Debugging time evolution

Lucas Myers

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1 Checking periodically perturbed free energy

To check that the configuration force matches with the energy, we check the energy for a uniform configuration first, and then for a small perturbation. The latter check gives an estimate for the configuration force by the functional Taylor series expansion:

$$F(Q + \delta Q) = \int_{\Omega} \left(f(Q) + \left. \frac{\delta f}{\delta Q} \right|_{Q} : \delta Q + \mathcal{O}\left(|\delta Q|^{2} \right) \right) dV \tag{1}$$

with

$$\frac{\delta F}{\delta Q} = \int_{\Omega} \left(\frac{\partial f}{\partial Q} - \nabla \cdot \frac{\partial f}{\partial \left(\nabla Q \right)} \right) dV \tag{2}$$

Here Q is given by:

$$Q = S_0 \begin{bmatrix} \frac{2}{3} & 0 & 0\\ 0 & -\frac{1}{3} & 0\\ 0 & 0 & -\frac{1}{3} \end{bmatrix}$$
 (3)

with

$$\delta Q = S_0 \begin{bmatrix} 0 & \epsilon \sin kx & 0\\ \epsilon \sin kx & 0 & 0\\ 0 & 0 & 0 \end{bmatrix}$$
 (4)

This corresponds (to first order) to a perturbation of the director so that the total configuration $(Q + \delta Q)$ is a uniform-S configuration with director $\mathbf{n} = (1, \epsilon \sin kx, 0)$.

Here, the nondimensional free energy is given by:

$$F = \int_{\Omega} \left(-\frac{1}{2}\alpha Q : Q + \left(\log 4\pi - \log Z + \Lambda : \left(Q + \frac{1}{3}I \right) \right) + \frac{1}{2} \left| \nabla Q \right|^2 \right) dV \tag{5}$$

Then we can calculate F(Q) partially analytically and partially numerically. Of course the elastic part is zero, and we calculate:

$$-\frac{1}{2}\alpha Q: Q = -S_0^2 \frac{\alpha}{3} = -1.2153600266666669$$
 (6)

with $S_0=0.6751$ and $\alpha=8.0$. Further, we can calculate numerically (see calc_lambda program with generate_periodic_Q_tensors.py) Λ and Z for $S_0=0.6751$:

Z = 3.87017170996747

$$\Lambda = \begin{bmatrix}
3.6005952163635766 & 0 & 0 \\
0 & -1.8002976081817883 & 0 \\
0 & 0 & -1.8002976081817883
\end{bmatrix}$$
(7)

Then we may calculate F by multiplying this energy density by the size of the domain. The uniform configuration check is then just to compare that number with the energy output from the simulation for a uniform configuration with the same S_0 and α values and domain size. Calculating the energy density from the mean-field interaction gives:

$$f_{\text{entropy}} = 3.60848720197831 \tag{8}$$

Running the simulation and calculating the free energy over the domain gives:

$$F_{\text{mean field}} = -11.99497833$$
 (9)

and calculating out the energy density from the entropy term gives:

$$F_{\text{entropy}} = 35.61419603$$
 (10)

Comparing these values to the energy density multiplied by $(3.1415926 \times 3.1415926)$ (the domain size) we get similar results.