piejak have presented three methods to determine the true power, a subtractive method, a phase shift method, and an integration method [181]. All three methods give the same results. In the phase shift method the discharge power is determined from the discharge voltage and current and the phase shift  $\phi$  between them. In the integration method the product of voltage and current waveform is integrated over one RF cycle. The subtractive method was adapted from Horwitz [182], and is easiest in use. The principle is to measure the power delivered to the system including the tuned matching network in the case that the discharge is on ( $P_{tot}$  and in the case that it is off, i.e. when the system is evacuated ( $P_{vac}$ ), with the constraint that in both cases the  $P_{tot}$  and  $P_{vac}$  are measured for the same electrode voltage  $V_{pp}$ . The matcher efficiency [181] or power transfer efficiency  $\eta_P$  [183] then is defined as:

$$\eta_P = \left( \frac{P_{tot} - P_{vac}}{P_{tot}} \right)_{V_{pp}=\text{constant}} = \frac{P_{eff}}{P_{tot}} \quad (14)$$

where  $P_{eff}$  is the effective power dissipated in the discharge. The power transfer efficiency  $\eta_P$  has been reported to vary between 40 and 90% as function of pressure [181] and excitation frequency [183]. Both the subtractive and the phase shift method are valid as long as the RF waveform remains sinusoidal [184]. For asymmetrical discharges the integral method is to be used.

### III.B.4. Paschen law

In order to sustain a discharge between two parallel plate electrodes the product of pressure  $p$  and interelectrode distance  $L$  have to satisfy the Paschen law for the used gas mixture. This law relates the  $pL$ -product and the breakdown voltage  $V_b$  (or minimum sustaining voltage) of the discharge:

$$V_b = \frac{C_1 p L}{C_2 + \ln p L} \quad (15)$$

with  $C_1$  and  $C_2$  constants that are dependent on the gas mixture. For large  $pL$ -products the breakdown voltage is proportional to  $pL$ . For most gases the minimum breakdown

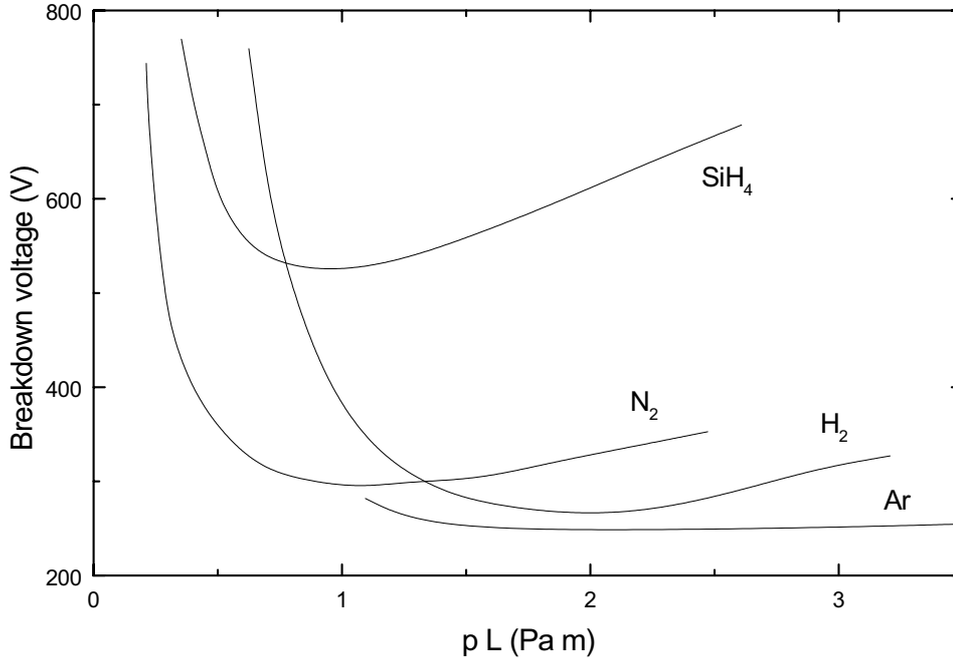


FIG. 12. Measured Paschen's law for silane, hydrogen, nitrogen and argon. (From K. M. H. Maessen, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1988).

voltage is between 100 and 500 V for  $pL$ -products in the range of 0.1-10 Torr·cm [117]. Measurements on argon, silane, hydrogen, and nitrogen [185,186] are shown in Fig. 12. Note that 1 Torr·cm equals 1.33 Pa·m.

At low  $pL$  values the energy of the electrons is limited by collisions with the electrodes, and a large voltage is required to compensate the energy lost to the walls. At high  $pL$  values electrons will collide with gas phase species, and the voltage again is large to compensate for the energy lost to collisions within the plasma, which however promotes polymerization or powder formation. Usually one employs a voltage just above the Paschen minimum to be able to deposit device quality material. Interelectrode distances typically are between 10 and 50 mm. Ishihara *et al.* [187] have reported only monohydride bonds to be present in the material for a  $pL$ -product of about 1 Torr·cm. Dihydride bonds already appear above a  $pL$ -product of 1.5 Torr·cm.

### III.C. Plasma chemistry

For low-pressure plasmas containing mainly inert gases the electrons can be characterized by a Maxwellian Electron Energy Distribution Function (EEDF), however deviations have been reported [188]. The shape of the EEDF depends on the energy lost via electron collisions and the energy gained by the time-dependent electric field in the plasma. Adding silane and/or hydrogen to the plasma results in a non-Maxwellian EEDF [189]. Electrons determine the plasma characteristics as they are responsible

for elastic collisions and inelastic collisions, that lead to rovibrational and electronic excitation, ionization, and dissociation of the neutral gas molecules. In elastic collisions the internal energy of the neutrals are unchanged, whereas the energy, momentum, and direction of the electrons undergo large changes. In inelastic collisions the energy that is transferred from the electron to the neutral species causes many processes to occur, such as dissociation, (dissociative) ionization, (dissociative) attachment, recombination, vibrational, electronic, and dissociative excitation. As a result of vibrational excitation of the molecules in the plasma, the collisions between these excited molecules allow for redistribution of vibrational states, charge exchange between molecules, excited molecules, dissociation, and dissipation of vibrational energy to walls of the system. Each of these processes is characterized by its own (energy-dependent) cross-section  $\sigma(E)$ . The rate constant  $k$  of the collisional processes are proportional to the product of EEDF ( $f(E)$ ) and cross-section, according to  $k = \int E^{1/2} f(E) \sigma(E) dE$ . The onset of these processes starts at 10 eV or higher, while the EEDF peaks around 1-2 eV. Hence, the tail of the EEDF mainly determines which processes occur.

### III.C.1. Species considered

In a silane/hydrogen discharge the feedstock gases  $\text{SiH}_4$  and  $\text{H}_2$  take part in all the processes that occur. A large number of reactions have been proposed (see e.g. Kushner [190]). Nienhuis *et al.* [191] have performed a sensitivity analysis in their self-consistent fluid model, from which a minimum set of reactions have been extracted for a typical low-pressure RF-discharge. The set presented here will be used in the plasma models presented in subsequent sections. Table 2 and Tables 3 list these reactions. The review articles on silane chemistry by Perrin *et al.* [192] and on hydrogen by Phelps [193] and Tawara *et al.* [194] have been used. The electron collision data are compiled in Fig. 13 [189].

### III.C.2. Electron impact collisions

Dissociation by electron impact of the gases  $\text{SiH}_4$  and  $\text{H}_2$  will create reactive species (radicals) and/or neutrals ( $\text{Si}_2\text{H}_6$  and even higher order silanes [195–198]). Atomic hydrogen is an important particle because it is formed nearly in all electron impact collisions and the H-abstraction reaction [199,200] of (di-)silane is an important process as results from the sensitivity study. Dissociation of  $\text{SiH}_4$  can create different  $\text{SiH}_x$  (with  $x = 0, 1, 2, 3$ ) radicals. Only silylene ( $\text{SiH}_2$ ) and silyl ( $\text{SiH}_3$ ) radicals are included in the plasma chemistry, because these two radicals are considered to be the most important ones [190]. The  $\text{SiH}_3$  radicals are assumed to be responsible for the film growth, they are so-called “growth precursors” and the  $\text{SiH}_2$  radicals are assumed to be precursors to form higher silanes, i.e.,  $\text{Si}_n\text{H}_{2n+2}$ . The H-abstraction reaction is important and it is therefore necessary to include the  $\text{Si}_n\text{H}_{2n+1}$  radical, if  $\text{Si}_n\text{H}_{2n+2}$  is included. Polymers larger than  $\text{Si}_8\text{H}_{18}$  can be omitted [191]. Dissociation of  $\text{Si}_2\text{H}_6$  produces mainly  $\text{SiH}_3$  and  $\text{Si}_2\text{H}_5$ . The latter is also assumed to be a growth precursor.

The total dissociation cross sections of silane and disilane have been taken from Perrin *et al.* [201]. An uncertainty in the present knowledge of the silane chemistry is

TABLE 2. Electron impact collisions. (Adapted from G. J. Nienhuis, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

acronym	electron impact reaction	threshold energy (eV)	reference	comment
SE1	$\text{SiH}_4 + e^- \rightarrow \text{SiH}_2^+ + 2 \text{H} + 2 e^-$	11.9	[206]	Lumped
S <sub>2</sub> E1	$\text{Si}_2\text{H}_6 + e^- \rightarrow \text{Si}_2\text{H}_4^+ + 2 \text{H} + 2 e^-$	10.2	[206]	Lumped
SV1	$\text{SiH}_4 + e^- \rightarrow \text{SiH}_4^{(1-3)} + e^- \rightarrow \text{SiH}_4 + e^-$	0.271	[212]	Vib. ex. 1-3
SV2	$\text{SiH}_4 + e^- \rightarrow \text{SiH}_4^{(2-4)} + e^- \rightarrow \text{SiH}_4 + e^-$	0.113	[212]	Vib. ex. 2-4
SE2	$\text{SiH}_4 + e^- \rightarrow \text{SiH}_2 + 2 \text{H} + e^-$	8.3	[199,201,202]	Fr. 83%
SE3	$\text{SiH}_4 + e^- \rightarrow \text{SiH}_3 + \text{H} + e^-$	8.3	[199,201,202]	Fr. 17%
S <sub>2</sub> E2	$\text{Si}_2\text{H}_6 + e^- \rightarrow \text{SiH}_3 + \text{SiH}_2 + \text{H} + e^-$	7.0	[201]	Fr. 91 % [204]
SE4	$\text{SiH}_4 + e^- \rightarrow \text{SiH}_3^- + \text{H}$	5.7	[208]	Lumped
HE1	$\text{H}_2 + e^- \rightarrow \text{H}_2^+ + 2 e^-$	15.7	[207]	Lumped
HV	$\text{H}_2^{v=0} + e^- \rightarrow \text{H}_2^{v \neq 0} + e^- \rightarrow \text{H}_2^{v=0} + e^-$	0.54	[213]	$v = 1$
		1.08		$v = 2$
		1.62		$v = 3$
HE2	$\text{H}_2 + e^- \rightarrow \text{H} + \text{H} + e^-$	8.9	[205]	$b^3\Sigma_u^+$ ex.

TABLE 3. Chemical reactions. (Adapted from G. J. Nienhuis, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

acronym	chemical reaction	reaction rate ( $\text{cm}^3\text{s}^{-1}$ )	reference	comment
HS	$\text{H} + \text{SiH}_4 \rightarrow \text{SiH}_3 + \text{H}_2$	$2.8 \times 10^{-11}$	[192,199]	
		$\times \exp(-1250/T_{gas})$	[192,199]	
HS <sub>2</sub> 1	$\text{H} + \text{Si}_2\text{H}_6 \rightarrow \text{Si}_2\text{H}_5 + \text{H}_2$	$1.6 \times 10^{-10}$	[192,199,200]	
		$\times \exp(-1250/T_{gas})$	[192,199,200]	
HS <sub>2</sub> 2	$\text{H} + \text{Si}_2\text{H}_6 \rightarrow \text{SiH}_3 + \text{SiH}_4$	$0.8 \times 10^{-10}$	[192,199,200]	
		$\times \exp(-1250/T_{gas})$	[192,199,200]	
HS <sub>n</sub>	$\text{H} + \text{Si}_n\text{H}_{2n+2} \rightarrow \text{Si}_n\text{H}_{2n+5} + \text{H}_2$	$2.4 \times 10^{-10}$	[192]	Est.,
		$\times \exp(-1250/T_{gas})$		$n > 2$
H <sub>2</sub> S1	$\text{H}_2 + \text{SiH}_2 \rightarrow \text{SiH}_4$	$3.0 \times 10^{-12}$	[192]	$p_0$ (Pa)
		$\times [1 - (1 + 0.00023p_0)^{-1}]$		
SS1	$\text{SiH}_4 + \text{SiH}_2 \rightarrow \text{Si}_2\text{H}_6$	$2.0 \times 10^{-10}$	[192]	$p_0$ (Pa)
		$\times [1 - (1 + 0.0032p_0)^{-1}]$		
S <sub>2</sub> S1	$\text{Si}_2\text{H}_6 + \text{SiH}_2 \rightarrow \text{Si}_3\text{H}_8$	$4.2 \times 10^{-10}$	[192]	$p_0$ (Pa)
		$\times [1 - (1 + 0.0033p_0)^{-1}]$		
S <sub>n</sub> S	$\text{Si}_n\text{H}_{2n+2} + \text{SiH}_2 \rightarrow \text{Si}_{n+1}\text{H}_{2n+4}$	$4.2 \times 10^{-10}$	[192]	Est.,
		$\times [1 - (1 + 0.0033p_0)^{-1}]$		$p_0$ (Pa), $n > 2$
SS2	$\text{SiH}_3 + \text{SiH}_3 \rightarrow \text{SiH}_4 + \text{SiH}_2$	$1.5 \times 10^{-10}$	[192,215]	
SS3	$\text{SiH}_3^- + \text{SiH}_2^+ \rightarrow \text{SiH}_3 + \text{SiH}_2$	$1.2 \times 10^{-7}$	[214]	
S <sub>2</sub> S2	$\text{SiH}_3^- + \text{Si}_2\text{H}_4^+ \rightarrow \text{SiH}_3 + 2 \text{SiH}_2$	$1.0 \times 10^{-7}$	[214]	
H <sub>2</sub> S2	$\text{SiH}_3^- + \text{H}_2^+ \rightarrow \text{SiH}_3 + \text{H}_2$	$4.8 \times 10^{-7}$	[214]	
S <sub>2</sub> S <sub>2</sub>	$\text{Si}_2\text{H}_5 + \text{Si}_2\text{H}_5 \rightarrow \text{Si}_4\text{H}_{10}$	$1.5 \times 10^{-10}$	[192]	Est.

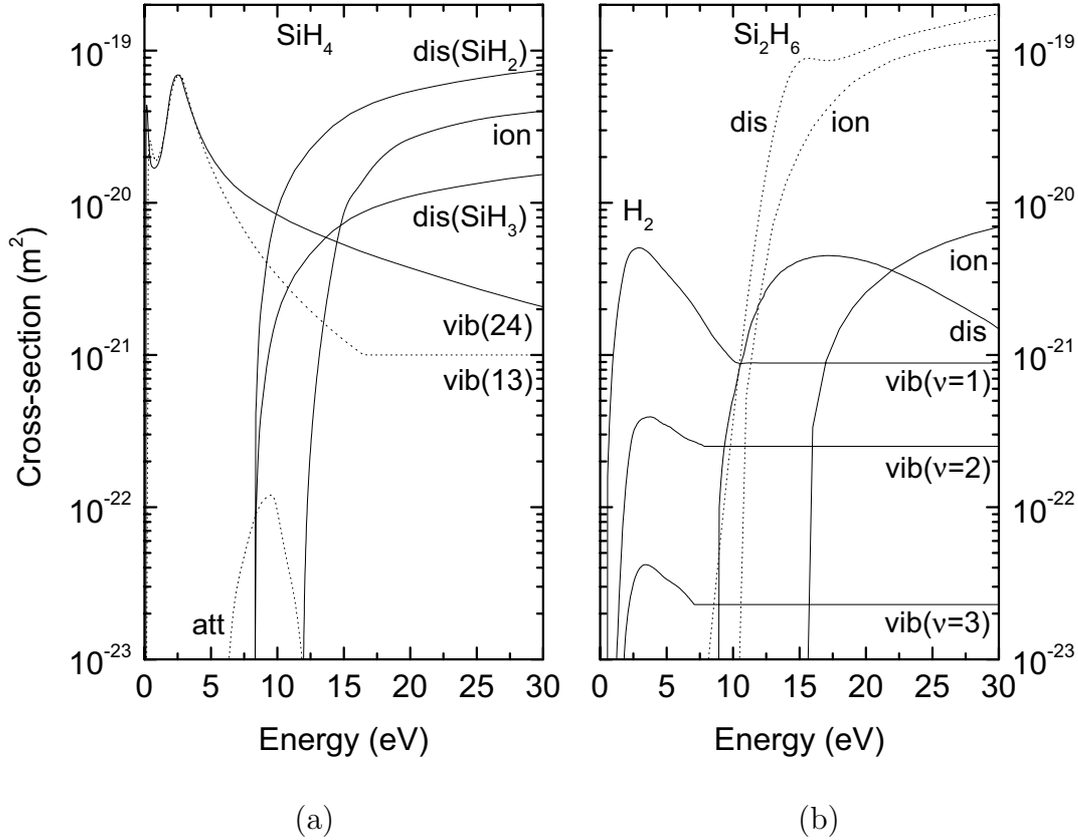


FIG. 13. Cross-sections of the electron collisions for  $\text{SiH}_4/\text{H}_2$ . (a) cross-sections for  $\text{SiH}_4$ , (b) cross-sections for  $\text{Si}_2\text{H}_6$  (dotted lines) and  $\text{H}_2$  (solid lines). Abbreviations are: 'ion' = ionization, 'dis' = dissociation, 'vib' = vibrational excitation, 'att' = attachment. See Tab. 2 for details and references. (Adapted from G. J. Nienhuis, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

the branching ratio of the silane dissociation channels [192]. Here, the branching ratio is taken from Doyle *et al.* [197], who suggest to use the branching ratio determined by Perkins *et al.* [202] for photolysis, i.e., a branching of 83 % into  $\text{SiH}_2$  and 17 % into  $\text{SiH}_3$  (reactions SE2 and SE3 in Tab. 2). However, Perrin *et al.* [192] suggest that the formation of  $\text{SiH}$  is also of importance, at low electron energies (9.9 eV): it should be at least 20 %. In contrast, Hertl *et al.* [203] have measured  $\text{SiH}$  densities in typical  $\text{SiH}_4/\text{H}_2$  discharges, and they show that the  $\text{SiH}$  density is at least one order of magnitude lower than the  $\text{SiH}_2$  density. Therefore, the  $\text{SiH}$  radical is not included. The fragmentation pattern of disilane dissociation (reaction  $\text{S}_2\text{E2}$  in Tab. 2) is taken from Doyle *et al.* [204], which is 91 %  $\text{SiH}_3 + \text{SiH}_2 + \text{H}$  and 9 %  $\text{H}_3\text{SiSiH} + 2\text{H}$ . As a simplification the second branching path is neglected. Dissociation of  $\text{H}_2$  by electron impact is included [205].

Ionization of  $\text{SiH}_4$ ,  $\text{Si}_2\text{H}_6$ , and  $\text{H}_2$  can produce a large number of different positive ions [206,207]. In order to simplify all the ionization cross sections of each background neutral leading to different ions have been lumped and attached to the ion which is most probable to be formed: for  $\text{SiH}_4$ ,  $\text{Si}_2\text{H}_6$ , and  $\text{H}_2$  these ions are  $\text{SiH}_2^+$ ,  $\text{Si}_2\text{H}_4^+$ , and

$\text{H}_2^+$ , respectively (reactions SE1, S<sub>2</sub>E1, and HE1 in Tab. 2).

Similar to the lumping of the positive ionization cross sections, the dissociative attachment cross sections of each background neutral leading to different anions have been lumped to one cross section (reaction SE4 in Tab. 2). The most abundant anions of  $\text{SiH}_4$ , and  $\text{H}_2$  are  $\text{SiH}_3^-$  and  $\text{H}^-$ , respectively [208,209]. Since calculations have shown that dissociative attachment of  $\text{H}_2$  has a negligible influence, it is not included. The sum of the dissociative attachment cross sections of  $\text{Si}_2\text{H}_6$  is comparable in magnitude to that of  $\text{SiH}_4$  [210]. However, as the partial pressure of  $\text{Si}_2\text{H}_6$  is at most a tenth of the partial pressure of  $\text{SiH}_4$ , the dissociative attachment of  $\text{Si}_2\text{H}_6$  is neglected.

Vibrational excitation by electron impact of the background neutrals is an important process, because it accounts for a major energy loss for the electrons (reactions SV1 ( $\text{SiH}_4$  stretching mode), SV2 ( $\text{SiH}_4$  bending mode), and HV in Tab. 2). Moreover, the density of the vibrationally excited molecules has been reported to be important [211]. However, information about reaction coefficients of vibrationally excited molecules is scarce [192]. Here, only the vibrational excitation of  $\text{SiH}_4$  and  $\text{H}_2$  are included [212,213].

### III.C.3. Chemical reactions

Table 3 lists the set of chemical reactions, with reaction rates and references. The rate constants for mutual neutralization of a positive ion and  $\text{SiH}_3^-$  (reactions SS3, S<sub>2</sub>S2, and H<sub>2</sub>S2 in Tab. 3) have been taken from a scaling formula from Hickman [214], where the electron affinities are taken from Perrin *et al.* [192] and the ion temperature is assumed equal to the gas temperature. The end products of the mutual neutralization reaction of  $\text{SiH}_3^-$  and  $\text{Si}_2\text{H}_4^+$  are taken  $\text{SiH}_3$  and  $2\text{SiH}_2$  (not  $\text{SiH}_3$  and  $\text{S}_2\text{H}_4$ ), hence no extra species (i.e.,  $\text{Si}_2\text{H}_4$ ) have to be introduced. The radical-radical reaction  $\text{SiH}_3 + \text{SiH}_3$  can lead to  $\text{Si}_2\text{H}_6$  or  $\text{SiH}_2 + \text{SiH}_4$  [215]. However,  $\text{Si}_2\text{H}_6$  will be an excited association product and collisional de-excitation will only occur at high pressures [192]. This excited  $\text{Si}_2\text{H}_6$  is likely to dissociate into  $\text{SiH}_2$  and  $\text{SiH}_4$  and therefore the only end product of  $\text{SiH}_3 + \text{SiH}_3$  is assumed to be  $\text{SiH}_2 + \text{SiH}_4$ .

Chemical reactions of positive ions with background neutrals [193,216], e.g.,  $\text{SiH}_2^+ + \text{SiH}_4 \rightarrow \text{SiH}_3^+ + \text{SiH}_3$  or  $\text{H}_2^+ + \text{H}_2 \rightarrow \text{H}_3^+ + \text{H}$  have not been included in the model. The reaction coefficients of these reactions are of equal magnitude as some radical-neutral reactions, but the densities of the ions are much lower than the radical densities making ion-neutral reaction of negligible influence on the partial pressure of the background neutrals and the growth rate. For a detailed analysis of the densities and fluxes of ions, the lumped sum of ion densities should be unraveled again.

### III.C.4. Plasma-wall interaction

The plasma-wall interaction of the neutral particles is described by a so-called sticking model [136,137]. In this model only the radicals react with the surface while non-radical neutrals ( $\text{H}_2$ ,  $\text{SiH}_4$ , and  $\text{Si}_n\text{H}_{2n+2}$ ) are reflected into the discharge. The surface reaction and sticking probability of each radical must be specified. The nature (material, roughness) and the temperature of the surface will influence the surface reaction probabilities. Perrin *et al.* [136] and Matsuda *et al.* [137] have shown that the surface reaction coefficient  $\beta$  of  $\text{SiH}_3$  is temperature independent at a value of  $\beta = 0.26 (\pm 0.05)$

at a growing *a*-Si:H surface in a temperature range from room temperature to 750 K. The surface sticking coefficient  $s$  of  $\text{SiH}_3$  is  $s = 0.09 (\pm 0.02)$  in a temperature range from room temperature to 575 K. Above 575 K the  $s/\beta$  ratio increases to 1 (saturation) at a temperature of 750 K. Note, that typical deposition temperatures are below 575 K.

Information about the surface reaction coefficients of  $\text{Si}_n\text{H}_{2n+1}$ , where  $n > 1$ , radicals is scarce. Because of the fact that the structure of these radicals is similar to that of  $\text{SiH}_3$ , the same surface reaction coefficients are used. It is assumed that if  $\text{Si}_n\text{H}_{2n+1}$  radicals recombine at the surface with a hydrogen atom, a  $\text{Si}_n\text{H}_{2n+2}$  neutral is formed which is reflected into the discharge. Another possibility is the surface recombination of  $\text{Si}_n\text{H}_{2n+1}$  radicals with physisorbed  $\text{Si}_m\text{H}_{2m+1}$  radicals at the surface. Matsuda *et al.* [137] have shown that the probability of surface recombination of  $\text{SiH}_3$  with physisorbed  $\text{SiH}_3$  decreases with increasing substrate temperature. Doyle *et al.* [204] concluded that at a typical substrate temperature of 550 K,  $\text{SiH}_3$  radicals mainly recombine with physisorbed H atoms.

For the  $\text{SiH}_2$  radicals the surface reaction coefficients have been taken as  $s = \beta = 0.8$  [192]. This sticking coefficient is large because there is no barrier for insertion of this species into the *a*-Si:H surface. Kae-Nune *et al.* [217] specify a surface recombination probability of about 1 for atomic hydrogen on an *a*-Si:H surface during deposition which results mainly in recombination of H with an H-atom bounded to the surface.

If a  $\text{Si}_n\text{H}_{2n+1}$  radical sticks to the surface, not all hydrogen will be incorporated in the layer and therefore molecular hydrogen will be reflected into the discharge. The amount of molecular hydrogen thus reflected into the discharge is taken such that the percentage of hydrogen in the deposited layer is 10 %. The growth rate is calculated by summing the flux of Si containing neutrals and ions toward the electrodes and assuming a constant density of  $2.2 \text{ g/cm}^3$  of the deposited layer [218].

### III.C.5. Transport coefficients

The diffusion constant  $D_j$  of neutral particle  $j$  is calculated in two steps. First, the binary diffusion coefficient in each of the background gas species ( $\text{SiH}_4$ ,  $\text{Si}_2\text{H}_6$ ,  $\text{H}_2$ ) is calculated, following Perrin *et al.* [192]. Then  $D_j$  is approximated using Blancs law [219]:

$$\frac{p_{tot}}{D_j} = \sum_{i=\text{background}} \frac{p_i}{D_{ij}} \quad (16)$$

where  $p_{tot}$  is the total pressure (Pa) and  $p_i$  is the partial pressure of background species  $i$ .

The ion mobility coefficients  $\mu_j$  are calculated similarly. First, the ion mobility of ion  $j$  in background neutral  $i$  is calculated using the low E-field Langevin mobility expression [219]. Then Blancs law is used to calculate the ion mobility in the background mixture. The ion diffusion coefficient is calculated with Einsteins relation

$$D_i = \frac{k_B T_{ion}}{e} \mu_i \quad (17)$$

where  $T_{ion}$  is the ion temperature, which is assumed to be equal to the gas temperature  $T_{gas}$ .

## IV. PLASMA MODELING

All properties of *a*-Si:H are a result of operating conditions of the discharge (pressure, RF frequency, RF power, gas mixture, geometry). Optimization of properties mostly is done empirically, due to the complex chemistry of the silane/hydrogen discharge, including plasma-wall (=substrate) interaction. Nevertheless, serious attempts have been made to model the discharge, thus providing knowledge on chemistry and the deposition process, with an ultimate goal to find the optimum parameter space of the specific reactor system used.

Various types of models have been used to describe and study glow discharges. These models differ in the approximations that are made (e.g., the dimension, self-consistency, fluid or kinetic approach, and the number of chemical processes), which will influence the physical relevance of the modeling results. However, a higher accuracy or fewer approximations will require a larger computational effort. Much effort has been put in the development of self-consistent models for glow discharges. Examples are fluid models [220,221], Particle In Cell/Monte Carlo (PIC/MC) [222–224] and hybrid models [190,225,226]. More extensive overviews have been presented elsewhere [195,196,227,228]. Most of these self-consistent models have been used for discharges with a relatively simple chemistry and no plasma-wall interaction of the neutrals. Other models include the silane/hydrogen chemistry and the surface interaction to a relatively high degree of completeness, but these models are not self-consistent with respect to the rates of the electron impact collisions (i.e., ionization, dissociation, excitation, and attachment), for instance Perrin *et al.* [211]. Models developed for plug-flow silane/hydrogen discharges [190,195,196] are self-consistent with respect to the silane/hydrogen chemistry, the electron impact collisions, the plasma-wall interaction, and the transport phenomena. However, in this type of models the composition of the neutral background gas is equal to the gas feedstock composition, that is, the depletion of silane will be low.

### IV.A. 1D fluid discharge model

#### IV.A.1. Model description

Nienhuis *et al.* [189,191] have developed a self-consistent fluid model which describes the electron kinetics, the silane/hydrogen chemistry, as well as the deposition process of a perfectly stirred reactor. Due to the discharge processes, the composition of the background neutrals will differ from the composition of the feedstock gases, because hydrogen and higher order silanes are formed, while silane is consumed. Fluid models describe the discharge by a combination of balances for the particle, momentum, and energy densities of the ions, electrons, and neutrals obtained from moments of the Boltzmann transport equation, see e.g. [174,229,230]. A limitation of a fluid model is the requirement that the mean free path of *all* particles must be less than the characteristic dimensions of the discharge. The balance equations are coupled to the Poisson equation to calculate the electric field, where only electrostatic forces are taken into account. Solving these equations yields the behaviour in space and time of the particle densities and the electric potential during one cycle of the periodic steady state RF discharge.

Rates for electron impact collisions as well as the electron transport coefficients depend on the electron energy distribution function (EEDF). The collision rates, the electron transport coefficients, and the average electron energy are obtained by solving the Boltzmann equation for the EEDF. This EEDF has been expanded in two terms with respect to the velocity [231]. The EEDF is calculated as a function of the electric field for a given composition of the neutral background density. A look-up table has been constructed to obtain the collision rates and electron transport coefficients as functions of the average electron energy which are used in the fluid model. The deposition process is modeled by the use of sticking coefficients, which actually specify the boundary conditions of the particle density balance equations.

The 1D fluid model describes a discharge created and sustained between two parallel plates, where the time-dependent discharge characteristics only vary along the  $z$ -axis, i.e., the direction normal to the plates. Plasma parameters such as the DC self-bias voltage, uniformity of the deposition, and radial transport cannot be studied. Self-consistent models in more than one dimension can model these transport phenomena more appropriately [220,221,232]. In reality radial dependencies do exist in the discharge, due to radial transport processes. Silane is introduced and is transported from the inlet into the discharge, where it may be transformed into other species via chemical reactions. This new species may be transported outside the discharge toward the pump-outlet. The model described here corrects for this by means of additional source terms. The reactor is divided into two volumes, the discharge volume (defined as the volume between the two electrodes), and the discharge-free volume, see also Fig. 4a. The charged particles and the radicals are confined to the discharge volume. Radical reaction times or diffusion times are in the millisecond range, and shorter than the diffusion time to the outside wall, and shorter than the residence time ( $\sim 1$  sec) in the reactor. Hence, radial dependences are neglected, and the reactor is considered perfectly stirred. The radical densities are spatially resolved in the  $z$ -direction, while the non-radical neutral densities are assumed to be homogeneous throughout the whole reactor volume. In the remaining discharge-free volume only non-radical neutrals are considered.

The main reason for using a one-dimensional model is the reduction of the computational effort compared to the higher dimensional models. This reduction to one dimension is acceptable since it is possible to study the sustaining mechanisms and the chemistry in the discharge with a one-dimensional self-consistent model.

To verify the model and to establish in which process parameter space the model is valid, a comparison is made with experimental data. These experimental data are the partial pressures of silane, disilane, hydrogen, and the growth rate, obtained during deposition of amorphous silicon. Data are compared for various combinations of the total pressure in the reactor, the electrical power, and the frequency of the power source.

A sensitivity study of the influence of the elementary data (i.e., reaction coefficients, cross sections, and transport coefficients) has been performed to determine the importance of specific elementary data, so to guide further research in this area [189].

## IV.A.2. Particle balances

For each non-radical neutral  $j$  in the discharge volume  $V_D$  the balance equation of their total number  $N_j^D$  can be written as:

$$\frac{\partial N_j^D}{\partial t} = \int_D S_{\text{reac},j} dV + Q_j^{F \rightarrow D} \quad (18)$$

where the source term  $S_{\text{reac},j}$  represents the creation or destruction of corresponding species  $j$  by electron impact collisions and/or by chemical reactions, e.g.,  $A+B \rightarrow C+D$ , where  $A$ ,  $B$ ,  $C$ , and  $D$  indicate the reaction species. The source term for the particles in such a reaction is

$$S_{\text{reac},C} = S_{\text{reac},D} = -S_{\text{reac},A} = -S_{\text{reac},B} = n_A n_B k_r \quad (19)$$

where  $k_r$  is the reaction rate. For electron collisions the reaction rate depends on the local (averaged) electron energy, see Fig. 13 and Tab. 2. Other chemical reaction rate coefficients are constant, see Tab. 3. The term  $Q_j^{F \rightarrow D}$  represents the total number of neutrals  $j$  per second that are transported from the discharge free ( $F$ ) volume into the discharge ( $D$ ) volume.

For each non-radical neutral  $i$  in the discharge-free volume  $V_F$  the balance equation of their total number  $N_i^F$  can be written as:

$$\frac{\partial N_j^F}{\partial t} = Q_j^{\text{in}} + Q_j^{\text{pump}} - Q_j^{F \rightarrow D} \quad (20)$$

where the term  $Q_j^{\text{in}}$  represents the number of particles that enter the reactor volume per second as feedstock gas. The term  $Q_j^{\text{pump}}$  represents the number of pumped particles per second, and is given by:

$$Q_j^{\text{pump}} = -\frac{N_j^F}{\tau} \quad (21)$$

with  $\tau$  the average residence time, which is computed such that the total pressure in the discharge  $p_{\text{tot}}$ , given by the ideal gas law, equals the preset pressure.

Only the steady-state situation is considered, therefore the partial time derivatives are set to zero. Then, combining the above equations and solving for  $N_j^F$  yields:

$$n_j = \frac{\tau}{V_F} \left( Q_j^{\text{in}} + \int_D S_{\text{reac},j} dV \right) \quad (22)$$

For each particle  $j$  in the discharge volume (electrons, ions, and radicals) the density balance can be written as:

$$\frac{\partial n_j}{\partial t} + \frac{\partial \Gamma_j}{\partial z} = S_{\text{reac},j} \quad (23)$$

where  $n_j$  is the density and  $\Gamma_j$  is the flux of particle  $j$ .

#### IV.A.3. Particle fluxes

In the fluid model the momentum balance is replaced by the drift-diffusion approximation, where the particle flux  $\Gamma$  consists of a diffusion term (caused by density gradients) and a drift term (caused by the electric field),

$$\Gamma_j = \mu_j n_j E - D_j \frac{\partial n_j}{\partial z} \quad (24)$$

where  $\mu_j$  and  $D_j$  are the mobility and diffusion transport coefficients, respectively, which are charge-sign dependent. Electron and negative ion mobilities are negative; the mobilities for positive ions is positive. For the neutral particles the mobility is set equal to zero. The use of the drift-diffusion approximation is allowed when two conditions are satisfied. The first condition is that the characteristic time between momentum transfer collisions (of the charged particles) is much smaller than the RF period. Or, in other words, the drift-diffusion approximation assumes that a charged particle reacts instantaneously to a change of the electric field, because the moment equations of the fluid model are simplified by neglecting the particle inertia. The typical momentum transfer frequency for electrons is about 100 MHz. For ions the characteristic momentum transfer frequency is only a few MHz. To use the drift-diffusion approximation for RF frequencies higher than a few MHz the electric field is replaced by an effective electric field [233], which accounts for the fact that the ions do not respond instantaneously to the actual electric field, unless the momentum transfer frequency is much larger than the applied frequency.

The second condition for the use of the drift-diffusion approximation specifies that the mean free path of these collisions is much smaller than the characteristic gradient lengths in the discharge. In other words the motion of the particles must be collision dominated. The mean free path of the electrons is the largest of all particles and their description will become less accurate in the low pressure regime, i.e., at pressures lower than about 10 Pa. The cross-section for momentum transfer for the electrons in silane is about  $10^{-19} \text{ m}^2$  [212]. At 10 Pa, this gives a mean free path of about  $5 \times 10^{-3} \text{ m}$ , which is of the order of the length scale of the electron density gradients in the discharge. For ions also this pressure limit is valid.

#### IV.A.4. Poisson equation

The electric field  $E$  and potential  $V$  are calculated using the Poisson equation,

$$\frac{\partial^2 V}{\partial z^2} = - \frac{e}{\epsilon_0} \left( \sum_{i=\text{ions}} q_i n_i - n_e \right); \quad E = - \frac{\partial V}{\partial z} \quad (25)$$

with  $e$  the elementary charge,  $\epsilon_0$  the vacuum permittivity,  $n_e$  the electron density,  $n_i$  the ion density, and  $q_i$  the sign of the ion charge  $i$ , i.e., +1 or -1.

#### IV.A.5. Energy balances

The electron energy density  $w_e = n_e \epsilon$  (i.e., the product of the electron density  $n_e$  and average electron energy  $\epsilon$ ) is calculated self-consistently from the second moment of the Boltzmann equation:

$$\frac{\partial w_e}{\partial t} + \frac{\partial \Gamma_w}{\partial z} = -e\Gamma_e E + S_w \quad (26)$$

where  $\Gamma_w$  is the electron energy density flux:

$$\Gamma_w = \frac{5}{3} \left( \mu_e w_e E - D_e \frac{\partial w_e}{\partial z} \right) \quad (27)$$

and  $\mu_e$  and  $D_e$  are the electron mobility and electron diffusion coefficients. The term  $S_w$  in the electron energy balance equation is the loss of electron energy due to electron impact collisions. The various source terms for electron impact collisions depend on the local average electron energy  $\epsilon$ , which is calculated by dividing the energy density  $w_e$  by the electron density  $n_e$ . The energy loss due to elastic collisions of electrons with neutrals [234], amounts to only about 1 % of the total energy dissipation at typical discharge conditions and this term is therefore neglected.

No energy balances for the ions and neutrals are included. The energy balance for neutrals is complex due to the large range of possible energy exchange reactions in a typical discharge. Moreover, most of the reaction coefficients are unknown [211]. It is assumed that the ions have nearly the same energy as the neutral atoms, because of the efficient energy transfer between ions and neutrals due to their nearly equal mass. Further, the energy gain of the ions from the electric field is low because of the high inertia of the ions; the ions cannot respond to the fast oscillations of the electric field. This assumption is correct for the bulk of the discharge, where the time averaged electric field is nearly zero. However, the time averaged electric field in the sheaths is *not* negligible and will increase the energy of the ions far above the thermal energy of the neutrals, as is experimentally shown for silane/hydrogen plasmas [235,236].

#### IV.A.6. Boundary conditions

In order to solve the differential equations, the boundary conditions of the potential and density profiles must be specified. The potential at the grounded electrode ( $z = L$ , radius  $R$ ) is set equal to zero. The potential at the driven electrode ( $z = 0$ ) is set equal to

$$V(z = 0, t) = V_{RF} \sin(2\pi\nu_{RF}t) \quad (28)$$

where  $V_{RF}$  is the amplitude of the potential and  $\nu_{RF}$  is the frequency of the oscillating potential. The value of  $V_{RF}$  is adjusted until the dissipated power in the discharge equals the preset electrical power. The dissipated power (expressed in W) is given by the ohmic heating of the charged particles:

$$P_{dis} = \frac{e\pi R^2}{\tau_{RF}} \sum_{\substack{j=\text{charged} \\ \text{particles}}} \int_0^{\tau_{RF}} \int_0^L E(z, t) \Gamma_j(z, t) dz dt \quad (29)$$

where  $\tau_{RF}$  is the RF-period.

As the electric field always points in the direction of the electrode, the densities of the electrons and negative ions are set equal to zero at the electrode. It is assumed that

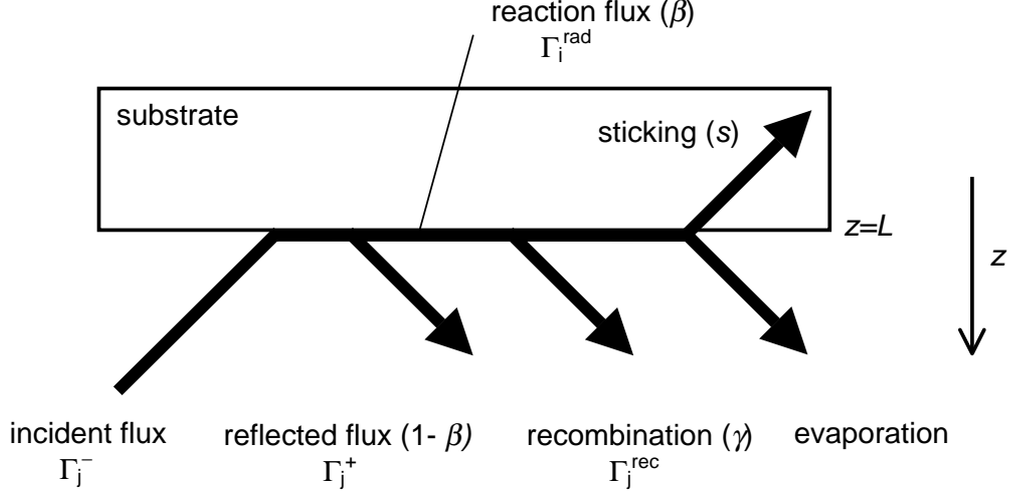


FIG. 14. Schematic representation of plasma-wall interaction for neutrals. (Adapted from G. J. Nienhuis, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

the ion flux at the electrodes has only a drift component, i.e., the density gradient is set equal to zero. The conditions in the sheath, which depend on pressure, voltage drop, and sheath thickness, are generally such that secondary electrons (created at the electrodes as a result of ion impact) will only ionize at most a few molecules, so no ionization avalanches will occur. Therefore, secondary electrons can be neglected.

The boundary conditions for the density balance equation of neutrals are given by the plasma-wall interaction (for instance by deposition processes), which is described by a sticking model, see Fig. 14. This surface interaction is modeled by using surface reaction coefficients  $\beta_j$ , which specify the probability that neutral  $j$  will react at the surface. The coefficient  $R_j = 1 - \beta_j$  represents the fraction of the incident neutrals which is reflected into the gas phase keeping their initial identity. It is assumed that the non-radical neutrals do not react at the wall, whereas the radical neutrals may react at the surface, and form a non-radical neutral, or contribute to  $\alpha$ -Si:H layer formation.

For the radical neutrals, boundary conditions are derived from diffusion theory [237,238]. One-dimensional particle diffusion is considered in gas close to the surface at which radicals react (Fig. 14). The particle fluxes in both  $z$  directions can be written as:

$$\Gamma_j^- = \frac{1}{4}n_j v_{th,j} + \frac{D}{2} \frac{\partial n_j}{\partial z} \quad (30)$$

$$\Gamma_j^+ = \frac{1}{4}n_j v_{th,j} - \frac{D}{2} \frac{\partial n_j}{\partial z} \quad (31)$$

where  $v_{th,j}$  is the thermal velocity of neutral  $j$ , given by  $v_{th,j} = \sqrt{8k_B T_{gas}/\pi m_j}$ . The first terms in Eq. (31) are the random contributions, and the second terms are the contributions due to the gradient in the concentration of diffusion particles. The resulting net flux  $\Gamma_j$  is given by  $\Gamma_j = \Gamma_j^+ - \Gamma_j^-$ . The fraction  $R_j$  of the flux incident on the wall ( $\Gamma_j^-$ ) is equal to the flux coming from the wall,  $\Gamma_j^+ = R_j \Gamma_j^-$ , when no radicals are produced at the wall. The boundary condition of radical  $j$ , that reacts at the surface

of the electrode then is given by:

$$\Gamma_j^{rad} = -D_j \left. \frac{\partial n_j^{rad}}{\partial z} \right|_{z=0} = -\frac{\beta_j}{1-\beta_j/2} \frac{v_{th,j}}{4} n_j^{rad} \Big|_{z=0} \quad (32)$$

where  $\Gamma_j^{rad}$ ,  $n_j^{rad} \Big|_{z=0}$ , and  $m_j$  are the flux of radicals  $j$  reacting at the surface, their density at the surface and their mass, respectively.

The recombination probability  $\gamma_{ij}$  is defined as the probability that a neutral radical  $j$  reacts on the surface, forming a volatile stable product, i.e., a non-radical neutral  $i$ , which is reflected into the discharge (e.g.  $\text{SiH}_3(gas) + \text{H}_{(surface)} \rightarrow \text{SiH}_4(gas)$ ). This causes a flux into the discharge:

$$\Gamma_j^{rec} = \sum_{j \neq i} \frac{\gamma_{ij}}{\beta_j} D_j \left. \frac{\partial n_j^{rad}}{\partial z} \right|_{z=0} \quad (33)$$

The sticking coefficients  $s_i = \beta_i - \gamma_i$  (where  $\gamma_i = \sum_{j \neq i} \gamma_{ij}$ ) is the probability that a neutral  $i$  sticks to the surface. In case of silane radicals  $\text{Si}_n\text{H}_m$ , this means film growth by Si incorporation. At the same time atomic hydrogen is released or evaporated by the layer. The evaporation flux is chosen such that the amount of hydrogen in the layer is 10 %, i.e., the typical device quality hydrogen content (the model of a hydrogen rich overlayer is neglected).

For each neutral particle the boundary conditions are formulated, which leads to a coupled set of equations which is solved to obtain the boundary conditions for all neutral density balance equations.

#### IV.A.7. Electron energy distribution function

In order to calculate the rates for electron impact collisions and the electron transport coefficients (mobility  $\mu_e$  and diffusion coefficient  $D_e$ ) the electron energy distribution function has to be known. This EEDF,  $f(\vec{r}, \vec{v}, t)$ , specifies the number of electrons at position  $\vec{r}$ , with velocity  $\vec{v}$  at time  $t$ . The evolution in space and time of the EEDF in the presence of an electric field is given by the Boltzmann equation [231]:

$$\frac{\partial f}{\partial t} + \vec{\nabla}_{\vec{r}} \cdot (\vec{v}f) - \vec{\nabla}_{\vec{v}} \cdot \left( \frac{e}{m_e} \vec{E}(\vec{r}, t) f \right) = \left( \frac{\partial f}{\partial t} \right)_{coll} \quad (34)$$

where  $(\partial f / \partial t)_{coll}$  is the collision term. Some approximations are to be made, as this equation is complex (details can be found in [189]).

First, it is assumed that the EEDF is spatially and temporally constant, which is allowed if the energy relaxation time of the EEDF is much shorter than the RF-cycle duration, and if the relaxation length is much smaller than the typical gradient scale length. This assumption implies a spatially and temporally constant electric field. It reduces the Boltzmann equation to a problem exclusively in the velocity space. In the collision term only electron-neutral collisions are considered, because RF plasmas are weakly ionized. The inelastic collisions considered are ionization, dissociation, excitation, and attachment (see also Tab 2). The crude approximation is relaxed in the fluid model by the introduction of the spatial and temporal dependency of the average electron energy. The collision rates, transport coefficients, and average electron

energy are calculated in a range of electric fields  $E$  for a given composition of the background neutrals. The results are then tabulated to obtain a look-up table of the rates and transfer coefficients as a function of the averaged electron energy. In the fluid model the (time and space dependent) average electron energy is calculated with the energy balance equation. The validity of this assumption has been demonstrated by [175,221,230,234,239].

Second, a further simplification of the Boltzmann equation is the use of the two term spherical harmonic expansion [231] for the EEDF (also known as the Lorentz approximation), both in the calculations and in the analysis in the literature of experimental data. This two term approximation has also been used by Kurachi and Nakamura [212] to determine the cross-section for vibrational excitation of  $\text{SiH}_4$  (see Tab. 2). Due to the magnitude of the vibrational cross section at certain electron energies relative to the elastic cross sections and the steep inclination of the vibrational cross section, the use of this two term approximation is of variable accuracy [240]. A Monte Carlo calculation is in principle more accurate, because in such a model the spatial and temporal behavior of the EEDF can be included. However, a Monte Carlo calculation has its own problems such as the large computational effort needed to reduce statistical fluctuations.

The method to calculate the rates of electron impact collisions as used in this model has been studied by Meijer *et al.* [230] and has also been used by others [221,234].

#### IV.A.8. Calculation scheme

The system of equations of the fluid model as well as the Boltzmann equation of the EEDF must be untangled numerically. The balance equations are spatially discretized (129 grid points in this study) using the Sharfetter-Gummel exponential scheme [241]. An implicit second order method is used to numerically treat the time evolution of the balance equations [220]. The electron transport equation is solved simultaneously with Poisson's equation to avoid numerical instabilities. A Newton method is used to solve the resulting set of nonlinear equations. This approach limits the required number of time discretization steps (80 time steps per RF-cycle in this study). Doubling the number of grid points and time steps resulted in a maximum relative change of the partial pressure of  $\text{SiH}_4$ ,  $\text{H}_2$ , and  $\text{Si}_2\text{H}_6$  and the deposition rate of  $10^{-2}$  and  $10^{-4}$ , respectively. A multigrid technique is used to enhance the convergence of the fluid model [242]. Further details about the numerical techniques of the fluid model and the Boltzmann solver can be found elsewhere [175,243].

The fluid model calculates the density profiles based on the electron impact collision rates calculated by the Boltzmann solver. This Boltzmann solver needs the densities of the background neutrals as input. Therefore, the fluid model and the Boltzmann solver are run alternately until the changes in the densities of the background neutrals are less than  $10^{-4}$ . Convergence of the fluid model is obtained when the relative change  $\epsilon_n$  of the discharge parameters (density profiles and potential profile) at the beginning of two subsequent RF-periods ( $n$  and  $n + 1$ ) is less than  $10^{-6}$ . To check whether this criterion is sufficient, some calculations were continued until  $\epsilon_n$  reached a value  $10^{-7}$ . In those cases, the partial pressures of  $\text{SiH}_4$ ,  $\text{H}_2$ , and  $\text{Si}_2\text{H}_6$  and the deposition rate differed relatively at most  $10^{-5}$  from the results obtained with  $\epsilon_n$  equal to  $10^{-6}$ .

## IV.A.9. Simulation results

The 1D fluid discharge model has been applied to the ASTER deposition system (see Sec. IID). The deposition reactor has an inner volume of 10 liter and an inner diameter of 20 cm. The upper electrode is grounded, c.f. Fig. 4a, and the powered electrode is located 2.7 cm lower. Other typical silane-hydrogen discharge parameters are summarized in Table 4.

Using these discharge settings in the model yields the following partial pressures for the three background neutral particles considered: 7.72 Pa for  $\text{SiH}_4$ , 11.51 Pa for  $\text{H}_2$ , and 0.55 Pa for  $\text{Si}_2\text{H}_6$ . With these partial pressures, and the data on electron collisions as given in Table 2, first the EEDF can be calculated, followed by the calculation of the electron transport coefficients and electron impact rates. From a comparison of a Maxwellian EEDF and the two term Boltzmann EEDF as calculated here, it follows that the Maxwellian EEDF underestimates the EEDF at lower energies and overestimates it at higher energies [189].

In Fig. 15 the electron impact rates for the various reactions (Tab. 2) are shown as function of the average electron energy. In a typical silane/hydrogen discharge the average electron energy amounts to about 5 eV. As can be seen in Fig. 15, at an energy of 5 eV the dissociation of silane and disilane, and the vibrational excitation of silane have the highest rates.

Ohmic heating of the electrons amounts to about 80 % of the plasma power. This dissipated power is mainly used for vibrational excitation of the molecules, in contrast to a inert gas discharge (e.g. Ar), where a much larger fraction of the power is used for the acceleration of the positive ions in the sheaths. Another remarkable outcome is the high density of negative ions ( $2.38 \times 10^{16} \text{ m}^{-3}$ ) compared to the electron density ( $7.19 \times 10^{14} \text{ m}^{-3}$ ). This is typical for an electronegative discharge, where the rate coefficient recombination of a negative ion with positive ions is low.

Figure 16 shows the calculated time averages densities of charged (Fig. 16a) and neutral species (Fig. 16b). The  $\text{SiH}_3^-$  density is the highest, in this case, followed by the  $\text{SiH}_2^+$  density: quasineutrality is preserved. It can be clearly seen that the  $\text{SiH}_3^-$  ions are confined in the discharge. The electrons are mobile and therefore much less confined.

TABLE 4. Typical settings for a silane/hydrogen discharge in the ASTER system.

parameter	value	unit
electrode spacing	0.027	m
electrode radius	0.08	m
reactor volume	0.010	$\text{m}^3$
silane flow	30.0	sccm
hydrogen flow	30.0	sccm
pressure	20.0	Pa
gas temperature	400	K
substrate temperature	550	K
RF-frequency	50	MHz
RF-power	5	W

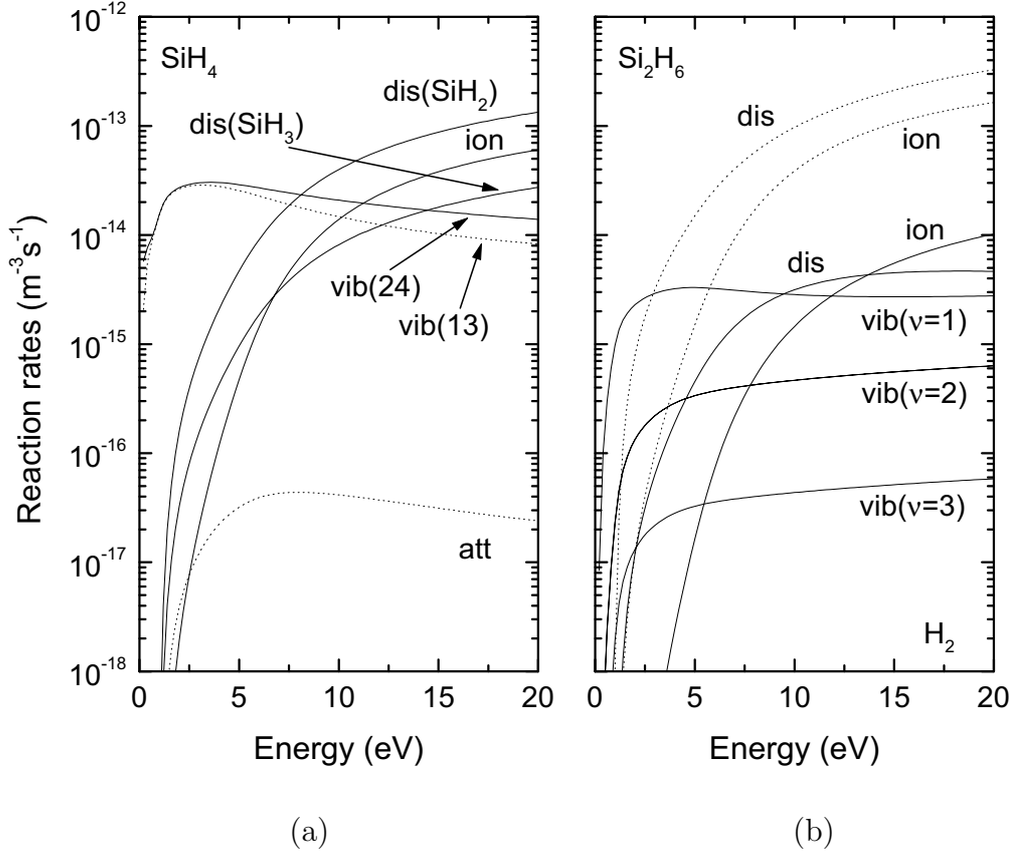


FIG. 15. Electron impact reaction rates as a function of the average electron energy. A 1:1 mixture of  $\text{SiH}_4$  and  $\text{H}_2$  was used, at a total pressure of 83 Pa. (a) reaction rates for  $\text{SiH}_4$ , (b) reaction rates for  $\text{Si}_2\text{H}_6$  (dotted lines) and  $\text{H}_2$  (solid lines). Abbreviations are: 'ion' = ionization, 'dis' = dissociation, 'vib' = vibrational excitation, 'att' = attachment. See Tab. 2 for details and references. (Adapted from G. J. Nienhuis, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

The temporal variation of the ion density profile is small, except in the sheaths. The mass of the ions is too high to respond instantaneously to the changing electric field. Ion transport is only influenced by the time averaged electric field. The total (positive) ion flux to the electrode is  $4.06 \times 10^{18} \text{ m}^{-3}\text{s}^{-1}$ .

From Fig. 16b it is clear that the density of the non-radical neutrals ( $\text{H}_2$ ,  $\text{SiH}_4$ ,  $\text{Si}_2\text{H}_6$ , and  $\text{Si}_3\text{H}_8$ ) is much higher than the radicals density ( $\text{SiH}_3$ ,  $\text{H}$ ,  $\text{Si}_2\text{H}_5$ , and  $\text{SiH}_2$ ). Moreover, from the decreasing density near the electrodes, the species that react can be identified. Note, that the  $\text{SiH}_3$  density is about one order of magnitude larger than the  $\text{SiH}_2$  density.

#### IV.A.10. Comparison with experiments

In this section the results of the 1D model are shown together with data taken from experiments in the ASTER system. Both partial pressures of the background neutrals

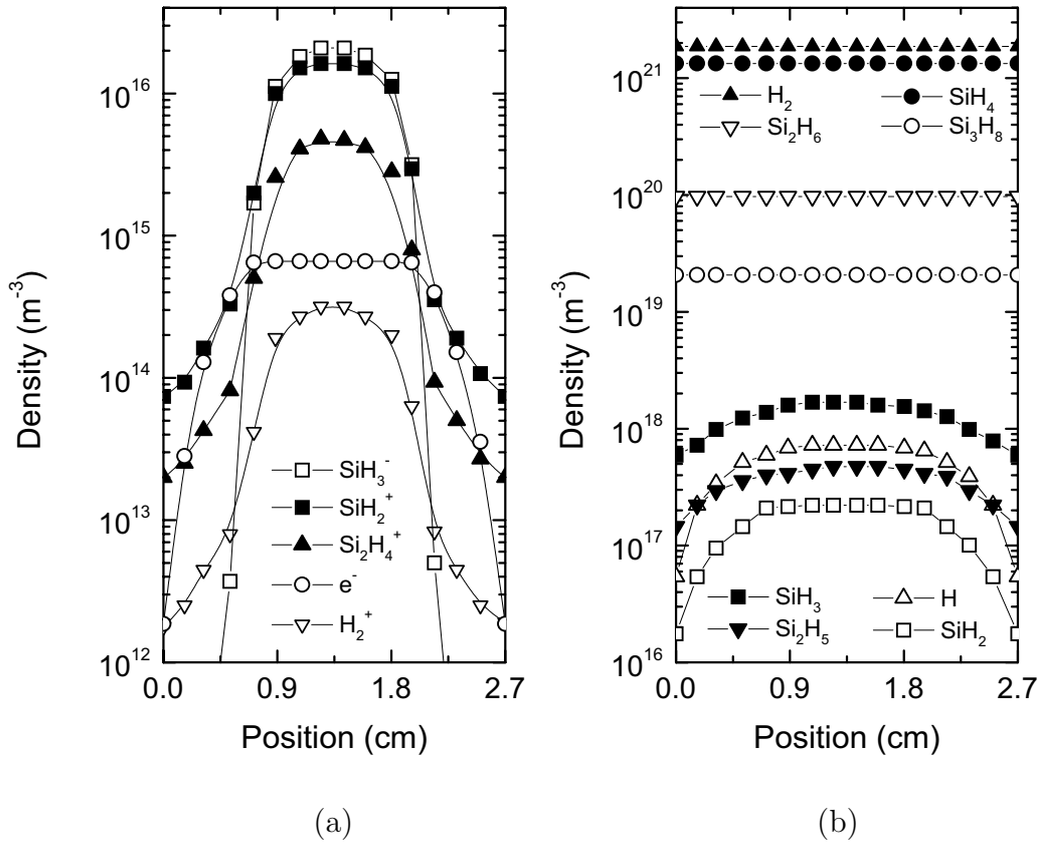


FIG. 16. Time averaged densities of (a) charged, and (b) neutral species, for the discharge settings given in Tab. 4. Note the difference in density-scales. (Compiled from G. J. Nienhuis, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

( $\text{H}_2$ ,  $\text{SiH}_4$ , and  $\text{Si}_2\text{H}_6$ ) as well as deposition rates are accessible in experiment and model. The comparison is made as a function of one of the discharge parameters: the RF-frequency, the RF-power, or total pressure in the reactor. Other process parameters have been kept constant, see Tab. 4.

Ofcourse, care should be taken when comparing the experimental data and the modeling results; the model is an approximation and the experiment has its uncertainties. The most important approximation is that the model is one-dimensional. A main uncertainty of the experiment is the relation between *source* power and *plasma* power. The source power is defined as the power delivered by the power generator. In experiments, the source power is a discharge setting (Tab. 4). The plasma power is the power dissipated by the plasma. As a rule, this plasma power is smaller than the source power, due to losses in the matching network, which matches the plasma impedance to the impedance of the source power generator, i.e.,  $50 \Omega$  [180]. From comparison of experimental data with modeling results (a.o. [220]) in a wide range of discharge settings it is inferred that the plasma power is approximately 50 % of the source power. This is a crude approximation, as this efficiency does depend on discharge settings and the resulting discharge impedance. Agreement, therefore, is not expected between experimental and modeling results. Tendencies, however, will be clear from the results.

Another important problem to address is the regime in which the discharge is operating, either the  $\alpha$ - or dust-free regime, or the  $\gamma'$ -regime, where dust plays an important role in the discharge [244] (see Sec. VI A 6). Dust particles consist of large clusters of Si and H atoms, and are negatively charged by attachment of free electrons. They can be considered as an additional wall surface. Consequently, the free electron density decreases. In order to maintain the discharge, i.e., the amount of ionization, the average electron energy has to increase. This also leads to an enhanced dissociation of  $\text{SiH}_4$  and thus to a larger radical production.

Generally, at low pressures and at low power levels the discharges operate in the  $\alpha$ -regime. At a certain critical pressure, which depends more or less upon the other discharge parameters, the discharge makes a transition from the  $\alpha$ - to the  $\gamma'$ -regime. Modeling does not take dust effects into account. The discharge settings used are interesting, because at increasing total pressure the discharge regime is changed from  $\alpha$  to  $\gamma'$ , at about a critical pressure of 30 Pa.

#### IV.A.10.a. Pressure variation

In Fig. 17 the effects of total pressure are shown, for the relative pressures, i.e., the ratio of the partial pressure and total pressure, of silane, hydrogen, and disilane (Fig. 17a) and the deposition rate (Fig. 17b). The RF-frequency is 50 MHz, and the plasma power is 5 W. The relative pressure of hydrogen slowly increases, and the relative pressure of silane slowly decreases, both in model as well as in experiment. This is caused by an increase in silane depletion at higher total pressures, which results from a higher power dissipation by the electrons. At 16 Pa the fraction of dissipated power that is used to heat electrons is 79 %, at 50 Pa this increases to 97 %. This increase leads to a higher dissociation rate of silane, and explains the increased depletion.

The relative pressure of disilane increases as a function of total pressure due to both the increased production of radicals, which is a result of increased dissociation of silane, as well as the shorter gas volume reaction times and longer diffusion times to the walls, which result from increasing the pressure.

Below 30 Pa the trends between model and experiment agree. The model slightly underestimates the silane partial pressure by about 10 %, and overestimates the hydrogen pressure by about 6 %. Possible causes for this discrepancy are the simple 1-D geometry of the model, the assumed relation between plasma power and source power, and the approximation for the pumping by using an average residence time. The discrepancy for the disilane data might a.o. be explained by questioning the dissociation branching ratio of silane. Layeillon *et al.* [195,196] have discussed this, and conclude that dissociation of silane should result in a larger fraction of  $\text{SiH}_3$ .

At about 30 Pa, the experimental deposition rate (Fig. 17b) clearly shows a sudden increase, which is not revealed by the model. In addition also the partial pressure silane decreases (and of hydrogen increases). These features have been observed in other experiments as well [245–248], and are caused by the transition from the  $\alpha$ - to the  $\gamma'$ -regime.

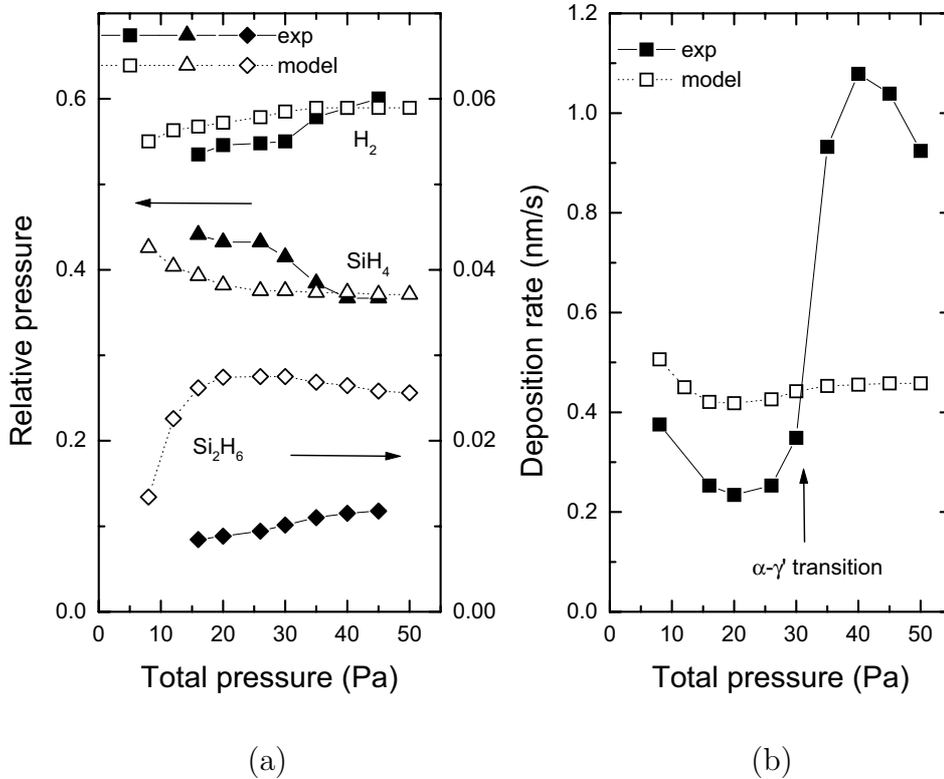


FIG. 17. The relative pressure (a) (i.e., the ratio of the partial pressure and total pressure) of  $H_2$ ,  $SiH_4$ , and  $Si_2H_6$  and the deposition rate (b) as a function of total pressure at an RF-frequency of 50 MHz and a plasma power of 5 W. Other discharge settings are given in Tab. 4. Modeling results are in dotted lines and open symbols, experimental data in solid lines and filled symbols. Note the sudden increase at 30 Pa, i.e., the transition from the  $\alpha$ - to the  $\gamma$ '-regime. (Compiled from G. J. Nienhuis, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

#### IV.A.10.b. RF-frequency variation

In Fig. 18 the effects of RF-frequency are shown, for the the partial pressures of silane, hydrogen, and disilane (Fig. 18a) and the deposition rate (Fig. 18b). The total pressure is 16 Pa, and the plasma power is 5 W. The discharge is in the  $\alpha$ -regime.

Both model and experiment show the same tendency: the dissociation of silane increases as a function of RF-frequency. This can be explained following the arguments from Heintze and Zedlitz [236]. They found that the power dissipated by the ions in the sheaths decreases as a function of RF-frequency. This is confirmed by the modeling results: at 13.56 MHz about 50 % of the power is used for the acceleration of the ions, whereas at 80 MHz this is reduced to about 15 %. Hence, at higher RF-frequencies more power is used for the heating of electrons, which in turn leads to a higher dissociation of silane. As a result, more hydrogen and silane radicals are produced, and more molecular hydrogen is formed. A saturation is observed, when all power is consumed by the electrons.

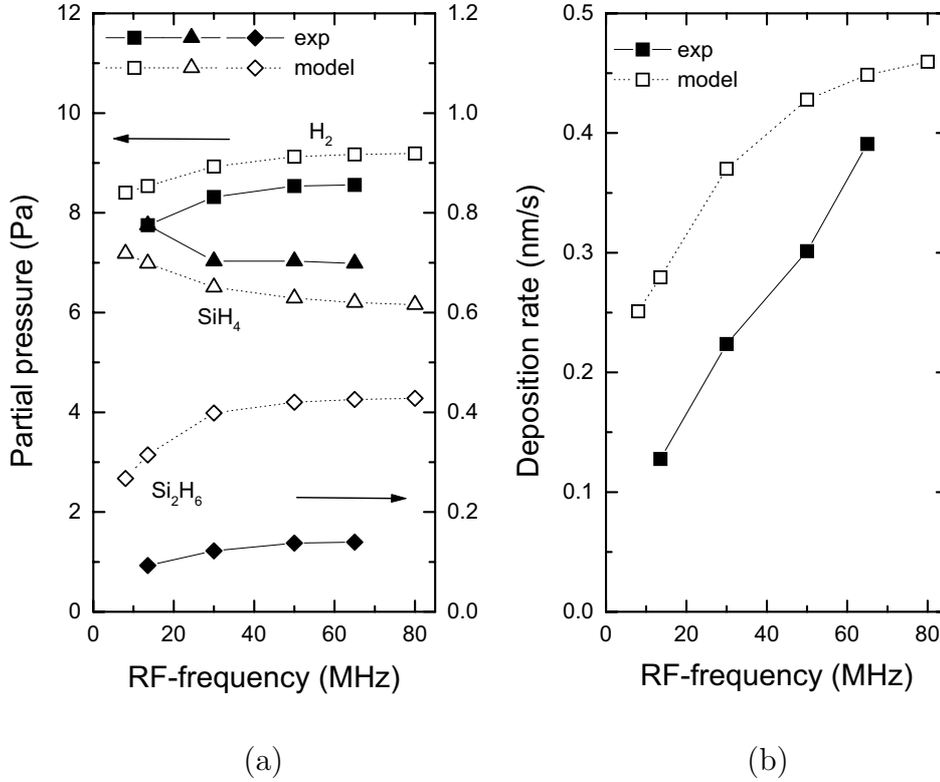


FIG. 18. The partial pressure (a) of H<sub>2</sub>, SiH<sub>4</sub>, and Si<sub>2</sub>H<sub>6</sub> and the deposition rate (b) as a function of RF-frequency at a total pressure of 16 Pa and a plasma power of 5 W. Other discharge settings are given in Tab. 4. At these settings the discharge is in the  $\alpha$ -regime. Modeling results are in dotted lines and open symbols, experimental data in solid lines and filled symbols. (Compiled from G. J. Nienhuis, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

The experimentally found linear increase of the deposition rate as a function of frequency is not seen in the modeling results, see Fig. 18b. This linear increase has also been measured by others [119,120,249], up to an RF-frequency of 100 MHz. Howling *et al.* [250] have measured this linear relationship, while taking special care that the effective power is constant as a function of frequency.

A possible explanation for the difference in tendencies of the deposition rate between experiment and model is that in the model the surface reaction and sticking coefficients of the radicals are taken to be independent of the discharge characteristics. However, these surface reaction coefficients may be influenced by the ions impinging on the surface [251]. An impinging ion may create an active site (or dangling bond) at the surface, which enhances the sticking coefficient. Recent experiments by Hamers *et al.* [163] corroborate this: the ion flux increases as a function of RF-frequency. However, Sansonnens *et al.* [252] show that the increase of deposition rate cannot be explained by the influence of ions only.

The discrepancy may also be caused by the approximations in the calculation of the electron energy distribution function. This EEDF is obtained by solving the two term Boltzmann equation, assuming full relaxation during one RF-period. When the RF-

frequency becomes comparable to the energy loss frequencies of the electrons, it is not correct to use the time independent Boltzmann equation to calculate the EEDF [253]. The saturation of the growth rate in the model is not caused by the fact that the RF-frequency approaches the momentum transfer frequency  $\nu_{me}$  [254]. This would lead to a less effective power dissipation by the electrons at higher RF-frequencies and thus to a decrease of the deposition rate at high frequencies compared to lower frequencies.

Another possible explanation is the approximation that the vibrationally excited silane molecules are treated in the model as ground state molecules. This is based on the assumption that collisional de-excitation in the gas phase and at the walls is not fast enough. The densities of vibrationally excited molecules is thus neglected. The fraction of the total power used for vibrationally exciting silane molecules increases as a function of RF-frequency, which may result in a non-negligible excited state density. Inclusion of vibrationally excited silane molecules leads to a higher radical production and deposition rate. Vibrationally excited silane molecules have a lower threshold for electron dissociation than a ground state molecule. Also, multivibrational excitation may lead to homogeneous pyrolysis ( $e + \text{SiH}_4 \rightarrow \text{SiH}_4^{vib} \rightarrow \text{SiH}_2 + \text{H}_2$ ). Perrin *et al.* [192,211] have ruled out this pyrolysis channel, because of the high rates of relaxation for vibrationally excited silane molecules, but this observation was made for an RF-frequency of 13.56 MHz. In addition, vibrationally excited silane molecules have a drastically increased cross-section for dissociative attachment [192,255,256] which increases the production of negative ions and the loss of electrons. In order to sustain the discharge a higher average electron energy is required, which also leads to a higher dissociation.

#### IV.A.10.c. Plasma power variation

In Fig. 19 the effects of RF-power are shown, for the partial pressures of silane, hydrogen, and disilane (Fig. 19a) and the deposition rate (Fig. 19b). The total pressure is 40 Pa, and the RF-frequency 50 MHz. The discharge is in the  $\gamma'$ -regime. Results are shown as a function of the supposed effective RF-power, i.e., 50 % of the power set at the power source.

The decrease of the silane partial pressure and the concomitant increase of the hydrogen partial pressure as a function of plasma power can be understood in terms of the increased electron density and electron energy. Both lead to a higher dissociation of silane and hydrogen. The silane radicals and atomic hydrogen thus created diffuse to the electrodes, where deposition takes place and molecular hydrogen is formed.

The partial pressure of disilane as a function of plasma power first increases due to the higher production of silane radicals. Above a certain power the increase in disilane dissociation is higher than the increase in the disilane production, hence the disilane partial pressure decreases again. A similar behavior has been observed by Kae-Nune *et al.* [217].

In Fig. 19b the deposition rate is seen to increase as a function of plasma power. Andújar *et al.* [246] also measured an increase of the deposition rate as a function of the total power. They found that the relation between deposition rate and electrical power depends on the pressure. At pressures lower than about 20 Pa, their measured deposition rate increases linearly with power, which is the same relation as shown by the 1D discharge model. However, at pressures higher than 20 Pa, there is a saturation

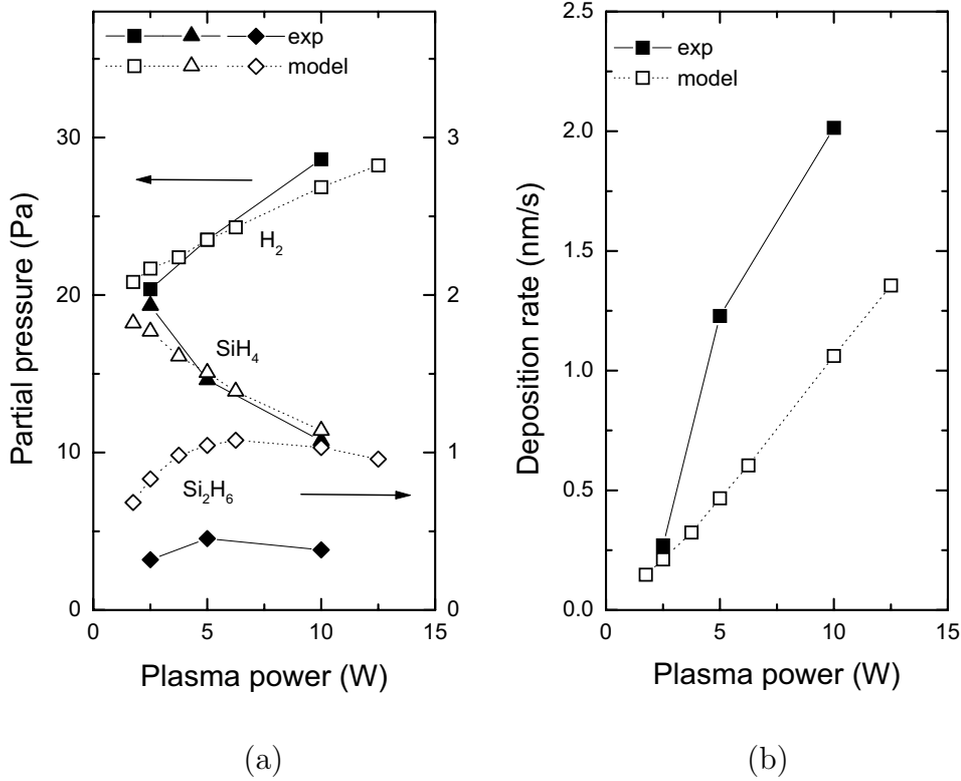


FIG. 19. The partial pressure (a) of H<sub>2</sub>, SiH<sub>4</sub>, and Si<sub>2</sub>H<sub>6</sub> and the deposition rate (b) as a function of plasma power at a total pressure of 40 Pa and an RF-frequency of 50 MHz. Other discharge settings are given in Tab. 4. At these settings the discharge is in the  $\gamma'$ -regime. Modeling results are in dotted lines and open symbols, experimental data in solid lines and filled symbols. (Compiled from G. J. Nienhuis, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

of the deposition rate as a function of the power, which is similar to the experiments shown here (Fig. 19b). Andújar *et al.* [246] conclude that at 20 Pa the discharge regime is changed from the  $\alpha$ - to the  $\gamma'$ -regime. This value of 20 Pa depends on discharge parameters.

#### IV.A.10.d. Sensitivity analysis

A possible cause for the discrepancy between experiment and model is an error in the elementary data (reaction coefficients, cross sections, and transport coefficients) which are obtained from the literature. With a sensitivity study it is possible to identify the most important processes [189].

