Fluctuation-induced Non-equilibrium Transition in a Liquid-Crystal Metastable System

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Abstract

The research herein studies the Langevin dynamics allowing for an exchange of energy between liquid crystals and the thermal environment. This dynamics leads to fluctuation and dissipation behaviors in the motions of liquid crystals, and therefore drives the system toward non-equilibrium evolitional processes. In particular, for the operations of liquid-crystal metastable systems, the fluctuation could allow an excitation (non-equilibrium) transition against energy barriers to the globally-stable state. Implemented with an actual case of liquid crystal $\pi$ configuration, this work statistically studies the non-equilibrium metastable transitions and shows the dependence of the transition-time on the correlations (of fluctuations).

Keywords: fluctuation, dissipation, metastable, $\pi$ configuration, Langevin equation
I. INTRODUCTION

In continuum models of liquid crystals [1-4], which are based on phenomenological elastic constants and hydrodynamic transport coefficients, the equations of motion for the liquid crystal director are obtained by minimizing the free energy of the system. However, in liquid-crystal metastable systems [5], the energy-minimization process probably causes the liquid crystal state to be in a nearby local minimum of energy (weakly stable state) and prohibits the barrier transition to the observable global minimum state (strongly stable state). This process could result in a deviation between theoretical predictions and observations in the metastable systems. Accordingly, this paper considers the non-equilibrium Langevin dynamics [6-8] which allows an energy exchange between liquid crystals and the thermal environment, and introduces fluctuation $\xi$ (absorbing energy) and dissipation $\propto \gamma$ (emitting energy) behaviors associated with the motions of liquid crystals. In principle, this is related to the fundamental fluctuation-dissipation relation $\langle \xi(t_1)\xi(t_0) \rangle \propto \gamma(t_1 - t_0)$, and exhibits noticeable influences in some liquid crystal systems. For instances, Barbero indicated that thermal fluctuations can lead to a re-normalization of the anchoring energy of nematic liquid crystals [9]. Galatola studied the Freedericksz transitions in nematic liquid crystals by thermal fluctuation approach [10]. Kelly showed the director fluctuations in nematic droplet with spherical harmonics method [11]. Ruhwandl presented the topological defects around a spherical colloidal particle by Monte Carlo simulation [12]. Relberg observed the spatial and temporal correlations of thermal fluctuations in liquid crystal systems [13]. Maclennan indicated a direct observation on the fluctuation of the liquid crystal director in his work [14]. In this work, we mainly introduce the influences of thermal fluctuations by Langevin dynamics to study the complete metastable transitions $V \rightarrow (T, H) \rightarrow H$ in liquid-crystal $\pi$ configurations [15], which still remains indefinite in real three-dimensional systems as we know.

Based on Frank-Oseen elastic theory [16], we consider the spatiotemporal-correlated fluctuations $\xi$ as well as the corresponding dissipations $\propto \gamma$ for the liquid crystal directors in Langevin dynamics, since (I) the common non-correlated fluctuations $\xi_n$ conditioned by $\langle \xi_n(t_1)\xi_n(t_0) \rangle \propto \gamma k_BT \delta(t_1 - t_0)$ will cause a non-physical/ divergent quantity in ultra-small time scale $t_1 - t_0 \approx 0$, while (II) the spatiotemporal-correlated fluctuations $\xi$ offers a more real algorithm to describe the experimental observations [13].
Definitely, we consider the fluctuations $\xi$ related to an exponential correlation behavior 

$$\langle \xi(r_1, t_1)\xi(r_0, t_0) \rangle \propto e^{-\frac{|r_1-r_0|}{\tau}}e^{-\frac{|t_1-t_0|}{\lambda_{z}}}$$

in the 1+1 (spatial+temporal) dimensions via the Ornstein-Uhlenbeck process[18], and thereby characterize the fluctuation behavior by correlation time $\tau$ and length $\lambda$ as experimental observations[13]. Implemented with an actual case of the metastable systems in the liquid-crystal $\pi$ configuration[15], this paper fulfills a statistical numerical analysis over an ensemble of systems with 1000 identical iterations, and elucidates the non-equilibrium metastable transitions.

This article is organized as follows. Section 2 reviews Frank-Oseen elastic free energy using Q tensor representation[16], and structures the fundamental equation of motion for liquid crystal directors. Section 3 further introduces the Ornstein-Uhlenbeck process[18] for the Langevin dynamics to consider an actual spatiotemporal-correlated fluctuations $\xi(r, t)$. An update formula of directors for the numerical calculation is then identified. Section 4 numerically analyzes the influences of fluctuations and dissipations in the liquid crystal $\pi$ configuration, and depicts the associated non-equilibrium metastable transitions, i.e., about how a initial $V$ state transit through the locally-stable $T$ and finally into the globally-stable $H$ state in the real three-dimensional systems. The main conclusions are then discussed in section 5. Two more experiments as well as the corresponding theoretical works are appendixed and show the influences of fluctuations and dissipations even in usual cells of Liquid-Crystal Displays (LCDs).

