# Allen-Cahn Phase Field Model

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This document explains the Allen-Cahn phase-field model and demonstrates its application in both single crystal growth and multi-grain growth through Python codes.

## 1 Single Crystal Growth Model

#### 1.1 Model Formulation

The Allen-Cahn equation stems from minimizing the following free energy functional (1):

$$F[\phi] = \int \left(\frac{\epsilon^2}{2} |\nabla \phi|^2 + \frac{1}{4} (\phi^2 - 1)^2\right) d\mathbf{x} \tag{1}$$

- The first term penalizes sharp interfaces (gradient energy).
- The second term is a double-well potential, with minimum values at  $\phi = \pm 1$ .

The time evolution is given by:

$$\frac{\delta\phi}{\delta t} = -M \frac{\delta F}{\delta \phi} = M \left[ \epsilon^2 \Delta \phi - (\phi^3 - \phi) \right]$$
 (2)

where M is the mobility,  $\epsilon$  is the gradient coefficient, and  $\Delta$  is the Laplacian operator.

#### 1.2 Python Implementation

```
1 import numpy as np
2 import matplotlib.pyplot as plt
3 # Simulation parameters
_{4} N = 128
                                 # The number of grid points
5 dx = 1.0
                                 # Size between grid points
6 dt = 0.01
                                 # Time step
7 \text{ nsteps} = 2500
                                 # The number of steps
8 grad_coeff = 0.1
                                 # Gradient coefficient
9 mobility = 5.0
                                 # Mobility
10 radius = 5
                                 # Seed radius
interval = 100
                                 # Plot update interval
```

```
12
# PBC options (Periodic Boundary Conditions)
14 pbc_x = True
15 pbc_y = True
17 # Initialization
18 phi = np.zeros((N, N))
19 x = np.arange(N)
y = np.arange(N)
X, Y = np.meshgrid(x, y)
22 center = N // 2
23 mask = (X - center)**2 + (Y - center)**2 <= radius**2
24 \text{ phi}[mask] = 1.0
26 def laplacian(phi):
      # X direction calculation using periodic boundary conditions if
27
       enabled
       if pbc_x:
28
          phi_xp = np.roll(phi, -1, axis=0)
29
          phi_xm = np.roll(phi, 1, axis=0)
30
31
          phi_xp = np.zeros_like(phi)
32
          phi_xm = np.zeros_like(phi)
33
           phi_xp[:-1, :] = phi[1:, :]
34
           phi_xm[1:, :] = phi[:-1, :]
35
      # Y direction calculation using periodic boundary conditions if
36
       enabled
37
      if pbc_y:
          phi_yp = np.roll(phi, -1, axis=1)
38
          phi_ym = np.roll(phi, 1, axis=1)
39
40
          phi_yp = np.zeros_like(phi)
41
          phi_ym = np.zeros_like(phi)
42
           phi_yp[:, :-1] = phi[:, 1:]
43
          phi_ym[:, 1:] = phi[:, :-1]
44
45
      return (phi_xp + phi_xm + phi_yp + phi_ym - 4 * phi)/(dx * dx)
46
47 # Plotting settings (pcolormesh, colorbar range [-1, 1])
48 plt.ion() # Turn on interactive mode
49 fig, ax = plt.subplots()
c = ax.pcolormesh(phi, cmap='RdBu', vmin=-1, vmax=1)
plt.colorbar(c, ax=ax)
for step in range(nsteps+1):
      lap = laplacian(phi)
54
      phi += dt * mobility * (grad_coeff * lap - (phi**3 - phi))
55
      # Update the plot every {interval} steps
56
57
      if step % interval == 0:
          ax.clear()
58
           c = ax.pcolormesh(phi, cmap='RdBu', vmin=-1, vmax=1)
          plt.title(f"Time: {step} steps")
60
          plt.pause(0.01)
61
63 plt.ioff()
64 plt.show()
```

Listing 1: Single Grain Growth Code

## 2 Multi-Grain Growth Model

#### 2.1 Model Formulation

For multi-grain growth, the order parameter is vector-valued.

$$\vec{\phi} = (\phi_1, \dots, \phi_{N_{\mathbf{g}}}) \tag{3}$$

where each  $\phi_i$  represents the *i*-th grain and  $N_{\rm g}$  is the number of grains. The free energy functional is extended:

$$F[\vec{\phi}] = \sum_{i=1}^{N_g} \left[ \int \left( \frac{\epsilon^2}{2} |\nabla \phi_i|^2 + \frac{1}{4} (\phi_i^2 - 1)^2 \right) d\mathbf{x} + \int \left( \lambda \sum_{i < j} \phi_i^2 \phi_j^2 \right) d\mathbf{x} \right]$$
(4)

where  $\epsilon$  is the gradient coefficient and  $\lambda$  is the penalty coefficient. The Allen-Cahn equations for each grain become:

$$\frac{\partial \phi_i}{\partial t} = M \left[ \epsilon^2 \Delta \phi_i - (\phi_i^3 - \phi_i) - 2\lambda \phi_i \sum_{j \neq i} \phi_j^2 \right]$$

• The penalty term  $(\lambda \sum_{i < j} \phi_i^2 \phi_j^2)$  increases the energy when different grains overlap, thereby reducing field overlap.

