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# Refractive Index Variation of Magnetron-Sputtered a- $Si_{1-x}Ge_x$ by "One-Sample Concept" Combinatory

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**Abstract:** Gradient a-Si<sub>1-x</sub>Ge $_x$  layers have been deposited by "one-sample concept" combinatorial direct current (DC) magnetron sputtering onto one-inch-long Si slabs. Characterizations by electron microscopy, ion beam analysis and ellipsometry show that the layers are amorphous with a uniform thickness, small roughness and compositions from x=0 to x=1 changing linearly with the lateral position. By focused-beam mapping ellipsometry, we show that the optical constants also vary linearly with the lateral position, implying that the optical constants are linear functions of the composition. Both the refractive index and the extinction coefficient can be varied in a broad range for a large spectral region. The precise control and the knowledge of layer properties as a function of composition is of primary importance in many applications from solar cells to sensors.

Keywords: SiGe; spectroscopic ellipsometry; optical properties

#### 1. Introduction

SiGe thin film research is largely motivated by diverse applications in various semiconductor devices [1,2], such as solar cells, thin film transistors [3], Schottky diodes [4], temperature sensors [5], detectors and bolometers [6,7]. However, the effect of the full-scale composition on the properties of thin SiGe films has not been investigated thoroughly; in most of the reports, high film deposition rates have been chosen as a rule instead. The majority of these publications deal with a relatively low Ge concentration [8] and only these films have provided opto-electronic properties suitable for application in device structures. Beyond the special applications, the structure and electrical properties of SiGe alloys have been investigated [9], also for a-Si<sub>1-x</sub>Ge<sub>x</sub> films over the entire range of  $0 \le x \le 1$  [10].

The actuality of our work is given by the fact that mid-infrared (IR) photonic circuits (mid-IR PCs) have recently gained significance due to the huge number of applications projected over the 2–20  $\mu$ m wavelength range [6,11]. The Si-based technologies have shown a great potential for the implementation of new mid-IR PCs, following a real trend motivated by the large supply of devices already demonstrated at telecom wavelengths [12]. The implementations of SiGe-on-Si is particularly interesting since it allows the fine control of material properties such as the bandgap or the refractive index by balancing the Ge concentration in the alloy, while extending the operation range up to at least 8  $\mu$ m (wavelength of Si absorption onset)—with the possibility of further extending it up to 14  $\mu$ m (Ge transparency cutoff wavelength) for devices with low Si concentration [12,13]. Moreover, Ge presents higher nonlinear properties than Si, hence, preparing Ge-rich structures is a convenient choice to develop nonlinear optical devices in the mid-IR spectral range [14].

Besides chemical vapour deposition (CVD), numerous preparation methods have been used to elaborate this alloy to make a competitive material, however, physical vapor deposition including direct

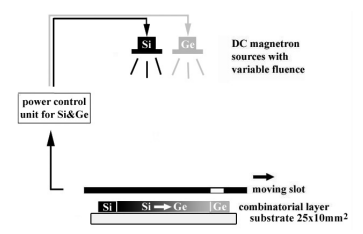
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current (DC) magnetron sputtering method is not extensively used. Two parameters are of primary importance: the low temperature and the high rate of deposition. This technique has the advantages of simplicity, possibility of separating material components, and of relatively high deposition rates. Materials libraries are deposited mostly by space resolved combinatory using DC magnetron sputtering from two to four targets arranged in a static set-up. The targets are arranged at certain distances from each other above a large substrate or a number of distinct substrates covering the exposed area. As a result of the static arrangements, composition spread alloy films are obtained in which the gradient is not constant and the concentration range is limited; usually, does not cover 0–100% of the components [15–17]. In this study, we present the preparation of non-hydrogenated gradient SiGe films by DC magnetron sputtering over a surface length of 2 cm. This is a second stage in the development of a combinatorial method [18,19] utilizing the "one sample concept" demonstrated for transmission electron microscopy (TEM) characterizations in smaller samples like TEM grids [20]. The thin film samples have been deposited with different, gradually varying compositions in the whole range and at preparation conditions identical for Si and Ge.