To calculate the sensitivity with respect to one of the elementary data, its value is changed slightly and the resulting change in the plasma characteristic (i.e., partial pressures of SiH<sub>4</sub>, Si<sub>2</sub>H<sub>6</sub>, and H<sub>2</sub>, and the deposition rate) is calculated. A measure  $\Lambda = \Delta Y / \Delta X$  of the sensitivity is the relative change  $\Delta Y$  of the plasma characteristic

$Y$  divided by the relative change  $\Delta X$  of the elementary data  $X$ . A sensitivity study has been made for a discharge at a pressure of 16 Pa, 5 W effective power, and RF-frequency of 50 MHz. The partial pressures of  $\text{SiH}_4$  and  $\text{H}_2$  are not sensitive ( $-0.01 < \Lambda < +0.01$ ) to any change in the chemical data. The partial pressure of  $\text{Si}_2\text{H}_6$  is most sensitive to the reactions  $\text{SiH}_4 + \text{SiH}_2$  ( $\Lambda = +0.18$ ) and  $\text{Si}_2\text{H}_6 + \text{SiH}_2$  ( $\Lambda = -0.14$ ) and the dissociation branching ratio  $\text{SiH}_3/\text{SiH}_2$  of  $\text{SiH}_4$  ( $\Lambda = -0.20$ ). The correctness of the silane dissociation branching ratio has also been discussed by Layeillon *et al.* [195,196]. They concluded that the branching ratio should result in a larger fraction of  $\text{SiH}_3$  in order to improve modeling results. This is also observed in our study. The growth rate is most sensitive to the surface reaction ( $\Lambda = -0.20$ ). and sticking probability ( $\Lambda = +0.29$ ) of  $\text{SiH}_3$  and to the diffusion coefficients of  $\text{SiH}_2$  ( $\Lambda = +0.12$ ) and H ( $\Lambda = -0.12$ ).

Nienhuis [189] has used a fitting procedure of the seven most sensitive elements in the elementary data (reactions  $\text{SiH}_4 + \text{SiH}_2$  and  $\text{Si}_2\text{H}_6 + \text{SiH}_2$ , dissociation branching ratio of  $\text{SiH}_4$ , surface reaction coefficient and sticking probability of  $\text{SiH}_3$  and diffusion coefficient of  $\text{SiH}_2$  and H). In order to reduce the discrepancy between model and experiment significantly, some values had to be changed with more than two orders of magnitude, which clearly is unrealistic. Uncertainties in literature never are larger than 50 %.

## IV.B. 2D fluid discharge model

### IV.B.1. Model description

In the previous section a 1D discharge model was formulated and results were compared to experimental data. Although this discharge model is sophisticated, there are some disadvantages related to this low dimensionality. Therefore, this model is extended to a 2D discharge model, the extension being only in geometry.

The self-consistent fluid model was developed by Nienhuis [189]. This model is two dimensional (cylindrically symmetric). It is an extension purely in geometry of a previously published one-dimensional model [191]. Only the important aspects of the model are summarized here. A fluid model consists of balance equations for the densities of the various species in the discharge (i.e. electrons, positive and negative ions and neutrals) and an energy balance equation for the electrons. The heavy species (ions and neutrals) are assumed to have a uniform temperature close to the wall temperature, because of the good energy transfer both between the heavy species and between the heavy species and the wall. Heating or cooling of the electrodes is not considered. The electric field in the discharge is calculated using the Poisson equation. For the charged particle fluxes the drift diffusion approximation is used, with the actual electric field for the electrons and an effective field for the ions, in order to account for their inertia [233]. In the 1D model [191] the non-radical neutrals were treated with a model based on perfect mixing between the discharge volume and the discharge-free volume of the reactor vessel. Here, diffusive transport of all neutrals is accounted for, while modeling only the discharge volume.

In a fluid model the correct calculation of the source terms of electron impact collisions (e.g. ionization) is important. These source terms depend on the electron energy

distribution function (EEDF). In the 2D model described here, the source terms as well as the electron transport coefficients are related to the average electron energy and the composition of the gas by first calculating the EEDF for a number of values of the electric field (by solving the Boltzmann equation in the two-term approximation) and constructing a look-up table.

A sticking model is used for the plasma-wall interaction [137]. In this sticking model each neutral particle has a certain surface reaction coefficient which specifies the probability that the neutral reacts at the surface when hitting it. In case of a surface reaction two events may occur. The first event is sticking which in the case of a silicon containing neutral leads to deposition. The second event is recombination in which the radical recombines with a hydrogen atom at the wall and is reflected back into the discharge.

The chemistry (i.e. the species and reactions included) is the same as described for the 1D model. However, here the higher order silanes  $\text{Si}_n\text{H}_{2n+2}$  and silane radicals  $\text{Si}_n\text{H}_{2n+1}$  are limited to  $n < 4$  to reduce the computational effort. Thus,  $\text{Si}_3\text{H}_7$  and  $\text{Si}_3\text{H}_8$  are representative for all silanes with  $n > 2$ . The formation of powder (large silane clusters) is not taken into account in this model. The discharge settings for the calculations shown here are a total pressure of 20 Pa, a power input of  $250 \text{ Wm}^{-2}$ , an RF frequency of 50 MHz and an inlet flow of 30 sccm of  $\text{SiH}_4$  and 30 sccm of  $\text{H}_2$ . This parameter set is chosen because it results in a situation where most of the silane is consumed in a large reactor. This situation is required for economic reasons in industrial applications.

#### IV.B.2. Comparison with experiments, small reactor

The 2D model has been validated and compared with the 1D model, and the experimental data shown in the previous section. Details can be found in [189]. It was found that the partial pressures (averaged in case of the 2D model) differed only by at most 9 %. On the symmetry axis in the 2D model the partial pressures of  $\text{SiH}_4$ ,  $\text{H}_2$ , and  $\text{Si}_2\text{H}_6$  are 7.77 Pa, 11.23 Pa, and 0.64 Pa, respectively. For the 1D model the values are 7.72 Pa, 11.51 Pa, and 0.55 Pa, respectively. Averaged over the whole discharge the 2D model calculates the partial pressures of  $\text{SiH}_4$ ,  $\text{H}_2$ , and  $\text{Si}_2\text{H}_6$  to be 8.05 Pa, 11.16 Pa, and 0.56 Pa, respectively, which is closer to the experimental data (see Fig. 17a). A partial pressure profile of  $\text{SiH}_4$  as calculated with the 2D model for both the volume between the electrodes (discharge volume) and the volume next to the electrodes (discharge-free volume) is shown in Fig. 20. The dimensions are chosen such to mimic the experimental reactor, with an electrode distance  $L$  of 2.7 cm (axial coordinate  $z$ ) and electrode radius  $R$  of 7.6 cm (radial coordinate  $r$ ). The inlet of  $\text{SiH}_4$  is a (cylinder symmetric) ring inlet, shown at the top left of the figure, where the partial pressure is at its highest value. The pump outlet is at the bottom left. It can be seen that the silane partial pressure only varies only 10 % as function of the axial position, and it is decreasing slightly towards the axis of the discharge. This reflects the consumption of silane. Most of the silane flows from the inlet to the outlet, only a small amount diffuses to the discharge region. Similar profiles for the partial pressures of hydrogen and disilane are found.

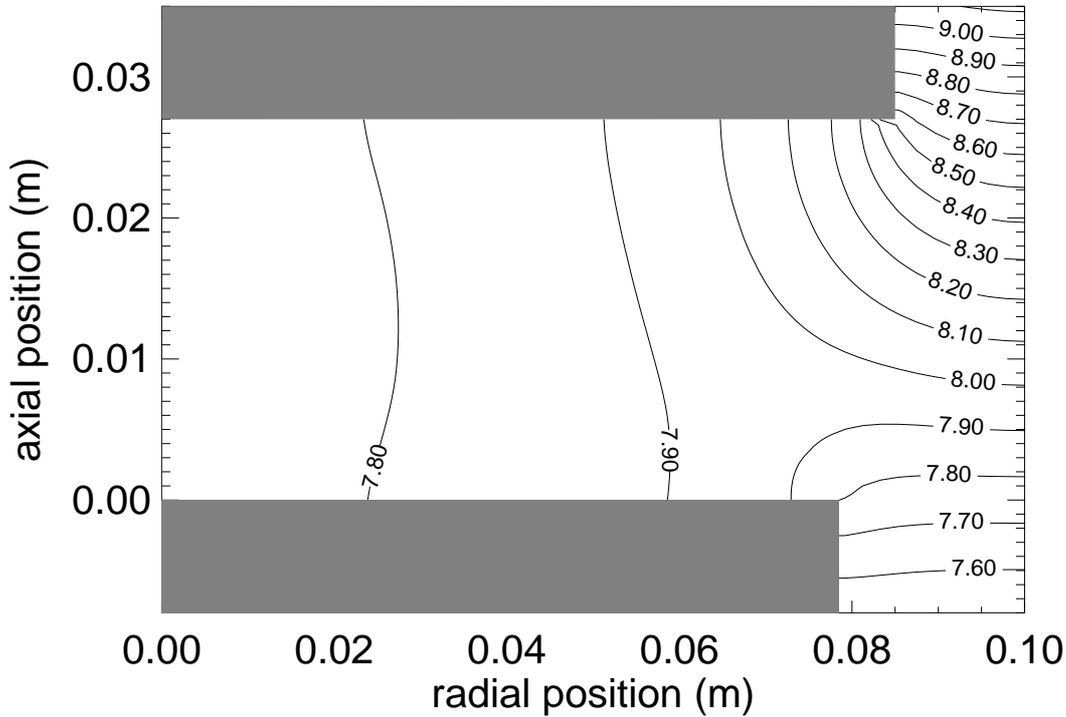


FIG. 20. The partial pressure profile  $\text{SiH}_4$  calculated with the 2D model. Discharge settings are: a total pressure of 20 Pa, a power of 5 W, and an RF-frequency of 50 MHz. (From G. J. Nienhuis, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

Also other discharge characteristics calculated with the 2D model differ only slightly with the ones calculated with the 1D model. The calculated growth rate is 0.419 nm/s versus 0.426 for the 1D model. Electron and ion densities are lower for the 2D model: the electron density is  $5.78 \times 10^{14} \text{ m}^{-3}$  (2D), and  $7.19 \times 10^{14} \text{ m}^{-3}$  (1D), the negative ion density is  $2.00 \times 10^{16} \text{ m}^{-3}$  (2D), and  $2.38 \times 10^{16} \text{ m}^{-3}$  (1D). The ion fluxes also are lower: the ion flux to the grounded electrode is  $3.38 \times 10^{18} \text{ m}^{-2}\text{s}^{-1}$  (2D), and  $4.06 \times 10^{18} \text{ m}^{-2}\text{s}^{-1}$  (1D), the ion flux to the powered electrode is  $3.82 \times 10^{18} \text{ m}^{-2}\text{s}^{-1}$  (2D), and  $4.06 \times 10^{18} \text{ m}^{-2}\text{s}^{-1}$  (1D). This is due to the radial losses in the 2D model. The difference in ion fluxes to the grounded and powered electrode result in a DC self bias voltage of -13.96 V, whereas the experimentally  $V_{DC}$  is -22 V.

From the profile of the average electron energy it is seen that a relatively high average electron energy exists near the edge of the powered electrode ( $z = 2 - 5 \text{ mm}$ ,  $r = 7 \text{ cm}$ ), which is independent of the phase in the RF-cycle [189]. This is caused by the high electric field between the outside of the powered electrode and the grounded shield around it (known as the dark space shield).

The partial pressure profile of  $\text{SiH}_3$  shows a maximum between the electrodes, much like the 1D profile (see Fig. 16). The  $\text{SiH}_2$  partial pressure profile shows a maximum towards the edge of the discharge volume ( $z \approx 8 \text{ mm}$ ,  $r \approx 6.5 \text{ cm}$ ). Both the  $\text{SiH}_3$  and  $\text{SiH}_2$  source term also peak in this region. The high average electron energy in this region of the discharge is responsible for the formation of these radials by electron

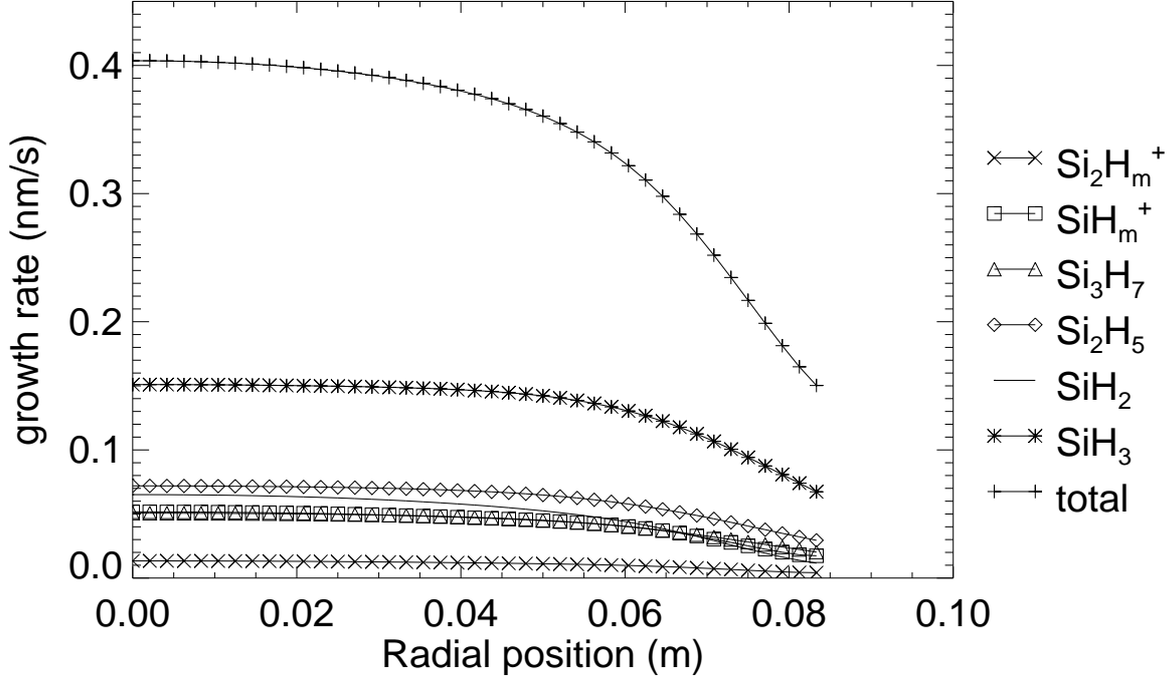


FIG. 21. Deposition rate at the grounded electrode with the contribution of each silicon containing species. Discharge settings are: A total pressure of 20 Pa, a power of 5 W, and an RF-frequency of 50 MHz. (From G. J. Nienhuis, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

dissociation of silane.

The total deposition rate at the grounded electrode and the contribution of each individual species to the total deposition rate is shown in Fig. 21 as a function of radial position. The deposition rate is related to the flux into the surface of silicon containing species,  $\text{SiH}_3$ ,  $\text{SiH}_2$ ,  $\text{Si}_2\text{H}_5$ ,  $\text{Si}_3\text{H}_7$ ,  $\text{SiH}_m^+$ , and  $\text{Si}_2\text{H}_m^+$ . For the discharge conditions used here it follows that  $\text{SiH}_3$  is the radical with the highest contribution to the deposition rate, 37 %. Other radicals contribute 16 % ( $\text{SiH}_2$ ) 17 % ( $\text{Si}_2\text{H}_5$ ), and 13 % ( $\text{Si}_3\text{H}_7$ ). The total contribution of ions to the deposition rate is 17 %, i.e., 13 % for  $\text{SiH}_m^+$ , and 4 % for  $\text{Si}_2\text{H}_m^+$ .

The deposition rate is not uniform over the grounded electrode. A common requirement for large scale reactors is a 5 % difference in deposition rate over a certain electrode area, preferably the complete electrode. Using this, the 2D model would predict a useful electrode area of about 30 %, i.e., an electrode radius of about 4 cm.

In the ASTER reactor deposition experiments were performed in order to compare with the 2D model results. Normalized deposition rates are plotted in Fig. 22 as a function of radial position for data taken at 25 Pa and 18 Pa. The deposition takes place on a square glass plate. For each pressure two profile measurements were performed, each profile perpendicular to the other. A clear discrepancy is present. The use of the simplified deposition model is and explanation for this. Another 2D fluid model recently

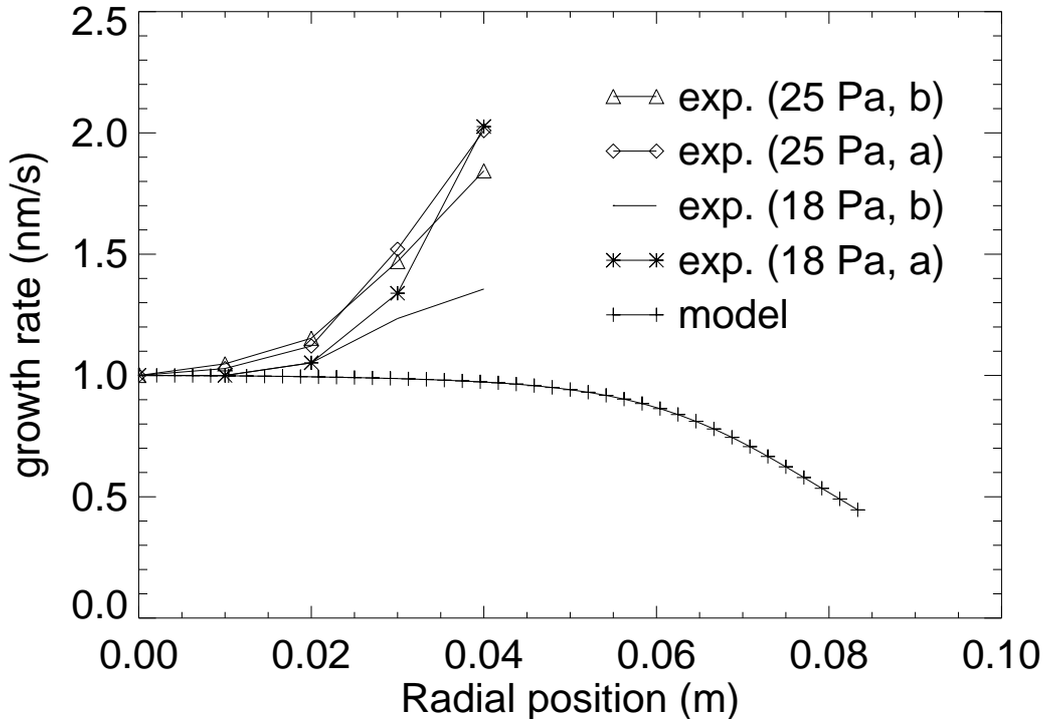


FIG. 22. Normalized deposition rate at the grounded electrode of the 2D model compared to experiments. Discharge settings for the model are: a total pressure of 20 Pa, a power of 5 W, and an RF-frequency of 50 MHz. The experiments are performed at 18 and 25 Pa. The measurements are done for two perpendicular directions,  $x$  and  $y$ . (From G. J. Nienhuis, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

also shows discrepancies between the measured and calculated deposition rate [257], which are attributed to the relative simplicity of the deposition model.

Another explanation for the observed discrepancy might be the influence of the electrode material. In all model calculations it is assumed that the potential at the grounded electrode is zero. In reality, as one generally uses insulating substrates such as glass, other boundary conditions are needed. Nienhuis [189] has reformulated boundary conditions taking into account the actual square  $10 \times 10 \text{ cm}^2$  glass substrate mounted on the circle symmetric grounded electrode. It was found that the deposition rate at the center of the grounded electrode (covered with the glass) was reduced by about 15 %. Going towards the edges of the glass, the deposition rate increases upto the level that is calculated for the full metal electrode. For larger radial positions the deposition rate equals the one for the full metal electrode. In addition, calculation of the contribution of individual species to the deposition rate show that the contributions of  $\text{SiH}_3$  and  $\text{SiH}_2$  increase around the glass/metal transition. It is concluded that the effect of a mixed boundary condition may at least partially explain the observed increase of deposition rate towards the edges of the glass.

### IV.B.3. Comparison with experiments, large reactor

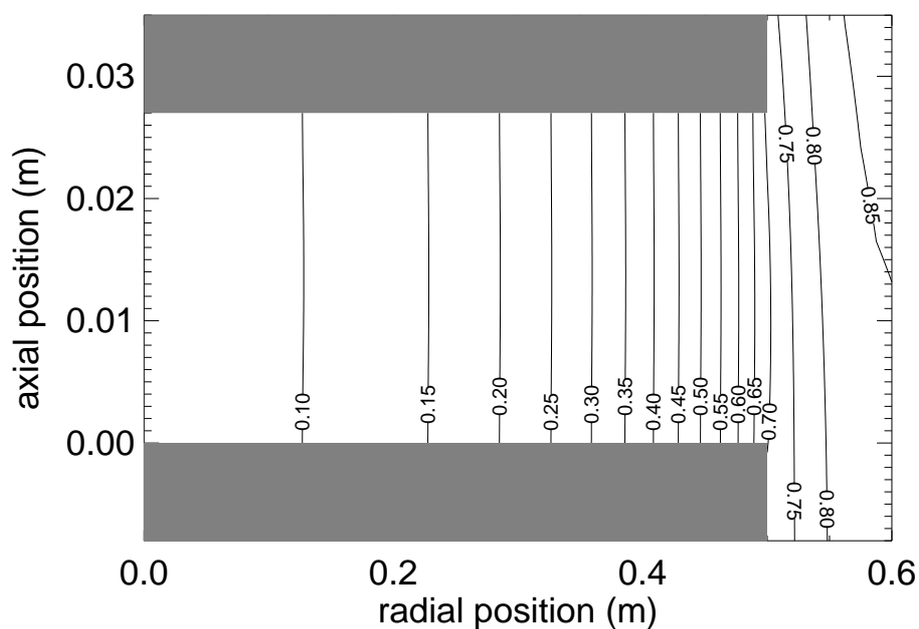
In a small reactor the depletion of silane is usually low and consequently, the background gas composition can be considered homogeneous. In a large reactor the composition varies throughout the discharge volume, as a result of the much higher consumption of silane. For commercial reasons the flow of silane is kept as low as possible, so as to obtain full consumption, however, this may lead to loss of homogeneity. Large reactor systems have recently been modeled using the 2D model by Nienhuis and Goedheer [232]. A reactor is modeled with identical electrode spacing as above (2.7 cm), but with electrode radius of 50 cm and reactor radius 60 cm, which are typical dimensions used in reactors for industrial applications [252]. Two types of gas inlet are modeled, a showerhead, i.e., a powered electrode with small pores through which the gas is introduced [258], and a ring inlet, i.e., an inlet through the ring-shaped area between the grounded electrode and the reactor wall. In both cases, pumping occurs through the area between the powered electrode and the wall.

The discharge settings for the calculations shown here are a total pressure of 20 Pa, a power input of  $250 \text{ Wm}^{-2}$ , an RF frequency of 50 MHz and an inlet flow of 30 sccm of  $\text{SiH}_4$  and 30 sccm of  $\text{H}_2$ . The resulting silane pressure distribution is shown in Fig. 23 as a function of radial position for both the ring inlet (Fig. 23a) and the showerhead inlet (Fig. 23b). In case of the ring inlet the silane pressure distribution is very inhomogeneous, and relatively homogeneous for the showerhead inlet. A strong depletion of silane is seen for the ring inlet going from the outside to the center of the discharge. In fact, the discharge consists of an almost pure hydrogen discharge in the center and a mixture of silane and hydrogen towards the edges. For the showerhead inlet the situation is reverse, albeit the effect is much smaller.

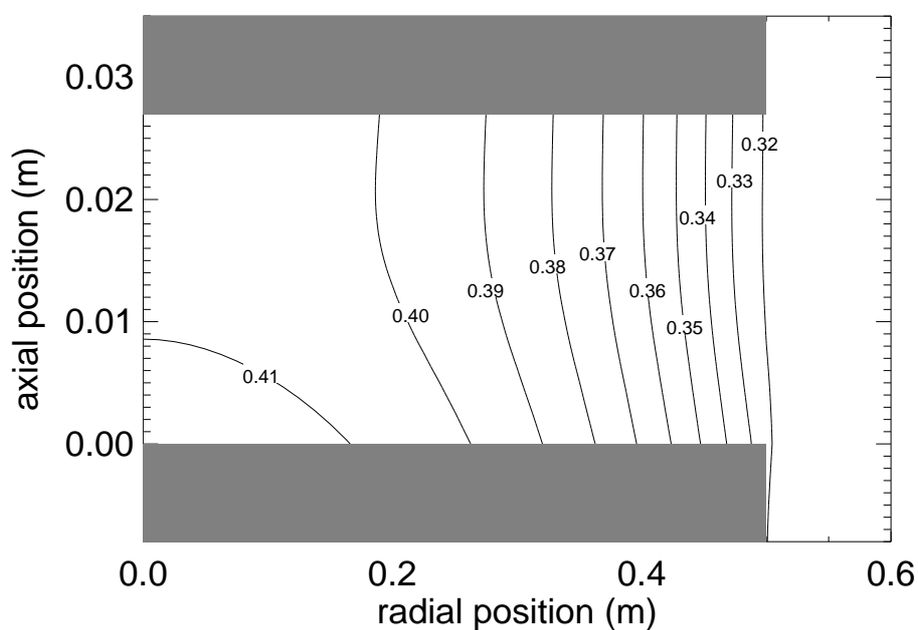
The depletion of silane has large consequences for the uniformity of the deposition, as is shown in Fig. 24. In the case of the showerhead the deposition is very homogeneous, and decreases slowly going from the center to the edge of the discharge, which can be related to the small decrease in silane partial pressure as seen in Fig. 23b. The ring inlet shows a large increase in the deposition rate, when going from the electrode center to the outside. This reflects the high silane consumption, in the opposite direction. The sharp increase of the deposition rate at the very edge of the electrode has also been observed experimentally [259].

The large depletion of silane has consequences for the chemistry. In the electron impact dissociation of silane the two radicals  $\text{SiH}_2$  and  $\text{SiH}_3$  are formed with branching ratio 83 %/17 % [197]; the other product is H (two or one, respectively). When there is enough silane available, the  $\text{SiH}_2$  radical is lost by the insertion reaction  $\text{SiH}_2 + \text{SiH}_4 \rightarrow \text{Si}_2\text{H}_6$ , and via the H-abstraction reaction  $\text{H} + \text{SiH}_4 \rightarrow \text{SiH}_3 + \text{H}_2$  the radical  $\text{SiH}_3$  is produced. Deposition of  $\text{SiH}_2$  and  $\text{SiH}_3$  is the loss process at the walls.

The loss and production depend on silane partial pressure. In case of a relative high partial pressure of silane most  $\text{SiH}_2$  radicals react with silane before reaching the wall. As also the production of  $\text{SiH}_3$  radical is high, the  $\text{SiH}_3$  radical is the main precursor for deposition. This is the case in the small reactor, see Fig. 21. In case of a relative low partial pressure, the reaction time of  $\text{SiH}_2$  with  $\text{SiH}_4$  is longer than the diffusion time to the wall. Together with the lower production of  $\text{SiH}_3$  this results in  $\text{SiH}_2$  to be the main precursor for deposition, see Fig. 23. Depending on the silane partial pressure the main precursor for deposition is different. This will have serious consequences for

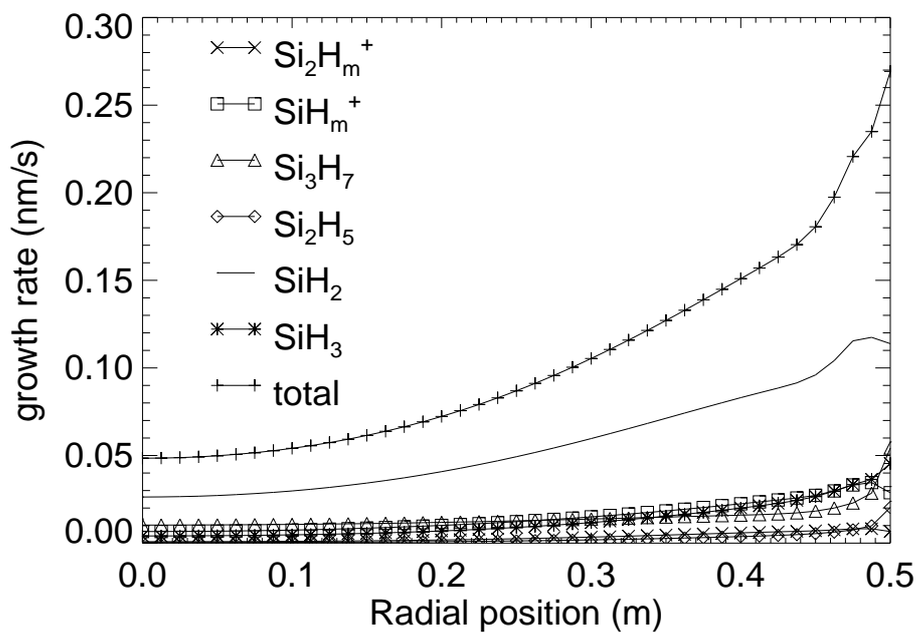


(a)

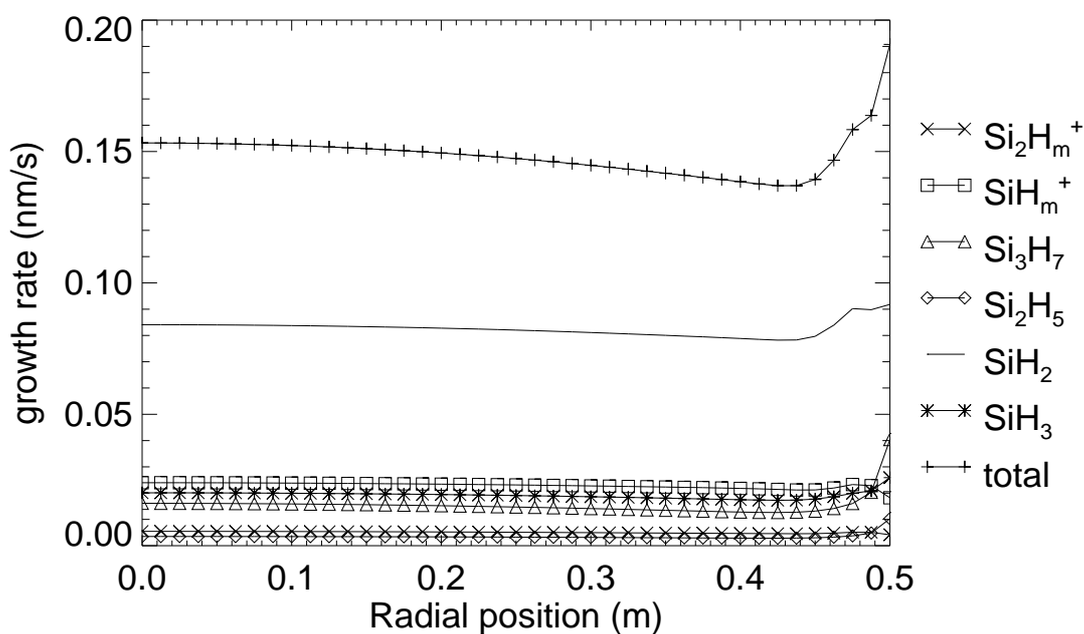


(b)

FIG. 23. The partial pressure of silane (Pa) for a large scale reactor with (a) ring inlet, and (b) showerhead inlet. Note the difference in axial and radial scale. Discharge settings are: A total pressure of 20 Pa, a power of 5 W, and an RF-frequency of 50 MHz. (From G. J. Nienhuis, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).



(a)



(a)

FIG. 24. The deposition rate at the grounded electrode with the contribution of each silicon containing species, for a large scale reactor with (a) ring inlet, and (b) showerhead inlet. Note the difference in uniformity. Discharge settings are: A total pressure of 20 Pa, a power of 5 W, and an RF-frequency of 50 MHz. (From G. J. Nienhuis, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

the quality of the deposited material. The  $\text{SiH}_2$  radical has a much larger sticking probability than the  $\text{SiH}_3$  radical. The structure of a layer deposited with  $\text{SiH}_2$  will be different from a layer deposited with  $\text{SiH}_3$ .  $\text{SiH}_2$  leads to columnar deposition, which is unwanted. So-called device quality material is obtained when deposition occurs via the  $\text{SiH}_3$  radical. Scaling-up of a small to a large reactor therefore is not straightforward.

#### IV.C. Particle-In-Cell discharge models

##### IV.C.1. Particle-in-Cell model principles

In a complete model for the description of plasma deposition of *a*-Si:H should include the kinetic properties of ion, electron and neutral fluxes towards the substrate and walls. The Particle-In-Cell/Monte Carlo (PIC/MC) model is known to provide a suitable way to study the electron and ion kinetics. Essentially, the method consists of the simulation of a (limited) number of computer particles, each of which represents a large number of physical particles (ions and electrons). The movement of the particles is simply calculated from the Newton laws of motion. Within the PIC method the movement of the particles and the evolution of the electric field is followed in finite time steps. In each calculation cycle first the forces on each particle due to the electric field are determined. Then the equations of motion for electrons and ions are integrated. Subsequently collisional processes are treated by the Monte-Carlo method. The chance for a collision for each particle during one time step is calculated using the cross sections for the various processes, and random number generation controls the actual occurrence of the collision. Finally the particle distribution is interpreted into electron and ion densities, and the Poisson equation for the electric field is solved. The movement of ions and electrons in an electric field which is itself directly determined by the particles densities (via the Poisson equation), is referred to as the consistent solution of the electric field.

Despite the simplifications and the incomplete information with respect to the cross sections, the PIC method gives remarkably good results in RF modeling [260,261]. The advantage of a fully kinetic description is correlated with the disadvantage of very time consuming computations. The PIC method allows only for a fully explicit treatment of the particle transport. Numerical instabilities due to large particle velocities and the coupling between particle densities and the electric field can only be avoided by reducing the time step in the simulation, so increasing the computational effort. Large numerical fluctuations in the calculated quantities as a result of the low number of computer particles (low with respect to the real, physical number for ions and electrons) can also not be avoided in PIC simulations.

Some authors have made their code for PIC calculations in RF discharges available for a larger public. In particular Birdsall *et al.* [261–263] developed the XPDP1 code, which is an implementation of the PIC method for RF discharges, written in C for use in UNIX/X-window environments. This XPDP1 code is developed for simulations in one-dimensional RF discharges with plasmas consisting of one type of ion (argon, hydrogen), which have as collisional processes: elastic collisions of electrons and ions with background-gas neutrals, excitation and ionization. The code is built up in a modular way, which makes it easy to modify or add parts of the code, i.e., to make the code suitable for use with other gases with more collisional processes. The code is intended

to run interactively on workstations or PCs using X-windows, and communicates with the outside world via windows and dialog boxes.

The  $\text{SiH}_4/\text{H}_2$  discharge implies the addition of extra processes (ionization, dissociation and attachment for the electrons, recombination for the ions) in the collision component. This resulted in major changes in this part of the code [264,265] Especially the incorporation of collision processes with target particles, which do not form a constant background density, but have a space and time varying density, needed some significant modifications. They were necessary to handle ion-ion-recombination, but may also be used in future, e.g. for ion-electron collisions, and for electron-radical collisions. In the original code partial collision chances were determined via cross sections, in the modified one inverse free paths had to be applied. The cross sections for the electrons were obtained from the literature [266]. The cross sections for ion-neutral scattering had to be estimated. For the energy dependency of the scattering cross section an ion-induced dipole interaction has been assumed [267]. The recombination cross section is directly estimated from rate coefficients [190], and results in a total recombination sink (i.e. integrated over velocity space) identical to the fluid one.

#### *IV.C.2. Hybrid PIC/MC-fluid model*

As a first attempt to modify the code to be able to run simulations on  $\text{SiH}_4/\text{H}_2$  discharges, a hybrid PIC/MC-fluid code was developed [264,265] It turned out in the simulations of the silane/hydrogen discharge that the PIC/MC methods is computationally too expensive to allow for extensive parameter scans. The hybrid code combines the PIC/MC method and the fluid method. The electrons in the discharge were handled by the fluid method, and the ions by the PIC/MC method. In this way a large gain in computational effort is achieved, whereas still kinetic information of the ions is obtained.

The fluid model is a description of the RF discharge in terms of averaged quantities [268,269]. Balance equations for particle, momentum and/or energy density are solved consistently with the Poisson equation for the electric field. Fluxes described by drift and diffusion terms may replace the momentum balance. In most cases for the electrons both the particle density and the energy are incorporated, whereas for the ions only the densities are calculated. If the balance equation for the averaged electron energy is incorporated, the electron transport coefficients and the ionization, attachment and excitation rate can be handled as functions of the electron temperature instead of the local electric field.

Especially for the electrons the fluid model has the advantage of a lower computational effort, compared to the PIC/MC method. Their low mass (i.e. high values of the transport coefficients) and, consequently, high velocities, gives rise to small time steps in the numerical simulation ( $v\Delta t < \Delta x$ ) if a so-called explicit method is used. This restriction is easily eliminated within the fluid model by use of an implicit method. Also, the electron density is strongly coupled with the electric field, which results in numerical instabilities. This requires a simultaneous, implicit solution of the Poisson equation for the electric field and the transport equation for the electron density. This simultaneous, implicit solution for the electric field and the electron density can be deployed within the fluid model and gives a considerable reduction of computational

effort as compared to a non-simultaneous solution procedure [179]. Within the PIC method only fully explicit methods can be applied.

The disadvantage of the fluid model is that no kinetic information is obtained. Also transport (diffusion, mobility) and rate coefficients (ionization, attachment) are needed, which can only be obtained from experiments or from kinetic calculations in simpler settings (eg. Townsend discharges). Experimental data on silane/hydrogen Townsend discharges are hardly available, therefore PIC/MC simulations have been performed.

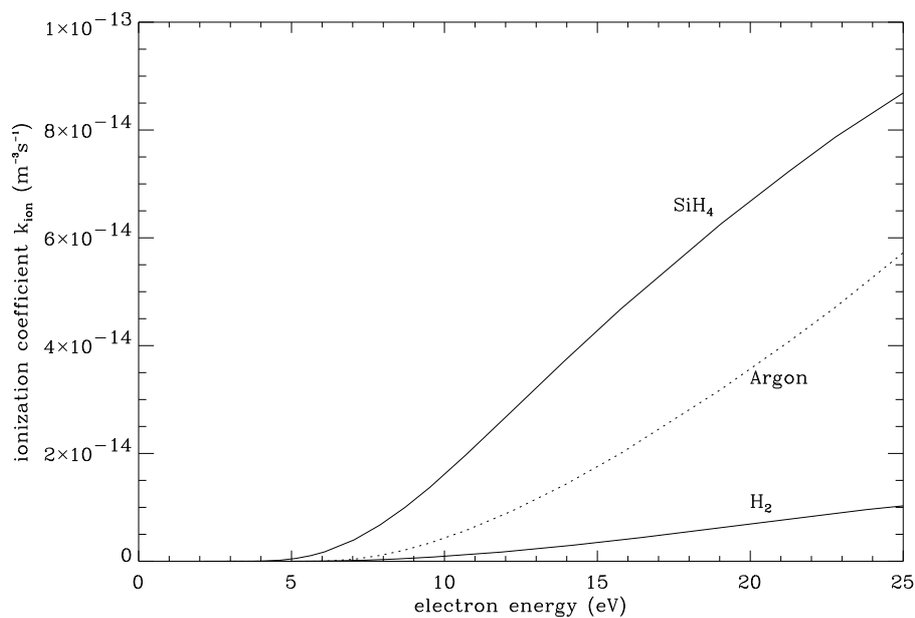
The quasi-Townsend discharge is an infinitely-extended plasma, in which the electrons move in a constant, spatially homogeneous electric field  $E$ . The electrons collide with neutrals in an background gas of infinite extension. The constant electric field replaces the solution of the Poisson equation and makes the PIC method applicable for the electrons within reasonable computational limits. During the particle simulation in the quasi-Townsend discharge, averaged values, over all electron particles  $i$ , of the energy  $\epsilon_i$ , diffusion constant  $D_i$ , mobility  $\mu_i$ , ionization rate  $k_{ion,i}$ , and attachment rate  $k_{att,i}$  are calculated. This results in values for the transport and rate coefficients: the electron transport coefficients,  $D_e$  and  $\mu_e$ , ionization and attachment rates,  $k_{ion}$  en  $k_{att}$ , averaged electron energy  $\langle \epsilon \rangle = 3k_B T_e/2$ . All parameters are obtained for several values of the electric field  $E$ .

Subsequently, because the electron temperature is known as a function of the electric field  $E$ , the temperature  $T_e$  can be used in stead of  $E$  as a parameter for coefficients and rates by elimination of the electric field  $E$ . Thus, the coefficients are available both as a function of  $E$  and of  $T_e$ . Both the local electric field [225,269] and the electron temperature [239,268] have been used as parameter in fluid modeling.

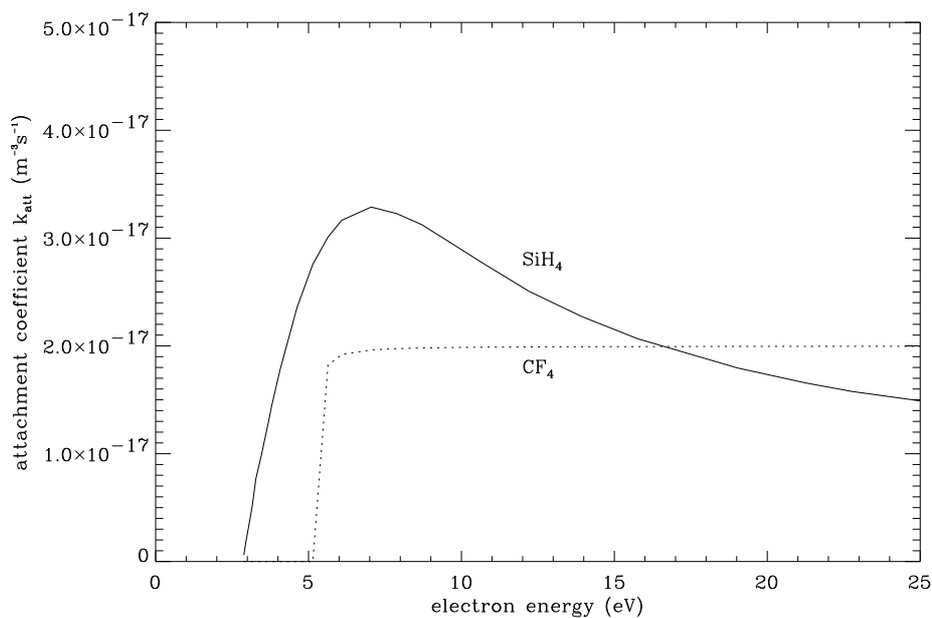
A Townsend PIC simulation was performed for several gas mixtures ( $\text{SiH}_4:\text{H}_2 = 100:0, 50:50, 20:80, \text{ and } 0:100$ ) [264,265]. It was found that the dependency of the electron temperature on the electric field strongly depends on the mixture ratio. As a result the rate coefficient parametrized by the electric field,  $k = k(E)$ , also strongly depends on the mixture ratio. In contrast it was found that the rate as a function of the electron energy,  $k = k(T_e)$ , shows only minor changes ( $\leq 2\%$ ) for different mixing ratios. In Fig 25 the ionization (Fig 25a) and attachment (Fig 25b) rates are given. Here, also two often used coefficients in RF fluid modeling are shown as a reference; an argon ionization rate derived from Townsend experiments (Fig 25a), and a  $\text{CF}_4$  attachment coefficient [179] (Fig 25b).

For the following reasons the choice of the temperature  $T_e$  is more appropriate [268,270]: a) the high-energy tail of the energy distribution function of the electrons determines ionization/attachment; b) both in pure silane and hydrogen and in mixtures,  $T_e$  is a good parameter for the distribution function; c)  $T_e$  as a function of the electric field varies strongly with the mixture ratio  $\text{SiH}_4/\text{H}_2$ .

Methods which compensate for nonequilibrium effects in the situation of  $E$ -parametrized coefficients are very complicated, and are sometimes not firmly grounded. Because the electron temperature gives also reasonable results without correction methods the rate and transport coefficients were implemented as a function of the electron energy, as obtained from the PIC calculations presented in Fig 25.



(a)



(b)

FIG. 25. Simulation results for quasi-Townsend discharges, (a) ionization coefficients for  $\text{SiH}_4$ ,  $\text{H}_2$  and Ar. (b) attachment coefficients for  $\text{SiH}_4$  and  $\text{CF}_4$ .

#### IV.C.3. Simulation of RF discharges in silane/hydrogen with the hybrid model

In the deposition process of amorphous silicon the RF frequency plays an important role. Therefore, in the simulations especially the frequency dependency was studied. Calculations were performed for frequencies of  $\nu_{RF} = 10, 40,$  and  $80$  MHz.

TABLE 5. Simulation results for the hybrid PIC/MC-fluid model for a SiH<sub>4</sub>/H<sub>2</sub> discharge driven with a power of 12 W.

$p$ (mTorr)	$\nu_{RF}$ (MHz)	Density ( $10^{16} \text{ m}^{-3}$ )				Power (W)				$E_{max}$ ( $10^4 \text{ V/m}$ )	Outflux ( $10^{18} \text{ m}^{-2}\text{s}^{-1}$ )	
		$e$	SiH <sub><i>m</i></sub> <sup>+</sup>	SiH <sub><i>m</i></sub> <sup>-</sup>	H <sub>2</sub> <sup>+</sup>	$e$	SiH <sub><i>m</i></sub> <sup>+</sup>	SiH <sub><i>m</i></sub> <sup>-</sup>	H <sub>2</sub> <sup>+</sup>		SiH <sub><i>m</i></sub> <sup>+</sup>	H <sub>2</sub> <sup>+</sup>
250	10	1.8	6.3	4.7	0.18	1.3	7.2	0.0	3.5	11	15	1.1
250	40	4.8	9.4	4.9	3.2	2.7	5.9	0.0	3.4	8.5	28	2.8
250	80	6.9	10.1	4.5	3.9	3.7	5.0	0.0	3.3	7.1	36	3.9
150	10	1.2	6.9	5.7	0.048	1.1	7.5	0.0	3.4	9.4	14	0.9

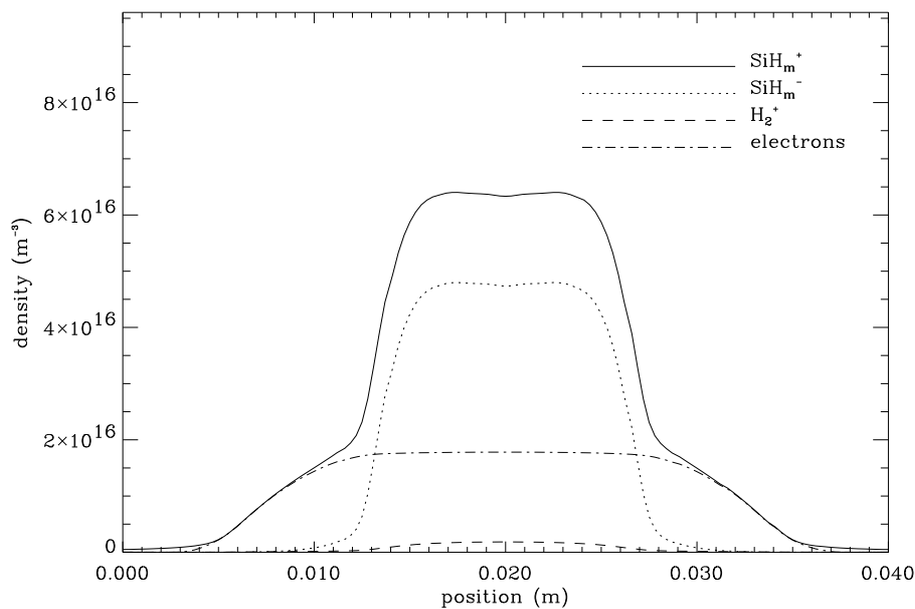
The SiH<sub>4</sub>/H<sub>2</sub> mixture was 20%/80% in all simulations. As most experiments in the ASTER deposition system are performed at a constant total power also the calculations were performed at constant power instead of a given RF voltage. This is especially important, because the power has a large influence on the ion energy spectra at the electrodes. Calculations were performed at a power of  $P = 12$  W, with an electrode area of  $0.010 \text{ m}^2$  and an electrode distance of  $0.04 \text{ m}$ . The pressure was  $p = 250 \text{ mTorr}$ . Also a calculation was performed at  $p = 150 \text{ mTorr}$  for the standard RF frequency of  $10 \text{ MHz}$ .

In Fig. 26 density profiles for two frequencies,  $10 \text{ MHz}$  (Fig. 26a) and  $80 \text{ MHz}$  (Fig. 26b) are shown. The density of the species (electrons, positive ions, negative ions) and power dissipated in these species is presented in Tab. 5, as well as the maximum electric field over the sheath and the outflux of positive ions.

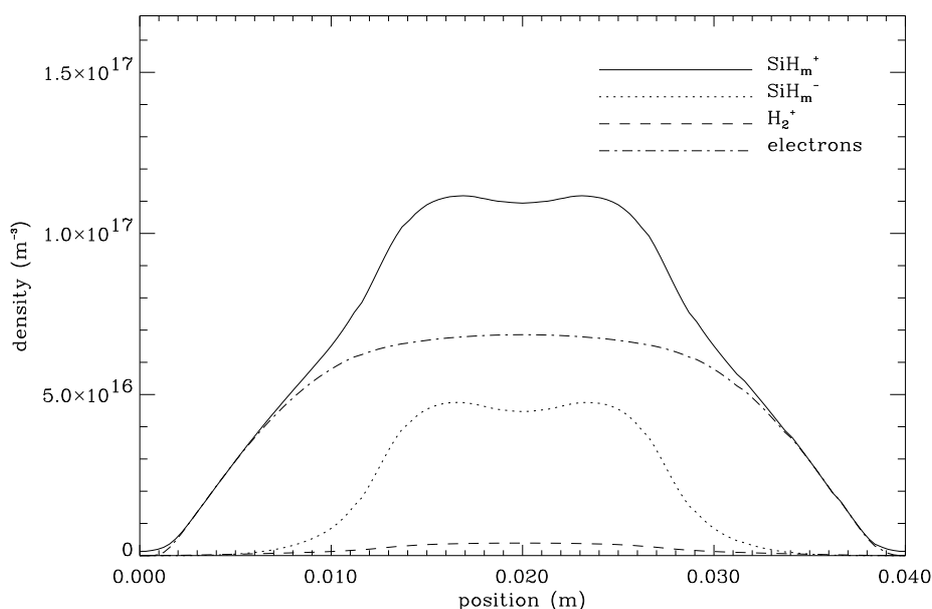
It can be clearly observed that the electron density increases with frequency. The increase of frequency at a constant power results in lower RF voltages and, consequently, lower electric fields in the sheath. The maximum electric field  $E_{max}$  in the sheath diminishes with increasing frequency. The field decrease is too small to prevent the displacement current,  $\epsilon_0 \partial E / \partial t = \epsilon_0 \nu_{RF} \partial E / \partial \tau$ , with  $\tau = t / \tau_{RF}$ , to increase. As a result the electron conduction current in presheath and bulk increases and the heating of electrons becomes higher, whereas the lower potential drop over the sheath causes a lower power dissipation into the ions. The increase of the electron heating results in a higher ionization of silane (SiH<sub>*m*</sub><sup>+</sup>) and hydrogen (H<sub>2</sub><sup>+</sup>). This production is partly compensated for by higher transport losses (outfluxes). The net result is an increasing electron and positive ion density with increasing frequency. On the other hand the negative ion density decreases due to the enhanced recombination between positive ions and SiH<sub>*m*</sub><sup>-</sup>. As a result of the different curves of  $k_{ion}(\epsilon)$  for H<sub>2</sub> and SiH<sub>4</sub> the ionization of H<sub>2</sub> enhances relatively more due to the increasing electron heating than that of SiH<sub>4</sub>. The increased ion fluxes lead to a shrinking of the sheath.

A further result from the increase of power dissipation in the electrons has consequences for the plasma chemistry. Besides the increased ion densities also the production of radicals will be increased, which may lead to higher deposition rates.

It is observed that a decrease of the pressure (from  $p = 250 \text{ mTorr}$  to  $150 \text{ mTorr}$ ) mainly results in a decrease of the densities due to higher transport losses, and in an extension of the sheath due to a higher ion mobility. The electric field and the electron heating diminish slightly for lower pressure. The electron and H<sub>2</sub><sup>+</sup> density, and consequently the H<sub>2</sub><sup>+</sup> outflux are much more influenced by the pressure decrease than the SiH<sub>*m*</sub><sup>+</sup> and SiH<sub>*m*</sub><sup>-</sup> ion densities and the SiH<sub>*m*</sub><sup>+</sup> outflux.



(a)



(b)

FIG. 26. Density profiles in a  $\text{SiH}_4/\text{H}_2$  discharge at 250 mTorr for an RF frequency of 10 MHz (a) and 80 MHz (b).

The ion energy distribution of  $\text{SiH}_m^+$  is presented in Fig 27 for 10 MHz and 80 MHz. Because the total power deposition was kept constant at 12 W, they give insight in the positive ion behavior in deposition experiments. It is seen that both for silane ions (Fig 27) and hydrogen ions (not shown) the ion energy distribution shifts to lower energies for increasing frequency, due to the decreasing electric field and sheath thickness. The saddle structure due to the RF frequency, which can be seen at very low pressures

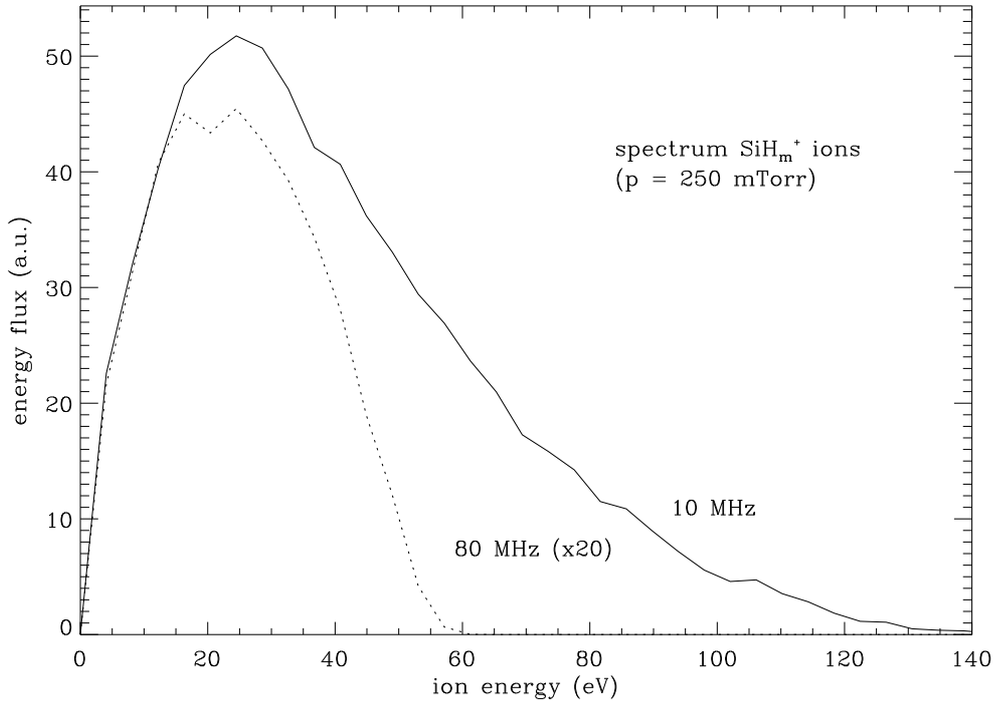


FIG. 27. Ion energy distribution of  $\text{SiH}_m^+$  ions in a  $\text{SiH}_4/\text{H}_2$  discharge at 250 mTorr for an RF frequency of 10 MHz (solid line) and 80 MHz (dotted line).

(< 10 mTorr) [271], has totally disappeared.

For one specific set of discharge parameters a comparison between the hybrid approach and a full PIC/MC method showed that the spectra and the ion densities of the hybrid model showed some deviations with respect to those of the full particle simulation. Nevertheless, due to the computational advantages the hybrid model is appropriate to get insight in plasma behavior under varying parameters, especially with respect to the ion energy distribution at the deposition layer.

#### IV.C.4. Simulation of RF discharges in silane/hydrogen with a full PIC/MC model

The full PIC/MC method has not been used frequently to study  $\text{SiH}_4/\text{H}_2$  discharges [223] because of the large amount of computer time required. Yan and Goedheer [224] have developed a 2D PIC/MC code with some procedures to speed up the calculation. In the code, electrons, positive and negative ions are described in a kinetic way. The capacitively coupled RF discharge in a mixture of  $\text{SiH}_4/\text{H}_2$  has been simulated for pressures below 300 mTorr and frequencies from 13.56 to 65 MHz. Yan and Goedheer [224] studied the effects of frequency on 1) the electron energy distribution (EEDF) in the discharge area, 2) the ion energy distribution (IED) at the substrate, 3) the ion flux to the substrate, and 4) the power dissipation at a constant power density.

The PIC/MC method is based on a kinetic description of the particle motion in velocity position space. Positive and negative ‘superparticles’ move in the self-consistent

electric field they generate. The electric field is obtained by solving the Poisson equation. A Monte Carlo formalism is used to describe the effect of collisions. The displacement between collisions is ruled by Newton's law. There are four kinds of charged particles in the 2D model that is used by Yan and Goedheer [224]: electrons, positive ions ( $\text{SiH}_3^+$ ,  $\text{H}_2^+$ ) and negative ions ( $\text{SiH}_3^-$ ). The negative hydrogen ion ( $\text{H}^-$ ) is not considered because its density is much lower than the other ion densities. For a computationally stable simulation the distance between adjacent points of the grid on which the Poisson equation is solved is taken to be less than several Debye lengths and the product of the timestep and the electron plasma frequency is taken to be much smaller than one. The number of superparticles for each species is around  $10^4$ . The superparticle size for species with a low density (electrons and  $\text{H}_2^+$ ) is between  $10^6$  and  $10^7$ ; for species with a high density ( $\text{SiH}_3^+$  and  $\text{SiH}_3^-$ ) it is between  $10^8$  and  $10^9$ .

The simulation of an electronegative gas discharge converges much more slowly than that of an electropositive discharge. This is mainly caused by the slow evolution of the negative ion density, which only depends on the attachment (to create negative ions) and ion-ion recombination (to annihilate negative ions), both processes with a very small cross section. In addition to the common procedures adopted in the literature [222,223,272,273], such as the null collision method, and different superparticle sizes and time steps for different types of particle, two other procedures were used to speed up the calculation [224].

The first procedure is to use the rate equilibrium equations for electrons and ions at the quasi-steady state for a new guess of the particle densities. In the final quasi-steady balances of the positive ions and negative ions integrated over the whole discharge region will be reached, i.e., the balance for the creation of positive ions ( $\text{SiH}_3^+$ ) by ionization and their loss to the reactor boundary and by recombination, and the balance for the creation of negative ions by attachment and their loss by recombination. These balances are used in the following way. After a number of cycles, the average ionization, recombination and attachment rates are known, as well as the average loss of positive ions to the wall. Also the deviation from the balances is known. The new densities of electrons, positive and negative ions can be guessed by combination of charge neutrality with the balance equations in a global model. These densities will be different from the values obtained in the previous simulation and the number of simulation particles thus has to be adapted. The easiest way to implement the changes is through a change of the size of the superparticles. To do this, only the interior part of the discharge is used, where the negative ions are present. The changes are spatially distributed according to the profile of the negative ions. After the correction the standard PIC/MC calculation is restarted. This procedure is repeated several times during a full simulation and reduces the computational effort considerably.

The second procedure is to use a low number of large particles in the beginning of the simulation. The results of that are used as the initial conditions for a simulation with a large number of small particles. Thus, in the final results, the statistical fluctuations are reduced. Also results from a previous simulation can be used as initial conditions for a new simulation with changed parameters, which saves a large amount of time during parameter scans.

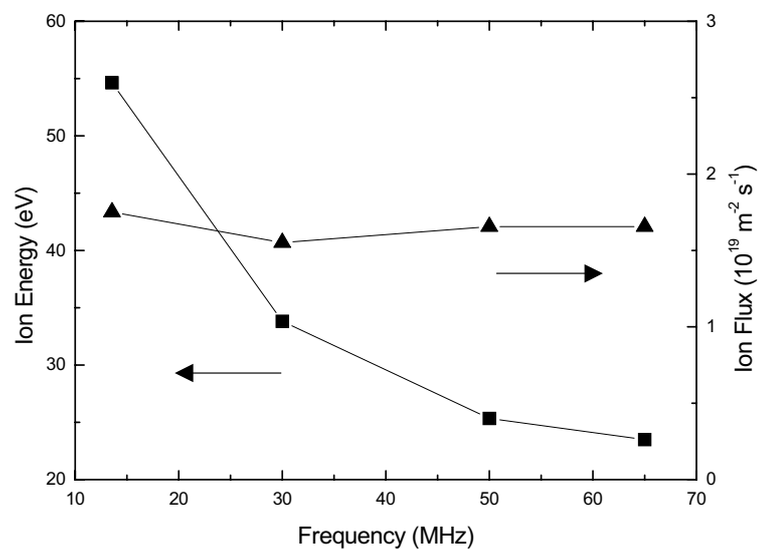
Most of the electron and ion impact reactions that are included in this code are listed in Tab. 2. Added to this are the elastic collisions  $\text{SiH}_4 + e$  [274] and  $\text{H}_2 + e$  [275], the recombination of  $\text{SiH}_3^- + \text{SiH}_3^+ \rightarrow 2 \text{SiH}_3$  [214], and the charge exchange  $\text{H}_2^{+*} +$

$\text{H}_2 \rightarrow \text{H}_2^+ + \text{H}_2^*$  [193]. Note that disilane is not included. The cross sections are taken from the references in Tab. 2. Ion-neutral elastic collision cross sections are obtained according to the Langevin formula [192]. The total number of ion-ion recombinations in the discharge is obtained from the integral of the product of the recombination rate coefficient  $k_{rec}$  and the ion density profiles. The recombination event is attributed to the negative ion; it recombines with the nearest positive ion. In this model the focus is on the energy dissipation, EEDF, IED and ion flux to the wall but not on the details of chemical reactions. Chemical reactions are not included in our code. The experiments described above showed that the background pressure and the gas composition can be regarded as being constant for these dimensions. In the simulation, the pressure and the gas composition are assumed to be constant.

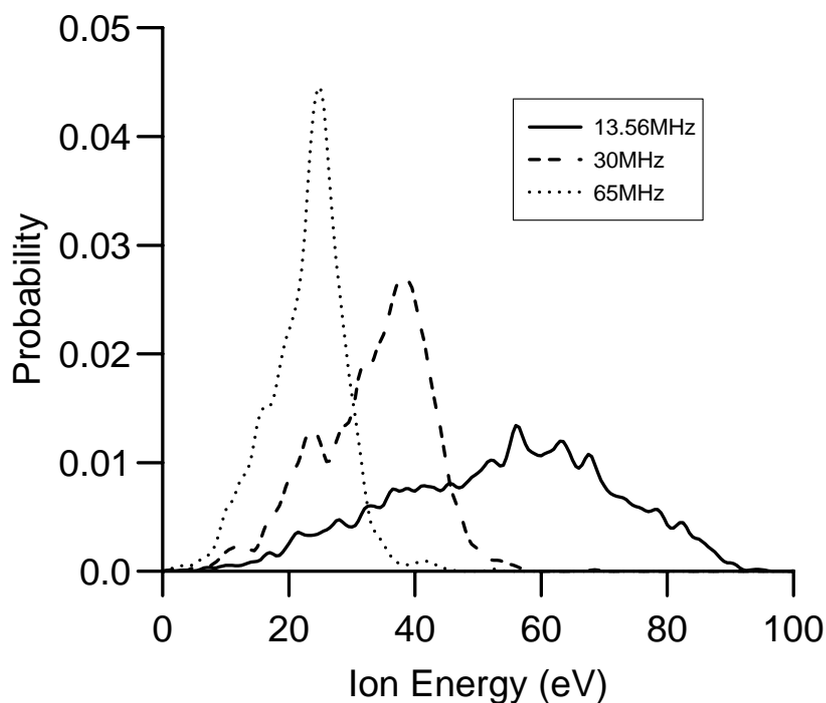
The frequency effects are studied in the cylinder geometry ( $R = 0.08$  m,  $L = 0.027$  m) at a constant power of 25 W, which corresponds to a volume power density  $46$  mW  $\text{cm}^{-3}$ . The pressure is 120 mTorr with 45 %  $\text{SiH}_4$  and 55 %  $\text{H}_2$ . It is found that the RF voltage at this power scales with the frequency as  $V_{RF}\nu_{RF}^{1.15} = C$ , with  $C$  a constant. Because the induced displacement current increases with the frequency, a lower RF voltage is needed at high frequencies to keep the same power density. This relation can explain the increase of the deposition rate with frequency at a constant RF voltage [276] because in that case the dissipated power increases strongly.

The simulations show further that the average electron energy decreases slightly as a function of frequency, from about 5.8 to 5 eV, going from 13.56 to 65 MHz, respectively. The low voltage applied at high frequencies leads to a weaker sheath electric field. Consequently positive ions are pulled out less efficiently, so their density increases. Due to the charge neutrality in the bulk, the electron density also increases, from  $1.8 \times 10^{15}$   $\text{m}^{-3}$  at 13.56 MHz to  $3.2 \times 10^{15}$   $\text{m}^{-3}$  at 65 MHz. On the other hand, the current density  $j \approx e\mu_e E$  in the bulk is constant because of the constant power. Therefore a higher electron density leads to a weaker electric field in the bulk, which is responsible for heating electrons. This causes the decrease of the average energy. The explanation of the increase of the deposition rate with frequency as observed by Curtins *et al.* [277] is that there is a relatively larger population of high-energy electrons. These high-energy electrons would yield a higher ionization and dissociation rate. The simulation described here shows that the density of low-energy electrons increases strongly while the density of high-energy electrons even goes down after combining the high-frequency effects on the EEDF and on the electron density. Stochastic heating [278] does not seem to lead to an increase of the electron energy, probably because its effect is cancelled by the decreasing voltage and by the efficient energy loss in vibrational excitation.

The flux of  $\text{SiH}_3^+$  to the substrate and the average ion energy in this flux at different frequencies are shown in Fig. 28a. The ion flux and ion energy are sampled over one quarter of the electrode radius around the centre of the grounded electrode, which avoids the influence of the edge. The average ion energy decreases quickly with increasing frequency due to the decrease of the RF voltage. The ion flux to the wall remains approximately the same ( $1.7 \times 10^{19}$   $\text{m}^{-2}\text{s}^{-1}$ ) when the frequency is varied. This result is quite different from the results presented by Heintze and Zedlitz [279], where the ion flux showed a significant increase with frequency. Their results, however, are not in agreement with the Bohm criterion because the electron density did not change much while the ion flux changed one order of magnitude [224]. This may be caused by dust formation. Another inconsistency is that the dissociation rate measured by



(a)



(b)

FIG. 28. The effect of RF frequency on (a) ion energy and ion flux to the substrate, and (b) the ion energy distribution. The power is 25 W, and the pressure is 120 mTorr. (From M. Yan and W. J. Goedheer, *Plasma Sources Sci. Technol.* **8**, 349 (1999).)

the same authors [280] only increased by a factor of 1.5 when the frequency was increased from 50 to 250 MHz. The dissociation cross section is larger than the ionization cross section and with lower threshold, see also Fig. 13. This implies that dissociation has higher probability than ionization. The obvious increase in ion flux should therefore be accompanied by at least the same amount of dissociation.

The IED of  $\text{SiH}_3^+$  on the substrate at different frequencies is shown in Fig. 28b. With increasing frequency, the energy distribution profile becomes more peaked. It is attributed to a decrease in the number of collisions experienced by ions, because at higher frequencies the lower applied RF voltage leads to a narrower sheath. The same reason obviously causes the maximum ion energy to decrease with increasing frequency. This result is consistent with the evolution of the estimated maximum ion energy by Dutta *et al.* [281]. But in other experiments by Hamers [163] it was found that the maximum ion energy did not change much.

The power dissipated at two different frequencies has been calculated for all reactions, and is compared to the energy loss to the walls. It is shown that at 65 MHz the fraction of power lost to the boundary decreases by a large amount compared to the situation at 13.56 MHz [224]. In contrast the power dissipated by electron impact collision increases from nearly 47 % to more than 71 %, among which vibrational excitation increases by a factor of 2, dissociation increases by 45 % and ionization stays approximately the same, which agrees with the product of the ionization probability per electron, with the electron density, and also with the ion flux as shown before. The vibrational excitation energy thresholds (0.11 and 0.27 eV) are much smaller than the dissociation (8.3 eV) and ionization (13 eV) ones, and the vibrational excitation cross sections are large too. The reaction rate of processes with a low energy threshold therefore increases more than those with a high threshold.

The simulated rate of vibrational excitation at 30 MHz is already twice the one at 13.56 MHz. At 65 MHz, this rate is three times the rate at 13.56 MHz. The dissociation and ionization rates do not change much. The increase of the vibrational excitation leads to an increase of the density of excited  $\text{SiH}_4$  molecules. This agrees with an  $\text{N}_2$  discharge experiment where the vibrational temperature of  $\text{N}_2$  was observed to rise with the applied frequency [282]. The excited  $\text{SiH}_4$  can participate in various reactions. For example one can speculate that the excited  $\text{SiH}_4$  molecules could enhance pyrolysis ( $\text{SiH}_4 \rightarrow \text{SiH}_2 + \text{H}_2$ ) [283], and consequently contribute to the deposition rate.  $\text{SiH}_2$  can further dissociate to  $\text{SiH}$  and it was observed that the emission of  $\text{SiH}$  shows a similar increased behaviour as the deposition rate [250]. This explanation of an increased deposition via excited  $\text{SiH}_4$  molecules, however, is not supported by the results obtained when the pressure is varied. Simulations were performed at 13.56 MHz and a power of 16 W at pressures ranging from 150 down to 90 mTorr. Although the behavior of the vibrational excitation, dissociation and ionization are quite similar to the behaviour as a result of frequency variation, the deposition rate in the experiments remains the same at low pressures [163,245]. At high pressures the discharge will reach the dusty or  $\gamma'$  regime which is not discussed here. This implies that the vibrational excitation does not influence the deposition directly.

The IED reacts quite differently to pressure changes than to frequency changes. Although the IEDs are somewhat different due to the higher number of collisions in the sheath at higher pressures, they cover approximately the same energy range. It has been reported by Dutta *et al.* [284] that the quality of the deposited material is better

at high frequency than that at lower frequency because the intrinsic stress goes down when the frequency is increased. Since the displacement energy of atoms in the solid is about 20 eV [285], there most likely is an energy range of ions which effectively assists the radical deposition on the surface by influencing the reactivity of the films. At high frequencies more ions are in the effective (lower) energy range than that at a lower frequency (Fig. 28b). This could well cause the higher deposition rate. At different pressures, the energy spectrum of the ions is almost the same, and hence the deposition rate is the same. Chemical reactions are not included here, so the influence of changes in the radical fluxes and their influence on the deposition process is not studied. The simulation and experimental results [247,191], however, indicate that the compositions of the  $\text{SiH}_4/\text{H}_2$  mixture does not change dramatically.

## V. PLASMA ANALYSIS

In order to relate material properties with plasma properties, several plasma diagnostic techniques are used. The main techniques for the characterization of silane/hydrogen deposition plasmas are optical spectroscopy, electrostatic probes, mass spectrometry, and ellipsometry [117,286]. Optical emission spectroscopy (OES) is a noninvasive technique and has been developed for identification of Si, SiH, Si<sup>+</sup>, and H<sup>+</sup> species in the plasma. Active spectroscopy, such as Laser Induced Fluorescence (LIF) also allows for the detection of radicals in the plasma. Mass spectrometry enables the study of ion and radical chemistry in the discharge, either *ex situ* or *in situ*. The Langmuir probe technique is simple and very suitable for measuring plasma characteristics in non-reactive plasmas. In case of silane plasma it can be used, but it is difficult. Ellipsometry is used to follow the deposition process *in situ*.

### V.A. Optical emission

A plasma emits light, as a result of excited species that undergo a radiative transition from a high to a lower level state. By measuring the wavelengths and intensities of the emitted spectral lines, it is possible to identify and quantify neutrals and ions in the plasma. This spectral fingerprint allows for the study of physical and chemical processes that occur in the plasma. Optical emission spectroscopy (OES) is a nonintrusive technique, which can provide spatial and temporal information. By its nature, OES is limited to the detection of light emitting species.

Optical emission spectra nowadays are simply measured using a fiber optic cable that directs the plasma light to a monochromator, which is coupled to a photodetector. By rotating the prism in the monochromator a wavelength scan of the emitted light can be obtained. Alternatively, an optical multichannel analyzer can be used to record (parts of) an emission spectrum simultaneously, allowing for much faster acquisition. A spectrometer resolution of about 0.1 nm is needed to identify species. Quantitative analysis of emission spectra is difficult. As a first assumption the emission intensity from a specific species is proportional to its concentration, but the proportionality constant depends a.o. on electron density and quantum efficiency of the excitation process. As the electron density depends on RF power and gas mixture, and the quantum efficiency on total pressure and gas mixture, the proportionality constant never really is constant [117].

Optical emission is a result of electron impact excitation or dissociation, or ion impact. As an example, the SiH radical is formed by electron impact of silane which yields an excited or superexcited silane molecule ( $e^- + \text{SiH}_4 \rightarrow \text{SiH}_4^* + e^-$ ). The excess energy in SiH<sub>4</sub><sup>\*</sup> is released into the fragments:  $\text{SiH}_4^* \rightarrow \text{SiH}^* + \text{H}_2 + \text{H}$ . The excited SiH<sup>\*</sup> spontaneously release their excess energy by emitting a photon at a wavelength around 414 nm, the blueish color of the silane discharge. In addition, the emission lines from Si<sup>\*</sup>, H<sup>\*</sup>, and H<sub>2</sub><sup>\*</sup> have also been observed at 288, 656, and 602 nm, respectively.

Matsuda and Hata [287] have argued that the type of species that are detectable using OES only form a very small part (< 0.1 %) of the total amount of species present in typical silane deposition conditions. From the emission intensities of Si<sup>\*</sup> and SiH<sup>\*</sup> the number density of these excited states was estimated to be between 10<sup>6</sup> and 10<sup>7</sup>

$\text{cm}^{-3}$ , on the basis of their optical transition probabilities. These values are much lower than radical densities,  $10^{12} \text{ cm}^{-3}$ . Hence, these species are not considered to partake in the deposition. However, a clear correlation between emission intensity of  $\text{Si}^*$  and  $\text{SiH}^*$  and the deposition rate has been observed [288]. From this it can be concluded that the emission intensity of  $\text{Si}^*$  and  $\text{SiH}^*$  is proportional to the concentration of deposition precursors. As the  $\text{Si}^*$  and  $\text{SiH}^*$  excited species are generated via a one-electron impact process, also the deposition precursors are generated via such a process [123]. Hence, for the characterization of deposition discharges information from OES experiments can be used when these common generation mechanisms exist [286].

Laser Induced Fluorescence (LIF) has been used to determine spatial profiles of  $\text{SiH}$  and  $\text{Si}$  radicals at various deposition conditions. These radicals are excited by a tuned pulsed laser, and the fluorescent intensities are a measure of the density of the particle studied. Spatial profiles can be obtained by translating incident laser beam and detection optics. LIF is highly selective and highly sensitive, however it is also a complex technique. Mataras *et al.* [289,290] have compared OES and LIF profiles of  $\text{SiH}^*$  and  $\text{SiH}$ . From the differences observed it is concluded that the generation processes of these radicals are different. The  $\text{SiH}^*$  concentration is related to the magnitude of both cathode and anode sheath potentials, whereas the  $\text{SiH}$  concentration is influenced by the density of energetic electrons in the plasma bulk. As the generation of  $\text{SiH}_2$  and  $\text{SiH}_3$  radicals follow similar pathways, it is inferred that the LIF profile of  $\text{SiH}$  is a good measure of the generation profile of  $\text{SiH}_2$  and  $\text{SiH}_3$  too [286]. This seems to be confirmed by data obtained by infrared laser absorption spectroscopy (IRLAS) [215,291,292], where the  $\text{SiH}_3$  profile is measured to be similar to the LIF  $\text{SiH}$  profile.

Profiles of  $\text{SiH}$  measured by OES as a function of RF voltage have been reported by Bohm and Perrin [184]. In their nearly symmetrical discharge a clear relation exists between  $\text{SiH}$  emission intensity and RF voltage, at low pressure (55 mTorr). Further, the  $\text{SiH}$  profiles are 'M'-shaped, i.e., they peak around the time-averaged sheath boundary, and are lower in the center of the plasma, reflecting the time-averaged electron density profile. Going to high pressures (180 mTorr) the so-called  $\alpha$ - $\gamma'$  transition occurs, and the  $\text{SiH}$  emission intensity is highest at the center of the discharge. LIF experiments at these pressures are impossible due to the presence of powder which causes strong light scattering.

### V.B. Electrostatic probes

An electrostatic of Langmuir probe is a metallic insulated electrode, except at the tip, which is introduced into the plasma. The metal mostly used is tungsten, because of its high melting point. The insulator of the probe should be chemically inert. The probe is inserted into a discharge chamber via an electrically insulating vacuum seal, with which the probe is either fixed or movable, mostly in one direction, via a bellows. This *in situ* technique locally perturbs the plasma, by changing the electric field. The magnitude of the disturbance depends on the dimensions of the probe relative to the Debye length  $\lambda_D$ , which in turn depends on the plasma properties. Usually  $\lambda_D$  is in the  $\mu\text{m}$ -range, and probe diameters in the mm-range, so disturbances are very small.

Employing a Langmuir probe it is possible to determine plasma density ( $n_i$  and  $n_e$ ), electron temperature  $T_e$ , plasma potential  $V_{pl}$ , and floating potential

$V_{fl}$  [134,293,117,294]. It is often used as a complementary technique to optical and/or mass spectrometry. A Langmuir probe is operated by applying an external variable potential (usually between -100 and +100 V) and measuring the current (mA-range) through the probe. The sweep rate usually is 1-10 V/ $\mu$ s and multiple I-V characteristics are acquired and averaged. From such an averaged I-V characteristic all plasma parameters mentioned above can be in principle be determined.