II. MODELING LIQUID CRYSTAL DIRECTOR

In elastic continuum theory, the local aligned sample may be regarded as a single liquid crystal, in which the molecules are on the average aligned along the direction defined by a director $\vec{n}$. The Frank-Oseen strain free energy density using tensor representation for liquid crystal directors can then be described by[16]:

$$f_s = \frac{1}{12}(K_{33} - K_{11} + 3K_{22})\frac{G^{(2)}}{S^2} + \frac{1}{2}(K_{11} - K_{22})\frac{G^{(2)}_2}{S^2} + \frac{1}{4}(K_{33} - K_{11})\frac{G^{(3)}_6}{S^3} + q_0K_{22}\frac{G^{(2)}_4}{S^2},$$

(1)
with

\[ G^{(2)}_1 = Q_{jk,l} Q_{jk,l} \]
\[ G^{(2)}_2 = Q_{jk,k} Q_{jl,l} \]
\[ G^{(3)}_6 = Q_{jk} Q_{lm,j} Q_{lm,k} \]

in which

\[ Q_{jk} = S \left( n_j n_k - \frac{1}{3} \delta_{jk} \right), \]
\[ Q_{jk,l} = \frac{\partial Q_{jk}}{\partial l}, \quad j, k, l \in \{x, y, z\}. \]

Here, \( K_{11}, K_{22}, \) and \( K_{33} \) are the splay, twist, and bend elastic constants, respectively. \( q_0 = 2\pi/p_0 \) associates the pitch \( p_0 \) of chiral liquid crystals, e.g. Cholesterics. Term Q is the tensor order parameter, and \( \vec{n} \) is the liquid crystal director with unit length. \( S \) is the scalar order parameter, and indicates the ideal isotropic and nematic phases by the values \( S = 0 \) and \( S = 1 \), respectively. \( \varepsilon_{jkl} \) is the Levi-Civita symbol. \( \delta_{jk} \) represent the Kronecker delta. Moreover, the convention of summing over repeated indices is used herein. The electric free energy density can be expressed in terms of the Q tensor as

\[ f_e = \frac{1}{2} \epsilon_0 \varepsilon V_j V_j + \frac{1}{2} \epsilon_0 \Delta \epsilon V_k V_k \frac{Q_{jk}}{S}, \]

with

\[ \tau = \frac{2\epsilon_\perp + \epsilon_\parallel}{3}, \quad \Delta \epsilon = \epsilon_\parallel - \epsilon_\perp, \]
\[ V_j = \frac{\partial V}{\partial j}, \quad j \in \{x, y, z\}. \]

Here, \( \epsilon \) is the LC dielectric tensor with elements \( \epsilon_\perp \) and \( \epsilon_\parallel \) for dielectric coefficients perpendicular and parallel to the LC director.

To solve equilibrium states of liquid crystal directors with applied potentials, the system free energy, which is obtained by integrating the Gibbs free energy density \( f_g = f_s - f_e \) over the volume, is minimized. Accordingly, the Euler-Lagrange equations for electric potentials and director components under condition \( |\vec{n}| = 1 \) can thus be obtained

\[ 0 = -[f_g]_{V} = \partial_i \left[ (\varepsilon_\perp \delta_{ij} + (\varepsilon_\parallel - \varepsilon_\perp) n_i n_j) \partial_j V \right], \]
\[ 0 = -[f_g]_{n_i} + \lambda n_i \quad i \in \{x, y, z\}, \]
Here, $\lambda$ is the Lagrange multiplier for constraint $|\vec{n}| = 1$. By applying the chain rule $[f_g]_{ni} = [f_g]_{Q_{jk}} \frac{\partial Q_{jk}}{\partial n_i} = [f_g]_{Q_{jk}} \cdot S (n_j \delta_{ki} + n_k \delta_{ji})$, we obtain the alternative definitions in tensor representation for Equation (10)

\[
[f_g]_{nx} = 2n_x S [f_g]_{Q_{xx}} + n_y S ([f_g]_{Q_{xy}} + [f_g]_{Q_{yx}}) + n_z S ([f_g]_{Q_{xz}} + [f_g]_{Q_{zx}}),
\]

(11)

\[
[f_g]_{ny} = 2n_y S [f_g]_{Q_{yy}} + n_x S ([f_g]_{Q_{xy}} + [f_g]_{Q_{yx}}) + n_z S ([f_g]_{Q_{yz}} + [f_g]_{Q_{zy}}),
\]

