1

Liquid Crystal Physics

1.1 Introduction

Liquid crystals are mesophases between crystalline solid and isotropic liquid [1–3]. The constituents are elongated rod-like (calamitic) or disk-like (discotic) organic molecules as shown in Figure 1.1. The size of the molecules is typically a few nanometers (nm). The ratio between the length and the diameter of the rod-like molecules or the ratio between the diameter and the thickness of disk-like molecules is about 5 or larger. Because the molecules are non-spherical, besides positional order, they may possess orientational order.

Figure 1.1(a) shows a typical calamitic liquid crystal molecule. Its chemical name is 4′-n-Pentyl-4-cyano-biphenyl and is abbreviated as 5CB [4,5]. It consists of a biphenyl, which is the rigid core, and a hydrocarbon chain which is the flexible tail. The space-filling model of the molecule is shown in Figure 1.1(c). Although the molecule itself is not cylindrical, it can be regarded as a cylinder, as shown in Figure 1.1(e), in considering its physical behavior, because of the fast rotation (on the order of $10^{-9}$ s) around the long molecule axis due to thermal motion. The distance between two carbon atoms is about 1.5 Å; therefore the length and the diameter of the molecule are about 2 nm and 0.5 nm, respectively. The molecule shown has a permanent dipole moment (from the CN head), but it can still be represented by the cylinder whose head and tail are the same, because in non-ferroelectric liquid crystal phases, the dipole has equal probability of pointing up or down. It is necessary for a liquid crystal molecule to have a rigid core(s) and flexible tail(s). If the molecule is completely flexible, it will not have orientational order. If it is completely rigid, it will transform directly from isotropic liquid phase at high temperature to crystalline solid phase at low temperature. The rigid part favors both orientational and positional order while the flexible part disfavors them. With balanced rigid and flexible parts, the molecule exhibits liquid crystal phases.
Figure 1.1(b) shows a typical *discotic* liquid crystal molecule [6]. It also has a rigid core and flexible tails. The branches are approximately on one plane. The space-filling model of the molecule is shown in Figure 1.1(d). If there is no permanent dipole moment perpendicular to the plane of the molecule, it can be regarded as a disk in considering its physical behavior as shown in Figure 1.1(f) because of the fast rotation around the axis which is at the center of the molecule and perpendicular to the plane of the molecule. If there is a permanent dipole moment perpendicular to the plane of the molecule, it is better to visualize the molecule as a bowl, because the reflection symmetry is broken and all the permanent dipoles may point in the same direction and spontaneous polarization occurs. The flexible tails are also necessary, otherwise the molecules form a crystal phase where there is positional order.

Figure 1.1 Calamitic liquid crystal: (a) chemical structure, (c) space-filling model, (e) physical model. Discotic liquid crystal: (b) chemical structure, (d) space-filling mode, (f) physical model.
The variety of phases that may be exhibited by rod-like molecules are shown in Figure 1.2. At high temperature, the molecules are in the isotropic liquid state where they do not have either positional or orientational order. The molecules can easily move around, and the material can flow like water. The translational viscosity is comparable to that of water. Both the long and short axes of the molecules can point in any direction.

When the temperature is decreased, the material transforms into the nematic phase, which is the most common and simplest liquid crystal phase, where the molecules have orientational order but still no positional order. The molecules can still diffuse around, and the translational viscosity does not change much from that of the isotropic liquid state. The long axis of the molecules has a preferred direction. Although the molecules still swivel due to thermal motion, the time-averaged direction of the long axis of a molecule is well defined and is the same for all the molecules at macroscopic scale. The average direction of the long molecular axis is denoted by \( \hat{n} \) which is a unit vector called the liquid crystal director. The short axes of the molecules have no orientational order in a uniaxial nematic liquid crystal.

When the temperature is decreased further, the material may transform into the Smectic-A phase where, besides the orientational order, the molecules have partial positional order, i.e., the molecules form a layered structure. The liquid crystal director is perpendicular to the layers. Smectic-A is a one-dimensional crystal where the molecules have positional order in the layer normal direction. The cartoon shown in Figure 1.2 is schematic. In reality, the separation between neighboring layers is not as well defined as that shown by the cartoon. The molecule number density exhibits an undulation with the wavelength about the molecular length. Within a layer, it is a two-dimensional liquid crystal in which there is no positional order, and the molecules can move around. For a material in poly-domain smectic-A, the translational viscosity is significantly higher, and it behaves like a grease. When the temperature is decreased further, the material may transform into the smectic-C phase, where the liquid crystal director is no longer perpendicular to the layer but tilted.

At low temperature, the material is in the crystal solid phase where there are both positional and orientational orders. The translational viscosity becomes infinitely high and the molecules (almost) do not diffuse anymore.

Liquid crystals get the ‘crystal’ part of their name because they exhibit optical birefringence as crystalline solids. They get the ‘liquid’ part of their name because they can flow and do not support shearing as regular liquids. Liquid crystal molecules are elongated and have different
molecular polarizabilities along their long and short axes. Once the long axes of the molecules orient along a common direction, the refractive indices along and perpendicular to the common direction are different. It should be noted that not all rod-like molecules exhibit all the liquid crystal phases. They may exhibit some of the liquid crystal phases.

Some of the liquid crystal phases of disk-like molecules are shown in Figure 1.3. At high temperature, they are in the isotropic liquid state where there are no positional and orientational orders. The material behaves in the same way as a regular liquid. When the temperature is decreased, the material transforms into the nematic phase, which has orientational order but not positional order. The average direction of the short axis perpendicular to the disk is oriented along a preferred direction, which is also called the liquid crystal director and denoted by a unit vector $\mathbf{n}$. The molecules have different polarizabilities along a direction in the plane of the disk and along the short axis. Thus the discotic nematic phase also exhibits birefringence as crystals.

When the temperature is decreased further, the material transforms into the columnar phase where, besides orientational order, there is partial positional order. The molecules stack up to form columns. Within a column, it is a liquid where the molecules have no positional order. The columns, however, are arranged periodically in the plane perpendicular to the columns. Hence it is a two-dimensional crystal. At low temperature, the material transforms into the crystalline solid phase where the positional order along the columns is developed.

The liquid crystal phases discussed so far are called thermotropic liquid crystals and the transitions from one phase to another are driven by varying temperature. There is another type of liquid crystals, called lyotropic liquid crystals, exhibited by molecules when they are mixed with a solvent of some kind. The phase transitions from one phase to another phase are driven by varying the solvent concentration. Lyotropic liquid crystals usually consist of amphiphilic molecules that have a hydrophobic group at one end and a hydrophilic group at the other end and the water is the solvent. The common lyotropic liquid crystal phases are micelle phase and lamellar phase. Lyotropic liquid crystals are important in biology. They will not be discussed in this book because the scope of this book is on displays and photonic devices.

Liquid crystals have a history of more than 100 years. It is believed that the person who discovered liquid crystals is Friedrich Reinitzer, an Austrian botanist [7]. The liquid crystal phase observed by him in 1888 was a cholesteric phase. Since then, liquid crystals have come a long way and become a major branch of interdisciplinary sciences. Scientifically, liquid crystals are important because of the richness of structures and transitions. Technologically, they have won tremendous success in display and photonic applications [8–10].

**Figure 1.3** Schematic representation of the phases of disk-like molecules.
1.2 Thermodynamics and Statistical Physics

Liquid crystal physics is an interdisciplinary science: thermodynamics, statistical physics, electrodynamics, and optics are involved. Here we give a brief introduction to thermodynamics and statistical physics.

1.2.1 Thermodynamic laws

One of the important quantities in thermodynamics is entropy. From the microscopic point of view, entropy is a measurement of the number of quantum states accessible to a system. In order to define entropy quantitatively, we first consider the fundamental logical assumption that for a closed system (no energy and particles exchange with other systems), quantum states are either accessible or inaccessible to the system, and the system is equally likely to be in any one of the accessible states as in any other accessible state [11]. For a macroscopic system, the number of accessible quantum states $g$ is a huge number ($\approx 10^{23}$). It is easier to deal with $\ln g$, which is defined as the entropy $\sigma$:

$$\sigma = \ln g$$ (1.1)

If a closed system consists of subsystem 1 and subsystem 2, the numbers of accessible states of the subsystems are $g_1$ and $g_2$, respectively. The number of accessible quantum states of the whole system is $g = g_1 g_2$ and the entropy is $\sigma = \ln g = \ln(g_1 g_2) = \ln g_1 + \ln g_2 = \sigma_1 + \sigma_2$.

Entropy is a function of the energy $u$ of the system $\sigma = \sigma(u)$. The second law of thermodynamics states that for a closed system, the equilibrium state has the maximum entropy. Let us consider a closed system which contains two subsystems. When two subsystems are brought into thermal contact (energy exchange between them is allowed), the energy is allocated to maximize the number of accessible states, that is, the entropy is maximized. Subsystem 1 has the energy $u_1$ and entropy $\sigma_1$; subsystem 2 has the energy $u_2$ and entropy $\sigma_2$. For the whole system, $u = u_1 + u_2$ and $\sigma = \sigma_1 + \sigma_2$. The first law of thermodynamics states that energy is conserved, that is, $u = u_1 + u_2 = \text{constant}$. From the second law of thermodynamics, for any process inside the closed system, $\delta u = \delta u_1 + \delta u_2 = 0$. From the first law of thermodynamics, for any process, we have $\delta \sigma = \delta \sigma_1 + \delta \sigma_2 \geq 0$. When the two subsystems are brought into thermal contact, at the beginning, energy flows. For example, an amount of energy $|\delta u_1|$ flows from subsystem 1 to subsystem 2, $\delta u_1 < 0$ and $\delta u_2 = -\delta u_1 > 0$, and $\frac{\partial \sigma}{\partial u_2} = \frac{\partial \sigma_1}{\partial u_2} + \frac{\partial \sigma_2}{\partial u_2} = \frac{\partial \sigma_1}{\partial u_2} + \frac{\partial \sigma_2}{\partial u_2} \geq 0$. When equilibrium is reached, the entropy is maximized and $\frac{\partial \sigma_1}{\partial u_1} - \frac{\partial \sigma_2}{\partial u_2} = 0$, that is, $\frac{\partial \sigma_1}{\partial u_1} = \frac{\partial \sigma_2}{\partial u_2}$. We know that when two systems reach equilibrium, they have the same temperature. Accordingly the fundamental temperature $\tau$ is defined by

$$1/\tau = \left(\frac{\partial \sigma}{\partial u}\right)_{N,V}.$$ (1.2)

Energy flows from a high temperature system to a low temperature system. The conventional temperature (Kelvin temperature) is defined by

$$T = \tau/k_B.$$ (1.3)
where \( k_B = 1.381 \times 10^{-23} \text{ Joule/Kelvin} \) is the Boltzmann constant. Conventional entropy \( S \) is defined by

\[
\frac{1}{T} = \frac{\partial S}{\partial u}.
\]

Hence

\[
S = k_B \sigma.
\]

### 1.2.2 Boltzmann Distribution

Now we consider the thermodynamics of a system at a constant temperature, that is, in thermal contact with a thermal reservoir. The temperature of the thermal reservoir (named B) is \( \tau \). The system under consideration (named A) has two states with energy 0 and \( \epsilon \), respectively. A and B form a closed system, and its total energy \( u = u_A + u_B = u_o = \text{constant} \). When A is in the state with energy 0, B has the energy \( u_o \), the number of accessible states:

\[
g_1 = g_A \times g_B = 1 \times g_B(u_o) = g_B(u_o).
\]

When A has the energy \( \epsilon \), B has the energy \( u_o - \epsilon \), the number of accessible states is

\[
g_2 = g_A \times g_B = 1 \times g_B(u_o - \epsilon) = g_B(u_o - \epsilon).
\]

For the whole system, the total number of accessible states is

\[
G = g_1 + g_2 = g_B(u_o) + g_B(u_o - \epsilon).
\]

\((A + B)\) is a closed system, and the probability in any of the \( G \) states is the same. When the whole system is in one of the \( g_1 \) states, A has the energy 0. When the whole system is in one of the \( g_2 \) states, A has the energy \( \epsilon \). Therefore the probability for A in the state with energy 0 is

\[
P(0) = \frac{g_1}{g_1 + g_2} = \frac{g_B(u_o)}{g_B(u_o) + g_B(u_o - \epsilon)}.
\]

The probability for A in the state with energy \( \epsilon \) is

\[
P(\epsilon) = \frac{g_2}{g_1 + g_2} = \frac{g_B(u_o - \epsilon)}{g_B(u_o) + g_B(u_o - \epsilon)}.
\]

From the definition of entropy, we have

\[
g_B(u_o) = e^{\sigma_B(u_o)}
\]

and

\[
g_B(u_o - \epsilon) = e^{\sigma_B(u_o - \epsilon)}.
\]

Because \( \epsilon \ll u_o \), \( \sigma_B(u_o - \epsilon) \approx \sigma_B(u_o) \approx \frac{\partial \sigma_B}{\partial u_B} \epsilon = \sigma_B(u_o) - \frac{1}{\tau} \epsilon \). Therefore we have

\[
P(0) = \frac{e^{\sigma_B(u_o)}}{e^{\sigma_B(u_o)} + e^{\sigma_B(u_o) - \epsilon/\tau}} = \frac{1}{1 + e^{-\epsilon/\tau k_B T}} \]

\[
P(\epsilon) = \frac{e^{\sigma_B(u_o) - \epsilon/\tau}}{e^{\sigma_B(u_o)} + e^{\sigma_B(u_o) - \epsilon/\tau}} = \frac{e^{-\epsilon/\tau}}{1 + e^{-\epsilon/\tau k_B T}}
\]

\[
\frac{P(\epsilon)}{P(0)} = e^{-\epsilon/k_B T}.
\]

For a system having \( N \) states with energies \( \epsilon_1, \epsilon_2, \ldots, \epsilon_i, \epsilon_{i+1}, \ldots, \epsilon_N \), the probability for the system in the state with energy \( \epsilon_i \) is

\[
P(\epsilon_i) = e^{-\epsilon_i/\tau} \sum_{j=1}^{N} e^{-\epsilon_j/k_B T}.
\]
The *partition function* of the system is defined as

\[ Z = \sum_{i=1}^{N} e^{-\epsilon_i/k_BT}. \]  

(1.11)

The *internal energy* (average energy) of the system is given by

\[ U = \langle \epsilon \rangle = \sum_i \epsilon_i P(\epsilon_i) = \frac{1}{Z} \sum_i \epsilon_i e^{-\epsilon_i/k_BT}. \]  

(1.12)

Because \( \frac{\partial Z}{\partial T} = \sum_i \left( \frac{\epsilon_i}{k_BT^2} \right) e^{-\epsilon_i/k_BT} = \frac{1}{k_BT^2} \sum_i \epsilon_i e^{-\epsilon_i/k_BT} \),

\[ U = \frac{k_BT^2}{Z} \frac{\partial Z}{\partial T} = k_BT^2 \frac{\partial (\ln Z)}{\partial T}. \]  

(1.13)

### 1.2.3 Thermodynamic quantities

As energy is conserved, the change of the internal energy \( U \) of a system equals the heat \( dQ \) absorbed and the mechanical work \( dW \) done to the system, \( dU = dQ + dW \). When the volume of the system changes by \( dV \) under the pressure \( P \), the mechanical work done to the system is given by

\[ dW = -PdV. \]  

