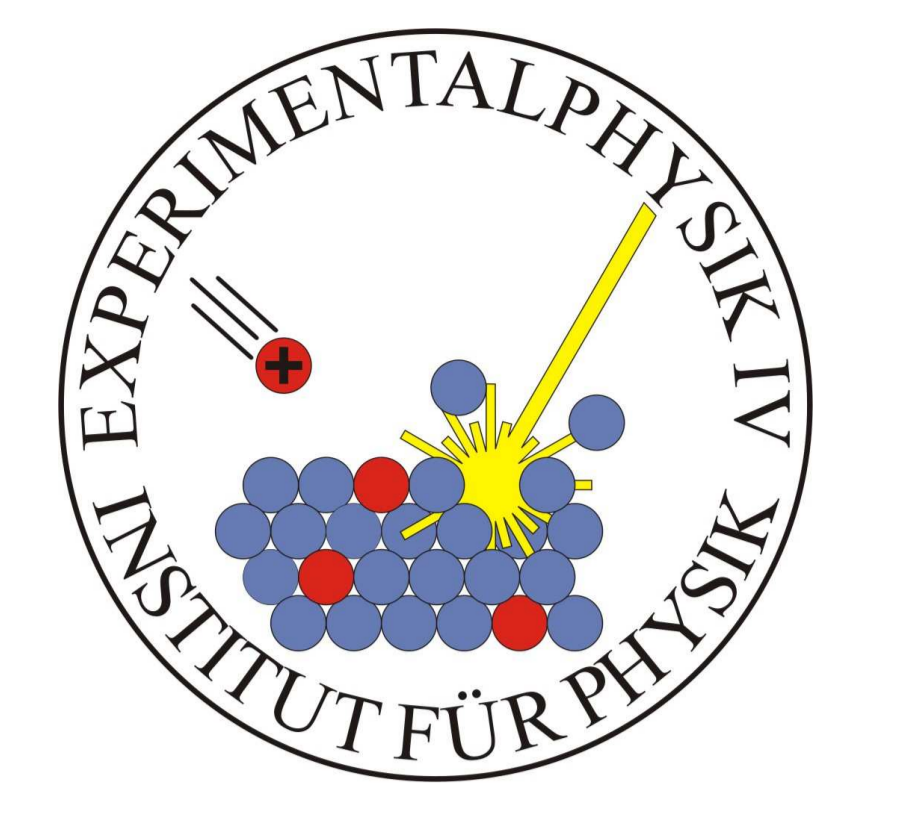


Monte Carlo simulation study of a selforganisation process leading to ordered precipitate structures

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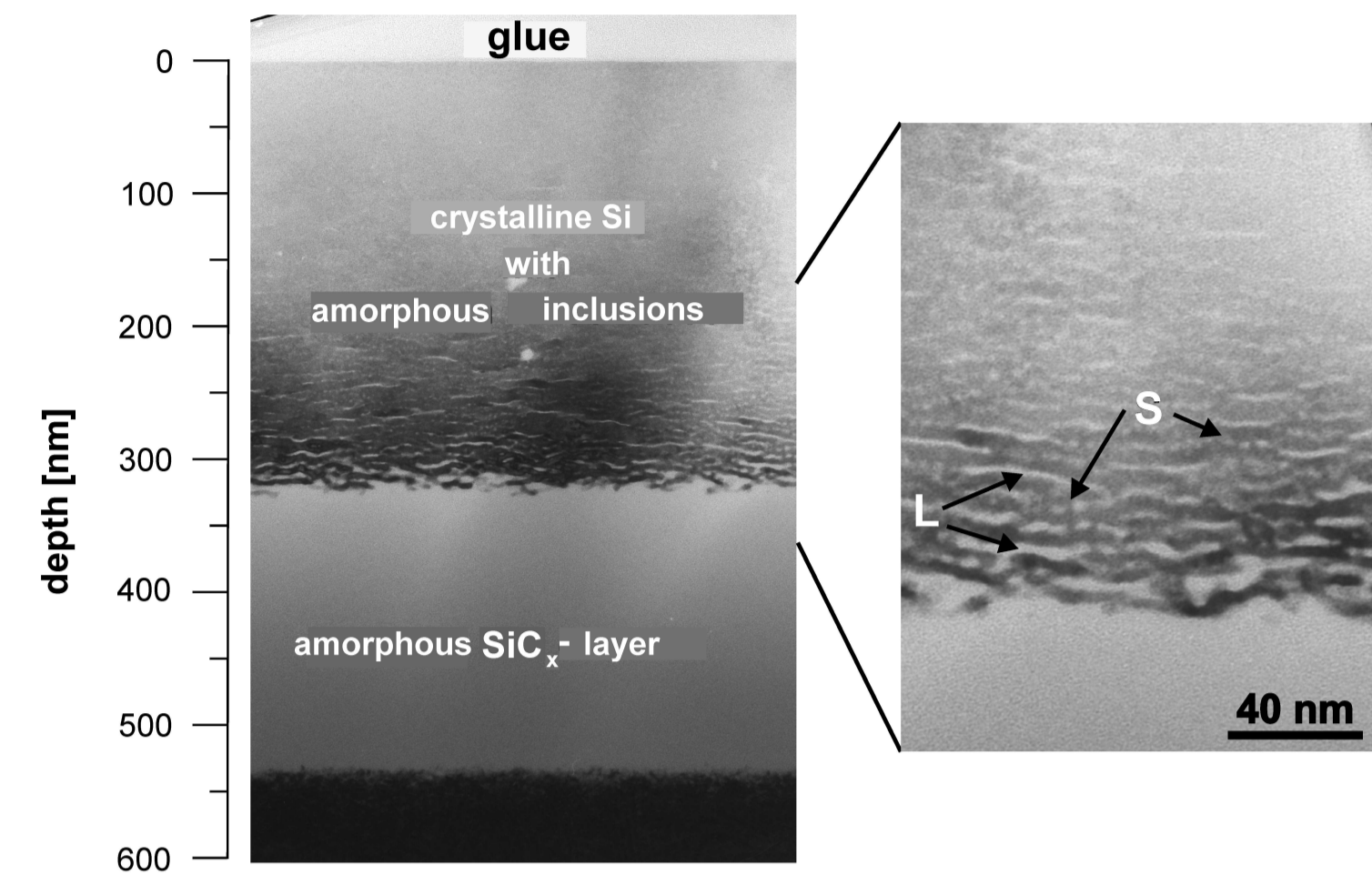
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Motivation

