Dynamical Landau-de Gennes theory for electrically-responsive liquid crystal networks

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I. INTRODUCTION

Liquid crystal networks (LCNs), such as nematic elastomers, combine the orientational properties of liquid crystals with the elastic properties of polymer networks by incorporating liquid crystalline mesogens into the cross-linked polymer matrix [1]. The resulting coupling between the strain imposed on the material and the orientation of the liquid crystalline mesogens it contains introduces novel features that cannot entirely be traced back to either of the LCN constituents [2]. Striking examples include large spontaneous deformations upon a temperature-induced phase transition of the liquid crystalline mesogens [3] and so-called “(semi-)soft” elasticity, i.e., macroscopic deformation at (almost) no energetic cost by reorientation of the liquid crystalline mesogens [4].

The theoretical description of these materials was pioneered by Warner and coworkers, who proposed a framework combining classical elasticity theory with the Landau–de Gennes theory for liquid crystals [5,6]. This was later expanded upon to illustrate, among other things, the occurrence of phase transitions and instabilities in LCNs [3,7,8], how these materials respond to electric fields [9], and why they display “soft” deformation modes if alignment, and the corresponding symmetry-breaking, arises spontaneously [10–13]. Conversely, this view of “soft” deformations has been challenged in the case of pre-stretched elastomers, where symmetry arguments no longer apply, and in which case it is argued that the reduced elastic modulus results entirely from nonlinear elasticity [14–19]. Warner and Terentjev provide a comprehensive overview of the most important results in their monograph [20].

A deeper understanding of LCNs has led to the rapid development toward industrial applications of these materials in (soft) actuators [21,22], surgical interventions [23–25] and 3D printing shape-memory liquid crystal elastomers [26]. Most of these exploit the susceptibility of the mesogens to external stimuli such as heat or electric fields [27]. Recently, LCNs were also recognised as a prime candidate for stimuli-responsive coatings due to their tunable dynamical behavior, with envisaged applications in the fields of haptic feedback, self-cleaning surfaces and finely controlled pixel-like deformations (so-called voxels) [28,29]. Desirable properties for these applications are a swift response and large susceptibility to the applied stimulus.

Considerable work has already been done to develop photoresponsive coatings [30,31], containing azobenzene moieties that undergo a trans-to-cis interconversion under illumination of appropriate wavelengths, as well as thermoresponsive coatings based on the isotropic-nematic phase transition of the liquid crystal network [32–34]. Although such coatings provide great versatility due to the possibility of remote and dynamical actuation, the range of pre-programmed topographical profiles and the timescale of deformation remain limited. Recent work by Liu et al. proposes an electrically-responsive LCN coating in an attempt to mitigate these limitations [35], further paving the way toward dynamical, finely-resolved control of topographic profiles. In

Guido L. A. Kusters,1,* Inge P. Verheul,2 Nicholas B. Tito,3 Paul van der Schoot,1 and Cornelis Storm1,4

1Department of Applied Physics, Eindhoven University of Technology, The Netherlands
2Department of Mathematics and Computer Science, Eindhoven University of Technology, The Netherlands
3Electric Ant Lab, Amsterdam, The Netherlands
4Institute for Complex Molecular Systems, Eindhoven University of Technology, The Netherlands

*Corresponding author.

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their work. Liu et al. employ a transparent LCN film, superimposed on an array of interdigitated comb electrodes. The LCN consists, in addition to the polymer component, in large part of two species of liquid-crystal mesogen, namely (i) relatively immobile mesogens incorporated into the polymer network as permanent crosslinks and (ii) end-on grafted side-group mesogens. The latter of these have attached to them a cyano group resulting in a strong permanent dipole moment, whereas the former do not carry a permanent dipole moment. Both mesogen species are present in approximately equal amounts and are bound exclusively to the polymer component, rather than to each other. The mesogens are prepared with homeotropic alignment in the film, leading to a pre-aligned network, dense in mesogenic component. Upon in-plane actuation with an AC electric field in between the electrodes, perpendicular to the director field of the mesogens, the authors report (i) the fast and reversible formation of surface corrugations pre-programmed by the electrode placement, (ii) a surface height modulation of up to several percent in magnitude, and (iii) a clear dependence of the modulation magnitude on the driving frequency. Thus, based on purely in-plane stimuli an out-of-plane expansion of the LCN is observed. Follow-up experiments by Van der Kooij et al. on the same experimental system have since shown that there is a characteristic plasticization time associated with the initial macroscopic deformation of the LCN

Although one might expect electrothermal or electrostrictive effects to play an important role here, it can be shown that neither can adequately account for the magnitude of the observed modulation [35–38]. Instead, Liu et al. rationalise the observed behavior by remarking that, upon in-plane actuation by the electric field, the end-grafted liquid crystalline mesogens reorient slightly away from their as-prepared homeotropic alignment, increasing their mutual excluded volume with the mesogens that cannot reorient as easily due to their incorporation in the background polymer network. This increases the volume unavailable to the centers of mass of the mesogens. Indeed, if the two elongated mesogens are oriented perpendicularly to each other, then they effectively occupy more volume than if their orientations are parallel, as was already argued by Lars Onsager in the 1940s [39].

The physical picture that emerges is one of molecular voids opening up following mesogen reorientation, an overall volume increase, and subsequent viscoelastic relaxation of that volume increase as a function of time by the filling in of the voids by nonmesogenic (polymeric) material. These molecular voids are often referred to as “free volume,” i.e., the total system volume minus the total hard-core volume of the molecules comprising the film [35,36,40–42]. The result of switching on an electric field perpendicular to the director field of the mesogens is a transient macroscopic, transverse expansion of the film by virtue of the high density of mesogenic component in the LCN. Interestingly, an oscillating electric field produces the largest expansion, which arguably ties in with suppressing the viscoelastic relaxation of the host polymer matrix.

Although the coarse-grained molecular dynamics (MD) simulations of Liu et al. [35], which probe the orientational properties of the LCN on the molecular scale, make it plausible that excluded volume indeed governs LCN expansion [35], a comprehensive theoretical framework remains lacking. In particular, such a framework may prove invaluable in informing on rational design strategies, by identifying experimentally relevant parameters dictating the effectiveness of the LCN modulation, quantifying the qualitative understanding we currently have based on simulations, and minimizing costly trial and error in experiments. This last point is especially pressing, given the fact that research on the proposed setup is ongoing and topical [36].

For this purpose we construct a time-dependent Landau–de Gennes-type theory for bulk electrically responsive LCNs; investigating the formation of spatial patterns only becomes sensible if the bulk behavior is well-understood, and so is left for future work. After noting the failure of the conventional (neo-)classical theory of nematic elastomers to capture the experimental phenomenology, we instead take the connection between excluded volume and the LCN expansion as the basis for our description. We design the model under the aspect of completeness, rather than simplicity, in that we do not “strip down” the theory to its bare minimum but aim for a comprehensive description of simulation and experimental results. We subsequently compare the resulting static and dynamic response with results from MD simulations building on earlier work of one of us [35], which show qualitative agreement. We demonstrate that the dynamic response to oscillating electric fields is characterized by two resonances, and that the timescales governing the LCN actuation can be influenced by varying experimental parameters, such as the aspect ratio of the mesogens, the cross-linking fraction of the network, and the degree of initial orientational order cross-linked into the liquid crystal network. Finally, we link back to the work of Liu et al. [35] and detail how our theory explains some of their major findings.

The remainder of this paper is structured as follows. In Sec. II, we construct an equilibrium model for our bulk LCN and carry out MD simulations for comparison, focusing on the static response of the LCN. We illustrate the qualitative agreement between the two. Subsequently, we extend our model to a time-dependent description in Sec. V, and qualitatively validate the resulting dynamic response to turning on a DC electric field by comparing with our simulations. We find that there is a characteristic plasticization time associated with the macroscopic deformation of the LCN, in accordance with experimental findings [36], which we argue to depend on the aspect ratio of the mesogens, the cross-linking fraction of the network, and the degree of initial orientational order cross-linked into the liquid crystal network. Next, in Sec. VI, we demonstrate that the dynamic response to oscillating electric fields is characterized by two resonances, corresponding to two distinct modes of mesogen reorientation, and we find that the associated timescales vary with the experimental parameters in the same manner as the plasticization time does. We use this qualitative picture to explain how the expansion of the LCN varies with the electric field strength and driving frequency in the simulations and in the experiments carried out by Liu et al. [35]. Finally, in Sec. VII, we summarize our most important findings, and we provide suggestions for simulations and experiments to further pave the way toward industrial applications.
II. MODEL INGREDIENTS

We describe our LCN by means of a Landau–de Gennes-type theory, rooted in the symmetry of the underlying liquid crystal. This provides a convenient avenue for constructing a qualitative theory, without the need for in-depth knowledge of the microscopic properties of the system at hand. Since we aim for a bulk description we disregard spatial inhomogeneities, and thus the theory operates on the level of the correlation length of the system. Although the obvious approach to modeling the LCN would be to subsequently connect to classical elasticity theory, following Warner and Terentjev [20], it turns out that such a model fails to reproduce the experimental findings. Indeed, experiments show a marked volume expansion, whereas the classical theory of nematic elastomers predicts approximately volume preserving deformation. We refer to Ref. [43] for further details. This suggests the experimentally observed volume response does not follow from the usual rubber elastic behavior, and it motivates us to construct a model description based on the interplay between excluded volume generation and the LCN expansion, rather than on (nematic) rubber elasticity.

As we are interested in explicitly modeling the volume increase of the LCN upon actuation with an electric field, the proper statistical ensemble for our theory is the isothermal-isobaric ensemble. Although one would then expect the relevant thermodynamic potential to be the Gibbs free-energy density, this cannot hold true in our case because we must treat the relative volume change of the LCN as a proper statistical ensemble. Although one would then expect the relative volume change to be a minimum with respect to this relative volume change, which we denote \( \eta \equiv (V - V_0)/V_0 \), with \( V \) the system volume and \( V_0 \) the initial volume prior to any deformation, in thermodynamic equilibrium. With this stipulation in mind, the relevant thermodynamic potential becomes the Gibbs free energy per unit reference volume, which we denote \( g \) [44]. In what follows we do not write the usual pressure-volume contribution to this free-energy density, as we are solely interested in the case of the absence of any excess pressure \( p \), implying \( p = 0 \).