(1.14)

When there is no mechanical work, the heat absorbed equals the change of internal energy. From the definition of temperature \( 1/T = (\frac{\partial S}{\partial U})_V \), the heat absorbed in a reversible process at constant volume is

\[ dU = dQ = TdS. \]  

(1.15)

When the volume is not constant, then

\[ dU = TdS - PdV. \]  

(1.16)

The derivatives are

\[ T = \left( \frac{\partial U}{\partial S} \right)_V, \]  

(1.17)

\[ P = -\left( \frac{\partial U}{\partial V} \right)_S. \]  

(1.18)
The internal energy $U$, entropy $S$, and volume $V$ are extensive quantities, while temperature $T$ and pressure $P$ are intensive quantities. The enthalpy $H$ of the system is defined by

$$H = U + PV. \quad (1.19)$$

Its variation in a reversible process is given by

$$dH = dU + d(PV) = (TdS - PdV) + (PdV + VdP) = TdS + VdP. \quad (1.20)$$

From this equation, it can be seen that the physical meaning of enthalpy is that in a process at constant pressure ($dP = 0$), the change of enthalpy $dH$ is equal to the heat absorbed $dQ = (TdS)$. The derivatives of the enthalpy are

$$T = \left( \frac{\partial H}{\partial S} \right)_P, \quad (1.21)$$

$$V = \left( \frac{\partial H}{\partial P} \right)_S. \quad (1.22)$$

The Helmholtz free energy $F$ of the system is defined by

$$F = U - TS. \quad (1.23)$$

Its variation in a reversible process is given by

$$dF = dU - d(TS) = (TdS - PdV) - (TdS + SdT) = -SdT - PdV. \quad (1.24)$$

The physical meaning of Helmholtz free energy is that in a process at constant temperature ($dT = 0$), the change of Helmholtz free energy is equal to the work done to the system. The derivatives are

$$S = - \left( \frac{\partial F}{\partial T} \right)_V, \quad (1.25)$$

$$P = - \left( \frac{\partial F}{\partial V} \right)_T. \quad (1.26)$$

The Gibbs free energy $G$ of the system is defined by

$$G = U - TS + PV. \quad (1.27)$$

The variation in a reversible process is given by

$$dG = dU - d(TS) - d(PV) = -SdT + VdP. \quad (1.28)$$
In a process at constant temperature and pressure, Gibbs free energy does not change. The derivatives are

\[ S = -\left( \frac{\partial G}{\partial T} \right)_P, \quad (1.29) \]

\[ V = \left( \frac{\partial G}{\partial P} \right)_T. \quad (1.30) \]

The Helmholtz free energy can be derived from the partition function. Because of Equations (1.13) and (1.25),

\[ F = U - TS = k_B T^2 \frac{\partial (\ln Z)}{\partial T} + T \left( \frac{\partial F}{\partial T} \right)_V \]

\[ F - T \left( \frac{\partial F}{\partial T} \right)_V = -T^2 \left\{ \frac{1}{T} \left( \frac{\partial F}{\partial T} \right)_V + F \left[ \frac{\partial (\ln Z)}{\partial T} \right]_V \right\} = -T^2 \left[ \frac{\partial (\ln Z)}{\partial T} \right]_V = k_B T^2 \frac{\partial (\ln Z)}{\partial T}. \]

Hence

\[ F = -k_B T \ln Z = -k_B T \ln \left( \sum_i e^{-\epsilon_i/k_B T} \right). \quad (1.31) \]

From Equations (1.11), (1.25) and (1.31), the entropy of a system at a constant temperature can be calculated:

\[ S = -k_B < \ln \rho > = -k_B \sum_i \rho_i \ln \rho_i \quad (1.32) \]

### 1.2.4 Criteria for thermodynamical equilibrium

Now we consider the criteria which can used to judge whether a system is in its equilibrium state under given conditions. We already know that for a closed system, as it changes from a non-equilibrium state to the equilibrium state, the entropy increases,

\[ \delta S \geq 0. \quad (1.33) \]

It can be stated in a different way that for a closed system the entropy is maximized in the equilibrium state.

In considering the equilibrium state of a system at constant temperature and volume, we construct a closed system which consists of the system (subsystem 1) under consideration and a thermal reservoir (subsystem 2) with the temperature \( T \). When the two systems are brought into thermal contact, energy is exchanged between subsystem 1 and subsystem 2. Because the whole system is a closed system, \( \delta S = \delta S_1 + \delta S_2 \geq 0 \). For system 2, \( 1/T = \left( \frac{\partial S_2}{\partial U_2} \right)_V \), and
therefore $\delta S_2 = \delta U_2 / T$ (this is true when the volume of subsystem is fixed, which also means that the volume of subsystem 1 is fixed). Because of energy conservation, $\delta U_2 = - \delta U_1$. Hence $\delta S = \delta S_1 + \delta S_2 = \delta S_1 + \delta U_2 / T = \delta S_1 - \delta U_1 / T \geq 0$. Because the temperature and volume are constant for subsystem 1, $\delta S_1 - \delta U_1 / T = (1/T) \delta (TS_1 - U_1) \geq 0$, and therefore

$$\delta (U_1 - TS_1) = \delta F_1 \leq 0.$$  \hspace{1cm} (1.34)

At constant temperature and volume, the equilibrium state has the minimum Helmholtz free energy.

In considering the equilibrium state of a system at constant temperature and pressure, we construct a closed system which consists of the system (subsystem 1) under consideration and a thermal reservoir (subsystem 2) with the temperature $T$. When the two systems are brought into thermal contact, energy is exchanged between subsystem 1 and subsystem 2. Because the whole system is a closed system, $\delta S = \delta S_1 + \delta S_2 \geq 0$. For subsystem 2, because the volume is not fixed, and mechanical work is involved, $\delta U_2 = T \delta S_2 - P \delta V_2$, that is, $\delta S_2 = (\delta U_2 + P \delta V_2) / T$. Because $\delta U_2 = - \delta U_1$ and $\delta V_2 = - \delta V_1$, $\delta S = \delta S_1 + (\delta U_2 + P \delta V_2) / T = \delta S_1 - (\delta U_1 + P \delta V_1) / T = (1/T) \delta (TS_1 - U_1 - PV_1) \geq 0$. Therefore

$$\delta (U_1 + PV_1 - TS_1) = \delta G_1 \leq 0.$$  \hspace{1cm} (1.35)

At constant temperature and pressure, the equilibrium state has the minimum Gibbs free energy. If electric energy is involved, then we have to consider the electric work done to the system by external sources such as a battery. In a thermodynamic process, if the electric work done to the system is $dW_e$, $\delta S \geq \frac{dQ}{T} - \frac{dU - dW_m - dW_e}{T} = \frac{dU + PdV - dW_e}{T}$. Therefore at constant temperature and pressure

$$\delta (U - W_e + PV - TS) = \delta (G - W_e) \leq 0.$$  \hspace{1cm} (1.36)

In the equilibrium state, $G - W_e$ is minimized.

1.3 Orientational Order

Orientational order is the most important feature of liquid crystals. The average directions of the long axes of the rod-like molecules are parallel to each other. Because of the orientational order, liquid crystals possess anisotropic physical properties, that is, in different directions they have different responses to external fields such as electric field, magnetic field and shear. In this section, we will discuss how to specify quantitatively orientational order and why rod-like molecules tend to parallel each other.

For a rigid elongated liquid crystal molecule, three axes can be attached to it to describe its orientation. One is the long molecular axis and the other two axes are perpendicular to the long molecular axis. Usually the molecule rotates fast around the long molecular axis. Although the molecule is not cylindrical, if there is no hindrance in the rotation in nematic phase, the fast rotation around the long molecular axis makes it behave as a cylinder. There is no preferred direction for the short axes and thus the nematic liquid crystal is usually uniaxial. If there is
hindrance in the rotation, the liquid crystal is biaxial. Biaxial nematic liquid crystal is a long-sought material. A lyotropic biaxial nematic phase has been observed [12]. A thermotropic biaxial nematic phase is still debatable, and it may exist in systems consisting of bent-core molecules [13,14]. Also the rotation symmetry around the long molecular axis can be broken by confinements. In this book, we deal with uniaxial liquid crystals consisting of rod-like molecules unless otherwise specified.

1.3.1 Orientational order parameter

In uniaxial liquid crystals, we have only to consider the orientation of the long molecular axis. The orientation of a rod-like molecule can be represented by a unit vector \( \hat{a} \) which is attached to the molecule and parallel to the long molecular axis. In the nematic phase, the average directions of the long molecular axes are along a common direction: the liquid crystal director denoted by the unit vector \( \hat{n} \). The orientation of \( \hat{a} \) in 3-D can be specified by the polar angle \( \theta \) and the azimuthal angle \( \phi \) where the \( z \) axis is chosen parallel to \( \hat{n} \) as shown in Figure 1.4. In general, the orientational order of \( \hat{a} \) is specified by an orientational distribution function \( f(\theta, \phi) \).

\[
f(\theta, \phi) \, d\Omega = \sin \theta \, d\theta \, d\phi
\]

is the probability that \( \hat{a} \) orients along the direction specified by \( \theta \) and \( \phi \) within the solid angle \( d\Omega \). In isotropic phase, \( \hat{a} \) has equal probability of pointing any direction and therefore \( f(\theta, \phi) = \text{constant} \). For uniaxial liquid crystals, there is no preferred orientation in the azimuthal direction, and then \( f = f(\theta) \), which depends only on the polar angle \( \theta \).

Rod-like liquid crystal molecules may have permanent dipole moments. If the dipole moment is perpendicular to the long molecule axis, the dipole has equal probability of pointing along any direction because of the fast rotation around the long molecular axis in uniaxial liquid crystal phases. The dipoles of the molecules cannot generate spontaneous polarization. If the permanent dipole moment is along the long molecular axis, the flip of the long molecular axis is much slower (of the order of \( 10^{-5} \) s); the above argument does not hold. In order to see the orientation of the dipoles in this case, we consider the interaction between two dipoles [15]. When one dipole is on top of the other dipole, if they are parallel, the interaction energy is low and thus parallel orientation is preferred. When two dipoles are side by side, if they are anti-parallel, the interaction energy is low and thus anti-parallel orientation is preferred. As we know, the molecules cannot penetrate each other. For elongated molecules, the distance between the two dipoles when on top of each other is farther than that when the two dipoles...

Figure 1.4 Schematic diagram showing the orientation of the rod-like molecule.
are side by side. The interaction energy between two dipoles is inversely proportional to the cubic power of the distance between them. Therefore anti-parallel orientation of dipoles is dominant in rod-like molecules. There are the same number of dipoles aligned parallel to the liquid crystal director \( \mathbf{\hat{n}} \) as the number of dipoles aligned anti-parallel to \( \mathbf{\hat{n}} \). The permanent dipole along the long molecular axis cannot generate spontaneous polarization. Thus even when the molecules have permanent dipole moment along the long molecule axes, they can be regarded as cylinders whose top and end are the same. It can also be concluded that \( \mathbf{\hat{n}} \) and \( -\mathbf{\hat{n}} \) are equivalent.

An order parameter must be defined in order to specify quantitatively the orientational order. The order parameter is usually defined in such a way that it is zero in the high temperature unordered phase and non-zero in the low temperature ordered phase. By analogy with ferromagnetism, we may consider the average value of the projection of \( \mathbf{\hat{a}} \) along the director \( \mathbf{\hat{n}} \):

\[
< \cos \theta > = \frac{1}{\pi} \int_0^\pi \cos \theta f(\theta) \sin \theta d\theta = \frac{1}{\pi} \int_0^\pi f(\theta) \sin \theta d\theta,
\]

where \(< \> \) indicates the average (temporal and spatial averages are the same), and \( \cos \theta \) is the first Legendre polynomial. In isotropic phase, the molecules are randomly oriented, \(< \cos \theta > \) is zero. We also know that in nematic phase the probabilities that the molecule orients at the angles \( \theta \) and \( \pi - \theta \) are the same, that is, \( f(\theta) = f(\pi - \theta) \), therefore \(< \cos \theta > = 0 \), and is not a good choice for the orientational order parameter. Next let us try the average value of the second Legendre polynomial for the order parameter:

\[
S = < P_2(\cos \theta) > = < \frac{1}{2} (3 \cos^2 \theta - 1) > = \frac{1}{\pi} \int_0^\pi (3 \cos^2 \theta - 1) f(\theta) \sin \theta d\theta = \frac{1}{\pi} \int_0^\pi f(\theta) \sin \theta d\theta
\]

In the isotropic phase, as shown in Figure 1.5(b), \( f(\theta) = c \), a constant. \[\frac{1}{\pi} \int_0^\pi (3 \cos^2 \theta - 1) f(\theta) \sin \theta d\theta = \frac{1}{\pi} \int_0^\pi (3 \cos^2 \theta - 1) c \sin \theta d\theta = 0. \] In nematic phase, \( f(\theta) \) depends on \( \theta \). For a perfectly ordered nematic phase as shown in Figure 1.5(d), \( f(\theta) = \delta(\theta) \), where \( \sin \theta \delta(\theta) = \infty \) when \( \theta = 0 \), \( \sin \theta \delta(\theta) = 0 \) when \( \theta \neq 0 \), and \( \int_0^\pi \delta(\theta) \sin \theta d\theta = 1 \), the order parameter is \( S = (1/2)(3 \cos^2 0 - 1) = 1 \). It should be pointed out that the order parameter can be positive or negative. For two order parameters with the same absolute value but different signs, they correspond to different states. When the molecules all lie in a plane but randomly orient in the plane, as shown in Figure 1.5(a), the distribution function is \( f(\theta) = \delta(\theta - \pi/2) \), where \( \delta(\theta - \pi/2) = \infty \) when \( \theta = \pi/2 \), \( \delta(\theta - \pi/2) = 0 \) when \( \theta \neq \pi/2 \), and \( \int_0^\pi \delta(\theta - \pi/2) \sin \theta d\theta = 1 \), the order
parameter is \( S = (1/2)[3 \cos^2(\pi/2) - 1]/1 = -0.5 \). In this case, the average direction of the molecules is not well defined. The director \( \mathbf{n} \) is defined by the direction of the uniaxial axis of the material. Figure 1.5(c) shows the state with the distribution function \( f(\theta) = (35/16)[\cos^4\theta + (1/35)] \), which is plotted vs. \( \theta \) in Figure 1.5(e). The order parameter is \( S = 0.5 \). Many anisotropies of physical properties are related to the order parameter and will be discussed later.

### 1.3.2 Landau–de Gennes theory of orientational order in nematic phase

Landau developed a theory for second-order phase transition [16], such as from diamagnetic phase to ferromagnetic phase, in which the order parameter increases continuously from zero as the temperature is decreased across the transition temperature \( T_c \) from the high temperature disordered phase to the low temperature ordered phase. For a temperature near \( T_c \), the order is very small. The free energy of the system can be expanded in terms of the order parameter.