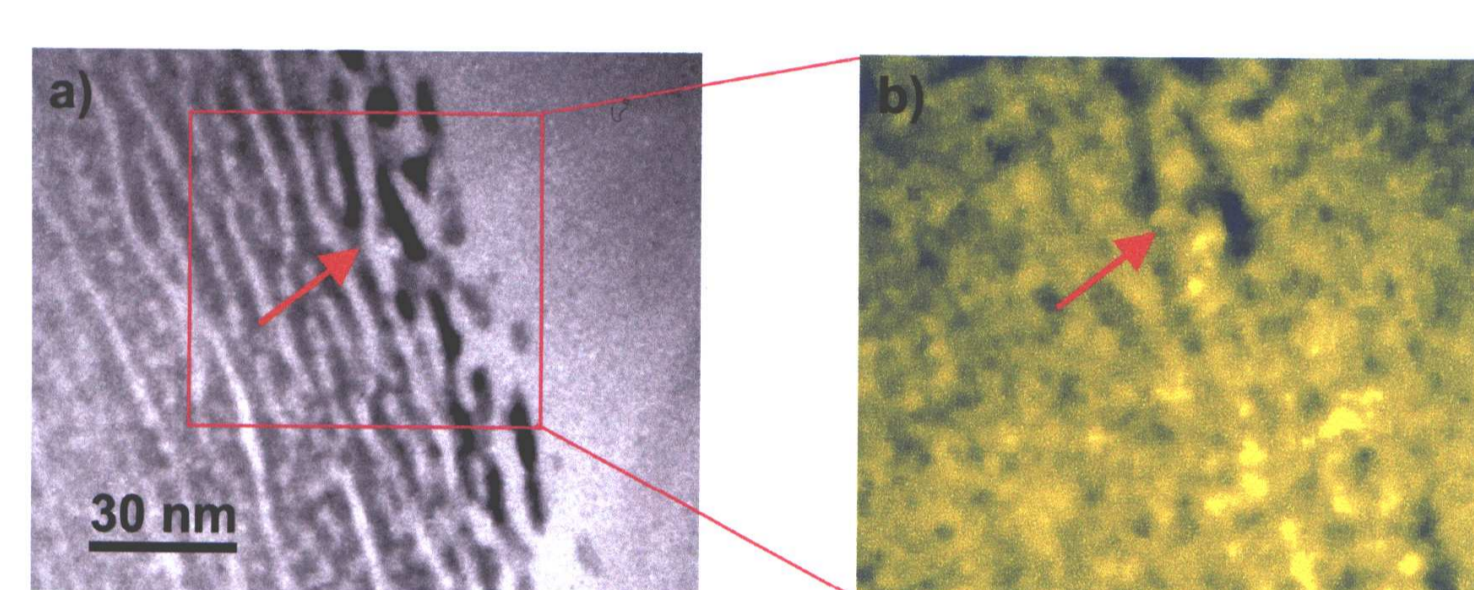
Experimentally observed selforganisation process at high-dose carbon implantations under certain implantation conditions.