The material for which we aim to construct this thermodynamic potential consists of three key components, those being the polymer component and two distinct species of mesogen schematically shown in Fig. 1, with the polymer component indicated in blue (dark gray). To ensure the system is as close to being a monodomain as possible, in the experiments that inspired our work [35] the LCN is synthesized from (di)acrylates of the forms =polymer-mesogen-polymer= and =polymer-mesogen, which precludes macroscopic phase separation of the mesogens and the polymer component. This, in part, motivates our bulk description from an experimental point of view.

The focus of the model is explicitly on the two different mesogen species—recall that the LCN is dense in mesogenic component, and so effectively liquid crystalline itself—while we largely neglect the viscoelastic properties of the polymer network that experiments have shown result in the relaxation of changes in the LCN volume [35,36]; we return to this below when we investigate the dynamical behavior of the model.

One of the mesogen species incorporated in the LCN is crosslinked into the polymer network on both ends, as indicated in orange (gray) in Fig. 1, meaning such mesogens are immobile and can because of that not significantly reorient in response in an externally applied electric field. The other mesogen species is grafted end on onto the polymer network, as indicated in green (light gray) in Fig. 1, resulting in much more mobile dangling side groups. This species has on top of that attached to it a permanent dipole moment, ensuring that it interacts more strongly with electric fields than the cross-linked mesogens do. This suggests that the cross-linked mesogens cannot respond strongly to any applied external electric field, justifying our assumption that only the dipolar mesogens effectively interact with electric fields. The interaction free energy of the dangling mesogens is quadratic in the electric field \( E \) for nematic liquid crystals, by virtue of the inversion symmetry of the nematic director. This proportionality applies even for a collection of mesogens with a permanent dipole moment, which formally breaks this inversion symmetry, provided that the electric field is sufficiently weak, and the nematic symmetry is not broken by polarization of the mesogens [45].

In addition to the different response of the two types of mesogen to electric fields, the model must also reflect the different ways in which the two mesogen species are connected to the polymer network. In particular, if the cross-linked mesogens reorient, for example, due to excluded-volume interactions with neighboring dipolar mesogens, they are subject to an elastic restoring force from the polymer network. In contrast, reorientation of the dipolar mesogens is impeded much less significantly by the polymer network, because these mesogens are only connected to the network by one end. To capture this asymmetry we assume that the dipolar mesogens can reorient freely in the polymer network, whereas the cross-linked mesogens are subject to a harmonic spring potential centered about the orientations
that were cross-linked into the LCN during its preparation in the absence of an electric field. Here, we refer specifically to the degree of orientational order to which the cross-linked mesogens have relaxed under the experimental conditions [20].

The presence of two distinct mesogen species already suggests we need, at the least, three order parameters in our theory to model the volume expansion of the LCN resulting from any changes in the orientations of the mesogens in response to the electric field. That is, in addition to the aforementioned order parameter \( \eta \) to keep track of the relative volume change, or, equivalently, the relative density change, of the LCN, we also require two order parameters to inform on the orientational properties of the cross-linked and dipolar mesogens, respectively. The most general way to express such orientational order is by means of the symmetric and traceless tensor nematic order parameter \( Q_{ij} = (u_iu_j - \delta_{ij}/3) \), with \( u \) the mesogen orientation unit vector and \( \delta_{ij} \) the Dirac \( \delta \). This suggests the introduction of two tensor order parameters, \( Q_{ij}^{(c)} \) and \( Q_{ij}^{(d)} \), corresponding to the cross-linked and dipolar mesogens, respectively, to describe the orientational order of the different mesogen species.

In what follows we assume that the cross-linked mesogens, due to their inability to strongly respond to any applied electric field, remain approximately uniaxial and along their original director even if we apply the electric field perpendicular to the director; below we illustrate that our MD simulations justify this approximation. We then recover the uniaxial tensor order parameter \( Q_{ij}^{(c)} = S_c(\delta_{ij} - \delta_{ij}/3) \) for the cross-linked mesogens, where we fix the director along the \( y \) axis and \( S_c \equiv \frac{1}{2}(\cos^2 \theta - 1) \) denotes the scalar nematic order parameter, with \( \theta \) the angle a test mesogen makes with the director field and \( \langle \ldots \rangle \) an angular average. It is clear, however, that the dipolar mesogens produce, in general, a biaxial configuration if we apply the electric field perpendicular to the director. Although the usual approach would be to then write down the corresponding tensor nematic order parameter in the biaxial form \( Q_{ij}^{(d)} = S_d(\delta_{ij} - \delta_{ij}/3) + P/3(\delta_{i1}\delta_{j1} - \delta_{i2}\delta_{j2}) \), where the electric field along the \( x \) axis produces a degree of biaxiality \( P = \frac{1}{2}(\sin^2 \theta \cos 2\phi) \), with \( \phi \) the angle a test mesogen makes with the electric field in the \( x-z \) plane, our MD simulations motivate us to take a different approach here.

From our MD simulations we learn that upon application of an electric field perpendicular to the director field in which both mesogen species are prepared, a clear division arises between dipolar mesogens that align with the director field and dipolar mesogens that align with the electric field. This is presumably due to the presence of the cross-linked mesogens, which remain ordered predominantly along the director field, justifying our approximation of uniaxial order for this mesogen species. The excluded-volume interactions with these mesogens must then be overcome by the dipolar mesogens to align with the electric field. This finding motivates us to, instead of describing the biaxial order of the dipolar mesogens collectively, separately describe the uniaxial order of the collection of director-field-aligned dipolar mesogens and the collection of electric-field-aligned dipolar mesogens. Note that since we consider the director and the axis of biaxial order as fixed, this does not in itself increase the rotational degrees of freedom of the LCN.

We model the above by associating the collection of nematic-director-field-aligned mesogens with a fraction \( 1 - f^2 \) of the dipolar mesogens and the collection of electric-field-aligned dipolar mesogens with the remaining fraction \( f^2 \). Treating \( f^2 \) as an order parameter in our model then allows us to interpolate between these different populations of dipolar mesogen, in effect signifying a bistable equilibrium between the two. Here, the use of the quadratic form \( f^2 \geq 0 \) ensures this fraction is always nonnegative; the constraint \( f^2 \leq 1 \) we enforce by hand. We remark that, of course, taken together these populations of dipolar mesogens again exhibit biaxial order; the main advantage of treating this biaxial order as a bistable equilibrium is the ability to also capture the interpopulation between the two populations, i.e., the reorientation of the dipolar mesogens, which our MD simulations indicate to be important.

The above suggests that we need not use two tensor nematic order parameters, \( Q_{ij}^{(c)} \) and \( Q_{ij}^{(d)} \), the latter of which is in general biaxial, to describe the orientational order of the cross-linked and dipolar mesogens, respectively. Instead, we use three uniaxial tensor nematic order parameters for the orientational order of the cross-linked mesogens, the collection of director-field-aligned dipolar mesogens and the collection of electric-field-aligned dipolar mesogens. Relabeling for ease of reference, we denote these \( Q_{ij}^{(1)} \), \( Q_{ij}^{(2)} \), and \( Q_{ij}^{(3)} \), respectively. The former two have their director along the \( y \) axis, along which both mesogen species are prepared, whereas the director of the latter lies along the \( x \) axis, along which the electric field is applied. We denote the corresponding scalar nematic order parameters, as measured along their respective directors, \( S_1, S_2, \) and \( S_3 \), which are schematically shown in Fig. 2.

Although this implies that we can capture the orientational order of the cross-linked and dipolar mesogens as \( Q_{ij}^{(c)} = \)
and \( Q_{i,j}^{(1)} \) and \( Q_{i,j}^{(2)} = (1 - f^2)Q_{i,j}^{(2)} + f^2 Q_{i,j}^{(3)} \), respectively, and the global orientational order as \( Q_i = \langle Q_i^{(c)} + Q_i^{(d)} \rangle / 2 \) on account of there being equal amounts of both in the network, we will not construct our Landau theory in terms of the invariants of these tensors. The reason for this is that since these tensor nematic order parameters are defined along different axes, their explicit combination turns out to introduce unphysical couplings. In particular, coupling terms promoting cross-hatched, i.e., perpendicular, orientational order of the different mesogen species and populations arise in the nematic phase, in which we are interested [43]. The alternative we follow here is to construct our model in terms of the invariants of the individual tensor nematic order parameters of the different mesogen species and populations we distinguish, i.e., \( Q_{i,j}^{(1)} \), \( Q_{i,j}^{(2)} \), and \( Q_{i,j}^{(3)} \) for the cross-linked mesogens, the director-field-aligned dipolar mesogens and the electric-field-aligned dipolar mesogens. A similar division of different types of nematogens into populations was in fact already used by Onsager as early as 1949 in his seminal work [39].

Clearly, the invariants that comprise our model must also give rise to interactions between the different mesogen species and order parameters. Broadly speaking, these interactions come in two flavors: mesogen-mesogen interactions and mesogen-volume interactions. The first of these couple the orientational properties of the cross-linked mesogens to those of the dipolar mesogens via excluded-volume-like interactions. This has the effect of promoting the alignment between the different mesogen species, as their mutual excluded volume is smallest when they align. Assuming identical rod-like shapes for both mesogens with length \( L \) and diameter \( D \), the mutual excluded volume of two mesogens is proportional to \( L^2 D + O(LD^2) \), we expect the same dependence on the dimensions of the mesogens for this interaction [39]. A similar argument can be made if we increase the cross-linking fraction of the network, while keeping the total amount of mesogens constant, i.e., if we increase the fraction of mesogens that are fully cross-linked into the polymer network at the expense of dipolar mesogens. Then, any dipolar mesogen will have an increased number of immobile cross-linked mesogens as its neighbors. This makes it increasingly difficult for the electric field to overcome the excluded-volume interactions of the dipolar mesogen with its neighbors to induce reorientation. Thus, the excluded-volume mesogen-mesogen interactions must be stronger in this case, at least regarding the reorientation of the dipolar mesogens. We remark that the reverse effect of any cross-linked mesogen interacting with a decreasing number of dipolar mesogens is much less significant. The reason for this is that these mesogens are fully cross-linked into the polymer network and cannot respond strongly to the applied electric field.

