Project 2

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Monte Carlo (MC) technique is becoming a very effective simulation method for prediction and analysis of the grain growth kinetics at mesoscopic level. The concept behind the Monte Carlo method in grain growth simulation is both simple and fascinating: its only basis is the thermodynamic of atomic interactions. There are no other experimental or theoretical inferences, nor mathematical approximations. The first step is to represent the material as a 2D or 3D matrix, in which each site corresponds to a surface or volume element. The content of each element represents its crystallographic orientation. During classes class algoritm was modified in accordance with the given guidelines. Additionally, new functionalities was made, such as: dual phase microstructures with CA simple growth and Monte Carlo; energy distribution and recrystalization.

Technology

Programming language: Python Language version: 3.7 Visualization library: Tkinter

User interface

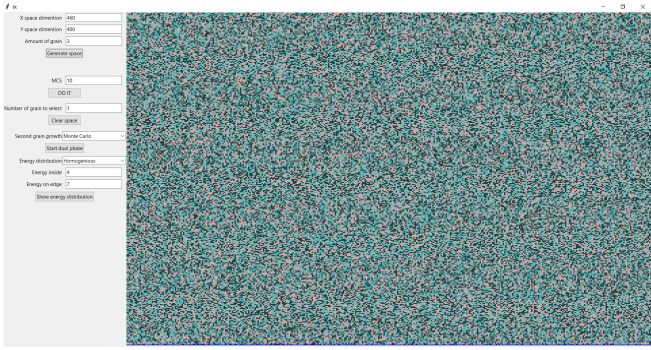


Image 1. User interface

In the Image 1 we can see main app view. On the left, there is main menu where we can start manipulate our simulation. On the right we have the main part of the application – microstructure visualization. The menu contain field with space generation, dualphase, and energy distribution.

App operating

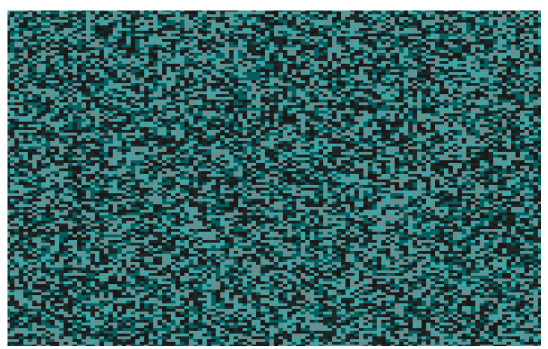


Image 2. Monte Carlo initialization – 130x130 field with 5 grains.



Image 3. Monte Carlo 130x130 field with 5 grains after 60 iterations.

Full example of all program options

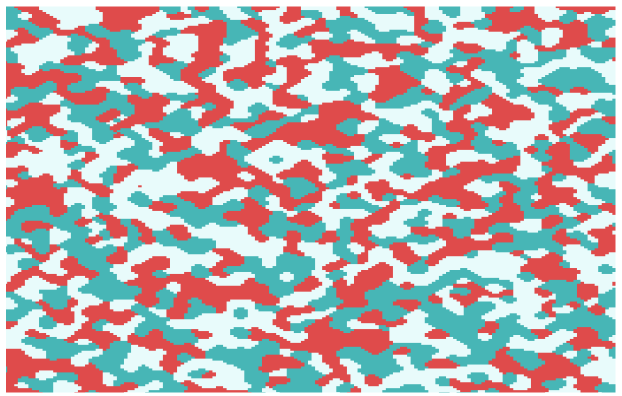


Image 4. Generate microstructure



Image 5. Grain selection

After structure generation (MC or CA) user can choose parameters of new substructures generation. The dual phase structure can be generated using CA or MC method.

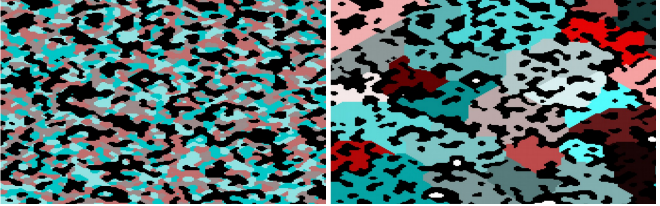


Image 6. Dualphase Monte Carlo

Image 7. Dualphase cellular automaton

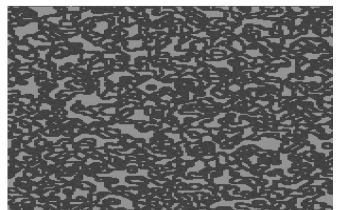


Image 8. Heterogenous energy distribution after Monte Carlo

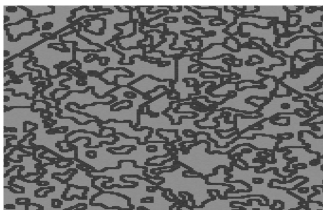


Image 9. Heterogenous energy distribution after cellular automation

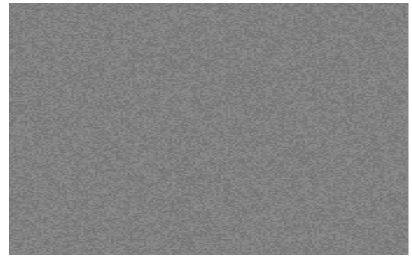
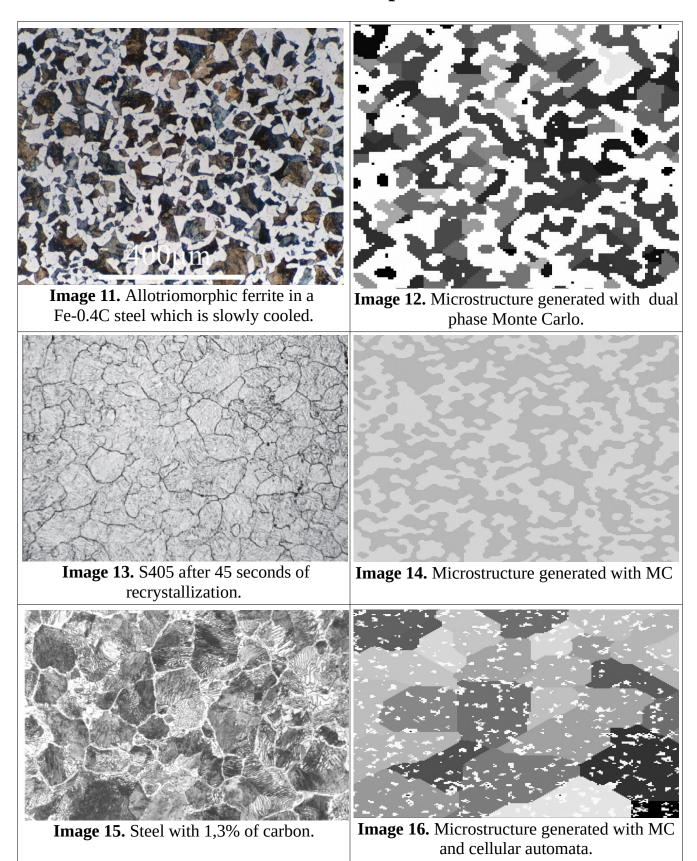


Image 10. Homogenous energy distribution

Real example



Conclusions

Simulations give results which are similar to real microstructures. Application can be used to solve real problems, is expandable if new functionalities will be needed, Monte Carlo method is slower than CA but its possible to imporving some parts of algorithm. Monte Carlo method is working completely random. The most important thing is amount of iterations and good random number generator. It is hard to generate big grains using that method.