- Regularly spaced, nanometric spherical and lamellar amorphous inclusions at the upper a/c interface



Cross-section TEM bright-field images:
 $180\text{ keV } C^+ \rightarrow Si$, $T_i = 150^\circ\text{C}$, Dose: $4.3 \times 10^{17} \text{ cm}^{-2}$
 Amorphous inclusions appear white on darker backgrounds
 L: amorphous lamellae, S: spherical amorphous inclusions

- Carbon accumulation in amorphous volumes



Bright-field TEM image and respective EFTEM C map:
 $180\text{ keV } C^+ \rightarrow Si$, $T_i = 200^\circ\text{C}$, Dose: $4.3 \times 10^{17} \text{ cm}^{-2}$
 yellow/blue: high/low concentrations of carbon

Similarly ordered precipitate nanostructures also observed for a number of ion/target combinations for which the material undergoes drastic density change upon amorphisation.

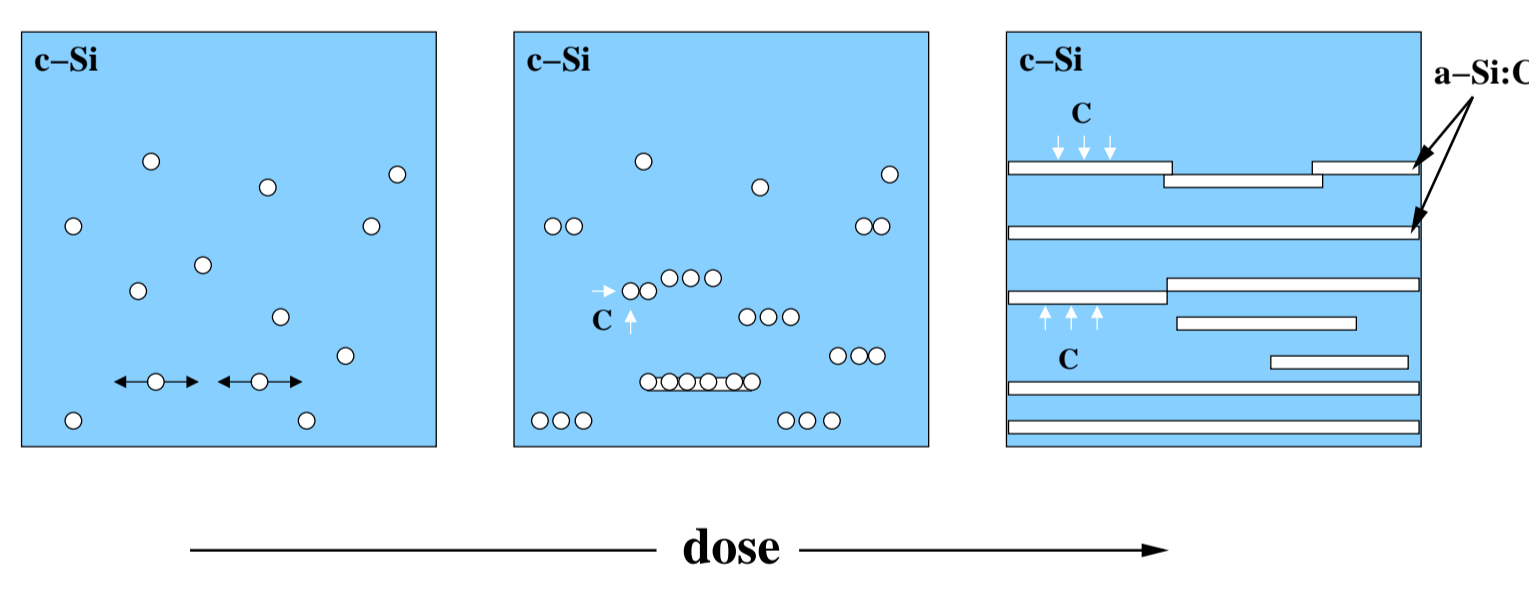
A. H. van Ommen, Nucl. Instr. and Meth. B 39 (1989) 194.