The transition from water to ice at 1 atmosphere pressure is a first-order transition, and the latent heat is about 100 J/g. The isotropic–nematic transition is a weak first-order transition because the order parameter changes discontinuously across the transition but the latent heat is only about 10 J/g. De Gennes extended Landau’s theory into isotropic–nematic transition.
because it is a weak first-order transition \[1,17\]. The free energy density \( f \) of the material can be expressed in terms of the order parameter \( S \),

\[
f = \frac{1}{2} a(T-T^*)S^2 - \frac{1}{3} bS^3 + \frac{1}{4} cS^4 + \frac{1}{2} L(\nabla S)^2, \tag{1.39}
\]

where \( a, b, c \) and \( L \) are constants and \( T^* \) is the virtual second-order phase transition temperature. The last term is the energy cost when there is a variation of the order parameter in space, and here we will consider only the uniform order parameter case. There is no linear term of \( S \), which would result in a non-zero order parameter at any temperature; \( a \) is positive, otherwise \( S \) will never be 0 and the isotropic phase will not be stable at any temperature. A significant difference between the free energy here and that of a magnetic system is the cubic term. In a magnetic system, the magnetization \( m \) is the order parameter. For a given value of \( |m| \), there is only one state, and the sign of \( m \) is decided by the choice of the coordinate. The free energy must be the same for a positive \( m \) and a negative \( m \), and therefore the coefficient of the cubic term must be zero. For nematic liquid crystal, positive and negative values of the order parameter correspond to two different states, and the corresponding free energies can be different, and therefore \( b \) must be not zero. \( b \) must be positive because at sufficiently low temperatures positive-order parameters have the global minimum free energies. We also know that the maximum value of \( S \) is 1. The quadratic term with a positive \( c \) prevents \( S \) from exploding. The values of the coefficients can be estimated in the following way: the energy of the intermolecular interaction between molecules associated with orientation is about \( k_BT = 1.38 \times 10^{-23} \text{(J/K)} \times 300 \text{ K} \approx 4 \times 10^{-21} \text{ J} \) and the molecular size is about 1 nm, \( f \) is the energy per unit volume, and therefore \( Ta \) (or \( b \) or \( c \)) \(- k_BT \text{volume of one molecule} \approx 4 \times 10^{-21} \text{ joule/(10}^{-9}\text{m})^3 \approx 1.6 \text{J/m}^3 \).

For a given temperature, the order parameter \( S \) is found by minimizing \( f \),

\[
\frac{\partial f}{\partial S} = a(T-T^*)S - bS^2 + cS^3 = [a(T-T^*) - bS + cS^2]S = 0. \tag{1.40}
\]

There are three solutions:

\[
S_1 = 0,
\]

\[
S_2 = \frac{1}{2c} \left[ b + \sqrt{b^2 - 4ac(T-T^*)} \right],
\]

\[
S_3 = \frac{1}{2c} \left[ b - \sqrt{b^2 - 4ac(T-T^*)} \right].
\]

\( S_1 = 0 \) corresponds to the isotropic phase and the free energy is \( f_1 = 0 \). The isotropic phase has the global minimum free energy at high temperature. It will be shown that at low temperature \( S_2 \) has the global minimum free energy \( f_2 = \frac{1}{2} a(T-T^*)S_2^2 - \frac{1}{3} bS_2^3 + \frac{1}{4} cS_2^4 \). And \( S_3 \) has a local maximum free energy. At the isotropic–nematic phase transition temperature \( T_{NI} \), the order parameter is \( S_c = S_{2c} \), and \( f_2(S_2 = S_c) = f_1 = 0 \), that is,

\[
\frac{1}{2} a(T_{NI}-T^*)S_c^2 - \frac{1}{3} bS_c^3 + \frac{1}{4} cS_c^4 = 0. \tag{1.41}
\]
From Equation (1.40), at this temperature, we also have

\[ a(T_{NI} - T^*) - bS_c + cS_c^2 = 0. \quad (1.42) \]

From the two equations above, we can obtain

\[ a(T_{NI} - T^*) - \frac{1}{3} bS_c = 0. \]

Therefore

\[ S_c = \frac{3a}{b} (T_{NI} - T^*). \quad (1.43) \]

Substitute Equation (1.43) into Equation (1.42), we will get the transition temperature

\[ T_{NI} = T^* + \frac{2b^2}{9ac}, \quad (1.44) \]

and the order parameter at the transition temperature

\[ S_c = \frac{2b}{3c}, \quad (1.45) \]

For liquid crystal 5CB, the experimentally measured order parameter is shown by the solid circles in Figure 1.6(a) [6]. In fitting the data, the following parameters are used: \( a = 0.023 \sigma \text{ J/K} \cdot \text{m}^3 \), \( b = 1.2 \sigma \text{ J/m}^3 \) and \( c = 2.2 \sigma \text{ J/m}^3 \), where \( \sigma \) is a constant which has to be determined by latent heat of the isotropic–nematic transition.

Because \( S \) is a real number in the region from \(-0.5\) to \(1.0\), when \( T - T^* > b^2/4ac \), that is, when \( T - T_{NI} > b^2/4ac - 2b^2/9ac = b^2/36ac \), \( S_2 \) and \( S_3 \) are not real. The only real solution is \( S = S_1 = 0 \), corresponding to the isotropic phase. When \( T - T_{NI} < b^2/36ac \), there are three solutions. However, when \( 0 < T - T_{NI} \leq b^2/36ac \), the isotropic phase is the stable state because its free energy is still the global minimum, as shown in Figure 1.6(b). When \( T - T_{NI} \leq 0 \), the nematic phase with the order parameter \( S = S_2 = \left( b + \sqrt{b^2 - 4ac(T - T^*)} \right)/2c \) is the stable state because its free energy is the global minimum.

In order to see clearly the physical meaning, let us plot \( f \) vs. \( S \) at various temperatures as shown in Figure 1.7. First we consider what occurs with decreasing temperature. At temperature \( T_1 = T_{NI} + b^2/36ac + 1.0^\circ\text{C} \), the curve has only one minimum at \( S = 0 \), which means that \( S_1 = 0 \) is the only solution, and the corresponding isotropic phase is the stable state. At temperature \( T_3 = T_{NI} + b^2/36ac - 0.5^\circ\text{C} \), there are two local minima and one local maximum, where there are three solutions: \( S_1 = 0 \), \( S_2 > 0 \), and \( S_3 > 0 \). Here, \( S_1 = 0 \) corresponds to the global minimum and the isotropic phase is still the stable state. At \( T_4 = T_{NP} \), the free energies of the isotropic phase with the order parameter \( S_1 \) and the nematic phase with the order parameter \( S_2 \) become the same; phase transition takes place and the order parameter changes discontinuously from \( 0 \) to \( S_c = 2b/3c \). This is a first-order transition. It can be seen from the figure that at this
temperature there is an energy barrier between $S_1$ and $S_2$. The height of the energy barrier is $b^4/81c^3$. If the system is initially in the isotropic phase and there are no means to overcome the energy barrier, it will remain in the isotropic phase at this temperature. As the temperature is decreased, the energy barrier is lowered. At $T_5 = T_{NI} - 3^\circ C$, the energy barrier is lower. At $T_6 = T^*$, the second-order derivative of $f$ with respect to $S$ at $S_1 = 0$ is

$$\left. \frac{\partial^2 f}{\partial S^2} \right|_{S=0} = a(T - T^*) - 2bS + 3cS^2 \bigg|_{S=0} = a(T - T^*) = 0.$$ 

$S_1$ is no longer a local minimum and the energy barrier disappears. $T^*$ is therefore the supercooling temperature below which the isotropic phase becomes absolutely unstable. At this temperature, $S_1 = S_3$. At $T_7 = T^* - 2^\circ C$, there are two minima located at $S_2(>0)$ and $S_3(<0)$ (the minimum value is slightly below 0), and a maximum at $S_1 = 0$.

**Figure 1.6** (a) The three solutions of order parameter as a function of temperature, (b) the corresponding free energies as a function of temperature, in Landau–de Gennes theory.
Now we consider what occurs with increasing temperature. If initially the system is in the nematic phase, it will remain in this phase even at temperatures higher than \( T_{NI} \) and its free energy is higher than that of the isotropic phase, because there is an energy barrier preventing the system to transform from the nematic phase to the isotropic phase. The temperature \( T_2 \) (superheating temperature) at which the nematic phase becomes absolutely unstable can be found by

\[
\frac{\partial^2 f}{\partial S^2} \bigg|_{S_2} = a(T_2 - T^*) - 2bS_2 + 3cS_2^2 = 0.
\]

Using \( S_2 = \frac{1}{2c} \left[ b + \sqrt{b^2 - 4ac(T_2 - T^*)} \right] \), we can get \( T_2 = T_{NI} + \frac{b^2}{36ac} \).

In reality, there are usually irregularities, such as impurities and defects, which can reduce the energy barrier against the isotropic–nematic transition. The phase transition takes place before the thermodynamic instability limits (supercooling or superheating temperature). Under an optical microscope, it is usually observed that with decreasing temperature nematic ‘islands’ are initiated by irregularities and growing out the isotropic ‘sea’ and with increasing temperature isotropic ‘lakes’ are produced by irregularities and grow on the nematic ‘land.’ The irregularities are called nucleation seeds and the transition is a nucleation process. In summary, the nematic–isotropic transition is a first-order transition; the order parameter changes discontinuously; there is an energy barrier against the transition and the transition is a nucleation process; there are superheating and supercooling. In second-order transition, there is no energy barrier and the transition occurs simultaneously everywhere at the transition temperature (the critical temperature).

There are a few points worth mentioning in Landau–de Gennes theory. It works well at temperatures near the transition. At temperatures far below the transition temperature, the order parameter increases without limit with decreasing temperature, and the theory does not work well because we know that the maximum order parameter should be 1. In Figure 1.6, the
parameters are chosen in such a way that the fitting is good for a relatively wide range of temperatures. \( T_{NI} - T^* = 2b^2/\theta ac = 6.3^\circ C \), which is much larger than the value (~1 \(^\circ C\)) measured by light scattering experiments in isotropic phase [18]. There are fluctuations in orientational order in the isotropic phase, which results in a variation of refractive index in space and causes light scattering. The intensity of the scattering light is proportional to \( 1/(T - T^*) \).

### 1.3.3 Maier–Saupe theory

In the nematic phase, there are interactions, such as the Van der Waals interaction, between the liquid crystal molecules. Because the molecular polarizability along the long molecular axis is larger than that along the short transverse molecular axis, the interaction is anisotropic and results in the parallel alignment of the rod-like molecules. In the spirit of the mean field approximation, Maier and Saupe introduced an effective single molecule potential \( V \) to describe the intermolecular interaction [19,20]. The potential has the following properties. (1) It must be a minimum when the molecule orients along the liquid crystal director (the average direction of the long molecular axis of the molecules). (2) Its strength is proportional to the order parameter \( S = <P_2(\cos \theta)> \) because the potential well is deep when the molecules are highly orientationally ordered and vanishes when the molecules are disordered. (3) It ensures that the probabilities for the molecules pointing up and down are the same. The potential in Maier–Saupe theory is given by

\[
V(\theta) = -\nu S \left( \frac{3}{2} \cos^2 \theta - \frac{1}{2} \right),
\]

where \( \nu \) is an interaction constant of the order of \( k_BT \) and \( \theta \) is the angle between the long molecular axis and the liquid crystal director as shown in Figure 1.4. The probability \( f \) for the molecule orienting along the direction with the polar angle \( \theta \) is governed by the Boltzmann distribution:

\[
f(\theta) = e^{-V(\theta)/k_BT} \int_0^\pi e^{-V(\theta)/k_BT} \sin \theta d\theta
\]

The single molecule partition function is

\[
Z = \int_0^\pi e^{-V(\theta)/k_BT} \sin \theta d\theta.
\]

From the orientational distribution function we can calculate the order parameter:

\[
S = \frac{1}{Z} \int_0^\pi P_2(\cos \theta) e^{-V(\theta)/k_BT} \sin \theta d\theta = \frac{1}{Z} \int_0^\pi P_2(\cos \theta) e^{SP_2(\theta)/k_BT} \sin \theta d\theta
\]
Introduce a normalized temperature $\tau = k_B T / v$. For a given value of $\tau$, the order parameter $S$ can be found by numerically solving Equation (1.50). An iteration method can be used in the numerical calculation of the order parameter: (1) use an initial value for the order parameter, (2) substitute into the right side of Equation (1.50), and (3) calculate the order parameter. Use the newly obtained order parameter to repeat the above process until a stable value is obtained. As shown in Figure 1.8(a), there are three solutions: $S_1$, $S_2$, and $S_3$. In order to determine which is the actual solution, we have to examine the corresponding free energies. The free energy $F$ has two parts: $F = U - T E_n$, where $U$ is the intermolecular interaction energy and $E_n$ is the

**Figure 1.8**  (a) The three solutions of order parameter as a function of the normalized temperature in Maier–Saupe theory. The solid circles represent the experimental data. (b) The normalized free energies of the three solutions of the order parameter.
entropy. The single molecular potential describes the interaction energy between one liquid crystal molecule and the rest of the molecules of the system. The interaction energy of the system with \( N \) molecules is given by

\[
U = \frac{1}{2} N \langle V \rangle = \frac{N}{2Z} \int_0^\pi V(\theta) e^{-V(\theta)/k_BT} \sin \theta d\theta
\]  

where the factor 1/2 avoids counting the intermolecular interaction twice. The entropy is calculated by using Equation (1.32):

\[
S_n = -Nk_B \ln f = - \frac{Nk_B}{Z} \ln [f(\theta)] e^{-V(\theta)/k_BT} \sin \theta d\theta
\]

From Equation (1.48) we have

\[
\ln[f(\theta)] = -\frac{V(\theta)}{k_BT} - \ln Z, \quad \text{and therefore} \quad S_n = \frac{N}{T} \langle V \rangle + Nk_B \ln Z \quad \text{and the free energy is}
\]

\[
F = U - TEn = -Nk_BT \ln Z - \frac{1}{2} N \langle V \rangle.
\]  

From Equations (1.47) we have \( \langle V \rangle = -V(\theta)/k_BT - \ln Z \) and therefore

\[
F = U - TEn = -Nk_BT \ln Z + \frac{1}{2} N\nu S^2
\]  

Although the second term of the above equation looks abnormal, this equation is correct and can be checked by calculating the derivative of \( F \) with respect to \( S \):

\[
\frac{\partial F}{\partial S} = -Nk_BT \frac{\partial \ln Z}{\partial S} - \frac{1}{2} N \frac{\partial \langle V \rangle}{\partial S} = - \frac{Nk_BT}{Z} \frac{\partial \ln Z}{\partial S} + N\nu S
\]

Letting \( \frac{\partial F}{\partial S} = 0 \), we have

\[
S = \frac{k_BT}{\nu Z} \frac{\partial Z}{\partial S} = \frac{k_BT}{\nu Z} \int_0^\pi -\frac{1}{k_BT} \frac{\partial V}{\partial \theta} e^{-V(\theta)/k_BT} \sin \theta d\theta = \int_0^\pi P_2(\cos \theta) e^{vS \theta} \sin \theta d\theta,
\]

which is consistent with Equation (1.50). The free energies corresponding to the solutions are shown in Figure 1.8(b). The nematic–isotropic phase transition temperature is \( \tau_{NI} = 0.22019 \). For temperature higher than \( \tau_{NI} \), the isotropic phase with the order parameter \( S = S_1 = 0 \) has lower free energy and thus is stable. For temperature lower than \( \tau_{NI} \), the nematic phase with the order parameter \( S = S_2 \) has lower free energy and thus is stable. The order parameter jumps from 0 to \( S_2 = 0.4289 \) at the transition.