When a probe is inserted into a plasma it will experience electrons and ions colliding with its tip. Due to the high mean speed of electrons the flow of electrons is higher than the flow of ions. Consequently, the tip will charge up negatively, until the electrons are repelled and the net current then is zero. The probe potential then is the floating potential,  $V_{fl}$ . The electron current density  $J_e$  then balances the ion current density  $J_i$ . At potentials lower than  $V_{fl}$  the ion current cannot increase further, in fact only ions are collected from the plasma, and the ion saturation current  $I_{si}$  is measured. The plasma potential  $V_{pl}$  is defined as the potential at which all electrons arriving near the probe are collected, and the probe current equals the electron current. Note that the plasma assumed the plasma potential in the absence of a probe, hence probe perturbation at  $V_{pl}$  is minimal. At potentials higher than  $V_{pl}$  the electron saturation current  $I_{se}$  is measured.

The I-V characteristic thus is divided into three regions: I)  $V < V_{fl}$  II)  $V_{fl} < V < V_{pl}$  and III)  $V > V_{pl}$ . In region I the ion saturation current dominates, in region III the electron saturation current. They are derived to be [117,294]:

$$I_{is} = 0.4n_i e A_p \sqrt{\frac{2k_B T_e}{m_i}} \quad (35)$$

$$I_{es} = n_e e A_p \sqrt{\frac{8k_B T_e}{m_e}} \sqrt{1 + \frac{eV}{k_B T_e}} \quad (36)$$

with  $A_p$  the probe area. The expression for  $I_{es}$  is derived for a cylindrical probe tip.

In the intermediate region II the electron retarding current is given by, assuming a Maxwellian EEDF:

$$I_{er} = n_e e A_p \sqrt{\frac{k_B T_e}{2\pi m_e}} \exp\left(\frac{eV}{k_B T_e}\right) \quad (37)$$

The electron temperature is derived from the I-V characteristic by first subtracting the measured ion current from the measured current in region II. From the slope of the logarithm of  $I_{er}$  versus  $V T_e$  is determined (Eq. (37)). Plotting the square of the probe current versus voltage in region III yields a straight line (Eq. (36)), and the electron density is derived from its slope (and the known  $T_e$ ). The ion density then is easily calculated with Eq. (35). The plasma potential is determined from the intersection between the extrapolated currents from region II and III. The floating potential is the potential at which the current is zero. Other methods for the extraction of plasma properties from I-V data, not assuming a Maxwellian EEDF, are described by Awakowicz [294].

In RF discharges one has to compensate for the RF voltage component across the probe sheath. This is done with a pick-up element, situated close to the probe tip, which should be in the plasma bulk as well. Compensating electronics within the probe are usually set to work at a prescribed RF-frequency, and cannot be used at other frequencies.

In the ASTER system a data series is measured for an argon and a hydrogen plasma running at 13.56 MHz, in which the power (5-30 W) and pressure (5-50 Pa) are varied [265,295]. The probe-tip is positioned exactly between powered and grounded electrode, at the center of the discharge.

The plasma potential determined from I-V data for argon increases linearly from about 24 to 27 V with increasing power at a pressure of 0.05 mbar. At the highest pressure of 0.35 mbar these values have shifted downwards by about 2 V only. For the hydrogen discharge a similar behaviour is observed, with an increase from 25 to 32 V with power, at 0.10 mbar. These values shift by about 3 V, going to a pressure of 0.5 mbar. The variation of the plasma potential in the case of hydrogen is larger than in the case of argon. These difference in behaviour is also seen in the electron temperature and the electron and ion density. In the case of argon,  $T_e$  varies between 1.8 and 2.8 eV. The effect of pressure variation is larger than the effect of power variation. At low pressure (0.05 mbar)  $T_e$  is around 2.5 eV, whereas at high pressure (0.35 mbar)  $T_e$  is around 2 eV. In the case of hydrogen,  $T_e$  varies between 1.5 and 4.2 eV. At low pressure (0.10 mbar)  $T_e$  varies slightly around 4 eV, whereas at high pressure (0.5 mbar)  $T_e$  changes from 4 eV at low power (5 W) to 1.5 eV at high power (30 W).

The electron and ion densities for argon and hydrogen are shown in Fig. 29 and Fig. 30, respectively. The absolute values of  $n_i$  for argon (Fig. 29b) are about 5-8 times larger than the ones for hydrogen (Fig. 30b). Further, the increase of  $n_i$  as a function of applied power is only slightly pressure-dependent in the case of argon. This is much clearer in the case of hydrogen. Similar observations can be made for the electron density.

Employing a extendable bellows the probe-tip can be moved towards the edge of the discharge, i.e. over a distance of about 5 cm. Measurements performed in an argon discharge at 0.15 mbar at three different power levels show that  $V_{pl}$ ,  $T_e$ ,  $n_e$ , and  $n_i$  do not vary within 5-10 %. This shows that the discharge is homogeneous.

The Langmuir probe method is easily employed in discharges, however especially in deposition plasmas care has to be taken in interpreting I-V data. The presence of negative ions in silane plasmas complicates the analysis. Their contribution should be taken into account. Further, (thin) insulating layers may form on the probe which can result in short circuiting of the probe. This contamination effects distort the probe characteristics, especially around  $V_{pl}$ . The probe surface may be cleaned *in situ* by ion bombardment, i.e., by biasing the probe to a large negative potential in a *non*-depositing plasma such as argon. Melting of the deposited contaminations off of the probe should be possible by passing a high electron current through the probe.

Probe measurements in silane discharges have been reported [296,297]. Apparently, no difficulties were experienced, as the deposited amorphous silicon layer on the tip were sufficiently photoconductive. For typical silane discharge conditions values for  $T_e$  are found to be between 2 and 2.5 eV. Electron densities are around  $1 \times 10^9 \text{ cm}^{-3}$  [296]. Probe measurement in the ASTER system failed due to strong distortions of the probe current, even after following cleaning procedures.

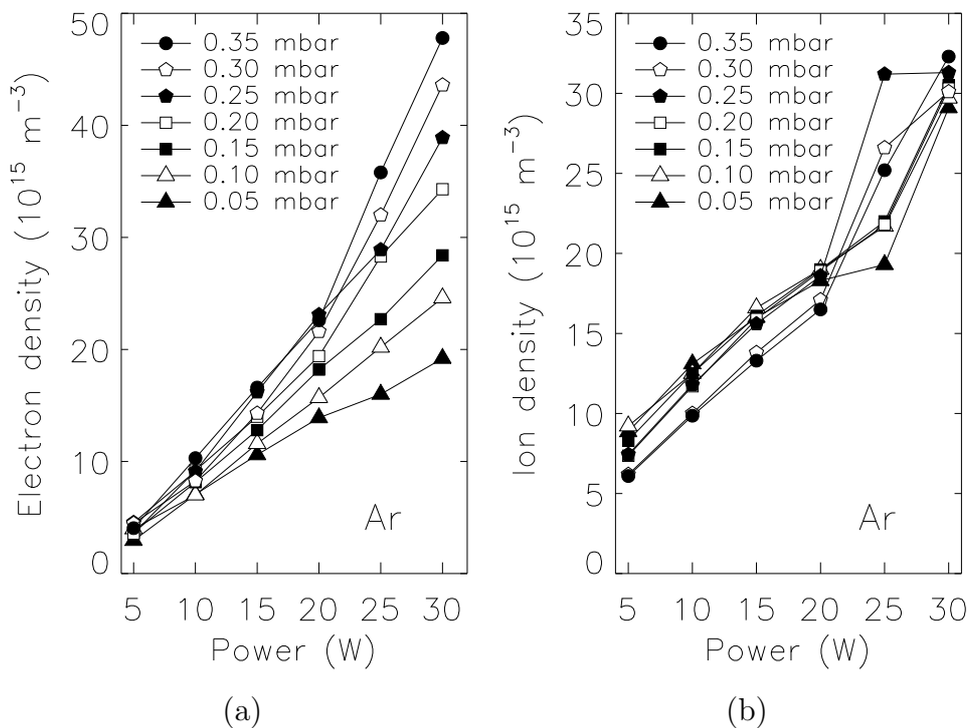


FIG. 29. Electron (a) and ion (b) density as a function of power for different pressures in case of an argon discharge.

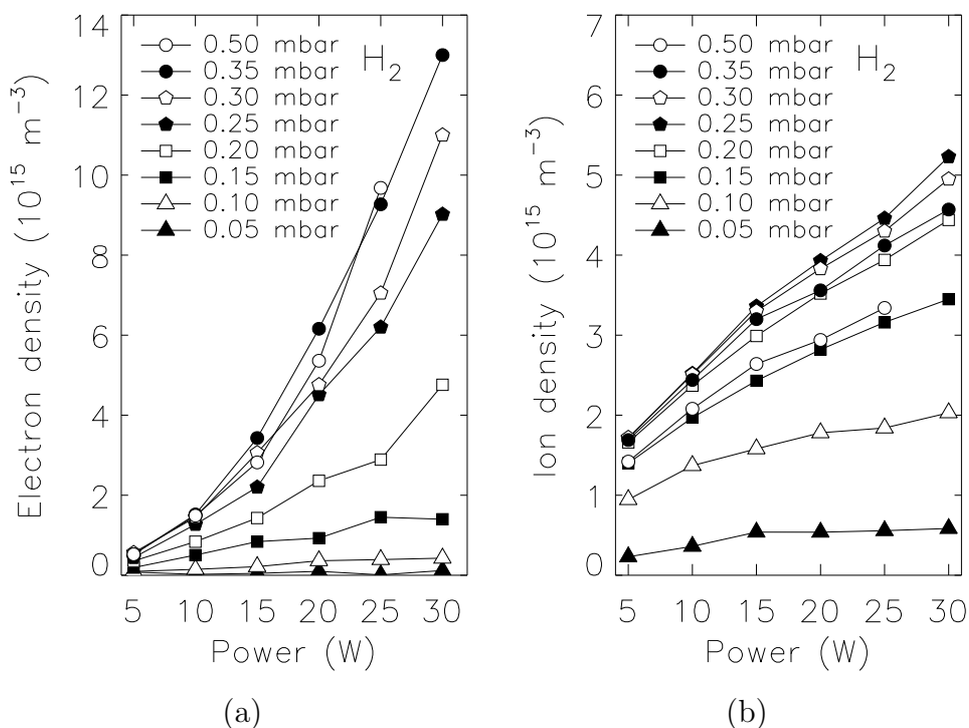


FIG. 30. Electron (a) and ion (b) density as a function of power for different pressures in case of an hydrogen discharge.

### V.C. Mass spectrometry

Mass spectrometry is used to obtain information on the neutral and ionic composition of the deposition discharge. Three modes of sampling are commonly used, i.e., analysis of discharge products downstream, line-of-sight sampling of neutrals, and direct sampling of ions from the discharge. Mass spectrometry is a simple method to measure and quantify neutral species in a silane discharge, although calibrations are required. For ionic species, mass spectrometry is the only method available. A quadrupole mass filter is a proven instrument essential in mass spectrometry of low pressure discharges.

#### V.C.1. Analysis of neutrals

The analysis of the neutral gas composition in a discharge yields useful information on the mechanisms and kinetics of silane dissociation. However, it should be borne in mind that with mass-spectrometric analysis one only detects the final products of a possible long chain of reactions.

The partial pressures of the stable neutral molecules in the discharge (silane, hydrogen, disilane, trisilane) can be measured by a quadrupole mass spectrometer (QMS). The QMS usually is mounted in a differentially pumped chamber, which is connected to the reactor via a small extraction port [286]. In the ASTER system a QMS is mounted on the reactor which is used for intrinsic material deposition. The QMS background pressure (after proper bake-out) is between  $10^{-12}$  and  $10^{-13}$  mbar. The controllable diameter in the extraction port is adjusted such that during discharge operation the background pressure never exceeds  $10^{-11}$  mbar.

The gas that enters the QMS is ionized by electron impact at a factory-preset electron energy of 70 eV (or 90 eV), and subsequently mass-analyzed. The ion currents at different mass over charge ( $m/e$ ) ratio need to be converted to partial pressures by careful calibrations, as reported by Hamers [163]. Gas X is admitted to the reactor and the corresponding pressure  $p$  is measured, as well as the ion current  $I$  at a specific  $m/e = \mu_X$  ratio of the gas, e.g., for argon  $m/e = 40$  a.m.u. The background signal at the same  $m/e = \mu_X$ , that results from residual gases in the QMS chamber is subtracted from the measured ion current. The calibration factor  $\gamma_X$  is defined as the ratio between the corrected ion current and the reactor pressure. Calibrations are performed typically at three pressures before and after measurements on the discharge, as the sensitivity of the channeltron in the QMS may rapidly change. The calibrations are performed with hydrogen, argon, silane, and disilane, with the main contributions at  $m/e = 2, 40, 30,$  and  $60$  a.m.u., respectively.

In the deposition of a-Si:H the dissociation of the process gas silane leads to the formation of hydrogen, disilane, trisilane, higher order silanes, and a solid film. Silane is depleted. From the consumption of silane, one may estimate the deposition rate of the solid film. The difference in the silane partial pressure between the 'discharge-on' and the 'discharge-off' state is often used as an indication of the amount of silane that is depleted [298–300]. However, the relative change in pressure in general is not equal to the relative change in flow, due to gas-dependent conductances [301,163] (see also Sec. II D). E.g., in the ASTER deposition system equal partial pressures of silane and hydrogen are observed for a flow ratio of silane to hydrogen of about 1.25. Hence, another way of

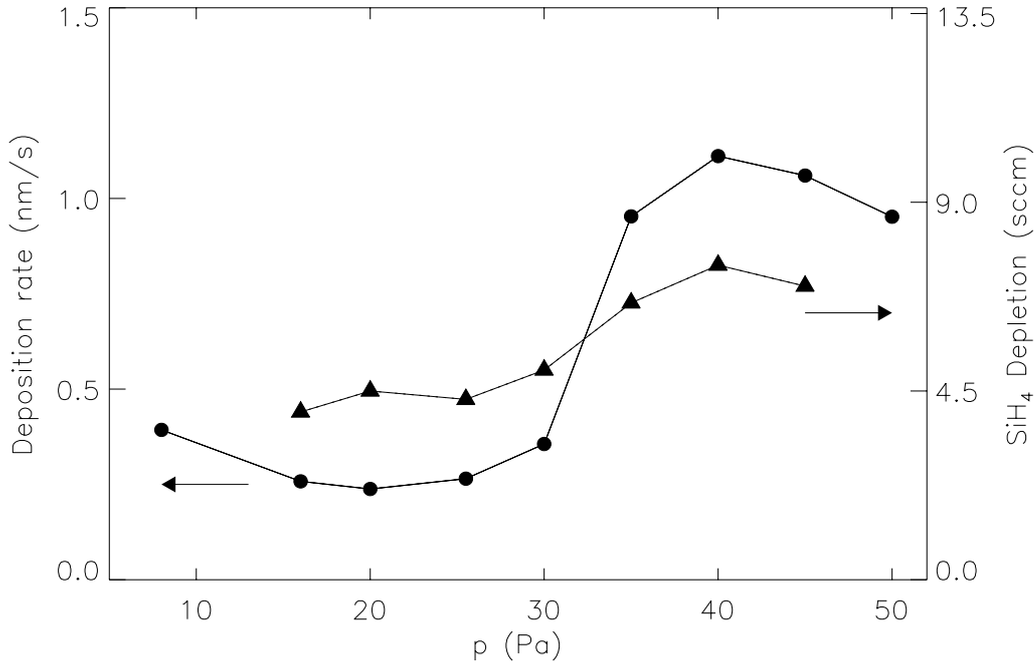


FIG. 31. The deposition rate and the corresponding silane depletion as a function of the total process pressure. Other conditions are 50 MHz, 30:30 SiH<sub>4</sub>:H<sub>2</sub>. (From E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

determination of silane depletion is used [301,163]. First, the partial pressures of silane and hydrogen are measured in the 'discharge-on' state. Subsequently, the discharge is switched off, and the silane and hydrogen flow rates are adjusted such that the same partial pressures as in the 'discharge-on' state are reached. These flow rates then are the flow rates of silane and hydrogen that leave the reactor in the 'discharge-on' state. The silane depletion  $\Delta Q_{\text{SiH}_4}$  is the difference between the admitted silane flow rate and the flow rate of silane, that leaves the reactor, when in the 'discharge-on' state. Additionally, an estimate can be made of the deposition rate, under the assumptions that all depleted silane is used for the deposition (no large amount of higher order silanes nor dust is formed). Also the deposition is assumed to occur homogeneous over the surface of the reactor. The estimated deposition rate is  $r_d^{est} = \Delta Q_{\text{SiH}_4} / k_B T_0 \rho A_r$ , with  $\rho$  the atomic density of the amorphous silicon network ( $5 \times 10^{22} \text{ cm}^{-3}$ ),  $T_0$  the standard temperature (300 K), and  $A_r$  the reactor surface ( $0.08 \text{ m}^2$ ). This yields an estimated deposition rate of 0.11 nm/s per sccm SiH<sub>4</sub> depletion [163]. In Fig. 31 the deposition rate is compared to the corresponding silane depletion as a function of process pressure, for a silane/hydrogen discharge at 50 Mhz and 10 W. For these conditions the  $\alpha$ - $\gamma'$  transition occurs at 30 Pa: the deposition rate is increased by a factor of 4, whereas the depletion is increased only by a factor of 2. The right-hand axis is scaled using the value of 0.11 nm/s/sccm. In the  $\alpha$  regime the depositon rate is underestimated, whereas in the  $\gamma'$  regime it is overestimated. This is in part due to the assumed uniformity of deposition. In the  $\alpha$  regime the deposition rate at the center of the substrate is lower

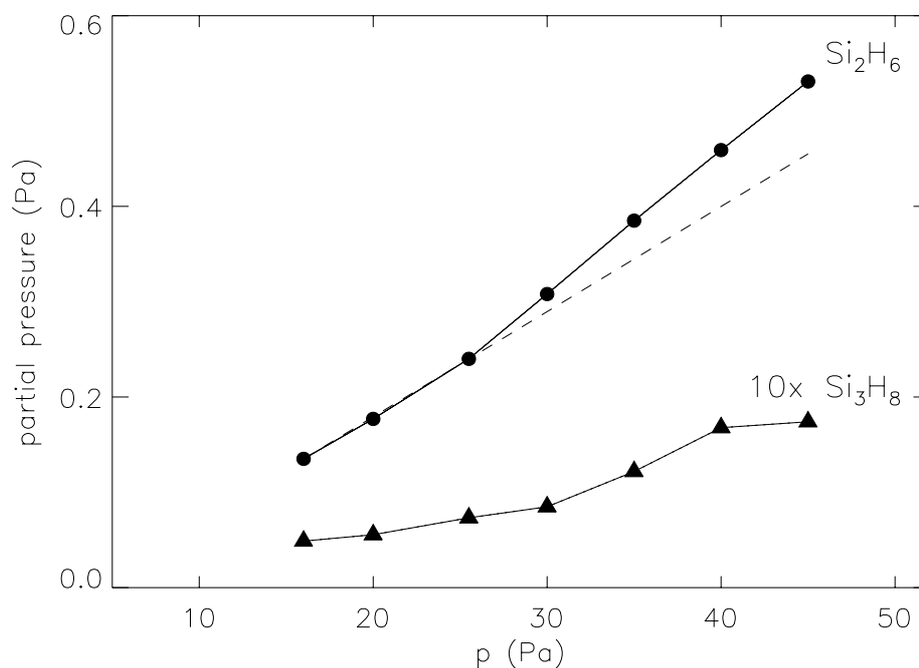


FIG. 32. The partial pressures of disilane and trisilane. The dashed line is an extrapolation of the disilane partial pressure in the  $\alpha$ -regime. (From E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

than at the edges, in the  $\gamma'$  regime this is reversed. Moreover, from the low DC bias voltages observed in the  $\gamma'$  regime it is inferred that the discharge is more confined.

The partial pressures of disilane and trisilane are shown in Fig. 32, for the same process conditions as in Fig. 31. Both partial pressures increase as a function of pressure. Around the  $\alpha$ - $\gamma'$  transition at 30 Pa the disilane partial pressure increases faster with increasing pressure, as can be seen from the deviation from the extrapolated dashed line. The disilane partial pressure amounts to about 1 % of the total pressure, and the trisilane partial pressure is more than an order of magnitude lower. Apparently, in the  $\gamma'$  regime the production of di- and trisilane is enhanced.

Not only the silane depletion, but also the hydrogen production can be used to obtain information on the reaction products of the decomposition process. In Fig. 33 the silane depletion and the corresponding hydrogen production is shown for a number of experiments, with process parameters such to cover both the  $\alpha$ - and the  $\gamma'$ -regime [163,301]. A clear correlation exists. In addition, the dotted line in Fig. 33 relates the hydrogen production and silane depletion in the case that only  $a$ -Si:H is formed with 10 at.% H, i.e.,  $a$ -Si:H<sub>0.1</sub> and H<sub>2</sub>. The dashed line represents the case where 30 % of the silicon from the consumed amount of silane would leave the reactor as disilane, instead of being deposited as  $a$ -Si:H<sub>0.1</sub>. The data can well be explained by these two cases. In fact, it is an indication that part of the consumed silane leaves the reactor with a higher amount of H per Si-atom than  $a$ -Si:H, which may be caused by the formation of higher silanes and/or powders.

Mass-spectrometric research on silane decomposition kinetics has been performed

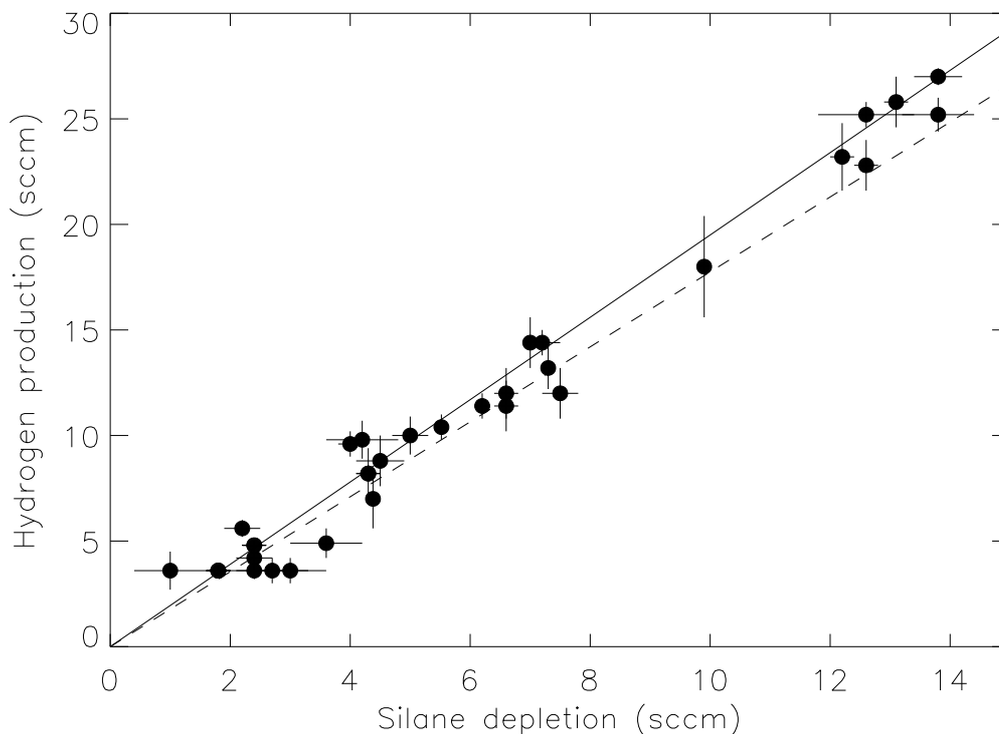


FIG. 33. The depletion of silane and the corresponding production of hydrogen for several process conditions, covering both the  $\alpha$  and the  $\gamma'$  regime. The dotted line represents the case where all the consumed silane is converted into  $\alpha$ -Si:H<sub>0.1</sub> and 1.95 H<sub>2</sub>. The dashed line represents the case where 30 % of the consumed silane is converted into disilane instead of being deposited. (From E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

for flowing [298,302–306] and static discharges [197,307]. In a DC discharge of silane it is found that the reaction rate for the depletion of silane is a linear function of the DC current in the discharge, which allows to determine a first-order reaction mechanism in electron density and temperature [302,304]. For an RF discharge, similar results are found [303,305]. Also, the depletion and production rates were found to be temperature-dependent [306]. Further, the depletion of silane and the production of disilane and trisilane are found to depend on dwell time in the reactor [298]. The increase of di- and trisilane concentration at short dwell times ( $< 0.5$  s) corresponds to the decrease of silane concentration. At long dwell times, the decomposition of di- and trisilane produces a much stronger decrease in silane concentration. Moreover, a strong correlation is observed between deposition rate and di- and trisilane concentration. This has led to a proposal [283,298,308] of the main electron impact dissociation channel of silane, i.e.,  $e + \text{SiH}_4 \rightarrow \text{SiH}_2 + \text{H}_2 + e$ . The higher silanes are partially decomposed into radicals, that are responsible for deposition [308].

In a static discharge first silane is introduced and subsequently all valves are closed. Then the discharge is ignited, and the mass-spectrometer signals for silane ( $m/e=30$  a.m.u.), disilane ( $m/e = 62$  a.m.u.), trisilane ( $m/e = 92$  a.m.u.), and tetrasilane ( $m/e = 120 - 128$  a.m.u.) are followed as a function of time [197]. Rate equations can be

formulated that explain the decrease in silane concentration and the increase in di-, tri-, and tetrasilane. From these data another proposal [197] of the main electron impact dissociation channel of silane is formulated:  $e + \text{SiH}_4 \rightarrow \text{SiH}_2 + 2 \text{H} + e$ , in agreement with others [201,309,310]. The abstraction reaction  $\text{H} + \text{SiH}_4 \rightarrow \text{SiH}_3 + \text{SiH}_3$  explains the large production of  $\text{SiH}_3$  radicals in this case. The discrepancy as is reflected in the hydrogen production (either H or  $\text{H}_2$ ) cannot be resolved solely by mass-spectrometric data.

### V.C.2. Analysis of radicals

It is possible but difficult to directly detect neutral radicals produced in a silane discharge by mass spectrometry. To discriminate between the radicals and the much more abundant neutrals, ionization of the species by low-energy electrons can be used [307], and the term threshold ionization mass spectrometry (TIMS) is used to denote this technique. This method is based on the principle of ionization of neutrals by electrons that have a well defined energy. These ionized neutrals then are detected by means of a quadrupole mass spectrometer. In general, the radical A can be detected when the electron impact ionization process leads to the formation of an ion  $\text{A}^+$ . In order for this to occur, the energy of the electrons should be larger than the ionization potential  $E_i$ . Also, dissociative ionization of a molecule containing A may lead to  $\text{A}^+$ , following  $\text{AB} + e^- \rightarrow \text{A}^+ + \text{B} + 2e^-$ . Here, the energy of the electrons should be larger than the appearance potential  $E_a$ . It holds that  $E_a > E_i$ , and  $E_a \approx E_i + E_{\text{A-B}}$ , with  $E_{\text{A-B}}$  is the energy of the A-B bond [311]. When the electron energy is larger than  $E_a$  the dissociative ionization process dominates, because the density of the molecule AB is much larger than the radical A. When the electron energy has a value between  $E_i$  and  $E_a$ , then only the radicals are ionized, and one is able to determine the radical density.

In silane discharges, one observes the following: when the discharge is *off*, the mass spectrometric signal at  $m/e = 31$  a.m.u. ( $\text{SiH}_3^+$ ) as a function of electron energy is due to dissociative ionization of  $\text{SiH}_4$  in the ionizer of the QMS, with an ionization potential of 12.2 eV [312]. The signal with the discharge *on* is due to ionization of the radical  $\text{SiH}_3$  plus the contribution from dissociative ionization of silane (the ions from the discharged are repelled by applying a positive voltage on the extraction optics). The appearance potential of the  $\text{SiH}_3$  radical is 8.4 eV [312], and therefore a clear difference between discharge *on* and *off* is observed. The corresponding threshold energies for  $\text{SiH}_2$  are 11.9 and 9.7 eV. (these four numbers are also given by Kae-Nune *et al.* [311], i.e., 12.0, 8.0, 11.5, and 9.0 eV, respectively). The net radical contribution to the ion signal is given by [311]:

$$\Delta I(\text{A}^+) = (I_{on} - I_{bg} - I_{ions}) - (I_{off} - I_{bg}) \frac{n_{\text{SiH}_4,on}}{n_{\text{SiH}_4,off}} \quad (38)$$

where  $I_{on}$  ( $I_{off}$ ) is the signal when the discharge is on (off),  $I_{bg}$  the background signal, and  $I_{ions}$  the signal due to ions (which should be zero). The term  $n_{\text{SiH}_4,on}/n_{\text{SiH}_4,off}$  accounts for the depletion of the density of  $\text{SiH}_4$ , which can be obtained from the ratio  $I_{on}/I_{off}$  at high electron energies. The concentration of radical  $\text{SiH}_x$  can be expressed as [311]:

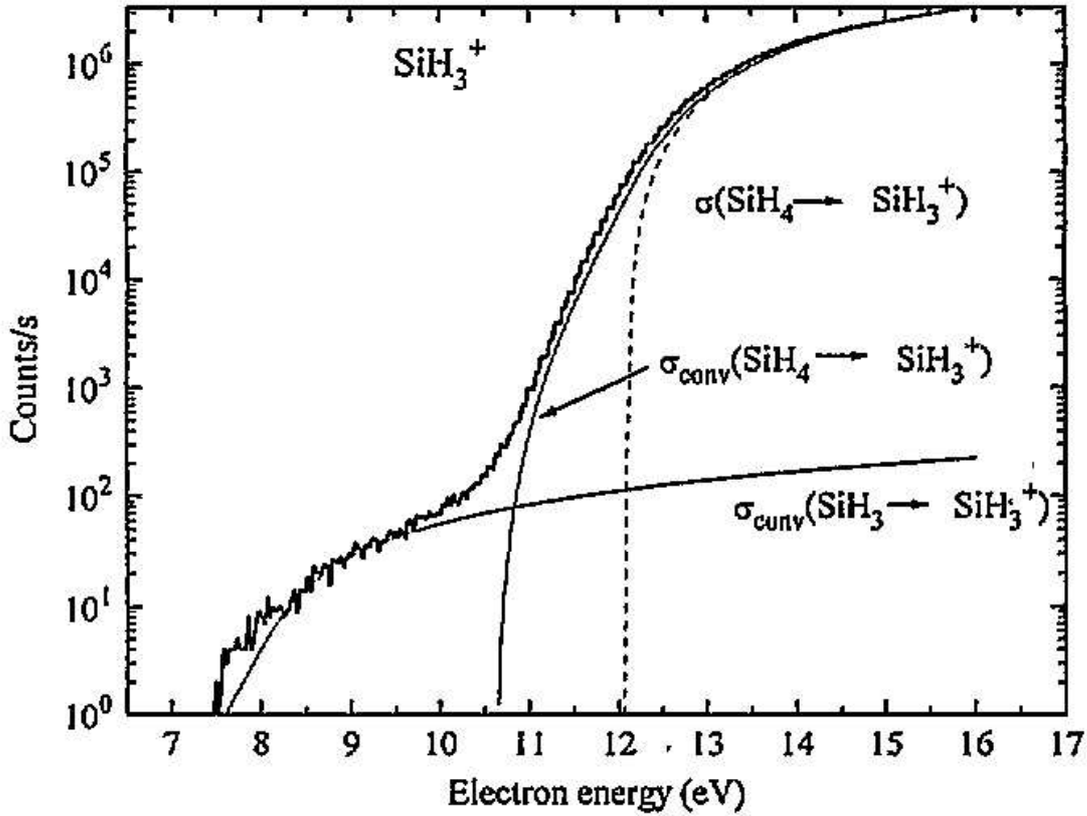


FIG. 34. Measured and fitted curves for the ionization of the  $\text{SiH}_3$  radical. (From P. Kae-Nune, J. Perrin, J. Guillon, and J. Jolly, *Plasma Sources Sci. Technol.* **4**, 250 (1995).)

$$\frac{n(\text{SiH}_x)}{n(\text{SiH}_4)} = C \frac{Q(\text{SiH}_4 \rightarrow \text{SiH}_x^+, E_2 > E_a)}{I(\text{SiH}_x^+, E_2 > E_a)} \frac{1}{E_1'' - E_1'} \int_{E_i - E_1'}^{E_1'' < E_a} \frac{\Delta I(\text{SiH}_x^+, E)}{Q(\text{SiH}_x \rightarrow \text{SiH}_x^+, E)} dE \quad (39)$$

where  $I(\text{SiH}_x^+, E_2 > E_a)$  is the signal due to dissociative ionization of  $\text{SiH}_4$ ,  $\Delta I(\text{SiH}_x^+, E)$  the net radical signal (Eq. (38)),  $Q(\text{SiH}_4 \rightarrow \text{SiH}_x^+, E_2 > E_a)$  the electron impact cross section of the dissociative ionization of  $\text{SiH}_4$ ,  $Q(\text{SiH}_x \rightarrow \text{SiH}_x^+, E)$  the electron impact cross section of the ionization of the radical  $\text{SiH}_x$ , and  $C$  a geometrical correction factor. Clearly, the knowledge of the electron impact cross sections is essential for the exact determination of the radical densities. Kae-Nune *et al.* [311] have used data for deuterated methane radicals ( $\text{CD}_3$  and  $\text{CD}_2$  [313]) to determine by scaling the  $Q(\text{SiH}_3 \rightarrow \text{SiH}_3^+, E)$  and  $Q(\text{SiH}_2 \rightarrow \text{SiH}_2^+, E)$ .  $Q(\text{SiH} \rightarrow \text{SiH}^+, E)$  is determined by taking the average of  $Q(\text{SiH}_2 \rightarrow \text{SiH}_2^+, E)$  and the measured  $Q(\text{Si} \rightarrow \text{Si}^+, E)$  [314]. In Fig. 34 the measured  $\text{SiH}_3^+$  signal is compared to fitted curves of radical and dissociative ionization [311].

Nowadays, a commercially available quadrupole mass spectrometer system (Hiden EQP300) is used by many groups, with which detection of radicals, but also of ions (see Sec. VC3), is possible [315]. Kae-Nune *et al.* [198,217,311] have mounted this QMS system in the grounded electrode of their parallel plate PECVD reactor. Neutral

(and ionic) species are sampled from the discharge via a 0.3 mm diameter hole in the grounded electrode. Two-stage differential pumping ensures a low background pressure ( $< 10^{-6}$  Torr) in the ionization chamber of the QMS.

With this set-up a series of measurements was taken in which the power was varied from 5 to 30 W. The pressure was low, 0.06 Torr, while the temperature was 250°. The  $\text{SiH}_x$  ( $0 < x < 3$ ) radical densities near the surface were measured, as a function of effective power delivered to the discharge. It was found that the  $\text{SiH}_3$  radical is the most abundant one, i.e, with a concentration varying between 1.8 and 3.2  $10^{11}$   $\text{cm}^{-3}$ . The  $\text{SiH}_2$  radical concentration was about a factor of 20 lower. Both radical concentrations increase with increasing power.

From the radical concentrations it is possible to derive the radical flux to the surface:

$$\Phi_{\text{SiH}_x} = n_{\text{SiH}_x} \frac{v_{\text{SiH}_x}}{4} \frac{\beta_{\text{SiH}_x}}{1 - \frac{1}{2}\beta_{\text{SiH}_x}} \quad (40)$$

where  $v_{\text{SiH}_x}$  is the thermal velocity ( $= \sqrt{8k_B T / \pi m}$  of the radical  $\text{SiH}_x$ ,  $\beta_{\text{SiH}_x}$  the surface reaction probability (see also Fig. 14 in Sec. IV A 6). The term  $\beta$  represents the adsorbed fraction of radicals, which either stick (sticking probability  $s_{\text{SiH}_x}$ ) or recombine as  $\text{SiH}_4$  or  $\text{Si}_2\text{H}_6$  to the gas phase (recombination probability  $\gamma_{\text{SiH}_x}$ ). By definition,  $\beta = s + \gamma$ . An amount  $r = 1 - \beta$  is reflected from the surface directly.

The contribution of radical  $\text{SiH}_x$  to the total deposition rate follows from:

$$R_{d,\text{SiH}_x} = \Phi_{\text{SiH}_x} \frac{s_{\text{SiH}_x}}{\beta_{\text{SiH}_x}} \frac{M_{\text{Si}}}{N_A \rho_{a-\text{Si:H}}} \quad (41)$$

where  $N_A$  is Avogadro's number,  $M_{\text{Si}}$  the molar mass of silicon, and  $\rho_{a-\text{Si:H}}$  the mass density of  $a$ -Si:H (2.2  $\text{g}/\text{cm}^3$ ). For  $\text{SiH}_3$  Matsuda *et al.* have determined  $\beta$  and  $s$  [137]:  $\beta_{\text{SiH}_3} = 0.26$  and  $s_{\text{SiH}_3} = 0.09$ . For  $\text{SiH}_2$ ,  $\text{SiH}$ , and  $\text{Si}$  large  $\beta$  and  $s \approx \beta$  is assumed, i.e.,  $\beta_{\text{SiH}_2} = s_{\text{SiH}_2} = 0.8$  [311],  $\beta_{\text{SiH}} = s_{\text{SiH}} = 0.95$  [316], and  $\beta_{\text{Si}} = s_{\text{Si}} = 1$  [311]. Using these values for  $\beta$  and  $s$ , Kae-Nune *et al.* [311] have determined the contribution of radicals to the deposition rate. At low power levels, the deposition rate can fully be accounted for by the  $\text{SiH}_3$  and  $\text{SiH}_2$  radicals in a 60:40 ratio. At high power levels also the other radicals become important. At their highest power level of about 50  $\text{mW}/\text{cm}^2$ , the sum of all radical contributions to the deposition rate is only 65%, indicating that dimer and trimer radicals, as well as ions, also contribute to the deposition.

Using the same threshold ionization mass spectrometry set-up, Perrin *et al.* [317] have measured the temporal decay of radical densities in a discharge afterglow. From these experiments the coefficient  $\beta$  for the radical  $\text{SiH}_3$  has been determined to be 0.28, which is in agreement with already known results from other (indirect) experimental approaches [136,137,318]. For the  $\text{Si}_2\text{H}_5$  radical  $\beta$  is determined to be between 0.1 and 0.3. The coefficient  $\beta$  for atomic hydrogen on  $a$ -Si:H lies between 0.4 and 1, and is thought to mainly consist of surface recombination as  $\text{H}_2$ .

### V.C.3. Analysis of ions

Detection of ions from a discharge is done by direct sampling through an orifice. In order to extract the ions collisionless the dimensions of the sampling orifice should be smaller than the sheath thickness, and are typically of the order of 100  $\mu\text{m}$ . Moreover,

the detected ions and their energies are only representative of the plasma bulk situation, when the sheath is collisionless, i.e., at low pressures [286]. One generally is interested in the interaction of ions with a growing surface, hence normal operating pressures are used.

In RF discharges of silane  $\text{SiH}_3^+$  usually is the most abundant ion, but others ( $\text{SiH}_2^+$ , and  $\text{Si}_n\text{H}_m^+$ , with  $1 < n < 9$ ) also are present [319]. The relative abundance of  $\text{SiH}_3^+$  ions increases with increasing pressure, while that of  $\text{SiH}_2^+$  decreases [319]. The ionization cross section of  $\text{SiH}_2^+$  is higher than that of  $\text{SiH}_3^+$  [320], but  $\text{SiH}_2^+$  is lost via the reaction  $\text{SiH}_2^+ + \text{SiH}_4 \rightarrow \text{SiH}_3^+ + \text{SiH}_3$  [305]. At pressures lower than 0.1 Torr  $\text{SiH}_2^+$  becomes the dominant ion.

The ion clusters ( $\text{Si}_2\text{H}_m^+$ ,  $\text{Si}_3\text{H}_n^+$ , ...) are also present and larger clusters (positive and neutral) can be formed through reactions with silane molecules. Negative ions have been detected [321], which are responsible for the powders in the discharge [322].

Ion energy distributions (IEDs) are measured by several groups [323–326]. The reliability of IEDs depends strongly on the knowledge of the transmission function of the instruments, which most likely is energy-dependent. Improper adjustment of the various potential levels throughout the instruments can result in 'ghost' structures in the IED, and they reflect the physics of the instrument rather than of the discharge. Hamers *et al.* [161,163] have presented a method to determine the proper transmission function of an electrostatic lens system of a commercially available ion energy and mass spectrometer, the Hiden EQP [315]. This system is used by many groups to study plasmas [160,198,321,327,328]. It is very versatile with respect to the control of voltages on the many electrostatic lenses.

The EQP is mounted in a plasma reactor identical to the ones in ASTER, see Fig. 35. The plasma is generated between the two parallel electrodes. In the grounded electrode, where normally the substrate is placed for the deposition of *a*-Si:H, a small orifice is located to sample particles that arrive from the discharge. The orifice is made in a 20  $\mu\text{m}$  thick stainless steel foil, and has a diameter of 30  $\mu\text{m}$ . The foil is integrated in a stainless steel flange. The ratio of the thickness of the foil and the diameter of the orifice results in a physical acceptance angle of  $55^\circ$  with respect to the normal of the electrode. The small size of the orifice compared to the sheath thickness and the mean free path of the particles in the plasma ensures that the orifice does not influence the discharge [323]. Using this orifice in deposition plasmas leads to deposition of *a*-Si:H on the surface of the orifice, but also on its inner sides, thereby narrowing the sampling diameter. This can be observed by monitoring the pressure rise in the mass spectrometer during experiments. Typically, a 50% reduction in orifice area takes 3 hours. The *a*-Si:H film on the orifice then needs to be etched away, which is done *ex situ* by a KOH etch.

The design of the instrument, together with the pumping capacity ensures a low background pressure ( $< 10^{-9}$  mbar). Under process conditions the pressure directly behind the orifice is about a factor of  $10^5$  lower than the process pressure, in the mass filter even a factor of  $10^6$ . The mean free path of particles that have entered the EQP therefore is several meters.

The ion optics (IO in Fig. 35) consists of a number of electrostatic lenses, that direct the sampled ions through the drift tube (DT) to the electrostatic analyzer (ESA). The ESA transmits only those ions with a specific energy over charge ratio  $\epsilon_{pass}$ . An ESA has a constant relative energy resolution  $\Delta\epsilon/\epsilon_{pass}$ . Because of the fact that this ESA

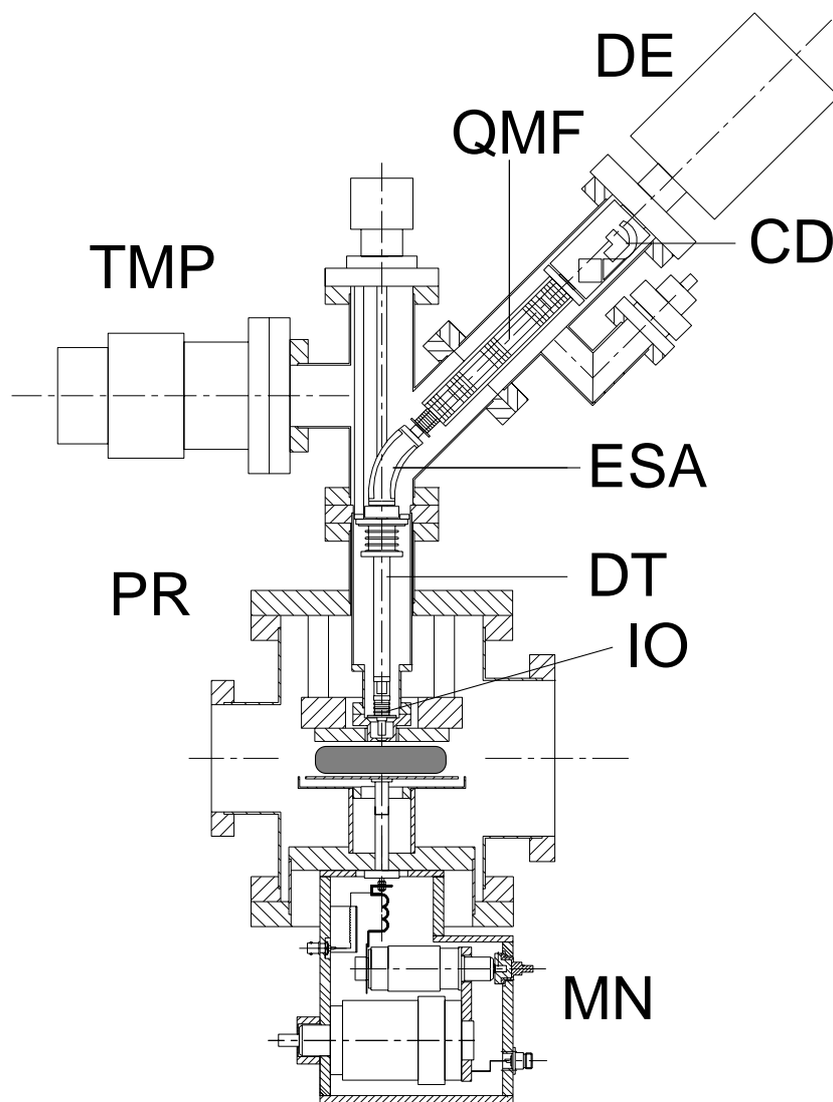


FIG. 35. Vertical cross-section of the reaction chamber equipped with the mass spectrometer system. Indicated are: (QMF) the quadrupole mass filter, (ESA) the electrostatic analyzer, (CD) the channeltron detector, (DE) the detector electronics, (DT) the drift tube, (IO) the ion optics, (TMP) the turbo molecular pump, (PR) the plasma reactor, and (MN) the matching network.

operates at a constant  $\Delta\epsilon$ , the pass energy  $\epsilon_{pass}$  is constant. Here,  $\epsilon_{pass} = 40$  eV, and  $\Delta\epsilon = 1.5$  eV (FWHM). The measurement of IEDs requires that sampled ions are to be decelerated or accelerated to this pass energy. This is also done in the ion optics part. The other very important function of the ion optics is to shape the ion beam. Voltages set on the various lenses should be set such as to avoid chromatic aberration, which causes energy dependent transmission of ions in the instrument, and as a result

erroneous IEDs [161,163]. The correct lens settings haven been found by simulations of ion trajectories in the EQP using the simulation program SIMION [329]. In addition, an experimental method to find the correct settings was presented as well [161,163].

The acceptance angle depends on the lens settings and on the ion energy. Low energy ions are deflected in the instrument more towards the optical axis than high energy ions, which results in a larger acceptance angle. The acceptance angle varies between about  $6^\circ$  and  $1^\circ$  for ion energies between 1 and 100 eV [161,163]. For lower energies the acceptance angle can reach values above  $20^\circ$ , which causes drastic increases of the low energy part of the measured IED.

The measured ion energy distributions are affected by the processes which ions experience during their passage through the plasma sheath. In a collisionless situation, the IED is solely determined by the RF modulation of the plasma potential. The extent to which an ion will follow these modulations depends on its mass. Electrons are able to follow the electric field variations instantaneously, while ions experience a time-averaged electric field. The energy of an ion that arrives at the electrode depends on the value of the plasma potential at the time that the ion entered the sheath. Hence, the ion energy is modulated in much the same way that the plasma potential is modulated. Many ions arrive at the electrode with energies close to the extreme values of the harmonically varying voltage, and this results in a saddle-structure in the IED, in which the two peaks reflect the minimum and maximum plasma potential [134]. The heavier the ion, the more it experiences a time-averaged plasma potential, and the saddle-structure is narrower.

Elastic collisions and chemical reactions in the sheath lead to a broad angular distribution of the ion velocity at the electrode. Examples for  $\text{SiH}_3^+$  and  $\text{SiH}_2^+$  are the  $\text{H}^-$  abstraction reaction  $\text{SiH}_3^+ + \text{SiH}_4 \rightarrow \text{SiH}_4 + \text{SiH}_3^+$  [330] and  $\text{SiH}_2^+ + \text{SiH}_4 \rightarrow \text{SiH}_3^+ + \text{SiH}_3$  [331]. Due to the small acceptance angle of the EQP, these processes are only in part reflected in the IED [163,332].

#### *V.C.3.a. Charge exchange*

Much more important for the shape of the IED are charge exchange processes. In a charge exchange process an electron is exchanged between a neutral and an ion. In the plasma sheath a charge exchange process between a neutral of thermal energy and a fast ion lead to the formation of thermal ions and fast neutrals. These newly created thermal ions are accelerated towards the electrode, and the kinetic energy that they gain depends on the phase in the RF period and their position at the time of creation. This leads to distinct peaks in the IED, and the origin of these peaks can be explained as follows [325]. Electrons respond to the RF modulation and move to and from the electrodes. The movement of the electron front is a function of the phase  $\phi$  in the RF period. When an ion is created (by a charge exchange process) behind the electron front, i.e., in the plasma bulk, it will not experience an electric field. Once the electron front moves inwards to the plasma bulk, the ion will be accelerated to the electrode. Depending on the place of creation  $x$  the ion will be able to reach the electrode before the electron front is at the position of the ion again. Once in the plasma bulk, the ion will not be accelerated, until the electron front moves inwards again. This may be repeated several times. Wild and Koidl [325] have presented a description of the origin of the maxima in IEDs. They argue that these maxima only occur if the ion energy  $\epsilon$

at the electrode is independent of the phase of creation  $\phi$  and the position of creation  $x$  of the newly formed ion. Two conditions are to be met:  $d\epsilon/d\phi = 0$  and  $d\epsilon/dx = 0$ . As shown by Wild and Koidl [325] the energy  $\epsilon$  varies between  $\epsilon_{min}$  and  $\epsilon_{max}$ , and for every  $x$   $\epsilon_{min}$ ,  $\epsilon_{max}$ , and  $\epsilon_{\phi=0}$  fulfill the condition  $d\epsilon/d\phi = 0$ . Further,  $\epsilon_{\phi=0}$  exhibits a series of extrema as a function of  $x$  and is at these extrema alternately equal to  $\epsilon_{min}$  and  $\epsilon_{max}$ . At these extrema both conditions are met, and a saddle structure is observed in the IED.

In addition, ions that arrive with the mean energy of the saddle structure have reached the electrode after an integer number of periods. Thus, ions in successive charge exchange saddle structures have needed an subsequent number of periods to reach the electrode, on average. Hamers has very nicely illustrated this by performing Monte Carlo simulations of charge exchange processes [163], using a time- and position-dependent electric field, which is calculated from the sheath model as described by Snijkers [333]. About 300,000  $\text{SiH}_2^+$  ions were followed. The ion density at the electrode was  $3 \times 10^8 \text{ cm}^{-3}$ , the plasma potential 25 V, and the excitation frequency 13.56 MHz. The main saddle structure is found to be around 25 V (23–27 V). In the IED 4 charge exchange peaks are clearly distinguished, at about 2, 7.5, 14, and 19 eV. From a comparison with the transit time, i.e. the time an ion needs to cross the sheath, it is clear that the peaks in the IED correspond to ions that needed an integer amount of periods to cross the sheath. The first peak at 2 eV corresponds to ions that needed 1 period, and so on. In general, ions in the charge exchange peak number  $p$  needed  $p$  periods to cross the sheath.

The energy position of the peak  $\epsilon_p$  and the corresponding  $p$  periods of time  $T$  is used by Hamers *et al.* [163,332] to reconstruct the time-averaged potential profile  $V(x)$  in the sheath. An ion that arrives with energy  $\epsilon_p$  at the electrode has a velocity that follows from  $\frac{1}{2}mv^2 = \epsilon_p - eV(x)$ . With  $v = dx/dt$  one derives

$$\int_0^{x_p} \sqrt{\frac{m/2}{\epsilon_p - eV(x)}} dx = \int_0^{pT} dt = pT \quad (42)$$

where  $x_p$  is the mean position of creation of the ions that arrive at the electrode with energy  $\epsilon_p$ . The unknown  $V(x)$  now is found by further assuming that  $V(x)$  can be represented by a parabolic function [328,333]. A parabolic potential profile corresponds to a constant (net) charge carrier density  $n$  as a function of distance  $x$  in the sheath, according to  $-\vec{\nabla}^2 V = \vec{\nabla} \cdot \vec{E} = en/\epsilon_0$ . The electric field  $E(x)$  in the sheath then is a linear function of  $x$ :

$$E(x) = -E_0 + \frac{enx}{\epsilon_0} \quad (43)$$

with  $E_0$  the electric field at the electrode ( $x = 0$ ). Integrating  $E(x)$  from the electrode ( $V = 0$ ) over a distance  $x$  gives the potential:

$$V(x) = E_0x + \frac{en}{2\epsilon_0}x^2 \quad (44)$$

The sheath thickness  $d_s$  is found from the position where the electric field vanishes:  $d_s = \epsilon_0 E_0 / (en)$ . Here the potential is equal to the time-averaged plasma potential  $V_{pl}$ . We then find with Eq. (44) for the electric field at the electrode  $E_0$ :

$$E_0 = \sqrt{\frac{2enV_{pl}}{\epsilon_0}} \quad (45)$$

and for the sheath thickness  $d_s$ :

$$d_s = \sqrt{\frac{2\epsilon_0 V_{pl}}{en}} \quad (46)$$

Rewriting Eq. (44) yields

$$V(x) = V_{pl} \left[ \frac{2x}{d_s} - \left( \frac{x}{d_s} \right)^2 \right] \quad (47)$$

Substitution of  $V(x)$  in Eq. (42) and solving for  $\epsilon_p$  then yields

$$\epsilon_p = eV_{pl} \left( 1 - \left[ \cosh \left( pT \sqrt{\frac{e^2 n}{m\epsilon_0}} \right) \right]^{-2} \right) \quad (48)$$

The energy position  $\epsilon_p$  of peak number  $p$  in the IED of an ion with mass  $m$  is seen to be dependent on the plasma potential  $V_{pl}$ , the RF period  $T$ , and the ion plasma frequency ( $\omega_i = \sqrt{e^2 n / (m\epsilon_0)}$ ). Equation (48) can be used to determine the (net) charge carrier density in the sheath and the time-averaged potential  $V_{pl}$  from measured IEDs. The mean position  $x_p$  follows from combining Eq. (47) and Eq. (48):

$$x_p = d_s \left( 1 - \left[ \cosh \left( pT \sqrt{\frac{e^2 n}{m\epsilon_0}} \right) \right]^{-1} \right) \quad (49)$$

In silane discharges several ions are observed to be involved in a charge exchange process, and therefore maxima in their ion energy distribution at distinct energies are observed. The charge carrier density and the plasma potential that result from the fit of the IED allow for the quantification of the related parameters sheath thickness and ion flux. This method has been used to relate the material quality of a-Si:H to the ion bombardment [301,332], see also Sec. VI B 3.

In the following, IEDs measured in silane/argon and silane/hydrogen discharges are shown, and  $V_{pl}$  and  $n$  are determined from fitting the data using Eq. (48). In fact,  $V_{pl}$  is determined from an IED that is not affected by collisions in the sheath; in the IED at  $m/e = 60$  a.m.u., corresponding to  $\text{Si}_2\text{H}_4^+$ , is the only peak observed in that IED. It should be noted that in general the width of the saddle structure is smaller than the energy resolution (1.5 eV) of the instrument, and therefore cannot be distinguished.

### V.C.3.b. Silane/argon discharges

Silane/argon discharges are very illustrative for no less than four different ions have charge exchange maxima in their ion energy distribution. In Fig. 36 IEDs are shown that were measured in a plasma that was created in a mixture of 13 sccm Ar, 13 sccm  $\text{SiH}_4$ , at a pressure of 0.1 mbar. The applied RF frequency was 13.56 MHz, at a power of 10 W. The DC self bias voltage that developed was -135 V. The substrate

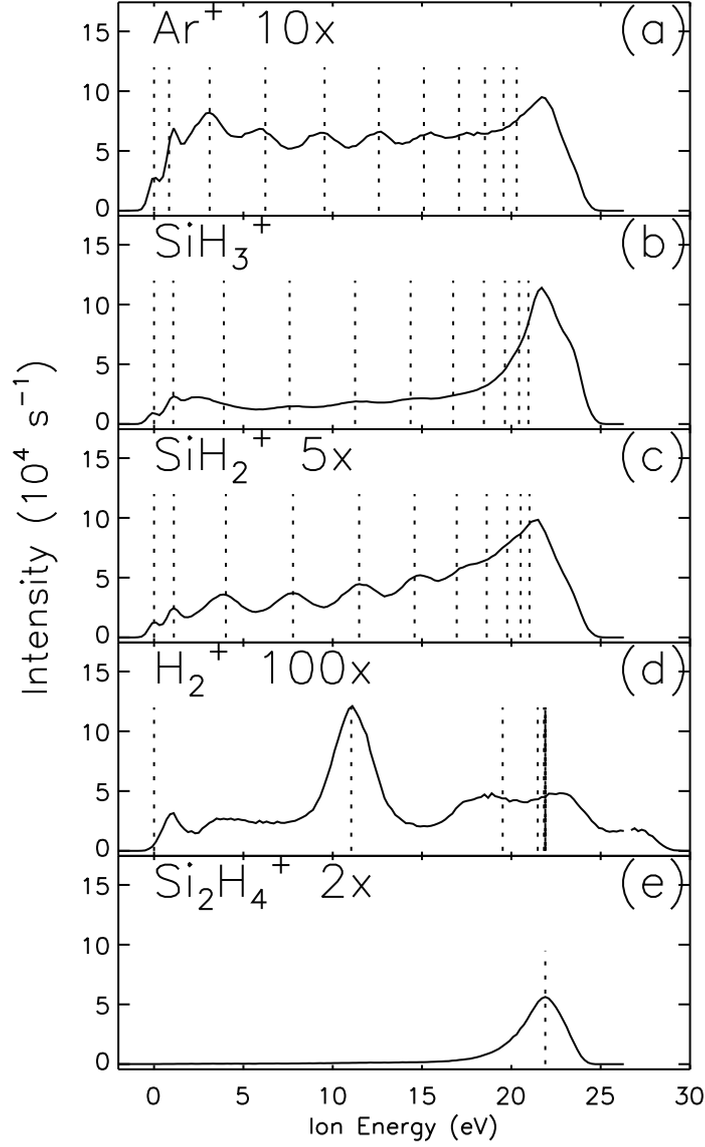


FIG. 36. The measured ion energy distributions of (a)  $\text{Ar}^+$ , (b)  $\text{SiH}_3^+$ , (c)  $\text{SiH}_2^+$ , (d)  $\text{H}_2^+$ , (e)  $\text{Si}_2\text{H}_4^+$  from a  $\text{SiH}_4/\text{Ar}$  plasma. The conditions are 13 sccm  $\text{SiH}_4$ , 13 sccm Ar, 13.56 MHz, 10 W, 0.2 mbar,  $250^\circ\text{C}$ ,  $V_{DC} = -135$  V. The dotted lines in (a), (b), (c), and (d) indicate the position of the first 10 charge exchange peaks based on a plasma potential of 21.8 V and a charge carrier density  $n$  of  $1.66 \times 10^8 \text{ cm}^{-3}$ . The dotted line in the IED of  $\text{Si}_2\text{H}_4^+$  (e), indicates the value of the plasma potential. Note the different scaling factors of the various IEDs. (From E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

temperature was  $250^\circ\text{C}$ . At least the first six peaks in the IED of  $\text{Ar}^+$  and the first five peaks in the IED of  $\text{SiH}_2^+$  can clearly be discerned. From the IED of  $\text{Si}_2\text{H}_4^+$  (Fig. 36e) a time-average plasma potential  $V_{pl}$  of  $(21.8 \pm 0.3)$  V is derived. Using now only *one* fitting parameter, i.e., the charge carrier density  $n$ , all distinct peaks in the IEDs of  $\text{Ar}^+$ ,  $\text{SiH}_3^+$ ,  $\text{SiH}_2^+$ , and  $\text{H}_2^+$  can be fitted with Eq. (48). Performing the fit yields  $n = (1.66 \pm 0.05) \times 10^8 \text{ cm}^{-3}$ . The energy positions are fitted well, as can be seen by comparing the data in Fig. 36 with the dotted vertical lines. The sheath thickness as calculated from Eq. (46) is  $(3.81 \pm 0.05)$  mm. The first charge exchange peak in the  $\text{Ar}^+$  IED originates at a distance of  $100 \mu\text{m}$  from the electrode, as calculated from Eq. (49).

The sixth peak, which is still visible, originates at a distance of about half the sheath thickness from the electrode.

It is observed that the  $\text{SiH}_3^+$  ion is the most abundant ion, see Fig. 36b. The peak around 23 V is the main saddle structure and is asymmetric due to collision processes.  $\text{SiH}_2^+$ , see Fig. 36(c), exhibits a 5 times lower peak intensity. The first five charge exchange maxima in the IED of  $\text{SiH}_2^+$  (Fig. 36c) are clearly visible. The IED intensity is about 5 times lower compared to  $\text{SiH}_3^+$ . Although the  $\text{SiH}_2^+$  ion is the main product of dissociative ionization of  $\text{SiH}_4$  [320], the measured IED intensity of  $\text{SiH}_3^+$  is always larger. The  $\text{H}^-$  transfer reaction causes  $\text{SiH}_3^+$  to be more abundant than  $\text{SiH}_2^+$ . Also in the IED of  $\text{Ar}^+$  (Fig. 36a) about six of the charge exchange maxima are clearly visible. The broad overall IED shows that there are relatively many ions which underwent a charge exchange reaction in the sheath. Even  $\text{H}_2^+$  (Fig. 36d) IEDs can be measured. The low intensity is due to the low partial pressure of hydrogen (it is created by silane dissociation only), and to the fact that the ionization potential of hydrogen is, like that of argon, higher than the one of silane. Therefore the number of created hydrogen ions will be low. The maximum in the IED at 11 eV is the first charge exchange maximum of  $\text{H}_2^+$ . The energy of this maximum is always somewhat lower than the energy of the fourth charge exchange maximum in the  $\text{SiH}_2^+$  IED. This is due to the almost four times higher plasma frequency of the  $\text{H}_2^+$  ion ( $m=2$  amu) with respect to the  $\text{SiH}_2^+$  ion ( $m=30$  amu). The  $\text{H}_2^+$  saddle structure is broader than the saddle structures of the other ions, due to the larger ion plasma frequency of  $\text{H}_2^+$ . The ions with an energy larger than 25 eV are part of the high energy side of the main saddle structure.

$\text{Si}_2\text{H}_4^+$  is an ion that is created in the plasma by polymerization reactions. Several pathways may lead to this ion. The first pathway is the dissociative ionization of  $\text{Si}_2\text{H}_6$ , that is formed in a radical-neutral reaction. The second pathway is the direct formation in the ion-molecule reaction [192]:  $\text{SiH}_2^+ + \text{SiH}_4 \rightarrow \text{Si}_2\text{H}_4^+ + \text{H}_2$ .

### V.C.3.c. Silane/hydrogen discharges

The IEDs of several ions in a typical silane/hydrogen plasma are shown in Fig. 37. The plasma was created in a mixture of 26 sccm  $\text{SiH}_4$ , 24.5 sccm  $\text{H}_2$ , at a pressure of 0.2 mbar. The applied RF frequency is 13.56 MHz, at a power of 10 W. The self bias voltage that developed was -118 V. The substrate temperature was 250 °C. The discharge is in the  $\alpha$  regime. The IED of  $\text{H}_2^+$  and  $\text{SiH}_2^+$  show distinct peaks. The position of the peaks are again described with the plasma potential, as is deduced from the  $\text{Si}_2\text{H}_4^+$  IED, and the charge carrier density  $n$ . The value for  $V_{pl}$  is  $(23.8 \pm 0.2)$  V and the value for  $n$  is  $(1.11 \pm 0.04) \times 10^8 \text{ cm}^{-3}$ . The sheath thickness is  $(4.88 \pm 0.10)$  mm. Under these and under most other conditions, the  $\text{SiH}_3^+$  IED shows no distinguishable charge exchange maxima.

The measured intensity in the  $\text{SiH}_3^+$  IED again is higher than the intensity of the  $\text{SiH}_2^+$  IED. There are two distinct peaks at low energies in the  $\text{SiH}_3^+$  IED and such peaks are observed in most  $\text{SiH}_3^+$  IEDs. Their energy positions can not be explained by charge exchange processes. The intensity in the  $\text{H}_2^+$  IED is higher than in the silane-argon plasma, simply because the partial pressure of hydrogen in this silane/hydrogen mixture is higher.

Charge exchange maxima are in general pronounced in the  $\text{SiH}_2^+$  IED in discharges at 13.56 MHz. The successive charge exchange peaks tend to overlap each other at

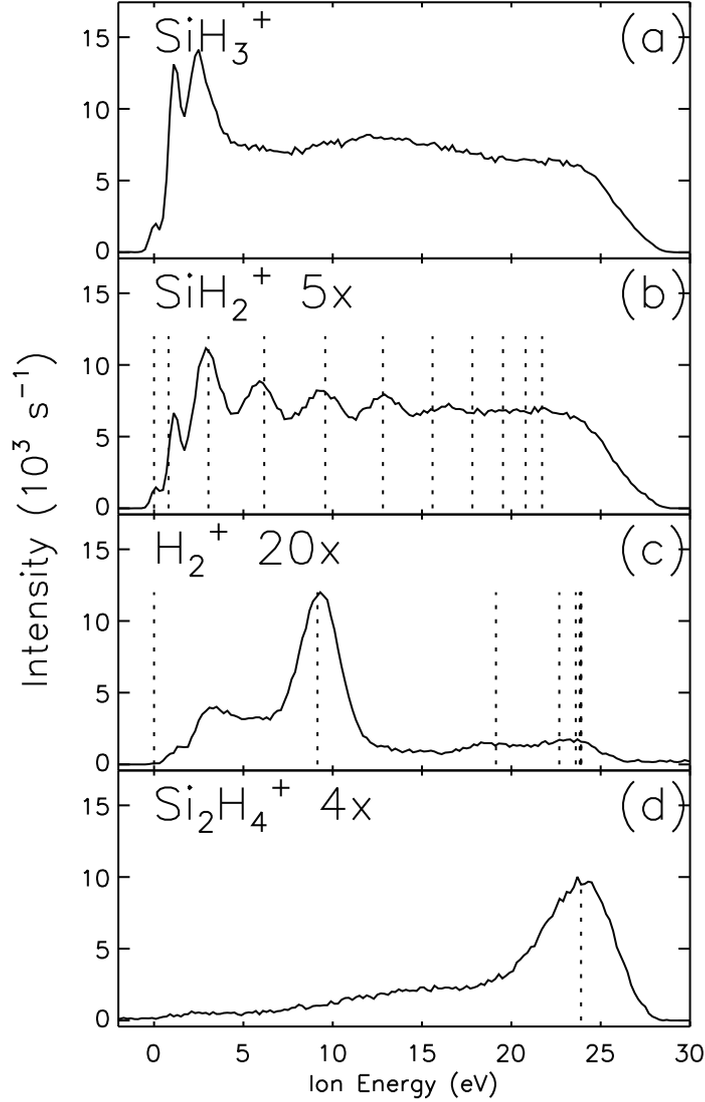


FIG. 37. The ion energy distributions of (a)  $\text{SiH}_3^+$ , (b)  $\text{SiH}_2^+$ , (c)  $\text{H}_2^+$ , (d)  $\text{Si}_2\text{H}_4^+$  from a  $\text{SiH}_4\text{-H}_2$  plasma. The conditions are 26 sccm  $\text{SiH}_4$ , 24.5 sccm  $\text{H}_2$ , 13.56 MHz, 10 W, 0.2 mbar,  $250^\circ\text{C}$ ,  $V_{DC} = -118$  V. The dotted lines in (b) and (c) indicate the position of the first 10 charge exchange peaks based on a plasma potential of 23.8 V and a charge carrier density  $n$  of  $1.11 \times 10^8 \text{ cm}^{-3}$ . The dotted line in the IED of  $\text{Si}_2\text{H}_4^+$  (d), indicates the value of the plasma potential. Note the different scaling factors of the various IEDs. (From E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

higher frequencies. The IED of  $\text{H}_2^+$  exhibits charge exchange peaks which are still well separated at higher frequencies due to the almost four times higher ion plasma frequency of  $\text{H}_2^+$  compared to the one of  $\text{SiH}_2^+$ .

Very different IEDs can be observed in discharges that are operated in the  $\gamma'$  regime, and that contain powder. The discharge, in this case, is created in a mixture of 30 sccm  $\text{SiH}_4$ , and 30 sccm  $\text{H}_2$  at a pressure of 0.60 mbar. The RF frequency is 13.56 MHz, and the applied power is 33.7 W. The DC self-bias voltage is -32.5 V. The substrate temperature is  $250^\circ\text{C}$ . The measured IEDs are shown in Fig. 38. The plasma potential is

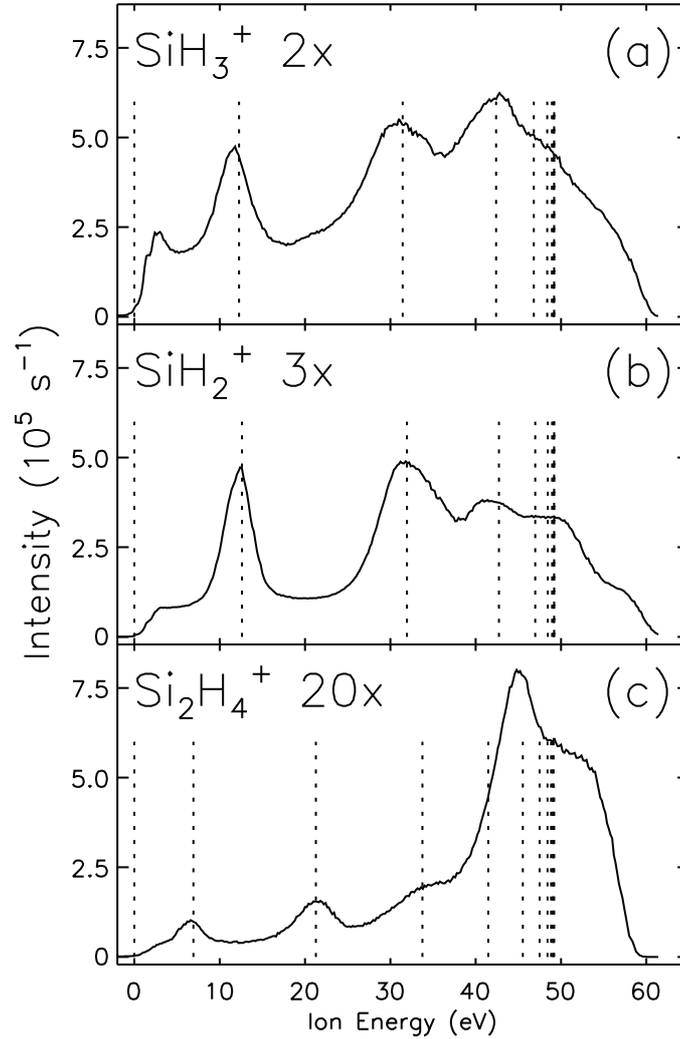


FIG. 38. The ion energy distributions of (a)  $\text{SiH}_3^+$ , (b)  $\text{SiH}_2^+$ , and (c)  $\text{Si}_2\text{H}_4^+$  from a  $\text{SiH}_4\text{-H}_2$  plasma in the  $\gamma'$  regime. The dotted lines in (a), (b), and (c) indicate the position of the first 10 charge exchange peaks based on a plasma potential of 49.2 V and a charge carrier density  $n$  of  $9.9 \times 10^8 \text{ cm}^{-3}$ . Note the different scaling factors of the various IEDs. (From E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

determined from the IED of  $\text{Si}_2\text{H}_4^+$ , and is  $49.2 \pm 0.5 \text{ V}$ , which is about twice the value in discharges operated in the  $\alpha$  regime. The charge carrier density is  $(9.9 \pm 0.3) \times 10^8 \text{ cm}^{-3}$ . The sheath thickness is  $(2.34 \pm 0.05) \text{ mm}$ .

The peaks in both the  $\text{SiH}_2^+$  (Fig. 38a) and the  $\text{SiH}_3^+$  (Fig. 38b) ion energy distribution are well separated and the intensity in the region between the charge exchange peaks is relatively low. The IED of  $\text{Si}_2\text{H}_4^+$  shows also charge exchange maxima in the low energy side of the IED, as well as an asymmetrical main saddle structure around 50 eV.

The sheath characteristics of the argon/silane and the two silane/hydrogen discharges of which the IEDs were shown above are summarized in Tab. 6. The sheath characteristics of the 10 W silane/argon discharge and the 10 W silane/hydrogen discharge in the  $\alpha$  regime are rather similar. The high power, high pressure discharge ( $\gamma'$

TABLE 6. Sheath properties of an argon/silane and two silane/hydrogen discharges ( $\alpha$  and  $\gamma'$  regime); plasma potential  $V_{pl}$ , charge carrier density  $n$ , sheath thickness  $d_s$ , electric field at the electrode  $E_0$ , ion flux  $\Gamma_{max}$ , and ion energy flux  $(\epsilon\Gamma)_{max}$ . (Compiled from E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

Property	Ar/SiH <sub>4</sub>	H <sub>2</sub> /SiH <sub>4</sub> ( $\alpha$ )	H <sub>2</sub> /SiH <sub>4</sub> ( $\gamma'$ )	Unit
$V_{pl}$	21.8±0.3	23.8±0.2	49.2±0.5	V
$n$	1.66±0.05	1.11±0.04	9.9±0.3	10 <sup>8</sup> cm <sup>-3</sup>
$d_s$	3.81±0.05	4.88±0.10	2.34±0.05	mm
$E_0$	11.4±0.2	9.8±0.2	42.0±0.8	kV/m
$\Gamma_{max}$	1.93±0.05	1.35±0.04	17.3±0.7	10 <sup>18</sup> m <sup>-2</sup> s <sup>-1</sup>
$(\epsilon\Gamma)_{max}$	6.7±0.2	5.2±0.2	136±4	W m <sup>-2</sup>

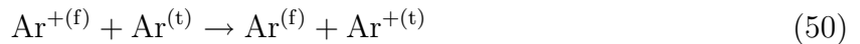
TABLE 7. Charge exchange reactions of Ar<sup>+</sup> and H<sub>2</sub><sup>+</sup> with SiH<sub>4</sub>. The cross sections ( $\sigma$ ) for reactions with Ar<sup>+</sup> are determined at thermal energies of the reactants [334], the cross sections for reactions with H<sub>2</sub><sup>+</sup> are determined at a kinetic energy of the reactants of 1 eV [335]. (Compiled from E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

Reaction	$\sigma$ (10 <sup>-16</sup> cm <sup>2</sup> )
Ar <sup>+</sup> + SiH <sub>4</sub> → Ar + Si <sup>+</sup> + 2H <sub>2</sub>	0.2
Ar <sup>+</sup> + SiH <sub>4</sub> → Ar + SiH <sup>+</sup> + H <sub>2</sub> + H	0.3
Ar <sup>+</sup> + SiH <sub>4</sub> → Ar + SiH <sub>2</sub> <sup>+</sup> + H <sub>2</sub>	0.4
Ar <sup>+</sup> + SiH <sub>4</sub> → Ar + SiH <sub>3</sub> <sup>+</sup> + H	2
H <sub>2</sub> <sup>+</sup> + SiH <sub>4</sub> → H <sub>2</sub> + Si <sup>+</sup> + 2H <sub>2</sub>	2
H <sub>2</sub> <sup>+</sup> + SiH <sub>4</sub> → H <sub>2</sub> + SiH <sup>+</sup> + H <sub>2</sub> + H	2
H <sub>2</sub> <sup>+</sup> + SiH <sub>4</sub> → H <sub>2</sub> + SiH <sub>2</sub> <sup>+</sup> + H <sub>2</sub>	11
H <sub>2</sub> <sup>+</sup> + SiH <sub>4</sub> → H <sub>2</sub> + SiH <sub>3</sub> <sup>+</sup> + H	34

regime) has a twice larger plasma potential, a twice smaller sheath distance, a four times larger electric field  $E_0$ , and a much larger ion flux and ion energy flux (see Sec. VI B 2) compared to the discharge in the  $\alpha$  regime.

#### V.C.3.d. Charge exchange reactions

The charge exchange reactions between Ar and Ar<sup>+</sup>, and between H<sub>2</sub> and H<sub>2</sub><sup>+</sup> are well known [325,328]:

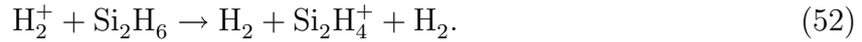


where ( $f$ ) and ( $t$ ) denote a fast ion or neutral and an ion or neutral with thermal energy. These reactions are so-called symmetrical resonant charge exchange reactions as they occur between an ion and the corresponding neutral.

The ionization energies of hydrogen (15.4 eV) and argon (15.8 eV) are higher than those of silane (11.6 eV) and disilane (9.9 eV). Therefore, ion-molecule reactions of  $\text{H}_2^+$  or  $\text{Ar}^+$  with silane or disilane will result in electron transfer from silane to the ion [190], which leads to dissociative ionization of silane. These known asymmetric charge exchange reactions result in  $\text{SiH}_x^+$  ( $0 < x < 3$ ) ions with thermal energy in the sheath, see Tab. 7 [334,335]. The charge exchange process of  $\text{H}_2^+$  with  $\text{SiH}_4$  is a major loss process for the hydrogen ions in typical silane plasmas.

The cross sections of these asymmetric charge exchange reactions are in general lower than the cross sections of resonant charge exchange processes [134]. For comparison, the cross section of the  $\text{Ar}^+$ -Ar charge exchange reaction is  $4 \times 10^{-15} \text{ cm}^2$ , about one order of magnitude larger than the reactions listed in Tab. 7. The total cross section of charge exchange reactions between  $\text{Ar}^+$  and  $\text{SiH}_4$  is much smaller than the total cross section of charge exchange reactions between  $\text{H}_2^+$  and  $\text{SiH}_4$ .