(12)

\[
[f_g]_{nz} = 2n_z S [f_g]_{Q_{zz}} + n_x S ([f_g]_{Q_{xz}} + [f_g]_{Q_{zx}}) + n_y S ([f_g]_{Q_{yz}} + [f_g]_{Q_{zy}})
\]

(13)

Solving Equations (9) and (10) simultaneously, the potential distribution and the director configuration for equilibrium status can now be evaluated. It is noted that, in general, $[f_g]_{Q_{jk}}$ is not equal to $[f_g]_{Q_{kj}}$ for cases of $j \neq k$. For the analyses of dynamic time evolutions on the liquid-crystal directors, however, we consider the time-dependent/non-equilibrium Langevin equation. In the Langevin dynamics which includes the stochastic torque (fluctuation) $\sigma \zeta$ as well as the dissipation torque $\kappa$ for directors, the general rotational Langevin equation under a external torque $F$ can be read as [17]:

\[
\kappa \frac{\partial}{\partial t} n_i = -F_i + \sigma \zeta,
\]

(14)

Here the constant $\sigma$ is related to $\kappa$ by Einstein’s relation $\sigma = \sqrt{2\kappa k_B T}$, and $\zeta$ is the normalized white noise with dimension $s^{-1/2}$. For the liquid-crystal system, to simply evaluate the dissipative torque for the liquid crystal director, the volume of the liquid crystal director is treated as an effective sphere with diameter $d$, and thereby the director will suffer a dissipative torque $\kappa = \pi \gamma d^3$ by Stokes relation. An arranged Langevin equation corresponding to Equation (10) is then obtained:

\[
\gamma \frac{\partial}{\partial t} n_i = -[f_g]_{ni} + \frac{\sigma \zeta}{\pi d^3} \\
\equiv -[f_g]_{ni} + \xi,
\]

(15)

Details for this part will be discussed in the next section. Moreover, the equation of motion for common use [16] can be estimated by considering the statisticly steady director $\langle n_i \rangle$, i.e.
ignore the (temperature-dependent) thermal fluctuations $\xi$

$$\gamma \frac{\partial}{\partial t} \langle n_i \rangle = -[f_g]_{\langle n_i \rangle},$$

(16)

with an effective order parameter $0 < S < 1$ in $[f_g]_{\langle n_i \rangle}$. Note that the function of the Lagrange multiplier $\lambda$ has been replaced here by re-normalizing the director $|\vec{n}| = 1$ at each time step.

III. SPATIOTEMPORAL CORRELATED FLUCTUATION FUNCTIONS

Based on the above-mentioned Langevin dynamics for liquid-crystal systems as the Equation (15), we now detail the spatiotemporal-correlated stochastic torque (fluctuation) $\xi$ as well as the corresponding dissipations for the liquid-crystal directors. Mainly we consider the Ornstein-Uhlenbeck process in 1+1 dimensions to generate the spatiotemporal-correlated stochastic torque $\xi(z, t)$ with:

$$\langle \xi(z, t) \rangle = 0$$

$$\langle \xi(z, t)\xi(z_0, t_0) \rangle = \frac{\gamma k_B T}{2\pi d^2 \lambda_z} e^{-[|z-z_0|/\lambda_z]} e^{-[|t-t_0|/\tau]}$$

(18)

Here, $\lambda_z$ and $\tau$ are the characteristic correlation lengths in the spatial ($z$) and temporal ($t$) dimensions, respectively. $k_B$ is the Boltzmann constant with a value of $1.38 \times 10^{-23}$ NmK$^{-1}$, $d$ is the effective size of the liquid crystal directors, and $T$ is the temperature of thermal environment. The Equation (18) can be further understood by its power spectrum $P(z_0, \omega) = \frac{\gamma k_B T}{\pi d^2 \lambda_z} \frac{1}{1+\tau^2\omega^2}$ which results from the temporal Fourier transform of the correlation function $\langle \xi(z_0, t)\xi(z_0, 0) \rangle$. It is indicated that a longer correlation time $\tau$ suppresses the higher-frequency (higher-energy) energy exchanges and leads to a smaller fluctuation amplitude $[\frac{\gamma k_B T}{2\pi d^2 \lambda_z} \frac{1}{\tau}]^{\frac{1}{2}}$. Likewise for the spatial Fourier transform of $\langle \xi(z, t_0)\xi(0, t_0) \rangle$, i.e. $P(k, t_0) = \frac{\gamma k_B T}{\pi d^2 \tau} \frac{1}{1+\lambda_z^2 k^2}$, a longer correlation length $\lambda_z$ implies the smaller-momentum (longer-wavelength) energy exchanges, and causes a longer-ranged but smaller fluctuation strength $[\frac{\gamma k_B T}{2\pi d^2 \tau} \frac{1}{\lambda_z}]^{\frac{1}{2}}$.