In the Maier–Saupe theory there are no fitting parameters. The predicted order parameter as a function of temperature is universal, and agrees qualitatively – but not quantitatively – with
experimental data. This indicates that higher-order terms are needed in the single molecule potential, that is,

$$V(\theta) = \sum_i \left[ -v_i <P_i(\cos \theta) >P_i(\cos \theta) \right]$$

(1.55)

where $P_i(\cos \theta)$ ($i = 2, 4, 6, \ldots$) are the $i$th-order Legendre polynomial. The fitting parameters are $v_i$. With higher-order terms, better agreement with experimental results can be achieved.

The Maier–Saupe theory is very useful in considering liquid crystal systems consisting of more than one type of molecules, such as mixtures of nematic liquid crystals and dichroic dyes. The interactions between different molecules are different and the constituent molecules have different order parameters.

None of the theories discussed above predicts well the orientational order parameter for temperatures far below $T_{NI}$. The order parameter as a function of temperature is better described by the empirical formula [21]

$$S = \left( 1 - \frac{0.987V^2}{T_{NI}V_{NI}^2} \right)^{0.22},$$

(1.56)

where $V$ and $V_{NI}$ are the molar volumes at $T$ and $T_{NI}$, respectively.

### 1.4 Elastic Properties of Liquid Crystals

In nematic phase, the liquid crystal director $\mathbf{n}$ is uniform in space in the ground state. In reality, the liquid crystal director $\mathbf{n}$ may vary spatially because of confinements or external fields. This spatial variation of the director, called the deformation of the director, costs energy. When the variation occurs over a distance much larger than the molecular size, the orientational order parameter does not change, and the deformation can be described by a continuum theory in analogue to the classic elastic theory of a solid. The elastic energy is proportional to the square of the spatial variation rate.

#### 1.4.1 Elastic properties of nematic liquid crystals

There are three possible deformation modes of the liquid crystal director as shown in Figure 1.9. Choose the cylindrical coordinate such that the $z$ axis is parallel to the director at the origin of the coordinate: $\mathbf{n} (0) = \hat{z}$. Consider the variation of the director at an infinite small distance away from the origin. When moving along the radial direction, there are two possible modes of variation: (1) the director tilts toward the radial direction $\hat{\rho}$, as shown in Figure 1.9(a), and (2) the director tilts toward the azimuthal direction $\hat{\phi}$, as shown in Figure 1.9(b). The first mode is called splay, where the director at $(\delta \rho, \phi, z = 0)$ is

$$\mathbf{n} (\delta \rho, \phi, z = 0) = \delta n_\rho (\delta \rho) \hat{\rho} + [1 + \delta n_z (\delta \rho)] \hat{z},$$

(1.57)
where $\delta n_\rho \ll 1$ and $\delta n_z \ll 1$. Because $|\vec{n}|^2 = n_\rho^2 + n_\phi^2 + n_z^2 = (\delta n_\rho)^2 + (1 + \delta n_z)^2 = 1$, therefore $\delta n_z = -(\delta n_\rho)^2/2$, $\delta n_z$ is a higher-order term and can be neglected. The spatial variation rate is $\partial n_\rho / \partial \rho$ and the corresponding elastic energy is

$$f_{\text{splay}} = (1/2)K_{11}(\partial n_\rho / \partial \rho)^2,$$

(1.58)

where $K_{11}$ is the splay elastic constant. The second mode is called twist, where the director at $(\delta \rho, \phi, z = 0)$ is

$$\vec{n}(\delta \rho, \phi, z = 0) = \delta n_\phi(\delta \rho)\hat{\phi} + [1 + \delta n_z(\delta \rho)]\hat{z},$$

(1.59)

where $\delta n_\phi \ll 1$ and $\delta n_z = -(\delta n_\phi)^2/2$, a higher-order term which can be neglected. The spatial variation rate is $\partial n_\phi / \partial \rho$ and the corresponding elastic energy is

$$f_{\text{twist}} = (1/2)K_{22}(\partial n_\phi / \partial \rho)^2$$

(1.60)

where $K_{22}$ is the twist elastic constant.

When moving along the $z$ direction, there is only one possible mode of variation, as shown in Figure 1.9(c), which is called bend. The director at $(\rho = 0, \phi, \delta z)$ is

$$\vec{n}(\rho = 0, \phi, \delta z) = \delta n_\rho(\delta z)\hat{\rho} + [1 + \delta n_z(\delta z)]\hat{z}$$

(1.61)
where $\delta n_\rho \ll 1$ and $\delta n_\phi = -(\delta n_\phi)^2/2$, a higher-order term which can be neglected. Note that when $\rho = 0$, the azimuthal angle is not well defined and we can choose the coordinate such that the director tilts toward the radial direction. The corresponding elastic energy is
\[ f_{\text{bend}} = \frac{1}{2} K_{33} \left( \partial n_\rho / \partial z \right)^2, \tag{1.62} \]

where $K_{33}$ is the bend elastic constant. Because $\delta n_\rho$ is a higher-order term, $\partial n_\rho / \partial \rho \approx 0$ and $\partial n_\rho / \partial \rho = 0$. Recall $\nabla \cdot \vec{n} \big|_{\rho=0, z=0} = (1/\rho) \partial (\rho n_\rho) / \partial \rho + (1/\rho) \partial n_\phi / \partial \phi + \partial n_\rho / \partial z \approx \partial n_\rho / \partial \rho + \partial n_\rho / \partial n_\phi$. Because $\partial n_\rho / \partial \rho$ is finite and $\delta n_\rho \ll 1$, then $\nabla \cdot \vec{n} \big|_{\rho=0, z=0} = \partial n_\rho / \partial \rho$. The splay elastic energy can be expressed as $f_{\text{splay}} = (1/2) K_{11} (\nabla \cdot \vec{n})^2$. Because $n \cdot \nabla \times \vec{n} \big|_{\rho=0, z=0} = (\nabla \times \vec{n})_\rho = (\nabla \times \vec{n})_\phi = \partial n_\rho / \partial z$, the bend elastic energy can be expressed as $f_{\text{bend}} = (1/2) K_{33} (\vec{n} \times \nabla \times \vec{n})^2$. Putting all the three terms together, we get the elastic energy density:
\[ f_{\text{ela}} = \frac{1}{2} K_{11} (\nabla \cdot \vec{n})^2 + \frac{1}{2} K_{22} (\vec{n} \cdot \nabla \times \vec{n})^2 + \frac{1}{2} K_{33} (\vec{n} \times \nabla \times \vec{n})^2 \tag{1.63} \]

This elastic energy is often referred to as the Oseen–Frank energy, and $K_{11}$, $K_{22}$, and $K_{33}$ are referred to as the Frank elastic constants, because of his pioneering work on the elastic continuum theory of liquid crystals [22]. The value of the elastic constants can be estimated in the following way. When a significant variation of the director occurs in a length $L$, the angle between the average directions of the long molecules axes of two neighboring molecules is $a L$, where $a$ is the molecular size. When the average direction of the long molecules axes of two neighboring molecules are parallel, the intermolecular interaction energy between them is a minimum. When the angle between the average directions of the long molecular axes of two neighboring molecules makes the angle of $a L$, the intermolecular interaction energy increases $(a L)^2 u$, where $u$ is the intermolecular interaction energy associated with orientation and is about $k_B T$. The increase of the interaction energy is the elastic energy, that is, $(a L)^2 u = K_{11} (\nabla \vec{n})^2 \times \text{molecular volume} = K_{11} \left( \frac{1}{6} \right)^2 a^3$. Therefore $K_{11} = \frac{u}{a} \approx 1.38 \times 10^{-23} \text{J/K} \times 300 \text{K} / (10^{-9} \text{m}) = 4 \times 10^{-12} \text{N}$. Experiments show that usually the bend elastic constant $K_{33}$ is the largest and twist elastic constant $K_{22}$ is the smallest. As an example, at room temperature the liquid crystal 5CB has these elastic constants: $K_{11} = 0.64 \times 10^{-11} \text{N}$, $K_{22} = 0.3 \times 10^{-11} \text{N}$, and $K_{33} = 1 \times 10^{-11} \text{N}$.

The elastic constants depend on the product of the order parameters of two neighboring molecules. If one of the molecules had the order of 0, the second molecule can orient along any direction with the same inter-molecular interaction energy even if it has non-zero order parameter. Therefore the elastic constants are proportional to $S^2$. When the temperature changes, the order parameter will change and so will the elastic constants.

It is usually adequate to consider the splay, twist, and bend deformations of the liquid crystal director in determining the configuration of the director, except in some cases where the surface-to-volume ratio is high, and a further two terms, called divergence terms (or surface terms), may have to be considered. The elastic energy densities of these terms are given by $f_{33} = -K_{23} \nabla \cdot (\vec{n} \nabla \cdot \vec{n})$ and $f_{33} = -K_{33} \nabla \cdot (\vec{n} \nabla \cdot \vec{n} + \vec{n} \nabla \times \vec{n})$, respectively [23]. The volume integral of these two terms can be changed to surface integral because of the Gauss theorem.
1.4.2 Elastic properties of cholesteric liquid crystals

So far we have considered liquid crystals consisting of molecules with reflection symmetry. The molecules are the same as their mirror images, and are called achiral molecules. 5CB shown in Figure 1.1(a) is an example of an achiral molecule. Now we consider liquid crystals consisting of molecules without reflection symmetry. The molecules are different from their mirror images and are called chiral molecules. One such example is CB15 shown in Figure 1.10(a). It can be regarded as a screw, instead of a rod, considering its physical properties. After considering the symmetry that \( n \) and \(-n\) are equivalent, the generalized elastic energy density is

\[
f_{\text{ela}} = \frac{1}{2} K_{11} (\nabla \cdot n)^2 + \frac{1}{2} K_{22} (n \cdot \nabla \times n + q_o)^2 + \frac{1}{2} K_{33} (n \times \nabla \times n)^2, \tag{1.64}
\]

where \( q_o \) is the chirality, and its physical meaning will be discussed in a moment. Note that \( \nabla \times n \) is a pseudo-vector which does not change sign upon reflection symmetry operation, and \( n \cdot \nabla \times n \) is a pseudo-scalar which changes sign upon reflection symmetry operation. Upon reflection symmetry operation, the elastic energy changes to

\[
f'_{\text{ela}} = \frac{1}{2} K_{11} (\nabla \cdot n)^2 + \frac{1}{2} K_{22} (-n \cdot \nabla \times n + q_o)^2 + \frac{1}{2} K_{33} (-n \times \nabla \times n)^2 \tag{1.65}
\]

If the liquid crystal molecule is achiral, and thus has reflection symmetry, the system does not change and the elastic energy does not change upon reflection symmetry operation. It is required that \( f_{\text{ela}} = f'_{\text{ela}} \), then \( q_o = 0 \). When the liquid crystal is in the ground state with the minimum free energy, \( f_{\text{ela}} = 0 \), and this requires \( \nabla \cdot n = 0 \), \( n \cdot \nabla \times n = 0 \), and \( n \times \nabla \times n = 0 \). This means that in the ground state, the liquid crystal director \( n \) is uniformly aligned along one direction.

If the liquid crystal molecule is chiral, and thus has no reflection symmetry, the system changes upon a reflection symmetry operation. The elastic energy may change. It is no longer required that \( f_{\text{ela}} = f'_{\text{ela}} \), but \( q_o \) may not be zero. When the liquid crystal is in the ground state with the minimum free energy, \( f_{\text{ela}} = 0 \), and this requires \( \nabla \cdot n = 0 \), \( n \cdot \nabla \times n = -q_o \), and \( n \times \nabla \times n = 0 \). A director configuration which satisfies the above conditions is

\[
n_x = \cos(q_o z), \quad n_y = \sin(q_o z), \quad n_z = 0; \tag{1.66}
\]

Figure 1.10  (a) Chemical structure of a typical chiral liquid crystal molecule; (b) physical model of a chiral liquid crystal molecule.
this is schematically shown in Figure 1.11. The liquid crystal director twists in space. This type of liquid crystal is called a cholesteric liquid crystal. The axis around which the director twists is called the helical axis and is chosen to be parallel to the z axis here. The distance $P_o$ over which the director twists $360^\circ$ is called the pitch and is related to the chirality by

$$P_o = \frac{2\pi}{q_o}.$$  \hspace{1cm} (1.67)

Depending on its chemical structure, the pitch of a cholesteric liquid crystal could take any value in the region from a few tenths of a micron to infinitely long. The periodicity of a cholesteric liquid crystal with the pitch $P_o$ is $P_o/2$, because $\hat{n}$ and $-\hat{n}$ are equivalent. Cholesteric liquid crystals are also called chiral nematic liquid crystals and denoted as N*. Nematic liquid crystals can be considered as a special case of cholesteric liquid crystals with an infinitely long pitch.

In practice, a cholesteric liquid crystal is usually obtained by mixing a nematic host and a chiral dopant. The pitch of the mixture is given by

$$P = \frac{1}{(HTP) \cdot x}.$$  \hspace{1cm} (1.68)
where $x$ is the concentration of the chiral dopant and (HTP) is the *helical twisting power* of the chiral dopant, which is mainly determined by the chemical structure of the chiral dopant and depends slightly on the nematic host.

### 1.4.3 Elastic properties of smectic liquid crystals

Smectic liquid crystals possess partial positional orders besides the orientational order exhibited in nematic and cholesteric liquid crystals. Here we only consider the simplest case: smectic-A. The elastic energy of the deformation of the liquid crystal director in smectic-A is the same as in nematic. In addition, the dilatation (compression) of the smectic layer also costs energy, which is given by [23]

$$f_{\text{layer}} = \frac{1}{2} B \left( \frac{d-d_o}{d_o} \right)^2,$$

where $B$ is elastic constant for the dilatation of the layer and is referred to as the Young modulus, $d_o$ and $d$ are the equilibrium layer thickness (the periodicity of the density undulation) and the actual layer thickness of the smectic layer, respectively. The typical value of $B$ is about $10^6 - 10^7$ joule/m$^3$, which is $10^3 - 10^4$ smaller than that in a solid. In a slightly deformed smectic-A liquid crystal, consider a closed loop as shown in Figure 1.12. The total number of layers traversed by the loop is zero, which can be mathematically expressed as $\oint n \cdot dl = 0$. Using the Stokes theorem, we have $\int \nabla \times \tilde{n} \cdot d\tilde{s} = \oint \tilde{n} \cdot d\tilde{l} = 0$. Therefore in smectic-A we have

$$\nabla \times \tilde{n} = 0,$$

which ensures that $\tilde{n} \cdot \nabla \times \tilde{n} = 0$ and $\tilde{n} \times \nabla \times \tilde{n} = 0$. The consequence is that twist and bend deformation of the director are not allowed (because they significantly change the layer thickness and cost too much energy). The elastic energy in a smectic-A liquid crystal is

$$f_{\text{elas}} = \frac{1}{2} K_{11} (\nabla \cdot \tilde{n})^2 + \frac{1}{2} B \left( \frac{d-d_o}{d_o} \right)^2.$$

Some chiral liquid crystals, as temperature is decreased, exhibit the mesophases: isotropic $\rightarrow$ cholesteric $\rightarrow$ smectic-A. Because of the property shown by Equation (1.70), there is no

![Figure 1.12](image_url)  
*Schematic diagram showing the deformation of the liquid crystal director and the smectic layer in the smectic-A liquid crystal.*
spontaneous twist in smectic-A. To express this in another way, the pitch in smectic-A is infinitely long. In the cholesteric phase, as the temperature is decreased toward the cholesteric–smectic-A transition, there is a pretransitional phenomenon that smectic-A order forms in short space-scale and time scale due to thermal fluctuation. This effect causes the pitch of the cholesteric liquid crystal to increase with decreasing temperature and diverges at the transition temperature as shown in Figure 1.13. As will be discussed later, a cholesteric liquid crystal with the pitch $P$ exhibits Bragg reflection at the wavelength $\lambda = \bar{n}P$ where $\bar{n}$ is the average refractive index of the material. If $\lambda = \bar{n}P$ is in the visible light region, the liquid crystal reflects colored light. When the temperature is varied, the color of the liquid crystal changes. Such cholesteric liquid crystals are known as thermochromic cholesteric liquid crystals [24]. As shown in Figure 1.13, the reflected light is in the visible region for temperature in the region $T_1$ to $T_2$. There are liquid crystals with $\Delta T = T_1 - T_2$ about 1 degree. If there are two thermochromic cholesteric liquid crystals with different cholesteric–smectic-A transition temperatures, mixtures with different concentrations of the two components will exhibit color reflection at different temperatures. This is how thermochromic cholesteric liquid crystals are used to make thermometers.

