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Si-nanocrystal-based LEDs fabricated by ion implantation and plasma-enhanced chemical vapour deposition

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Abstract

An in-depth study of the physical and electrical properties of Si-nanocrystal-based MOSLEDs is presented. The active layers were fabricated with different concentrations of Si by both ion implantation and plasma-enhanced chemical vapour deposition.

Devices fabricated by ion implantation exhibit a combination of direct current and field-effect luminescence under a bipolar pulsed excitation. The onset of the emission decreases with the Si excess from 6 to 3 V. The direct current emission is attributed to impact ionization and is associated with the reasonably high current levels observed in current–voltage measurements. This behaviour is in good agreement with transmission electron microscopy images that revealed a continuous and uniform Si nanocrystal distribution. The emission power efficiency is relatively low, \( \sim 10^{-3}\% \), and the emission intensity exhibits fast degradation rates, as revealed from accelerated ageing experiments.

Devices fabricated by chemical deposition only exhibit field-effect luminescence, whose onset decreases with the Si excess from 20 to 6 V. The absence of the continuous emission is explained by the observation of a 5 nm region free of nanocrystals, which strongly reduces the direct current through the gate. The main benefit of having this nanocrystal-free region is that tunnelling current flow assisted by nanocrystals is blocked by the SiO2 stack so that power consumption is strongly reduced, which in return increases the device power efficiency up to 0.1%. In addition, the accelerated ageing studies reveal a 50% degradation rate reduction as compared to implanted structures.

1. Introduction

Monolithic silicon light-emitting diodes (LEDs) and lasers are expected to dramatically evolve silicon technology by opening up new areas of application for Si never thought possible in the past [1–5]. With the aim of improving the radiative recombination rate in Si, Si nanocrystals (Si-nc) have been object of extensive research since the electron and hole wavefunctions are confined into a small volume. Nowadays, systems composed of Si-nc embedded in SiO2 (silicon dioxide) represent a promising alternative thanks to their chemical stability and tunable emission in the red–infrared range (from 1.4 to 1.7 eV).
In this work, we establish a correlation between the properties of LEDs fabricated under similar conditions. The and ion implantation and the electrical and electro-optical consumption, leading to higher emission efficiencies.

(ii) the achievement of uniform Si profiles in thick oxide layers is limited by the energy required to perform deep implantation processes. By PECVD, the fabrication of thick SRO layers is possible. The plasma-enhanced process stands out against the different deposition techniques as thicker layers can be rapidly grown at low temperatures. However, as in the case of the ion implantation technique, the concentration of defects must be controlled, a process that strongly depends on the deposition rate.

In recent years, the emission from Si-nc-based materials under electrical pumping has been widely reported [6–9], mostly under DC (direct current) polarization. The application process were previously simulated by SRIM (stopping and range of ions in matter) calculations [12]. The phase separation and growth of the Si-nc was achieved by subjecting the SRO layers to high temperature annealing at 1100°C in N₂ atmosphere for 4 h.

Ellipsometry measurements were performed to evaluate the thickness and the refractive index of the different active layers using a Rudolph Research AutoEl IV ellipsometer, operating at 632.8 nm under an incidence angle of 70°. XPS (x-ray photoelectron spectroscopy) measurements were performed in order to study the composition profiles. The cross-section images of the samples and the size distribution of Si nanocrystals were obtained by EFTEM (energy-filtered transmission electron microscopy). Photoluminescence (PL) spectra were measured by exciting the samples with an He–Cd laser at 325 nm. The detection set-up was composed of a Hamamatsu thermoelectrically cooled H7422-50 photomultiplier and a Stanford Research SR430 photon-counting optic modulator with ~0.195 µs characteristic rise time. The emission transients were detected with a photon-counting set-up composed of a Hamamatsu thermoelectrically cooled H7422-50 photomultiplier and a Stanford Research SR430 multi-channel scaler/averager.

In the next stage, Si-nc-based metal oxide semiconductor (MOS) structures were fabricated and characterized. Tables 3 and 4 show some experimental data of PECVD and implanted devices, respectively. PECVD devices are labelled as PD and implanted ones are labelled as ID. Note that, in this case, the fabrication parameters are slightly different from those observed in tables 1 and 2. This new choice meets the necessity of improving the conductivity of the active layers in order to facilitate the emission under electrical pumping.