### 2.2 Python Implementation

```
import numpy as np
  import matplotlib.pyplot as plt
_{4} # Simulation parameters
5 N = 128
                               # The number of grid points
6 num_grains = 5
                               # The number of grains
7 dx = 1.0
                               # Size between grid points
8 dt = 0.01
                               # Time step
9 nsteps = 2500
                               # The number of steps
10 grad_coeff = 0.1
11 p_coeff = 2.0
                               # Gradient coefficient
                               # Penalty term coefficient
mobility = 5.0
                               # Mobility
13 radius = 5
                               # Seed radius
14 interval = 100
                               # Update interval for plotting
# PBC options (Periodic Boundary Conditions)
17 pbc_x = True
18 pbc_y = True
# phi (N x N x num_grains)
phi = np.zeros((N, N, num_grains))
22 X, Y = np.meshgrid(np.arange(N), np.arange(N))
# Initialization: set the initial seed for each grain
for i in range(num_grains):
x0 = np.random.randint(0, N)
```

```
y0 = np.random.randint(0, N)
27
      mask = (X - x0)**2 + (Y - y0)**2 \le radius**2
28
      phi[mask, i] = 1.0
29
30
31 def laplacian(phi_2d):
      # X direction calculation using periodic boundary conditions if
32
       enabled
      if pbc_x:
33
          phi_xp = np.roll(phi_2d, -1, axis=0)
35
          phi_xm = np.roll(phi_2d, 1, axis=0)
36
37
          phi_xp = np.zeros_like(phi_2d)
          phi_xm = np.zeros_like(phi_2d)
38
          phi_xp[:-1, :] = phi_2d[1:, :]
39
          phi_xm[1:, :] = phi_2d[:-1, :]
40
      # Y direction calculation using periodic boundary conditions if
41
       enabled
      if pbc_y:
42
          phi_yp = np.roll(phi_2d, -1, axis=1)
43
          phi_ym = np.roll(phi_2d, 1, axis=1)
44
45
          phi_yp = np.zeros_like(phi_2d)
46
          phi_ym = np.zeros_like(phi_2d)
47
48
          phi_yp[:, :-1] = phi_2d[:, 1:]
          phi_ym[:, 1:] = phi_2d[:, :-1]
49
      return (phi_xp + phi_xm + phi_yp + phi_ym - 4 * phi_2d) / (dx *
50
       dx)
51
# Plotting settings (pcolormesh, colorbar range [-1, 1])
# (2x3 subplots: 5 grains plots + combined view)
fig, axs = plt.subplots(2, 3, figsize=(15, 8))
56 cmap_combined = plt.get_cmap('tab10', num_grains) # Grain coloring
57
58 for i in range(num_grains):
59
      ax = axs.flatten()[i]
      c = ax.pcolormesh(phi[:, :, i], cmap='RdBu', vmin=-1, vmax=1)
60
61
      ax.set_title(f'Grain {i+1}')
      plt.colorbar(c, ax=ax)
62
63
64 # Initializing combined view
ax_{combined} = axs[1, 2]
cmap = plt.get_cmap('tab10', num_grains)
grain_colors = [np.array(cmap(i)[:3]) for i in range(num_grains)]
68 combined_rgb = np.ones((N, N, 3)) # Initialize with white
      background (RGB 3-channel, float)
69 for i in range(num_grains):
      combined_rgb += np.expand_dims(phi[:, :, i], axis=2) *
      grain_colors[i]
im_combined = ax_combined.imshow(combined_rgb,
                                    interpolation='nearest',
                                    extent=[0, N, 0, N])
74 ax_combined.set_title('Combined Grains')
75
for step in range(nsteps):
    lap = np.zeros_like(phi)
for i in range(num_grains):
```

```
lap[:, :, i] = laplacian(phi[:, :, i])
79
80
       # Update each grain
81
       total_phi_sq = np.sum(phi**2, axis=2)
82
       for i in range(num_grains):
83
            penalty = 2 * p_coeff * phi[:, :, i] * (total_phi_sq - phi
84
        [:, :, i]**2)
       phi[:, :, i] += dt * mobility * (grad_coeff * lap[:, :, i]
- (phi[:, :, i]**3 - phi[:, :, i]) - penalty)
85
86
       # Update every {interval}
87
        if step % interval == 0:
88
            # Update each grain plot
89
90
            for i in range(num_grains):
                ax = axs.flatten()[i]
91
                ax.clear()
92
                c = ax.pcolormesh(phi[:, :, i], cmap='RdBu', vmin=-1,
93
       vmax=1)
94
                ax.set_title(f'Grain {i+1} (Step {step})')
95
            # Update combined view
96
            #(Combine each grain's field weighted by its respective
97
       color
98
            combined_rgb = np.ones((N, N, 3))
            for i in range(num_grains):
99
                combined_rgb -= np.expand_dims(phi[:, :, i], axis=2) *
100
       grain_colors[i]
            combined_rgb = np.clip(combined_rgb, 0, 1)
101
102
            im_combined.set_data(combined_rgb)
104
            ax_combined.set_title(f'Combined Grains (Step {step})')
            plt.pause(0.01)
105
106
plt.ioff()
108 plt.show()
```

Listing 2: Multi Grain Growth Code