To generate this real-space fluctuation function $\xi(z, t)$ to evaluate the motion of liquid crystals in Equation (15), an exact recursion relation with temporal spacing $\Delta_t$ and spatial spacing $\Delta_z$ in the finite difference method is applied:
\[ \xi(z, t) = -\exp\left(-\frac{\Delta z}{\lambda_z} - \frac{\Delta t}{\tau}\right) \xi(z - \Delta z, t - \Delta t) \]
\[ + \exp\left(-\frac{\Delta z}{\lambda_z}\right) \xi(z - \Delta z, t) \]
\[ + \exp\left(-\frac{\Delta t}{\tau}\right) \xi(z, t - \Delta t) \]
\[ + \left[1 - \exp\left(-\frac{2\Delta z}{\lambda_z}\right)\right]^{1/2} \]
\[ \times \left[1 - \exp\left(-\frac{2\Delta t}{\tau}\right)\right]^{1/2} A\eta(z, t) \] (19)

Here, \( \eta(z, t) \) is spatiotemporal Gaussian white noise with \( \langle \eta(z, t)\eta(z', t') \rangle = \delta_{z,z'}\delta_{t,t'} \), and \( A \) is the fluctuation amplitude \( \left[\frac{k_BT}{2\pi d\tau\lambda_z}\right]^\frac{1}{2} \). Due to the exponentially decaying correlation for spatiotemporal parameters and the characteristic of white noise function \( \langle \eta(z, t) \rangle = 0 \) in Equation (19), \( \xi(z, t) \) is independent of its initial value and evidently meets the condition in Equation (17). For the condition of fluctuation correlations \( \langle \xi(z, t)\xi(z_0, t_0) \rangle \) in Equation (18), numerical statistics averaged over \( 10^3 \) iterations (each iteration includes the data number \( N = 100 \) in spatial/temporal space) are fulfilled to verify the validity of Equation (19). The numerical results by Equation (19) are shown to compare well with the analytical results of Equation (18) in Figures 1(b) and 2(b) for temporal and spatial arguments, respectively. Figure 1(a) shows \( \xi(z_0, t) \) of real temporal space in one of the iterations and characterizes a \( \tau \)-period fluctuation function. Figure 2(a) then presents \( \xi(z, t_0) \) of real spatial space with coherent length \( \lambda_z \).

Implemented with Equation (19) for the real-space fluctuation function \( \xi(z, t) \), an updated formula by discretizing Equation (15) can be obtained:

\[ n_i(z, t + \Delta t) - n_i(z, t) = \frac{\Delta t}{\gamma} \left(-[f_g]_{n_i} + \xi(z, t)\right), \quad i \in \{x, y, z\} \] (20)

for the equation of directors in 1+1 dimension systems. This algorithm includes the influences of thermal baths and supplies a theoretical foundation for non-equilibrium liquid crystal dynamics.
FIG. 1: Analyses of numerical fluctuation function $\xi$ for fixed spatial state: $z = z_0$ and $\lambda_z = 2\Delta_z$. (a) Fluctuation function in real temporal space for $\tau = \Delta t$ (thin red line) and $\tau = 5\Delta t$ (bold blue line). (b) Temporal correlation function for $\tau = \Delta t$ (symbol □) and $\tau = 5\Delta t$ (symbol ○). Solid lines indicate the corresponding analytical results.

FIG. 2: Analyses of numerical fluctuation function $\xi$ for fixed temporal state: $t = t_0$ and $\tau = 2\Delta t$. (a) Fluctuation function in real spatial space for $\lambda_z = 2\Delta_z$ (thin red line) and $\lambda_z = 8\Delta_z$ (bold blue line). (b) Spatial correlation function for $\lambda_z = 2\Delta_z$ (symbol □) and $\lambda_z = 8\Delta_z$ (symbol ○). Solid lines indicate the corresponding analytical results.