To establish a reliable comparison, deposited and implanted

<table>
<thead>
<tr>
<th>Table 1. PECVD samples.</th>
<th>Thickness (nm)</th>
</tr>
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<tbody>
<tr>
<td>Sample</td>
<td>[Si]ex at.%</td>
</tr>
<tr>
<td>P1 22.00</td>
<td>5.39 ± 1.64</td>
</tr>
<tr>
<td>P2 9.17</td>
<td>10.92 ± 1.41</td>
</tr>
<tr>
<td>P3 5.50</td>
<td>16.94 ± 1.37</td>
</tr>
<tr>
<td>P4 3.60</td>
<td>26.10 ± 1.38</td>
</tr>
<tr>
<td>P5 2.20</td>
<td>42.06 ± 1.33</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 2. Ion-implanted samples.</th>
<th>Thickness (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample</td>
<td>Dose (×10¹⁶ cm⁻²)</td>
</tr>
<tr>
<td>-------</td>
<td>------------------</td>
</tr>
<tr>
<td>I1</td>
<td>0.6</td>
</tr>
<tr>
<td>I2</td>
<td>2.4</td>
</tr>
<tr>
<td>I3</td>
<td>1.2</td>
</tr>
<tr>
<td>I4</td>
<td>2.4</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Table 3. MOS devices (PECVD).</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample</td>
</tr>
<tr>
<td>-------</td>
</tr>
<tr>
<td>PD1</td>
</tr>
<tr>
<td>PD2</td>
</tr>
<tr>
<td>PD3</td>
</tr>
<tr>
<td>PD4</td>
</tr>
<tr>
<td>PD5</td>
</tr>
</tbody>
</table>
The hydrogen comes from the decomposition of silane (SiH₄) and C using an Agilent B1500 semiconductor device analyser. 1 (liquid-nitrogen) cooled CCD camera and an Oriel MS257 measured by means of a Princeton Instruments 100B LN counting set-up described above. The EL spectra were resolved electroluminescence was detected with the photon-taking by polarizing with variable width and frequency bipolar and optical transmission measurements were performed with so, poly-Si layers were deposited onto fused silica substrates of the polycrystalline silicon layers were measured. To do the top electrode on the luminescence, transmittance spectra noise ratio of EL. In order to characterize the influence of the N₂O precursor.

These layers are also characterized by a relatively high feature for the achievement of uniform Si-nc distributions. flat Si concentration profiles in all the SRO layers, a desirable samples with high Si contents. XPS profiling revealed quite quite nanocrystals of about 4 nm. An important result depicted in table 2. The thickness was determined by ellipsometry and the Si excess of the different layers are indicated

Table 4. MOS devices (ion-implanted).

<table>
<thead>
<tr>
<th>Sample</th>
<th>Dose (×10¹⁶ cm⁻²)</th>
<th>Ion energy (keV)</th>
<th>[Si]ex (at.%)</th>
<th>Thickness (nm)</th>
<th>Cmax (pF)</th>
<th>Nₐ (cm⁻²)</th>
<th>Charged nc (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ID1</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>41.1 ± 0.3</td>
<td>215.51</td>
<td>8.456 × 10⁸</td>
<td>—</td>
</tr>
<tr>
<td>ID2</td>
<td>1.2</td>
<td>25</td>
<td>—</td>
<td>46.3 ± 0.5</td>
<td>277.88</td>
<td>1.199 × 10¹²</td>
<td>16.2</td>
</tr>
<tr>
<td></td>
<td>5.0</td>
<td>50</td>
<td>9.6</td>
<td>49.4 ± 0.2</td>
<td>294.78</td>
<td>1.509 × 10¹²</td>
<td>19.3</td>
</tr>
<tr>
<td>ID3</td>
<td>1.4</td>
<td>25</td>
<td>10.5</td>
<td>48.8 ± 0.2</td>
<td>320.51</td>
<td>9.017 × 10¹¹</td>
<td>12.0</td>
</tr>
<tr>
<td>ID4</td>
<td>1.8</td>
<td>25</td>
<td>11.7</td>
<td>47 ± 3</td>
<td>308.60</td>
<td>4.167 × 10¹¹</td>
<td>5.7</td>
</tr>
<tr>
<td></td>
<td>7.2</td>
<td>50</td>
<td>11.7</td>
<td>46 ± 3</td>
<td>315.74</td>
<td>3.558 × 10¹¹</td>
<td>4.4</td>
</tr>
<tr>
<td>ID5</td>
<td>1.8</td>
<td>25</td>
<td>—</td>
<td>50 ± 4</td>
<td>315.74</td>
<td>3.558 × 10¹¹</td>
<td>—</td>
</tr>
<tr>
<td>ID6</td>
<td>2.0</td>
<td>25</td>
<td>—</td>
<td>50 ± 4</td>
<td>315.74</td>
<td>3.558 × 10¹¹</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td>8.3</td>
<td>50</td>
<td>12.6</td>
<td>—</td>
<td>—</td>
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<td>—</td>
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</tbody>
</table>