1.5 Response of Liquid Crystals to Electromagnetic Fields

Liquid crystals are anisotropic dielectric and diamagnetic media [1,25]. Their resistivities are very high ($\sim 10^{10} \Omega \cdot \text{cm}$). Dipole moments are induced in them by external fields. They have different dielectric permittivities and magnetic susceptibilities along the directions parallel to and perpendicular to the liquid crystal director.

1.5.1 Magnetic susceptibility

We first consider magnetic susceptibility. Because the magnetic interaction between the molecules is weak, the local magnetic field on the molecules is approximately the same as the externally applied magnetic field. For a uniaxial liquid crystal, the molecule can be regarded as a cylinder. When a magnetic field $\vec{H}$ is applied to the liquid crystal, it has different responses
to the applied field, depending on the angle between the long molecular axis $\vec{a}$ and the field $\vec{H}$. The magnetic field can be decomposed into a parallel component and a perpendicular component, as shown in Figure 1.14. The magnetization $\vec{M}$ is given by

$$\vec{M} = N\kappa / (\vec{a} \cdot \vec{H}) \vec{a} + N\kappa_\perp \left[ \vec{H} - (\vec{a} \cdot \vec{H}) \vec{a} \right]$$

$$= N\kappa_\perp \vec{H} + N\Delta \kappa (\vec{a} \cdot \vec{H}) \vec{a}$$

$$= N\kappa_\perp \vec{H} + N\Delta \kappa (\vec{a} \vec{a}) \cdot \vec{H},$$

where $N$ is the molecular number density, $\kappa_\parallel$ and $\kappa_\perp$ are molecular magnetic polarizabilities along and perpendicular to the long molecular axis, respectively, and $\Delta \kappa = \kappa_\parallel - \kappa_\perp$. Expressed in matrix form, Equation (1.72) changes to

$$\vec{M} = N \begin{pmatrix} \kappa_\perp + \Delta \kappa a_x & \Delta \kappa a_y & \Delta \kappa a_z \\ \Delta \kappa a_x & \kappa_\parallel + \Delta \kappa a_y & \Delta \kappa a_z \\ \Delta \kappa a_y & \Delta \kappa a_z & \kappa_\parallel + \Delta \kappa a_x \end{pmatrix} \cdot \vec{H} = N \begin{pmatrix} \kappa_\perp \\ \kappa_\parallel \end{pmatrix} \cdot \vec{H},$$

$$\quad$$

(1.73)

where $a_i (i = x, y, z)$ are the projection of $\vec{a}$ in the $x$, $y$, and $z$ directions in the lab frame whose $z$ axis is parallel to the liquid crystal director, and $a_z = \cos \theta$, $a_x = \sin \theta \cos \phi$, and $a_y = \sin \theta \sin \phi$. The molecule swivels because of thermal motion. The averaged magnetization is $\vec{M} = N < \kappa > \cdot \vec{H}$. For a uniaxial liquid crystal, recall $< \cos^2 \theta > = (2S + 1)/3$, $< \sin^2 \theta > = (2 - 2S)/3$, $< \sin^2 \phi > = < \cos^2 \phi > = 1/2$, and $< \sin \phi \cos \phi > = 0$. Therefore

$$< \kappa > = \begin{pmatrix} \kappa_\perp + \frac{1}{3}(1-S)\Delta \kappa & 0 & 0 \\ 0 & \kappa_\parallel + \frac{1}{3}(1-S)\Delta \kappa & 0 \\ 0 & 0 & \kappa_\perp + \frac{1}{3}(2S+1)\Delta \kappa \end{pmatrix}.$$
Because $\vec{M} = \vec{\chi} \cdot \vec{H}$, the magnetic susceptibility tensor is

$$\vec{\chi} = \begin{pmatrix} \chi_\perp & 0 & 0 \\ 0 & \chi_\perp & 0 \\ 0 & 0 & \chi_{//} \end{pmatrix} = N \begin{pmatrix} \kappa_\perp + \frac{1}{3}(1-S)\Delta\kappa & 0 & 0 \\ 0 & \kappa_\perp + \frac{1}{3}(1-S)\Delta\kappa & 0 \\ 0 & 0 & \kappa_\perp + \frac{1}{3}(2S+1)\Delta\kappa \end{pmatrix}. \quad (1.75)$$

The anisotropy is

$$\Delta \chi = \chi_{//} - \chi_\perp = N\Delta\kappa S. \quad (1.76)$$

For most liquid crystals, $\chi_{//}$ and $\chi_\perp$ are negative and small ($\sim 10^{-5}$ in SI units). $\Delta \chi$ is usually positive. From Equation (1.75) it can be seen that $(2\chi_\perp + \chi_{//})/3 = N(3\kappa_\perp + \Delta\kappa)/3 = N(2\kappa_\perp + \kappa_{//})/3$, which is independent of the order parameter. The quantity $(2\chi_\perp + \chi_{//})/3N$ does not change discontinuously when crossing the isotropic–nematic transition.

### 1.5.2 Dielectric permittivity and refractive index

When an electric field is applied to a liquid crystal, it will induce dipole moments in the liquid crystal. For a uniaxial liquid crystal, the molecule can be regarded as a cylinder, and it has different molecular polarizabilities along and perpendicular to the long molecular axis $\vec{a}$. Similar to the magnetic case, when a local electric field $\vec{E}_{\text{loc}}$ (also called internal field) is applied to the liquid crystal, the polarization (dipole moment per unit volume) is given by

$$\vec{P} = N\alpha_{//}(\vec{a} \cdot \vec{E}_{\text{loc}}) \vec{a} + N\alpha_\perp \left[ E_{\text{loc}} - (\vec{a} \cdot \vec{E}_{\text{loc}}) \vec{a} \right]$$

$$= N\alpha_{//}E_{\text{loc}} + N\Delta\alpha (\vec{a} \cdot \vec{E}_{\text{loc}}) \vec{a}$$

$$= N\alpha_{//}E_{\text{loc}} + N\Delta\alpha (\vec{a} \cdot \vec{E}_{\text{loc}}) \vec{E}_{\text{loc}}, \quad (1.77)$$

where $N$ is the molecular density, $\alpha_{//}$ and $\alpha_\perp$ are the molecular polarizabilities along and perpendicular to the long molecular axis, respectively, and $\Delta\alpha = \alpha_{//} - \alpha_\perp$. Different from the magnetic case, the dipole–dipole interaction between the molecules are strong or, stated in another way, the local electric field on a molecule is the sum of the externally applied electric field and the electric field produced by the dipole moment of other molecules. We can approach this problem in the following way. Imagine a cavity created by removing the molecule under consideration, as shown in Figure 1.15. The macroscopic field $\vec{E}$ is the sum of the field $\vec{E}_{\text{self}}$ produced by the molecule itself and the field $\vec{E}_{\text{else}}$, which is the local field $\vec{E}_{\text{local}}$, produced by the external source and the rest of the molecules of the system:

$$\vec{E} = \vec{E}_{\text{self}} + \vec{E}_{\text{else}} = \vec{E}_{\text{self}} + \vec{E}_{\text{local}} \quad (1.78)$$
In order to illustrate the principle, let us first consider an isotropic medium. The cavity can be regarded as a sphere. The field $E$ is produced by the dipole moment inside the sphere, which can be calculated in the following way. In the calculation of the field, the dipole moment can be replaced by the surface charge produced by the dipole moment on the surface of the sphere. The surface charge density is $\sigma = \frac{P}{C_1 \text{m}}$. The field produced by the surface charge is $E_{\text{self}} = -\frac{P}{3\varepsilon_o}$. The local field is $E_{\text{local}} = E + \frac{P}{3\varepsilon_o}$. Hence the polarizability is

$$P = N\alpha E_{\text{loc}} = N\alpha \left( E + \frac{P}{3\varepsilon_o} \right), \quad (1.79)$$

$$\frac{P}{E_{\text{loc}}} = \frac{N\alpha E}{1 - N\alpha/3\varepsilon_o}. \quad (1.80)$$

The electric displacement $D = \varepsilon_o e E = \varepsilon_o E + P$, where $\varepsilon_o = 8.85 \times 10^{12} \text{N}/\text{V}^2$ is the permittivity of vacuum, and $\varepsilon$ is the (relative) dielectric constant, which is given by

$$\varepsilon = 1 + \frac{P}{\varepsilon_o E} = 1 + \frac{N\alpha/\varepsilon_o}{1 - N\alpha/3\varepsilon_o}, \quad (1.81)$$

$$\frac{\varepsilon - 1}{\varepsilon + 2} = \frac{1}{3\varepsilon_o} N\alpha, \quad (1.82)$$

which is called the Clausius–Mossotti relation. At optical frequency, the refractive index $n$ is given by $n^2 = \varepsilon$, and therefore

$$\frac{n^2 - 1}{n^2 + 2} = \frac{1}{3\varepsilon_o} N\alpha, \quad (1.83)$$

which is called the Lorentz–Lorenz relation. The local field is related to the macroscopic field by

$$E_{\text{local}} = E + \frac{P}{3\varepsilon_o} = E + \frac{N\alpha/3\varepsilon_o}{1 - N\alpha/3\varepsilon_o} E = \frac{1}{1 - N\alpha/3\varepsilon_o} E = K E, \quad (1.84)$$

where the defined $K = 1/(1 - N\alpha/3\varepsilon_o)$ is called the internal field constant.

![Figure 1.15](image-url) Schematic diagram showing how the macroscopic field is produced in the medium.
Liquid crystals are anisotropic. In them the local field $\tilde{E}_{\text{local}}$ depends on the macroscopic field $\tilde{E}$ as well as the angles between $\tilde{E}$ and the long molecular axis $\tilde{a}$ and the liquid crystal director $\tilde{n}$. They are related to each other by

$$\tilde{E}_{\text{local}} = \tilde{K} \cdot \tilde{E},$$  \hspace{1cm} (1.85)

where $\tilde{K}$ is the internal field tensor which is a second-rank tensor. After taking account of the internal field tensor and the thermal motion of the molecules, the polarization is

$$\tilde{P} = N\alpha_\perp < \tilde{K} > + N\Delta \alpha < \tilde{K} \cdot (\tilde{a} \tilde{a}) > \cdot \tilde{E}.$$ \hspace{1cm} (1.86)

The macroscopic dielectric tensor is

$$\tilde{\varepsilon} = I + \frac{N}{\varepsilon_0} \left[ \alpha_\perp < \tilde{K} > + \Delta \alpha < \tilde{K} \cdot (\tilde{a} \tilde{a}) > \right].$$ \hspace{1cm} (1.87)

In a material consisting of non-polar molecules, the induced polarization is contributed by two parts: (1) electronic polarization, $P_{\text{electronic}}$, which comes from the deformation of the electron clouds of the constituting atoms of the molecule, (2) ionic polarization, $P_{\text{ionic}}$, which comes from the relative displacement of the atoms constituting the molecule. For a material consisting of polar molecules, there is a third contribution: dipolar polarization, $P_{\text{dipolar}}$, which comes from the reorientation of the dipole. These contributions to the molecular polarizability depend on the frequency of the applied field. The rotation of the molecule is slow and therefore the dipole-orientation polarization can only contribute up to a frequency of the order of MHz. The vibration of atoms in molecules is faster and the ionic polarization can contribute up to the frequency of infrared light. The motion of electrons is the fastest and the electronic polarization can contribute up to the frequency of UV light. In relation to their magnitudes, the order is $P_{\text{electronic}} < P_{\text{ionic}} < P_{\text{dipolar}}$.

At optical frequencies, only the electronic polarization contributes to the molecular polarizability, which is small, and the electric field is usually low. De Jeu and Bordewijk experimentally showed that: (1) $(2\varepsilon_\perp + \varepsilon_\parallel)/3\rho$ is a constant through the nematic and isotropic phases [25,26], where $\rho$ is the mass density, (2) the dielectric anisotropy $\Delta \varepsilon = \varepsilon_\parallel - \varepsilon_\perp$ is directly proportional to the anisotropy of the magnetic susceptibility. Based on these facts, it was concluded that $\tilde{K}$ is a molecular tensor independent of the macroscopic dielectric anisotropy. In the molecular principal frame $\eta\xi\zeta$ with the $\xi$ axis parallel to the long molecular axis $\tilde{a}$, $\tilde{K}$ has the form

$$\tilde{K} = \begin{pmatrix} K_\perp & 0 & 0 \\ 0 & K_\parallel & 0 \\ 0 & 0 & K_{\parallel} \end{pmatrix}.$$ \hspace{1cm} (1.88)

Next we need to find the form of $\tilde{K}$ in the lab frame $xyz$ with the $z$ axis parallel to the liquid crystal director $\tilde{n}$. Because of the axial symmetry around $\tilde{a}$, we only need to consider the transformation of the matrix between the two frames as shown in Figure 1.16. The frame $\eta\zeta\xi$ is
achieved by first rotating the frame \(xyz\) around the \(z\) axis by the angle \(\phi\) and then rotating the frame around the \(\varsigma\) axis by the angle \(\theta\). The rotation matrix is

\[
\begin{pmatrix}
\cos \theta \cos \phi & -\sin \phi & \sin \theta \cos \phi \\
\cos \theta \sin \phi & \cos \phi & \sin \theta \sin \phi \\
-\sin \theta & 0 & \cos \theta
\end{pmatrix}.
\] (1.89)

The reserve rotation matrix is

\[
\begin{pmatrix}
\cos \theta \cos \phi & \cos \theta \sin \phi & -\sin \theta \\
-\sin \phi & \cos \phi & 0 \\
\sin \theta \cos \phi & \sin \theta \sin \phi & \cos \theta
\end{pmatrix}.
\] (1.90)