E. D. Specht et al., Nucl. Instr. and Meth. B 84 (1994) 323.

M. Ishimaru et al., Nucl. Instr. and Meth. B 166-167 (2000) 390.

Model

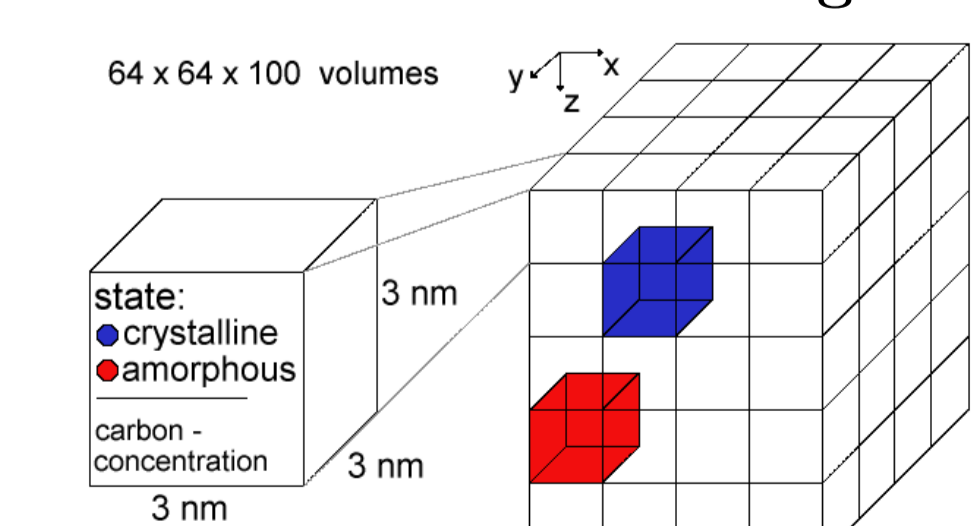
Model schematically displaying the formation of ordered lamellae with increasing dose.



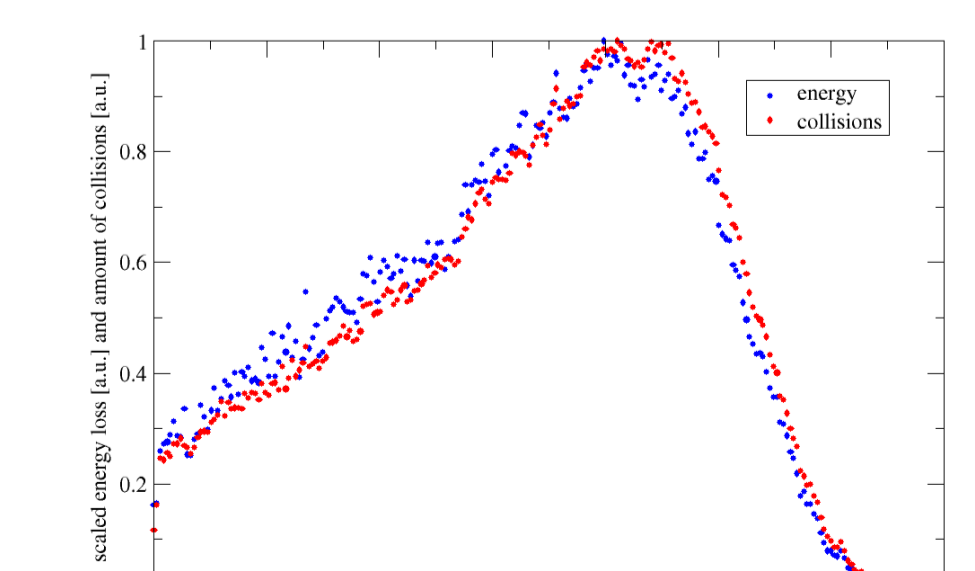
- Supersaturation of C in c-Si
 → Carbon induced nucleation of spherical SiC_x -precipitates
- High interfacial energy between $3C - SiC$ and c-Si
 → Amorphous precipitates
- 20 – 30% lower silicon density of a- SiC_x compared to c-Si
 → Lateral strain (black arrows)
- Implantation range near surface
 → Relaxation of vertical strain component
- Reduction of the carbon supersaturation in c-Si
 → Carbon diffusion into amorphous volumina (white arrows)
- Remaining lateral strain
 → Strain enhanced lateral amorphisation
- Absence of crystalline neighbours (structural information)
 → Stabilisation of amorphous inclusions against recrystallisation

Simulation

Discretisation of the target



TRIM collision statistics



- divided into cells with a cube length of 3 nm
- periodic boundary conditions in x,y-direction

⇒ identical depth profiles for number of collisions per depth and nuclear stopping power
 ⇒ mean constant energy loss per collision

Simulation algorithm

The simulation algorithm consists of the following three parts looped s times corresponding to a dose $D = s/(64 \times 64 \times (3\text{ nm})^2)$:

1. Amorphisation/Recrystallisation

- random numbers distributed according to the nuclear energy loss to determine the volume in which a collision occurs
- compute local probability for amorphisation:

$$p_{c \rightarrow a}(\vec{r}) = p_b + p_{ccC}(\vec{r}) + \sum_{\text{amorphous neighbours}} \frac{p_{scC}(\vec{r'})}{(r - r')^2}$$

and recrystallisation:

$$p_{a \rightarrow c}(\vec{r}) = (1 - p_{c \rightarrow a}(\vec{r})) \left(1 - \frac{\sum_{\text{direct neighbours}} \delta(\vec{r'})}{6} \right),$$

$$\delta(\vec{r}) = \begin{cases} 1 & \text{if volume at position } \vec{r} \text{ is amorphous} \\ 0 & \text{otherwise} \end{cases}$$

- loop for the mean amount of hits by the ion

Three contributions to the amorphisation process controlled by:

- p_b normal 'ballistic' amorphisation
- p_{cc} carbon induced amorphisation
- p_s stress enhanced amorphisation

2. Carbon incorporation

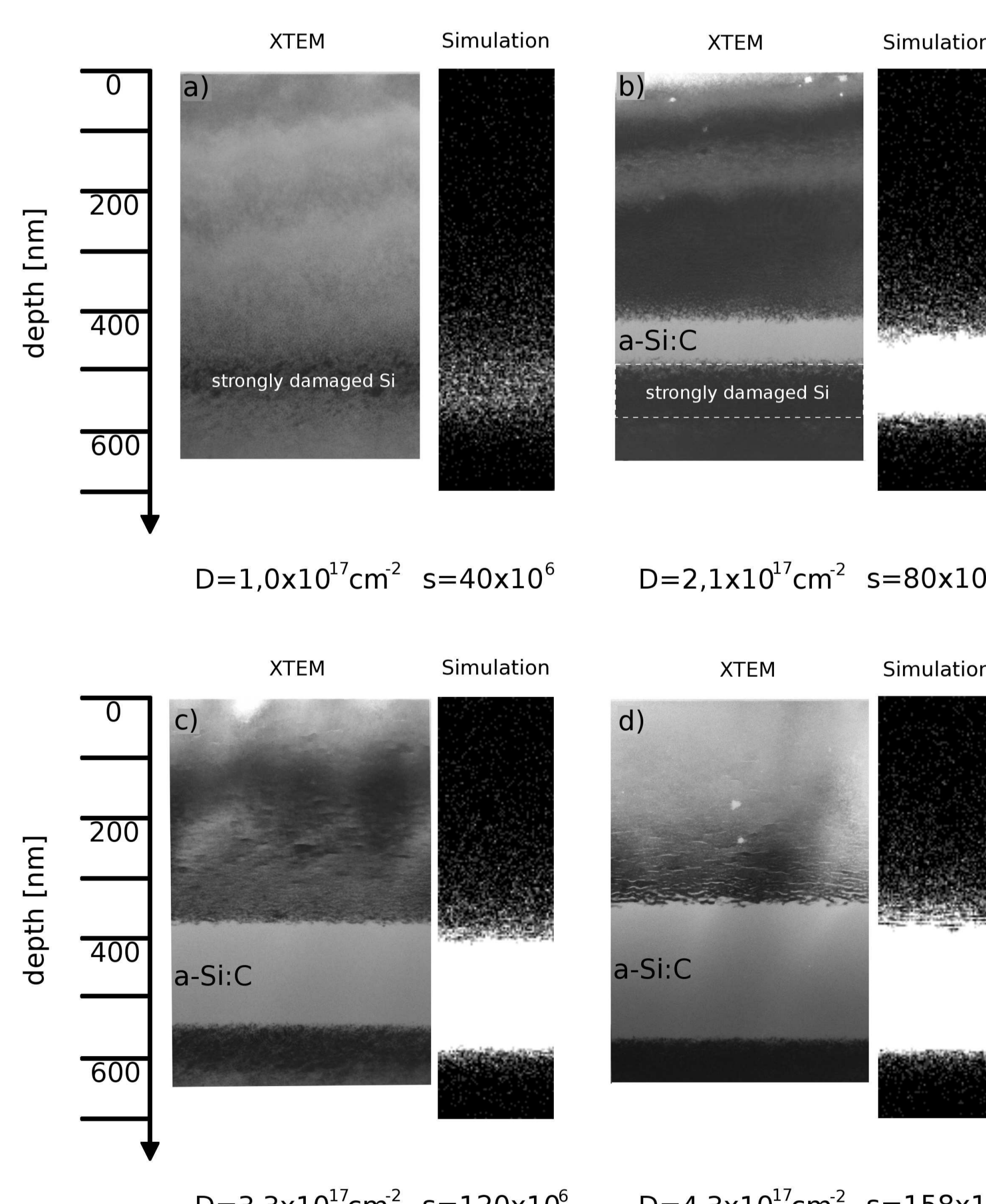
- random numbers distributed according to the implantation profile to determine the incorporation volume
- increase the amount of carbon atoms in that volume

3. Diffusion/Sputtering

Simulation parameters d_v , d_r and n control the diffusion and sputtering process.