3. Structural characterization

3.1. PECVD

Table 1 shows, apart from the nominal [N₂O]/[SiH₄] ratio and the Si excess yield by XPS measurements, the thickness of as-deposited and annealed PECVD layers measured by ellipsometry. After high temperature annealing (1250 °C), the ellipsometry measurements revealed an important reduction in the layer thickness due to H desorption and atomic redistribution. In good agreement with the flat Si profile observed by XPS, the Si-nc distribution is quite uniform, with a mean size of ∼3.60 nm (standard deviation of σ = ±0.50 nm), a density of nanocrystals of about 9.0 × 10¹⁷ cm⁻³ and an average distance between the closest nanocrystals of about 4 nm. An important result depicted in figure 1(a) is the appearance of a ∼5 nm Si-nc-free region close to the substrate, which has an important impact on device performance (as discussed hereafter). This absence of nanoparticles is attributed either to a diffusion of Si atoms to the substrate during the annealing and to the characteristic deposition instabilities at the first stages of the PECVD process. On the other hand, it is worth noting the strong oxidation of the surface (∼15 nm), which is probably induced by the high porosity of the as-deposited layers. This layer will be eliminated for device fabrication.

3.2. Ion implantation

Implanted samples were characterized in a similar way. The thickness and the Si excess of the different layers are indicated in table 2. The thickness was determined by ellipsometry and the content of Si was extracted from the XPS profiles. Similarly to the PECVD samples, a flat profile of silicon is observed in the active layer in agreement with SRIM simulations. It is remarkable the significant content of nitrogen

structures were prepared with similar thicknesses, about 45–50 nm. N-doped semitransparent polycrystalline silicon (poly-Si) ~350 nm thick was deposited as a top contact layer and aluminium pads were placed to facilitate the probe connection. Before this step and only for the implanted structures, the nitride buffer was chemically etched. Even though the devices were prepared with different contact areas (see figure 3), only those with larger electrodes areas were studied (2.3 × 10⁻³ and 9.6 × 10⁻³ cm²), since they exhibit the best signal-to-noise ratio of EL. In order to characterize the influence of the top electrode on the luminescence, transmittance spectra of the polycrystalline silicon layers were measured. To do so, poly-Si layers were deposited onto fused silica substrates and optical transmission measurements were performed with a Shimadzu UV-2101 PC spectrometer. The EL data were taken by polarizing with variable width and frequency bipolar square waves from an Agilent 8114A Pulse Generator. Time-resolved electroluminescence was detected with the photocounting set-up described above. The EL spectra were measured by means of a Princeton Instruments 100B LN (liquid-nitrogen) cooled CCD camera and an Oriel MS257 1/4 m monochromator. Quasi-static I–V (current–voltage) and C–V (capacitance–voltage) characteristics were measured using an Agilent B1500 semiconductor device analyser.

Figure 1. EFTEM profile images of sample P2 (a) and sample I3 (b).
in the SRO layer, diffused from the buffer layer during implantation.