FIG. 3: Liquid crystal configurations and the corresponding Gibbs free energy in $\pi$ configuration: bend ($V$), twist ($T$), and homogenous ($H$) equilibrium states.
IV. NUMERICAL ANALYSES OF NON-EQUILIBRIUM DYNAMICS

To concretely understand the influence of the thermal fluctuations on the liquid-crystal metastable systems, we now consider an actual case with $\pi$ configuration $[15]$. As shown in Figure 3, with the Frank-Oseen elastic theory, bend ($V$), twist ($T$), and homogenous ($H$) states form a metastable status. A transition barrier exists between the $T$ and $H$ states, leading to a local minimum localization on the $T$ state for the time evolution from the $V$ state. (Here the red, green, and blue arrows indicate the spatial $\hat{x}$, $\hat{y}$, and $\hat{z}$ directions, respectively.) In this work, we mainly explain how the initial $V$ state can transit through the locally-stable $T$ state, and finally into the globally-stable $H$ in the long run as experimental observations. To statistically evaluate the influences of the fluctuation and dissipation, we consider an ensemble ($10^3$ iterations) of liquid crystal cells for each set-up conditions of parameters. The associated parameters are defined as: $\gamma = 50mPa \cdot sec$, $K_{11} = 13.6pN$, $K_{22} = 10.0pN$, $K_{33} = 14.7pN$, $\varepsilon_\parallel = 10$, $\varepsilon_\perp = 5$, and $T = 300K$. The thickness of the cell is set as $4.5 \mu m$. No chiral agent is added. The pre-tilt angle between the liquid crystal directors and the alignment (green) layers is set to $5^\circ$. For the finite difference algorithm, the spatial grid is $\Delta z = 0.15\mu m$ and the temporal grid is $\Delta t = 0.01ms$. Besides, the effective director size $d$ is treated as about $10^1 - 10^2 nm$ for this study, which is $1 - 2$ order larger than the liquid crystal molecules. For each iteration of the ensemble, the $V$ state is initialized by applying a bias $15V$, which lasts for a long enough period ($200ms$ in our cases), and the voltage is turned off at $t \geq 0$ in order to observe the evolution of the liquid crystals. Figure 4(a) illustrates the transition rate $\Gamma_H = 1/\langle t_{V\rightarrow H} \rangle$ versus fluctuation amplitude $A$, where $\langle t_{V\rightarrow H} \rangle$ represents the average time of $V \rightarrow H$ transitions of the ensemble. In the Figure 4(a), in the absence of the thermal bath (at $A = 0Pa$), the process of minimizing-energy as the Equation (16) drives the dynamics of the liquid crystals, and consequently the initial $V$ states in all of the 1000 iterations are found to follow the free-energy-decreasing trace into the locally-stable $T$ state as the blue solid-line segment form $V$ to $T$ in the Figure 4(b). No iteration of transition from $V$ to $H$ is found in our simulation in this case. This results in the infinite transition time $\langle t_{V\rightarrow H} \rangle = \infty$ for $V$ to $H$ state, or the zero transition rate $\Gamma_H = 1/\langle t_{V\rightarrow H} \rangle = 0$ as in Figure 4(a). (Note that Figure 4(a) shows the log-scale plot for clear appearance, and the infinite time is realized by a cur-off time $10^5ms$ in real numerical calculations.) As the fluctuation amplitude $A$ increases ($A \neq 0Pa$), the
liquid crystals contain not only the elastic free energy but also the finite fluctuation energy; Or alternatively, the liquid crystals suffer an extra stochastic torque $\xi$ in addition to the fundamental elastic torques. This allows the initial $V$ state in some/few iterations of the 1000 iterations have the possibility to deviate from the free-energy-decreasing trace, and are found to proceed another non-equilibrium path (as the green dashed curve in Figure 4(b)) to reach the final $H$ states, although the paths of the most iterations still remain on the free-energy-decreasing trace (red dashed-dotted curve in Figure 4(b)) to be localized at $T$ states in our calculations. This causes the transition time $\langle t_{V\rightarrow H} \rangle$ of some of the 1000 iterations to be finite and thereby the finite statistic transition rate $\Gamma_H = 1/\langle t_{V\rightarrow H} \rangle$ as in Figure 4(a). Consequently, it is found that larger fluctuation amplitude $A$ can cause more iterations of the ensemble to proceed the non-equilibrium path ($V \rightarrow H$; green dashed curve in Figure 4(b)) and raise the transition rate $\Gamma_H = 1/\langle t_{V\rightarrow H} \rangle$ as in Figure 4(a). Here, Figure 4(b) illustrates quantitatively the energy evolution traces of the transitions $V \rightarrow T$ and $V \rightarrow H$ for $A = 100 \text{ Pa}$, $\tau = \Delta_t$, and $\lambda_z = \Delta_z$. For the large fluctuation amplitude ($A \rightarrow \infty$ Pa), however, the overwhelming fluctuation torque leads to strong-fluctuated liquid crystal orientations, and thereby the liquid crystal configurations ($V$, $T$, and $H$) cannot be well identified. This is far away from phenomenological observations and is beyond the scope of this work. In addition to these analyses of the transition dynamics on the aspect of time and energy as in Figures 4(a) and 4(b), respectively, We further analyze the corresponding time evolutions of the directors to help to understand the transition dynamics.