- every d_v steps transfer of a fraction d_r of carbon atoms from crystalline volumina to an amorphous neighbour volume
- remove 3 nm surface layer after n loops, shift remaining cells 3 nm up and insert an empty, crystalline 3 nm bottom layer

Comparison of experiment and simulation



Simulation parameters:

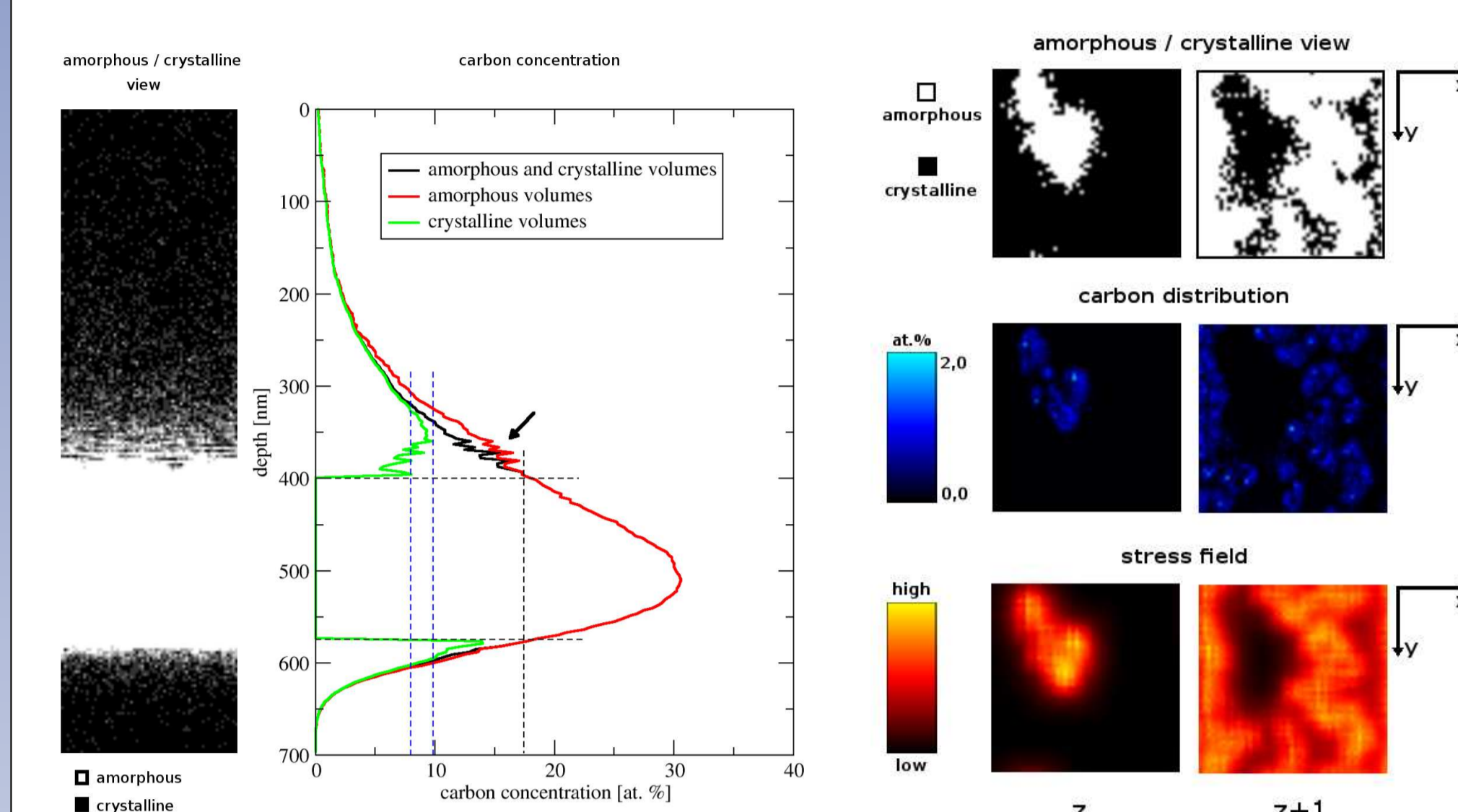
$$p_b = 0.01, p_c = 0.001 \times (3\text{ nm})^3, p_s = 0.0001 \times (3\text{ nm})^5, d_r = 0.05, d_v = 1 \times 10^6$$