EFTEM images were taken in order to study the characteristics of the Si-nc distribution. Figure 1(b) shows an example of Si-nc spatial distribution. The cross-section view corresponds to sample I3. The distribution is continuous and uniform, unlike PECVD samples where a thin depletion of Si-nc was observed. An average size of $\sim 3.10$ nm (standard deviation $\sigma = \pm 0.47$ nm) was measured with an Si-nc density of about $1.6 \times 10^{18}$ cm$^{-3}$ and an average distance between closest neighbours of $\sim 3$ nm. Note that, in spite of the similar Si contents in samples I3 and P2, the mean Si-nc size is slightly different ($\sim 3.10$ nm in sample I3 as opposed to $\sim 3.60$ nm in sample P2) which is, in some way, compensated by the different Si-nc densities observed. The $\sim 5$ nm SiO$_2$ layer observed at the surface is a signature of oxidation by air exposure, probably induced by the implantation process. Also in this case, prior to device fabrication this layer is removed.

4. Emission properties

Figure 2 compares the PL spectrum of samples P2 and I3. The emission, placed in the range of 1.4–1.7 eV, correlates with the typical luminescence in Si-nc/SiO$_2$ systems. The inset shows the dependence of the PL intensity in deposited and implanted samples as a function of the Si excess.

The inset of figure 2 depicts the PL intensity of implanted and PECVD samples for different Si excesses. The PL intensity of the implanted sample (I3) is lower than that of the PECVD sample (P2) for the same Si excess. This is due to the lower density of Si-nc in sample I3, which leads to a lower number of absorbing states and thus, a lower PL intensity.

In order to calculate the absorption cross section of the different samples (implanted and PECVD), time-resolved measurements were performed. The PL transients were modelled by means of stretched exponential functions [16], which led, in both cases, to emission lifetimes in the range of $\sim 10$–$\sim 45$ $\mu$s with values of $\beta$ in the range of 0.60–0.70. With respect to the absorption cross sections, our calculations reveal values of $\sigma_{abs}$ of the order of $10^{-16}$ cm$^2$ for both kinds of samples, in good accordance with values typically reported [17].

5. Device characterization

Metal-oxide-semiconductor structures with different active areas were fabricated by means of a standard photolithographic process. Figure 3(a) is a top-view image of the device. A basic cross-section scheme of these devices can be seen in figure 3(b). The top electrode is a 350 nm semitransparent poly-Si with an average transmittance of about 50% in the range of interest. The active layers typically have $\sim 45$–50 nm and have been prepared with different Si contents, taking into account the results depicted in the inset of figure 2 and bearing in mind the necessity of improving the conductivity of the active layers. In order to maximize the signal-to-noise ratio of the electroluminescence trace, this work is mainly focused on devices with larger electrode areas ($9.6 \times 10^{-3}$ and $2.3 \times 10^{-3}$ cm$^2$).

5.1. PECVD

Table 3 summarizes some basic characteristics of MOS structures fabricated by PECVD.

Forward–backward $C–V$ characteristics were measured (see the example in the inset of figure 4) in order to obtain the density of trapped charge inside the SRO layer, $N_t$. To ensure an effective carrier injection, the voltage sweeps were adjusted for each sample taking into account its respective onset of emission and breakdown voltage (if any). The results are summarized in table 3. The trapped charge is calculated...
from the shifts of the flat-band voltage, $\Delta V_{FB}$, measured from a reference $C-V$ curve of a fresh device:

$$N_t = \frac{C_{max} \Delta V_{FB}}{eS}$$

where $C_{max}$ represents the maximum capacitance of the structure (capacitance in accumulation regime), $e$ is the electron charge and $S$ is the device area. The values of $N_t$ are indicated in table 3. The density of trapped charge shows a significant increase with the Si content, which is due to the concurrence of the following features: (1) the enhanced conductivity of the SRO layers and (2) the increase of the average Si-nc size that facilitates the carrier trapping.