In the lab frame \(\vec{K}\) has the form

\[
\vec{K} = R \cdot \begin{pmatrix} K_\perp & 0 & 0 \\ 0 & K_\perp & 0 \\ 0 & 0 & K_{//} \end{pmatrix} \cdot R^{-1}
\]

\[
= \begin{pmatrix} K_\perp + \Delta K \sin^2 \theta \cos^2 \phi & \Delta K \sin^2 \theta \sin \phi \cos \phi & \Delta K \sin \theta \cos \theta \cos \phi \\ \Delta K \sin^2 \theta \sin \phi \cos \phi & K_\perp + \Delta K \sin^2 \theta \sin^2 \phi & \Delta K \sin \theta \cos \theta \sin \phi \\ \Delta K \sin \theta \cos \theta \cos \phi & \Delta K \sin \theta \cos \theta \sin \phi & K_{//} + \Delta K \cos^2 \theta \end{pmatrix},
\] (1.91)

where \(\Delta K = K_{//} - K_\perp\), and \(\vec{a} \vec{a}\) has the form

Figure 1.16 Schematic diagram showing the transformation between the molecular principal frame \(\eta\varsigma\xi\) and the lab frame \(xyz\).
\[
\vec{a}\vec{a} = \begin{pmatrix}
\sin \theta \cos \phi \\
\sin \theta \sin \phi \\
\cos \theta
\end{pmatrix}
\begin{pmatrix}
\sin \theta \cos \phi \sin \theta \sin \phi \cos \theta \\
\sin \theta \sin \phi \cos \phi \sin \theta \sin \phi \cos \theta \\
\sin \theta \cos \cos \phi \sin \theta \sin \phi \cos \phi \\
\sin \theta \cos \cos \phi \sin \theta \sin \phi \cos \phi \\
\cos \theta
\end{pmatrix},
\]
(1.92)

and \(\vec{a}\vec{a} \cdot \vec{K}\) has the form

\[
\vec{a}\vec{a} \cdot \vec{K} = \begin{pmatrix}
K_{\parallel} \sin^2 \theta \cos^2 \phi & K_{\parallel} \sin^2 \theta \sin \phi \cos \phi & K_{\parallel} \sin \theta \cos \theta \cos \phi \\
K_{\parallel} \sin^2 \theta \sin \phi \cos \phi & K_{\parallel} \sin^2 \theta \sin^2 \phi & K_{\parallel} \sin \theta \cos \theta \sin \phi \\
K_{\parallel} \sin \theta \cos \cos \phi & K_{\parallel} \sin \theta \sin \phi \cos \phi & K_{\parallel} \cos \theta
\end{pmatrix},
\]
(1.93)

Recall that \(< \cos^2 \theta > = (2S + 1)/3, < \sin^2 \theta > = (2 - 2S)/3, < \sin^2 \phi > = < \cos^2 \phi > = 1/2, < \cos \theta > = < \sin \phi > = < \cos \phi > = < \sin \phi \cos \phi > = 0\), therefore their averaged values are

\[
< \vec{K} > = \begin{pmatrix}
K_{\perp} + \Delta K(1 - S)/3 & 0 & 0 \\
0 & K_{\perp} + \Delta K(1 - S)/3 & 0 \\
0 & 0 & K_{\perp} + \Delta K(2S + 1)/3
\end{pmatrix},
\]
(1.94)

\[
< \vec{a}\vec{a} \cdot \vec{K} > = \begin{pmatrix}
K_{\parallel}(1 - S)/3 & 0 & 0 \\
0 & K_{\parallel}(1 - S)/3 & 0 \\
0 & 0 & K_{\parallel}(2S + 1)/3
\end{pmatrix},
\]
(1.95)

Therefore

\[
\vec{\varepsilon} = \vec{I} + \frac{N}{\varepsilon_0} \left[ \alpha_{\perp} < \vec{K} > + \Delta \alpha < \vec{K} \cdot (\vec{a}\vec{a}) > \right].
\]

The anisotropy is

\[
\Delta \varepsilon = \varepsilon_{\parallel} - \varepsilon_{\perp} = \frac{N}{\varepsilon_0} (\alpha_{\parallel} K_{\parallel} - \alpha_{\perp} K_{\perp}) S,
\]
(1.97)
which is linearly proportional to the order parameter $S$. In terms of the refractive indices, Equation (1.97) becomes

$$n_{xx}^2 - n_{\perp}^2 = 2\tilde{n}\Delta n = \frac{N}{\varepsilon_0} (\alpha_{xx}/K_{xx} - \alpha_{\perp} K_{\perp}) S,$$

where $\tilde{n} = (n_{xx} + n_{\perp})/2$ and $\Delta n = (n_{xx} - n_{\perp})$. Approximately, the birefringence $\Delta n$ is linearly proportional to the order parameter. For most liquid crystals, $\tilde{n} \sim 1.5$–2.0 and $\Delta n \sim 0.05$–0.3.

The electronic polarization may be treated by using classical mechanics, where the system is regarded as a simple harmonic oscillator. There are three forces acting on the electron: (1) elastic restoring force $-Kx$, where $K$ is the elastic constant and $x$ is the displacement of the electron from its equilibrium position, (2) viscosity force $-\gamma \partial x/\partial t$, and (3) the electric force $-eE_o e^{i\omega t}$, where $E_o$ and $\omega$ are the amplitude and frequency of the applied electric field, respectively. The dynamic equation is

$$m \frac{d^2 x}{dt^2} = -Kx - eE_o e^{i\omega t} - \frac{\gamma}{m} \frac{\partial x}{\partial t}.$$  \hspace{1cm} (1.98)

The solution is $x = x_o e^{i\omega t}$ and the amplitude of the oscillation is

$$x_o = \frac{-eE_o}{m(\omega_o^2 - \omega^2) + i\gamma \omega},$$  \hspace{1cm} (1.99)

where $\omega_o = \sqrt{k/m}$ is the frequency of the oscillator (the frequency of the transition dipole moment in quantum mechanics). The induced dipole moment is $p = -ex_o$. The molecule polarizability is

$$\alpha = p/E_{loc} = \frac{e^2}{m} \frac{(\omega_o^2 - \omega^2)}{(\omega_o^2 - \omega^2)^2 + (\gamma \omega/m)^2} - \frac{i}{m} \frac{\gamma \omega/m}{(\omega_o^2 - \omega^2)^2 + (\gamma \omega/m)^2},$$  \hspace{1cm} (1.100)

which is a complex number, and the imaginary part corresponds to absorption. When the frequency of the light is far away from the absorption frequency $\omega_o$, the absorption is negligible, $\alpha = p/E_{loc} = (e^2/m)(\omega_o^2 - \omega^2)$. The refractive index is

$$n^2 \propto \alpha \propto \frac{1}{(\omega_o^2 - \omega^2)} = \frac{1}{[(2\pi/C\lambda_o)^2 - (2\pi/\lambda)]^2} = \frac{C^2 \lambda_o^2}{4\pi^2 \lambda^2 - \lambda_o^2},$$

as expressed in Sellmeier’s equation

$$n^2 = 1 + \frac{H\lambda_o^2}{\lambda^2 - \lambda_o^2},$$  \hspace{1cm} (1.101)

where $H$ is a constant. When $\lambda$ is much longer than $\lambda_o$, expanding the above equation, we have

$$n \approx A + \frac{B}{\lambda^2} + \frac{C}{\lambda^4},$$  \hspace{1cm} (1.102)
This is Cauchy’s Equation. The refractive index increases with decreasing wavelength. For liquid crystals, along different directions with respect to the long molecular axis, the molecular polarizabilities are different. Also along different directions, the frequencies of the transition dipole moments are different, which results in dichroic absorption: when the electric field of light is parallel to the transition dipole moment, light is absorbed, but when the electric field is perpendicular to the transition moment, light is not absorbed. Positive dichroic dyes have transition dipole moments parallel to the long molecular axis while negative dichroic dyes have transition dipole moments perpendicular to the long molecular axis.

At DC or at low frequency applied electric fields, for liquid crystals of polar molecules, the dipolar polarization is dominant. For a liquid crystal with a permanent dipole moment \( \vec{p} \), the polarization is now given by

\[
P = N\alpha_\perp < \vec{K} > \cdot \vec{E} + N\Delta\alpha < \vec{K} \cdot (\vec{a} \vec{a}) > \cdot \vec{E} + N < \vec{\rho} >.
\]  

The macroscopic dielectric tensor is

\[
\tilde{\varepsilon} = \tilde{I} + \frac{N}{\varepsilon_0} \left[ \alpha_\perp < \vec{K} > + \Delta\alpha < \vec{K} \cdot (\vec{a} \vec{a}) > + < \vec{\rho} > \vec{E} / E^2 \right].
\]

The energy of the dipole in the directing electric field \( \vec{E}_d \) is \( u = -\vec{\rho} \cdot \vec{E}_d \). The directing field \( \vec{E}_d \) is different from the local field \( \vec{E}_{loc} \) because the dipole polarizes its surroundings, which in turn results in a reaction field \( \vec{E}_r \) at the position of the dipole. As \( \vec{E}_r \) is always parallel to the dipole, it cannot affect the orientation of the dipole. As an approximation, it is assumed that \( \vec{E}_d = \vec{d} \cdot \vec{E} \), where \( \vec{d} \) is a constant. Usually the dipole moment \( \vec{p} \) is about \( 1 \text{e} \times 1 \text{Å} = 1.6 \times 10^{-19} \text{C} \times 10^{-10} \text{m} \). At room temperature (\( T \sim 300 \text{K} \)) and under the normal strength field \( E \sim 1 \text{V/μm} = 10^6 \text{V/m} \), \( pE/3k_BT \ll 1 \). Consider a liquid crystal molecule with a permanent dipole moment making the angle \( \beta \) with the long molecular axis. In the molecular frame \( \eta\xi\zeta \), the components of \( \vec{p} \) are \( (p\sin \beta \cos \psi, p\sin \beta \sin \psi, p\cos \beta) \), as shown in Figure 1.17. Using the

![Figure 1.17](image-url)
rotation matrix given by Equation (1.90), we can calculate the components of $\bar{p}$ in the lab frame $xyz$:

$$
\bar{p} = \begin{pmatrix}
\cos \phi & -\cos \theta \sin \phi & -\sin \theta \\
\sin \phi & \cos \theta \cos \phi & \sin \theta \cos \phi \\
0 & -\sin \theta & \cos \theta
\end{pmatrix}
\begin{pmatrix}
\sin \beta \\
\sin \beta \sin \psi \\
\cos \beta
\end{pmatrix}
\begin{pmatrix}
\sin \beta \cos \psi \\
\sin \beta \sin \psi \\
\cos \beta
\end{pmatrix}.
$$

(1.105)

When the applied field is parallel to $\vec{n}$, $\vec{E} = E_{/\parallel} \vec{z}$, the projection of the dipole along the applied field is

$$
p_{/\parallel} = -p(\sin \beta \sin \psi \sin \theta + \cos \beta \cos \theta),
$$

(1.106)

and the energy is

$$
u = -dp(\cos \beta \cos \theta - \sin \beta \sin \psi \sin \theta)E_{/\parallel}.
$$

(1.107)

The average value of the projection is

$$
\langle p_{/\parallel} \rangle = \frac{\int (p \cos \beta \cos \theta - p \sin \beta \sin \psi \sin \theta)e^{-u/k_BT - V(\theta)/k_BT} \sin \theta d\theta d\phi d\psi}{\int e^{-u/k_BT - V(\theta)/k_BT} \sin \theta d\theta d\phi d\psi}.
$$

(1.108)

Because $-u \ll k_BT$, $e^{-u/k_BT} \approx (1 - u/k_BT)$, then

$$
\langle p_{/\parallel} \rangle = \frac{dE_{/\parallel}}{k_BT} \int (p \cos \beta \cos \theta - p \sin \beta \sin \psi \sin \theta)^2 e^{-V(\theta)/k_BT} \sin \theta d\theta d\phi d\psi
$$

$$
= \frac{dE_{/\parallel}p^2}{k_BT} < (\cos^2 \beta \cos^2 \theta + \sin^2 \beta \sin^2 \theta \sin^2 \psi - \sin \beta \cos \beta \sin \theta \cos \theta \sin \psi) >
$$

Because $\langle \sin^2 \psi \rangle = 1/2$, $\langle \sin \psi \rangle = 0$, $\langle \cos^2 \theta \rangle = (2S + 1)/3$ and $\langle \sin^2 \theta \rangle = (2 - 2S)/3$,

$$
\langle p_{/\parallel} \rangle = \frac{dE_{/\parallel}p^2}{3k_BT} \left[ \cos \beta (2S + 1) + \sin \beta (1 - S) \right] = \frac{dE_{/\parallel}p^2}{3k_BT} \left[ 1 - (1 - 3\cos^2 \beta)S \right]
$$

(1.109)

From Equations (1.96), (1.104), and (1.109), we have

$$
\varepsilon_{/\parallel} = 1 + \frac{N}{3\varepsilon_o} \left\{ \alpha_{/\parallel} K_{/\parallel} (2 - 2S) + \alpha_{/\parallel} K_{/\parallel} (1 + 2S) + \frac{dp^2}{k_BT} \left[ 1 - (1 - 3\cos^2 \beta)S \right] \right\}.
$$

(1.110)
Note that $\alpha_\parallel$ and $\alpha_\perp$ are the molecular polarizabilities contributed by the electronic and ionic polarizations.

When the applied field is perpendicular to $\vec{n}$, say $\vec{E} = E_\perp \hat{x}$, the projection of the dipole along the applied field is

$$p_\perp = p (\sin \beta \cos \psi \cos \phi - \sin \beta \sin \psi \cos \theta \sin \phi - \cos \beta \sin \theta \sin \phi),$$

and the energy is

$$u = -dp (\sin \beta \cos \psi \cos \phi - \sin \beta \sin \psi \cos \theta \sin \phi - \cos \beta \sin \theta \sin \phi) E_\perp.$$

The average value of the projection is

$$<p_\perp> = \frac{dE_\perp p^2}{k_B T} \left[ \frac{1}{4} \sin^2 \beta + \frac{1}{4} \sin^2 \beta \frac{(2S+1)}{3} + \cos^2 \beta \frac{(1-S)}{3} \right] = \frac{dE_\perp p^2}{3k_B T} \left[ 1 + \frac{1}{2} (1-3 \cos^2 \beta) S \right].$$

From Equations (1.96), (1.104) and (1.112), we have

$$\varepsilon_\perp = 1 + \frac{N}{3 \varepsilon_o} \left\{ \alpha_\perp K_\perp (2 + S) + \alpha_\parallel K_\parallel (1 - S) + \frac{dp^2}{k_B T} \left[ 1 + \frac{1}{2} (1-3 \cos^2 \beta) S \right] \right\},$$

The dielectric anisotropy is

$$\Delta \varepsilon = \varepsilon_\parallel - \varepsilon_\perp = \frac{N}{\varepsilon_o} \left[ (\alpha_\parallel K_\parallel - \alpha_\perp K_\perp) - \frac{dp^2}{2k_B T} (1-3 \cos^2 \beta) \right] S,$$

which is proportional to the order parameter $S$. The contribution of induced polarization (electronic and ionic polarizations) changes with temperature as $S$, while the contribution of the orientation polarization changes with temperature as $S/T$. When the angle between the permanent dipole and the long molecular axis is $\beta = 55^\circ$, $(1-3 \cos^2 \beta) = 0$, the orientation polarization of the permanent dipole does not contribute to $\Delta \varepsilon$.