Conclusion:

- Simulation in good agreement with experimentally observed formation and growth of the continuous amorphous layer
- Lamellar precipitates and their evolution at the upper a/c interface with increasing dose is reproduced

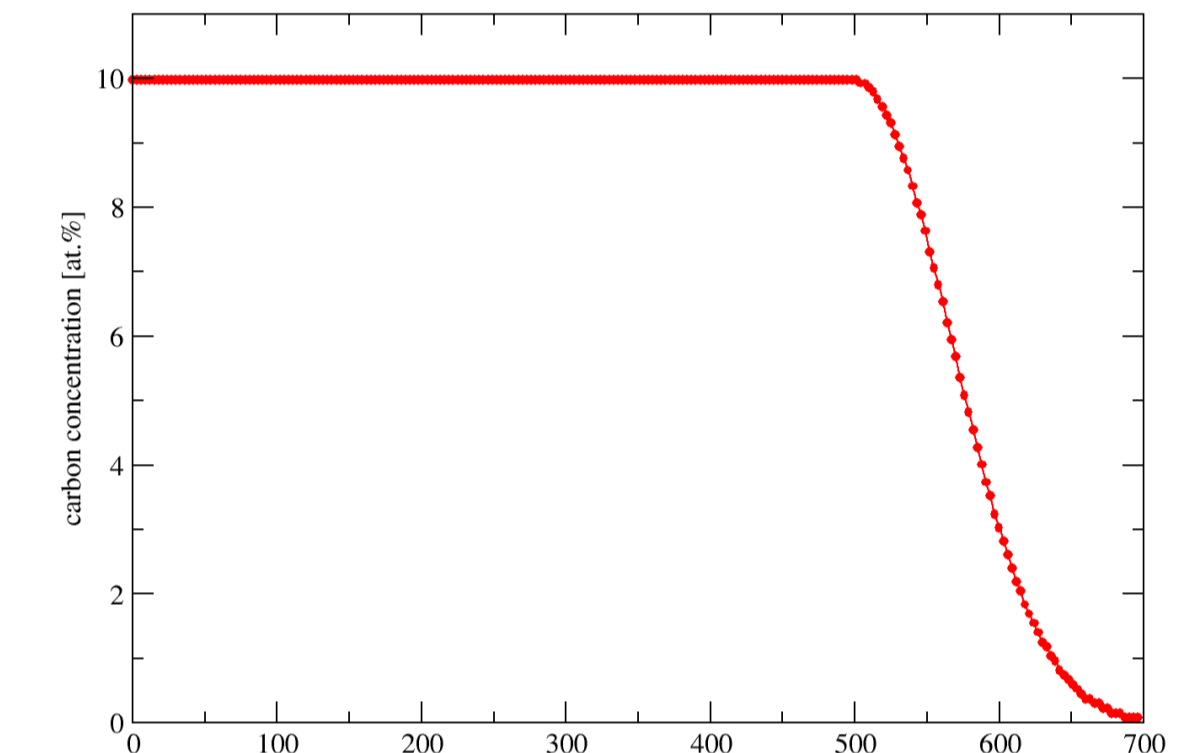
Simulation is able to model the whole depth region affected by the irradiation process

Structural/compositional information



- Fluctuation of the carbon concentration in the region of the lamellae
- Saturation limit of carbon in c-Si under given implantation conditions between 8 and 10 at.%
- Complementarily arranged and alternating sequence of layers with high and low amount of amorphous regions
- Carbon accumulation in the amorphous phase

Recipe for thick films of ordered lamellae



Prerequisites:

Crystalline silicon target with a nearly constant carbon concentration at 10 at.% in a 500 nm thick surface layer