The discharge in which the charge exchange peaks of  $\text{Si}_2\text{H}_4^+$  are observed, is in the  $\gamma'$  regime. A considerable amount of silane will be depleted and gas phase polymerizations occur, which means that  $\text{Si}_2\text{H}_6$  is likely to be present in large high quantities. Therefore, Hamers *et al.* [235] have suggested that  $\text{Si}_2\text{H}_6$  is involved in the charge exchange reaction where  $\text{Si}_2\text{H}_4^+$  is created:



### V.D. Ellipsometry

Another *in situ* technique that is used to study deposition in real time is ellipsometry. This optical technique is non-invasive and does not perturb the discharge. In ellipsometry one measures the amplitude and phase of the reflected light from a surface. Spectroscopic ellipsometry (SE) allows for the wavelength-dependent measurement of the dielectric function of the material, at sub-monolayer sensitivity [336]. Spectroscopic phase-modulated ellipsometry (SPME) gives the possibility of fast data recording, and therefore is exploited for kinetic studies of the deposition of *a*-Si:H [337]. Spectroscopic ellipsometry is very sensitive tool to study to surface and interface morphology in the UV-visible range. The use of IR-light allows for the identification of Si-H bonding configurations [338].

Polarization modulation can be done by rotating the polarizer as is done in rotating-element ellipsometers [339] or by using photoelastic devices [337]. Data acquisition nowadays is fast which makes real-time measurements of film and interface formation possible: a full spectrum ranging from 1.5 to 5 eV can be measured in less than a second [340–342].

In a typical ellipsometry experiment a sample is irradiated with polarized light, that subsequently is reflected from the sample surface, and detected after passing an analyzer. The ratio  $\rho$  of complex reflectances for perpendicularly (*s*) and parallelly (*p*) polarized light usually is represented as follows:

$$\rho = \frac{r_p}{r_s} = \tan \Psi \exp(i\Delta) \quad (53)$$

with  $r_p$  and  $r_s$  the complex reflectances for p- and s-polarized light, and  $\Psi$  and  $\Delta$  two convenient angles. This ratio can directly be related to the complex dielectric function

$\epsilon = \epsilon_1 + i\epsilon_2$ , in the special case of light incident with an angle  $\phi$  on the surface of a semi-infinite medium [336]:

$$\epsilon = \sin^2 \phi \left( 1 + \tan^2 \phi \left[ \frac{1 - \rho}{1 + \rho} \right]^2 \right) \quad (54)$$

In case of multilayer structures much more complex expressions result.

In the IR one usually presents spectroellipsometric measurements using the optical density  $D = \ln(\bar{\rho}/\rho)$ , where  $\bar{\rho}$  refers to the substrate before deposition of a film [343]. The presence of a vibrational mode (rocking, bending, stretching) of a silicon-hydrogen bond, for instance, is revealed by a peak at the corresponding energy of that mode in the real part of  $D$  versus energy (in  $\text{cm}^{-1}$ ). The contribution of a specific vibration mode to the dielectric function of the film is estimated from the Lorentz harmonic-oscillator expression [344]:

$$\epsilon = \epsilon_\infty + \frac{F}{\omega_0^2 - \omega^2 - i\Gamma\omega} \quad (55)$$

with  $\omega_0$  the frequency of the mode,  $F$  its oscillator strength, and  $\Gamma$  a damping constant. The oscillator strength can be expressed as  $F = \omega_0^2(\epsilon_0 - \epsilon_\infty)$ , where  $\epsilon_0$  and  $\epsilon_\infty$  are the high- and low-frequency dielectric constants, respectively. The vibration mode of a chemical bond will show up as a maximum or minimum in the real part of  $D$  versus energy, depending on the dielectric function of the substrate.

The interpretation of ellipsometric data is based on the description of the optical properties of the material under study. Effective medium theories (EMTs) are widely used in this respect as they can be applied when materials are heterogeneous and when the size of inhomogeneities are small compared to the wavelength of the light, so that scattering of light can be neglected. In *a*-Si:H the presence of voids will have an effect on the dielectric function. Also, during deposition the surface will roughen more, or less, depending on deposition conditions, which will have a perturbing effect on the polarization of the incident light. An optical model of a substrate with an *a*-Si:H film could consist of a substrate, an *a*-Si:H film with voids, and a surface layer of *a*-Si:H with a large number of voids, which reflects the surface roughness.

The general form that is used in EMTs to describe the dielectric function  $\epsilon$  of an *a*-Si:H layer with a certain fraction of voids is given by [345]:

$$\epsilon = \frac{\epsilon_a \epsilon_v + \kappa \epsilon_h (f_a \epsilon_a + f_v \epsilon_v)}{\kappa \epsilon_h + (f_a \epsilon_v + f_v \epsilon_a)} \quad (56)$$

where  $\kappa$  is the screening parameter ( $\kappa = (1/q) - 1$  and  $0 \leq q \leq 1$ ),  $\epsilon_h$  is the host dielectric function,  $\epsilon_a$  and  $f_a$  are the dielectric function and volume fraction of the *a*-Si:H component, and  $\epsilon_v$  and  $f_v$  are the dielectric function and volume fraction of the void component. Several approximations can be made. At the end of the seventies Aspnes *et al.* [346] found that the Bruggeman approximation provided the best description for the data available. Recently Fujiwara *et al.* [345], with new spectroellipsometric data confirmed this fact. The Bruggeman approximation [61] defines  $\epsilon_h = \epsilon$ . For spherical inclusions  $q = \frac{1}{3}$ , i.e.,  $\kappa = 2$ , and the dielectric function is isotropic. The effect of voids present in the *a*-Si:H layer further simplifies Eq. 56, by taking  $\epsilon_v = 1$ , and substituting  $f_a = 1 - f_v$ .

In a real-time spectroellipsometric measurement in which the kinetics of *a*-Si:H deposition is studied, trajectories are recorded in the  $\Delta$ - $\Psi$  plane at various photon energies between 2 and 4 eV. These trajectories can be simulated, and fitted to models that represent the growing *a*-Si:H layer. Canillas *et al.* [347] have made a detailed study of the deposition of the first few layers of *a*-Si:H on a NiCr/glass substrate. Similar results are obtained for a *c*-Si substrate. They have proposed several models to explain the data. One possible model is the hemispherical nucleation model, that describes a hexagonal network of spherical *a*-Si:H nuclei located at an average distance  $d$  between them. The growth is represented by an increase in the radii of the nuclei, until these nuclei make contact. This results in an *a*-Si:H layer with a large void fraction, of  $f_v = 0.39$  [286]. From the start of the deposition ( $f_v = 1$ ) to the point where the nuclei make contact ( $f_v = 0.39$ ), the dielectric function is calculated using the Bruggeman EMT, and a  $\Delta$ - $\Psi$  trajectory can be simulated. In another model, the columnar nucleation model, columns of *a*-Si:H material start to grow from the initial hexagonal network, where the growth is represented by an increase in column radius and height. Again,  $f_v$  is calculated, and the dielectric function can be deduced from that. A further refinement is that the columns coalesce, and form a dense layer, with on top a layer of a certain thickness, that consists of free standing columns. This extra layer of typically 5 nm in thickness represents the roughness, and is larger for larger film thicknesses, for typical deposition conditions. A recent study supports the concept of coalescing 3D-islands of *a*-Si:H [340]: directly from the start of the deposition the thickness of the surface roughness layer increases upto 2 nm. At that thickness the islands start to combine into a continuous layer, and the thickness of the surface roughness layer decreases to remain constant at 1.5 nm.

Andújar *et al.* [246] have used SE to study the effects of pressure and power on the properties of *a*-Si:H. In their system the  $\alpha$ - $\gamma'$  transition occurs just above 0.2 mbar, at a power of 10 W. They found that the density of the material is decreased, going from the  $\alpha$  to the  $\gamma'$  regime, as is deduced from the decrease in the imaginary part of the dielectric function  $\epsilon_2$ .

Collins *et al.* [342] have used SE to study the effects of hydrogen dilution. On native oxide/*c*-Si substrates they have found that the thickness of the roughness layer is nearly constant at 1.5-2 nm in case of deposition with pure silane for a total deposited layer thickness up to 1000 nm. At a dilution factor  $R = [\text{H}_2]/[\text{SiH}_4]$  of 10, the thickness of the roughness layer decreases from 1.5 to just below 1 nm. In case of  $R = 20$ , a similar decrease is observed (with a minimum of 0.3 nm at a bulk layer thickness of 20 nm), however followed by a steep increase to 4.5 nm at a bulk layer thickness of 100 nm, which is consistent with the formation of a microcrystalline film. At values above  $R = 25$ , the deposited layer is microcrystalline. These deposition experiments have also been carried out starting on a 300 nm thick *a*-Si:H film, prepared with  $R = 0$ . Here, the amorphous to microcrystalline transition, as evidenced by the steep increase of the thickness of the roughness layer, occurs at 200 nm for  $R = 15$  and at 60 nm for  $R = 30$ . Based on these observations a phase-diagram was constructed, showing the amorphous to microcrystalline transition as a function of bulk layer thickness and the dilution factor  $R$ . The importance of this phase diagram lies in the fact that a deposition process that is close to the amorphous-microcrystalline transition yields material that has excellent opto-electronic properties [348–352].

Spectroscopic ellipsometry has also been applied in the characterization of compo-

sitionally graded  $a$ -SiC:H alloys [353], where the flow ratio  $z = [\text{CH}_4]/([\text{CH}_4 + \text{SiH}_4])$  was varied during deposition. The analysis of the SE-data of the graded layer prompted the development of a new four-medium model, which consists of the ambient, a surface roughness layer, an outerlayer, and the pseudosubstrate. A virtual interface approximation is applied at the boundary at the interface between outerlayer and pseudosubstrate. This model gives the near-surface carbon content, and the instantaneous deposition rate, as well as the thickness of the roughness layer. Besides  $a$ -Si:H/ $a$ -SiC:H interfaces also  $p/i$  interfaces have been characterized, which is possible due to the fact that the dielectric function changes at the interface [354,355].

The use of IR in spectroscopic ellipsometry allows for the investigation of the hydrogen incorporation in  $a$ -Si:H. The identification of vibrational modes in nm-thin films is difficult, due to the weak signals. Nevertheless, Blayo and Drévilon have shown that mono-hydride and di-hydride bonding configuration can be discerned in 0.5-1 nm thick  $a$ -Si:H layers [338], using IR phase-modulated ellipsometry (IRPME). In *in situ* IRPME studies the real part of  $D$  is presented in the stretching mode region of the IR, i.e., 1900-2200  $\text{cm}^{-1}$ . Three stretching modes can be revealed in this range, typically around 2000  $\text{cm}^{-1}$ , 2100  $\text{cm}^{-1}$ , and 2160  $\text{cm}^{-1}$ , corresponding to SiH, SiH<sub>2</sub>, and SiH<sub>3</sub> bonding configurations. From the evolution of the SiH- and the SiH<sub>2</sub> stretching mode with increasing deposited thickness, it was found [356] that the deposited layer is built up from two layers. The bottom layer contains SiH bonds, and linearly increases in thickness with deposition time. The top layer has an about constant thickness of 1-2 nm, and contains SiH<sub>2</sub> bonds. At the start of the deposition a layer consisting of SiH<sub>2</sub> is formed, and its thickness increases with time. At about a thickness of 1 nm, another layer between substrate and the SiH<sub>2</sub> containing layer is formed, which contains SiH. The deposition proceeds as the thickness of this latter layer increases with time. Attenuated total reflection IR spectroscopy has revealed that SiH<sub>2</sub> and SiH<sub>3</sub> surface modes are present even before the SiH<sub>2</sub>-containing layer is formed [340].

The position of the SiH and SiH<sub>2</sub> stretching modes varies with the thickness of the deposited layer. In the first few monolayers the SiH peak shifts upwards by about 20  $\text{cm}^{-1}$ , as a result of the very high hydrogen content [340]. At the end of the coalescence phase, the SiH peak is at 1995  $\text{cm}^{-1}$ , and its position does not change anymore. The substrate used was c-Si with a native oxide layer. In contrast, another study shows an increase in the SiH peak position from 1960  $\text{cm}^{-1}$  to 1995  $\text{cm}^{-1}$ , together with an increase in the SiH<sub>2</sub> peak position from 2060  $\text{cm}^{-1}$  to 2095  $\text{cm}^{-1}$ , for thicknesses of the deposited layer between 0 and 500 nm [344]. Moreover, these shifts depend on the deposition temperature, and on the substrate. These findings were related to the change in disorder as a function of deposited thickness. The disorder, characterized by the width of the Raman TO-peak, decreased with increasing thickness.

## VI. RELATION BETWEEN PLASMA PARAMETERS AND MATERIAL PROPERTIES

### VI.A. External parameters

Luft and Tsuo have presented a qualitative summary of the effects of various plasma parameters on the properties of the deposited  $a$ -Si:H [6]. These generalized trends are very useful in designing deposition systems. It should be borne in mind, however, that for each individual deposition system the optimum conditions for obtaining device quality material have to be determined by empirical fine-tuning. The most important external controls that are available for tuning the deposition process are the power (or power density), the total pressure, the gas flow(s), and the substrate temperature. In the following the effects of each parameter on material properties will be discussed.

#### VI.A.1. Plasma power

When the plasma power (density) is increased the deposition rate is increased monotonically up to the point where the gas flow rate becomes the limiting factor [81,357] (see Fig. 39). This deposition rate increase comes with a number of disadvantages, including poor film quality and powder formation. At low power levels a large part of the  $\text{SiH}_4$  remains undissociated [303], and the corresponding films contain only silicon monohydrides, inferred from FTIR measurements. At high power levels the  $\text{SiH}_4$  is strongly dissociated, and the films contain a large amount of  $\text{SiH}_2$  [358]. The microstructure parameter  $R^*$  is high. The high power levels also lead to columnar microstructure, which is accompanied by high spin densities [359,360]. Experimental results obtained in ASTER show similar trends, as is shown in Sec. VI B 3.

The electrical properties (dark- and photoconductivity) are reported to first decrease and then increase upon increasing power [361]. The optical band gap increases with increasing power due to the increase of the hydrogen content [63,82,362,363]. However, at very high power levels, microcrystalline silicon is formed [364], which causes the hydrogen content (and, consequently, the band gap) to decrease.

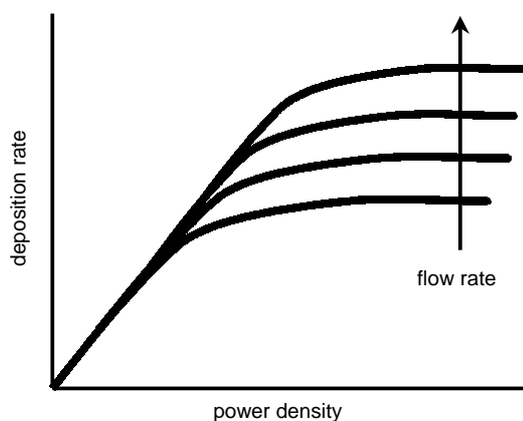


FIG. 39. Schematic representation of the influence of power density and flow rate on the deposition rate. (After A. Matsuda, *J. Vac. Sci. Technol. A* **16**, 365 (1998).)

As a result of deterioration of film properties with increasing power levels, to obtain device quality material the use of low power is required, albeit with a concomitant low deposition rate. The increase of deposition rate while unaltering the device quality material properties is a large research challenge.

#### *VI.A.2. Total pressure*

The deposition rate as a function of total pressure shows two different dependencies. In the low-pressure regime or supply-limited regime the deposition rate is proportional to the pressure, while in the power-limited regime it is constant [365]. At low pressure the film quality is not that good. The supply of  $\text{SiH}_3$  radicals, needed for high quality films, is depleted easily, and  $\text{SiH}_2$  and  $\text{SiH}$  radicals can reach the substrate. Moreover, ion bombardment may be too severe. Therefore, higher pressure levels are required. Usually the operating pressure (in combination with electrode distance and power level) is set such that one works just above the Paschen curve (see Sec. III B 4). The pressure should not be too high, otherwise gas-phase polymerization occurs [55]. These yellow powders (which turn white upon oxygen exposure) are to be avoided, as they increase the downtime of deposition systems due to clogging of the pumps. At normal working pressures it has been found that the hydrogen content and dihydride content increase as a function of pressure [81].

#### *VI.A.3. Gas flow*

The gas flow rate is presented usually as a deposition parameter, however it is much more instructive to report gas residence time [6], which is determined from the flow rate and the geometry of the system. The residence time is a measure of the probability of a molecule to be incorporated into the film. The gas depletion, which is determined by the residence time, is a critical parameter for deposition. At high flow rates, thus low residence times and low depletion [303], the deposition rate is increased [357,365] (see Fig. 39) and better film quality is obtained, as is deduced from low microstructure parameter values [366].

#### *VI.A.4. Hydrogen dilution*

The addition of a hydrogen flow to the silane flow often is used to improve the material quality of the deposited film. Moderate hydrogen dilution of silane ( $0.15 \leq [\text{SiH}_4]/([\text{SiH}_4]+[\text{H}_2]) \leq 1$ ) has been found to yield a lower optical band gap, a lower activation energy, a lower total hydrogen content, a lower microstructure factor, and a higher photoresponse [15,82,367–369]. Unfortunately, also the deposition rate is reduced. Device quality material is obtained for values of  $[\text{SiH}_4]/([\text{SiH}_4]+[\text{H}_2])$  between about 0.3 and 0.7. Moreover, an improvement of the uniformity of deposition over a  $10 \times 10 \text{ cm}^2$  substrate area has been found as well [370]: for  $[\text{SiH}_4]/([\text{SiH}_4]+[\text{H}_2]) = 1/3$  the variation in thickness amounted to only 1.5%.

In the intermediate regime of hydrogen dilution ( $0.05 \leq [\text{SiH}_4]/([\text{SiH}_4]+[\text{H}_2]) \leq 0.15$ ) the hydrogen content increases again, as well as the band gap, while the microstructure

factor remains low [369]. In this regime wide band gap  $a$ -Si:H can be obtained with better optoelectronic properties than  $a$ -SiC:H.

Highly diluted silane ( $[\text{SiH}_4]/([\text{SiH}_4]+[\text{H}_2]) \leq 0.05$ ) causes the deposited films to be crystalline rather than amorphous, due to selective etching by atomic hydrogen of strained and weak bonds [371,372]. Interestingly, using hydrogen dilution to deposit  $a$ -Si:H films "on the edge of crystallinity" [154,349] resulted in an improved stability. These materials are also referred to as polymorphous silicon (pm-Si) [373,374].

The admixture of a small amount of silane to a pure hydrogen plasma drastically changes the plasma potential, the DC self-bias voltage and the charge carrier density. Hamers [163] has found a doubling of the carrier density by changing  $\text{SiH}_4/(\text{SiH}_4+\text{H}_2)$  from 0 to 0.1. The DC self bias becomes more negative by 25%, and the plasma potential lowers by about 10-15%. Between values from 0.1 to 0.5 these parameters remain about constant. The parameters change again when  $\text{SiH}_4/(\text{SiH}_4+\text{H}_2)$  is larger than 0.5.

#### VI.A.5. Substrate temperature

The substrate temperature is a very important deposition parameter, as it directly affects the kinetics of ad- and desorption of growth precursors, surface diffusion, and incorporation. Actual substrate temperatures may differ from substrate heater setpoints. Calibration of temperature readings are needed, so as to report the correct substrate temperature.

The deposition rate is nearly independent of temperature, while the total hydrogen content, the microstructure parameter, and the disorder decrease with increasing temperature [375], see also Fig. 40 [84,85]. As a consequence, the optical band gap decreases as well. An optimum deposition temperature around 250 °C exists, and the material contains only SiH bonds. At higher temperatures the hydrogen evolution from the film causes the lowering of the hydrogen content, and leads to higher defect densities. The photoconductivity and photoresponse also have an optimum value around 250 °C.

#### VI.A.6. $\alpha$ - $\gamma'$ transition

As was shown above, the properties of the discharge will change with process parameters such as the process pressure, the RF power and the excitation frequency. Two different plasma regimes can be distinguished: the  $\alpha$  and the  $\gamma'$  regime [184,244–246,248]. In the transition from the  $\alpha$  regime to the  $\gamma'$  regime a change occurs in plasma properties. This includes the plasma impedance, the optical emission from the plasma [184], and the DC self-bias voltage at the powered electrode [245] in a reactor with electrodes of unequal size.

An increase in the deposition rate of the film and a change of the film properties is observed near the transition from the  $\alpha$  to the  $\gamma'$  regime [245]. As an example, Fig. 41 shows the deposition rate as a function of pressure, for various hydrogen dilution ratios [376]. It can be seen that higher hydrogen dilution pushes the  $\alpha$ - $\gamma'$  transition away, as it were. Other examples are shown in Fig. 46 and Fig. 64. The transition is assumed to be initiated by the formation of larger particles in the gas phase [246,322].

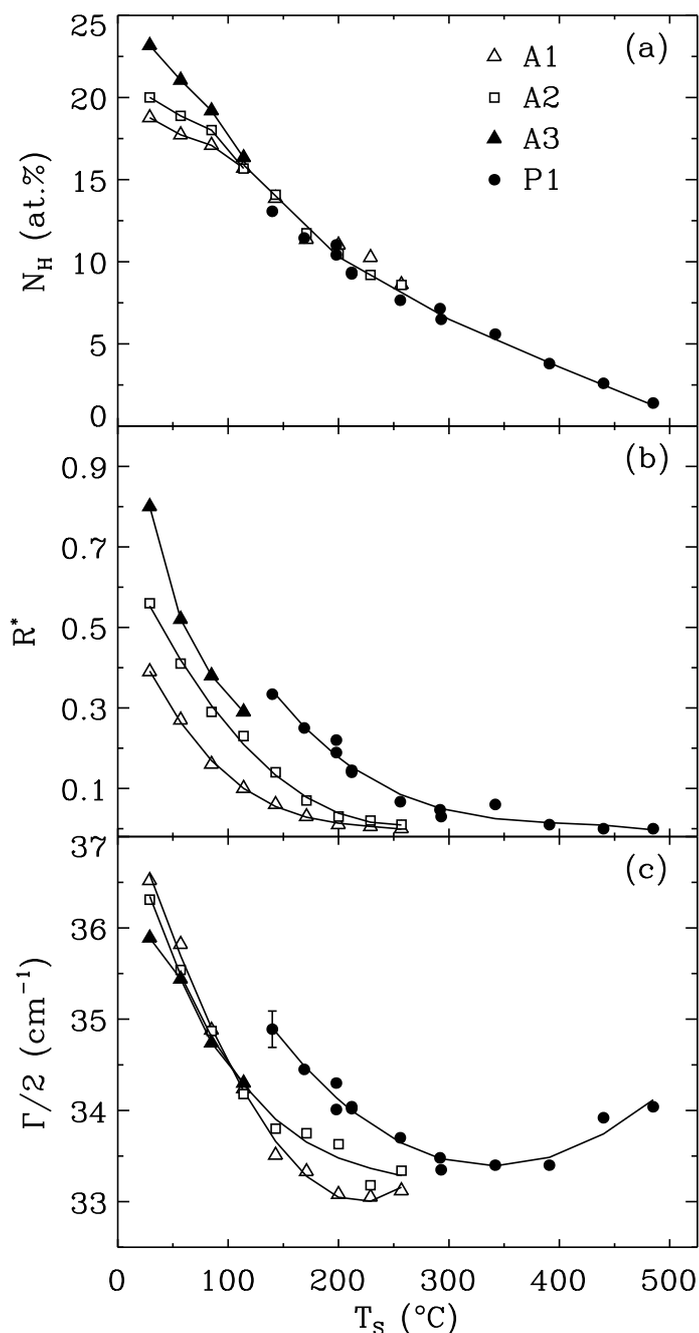


FIG. 40. The influence of deposition temperature on (a) the hydrogen concentration, (b) the microstructure parameter, and (c) the Raman half-width  $\Gamma/2$ . The labels A and P refer to the ASTER and the PASTA deposition system. Series A1 was prepared from a  $\text{SiH}_4/\text{H}_2$  mixture at 0.12 mbar. Series A2 and A3 were deposited from undiluted  $\text{SiH}_4$  at 0.08 and 0.12 mbar, respectively. Series P1 was deposited from undiluted  $\text{SiH}_4$ . (From A. J. M. Berntsen, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

Negative ions [377] are trapped in the plasma bulk, and recombine with silane radicals to become large negative ions (or small clusters). The charge of these particles can fluctuate, and can become positive. The clusters coalesce, and when the size of the coagulates is sufficiently large the charge builds up and remains negative. They grow

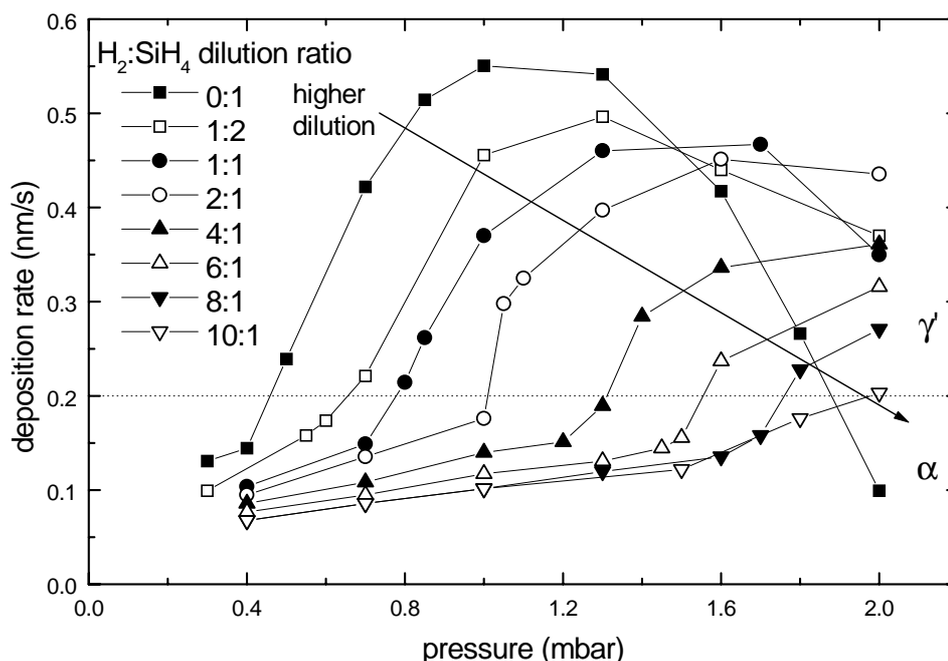


FIG. 41. The deposition rate as a function of deposition pressure for different ratios of hydrogen dilution ( $0:1 < \text{H}_2:\text{SiH}_4 < 10:1$ ). The deposition temperature was  $240^\circ\text{C}$ , the  $\text{SiH}_4$  flow amounted to 600 sccm and the excitation frequency was 13.56 MHz. The arrow indicates increasing hydrogen dilution. The  $\alpha$  regime is below the dotted line, the  $\gamma'$  above it. (After R. B. Wehrspohn, S. C. Deane, I. D. French, I. Gale, J. Hewett, M. J. Powell, and J. Robertson, *J. Appl. Phys.* **87**, 144 (2000).)

further by deposition of  $\alpha$ -Si:H on their surface. These particles are often referred to as powder or dust [378]. Due to the large amount of negative charge (up to hundreds of electrons) on this powder, the free electron density drops. Therefore, the electron temperature has to increase in order to sustain the discharge. The discharge becomes more resistive, causing more efficient power coupling into the plasma [322]. Most of these phenomena are observed in discharges with the conventional 13.56 MHz excitation frequency, but can also be observed at higher excitation frequencies [301].

Interestingly, it has been argued that nanoparticulate formation might be considered as a potential for obtaining new silicon films [379]. The nanoparticles can be crystalline and this fact prompted a new line of research [380–383]. If the particles that are suspended in the plasma are irradiated with, e.g., an Ar laser (488 nm) photoluminescence is observed when they are crystalline [384]. The broad spectrum shifts to the red, due to quantum confinement. Quantum confinement enhances the band gap of material when the size of the material becomes smaller than the radius of the Bohr exciton [385,386]. The broad PL spectrum shows that a size distribution of nanocrystals exist, with sizes lower than 10 nm.

Moreover, it was found that incorporation of nanoparticles of about 8 nm in diameter in  $\alpha$ -Si:H lead to improved properties, the most important one being the enhanced stability against light soaking and thermal annealing [387]. A later study revealed a typical crystallite size of 2–3 nm, with a hexagonal close packed structure [388]. Also diamond structures can be observed [389]. Hence the name polymorphous silicon is justified.

In view of the above, instead of avoiding powder formation regimes in discharges, careful powder management involving optimization of the ratio of radicals and silicon nanoparticles arriving on the substrate has been proposed [379].

#### VI.A.7. Other deposition parameters

Depending on the flow pattern in the reactor, the depletion of gases can cause nonuniform deposition across the surface of the substrate. As is depicted in Fig. 4, the gas usually is introduced at the top of the cylindrically symmetric reactor, and it flows into (and out of) the discharge region from the sides. Many other configurations exist, e.g., a radial flow reactor, where the gas flow is introduced underneath the powered electrode, and pumped away through the center of this electrode. A reverse radial flow reactor, where the gas flow is introduced from the center of the powered electrode, and pumped away at the sides has also been proposed [117,370]. The best solution to overcome depletion is to directly inject the gas via a showerhead at the grounded electrode [173].

Meiling *et al.* [82] have investigated the effect of electrode shape. They found that in the case of a ring electrode less dense material with a higher band gap was obtained than in the case of a flat-plate electrode. In addition, the ring electrode induces an extra non-uniformity in thickness, as a result of the non-uniform electric field between the powered and grounded electrode. Further, the area ratio of grounded to powered electrode in the case of the ring electrode is much higher, which yields higher DC bias voltages, and higher ion bombardment energies. Hence, a ring electrode should not be used.

Many different substrates are used for *a*-Si:H deposition. Usually Corning 7059 glass [390] and crystalline silicon are used for materials research purposes, as both have similar thermal expansion coefficients as *a*-Si:H. Devices are mostly made on Transparent Conductive Oxide (TCO)-coated glass. As TCO coatings one uses indium tin oxide (ITO), fluorine-doped tin oxide ( $\text{SnO}_2\text{:F}$ ), and zinc oxide (ZnO). Ion bombardment may lead to the reduction of the ITO coating [391], while ZnO-coated glass is much more resistant to this. Polymer films have been used as substrates for flexible solar cell structures. They require lower deposition temperatures than glass or stainless steel.

It has been found that various material properties are thickness-dependent. Raman experiments show a dependence on the type of substrate (glass, c-Si, stainless steel, ITO/glass) and on the thickness (up to 1  $\mu\text{m}$ ) of the films [392,393]. Recent Transmission Electron Microscopy (TEM) results also show this [394]. This is in contrast to other results, where these effects are negligible for thicknesses larger than 10 nm [395,396], which is also confirmed by ellipsometry [397] and infrared absorption [398] studies.

### VI.B. Internal parameters

#### VI.B.1. The role of ions

There are many effects possible due to the interaction of ions with a surface during deposition, e.g., the enhancement of adatom diffusion, ion-induced desorption, dis-

placement of surface or sub-surface atoms, sticking or sub-surface trapping of the ions, sputtering, and implantation in sub-surface layer (subplantation). Light and slow ions are able to excite SiH surface bonds vibrationally, which may enhance the surface migration or desorption of weakly physisorbed species, such as SiH<sub>3</sub>. For the nonmobile chemisorbed species, such as SiH<sub>2</sub>, to be desorbed, a higher ion mass and/or energy is needed. At the same time, however, this may cause subplantation and resulting collision cascades or thermal spikes, which will lead to local defects, and poor electronic properties [173].

Several views exist on the exact processes that occur during the deposition of *a*-Si:H. Ganguly *et al.* [399] explain the growth of *a*-Si:H using a surface diffusion model, in which ions are ignored. Perrin [211] has formulated a model of the RF discharge, in which also ions are not thought to be of importance to the growth. On the other hand, Vepřek *et al.* [400] introduce the term ion induced dehydrogenation, and Heintze *et al.* [249,280] show the importance of ion fluxes in plasmas, which are excited with very high frequency (VHF) electrode voltages. In addition, molecular dynamics studies of low-energy (10 eV) particle bombardment on crystalline silicon surfaces show a.o. surface dimer bond breaking [401] and the formation of interstitials [402]. Few measurements are available on the ion energies and fluxes on the sheath of a discharge under typical deposition conditions [280,403–405]. The influence of ion bombardment on defect density [405], mobility [404], and electronic properties [403,406] has been reported. The dependence of stress on the ion bombardment has been suggested by many authors [284,376,407,408]. In the field of PECVD of amorphous or diamond-like carbon, it is assumed that ions are of prime importance for the formation of dense material [409]. Others also stress the importance of ion bombardment on film properties [410–412]. In addition, in electron beam deposition of amorphous silicon it was shown that the formation of microvoids is inhibited, when Ar<sup>+</sup> assistance is used during deposition [413]. It was found that the energy per atom ratio is the decisive parameter instead of the ion per atom ratio. Supplying 12 eV per Ar<sup>+</sup> ion is sufficient for annihilation of all microvoids. As a further example, calculation of projected ranges of low-energy particles in a solid can be performed with the Monte-Carlo simulation program TRIM (TRansport of Ions in Matter) developed and supplied by Ziegler *et al.* [414]. The results of simulations with TRIM92 (TRIM Version 1992) of hydrogen and silicon ions incident on *a*-Si:H ( $C_H = 0.1$ ) with an energy of 10 and 50 eV, respectively, are presented in Fig. 42. The projected ranges of hydrogen (1 nm at 10eV, 2.5 nm at 50 eV) are much larger than the ones of silicon (0.3 nm at 10eV, 0.75 nm at 50 eV). Clearly, these low energy ions influence the surface and subsurface layers of the deposited film.

It is difficult to solely assess the effect of ion bombardment on material properties. An external bias can be applied to the discharge in an attempt to study the effect of varying acceleration voltages over the sheath. However, this modifies the whole potential profile, and consequently the discharge chemistry. Conclusions drawn from such experiments obviously are misleading. It has been demonstrated in a multipole discharge at low pressure (< 5 mTorr) that the ion flux can be controlled without disturbing the plasma properties and radical flux [358,415–418]. At such low pressure the ion flux is monoenergetic and the relative reactive ion flux ( $\Gamma_+/\Gamma_0$ ) can be as large as 0.8. Perrin [173] has summarized the effects of ion bombardment for  $\Gamma_+/\Gamma_0 \approx 0.2$  and ion energy up to 150 eV: the deposited films show a densification of the microstructure, a disappearance of columnar growth, and a reduction of surface roughness. Further,

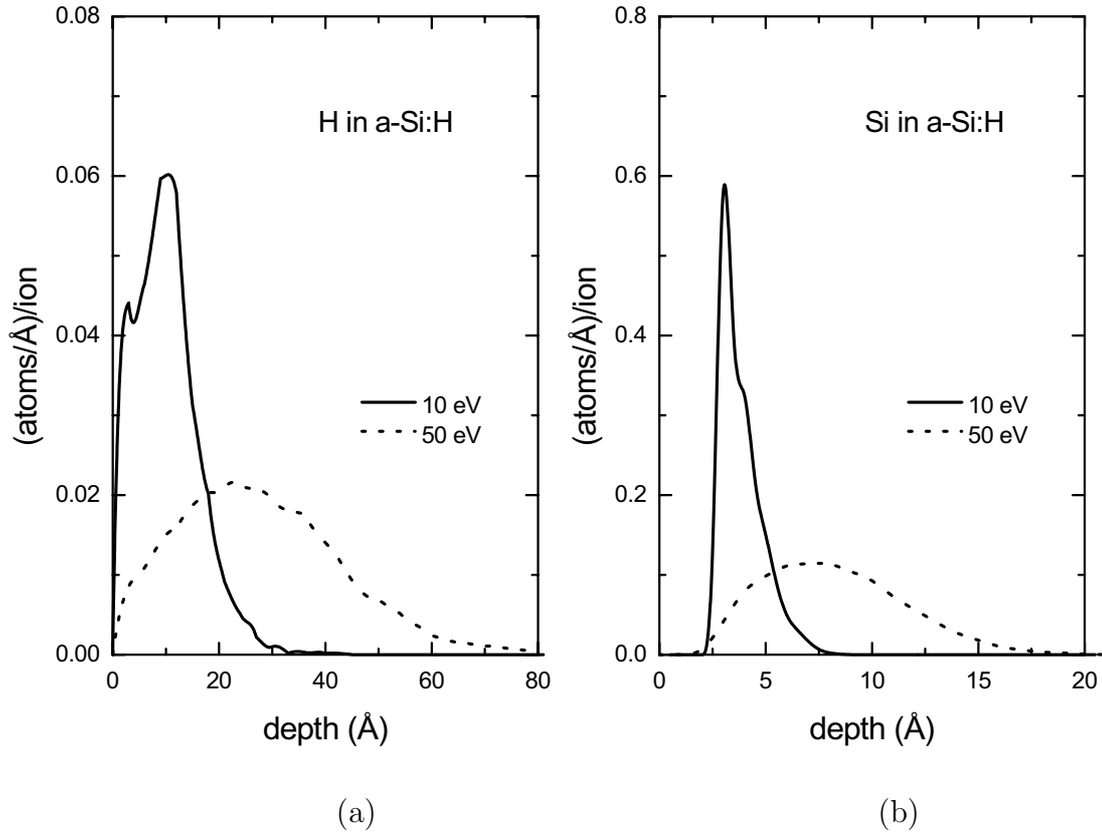


FIG. 42. Simulated depth distributions of hydrogen (a) and silicon (b) ions incident on *a*-Si:H with an energy of 10 and 50 eV. Note the difference in depth-scale. Simulations were performed with TRIM92.

the monohydride bonds in the network are favoured over dihydride bonds, i.e.  $R^*$  is lowered to about zero, and the compressive stress present in the material is increased. Also the optoelectronic properties are improved.

In an RF discharge the relative reactive ion flux is much lower (0.03 [358]) and the IED is considerably broadened. Kae-Nune *et al.* [311] have estimated from measurements on radicals, that ions could account for about 10% of the deposition rate. A straightforward method to directly determine the contribution of ions to the deposition process has been presented by Hamers *et al.* [163,419]. The ions and radicals are separated by exploiting the difference in angular distribution of their velocities. As ions are accelerated towards the electrode, their angular distribution of ion velocities will be narrow. Radicals diffuse through the sheath, and their angular velocity distribution will be isotropic. If one places an aperture in front of a substrate, the ions will mainly reach the substrate directly behind the aperture, while radicals will spread over a much larger area on the substrate. A similar configuration has been used to determine the reaction probability of radicals [420]. The effect is clearly demonstrated in Fig. 43: the thickness (deposition rate) is enhanced by about a factor of two, just behind the aperture. The deposition conditions were chosen such as to yield good quality material. From geometrical modeling it was found that under these discharge conditions the contribution of ions to the deposition was 10% [419], which confirms the estimate from Kae-Nune *et al.* [311].

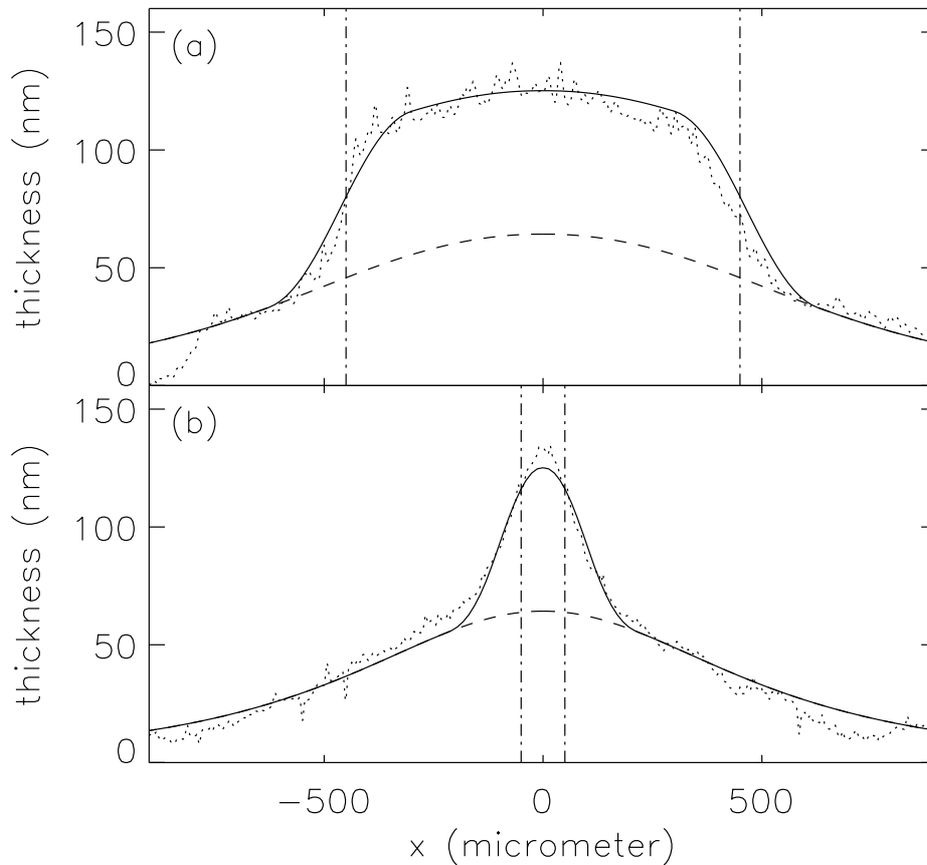


FIG. 43. Measured and simulated thickness profiles on the substrate behind the slit for a film deposited under discharge conditions that typically yields good material properties: (a) along the length of the slit, (b) across the slit. The vertical dashed-dotted lines indicate the boundaries of the apertures. The dotted lines represent the measured profiles, the solid lines are the simulated profiles. The dashed lines are the simulated deposition profiles of the radicals. (From E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

### VI.B.2. The ion flux

In order to study the influence of ions on the deposition process, a reliable quantification of the ion flux and energy is imperative. This flux cannot be determined directly from the detected number of ions in an IED as measured by means of QMS, for three reasons [332]. First, the orifice size decreases during subsequent measurements due to deposition of a-Si:H on the edges of the orifice. Second, due to the limited acceptance angle of the mass spectrometer system, only a fraction of the ions that arrive at the substrate is actually detected. This fraction depends on the type and number of interactions that an ion experiences while traversing the sheath, and also on the ion species itself. In addition, this implies that the mean kinetic energy of the impinging ions cannot be determined. Third, the ion flux from a typical silane plasma consists of many different ions. All ions must be taken into account in the quantification of the ion flux.

Another approach is needed [301,163,332], and follows from the definition of the ion

flux  $\Gamma$ , i.e. the product of the density of the ions  $n_i$  and their mean velocity  $\bar{v}_i$ :  $\Gamma = n_i \bar{v}_i$ . The contribution of electrons to the time averaged charge carrier concentration at the electrode  $n$  can be neglected, as the electron current towards the electrode is peaked in a very short time during the RF period. We thus have  $n_i = n$ . A further assumption is that there are no collisions in the sheath. As a result of this, the velocity  $\bar{v}_i$  is equal to the maximum velocity,  $v_{max}$ , which the ions gain. This velocity  $v_{max}$  follows from the time-averaged plasma potential  $eV_{pl}$ :  $v_{max} = \sqrt{2eV_{pl}/\bar{m}}$ , with  $\bar{m}$  the average mass of the ions. If the ions reach the electrode without collisions in the sheath, the IED will show a distinct peak at  $eV_{pl}$ , see e.g. the IED of  $\text{Si}_2\text{H}_4^+$  in an  $\alpha$  silane discharge (Fig. 37).

The maximum ion flux  $\Gamma_{max}$  at the electrode is thus estimated to be:

$$\Gamma_{max} = nv_{max} = n\sqrt{\frac{2eV_{pl}}{\bar{m}}} \quad (57)$$

The maximum ion *kinetic* energy flux of ions  $(\epsilon\Gamma)_{max}$  ( $\text{Wm}^{-2}$ ) can be defined as:

$$(\epsilon\Gamma)_{max} = eV_{pl}\Gamma_{max} \quad (58)$$

where  $\epsilon$  denotes the ion energy.

The 'collisionless sheath' assumption leads to an overestimation of the velocity. The use of an average mass  $\bar{m}$ , which is taken to be 31 a.m.u., i.e., the mass of  $\text{SiH}_3^+$ , in addition may lead to an overestimation of the velocity, as ions with more than one silicon atom are present, albeit in lower concentrations. Hamers *et al.* [163,301,332] have argued that the velocity of the ions (and the flux) is overestimated by at most a factor of two. Also the ion kinetic energy flux overestimates the real energy flux, if collisions take place in the sheath. However, in many collisions the kinetic energy is distributed over the different particles. For instance, in the situation of a charge exchange process, the original ion is neutralized, but retains its kinetic energy. The newly formed ion will gain the energy which the neutralized ion would have gained in the rest of the trajectory. Both particles will transport the total amount of kinetic energy corresponding to the plasma potential to the surface [163].

In order to express the importance of the ions to the growth process quantitatively, two related quantities can be defined: the fraction of arriving *ions* per deposited *atom*,  $R_i$ , and the kinetic energy transferred by *ions* per deposited *atom*,  $E_{max}$ . These quantities are used in ion beam assisted deposition in order to relate material properties to ion flux and energy [421]. Their definition is:

$$R_i = \frac{\Gamma_{max}}{\rho r_d} \quad (59)$$

$$E_{max} = \frac{(\epsilon\Gamma)_{max}}{\rho r_d} \quad (60)$$

where the quantity of deposited atoms per unit of time and unit of area is  $\rho r_d$ , with  $\rho$  the atomic density of the amorphous network ( $5 \times 10^{22} \text{ cm}^{-3}$ ) and  $r_d$  the growth rate.

### VI.B.3. Relation between ion flux and material quality

A systematic study of the role of the ions in the deposition process and their influence on the quality of the layers has been performed by Hamers *et al.* [163,301,332] in the

ASTER deposition system. More specifically, a study has been made on the relation between the plasma parameters and the material properties in both the  $\alpha$ - and the  $\gamma'$ -regime at typical deposition conditions. Here, the results for power and pressure variation are summarized. Details can be found elsewhere [163,301,332].

First, results on power variation are described, for two different discharges: 1) a silane/hydrogen discharge in the powder-free  $\alpha$  regime, and 2) a silane/argon discharge in both the  $\alpha$  and the powder producing  $\gamma'$  regime.

The process conditions of the series in silane/hydrogen are an excitation frequency of 13.56 MHz, a pressure of 0.20 mbar, gas flows of 30 sccm  $\text{SiH}_4$  and 30 sccm  $\text{H}_2$ , and a substrate temperature of 250 °C. The power is varied between 5 and 20 W. It is found that the absolute magnitude of the DC self-bias voltage increases linearly with increasing power, the trend of the ion flux is similar ( $0.8 \times 10^{18} \text{ m}^{-2}\text{s}^{-1}$  at 5 W, and  $2.1 \times 10^{18} \text{ m}^{-2}\text{s}^{-1}$  at 20 W). This linear relationship between  $V_{DC}$  and growth rate has been observed earlier [15,151,422], and in practice can even be used as a calibration curve. The plasma potential is 22.7 V at 5 W and is only 3 V higher at 20 W. In this series only two films have been deposited, one at 10 W and one at 15 W. The refractive index in both cases is 4.33, i.e. representative for good material. The fractions of arriving ions per deposited atom,  $R_i$ , as defined in Sec. VIB 2 are 0.26 and 0.20, respectively, while the kinetic energy transferred by ions per deposited atom,  $E_{max}$  is about 5 eV, in both cases.

The process conditions of the series in silane/argon are an excitation frequency of 50 MHz, a pressure of 0.40 mbar, gas flows of 30 sccm  $\text{SiH}_4$  and 60 sccm Ar and a substrate temperature of 250 °C. The power is varied between 3 and 50 W. Plasma parameters deduced from measurements of the IED and material properties are shown in Fig. 44 and Fig. 45. The discharge changes from the  $\alpha$  regime to the  $\gamma'$  regime around 10 W, which can best be observed by the decrease of the DC self bias and an increase in sheath thickness, see Fig. 44a and Fig. 44b. The bias is very low in the  $\gamma'$  regime, typically between -2 and -1 V, indicative of a more symmetrical discharge. The plasma potential increases with increasing power and tends to saturate at the highest powers. Visual inspection of the discharge shows that the optical emission from the plasma becomes brighter and more homogeneously distributed, when going from the  $\alpha$  to the  $\gamma'$  regime. In the  $\alpha$  regime the emission intensity is the highest at the boundary between plasma bulk and sheaths.

The growth rate, plotted in Fig. 44d, increases with increasing power, which is ascribed to the higher degree of dissociation in the discharge. The growth rate reaches a maximum around 30 W, and decreases somewhat at higher powers: the growth rate then is limited by the flow rate. It has been estimated that about 60% of the silane is used for deposition [163], the rest may well (partially) partake in the processes that lead to the creation of powder. These yellow powder is always observed in the reactor after the experiments performed in the  $\gamma'$  regime. The powder is always found outside the plasma confinement, and never on the substrate.

At the transition from the  $\alpha$  to the  $\gamma'$  regime the enhancement of the growth rate (Fig. 44d) is larger than the change in ion flux (Fig. 44c), as shown in Fig. 45a, where  $R_i$  is plotted. A value of  $R_i$  of 0.25 or larger is commonly found in the  $\alpha$  regime, whereas  $R_i$  is typically 0.10 or lower in the  $\gamma'$  regime. In addition, the amount of kinetic ion energy per deposited atom  $E_{max}$  shows a minimum of about 2 eV between 10 and 20 W, see Fig. 45b.

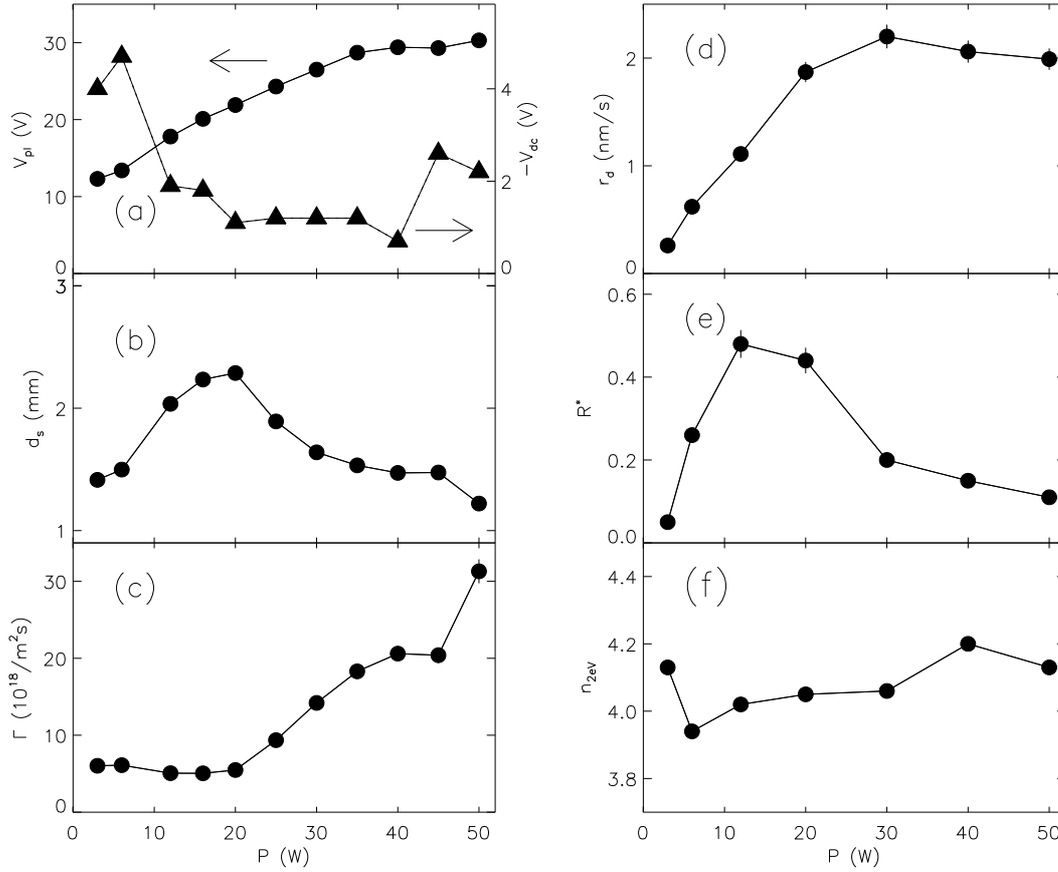


FIG. 44. Plasma parameters as deduced from the IEDs and material properties as a function of power delivered to the  $\text{SiH}_4/\text{Ar}$  discharge at an excitation frequency of 50 MHz and a pressure of 0.4 mbar: (a) the plasma potential  $V_{pl}$  (circles) and DC self bias  $V_{DC}$  (triangles), (b) the sheath thickness  $d_s$ , (c) the maximum ion flux  $\Gamma_{max}$ , (d) the growth rate  $r_d$ , (e) the microstructure parameter  $R^*$ , and (f) the refractive index  $n_{2eV}$ . (Compiled from E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

The microstructure parameter is low in the material deposited at the lowest power, (Fig. 44e), while it increases rapidly with increasing power up to 20 W, and then decreases again with further increasing power. The opposite holds for the refractive index (Fig. 44f), although less clear. A high value of the microstructure indicates a relatively large fraction of Si-H<sub>2</sub> bonds in the material, corresponding to an open material structure, and a low refractive index.

From a comparison between the behaviour of the microstructure parameter  $R^*$  (Fig. 44e) and the ion kinetic energy per deposited atom  $E_{max}$  (Fig. 45b) it can be concluded that a one-to-one relation between the relative strength of the ion bombardment, expressed in terms of  $E_{max}$ , and the microstructure parameter appears to exist. This has also been suggested by others [246,422].

Both the dark- and photoconductivity of the sample deposited at 40 W have been measured. The photoconductivity has been measured under AM1.5 conditions. The dark- and photoconductivity of the material deposited at 40 W are  $1 \times 10^{-10} \Omega^{-1}\text{cm}^{-1}$  and  $1.2 \times 10^{-4} \Omega^{-1}\text{cm}^{-1}$ , respectively. These values are indicative of good electronic

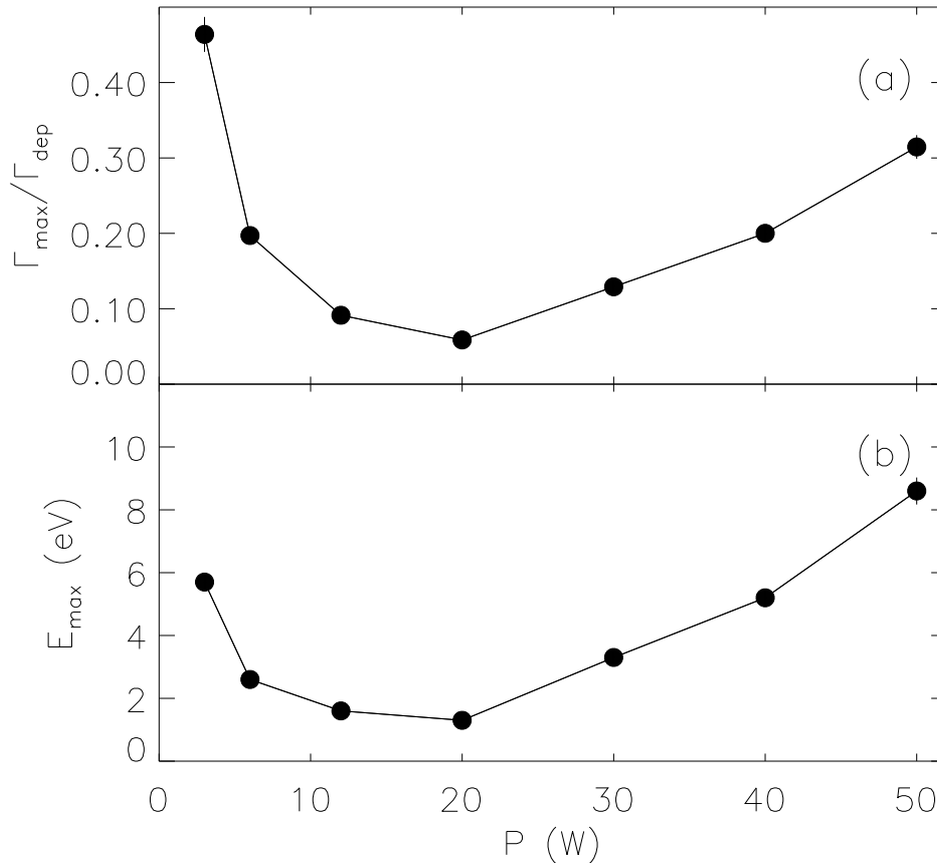


FIG. 45. The ratio of the ion flux and the deposition flux  $R_i$  (a) and the ratio of the energy flux and the deposition flux  $E_{max}$  (b) as a function of power delivered to the  $\text{SiH}_4/\text{Ar}$  discharge at an excitation frequency of 50 MHz and a pressure of 0.4 mbar. (Compiled from E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

properties, e.g. a low defect density and a high carrier mobility. The improvement of electrical properties of the deposited films at high RF power densities has also been reported by Nishikawa *et al.* [361].

The influence of power variation on the material properties is in agreement with the trends observed by Andújar *et al.* [246], who studied the  $\alpha$ - $\gamma'$  transition in pure silane discharges at 13.56 MHz. Further, this has also been observed for pure silane discharges at 50 MHz have been observed by Meiling *et al.* [423].

The  $\alpha$ - $\gamma'$  transition can also be induced by changing the process pressure. At low pressures the discharge is in the  $\alpha$  regime. At a certain pressure it changes to the  $\gamma'$  regime by increasing the pressure with only 0.02-0.04 mbar. The experiments presented here span a pressure range from 0.08 to 0.5 mbar, and, as above, plasma properties that are deduced from IED measurements are compared to material properties. The other process conditions are an excitation frequency of 50 MHz, a power level of 10 W, gas flows of 30 sccm  $\text{SiH}_4$  and 30 sccm  $\text{H}_2$  and a substrate temperature of 250 °C. Data are summarized in Fig. 46 and Fig. 47. The self-bias voltage decreases quite rapidly in magnitude with increasing pressure, as shown in Fig. 46a. The plasma potential slowly increases with pressure up to a pressure of 0.20 mbar and decreases towards the highest pressures (Fig. 46a). As both DC self bias and plasma potential decrease, also

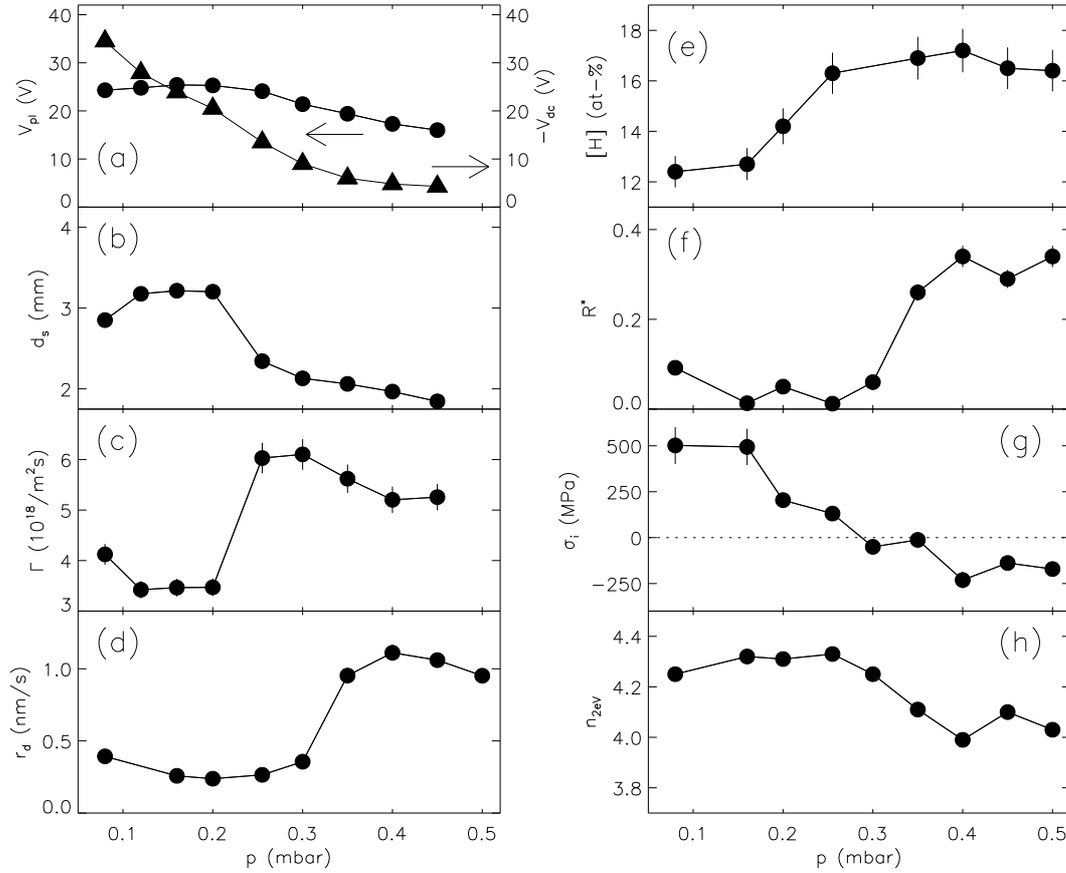


FIG. 46. Plasma parameters as deduced from the IEDs and material properties as a function of process pressure of a  $\text{SiH}_4/\text{H}_2$  discharge at an excitation frequency of 50 MHz and a power of 10 W: (a) the plasma potential  $V_{pl}$  (circles) and DC self bias  $V_{DC}$  (triangles), (b) the sheath thickness  $d_s$ , (c) the maximum ion flux  $\Gamma_{max}$ , (d) the growth rate  $r_d$ , (e) the hydrogen content, (f) the microstructure parameter  $R^*$ , (g) the internal stress  $\sigma$ , and (h) the refractive index  $n_{2eV}$ . (Compiled from E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

$V_{rf}$  will decrease [134]. Such a decrease has been measured by others [184,245]. The sheath thickness falls from around 3 to 2 mm around 0.2 mbar (Fig. 46b). The ion flux (Fig. 46c) is rather constant in the region up to 0.2 mbar, whereas it is nearly doubled at the high pressure side. A slight decrease of the ion flux in the  $\alpha$  regime has been observed in 13.56 MHz discharges by Roca i Cabarrocas [403]. The growth rate increases with about a factor of 4 upon a pressure change from 0.2 to 0.3 mbar (Fig. 46d). The fraction  $R_i$  is shown in Fig. 47a to be around 0.25 in the  $\alpha$  regime, whereas it drops to about 0.10 at pressures higher than 0.3 mbar, due to the fact that the deposition rate increases a factor of 2.5 more than the ion flux. Since the plasma potential decreases with increasing pressure, the ions arriving at the film surface are expected to have a lower energy per ion in the  $\gamma'$  regime than in the  $\alpha$  regime. This is clearly seen in Fig. 47b, where  $E_{max}$  changes from about 5 eV at pressures below 0.3 mbar to values of about 2 eV at the higher pressures.

As can most clearly be seen in Fig. 46d, the  $\alpha$ - $\gamma'$  transition occurs at a pressure of about 0.3 mbar for these experimental conditions. The impedance of the plasma, as

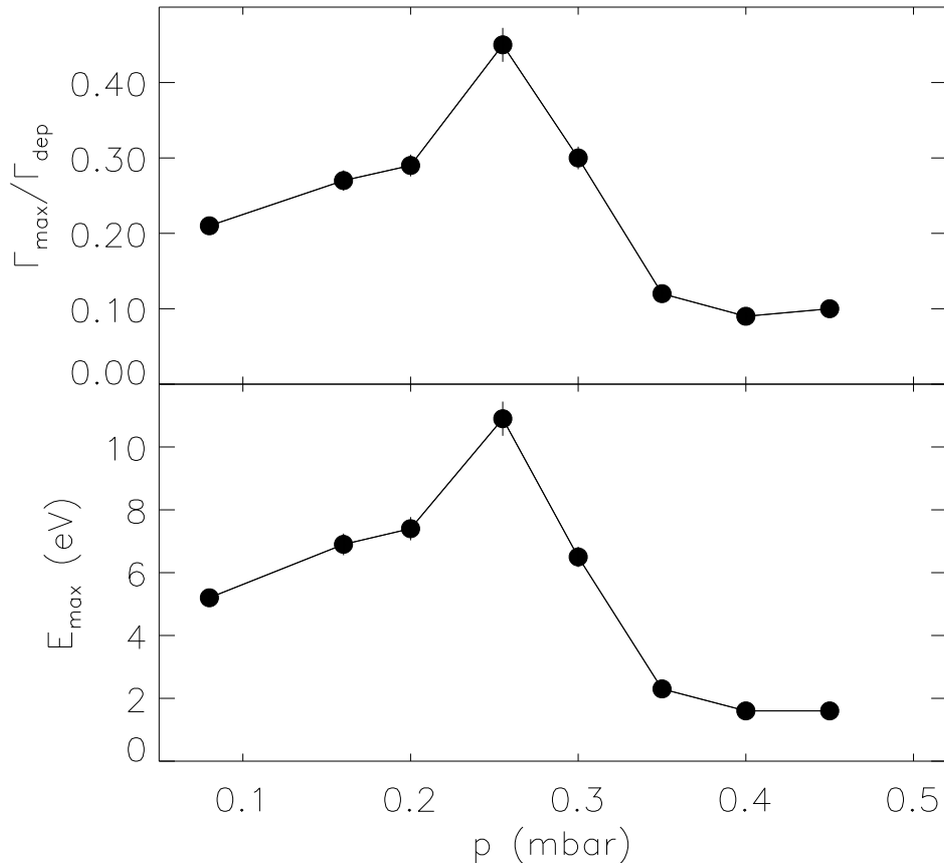


FIG. 47. The ratio of the ion flux and the deposition flux  $R_i$  (a) and the ratio of the energy flux and the deposition flux  $E_{max}$  (b) as a function of process pressure of a  $\text{SiH}_4/\text{H}_2$  discharge at an excitation frequency of 50 MHz and a power of 10 W. (Compiled from E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

well as the optical emission from the plasma change in going through the transition. The depletion of  $\text{SiH}_4$  during deposition was already shown and compared with the deposition rate in Fig. 31. The effect of the  $\alpha$ - $\gamma'$  transition on the partial pressures of disilane and trisilane was already presented in Fig. 32, see Sec. VC 1. The amount of silicon in these higher mass neutral species is only about five percent of the total amount of silicon in  $\text{SiH}_4$ , as can be concluded from the measurements of the partial pressures of these gases. The transition however effects the production of these higher silanes. The increase of the partial pressures is larger than the increase in depletion. This means that the amount of these higher silanes produced per consumed quantity of silane increases at the higher pressures.

The material properties are also effected when going through the  $\alpha$ - $\gamma'$  transition. The hydrogen content is about 12% up to 0.2 mbar, and increases to over 16% at higher pressures (Fig. 46e). The increase in the hydrogen content occurs at a lower pressure than the increase in deposition rate, much like the behaviour of the ion flux. The microstructure parameter is lower than 0.1 up to a pressure of 0.3 mbar, and is around 0.3 above this pressure (Fig. 46f). The internal stress in the layers is about 500 MPa compressive at low pressures and changes to a tensile stress around 0.3 mbar, and remains tensile at 200 MPa at higher pressures (Fig. 46g). The refractive index is

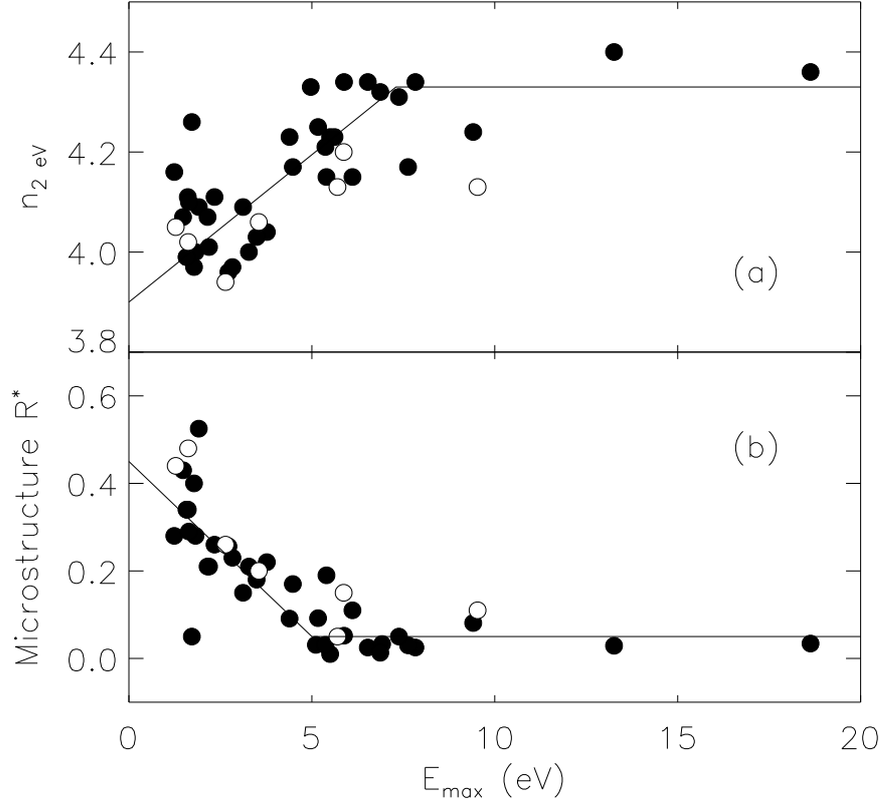


FIG. 48. Material properties as function of  $E_{\max}$  at 250°C: (a) refractive index  $n_{2\text{eV}}$ , and (b) microstructure parameter  $R^*$ . The closed circles represent pure silane and silane-hydrogen plasmas. The open circles refer to silane-argon plasmas. Lines are to guide the eye. (Adapted from E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

around 4.30 up to a pressure of 0.3 mbar, but decreases towards the highest pressures (Fig. 46h). In summary, the material becomes of poorer quality at higher pressures. The  $\gamma'$  regime is associated with the formation of particulates in the plasma [322]. However, these particles are assumed not to be incorporated in the film and thus not to be the direct cause of the different material properties.

In the above presented range of pressure variation, Hamers [163] also has studied the influence of the substrate temperature on the plasma and the material. It was found that in the temperature range of 200 to 300°C the trends of the bias voltage, the plasma potential and the growth rate as function of pressure all are the same, while the absolute magnitude depends on the temperature. The trends in material properties are similar to the ones reported above: at a temperature of 200°C the material quality is worse than at higher temperatures. The  $\alpha$ - $\gamma'$  transition occurs at a lower pressure than at a temperature of 250°C. This has been observed before [248].

In all the experiments reported here it is observed that the value of  $R_i$  is rather constant at a value of 0.25 in the  $\alpha$  regime, whereas in the  $\gamma'$  regime the value of  $R_i$  typically amounts to 0.10. In other words, if all the Si atoms that originate from ions contribute to the deposition, the contribution of ions to the deposition in the  $\alpha$  regime is 25%, and in the  $\gamma'$  regime 10%. The observation that these values are rather constant in each regime strongly indicates that the deposition rate is limited by the ion

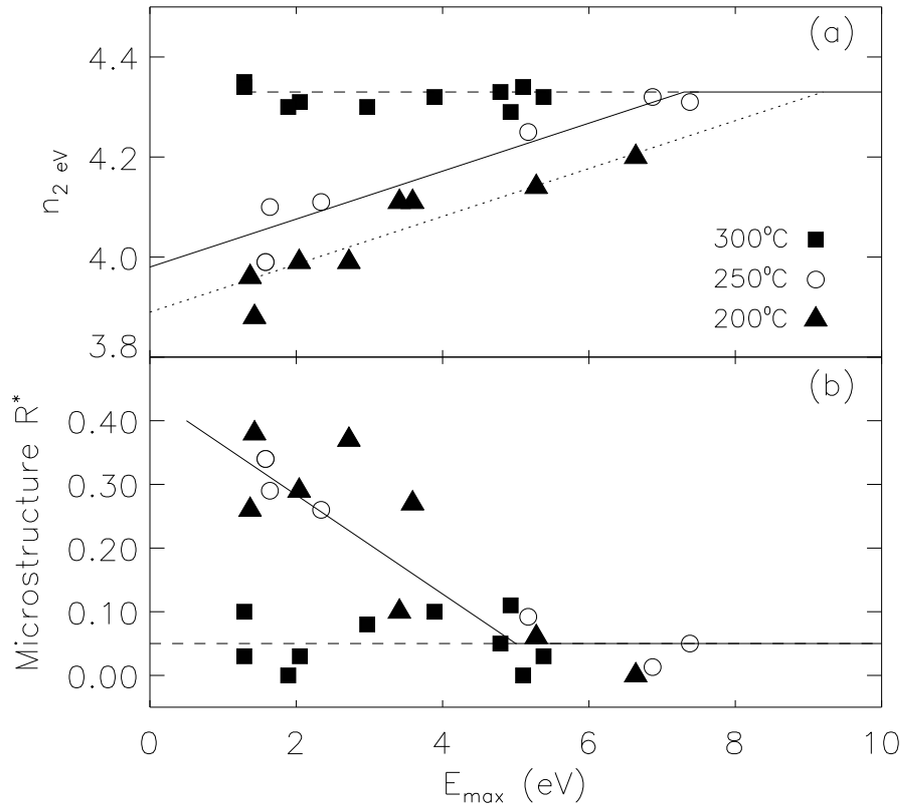


FIG. 49. Material properties as function of  $E_{max}$  at three temperatures: (a) refractive index  $n_{2eV}$ , and (b) microstructure parameter  $R^*$ . Lines are to guide the eyes. (Adapted from E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

flux.

In an attempt to relate ion bombardment with material structure it is very illustrative to correlate the refractive index  $n_{2eV}$  and the microstructure parameter  $R^*$  with the kinetic ion energy per deposited atom  $E_{max}$ . The data presented above for various discharges (pure silane, silane/argon, and silane/hydrogen) at a temperature of 250°C are thus summarized in Fig. 48. It is very clear that the structural properties of the layers are poor (small  $n_{2eV}$  and large  $R^*$ ) for small  $E_{max}$ . The structural properties improve rapidly with increasing  $E_{max}$  up to a value of about 5 eV. All samples with  $E_{max}$  below 5 eV are deposited in the  $\gamma'$  regime. Above 5 eV a dense network with only a small fraction of Si-H<sub>2</sub> bonds, is produced. The structural properties do not change further with increasing  $E_{max}$ .

The role of the substrate temperature can be inferred from a plot of  $n_{2eV}$  and  $R^*$  versus  $E_{max}$  at the three temperatures mentioned above, i.e., 200, 250, and 300°C, see Fig. 49. At a substrate temperature of 200°C the refractive index is lower at every  $E_{max}$  than at a substrate temperature of 250°C. Further, the threshold at which dense material is obtained, is observed to be a few eV higher than at 250°C. The refractive index at 300°C is high and independent of  $E_{max}$ . The microstructure parameter  $R^*$  as a function of  $E_{max}$  behaves similarly for material deposited at 200 and at 250°C. At 300°C the value of  $R^*$  is less than 0.1 and independent of  $E_{max}$ . It is noteworthy to show the relation between the internal stress and  $E_{max}$ , as a function of temperature

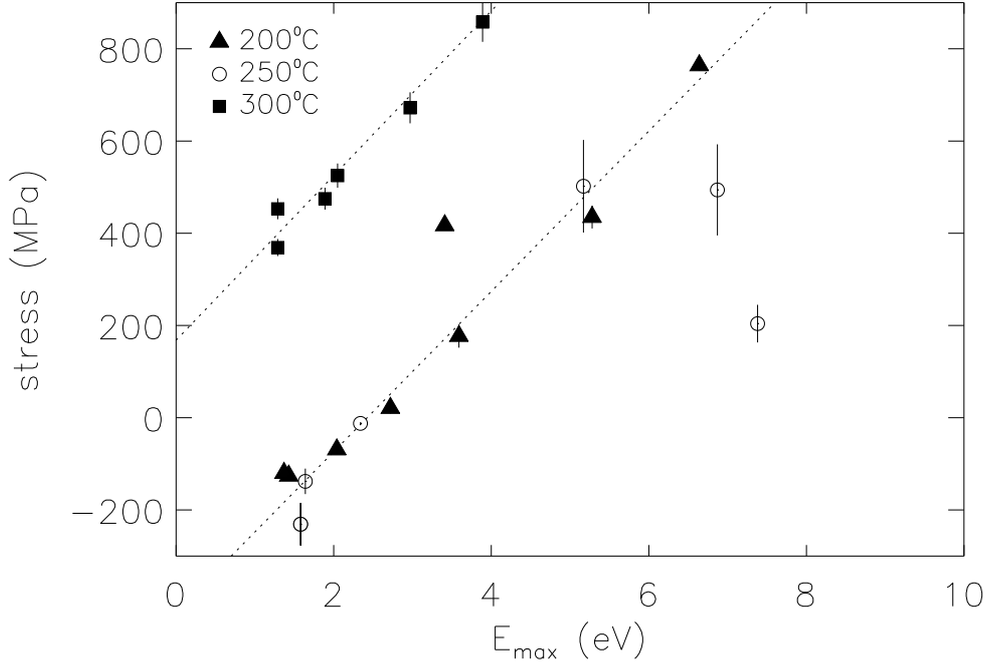


FIG. 50. Internal stress as a function of  $E_{max}$  at three temperatures, for the pressure series (46) in the  $\text{SiH}_4/\text{H}_2$  discharge running at an excitation frequency of 50 MHz and a power of 10 W. (From E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

(Fig 50). The stress is linearly dependent on  $E_{max}$  (between 1 and 7 eV) with a slope of about 175 MPa/eV for material deposited at 200 and at 250°C. For the 300°C data series this relation is shifted upwards by about 600 MPa, and at values of the stress larger than about 1000 MPa ( $E_{max}$  was about 4.5 eV) the deposited layers started to peel off from the silicon substrate directly after exposure to air. In summary, the ion bombardment clearly is needed to create or promote a dense amorphous network, at temperatures up to 250°C. At higher temperatures energetic ion bombardment results in too high intrinsic stress.