The permanent dipole moment is fixed on the molecule. Thus the molecule has to reorient in order to contribute to the dielectric constants. Qualitatively speaking, only when the frequency of the applied field is lower than a characteristic frequency $\omega_\parallel$, can the molecule rotate to follow the oscillation of the applied field and therefore to contribute to the dielectric constants. For rod-like liquid crystal molecules, it is easier to spin around the long molecular axis than to rotate around a short molecular axis. Therefore the characteristic frequency $\omega_\perp$ for $\varepsilon_\perp$ is higher than the characteristic frequency $\omega_\parallel$ for $\varepsilon_\parallel$. For molecules on which the angle $\beta$ between the permanent dipole and the long molecular axis is very small, $\Delta \varepsilon$ is always positive at all frequencies. For molecules with large permanent dipole moment $p$ and large $\beta$, $\Delta \varepsilon$ is negative at low frequencies. For molecules with large permanent dipole moment $p$ and intermediate $\beta$, $\Delta \varepsilon$ is positive at low frequencies, then changes to negative when the frequency is increased above a crossover frequency $\omega_c$. The crossover frequency is in the region from a few kHz to a few tens of kHz. At infrared light or higher frequencies, the dipolar polarization does not contribute anymore, and $\Delta \varepsilon$ is always positive.
1.6 Anchoring Effects of Nematic Liquid Crystal at Surfaces

In most liquid crystal devices, the liquid crystals are sandwiched between two substrates coated with alignment layers. In the absence of externally applied fields, the orientation of the liquid crystal in the cell is determined by the anchoring condition of the alignment layer [26–28].

1.6.1 Anchoring energy

Consider an interface between a liquid crystal \((z > 0)\) and an alignment layer \((z < 0)\), as shown in Figure 1.18. For a liquid crystal molecule on the interface, some of the surrounding molecules are liquid crystal molecules and the other surrounding molecules are alignment layer molecules. The potential for the molecule’s orientation is different from that of the liquid crystal in the bulk, where all the surrounding molecules are liquid crystal molecules. At the interface, the orientational and positional orders may be different from those in the bulk. Here we only discuss the anisotropic part of the interaction between the liquid crystal molecule and the alignment layer molecule. The liquid crystal is anisotropic. If the alignment layer is also anisotropic, then there is a preferred direction – referred to as the easy axis – for the liquid crystal director at the interface, as shown in Figure 1.18. The interaction energy is a minimum when the liquid crystal director is along the easy axis. The \(z\) axis is perpendicular to the interface and pointing toward the liquid crystal side. The polar angle and azimuthal angle of the easy axis are \(\theta_o\) and \(\phi_o\), respectively. If \(\theta_o = 0^\circ\), the anchoring is referred to as homeotropic. If \(\theta_o = 90^\circ\) and \(\phi_o\) is well defined, the anchoring is termed homogeneous. If \(\theta_o = 90^\circ\) and there is no preferred azimuthal angle, the anchoring is called planar. If \(0 < \theta_o < 90^\circ\), the anchoring is referred to as tilted.

When the liquid crystal director \(\hat{n}\) is aligned along the direction specified by the polar angle \(\theta\) and azimuthal angle \(\phi\), the anisotropic part of the surface energy – referred to as the anchoring energy function – of the liquid crystal is \(f_s = f_s(\theta, \phi)\). When \(\theta = \theta_o\) and \(\phi = \phi_o\), \(f_s\) has the minimum value of 0, and thus \(\partial f_s / \partial \theta |_{\theta = \theta_o} = 0\) and \(\partial f_s / \partial \phi |_{\phi = \phi_o} = 0\).

The materials above and below the interface are different, and there is no reflection symmetry about the interface. If \(\theta_o \neq 0\), the anchoring energy does not have azimuthal rotational symmetry.

Figure 1.18 Schematic diagram showing the easy direction of the surface anchoring and the deviation of the liquid crystal director.
symmetry around the easy direction. Therefore the anchoring energies are different for deviation in polar angle and azimuthal angle. For small deviations, in the Rapini–Papoular model \[29\], the anchoring energy function can be expressed as

\[
fs = \frac{1}{2} W_p \sin^2 \beta_p + \frac{1}{2} W_a \sin^2 \beta_a,
\]

where \(\beta_p\) and \(\beta_a\) are the angles between \(\hat{n}\) and the easy axis when \(\hat{n}\) deviates from the easy axis in the polar angle direction and the azimuthal angle direction, respectively. \(W_p\) and \(W_a\) are the polar and azimuthal anchoring strengths, respectively. For small values of \(\theta - \theta_o\) and \(\phi - \phi_o\), we have the approximations \(\sin^2 \beta_p = \sin^2 (\theta - \theta_o)\) and \(\sin^2 \beta_a = \sin^2 (\phi - \phi_o)\sin^2 \theta_o\). Therefore the anchoring energy function is

\[
fs = \frac{1}{2} W_p \sin^2 (\theta - \theta_o) + \frac{1}{2} W_a \sin^2 \theta_o \sin^2 (\phi - \phi_o).
\]

Equations (1.116) and (1.117) are valid only for small deviations. For large deviations, quartic terms must be included. The anchoring strengths can be determined experimentally, which will be discussed in Chapter 5.

### 1.6.2 Alignment layers

Homogeneous anchoring can be achieved by mechanically rubbing the surface of the substrate, such as glass, of the liquid crystal cell with a cotton ball or cloth. The rubbing creates micro-grooves along the rubbing direction in the form of ridges and troughs, as shown in Figure 1.19(a). When the liquid crystal is aligned parallel to the grooves, there is no orientation deformation. If the liquid crystal were perpendicular to the grooves, there would be orientation deformation which costs elastic energy. Therefore the liquid crystal will be homogeneously aligned along the grooves (the rubbing direction). The problem with alignment created in this way is that the anchoring strength (~10\(^{-5}\) J/m\(^2\)) is weak. The widely used homogeneous alignment layers are rubbed polyimides. The rubbing not only creates the micro-grooves but also aligns the polymer chains. The intermolecular interaction between the liquid crystal and the aligned polymer chains also favors the parallel alignment and thus increases the anchoring energy. The anchoring strength can become as high as 10\(^{-3}\) J/m\(^2\). Furthermore, pretilt angle of a few degrees can be generated. Homogeneous anchoring can also be achieved by using obliquely evaporated SiO film.

Homeotropic anchoring can be achieved using monolayer surfactants such as lecithin and silane. The polar head of the surfactant is chemically attached to the glass substrate, and the hydrocarbon tail points out and perpendicular to the surface, as shown in Figure 1.19(b).
The intermolecular interaction between the surfactant and the liquid crystal promotes the homeotropic alignment.

1.7 Liquid crystal director elastic deformation

When a nematic liquid crystal is in the ground state the direction of the liquid crystal director $\mathbf{n}$ is uniform in space. When $\mathbf{n}$ is deformed, there will be elastic energy. Although liquid crystal director deformations cost energy, they do occur in reality because of surface anchoring, spatial confinements, impurities, irregularities, and externally applied fields [1,23]. In this section, we consider possible director deformations, associated elastic energies, and transformation between deformations.

1.7.1 Elastic deformation and disclination

We consider the possible deformations of a nematic liquid crystal confined between two parallel substrates with tangential anchoring condition (parallel to the substrates but no preferred direction on the plane of the substrate). We use the one elastic constant approximation ($K_{11} = K_{22} = K_{33} = K$), the elastic energy is given by

$$f = \frac{1}{2} K (\nabla \cdot \mathbf{n})^2 + \frac{1}{2} K (\mathbf{n} \cdot \nabla \times \mathbf{n})^2 + \frac{1}{2} K (\mathbf{n} \times \nabla \times \mathbf{n})^2 + \frac{1}{2} K (\nabla \cdot \mathbf{n})^2 + \frac{1}{2} K (\nabla 	imes \mathbf{n})^2. \quad (1.117)$$

Figure 1.19 Schematic diagram of alignment layers: (a) homogeneous alignment layer, (b) homeotropic alignment layer.

The intermolecular interaction between the surfactant and the liquid crystal promotes the homeotropic alignment.
Because of the anchoring condition, the liquid crystal director is parallel to the plane of the substrate and depends on the coordinates $x$ and $y$ on the plane. The liquid crystal director is described by

$$\vec{n} = \cos \theta(x,y)\hat{x} + \sin \theta(x,y)\hat{y}, \quad (1.118)$$

where $\theta$ is the angle between the director and the $x$ axis. Substituting Equation (1.118) into Equation (1.117) we get

$$f = \frac{1}{2}K \left( \frac{\partial(\cos \theta)}{\partial x} + \frac{\partial(\sin \theta)}{\partial y} \right)^2 + \frac{1}{2}K \left[ \left( \frac{\partial(\sin \theta)}{\partial x} - \frac{\partial(\cos \theta)}{\partial y} \right) \right]^2 = \frac{1}{2}K \left[ \left( \frac{\partial \theta}{\partial x} \right)^2 + \left( \frac{\partial \theta}{\partial y} \right)^2 \right]. \quad (1.119)$$

Using the Euler–Lagrange method (which will be presented in Chapter 5) to minimize the free energy we get

$$\frac{\partial^2 \theta}{\partial x^2} + \frac{\partial^2 \theta}{\partial y^2} = 0. \quad (1.120)$$

When we switch from Cartesian coordinates to cylindrical coordinates, the above equation becomes

$$\frac{1}{r} \frac{\partial}{\partial r} \left( r \frac{\partial \theta}{\partial r} \right) + \frac{1}{r^2} \frac{\partial^2 \theta}{\partial \phi^2} = 0. \quad (1.121)$$

We consider the case where $\theta$ only depends on the azimuthal angle $\phi$, but not the radius $r$. The solution to the above equation is

$$\theta = S\phi + \theta_o = S\tan^{-1}(x/y) + \theta_o. \quad (1.122)$$

When we go around one complete circle, the azimuthal angle changes by $2\pi$, the liquid crystal director must be in the same direction. Furthermore $\vec{n}$ and $-\vec{n}$ are equivalent. Therefore it is required that

$$\theta(\phi = 2\pi) - \theta(\phi = 0) = (S \cdot 2\pi + \theta_o) - (S \cdot 0 + \theta_o) = 2\pi S = \pm m \pi, \quad S = \pm m/2, \quad (1.123)$$

where $m = 0, 1, 2, \ldots$. The liquid crystal director configurations for the deformations with variety of $S$ and $\theta_o$ values are shown in Figure 1.20. In the center there is a singularity. In the liquid crystal cell the singularity goes from one surface to the other surface of the cell. Thus is it a line singularity and is called disclination. $S$ is called the strength of the disclination. For a positive $S$, the liquid crystal director rotates counterclockwise when the azimuthal angle changes counterclockwise. For a negative $S$, the liquid crystal director rotates clockwise when the azimuthal angle changes counterclockwise. The elastic energy density of the deformation is given by

$$f = \frac{1}{2}K \left[ \left( \frac{\partial \tan^{-1}(x/y)}{\partial x} \right)^2 + \left( \frac{\partial \tan^{-1}(x/y)}{\partial y} \right)^2 \right] = \frac{K S^2}{2r^2}. \quad (1.124)$$
The elastic energy is proportional to the square of the strength of the disclination and increases when approaching the singularity.

1.7.2 Escape of liquid crystal director in disclinations

For a disclination with the strength $S$, as one approaches the center of the disclination, the elastic energy diverges, as shown by Equation (1.124). In reality this will not occur. The liquid crystal will transform either into isotropic phase at the center of the disclination or a different deformation where there is no singularity. Here we only discuss the cases of cylindrical confinements (two-dimensional confinement) where it is possible to obtain analytical solutions. The mechanism of liquid crystal director escape in spherical confinement (three-dimensional confinement) is similar to that of two-dimensional.

1.7.2.1 Escape to isotropic phase

We consider a nematic liquid crystal confined in a cylinder with a radius of $R$. The anchoring condition on the surface of the cylinder is perpendicular, as shown in Figure 1.21. The liquid crystal director aligned along the radial axis direction, as shown in Figure 1.21(a), and is described by $\vec{n} = \hat{r}$. The elastic deformation of the liquid crystal director is splay with the strength of $S = 1$. The elastic energy is

$$ f = \frac{1}{2} K_{11} \frac{1}{r^2}. $$

(1.125)
When $r \to 0$, $f \to \infty$. In order to avoid the divergence of the elastic energy, the liquid crystal transforms into the isotropic phase inside a core with the radius $R_c$, as shown in Figure 1.21(a) [24]. The radius of the isotropic core is given by

$$\frac{1}{2} K_{11} \frac{1}{R_c^2} = \Delta f = f_I - f_N,$$

$$R_c = \sqrt{K_{11}/(2\Delta f)},$$

where $\Delta f$ is the free energy difference between the isotropic phase and the nematic phase. At temperature $T$, the free energy difference can be estimated by

$$\Delta f = f_I - f_N \approx \left[ \frac{\partial f_I}{\partial T} \right]_{T_N} - \left[ \frac{\partial f_N}{\partial T} \right]_{T_N} (T - T_N) = \left[ -\frac{S_I}{T} + \frac{S_N}{T} \right] (T - T_N) = \frac{L}{T_N} (T_N - T),$$

where $S_I$ and $S_N$ are the entropies of the isotropic and nematic phases, respectively, and $L$ is the latent heat per unit volume of the nematic-isotropic transition. A typical latent heat is 100 cal/mole = 100 × 4.18 J/mole. The density of the liquid crystal is about 1 g/cm$^3$. One mole of liquid
crystal weighs about 300 g and occupies the volume \( 300 \text{ g/(1 g/cm}^3 \text{)} = 300 \text{ cm}^3 \). The latent heat is about \( L = 418 \text{ J/cm}^3 \approx 1.4 \text{ J/cm}^3 \). If the transition temperature \( T_{NI} \) is 300 K and the splay elastic constant \( K_{11} \) is \( 10^{-11} \text{ N} \), from Equations (1.126) and (1.127) we can calculate the radius of the isotropic core at 10 K between the transition temperature:

\[
R_c = \sqrt{10^{-11} \text{ N}/[2 \times 1.4 (\text{J/cm}^3) \times 10 \text{ K}/300 \text{ K}]} \approx 10 \text{ nm}
\]

(1.128)

The total free energy per unit length of the configuration with the isotropic core is

\[
F_{IC} = \int_0^R \left[ \frac{1}{2} K_{11} \frac{1}{r^2} 2 \pi r dr + \pi R^2 \Delta f = \pi K_{11} \ln \left( \frac{R}{R_c} \right) + \frac{1}{2} \pi K_{11} \pi \ln \left( \frac{R}{2 \Delta f} \right)^{1/2} + \frac{1}{2} \pi K_{11} \right].
\]

(1.129)

The total free energy increases with the radius of the cylinder. Note that we neglect the surface energy of the interface between the nematic liquid crystal and the isotropic core.