The second type of interaction between order parameters we incorporate couples the orientational properties of the mesogens to the volume increase of the LCN. This interaction is closely related to the excluded-volume interactions discussed above, as in sufficiently dense mesogenic systems the excluded volume generated upon mesogen reorientation contributes to the total volume of the system. To reflect this, we associate configurations in which the different mesogen species become increasingly aligned with a volume decrease, whereas the volume increases if the mesogens become orientationally disordered.

Taken together, the above ingredients define a Gibbs free energy per unit reference volume for our model as a function of the following order parameters: (i) the relative volume change of the LCN, \( \eta \), (ii) the tensor nematic order parameters \( Q_{i,j}^{(1)} \), \( Q_{i,j}^{(2)} \), and \( Q_{i,j}^{(3)} \) for the cross-linked mesogens, the director-field-aligned dipolar mesogens and the electric-field-aligned dipolar mesogens, and (iii) the population fraction \( f^2 \). Note that since the tensor nematic order parameters are all presumed to be uniaxial, with fixed directors, all information on the magnitude of orientational order is encapsulated in the associated scalar nematic order parameters \( S_1 \), \( S_2 \), and \( S_3 \) in terms of which we express our model below.

### III. Theory

Taking all the contributions described in the section above into account, our model Gibbs free energy per unit reference volume becomes

\[
g = \sum_{i=1}^{12} g_i, \tag{1}
\]

where we will discuss the form of each contribution \( g_i \) in turn. The first three contributions correspond simply to the free energy associated with each mesogen species or population, according to

\[
g_1 = g_{\text{LdG}}(S_1), \quad g_2 = (1 - f^2)g_{\text{LdG}}(S_2), \quad g_3 = f^2 g_{\text{LdG}}(S_3). \tag{2}
\]

Here,

\[
g_{\text{LdG}}(S_i) = \frac{1}{4} A S_i^2 - \frac{1}{4} B S_i^3 + \frac{1}{4} C S_i^4, \quad i = 1, 2, 3, \tag{3}
\]

denotes the standard Landau–de Gennes free-energy density for uniaxial nematic liquid crystals, with \( A, B, C \) phenomenological coefficients, as follows from a power series expansion in terms of \( \text{Tr} Q^{(ij)} \), where \( \text{Tr} \) denotes the trace operator and the series is truncated after \( n = 4 \) [27,46].

Although the phenomenological coefficient \( A \) is generally taken to depend explicitly on temperature, to reflect the temperature-induced isotropic-nematic phase transition, we are currently not interested in modeling this dependence. In fact, prior simulations and experiments suggest the temperature-dependent behavior of the LCN is significantly more complicated than can be expected from a simple Landau theory [35,36]; our own MD simulations confirm this. Instead, we choose \( A \) such that the configuration in which the LCN is prepared, given by \( S_1 = S_2 = S_3 = 0 \), \( f^2 = 0 \), and \( \eta = 0 \) minimizes the total free-energy Eq. (1) in the absence of an electric field. Here, \( S_i \) denotes the as-prepared scalar nematic order parameter, the value of which we specify further below by comparing with the simulations, and the value of \( S_3 \) matters not as there are no electric-field-aligned dipolar mesogens in the absence of an electric field. As will become apparent below, this statement is identical to choosing \( A \) such that \( S_0 \) minimizes Eq. (3), since the remaining contributions to the free energy are automatically minimized under these conditions. This effectively eliminates \( A \) from our model description in favor of \( S_0 \).
We stress that the above should not be interpreted as us neglecting entropy by making the temperature dependence of our theory implicit; we are just not interested in probing this explicitly. The effect is merely cosmetic: all parameters in our theory carry, to a greater or lesser extent, a temperature dependence, and so have associated with them, in a coarse-grained fashion, entropy.

Finally, note that in Eq. (2) the free-energy contribution of Eq. (3) is multiplied by factors of \((1 - f^2)\) and \(f^2\) for the free energy associated with the director-field-aligned dipolar mesogens \(g_2\) and the electric-field-aligned dipolar mesogens \(g_3\), ensuring only the fraction of dipolar mesogens that is in a given population contributes to its free energy. As we presume there to be equal amounts of the cross-linked and dipolar mesogens, reflecting the experimental work of Liu et al. [35], their free energies bear equal weight.

The next contribution encodes the interaction of the dipolar mesogens with the electric field, which, on symmetry grounds, must be proportional to the product of the electric field \(\mathbf{E}\) with the electric susceptibility tensor \(\Sigma\) of said mesogens, according to \(-\mathbf{E} \cdot \Sigma \cdot \mathbf{E}\). Since the electric susceptibility of the dipolar mesogens is closely related to their orientational order, this contribution takes the form [27,46]

\[ g_4 = -f^2 HS_3, \tag{4} \]

where \(H \propto |\mathbf{E}|^2\) acts only on the electric-field-aligned dipolar mesogens, not the director-field-aligned dipolar mesogens. This is because the electric field is applied perpendicular to the director field, and so the associated contribution to the free energy is small due to the low susceptibility of the mesogens along this axis. For this reason \(H\) actually represents a relative value, rather than an absolute one. This approximation is also in line with the philosophy of the model, in which we have two populations of dipolar mesogens, which order along perpendicular axes. Indeed, a similar coupling between \(S_2\) and \(H\) would promote orientational order of the electric-field-aligned dipolar mesogens lies along the electric field, along the \(x\) axis, rather than the director field, along the \(y\) axis, yielding \(\text{Tr}((\mathbf{Q}^{(1)}) - (\mathbf{Q}^{(3)}))^2\) instead.

We point out that we do not explicitly incorporate excluded-volume-like interactions between the different populations of dipolar mesogen because this would, again, promote orientational order of these populations perpendicular to their assigned director, undermining the philosophy of the model. Instead, the excluded-volume-like interactions between the different populations of dipolar mesogen are mediated by the population fraction \(f^2\), such that if the orientational mismatch between the two becomes sufficiently stringent, the population fraction adjusts to mitigate this. This feature is already included qualitatively in Eq. (6). The sceptical reader may take comfort in the fact that including a cross term between the different populations of dipolar mesogens does not significantly alter the qualitative behavior of the model, provided that the associated coupling constant remains moderate (results not shown).

As already announced, the interaction between the orientational order of the mesogens and the relative volume increase, or, equivalently, the density decrease, of the LCN is closely related to the excluded volume of the mesogens in our model. However, instead of penalizing misalignment between the mesogens, we now must penalise the volume of the LCN when the mesogens become increasingly aligned. Thus, instead of constructing the invariants of the difference between tensor order parameters we now take their sums \(\text{Tr}((\mathbf{Q}^{(1)}) + (\mathbf{Q}^{(2)}))^2\) and \(\text{Tr}((\mathbf{Q}^{(1)}) + (\mathbf{Q}^{(3)}))^2\), resulting in the contributions

\[ g_8 = (1 - f^2)\lambda(S_1 + S_2)^2, \]

\[ g_9 = f^2\xi\eta(S_1^2 - S_1S_3 + S_3^2), \]

\[ g_{10} = -\xi\eta(4S_0^2). \tag{7} \]

Here, we have multiplied the various contributions by the volume order parameter \(\eta\) to ensure the proper effect on the...
system volume, $ξ$ is a phenomenological coefficient and we have again separated the contributions associated with the different populations of dipolar mesogens apparent from $g_8$ and $g_9$. We ignore explicit cross terms, as we did before, to ensure the model remains internally consistent. The final contribution shown in Eq. (7) ensures the orientational order of the mesogens is measured relative to the initial configuration by subtracting from the contributions $g_8$ and $g_9$ their counterparts with $S_1 = S_2 = S_0$ and $f^2 = 0$.

Finally, we must ensure that the free energy of Eq. (1) is bounded from below, which leads us to add the additional terms

$$g_{i1} = \frac{1}{2} B_i f^4, \quad g_{i2} = \frac{1}{2} B_0 \eta^2,$$

representing bulk-modulus-like terms in the population fraction $f^2$ and the relative volume fraction $η$. Note that, technically, the former is not required to keep the free energy bounded from below since we constrain $0 \leq f^2 \leq 1$. However, we do need it to allow $f^2$ to take intermediate values between 0 and 1 upon minimizing Eq. (1) with respect to all order parameters.

Although the model we present here focuses on a bulk system, the same framework can be used to analyze the formation of spatial patterns, by including (square) gradient terms in all order parameters (cross terms vanish, to lowest order, on symmetry grounds). We have carried out a preliminary analysis on a nonlocal extension of the current theory, which shows that free volume permeates the LCN from the top in the thin-film geometry used in the experiments of Liu et al. [35]; this is in accordance with additional MD simulations we have carried out. We hope to present our results in a future publication [47].

It is clear that the model outlined above in principle requires for a very large parameter space to be explored, potentially giving rise to very rich behavior (see also below). However, the parameter space can be significantly reduced by scaling the theory to make it dimensionless. To this end, we introduce the set of scaled scalar nematic order parameters $s_i = S_i/(B_i^2/2C)$, with $i = 1, 2, 3$, and the scaled free volume order parameter $\tilde{\eta} = \eta/(\tilde{\xi}^2 B_0 C)$, where we leave the population fraction $f^2$ unchanged; these order parameters are subject to the scaled electric field strength $h = H/(B_i^2/2C^2)$. This procedure reduces the set of phenomenological parameters characterizing our model to the parameter combinations $\tilde{k} = k/(B_i^2/C)$, associated with the elastic energy of the cross-linked mesogens to the polymer network, $\tilde{\lambda} = \lambda/(B_i^2/C)$, associated with the excluded-volume interactions between the different mesogen species, $\tilde{\xi} = \xi^2/(B_0 C)$, associated with the coupling between the orientational order of the mesogens and the LCN volume, $\tilde{B}_i = B_i/(B_i^2/C)$, corresponding to the bulk-modulus free-energy penalty associated with the population fraction $f^2$, and $\tilde{s}_0 = S_0/(B/2C)$, which represents the initial degree of orientational order, crosslinked into the network. Recall that we eliminate the phenomenological coefficient $A$ from the model description by demanding that the initial configuration, with $S_1 = S_2 = S_0$, $f^2 = 0$ and $\eta = 0$, minimizes the free-energy density per unit reference volume Eq. (1) in the absence of an electric field, implying that $A = B S_0 - C S_0^2$. The result of this scaling procedure is a universal curve for the theory in terms of the scaled order parameters $s_1, s_2, s_3, f^2$, and $\tilde{\eta}$, dependent only on the set of scaled parameters $\tilde{k}, \tilde{\lambda}, \tilde{\xi}, \tilde{B}_1$, and $s_0$, as well as the scaled electric field strength $h$. We stress that this already significantly reduces the model parameter space, and refer to Appendix B for a discussion on a fully "striped-down" version of the model that captures the most important physics and results, though it lacks the more subtle aspects of the full theory required to complete the picture.