The density of trapped charge, in combination with the Si-nc density obtained by EFTEM, leads to a rough estimation of the percentage of charged nanocrystals. This parameter is rather an upper bound to the ratio of excitable Si-nc and, in some way, can be considered as a marker of the room available to further improve the excitation efficiency, since it also indicates the ratio of non-active Si-nc. The calculation is performed under the assumption that most of the trapped charge is associated with the Si-nc, which is justified by the low value of $N_t$ observed in pure oxide structures (device PD1). Nevertheless, it is also true that the addition of the Si excess causes the generation of defects and traps not observed in SiO$_2$ that probably leads to an overestimation of the number of active Si-nc. Figure 4 shows the results of these calculations. For low Si contents, i.e. structures PD2 and PD3, the thin zone depleted of Si-nc strongly limits the carrier injection from the substrate, leading to occupation ratios of $\sim 0.4\%$ and $\sim 8.9\%$, respectively. As the Si excess further increases (up to $\sim 13.7\%$, sample PD4), there is a significant increase in the number of carriers injected from the substrate. In this case, the higher Si concentration facilitates the tunnelling from the substrate at lower voltages, leading to the highest percentages of trapped charge in devices PD4 and PD5 with values close to $\sim 90\%$. This behaviour is very similar to that observed when the tunnel oxide thickness is modified, as reported in reference [18].

In order to study the conduction mechanisms, quasi-static current–voltage characteristics were acquired under forward–backward voltage sweeps. In all the PECVD structures, the current is, in the range of interest (emission range), well fitted by Fowler–Nordheim (FN) tunnelling with injection barrier heights that decrease with the Si content: from 2.3 eV in sample PD2 (7.9%) to 0.9 eV in sample PD5 (18.8%). In accordance with this, the onset of the FN injection also exhibits significant reduction with the Si content: from 23 V in sample PD2 to 6–7 V in sample PD2.

Figure 5 summarizes the behaviour of the field-effect luminescence in PECVD structures. Figure 5(a) shows the time-resolved EL of sample PD3 under application of square wave voltages with an amplitude of 25 V, whereas figures 5(b) and (c) represent the PL and EL spectra of similarly prepared samples, respectively. The onset of the emission decreases with the Si content from $\sim 20$ V, in structure PD2 (7.9%), to $\sim 6$ V, in sample PD5 (18.8%), as depicted in figure 6(a). It should be noted the good accordance between the onsets of emission and FN injection, which clearly links the field-effect emission of PECVD samples to this injection mechanism. The field-effect mechanism entails the sequential injection of electrons and holes from the substrate, in a process assisted by the Coulomb field attraction from the previously stored carriers. Note that, during the second carrier injection, a sizeable percentage of the trapped charge tunnels back to the substrate and therefore the achievement of an efficient emission depends on both the retention time of the first carrier injected and the tunnelling time of the incoming second carrier. Due to the lower barrier for electrons, the migration of the stored charge is more marked when holes are injected into electron-charged Si-nc, which results in a weaker emission compared to the
case of electron injection into hole-charged nanocrystals [22]. This behaviour is clearly seen in figure 5(a), where the EL intensity of the peaks differs by approximately one order of magnitude. The hole-triggered peak is slightly faster in duration, even though the difference with the peak associated with the electron injection into hole-charged Si-nc does not appear to be significant. In any case, both transients are faster than those observed under optical excitation. The emission decays are well fitted by stretched exponential functions with time constants slightly shorter than in the case of PL, ranging from $\sim 5$ to $\sim 20$ $\mu$s, with $\beta \sim 0.5$. Operating under a field-effect mechanism, the absorption cross sections are of the order of $\sim 10^{-11}$ cm$^2$, approximately three orders of magnitude larger than values usually reported under DC excitation [19]. Some authors have pointed out [20, 21] that typically reported values of $\sim 10^{-14}$ cm$^2$ should be interpreted as a lower bound, since leakage currents not contributing to the emission are unavoidably included in the calculations. In addition, in reference [20], Carreras et al reported a theoretical estimate of the field-effect absorption cross section of about $10^{-12}$ cm$^2$. In the light of this, our results should be considered as the consequence of a strong reduction of the leakage current (thanks to the AC polarization) that leads to a better profit of the injected charge. The comparison of PL and EL spectra reveals modulation of the emission due to interference phenomena inside the poly-Si electrode. This effect is clearly seen if one compares the measured EL spectrum with the one obtained by combining the PL emission and the electrode transmittance represented in 5(b). The good match between both spectra (see figure 5(c)) underlines the relevant role played by the top contact on the emission properties.