Silicon-based SiGe thin films have been demonstrated for microelectronics, the electronic properties of which are well known. The optical properties of crystalline alloys have also been determined for certain compositions [21,22]. Our objective is to establish the relation between the refractive index and the composition of the layers deposited by the DC magnetron technique. In this article, we show that not only the strictly controlled preparation of a-Si<sub>1-x</sub>Ge<sub>x</sub> films is possible over the entire range of  $0 \le x \le 1$  using magnetron sputtering over a length of 2 cm, but the composition, and even more importantly the refractive index (n) and extinction coefficient (k) all show a linear dependence on the position in most of the wavelength range. Using optical measurements by ellipsometry, we demonstrate that our combinatorial setup is suitable for a comprehensive optical characterization of a-Si<sub>1-x</sub>Ge<sub>x</sub> films with accurate spatial and compositional control at a high resolution.

# 2. 'One-Sample' Combinatory

The combinatorial a-Si<sub>1-x</sub>Ge<sub>x</sub> samples with gradient composition ranging in  $0 \le x \le 1$  were deposited onto  $25 \times 10$  mm<sup>2</sup> size Si slabs by DC magnetron sputtering. We followed the "one sample concept" combinatory in order to cover the entire composition range of a binary film within a single specimen. This is implemented by means of a scaled-up device originally constructed for the preparation of 3-mm diameter micro-combinatorial TEM samples [20]. The present arrangement (Figure 1) moves a shutter with a  $2 \times 10$  mm<sup>2</sup> slot in fine steps above the substrate while the power of the two magnetron sources is regulated in sync with the position of the slot.



**Figure 1.** Set-up used for the combinatorial deposition of the a-Si<sub>1-x</sub>Ge<sub>x</sub> layer.

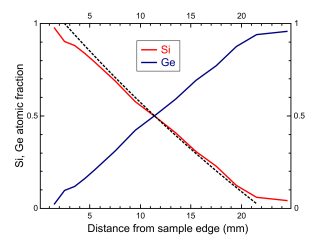
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Using the above method, a combinatorial sample is deposited in form of a 10 mm wide and 25 mm long layer track including a gradient section of 20 mm in length. The gradient part is enclosed in between 2.5 mm long sections of exclusively Si and Ge. The scheme in Figure 1 depicts the experimental arrangement and the construction of the combinatorial specimen.

A Si-Ge combinatorial sample was deposited as follows: the vacuum system was evacuated to a base pressure of  $5 \times 10^{-6}$  Pa. The substrate was mounted in the combinatorial device and it was load-locked into the chamber. Ar sputtering gas of partial pressure of  $2.5 \times 10^{-1}$  Pa was introduced. The Si and Ge targets were cleaned, behind closed shutters, for five minutes applying 340 W and 100 W magnetron power, respectively. These values were selected by preliminary measurements as maximum (100%) of the regulated power of the individual sources, that provided, equally, 0.45 nm/s deposition rates. Subsequently, the slot's movement of the combinatorial device was started, the shutter of the Si source was opened and Si was deposited at 100% power through the moving slot onto the substrate. This provided the Si section of the combinatorial track. In due time, the power of the Ge source was set to zero and its shutter was opened. It was followed by the gradual increase of the power of the Ge source, simultaneously, with the decrease of the power of the Si source, so the binary, gradient section of the combinatorial sample was deposited. As the power of the Si source arrived to zero and, simultaneously, that of the Ge arrived to its maximum, the Si source was switched off and, finally, the Ge section of the sample was deposited at 100% power of the Ge source.