Although the parameter space available to the scaled theory is still significant, extensive numerical tests, carried out by numerically minimizing the scaled free energy per unit reference volume Eq. (1) using the “lsolve” root-finding algorithm in the programming language Python, show that there is a broad range of parameter values that produce the key features of the results of both our MD simulations and the experiments (results not shown). We point out that in carrying out the minimization procedure, we fix the value of order parameters that exceed their permitted value range, i.e., $0 \leq f^2 \leq 1$. Although a similar constraint applies to the scalar nematic order parameter $-1/2 \leq S_i \leq 1$, with $i = 1, 2, 3$, we cannot explicitly enforce this constraint for their scaled counterparts $s_i$, since the effective values of the scaling coefficients $B$ and $C$ are unknown for the LCN under consideration. For the purpose of this paper we focus on parameter values representative of the regimes described above.

Before we present the corresponding equations of state, we first pause to elucidate the simulation protocol for our coarse-grained MD simulations, as these will serve as a reference for the theory in the remainder of this paper. In our simulations, we work in terms of the fundamental Lennard-Jones units for distance $\lambda$, energy $\mathcal{E}$, mass $M$ and time $\tau = \sqrt{\lambda M D^2/\mathcal{E}}$. The electric field strength is expressed in terms of $\sqrt{\mathcal{E}/(4\pi \epsilon_0 \lambda D^3)}$, with $\epsilon_0$ the vacuum permittivity.

We use the HOOMD-blue package (v2.1.1) to simulate the liquid crystal network within the isothermal-isobaric (NpT) ensemble [48,49], using periodic boundary conditions. Sample snapshots of the simulation model are shown in Fig. 1. As stated above, these simulations aim only to examine bulk behavior of the LCN. Thus, the simulation box size need only be chosen to be sufficiently large to capture the correlation length for structural perturbations (primarily void creation) caused by actuation of a given dipolar mesogen. This length scale is rather small, being only on the order of several mesogens in size at most. Our chosen simulation box size encompasses hundreds of these correlation volumes, as visually evident in Fig. 1.

As visualized in Fig. 1, we represent the polymer mainchain [blue (dark gray)] as strings of particles, held together by harmonic bonds

$$U_{\text{bond}}(r) = \frac{1}{2} k (r - r_0)^2,$$

with $k = 1110 \mathcal{E}/D^2$ and $r_0 = 0.9 \lambda$ [50], acting on their distance of separation $r$. In addition, we represent the crosslinked mesogens [orange (gray)] and the dipolar mesogens [green (light gray)] as collections of five particles held together by stiff harmonic bonds with $k = 5000\mathcal{E}/D^2$ and $r_0 = 0.5 \lambda$, and kept in a rodlike shape by harmonic potentials,

$$U_{\text{ang}}(\theta) = \frac{1}{2} k_0 (\theta - \pi)^2,$$
with \( k_0 = 200 \, E/\text{rad}^2 \), acting on the angle \( \theta \) between three particles. Note that the stiff rodlike connectivity for the Lennard-Jones beads composing each mesogen results in an effective angle-dependent mesogen-mesogen intermolecular potential, which stabilizes the nematic phase.

Our simulations contain 70 polymer main-chains, each 50 particles long, 1166 mesogens of both species, and we incorporate no explicit or implicit solvent. Finally, all particles interact with each other by means of Lennard-Jones interactions,

\[
U_{\text{LJ}}(r) = 4\epsilon \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^6 + U_{\text{LJ,0}},
\]

with \( \epsilon = 1 \, E \) and \( \sigma = 1 \, D \), which are cut off at \( r_c = 2.5 \, D \), \( U_{\text{LJ,0}} \) is chosen such that \( U_{\text{LJ}}(r_c) = 0 \).

To generate an initial condition for our simulated LCN, a virtual polymerization of the initial ingredients (polymer main chains + mesogens) is carried out in situ. The polymerization kinetics are irrelevant for our purposes here; the objective is only to obtain a representative (in terms of a realistic density and topology) cross-linked LCN by the end of the procedure, for use in the subsequent perturbation simulations.

The in situ polymerization is done by assigning binding sites to the polymer main-chain particles [shown as green (light gray) dots in Fig. 1], and assigning to the terminal particles of both mesogen species the binding partner [shown as orange (gray) dots in Fig. 1]. The cross-linked mesogens contain binding partners on either end, whereas the dipolar mesogens only contain a binding partner on one end. To enforce one-to-one binding of binding sites and binding partners, as we discussed is the case in the experiments of Liu et al. [35], we introduce first a repulsive inverse power potential,

\[
U_{\text{pow}}(r) = 4\epsilon \left( \frac{\sigma}{r} \right)^{12} + U_{\text{pow,0}},
\]

which is cut off at \( r_c = 2.5 \, D \), and \( U_{\text{pow,0}} \) is chosen such that \( U_{\text{pow}}(r_c) = 0 \). This potential acts on both binding sites and pairs of binding partners [both with \( \sigma = 0.9 \, D \) and \( \epsilon = 1 \, E \)], as well as on pairs containing both a binding site and a binding partner and pairs of the Lennard-Jones particles themselves [both with \( \sigma = 0.6 \, D \) and \( \epsilon = 1 \, E \)]. We next add a strongly attractive Gaussian potential,

\[
U_{\text{Gauss}}(r) = \epsilon_{\text{site}} \exp \left[ \frac{1}{2} \left( \frac{r}{\sigma_{\text{site}}} \right)^2 \right].
\]

with \( \epsilon_{\text{site}} = -500 \, E \), \( \sigma_{\text{site}} = 0.2 \, D \), and cutoff length \( r_c = 3 \, D \), acting only on pairs containing both a binding site and a binding partner. Finally, to achieve the desired degree of mesogen alignment during crosslinking, equal and opposite unity charges \( q = \pm \sqrt{4\pi \epsilon_0 E/2D} \) are added to the ends of both the cross-linked and the dipolar mesogens, and a dummy electric field \( E \) is applied along the \( x \) axis. The electric field strength can be varied to change the degree of mesogen alignment and the charges serve only to couple to the electric field; no Coulombic interactions between the dipoles are taken into account.

We then crosslink the system in the presence of the dummy electric field by repeatedly performing short simulations in the \( NVE \) ensemble and permanently connecting overlapping binding sites and partners. This process is repeated until a desired fraction of the binding sites (85%, in our case) has been permanently bound to a partner. Here, the stoichiometry is such that there are on average two bonds to a mesogen of either species for every three particles on the polymer main-chain, emulating the experimental system, dense in mesogenic component. Following this, all binding sites and partners, as well as the associated potentials \( U_{\text{pow}} \) and \( U_{\text{Gauss}} \), are removed from the simulation.

After crosslinking, the LCN is equilibrated in the \( NpT \) ensemble in the presence of the dummy electric field, using the Martyna-Tobias-Klein barostat-thermostat [51], with temperature and pressure coupling constants set to unity. We use \( T = 0.35 \, E/k_B \) for the temperature, \( p = 1 \, E/D^3 \) for the pressure, \( dt = 0.001 \, T \) for the time step size, and we equilibrate for \( 10^4 \) time steps. Subsequently, the charges are removed from the cross-linked mesogens, which are inert in our simulations, and we carry out a second \( NpT \) equilibration for \( 10^4 \) time steps in the absence of the dummy electric field. The “built-in” nematic order persists due to two key factors: (i) the rather high volume fraction of mesogens relative to polymer segments, and (ii) the existence of steric constraints, stemming from the dense connectivity of the network. This yields the initial configuration of the LCN for the simulations presented in this paper, for which we use a total simulation time of \( 300 \, T \). We remark that we ensure that the simulation results are not particular to a specific realization by carrying out the procedure described above multiple times and each time introducing random distortions (permanent bonds) during the crosslinking process, resulting in different random network topologies for the LCN. For further details regarding the simulation protocol, we refer to Refs. [35,52].

This concludes a discussion of our theory. We now present our equations of state.

### IV. EQUATIONS OF STATE

Figure 3 shows the results of a numerical minimization of Eq. (1), scaled as described in the previous section, compared with the phase diagram we obtain from our MD simulations. The figure illustrates good qualitative agreement between theory (a, c), where we show a single universal curve for the chosen parameter values, and simulations (b, d), where we show results for three randomly polymerized network topologies. Indeed, focusing first on the orientational properties of the LCN (a, b), we find for increasing electric field strength: (i) no to weak response to variations in the electric field strength for both mesogen species below a critical field strength, (ii) a critical field strength beyond which the response to variations in the electric field strength is strong, and (iii) saturation of the orientational order. This last regime is colored green (gray) in Fig. 3 and corresponds to a saturated population of field-aligned dipolar mesogens, i.e., \( f^2 \leq 1 \) by hand. Recall that a similar constraint for the scalar nematic order parameters \( s_i \), with \( i = 1, 2, 3 \), cannot be enforced because these have been scaled with coefficients of which the values are unknown.