#### VI.B.4. Ion-surface interactions

The different processes in which the ions might well be involved at the surface can be indentified on the basis of their chemical reactivity at the surface and the amount of energy they transport towards the surface.

It is assumed that  $\text{SiH}_3$  radicals are responsible for the production of deposition sites at the hydrogen-covered surface, when no ions are present [138]. If ions contribute to the deposition process, a higher deposition rate has been reported, see e.g. [419]. The fraction of arriving ions per deposited silicon atom varies between 0.1 and 0.3, depending on the discharge regime, see above. Assuming equal sticking probabilities of ions and radicals (0.3 for the  $\text{SiH}_3$  radical [192]), up to about 10% of the deposition is attributed directly to the ions. This value has also been found by Kae-Nune *et al.* [311]. On the other hand, ions can penetrate into the sub-surface layers and hence,

their sticking probability will be closer to one. In addition, direct incorporation of ions in the amorphous network can only explain partly the increase in deposition rate. The creation of deposition sites by the ions possibly is of greater importance. It has been suggested that a direct relation between the ion flux and the deposition rate exists, if the availability of deposition sites is the limiting factor in the deposition process [236,400,424]. Also, partial dissociation of a polyatomic ion upon impact may occur [412,425–427]. The dissociation of an hydrogen containing ion may lead to the production of more than one hydrogen atom per ion. These atoms create molecular hydrogen and dangling bonds, upon reaction with hydrogen bonded to the surface. The probability  $\gamma$  of the recombination process of atomic hydrogen has been estimated to be between 0.4 and 1 [317].

In discharges operating in the  $\gamma'$  regime large ions consisting of several ions are present, in contrast to discharges in the  $\alpha$  regime. In both regimes the same relative amount of ions contributes to the deposition, as is deduced from the measured amount of radicals that contribute to the deposition, which is regime-independent [317]. These larger ions in the  $\gamma'$  regime are thought to create more deposition sites per arriving ion, compared to the smaller ions in the  $\alpha$  regime [163]. This results in a lower number of ions per deposited atom in the  $\gamma'$  regime.

The kinetic ion energy flux,  $(\epsilon \Gamma)_{max}$ , which is typically  $20 \text{ Wm}^{-2}$  [163,301], will raise the substrate temperature by only a few degrees. Therefore, the influence of ions will be limited to the location of impact. Furthermore, typical ion energies are below the sputtering threshold of silicon [134].

Enhancement of surface diffusion of the growth precursors is considered as one of the beneficial effects of ion bombardment [246,428]. The potential energy of ions, which is released when the ion is neutralized, is typically 10 eV. This energy can be a substantial fraction of the total energy transferred. The release of this ionization energy is sufficient to excite atoms into excited electronic states, thereby weakening their bonds and enhancing their mobilities [429].

The influence of ion bombardment on the structure of deposited materials has been studied by Müller [428,430] by performing Molecular Dynamics calculations on ion assisted vapor phase growth of nickel. Nickel, of course, is very different from hydrogenated amorphous silicon, but the resemblance of both deposition processes and some material properties is remarkable. In both cases, the growth precursors stick to the surface or diffuse along the surface, and the material can have a columnar-like structure or a structure with voids. Ion bombardment during the deposition of Ni promotes the formation of a dense film. The ion energies and fluxes in the Molecular Dynamics studies were similar to those in typical *a*-Si:H deposition conditions. Two important processes modify the structure of the nickel film, i.e., forward sputtering, and diffusion enhancement of surface species. Similar processes are likely to occur in *a*-Si:H deposition. In the deposition of amorphous germanium by ion assisted electron-beam evaporation, it was also found that a kinetic energy of about 5 eV per deposited atom is needed to produce a dense amorphous germanium network, irrespective of the amount of kinetic energy of the ions: the effect of 5 ions with 20 eV was the same as of 1 ion of 100 eV [431]. These two additional examples clearly indicate the beneficial role that ion bombardment apparently has on the density of the deposited film.

In the formation of a hydrogenated amorphous silicon network, the presence of the bonded hydrogen must be taken into account, as it was shown above that both the

hydrogen content and the microstructure parameter are influenced by the ion bombardment. The high value of the microstructure parameter and the concomitant low density of the material for low values of  $E_{max}$  at 250°C (Fig. 48) indicate that the cross-linking process is not sufficiently fast to create a dense network. This cross-linking process is activated by the locally released energy of chemisorption [432,433], and involves the temperature activated evolution of H<sub>2</sub> from the surface. Hydrogen desorption due to cross-linking, which is known to occur from thermal desorption measurements [434], starts to become important at temperatures higher than 300°C. At this temperature a drastic change in the importance of ion bombardment for achieving a dense network was observed (see Fig. 49). At lower deposition temperatures the cross-linking process during the deposition is temperature limited and the kinetic energy of the ions is needed to achieve the cross-linking process. If the amount of ion kinetic energy is insufficient, cross-linking will be incomplete and will result in an open network structure with a large amount of Si-H<sub>2</sub> bonds. At larger values of the ion kinetic energy the cross-linking is stimulated and a dense network is formed. Ion bombardment rather than the temperature induces cross-linking, as the microstructure parameter, the stress, and (to a lesser extent) the refractive index, have the same dependence on the  $E_{max}$  at both 200 and 250°C. At 300°C the thermally activated cross-linking process is becoming important.

Tensile stress can be related to the presence of voids in the material. The large microstructure parameter at low  $E_{max}$  values is indicative of voids. A kinetic energy of the ions of typically 20 eV is enough to implant part of these ions a few monolayers below the surface of the growing film [400]. The extra Si atoms deposited in the layer at this depth will cause a compressive stress. Light hydrogen ions, which are less abundant than e.g. SiH<sub>3</sub><sup>+</sup> ions, penetrate deeper into the material (see also TRIM simulation results in Fig. 42). The stress in the films deposited at 200 and 250° depends similarly on  $E_{max}$ . At these temperatures the ion energy is needed to densify and to cross-link the network. At a temperature of 300° the stress is much higher, which is explained by the fact that the ions reach an already dense network and some will be implanted, resulting in excessive stress in the material.

## VII. DEPOSITION MODELS

A conceptual view of the processes that occur in PECVD of  $a$ -Si:H has been given by Perrin [173], and is shown in a slightly adapted form in Fig. 51. Primary gas phase reactions are electron-impact excitation, dissociation, and ionization of the source gas molecules ( $\text{SiH}_4$ ), thereby producing radicals, and positive and negative ions. Secondary reactions in the gas phase between (charged) molecules and radicals produce other species. Diffusion of reactive neutral species leads to material deposition. Positive ions are accelerated to the substrate, and bombard the growing film. Negative ions are confined in the bulk of the discharge, which can lead to particulate formation. On the surface species diffuse to growth sites, and contribute to the film after sticking to the underlying material. Also, surface species recombine with other species and desorb from the surface (see also Fig. 14). Sub-surface reactions are the release of hydrogen from this hydrogen-rich layer and the relaxation of the silicon network.

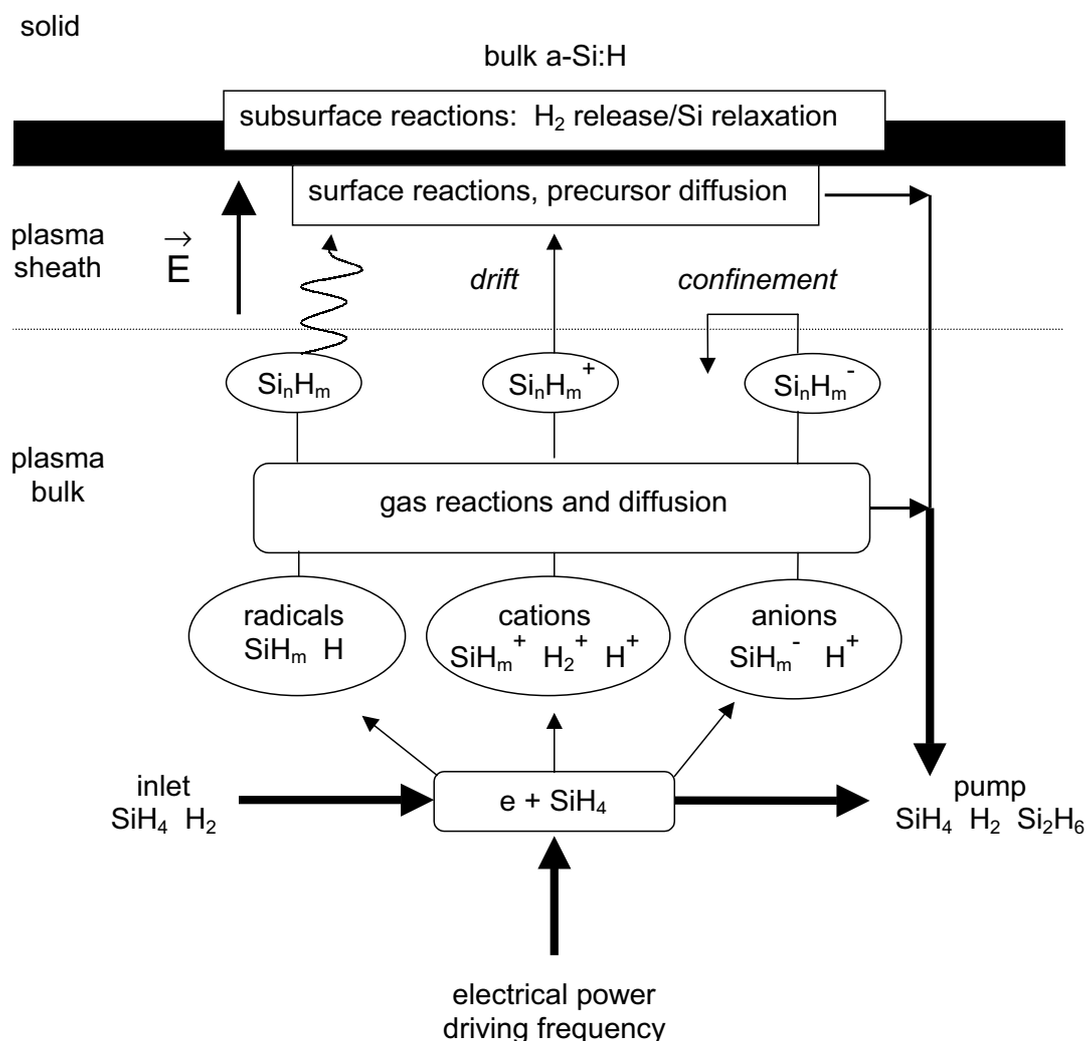


FIG. 51. Schematic representation of the processes occurring in a  $\text{SiH}_4/\text{H}_2$  discharge and the various particles present in the energy and material balances. (After J. Perrin, in *Plasma Deposition of Amorphous Silicon-Based Materials*, edited by G. Bruno, P. Capezzuto, and A. Madan (Academic Press, Boston, MA, U.S.A., 1995), Chap. 4, p. 177.).

Nowadays, it is generally accepted that 'device quality' *a*-Si:H is obtained under PECVD conditions where the SiH<sub>3</sub> radical is the predominant growth precursor [123,137,192]. At the typical deposition temperatures and pressures the *a*-Si:H surface is almost completely terminated by hydrogen atoms. The SiH<sub>3</sub> radical is thought to physisorb on this hydrogen terminated surface, before it is incorporated into the film [136,137,317,435–438]. The excess hydrogen is eliminated from the hydrogen-rich (40–50 %) sub-surface, so as to become 'bulk' *a*-Si:H, with a hydrogen content of about 10 %.

Robertson has summarized the three recent classes of models of *a*-Si:H deposition [439]. In the first one, proposed by Ganguly and Matsuda [399,440], the adsorbed SiH<sub>3</sub> radical reacts with the hydrogen-terminated silicon surface by abstraction or addition, which creates and removes dangling bonds. They further argue that these reactions determine the bulk dangling bond density, as the surface dangling bonds are buried by deposition of subsequent layers to become bulk defects.

The second class of models was formulated by Winer [441] and Street [442,443]. Here the notion that hydrogen atoms are more mobile than silicon atoms forms the basis of the model. The silicon network is fixed up to a temperature near the crystallization temperature (650 °C). Hydrogen is mobile above a temperature of about 200 °C and the motion of hydrogen allows for the interconversion of defects, weak bonds, and strong bonds. These processes occur in the sub-surface layer, and surface processes are not explicitly included. In this model, the optimum deposition temperature is the one at which hydrogen atoms can diffuse one atomic spacing as to remove the weak bonds. The weak bond distribution can only be changed irreversibly after deposition, while the defect distribution can be changed reversibly, as described by the defect pool models [444–446].

In the third class of models, computer simulations try to fully incorporate all processes in the discharge, the interaction of created species in the discharge with the wall (i.e., the substrate), and the network formation [190,191,232,447–449]. These models to date do not treat the formation of disorder or defects, but aim at the understanding of the deposition rate, hydrogen content, and other macroscopic properties in relation to the discharge conditions (see also Sec. IV).

Robertson has combined and extended the first and second class of models [439], by focussing on the origin of weak bonds. A surface adsorption model is used to describe surface coverage of the silyl radical SiH<sub>3</sub>. Then, the processes that cause hydrogen to be expelled from the sub-surface layer lead to the formation of weak bonds, which are frozen in. The dangling bond defects arise from these weak bonds by the defect pool process. In the following, a description is given of the Robertson deposition model (details can be found in [439,450]).

### VII.A. Surface adsorption

On the hydrogen-terminated surface, the surface species are mainly  $\equiv\text{SiH}$ , but at lower temperature ( $< 200$  °C)  $=\text{SiH}_2$  and  $-\text{SiH}_3$  become increasingly important. The SiH<sub>2</sub> and SiH<sub>3</sub> radicals react differently with this surface. The SiH<sub>2</sub> radical can insert directly into Si-H or Si-Si bonds. The sticking coefficient is close to 1, which leads to columnar growth (physical vapour deposition [451]). Also, the SiH<sub>2</sub> can react with SiH<sub>4</sub>

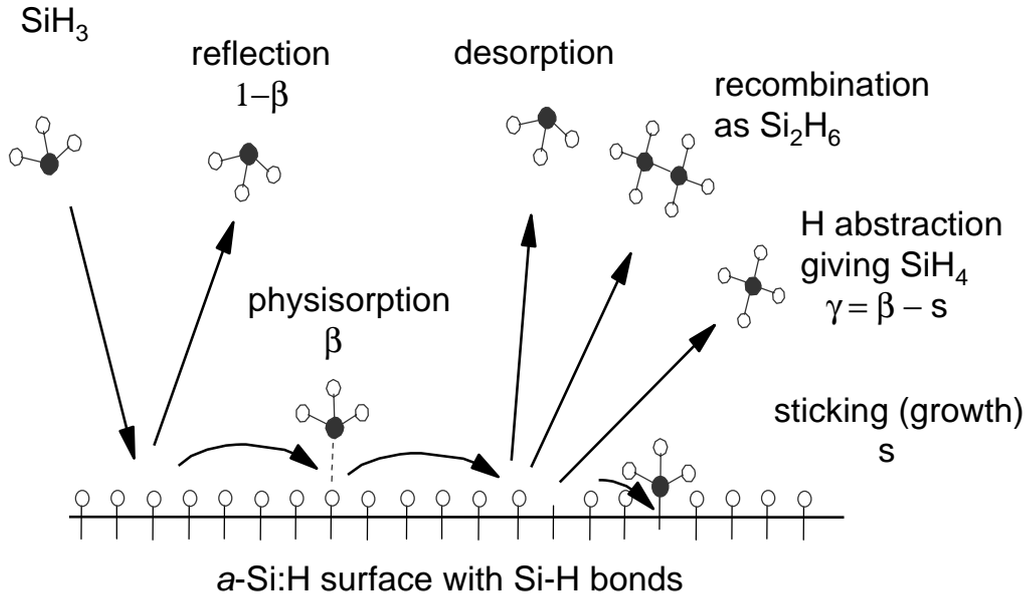


FIG. 52. Schematic representation of the processes that a  $\text{SiH}_3$  may undergo at a hydrogen-terminated  $a\text{-Si:H}$  surface. (After J. Robertson, *J. Appl. Phys.* **87**, 2608 (2000).)

TABLE 8. Reactions of surface sites  $\equiv\text{SiH}$ ,  $\equiv\text{Si}\cdot$ , and  $\equiv\text{SiHSiH}_3$  with the radical  $\cdot\text{SiH}_3$ , with corresponding rate coefficients.

acronym	reaction	rate coefficient
R1	$\equiv\text{SiH} + \cdot\text{SiH}_3 \rightarrow \equiv\text{SiHSiH}_3$	$s_1$
R2	$\equiv\text{SiHSiH}_3 \rightarrow \equiv\text{SiH} + \cdot\text{SiH}_3$	$\nu_d$
R3	$\equiv\text{SiHSiH}_3 + \equiv\text{SiH} \rightarrow \equiv\text{SiH} + \equiv\text{SiHSiH}_3$	$\nu_h$
R4	$\equiv\text{SiH} + \cdot\text{SiH}_3 \rightarrow \equiv\text{Si}\cdot + \text{SiH}_4$	$\nu_a$
R5	$\equiv\text{Si}\cdot + \cdot\text{SiH}_3 \rightarrow \equiv\text{Si-SiH}_3$	$s_0$
R6	$2 \equiv\text{SiHSiH}_3 \rightarrow 2 \equiv\text{SiH} + \text{Si}_2\text{H}_6$	$\nu_r$
R7	$2 \equiv\text{SiH} \rightarrow 2 \equiv\text{Si}\cdot + \text{H}_2$	$\nu_b$

to form  $\text{Si}_2\text{H}_6$ . The  $\text{SiH}_3$  radical cannot react directly with the surface. It has a small sticking coefficient, and it diffuses over the surface to find a reactive site. This reactive site has been suggested to be a dangling bond at kink and step edges [452]. As a result, this leads to conformal coverage (chemical vapour deposition regime [451]). The  $\text{SiH}_3$  radical is physisorbed onto the surface [436]. A contrasting view has been proposed by Von Keudell and Abelson [453], who conclude from their experiments that  $\text{SiH}_3$  can insert into strained Si-Si bonds on the surface to form a five-fold coordinated bond. This was also observed by Ramalingam *et al.* [454] in molecular-dynamics simulations of  $\text{SiH}_3$  impinging on a  $(2 \times 1)$  Si(100) reconstructed surface. Nevertheless, as these observations do not as yet explain film deposition, they are considered at least questionable.

The possible reactions with a  $\equiv\text{SiH}$  surface site are schematically shown in Fig. 52, and summarized together with their reaction rates in Tab. 8. As a simplification, the

surface site is  $\equiv\text{SiH}$ , the dangling bond is  $\equiv\text{Si}\cdot$ , and the adsorbed  $\text{SiH}_3$  radical is  $\equiv\text{SiHSiH}_3$ . The possible reactions include adsorption (R1 in Tab. 8), desorption (R2), hopping to an adjacent  $\equiv\text{SiH}$  site (R3), abstraction of an hydrogen atom from  $\equiv\text{SiH}$  to create  $\equiv\text{Si}\cdot$  (R4), addition to  $\equiv\text{Si}\cdot$  (R5), surface recombination with another  $\text{SiH}_3$  radical to form a desorbing  $\text{Si}_2\text{H}_6$  molecule (R6), and elimination of hydrogen molecules from two adjacent  $\equiv\text{SiH}$  sites to yield surface dangling bonds (R7).

Several simplifications are further made [439]: 1) the  $\text{SiH}_3$  adsorption (R1) is temperature independent, and  $s_1$  is constant, 2) the  $\text{SiH}_3$  addition to a dangling bond is temperature independent, and  $s_0 = 1$ , however, the  $\text{SiH}_3$  must hop to this dangling bond site, from a neighbouring site, so the net rate is  $v_h s_0$ , 3) the abstraction reaction (R4) is faster than the desorption reaction (R2), so the latter is ignored, and 4) the recombination reaction R6 is ignored.

Perrin has defined the fractional coverages of the surface sites as  $\theta_1$ ,  $\theta_0$ , and  $\theta_4$ , for  $\equiv\text{SiH}$ ,  $\equiv\text{Si}$ , and  $\equiv\text{SiHSiH}_3$ , respectively [317], with  $\theta_1 + \theta_0 + \theta_4 = 1$ . These three fractional coverages can be found by expressing them as rates in terms of creation and destruction. This gives, with  $\phi$  the incident  $\text{SiH}_3$  flux density per surface site [439]:

$$\frac{d\theta_0}{dt} = \nu_0\theta_4(1 - \theta_0) - \nu_4\theta_4\theta_0 + \nu_b(1 - \theta_0) \quad (61)$$

$$\frac{d\theta_4}{dt} = \phi - \nu_a\theta_4(1 - \theta_0) - \nu_h\theta_4\theta_0 \quad (62)$$

In the steady state, and ignoring multiple hopping the coverages are approximated to be:

$$\theta_0 \approx \frac{\nu_a}{\nu_h} + \frac{1}{1 + \frac{\phi}{\nu_b}} \quad (63)$$

$$\theta_4 \approx \frac{\phi}{2\nu_a \left(1 + \frac{\nu_b}{\phi}\right)} \quad (64)$$

At temperatures lower than 350-400 °C the expressions for  $\theta_0$  and  $\theta_4$  reduce to  $\theta_0 \approx \nu_a/\nu_h$  and  $\theta_4 \approx \phi/2\nu_a$ . The residence time of  $\text{SiH}_3$  radicals at the surface  $\tau_4$  is given by the coverage  $\theta_4$  divided by the arrival flux  $\phi$ :  $\tau_4 \approx 1/2\nu_a$ . The  $\text{SiH}_3$  surface diffusion length  $L = \sqrt{D\tau_4}$ , where  $D = \nu_h a^2$  is the  $\text{SiH}_3$  diffusion coefficient, with  $a$  the Si-Si bond length. We thus have  $L = a\sqrt{\nu_h/2\nu_a} = a/\sqrt{\theta_0}$ .

The reaction rates  $\nu_i$  ( $i = h, a, b$ ) for each reaction are thermally activated as  $\nu_i = \nu_{i0} \exp(-E_i/k_B T)$ . Robertson has critically assessed the values  $\nu_{i0}$  and  $E_i$  for each reaction [439], and used for the model  $\nu_{h0} = 10^{13} \text{ s}^{-1}$ ,  $E_h = 0.2 \text{ eV}$ ,  $\nu_{a0} = 3 \times 10^{11} \text{ s}^{-1}$ ,  $E_a = 0.4 \text{ eV}$ ,  $\nu_{b0} = 10^{13} \text{ s}^{-1}$ , and  $E_b = 2.2 \text{ eV}$ . These values give a surface dangling bond density that increases monotonically with temperature from about  $10^{-5}$  at room temperature to about  $10^{-3}$  at 400 °C. Thermal desorption of hydrogen then sets in, and the surface dangling bond density is suddenly increased and approaches unity around 600 °C. This increase in  $\theta_0$  is due to the fact that  $E_a > E_h$ . The critical parameter is the difference  $E_a - E_h$ , not their absolute values. At typical deposition temperatures of 200-250 °C the surface dangling bond density is about  $3 \times 10^{-4}$ , which is two orders of magnitude larger than the bulk dangling bond density. The  $\text{SiH}_3$  surface diffusion length decreases from about 30 nm at room temperature to about 8 nm at 250 °C.

These findings from Robertson [439,450] are in contrast to the ones from the model proposed by Ganguly and Matsuda [399,440]. They found a minimum in the surface

dangling bond density, and a maximum in the surface diffusion coefficient, simultaneously occurring at about 250 °C ( $\theta_0 \approx 10^{-5}$ ,  $L \approx 400$  nm). This would explain the optimum in material quality around 250 °C. Surface defect densities have been measured during deposition by *in situ* electron spin resonance by Yamasaki *et al.* [455]. They found that the surface defect density was much higher than the bulk defect density. Therefore, the concept of surface dangling bonds that are buried by deposition of subsequent layers is questionable. Robertson argues that as from the hydrogen rich *a*-Si:H surface layer most of the hydrogen is eliminated about half to the bonds is changed, from Si-H to Si-Si. It would therefore be very unlikely that this enormous change in bonding configurations would leave a surface dangling bond density of  $10^{-6}$  unchanged [450].

### VII.B. Solubility of hydrogen in silicon

The elimination of excess hydrogen from the surface layer seems counterintuitive, the hydrogen leaves the *a*-Si:H network towards the surface, where the hydrogen content is larger. This can be understood by considering the thermodynamics of hydrogen in silicon [439]. In general, the Gibbs free energy of a mixture of two components *A* and *B*, with mole fractions  $x_A$  and  $x_B$  (with  $x_A + x_B = 1$ ) is given as [456]:

$$G = x_A G_A + x_B G_B + x_A x_B \Omega_{AB} + RT(x_A \ln x_A + x_B \ln x_B) \quad (65)$$

with  $G_A$  and  $G_B$  the Gibbs free energies of the pure species,  $\Omega_{AB}$  the heat of mixing, and  $RT(x_A \ln x_A + x_B \ln x_B)$  the entropy of mixing. The chemical potential  $\mu_A$  is given by:

$$\mu_A = \left( \frac{\partial G}{\partial x_A} \right)_{T,B} = G_A + \Omega_{AB}(1 - x_A)^2 + RT \ln x_A \quad (66)$$

and similarly for  $\mu_B$ . If  $\Omega_{AB}$  is negative, mixing is favoured. If  $\Omega_{AB}$  is positive, two stable mixtures (*A*-rich and *B*-rich) exist with a miscibility gap in between. Any mixture with a composition between the two stable phases will decrease its free energy by decomposing into an *A*-rich and a *B*-rich phase. Such a process is known as spinoidal decomposition, and in this process species diffuse up a concentration gradient [456].

In the case of *a*-Si:H, it turns out that  $\Omega_{AB}$  is positive [439]. This can be inferred from the bond energies of Si-Si (2.35 eV), Si-H (3.3 eV), and H-H (4.5 eV). A Si-H network is thus unstable compared to pure Si and H<sub>2</sub>. A mixture of Si-Si and Si-H bonds will be driven towards their two stable mixture phases, which leads to the solubility limit of H in Si. Acco *et al.* [69,70] have determined this limit to be about 4%. Excess hydrogen in *a*-Si:H forms SiH<sub>2</sub> or SiH around microvoids.

### VII.C. Elimination of hydrogen from *a*-Si:H

Successive burial of hydrogen-rich surface layers leads to the formation of the *a*-Si:H material. The large amount of hydrogen at the surface is to saturate the surface dangling bonds. The much lower hydrogen content in the bulk is due to the solubility limit of hydrogen in silicon.

The hydrogen content in the bulk  $a$ -Si:H depends on deposition temperature and plasma conditions, e.g. [375,457] (see also Sec. VIII A). In the  $\gamma'$ -regime the hydrogen content increases considerably with decreasing deposition temperature, while in the  $\alpha$ -regime the hydrogen content also increases with decreasing temperature, but this increase sets in at lower temperatures. Robertson has defined the hydrogen elimination rate  $R_H$ , expressed in the number of hydrogen atoms per unit cell, as the sum of a thermal  $R_T$  and an athermal, plasma-driven  $R_P$  elimination process [450]:  $R_H = R_T + R_P$ . Also,  $R_H$  equals the growth rate  $r_d$  multiplied by the fractional loss of hydrogen from the surface layer to the bulk:  $R_H = r_d(C_{H,s} - C_{H,b})$ , where  $C_{H,s}$  and  $C_{H,b}$  are the hydrogen concentration at the surface and in the bulk, respectively.

### VII.C.1. Thermal reactions

The thermal processes can be derived starting with the behaviour of hydrogen content as a function of reciprocal temperature [439,450]. There are two regimes: a high temperature regime ( $T_s > 400$  °C), characterized with an apparent activation energy of about 1.6 eV, and a low temperature regime ( $T_s < 250$  °C) with an apparent activation energy of about 0.15 eV. Various hydrogen elimination processes are summarized in Fig. 53 [439].

In the high-temperature regime it is most likely that hydrogen is eliminated to form  $H_2$  at the surface, with an activation energy of 2.1 eV (Fig. 53a). This is similar to H evolution from crystalline silicon [439,458]. At intermediate temperatures ( $250$  °C  $< T_s < 400$  °C) the elimination of hydrogen occurs via diffusion of atomic hydrogen towards the surface with an activation energy of 1.5 eV to recombine as  $H_2$ , i.e., the standard effusion process [459–461]. This is described with the hydrogen density of states (HDOS) diagram [442,443,446,459,462], see Fig. 53b. The HDOS diagram signifies the energy of a hydrogen atom in  $a$ -Si:H compared to that of a free hydrogen atom in vacuum. A hydrogen bound as Si-H has an energy of -3.3 eV, with respect to the vacuum level, while hydrogen at a Si-Si bond center (BC in Fig. 53b) has an energy of about -1 eV. In  $a$ -Si:H hydrogen is more stable at the bond center of a weak bond, which is represented as a tail in the HDOS below -1 eV. In the HDOS there exists a mobility edge ( $H_M$ ), above which the hydrogen states form a percolation path. Hydrogen diffusion occurs by excitation from  $\mu_H$  to  $H_M$ . The difference  $\mu_H - H_M$  amounts to 1.5 eV, i.e., less than 2.2 eV, but much larger than the apparent activation energy of 0.15 eV.

Robertson proposed that the local bond rearrangement close to the surface can account for this low activation energy [439]. It is assumed that hydrogen exists largely in pairs, analogous to the  $H_2^*$  in crystalline silicon [463]. This  $H_2^*$  consists of two Si-H bonds in the same direction [464,465]. It can undergo a local rearrangement with one hydrogen atom passing through two bond centre positions, at a cost of 1 eV [245]. Another local rearrangement is possible in which the H passes through the bond centre to combine with the other hydrogen atom to form the interstitial molecule  $H_2$  (Fig. 53c). The Si-Si bond centre site is the transition state of the reaction. As hydrogen is more strongly bound at a weak Si-Si bond than at a normal Si-Si bond [466,467], the energy barrier is lowered. As a consequence, the elimination of hydrogen to form  $H_2$  is eased in the presence of weak bonds.

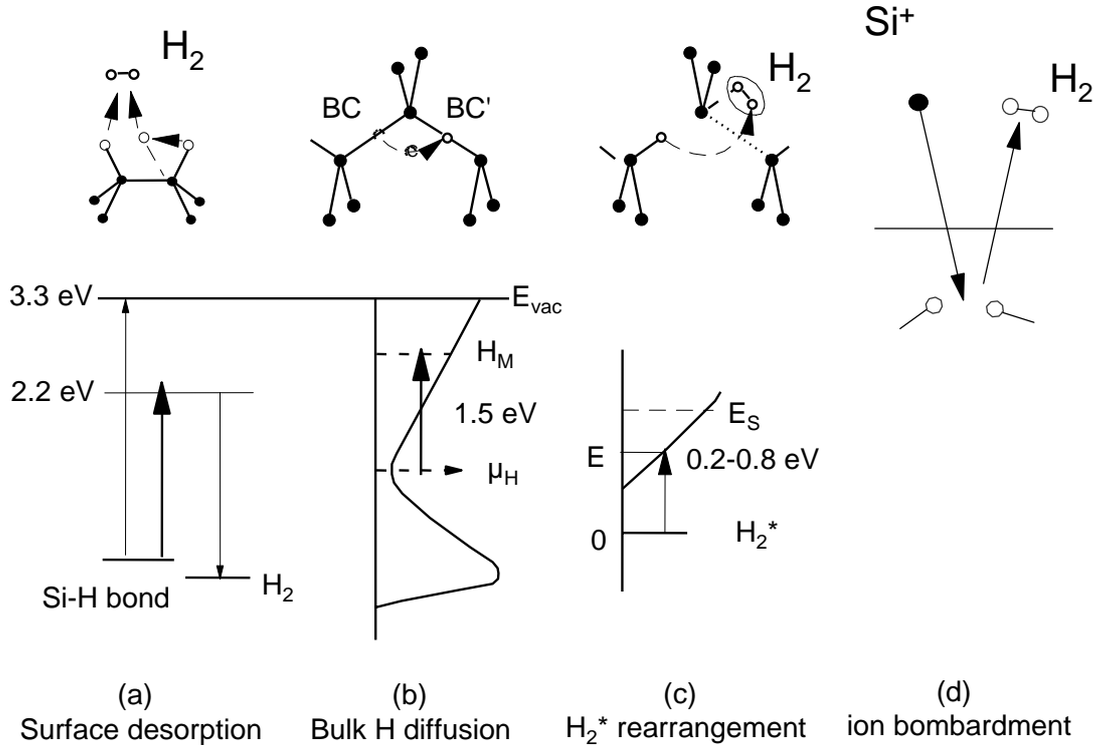


FIG. 53. Possible hydrogen elimination processes: (a) surface loss from adjacent Si-H groups by pre-pairing on a single Si-site, (b) bulk diffusion of atomic H, by activation to a H mobility level  $\mu_M$ , as described on the hydrogen density of states (HDOS), (c) rearrangement of  $H_2^*$ , and the lowering of the barrier energy below the normal value  $E_s$  by excitation to a weak bond expressed on a HDOS, (d) subsurface ion induced displacement and desorption. (After J. Robertson, *J. Appl. Phys.* **87**, 2608 (2000).)

An alternative hydrogen elimination process has been proposed by Severens *et al.* [432,468]. They argue that hydrogen can be eliminated by a cross-linking step [469] immediately after a physisorbed  $SiH_3$  radical has chemisorbed on a surface dangling bond. This cross-linking probability is thermally activated, but due to the energy released at the chemisorption of  $SiH_3$  (about 2 eV) the activation is quite low [432]. If this cross-linking process does not occur immediately after chemisorption, it is assumed that hydrogen is incorporated into the film. An activation energy of 0.15 eV is deduced from fitting hydrogen incorporation data [468].

A detailed description of the local bond rearrangement has been derived [439], using the concept of the HDOS with a low energy tail that corresponds to the H present at weak Si-Si bonds. The width of this tail is  $2E_{v0}$ , i.e., twice the width of the valence band tail in the electronic density of states, which in turn is about equal to the Urbach energy  $E_0$  [442,443]. The HDOS then is [439]:

$$N(E) = \frac{N_0}{2E_{v0}} \exp\left(\frac{E - E_s}{2E_{v0}}\right) \quad (67)$$

with  $E_s$  at the top of the HDOS, i.e., the energy barrier for rearrangement of the ideal  $H_2^*$  (1 eV). The rearrangement occurs by activation of H to a state  $E$  in the tail. The

reaction rate in atoms per unit surface area is the product of this HDOS, the Maxwell Boltzmann factor, and an attempt frequency  $\nu_1$  ( $\sim 10^3 \text{ s}^{-1}$ ), and is expressed in the low temperature regime as [439]:

$$R_T = \frac{\nu_1 k_B T}{2E_{v0}} \exp\left(-\frac{E_s}{2E_{v0}}\right) \quad (68)$$

The values obtained from this equation compare well with the experimental values of  $E_{v0}$  as a function of temperature.

### VII.C.2. Athermal reactions

Next, athermal, plasma driven, hydrogen elimination reactions are considered. Ions play an important role in the deposition. They can account for 10-25 % to the total deposition rate [301,311], depending on the regime of the discharge. Moreover, the energy that they carry to the (sub)surface [163] can be used to overcome energy barriers (see also Sec. VIB 1). Ions penetrate the film and eliminate hydrogen by displacing hydrogen atoms from subsurface Si-H bonds. They subsequently recombine as hydrogen molecules and effuse back to the surface (see Fig. 53d). The displacement yield  $Y_D$  is given by the modified Kirchin-Pease equation  $Y_D = E_i/2E_D$ , where  $E_i$  is the ion energy and  $E_D$  the displacement energy [412]. Here,  $E_D$  is equal to the Si-H bond energy, i.e., 3.3 eV. In the  $\alpha$ -regime the contribution of ions to the deposition rate is 25 % [301], and a typical average ion energy is 10 eV (see Sec. VIB 1). This gives a displacement yield  $Y_D$  of about 40 % of the deposition rate. This illustrates the beneficial role of ions in improving the material quality [284,301,358,417,470]. For the  $\gamma'$  regime about 10 % of the ions contribute to the deposition rate, and typical average ion energies are 20-30 eV [301]. This gives a displacement energy  $Y_D$  of about 30-45 % of the deposition rate. In this regime average ion energies are larger than the bulk displacement energy of silicon of 22 eV [285,376], which deteriorates the material quality.

This athermal dehydrogenation process reduces the excess weak bond density associated with a high polymeric content, rather than that ions densify the material by removing polymeric  $\text{SiH}_x$  groups. Especially low energy ions are effective at dehydrogenation due to the fact that the displacement energy  $E_d$  is low. This favours the use of e.g. Very High Frequency (VHF) PECVD reactors, where the ion energies are much lower and the ion densities much higher than at 13.56 MHz.

Further, atomic hydrogen in hydrogen-diluted discharges can contribute to abstraction of surface and subsurface hydrogen [471]. The atomic hydrogen reduces the fraction of higher  $\text{SiH}_x$  groups [466]. Only a small amount is incorporated directly [472]. Atomic hydrogen does not reduce the surface hydrogen content below 50 %, because this would result in surface dangling bonds that would be passivated by incident hydrogen [473]. Atomic hydrogen can also diffuse into the  $a$ -Si:H with an activation energy of 0.4-0.7 eV [461], which allows subsurface hydrogen abstraction. However, this would be less important at low temperatures due to the considerable activation energy.

### VII.D. Dangling bond and weak bond density

The hydrogen elimination processes described above create the distribution of weak bonds. The dangling bond distribution is formed from this by interconversion from

weak bonds to dangling bonds. For substrate temperatures above the hydrogen equilibration temperature of 200 °C, hydrogen is mobile, and the defect pool operates. This allows the weak bond and dangling bond distributions to equilibrate [439]. At temperatures below 200 °C hydrogen motion is inhibited, and equilibration should not be possible. Nevertheless, Stutzmann [474] reported that the dangling bond density as a function of temperature shows the the same behaviour as the weak bond density, for all temperatures. This suggests that a defect pool of sorts operates below 200 °C in the surface layer [439].

It is assumed that a large athermal, plasma driven, hydrogen flux is present in the hydrogen elimination zone at temperatures below 200 °C, which mediates the equilibration of the dangling bond and weak bond distribution. The dangling bond density and weak bond density thus remain linked in the elimination zone by a largely temperature-independent relationship, given by the defect pool [445,446]. The dangling bond density in the elimination layer becomes the bulk defect density as the layer is buried deeper, and becomes frozen in. This proposed process allows the weak bond density to determine the dangling bond density at all temperatures [439].

## VIII. MODIFICATIONS OF PECVD

Of the great number of possible and reported modifications, in this section only the most important ones are described, i.e., the use of higher excitation frequency (Very High Frequency, VHF), the use of gas flow modulation (known as chemical annealing or layer-by-layer deposition), and the use of RF modulation.

### VIII.A. VHF

#### *VIII.A.1. General*

Deposition of hydrogenated amorphous silicon employing the VHF-PECVD technique (typical frequency range 20-110 MHz) has been reported to yield an increase in deposition rate by one order of magnitude with respect to the conventional used frequency of 13.56 MHz [16,146,250,280], without adversely affecting material quality [183,280,475]. This is of great importance for lowering the production cost of *a*-Si:H solar cells.

The explanation of the very high frequency (VHF) influence on the deposition rate is still a topic of discussion. It has been proposed theoretically that the high energy tail in the Electron Energy Distribution Function is increased with an increasing ratio  $\Omega/\nu$  of the excitation frequency and the energy collision frequency [276,476]. This leads to an increased ionization rate [146,250,280]. However, mass spectrometry results on the decomposition of silane as a function frequency show that the increase of the deposition rate cannot solely be attributed to the enhancement of radical production [119,120]. It has also been found or deduced that the flux of ions towards the surface is increased with increasing frequency [146,183,279,284] while at the same time the ion energy is decreased. This low-energy, high flux ion bombardment enhances surface mobilities of adsorbed species. It was shown in Sec. VI B 3 that a kinetic energy of 5 eV per deposited atom is needed to produce good quality *a*-Si:H [301].

Due to the fact that the wavelength of the radio-frequency signal is of the order of the substrate dimensions (3 m at 100 MHz), it can be expected that uniform deposition is more difficult at these high frequencies [477]. In fact, a practical optimum frequency is used around 60-70 MHz [478,479], which provides a good compromise between high deposition rate and attainability of uniform deposition. Further, the use of a distributed RF electrode network where all nodes have identical amplitude and phase improves the homogeneity of deposition [480].

#### *VIII.A.2. Optimization of deposition conditions*

A change of excitation frequency prompts for optimization of other deposition parameters, such as pressure, power, and geometry. Chatham and Bhat have shown that the maximum of SiH\* emission at 414 nm is shifted towards lower pressure upon increasing the frequency [146]. This means that the maximum in radical production also shifts to lower pressure. Experimental results obtained in the ASTER deposition system corroborate this [247].

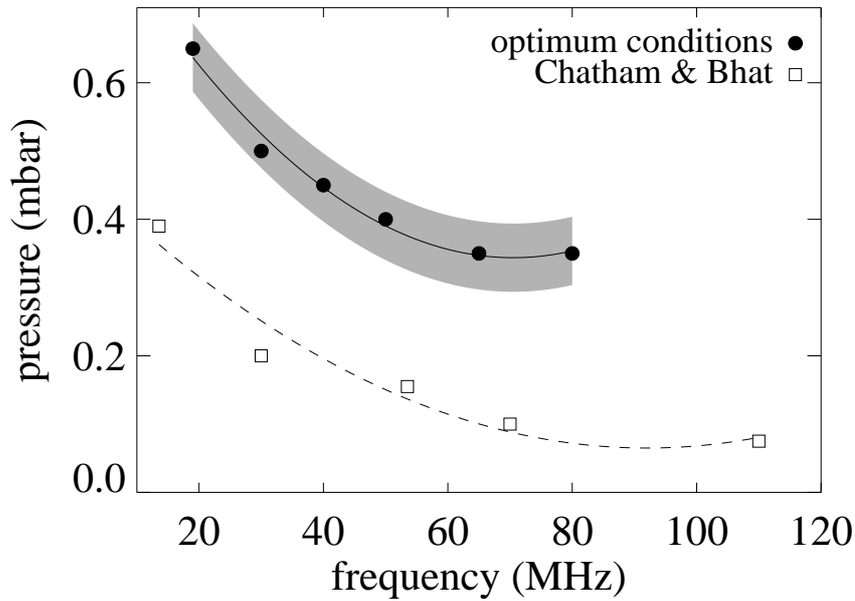


FIG. 54. Optimum values for pressure and frequency for uniform (thickness variation  $< 5\%$ ) film deposition. The shaded area signifies the process window. Also shown are similar data by Chatham and Bhat [146].

A 20-fold increase in deposition rate was reported for an increase in frequency from 13.56 to 100 MHz [119,120]. Here the electrode distance was about 4 cm, while the pressure was 0.12 mbar. A reasonable thickness uniformity was observed. For a smaller electrode distance (2.7 cm) higher pressures are needed in order to maintain good homogeneity in thickness. This is demonstrated in Fig. 54, where the conditions to obtain a homogeneity deviating less than 5% over a  $10 \times 10 \text{ cm}^2$  area are presented. The power density in this case was  $57 \text{ mW/cm}^2$ . The shaded area represents the process window. It is clear that for this electrode distance the homogeneity requirement can be met, if one decreases the pressure at a concomitant increase in frequency. This observation also explains the improvement in homogeneity as reported by Howling *et al.* [250]: at constant pressure the deposited layers become more homogeneous upon increasing the frequency. A similar effect was observed by Chatham and Bhat [146]. For comparison, their data are also shown in Fig. 54. The electrode distance in their work amounted to about 1 cm, while the reported power density was  $31 \text{ mW/cm}^3$ .

The fact that pressure and frequency are related in order to obtain homogeneous layers leads to the suggestion that the sheath thickness  $d_s$  between the grounded electrode and plasma bulk plays an important role. It is known, that an increase of frequency yields a decrease in sheath width, while a decrease in pressure enlarges it. The combination of these effects suggests that an optimum sheath width is required.

The dependence of sheath thickness versus frequency or pressure has been reported by several authors [179,328,481–485]. Their results are based on both theoretical studies concerning argon and experimental studies concerning argon, helium, and silane, using parallel plate configurations. It was found in these studies that  $d_s \propto p^a$ , where  $a$  ranges from -0.3 to -0.55 and  $d_s \propto \omega^b$ , where  $b$  ranges from -0.5 to -1.0. The data represented in Fig. 46 and Fig. 63 give  $a = -0.28 \pm 0.08$  and  $b = -0.26 \pm 0.02$ .

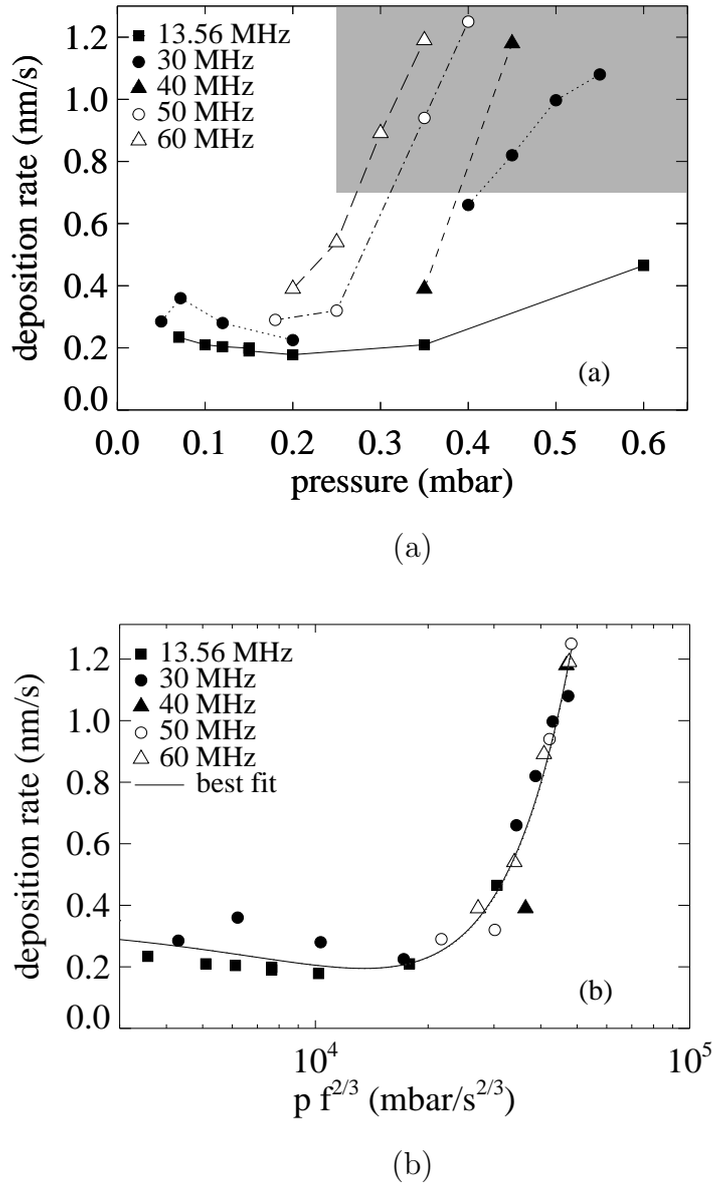


FIG. 55. (a) Deposition rate (nm/s) as function of pressure at various frequencies. The shaded area shows the 5 % uniformity conditions; (b) Best fit of deposition rate versus  $pf^{2/3}$ , clearly showing the  $\alpha$ - $\gamma'$  transition.

Fitting the data in Fig. 54 using the relationship  $p^a f^b = c$ , with  $c$  a constant which could represent  $d_s$ , shows that several combinations of the powers  $a$  and  $b$  can be used, as long as the power factors  $a$  and  $b$  are related as  $b \sim 0.45a$ . These fit results are inconsistent with the above mentioned values for  $a$  and  $b$ , hence the sheath thickness probably is not constant. This was also deduced from direct measurement of the IEDs [235,486,487]. In contrast, a recently developed method to determine the sheath thickness directly from deposition experiments [150,488] revealed that the value of  $d_s$  is constant for the conditions where  $p\omega^{1/2}$  is constant [487].

The deposition rate as function of pressure at various frequencies and at a power density of 57 mW/cm<sup>2</sup> is shown in Fig. 55a. The shaded area represents the conditions

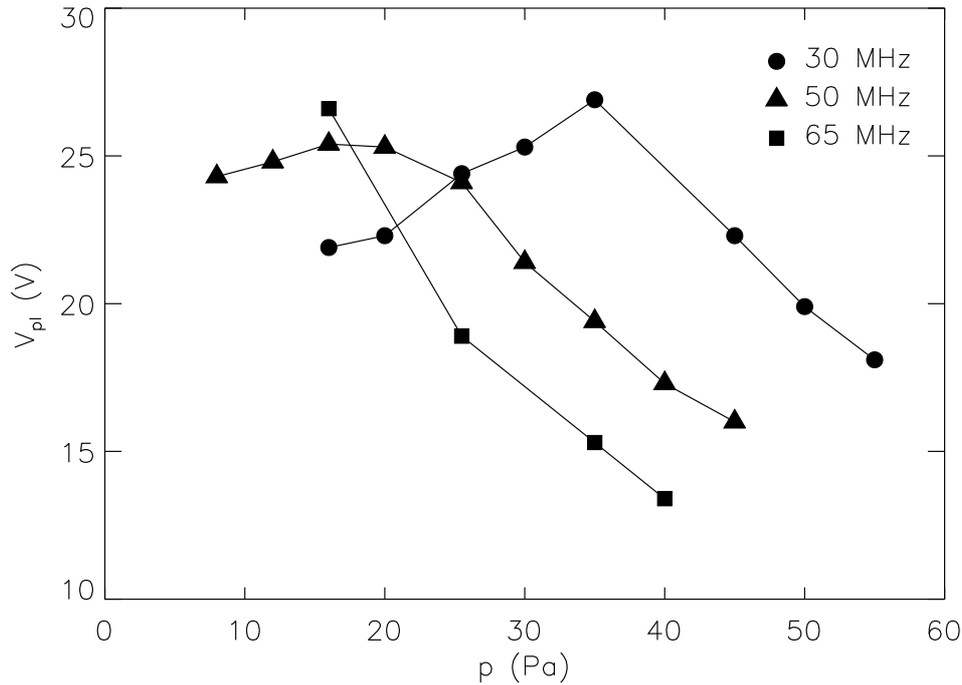


FIG. 56. The plasma potential as a function of pressure for a frequency of 30, 50, and 65 MHz. (From E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

for which uniform deposition is observed. At all frequencies the deposition rate at low pressure is lower than at high pressure by a factor of 5 to 6. This effect is known, and it is explained by the transition from the so-called  $\alpha$ - to the  $\gamma$ -regime of the plasma [245]. It follows from the figure that the transition region shifts to lower pressure with higher frequencies. Fitting the deposition rate as a function of the scaling parameter  $p^a f^b$  shows that a good fit is obtained for  $a = 1$  and  $b = 2/3$ , see Fig. 55b. This holds for all frequencies. The same behaviour has been found for the plasma potential [163]. As is illustrated in Fig. 56, the plasma potential slightly increases in the low pressure  $\alpha$  regime, but clearly decreases with increasing pressure at all three frequencies. The pressure at which the plasma potential starts to decrease occurs at a lower pressure at the higher frequencies. The rapid increase of deposition rate as a function of frequency, as reported in literature, e.g. [16,119,120,280,475], is observed in the  $\alpha$ -regime. In or near the  $\gamma$ -regime the enhancement in deposition rate as a function of frequency is not pronounced. This is illustrated in Fig. 57 by results at three power densities. The Chatham-and-Bhat data [146] also do show only a slight increase with frequency.

In Fig. 58 optical and structural material properties are shown. The refractive index at 2.07 eV or 600 nm,  $n_{2.07 \text{ eV}}$  decreases as function of applied RF power  $P$ , more or less independent on frequency (Fig 58a). The absorption coefficient at 2.07 eV behaves similarly. Values are around  $2$  to  $3 \times 10^4 \text{ cm}^{-1}$ . The cubic band gap  $E_g$  increases with power, also not dependent on frequency (Fig 58b). The hydrogen content  $C_H$  increases from 13 at.% at low power to 19 at.% at high power, again nearly independent on frequency (Fig 58c). The microstructure parameter  $R^*$  increases with power, with only a slight dependence on frequency (Fig 58d). The character of the intrinsic stress is

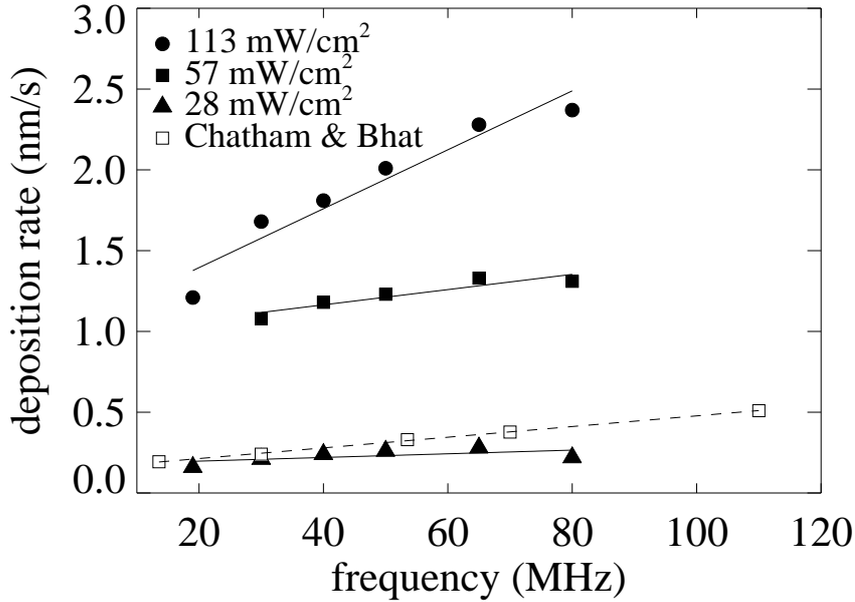


FIG. 57. Deposition rate (nm/s) as function of frequency at optimum pressure at each frequency (see Fig. 54 at three power densities. Also shown are data by Chatham and Bhat [146].

changed from compressive (about 600 MPa) at low power to tensile (100 MPa) at high power. Only a slight influence of frequency is observed, the dependence on power is much clearer. Small Angle X-ray Spectroscopy (SAXS) data show that voids are absent in the samples deposited at low power densities [66]. For higher powers voids are observed, and there appears to be a sudden change between 27 and 58 mW/cm<sup>2</sup>.

Electrical data are shown in Fig. 59 as a function of deposition rate for all frequencies, using the relation between deposition rate and power density as depicted in Fig. 54. Both dark and photo conductivity decrease exponentially with increasing deposition rate. The data in this range of deposition rates can be fitted with  $\sigma_d = 9 \times 10^{-12} \exp(-1.5r_d)$  and  $\sigma_{ph} = 8 \times 10^{-6} \exp(-0.5r_d)$ , with  $\sigma_d$  and  $\sigma_{ph}$  expressed in  $\Omega^{-1}\text{cm}^{-1}$ , and  $r_d$  in nm/s. Consequently the photoresponse  $\sigma_{ph}/\sigma_d$  increases with deposition rate as about  $10^6 \exp(r_d)$ . Activation energies amounted to typically 0.7-1.0 eV. From Thermally Stimulated Conductivity (TSC) measurements [489-492] a midgap density of states (DOS) of  $1.5 \times 10^{16} \text{ cm}^{-3}\text{eV}^{-1}$  is determined. The  $\mu\tau$ -product at 300 K is  $9 \times 10^{-5} \text{ cm}^2\text{V}^{-1}$ . Both DOS and  $\mu\tau$ -product are independent on frequency.

Summarising the results, it is clear that most of the material properties are not strongly dependent on frequency in the range of 30 to 80 MHz. The effect of power density is much more important. The material deposited at the lowest power is of ‘device quality’. Using larger power densities results in less dense material. This can be inferred from the decrease of the refractive index and the increase of the microstructure parameter, in combination with the increase in hydrogen content.

Perrin *et al.* [245] have shown that the increase in deposition rate is more pronounced above a certain threshold power density. This power density is close to the lowest power density in the experiments shown here. The deposition rate is probably the main parameter causing the observed difference between the low-power and high-power material and the deterioration of layers at higher power levels. This confirms the known observation that in general, the quality of material deposited in the  $\gamma'$ -regime is worse

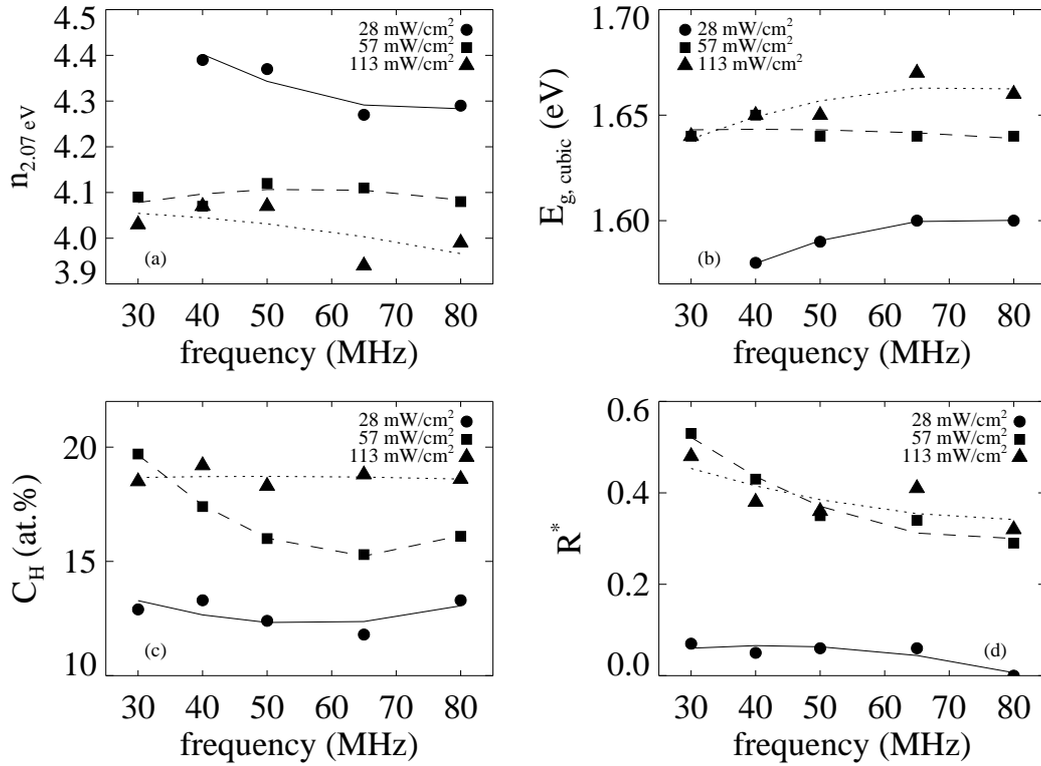


FIG. 58. (a) Refractive index  $n_{2.07 \text{ eV}}$ , (b) cubic band gap  $E_g$ , (c) hydrogen content  $C_H$  (at.%) and (d) microstructure parameter  $R^*$  as function of frequency at optimum pressures (see Fig. 54) at three power densities.

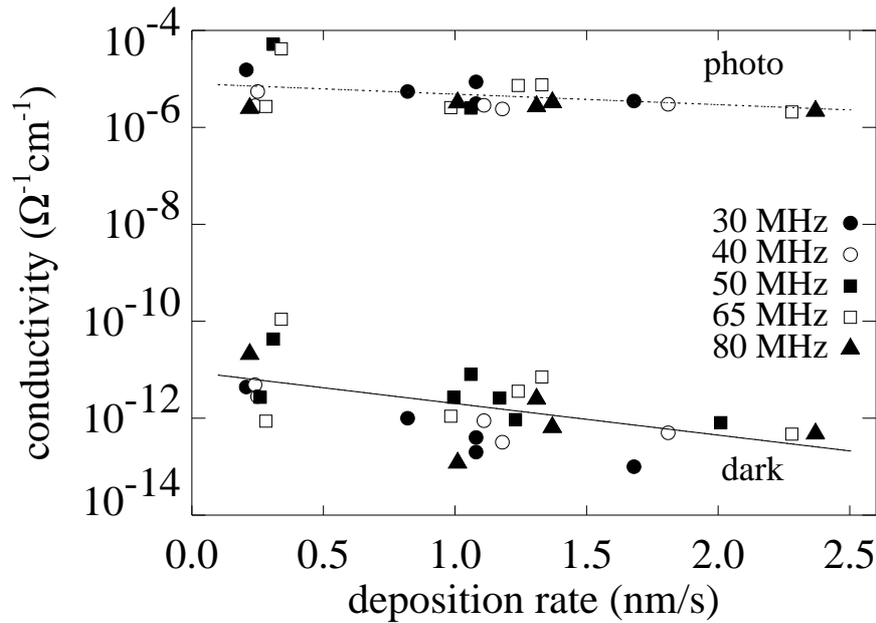


FIG. 59. Dark ( $\sigma_d$ ) and photoconductivity ( $\sigma_{ph}$ ) as function of deposition rate for all frequencies.

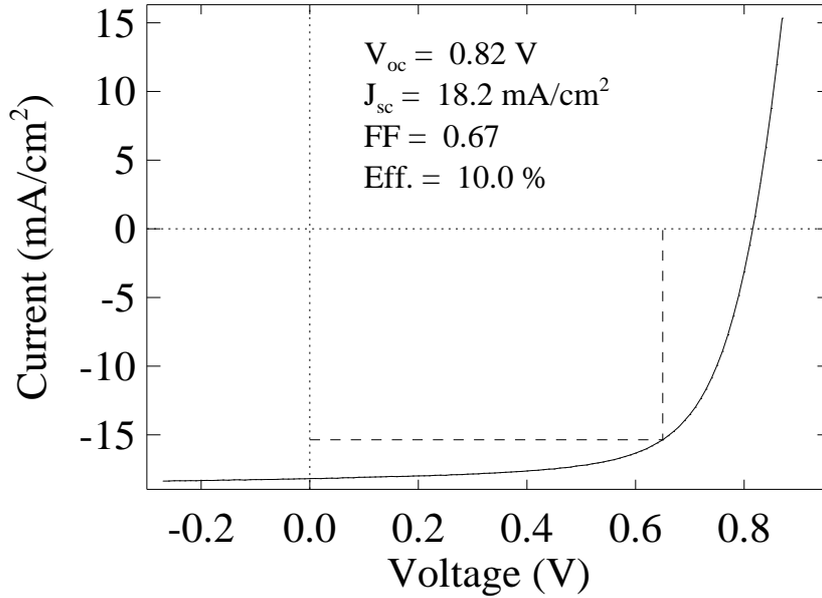


FIG. 60. Current-voltage characteristics of a solar cell made at 65 MHz and 42 mW/cm<sup>2</sup>. The dashed line indicates the maximum power point.

as compared to  $\alpha$ -regime material [245].

The presented material quality results are similar to results reported by others [119,120,280,475], although those frequency series were obtained in the  $\alpha$ -regime at constant low pressures. Here, good quality homogeneous layers were deposited *at* the  $\alpha$ - $\gamma'$  transition region. Especially the low power results compare well to results by others.

The intrinsic material fabricated at the frequencies reported above was incorporated in  $p^+i-n^+$  solar cells [493]. The  $p$  and  $n$ -layer were prepared by the conventional 13.56 MHz discharge. The ‘device quality’ films indeed yield good solar cells, of 10 % efficiency, as is shown in Fig. 60. This cell is manufactured with a 500 nm thick  $i$ -layer made at 65 MHz with a power density of 42 mW/cm<sup>2</sup>, resulting in excellent properties. The deposition rate still is 2-3 times higher than the conventional 13.56 MHz, i.e. 0.2–0.5 nm/s versus 0.1–0.15 nm/s for 13.56 MHz [494]. Similar results were reported by Jones *et al.* [495].

The effects of power density on the material quality are also reflected in the cell quality, e.g. the efficiency. Figure 61a shows a decrease in efficiency as function of power density or deposition rate. This is also seen in the spectral response measurements (Fig 61b). A shift in the red from high (700 nm) to lower (650 nm) wavelength is observed if one compares cells deposited at low and high power densities. This can be explained by the fact that the material is less dense, with a higher hydrogen content, yielding a larger band gap. The observed decrease in the blue part of the curve as a function of power density may probably be due to changes in the  $p$ - $i$  interface, which are induced by a different ion bombardment condition.

The differences in material properties are also reflected in the degradation behaviour of the cells. Figure 62 shows the normalized efficiencies as a function of illumination time. The efficiencies of the cells with the  $i$ -layer deposited at low

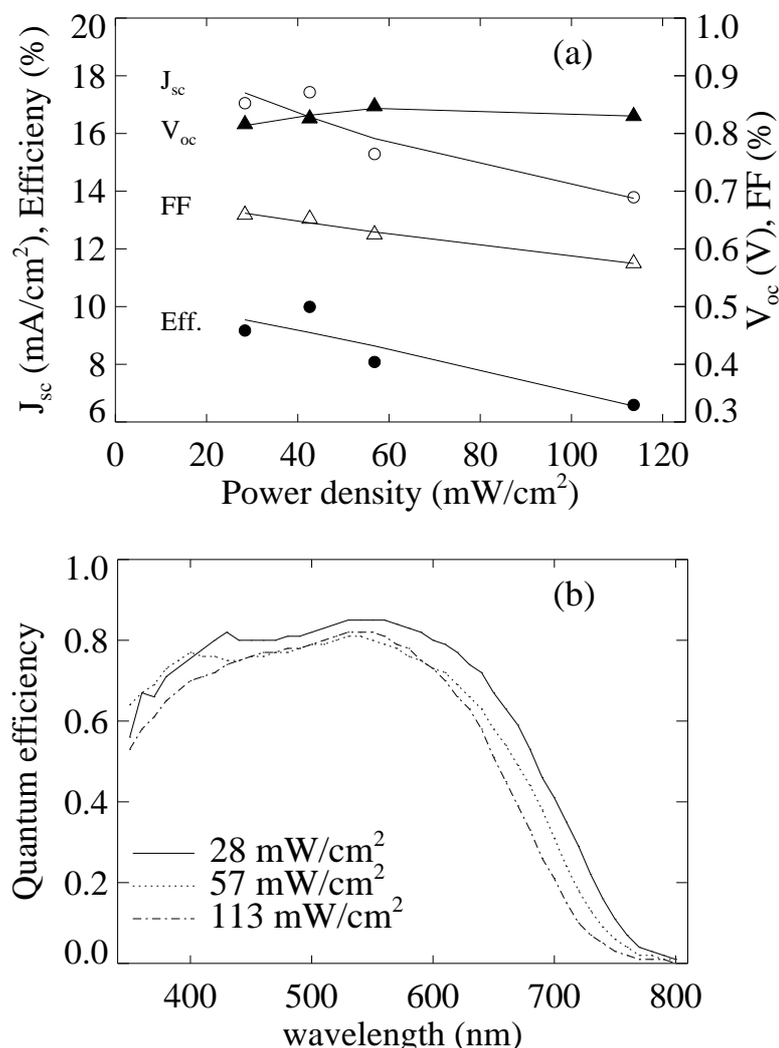


FIG. 61. Solar cell performance parameters as function of power density for cells made at 65 MHz: (a)  $J_{sc}$ , efficiency,  $V_{oc}$ , and  $FF$ ; (b) Spectral response.

power densities stabilize at around 60 % of their initial value, while the cells with the  $i$ -layer deposited at high power densities stabilize at 40 %. This correlates with the hydrogen content of the  $i$ -layers, which is around 12 % and 19 % for the low and the high power densities (see Fig. 58) and which is indicative for the low density material at obtained at these high power densities.

Also a frequency dependence was observed, especially in the  $V_{oc}$ , which was not expected from the material study. As *only* the deposition conditions of the  $i$ -layers are varied, a change at the  $p$ - $i$  interface must be responsible for the change in  $V_{oc}$ . The lower value of  $V_{oc}$  at low frequency was attributed to the difference in ion bombardment at the  $p$ - $i$  interface [493].

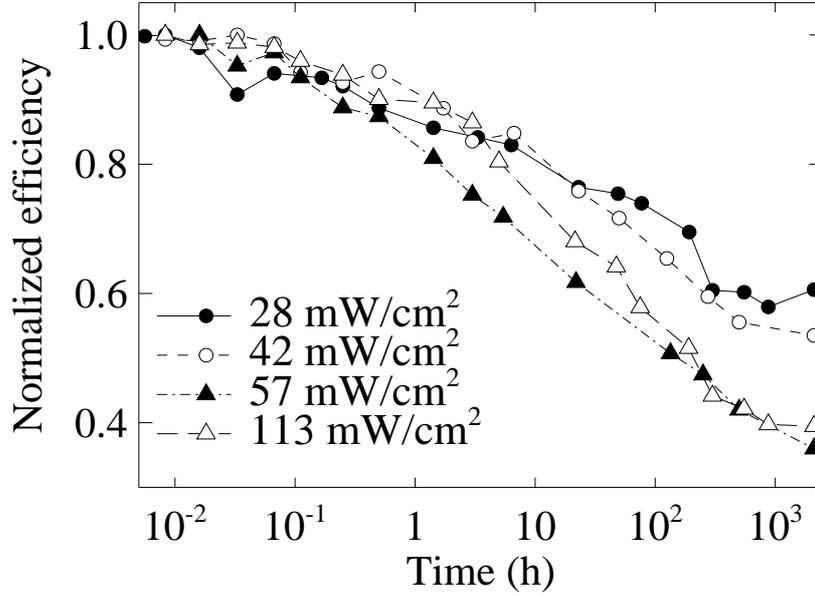


FIG. 62. Normalized solar cell efficiency as a function of illumination time for different power densities as obtained by continuous illumination of  $1000 \text{ W/m}^2$  AM1.5 light. The initial efficiencies of the four cells were 9, 10, 8, and 6% for 28, 42, 57,  $113 \text{ mW/cm}^2$  power density, respectively.

### VIII.A.3. Discharge analysis

In the ASTER deposition system, experiments have been carried out in which the excitation frequency was varied between 13.56 and 65 MHz. The other process conditions were kept constant at a power of 10 W, a pressure of 0.16 mbar, gas flows of 30 sccm  $\text{SiH}_4$  and 30 sccm  $\text{H}_2$  and a substrate temperature of  $250 \text{ }^\circ\text{C}$ . As above, plasma properties that are deduced from IED measurements are compared to material properties in Fig. 63. The IEDs of  $\text{SiH}_2^+$  at four frequencies are shown in Fig. 64.

The plasma potential is about 25 V (Fig. 63a). This value of the plasma potential is typical for the silane plasmas in the *asymmetric* RF capacitively coupled reactors as used in the ASTER deposition system and is also commonly found in argon or hydrogen plasmas [170,327,280]. From the considerable decrease of the DC self bias as a function of frequency (Fig. 63a) it is inferred that the potential drop over the sheath of the grounded electrode ( $V_{pl}$ ) and the one over the sheath of the powered electrode ( $V_{pl} - V_{DC}$ ), become comparable in magnitude. Hence, the discharge is becoming more symmetric with increasing frequency.

In contrast, Heintze and Zedlitz [236] also presented data on the plasma potential as function of frequency in silane plasmas: the plasma potential varies from about 27 V at 35 MHz to about 20 V at 180 MHz. Moreover, Dutta *et al.* [284] used a *symmetric* RF capacitively coupled reactor and estimated the plasma potential in their system from the applied voltage at the powered electrode. A decrease of the plasma potential from 45 V at 13.56 MHz to only 15 V at 70 MHz is observed. This difference in behaviour is thought to be solely due to the different reactor geometries.

The charge carrier density in the sheath increases by a factor of 3, when increasing the excitation frequency from 13.56 to 65 MHz. As a consequence, the sheath thickness

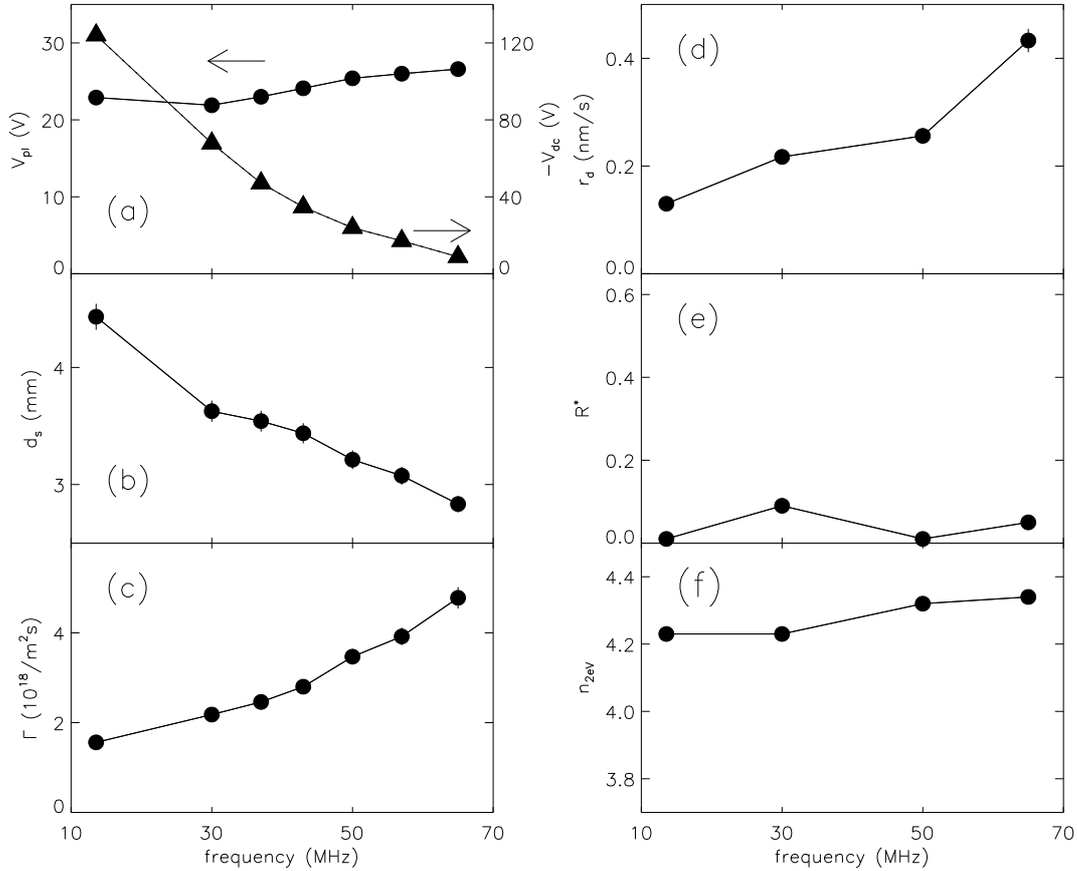


FIG. 63. Plasma parameters as deduced from the IEDs and material properties as a function of excitation frequency of to the  $\text{SiH}_4/\text{H}_2$  discharge at a power of 10 W and a pressure of 0.16 mbar: (a) the plasma potential  $V_{pl}$  (circles) and DC self bias  $V_{DC}$  (triangles), (b) the sheath thickness  $d_s$ , (c) the maximum ion flux  $\Gamma_{max}$ , (d) the growth rate  $r_d$ , (e) the microstructure parameter  $R^*$ , and (f) the refractive index  $n_{2eV}$ . (Compiled from E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

$d_s$  decreases from 4.4 mm at 13.56 MHz to 2.8 mm at 65 MHz (Fig. 63b), which results in a reduced number of collisions in the sheath. This is nicely illustrated in Fig. 64, where the IEDs of  $\text{SiH}_2^+$  are shown for several frequencies. Clearly, the contribution of low energy ions decreases upon increasing the frequency.

The increase in the deposition rate  $r_d$  (Fig. 63d) corresponds to the increase in the ion flux (Fig. 63c): the fraction of arriving ions per deposited atom  $R_i$  is constant at about 0.25. Such observations have also been reported by Heintze and Zedlitz [236], who furthermore suggested that the deposition rate may well be controlled by the ion flux. The kinetic ion energy per deposited atom  $E_{max}$  is also constant and amounts to about 5 eV. As was shown above (Sec. VIB3) the material quality as reflected in the refractive index  $n_{2eV}$  (Fig. 63e) and the microstructure parameter  $R^*$  (Fig. 63f) is good:  $n_{2eV}$  is around 4.25 and  $R^*$  is low ( $< 0.1$ ). The depletion of the silane stays constant at a value of  $4.0 \pm 0.4$  sccm in this frequency range. The partial pressures of silane, hydrogen, disilane ( $1.3 \times 10^{-3}$  mbar) and trisilane ( $2 \times 10^{-5}$  mbar) in the plasma are also constant

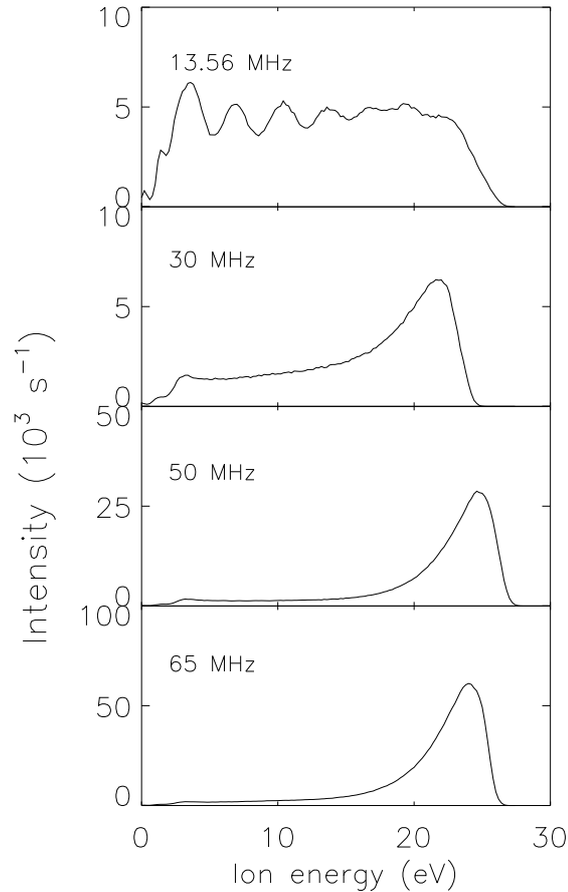


FIG. 64. The ion energy distributions of  $\text{SiH}_2^+$  at several frequencies. (From E. A. G. Hamers, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

with frequency. Similar results have been obtained by Heintze and Zedlitz [280]. As a result of varying the frequency from 50 to 250 MHz, the dissociation rate only increased by a factor of 1.5, while the deposition rate on the electrodes increased by nearly an order of magnitude.