Figure 6(b) shows the EL and PL intensities as a function of the Si content. The EL corresponds to a fixed current density of $\sim 10 \, \mu$A cm$^{-2}$. In accordance with the quantum confinement theory (QCT), PL and EL intensities exhibit similar behaviours, even though the EL intensity peaks at higher Si contents than in the case of the emission under optical pumping, $\sim 13.7\%$ versus $\sim 11.2\%$. This difference is due to the fact that, under electrical pumping, an optimum emission requires an additional amount of Si in order to improve carrier injection. The device breakdown also depends on the Si excess, since it is related to the current level through the device. The samples with large Si contents, PD5 and PD6, break down at about $14$ V and $22$ V respectively, whereas the samples with low Si concentrations do not show an observable breakdown, even for voltages higher than $30$ V.

5.2. Ion implantation

The characteristics of the different devices fabricated by ion implantation are presented in figure 4. $C–V$ characteristics were measured in order to obtain the density of trapped charge, $N_t$. The inset of figure 7 shows a typical $C–V$ curve of implanted devices. In general, the capacitance hysteresis is characterized by a sizeable trapping of electrons that is related to an easier carrier injection (if compared with PECVD), as demonstrated below. The density of trapped charge is calculated from the capacitance hysteresis cycle with a similar procedure than in the case of PECVD samples. The calculations reveal a maximum value of $N_t$ in structures with low Si concentrations. In those structures (ID2 and ID3), typical densities of $\sim 1.0–1.5 \times 10^{12}$ cm$^{-2}$ are observed that represent percentages of charged nanocrystals of about 16–19%. As the Si concentration increases, there is a progressive reduction of trapped charge (see figure 7) probably related to the increased conductivity and the subsequent carrier migration. The trapped charge decreases down to $\sim 4 \times 10^{11}$ cm$^{-2}$ in structure ID6, which corresponds to an occupation ratio of $\sim 4.4\%$. In order to study the conduction mechanisms, $I–V$ characteristics were acquired. The current is, in general, higher than in the case of PECVD structures due to the continuous and uniform Si-nc distribution. The $I–V$ curves are well fitted by FN expressions with low injection barriers (if compared with PECVD), as demonstrated below. The density of trapped charge is calculated from the capacitance hysteresis cycle with a similar procedure than in the case of PECVD samples. The calculations reveal a maximum value of $N_t$ in structures with low Si concentrations. In those structures (ID2 and ID3), typical densities of $\sim 1.0–1.5 \times 10^{12}$ cm$^{-2}$ are observed that represent percentages of charged nanocrystals of about 16–19%. As the Si concentration increases, there is a progressive reduction of trapped charge (see figure 7) probably related to the increased conductivity and the subsequent carrier migration. The trapped charge decreases down to $\sim 4 \times 10^{11}$ cm$^{-2}$ in structure ID6, which corresponds to an occupation ratio of $\sim 4.4\%$. In order to study the conduction mechanisms, $I–V$ characteristics were acquired. The current is, in general, higher than in the case of PECVD structures due to the continuous and uniform Si-nc distribution. The $I–V$ curves are well fitted by FN expressions with low injection barriers (if compared with PECVD), as demonstrated below. The density of trapped charge is calculated from the capacitance hysteresis cycle with a similar procedure than in the case of PECVD samples. The calculations reveal a maximum value of $N_t$ in structures with low Si concentrations. In those structures (ID2 and ID3), typical densities of $\sim 1.0–1.5 \times 10^{12}$ cm$^{-2}$ are observed that represent percentages of charged nanocrystals of about 16–19%. As the Si concentration increases, there is a progressive reduction of trapped charge (see figure 7) probably related to the increased conductivity and the subsequent carrier migration. The trapped charge decreases down to $\sim 4 \times 10^{11}$ cm$^{-2}$ in structure ID6, which corresponds to an occupation ratio of $\sim 4.4\%$.
the negative half-cycle of the driving signal. This component is approximately one order of magnitude larger than the field-effect luminescence (for driving signals of $\sim 1$ kHz) and is likely related to the continuous Si-nc distribution observed in implanted structures. The inset of figure 8(b) compares the average current through the SRO layer with the intensity of the DC EL; both magnitudes as a function of the driving voltage. The good accordance between both curves suggests that impact ionization is the mechanism responsible for the DC emission. Bipolar injection from opposite electrodes could also be possible and could perfectly explain the DC component observed, since it is only present during the negative half-cycle of the gate bias. However, this possibility is less probable due to the thickness of the layers.