The same qualitative features hold for the concomitantly generated free volume (c, d), albeit with a decrease in volume.
in the final regime rather than the saturation of the orientational order. This last feature, which gives rise to a maximum in the free volume curve, follows directly from the orientational order at the corresponding electric field strength: the maximum occurs at the point of maximum global orientational disorder. We attribute the slight decrease in volume we find in our simulations at very weak electric field strengths (d) to the initial configuration in our simulations corresponding to a local minimum of the free energy, rather than the global free-energy minimum. This is because relaxation of the LCN to thermodynamic equilibrium occurs on timescales slower than those accessible in our simulations; prior simulations and experiments also hint at this slow relaxation [35,36]. If we subsequently apply a weak electric field to this initial configuration, which is stable on the simulation timescales in the absence of an electric field, then we help the system to relax to a configuration with a lower free energy by decreasing its volume slightly. Noting that we do not aim to capture this last feature explicitly within the model, we find that the mechanism of free volume generation, on which the theoretical model is based, qualitatively explains simulation results and shows good qualitative agreement.

Although the static behavior of the model we show here serves as a proof of concept, supporting the role of excluded volume in driving the LCN to expand upon actuation, we still lack some key features of the simulations and the experiments. Most notably, the static model neglects the viscoelastic relaxation of the polymer network, which ensures the volume of the LCN relaxes as a function of time, if we apply a constant electric field. Accordingly, to achieve a steady-state volume increase of the LCN in the simulations and the experiments, alternating electric fields are used to continually actuate the LCN before it relaxes viscoelastically [35,36]. In what follows, we will be interested in how our model parameters, and in particular those that are experimentally accessible, influence the response of the LCN to alternating electric fields. To this end, we dedicate the next section to dynamics, where we first ensure our model dynamics properly reflects the viscoelastic relaxation of the LCN in response to the application of a constant electric field.

V. DYNAMICS

The equilibrium model we describe in the previous sections can also be used to investigate the dynamical behavior of the LCN, by complementing it with a set of differential equations governing the temporal behavior of the order parameters. To this end, however, it is important to recall that the model, represented by Eq. (1), neglects the viscoelastic relaxation of the LCN, which is known to play an integral role in the volume expansion dynamics observed in the simulations and the experiments [35,36]. This viscoelastic relaxation is also the reason that AC actuation is needed to achieve steady-state LCN modulation in the simulations and the experiments, as the response following DC actuation simply relaxes. Thus, to connect with simulations and experiments, the dynamical equations characterizing the model must be adapted to account for this. For the nonconserved order parameters under consideration, the simplest form the associated differential equations can take describes relaxational dynamics [53], according to

$$\partial_t \psi = -\Gamma_\psi \left( \frac{\partial g}{\partial \psi} + \gamma \eta(\tau)\phi^2 \delta_{\psi, \eta} \right) + \theta_\psi \cdot \text{(14)}$$

Here, $\tau$ denotes time, $\psi$ represents the relevant order parameter, i.e., $S_1$, $S_2$, $S_3$, $f$ or $\eta$, $\Gamma_\psi$ denotes the associated kinetic coefficient, which “hides” the dissipative physics that facilitate decreasing the (free) energy upon varying the relevant order parameter, and $\theta_\psi$ represents a Gaussian noise term, which ensures that the fluctuation-dissipation theorem is satisfied [54]. The first term in brackets represents the mean-field dynamics following from the free-energy density shown in Eq. (1), whereas the second term effectively corrects for the viscoelastic relaxation of the LCN, i.e., it ensures the free volume order parameter $\eta$ relaxes as a function of time. Here $\gamma$ is a phenomenological constant, $\langle \tau \rangle_\eta$ denotes the ensemble-
averaged time the free volume has existed for, and \( \delta_{\psi, \eta} \) the Kronecker \( \delta \). Note that this last term provides an effective dynamical coupling between the different order parameters that is of a non-free-energetic nature.

Formally, the inclusion of this relaxational term implies that at thermodynamic equilibrium there cannot exist any nonzero free volume. However, by virtue of the aforementioned slow relaxation of the LCNs under consideration [35,36], we do not expect full relaxation to thermodynamic equilibrium to occur on the available timescales. Instead, in our MD simulations the temporal evolution of the system stops upon reaching some locally stable configuration. A similar construction for the scaled theory can be achieved by stopping the model dynamics after some time \( t_{\text{stop}} \), shorter than the timescale required for full relaxation to thermodynamic equilibrium, has passed. This construction can be used to show that the qualitative features of the phase diagram in Fig. 3 remain unchanged following the addition of free volume relaxation, provided that the system cannot fully relax to thermodynamic equilibrium on the used timescales [43].

In addition, although the Gaussian noise term \( \theta_{\eta} \) plays an integral part in the dynamics of some systems [55–59], here we limit ourselves to mean-field dynamics by setting \( \theta_{\eta} = 0 \). This is justified, as the phase diagram we report on Appendix A for the full scaling procedure. In addition, although the Gaussian noise term \( \theta_{\psi} \) plays an integral part in the dynamics of some systems [55–59], here we limit ourselves to mean-field dynamics by setting \( \theta_{\psi} = 0 \). This is justified, as the phase diagram we report on Appendix A for the full scaling procedure. In this case the instability plasticizing the LCN is forced, such that we are unable to temporally resolve the existence of a plasticization time. The importance of such fluctuations in plasticizing the LCN is illustrated in the paragraphs below, where we more strongly connect our theoretical findings regarding this plasticization time with experiments.

To further substantiate the experimental relevance of the predicted plasticization time, and to illustrate the importance of fluctuations in plasticizing the LCN, we estimate the short-time relaxation dynamics by linearizing the dynamical Eq. (14) in the observables \( S_1, S_2, S_3, f^2 \), and \( \eta \). That is, we expand about the initial configuration \( S_1 = S_0 + \epsilon \delta S_1, S_2 = S_0 + \epsilon \delta S_2, S_3 = S_0 + \epsilon \delta S_3, f^2 = f^2(\delta f)^2, \) and \( \eta = \epsilon \delta \eta \), with \( \epsilon \ll 1 \), and solve the set of dynamical equations up to \( O(\epsilon) \). The resulting exponential solution for the scaled free volume \( \tilde{\eta} \) is indicated by the dashed curve in Fig. 4. We find that the plasticization time roughly corresponds to the time required to saturate the field-aligned population of dipolar mesogens \( f^2 = 1 \) in the short-time approximation. The estimate then reads

\[
\tau_p \approx \frac{1}{\Gamma_f S_0} \frac{1}{2H - 3\lambda S_0} \log \delta f(0),
\]

where \( \delta f(0)^2 \ll 1 \) denotes the initial fraction of field-aligned dipolar mesogens. From the divergence of Eq. (15) at \( H = 3\lambda S_0/2 \) it is apparent that we recover a critical field strength, as was also the case for the phase diagram in Fig. 3. Above this critical field strength \( \tau_p > 0 \), and so fluctuations in \( f^2 \) spontaneously grow under the influence of the electric field. Conversely, below the critical field strength \( \tau_p < 0 \), indicating that fluctuations relax as a function of time.
Notably, Eq. (15) explicitly shows the dependence of the plasticization time on experimentally accessible parameters. Although the dependence on the electric field strength $H$ is straightforward to interpret, by noting that it exerts a torque on the dipolar mesogens to induce reorientation, Eq. (15) also suggests that decreasing the orientational order cross-linked into the network, $S_0$, generally decreases the plasticization time. Similarly, the dependence on the parameter $\lambda$, which scales with the magnitude of excluded volume interactions, suggests that a decrease in plasticization time can be achieved by decreasing the liquid crystalline mesogen aspect ratio or the cross-linking fraction.

Finally, Eq. (15) clearly indicates that some initial disorder is important in driving free volume generation, as for a perfectly aligned initialization $\delta f(0) \rightarrow 0$ the plasticization time diverges $\tau_p \rightarrow \infty$. Although in experiments and simulations the initial configuration is naturally characterised by small random fluctuations of the order parameters around the equilibrium values, in the theory we enforce this by setting $\delta f(0) > 0$.

This establishes the dynamical response of our model LCN upon the application of a constant electric field. Below we extend this discussion to alternating electric fields, which are of most direct interest for future experiments and applications.

VI. AC ACTUATION

The characteristic plasticization time is also relevant when applying an AC electric field $H \propto (|E|^2 \cos 2\pi \omega t)^{\frac{1}{2}}$, with $\omega$ the driving frequency and $r$ the time, as illustrated by Fig. 5. Figure 5(a) shows, for a given electric field strength, how the steady-state free volume varies as a function of the driving frequency. This yields two temporally separated resonances for free volume generation: a low-frequency resonance (blue circles) and a high-frequency resonance (red squares).

Figure 5(b) shows how these resonance frequencies vary as a function of the electric field strength (colored symbols), as compared to the numerically evaluated plasticization time [solid black curve; the dashed line shows the estimate Eq. (15)]. We again observe no response below the critical field strength, followed by two distinct regimes: “low” driving frequencies dominate at low field strengths (blue circles) and “high” driving frequencies dominate at high field strengths (red squares). Note that we do not show the low-frequency resonance for high electric field strengths because it is completely dominated by the high-frequency resonance there, i.e., it is completely absorbed into the high-frequency resonance and no longer detectable. The same holds for the high-frequency resonance at low electric field strengths.

The existence of two resonances implies two coupled processes must underlie our findings: mesogen reorientation and the response of the volume to this reorientation. This claim is supported by the fact that the two distinct resonances persist even if we fix the nematic scalar order parameters to their as-prepared values $S_1 = S_2 = S_3 = S_0$, indicating the fraction $f^2$ and the free volume order parameter $\eta$ dictate the most important dynamics. In fact, some of the most salient results we find can be reproduced based on a “minimal” version of our theory, governed only by the order parameters $f^2$ and $\eta$, as presented in Appendix B. Although the nonlinear nature of the model, even if stripped down to its basics, has not allowed us to analytically estimate the resonance frequencies in a meaningful way, a qualitative comparison of the resonance frequencies with the plasticization time yields additional insights. Indeed, despite the low-frequency resonance occurring on slightly slower timescales than the plasticization time, and the high-frequency resonance occurring on significantly faster timescales, both exhibit the qualitative trends apparent from Eq. (15) upon variation of the model parameters. This provides an experimental handle to influence the resonant driving frequency.