From the constant depletion, based on a homogeneous deposition throughout the reactor a frequency-independent deposition rate is deduced of 0.44 nm/s [163]. At 13.56 MHz the observed deposition rate is a factor of 3 lower, whereas at 65 MHz the measured deposition rate equals the estimated value of 0.44 nm/s. This suggests that the increased deposition rate at higher frequencies, as measured at the center of the grounded electrode, is partly due to a more homogeneous deposition profile throughout the reactor at higher frequencies [163,280]. This suggestion is further supported by the fact that the discharge is more symmetric at higher frequencies, as deduced from the low DC self bias at high frequencies.

### VIII.B. Chemical annealing

It was recognized by, e.g., Tsai *et al.* [496] that high hydrogen dilution resulted in microcrystalline films as a result of atomic hydrogen that preferentially etches weak

bonds during deposition. This prompted the use of fluorine-containing precursors, with which enhanced etching is achieved during deposition. Etching by means of periodically interrupting the deposition process has been reported by Tsuo *et al.* [497], who used a constant  $\text{XeF}_2$  flow and a periodic  $\text{SiH}_4$  flow. The use of an RF hydrogen discharge for the etching cycles and a diluted silane discharge for the deposition cycles was used by Xu, as reported by Luft and Tsuo [6]. With both methods good material is obtained.

The "chemical annealing" technique was proposed by Shirai *et al.* [17,471,498], as a means to tackle the metastability problem in *a*-Si:H solar cells. It aims at the lowering of the hydrogen content in *a*-Si:H, by alternating deposition and exposure to atomic hydrogen. It was demonstrated that a lower hydrogen content indeed leads to more stable material. A two-discharge zone apparatus was used by Asano *et al.* [499,500]. In one zone  $\text{SiH}_4$  and  $\text{H}_2$  were mixed to deposit *a*-Si:H, while in the other zone pure  $\text{H}_2$  was used for the etching process. The substrate was rotated between the two zones. Chemical annealing alternatively is termed ADHT (Alternately repeating Deposition and Hydrogen plasma Treatment) and LBL (Layer-By-Layer).

The application of this technique in conventional RF deposition equipment revealed the possibility of producing microcrystalline silicon at relatively low power densities and temperatures [501]. Depending on deposition conditions, i.e. deposition cycle time and hydrogen exposure time and the power density, different microcrystalline fractions are found, see [251,165,502].

As an example the Raman spectra of three films deposited in the ASTER deposition system are shown in Fig. 65, for which only the hydrogen exposure time was varied, 10–30 s. In the deposition cycle a 1:1 mixture of silane and hydrogen was used. Conditions were chosen such that device-quality material is obtained [370]. The duration of the deposition cycle was kept constant at 10 s. At the start of the  $\text{H}_2$ -plasma cycle the  $\text{SiH}_4$  flow was stopped and extra  $\text{H}_2$  was introduced, to keep the pressure constant. A practical problem to be addressed is the matching of the discharge when changing from  $\text{H}_2$  to  $\text{SiH}_4$ , and vice versa. A sudden change in DC self bias is observed, as well as a change in reflected power. Fast automated matching networks are to be used to minimize these effects.

The broad TO peak, characteristic for *a*-Si:H is observed at  $480\text{ cm}^{-1}$ , while the crystalline peak is at  $520\text{ cm}^{-1}$ . The ratio between integrated peak intensities clearly depends on hydrogen exposure time. The hydrogen content linearly increases with exposure time, where the extra hydrogen is present in  $\text{SiH}_2$  bonds. These effects were thought to originate from the hydrogen bombardment during the hydrogen exposure cycle [165].

Layadi *et al.* have shown using *in situ* spectroscopic ellipsometry that both surface and sub-surface processes are involved in the formation of  $\mu\text{c-Si}$  [502,503]. In addition, it was shown that the crystallites nucleate in the highly porous layer below the film surface [502,504], as a result of energy released by chemical reactions [505,506] (chemical annealing). In this process four phases can be distinguished, incubation, nucleation, growth, and steady state [507]. In the incubation phase, the void fraction increases gradually, while the amorphous fraction decreases. Crystallites start to appear, when the void fraction reaches a maximum (at about a thickness of 10 nm). Subsequently, the crystalline fraction increases exponentially in the nucleation phase. Then, a linear increase in the crystalline fraction sets in, corresponding to the growth of crystallites in the whole film. At a certain thickness, the crystalline fraction remains constant, and

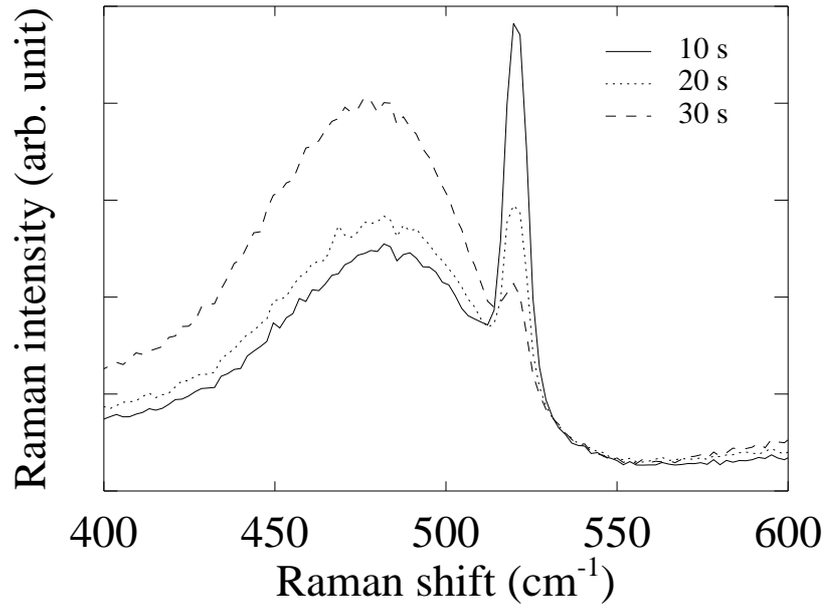


FIG. 65. Raman spectra for films deposited employing the chemical annealing technique for different hydrogen exposure times.

a steady state situation is reached. *In situ* conductivity measurements revealed that the dark conductivity increases by more than an order of magnitude at the onset of the growth phase [507]. These films have been employed in Thin Film Transistors, and an excellent stability of the TFTs was demonstrated [508].

It was argued by Saito *et al.* [509] that a hydrogen plasma treatment as used in the LBL technique causes "chemical transport". In this case the *a*-Si:H film deposited on the cathode prior to the hydrogen treatment is etched and transferred to the anode. Hence, in the hydrogen plasma silane related molecules and radicals are present. In fact, material is deposited on the anode under identical conditions as in the case of high dilution of silane with hydrogen.

### VIII.C. RF modulation

Modulation of the RF excitation has been used in an attempt to increase the deposition rate. Increasing the gas pressure or raising the power generally leads to dust formation and deterioration of material properties. To overcome this problem, one can pulse the plasma by modulating the RF signal in amplitude by a square wave. Depending on the regime ( $\alpha$  or  $\gamma'$ ) different effects are observed.

A considerable decrease both in the deposition rate and in the powder formation was found, when the 13.56 MHz excitation was modulated by a square wave (SQWM) of 2 Hz [510,511]. In the  $\gamma'$  regime, Biebericher *et al.* [512] have observed a decrease in deposition rate from 1.0 nm/s in a continuous-wave (CW) 50 MHz SiH<sub>4</sub>/H<sub>2</sub> plasma, to 0.2 nm/s in a similar (i.e. with the same *average* power of 10 W) plasma, modulated by a frequency of 100 Hz.

When modulating the power in the  $\alpha$  regime, an increase in deposition rate has been observed by SQWM with an optimum at a frequency of about 100 kHz [513–515].

It was reported that the time-averaged electron density for a modulated discharge was higher than in the case of a continuous discharge [18]. Moreover, the amount of dust generated in the discharge was much lower, due to the low density of negative ions.

At the start of each modulation pulse, a sharp peak in optical emission is seen. Similar  $\text{SiH}^*$  emission peaks in pulsed plasmas have been found by Scarsbrook *et al.* [516] and Howling *et al.* [321]. The sharp peak was claimed to be caused by a pulse of high-energy electrons. Overzet and Verdeyen [517] measured electron densities, at a 2.9 MHz excitation frequency and modulation frequencies up to 20 kHz. The optical emission of a SQWM argon plasma was measured by Booth *et al.* [518], who also performed particle-in-cell modeling.

Here, results are shown from experiments performed in ASTER, reported by Biebericher *et al.* [512,519]. A  $\text{SiH}_4/\text{H}_2$  (50:50 flow ratio, total flow 60 sccm) plasma was generated at an RF excitation frequency of 50 MHz. The substrate temperature was 250°C. The RF signal was amplitude modulated (AM) by a square wave. The modulation frequency has been varied in a range of 1-400 kHz. The modulation depth was always 90%. The duty cycle was fixed at 50%. The pressure amounted to 0.2 mbar and the *average* power was kept at 10 W. With a duty cycle of 50%, this leads to a power of 20 W during the 'plasma-on' period.

The measured dependence of deposition rate on the modulation frequency is shown in Fig. 66a. It can be clearly seen that the deposition rate at the optimum of 100 kHz is about three times higher than the deposition rate of a CW discharge of 10 W (dotted line). In addition, there is a clear correlation between deposition rate and the time-averaged optical emission intensity of the pulsed discharge. This average intensity is related to the electron temperature: a higher light intensity can be attributed to an increase in the electron temperature. This results in an increased production of radicals, which is reflected in an increased deposition rate.

The spectrally integrated (185-820 nm) emission of a 50 MHz discharge, modulated by 50 kHz, is shown in Fig. 66b. Similar results are obtained for the  $\text{SiH}^*$  emission. The peak appearing at the onset is generated by the plasma. The rise time of about 1  $\mu\text{s}$  and the FWHM of the peak of 6  $\mu\text{s}$  are much longer than the 40 ns detection limit of the electronics. When the power is turned off, the light intensity decreases very fast and the plasma fully extinguishes.

Upon changing the modulation frequency it was observed that only the length of the plateau varied, whereas the height and FWHM of the peak at plasma onset remained almost constant. Therefore, the relative importance of the peak in the time-averaged light intensity is higher at higher modulation frequencies, and consequently, the average light intensity increases with increasing modulation frequency. This holds up to the observed optimum of 100 kHz. At this frequency the 'plasma-on' period is 5  $\mu\text{s}$ , i.e. about equal to the FWHM of the peak. A further increase in modulation frequency leads to a reduction of the peak and the average light intensity.

In order to be able to explain the observed results plasma modeling was applied. A one-dimensional fluid model was used, which solves the particle balances both for the charged and neutral species, using the drift-diffusion approximation for the particle fluxes, the Poisson equation for the electric field, and the energy balance for the electrons [191] (see also Sec. IV A).

The calculated production of  $\text{SiH}_3$  radicals during two modulation cycles at a modulation frequency of 50 kHz is also shown in Fig. 66b. A good correlation with the

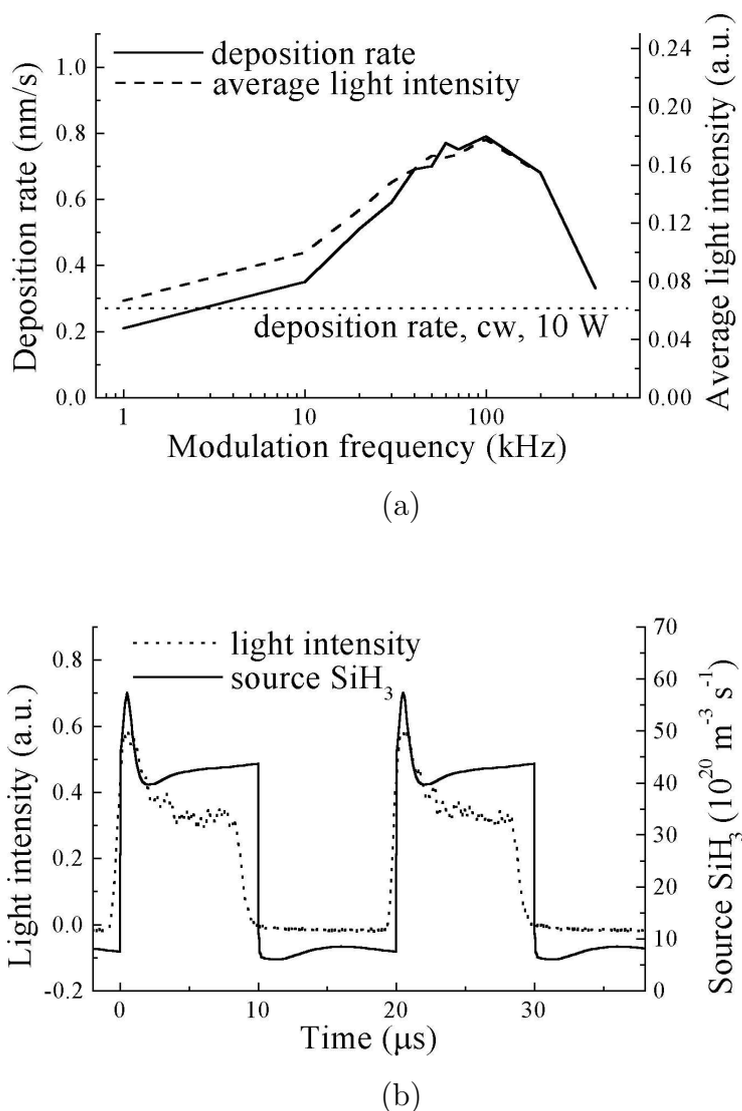


FIG. 66. Effects of modulating the RF excitation frequency: (a) deposition rate and average light intensity as a function of the modulation frequency, with the deposition rate at CW conditions indicated by the dotted line, (b) measured spectrally integrated emission and calculated production of  $\text{SiH}_3$  radicals as a function of time, at a modulation frequency of 50 kHz and a 50% duty cycle. (From A. C. W. Biebericher, J. Bezemer, and W. F. van der Weg, *Appl. Phys. Lett.* **76**, 2002 (2000).)

behaviour of average light intensity versus time is observed, including the sharp peak at plasma onset. The shape of the curve is determined by the instantaneous sharp rise of dissociation and by chemical reactions. The non-zero production of  $\text{SiH}_3$  radicals during the 'plasma-off' period is due to chemical reactions.  $\text{SiH}_3$  is not only produced by dissociation of  $\text{SiH}_4$  through electron impact, but also by an abstraction reaction of  $\text{SiH}_4$  and H. These hydrogen atoms can originate from the dissociation of  $\text{SiH}_4$  into  $\text{SiH}_2$  and  $2\text{H}$ .

The behaviour of the electrons as a function of time can explain the observed results. Simulation results for a discharge operated at a pressure of 0.267 mbar, a frequency of

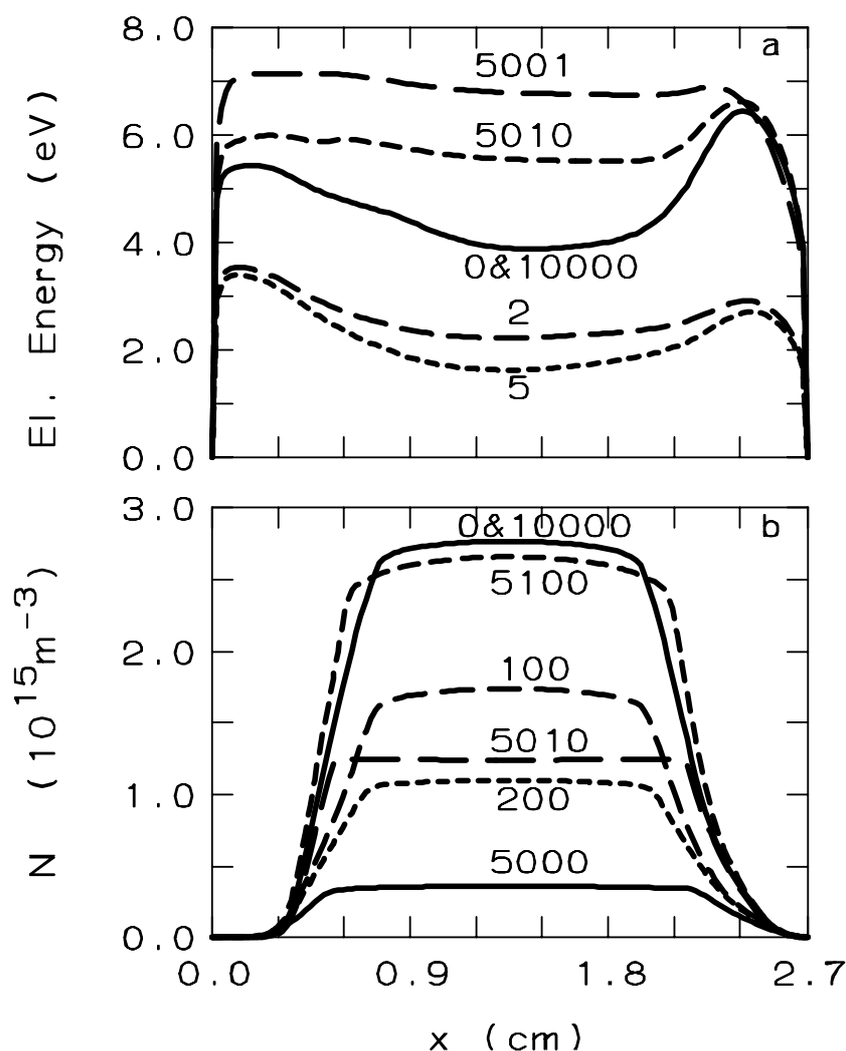


FIG. 67. The development in time of the average electron energy (a) and of the density of electrons (b) between the electrodes in the case of a 50 MHz plasma and a modulation frequency of 5 kHz. The RF electrode is located at  $x = 0$ , and the grounded electrode at  $x = 2.7$  cm. The numbers near the curves in both graphs represent the number of RF cycles. One RF cycle  $\equiv 20$  ns (50 MHz). The plasma is turned off at cycle 0, turned on at cycle 5000, and turned off again at cycle 10000. (From A. C. W. Biebericher, J. Bezemer, and W. F. van der Weg, *Appl. Phys. Lett.* **76**, 2002 (2000).)

50 MHz, an electrode distance of 27 mm, a voltage modulated between 134 V (power 20 W) and 40 V (power  $< 1$  W), and a modulation frequency of 5 kHz are shown in Fig. 67. In this figure the electron energy and density are shown as a function of the distance between the electrodes. The numbers indicate the numbers of passed RF cycles. A modulation frequency of 5 kHz corresponds to 10000 RF cycles. The plasma is turned off at cycle 0 and is turned on at cycle 5000. It can be seen in Fig. 67a that the average electron energy decreases very fast. This is consistent with the observed decay of the emission of the discharge, see Fig. 66b. The electron density also decreases, albeit with a longer decay time. The electron density is about halved after 100 cycles. Thus,

very soon after shut-off, within a few cycles, no new radicals are produced, apart from those generated in neutral-neutral reactions. The electron (and ion) density decreases by the loss due to ambipolar diffusion to a value one order of magnitude lower than the value at cycle 0. When the power is switched on at cycle 5000 the electron energy increases to a value nearly two times larger than the value at cycle 0 (and 10000). This fast increase is attributed to the low electron density at cycle 5000, that absorbs the power. The increase in electron energy results in an even steeper increase in reaction rates per electron, due to the energy-dependence of the cross-sections. Thus, a peak in light intensity as well as a peak in  $\text{SiH}_3$  radical production is observed. This leads to an increase of the average deposition rate to a value above the rate corresponding to the average deposited power (10 W), cf. Fig. 66a. At a low modulation frequency (1 kHz) the decay of the profiles and the overshoot of the rates cover only a small fraction of the modulation period. In other words, the discharge is in a quasi steady state with respect to the applied power, and the deposition rate is only slightly above the corresponding deposition rate at CW conditions. As the modulation frequency increases, the overshoot becomes relatively more important, which leads to an increase in deposition rate. When the period of the modulation becomes shorter than the decay time of the electron density, the power is distributed over too many electrons to yield considerable heating. The heating becomes less and less radicals are produced, as is shown in Fig. 66a. It is therefore concluded that an optimum in the modulation frequency exists, which corresponds to the decay time of the electron density. From Fig. 67b it can be inferred that this optimum should be around 100 kHz, as at 200 cycles the electron density is reduced to  $1/e$  of its value at cycle 0, i.e. the lifetime is 200 RF cycles, or 4  $\mu\text{s}$ .

Another beneficial effect of modulation of the discharge is the observed improvement of uniformity of deposition. At CW discharge conditions that normally result in thickness variations of 70%, modulation lowers the variation to about 10% [519]. This was attributed to the fact that, as a result of modulation, reactions are not confined to the sheaths, and radicals are produced in the entire volume between the electrodes.

Madan *et al.* [515] have presented the effect of modulation on the properties of the material (dark- and photoconductivity) and of solar cells. They also observe an increase in deposition rate as a function of modulation frequency (up to 100 kHz) at an excitation frequency of 13.56 MHz, in their PECVD system [159]. The optimum modulation frequency was 68 kHz, which they attribute to constraints in the matching networks. Increasing the deposition rate in CW operation of the plasma by increasing the RF power leads to worse material. Modulation with a frequency large than 60 kHz results in an improved material quality, for material deposited with equal deposition rates. This is also seen in the solar cell properties. The intrinsic  $a\text{-Si:H}$  produced by RF modulation was included in standard  $p\text{-i-n}$  solar cells, without buffer or graded interface layers. For comparison also solar cells employing layers that were deposited under CW conditions were made. At a low deposition rate of about 0.2 nm/s CW solar cell parameters are:  $V_{oc} = 0.83$  V,  $J_{sc} = 15.8$  mA/cm<sup>2</sup>, FF = 0.67, and  $\eta = 8.8$  % (at AM1.5 illumination conditions). Solar cells produced with the intrinsic layer deposited at 68 kHz modulation frequency are of the same quality:  $V_{oc} = 0.81$  V,  $J_{sc} = 17.1$  mA/cm<sup>2</sup>, FF = 0.63, and  $\eta = 8.7$  %.

Biebericher *et al.* [519] report that beyond a modulation frequency of about 40 kHz in their 50 MHz discharge the material is slightly worse than at standard CW conditions.

However, they argue that due to the enhancement in deposition rate, lower partial pressures of silane can be used, which results in more efficient silane consumption. Indeed, they have shown that lowering the gas flows by a factor of three leads to a reduction in deposition rate of only a factor of 1.5 at 100 kHz modulation frequency [520]. Clearly, the efficiency of silane consumption is of great importance in production.

## IX. HOT WIRE CHEMICAL VAPOUR DEPOSITION

### IX.A. General description

The Hot-Wire Chemical Vapour Deposition (HWCVD) method has been introduced already in 1979 by Wiesmann *et al.* [521]. The principle of HWCVD is based on the thermal decomposition of silane at a heated tungsten (or tantalum) filament, foil, or grid. At temperatures of 1400-1600 °C the silane is decomposed into a gaseous mixture of silicon and hydrogen atoms. Matsumura further developed HWCVD but termed it catalytical CVD (CTL-CVD) [522-524], as he showed that the decomposition of silane at a heated tungsten filament is a catalytic process. The use of a much higher pressure ( $> 0.1$  mbar) compared to the one Wiesmann *et al.* used ( $< 5 \times 10^{-4}$  mbar) lead to a high deposition rate (0.5 nm/s). Doyle *et al.* showed that the silicon atoms deposited on the filament are then thermally evaporated onto the substrate that is located within a few cm from the filament, hence the term evaporative surface decomposition [525]. At the same time, thermal and catalytic dissociation of silane and hydrogen lead to the generation of hydrogen atoms at the filament surface [526].

Device quality *a*-Si:H made by HWCVD (as they termed it) was first reported by Mahan *et al.* [19,527]. They obtained *a*-Si:H with hydrogen concentrations as low as 1%. Deposition rates as high as 5 nm/s [528] and 7 nm/s [529] have been achieved while the *a*-Si:H was of high quality. In order to obtain device quality material it was shown by Doyle *et al.* [525] that the radicals that are generated at the filament (atomic Si and atomic H) must react in the gas phase to yield a precursor with high surface mobility. Hence, the mean free path of silane molecules should be smaller than the distance between filament and substrate  $d_{fs}$ . Too many reactions between radicals and silane molecules, however, result in worse material. In fact, optimal film properties are found for values of  $p d_{fs}$  of about 0.06 mbar cm [530,531].

The combination of high deposition rate and the ability to produce device quality material is of particular interest for solar cell production and thin-film transistor (TFT) fabrication [532-535]. Further, the low hydrogen content was expected to yield improved stability against light-induced degradation [527], as the Staebler-Wronski effect is related to the hydrogen content in the material (see also Sec. IB5). This was demonstrated by Crandall *et al.* [536], who incorporated an HWCVD-deposited layer in solar cells, and observed a reduced degradation upon light soaking as compared to devices with a conventional PECVD layer. In these "hybrid" solar cells only the intrinsic layer was made by HWCVD, all other (doped) layers were deposited by PECVD. Wang *et al.* [537] have reported solar cells which were completely made using HWCVD.

HWCVD-deposited *a*-Si:H layers have also successfully been used as the semiconductor layer in inverted-staggered TFTs [538,539]. Moreover, it was demonstrated that these TFTs have excellent electrical properties and do not suffer from a shift of the threshold voltage upon prolonged application of a gate-voltage. As in a TFT the metastable character of *a*-Si:H manifests itself as a reversible threshold-voltage shift upon prolonged application of gate-voltage (see Sec. XIB), these HWCVD-TFTs are stable [538,540].

Dilution of silane with hydrogen leads to the formation of polycrystalline silicon films (poly-Si:H) that still contain a low amount ( $< 1\%$ ) of hydrogen [529,541-544]. This is believed to occur as a result of increased etching by atomic hydrogen. Molecular

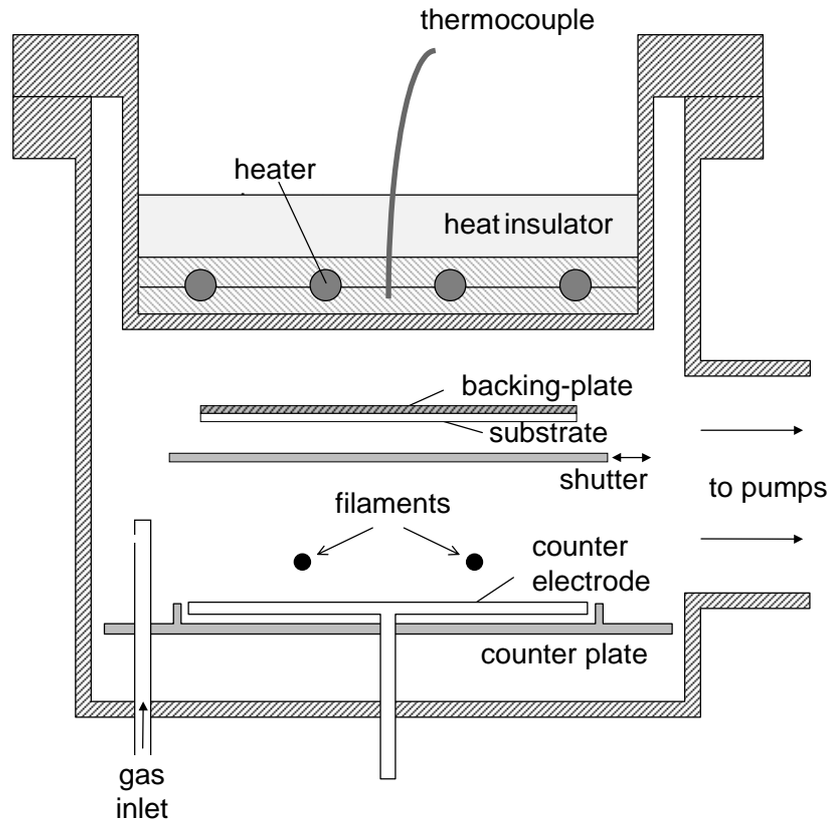


FIG. 68. Cross-sectional view of a HWCVD deposition reactor. (From K. F. Feenstra, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

hydrogen is decomposed at the filament to reactive atomic hydrogen that is able to etch silicon atoms from disordered or strained bonding sites in the amorphous (sub)surface network. The amorphous network thus is transformed to a polycrystalline one, without amorphous tissue. Large grains are obtained (70 nm average grain size) and the crystalline fraction in these highly compact materials amounts to 95% [544]. The grain boundaries are passivated by the small amount of hydrogen.

Doping is achieved by adding dopant gases such as phosphine, diborane, or trimethylboron to the silane [545–547]. The doped material is of electronic quality comparable to PECVD doped material.

### IX.B. Experimental set-up

A cross-sectional overview of a HWCVD set-up is shown in Fig. 68 [548]. This particular reactor is part of a multichamber, ultrahigh vacuum deposition system, PASTA, described by Madan *et al.* [159]. The background pressure of the reactor is  $10^{-8}$  mbar. In this HWCVD reactor two coiled tungsten filaments are mounted parallel to each other in order to improve thickness uniformity. The filaments have a diameter of 0.5 mm, and a length of 15 cm. The distance between the filaments is 4 cm, while the distance from the filaments to the substrate ( $d_{fs}$ ) amounts to 3 cm. The filaments are heated by sending a DC current through the wires. Calibration of the filament tem-

perature as a function of DC current is needed, and can be done with a pyrometer. Temperatures exceeding about 1800 °C should be used as to avoid silicide formation on the filament. On the other hand, the temperature should not be so high that tungsten is evaporated from the filament and is incorporated in the film [549].

A shutter can be placed in front of the substrate to prevent unwanted deposition during filament pre-heating. The reaction gases are injected on the left side of the reactor, and directed towards the reactor wall in order to achieve more homogeneous gas flows. The (unreacted) gas is pumped out at the right side, so the gas flow is perpendicular to the filaments.

The substrate is radiatively heated by heaters that are placed outside of the vacuum. A backing plate ensures a laterally homogenous temperature profile. In the same chamber also PECVD can be carried out. The backing plate then is the grounded electrode, and the RF voltage is applied to the counter electrode.

The high filament temperature used causes additional radiative heating of the substrate [530,531]. Feenstra *et al.* [531,548] have developed a heat transport model of their set-up (Fig. 68). All heat exchange is assumed to occur via radiation. They found, e.g., that at a heater temperature of 100 °C and a filament temperature of 1900 °C the substrate temperature reached a value of 350 °C after a typical stabilization time of 25 minutes. With the shutter closed the stable substrate temperature was 50 °C lower. This can be exploited to preheat the substrate, when the shutter is closed, to the desired temperature. Just before the deposition is started, the filament temperature is set to its correct value, and the shutter can be opened. This procedure avoids any temperature drift during deposition.

### IX.C. Material properties and deposition conditions

With the set-up described above a series of depositions was carried out [531,548], in which the substrate temperature was varied between 125 and 650 °C, the pressure between 0.007 and 0.052 mbar, the gas flow rate between 15 and 120 sccm, and the dilution of the silane gas with hydrogen ( $[\text{SiH}_4]/([\text{SiH}_4]+[\text{H}_2])$ ) between 0.1 and 1. Under these conditions the deposition rate varied between 1 and 2.5 nm/s [531]. Molenbroek *et al.* [530] reported a variation of the deposition rate between 1 and 9 nm/s for similar process conditions, at a filament temperature of 2000 °C.

The variation of deposition temperature has similar effects on the material properties compared to PECVD deposited material. With increasing temperature (125-650 °C), the material becomes more dense (the refractive index extrapolated to 0 eV increases from 3.05 to 3.65), the hydrogen content is decreased (15 to 0.3 at.%), as well as the microstructure factor (0.4 to 0). The activation energy is 0.83 eV up to a deposition temperature of 500 °C. The dark and AM1.5 photoconductivity are about  $5 \times 10^{-11} \Omega^{-1}\text{cm}^{-1}$  and  $5 \times 10^{-6} \Omega^{-1}\text{cm}^{-1}$ , respectively, up to a deposition temperature of 500 °C, i.e., the photoresponse is  $10^5$ .

The deposition rate increases upon increasing the pressure. This is explained by noting that the impingement rate per unit area  $r_i$  of molecules on the filament is linearly dependent on the pressure as  $r_i = p/\sqrt{2\pi k_B T_g}$ , with  $T_g$  the gas temperature. However, as the pressure becomes higher, the collisional mean free path of the silane becomes smaller, and the silane supply to the filaments becomes restricted. Moreover,

the transport of deposition precursors to the substrate is restricted as well. The mean free path of silane was estimated to be 2.5 cm at a pressure of 0.02 mbar [531], i.e., the mean free path about equals the distance between filament and substrate. Indeed, a maximum in deposition rate is observed at this pressure. This corresponds to a value of  $p d_{fs}$  of 0.06 (cf. [530]). The microstructure parameter plotted as a function of  $p d_{fs}$  has a minimum around a value of  $p d_{fs}$  of  $0.6 \pm 0.2$  [530].

The silane flow rate determines to a large extent the possible deposition rate. If the supply of silane is too low, the environment in the direct vicinity of the filament will become depleted of silane. The generation of deposition precursors (and also the deposition rate) is limited by the silane supply. Above a certain flow, the dissociation reaction rate is the limiting factor, and as a consequence the deposition rate will remain constant upon a further increase of the silane flow.

Dilution of silane with hydrogen to moderate amounts ( $0.75 < [\text{SiH}_4]/([\text{SiH}_4]+[\text{H}_2]) < 1$ ) is not expected to have a large influence on material properties, as already in the pure silane case a large amount of atomic hydrogen is present. Every  $\text{SiH}_4$  molecule dissociates to give 4 H atoms [525]. Also, the mean free path of the atomic H is larger than the one of Si, i.e. about 8 cm at 0.02 mbar [531], and consequently all hydrogen will reach the substrate. It is found that the deposition rate depends linearly on the dilution fraction as it depends on the partial pressure of silane. Significant changes in the refractive index, the hydrogen content, the microstructure parameter, and electrical properties are observed for values of  $[\text{SiH}_4]/([\text{SiH}_4]+[\text{H}_2])$  lower than about 0.3-0.4. A closer look to this data reveals that moderate hydrogen dilution slightly improves the dark conductivity and photoresponse, whereas dilution fractions smaller than 0.3-0.4 lead to deterioration of the material. Similar trends were also observed by Bauer *et al.* [550] and Molenbroek *et al.* [528,530].

The common linear dependence of the band gap on the hydrogen content as is observed for PECVD deposited material is not observed for HWCVD deposited material. A large dataset is shown in Fig. 69, in which the relation between band gap and hydrogen content is compared for HWCVD and PECVD deposited material [531]. It is seen that the band gap of HWCVD deposited material follows a similar behaviour as a function of hydrogen content compared to the one of PECVD deposited material, for values of the hydrogen content of 7 % and higher: the band gap varies in a range between 1.7 and 1.85 eV [531]. The band gap of PECVD deposited materials decreases down to 1.5 eV at a hydrogen concentration of 2 %, whereas the band gap of HWCVD deposited material remains unchanged at 1.7 eV, even at hydrogen concentrations of 1 %. This difference in behaviour is due to the difference in the amount of voids present in the material. A large number of voids reduces the effective density of the material, and increases the average Si-Si bond distance. As a result, the band gap remains high. Crandall *et al.* [71] found that the amount of voids increased with decreasing hydrogen content in HWCVD deposited material. This observation is also supported by Small Angle X-ray Scattering (SAXS) [551] and Nuclear Magnetic Resonance (NMR) [552] data.

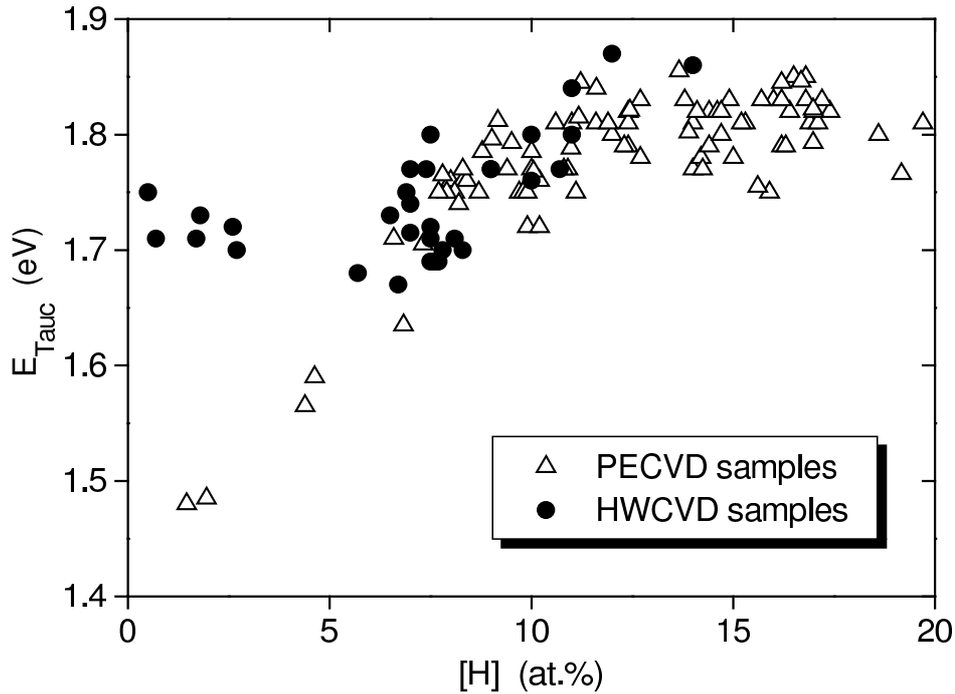


FIG. 69. Relation between the optical band gap (Tauc convention) and the hydrogen content in  $a$ -Si:H deposited by HWCVD (closed circles) and PECVD (open triangles). (From K. F. Feenstra, Ph. D. Thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998).

#### IX.D. Deposition model

The deposition mechanism in HWCVD of  $a$ -Si:H can be divided in three spatially separated processes. First, silane is decomposed at the tungsten filament. Second, during the diffusion of the generated radicals (Si, H) from the filament to the substrate, these radicals react with other gas molecules and radicals, and new species will be formed. Third, these species arrive at the substrate and contribute to the deposition of  $a$ -Si:H.

The filament material acts as a catalyser in the silane decomposition process. This is clear by comparing the Si-H bond energy (3-4 eV) in silane with the thermal energy at the filament temperature of 2000 °C (0.25 eV): they differ by an order of magnitude. The dissociation of silane proceeds in two steps [525]. First, the silane reacts with the tungsten to form a tungsten silicide  $W_xSi_y$  at the surface of the filament. Hydrogen atoms leave the surface before they can recombine to form molecular hydrogen. Subsequently, the silicon atoms evaporate from the surface, which is an activated process with an activation energy of 3.6 eV [525]. At a filament temperature of about 1700 °C the silane decomposition rate and the evaporation rate balance each other. For lower temperatures a tungsten silicon alloy is formed on the filament.

A fraction of the generated radicals react with gas molecules before they are able to reach the substrate directly. This fraction depends on the process pressure. A number of reactions may take place and only the exothermic ones are listed in Tab. 9, as endothermic reactions are not very likely to occur. As can be seen in Tab. 9, the Si and H radicals are highly reactive with silane. Most reactions will occur with the omnipresent silane. Quantum chemistry computations, confirming the findings of

TABLE 9. Exothermic gas phase reactions in the HWCVD process, their enthalpies, and reaction rates [192], estimated for typical operating conditions.

chemical reaction	enthalpy (eV)	reaction rate (cm <sup>3</sup> s <sup>-1</sup> )
H + SiH <sub>4</sub> → SiH <sub>3</sub> + H <sub>2</sub>	-0.54	2.68 × 10 <sup>-12</sup>
H + SiH <sub>3</sub> → SiH <sub>2</sub> + H <sub>2</sub>	-1.46	2 × 10 <sup>-11</sup>
SiH <sub>3</sub> + SiH <sub>3</sub> → SiH <sub>4</sub> + SiH <sub>2</sub>	-0.91	1.5 × 10 <sup>-10</sup>
Si + SiH <sub>4</sub> → Si <sub>2</sub> H <sub>2</sub> + H <sub>2</sub>	-0.34	3.5 × 10 <sup>-10</sup>
Si <sub>2</sub> H <sub>2</sub> + SiH <sub>4</sub> → Si <sub>3</sub> H <sub>4</sub> + H <sub>2</sub>	-0.4/-0.05	2 × 10 <sup>-10</sup>

Molenbroek *et al.* [530], have shown that the reaction of Si with SiH<sub>4</sub> proceeds via insertion of Si into the SiH<sub>4</sub> to form Si<sub>2</sub>H<sub>4</sub>, followed by a rearrangement of the thus formed species, and the elimination of H<sub>2</sub>, which yields Si<sub>2</sub>H<sub>2</sub> [553]. An increasing amount of SiH<sub>3</sub> will be created via the reaction H with SiH<sub>4</sub> upon increasing the pressure. Due to the SiH<sub>3</sub> homogeneous reaction SiH<sub>2</sub> is created, which at higher pressures may polymerize further to Si<sub>3</sub>H<sub>4</sub>. Holt *et al.* [554] have performed Monte Carlo simulations to study gas phase and kinetic processes in HWCVD. They showed that under the conditions for obtaining device quality material SiH<sub>3</sub> is the most abundant species at the substrate. At a pressure of about 0.08 mbar the fluxes of SiH<sub>3</sub>, Si<sub>2</sub>H<sub>2</sub>, and Si to the substrate were calculated to be 5 × 10<sup>17</sup>, 1 × 10<sup>16</sup>, and 5 × 10<sup>13</sup> cm<sup>-2</sup>s<sup>-1</sup>, respectively.

The HWCVD deposition process is more or less the same as for PECVD, and was described in Sec. VII. An important difference between the HWCVD and the PECVD process is the absence of ions, and the limited number of different species present in the gas phase. At low pressure atomic Si is the main precursor. This yields void-rich material with a high microstructure factor. Increasing the pressure allows gas phase reactions with Si and H to create more mobile deposition precursors (SiH<sub>3</sub>), which improves the material quality. A further increase leads to the formation of higher silanes, and consequently to a less dense film.

Using threshold ionization mass spectrometry and *in situ* ellipsometry, Schröder and Bauer [555] have shown that the Si<sub>2</sub>H<sub>4</sub> radical may well be the species responsible for deposition, rather than SiH<sub>3</sub>, as in PECVD. This larger and less mobile precursor is thought to be the cause of the observed differences in deposition conditions in HWCVD and PECVD to obtain device quality material.

## X. EXPANDING THERMAL PLASMA CHEMICAL VAPOUR DEPOSITION

### X.A. General description

The Expanding Thermal Plasma Chemical Vapour Deposition technique has been developed in the group of Schram [20,556] and has been used for the deposition of several materials, such as hydrogenated amorphous silicon [21] and carbon [557], but also for diamond [558]. The ETP-CVD technique is a remote-plasma deposition method. The generation of the plasma, the transport of the plasma and the deposition are geometrically separated. In remote-plasma CVD the substrate holder is not a necessary electrode for the plasma. The absence of direct plasma contact with the substrate allows for a better control of ion bombardment, which is advantageous. Also, the properties of the plasma can be varied independently, which makes optimization of the whole process simpler.

An argon/hydrogen plasma is created in a DC thermal arc (cascaded arc) operated at high pressure ( $\approx 0.5$  bar) [556,559,560] (the cascaded arc is also employed in IR ellipsometry, providing a well-defined source of intense IR radiation, see Sec. VD [343]). As the deposition chamber is at much lower pressure (0.1-0.3 mbar) a plasma jet is created, expanding into the deposition chamber. Near the plasma source silane is injected and the active plasma species dissociate the silane in radicals and ions. These species can deposit on the substrate that is positioned further downstream.

The main advantage of this technique is the very high deposition rate that can be obtained for the different materials. However, this high deposition rate may not always be compatible with good material quality. A large effort has been made and has resulted in the deposition of good quality *a*-Si:H at deposition rates as high as 10 nm/s [432,561,562].

### X.B. Apparatus, setup

The ETP-CVD set-up is schematically shown in Fig. 70. It consists of a thermal plasma source, a cascaded arc, and a low-pressure deposition chamber [563,564]. The cascaded arc consists of three cathodes, a stack of ten copper plates with a central bore of 4 mm, and an anode plate with a conical shaped nozzle [556]. The length of the cascade is about 10 cm. All parts are water-cooled. Pure argon or a argon/hydrogen mixture is introduced into the arc. The argon flow is varied between 1800 and 6000 sccm, and the hydrogen flow between 0 and 1200 sccm. The arc pressure is between 0.4 and 0.6 bar. The DC discharge is current controlled. The arc current and voltage range from 30-75 A and 45-120 V, respectively. Typical arc plasma parameters are a electron density and temperature of  $10^{16}$  cm<sup>-3</sup> and 1 eV [565], respectively.

The deposition chamber is a cylindrical vessel with a diameter of 50 cm and a length of 80 cm. At about 5 cm from the arc outlet silane can be introduced via an injection ring (7.5 cm diameter) that contains 8 holes of 1 mm diameter each. The distance between arc outlet and substrate amounts to 38 cm. The substrates are heated via the substrate holder, of which the temperature can be controlled between 100 and 500 °C. Samples can be loaded via a load lock equipped with a magnetic transfer arm. The

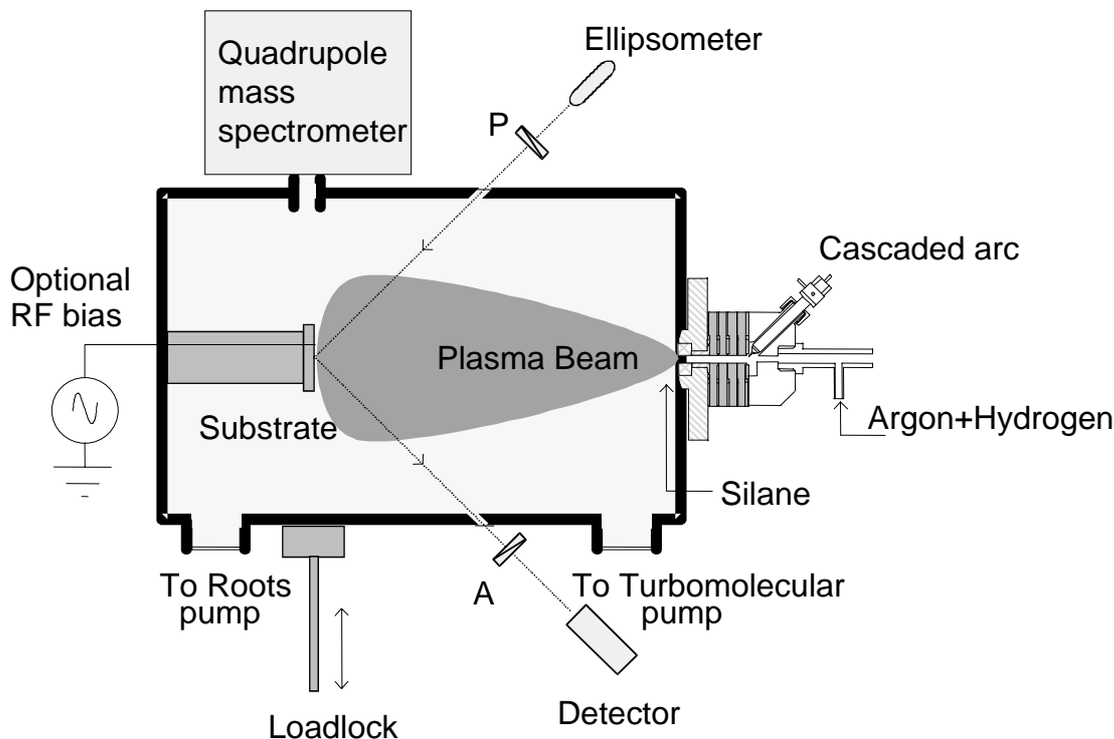


FIG. 70. Cross-sectional view of an expanding thermal plasma/ETP deposition reactor. (Courtesy of M.C.M. van de Sanden, Eindhoven University of Technology).

substrate can be optionally RF biased. A residual gas analyzer (QMS) and an ellipsometer complete the set-up. The typical pressure is in the range of 0.15-0.5 mbar. The deposition chamber has a volume of 180 liter. During processing it is pumped by a stack of two roots blowers and one fore-pump (total pumping capacity is about 1500 m<sup>3</sup>/h), otherwise it is pumped by a turbo pump (450 l/s) with which a base pressure of 10<sup>-6</sup> mbar is reached.

As a result of the large pressure gradient between arc and deposition chamber the plasma expands supersonically from the nozzle into the deposition chamber. After a stationary shock front at about 5 cm from the nozzle, the plasma expands subsonically. As a consequence, the electron density and temperature are drastically reduced [566]. In this downstream region the plasma is recombining, much like in an afterglow plasma. The low electron temperature in the region where the silane is introduced implies that electron-induced reactions are ineffective for the ionization and dissociation of silane. It is found that the admixture of H<sub>2</sub> to Ar in the arc is the cause for the strong reduction in the ion and electron density [567]. For pure Ar, the ion density is 10<sup>13</sup> cm<sup>-3</sup>, and exists mainly of Ar<sup>+</sup>. At high H<sub>2</sub> flows (1800 sccm), the ion density is about 4×10<sup>10</sup> cm<sup>-3</sup>, and mainly is H<sup>+</sup>. At low hydrogen flows, ArH<sup>+</sup> is the most abundant ion. The reason for this drastic change in density is the fact that fast molecular processes become an important recombination channel for the ions [560,568]. The atomic H concentration increases from about 2×10<sup>11</sup> cm<sup>-3</sup> at low (240 sccm) H<sub>2</sub> flow to about 2×10<sup>12</sup> cm<sup>-3</sup> at high (1800 sccm) H<sub>2</sub> flow. The arc changes basically from an argon ion source to an

atomic hydrogen source, when going from pure argon to admixture of a high amount of hydrogen.

### X.C. Material properties and deposition conditions

The deposition rate is found to be independent on the deposition temperature, for low (0.3 nm/s) to high (30 nm/s) rates [473]. The deposition rate and the silane consumption increase with increasing partial pressure of silane [569]. A higher arc current yields a higher silane depletion and deposition rate [563]. The bond-angle distortion, as determined from Raman spectroscopy measurements varied between 8.3° at low deposition rate to 9.1° at high deposition rate [570], indicating a slightly increased disorder.

The incorporated amount of hydrogen depends on the deposition temperature. The hydrogen content decreases with increasing temperature. For films deposited at a high deposition rate, the hydrogen content always is higher compared to the films that were deposited at a low deposition rate. The hydrogen content varies from 60 at.% at a deposition temperature of 100 °C to 10 at.% at a deposition temperature of 400 °C for high-rate deposited films. The corresponding values for low-rate deposited films are 22 at.% (100 °C) and 5 at.% (400 °C). The microstructure factor also decreases with increasing temperature. The refractive index increases with increasing temperature [468]. Both dark- and photoconductivity show an increase with increasing temperature, and have a maximum at 400-450 °C [571] of  $4 \times 10^{-6} \Omega^{-1} \text{cm}^{-1}$  and  $10^{-9} \Omega^{-1} \text{cm}^{-1}$ , respectively.

The deposition rate and the silane consumption decreases sharply by a factor of about four upon adding 60-120 sccm hydrogen to pure argon in the arc, as in shown in Fig. 71a. Further addition of hydrogen does not influence these parameters much [569,572]. Similar effects are observed for the refractive index, microstructure factor, activation energy, and dark- and photoconductivity, i.e. adding a small amount hydrogen leads to a large change, which is saturated upon further hydrogen admixture.

Using ion and threshold ionization mass spectrometry (TIMS), Langmuir probe measurements, and cavity ring down absorption spectroscopy (CRDS) the influence of hydrogen admixture on the film growth precursors has been investigated [567,569,573]. For the argon/hydrogen plasma, it was found that electron temperature and ion fluence decrease as a function of hydrogen flow. At low hydrogen flow electron temperature amounts to 0.3 eV, whereas at high hydrogen flow it is 0.1 eV. The ion fluence is decreased from about 120 sccm at low hydrogen flow to about 4 sccm at high flow.

Adding silane to the argon/hydrogen plasma leads to the formation of large and hydrogen-poor positive silicon ions [567]. With increasing flow the cluster size increases, in fact the higher silane clusters  $\text{Si}_n\text{H}_m^+$  ( $n > 5$ ) are much more abundant than the lower silane clusters. The total ion flux towards the substrate scales linearly with the silane flow, while it decreases with increasing hydrogen flow. The maximum contribution of the cationic clusters to film deposition ranges from 4-7 % for varying silane flows, and from 5-9% for varying hydrogen flows (Fig. 71b), and is rather independent of both the silane as well as the hydrogen flow [567].

As a consequence, the contribution of neutral species to the deposition is more than 90%, and is built up mainly from  $\text{SiH}_3$ . This contribution first increases at low hydrogen

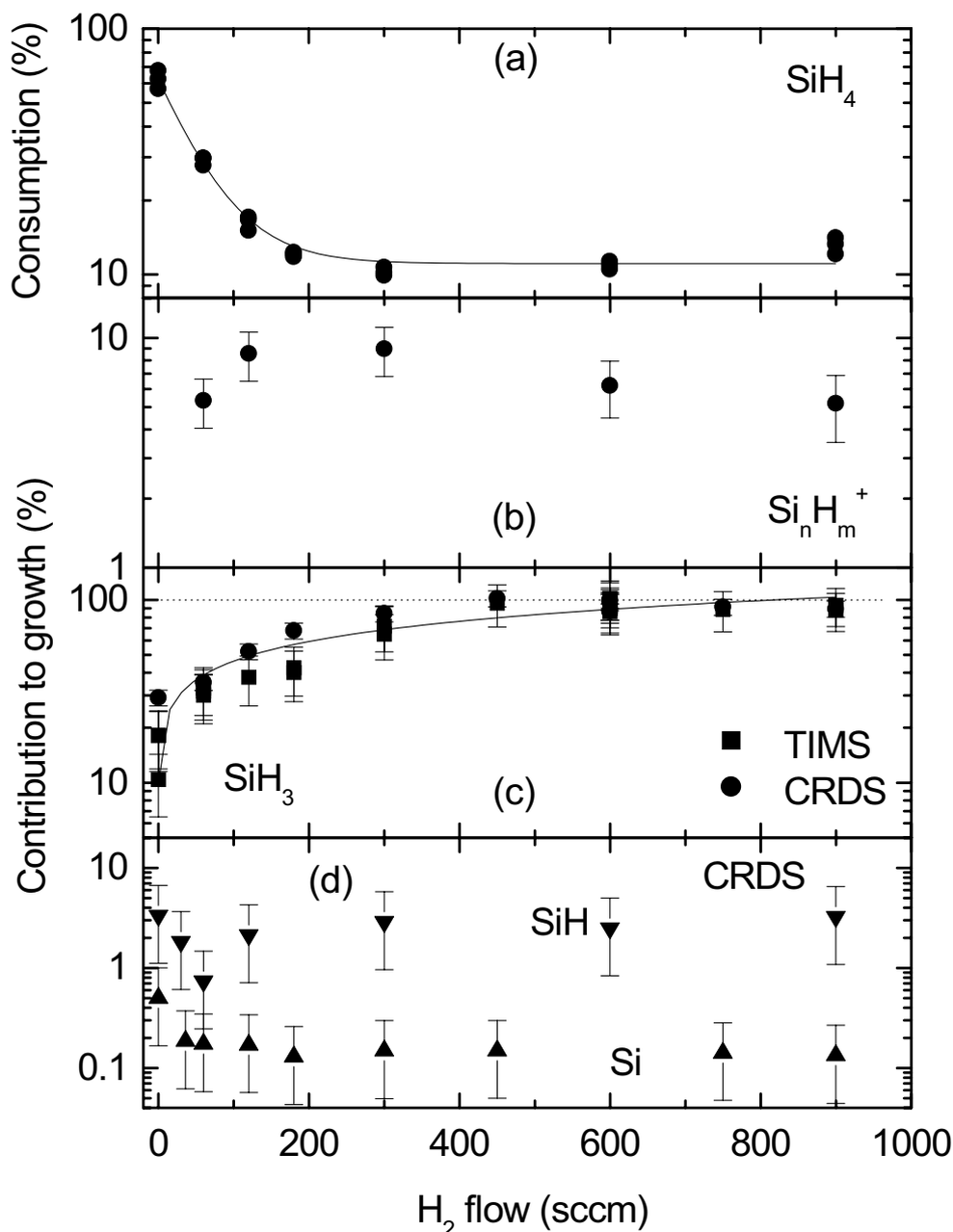


FIG. 71. The influence of hydrogen admixture on (a) the silane consumption, and the contribution of  $\text{Si}_n\text{H}_m^+$  (b),  $\text{SiH}_3$  (c),  $\text{SiH}$  (d), and  $\text{Si}$  (d) to the deposition, as measured by TIMS and CRDS. (From W. M. M. Kessels, Ph. D. Thesis, Technische Universiteit Eindhoven, Eindhoven, the Netherlands, 2000).

flow from about 20% to about 90% at intermediate hydrogen flow, and remains constant with further increase of the hydrogen flow (Fig. 71c). A contribution of  $\text{SiH}_2$  larger than 0.1% could only be measured at zero hydrogen flow, and amounted to about 5%. Other radicals, i.e.,  $\text{SiH}$  and  $\text{Si}$ , contribute about 2% and 0.2%, respectively, to the deposition (Fig. 71d), irrespective of hydrogen flow.

The optimized conditions for the deposition of good quality  $a\text{-Si:H}$  have been found to be: an argon flow of 3300 sccm, a hydrogen flow of 600 sccm, a silane flow of 600

sccm, an arc current of 45 A, an arc voltage of 140 V, an arc pressure of 0.4 bar, a process pressure of 0.2 mbar in the downstream region, and a substrate temperature of 400 °C [571,572]. At this high deposition temperature the hydrogen content is 6-7 at.%. Consequently the (cubic) band gap is low, 1.51 eV. Also the microstructure factor still is non-zero, i.e. 0.2, and especially the dark conductivity is a factor of 10 higher than needed for device quality material. The electron drift mobility is somewhat smaller than usual for device quality films, but the hole drift mobility is one order of magnitude larger. The activation energy, Urbach energy and defect density are 0.75 eV, 50-55 meV, and  $10^{16}\text{cm}^{-3}$ .

#### X.D. Deposition model

Kessels has summarized the information on the reactive species emanating from the Ar-H<sub>2</sub> operated cascaded arc [571], and has formulated a global reaction scheme. For a hydrogen flow between 0 and 120 sccm, the dissociation of silane is governed by dissociative charge transfer of Ar<sup>+</sup> with SiH<sub>4</sub>, which generates SiH<sub>*n*</sub><sup>+</sup> (*n* ≤ 3) ions: Ar<sup>+</sup> + SiH<sub>4</sub> → Ar + SiH<sub>*n*</sub><sup>+</sup> + ... Then, as the electron density is high, these reactions are quickly followed by dissociative recombination reactions with electrons, which generates SiH<sub>*m*</sub><sup>+</sup> (*m* ≤ 2) radicals: SiH<sub>*n*</sub><sup>+</sup> + *e* → SiH<sub>*m*</sub> + ... These highly reactive radicals react with silane molecules forming polysilane radicals: SiH<sub>*m*</sub> + SiH<sub>4</sub> → Si<sub>2</sub>H<sub>*p*</sub> + ... When the electron density is decreased as a result of the dissociative recombination reactions, ion-molecule reactions between silane ions and silane will become significant as well. This reaction (SiH<sub>*n*</sub><sup>+</sup> + SiH<sub>4</sub> → Si<sub>2</sub>H<sub>*m*</sub><sup>+</sup> + ...) leads to sequential ion-SiH<sub>4</sub> reactions (Si<sub>*p*</sub>H<sub>*q*</sub><sup>+</sup> + SiH<sub>4</sub> → Si<sub>*p+1*</sub>H<sub>*r*</sub><sup>+</sup> + ...) which generate the cationic clusters (up to Si<sub>10</sub>H<sub>*r*</sub><sup>+</sup> has been observed).

At a hydrogen flow larger than 300 sccm the ion fluence from the arc is greatly reduced. Reactions with atomic H emanating from the arc now are more effective. The dissociation of silane is governed by hydrogen abstraction, which generates the SiH<sub>3</sub> radical (H + SiH<sub>4</sub> → H<sub>2</sub> + SiH<sub>3</sub>). The small flow of H<sup>+</sup> from the arc is responsible for a charge exchange reaction with silane, that creates SiH<sub>*n*</sub><sup>+</sup> (*n* ≤ 3), which initiates sequential ion-molecule reactions as above.

The deposition of good quality *a*-Si:H using ETP-CVD is a result of the dominant presence of the SiH<sub>3</sub> radical under conditions of high hydrogen flow in the cascaded arc. The deposition process itself is more or less the same as for PECVD, and was described in Sec. VII. An important difference between the ETP-CVD and the PECVD process is the absence of ions with considerable energy (> 1 eV), and the amount of cationic clusters. Nevertheless, good quality *a*-Si:H is deposited at high deposition rates.

## XI. APPLICATIONS

The first practical device demonstrating the use of *a*-Si:H as a photovoltaic material was the 2.4% efficient solar cell reported by Carlson and Wronski [34]. Since then, interest in *a*-Si:H has been growing rapidly, more or less prompted by the many possible applications. Based on the photovoltaic effect *a*-Si:H is used in solar cells and optical sensors. Especially the possibility of uniform large-area deposition has been exploited in *a*-Si:H thin film transistors (TFT) that are used in controlling active-matrix liquid crystal displays (LCD). The high photoconductivity and fast photoresponse of *a*-Si:H is of great importance for use in large-size linear image sensors.

A selection of applications is presented in the following sections, i.e., solar cells, thin film transistors, light sensors (visible, IR, X-ray), and chemical sensors. Also light-emitting devices, in particular due to erbium incorporation in *a*-Si:H are presented. Finally, electrostatic loudspeakers in which an *a*-Si:H film is incorporated are described. Details of various applications described here, as well as many other applications, can be found in excellent (edited) books [4,5,11,13,574].

### XI.A. Solar Cells

#### XI.A.1. Operation principle

The operation principle of a solar cell is based on charge separation. Photogenerated electrons and holes must be spatially separated in order to contribute to the net current of the device. Charge separation is done by an internal electric field. In crystalline semiconductors this can be achieved by stacking a *p*-type doped material on an *n*-type doped material. The internal field at the *p-n* junction prevents the recombination of photogenerated electrons and holes. Once separated, however, recombination may occur, and is an important loss process in a solar cell [575]. In amorphous semiconductors a *p-n* junction hardly shows photovoltaic action. Photogenerated electrons and holes cannot diffuse over long distances, as the defect density in *p* and *n*-type doped material is high. Therefore an undoped (intrinsic) layer is to be introduced between *p* and *n* layer, with a thickness smaller than the mean free path of the slowest carriers, i.e. holes.

In the presence of an electric field the drift length is the product of the mobility-lifetime product and the electric field:  $\lambda_{mfp} = \mu\tau E$  [576]. With typical values of  $\mu\tau$  and  $E$  the mean free path usually by far exceeds the thickness of the solar cell, and virtually all photogenerated carriers can be collected. However, under certain operating conditions, field-free regions in the *i*-layer may exist, and the collection efficiency is decreased due to the fact that the diffusion lengths of the carriers is much smaller than the thickness of the solar cell [11,577].

The *p* and *n* layers provide the built-in potential and do not contribute to the collection of carriers, Therefore these layers need to be as thick as the depletion layer thickness (5-20 nm), as any additional thickness unnecessarily will decrease the collection efficiency by absorbing light. The intrinsic layer should be as thick as possible to absorb the maximum amount of photons, but the depletion width at operating conditions of about 1  $\mu\text{m}$  sets an upper limit. Typically, one uses a thickness of 500 nm.

A schematic cross-section of a *p-i-n* *a*-Si:H solar cell [11] is shown in Fig. 72a. In this so-called superstrate configuration (the light is incident from above) the material

onto which the solar cell structure is deposited, usually glass, also serves as a window to the cell. In a substrate configuration the carrier onto which the solar cell structure is deposited forms the back side of the solar cell. The carrier usually is stainless steel, but flexible materials such as metal-coated polymer foil (e.g. polyimide) or a very thin metal make the whole structure flexible [11].

The current-voltage ( $I - V$ ) characteristics of a solar cell is given by

$$I(V) = I_{ph}(V) - I_0 \left[ \exp \left( \frac{e(V + I(V)R_s)}{Ak_B T} \right) - 1 \right] - \frac{V + I(V)R_s}{R_{sh}} \quad (69)$$

where  $I(V)$  is the current  $I$  at voltage  $V$ ,  $I_{ph}(V)$  the photogenerated current,  $I_0$  the reverse saturation current,  $A$  the diode ideality factor,  $R_s$  the series resistance, and  $R_{sh}$  the shunt resistance parallel to the junction. Usually the current density  $J(V)$  rather than the current  $I(V)$  is shown, e.g., see Fig. 60. An  $I - V$  measurement is performed by connecting a variable load or by applying an external voltage [578], in the dark or under illumination with an AM1.5 spectrum. Four parameters determine the performance of a solar cell, i.e., the efficiency  $\eta$ , the fill factor  $FF$ , the open circuit voltage  $V_{oc}$  and the short circuit current  $I_{sc}$ . The efficiency is defined as the ratio of the maximum delivered power and the power of the incident light. The maximum power point is the point where  $I_{max}V_{max}$  has the maximum value. The open circuit voltage  $V_{oc}$  and the short circuit current  $I_{sc}$  are defined as the values of the voltage and current at  $I(V) = 0$  and  $V = 0$ , respectively. The fill factor  $FF$  is defined as the ratio of the maximum power and the product of  $I_{sc}$  and  $V_{oc}$ . Typical values of these parameters were shown in Fig. 60. A characteristic resistance  $R_{ch}$  is sometimes defined as  $V_{oc}/I_{sc}$  in order to assess the effect of  $R_s$  and  $R_{sh}$  on the fill factor. When  $R_s \ll R_{ch}$  or  $R_{sh} \gg R_{ch}$  little effect is seen on the fill factor [575].

The short circuit current is the product of the photon flux  $\Phi(\lambda)$  of the incident solar spectrum and the wavelength-dependent spectral response or collection efficiency  $Q(\lambda)$  integrated over all wavelengths:  $J_{sc} = \int \Phi(\lambda)Q(\lambda)d\lambda$  (see Fig. 61b). The collection efficiency is about 80% between 450 and 600 nm, demonstrating that there is little loss due to recombination (the  $i$ -layer is of 'device quality'). The decreasing collection efficiency at the red side is due to the decreasing absorption coefficient of  $a$ -Si:H. In the blue, the decreasing collection efficiency is due to absorption in the  $p$ -layer and/or buffer layer.

### XI.A.2. Single junction solar cells

A schematic cross-section of a single-junction  $p$ - $i$ - $n$   $a$ -Si:H solar cell in the superstrate configuration was shown in Fig. 72a. The front electrode consists of a transparent conductive oxide (TCO). Examples are tin oxide ( $\text{SnO}_2:\text{F}$ ), indium tin oxide (ITO,  $\text{In}_2\text{O}_3:\text{Sn}$ ), or doped zinc oxide ( $\text{ZnO}:\text{Al}$ ) [579]. A high transparency in the spectral range of the solar cell is required, as well as a low sheet resistance, to minimize series resistance losses. Further, the contact resistance with the  $p$ -layer should be low. The surface of the TCO should be textured as a means to reduce optical reflection and to increase the optical path length by scattering, thereby increasing the absorption in the active  $i$  layer [577,580,581]. Shunting paths and pinholes should be avoided by a proper morphology. The TCO should also be PECVD compatible, i.e., it should be highly resistant against the reducing H ambient during PECVD deposition.

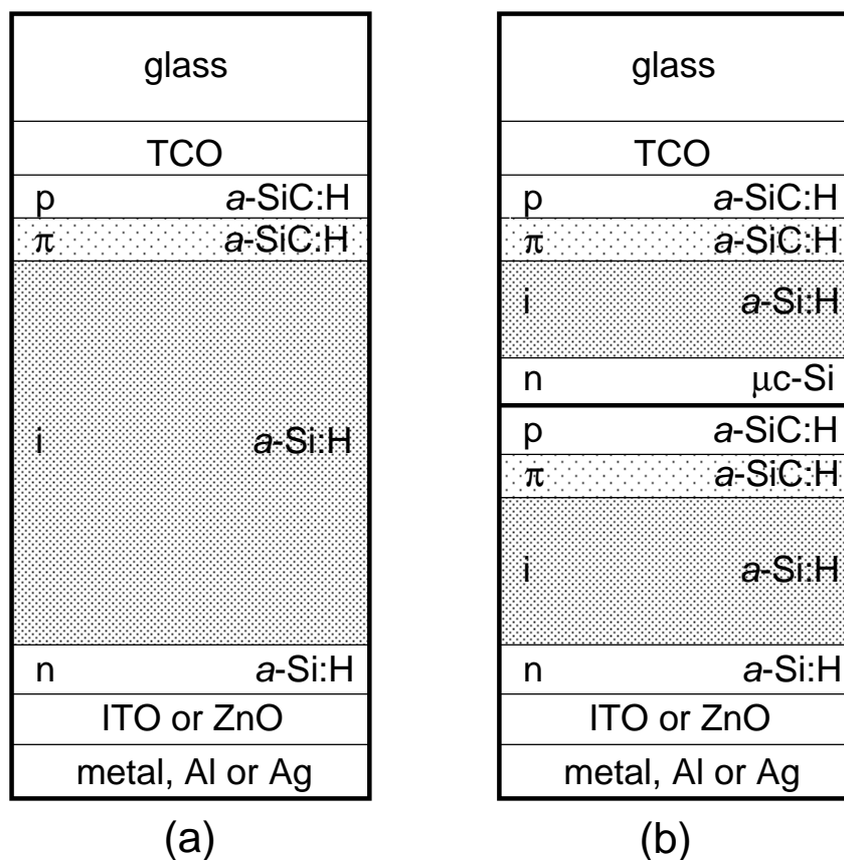


FIG. 72. Schematic cross-section of (a) a single junction  $p-i-n$   $a\text{-Si:H}$  superstrate solar cell and (b) a tandem solar cell structure. (From R. E. I. Schropp and M. Zeman, *Amorphous and Microcrystalline Silicon Solar Cells - Modeling, Materials and Device Technology* (Kluwer Academic Publishers, Boston, MA, U.S.A., 1998).).

ITO,  $\text{SnO}_2\text{:F}$ , and  $\text{ZnO:Al}$  share a high transparency ( $> 90\%$ ), a high bandgap ( $> 3.5$  eV), and a low sheet resistance ( $< 10 \Omega/\square$ ). Tin oxide is made by atmospheric CVD which inherently produces a native texture. Zinc oxide and ITO, both made by sputtering or evaporation, usually are flat. Zinc oxide can be etched by HCl to produce a rough surface [11]. Both tin oxide and zinc oxide are chemically resistant. ITO can be made chemically resistant by adding a thin titanium oxide layer as a protective coating [582]. Zinc oxide has a high contact resistance with doped  $a\text{-Si:H}$ , which can be circumvented by using a microcrystalline doped layer [583].

Absorption in the  $p$ -layer can be reduced by using a  $a\text{-SiC:H}$  alloy with a band gap of about 2 eV [584]. Carbon profiling within the  $p$ -layer further improves the window properties [585]. An intentionally graded  $p/i$  interface ('buffer layer') of 10 nm in thickness enhances the spectral response in the blue [125,494,586], which can be attributed to a reduced interface recombination.

The properties of the  $i/n$  interface and the  $n$ -layer are not that critical, as long as the conductivity is high. Introducing a TCO layer between  $n$ -layer and metal back contact enhances even further the light trapping. Absorption mainly is enhanced in the long wavelength region [587]. The optimum thickness is found to be 70 nm [588].

In the substrate configuration the stainless steel carrier is coated with a Ag/ZnO bilayer in order to enhance the back reflection of the back contact, see Fig. 73 [11]. An increase in  $J_{sc}$  of about 50% was achieved by Banerjee and Guha [589] by using a textured Ag/ZnO bilayer, that further enhances the optical path length, and consequently the absorption. As at this stage no  $a$ -Si:H has been deposited there are virtually no restrictions on process temperature.

After depositing the  $n$ -layer a similar  $i/n$  buffer layer can be deposited as in  $p-i-n$  cells. Band gap profiling, i.e., increasing the band gap towards the top  $p$ -layer, is much easier for  $n-i-p$  structures, as here simply decreasing the deposition temperature during deposition automatically yields a higher band gap [11]. In  $p-i-n$  cells deposition of the  $i$ -layer is from the  $p$ - to the  $n$ -layer. In this case, a decrease in the band gap is achieved by increasing the temperature during deposition, which may not always be desirable for the already deposited  $p$ -layer and  $p/i$  interface. Another advantage is that the most critical top layers are deposited the last.

The transparent top contact is deposited last, which imposes restrictions to the process temperature. Thermally evaporated ITO and ZnO deposited by Metal Organic CVD (MOCVD) are most suitable. At a typical thickness 70 nm the ITO serves as a good antireflection coating as well. Due to the somewhat high sheet resistance, a metal (Ag) grid is necessary to reduce the series resistance [11].

### XI.A.3. Multiple junction solar cells

The performance of  $a$ -Si:H can be improved by stacking two (or more) cells with a different optical band gap on top of each other, see Fig. 72b [11]. In such a tandem solar cell, the higher band gap material of which the top cell is made, collects the high energy photons, while the lower energy photons are collected in the lower cell. While theoretical efficiencies of single junction  $a$ -Si:H solar cells are around 15%, values above 20% are calculated for multijunction cells. A two-terminal (top- and bottom contact) double-junction cell consisting of a top cell with a band gap of 1.75 eV and a bottom cell with a band gap of 1.15 eV is predicted to have a conversion efficiency of 21%, while a triple stack of 2.0 eV/1.7 eV/1.45 eV cells is predicted to have an efficiency of 24% [590]. In the latter cell carbon alloying of  $a$ -Si:H would make the top cell, and germanium alloying the bottom cell. However, due to high defect densities present in  $a$ -SiC:H and  $a$ -SiGe:H other ways of obtaining good quality high and low band gap material were developed. Dilution of  $\text{SiH}_4$  with hydrogen in the regime  $0.05 \leq [\text{SiH}_4]/([\text{SiH}_4]+[\text{H}_2]) \leq 0.15$  yields material of better optoelectronic properties than  $a$ -SiC:H [369], and is therefore used for the top cell (Fig. 72b).

Hydrogen dilution also improves the quality of  $a$ -SiGe:H alloys [369]. A  $a$ -SiGe:H bottom cell must be designed to operate at long wavelengths, with relatively weakly absorbed light. The concept of band gap grading at the  $p/i$  interface as in  $a$ -Si:H cells was demonstrated to be inappropriate for  $a$ -SiGe:H cells [591]. Several band gap profiles were investigated. The best solar cell performance was obtained for a band gap that first decreased but then increased, see also Zimmer *et al.* [592].

A triple junction structure based on 1.79 eV, 1.55 eV, and 1.39 eV materials [593] was reported to have an initial efficiency of 14.6% and a stabilized (see Sec. XI A 4) efficiency of 12.1% [154]. In these series-connected structures the open circuit voltage

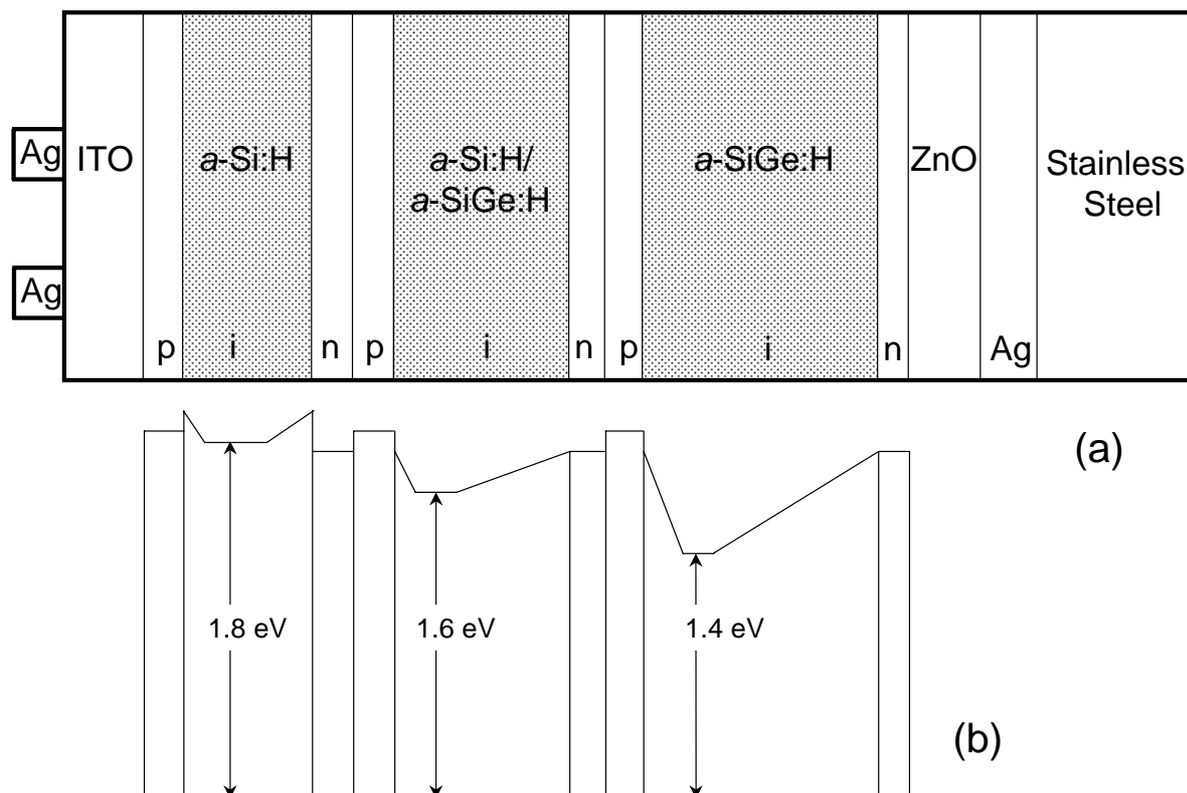


FIG. 73. Schematic cross-section of a triple junction  $a$ -Si:H substrate solar cell on stainless steel (a), and the corresponding schematic band diagram (b). (From R. E. I. Schropp and M. Zeman, *Amorphous and Microcrystalline Silicon Solar Cells - Modeling, Materials and Device Technology* (Kluwer Academic Publishers, Boston, MA, U.S.A., 1998).).

is the sum of the  $V_{oc}$ 's of the individual cells (here  $V_{oc} = 2.3$  V), while the currents are matched to be equal in all cells:  $J_{sc} = 7.56$  mA/cm<sup>2</sup>. The fill factor is = 0.7 [154]. A schematic cross-section of such a triple-junction  $p$ - $i$ - $n$   $a$ -Si(Ge):H solar cell structure in the substrate configuration is shown in Fig. 73 [11], together with the corresponding band diagram. This band diagram shows the intricate band gap profiling scheme that is needed obtain these high efficiency solar cells.