#### 3. Distribution of Si and Ge Atomic Fractions

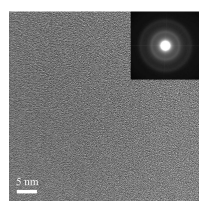
The atomic fractions of Si and Ge were measured by Rutherford backscattering spectrometry (RBS), as shown in Figure 2. The RBS analysis was performed in a scattering chamber with a two-axis goniometer connected to the custom-made 5-MV EG-2R Van de Graaff accelerator of the Wigner RMI Institute of the Hungarian Academy of Sciences. The 2-MeV  $^4$ He $^+$  analyzing ion beam was collimated with two sets of four-sector slits to the spot size of 0.5 mm  $\times$  0.5 mm, while the beam divergence was kept below  $0.06^{\circ}$ . The beam current was measured by a transmission Faraday cup [23]. Backscattered He $^+$  ions were detected using an ORTEC surface barrier detector (ORTEC, Illinois, USA). The energy resolution of the detection system was 20 keV. Spectra were recorded in Cornell geometry at scattering angle of  $\Theta = 165^{\circ}$  for two different sample tilt angles of  $7^{\circ}$  and  $60^{\circ}$ . For quantitative compositional analysis, both axial and planar channeling of the He $^+$  projectiles in the single-crystalline Si substrate were avoided. The measured data were evaluated with the spectrum simulation code named RBX [24].



**Figure 2.** Atomic fraction as a function of position along the center line of the combinatorial sample measured by Rutherford backscattering spectrometry. The black dotted line shows the distribution of the atomic fraction based on the calibration using energy dispersive spectrometry (EDS) in a scanning electron microscope.

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The crystalline quality of single crystalline samples can be qualified by RBS/channeling experiments if the sample is oriented along a major crystalline axis and the yield of the oriented spectrum is compared to the yield of the random, i.e., not oriented spectrum. In our case, oriented (channeling) RBS spectra were not measured, only random spectra were recorded. In the evaluation, we supposed random orientation of the SiGe layer. In this case, a quantitative analysis for the Si/Ge atomic fraction can be performed. On the other hand, selected area electron diffraction pattern represents an amorphous structure (see Figure 3).



**Figure 3.** High resolution TEM image of a 10 nm thick self supporting combinatorial SiGe sample taken for the composition of x = 0.5.

The RBS spectra have been recorded from edge to edge in the center line of the sample. Figure 2 reveals that the atomic fraction shows an exactly linear dependence on the position. The distribution is in a good agreement with the calibration based on electron microscopy measurement on a test sample as shown by the black dotted line in the figure.

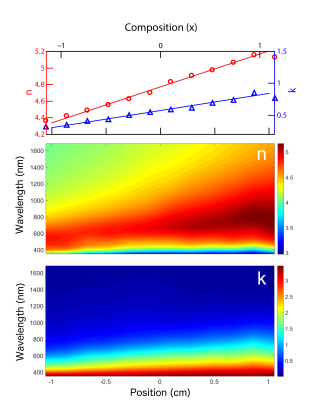
### 4. Distribution of the Optical Properties

The 25 mm  $\times$  10 mm combinatorial sample was scanned by a M-2000DI rotating compensator spectroscopic ellipsometer (Woollam Co. Inc., Lincoln, NE, USA). A focused spot has been moved parallel to the long edge along the center line of the wafer. The plane of incidence was parallel to the short edge. The angle of incidence was varied between 60° and 70°, and the corresponding size of the focused spot was 0.3 mm wide and 0.6–0.9 mm long. The measurement time was a few seconds for one spot and one angle of incidence. That provided high-resolution and high-accuracy of the data library with a reasonable acquisition time of the entire map (Figure 4).

In order to determine the parameters of the deposited a-Si<sub>1-x</sub>Ge<sub>x</sub> films, an optical model was constructed to calculate the ellipsometric angles from the model, and to fit the parameters of the model to obtain a good match between the measured and fitted ellipsometry spectra. Since the a-Si<sub>1-x</sub>Ge<sub>x</sub> film was deposited on a silicon wafer, the model uses a Si substrate assuming a vertically uniform structure, in which only a surface nanoroughness was taken into account by the usual model of mixing 50% layer material with 50% void. The dispersion of the layer was modeled using the Tauc–Lorentz (TL) approach [25]. In this parameterization, the dielectric function is described as a combination of a Lorentz oscillator with the three usual parameters (the broadening, the amplitude and the peak energy position) and an additional Tauc gap parameter. This approach allows that the total number of fitted parameters is kept at a reasonable value (thicknesses of the roughness layer and the film, and the TL parameters. The spectra measured at each position (each composition) have been fitted independently, by changing the above parameters to find the best match between the measured and calculated ellipsometry spectra. The spectra have been recorded at three angles of incidence providing six measured values at each wavelength, and a total of about 2000 measured values for each scanned position. Consequently, the fit was robust, avoiding local minima in the process, resulting in a smooth

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map of the optical spectra, as shown by Figure 4. Note the smooth changes of the spectra in Figure 4 in spite of the independent fit procedures at each position, which show the reliability of the fit procedure.