We further probe the origin of these two resonances by investigating the temporal evolution of the fraction of field-aligned dipolar mesogens $f^2$ [43], which indicates that in the low-frequency regime the dipolar mesogens follow the electric field. That is, for each cycle of the electric field, the dipolar mesogens completely reorient to align with the field when its magnitude is high, and fully relax when the field strength passes through zero. This behavior is combined with
the dynamics of the free volume order parameter $\eta$, which describes how the volume dynamically “wraps itself around” the space that is freed up by mesogen reorientation, to yield the resonance. Figure 5(c) schematically illustrates the described low-frequency resonance dynamics, where we show only the response of the mesogen orientation to the electric field for visual clarity. This visualizes the low-frequency resonance dynamics of the order parameter $f^2$, but not that of the free volume order parameter $\eta$.

Conversely, the high-frequency regime corresponds to frequencies too fast for the dipolar mesogens to follow. Instead, upon turning on the electric field, the dipolar mesogens start to reorient to align with the field. However, before the reorientation is complete, the electric field has already decreased in magnitude significantly and passes through zero, and the dipolar mesogens relax briefly in response to this. Then, as the oscillation continues, the electric field strength increases again, and in response the dipolar mesogens reorient a bit further than on the previous cycle. This “pumping” of the dipolar mesogens by the electric field continues until a steady state of reorientation and (partial) relaxation is achieved. Again, the resonance results from a combination of this behavior with the dynamics of the free volume order parameter $\eta$.

Figure 5(c) schematically illustrates the high-frequency resonance dynamics, where we again show only the response of the mesogen reorientation to the electric field, which visualizes the dynamics of the order parameter $f^2$, for visual clarity.

To more strongly link these different mechanisms to experimentally verifiable predictions, we subsequently perform a simultaneous sweep over both electric field strength and driving frequency, as Fig. 6 shows. Focusing first on Fig. 6(a), which shows a range of driving frequencies encompassing the resonances for both mechanisms, we again recover two regimes. At low field strengths low-frequency actuation yields the greatest free volume generation, whereas at high field strengths high-frequency actuation yields optimal results. We remark that Liu et al. have performed similar sweeps in their MD simulations, shown in Fig. 6(b) [35], which show the same separation into two distinct regimes as a function of field strength. This suggests that the competition between two different mechanisms for free volume generation also plays an important role in the MD simulations.

Figure 6(c), alternatively, shows the theoretical prediction for the same sweep but over a narrower range of driving frequencies. As for a sufficiently narrow range of driving frequencies one effectively probes only a single mechanism, the result shows a single, monotonic regime. The trend we observe here is in line with experimental results reported by Liu et al., shown in Fig. 6(d), suggesting that the experimentally used range of driving frequencies is too narrow to observe the different mechanisms for free volume generation that we predict. Note, in addition, that a qualitative comparison between Figs. 6(c) and 6(d) allows for a rough mapping of the model onto experimentally used electric field strengths.

Thus, from an experimental point of view the model predicts, on the one hand, that broadening the range of driving frequencies gives rise to nonmonotonic behavior as a function of the driving frequency. This leads to two distinct regimes dominated by different physics, namely reorientation and complete relaxation of the dipolar mesogens at low electric field strengths and “pumping” of the dipolar mesogens at high electric field strengths. On the other hand, the model, in accordance with MD simulations, suggests that increasing the electric field strength leads to saturation of the free volume generation at a given driving frequency.

VII. CONCLUSION AND OUTLOOK

In summary, we have constructed a Landau-type theory based on the principle of excluded-volume-driven free volume generation, which shows strong qualitative agreement with our own but also previous MD simulations. In particular, we show that under the influence of a static electric field the orientational order of both cross-linked and dipolar mesogens saturates gradually, and that this gives rise to a maximum in the associated free-volume curve. Subsequently extending the model to include dynamics, we find that the most important model features are captured in terms of the reorientation of the dipolar mesogens, expressed through the population fraction $f^2$, and the response of the free volume to this reorientation, expressed through the free volume order parameter $\eta$. Although a “stripped-down” version of the model based solely on these two order parameters, as detailed in Appendix B, fails to reproduce the aforementioned equations of state, it...
is sufficient to explain the predictions regarding dynamics discussed above in the context of the full theory. To sum these up, we predict: (i) the existence of a plasticization time associated with macroscopic volume gain; a finding reinforced by experiments [36], (ii) the inverse dependence of this plasticization time on the liquid crystalline mesogen aspect ratio, the cross-linking fraction and the initial orientational order cross-linked into the network, (iii) the existence of two resonances upon AC actuation of the LCN, which correspond to two competing mechanisms that operate at different frequencies and dominate at different electric field strengths, and (iv) the proportionality of both resonance frequencies to the liquid crystalline mesogen aspect ratio, the cross-linking fraction and the initial orientational order cross-linked into the network. Comparing our Landau-theoretical framework with the results reported by Liu et al. [35], we can (v) rationalize their observation of two distinct regimes in the MD simulation data, (vi) postulate that the absence of a second regime in their experiments is due to a limited range of driving frequencies, and (vii) predict that the AC response of the LCN saturates at high electric field strengths. Accordingly, we propose follow-up experiments in which the liquid crystalline mesogen aspect ratio, cross-linking fraction and the initial orientational order cross-linked into the network are varied, and the range of driving frequencies and field strengths widened. This would not only allow for a verification of the theory, but also potentially provide concrete experimental handles to optimize the design of LCNs in industrial applications, paving the way toward LCN-based devices in the fields of haptic feedback, self-cleaning surfaces and pattern formation.

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APPENDIX A: SCALING PROCEDURE

Throughout this paper we use the following scaling procedure. Although the scalings used for the static model are also given in the main text, here we briefly reiterate these for ease of reference. First, we introduce the substitutions

\[ s_i = S_i / S_+ , \quad i = 0, 1, 2, 3 \]
\[ a = A / A_+ , \]
\[ h = H / \overline{H} , \]
\[ \tilde{\eta} = \eta / (\xi B^2 / B_0 C^2) , \quad (A1) \]

where \( S_+ = B / 2C \) and \( A_+ = B^2 / 4C \) denote the spinodal of the nematic liquid crystal, and \( \overline{H} = B^3 / 27C^2 \) is the critical field strength. In addition, we scale the parameters \( \tilde{k} \equiv k / (B^2 / C) \), \( \tilde{\xi} \equiv \xi^2 / B_0 C \), and \( \tilde{\lambda} \equiv \lambda / (B^2 / C) \), which represent the magnitude of the effective coupling to the polymer network, the coupling to the free volume and the excluded-volume-like coupling terms. Finally, we scale the parameter \( \tilde{B}_1 \equiv B_1 / (B^4 / C^3) \), associated with the bulk-modulus-like term in \( f^2 \). As noted in the main text, we choose the phenomenological constant \( \alpha = 2s_0 - s_0^2 \) such that the initial configuration \( s_1 = s_2 = s_3 = s_0, f^2 = 0 \) and \( \eta = 0 \) coincides with the free-energy minimum in the absence of an electric field, \( h = 0 \). This eliminates \( \alpha \) as a parameter in our model in favor of \( s_0 \), and leaves us, for a given field strength \( h \), with the set of parameters \( k, \tilde{\xi}, \lambda, \tilde{B}_1 \), and \( s_0 \) characterizing the equilibrium properties of the model. The free-energy density \( g \) of Eq. (1) accordingly becomes \( g = \tilde{g}(\tilde{k}, \tilde{\xi}, \lambda, \tilde{B}_1, s_0) B^2 / C^3 \), with \( \tilde{g} \) a dimensionless free-energy density.

If we subsequently consider the mean-field dynamics as a function of the time \( \tau \), described by Eq. (14), reprinted here for ease of reference,

\[ \partial_\tau \psi = -\Gamma_\psi \left[ \frac{\partial \tilde{g}}{\partial \psi} + \gamma \eta \delta \psi, \eta \right] , \quad (A2) \]

with \( (\tau)_{\psi} \), the ensemble-averaged time the free volume has existed for and \( \delta \psi, \eta \), the Kronecker delta, then we introduce a new set of parameters: the kinetic coefficients \( \Gamma_\psi \), with \( \psi = S_1, S_2, S_3, f, \eta \), and the phenomenological constant \( \gamma \), which represents the viscoelastic relaxation of the LCN. Starting with the kinetic coefficients \( \Gamma_\psi \), we can choose a convenient scaling by recognizing that all order parameter scalings are of the form \( \psi = \psi / \psi_{\text{ref}} \), with \( \psi_{\text{ref}} = S_+, S_+, S_+, 1, \xi B^2 / B_0 C^2 \) indicating the relevant scaling. Writing the nondimensionalized free-energy density \( \tilde{g} = g C^3 / B^4 \) and scaling the time \( \tilde{\tau} = \tau / \tau_0 \) with some as of yet unspecified reference time \( \tau_0 \), then allows us to recast the set of dynamical Eq. (A2) in the dimensionless form

\[ \partial_\tilde{\tau} \tilde{\psi} = -\tau_0 \Gamma_\psi \frac{B^4}{\psi_{\text{ref}}^S} \left[ \frac{\partial \tilde{g}}{\partial \psi} + \gamma \eta \tilde{\eta} \tilde{\tau}_0 \frac{C^3}{B^4} \left( \tilde{\tau} \right) \delta \tilde{\psi}, \eta \right] . \quad (A3) \]

From Eq. (A3), we read off the effective kinetic coefficients \( \tilde{\Gamma}_\psi = \tau_0 \Gamma_\psi B^4 / \psi_{\text{ref}}^S C^3 \) that apply to the scaled order parameters \( \tilde{\psi} \). This suggests a sensible choice for the reference time is to set \( \tau_0 = \tilde{\xi}^2 / \Gamma_\psi B_0^2 C \), such that we have \( \tilde{\Gamma}_\psi = 1 \), i.e., we measure time relative to the dynamics of the scaled free volume order parameter \( \tilde{\eta} \). This leaves us with the following set of scaled kinetic coefficients:

\[ \tilde{\Gamma}_{S_i} = (\Gamma_S / \Gamma_\eta) / (C^2 B_0^2 / 4B^2 \tilde{\xi}^2) , \]
\[ \tilde{\Gamma}_{S_2} = (\Gamma_S / \Gamma_\eta) / (C^2 B_0^2 / 4B^2 \tilde{\xi}^2) , \]
\[ \tilde{\Gamma}_{S_3} = (\Gamma_S / \Gamma_\eta) / (C^2 B_0^2 / 4B^2 \tilde{\xi}^2) , \]
\[ \tilde{\Gamma}_f = (\Gamma_f / \Gamma_\eta) / (C^4 B_0^2 / B^4 \tilde{\xi}^2) , \]
\[ \tilde{\Gamma}_\eta = 1 . \quad (A4) \]

We remark that this timescale is identically used to scale the driving frequency \( \tilde{\omega} = \omega \tau_0 \) for AC actuation, and allows us to also evaluate the scaled phenomenological constant \( \tilde{\gamma} = \gamma / (\Gamma_\psi B_0^2 C / \tilde{\xi}^4) \), which completes the scaling procedure.