A critical part in the structure of multi junction solar cells is the  $n$ - $p$  junction that divides the individual cells. Here, electrons that are generated in the top cell (Fig. 72b) flow to the  $n$ - $p$  junction, where they must recombine with holes from the bottom cell. As a result of the very high doping levels used, recombination takes place via tunneling. When the recombination rate is not balanced with the supply of carriers, space charge accumulation will occur, which negatively influences the electric field in the adjacent cell that has the highest generation rate [11]. Often one of the layers of the tunnel junction is microcrystalline [594].

Another important issue in multi junction cells is current matching. The individual currents must exactly balance, otherwise a loss in efficiency occurs. A current mismatch can be easily revealed by measuring the spectral response [595]. If the currents are matched, then the quantum efficiency is flat over a wide range of wavelengths. If one

of the cells is limiting the current, then the observed quantum efficiency is not flat, and in fact is the quantum efficiency of the current-limiting cell.

Empirical optimization of the thicknesses of the individual cells within the structure is to be combined with computer modeling. A comprehensive model is required in which an optical and electrical model of *a*-Si:H are integrated [11]. Since such a model contains about 100 input parameters for a single junction solar cell a careful calibration procedure is needed to extract input parameters from measured layer properties [596,597]. Current matching in tandem cells was investigated using a model which not only well describes recombination via tunneling but also takes into account the optical enhancement due to the textured TCO [598]. It was found that for an *a*-Si:H/*a*-Si:H stack the optimum thickness of the bottom cell amounted to 300 nm, while for an *a*-Si:H/*a*-SiGe:H stack the optimum thickness of the bottom cell amounted to 150 nm. The optimum thickness of the top cell were 50 and 60 nm for the *a*-Si:H/*a*-Si:H and *a*-Si:H/*a*-SiGe:H stack, respectively [11,598].

#### XI.A.4. Stability

Illumination of solar cells causes a reduction of efficiency and fill factor, as a result of light-induced creation of defects (Staebler-Wronski effect, Sec. I B 5). This reduction is halted after several hundreds of hours illumination time. The reduction is correlated with solar cell thickness. A large intrinsic layer thickness leads to a large reduction of efficiency and fill factor compared to a small intrinsic layer thickness. The solar cell properties can be completely recovered by annealing at 150 °C. The open circuit voltage and short circuit current decrease only slightly.

Material properties cannot always be correlated with degradation behaviour [493,495,599,600]. Lee *et al.* [600] have shown the degradation kinetics of cells in which the intrinsic layer was deposited by highly diluting the silane. A clear correlation was observed between the decrease in efficiency and the increase of the  $\mu\tau$  product. Another example that demonstrates a correlation between degradation behaviour and material quality was shown in Fig. 62. Here the normalized cell efficiency as a function of illumination time was depicted for solar cells, where the intrinsic layer is deposited at different discharge power levels [493]. At the lower power levels the efficiency stabilizes at about 60% of its initial value, whereas at the higher power levels stabilization at 40% occurs. This has been attributed to the fact that at these higher power densities the density of the material is lower and the hydrogen content is higher compared to low power densities.

As a result of the creation of defects trapping of electrons or holes is enhanced. Thus, the created defects reduce the product  $\mu\tau$ , which suppresses charge collection and solar cell efficiency. The charge collection length, i.e., the average distance that a carrier travels before it is trapped, is defined as  $d_c = \mu\tau V_{bi}/d$ , with  $V_{bi}$  the built-in voltage (typically around 1.2 V [3]) and  $d$  the thickness of the cell. An empirical relationship between fill factor and charge collection length  $d_c$  was reported by Faughnan and Crandall [601]:

$$FF = 0.35 + 0.15 \ln \left( \frac{d_c}{d} \right) \quad (70)$$

Smith *et al.* [602] have derived the dependence of  $FF$  on illumination time. They combined the empirical relationship between  $FF$  and  $d_c$  (Eq. (70)) with the time dependence of the defect density ( $N_{db}(t) \sim G^{0.6}t^{1/3}$  [89]) and the relation between  $\mu\tau$  and defect density ( $N_{db} \sim 1/\mu\tau$  [3]). They arrived at

$$FF(t) = FF_i - \frac{k}{3} \log\left(\frac{t}{t_i}\right) \quad (71)$$

where  $FF_i$  is the initial  $FF$ ,  $k$  a kinetic constant for degradation, and  $t_i$  the initial time. Their data were fitted with  $FF_i = 0.68$ , and  $k = 0.25$ .

Catalano *et al.* [603] have introduced the device thickness in the empirical time-dependence of the efficiency:

$$\eta = \eta_i \left[ 1.1 - K \log\left(\frac{t}{t_i}\right) \right] \quad (72)$$

where  $\eta_i$  is the initial efficiency, and  $K$  a rate constant. The removal of defects was found to be a temperature activated process [604] represented by  $K = K' \exp(E_A/k_B T)$ , where  $K'$  is a constant proportional to thickness ( $\propto d^{0.54}$ ), and  $E_A$  an activation energy (0.2 eV [603]). At 35 °C Eq. (72) can be rewritten as [603]:

$$\eta = \eta_i \left[ 1.1 - 0.165d^{0.54} \log\left(\frac{t}{t_i}\right) \right] \quad (73)$$

From the proportionality of the rate constant with thickness it is clear that thick cells will degrade deeper than thin cells. However, due to the fact that the initial efficiency increases with thickness an optimum thickness of 200-300 nm is found [577].

The degradation of solar cell properties can be circumvented by proper device design [11]. The parameter of interest here is the electric field profile after degradation, which should be optimized for carrier collection. A thinner intrinsic layer combined with enhanced light confinement leads to a higher electric field after degradation. Also band gap profiling may assist the carrier transport in the low-field region that is present in the intrinsic layer after degradation [605]. Further hydrogen dilution during deposition of the intrinsic layer has been reported to improve the stability [606].

Individual cells in multi junction cells are more stable due to their reduced thickness compared to single junction cells. Moreover, as the amount of light that is absorbed in the bottom cell is reduced and as the degradation rate is reduced at lower light levels, the stability is improved [606]. Therefore the multi junction cell is expected to be more stable than a single junction cell of the same thickness, which indeed has been demonstrated [577,606–608].

Outdoor operating temperatures of solar cells are around 60°C. At this temperature already significant annealing of defects occurs [606], and the stability is better than at 25°. Interestingly, seasonal effects on the performance of solar cells show that the efficiency is higher in summer than in winter [609], which was attributed to annealing of defects [610]. However, it was reported that these seasonal effects are due to the spectral differences in summer and winter, rather than the increased operating temperature in summer compared to winter [611,612].

The research effort in many laboratories around the world on the optimization of laboratory-scale (1 cm<sup>2</sup> area) cells has lead to the development of large-area (1 m<sup>2</sup>) solar cell modules with an efficiency exceeding 10% [613,614]. In a module individual cells can be connected in series or parallel, depending on the desired output voltage and current. Details on this topic can be found elsewhere, e.g., [11,157,577,607,615,616].

## XI.B. Thin Film Transistors

### XI.B.1. Operation principle

The most common *a*-Si:H TFT structure is the so-called inverted staggered transistor structure [40], in which silicon nitride is used as the gate insulator. A schematic cross-section is shown in Fig. 74. The structure comprises an *a*-Si:H channel, a gate dielectric ( $\text{SiN}_x$ ) and source, drain, and gate contacts.

The operation principle of these TFTs is identical to that of the metal-oxide semiconductor field-effect transistor (MOSFET) [617,618]. When a positive voltage  $V_G$  is applied to the gate, electrons are accumulated in the *a*-Si:H. At small voltages these electrons will be localized in the deep states of the *a*-Si:H. The conduction and valence band at the  $\text{SiN}_x$ /*a*-Si:H interface bend down, and the Fermi level shifts upward. Above a certain threshold voltage  $V_{th}$  a constant proportion of the electrons will be mobile, and the conductivity is increased linearly with  $V_G - V_{th}$ . As a result the transistor switches on, and a current flows from source to drain. The source-drain current  $I_{SD}$  can be expressed as [619]:

$$I_{SD} = \mu_{FE} C_G \frac{W}{L} \left[ (V_G - V_{th}) - \frac{V_D}{2} \right] V_D \quad (74)$$

where  $\mu_{FE}$  is the effective carrier or field-effect mobility,  $C_G$  the gate capacitance,  $V_D$  the drain voltage, and  $W$  and  $L$  the channel width and length, respectively. Saturation of the source-drain current occurs when  $dI_{SD}/dV_D = 0$ , and then is expressed as:

$$I_{SD,sat} = \mu_{FE} C_G \frac{W}{2L} (V_G - V_{th})^2 \quad (75)$$

The maximum current in the on-state is about  $10^{-4}$  A, while the current at  $V_G = 0$  is  $10^{-11}$  A, which shows the large dynamic range. The threshold voltage is around 1 V, typically, and  $\mu_{FE}$  can amount to  $1 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$  [618].

At the interface of the nitride ( $E_g = 5.3 \text{ eV}$ ) and the *a*-Si:H the conduction and valence band line-up. This results in band offsets. These offsets have been determined experimentally: the conduction band offset is 2.2 eV, and the valence band offset 1.2 [620]. At the interface a small electron accumulation layer is present under zero gate voltage due to the presence of interface states. As a result band bending occurs. The voltage at which the bands are flat (the flat-band voltage  $V_{FB}$  is slightly negative.

### XI.B.2. Stability

TFTs are not perfectly stable. After prolonged time of application of a gate voltage, the threshold voltage starts to shift. Two mechanisms can account for this [621]. One is the charge trapping of electrons in the nitride, where the silicon dangling bond is the dominant electron trap. When electrons are transferred from the *a*-Si:H to the nitride they can be trapped close to the interface, and a layer of fixed charge results. This fixed charge  $Q_F$  reduces the charge induced in the *a*-Si:H and the transfer characteristic shifts by an amount  $Q_F/C_G$ . In the second mechanism deep defect states are created by the

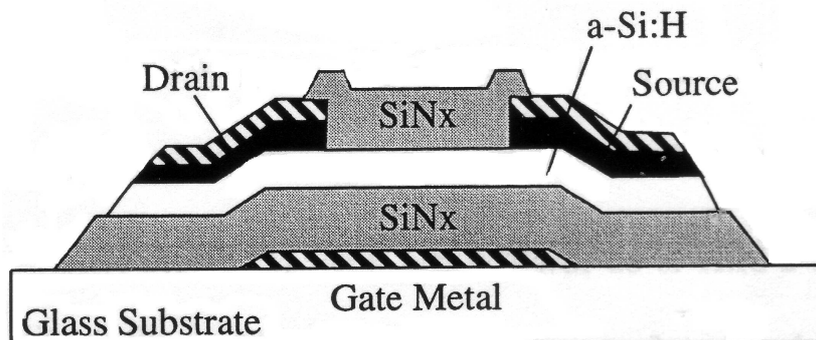


FIG. 74. Schematic cross-section of an inverted staggered TFT structure.

breaking of strained or weak Si-Si bonds. Electrons are trapped in these states and act as fixed negative charges, which results in band bending. This gives a decrease in source-drain current, or a positive shift of  $V_{th}$  of several volts. It has been found that the time dependence of the threshold current follows a stretched exponential behaviour [622]:

$$\Delta V_{th} = (V_{th}(\infty) - V_{th}(0)) \left[ \left( \exp \left( -\frac{t}{t_0} \right)^\beta \right) - 1 \right] \quad (76)$$

where  $V_{th}(\infty)$  and  $V_{th}(0)$  are the threshold voltages at infinity and at  $t = 0$ , respectively. The factor  $\beta$  has been related to the time-dependent ( $\sim t^{\beta-1}$ ) hydrogen diffusion coefficient in  $a$ -Si:H [623,624].

HWCVD-deposited  $a$ -Si:H layers incorporated in inverted-staggered TFTs have been reported to result in stable behaviour of the TFT [538,539]. The threshold voltage does not shift upon prolonged application of a gate-voltage. It has been suggested that not only the H content but rather the H bonding structure is responsible for the difference in metastable character for PECVD and HWCVD layers [94,96,97,625].

Another disadvantageous phenomenon in TFTs is the photoconductivity of  $a$ -Si:H [626]. Electrons and holes are photogenerated and recombine at the back surface (gate insulator). The photocurrent reduces the on/off ratio of the TFT. Illumination however, cannot always be avoided, e.g., in active matrix displays. A way of circumventing this is to make the  $a$ -Si:H as thin as possible.

### XI.B.3. Fabrication

The TFTs are made on transparent glass substrates onto which gate electrodes are patterned. Typically, the gate electrode is made of chromium. This substrate is introduced in a PECVD reactor, in which silane and ammonia are used for plasma deposition of  $\text{SiN}_x$  as the gate material. After subsequent deposition of the  $a$ -Si:H active layer, and the heavily doped  $n$ -type  $a$ -Si:H for the contacts, the devices are taken out of the reactor. Cr contacts are evaporated on top of the structure. The transistor channel is then defined by etching away the top metal and  $n$ -type  $a$ -Si:H. Special care must be taken as the etchant used for the  $n$ -type  $a$ -Si:H also etches the intrinsic  $a$ -Si:H. Finally the top passivation  $\text{SiN}_x$  is deposited in a separate run. This passivation layer is needed to protect the TFT during additional processing steps.

The thickness of the active layer is about 100-300 nm, while the source-drain distance (channel length) amounts to a few  $\mu\text{m}$ . The channel length is determined by the current requirements and usually exceeds 10  $\mu\text{m}$ . Other manufacturing schemes as well as alternative structure are described elsewhere [619,621]. Technology developments for the next generation TFTs that are to be used for high resolution displays have been summarized by Katayama [627].

#### *XI.B.4. Application in active matrix displays*

One of the most important applications of thin film transistors is in active matrix addressing of liquid crystal displays (AMLCD) [619,621]. The light transmission of a liquid crystal between two glass plates is controlled by the voltage on the pixel electrode. Nematic liquid crystals consist of rod-shaped molecules that align themselves parallel to an applied electrical field. When no field is present, they align themselves according to the surfaces in between which they are contained. By orienting the crystals a change in polarization of the light passing through the crystals can be achieved. In this way, the crystals act as light switches. The voltage on the pixel electrode is controlled by the transistors. These act as switches that pass the voltage on their source to the drain, that is the pixel electrode. When the gate is switched off, the pixel electrode is electrically isolated from the source, and the information remains on the pixel electrode. An  $n \times m$  matrix of source and gate lines is able to address  $n \times m$  pixels. This allows for high-definition large area television displays. As an example, a 640 by 480 pixel display with three subpixels for each colour consists of over one million transistors.

Active matrix addressing can also be used in printer heads [628] and linear sensor arrays [629]. The *a*-Si:H TFTs are used to address a linear array of output or input transducers, respectively. In the linear sensor array the *a*-Si:H photodiodes are used.

### **XI.C. Light sensors**

#### *XI.C.1. Linear arrays*

Linear arrays of *a*-Si:H photodiodes are widely used in optical page scanning applications such as fax machines and document scanners [630]. The large linear dimension of the array (as wide as the page to be scanned) allows a much simpler design. No optics are needed for image-size reduction, which is required when a CCD camera is used as the detecting element. Matrix addressing is similar to addressing in AMLCD technology, but confined to one dimension. Per pixel a *p-i-n* photodiode is connected to a transistor. During illumination of the pixel, charge is transferred to the bottom electrode, and accumulates when the TFT is off. Upon switching on of the TFT the charge will flow out and can be read by external electronic circuitry. In this way, the sensor integrates the signal during the time that it is not addressed. Read-out times are very short in comparison to accumulation times.

*XI.C.2. Photoreceptors*

The *a*-Si:H photoconductors can be used as the light-sensitive component in the electrophotographic or xerographic process [39,631,632]. Compared to conventional material such as As<sub>2</sub>Se<sub>3</sub>, CdS or ZnO *a*-Si:H is non-toxic and provides a hard surface. Typically, a three-layer structure is used on top of a substrate: glass/*n* *a*-Si:H/intrinsic *a*-Si:H/*a*-SiC:H.

The electrophotographic process consists of a number of steps [633]. First, the photoconductor is charged in the dark via a corona discharge. Then the photoconductor is illuminated such that the image to be copied is projected onto the surface. In the areas that were exposed by light the material becomes conductive and the charge will flow to the substrate. Thus, the non-illuminated parts still contain charge. The photoconductor then is moved under a bias electrode, and an electric field is present in the gap between electrode and charged surface. Now, (liquid) toner (ink) is introduced in this gap, and the pigment particles from the toner drift to the surface of the photoconductor due to the electrostatic force. The last step is the transfer of the toner to paper, where it is fused by heat or pressure to make the image permanent.

A charged photoconductor discharges slowly in the dark, as is evident from the small decrease in time of the surface voltage. The discharge during illumination is much faster. When the discharge is not complete a small residual voltage remains on the surface. A low dark current is required, which sets a limit on the dark photoconductivity of *a*-Si:H of  $10^{-12} \Omega^{-1}\text{cm}^{-1}$ . The residual voltage can be minimized by optimizing the  $\mu\tau$  product of the material.

The purpose of the *n*-type doped *a*-Si:H layer is to prevent injection of charge from the substrate into the photoconductor. Thus it serves as a blocking layer. Injection of surface charge into the photoconductor is prevented by the surface blocking layer. The *a*-SiC:H is of "low quality", with a high mid-gap density of states. Surface charge will be trapped in these mid-gap states.

The corona discharges produces oxygen ions and ozone that may react with the photoconductor [634]. As a means to circumvent possible degradation of the surface layer, an extra, protective thin layer was proposed, with high carbon content [101,635,636]. This would reduce silicon-oxygen reactions at the surface. Excellent electrophotographic characteristics have been obtained with a thin device comprising of a 0.1  $\mu\text{m}$  thick *n*-type *a*-Si:H layer, a 1.0  $\mu\text{m}$  intrinsic *a*-Si:H layer, a 0.1  $\mu\text{m}$  undoped *a*-SiC<sub>0.1</sub>:H layer, and a 0.014  $\mu\text{m}$  undoped *a*-SiC<sub>0.3</sub>:H layer [101].

*XI.C.3. Position sensor*

An *a*-Si:H based position sensor consists of an intrinsic film sandwiched between two transparent conductive electrodes [637]. Two line contacts on the top are perpendicular in direction to two bottom line contacts. When a light spot is incident on the device, carriers will be generated, and a photocurrent flows to the contacts. The contacts form resistive dividers, so that from the ratio of the photocurrents the position lateral position relative to the top or bottom contacts can be determined. The top contacts give the *x* position, and the bottom contacts the *y* position.

A *p-i-n* diode has been used on glass and on polyimide as a position sensitive detector [638,639]. The position of an incident light spot is measured by means of the lateral

photovoltage.

#### *XI.C.4. Colour sensors*

Similar to multi junction solar cells are colour detectors, that have been designed to detect two [640,641] or three [642–644] colours. A so-called adjustable threshold colour detector (ATCD) consists of an *a*-Si:H/*a*-SiC:H *p-i-n-i-p-i-n* stack [642]. This ATCD discriminates between the three fundamental colours, red, green and blue. As a result of proper thickness and band gap design, the blue is absorbed in the top *p-i-n* junction, the green in the middle *n-i-n* junction, and the red in the bottom *n-i-p* junction. A change in polarity and value of the voltage selects the colour that can be detected. The operation of the device is based on adjusting the photocurrent detection in the middle *n-i-n* junction.

A stack of TCO/*p-i-n-i-p*/TCO/*p-i-n*/metal has been designed as a three colour sensor [643,644]. An extra contact is made to the middle TCO. With appropriate band gaps the peak detection is at 450, 530, and 635 nm, for the blue, green, and red, respectively.

An active matrix of two-colour *a*-Si:H photodetectors has been reported [645], where a *n-i-p-i-n* switching device is stacked on a two-colour *p-i-n-i-p* structure.

#### *XI.C.5. IR sensor*

The absorption values of *a*-Si:H in the IR are much higher than of c-Si, due to transitions between extended states and deep levels. The IR-sensitivity increases with increasing defect density. However, as the collection efficiency decreases drastically with increasing defect density, photodiodes cannot be used. Another approach is based on the change in photocapacitance due to IR radiation. A structure similar to a *p-i-n* *a*-Si:H solar cell has been used as an infrared detector [646]. The intrinsic layer in fact is a so-called micro-compensated layer with very low concentrations of boron and phosphorous, but with a high defect density. In this layer the IR-sensitivity is enhanced, due to transitions between extended states and hole traps. The increase in trapped charge modifies the electric field in the device, which is observed as a change in capacitance. A high responsivity is observed between 800 and 1400 nm, as well as between 3500 and 4500 nm.

#### *XI.C.6. X-Ray sensor*

High energy radiation can be imaged with *a*-Si:H, either directly, or via a converter [3]. A thick film is required for direct detection, due to the weak interaction of the radiation with the material. A converter usually is a phosphor, which emits in the visible, and thin *a*-Si:H films are needed. X-rays with an energy up to 100 keV eject the electrons from the inner atomic core levels to high levels in the conduction band. The emitted electrons create electron-hole pairs due to ionization. These electron-hole pairs can be detected in the same way as in *p-i-n* photodiodes.

X-ray imagers consist of a phosphor in direct contact with the surface of an array of *a*-Si:H photodiodes. The device is a matrix-addressed array, in which each imaging

pixel consists of a photodiode and a TFT [647–649]. These X-ray imagers are very suitable for static and dynamic imaging in medical diagnosis.

### XI.D. Chemical sensors

#### XI.D.1. *pH-ISFET*

An ion-sensitive field effect transistor (ISFET) with a reference electrode is similar to a MOSFET, with the gate exposed to an electrolyte. Selected ion concentrations can be measured. The threshold voltage of an ISFET depends on the surface potential of the electrolyte/insulator, which in turn is dependent on temperature and on the pH of the electrolyte. The threshold voltage has been found to shift down linearly with about 0.5 V going from pH 7 to pH 1 [650]. Also a small drift in time is found of the gate voltage, and is small (1 mV/hr) for low pH and high (7 mV/hr) for high pH [651]. This is related to a small temporal change in capacitance of the insulator [652].

#### XI.D.2. *Hydrogen sensor*

A metal-insulator-semiconductor (MIS) device, where the palladium is used as the metal is sensitive to hydrogen. This sensitivity is a result of dissociation of hydrogen on the Pd surface, which induces an interface dipole layer that changes the barrier height and dark diode current (reverse bias current) [653]. Using a MIS structure with *a*-Si:H, i.e., glass/Cr/*n*-type *a*-Si:H/intrinsic *a*-Si:H/SiO<sub>x</sub>/Pd, Fortunato *et al.* [654] reported that the presence of 400 ppm of hydrogen leads to an increase of the reverse current of this device by three orders of magnitude. When these devices are illuminated during operation 400 ppm of hydrogen induces a shift of the open circuit voltage to lower values. Hence, these photochemical sensors can operate as a gas sensor either in the dark or under illumination.

### XI.E. Other applications

#### XI.E.1. *Loudspeakers*

The capability of depositing *a*-Si:H uniformly over large areas has been of special interest for application in electrostatic loudspeakers, where the *a*-Si:H layer is suitable for the retention layer on the vibrating foils in the loudspeaker. One aims at obtaining a large area thin film that can be either positively or negatively charged, by applying a high voltage, but at the same time can hold large surface charge densities, that do not displace laterally. Here, the high photosensitivity is not important, and in fact should be quenched for proper operation.

Electro-acoustic transducers can be divided into electrodynamic and electrostatic transducers. The most commonly used loudspeakers convert electrical energy to acoustic energy based on the electrodynamic principle, by driving an electromagnetic coil that activates a diaphragm. At low frequencies, however, the diaphragm acts as a high pass filter, with a cut-off in the low-frequency range. This cut-off frequency depends on

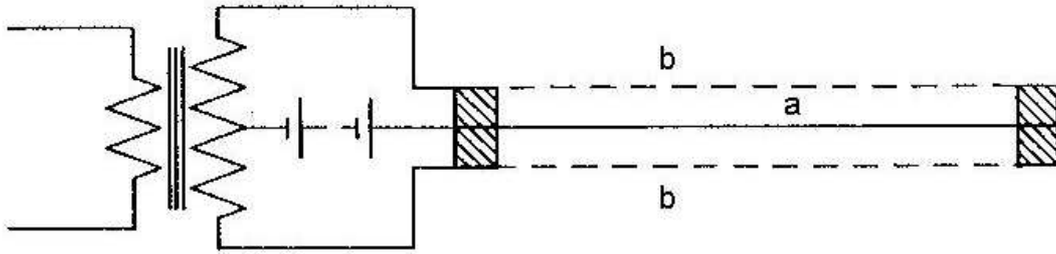


FIG. 75. Schematic drawing of an electrostatic loudspeaker. 'a' denotes the flexible thin  $a$ -Si:H (or graphite) coated foil, and 'b' denote the perforated flat fixed plates, respectively.

the diameter of the diaphragm. The mechanical construction of large-area loudspeakers adversely influences the amplitude and phase behaviour, especially at high frequencies. Therefore these drawback usually is overcome by using differently sized loudspeakers for different frequency ranges.

The electrostatic loudspeaker is used for high-end sound reproduction. The operation principle of an electrostatic loudspeaker is illustrated with Fig. 75. Essentially, the electrostatic loudspeaker is a capacitor with air as the dielectric. One of the electrode is a thin, electrically conducting foil (a in Fig. 75) and the other is a perforated flat fixed plate (b in Fig. 75) that also is conductive. The electrodes are oppositely charged by a high voltage, 5 kV. Sound is produced by superimposing audio-frequency voltages that drive the movement of the foil [655].

The following issues need to be addressed. The amplitude of the foil should be very small to avoid non-linear distortion. A large-area foil therefore is used. Further, the electric force between the electrodes can be as large as 20 kV/cm, which may lead to electrical breakthrough. Another issue is the presence of an electric field component over the surface of the foil, as a result of clamping the foil to the edges of the support. This electric field will vary with the amplitude of the foils, which leads to charge displacements and associated power dissipation. This effect especially may be present at low frequencies, where amplitudes are large.

The thin film should meet the following requirements. Areas in excess of 1 m<sup>2</sup> should be deposited continuously, reproducibly, and homogeneously. The use of plastic foils sets a limit to the deposition temperature. The thin film should be capable of holding large surface charge concentrations, and have a large lateral resistivity. It should be mechanically stable, capable of accommodating mechanical deformations, and resistant against humidity. Usually, the foils are coated with a graphite layer as the semiconducting thin film. However, these layers are frequently instable, and suffer from charge displacement effects, which eventually lead to electrical breakdown [656].

Prototype electrostatic loudspeakers where the graphite is replaced by  $a$ -Si:H have been made, where a mylar foil (area 10×10 cm<sup>2</sup>, thickness 6 μm) is used [657]. Deposition of the  $a$ -Si:H layer was carried out in the ASTER deposition system. Uniform deposition (standard deviation in thickness of 1.5 %) was achieved by diluting the SiH<sub>4</sub> with H<sub>2</sub> with SiH<sub>4</sub>:H<sub>2</sub> = 1:2. The deposition temperature was room temperature. The hydrogen content amounted to 18 at.%, and the band gap was 1.81 eV. The dark and AM1.5 photoconductivity were 7.5×10<sup>-9</sup> Ω<sup>-1</sup>cm<sup>-1</sup> and 1.8×10<sup>-8</sup> Ω<sup>-1</sup>cm<sup>-1</sup>, respectively. Note, that in practice the film would not be exposed to light.

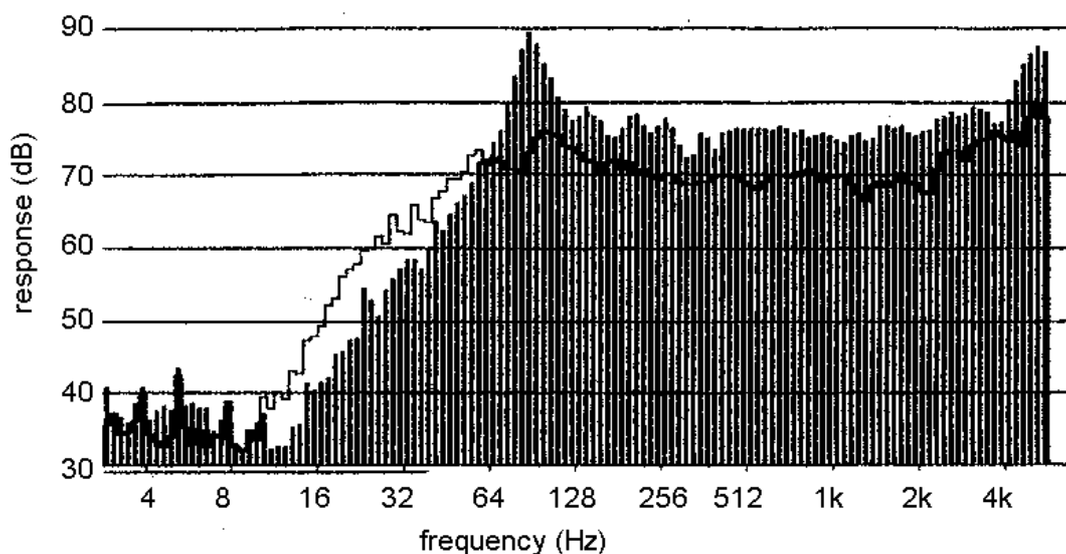


FIG. 76. Frequency response characteristics (dB versus frequency) of a conventional, graphite coated diaphragm (bar graph) and an *a*-Si:H coated diaphragm (line graph).

The frequency response characteristics of the foil with *a*-Si:H were measured in the range from 2.9 Hz to 5.46 kHz, by making use of generated pink noise with a continuous energy distribution. In Fig. 76 the frequency response characteristics of the *a*-Si:H coated foil are compared to a conventional graphite coated foil, of the same area. It can be concluded from this figure that the output of the *a*-Si:H coated loudspeaker is lower than that of the graphite coated one, from 60 Hz upwards. In spite of the lower efficiency in this range the *a*-Si:H coated loudspeaker outperforms the graphite coated one in the low range (10-60 Hz). The response is more extended towards low frequencies and the cut-off behaviour (dB/octave) also seems improved. The frequency response is stable, even after prolonged application of low frequency signals. The improvement was attributed to the reduced power dissipation at low frequencies due to the absence of charge displacement currents at the surface of the *a*-Si:H coated foil.

#### XI.E.2. Erbium in *a*-Si:H

Erbium, one of the rare-earth materials, has been very important in the development of optical communication technology in the past decades. The trivalent erbium ion ( $\text{Er}^{3+}$ ) has an incomplete  $4f$  shell which is shielded from the outside by  $5s$  and  $5p$  shells. Due to this, in erbium doped materials a sharp optical transition from the first excited state to the ground state of  $\text{Er}^{3+}$  is observed, which occurs at an energy of 0.8 eV. The corresponding wavelength is  $1.54 \mu\text{m}$ , which is of great importance as standard silica-based optical fibers have the highest transparency at this wavelength. Indeed, Er-doped silica optical fibers have been demonstrated to operate as optical amplifiers at  $1.54 \mu\text{m}$  [658,659]. Planar amplifiers, in which Er-doped channel waveguides are manufactured on a planar substrate have also been demonstrated [660]. These are of particular importance as they can be integrated with other waveguide devices on a single chip. If efficient light emission from Er-doped silicon could be possible, integration of optical and electrical functions on a single silicon chip would be within reach.

Due to small emission and absorption cross sections of  $\text{Er}^{3+}$  a high Er density is needed to reach reasonable values of optical gain. Typically Er densities are between 0.1 and 1.0 % ( $10^{19}$ - $10^{20}$   $\text{Er}/\text{cm}^3$ ). These values are far beyond the equilibrium solubility limits of Er in silicon. Therefore, non-equilibrium methods have to be used, such as ion implantation. Er implantation in crystalline silicon leads to amorphization, and additional annealing (600 °C) is required to recrystallize the silicon. Optical activation of the Er may even require subsequent annealing at higher temperatures. Impurities such as oxygen or carbon have been found to enhance the luminescence. More can be found in the reviews on Er in silicon, that have been published by Polman [661] and Priolo *et al.* [662,663].

In this section, the Er-doping of amorphous silicon is discussed. The problem of limited solubility of Er in crystalline silicon is circumvented. However, the electrical properties of pure a-Si are poor compared to c-Si. Therefore, hydrogenated amorphous silicon is much more interesting. Besides, the possibility of depositing a-Si:H directly on substrates, i.e., optical materials, would make integration possible. Both low-pressure chemical vapour deposition (LPCVD) [664] and PECVD [665,666] have been used to make the a-Si:H into which Er is implanted. In both methods oxygen intentionally is added to the material, to enhance the luminescence.

The a-Si:H is deposited on Si(100) by LPCVD from  $\text{SiH}_4$  and  $\text{N}_2\text{O}$  at 620 °C, with hydrogen and oxygen contents of 10 and 31 %, respectively. The a-Si:H layer was 340 nm thick. Erbium was implanted at 500 keV to a dose of  $1 \times 10^{15}$   $\text{Er}/\text{cm}^2$ . The peak concentration of Er at a depth of 150 nm was 0.2 %. Upon annealing at 400 °C room-temperature photoluminescence is observed [664], with the characteristic  $\text{Er}^{3+}$ -peak at 1.54  $\mu\text{m}$ . Temperature dependent luminescence measurements in the range from 77 K to 300 K, show quenching of the peak-luminescence by only a factor of three. In the case of Er-implanted crystalline silicon, this quenching is 10-100 times larger. Moreover, the luminescence intensity for the amorphous material is higher over the whole range of measured temperatures.

Luminescence lifetime measurements reveal a double exponential decay behaviour, with lifetimes around 160 and 800  $\mu\text{s}$ , independent of temperature in the range of 77-300 K. This behaviour differs from the lifetime quenching as observed for oxygen or nitrogen co-doped c-Si, where the lifetime is quenched by one to nearly three orders of magnitude [667,668]. This has been attributed to a nonradiative deexcitation process of excited  $\text{Er}^{3+}$  that occurs at higher temperatures and which depends on the band gap of the material [669]: the larger the band gap the less quenching should be observed. Indeed, the large band gap of the LPCVD a-Si:H of about 2 eV seems to inhibit quenching.

With PECVD a-Si:H is deposited on Corning glass substrate from  $\text{SiH}_4$  at 230 °C, employing the PASTA deposition system described by Madan *et al.* [159]. The hydrogen and oxygen contents were of 10 and 0.3 %, respectively. The presence of this background concentration of oxygen probably was due to postponed maintenance, however, it was advantageous for this particular experiment. The a-Si:H layer was 250 nm thick. Erbium was implanted at 125 keV to a dose of  $4 \times 10^{14}$   $\text{Er}/\text{cm}^2$ . The peak concentration of Er at a depth of 35 nm was 0.2 %. In some samples additional oxygen was implanted at 25 keV to a dose of  $7 \times 10^{15}$   $\text{Er}/\text{cm}^2$ , which resulted in an oxygen peak concentration of 1.7 %, that overlapped the Er depth profile. Upon annealing at 400 °C room-temperature photoluminescence is observed [665,666], with the

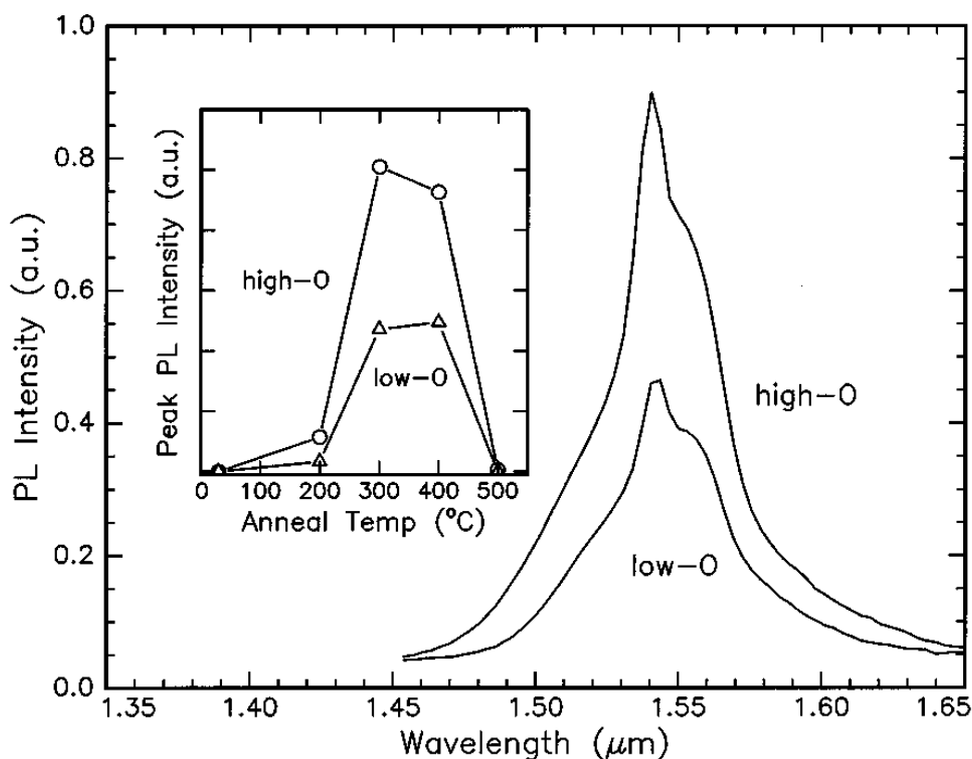


FIG. 77. Room-temperature photo-luminescence spectra of Er-implanted PECVD a-Si:H, annealed at 400 °C. The implantation energy and dose were 125 keV and  $4 \times 10^{14}$  Er/cm<sup>2</sup>, respectively, which resulted in peak concentration of 0.2 at.%. 'low-O' and 'high-O' denote a peak oxygen concentration in a-Si:H of 0.3 and 1.3 at.%, respectively. The inset shows the 1.54  $\mu\text{m}$  peak intensity as a function of annealing temperature, for both oxygen concentrations. (From J. H. Shin, R. Serna, G. N. van den Hoven, A. Polman, W. G. J. H. M. van Sark, and A. M. Vredenberg, *Appl. Phys. Lett.* **68**, 697 (1996).).

characteristic Er<sup>3+</sup>-peak at 1.54  $\mu\text{m}$ , as shown in Fig. 77. The inset shows the behaviour of the peak intensity as a function of annealing temperature. Temperatures between 300 and 400 °C clearly are optimum annealing temperatures. Both the as-implanted samples and the samples annealed at 500 °C show no erbium-related luminescence. In order to remove irradiation-induced effects high annealing temperatures are needed, however, this competes with the outdiffusion of hydrogen which is enhanced at higher temperatures.

Temperature dependent luminescence measurements in the range from 77 K to 300 K, show quenching of the peak-luminescence by a factor of about 15. A similar behaviour is observed in the lifetime quenching [665,666]. As the band gap of the PECVD a-Si:H is about 1.6 eV, nonradiative deexcitation of Er may occur at elevated temperatures. The amount of quenching lies inbetween that of c-Si and LPCVD a-Si:H, just as the band gap.

Electroluminescent devices were made to demonstrate the possible application of these a-Si:H materials. Er-doped *p-n* diodes in c-Si show electroluminescence, both in forward and reverse bias [670–672]. Under forward bias the electroluminescence signal is attributed to electron-hole recombination, which results in excitation of the Er<sup>3+</sup>. Under reverse bias, in particular beyond the Zener breakdown, the erbium is excited

by impact excitation by hot electrons that are accelerated across the junction.

The LPCVD *a*-Si:H material, incorporated in a device, shows electroluminescence only in reverse bias [673]. The mechanism is similar to the one for *c*-Si described above. The PECVD *a*-Si:H material was incorporated in a *p-i-n* solar cell structure, with a thickness of the intrinsic layer of 500 nm (see Sec. XIA). Oxygen was coimplanted at 80 keV ( $3.2 \times 10^{15}$  O/cm<sup>2</sup>) and at 120 keV ( $5.5 \times 10^{15}$  O/cm<sup>2</sup>) which resulted in a roughly constant oxygen concentration of 1.0 % at the Er projected range in the middle of the intrinsic *a*-Si:H layer. Electroluminescence is observed under forward bias [674].

Incorporation of Er *during* deposition of *a*-Si:H has also been achieved using molecular beam epitaxy [675], magnetron-assisted decomposition of silane [676], PECVD with a metalorganic source of solid erbium (tris(2,2,6,6-tetramethyl-2,5-heptadionato)Er(III) [677,678] and tris(hexafluoroacetylacetonate)Er(III)·1,2-dimethoxyethane [679]), electron cyclotron resonance PECVD in combination with sputtering of erbium [680], and catalytic chemical vapour deposition combined with pulsed laser-ablation [681]. Explanations of the quenching behaviour seem to be dependent on the sample preparation technique, however recently a comprehensive energy transfer model based on a Förster mechanism (resonant dipole coupling [682]) has been proposed [683].

### XI.E.3. Miscellaneous

Photodiode arrays have been used as retinal implants [684]. These arrays of *p-i-n* diodes are fabricated on a thin titanium layer bonded to a glass plate. The total thickness of this flexible structure amounts to 1.5 μm. The microphotodiode array (MPDA) is used to replace photoreceptors (rods and cones) that have become defective due some diseases.

Near-field optical microscopy (NSOM) [685,686] has been used as a tool for nanolithography on the submicron length scale. Here, the fiber optic probe is used as a light source exposing a resist by scanning the surface. Dithering (oscillating the probe tip parallel to the surface) is needed to keep a constant probe-sample distance. As NSOM is not diffraction limited it is in principle possible to generate features with smaller dimensions (< 100 nm [687,688]) than those that are achievable with far-field optical lithography. Patterning of *a*-Si:H involves oxidizing the surface by local removal of hydrogen passivation. The thus formed oxide layer is the mask for subsequent etching. Herndon *et al.* [689] have reported on NSOM patterning of PECVD deposited *a*-Si:H in open air, by using an Ar-ion laser as the light source. Subsequent etching in a hydrogen plasma revealed an etch selectivity of 500:1. The feature widths are found to be dependent on dither amplitude

A monolithic, amorphous-silicon-based, photovoltaic-powered electrochromic window has been reported by Gao *et al.* [690]. A thin, wide band gap *a*-SiC:H *n-i-p* solar cell on top of a glass substrate is used as a semitransparent power supply. An electrochromic device consisting of Li<sub>y</sub>WO<sub>3</sub>/LiAlF<sub>4</sub>/V<sub>2</sub>O<sub>5</sub> is deposited directly on top of the solar cell. The operation principle is based on the change of optical absorption by inserting and extracting Li<sup>+</sup> in and out of the WO<sub>3</sub> film. A prototype 16 cm<sup>2</sup> sized device is able to modulate the transmittance of the stack by more than 60% over a large part of the visible spectrum.

## XII. CONCLUSION

In this chapter the deposition of *a*-Si:H by PECVD was described. It covered material as well as discharge issues. It was tried to relate material and discharge properties in various ways. Plasma modeling provides a means to study in detail the physical and chemical processes that occur in the plasma. The presented models show a high degree of sophistication, but from the comparison with experimental data it is clear that especially the deposition model needs improvement. Also, a full 2D model most probably is not needed, as differences between 1D and 2D modeling results are not that large.

Plasma analysis reveals information on the products of chemical processes, and can excellently be used as a feedback to the plasma modeling. The role of ions has been thoroughly illustrated, and the important result that ion bombardment with moderate energy is beneficial for material quality has been quantified.

The many modifications to the conventional RF-PECVD method show that one still is trying to find methods that will in the end lead to improved material properties. This is especially the case for the intrinsic metastability of *a*-Si:H. In this respect, the stable material that is obtained at discharge conditions "at the edge of crystallinity" is very promising. Also, the quest for higher deposition rates while at least maintaining device quality material properties shows the industrial drive behind the research. Faster deposition simply allows for more solar cells produced in the same time.

In the three deposition methods PECVD, HWCVD, and ETP-CVD one can see certain similarities. In one region in the reactor system, the generation of growth precursors takes place. For PECVD this occurs in the discharge bulk, for HWCVD at/around the wire, and for ETP-CVD in the vicinity of the expanding beam. A transport region exists where precursors are transported to the substrate. For HWCVD and ETP-CVD this region is between wire/source and substrate. For PECVD this region is the sheath. The third region is the deposition region, in all cases the substrate. Looking at these techniques from this perspective, one can conclude that the deposition mechanism of *a*-Si:H is universal. The best material quality is obtained when the SiH<sub>3</sub> radical is abundantly present.

In the last part of this chapter a summary was given of some applications. These are the driving force for the large research effort that has existed and still continues to exist. From these applications and the many more that were not treated here, it will be clear that *a*-Si:H is a material that is omnipresent, and contributes greatly to our well-being.

### XIII. ACKNOWLEDGEMENTS

It goes without saying that the work presented here could not have been done without the help and support of many people. First, many thanks go to John Beze-mer, my supervisor, who taught me all there is to know about plasma deposition of *a*-Si:H and now enjoys his retirement. Second, I gratefully acknowledge the colleagues with whom I performed and discussed numerous deposition and plasma analysis experiments: Hans Meiling, Edward Hamers, Stefania Acco, Martin van de Boogaard, Arjan Berntsen, Ernst Ullersma, René van Swaaij, Meine Kars, Gerard van der Mark, and students Renée Heller, Remco van der Heiden, Pieter van de Vliet, Micha Kuiper, Tjitske Kooistra, Jan-Wijtse Smit, Edward Prendergast, and Tom Thomas. Further, I would gratefully like to thank the following persons for various types of continuous support during my active time of research on *a*-Si:H deposition: Wim Arnold Bik, Frans Habraken, Arjen Vredenberg, Dirk Knoesen, Bert Slomp, Albert Polman, Gerhard Landweer, Jeroen Daey Ouwens, Kees Feenstra, Maarten van Cleef, Jatindra Rath, Wim Sinke, Bert Slomp, Jeike Wallinga, Karine van der Werf, Ruud Schropp, and Werner van de Weg, who headed the group for over a decade.

I very much enjoyed the fruitful collaboration with Wim Goedheer and his group, i.e., Gert Jan Nienhuis, Peter Meijer, and Diederick Passchier, at the FOM Institute for Plasmaphysics "Rijnhuizen" and thank Wim Goedheer for supplying me with published and additional data used in the section on plasma modeling.

Also the bi-annual dutch meetings on solar cell research always allowed for many discussions with colleagues from Eindhoven Technical University (Daan Schram, Richard van de Sanden) and Delft Technical University (Miro Zeman, Wim Metselaar).

Further I acknowledge Hans Gerritsen and Gijs van Ginkel for allowing and stimulating me to write this chapter.

And last but not least I thank my wife Joni, son Guido, and daughter Sophie for the enormous joy they give me every day.

Parts of the work presented here were financially supported by the Netherlands Agency for Energy and the Environment (NOVEM), the Foundation for Fundamental Research on Matter (FOM), the Royal Netherlands Academy of Art and Sciences (KNAW), the Netherlands Technology Foundation (STW), and the Netherlands Organization for Scientific Research (NWO).

## XIV. REFERENCES

- 
1. J. I. Pankove, Ed., *Hydrogenated Amorphous Silicon, Semiconductors and Semimetals, Vol. 21, Parts A-D* (Academic Press, Orlando, FL, U.S.A., 1984).
  2. K. Tanaka, Ed., *Glow-Discharge Hydrogenated Amorphous Silicon* (KTK Scientific Publishers, Tokyo, Japan, 1989).
  3. R. A. Street, *Hydrogenated Amorphous Silicon, Cambridge Solid State Science Series* (Cambridge University Press, Cambridge, U.K., 1991).
  4. J. Kanicki, Ed., *Amorphous and Microcrystalline Semiconductor Devices – Optoelectronic devices* (Artech House, Norwood, MA, U.S.A., 1991).
  5. J. Kanicki, Ed., *Amorphous and Microcrystalline Semiconductor Devices – Materials and Device Physics* (Artech House, Norwood, MA, U.S.A., 1992).
  6. W. Luft and Y. S. Tsuo, *Hydrogenated amorphous silicon alloy deposition processes* (Marcel Dekker, Inc., New York, NY, U.S.A., 1993).
  7. R. F. Bunshah, Ed., *Handbook of Deposition Technologies for Films and Coatings – Science, Technology and Applications, Second Edition* (Noyes Publications, Park Ridge, NJ, U.S.A., 1994).
  8. G. Bruno, P. Capezzuto, and A. Madan, Eds., *Plasma Deposition of Amorphous Silicon-Based Materials* (Academic Press, Boston, MA, U.S.A., 1995).
  9. E. S. Machlin, *Materials Science in Microelectronics – The Relationships between Thin Film Processing and Structure* (Giro Press, Croton-on-Hudson, NY, U.S.A., 1995).
  10. E. S. Machlin, *Materials Science in Microelectronics – Volume 2, The Effects of Structure on Properties in Thin Films* (Giro Press, Croton-on-Hudson, NY, U.S.A., 1998).
  11. R. E. I. Schropp and M. Zeman, *Amorphous and Microcrystalline Silicon Solar Cells – Modeling, Materials and Device Technology* (Kluwer Academic Publishers, Boston, MA, U.S.A., 1998).
  12. T. M. Searle, Ed., *Properties of Amorphous Silicon and its Alloys, EMIS Datareview Series, No. 19* (IEE, London, U.K., 1998).
  13. R. A. Street, Ed., *Technology and Applications of Amorphous Silicon* (Springer, New York, NY, U.S.A., 2000).
  14. C. A. M. Stap, H. Meiling, G. Landweer, J. Bezemer, and W. F. van der Weg, in *Proceedings of the Ninth E.C. Photovoltaic Solar Energy Conference, Freiburg, F.R.G., 1989*, edited by W. Palz, G. T. Wrixon, and P. Helm (Kluwer Academic, Dordrecht, the Netherlands, 1989), p. 74.
  15. H. Meiling, Ph.D. thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1991.
  16. H. Curtins, N. Wyrsh, and A. V. Shah, *Electron. Lett.* **23**, 228 (1987).
  17. H. Shirai, J. Hanna, and I. Shimizu, *Jpn. J. Appl. Phys.* **30**, L881 (1991).
  18. J. T. Verdeyen, L. Beberman, and L. Overzet, *J. Vac. Sci. Technol. A* **8**, 1851 (1990).
  19. A. H. Mahan, J. Carapella, B. P. Nelson, R. S. Crandall, and I. Balberg, *J. Appl. Phys.* **69**, 6728 (1991).
  20. G. M. W. Kroesen, D. C. Schram, and M. J. F. van de Sande, *Plasma Chem. Plasma Proc.* **10**, 49 (1990).

21. G. J. Meeusen, E. A. Ershov-Pavlov, R. F. G. Meulenbroeks, M. C. M. van de Sanden, and D. C. Schram, *J. Appl. Phys.* **71**, 4156 (1992).
22. H. Overhof, *J. Non-Cryst. Solids* **227–230**, 15 (1998).
23. J. Tauc, Ed., *Amorphous and Liquid Semiconductors* (Plenum, New York, NY, U.S.A., 1974).
24. A. J. Lewis, G. A. N. Connell, W. Paul, J. R. Pawlik, and R. J. Temkin, in *Tetrahedrally Bonded Amorphous Semiconductors*, No. 20 in *AIP Conference Proceedings*, edited by M. H. Brodsky, S. Kirkpatrick, and D. Weaire (American Institute of Physics, New York, N.Y., U.S.A., 1974), p. 27.
25. E. C. Freeman and W. Paul, *Phys. Rev. B* **18**, 4288 (1978).
26. W. Paul and H. Ehrenreich, *Phys. Today* **38**, (8) 13 (1985).
27. H. F. Sterling and R. C. G. Swann, *Solid State Electron.* **8**, 653 (1965).
28. R. C. Chittick, J. H. Alexander, and H. F. Sterling, *J. Electrochem. Soc.* **116**, 77 (1969).
29. A. Triska, D. Dennison, and H. Fritzsche, *Bull. Am. Phys. Soc.* **20**, 392 (1975).
30. H. Fritzsche, in *Proceedings of the 7<sup>th</sup> International Conference on Amorphous and Liquid Semiconductors*, edited by W. E. Spear (CICL, Edinburgh, U.K., 1977), p. 3.
31. H. Fritzsche, *Mater. Res. Soc. Proc. Symp.* **609**, A17.1.1 (2000).
32. W. Spear and P. LeComber, *Solid State Commun.* **17**, 1193 (1975).
33. I. Solomon, *Mater. Res. Soc. Proc. Symp.* **609**, A17.3.1 (2000).
34. D. E. Carlson and C. R. Wronski, *Appl. Phys. Lett.* **28**, 671 (1976).
35. H. Okamoto, Y. Nitta, T. Yamaguchi, and Y. Hamakawa, *Sol. Energy Mater.* **3**, 313 (1980).
36. Y. Hamakawa, *Mater. Res. Soc. Proc. Symp.* **609**, A17.2.1 (2000).
37. Y. Kuwano, T. Imai, M. Ohnishi, and S. Nakano, in *Proceedings of the 14<sup>th</sup> IEEE Photovoltaic Specialists Conference, San Diego, 1980* (IEEE, New York, N.Y., U.S.A., 1980), p. 1408.
38. I. Shimizu, T. Komatsu, T. Saito, and E. Inoue, *J. Non-Cryst. Solids* **35 & 36**, 773 (1980).
39. S. Oda, Y. Saito, I. Shimizu, and E. Inoue, *Philos. Mag. B* **43**, 1079 (1981).
40. P. G. LeComber, W. E. Spear, and A. Ghaith, *Electron. Lett.* **15**, 179 (1979).
41. A. J. Snell, K. D. MacKenzie, W. E. Spear, P. G. LeComber, and A. J. Hughes, *Appl. Phys.* **24**, 357 (1981).
42. M. J. Powell, B. C. Easton, and O. F. Hill, *Appl. Phys. Lett.* **38**, 794 (1981).
43. R. L. Weisfield, *J. Non-Cryst. Solids* **164–166**, 771 (1993).
44. S. N. Kaplan, J. R. Morel, T. A. Mulera, V. Perez-Mendez, G. Schlurmacher, and R. A. Street, *IEEE Trans. Nuc. Sci.* **NS-33**, 351 (1986).
45. T. C. Chuang, L. E. Fennel, W. B. Jackson, J. Levine, M. J. Thompson, H. C. Tuan, and R. Weisfield, *J. Non-Cryst. Solids* **97 & 98**, 301 (1987).
46. L. E. Antonuk, C. F. Wild, J. Boundry, J. Jimenez, M. J. Longo, and R. A. Street, *IEEE Trans. Nuc. Sci.* **NS-37**, 165 (1990).
47. C. van Berkel, N. C. Bird, C. J. Curling, and I. D. French, *Mater. Res. Soc. Symp. Proc.* **297**, 939 (1993).
48. W. H. Zachariasen, *J. Am. Chem. Soc.* **54**, 3841 (1932).
49. D. E. Polk, *J. Non-Cryst. Solids* **5**, 365 (1971).
50. M. A. Paesler, D. E. Sayers, R. Tsu, and J. G. Hernandez, *Phys. Rev. B* **28**, 4550 (1983).
51. D. Beeman, R. Tsu, and M. F. Thorpe, *Phys. Rev. B* **32**, 874 (1985).
52. R. A. Street, *Phys. Rev. Lett.* **49**, 1187 (1982).

53. N. F. Mott, *Philos. Mag.* **19**, 835 (1969).
54. W. M. Arnold Bik, C. T. A. M. de Laat, and F. H. P. M. Habraken, *Nucl. Instrum. Methods Phys. Res. B* **64**, 832 (1992).
55. M. H. Brodsky, *Thin Solid Films* **40**, L23 (1977).
56. G. Lucovsky, R. J. Nemanich, and J. C. Knights, *Phys. Rev. B* **19**, 2064 (1979).
57. H. Shanks, C. J. Fang, L. Ley, M. Cardona, F. J. Demond, and S. Kalbitzer, *Phys. Status Solidi B* **100**, 43 (1980).
58. A. A. Langford, M. L. Fleet, B. P. Nelson, W. A. Lanford, and N. Maley, *Phys. Rev. B* **45**, 13367 (1992).
59. W. Beyer and M. S. Abo Ghazala, *Mater. Res. Soc. Symp. Proc.* **507**, 601 (1998).
60. A. H. Mahan, P. Raboisson, and R. Tsu, *Appl. Phys. Lett.* **50**, 335 (1987).
61. D. A. G. Bruggeman, *Ann. Phys. (Leipzig)* **24**, 636 (1935).
62. A. H. Mahan, D. L. Williamson, B. P. Nelson, and R. S. Crandall, *Sol. Cells* **27**, 465 (1989).
63. H. Meiling, M. J. van den Boogaard, R. E. I. Schropp, J. Bezemer, and W. F. van der Weg, *Mater. Res. Soc. Symp. Proc.* **192**, 645 (1990).
64. M. J. van den Boogaard, S. J. Jones, Y. Chen, D. L. Williamson, R. A. Hakvoort, A. van Veen, A. C. van der Steege, W. M. Arnold Bik, W. G. J. H. M. van Sark, and W. F. van der Weg, *Mater. Res. Soc. Symp. Proc.* **258**, 407 (1992).
65. M. J. van den Boogaard, Ph.D. thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1992.
66. S. Acco, D. L. Williamson, W. G. J. H. M. van Sark, W. C. Sinke, W. F. van der Weg, A. Polman, and S. Roorda, *Phys. Rev. B* **58**, 12853 (1998).
67. A. H. Mahan, D. L. Williamson, B. P. Nelson, and R. S. Crandall, *Phys. Rev. B* **40**, 12024 (1989).
68. P. C. Taylor, in *Semiconductors and Semimetals*, edited by J. I. Pankove (Academic, Orlando, Fla., U.S.A., 1984), Vol. 21C, p. 99.
69. S. Acco, D. L. Williamson, P. A. Stolk, F. W. Saris, M. J. van den Boogaard, W. C. Sinke, W. F. van der Weg, and S. Roorda, *Phys. Rev. B* **53**, 4415 (1996).
70. S. Acco, W. Beyer, E. E. Van Faassen, and W. F. van der Weg, *J. Appl. Phys.* **82**, 2862 (1997).
71. R. S. Crandall, X. Liu, and E. Iwaniczko, *J. Non-Cryst. Solids* **227–230**, 23 (1998).
72. S. Acco, D. L. Williamson, S. Roorda, W. G. J. H. M. van Sark, A. Polman, and W. F. van der Weg, *J. Non-Cryst. Solids* **227–230**, 128 (1998).
73. P. J. Zanzucchi, C. R. Wronski, and D. E. Carlson, *J. Appl. Phys.* **48**, 5227 (1977).
74. G. D. Cody, B. Abeles, C. Wronski, C. R. Stephens, and B. Brooks, *Sol. Cells* **2**, 227 (1980).
75. J. Tauc, in *Optical Properties of Solids*, edited by F. Abelès (North-Holland, Amsterdam, the Netherlands, 1972), Chap. 5, p. 277.
76. R. H. Klazes, M. H. L. M. van den Broek, J. Bezemer, and S. Radelaar, *Philos. Mag. B* **45**, 377 (1982).
77. T. Tiedje, J. M. Cebulka, D. L. Morel, and B. Abeles, *Phys. Rev. Lett.* **46**, 1425 (1981).
78. K. Pierz, B. Hilgenberg, H. Mell, and G. Weiser, *J. Non-Cryst. Solids* **97/98**, 63 (1987).
79. K. Pierz, W. Fuhs, and H. Mell, *Philos. Mag. B* **63**, 123 (1991).
80. N. Maley and J. S. Lannin, *Phys. Rev. B* **36**, 1146 (1987).
81. R. C. Ross and J. Jaklik, *J. Appl. Phys.* **55**, 3785 (1984).
82. H. Meiling, W. Lenting, J. Bezemer, and W. F. van der Weg, *Philos. Mag. B* **62**, 19

- (1990).
83. R. S. Crandall, *Sol. Cells* **24**, 237 (1988).
  84. A. J. M. Berntsen, W. F. van der Weg, P. A. Stolk, and F. W. Saris, *Phys. Rev. B* **48**, 14656 (1993).
  85. A. J. M. Berntsen, Ph.D. thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1993.
  86. W. E. Spear and P. G. LeComber, *Adv. Phys.* **26**, 811 (1977).
  87. D. L. Staebler and C. R. Wronski, *Appl. Phys. Lett.* **31**, 292 (1977).
  88. D. L. Staebler and C. R. Wronski, *J. Appl. Phys.* **51**, 3262 (1980).
  89. M. Stutzmann, W. B. Jackson, and C. C. Tsai, *Phys. Rev. B* **32**, 23 (1985).
  90. T. Kamei, N. Hata, A. Matsuda, T. Uchiyama, S. Amano, K. Tsukamoto, Y. Yoshioka, and T. Hirao, *Appl. Phys. Lett.* **68**, 2380 (1996).
  91. C. C. Tsai, J. C. Knights, R. A. Lujan, W. B., B. L. Stafford, and M. J. Thompson, *J. Non-Cryst. Solids* **59&60**, 731 (1983).
  92. S. Yamasaki and J. Isoya, *J. Non-Cryst. Solids* **164–166**, 169 (1993).
  93. P. Stradins and H. Fritzsche, *Philos. Mag. B* **69**, 121 (1994).
  94. C. Godet, P. Morin, and P. Roca i Cabarrocas, *J. Non-Cryst. Solids* **198–200**, 449 (1996).
  95. J. Daey Ouwens and R. E. I. Schropp, *Phys. Rev. B* **54**, 17759 (1996).
  96. S. Zafar and E. A. Schiff, *Phys. Rev. B* **40**, 5235 (1989).
  97. C. Manfredotti, F. Fizzotti, M. Boero, P. Pastorino, P. Polesello, and E. Vittone, *Phys. Rev. B* **50**, 18046 (1994).
  98. H. M. Branz, *Solid State Commun.* **105**, 387 (1998).
  99. R. Biswas, Q. Li, B. C. Pan, and Y. Yoon, *Mater. Res. Soc. Symp. Proc.* **467**, 135 (1997).
  100. D. A. Anderson and W. E. Spear, *Philos. Mag.* **35**, 1 (1977).
  101. R. A. C. M. M. van Swaaij, Ph.D. thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1994.
  102. R. A. C. M. M. van Swaaij, A. J. M. Berntsen, W. G. J. H. M. van Sark, H. Herremans, J. Bezemer, and W. F. van der Weg, *J. Appl. Phys.* **76**, 251 (1994).
  103. J. Chevallier, H. Wieder, A. Onton, and C. R. Guarnieri, *Solid State Comm.* **24**, 867 (1977).
  104. S. Wagner, V. Chu, J. P. Conde, and J. Z. Liu, *J. Non-Cryst. Solids* **114**, 453 (1989).
  105. C. M. Fortmann, in *Plasma Deposition of Amorphous Silicon-Based Materials*, edited by G. Bruno, P. Capezzuto, and A. Madan (Academic Press, Boston, MA, U.S.A., 1995), Chap. 3, p. 131.
  106. W. Paul, S. J. Jones, F. C. Marques, D. Pang, W. A. Turner, A. E. Wetsel, P. Wickboldt, and J. H. Chen, *Mater. Res. Soc. Symp. Proc.* **219**, 211 (1991).
  107. A. Matsuda and K. Tanaka, *J. Non-Cryst. Solids* **97**, 1367 (1987).
  108. G. Bruno, P. Capezzuto, G. Cicala, and F. Cramarossa, *J. Mater. Res.* **4**, 366 (1989).
  109. G. Bruno, P. Capezzuto, M. Losurdo, P. Manodoro, and G. Cicala, *J. Non-Cryst. Solids* **137&138**, 799 (1991).
  110. H. Herremans, W. Grevendonk, R. A. C. M. M. van Swaaij, W. G. J. H. M. van Sark, A. J. M. Berntsen, W. M. Arnold Bik, and J. Bezemer, *Philos. Mag. B* **66**, 787 (1992).
  111. M. P. Schmidt, I. Solomon, H. Tran-Quoc, and J. Bullot, *J. Non-Cryst. Solids* **77–78**, 849 (1985).
  112. I. Solomon, M. P. Schmidt, and H. Tran-Quoc, *Phys. Rev. B* **38**, 9895 (1988).
  113. M. N. P. Carreño, I. Pereyra, M. C. A. Fantini, H. Takahashi, and R. Landers, *J. Appl.*

- Phys. **75**, 538 (1994).
114. W. M. Arnoldbik and F. H. P. M. Habraken, Rep. Prog. Phys. **56**, 859 (1993).
  115. M. L. Oliveira and S. S. Camargo Jr., J. Appl. Phys. **71**, 1531 (1992).
  116. F. Evangelisti, J. Non-Cryst. Solids **164–166**, 1009 (1993).
  117. A. Grill, *Cold Plasma in Materials Fabrication – From Fundamentals to Applications* (IEEE Press, Piscataway, NJ, U.S.A., 1994).
  118. J. F. O'Hanlon, *A User's Guide to Vacuum Technology, Second Edition* (Wiley, New York, NY, U.S.A., 1989).
  119. J. Bezemer, W. G. J. H. M. van Sark, M. B. von der Linden, and W. F. van der Weg, in *Proceedings of the Twelfth E.C. Photovoltaic Solar Energy Conference, Amsterdam, the Netherlands, 1994*, edited by R. Hill, W. Palz, and P. Helm (H.S. Stephens & Associates, Bedford, U.K., 1994), p. 327.
  120. J. Bezemer and W. G. J. H. M. van Sark, in *Electronic, Optoelectronic and Magnetic Thin films, Proceedings of the 8th International School on Condensed Matter Physics (ISCMP), Varna, Bulgaria, 1994* (Wiley, New York, NY, U.S.A., 1995), p. 219.
  121. M. Böhm, A. E. Delahoy, F. B. Ellis Jr., E. Eser, S. C. Gau, F. J. Kampas, and Z. Kiss, in *Proceedings of the 18<sup>th</sup> IEEE Photovoltaic Specialists Conference, Las Vegas, 1985* (IEEE, New York, N.Y., U.S.A., 1985), p. 888.
  122. C. R. Dickson, J. Pickens, and A. Wilczynski, Solar Cells **19**, 179 (1986).
  123. K. Tanaka and A. Matsuda, Mater. Sci. Rep. **2**, 139 (1987).
  124. F. Jansen and D. Kuhman, J. Vac. Sci. Technol. **6**, 13 (1988).
  125. W. Y. Kim, H. Tasaki, M. Konagai, and K. Takahashi, J. Appl. Phys. **61**, 3071 (1987).
  126. A. Catalano and G. Wood, J. Appl. Phys. **63**, 1220 (1988).
  127. Y. Kuwano, M. Ohnishi, S. Tsuda, Y. Nakashima, and N. Nakamura, Jpn. J. Appl. Phys. **21**, 413 (1982).
  128. R. A. Street, J. Zesch, and M. J. Thompson, Appl. Phys. Lett. **43**, 672 (1983).
  129. M. Ohnishi, H. Nishiwaki, E. Enomoto, Y. Nakashima, S. Tsuda, T. Takahama, H. Tarui, M. Tanaka, H. Dojo, and Y. Kuwano, J. Non-Cryst. Solids **59/60**, 1107 (1983).
  130. S. Nakano, S. Tsuda, H. Tarui, T. Takahama, H. Haku, K. Watanabe, M. Nishikuni, Y. Hishikawa, and Y. Kuwano, Mater. Res. Soc. Symp. Proc. **70**, 511 (1986).
  131. S. Tsuda, T. Takahama, M. Isomura, H. Tarui, Y. Nakashima, Y. Hishikawa, N. Nakamura, T. Matsuoka, H. Nishiwaki, S. Nakano, M. Ohnishi, and Y. Kuwano, Jpn. J. Appl. Phys. **26**, 33 (1987).
  132. W. Luft, Ed., *Photovoltaic Safety*, Vol. 166 of *AIP Conference Proceedings* (American Institute of Physics, New York, N.Y., U.S.A., 1988), .
  133. P. J. Hargis Jr., K. E. Greenberg, P. A. Miller, J. B. Gerardo, J. R. Torczynski, M. E. Riley, G. A. Hebner, J. R. Roberts, J. K. Olthoff, J. R. Whetstone, R. J. Van Brunt, M. A. Sobolewski, H. M. Anderson, M. P. Splichal, J. L. Mock, P. Bletzinger, A. Garscadden, R. A. Gottscho, G. Selwyn, M. Dalvie, J. E. Heidenreich, J. W. Butterbaugh, M. L. Brake, M. L. Passow, J. Pender, A. Lujan, M. E. Elta, D. B. Graves, H. H. Sawin, M. J. Kushner, J. T. Verdeyen, R. Horwath, and T. R. Turner, Rev. Sci. Instrum. **65**, 140 (1994).
  134. B. Chapman, *Glow Discharge Processes – Sputtering and Plasma Etching* (Wiley, New York, NY, U.S.A., 1980).
  135. A. Gallagher, Mater. Res. Soc. Symp. Proc. **70**, 3 (1986).
  136. J. Perrin, Y. Takeda, N. Hirano, Y. Takeuchi, and A. Matsuda, Surf. Sci. **210**, 114 (1989).

137. A. Matsuda, K. Nomoto, Y. Takeuchi, A. Suzuki, A. Yuuki, and J. Perrin, *Surf. Sci.* **227**, 50 (1990).
138. A. Matsuda and K. Tanaka, *J. Appl. Phys.* **60**, 2351 (1986).
139. T. Ichimura, T. Ihara, T. Hama, M. Ohsawa, H. Sakai, and Y. Uchida, *Jpn. J. Appl. Phys.* **25**, L276 (1986).
140. T. Hamasaki, M. Ueda, A. Chayahara, M. Hirose, and Y. Osaka, *Appl. Phys. Lett.* **44**, 600 (1984).
141. T. Hamasaki, M. Ueda, A. Chayahara, M. Hirose, and Y. Osaka, *Appl. Phys. Lett.* **44**, 1049 (1984).
142. P. Roca i Cabarrocas, J. B. Chevrier, J. Huc, A. Lloret, J. Y. Parey, and J. P. M. Schmitt, *J. Vac. Sci. Technol. A* **9**, 2331 (1991).
143. M. Taniguchi, M. Hirose, T. Hamasaki, and Y. Osaka, *Appl. Phys. Lett.* **37**, 787 (1980).
144. M. Ohnishi, H. Nishiwaki, K. Uchihashi, K. Yoshida, M. Tanaka, K. Ninomiya, M. Nishikuni, N. Nakanura, S. Tsuda, S. Nakano, T. Yazaki, and Y. Kuwano, *Jpn. J. Appl. Phys.* **27**, 40 (1988).
145. F. N. Boulitrop, N. Proust, J. Magarino, E. Criton, J. F. Peray, and M. Dupre, *J. Appl. Phys.* **58**, 3494 (1985).
146. H. Chatham and P. K. Bhat, *Mater. Res. Soc. Symp. Proc.* **149**, 447 (1989).
147. H. Chatham, P. Bhat, A. Benson, and C. Matovich, *J. Non-Cryst. Solids* **115**, 201 (1989).
148. P. M. Meijer, J. D. P. Passchier, W. J. Goedheer, J. Bezemer, and W. G. J. H. M. van Sark, *Appl. Phys. Lett.* **64**, 1780 (1994).
149. W. J. Goedheer, P. M. Meijer, J. Bezemer, J. D. P. Passchier, and W. G. J. H. M. van Sark, *IEEE Trans. Plasma Sci.* **PS-23**, 644 (1995).
150. W. G. J. H. M. van Sark, H. Meiling, J. Bezemer, M. B. von der Linden, R. E. I. Schropp, and W. F. van der Weg, *Sol. Energy Mater. Sol. Cells* **45**, 57 (1997).
151. H. Meiling, W. G. J. H. M. van Sark, J. Bezemer, and W. van der Weg, *J. Appl. Phys.* **80**, 3546 (1996).
152. J. P. M. Schmitt, J. Meot, P. Roubeau, and P. Parrens, in *Proceedings of the Eighth E.C. Photovoltaic Solar Energy Conference, Florence, Italy, 1988*, edited by I. Solomon, B. Eguer, and P. Helm (Kluwer Academic, Dordrecht, the Netherlands, 1989), p. 964.
153. S. Tsuda, S. Sakai, and S. Nakano, *Appl. Surf. Sci.* **113/114**, 734 (1997).
154. J. Yang, A. Banerjee, S. Sugiyama, and S. Guha, *Appl. Phys. Lett.* **70**, 2975 (1997).
155. R. E. I. Schropp, C. H. M. van der Werf, M. Zeman, M. C. M. van de Sanden, C. I. M. A. Spee, E. Middelman, L. V. de Jonge-Meschaninova, A. A. M. van der Zijden, M. M. Besselink, R. J. Severens, J. Winkeler, and G. J. Jongerden, *Mater. Res. Soc. Symp. Proc.* **557**, 713 (1999).
156. S. Fujikake, K. Tabuchi, T. Yoshida, Y. Ichikawa, and H. Sakai, *Mater. Res. Soc. Symp. Proc.* **377**, 609 (1995).
157. T. Yoshida, S. Fujikake, S. Kato, M. Tanda, K. Tabuchi, A. Takano, Y. Ichikawa, and H. Sakai, *Sol. Energy Mater. Sol. Cells* **48**, 383 (1997).
158. E. Middelman, E. van Andel, P. M. G. M. Peters, L. V. de Jonge-Meschaninova, R. J. Severens, G. J. Jongerden, R. E. I. Schropp, H. Meiling, M. Zeman, M. C. M. van de Sanden, A. Kuipers, and C. I. M. A. Spee, in *Proceedings of the 2<sup>nd</sup> World Conference on Photovoltaic Energy Conversion, Vienna, Austria*, edited by J. Schmid, H. A. Ossensbrink, P. Helm, H. Ehmman, and E. D. Dunlop (European Commission Joint Research Centre, Ispra, Italy, 1998), p. 816.

159. A. Madan, P. Rava, R. E. I. Schropp, and B. von Roedern, *Appl. Surf. Sci.* **70/71**, 716 (1993).
160. W. G. J. H. M. van Sark, J. Bezemer, M. Kars, M. Kuiper, E. A. G. Hamers, and W. F. van der Weg, in *Proceedings of the Twelfth E.C. Photovoltaic Solar Energy Conference, Amsterdam, the Netherlands, 1994*, edited by R. Hill, W. Palz, and P. Helm (H.S. Stephens & Associates, Bedford, U.K., 1994), p. 350.
161. E. A. G. Hamers, W. G. J. H. M. van Sark, J. Bezemer, W. J. Goedheer, and W. F. van der Weg, *Int. J. Mass Spectrom. and Ion Processes* **173**, 91 (1998).
162. W. G. J. H. M. van Sark, J. Bezemer, and W. J. Goedheer, unpublished results.
163. E. A. G. Hamers, Ph.D. thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998.
164. H. Shirai, D. Das, J. Hanna, and I. Shimizu, *Appl. Phys. Lett.* **59**, 1096 (1991).
165. W. G. J. H. M. van Sark, J. Bezemer, P. G. van de Vliet, M. Kars, and W. F. van der Weg, in *Proceedings of the Twelfth E.C. Photovoltaic Solar Energy Conference, Amsterdam, the Netherlands, 1994*, edited by R. Hill, W. Palz, and P. Helm (H.S. Stephens & Associates, Bedford, U.K., 1994), p. 335.
166. H. Kausche, M. Möller, and R. Plättner, in *Proceedings of the Fifth E.C. Photovoltaic Solar Energy Conference, Athens, Greece, 1983* (Reidel, Dordrecht, the Netherlands, 1983).
167. A. Morimoto, M. Matsumoto, M. Kumeda, and T. Shimizu, *Jpn. J. Appl. Phys.* **29**, L1747 (1990).
168. A. Ricard, *Reactive Plasmas* (Société Française du Vide, Paris, France, 1996).
169. H. S. Butler and G. S. Kino, *Phys. Fluids* **6**, 1346 (1963).
170. K. Köhler, J. W. Coburn, D. E. Horne, E. Kay, and J. H. Keller, *J. Appl. Phys.* **57**, 59 (1985).
171. M. A. Liebermann and S. E. Savas, *J. Vac. Sci. Technol. A* **16**, 1632 (1990).
172. M. V. Alves, M. A. Liebermann, V. Vahedi, and C. K. Birdsall, *J. Appl. Phys.* **69**, 3823 (1991).
173. J. Perrin, in *Plasma Deposition of Amorphous Silicon-Based Materials*, edited by G. Bruno, P. Capezzuto, and A. Madan (Academic Press, Boston, MA, U.S.A., 1995), Chap. 4, p. 177.
174. P. M. Meijer and W. J. Goedheer, *IEEE Trans. Plasma Sci.* **PS-19**, 170 (1991).
175. P. M. Meijer, Ph.D. thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1991.
176. K. Köhler, D. E. Horne, and J. W. Coburn, *J. Appl. Phys.* **58**, 3350 (1985).
177. A. D. Kuypers and H. J. Hopman, *J. Appl. Phys.* **63**, 1894 (1988).
178. W. J. Goedheer, private communication.
179. J. D. P. Passchier, Ph.D. thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1994.
180. S. Rauf and M. J. Kushner, *J. Appl. Phys.* **83**, 5087 (1998).
181. V. A. Godyak and R. B. Piejak, *J. Vac. Sci. Technol. A* **8**, 3833 (1990).
182. C. M. Horwitz, *J. Vac. Sci. Technol. A* **1**, 1795 (1983).
183. F. Finger, U. Kroll, V. Viret, A. Shah, W. Beyer, X.-M. Tang, J. Weber, A. A. Howling, and C. Hollenstein, *J. Appl. Phys.* **71**, 5665 (1992).
184. C. Böhm and J. Perrin, *J. Phys. D: Appl. Phys.* **24**, 865 (1991).
185. K. M. H. Maessen, Ph.D. thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1988.
186. M. J. M. Pruppers, Ph.D. thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1988.
187. S. Ishihara, K. Masatoshi, H. Takashi, W. Kiyotaka, T. Arita, and K. Mori, *J. Appl. Phys.* **62**, 485 (1987).
188. V. A. Godyak and R. B. Piejak, *Phys. Rev. Lett.* **65**, 996 (1990).