1.7.2.2 Escape to bend

The second possibility to avoid the singularity at the center of the cylinder is to escape from the splay deformation to the bend deformation, as shown in Figure 1.21(b) [30–32]. The liquid crystal director tilts to the \( z \) direction and is given by

\[
\hat{n} = \sin \theta(r) \hat{r} + \cos \theta(r) \hat{z},
\]

(1.130)

where \( \theta \) is the angle between the liquid crystal director and the \( z \) axis. Under the approximation \( K_{33} = K_{11} \), the elastic energy density is

\[
f = \frac{1}{2} K_{11} \left[ \sin^2 \theta \frac{1}{r^2} + 2 \sin \theta \cos \theta \frac{1}{r} \frac{d \theta}{dr} + \left( \frac{d \theta}{dr} \right)^2 \right].
\]

(1.131)

The total elastic energy per unit length of the cylinder is

\[
F_B = 2 \pi \int_0^R f dr = \pi K_{11} \int_0^R \left[ \frac{\sin^2 \theta}{r} + 2 \sin \theta \cos \theta \frac{d \theta}{dr} + r \left( \frac{d \theta}{dr} \right)^2 \right] dr.
\]

(1.132)

Using Euler–Lagrange method to minimize the elastic energy, we obtain

\[
\frac{\delta (rf)}{\delta \theta} = \frac{\partial (rf)}{\partial \theta} \frac{d}{dr} \left( \frac{\partial (rf)}{\partial r} \right) = \frac{2 \sin \theta \cos \theta}{r} - 2 \frac{d \theta}{dr} - 2 r \frac{d^2 \theta}{dr^2} = 0,
\]

(1.133)
\[
\frac{d\theta}{dr} + r \frac{d^2\theta}{dr^2} = \frac{\sin \theta \cos \theta}{r},
\]
\[
\frac{d}{dr} \left( r \frac{d\theta}{dr} \right) = \frac{1}{r} \sin \theta \cos \theta.
\]  
(1.134)

Multiplying both sides of the above equation by \(r d\theta/dr\), we get
\[
\left( r \frac{d\theta}{dr} \right) d\left( r \frac{d\theta}{dr} \right) = \frac{1}{2} d\left( r \frac{d\theta}{dr} \right) = \frac{1}{2} \sin \theta \cos \theta \left( r \frac{d\theta}{dr} \right) = \frac{1}{2} d\left( \sin^2 \theta \right).
\]  
(1.135)

Integrating the above equation from 0 to \(r\), and using the boundary condition \(\theta(r=0) = 0\), we obtain
\[
r \frac{d\theta}{dr} = \sin \theta,
\]
\[
\frac{dr}{r} = \frac{d\theta}{\sin \theta}.
\]  
(1.136)

Integrating the above equation from \(r\) to \(R\), and using the boundary condition \(\theta(r=R) = \pi/2\), we obtain
\[
\theta = 2\tan^{-1} \left( \frac{R}{r} \right).
\]  
(1.137)

Substituting Equations (1.136) and (1.137) into Equation (1.132) we can get the total free energy,
\[
F_B = 3\pi K_{11},
\]  
(1.138)

which is independent of the radius \(R\).

In order to determine which escape will occur, we compare the total free energies of the two escapes. From Equations (1.129) and (1.138), we get
\[
\Delta F = F_B - F_{IC} = 3\pi K_{11} - \left[ K_{11} \pi \ln \left( \frac{R}{R_c} \right) + \frac{1}{2} \pi K_{11} \right] = \frac{5}{2} \pi K_{11} - K_{11} \pi \ln \left( \frac{R}{R_c} \right).
\]  
(1.139)

We can see that if
\[
\frac{5}{2} \pi K_{11} \leq K_{11} \pi \ln \left( \frac{R}{R_c} \right),
\]
\[
R \geq e^{5/2} R_c \approx 12 R_c.
\]  
(1.140)

The free energy of the state with the bend escape is lower than that of the state with the isotropic core. When the radius of the cylinder is small, it costs too much elastic energy for the liquid crystal to change from 0° at the center to 90° at the surface, and the liquid crystal system will have an isotropic core to avoid the divergence of the splay elastic energy. When the radius of
the cylinder is sufficiently large, the liquid crystal will escape from the splay state to the bend state to further reduce the free energy.

1.7.2.3 Escape to twist

For most liquid crystals, the twist elastic constant is smaller than the bend elastic constant. Therefore it is possible to reduce the total elastic energy by escaping from the bend deformation to the twist deformation as shown in Figure 1.21(c). The liquid crystal director is no longer on the \( r-z \) plane but twists out of the plane and is given by

\[
\hat{n} = \sin \theta(r) \hat{r} + \cos \theta(r) \sin \phi(r) \hat{\phi} + \cos \theta(r) \cos \alpha(r) \hat{z},
\]

where \( \theta \) is the angle of the liquid crystal director defined with respect to the \( \phi-z \) plane, and the twist angle \( \alpha \) is the angle of the projection of the liquid crystal director on the \( \phi-z \) plane defined with respect to the \( z \) axis. The divergence and curl of \( \hat{n} \) are

\[
\nabla \cdot \hat{n} = \frac{1}{r} \frac{\partial}{\partial r} (r \sin \theta) = \frac{\sin \theta}{r} + \cos \theta \frac{\partial \theta}{\partial r}
\]

\[
\nabla \times \hat{n} = \frac{1}{r} \frac{\partial}{\partial r} (r \cos \theta \sin \alpha) \hat{\phi} - \frac{\partial}{\partial r} (\cos \theta \cos \alpha) \hat{\phi}
\]

\[
= \left( \frac{1}{r} \cos \theta \sin \alpha - \sin \theta \sin \alpha \frac{\partial \theta}{\partial r} + \cos \theta \cos \alpha \frac{\partial \alpha}{\partial r} \right) \hat{\phi} + \left( \sin \theta \cos \alpha \frac{\partial \theta}{\partial r} + \cos \theta \sin \alpha \frac{\partial \alpha}{\partial r} \right) \hat{\phi}
\]

\[
(\nabla \times \hat{n})^2 = \left( \frac{1}{r} \cos \theta \sin \alpha \right)^2 + \left( \sin \theta \frac{\partial \theta}{\partial r} \right)^2 + \left( \cos \theta \frac{\partial \alpha}{\partial r} \right)^2
\]

\[
+2 \frac{1}{r} \cos \theta \sin \alpha \left( - \sin \theta \sin \alpha \frac{\partial \theta}{\partial r} + \cos \theta \cos \alpha \frac{\partial \alpha}{\partial r} \right)
\]

\[
\hat{n} \cdot \nabla \times \hat{n} = \left( \frac{1}{r} \cos \theta \sin \alpha - \sin \theta \sin \alpha \frac{\partial \theta}{\partial r} + \cos \theta \cos \alpha \frac{\partial \alpha}{\partial r} \right) \cos \theta \cos \alpha
\]

\[
+ \left( \sin \theta \cos \alpha \frac{\partial \theta}{\partial r} + \cos \theta \sin \alpha \frac{\partial \alpha}{\partial r} \right) \cos \theta \sin \alpha.
\]

The elastic energy density is

\[
f = \frac{1}{2} K_{11} (\nabla \cdot \hat{n})^2 + \frac{1}{2} K_{22} (\hat{n} \cdot \nabla \times \hat{n})^2 + \frac{1}{2} K_{33} \left[ (\nabla \times \hat{n})^2 - (\hat{n} \cdot \nabla \times \hat{n})^2 \right]
\]

\[
f = \frac{1}{2} K_{11} (\nabla \cdot \hat{n})^2 + \frac{1}{2} (K_{22} - K_{33}) (\hat{n} \cdot \nabla \times \hat{n})^2 + \frac{1}{2} K_{33} (\nabla \times \hat{n})^2.
\]
By introducing the twist, the bend elastic can be reduced, but the trade-off is that the twist elastic energy is increased. The change of the elastic energy density [Eq. (1.146) subtracted by Eq. (1.131)] is

$$\Delta f = \frac{1}{2}(K_{22} - K_{33})f_1 + \frac{1}{2}K_{33}f_2,$$

(1.147)

where

$$f_1 = \left[ \frac{1}{r} \cos \theta \sin \alpha - \sin \theta \sin \alpha \frac{\partial \theta}{\partial r} + \cos \theta \cos \alpha \frac{\partial \alpha}{\partial r} \right] \cos \alpha$$
$$+ \left( \sin \theta \cos \alpha \frac{\partial \theta}{\partial r} + \cos \theta \sin \alpha \frac{\partial \alpha}{\partial r} \right) \sin \alpha \cos^2 \theta,$$

(1.148)

$$f_2 = \left( \frac{1}{r} \cos \theta \sin \alpha \right)^2 + 2 \frac{1}{r} \cos \theta \sin \alpha \left( - \sin \theta \sin \alpha \frac{\partial \theta}{\partial r} + \cos \theta \cos \alpha \frac{\partial \alpha}{\partial r} \right) + \left( \cos \theta \frac{\partial \alpha}{\partial r} \right)^2.$$

(1.149)

The total change of the elastic energy per unit length of the cylinder is

$$\Delta F = 2\pi \left[ \int_0^R \Delta f dr \right] = \pi \left[ (K_{22} - K_{33})f_1 + K_{33}f_2 \right] r dr = \pi K_{22} \Delta F_1 - \pi K_{33} (\Delta F_1 - \Delta F_2).$$

(1.150)

The total free energy is decreased if

$$\frac{K_{22}}{K_{33}} \leq \frac{(\Delta F_1 - \Delta F_2)}{\Delta F_1}.$$

(1.151)

Instead of using the Euler–Lagrange method to find the exact solutions for $\theta$ and $\alpha$, we use approximations for them. For $\theta(r)$, we use the same solution found in the last section. For $\alpha(r)$, because the boundary conditions are $\alpha(r=0) = 0$, we use the approximation

$$\alpha = Ar^b.$$

(1.152)

We minimize $\Delta F$ with respect to the amplitude $A$ and power $b$ of the twist angle. It is numerically found that when $A = 0.34\pi$ and $b = 0.3$, $\Delta F_1 = 1.14$ and $\Delta F_2 = 0.75$. From Equation (1.151) we have

$$\frac{K_{22}}{K_{33}} \leq \frac{(1.14 - 0.75)}{1.14} \approx 0.34.$$

(1.153)

If the ratio between the twist and bend elastic constants is smaller than 0.34, the total elastic energy is reduced by escaping from the bend deformation to the twist deformation.
Homework Problems

1.1. Consider a nematic liquid crystal. The molecule can be regarded as a cylinder with the length of 2 nm and diameter of 0.5 nm. The molecule has a permanent dipole moment of $10^{-29}$ m · C at the center of the molecule. The interaction between the molecules comes from the interactions between the permanent dipoles. Calculate the interaction between two molecules in the following cases: (1) one molecule is on top of the other molecule and the dipoles are parallel, (2) one molecule is on top of the other molecule and the dipoles are anti-parallel, (3) the molecules are side by side and the dipoles are parallel, and (4) the molecules are side by side and the dipoles are anti-parallel.

1.2. Using Equations (1.11), (1.25), and (1.31), prove that the entropy of a system at a constant temperature is $S = -k_B \ln \rho > -k_B \sum_i \rho_i \ln \rho_i$.

1.3. Calculate the orientational order parameter in the following two cases. (1) The orientational distribution function is $f(\theta) = \cos^2 \theta$. (2) The orientational distribution function is $f(\theta) = \sin^2 \theta$. $\theta$ is the angle between the long molecular axis and the liquid crystal director.

1.4. Landau–de Gennes theory. For a liquid crystal with the parameters $a = 0.1319 \times 10^5$ J/K · m$^3$, $b = -1.836 \times 10^5$ J/m$^3$, and $c = 4.05 \times 10^5$ J/m$^3$. Numerically calculate the free energy as a function of the order parameter, and identify the order parameters corresponding to the maximum and minimum free energy at the following temperatures. (1) $T - T^* = 4.0^\circ$C, (2) $T - T^* = 3.0^\circ$C, (3) $T - T^* = 2.0^\circ$C, (4) $T - T^* = 1.0^\circ$C, (5) $T - T^* = 0.0^\circ$C, (6) $T - T^* = -10.0^\circ$C.

1.5. Maier–Saupe theory. Use Equation (1.50) to numerically calculate all the possible order parameters as a function of the normalized temperature $\tau = k_BT/v$, and use Equation (1.54) to calculate the corresponding free energy.

1.6. Use Maier–Saupe theory to study isotropic–nematic phase transition of a binary mixture consisting of two components A and B. For molecule A, when its long molecular axis makes the angle $\theta_A$ with respect to the liquid crystal director, the single molecular potential is $V_A(\theta) = -v_{AA}(1-x)S_A(\frac{1}{2} \cos^2 \theta_A - \frac{1}{4}) - v_{AB}xS_B(\frac{1}{2} \cos^2 \theta_A - \frac{1}{4})$. For molecule B, when its long molecular axis makes the angle $\theta_B$ with respect to the liquid crystal director, the single molecular potential is $V_B(\theta) = -v_{AB}(1-x)S_A(\frac{1}{2} \cos^2 \theta_B - \frac{1}{4}) - v_{BB}xS_B(\frac{1}{2} \cos^2 \theta_B - \frac{1}{4})$. $x$ is the molar fraction of component B. The interaction constants are $v_{BB} = 1.05v_{AA}$ and $v_{AB} = 0.95v_{AA}$. Express the normalized temperature by $\tau = k_BT/v_{AA}$. Assume that the two components are miscible at any fraction. Numerically calculate the transition temperature as a function of the molar fraction $x$.

1.7. Consider a nematic liquid crystal cell with the thickness of 10 μm. On the bottom surface the liquid crystal is aligned parallel to the cell surface, and on top of the top surface the liquid crystal is aligned perpendicular to the cell surface. Assume the tilt angle of the liquid crystal director changes linearly with the coordinate $z$, which is in the cell normal direction. Calculate the total elastic energy per unit area. The elastic constants of the liquid crystal are $K_{11} = 6 \times 10^{-12}$N, $K_{22} = 3 \times 10^{-12}$N and $K_{11} = 10 \times 10^{-12}$N.

1.8. The Cano-wedge method is an experimental technique to measure the pitch of cholesteric liquid crystals. It consists of a flat substrate and a hemisphere with a cholesteric liquid crystal sandwiched between them as shown in Figure 1.22(a). At the center, the spherical surface touches the flat surface. On both the flat and spherical surfaces there is a homogeneous alignment layer. The intrinsic pitch of the liquid crystal is $P_\varphi$. Because of the boundary
condition, the pitch of the liquid crystal is quantized to match the boundary condition. In region \( n \), \( h = n(P/2) \). In each region, on the inner side, the pitch is compressed, that is, \( P < P_o \), while on the outer side, the pitch is stretched, that is, \( P > P_o \). Between region \((n-1)\) and region \( n \), there is a disclination ring as shown in Figure 1.22(b). Find the square of the radius of the \( n \)th disclination ring \( r_n^2 \) as a function of the intrinsic pitch \( P_o \), the radius \( R \) of the hemisphere and the ring number \( n \). \( R \gg P_o \) and for small \( r \) only twist elastic energy has to be considered. Hint, \( r_n^2 \) vs. \( n \) is a straight line with a slope dependent on \( P_o \) and \( R \).

1.9. Consider a sphere of radius \( R \). The polarization inside the sphere is \( \vec{P} \). Calculate the electric field at the center of the sphere produced by the polarization. Hint, the polarization can be replaced by a surface charge whose density is given by \( \vec{P} \cdot \vec{n} \), where \( \vec{n} \) is the unit vector along the surface normal direction.

1.10. Using Equations (1.87), (1.91), and (1.93), calculate the dielectric tensor \( \bar{\varepsilon} \) in terms of the order parameter \( S \).

References