APPENDIX B: “MINIMAL” LANDAU–DE GENNES MODEL

1. “Minimal” model

The most intuitive understanding of our theory can be reached by considering a “minimal” model that contains the most important physics, reflects some of the most salient
results, but lacks the more subtle aspects of the full model needed to fully reconcile the theory with simulations and experiments. As alluded to in the main text, the crucial model ingredients are mesogen reorientational and the response of the volume to this reorientation. This suggests a “minimal” description of our model can be achieved as a function of only the population order parameter, \( f^2 \), and the free volume order parameter, \( \eta \), by fixing all scalar nematic order parameters to their as-prepared values \( S_1 = S_2 = S_3 = S_0 \). This significantly simplifies the model free-energy density (per unit reference volume) to read
\[
g - g_0 = f^2 \left( \frac{3}{2} \lambda S_0 - H \right) S_0 + \frac{1}{2} B_1 f^4 + \frac{1}{2} B_0 \eta^2 - 3 \xi S_0^2 \eta f^2, \tag{B1}\]
with \( g_0 \) an unimportant, for our purposes, reference value of the free energy encompassing the frozen-in degrees of freedom of our model. As discussed in the main text, \( \lambda \) and \( \xi \) are phenomenological coupling coefficients, and \( B_1 \) and \( B_0 \) are bulk-modulus-like coefficients. Note that in writing Eq. (B1), we have already reduced the number of model parameters to four, excluding the as-prepared scalar nematic order parameter \( S_0 \) and the electric field strength \( H \).

Crucially, the free-energy Eq. (B1) can be minimized analytically, resulting in the equilibrium solution \( f^2 = \eta = 0 \) for \( H < \frac{3}{2} \lambda S_0 \), and
\[
f^2 = S_0 \left( \frac{H - \frac{3}{2} \lambda S_0}{B_1 - 9 \xi^2 S_0^2 / B_0} \right), \quad \eta = 3 \xi S_0^3 \left( \frac{H - \frac{3}{2} \lambda S_0}{B_1 B_0 - 9 \xi^2 S_0^2} \right), \tag{B2}\]
for \( H > \frac{3}{2} \lambda S_0 \). This immediately makes apparent there exists a critical field strength \( H_{\text{crit}} = \frac{3}{2} \lambda S_0 \) below which there is no response to the electric field; this is identical to what we found in the main text. In addition, the field strength at which the population fraction saturates, \( f^2 = 1 \), evaluates to
\[
H_{\text{sat}} = \frac{3}{2} \lambda S_0 + \frac{B_0 B_1 - 9 \xi^2 S_0^2}{B_0 S_0}. \tag{B3}\]
Here, the first term clearly represents the critical field strength, whereas the second term comprises of both the additional field strength needed to saturate \( f^2 \) in the absence of a free-volume coupling, as well as the free-volume correction.

The above characteristics can be straightforwardly related to the free volume as well, by remarking that Eq. (B2) shows that the equilibrium behavior of \( \eta \) is enslaved to that of \( f^2 \), effectively decreasing the free-energy penalty associated with increasingly large electric-field-aligned populations of dipolar mesogens, \( f^2 > 0 \).

The observant reader will notice that—if the system is in thermodynamic equilibrium—the above allows us to “integrate out” the free volume degree of freedom. This yields a free-energy density expressed entirely in terms of the population fraction \( f^2 \), according to
\[
g \rightarrow g_{\text{ref},0} = f^2 \left( \frac{3}{2} \lambda S_0 - H \right) S_0 + \frac{1}{2} \left( B_1 - \frac{9 \xi^2 S_0^2}{B_0} \right) f^4, \tag{B4}\]
where the coefficient \( B_1 \) is now renormalized to account for free-volume effects. We remark that Eq. (B4) maps our “minimal” model onto the Landau theory for a simple Ising ferromagnet: the population fraction \( f^2 \) plays the role of the (squared) magnetization, and the electric field strength \( H \) plays the role of the temperature. Although our “minimal” model is a bit more complicated than this—we must still constrain the population fraction \( 0 > f^2 > 1 \), and it only makes physical sense to resolve the “squared magnetization” \( f^2 \) in lieu of the “magnetization” \( f \)—the analogy is not entirely unexpected; the phase behavior of a paranematic liquid crystal, if expanded around the critical field strength, can also be mapped exactly onto the Ising ferromagnet [60].

Despite the qualitative insight that Eq. (B4) provides into the equilibrium behavior of the “minimal” model, it is only valid under the express assumption of thermodynamic equilibrium, and so unsuitable for describing dynamics; the main focus of our work. As a result, to further reduce the parameter space, and with it the complexity, of the “minimal” model, the only avenue left to us is scaling the theory, identical to the procedure carried out in the main text (see Appendix A), to yield
\[
g \rightarrow g_0 = f^2 \left( \frac{3}{4} \lambda S_0 - \frac{1}{27} \eta \right) S_0 + \frac{1}{3} B_1 f^4 + \frac{1}{2} \xi S_0^2 \eta f^2. \tag{B5}\]
Here, the newly introduced tildes and lowercase symbols denote scaled quantities, and \( \xi \) replaces \( \eta \) and \( B_0 \) as a measure for the coupling strength between the orientational order of the mesogens and the LCN volume. The reader will note that the scaled model now features only two order parameters, \( f^2 \) and \( \eta \), three model parameters, \( \lambda, B_1, \xi \), as well as the (scaled) as-prepared scalar nematic order parameter, \( s_0 \), and the (scaled) electric field strength, \( \eta \). We remark that if we investigate dynamics using the “minimal” model, we must also add two scaled kinetic coefficients, \( \Gamma_f \) and \( \Gamma_\eta \), as well as the scaled phenomenological coefficient \( \gamma \), which addresses free-volume relaxation, to this list. Analogous to the procedure carried out in the main text, the corresponding mean-field dynamical equations read
\[
\partial_t f = -\Gamma_f \left[ \frac{\partial (\tilde{g} - \tilde{g}_0)}{\partial f} \right], \tag{B6}\]
\[
\partial_t \eta = -\Gamma_\eta \left[ \frac{\partial (\tilde{g} - \tilde{g}_0)}{\partial \eta} + \gamma \tilde{\eta} \langle \tau \rangle_{\tilde{g}} \right],
\]
with \( \bar{t} \) the (scaled) time and \( \langle \tau \rangle_{\tilde{g}} \) the ensemble-averaged (scaled) time (the (scaled) free volume has exist for). In what follows we detail the results that can be extracted from this scaled “minimal” theory, as compared with our full theory.

2. Results

To illustrate to what extent the “minimal” model captures our most important results, here we reproduce Figs. 3–6 of the main text based on the free-energy Eq. (B5). We discuss these figures sequentially below.

Figure 7 (cf. Fig. 3 in the main text) shows the equations of state of the “minimal” model (left), as compared with our MD simulations (right). As discussed above, the “minimal” model captures both the critical field strength and the saturation field strength predicted by the full theory. However, as can be seen from the Fig. 7(a), the “minimal” model lacks any response of the cross-linked mesogens (dashed curve), as well as the
gradual saturation of the orientational order of the dipolar mesogens (solid line), shown by the full theory and our MD simulations.

Moreover, both the full theory [Fig. 3(c) in the main text] and our MD simulations [Fig. 7(d)] predict the equilibrium volume of the LCN to decrease upon increasing the electric field strength in the high-field-strength regime where the orientational order saturates. This feature, which Fig. 7(d) shows is absent from the “minimal” model, yields a maximum in the free volume curve and results from the LCN becoming increasingly aligned along the electric-field axis, rather than becoming increasingly orientationally disordered. We remark that in the “minimal” model neither can occur beyond the saturation field strength, as we have frozen the degrees of freedom associated with the scalar nematic order parameters $S_1$, $S_2$, and $S_3$.

It is worth pointing out that fixing the scalar nematic order parameters $S_1 = S_2 = S_3 = 0$ only increases the free volume in the manner described above at high electric field strengths, where the electric-field-aligned dipolar mesogens tend to dominate the global degree of orientational disorder. At low electric field strengths (but above the critical field strength) the reverse effect occurs, as the slight changes in the scalar nematic order parameters that would be induced by the electric field tend to increase the global degree of orientational disorder in this regime. The “minimal” model misses out on this potential free volume gain by freezing the associated degrees of freedom.

Figure 8 (cf. Fig. 4 in the main text) shows the volume increase of the LCN as a function of time for the “minimal” model (a) and our MD simulations (b). The dashed line (a) indicates a short-time approximation obtained by linearizing the theory. Theory: we denote the scaled free volume order parameter $\tilde{\eta}$, the scaled time $\tilde{t} = t/(|\varepsilon|^2/\Gamma_1 B_0^2 C)$ and we apply a scaled electric field strength $\sqrt{\eta} = 18$. See Appendix A for the full scaling procedure for these, as well as for our model parameters. Parameter values used: $\tilde{\lambda} = 0.4$, $\tilde{\xi} = 0.03$, $\tilde{B}_1 = 5.0$, $\tilde{s}_0 = 3.0$, $\delta/f(0)^2 = 10^{-16}$ for the initial fraction of field-aligned dipolar mesogens, $\gamma = 0.001$ for the free volume relaxation parameter, $\Gamma_\eta = 0.01$ and $\Gamma_\eta = 1.0$ for the scaled kinetic coefficients. Simulations: we work in terms of fundamental Lennard-Jones units, at fixed temperature $T = 0.35$ and fixed pressure $p = 1.0$, with time steps $dt = 0.001$ and electric field strength $E = 30.0$; see the end of Sec. III in the main text for further details. (To be compared with Fig. 4 in the main text; the range plotted here for the scaled free volume order parameter $\tilde{\eta}$ is larger than in the main text.)