Figure 5 purposes to briefly characterize the liquid-crystal profiles of the above-mentioned transitions first, and the detailed time-evolution traces of the corresponding liquid-crystal directors are then depicted in Figure 6. The characteristic director profiles of the transition $V \rightarrow T$, which corresponds to the red dashed-dotted curve on Figure 4(b), is characterized on Figure 5(a). The characteristic director profiles of the transition $V \rightarrow H$, which corresponds to the green dashed curve on Figure 4(b), are found to have two possible time-evolution traces in our simulations and are characterized by $S_0$ and $S_1$ in Figures 5(b) and 5(c), respectively. In Figure 6(a) (corresponding to Figure 5(a)) the orientations of the individual liquid-crystal directors draws the bend ($V$) state at $t = 0$ and denoted by $(n_x, n_y)\rightarrow(0, 0)$, i.e. $|n_z| \rightarrow \sqrt{1 - n_x^2 - n_y^2} = 1$. By the time going, these directors then proceed (along the red dashed-dotted energy trace in Figure 4(b)) to the locally-stable $T$ state, and consequently describe the transition $V \rightarrow T$ over the long run. Here, an alternative anti-clockwise-twist
FIG. 4: (a) Transition rate of $V \rightarrow H$ as a function of fluctuation amplitude $A$ with characteristic correlation scale $\tau = \Delta t$, $\lambda_z = \Delta_z$ (mark □), $\tau = \Delta t$, $\lambda_z = 5\Delta_z$ (mark +), and $\tau = 5\Delta t$, $\lambda_z = \Delta_z$ (mark ♦). In which the horizontal axis corresponds to the $d = 93nm$, 47nm, and 31nm for $A = 50$, 100, and 150Pa, respectively for $\tau = \Delta t$, $\lambda_z = \Delta_z$, and $T = 300K$. And for the case of $\tau = \Delta t$, $\lambda_z = 5\Delta_z$ and $\tau = 5\Delta t$, $\lambda_z = \Delta_z$, $A = 50$, 100, and 150Pa corresponds to $d = 41nm$, 20nm, and 14nm, respectively. (b) Energy evolution traces of alternative non-equilibrium transitions for $V \rightarrow T$ (dashed-dotted curve) and $V \rightarrow H$ (dashed curve) for $A = 100$ Pa ($T = 300$ K with $d = 50$ nm), $\tau = \Delta t$, and $\lambda_z = \Delta_z$.

profiles is analogous to the presented clockwise-twist one, and is ignored in the context. Similar anti-clockwise/clockwise configurations occurred on the transition $V \rightarrow H$ is ignored below as well. Figures 6(b) and 6(c) numerically show two possible time-evolution traces of directors $(n_x, n_y)$ corresponding to the transitions $V \rightarrow S_0 \rightarrow H$ and $V \rightarrow S_1 \rightarrow H$, respectively. Note these two transitions ($V \rightarrow S_0 \rightarrow H$ and $V \rightarrow S_1 \rightarrow H$) show an indistinguishable energy trace as the green dashed curve in Figure 4(b) in our simulations. All these construct the fundamental understandings to help to study the complete $V \rightarrow H$ transition in real three-dimensional liquid-crystal π systems as below.

In the following section, we numerically investigate an extension study of the three-dimensional liquid-crystal metastable system to give a result corresponding to the actual experimental observations [15]. It is emphasized that the spatial/temporal grid size ($\Delta_z$ and $\Delta_t$) is herein assumed to be much larger than the coherent length/time ($\lambda_z$ and $\tau$), such that the fluctuations for each time step at the individual site are almost independent, i.e. $\xi(z,t) = A \eta(z,t)$ in Equation (19). This thereby allows a straightforward extension to the three-dimensional condition $\xi(x,y,z,t) = A' \eta(x,y,z,t)$ for the studied system. The
FIG. 5: Characteristics of liquid crystal configurations of alternative non-equilibrium transitions for (a) $V \rightarrow T$ process, (b) $V \rightarrow S_0 \rightarrow H$ process, (c) $V \rightarrow S_1 \rightarrow H$ process viewed from $+x$ (left) and $-y$ (right).