**Figure 4.** Spectra of the refractive index (n, color map on the middle graph) and the extinction coefficient (k, color map of the bottom graph) as a function of position along the center line of the wafer, calculated using the Tauc–Lorentz (TL) parameterization, as described in the text. The n and k spectra fitted by the TL model individually can be identified along the vertical axis of the bottom and middle maps, the values of which are color coded at the right hand side of the graphs. The composition of a-Si<sub>1-x</sub>Ge $_x$  changes from x = 0 to x = 1 over the lateral positions from -1 to 1, as shown in the horizontal axis at the top. The zero position corresponds to the composition of x = 0.5. The top graph shows n and k at the wavelength of 849.7 nm.

The measured  $\Psi$  and  $\Delta$  spectra fitted by the TL parameterization provide us not only with the thickness of the layer ( $\approx$ 100 nm) and the surface roughness (less than 3 nm) but also with n and k spectra at each spatial position of the sample. The maps in Figure 4 show those spectra as a function of the position along the center line of the wafer, with the zero position corresponding to the composition of x=0.5. To check the crystallinity of the layer, High Resolution Transmission Electron Microscope (HRTEM) and selected area electron diffraction (SAED) investigations were carried out by a 300-kV JEOL 3010 HRTEM (Japan Electron Optics Laboratory, Tokyo, Japan) with 0.17 nm point resolution. Both the granular image contrast and the diffuse rings of the inserted selected area electron diffraction pattern represent an amorphous structure (Figure 3).

The most important feature of the optical properties is that both n and k show a linear dependence on the position in the whole measured photon energy range, as shown by the color stripes with straight edges corresponding to large changes of the optical properties as a function of the photon energy. This is remarkable, because together with the RBS results of Figure 2 that indicates a linear dependence of the composition on the position (also shown in Ref. [20]), the linear dependence of n and k on the position proves that they also linearly depend on the composition ( $n_{Si_{1-x}Ge_x}(x) = (1-x) \times n_{Si} + x \times n_{Ge}$  and  $k_{Si_{1-x}Ge_x}(x) = (1-x) \times k_{Si} + x \times k_{Ge}$ , where  $n_{Si} = 4.33$ ,  $n_{Ge} = 5.33$ ,  $k_{Si} = 0.46$ ,  $k_{Ge} = 1.10$  for the wavelength of 849.7 nm). Both n and k span a wide range depending on the composition, indicating

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that these properties can well be controlled by properly adjusting x in the a-Si<sub>1-x</sub>Ge $_x$  composition. It is also important to emphasize that this behavior holds for a broad range of wavelengths. Note that the refractive index can also be modified by hydrogenation, which will be the target of our next study. In this work, we deal with non-hydrogenated amorphous Si and Ge (see also Ref. [26]).

## 5. Conclusions

It was shown that the "one-sample concept" combinatorial DC magnetron sputtering developed for transmission electron microscopy characterizations [20] can be up-scaled to a size of one inch, suitable for ellipsometry, with a linear spatial variation of the Si-Ge composition. The real and imaginary parts of the refractive index also show a linear dependence on both the lateral position and the composition in the investigated wavelength range of 350–1650 nm. Using the proposed concept of a:Si<sub>1-x</sub>Ge<sub>x</sub> DC magnetron combinatory, the refractive index and the extinction coefficient can be varied, on purpose, in a broad range at each wavelength.

**Author Contributions:** G.S. and M.S. prepared the samples. T.L., B.K. and Z.Z. made the measurements and the evaluations. P.P. and M.S. wrote the manuscript.

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