Figure 9 (cf. Fig. 5 in the main text) illustrates the response of the LCN as a function of the driving frequency of an alternating electric field $H \propto (|E| \cos 2\pi \omega t)^2$, with $\omega$ the driving frequency and $\tau$ the time. Figures 9(a) and 9(b) indicate that the “minimal” model, like the full theory, exhibits two resonance frequencies: a low-frequency resonance dominant at low electric field strengths (blue) and a high-frequency resonance dominant at high electric field strengths (red).
already anticipated and alluded to this in the main text by noting that the resonances are dictated by two coupled processes: mesogen reorientation and the response of the volume to this reorientation; these results support this claim.

In addition, as is the case for the full theory, the two resonance frequencies correspond to two distinct mechanisms for free volume generation, namely reorientation and complete relaxation of the dipolar mesogens [Fig. 9(c)] at low electric field strengths and “pumping” of the dipolar mesogens [Fig. 9(d)] at high electric field strengths.

Finally, it is understood that Fig. 9(a) depicts a frequency sweep at relatively low electric field strength, \( \sqrt{\eta} \approx 7.1 \), hence why the values shown for the free volume order parameter are slightly lower than those reported in the main text.

Figure 10 (cf. Fig. 6 in the main text) shows simultaneous sweeps over both the driving frequency and the electric field strength for the “minimal” model (a), (c), and for the MD simulations (b) and experiments (d) reported by Liu et al. [35].

Focusing first on Figs. 10(a) and 10(b), we conclude that both the “minimal” model and the MD simulations exhibit two distinct regimes: at low electric field strengths low driving frequencies yield the greatest volume change, whereas at high electric field strengths high driving frequencies yield the greatest volume change. This is consistent with the results obtained from the full theory, reported in the main text, and suggests that a competition between two distinct mechanisms also underlies the MD simulations.

Next, we investigate Figs. 10(c) and 10(d), which show a narrower frequency range for the “minimal” model [Fig. 10(c)], as compared with the experimental results of Liu et al. [Fig. 10(d)] [35]. Just as with the full theory discussed in the main text, the “minimal” model exhibits a single monotonic regime when probed on a sufficiently narrow frequency range, similar to the experiments shown in Fig. 10(d). Accordingly, the “minimal” model supports the hypothesis that the experimentally used range of driving frequencies is too narrow to observe the full behavior predicted by theory and simulations.

Another point of note for Fig. 10 is the saturation of the free volume at high electric field strengths apparent from

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FIG. 9. Resonance frequencies for steady-state free volume gain as a function of electric field strength (a), (b) and the corresponding microscopic mechanisms (c), (d). (a) Steady-state gain in the scaled free volume order parameter \( \tilde{\eta} \) as a function of the scaled driving frequency \( \tilde{\omega} = \omega |\sqrt{2/\Gamma_0 B_0^2 C}| \), for a scaled electric field strength \( \sqrt{\eta} = 7 \); the colored symbols denote resonance frequencies. (b) Scaled resonance frequencies \( \tilde{\omega} \) as a function of the scaled electric field \( \sqrt{\eta} \propto |E| \) in the regimes of low electric field strength (blue circles) and high electric field strength (red squares). The filled symbols correspond to the field strength used in panel (a). The black curve denotes the numerically evaluated plasticization time at the same electric field, with the corresponding estimate Eq. (15) (in the main text) indicated with the dashed line. See Appendix A for the full scaling procedure and see Fig. 8 for the used parameter values. Simulations are in terms of fundamental Lennard-Jones units, at fixed temperature \( T = 0.35 \) and fixed pressure \( p = 1 \), with time steps \( dt = 0.001 \), total simulation time \( t_{\text{sim}} = 10.0 \) and the driving frequency \( f \) in terms of \((100 \times dt)^{-1}\); see the end of Sec. III in the main text and Ref. [35] for further details. Panels (b) and (d) reprinted from Ref. [35] under a Creative Commons Attribution 4.0 International License http://creativecommons.org/licenses/by/4.0/. (To be compared with Fig. 6 in the main text; the range plotted here for the scaled free volume order parameter \( \tilde{\eta} \) is larger than in the main text.)

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FIG. 10. Steady-state free volume gain as a function of electric field strength and driving frequency for “minimal” theory (a), (c), MD simulations (b) and experiments (d). Theory: we denote the scaled free volume order parameter \( \tilde{\eta} \), the scaled driving frequency \( \tilde{\omega} = \omega |\sqrt{2/\Gamma_0 B_0^2 C}| \) and the scaled electric field strength \( \sqrt{\eta} \propto |E| \). See Appendix A for the full scaling procedure and see Fig. 8 for the used parameter values. Simulations are in terms of fundamental Lennard-Jones units, at fixed temperature \( T = 0.35 \) and fixed pressure \( p = 1 \), with time steps \( dt = 0.001 \), total simulation time \( t_{\text{sim}} = 10.0 \) and the driving frequency \( f \) in terms of \((100 \times dt)^{-1}\); see the end of Sec. III in the main text and Ref. [35] for further details. Panels (b) and (d) reprinted from Ref. [35] under a Creative Commons Attribution 4.0 International License http://creativecommons.org/licenses/by/4.0/. (To be compared with Fig. 6 in the main text; the range plotted here for the scaled free volume order parameter \( \tilde{\eta} \) is larger than in the main text.)
the “minimal” model [Figs. 10(a) and 10(c)] and the MD simulations [Fig. 10(b)]. This is again consistent with the full theory discussed in the main text, and supports the prediction that the experimentally measured modulation will saturate at electric field strengths higher than those shown in Fig. 10(d).

Finally, the range of values plotted for the scaled free volume order parameter in Fig. 10 is larger than that shown in the main text, because the “minimal” model predicts a greater free volume gain than the full theory for high electric field strengths, as discussed above.

3. Conclusion

In conclusion, we summarize the salient features of the MD simulations and the experiments, the predictions we provide in the main text, and which of these we can reproduce using the “minimal” model (for the remainder, the full theory is required). Starting with the response of the LCN to a static electric field, our MD simulations show that (i) the orientational order of both the cross-linked and the dipolar mesogens saturates gradually as a function of the electric field strength, and that (ii) this gives rise to a maximum in the associated free-volume curve. The “minimal” model fails to capture either of these features.

Upon investigating the dynamical response of the LCN, the “minimal model does, however, capture (iii) the existence of a plasticization time associated with macroscopic volume gain, which is observed in experiments but absent from our MD simulations. As is the case for the full theory, the “minimal” model provides an analytic estimate for this plasticization time [which remains unchanged, and is given by Eq. (15) in the main text], allowing us to predict that (iv) the plasticization timescales inversely with the liquid crystalline mesogen aspect ratio, the cross-linking fraction and the initial orientational order cross-linked into the network.

Under the influence of an oscillating electric field, the “minimal” model also predicts (v) the existence of two resonances, which correspond to two competing mechanisms that operate at different frequencies and dominate at different electric field strengths, and (vi) the proportionality of both resonance frequencies to the liquid crystalline mesogen aspect ratio, the cross-linking fraction and the initial orientational order cross-linked into the network. This is in accordance with the full theory.

Comparing our “minimal” model to the MD simulations reported by Liu et al. [35] [see Figs. 10(a) and 10(b)], we reproduce (vii) their observation of two distinct regimes, which we can rationalize in terms of the aforementioned competing mechanisms. Moreover, upon narrowing down the range of driving frequencies, we are left with only a single regime, in accordance with their experimental results [see Figs. 10(c) and 10(d)]. This leads us to postulate that (viii) the absence of a second regime in their experiments is due to a limited range of driving frequencies. Finally, (ix) the MD simulations by Liu et al. show a saturation of the free volume under the influence of an oscillating electric field, which our “minimal” model reproduces. This leads us to predict that (x) the AC response of the LCN saturates at high electric field strengths in experiments as well.

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the analogous expression would yield the proper behavior upon minimisation with respect to all order parameters.

If one were to write down the energy of an induced dipole due to the inversion symmetry of the nematic director $\mathbf{n} \rightarrow -\mathbf{n}$. Although for a mesogen with permanent dipole moment $\mathbf{p}$ the analogous expression would read $-\mathbf{p} \cdot \mathbf{E}$, averaging this over an ensemble of mesogens using a Boltzmann distribution this can again be written in the form $-\langle \mathbf{p} \cdot \mathbf{E} \rangle \propto |\mathbf{E}|^2$ for sufficiently weak electric fields [27,61,62].

If one were to write down the Gibbs free-energy density $\mathcal{G} = G/V$, with $G$ the total Gibbs free energy, then thermodynamic equilibrium would demand $\partial \mathcal{G} / \partial \eta = -G/(1 + \eta)$. This means the Gibbs free-energy density must take the form $\mathcal{G} = g/(1 + \eta)$, with $\partial g / \partial \eta = 0$. This indicates that, if one wishes to treat $\eta$ as a proper order parameter, the Gibbs free energy per unit reference volume, $g = G/V_0$, is the relevant thermodynamic potential. After all, one explicitly constructs the Landau theory to yield the proper behavior upon minimisation with respect to all order parameters.

The form $H \propto |\mathbf{E}|^2$ is appropriate for describing the free energy of an induced dipole due to the inversion symmetry of the nematic director $\mathbf{n} \rightarrow -\mathbf{n}$. Although for a mesogen with permanent dipole moment $\mathbf{p}$ the analogous expression would yield $-\mathbf{p} \cdot \mathbf{E}$, averaging this over an ensemble of mesogens using a Boltzmann distribution this can again be written in the form $-\langle \mathbf{p} \cdot \mathbf{E} \rangle \propto |\mathbf{E}|^2$ for sufficiently weak electric fields [27,61,62].