The transients of the DC contribution are approximately one order of magnitude longer than those of the field-effect emission. Decay times of up to 0.7 ms (structure ID6) with $\beta \sim 0.5$ were observed. These lifetimes were extracted by modelling the de-excitation at the rise edge of the gate bias with two exponentials: the first one corresponding to the fast AC transient and the second one associated with the continuous emission. The value of $\sigma_{\text{EL}}^{\text{abs}}$ was found around $3 \times 10^{-14}$ cm$^2$, in agreement with values usually reported from Si-nc-based devices under DC polarization [19].

With respect to the field-effect transients, it is clearly seen in figure 8(a) that the electron-triggered emission is larger in intensity than that observed when holes are injected into electron-charged Si-nc, even though the difference between these two peaks is not as significant as in deposited structures (due to some extent to the higher conductivity of the layer). The decay times of both peaks are very similar with values in the range of $\sim 20–30$ $\mu$s. These decays were extracted from stretched exponential fits with typical $\beta$ of about 0.5. Absorption cross sections of about $2 \times 10^{-12}$ cm$^2$ were obtained, approximately two orders of magnitude higher than that for the DC emission. This difference suggests that the field-effect mechanism takes best profit of the injected charge in comparison to the impact ionization mechanism. Note that in PECVD samples the value of $\sigma_{\text{EL}}^{\text{abs}}$ is one order of magnitude higher, which is attributed to their relatively low operation currents.

Figure 8(b) shows the EL spectra of the different implanted devices under 9 V square wave voltages. The apparent immobility of the emission peak with the Si content is related to the modulation induced by the transmittance of the poly-Si layer, as discussed above. Figure 9(a) represents the EL onset voltage as a function of the Si excess. The addition of Si favours the carrier injection and shifts the onset voltage to lower values. Nevertheless, this leads to higher current flows that unavoidably lead to lower breakdown voltages, as shown in figure 9(b). Note that also in the case of implanted MOSLEDs there is a good agreement between the onsets of emission and FN tunnelling. The integrated EL was studied in the different structures for a given current flow of $\sim 52$ $\mu$A cm$^{-2}$ (see figure 9(b)). The dependence of the emission on the Si excess is very similar to the results presented in the inset of figure 2, and is in good agreement with the QCT. The maximum emission is observed in device ID5 (11.7% of Si excess).

6. LED performances

In order to study the suitability of PECVD and implanted devices as candidates for an Si-nanocrystal-based LED implementation, we compare in this section the best operational performances of these structures. Figure 10(a) shows the response of the field-effect luminescence of samples ID5 and PD3 (with similar Si contents) to different driving frequencies. First, the curves show that, as the integration time is held constant, enhancement of the EL signal is initially observed due to the increasing number of integrated cycles [22]. The EL intensity peaks at a frequency that is determined by both the tunnelling dynamics and recombination
devices exhibit typical emission powers of amplitudes close to the breakdown voltage (if any). Implanted of carrier migration phenomena. Figure 10(c) compares the with the loss of quantum confinement and to the emergence PD5 is, in contrast to samples PD2 and PD3, associated intense emission is observed. The weak emission of sample (∼10−3 %) that rapidly decrease with the applied voltage as the increasing power consumption is not compensated by the gain in the output power. Thanks to the low current levels observed in PECVD samples, their values of power efficiency are typically two orders of magnitude higher than that observed in implanted structures, up to ∼0.1%. Also in this case, a fast reduction of the efficiency with the applied voltage is observed, mostly due to the strong increase of the FN current at higher fields. In the case of the MNOS structure, the nitride buffer reduces the device leakage current leading to a better profit of the injected charge that translates into power efficiencies of about 1% and output emission powers of ∼10 nW. It is worth noting the slow variation of the power efficiency with the gate bias in MNOS structures that is attributed to the increase of the FN onset [23].