189. G. J. Nienhuis, Ph.D. thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998.
190. M. J. Kushner, *J. Appl. Phys.* **63**, 2532 (1988).
191. G. J. Nienhuis, W. J. Goedheer, E. A. G. Hamers, W. G. J. H. M. van Sark, and J. Bezemer, *J. Appl. Phys.* **82**, 2060 (1997).
192. J. Perrin, O. Leroy, and M. C. Bordage, *Contrib. Plasma Phys.* **36**, 3 (1996).
193. A. V. Phelps, *J. Phys. Chem. Ref. Data* **19**, 653 (1990).
194. H. Tawara, Y. Itikawa, H. Nishimura, and M. Yoshino, *J. Phys. Chem. Ref. Data* **19**, 617 (1990).
195. L. Layeillon, P. Duverneuil, J. P. Couderc, and B. Despax, *Plasma Sources Sci. Technol.* **3**, 61 (1994).
196. L. Layeillon, P. Duverneuil, J. P. Couderc, and B. Despax, *Plasma Sources Sci. Technol.* **3**, 72 (1994).
197. J. R. Doyle, D. A. Doughty, and A. Gallagher, *J. Appl. Phys.* **68**, 4375 (1990).
198. P. Kae-Nune, J. Perrin, J. Guillon, and J. Jolly, *Jpn. J. Appl. Phys.* **33**, 4303 (1994).
199. E. R. Austin and F. W. Lampe, *J. Phys. Chem.* **81**, 1134 (1977).
200. T. L. Pollock, H. S. Sandhu, A. Jodhan, and O. P. Strausz, *J. Am. Chem. Soc.* **95**, 1017 (1973).
201. J. Perrin, J. P. M. Schmitt, G. De Rosny, B. Drévilion, J. Huc, and A. Lloret, *Chem. Phys.* **73**, 383 (1982).
202. G. C. A. Perkins, E. R. Austin, and F. W. Lampe, *J. Am. Chem. Soc.* **101**, 1109 (1979).
203. M. Hertl, N. Dorval, O. Leroy, J. Jolly, and M. Péalet, *Plasma Sources Sci. Technol.* **7**, 130 (1998).
204. J. R. Doyle, D. A. Doughty, and A. Gallagher, *J. Appl. Phys.* **71**, 4771 (1992).
205. A. G. Engelhardt and A. V. Phelps, *Phys. Rev.* **131**, 2115 (1963).
206. E. Krishnakumar and S. K. Srivastava, *Contrib. Plasma Phys.* **35**, 395 (1995).
207. H. Tawara and T. Kato, *At. Data Nucl. Data Tables* **36**, 167 (1987).
208. P. Haaland, *J. Chem. Phys.* **93**, 4066 (1990).
209. J. M. Wadehra and J. N. Bardsley, *Phys. Rev. Lett.* **41**, 1795 (1978).
210. E. Krishnakumar, S. K. Srivastava, and I. Iga, *Int. J. Mass Spectrom. Ion Processes* **103**, 107 (1991).
211. J. Perrin, *J. Phys. D: Appl. Phys.* **26**, 1662 (1993).
212. W. Kurachi and Y. Nakamura, *J. Phys. D: Appl. Phys.* **22**, 107 (1989).
213. H. Ehrhardt, L. Langhans, F. Linder, and H. Taylor, *Phys. Rev.* **173**, 222 (1968).
214. A. P. Hickman, *J. Chem. Phys.* **70**, 4872 (1979).
215. N. Itabashi, K. Kato, N. Nishiwaki, T. Goto, C. Yamada, and E. Hirota, *Jpn. J. Appl. Phys.* **28**, L325 (1989).
216. J. M. S. Henis, G. W. Steward, M. K. Tripodi, and P. P. Gaspar, *J. Chem. Phys.* **57**, 389 (1972).
217. P. Kae-Nune, J. Perrin, J. Jolly, and J. Guillon, *Surf. Sci.* **360**, L495 (1996).
218. T. Fuyuki, B. Allain, and J. Perrin, *J. Appl. Phys.* **68**, 3322 (1990).
219. E. W. McDaniel and E. A. Mason, *The Mobility and Diffusion of Ions in Gases* (Wiley, New York, NY, U.S.A., 1973).
220. J. D. P. Passchier and W. J. Goedheer, *J. Appl. Phys.* **74**, 3744 (1993).
221. J. P. Boeuf and L. C. Pitchford, *Phys. Rev. E* **51**, 1376 (1995).
222. C. K. Birdsall, *IEEE Trans. Plasma Sci.* **19**, 65 (1991).
223. J. P. Boeuf and P. Belenguer, *J. Appl. Phys.* **71**, 4751 (1992).
224. M. Yan and W. J. Goedheer, *Plasma Sources Sci. Technol.* **8**, 349 (1999).

225. N. Sato and H. Tagashira, IEEE Trans. Plasma. Sci, **PS-19**, 10102 (1991).
226. N. V. Mantzaris, A. Boudouvis, and E. Gogolides, J. Appl. Phys. **77**, 6169 (1995).
227. M. J. Kushner, Solid State Technol. **6**, 135 (1996).
228. G. G. Lister, Vacuum **45**, 525 (1994).
229. E. Gogolides and H. H. Sawin, J. Appl. Phys. **72**, 3971 (1992).
230. P. M. Meijer, W. J. Goedheer, and J. D. P. Passchier, Phys. Rev. A **45**, 1098 (1992).
231. I. P. Shkarofsky, T. W. Johnston, and M. P. Bachynski, *The Particle Kinetics of Plasmas* (Addison-Wesley, Reading, MA, U.S.A., 1966).
232. G. J. Nienhuis and W. J. Goedheer, Plasma Sources Sci. Technol. **8**, 295 (1999).
233. A. D. Richards, B. E. Thompson, and H. H. Sawin, Appl. Phys. Lett. **50**, 492 (1987).
234. D. P. Lymberopoulos and D. J. Economou, J. Phys. D **28**, 727 (1995).
235. E. A. G. Hamers, W. G. J. H. M. van Sark, J. Bezemer, W. F. van der Weg, and W. J. Goedheer, Mater. Res. Soc. Symp. Proc. **420**, 461 (1996).
236. M. Heintze and R. Zedlitz, J. Non-Cryst. Solids **198-200**, 1038 (1996).
237. E. W. McDaniel, *Collision Phenomena in Ionized Gases* (Wiley, New York, NY, U.S.A., 1964).
238. P. J. Chantry, J. Appl. Phys. **62**, 1141 (1987).
239. M. S. Barnes, T. J. Cotler, and M. E. Elta, J. Comput. Phys. **77**, 53 (1988).
240. W. L. Morgan, Plasma Chem. Plasma Process. **12**, 477 (1992).
241. D. L. Sharfetter and H. K. Gummel, IEEE Trans. Electron Devices **ED-16**, 64 (1969).
242. W. Hackbusch, *Multigrid Methods and Applications* (Springer, Berlin, Germany, 1985).
243. J. D. P. Passchier, Ph.D. thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1994.
244. A. A. Fridman, L. Boufendi, T. Hbid, B. V. Potapkin, and A. Bouchoule, J. Appl. Phys. **79**, 1303 (1996).
245. J. Perrin, P. Roca i Cabarrocas, B. Allain, and J.-M. Friedt, Jpn. J. Appl. Phys. **27**, 2041 (1988).
246. J. L. Andújar, E. Bertran, A. Canillas, C. Roch, and J. L. Morenza, J. Vac. Sci. Technol. A **9**, 2216 (1991).
247. W. G. J. H. M. van Sark, J. Bezemer, E. M. B. Heller, M. Kars, and W. F. van der Weg, Mater. Res. Soc. Symp. Proc. **377**, 3 (1995).
248. G. Oversluizen and W. H. M. Lodders, J. Appl. Phys. **83**, 8002 (1998).
249. M. Heintze and R. Zedlitz, Prog. Photovolt.: Res. and Appl. **1**, 213 (1993).
250. A. A. Howling, J.-L. Drier, C. Hollenstein, U. Kroll, and F. Finger, J. Vac. Sci. Technol. A **10**, 1080 (1992).
251. W. G. J. H. M. van Sark, J. Bezemer, and W. F. van der Weg, Surf. Coat. Technol. **74-75**, 63 (1995).
252. L. Sansonnens, A. A. Howling, and C. Hollenstein, Plasma Sources Sci. Technol. **7**, 114 (1998).
253. M. Capitelli, C. Gorse, R. Winkler, and J. Wilhelm, Plasma Chem. Plasma Process. **8**, 399 (1988).
254. H. Curtins, N. Wyrsh, N. Favre, and A. V. Shah, Plasma Chem. Plasma Process. **7**, 267 (1987).
255. L. A. Pinnaduwa, W. Ding, and D. L. McCorkle, Appl. Phys. Lett. **71**, 3634 (1997).
256. L. A. Pinnaduwa and P. G. Datskos, J. Appl. Phys. **81**, 7715 (1997).
257. O. Leroy, G. Gousset, L. L. Alves, J. Perrin, and J. Jolly, Plasma Sources Sci. Technol. **7**, 348 (1998).
258. J. P. M. Schmitt, Thin Solid Films **174**, 193 (1989).

259. L. Sansonnens, D. Franz, C. Hollenstein, A. A. Howling, J. Schmitt, E. Turlot, T. Emeraud, U. Kroll, J. Meier, and A. Shah, in *Proceedings of the 13th European Photovoltaic Solar Energy Conference, Nice, France*, edited by W. Freiesleben, W. Palz, H. A. Ossenbrink, and P. Helm (H.S. Stephens & Associates, Bedford, U. K., 1995), p. 276.
260. D. Vender and R. W. Boswell, *IEEE Trans. Plasma Sci.* **18**, 725 (1990).
261. C. K. Birdsall, *IEEE Trans. Plasma. Sci.*, **19**, 65 (1991).
262. C. K. Birdsall and A. B. Langdon, *Plasma Physics Via Computer Simulation* (Adam Hilger, Bristol, U.K., 1991).
263. V. Vahedi, *XPDP1 Manual* (UCLA Berkeley, Berkeley, CA, U.S.A., 1994).
264. J. D. P. Passchier, Technical Report No. GF95.08, Utrecht University, Utrecht, The Netherlands, 1995.
265. W. G. J. H. M. van Sark and J. Bezemer, Technical report, Utrecht University, Utrecht, The Netherlands, 1995.
266. Y. Ohmori, M. Shimozuma, and H. Tagashira, *J. Phys. D: Appl. Phys.* **19**, 1029 (1986).
267. E. A. Mason and E. W. McDaniel, *Transport Properties of Ions in Gases* (Wiley, New York, NY, U.S.A., 1988).
268. D. B. Graves, *J. Appl. Phys.* **62**, 88 (1987).
269. J.-P. Boeuf, *Phys. Rev. A* **36**, 2782 (1987).
270. E. Gogolides and H. H. Sawin, *J. Appl. Phys.* **72**, 3971 (1992).
271. A. Manenschijn and W. J. Goedheer, *J. Appl. Phys.* **69**, 2923 (1991).
272. V. Vahedi, C. K. Birdsall, M. A. Lieberman, G. Dipeso, and T. D. Rognlien, *Phys. Fluids B* **5**, 2719 (1993).
273. K. Nanbu and Y. Kitatani, *Vacuum* **47**, 1023 (1996).
274. A. K. Jain and D. G. Thompson, *J. Phys. B: At. Mol. Phys.* **20**, L389 (1987).
275. H. Tawara, Y. Itikawa, H. Nishimura, and M. Yoshino, *J. Phys. Chem. Ref. Data* **19**, 621 (1990).
276. M. R. Wertheimer and M. Moisan, *J. Vac. Sci. Technol. A* **3**, 2643 (1985).
277. H. Curtins, N. Wyrsh, M. Favre, and A. V. Shah, *Plasma Chem. Plasma Process.* **7**, 267 (1987).
278. V. A. Godyak, *Sov. Phys.-Tech. Phys.* **16**, 1073 (1973).
279. M. Heintze and R. Zedlitz, *J. Phys. D* **26**, 1781 (1993).
280. M. Heintze and R. Zedlitz, *J. Non-Cryst. Solids* **164–166**, 55 (1993).
281. J. Dutta, W. Bacsá, and C. Hollenstein, *J. Appl. Phys.* **72**, 3220 (1992).
282. P. Mérel, M. Chaker, M. Moisan, A. Ricard, and M. Tabbal, *J. Appl. Phys.* **72**, 3220 (1992).
283. S. Vepřek and M. G. J. Vepřek-Heijman, *Appl. Phys. Lett.* **56**, 1766 (1990).
284. J. Dutta, U. Kroll, P. Chabloz, A. Shah, A. A. Howling, J.-L. Dorier, and C. Hollenstein, *J. Appl. Phys.* **72**, 3220 (1992).
285. T. Takagi, *J. Vac. Sci. Technol. A* **2**, 282 (1983).
286. G. Turban, B. Drévilion, D. S. Mataras, and D. E. Rapakoulias, in *Plasma Deposition of Amorphous Silicon-Based Materials*, edited by G. Bruno, P. Capezzuto, and A. Madan (Academic Press, Boston, MA, U.S.A., 1995), Chap. 2, p. 63.
287. A. Matsuda and N. Hata, in *Glow-Discharge Hydrogenated Amorphous Silicon*, edited by K. Tanaka (KTK Scientific Publishers, Tokyo, Japan, 1989), Chap. 2, p. 9.
288. A. Matsuda, T. Kaga, H. Tanaka, L. Malhotra, and K. Tanaka, *Jpn. J. Appl. Phys.* **22**, L115 (1983).
289. D. Mataras, S. Cavadias, and D. Rapakoulias, *J. Appl. Phys.* **66**, 119 (1989).

290. D. Mataras, S. Cavadias, and D. Rapakoulias, *J. Vac. Sci. Technol. A* **11**, 664 (1993).
291. N. Itabashi, K. Kato, N. Nishiwaki, T. Goto, C. Yamada, and E. Hirota, *Jpn. J. Appl. Phys.* **27**, L1565 (1988).
292. N. Itabashi, N. Nishiwaki, M. Magane, S. Naito, T. Goto, A. Matsuda, C. Yamada, and E. Hirota, *Jpn. J. Appl. Phys.* **29**, L505 (1990).
293. N. Hershkowitz, M. H. Cho, C. H. Nam, and T. Intrator, *Plasma Chem. Plasma Process.* **8**, 35 (1988).
294. P. Awakowicz, *Materials Science Forum* **287-288**, 3 (1998).
295. R. van der Heijden, Technical Report No. GF96.07, Utrecht University, Utrecht, The Netherlands (unpublished), (1996).
296. E. R. Mosburg, R. C. Kerns, and J. R. Abelson, *J. Appl. Phys.* **54**, 4916 (1983).
297. G. Bruno, P. Capezzuto, G. Cicala, P. Manodoro, and V. Tassielli, *IEEE Trans. Plasma Sci.* **PS-18**, 934 (1990).
298. S. Veprek and M. Heintze, *Plasma Chem. Plasma Process.* **10**, 3 (1990).
299. J. Cárabe, J. J. Gandía, and M. T. Gutiérrez, *J. Appl. Phys.* **73**, 4618 (1993).
300. N. Spiliopoulos, D. Mataras, and D. E. Rapakoulias, *J. Electrochem. Soc.* **144**, 634 (1997).
301. E. A. G. Hamers, W. G. J. H. M. van Sark, J. Bezemer, H. Meiling, and W. F. van der Weg, *J. Non-Cryst. Solids* **226**, 205 (1998).
302. G. Nolet, *J. Electrochem. Soc.* **122**, 1030 (1975).
303. G. Turban, Y. Catherine, and B. Grolleau, *Thin Solid Films* **60**, 147 (1979).
304. J. J. Wagner and S. Vepřek, *Plasma Chem. Plasma Process.* **2**, 95 (1982).
305. G. Turban, Y. Catherine, and B. Grolleau, *Plasma Chem. Plasma Process.* **2**, 61 (1982).
306. P. A. Longeway, H. A. Weakliem, and R. D. Estes, *J. Appl. Phys.* **57**, 5499 (1985).
307. R. Robertson, D. Hils, H. Chatham, and A. Gallagher, *Appl. Phys. Lett.* **43**, 544 (1983).
308. M. Heintze and S. Vepřek, *Appl. Phys. Lett.* **54**, 1320 (1989).
309. G. Turban, Y. Catherine, and B. Grolleau, *Thin Solid Films* **77**, 287 (1981).
310. G. Turban, *Pure Appl. Chem.* **56**, 215 (1984).
311. P. Kae-Nune, J. Perrin, J. Guillon, and J. Jolly, *Plasma Sources Sci. Technol.* **4**, 250 (1995).
312. R. Robertson and A. Gallagher, *J. Appl. Phys.* **59**, 3402 (1986).
313. F. A. Baiocchi, R. C. Wetzels, and R. S. Freund, *Phys. Rev. Lett.* **53**, 771 (1984).
314. R. S. Freund, R. C. Wetzels, R. J. Shul, and T. R. Hayes, *Phys. Rev. A* **41**, 3575 (1990).
315. Hiden Analytical Ltd., 240 Europa Boulevard, Gemini Business Park, Warrington WA5 5UN, England.
316. P. Ho, W. G. Breiland, and R. J. Buss, *J. Chem. Phys.* **91**, 2627 (1989).
317. J. Perrin, M. Shiratani, P. Kae-Nune, H. Videlot, J. Jolly, and J. Guillon, *J. Vac. Sci. Technol. A* **16**, 278 (1998).
318. J. Perrin, C. Böhm, R. Eternadi, and A. Lloret, *Plasma Sources Sci. Technol.* **3**, 252 (1994).
319. G. Turban, Y. Catherine, and B. Grolleau, *Thin Solid Films* **67**, 309 (1980).
320. H. Chatham, D. Hils, R. Robertson, and A. Gallagher, *J. Chem. Phys.* **81**, 1770 (1984).
321. A. A. Howling, L. Sansonnens, J.-L. Dorier, and C. Hollenstein, *J. Appl. Phys.* **75**, 1340 (1994).
322. L. Boufendi and T. Bouchoule, *J. Vac. Sci. Technol. A* **14**, 572 (1996).
323. B. E. Thompson, K. D. Allen, A. D. Richards, and H. H. Sawin, *J. Appl. Phys.* **59**, 1890 (1986).

324. J. Liu, G. L. Huppert, and H. H. Sawin, *J. Appl. Phys.* **68**, 3916 (1990).
325. C. Wild and P. Koidl, *J. Appl. Phys.* **69**, 2909 (1991).
326. R. J. M. M. Snijkers, M. J. M. Sambeek, G. M. W. Kroesen, and F. J. de Hoog, *Appl. Phys. Lett.* **63**, 308 (1993).
327. J. K. Olthoff, R. J. van Brunt, S. B. Radovanov, J. A. Rees, and R. Surowiec, *J. Appl. Phys.* **75**, 115 (1994).
328. M. Fivaz, S. Brunner, W. Swarzenbach, A. A. Howling, and C. Hollenstein, *Plasma Sources Sci. Technol.* **4**, 373 (1995).
329. D. A. Dahl, J. E. Delmore, and A. D. Appelhans, *Rev. Sci. Instrum.* **61**, 607 (1990).
330. M. L. Mandich, W. D. Reents, and K. D. Kolenbrander, *J. Chem. Phys.* **92**, 437 (1990).
331. W. D. Reents and M. L. Mandich, *J. Chem. Phys.* **96**, 4429 (1992).
332. E. A. G. Hamers, J. Bezemer, H. Meiling, W. G. J. H. M. van Sark, and W. F. van der Weg, *Mater. Res. Soc. Symp. Proc.* **467**, 603 (1997).
333. R. J. M. M. Snijkers, Ph.D. thesis, Technische Universiteit Eindhoven, Eindhoven, the Netherlands, 1993.
334. E. R. Fisher and P. B. Armentrout, *J. Chem. Phys.* **93**, 4858 (1990).
335. W. N. Allen, T. M. H. Cheng, and F. W. Lampe, *J. Chem. Phys.* **66**, 3371 (1977).
336. R. M. A. Azzam and N. M. Bashara, *Ellipsometry and Polarized Light* (North Holland, Amsterdam, the Netherlands, 1977).
337. B. Dré villon, J. Perrin, R. Marbot, A. Violet, and J. L. Dalby, *Rev. Sci. Instrum.* **53**, 969 (1982).
338. N. Blayo and B. Dré villon, *Appl. Phys. Lett.* **59**, 950 (1991).
339. R. W. Collins, *Rev. Sci. Instrum.* **61**, 2029 (1990).
340. H. Fujiwara, Y. Toyoshima, M. Kondo, and A. Matsuda, *Phys. Rev. B* **60**, 13598 (1999).
341. R. Brenot, B. Dré villon, P. Bulkin, P. Roca i Cabarrocas, and R. Vanderhaghen, *Appl. Surf. Sci.* **154–155**, 283 (2000).
342. R. W. Collins, J. Koh, A. S. Ferlauto, P. I. Rovira, Y. Lee, R. J. Koval, and C. R. Wronski, *Thin Solid Films* **364**, 129 (2000).
343. B. Dré villon, *Prog. Cryst. Growth Charact. Mater.* **27**, 1 (1993).
344. R. Ossikovski and B. Dré villon, *Phys. Rev. B* **54**, 10530 (1996).
345. H. Fujiwara, J. Koh, P. I. Rovira, and R. W. Collins, *Phys. Rev. B* **61**, 10832 (2000).
346. D. E. Aspnes, J. B. Theeten, and F. Hottier, *Phys. Rev. B* **20**, 3292 (1979).
347. A. Canillas, E. Bertran, J. L. Andújar, and B. Dré villon, *J. Appl. Phys.* **68**, 2752 (1990).
348. Y. Lu, S. Kim, M. Gunes, Y. Lee, C. R. Wronski, and R. W. Collins, *Mater. Res. Soc. Symp. Proc.* **336**, 595 (1994).
349. D. V. Tsu, B. S. Chao, S. R. Ovshinsky, S. Guha, and J. Yang, *Appl. Phys. Lett.* **71**, 1317 (1997).
350. J. Koh, Y. Lee, H. Fujiwara, C. R. Wronski, and R. W. Collins, *Appl. Phys. Lett.* **73**, 1526 (1998).
351. S. Guha, J. Yang, D. L. Williamson, Y. Lubianiker, J. D. Cohen, and A. H. Mahan, *Appl. Phys. Lett.* **74**, 1860 (1999).
352. A. H. Mahan, J. Yang, S. Guha, and D. L. Williamson, *Phys. Rev. B* **61**, 1677 (2000).
353. H. Fujiwara, J. Koh, C. R. Wronski, and R. W. Collins, *Appl. Phys. Lett.* **70**, 2150 (1997).
354. H. Fujiwara, J. Koh, C. R. Wronski, and R. W. Collins, *Appl. Phys. Lett.* **72**, 2993 (1998).
355. H. Fujiwara, J. Koh, C. R. Wronski, and R. W. Collins, *Appl. Phys. Lett.* **74**, 3687

- (1999).
356. N. Blayo and B. Dré villon, *J. Non-Cryst. Solids* **137&138**, 771 (1991).
  357. A. Matsuda, *J. Vac. Sci. Technol. A* **16**, 365 (1998).
  358. B. Dré villon, J. Perrin, J. M. Siefert, J. Huc, A. Lloret, G. de Rosny, and J. P. M. Schmitt, *Appl. Phys. Lett.* **42**, 801 (1983).
  359. R. A. Street, J. C. Knights, and D. K. Biegelsen, *Phys. Rev. B* **19**, 3027 (1978).
  360. J. C. Knights and R. A. Lujan, *Appl. Phys. Lett.* **35**, 244 (1979).
  361. S. Nishikawa, H. Kakinuma, T. Watanabe, and K. Nihei, *Jpn. J. Appl. Phys.* **24**, 639 (1985).
  362. G. D. Cody, T. Tiedje, B. Abeles, B. Brooks, and Y. Goldstein, *Phys. Rev. Lett.* **47**, 1480 (1981).
  363. K. M. H. Maessen, M. J. M. Pruppers, J. Bezemer, F. H. P. M. Habraken, and W. F. van der Weg, *Mater. Res. Soc. Symp. Proc.* **95**, 201 (1987).
  364. K. Tanaka, K. Nakagawa, A. Matsuda, M. Matsumura, H. Yamamoto, S. Yamasaki, H. Okushi, and S. Izima, *Jpn. J. Appl. Phys.* **20**, 267 (1981).
  365. P. Sichanugrist, M. Konagai, and K. Takahashi, *Jpn. J. Appl. Phys.* **25**, 440 (1986).
  366. M. Hirose, *J. Phys. (Paris), Colloq.* **42**, C4 (1981).
  367. P. Chaudhuri, S. Ray, and A. K. Barua, *Thin Solid Films* **113**, 261 (1984).
  368. J. Shirafuji, S. Nagata, and M. Kuwagaki, *J. Appl. Phys.* **58**, 3661 (1985).
  369. S. Okamoto, Y. Hishikawa, and S. Tsuda, *Jpn. J. Appl. Phys.* **35**, 26 (1996).
  370. H. Meiling, R. E. I. Schropp, W. G. J. H. M. van Sark, J. Bezemer, and W. F. van der Weg, in *Proceedings of the Tenth E.C. Photovoltaic Solar Energy Conference, Lisbon, Portugal, 1991*, edited by A. Luque, G. Sala, W. Palz, G. Dos Santos, and P. Helm (Kluwer Academic, Dordrecht, the Netherlands, 1991), p. 339.
  371. J. P. M. Schmitt, *J. Non-Cryst. Solids* **59&60**, 649 (1983).
  372. C. C. Tsai, G. B. Anderson, and R. Thompson, *J. Non-Cryst. Solids* **137-138**, 637 (1991).
  373. P. Roca i Cabarrocas, S. Hamma, S. N. Sharma, G. Viera, E. Bertran, and J. Costa, *J. Non-Cryst. Solids* **227-230**, 871 (1998).
  374. C. Longeaud, J. P. Kleider, P. Roca i Cabarrocas, S. Hamma, R. Meaudre, and M. Meaudre, *J. Non-Cryst. Solids* **227-230**, 96 (1998).
  375. J. C. Knights, *Jpn. J. Appl. Phys.* **18**, 101 (1979).
  376. R. B. Wehrspohn, S. C. Deane, I. D. French, I. Gale, J. Hewett, M. J. Powell, and J. Robertson, *J. Appl. Phys.* **87**, 144 (2000).
  377. A. A. Howling, J.-L. Dorier, and C. Hollenstein, *Appl. Phys. Lett.* **62**, 1341 (1993).
  378. A. Bouchoule, Ed., *Dusty Plasmas: Physics, Chemistry, and Technological Impact in Plasma Processing* (Wiley, New York, NY, U.S.A., 1999).
  379. P. Roca i Cabarrocas, *J. Non-Cryst. Solids* **266-269**, 31 (2000).
  380. M. Otake, T. Kanai, T. Ifuku, H. Yajima, and B. Oda, *J. Non-Cryst. Solids* **198-200**, 875 (1996).
  381. D. M. Tanenbaum, A. L. Laracuente, and A. Gallagher, *Appl. Phys. Lett.* **68**, 705 (1996).
  382. P. Roca i Cabarrocas, *Mater. Res. Soc. Symp. Proc.* **507**, 855 (1998).
  383. E. Stoffels, W. W. Stoffels, G. M. W. Kroesen, and F. J. de Hoog, *Electron Technology* **31**, 255 (1998).
  384. C. Courteille, J.-L. Dorier, J. Dutta, C. Hollenstein, A. A. Howling, and T. Stoto, *J. Appl. Phys.* **78**, 61 (1995).
  385. S. Schmitt-Rink, D. A. B. Miller, and D. S. Chemla, *Phys. Rev. B* **35**, 8113 (1987).

386. A. J. Read, R. J. Needs, K. J. Nash, L. T. Canham, P. D. J. Calcott, and A. Qteish, *Phys. Rev. Lett.* **69**, 1232 (1992).
387. P. Roca i Cabarrocas, P. Gay, and A. Hadjadj, *J. Vac. Sci. Technol. A* **14**, 655 (1996).
388. A. Fontcuberta i Morral, R. Brenot, E. A. G. Hamers, R. Vanderhaghen, and P. Roca i Cabarrocas, *J. Non-Cryst. Solids* **266–269**, 48 (2000).
389. R. Butté, S. Vignoli, M. Meaudre, R. Meaudre, O. Marty, L. Saviot, and P. Roca i Cabarrocas, *J. Non-Cryst. Solids* **266–269**, 263 (2000).
390. Corning 7059 Material Information, Corning Glass Works, NY, U.S.A.
391. J. Wallinga, D. Knoesen, E. A. G. Hamers, W. G. J. H. M. van Sark, W. F. van der Weg, and R. E. I. Schropp, *Mater. Res. Soc. Symp. Proc.* **420**, 57 (1996).
392. M. B. Schubert and G. H. Bauer, in *Proceedings of the Twenty First IEEE Photovoltaic Specialists Conference, Kissimee, U.S.A., 1985* (IEEE, New York, N.Y., U.S.A., 1990), p. 1595.
393. Y. Hishikawa, M. Ohnishi, and Y. Kuwano, *Mater. Res. Soc. Symp. Proc.* **192**, 3 (1990).
394. A. Shin and S. Lee, *J. Non-Cryst. Solids* **260**, 245 (1999).
395. N. Maley and J. S. Lannin, *Phys. Rev. B* **31**, 5577 (1985).
396. A. J. M. Berntsen, W. G. J. H. M. van Sark, and W. F. van der Weg, *J. Appl. Phys.* **78**, 1964 (1995).
397. R. W. Collins and J. M. Cavese, *J. Appl. Phys.* **61**, 1869 (1987).
398. Y. Toyoshima, K. Arai, A. Matsuda, and K. Tanaka, *Appl. Phys. Lett.* **56**, 1540 (1990).
399. G. Ganguly and A. Matsuda, *J. Non-Cryst. Solids* **164–166**, 31 (1993).
400. S. Vepřek, F.-A. Sarrott, S. Rambert, and E. Taglauer, *J. Vac. Sci. Technol. A* **7**, 2614 (1989).
401. B. J. Garrison, M. T. Miller, and D. W. Bremmen, *Chem. Phys. Lett.* **146**, 513 (1988).
402. M. Kitabatake, P. Fons, and J. E. Greene, *J. Vac. Sci. Technol. A* **8**, 3726 (1990).
403. P. Roca i Cabarrocas, *Mater. Res. Soc. Symp. Proc.* **149**, 33 (1989).
404. G. Ganguly, T. Ikeda, I. Sakata, and A. Matsuda, *Mat. Res. Soc. Proc. Symp.* **420**, 347 (1996).
405. A. I. Kosarev, A. S. Smirnov, A. S. Abramov, A. J. Vinogradov, A. Y. Ustavshnikov, and M. V. Shutov, *J. Vac. Sci. Technol. A* **15**, 298 (1997).
406. A. S. Abramov, A. Y. Vinogradov, A. I. Kosarev, M. V. Shutov, A. S. Smimov, and K. E. Orlov, *Tech. Phys.* **43**, 180 (1998).
407. J. P. Harbison, A. J. Williams, and D. V. Lang, *J. Appl. Phys.* **55**, 946 (1984).
408. K. H. Müller, *J. Appl. Phys.* **62**, 1796 (1987).
409. J. Robertson, *J. Non-Cryst. Solids* **164–166**, 1115 (1993).
410. J. R. Abelson, *Appl. Phys. A* **56**, 493 (1993).
411. R. Biswas, *J. Appl. Phys.* **73**, 3295 (1993).
412. W. Möller, *Appl. Phys. A* **56**, 527 (1993).
413. M. Hoedemaker, J. van der Kuur, E. J. Melker, and B. J. Thijsse, *Nucl. Instr. Meth. B* **127/128**, 888 (1997).
414. J. F. Ziegler, J. P. Biersack, and U. Littmarck, *The Stopping and Range of Ions in Solids* (Pergamon, New York, N.Y., U.S.A., 1985).
415. B. Drévilion, J. Huc, and N. Boussarssar, *J. Non-Cryst. Solids* **59/60**, 735 (1983).
416. B. Drévilion and M. Toutlemonde, *Appl. Phys. Lett.* **59**, 950 (1985).
417. J. Perrin, Y. Takeda, N. Hirano, H. Matsuura, and A. Matsuda, *Jpn. J. Appl. Phys.* **28**, 5 (1989).
418. P. Roca i Cabarrocas, P. Morin, V. Chu, J. P. Conde, J. Z. Liu, H. R. Park, and S.

- Wagner, J. Appl. Phys. **69**, 2942 (1991).
419. E. A. G. Hamers, J. Bezemer, and W. F. van der Weg, Appl. Phys. Lett. **75**, 609 (1999).
420. D. A. Doughty, J. R. Doyle, G. H. Lin, and A. Gallagher, J. Appl. Phys. **67**, 6220 (1990).
421. J. K. Hirvonen, in *Materials and Processes for Surface and Interface Engineering*, edited by Y. Pauleau (Kluwer Academic Publishers, Dordrecht, The Netherlands, 1995), Chap. 9, p. 307.
422. J. E. Potts, E. M. Peterson, and J. A. McMillan, J. Appl. Phys. **52**, 6665 (1981).
423. H. Meiling, J. Bezemer, R. E. I. Schropp, and W. F. van der Weg, Mat. Res. Soc. Proc. Symp. **467**, 459 (1997).
424. S. Habermehl and G. Lucovsky, J. Vac. Sci. Technol. A **14**, 3024 (1996).
425. S. R. Kasi, H. Kang, C. S. Sass, and J. W. Rabalais, Surf. Sci. Rept. **10**, 1 (1989).
426. L. Hanley, H. Lim, D. G. Schultz, S. B. Wainhaus, P. de Sainte Claire, and W. L. Hase, Nucl. Instr. Meth. B **125**, 218 (1997).
427. H. Sugai, Y. Mitsuoka, and H. Toyoda, J. Vac. Sci. Technol. A **16**, 290 (1998).
428. K. H. Müller, Phys. Rev. B **35**, 7906 (1987).
429. J. W. Rabalais, A. H. Al-Bayati, K. J. Boyd, D. Marton, J. Kulik, Z. Zhang, and W. K. Chu, Phys. Rev. B **53**, 10781 (1996).
430. K. H. Müller, J. Appl. Phys. **59**, 2803 (1986).
431. J. E. Yehoda, B. Yang, K. Vedam, and R. Messier, J. Vac. Sci. Technol. A **6**, 1631 (1988).
432. R. J. Severens, M. C. M. van de Sanden, H. J. M. Verhoeven, J. Bastiaansen, and D. C. Schram, Mater. Res. Soc. Symp. Proc. **420**, 341 (1996).
433. M. C. M. van de Sanden, R. J. Severens, W. M. M. Kessels, F. van de Pas, L. van IJzendoorn, and D. C. Schram, Mater. Res. Soc. Proc. Symp. **467**, 621 (1997).
434. E. H. C. Ullersma, Ph.D. thesis, Utrecht University, Utrecht, the Netherlands, 1998.
435. R. Robertson and A. Gallagher, J. Chem. Phys. **85**, 3623 (1986).
436. A. Gallagher, J. Appl. Phys. **63**, 2406 (1988).
437. J. L. Guizot, K. Nomoto, and A. Matsuda, Surf. Sci. **244**, 22 (1991).
438. A. J. Flewitt, J. Robertson, and W. I. Milne, J. Appl. Phys. **85**, 8032 (1999).
439. J. Robertson, J. Appl. Phys. **87**, 2608 (2000).
440. G. Ganguly and A. Matsuda, Phys. Rev. B **47**, 3661 (1993).
441. K. Winer, Phys. Rev. B **41**, 7952 (1990).
442. R. A. Street, Phys. Rev. B **43**, 2454 (1991).
443. R. A. Street, Phys. Rev. B **44**, 10610 (1991).
444. R. A. Street and K. Winer, Phys. Rev. B **40**, 6236 (1989).
445. M. J. Powell and S. C. Deane, Phys. Rev. B **48**, 10815 (1993).
446. M. J. Powell and S. C. Deane, Phys. Rev. B **53**, 10121 (1996).
447. K. Gleason, K. S. Wang, M. K. Chen, and J. A. Reimer, J. Appl. Phys. **61**, 2866 (1987).
448. M. J. McCaughy and M. J. Kushner, J. Appl. Phys. **65**, 186 (1989).
449. S. Shirafuji, S. Nakajima, Y. F. Wang, T. Genji, and K. Tachibana, Jpn. J. Appl. Phys. **32**, 1546 (1993).
450. J. Robertson, Mater. Res. Soc. Proc. Symp. **609**, A1.4.1 (2000).
451. C. C. Tsai, J. C. Knights, G. Chang, and B. Wacker, J. Appl. Phys. **59**, 2998 (1986).
452. A. J. Flewitt, J. Robertson, and W. I. Milne, J. Non-Cryst. Solids **266–269**, 74 (2000).
453. A. von Keudell and J. R. Abelson, Jpn. J. Appl. Phys. **38**, 4002 (1999).
454. S. Ramalingam, D. Maroudas, and E. S. Aydill, J. Appl. Phys. **86**, 2872 (1999).
455. S. Yamasaki, T. Umeda, J. Isoya, and K. Tanaka, Appl. Phys. Lett. **70**, 1137 (1997).

456. D. A. Porter and K. E. Easterling, *Phase Transformations in Metals and Alloys* (Chapman & Hall, London, U.K., 1992).
457. P. Roca i Cabarrocas, Y. Bouizem, and M. L. Theye, *Philos. Mag. B* **65**, 1025 (1992).
458. C. M. Greenlief, S. M. Gates, and P. A. Holbert, *J. Vac. Sci. Technol. A* **7**, 1845 (1989).
459. W. B. Jackson, *J. Non-Cryst. Solids* **164–166**, 263 (1993).
460. W. Beyer, *J. Non-Cryst. Solids* **198–200**, 40 (1996).
461. W. Beyer and U. Zastrow, *Mater. Res. Soc. Symp. Proc.* **420**, 497 (1996).
462. C. G. Van de Walle, *Phys. Rev. B* **49**, 4579 (1994).
463. J. Robertson, C. W. Chen, M. J. Powell, and S. C. Deane, *J. Non-Cryst. Solids* **227–230**, 138 (1998).
464. K. J. Chang and J. D. Chadi, *Phys. Rev. Lett.* **62**, 937 (1989).
465. K. J. Chang and J. D. Chadi, *Phys. Rev. B* **40**, 11644 (1990).
466. Y. Miyoshi, Y. Yoshida, S. Miyazaki, and M. Hirose, *J. Non-Cryst. Solids* **198–200**, 1029 (1996).
467. J. Robertson and M. J. Powell, *Thin Solid Films* **337**, 32 (1999).
468. W. M. M. Kessels, R. J. Severens, M. C. M. van de Sanden, and D. C. Schram, *J. Non-Cryst. Solids* **227–230**, 133 (1998).
469. G. H. Lin, J. R. Doyle, M. Z. He, and A. Gallagher, *J. Appl. Phys.* **64**, 341 (1988).
470. P. Roca i Cabarrocas, *Appl. Phys. Lett.* **65**, 1674 (1995).
471. H. Shirai, J. Hanna, and I. Shimizu, *Jpn. J. Appl. Phys.* **30**, L679 (1991).
472. M. C. M. van de Sanden, W. M. M. Kessels, R. J. Severens, and D. C. Schram, *Plasma Phys. Controlled Fusion* **41**, A365 (1999).
473. M. C. M. van de Sanden, W. M. M. Kessels, R. J. Severens, , and D. C. Schram, *Plasma Phys. Control. Fusion* **41**, A365 (1999).
474. M. Stutzmann, *Philos. Mag. B* **60**, 531 (1989).
475. A. Shah, J. Dutta, N. Wyrsh, K. Prasad, H. Curtins, F. Finger, A. Howling, and C. Hollenstein, *Mater. Res. Soc. Symp. Proc.* **258**, 15 (1992).
476. C. M. Ferreira and J. Loureiro, *J. Phys. D* **17**, 1175 (1984).
477. J. Kuske, U. Stephan, O. Steinke, and S. Röhlecke, *Mater. Res. Soc. Symp. Proc.* **377**, 27 (1995).
478. U. Kroll, J. Meier, M. Goetz, A. Howling, J.-L. Dorier, J. Dutta, A. Shah, and C. Hollenstein, *J. Non-Cryst. Solids* **164–166**, 59 (1993).
479. H. Meiling, J. F. M. Westendorp, J. Hautala, Z. Saleh, and C. T. Malone, *Mater. Res. Soc. Symp. Proc.* **345**, 65 (1995).
480. S. Röhlecke, R. Tews, A. Kottwitz, and K. Schade, *Surf. Coat. Technol.* **74–75**, 259 (1995).
481. C. Beneking, *J. Appl. Phys.* **68**, 4461 (1990).
482. C. Beneking, *J. Appl. Phys.* **68**, 5435 (1990).
483. C. Beneking, F. Finger, and H. Wagner, in *Proceedings of the 11th European Photovoltaic Solar Energy Conference, Montreux, Switzerland*, edited by L. Guimarães, W. Palz, C. de Reyff, H. Kiess, and P. Helm (Harwood, Chur, Switzerland, 1993), p. 586.
484. M. J. Colgan, M. Meyyappan, and D. E. Murnick, *Plasma Sources Sci. Technol.* **3**, 181 (1994).
485. U. Kroll, Y. Ziegler, J. Meier, H. Keppner, and A. Shah, *Mater. Res. Soc. Symp. Proc.* **336**, 115 (1994).
486. E. A. G. Hamers, W. G. J. H. M. van Sark, J. Bezemer, W. J. Goedheer, and W. F. van der Weg, *Int. J. Mass Spectrom. and Ion Processes* **173**, 91 (1998).

487. W. G. J. H. M. van Sark, H. Meiling, E. A. G. Hamers, J. Bezemer, and W. F. van der Weg, *J. Vac. Sci. Technol. A* **15**, 654 (1997).
488. H. Meiling, W. G. J. H. M. van Sark, J. Bezemer, and W. van der Weg, *J. Appl. Phys.* **80**, 3160 (1996).
489. G. E. N. Landweer and J. Bezemer, in *Amorphous Silicon and Related Materials, Advances in Disordered Semiconductors 1*, edited by H. Fritzsche (World Scientific, Singapore, 1989).
490. M. Zhu, M. B. von der Linden, J. Bezemer, R. E. I. Schropp, and W. F. van der Weg, *J. Non-Cryst. Solids* **137–138**, 355 (1991).
491. M. Zhu, M. B. von der Linden, and W. F. van der Weg, *Mater. Res. Soc. Symp. Proc.* **336**, 449 (1994).
492. M. B. von der Linden, Ph.D. thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1994.
493. W. G. J. H. M. van Sark, J. Bezemer, and W. F. van der Weg, *J. Mater. Res.* **13**, 45 (1998).
494. R. E. I. Schropp, H. Meiling, W. G. J. H. M. van Sark, J. Stammeijer, J. Bezemer, and W. F. van der Weg, in *Proceedings of the Tenth E.C. Photovoltaic Solar Energy Conference, Lisbon, Portugal, 1991*, edited by A. Luque, G. Sala, W. Palz, G. Dos Santos, and P. Helm (Kluwer Academic, Dordrecht, the Netherlands, 1991), p. 1087.
495. S. J. Jones, D. L. Williamson, T. Liu, X. Deng, and M. Izu, *Mater. Res. Soc. Proc. Symp.* **609**, A7.4.1 (2000).
496. C. C. Tsai, R. Thompson, C. Doland, F. A. Ponce, G. B. Anderson, and B. Wacker, *Mater. Res. Soc. Symp. Proc.* **118**, 49 (1988).
497. Y. S. Tsuo, R. Weil, S. Asher, A. Nelson, Y. Xu, and R. Tsu, in *Proceedings of the 19<sup>th</sup> IEEE Photovoltaic Specialists Conference, New Orleans, 1987* (IEEE, New York, N.Y., U.S.A., 1987), p. 705.
498. H. Shirai, D. Das, J. Hanna, and I. Shimizu, *Appl. Phys. Lett.* **59**, 1096 (1991).
499. A. Asano, T. Ichimura, and H. Sakai, *J. Appl. Phys.* **65**, 2439 (1989).
500. A. Asano, *Appl. Phys. Lett.* **56**, 533 (1990).
501. G. Parsons, *IEEE Electron Device Lett.* **13**, 80 (1992).
502. N. Layadi, P. Roca i Cabarrocas, B. Drévillon, and I. Solomon, *Phys. Rev. B* **52**, 5136 (1995).
503. P. Roca i Cabarrocas, N. Layadi, B. Drévillon, and I. Solomon, *J. Non-Cryst. Solids* **198-200**, 871 (1996).
504. P. Roca i Cabarrocas, S. Hamma, A. Hadjadi, J. Bertomeu, and J. Andreu, *Appl. Phys. Lett.* **69**, 529 (1996).
505. N. Shibata, K. Fukada, H. Ohtoshi, J. Hanna, S. Oda, and I. Shimizu, *Mater. Res. Soc. Symp. Proc.* **95**, 225 (1987).
506. Y. H. Yang, M. Katiyar, G. F. Feng, N. Maley, and J. R. Abelson, *Appl. Phys. Lett.* **65**, 1769 (1994).
507. S. Hamma and P. Roca i Cabarrocas, *J. Appl. Phys.* **81**, 7282 (1997).
508. P. Roca i Cabarrocas, R. Brenot, P. Bulkin, R. Vanderhaghen, B. Drévillon, and I. French, *J. Appl. Phys.* **86**, 7079 (1999).
509. K. Saito, M. Kondo, M. Fukawa, T. Nishimiya, A. Matsuda, W. Futako, and I. Shimizu, *Appl. Phys. Lett.* **71**, 3403 (1997).
510. C. Anandan, C. Mukherjee, T. Seth, P. N. Dixit, and R. Bhattacharyya, *Appl. Phys. Lett.* **66**, 85 (1995).

511. C. Mukherjee, C. Anandan, T. Seth, P. N. Dixit, and R. Bhattacharyya, *Appl. Phys. Lett.* **68**, 194 (1996).
512. A. C. W. Biebericher, J. Bezemer, W. F. van der Weg, and W. J. Goedheer, *Appl. Phys. Lett.* **76**, 2002 (2000).
513. H. Kirimura, H. Maeda, H. Murakami, T. Nakahigashi, S. Ohtani, T. Tabata, T. Hayashi, M. Kobayashi, Y. Mitsuda, N. Nakamura, H. Kuwahara, and A. Doi, *Jpn. J. Appl. Phys.* **33**, 4389 (1994).
514. S. Morrison, J. Xi, and A. Madan, *Mater. Res. Soc. Symp. Proc.* **507**, 559 (1998).
515. A. Madan, S. Morrison, and H. Kuwahara, *Sol. Energy Mater. Sol. Cells* **59**, 51 (1999).
516. G. Scarsbrook, I. P. Llewellyn, S. M. Ojha, and R. A. Heinecke, *Vacuum* **38**, 627 (1988).
517. L. J. Overzet and J. T. Verdeyen, *Appl. Phys. Lett.* **48**, 695 (1986).
518. J. P. Booth, G. Cunge, N. Sadeghi, and R. W. Boswell, *J. Appl. Phys.* **82**, 552 (1997).
519. A. C. W. Biebericher, J. Bezemer, W. F. van der Weg, and W. J. Goedheer, *Mater. Res. Soc. Proc. Symp.* **609**, A4.1.1 (2000).
520. A. C. W. Biebericher, J. Bezemer, W. F. van der Weg, and W. J. Goedheer, unpublished results.
521. H. Wiesmann, A. K. Ghosh, T. McMahon, and M. Strongin, *J. Appl. Phys.* **50**, 3752 (1979).
522. H. Matsumura, *Jpn. J. Appl. Phys.* **25**, L949 (1986).
523. H. Matsumura, *Appl. Phys. Lett.* **51**, 804 (1987).
524. H. Matsumura, *J. Appl. Phys.* **65**, 4396 (1989).
525. J. R. Doyle, R. Robertson, G. H. Lin, M. Z. He, and A. Gallagher, *J. Appl. Phys.* **64**, 3215 (1988).
526. F. Jansen, I. Chen, and M. A. Machonkin, *J. Appl. Phys.* **66**, 5749 (1989).
527. A. H. Mahan and M. Vaněček, in *Amorphous Silicon Materials and Solar Cells*, No. 234 in *AIP Conference Proceedings*, Solar Energy Research Institute, edited by B. Stafford (American Institute of Physics, New York, N.Y., U.S.A., 1991), p. 195.
528. E. C. Molenbroek, A. H. Mahan, E. Johnson, and A. C. Gallagher, *J. Appl. Phys.* **79**, 7278 (1996).
529. M. Heintze, R. Zedlitz, H. N. Wanka, and M. B. Schubert, *J. Appl. Phys.* **79**, 2699 (1996).
530. E. C. Molenbroek, A. H. Mahan, and A. C. Gallagher, *J. Appl. Phys.* **82**, 1909 (1997).
531. K. F. Feenstra, R. E. I. Schropp, and W. F. van der Weg, *J. Appl. Phys.* **85**, 6843 (1999).
532. P. Papadopoulos, A. Scholz, S. Bauer, B. Schröder, and H. Öchsner, *J. Non-Cryst. Solids* **164–166**, 87 (1993).
533. B. P. Nelson, E. Iwaniczko, R. E. I. Schropp, H. Mahan, E. C. Molenbroek, S. Salamon, and R. S. Crandall, in *Proceedings of the Twelfth E.C. Photovoltaic Solar Energy Conference, Amsterdam, the Netherlands, 1994*, edited by R. Hill, W. Palz, and P. Helm (H.S. Stephens & Associates, Bedford, U.K., 1994), p. 679.
534. A. H. Mahan, E. Iwaniczko, B. P. Nelson, R. C. Reedy Jr., R. S. Crandall, S. Guha, and J. Yang, in *Proceedings of the 25<sup>th</sup> IEEE Photovoltaic Specialists Conference, Washington DC, 1996* (IEEE, New York, NY, U.S.A., 1996), p. 1065.
535. R. E. I. Schropp, K. F. Feenstra, E. C. Molenbroek, H. Meiling, and J. K. Rath, *Philos. Mag. B* **76**, 309 (1997).
536. R. S. Crandall, A. H. Mahan, B. Nelson, M. Vaněček, and I. Balberg, in *AIP Conference Proceedings*, edited by R. Noufi (American Institute of Physics, Denver, 1992), Vol. 268, pp. 81–87.

537. Q. Wang, E. Iwaniczko, Y. Xu, W. Gao, B. P. Nelson, A. H. Mahan, R. S. Crandall, and H. M. Branz, *Mater. Res. Soc. Proc. Symp.* **609**, A4.3.1 (2000).
538. H. Meiling and R. E. I. Schropp, *Appl. Phys. Lett.* **70**, 2681 (1997).
539. V. Chu, J. Jarego, H. Silva, T. Silva, M. Reissner, P. Brogueira, and J. P. Conde, *Appl. Phys. Lett.* **70**, 2714 (1997).
540. H. Meiling, A. M. Brockhoff, J. K. Rath, and R. E. I. Schropp, *J. Non-Cryst. Solids* **227-230**, 1202 (1998).
541. J. Cifre, J. Bertomeu, J. Puigdollers, M. C. Polo, J. Andreu, and A. Lloret, *Appl. Phys. A.* **59**, 645 (1994).
542. A. R. Middya, S. Hazra, S. Ray, C. Longeaud, and J. P. Kleider, in *Proceedings of the Thirteenth European Photovoltaic Solar Energy Conference, Nice, France, 1995*, edited by W. Freiesleben, W. Palz, H. A. Ossenbrink, and P. Helm (H.S. Stephens & Associates, Bedford, U.K., 1996), p. 679.
543. P. Brogueira, J. P. Conde, S. Arekat, and V. Chu, *J. Appl. Phys.* **79**, 8748 (1996).
544. J. K. Rath, H. Meiling, and R. E. I. Schropp, *Sol. Energy Mater. Sol. Cells* **48**, 269 (1997).
545. H. N. Wanka, R. Zedlitz, M. Heintze, and M. B. Schubert, in *Proceedings of the Thirteenth European Photovoltaic Solar Energy Conference, Nice, France, 1995*, edited by W. Freiesleben, W. Palz, H. A. Ossenbrink, and P. Helm (H.S. Stephens & Associates, Bedford, U.K., 1996), p. 1753.
546. V. Brogueira, P. Chu, A. C. Ferro, and J. P. Conde, *J. Vac. Sci. Technol. A* **15**, 2968 (1997).
547. P. Alpuim, V. Chu, and J. P. Conde, *Mater. Res. Soc. Proc. Symp.* **609**, A22.6.1 (2000).
548. K. F. Feenstra, Ph.D. thesis, Universiteit Utrecht, Utrecht, the Netherlands, 1998.
549. C. Horbach, W. Beyer, and H. Wagner, *J. Non-Cryst. Solids* **137&138**, 661 (1991).
550. S. Bauer, B. Schröder, and H. Oechsner, *J. Non-Cryst. Solids* **227-230**, 34 (1998).
551. A. H. Mahan, Y. Chen, D. L. Williamson, and G. D. Mooney, *J. Non-Cryst. Solids* **137-138**, 65 (1991).
552. Y. Wu, J. T. Stephen, D. X. Han, J. M. Rutland, R. S. Crandall, and A. H. Mahan, *Phys. Rev. Lett.* **77**, 2049 (1996).
553. R. P. Muller, J. K. Holt, D. G. Goodwin, and W. A. Goddard, III, *Mater. Res. Soc. Proc. Symp.* **609**, A6.1.1 (2000).
554. J. K. Holt, M. Swiatek, D. G. Goodwin, and H. A. Atwater, *Mater. Res. Soc. Proc. Symp.* **609**, A6.2.1 (2000).
555. B. Schröder and S. Bauer, *J. Non-Cryst. Solids* **266-269**, 1115 (2000).
556. A. T. M. Wilbers, G. M. W. Kroesen, C. J. Timmermans, and D. C. Schram, *Meas. Sci. Technol.* **1**, 1326 (1990).
557. J. W. A. M. Gielen, P. R. M. Kleuskens, M. C. M. van de Sanden, L. J. van IJzendoorn, D. C. Schram, E. H. A. Dekempeneer, and J. Meneve, *J. Appl. Phys.* **80**, 5986 (1996).
558. J. J. Beulens, A. J. M. Buuron, and D. C. Schram, *Surf. Coat. Technol.* **47**, 401 (1991).
559. M. J. de Graaf, R. J. Severens, R. P. Dahiya, M. C. M. van de Sanden, and D. C. Schram, *Phys. Rev. E* **48**, 2098 (1993).
560. R. F. G. Meulenbroeks, M. F. M. Steenbakkens, Z. Qing, M. C. M. van de Sanden, and D. C. Schram, *Phys. Rev. E* **49**, 2272 (1994).
561. W. M. M. Kessels, A. H. M. Smets, B. A. Korevaar, G. J. Adriaenssens, M. C. M. van de Sanden, and D. C. Schram, *Mater. Res. Soc. Symp. Proc.* **557**, 25 (1999).
562. B. A. Korevaar, G. J. Adriaenssens, A. H. M. Smets, , W. M. M. Kessels, H.-Z. Song,

- M. C. M. van de Sanden, and D. C. Schram, *J. Non-Cryst. Solids* **266–269**, 380 (2000).
563. M. C. M. van de Sanden, R. J. Severens, W. M. M. Kessels, R. F. G. Meulenbroeks, and D. C. Schram, *J. Appl. Phys.* **84**, 2426 (1998).
564. M. C. M. van de Sanden, R. J. Severens, W. M. M. Kessels, R. F. G. Meulenbroeks, and D. C. Schram, *J. Appl. Phys.* **85**, 1243 (1999).
565. J. J. Beulens, M. J. de Graaf, and D. C. Schram, *Plasma Sources. Sci. Technol.* **2**, 180 (1993).
566. M. C. M. van de Sanden, J. M. de Regt, and D. C. Schram, *Plasma Sources. Sci. Technol.* **3**, 501 (1994).
567. W. M. M. Kessels, C. M. Leewis, M. C. M. van de Sanden, and D. C. Schram, *J. Appl. Phys.* **86**, 4029 (1999).
568. R. F. G. Meulenbroeks, R. A. H. Engeln, M. N. A. Beurskens, R. M. J. Paffen, M. C. M. van de Sanden, J. A. M. van der Mullen, and D. C. Schram, *Plasma Sources Sci. Technol.* **4**, 74 (1995).
569. W. M. M. Kessels, M. C. M. van de Sanden, and D. C. Schram, *J. Vac. Sci. Technol. A* **18**, 2153 (2000).
570. W. M. M. Kessels, M. C. M. van de Sanden, R. J. Severens, L. J. van Ijzendoorn, and D. C. Schram, *Mater. Res. Soc. Symp. Proc.* **507**, 529 (1998).
571. W. M. M. Kessels, Ph.D. thesis, Technische Universiteit Eindhoven, Eindhoven, the Netherlands, 2000.
572. W. M. M. Kessels, R. J. Severens, A. H. M. Smets, B. A. Korevaar, G. J. Adriaenssens, D. C. Schram, and M. C. M. van de Sanden, *J. Appl. Phys.* **89**, 2404 (2001).
573. W. M. M. Kessels, J. P. M. Hoefnagels, M. G. H. Boogaarts, D. C. Schram, and M. C. M. van de Sanden, *J. Appl. Phys.* **89**, 2065 (2001).
574. H. S. Nalwa, Ed., *Handbook of Advanced Electronic and Photonic Materials and Devices* (Academic Press, Boston, MA, U.S.A., 2000).
575. M. A. Green, *Solar Cells, Operating Principles, Technology and System Applications* (University of New South Wales, Kensington, NSW, Australia, 1992).
576. M. Hack and M. Shur, *J. Appl. Phys.* **58**, 997 (1985).
577. A. Catalano, in *Amorphous and Microcrystalline Semiconductor Devices – Optoelectronic devices*, edited by J. Kanicki (Artech House, Norwood, MA, U.S.A., 1991), Chap. 2, p. 9.
578. A. Polman, W. G. J. H. M. van Sark, W. C. Sinke, and F. W. Saris, *Sol. Cells* **17**, 241 (1986).
579. H. Hartnagel, A. Dawar, A. Jain, and C. Jagadish, *Semiconducting Transparent Thin Films* (Institute of Physics Publishing, Bristol, U.K., 1995).
580. E. Yablonovitch and G. D. Cody, *IEEE Trans. Elec. Dev.* **ED-29**, 300 (1982).
581. H. W. Deckman, C. R. Wronski, H. Witzke, and E. Yablonovitch, *Appl. Phys. Lett.* **42**, 968 (1983).
582. J. Daey Ouwens, R. E. I. Schropp, J. Wallinga, W. F. van der Weg, M. Ritala, M. Leskelä, and Hyvärinen, in *Proceedings of the Twelfth E.C. Photovoltaic Solar Energy Conference, Amsterdam, the Netherlands, 1994*, edited by R. Hill, W. Palz, and P. Helm (H.S. Stephens & Associates, Bedford, U.K., 1994), p. 1296.
583. M. Kubon, E. Böhmer, F. Siebke, B. Rech, C. Beneking, and H. Wagner, *Sol. Energy Mater. Sol. Cells* **41/42**, 485 (1996).
584. Y. Tawada, H. Okamoto, and Y. Hamakawa, *Appl. Phys. Lett.* **39**, 237 (1981).
585. K. Miyachi, N. Ishiguro, T. Miyashita, N. Yanagawa, H. Tanaka, M. Koyama, Y. Ashida,

- and N. Fukuda, in *Proceedings of the 11th European Photovoltaic Solar Energy Conference, Montreux, Switzerland*, edited by L. Guimarães, W. Palz, C. de Reyff, H. Kiess, and P. Helm (Harwood, Chur, Switzerland, 1993), p. 88.
586. R. R. Arya, A. Catalano, and R. S. Oswald, *Appl. Phys. Lett.* **49**, 1089 (1986).
587. C. Beneking, B. Rech, T. Eickhoff, Y. G. Michael, N. Schultz, and H. Wagner, in *Proceedings of the Twelfth E.C. Photovoltaic Solar Energy Conference, Amsterdam, the Netherlands, 1994*, edited by R. Hill, W. Palz, and P. Helm (H.S. Stephens & Associates, Bedford, U.K., 1994), p. 683.
588. G. E. N. Landweer, B. S. Girwar, C. H. M. van der Werf, J. W. Metselaar, and R. E. I. Schropp, in *Proceedings of the Twelfth E.C. Photovoltaic Solar Energy Conference, Amsterdam, the Netherlands, 1994*, edited by R. Hill, W. Palz, and P. Helm (H.S. Stephens & Associates, Bedford, U.K., 1994), p. 1300.
589. A. Banerjee and S. Guha, *J. Appl. Phys.* **69**, 1030 (1991).
590. Y. Kuwano, M. Ohnishi, H. Nishiwaki, S. Tsuda, T. Fukatsu, K. Enomoto, Y. Nakashima, and H. Tarui, in *Proceedings of the 16<sup>th</sup> IEEE Photovoltaic Specialists Conference, San Diego, 1982* (IEEE, New York, N.Y., U.S.A., 1982), p. 1338.
591. S. Guha, J. Yang, A. Pawlikiewicz, T. Glatfelter, R. Ross, and S. R. Ovshinsky, *Appl. Phys. Lett.* **54**, 2330 (1989).
592. J. Zimmer, H. Stiebig, and H. Wagner, *J. Appl. Phys.* **84**, 611 (1998).
593. R. E. Rochelau, M. Tun, and S. S. Hegedus, in *Proceedings of the 26<sup>th</sup> IEEE Photovoltaic Specialists Conference, Anaheim, 1996* (IEEE, New York, NY, U.S.A., 1997), p. 703.
594. J. K. Rath, R. E. I. Schropp, and W. Beyer, *J. Non-Cryst. Solids* **227–230**, 1282 (1998).
595. J. Bruns, M. Choudhury, and H. G. Wagemann, in *Proceedings of the Thirteenth European Photovoltaic Solar Energy Conference, Nice, France, 1995*, edited by W. Freiesleben, W. Palz, H. A. Ossenbrink, and P. Helm (H.S. Stephens & Associates, Bedford, U.K., 1996), pp. 230–233.
596. M. Zeman, J. A. Willemen, S. Solntsev, and G. W. Metselaar, *Sol. Energy Mater. Sol. Cells* **34**, 557 (1994).
597. M. Zeman, R. A. C. M. M. van Swaaij, E. Schroten, L. L. A. Vosteen, and J. W. Metselaar, *Mater. Res. Soc. Symp. Proc.* **507**, 409 (1998).
598. M. Zeman, J. A. Willemen, L. L. A. Vosteen, G. Tao, and G. W. Metselaar, *Sol. Energy Mater. Sol. Cells* **46**, 81 (1997).
599. X. Xu, J. Yang, and S. Guha, *Appl. Phys. Lett.* **62**, 1399 (1993).
600. Y. Lee, L. Jiao, Z. Lu, R. W. Collins, and C. R. Wronski, *Sol. Energy Mater. Sol. Cells* **49**, 149 (1997).
601. B. W. Faughnan and R. Crandall, *Appl. Phys. Lett.* **44**, 537 (1984).
602. Z. E. Smith, S. Wagner, and B. W. Faughnan, *Appl. Phys. Lett.* **46**, 1078 (1985).
603. A. Catalano, R. Bennett, R. Arya, K. Rajan, and J. Newton, in *Proceedings of the 18<sup>th</sup> IEEE Photovoltaic Specialists Conference, Las Vegas, 1985* (IEEE, New York, N.Y., U.S.A., 1985), p. 1378.
604. M. Stutzmann, W. B. Jackson, and C. C. Tsai, *Phys. Rev. B* **34**, 63 (1986).
605. R. E. I. Schropp, A. Sluiter, M. B. von der Linden, and J. Daey Ouwens, *J. Non-Cryst. Solids* **164–166**, 709 (1993).
606. D. E. Carlson, K. Rajan, R. R. Arya, F. Willing, and L. Yang, *J. Mater. Res.* **13**, 2754 (1998).
607. Y. Hamakawa, W. Ma, and H. Okamoto, in *Plasma Deposition of Amorphous Silicon-Based Materials*, edited by G. Bruno, P. Capezzuto, and A. Madan (Academic Press,

- Boston, MA, U.S.A., 1995), Chap. 6, p. 283.
608. L. L. Kazmerski, *Ren. Sust. Energy Rev.* **1**, 71 (1997).
609. A. V. Shah, R. Platz, and H. Keppner, *Sol. Energy Mater. Sol. Cells* **38**, 501 (1995).
610. T. Strand, L. Mrig, R. Hansen, and K. Emery, *Sol. Energy Mater. Sol. Cells* **41/42**, 617 (1996).
611. R. Ruther and J. Livingstone, *Sol. Energy Mater. Sol. Cells* **36**, 29 (1996).
612. J. Merten and J. Andreu, *Sol. Energy Mater. Sol. Cells* **52**, 11 (1998).
613. J. Yang, A. Banerjee, T. Glatfelter, K. Hoffman, X. Xu, and S. Guha, in *Proceedings of the 24<sup>th</sup> IEEE Photovoltaic Specialists Conference, Waikoloa HI, 1994* (IEEE, New York, NY, U.S.A., 1995), p. 380.
614. S. Guha, J. Yang, A. Banerjee, T. Glatfelter, and S. Sugiyama, *Sol. Energy Mater. Sol. Cells* **48**, 365 (1997).
615. S. Kiyama, S. Nakano, Y. Domoto, H. Hirano, H. Tarui, K. Wakisaka, M. Tanaka, S. Tsuda, and S. Nakano, *Sol. Energy Mater. Sol. Cells* **48**, 373 (1997).
616. Y. Hamakawa, *Appl. Surf. Sci.* **142**, 215 (1999).
617. S. M. Sze, *Semiconductor Devices, Physics and Technology* (Wiley, New York, 1985).
618. M. J. Powell, *IEEE Trans. Elec. Dev.* **ED-36**, 2753 (1989).
619. K. Suzuki, in *Amorphous and Microcrystalline Semiconductor Devices – Optoelectronic devices*, edited by J. Kanicki (Artech House, Norwood, MA, U.S.A., 1991), Chap. 3, p. 77.
620. A. Iqbal, W. B. Jackson, C. C. Tsai, J. W. Allen, and C. W. Bates, *J. Appl. Phys.* **61**, 2947 (1987).
621. C. van Berkel, in *Amorphous and Microcrystalline Semiconductor Devices – Materials and Device Physics*, edited by J. Kanicki (Artech House, Norwood, MA, U.S.A., 1992), Chap. 8, p. 397.
622. W. B. Jackson and M. Moyer, *Mater. Res. Soc. Symp. Proc.* **118**, 231 (1988).
623. J. Kakalios, R. A. Street, and W. B. Jackson, *Phys. Rev. B* **59**, 1037 (1987).
624. W. B. Jackson, J. M. Marshall, and M. D. Moyer, *Phys. Rev. B* **39**, 1164 (1989).
625. A. M. Brockhoff, E. H. C. Ullersma, H. Meiling, F. H. P. M. Habraken, and W. F. van der Weg, *Appl. Phys. Lett.* **73**, 3244 (1998).
626. C. van Berkel and M. J. Powell, *J. Appl. Phys* **60**, 1521 (1986).
627. M. Katayama, *Thin Solid Films* **341**, 140 (1999).
628. H. C. Tuan, *J. Non-Cryst. Solids* **115**, 132 (1989).
629. H. Matsumura, H. Hayama, Y. Nara, and K. Ishibashi, *Jpn. J. Appl. Phys.* **20**, 311 (1980).
630. R. L. Weisfield, *IEEE Trans. Elec. Dev.* **ED-36**, 2935 (1989).
631. I. Shimizu, S. Shirai, and E. Inoue, *J. Appl. Phys.* **52**, 2776 (1981).
632. I. Shimizu, in *Semiconductors and Semimetals*, edited by J. I. Pankove (Academic, Orlando, Fla., U.S.A., 1984), Vol. 21D, p. 55.
633. R. Schaffert, *Electrophotography* (Focal, London, U.K., 1975).
634. J. Kodama, S. Araki, M. Kimura, and T. Inagaki, *Jpn. J. Appl. Phys.* **29**, L867 (1990).
635. R. A. C. M. M. van Swaaij, W. P. M. Willems, J. Bezemer, H. J. P. Lokker, and W. F. van der Weg, *Mater. Res. Soc. Symp. Proc.* **297**, 559 (1993).
636. R. A. C. M. M. van Swaaij, S. J. Elmer, W. P. M. Willems, J. Bezemer, J. M. Marshall, and A. R. Hepburn, *J. Non-Cryst. Solids* **164–166**, 533 (1993).
637. T. Takeda and S. Sano, *Mater. Res. Soc. Symp. Proc.* **118**, 399 (1988).
638. E. Fortunato, M. Vieira, I. Ferreira, C. N. Carvalho, G. Lavareda, and R. Martins,

- Mater. Res. Soc. Symp. Proc. **297**, 981 (1993).
639. E. M. C. Fortunato, D. Brida, I. M. M. Ferreira, H. M. B. Águas, P. Nunes, A. Cabrita, F. Guilianì, Y. Nunes, M. J. P. Maneira, and R. Martins, Mater. Res. Soc. Proc. Symp. **609**, A12.7.1 (2000).
640. H. Stiebig and N. Bohm, J. Non-Cryst. Solids **164–166**, 789 (1993).
641. R. Brüggemann, T. Neidlinger, and M. B. Schubert, J. Appl. Phys. **81**, 7666 (1997).
642. G. de Cesare, F. Irrera, F. Lemmi, and F. Palma, Appl. Phys. Lett. **66**, 1178 (1995).
643. M. Topič, F. Smole, J. Furlan, and W. Kusian, J. Non-Cryst. Solids **227–230**, 1326 (1998).
644. J. Krč, M. Topič, F. Smole, D. Knipp, and H. Stiebig, Mater. Res. Soc. Proc. Symp. **609**, A12.5.1 (2000).
645. D. Caputo and G. de Cesare, Sensors and Actuators A **78**, 108 (1999).
646. D. Caputo, G. de Cesare, A. Nascetti, F. Palma, and M. Petri, Appl. Phys. Lett. **72**, 1229 (1998).
647. R. A. Street, X. D. Wu, R. Weisfield, S. Ready, R. Apte, M. Nguyen, and P. Nylen, Nucl. Instrum. Methods Phys. Res. A **380**, 450 (1996).
648. R. A. Street, R. B. Apte, T. Granberg, M. P., S. E. Ready, K. S. Shah, and R. L. Weisfield, J. Non-Cryst. Solids **227–230**, 1306 (1998).
649. J.-P. Moy, Nucl. Instrum. Methods Phys. Res. A **442**, 26 (2000).
650. J. C. Chou, Y.-F. Wang, and J. S. Lin, Sensors and Actuators B **62**, 92 (2000).
651. J. C. Chou, H. M. Tsai, C. N. Shiao, and J. S. Lin, Sensors and Actuators B **62**, 97 (2000).
652. S. Jamash, S. D. Collins, and R. L. Smith, IEEE Trans. Elec. Dev. **ED-45**, 1239 (1998).
653. P. F. Ruths, S. Askok, S. J. Fonash, and J. M. Ruths, IEEE Trans. Elec. Dev. **ED-28**, 1003 (1981).
654. E. Fortunato, A. Malik, A. Sêco, I. Ferreira, and R. Martins, J. Non-Cryst. Solids **227–230**, 1349 (1998).
655. F. V. Hunt, *Electroacoustics, the Analysis of Transduction and Its Historical Background* (American Institute of Physics, New York, NY, U.S.A., 1982).
656. M. Smits, private communication.
657. R. E. I. Schropp, M. Smits, H. Meiling, W. G. J. H. M. van Sark, M. M. Boone, and W. F. van der Weg, Mater. Res. Soc. Symp. Proc. **219**, 519 (1991).
658. P. J. Mears, L. Reekie, I. M. Jauncey, and D. N. Payne, Electron. Lett. **23**, 1026 (1987).
659. E. Desurvire, R. J. Simpson, and P. C. Becker, Opt. Lett. **12**, 888 (1987).
660. G. N. van den Hoven, A. Polman, C. van Dam, J. W. M. van Uffelen, and M. K. Smit, Appl. Phys. Lett. **68**, 1886 (1996).
661. A. Polman, J. Appl. Phys. **82**, 1 (1997).
662. F. Priolo, G. Franzò, S. Coffa, and A. Carnera, Mat. Chem. Phys. **54**, 273 (1998).
663. F. Priolo, G. Franzò, S. Coffa, and A. Carnera, Phys. Rev. B **57**, 4443 (1998).
664. G. N. van den Hoven, J. H. Shin, A. Polman, S. Lombardo, and S. U. Campisano, J. Appl. Phys. **78**, 2642 (1995).
665. J. H. Shin, R. Serna, G. N. van den Hoven, A. Polman, W. G. J. H. M. van Sark, and A. M. Vredenberg, Appl. Phys. Lett. **68**, 697 (1996).
666. A. Polman, J. H. Shin, R. Serna, G. N. van den Hoven, W. G. J. H. M. van Sark, A. M. Vredenberg, S. Lombardo, and S. U. Campisano, Mater. Res. Soc. Symp. Proc. **422**, 239 (1996).
667. S. Coffa, G. Franzò, A. Polman, and R. Serna, Phys. Rev. B **49**, 16313 (1994).

668. P. G. Kik, M. J. A. de Dood, K. Kikoin, and A. Polman, *Appl. Phys. Lett.* **70**, 1721 (1997).
669. F. Priolo, G. Franzò, S. Coffa, A. Polman, S. Libertino, R. Barklie, and D. Carey, *J. Appl. Phys.* **78**, 3874 (1995).
670. G. Franzò, F. Priolo, S. Coffa, A. Polman, and A. Carnera, *Appl. Phys. Lett.* **64**, 2235 (1994).
671. B. Zheng, J. Michel, F. Y. G. Ren, L. C. Kimerling, D. C. Jacobson, and J. M. Poate, *Appl. Phys. Lett.* **64**, 2842 (1994).
672. J. Palm, F. Gan, B. Zheng, J. Michel, and L. C. Kimerling, *Phys. Rev. B* **54**, 17603 (1996).
673. S. Lombardo, S. U. Campisano, G. N. van den Hoven, and A. Polman, *J. Appl. Phys.* **77**, 6504 (1995).
674. L. H. Slooff, Master's thesis, FOM-Institute for Atomic and Molecular Physics, Amsterdam, the Netherlands, 1996.
675. J. Stimmer, A. Reittinger, J. F. Nützel, G. Abstreiter, H. Holzbrecher, and C. Buchal, *Appl. Phys. Lett.* **68**, 3290 (1996).
676. W. Fuhs, I. Ulber, G. Weiser, M. S. Bresler, O. B. Gusev, A. N. Kuznetsov, V. K. Kudoyarova, E. I. Terukov, and I. N. Yassievich, *Phys. Rev. B* **56**, 9545 (1997).
677. E. I. Terukov, O. I. Kon'kov, V. K. Kudoyarova, O. B. Gusev, and G. Weiser, *Semiconductors* **24**, 884 (1998).
678. E. I. Terukov, M. M. Kazanin, O. I. Kon'kov, V. K. Kudoyarova, K. V. Kougiya, Y. A. Nikulin, and Kazanskiĭ, *Semiconductors* **34**, 829 (2000).
679. V. B. Voronkov, V. G. Golubev, A. V. Medvedev, A. B. Pevtsov, N. A. Feoktistov, N. I. Gorshkov, and D. N. Suglobov, *Phys. Solid State* **40**, 1301 (1998).
680. J. H. Shin and M. Kim, *J. Vac. Sci. Technol. A* **17**, 3230 (1999).
681. A. Masuda, J. Sakai, H. Akiyama, O. Eryu, K. Nakashima, and H. Matsumura, *J. Non-Cryst. Solids* **226-229**, 136 (2000).
682. T. Förster, *Discuss. Faraday Soc.* **27**, 7 (1959).
683. H. Kühne, G. Weiser, E. I. Terukov, A. N. Kuznetsov, and V. K. Kudoyarova, *J. Appl. Phys.* **86**, 1896 (1999).
684. M. B. Schubert, A. Hierzenberger, H. J. Lehner, and J. H. Werner, *Sensors and Actuators A* **74**, 193 (1999).
685. E. Betzig and J. K. Trautmann, *Science* **257**, 189 (1992).
686. E. Betzig, P. L. Finn, and J. S. Weiner, *Appl. Phys. Lett.* **60**, 2484 (1992).
687. S. Madsen, M. Mullenborn, K. Birkelund, and F. Grey, *Appl. Phys. Lett.* **69**, 544 (1996).
688. S. Madsen, S. I. Bozhevolnyi, K. Birkelund, M. Mullenborn, J. M. Hvam, and F. Grey, *J. Appl. Phys.* **82**, 49 (1997).
689. M. K. Herndon, R. T. Collins, R. E. Hollingsworth, P. R. Larson, and M. B. Johnson, *Appl. Phys. Lett.* **74**, 141 (1999).
690. W. Gao, S. H. Lee, J. Bullock, Y. Xu, D. K. Benson, S. Morrison, and H. M. Branz, *Sol. Energy Mater. Sol. Cells* **59**, 243 (1999).