FIG. 6: Evolution traces of liquid crystal director ($n_x, n_y$) corresponding to (a) $V \rightarrow T$ process, (b) $V \rightarrow S_0 \rightarrow H$ process, (c) $V \rightarrow S_1 \rightarrow H$ process. In which the directors at $z = d/9$ (mark $\bullet$), $z = 2d/9$ (mark $+$), $z = 3d/9$ (mark $\times$), $z = 4d/9$ (mark $\circ$), $z = 5d/9$ (mark $\square$), $z = 6d/9$ (mark $\Diamond$), $z = 7d/9$ (mark $\triangleleft$), $z = 8d/9$ (mark $\triangleright$) are shown.

relevant parameters of the $\pi$ configuration are similar to the above-mentioned, except for $\Delta_z = 0.45 \mu m$ as well as the considerations of optical transmittances related to practical observations. The associated optical parameters are set as: the incident wavelength is 0.55 $\mu m$, ordinary refractive index of liquid crystals is $n_o = 1.5$, extraordinary refractive index of liquid crystals is $n_e = 1.58$, and the absorption axes of the upper and lower polarizers are located at $\phi = 45^\circ$, and $135^\circ$ in the $x - y$ plane, respectively. The numerical results for the metastable transition $V \rightarrow T, H$ without ($A' = 0$) and with ($A' = 100$) thermal couplings are shown in Figures 7 and 8, respectively. In the case of $A' = 0$ in Figure 7, the initial $V$ state follows the energy-minimizing process in the elastic continuum theory and leads to the localization at the $T$ state as described above. However, the observable $H$ state exhibits
FIG. 7: Optical transmittances of (a) the initial $V$ state at $t = 0\ ms$, (b) the stable $T$ state at $t = 1000\ ms$, and (c) the forbidden $H$ state for the $V \rightarrow T, H$ transition without thermal coupling $A' = 0$. The corresponding liquid crystal directors (LC profiles) at $z = d/2$ are showed as below plots (d), (e), and (f), respectively.

significant differences of liquid crystal configurations to the $T$ state in Figures 7(e) and 7(f) and is forbidden by the energy barrier (Figure 4(b)). In the case of $A' = 100$ in Figure 8, thermal fluctuations as well as the associated correlations offer the possibility of the non-equilibrium $V \rightarrow H$ transitions and cause the $T + H$ transient states in 100 $ms$ after the bias is dropped to zero (Figure 8(b)). Thereafter, the elastic dynamics describing the disclination motions [15] drives a final $H$ stable state as in Figure 8(c). This gives an explanation for the overall $V \rightarrow H$ process. Overall, with thermal fluctuations, a straightforward analysis corresponding to the observed state transitions of $\pi$-configuration in Reference [15] can be realized as in Figure 9.

V. CONCLUSIONS

In this work, based on the elastic continuum theory, we have introduced the Langevin algorithm with the Ornstein-Uhlenbeck process to realize the non-equilibrium dynamics of liquid crystal systems coupled weakly to the thermal environment. In which the Ornstein-
FIG. 8: Optical transmittances of (a) the initial $V$ state at $t = 0$ ms, (b) the transient $T + H$ state at $t = 100$ ms, and (c) the stable $H$ state at $t = 1000$ ms for the $V \rightarrow T, H$ transition with thermal coupling $A' = 100$. The corresponding liquid crystal directors (LC profiles) at $z = d/2$ are showed as below plots (d), (e), and (f), respectively.

FIG. 9: Transition processes of $\pi$-configuration under the bias sequence. In the field-on period, the homogenous (H) state transforms to the stochastic symmetric (Hs) and asymmetric (Ha) states, and further proceed to the bend (V) state under higher biases. In the field-off period, the bend (V) state first relaxes to the twist (T) and homogenous (H) states as described in the context, and recovers to the overall homogenous (H) configuration in the long run.
Uhlenbeck process supplies a connection to the natural spatial and temporal correlation scales of thermal fluctuations. All these allow one to simulate a non-equilibrium time evolution and give physical insights into the processes taking place in the studied systems. It is noted that another thermal hopping (hop over the potential barrier, i.e. a direct transition $T \rightarrow H$ in Figure 4(b)) dynamics can be set up into this system through strong coupling cases, but go beyond this study’s main goal. Definitely, this work provides one possible explanation for the complete transition $V \rightarrow H$ in the $\pi$ configurations and supplies the corresponding real-time simulation results for the experimental observations [15].