Based on these results, it is clear that, in order to achieve efficient devices, the fabrication of discontinuous Si-nc distributions appears to be beneficial, as it reduces the device power consumption. On the other hand, if we take into account the ratios of charged nanocrystals represented in figures 4 and 7, it is also clear that, apart from the reduction of the power consumption, the power efficiency can be improved by increasing the number of excited Si-nc. This should be done by accurately controlling the amount of defects and the size and position of the Si-nc distribution. In the light of our results, the combination of these two factors could enhance the power efficiency of Si-nc-based MOS devices up to 1–10% with typical powers in the range of ∼10–20 nW.

Implanted and PECVD structures were submitted to accelerated ageing for more than 24 h of continuous operation under application of 13 V and 30 V square waves, respectively, at a frequency of 1 kHz. Figure 11 shows the degradation of the emission for samples PD4 and ID6. The plot shows that sample ID6 exhibits an emission degradation rate of about 45% that contrasts with the lower value observed in sample PD4, about 20%. This difference correlates with the respective current levels observed in I–V characteristics. Both curves are well fitted by a logarithmic function within the experimental timescale [24], resulting in estimated lifetimes of at least several months.

7. Conclusions

The luminescence of Si-nc/SiO2 layers fabricated by PECVD and ion implantation has been thoroughly studied under optical and electrical excitation. The results have been correlated with an exhaustive characterization of the structural and electrical
properties. First, XPS and EFTEM measurements revealed that, in spite of the almost flat profile of Si excess, PECVD layers exhibited a ~5 nm Si-nc free region, attributed either to the diffusion of Si to the substrate or to the instabilities at the first stages of the deposition process. In implanted layers, the profile of added Si is also relatively flat (XPS) and translates into a uniform and continuous Si-nc distribution (EFTEM), which clearly underlines the suitability of the implantation process. We have also shown that, for similar Si contents, the average sizes of the Si-nc obtained by ion implantation are smaller than those observed in deposited layers, which leads to a higher Si-nc density and shorter interdot (interaction range) distances. The PL emission showed a good agreement between PECVD and implanted samples with maximum intensities of emission for layers with ~11% of Si excess. In both cases, the absorption cross sections are of the order of $10^{-16}$ cm$^2$.

EL measurements have been performed by polarizing with bipolar and monopolar square wave voltages. The EL spectra of implanted and PECVD structures reveal typical emission from Si-nc/SiO$_2$ systems that correlates with PL results. A strong modulation of the spectral shape and peak position is observed due to interference phenomena in the poly-Si top contact. The time-resolved EL revealed both AC and DC emission from implanted samples, in contrast to the measurements carried out in PECVD devices, where only transient emission was observed. The absence of emission under constant electrical excitation in the deposited layers is due to the discontinuous Si-nc distribution that hinders the DC flow through the structure. Moreover, this current reduction results in onset and breakdown voltages remarkably higher than in the case of implanted samples. The absorption cross sections of the implanted AC and DC emission are of the order of $10^{-12}$ and $10^{-14}$ cm$^2$, several orders of magnitude higher than those values obtained by photoexcitation. In the case of PECVD structures the value of $\sigma_{\text{abs}}^{\text{AC}}$ is approximately $10^{-11}$ cm$^2$. The maximum of the EL intensity is observed in implanted and PECVD structures with Si contents between 11–13%, apparently higher than that under optical excitation. We have also shown the limited frequency response of gate-biased Si-nc-based MOS structures, attributed to the characteristic radiative lifetimes of the Si nanoparticles. The maximum EL intensities obtained are ~0.3 nW for implanted devices and ~2 nW for PECVD. The efficiency of the implanted devices is of about $10^{-3}$%, two orders of magnitude lower than the one obtained for PECVD structures, 0.1%. This result tells us that, in terms of efficiency, a continuous distribution of Si-nc appears to be detrimental.

In order to evaluate the room available to further enhance the emission capabilities in these LEDs, a rough estimation of the electrically active Si-nc (percentage of charged nanocrystals) has been performed. We have shown that most of the structures (both PECVD and implanted) exhibit poor percentages of charged nanocrystals. With an accurate control of the amount of defects and the size and position of the Si-nc it could be possible to increase the excitation efficiency. Thus the optical output power could be enhanced up to ~20 nW, with typical power efficiencies in the range of 1–10%.

Finally, accelerated ageing studies reveal stronger degradation of the emission in implanted devices, which is attributed to their higher current levels in comparison to PECVD devices.

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