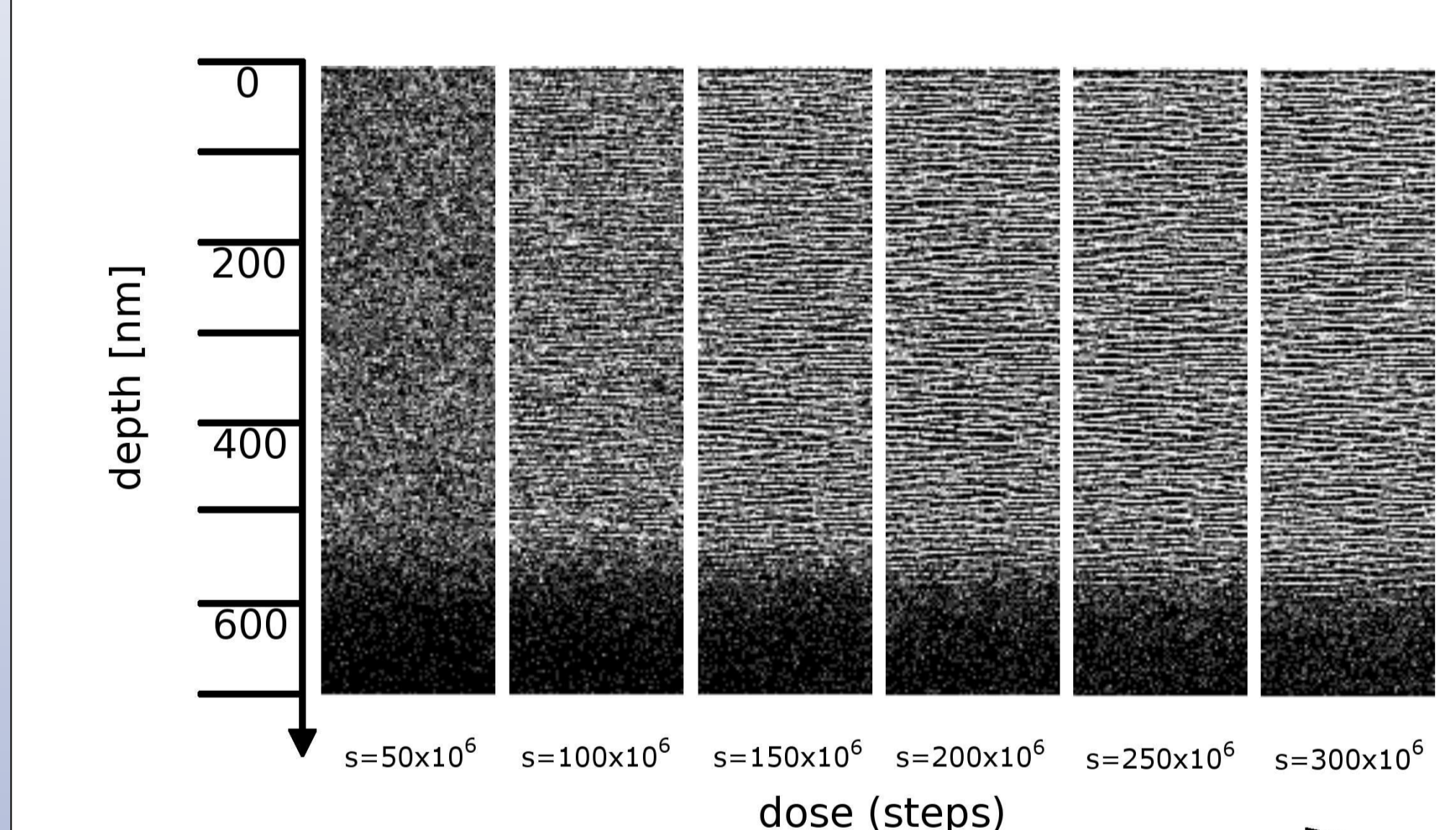
Creation:

- Multiple energy ($180\text{-}10\text{ keV}$) $C^+ \rightarrow Si$ implantation
- $T_i = 500^\circ\text{C}$, to prevent amorphisation

Stirring up:

- $2\text{ MeV } C^+ \rightarrow Si$ irradiation step at 150°C
- This does not significantly change the carbon concentration in the top 500 nm
- Nearly constant nuclear energy loss in the top 700 nm region

Result:



- Already ordered structures after 100×10^6 steps corresponding to a dose of $D = 2.7 \times 10^{17} \text{ cm}^{-2}$
- More defined structures with increasing dose

Starting point for materials showing strong photoluminescence

Dihu Chen et al. Opt. Mater. 23 (2003) 65.

Conclusions

- Observation of selforganised nanometric precipitates by ion irradiation
- Model proposed describing the selforganisation process
- Model implemented in a Monte Carlo simulation code
- Modelling of the complete depth region affected by the irradiation process
- Simulation is able to reproduce entire amorphous phase formation
- Precipitation process gets traceable by simulation
- Detailed structural/compositional information available by simulation
- Recipe proposed for the formation of thick films of lamellar structure

Literature

- F. Zirkelbach, M. Häberlen, J. K. N. Lindner, B. Stritzker. Comp. Mater. Sci. 33 (2005) 310.
 F. Zirkelbach, M. Häberlen, J. K. N. Lindner, B. Stritzker. Nucl. Instr. and Meth. B 242 (2006) 679.