Appendix A: Fluctuation-induced Vortex/antivortex in VA cells

In this appendix, we briefly show the fluctuation-induced (stochastic) vortex/antivortex through experimental and theoretical works, and the associated fluctuation amplitude $A'$ can be estimated as well. The used structure of the VA (vertical alignment) liquid crystal cells is depicted in Figure [10]. The associated LC parameters are defined as: $\gamma = 135 mPa \cdot sec$, $K_{11} = 13.6 pN$, $K_{22} = 10.0 pN$, $K_{33} = 14.7 pN$, $\varepsilon_|| = 3.6$, and $\varepsilon_\perp = 7.3$. Here, the fluctuation amplitude $A'$ are decided by fitting the density and period of vortexes/antivortexes with experimental observations, and obtained as $A' = 80 – 125 Pa$. These amplitude value correspond to the fluctuated angle of the director around $5^\circ$ for studied cases in our simulations. The pre-tilt angle between the liquid crystal directors and the alignment (xy) plane is set to $90^\circ$ as the experimental setup. For optical analyses, the incident light wavelength $\lambda$ is set as $0.55 \mu m$, $n_o = 1.5$, $n_e = 1.6$, and the absorption axes of the upper and lower polarizers are located at $\phi = 0^\circ$, and $90^\circ$ in the $x – y$ plane, respectively. Besides, a long enough pre-simulation (lasting 100 ms) is fulfilled to simulate the natural/initial LC status for the sequential numerical simulation. Numerical results as well as experimental observations for optical transmittances are illustrated in Figure [11] by time. It indicated that the thermal fluctuations can introduce stochastic vortex/antivortex as observed in experiments [11](a), while without fluctuations a regular time evolution of LC profiles is predicted as in Figure [11](c). Definitely, these allow a further analysis on the emergence and annihilation behaviors of vortex/antivortex associated the spatial and temporal correlations of fluctuations [9], while beyond the scope of this work.
FIG. 10: Structures of the VA liquid crystal cells with patterned ITO (indium tin oxide) electrodes. The cell volume is set to be \(50\,\text{um} \times 50\,\text{um} \times 4\,\text{um}\) in width\(\times\)length\(\times\)height \((x \times y \times z)\) dimensions.

FIG. 11: Optical transmittances of the VA cell under bias 5V in (a) experimental observations, (b) numerical analyses with fluctuations, and (c) numerical analyses without fluctuations. In which (a) and (b) show the comparable emergence and annihilation behaviors of vortex/antivortex by time. (c) shows the regular time evolutions of LC profiles on transmittances.

**Appendix B: Effects of Fluctuations on LC response time**

In this appendix, not only the fluctuation-induced vortex/antivortex as described at Appendix A, the thermal fluctuations are shown to have appreciable influences on the LC response behaviors (against biases). In the meantime, this influence supplies an alternative
method to quantitatively evaluate the fluctuation amplitude $A'$. The used structure of the MVA (multi-domain vertical alignment) liquid crystal cells is depicted in Figure (12), in which the upper ITO (indium tin oxide) has fine-slit patterns and the lower ITO stacks with protrusion layers for the tilted-alignment purposes. The associated parameters are referred to these at Appendix A; while the fluctuation amplitude $A'$ are decided to be 115 $Pa$ by fitting the response time as below. Numerical results as well as the experimental observations for the optical transmittance at $t=16$ ms and $V=5$ V are shown in Figure 13. Normalized transmittances as a function of time at biases $V=4$, 5, and 6 V are then depicted by numerical simulations with fluctuations in Figure 14(a), numerical simulations without fluctuations Figure 14(b), and experimental observations in Figures 14(a) and (b) for comparison. Compared to the common understandings: larger bias/field causes faster LC response in field-on process as shown in Figure 14(b) (solid curves; without fluctuations), thermal fluctuations as well as experimental observations give another story herein. Besides fast LC response in field-on process, large bias/field cause simultaneously dense vortex spots as shown in Figure 13(a). These observations agree with the numerical results if the fluctuations are included as Figure 13(b). Definitely, the sequential reliefs of the entangled vortexes could cost more time to approach the final stable status, and thereby could reduce the liquid crystal response time. This can be observed in Figure 14(a): compared between these curves with $V=4$ and 5 V, the response time with larger bias (5V) is indeed faster than that with lower bias (4V); however, compared between these curves with $V=5$ and 6V, the response time with larger bias (6V) is slower than that with lower bias (5V). In Figure 14(b), however, the numerical simulations without fluctuations could not help to understand these experimental observations. Substantially, this gives an example to indicate the significance of fluctuations in some real liquid crystal systems.

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FIG. 12: Structures of the MVA liquid crystal cells, in which the upper ITO has fine-slit patterns and the lower ITO stacks with protrusion layers. The cell volume is set to be $160\,\mu m \times 200\,\mu m \times 4\,\mu m$ in width×length×height ($x \times y \times z$) dimensions.

FIG. 13: Optical transmittances of the MVA cell with bias 5 V at t=16 ms by (a) experimental observations, (b) numerical simulations with fluctuations, and (c) numerical simulations without fluctuations. In which (a) and (b) show the stochastic vortex/antivortex spots. (c) shows the regular transmittance pattern.

FIG. 14: Normalized transmittances as a function of time at biases V=4,5,6 V for (a) numerical simulations with fluctuations, and (b) numerical simulations without fluctuations. The experimental results are added to both analyses for comparison.
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