Mode expansion for the density profiles of crystal–fluid interfaces: hard spheres as a test case

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Abstract
We present a technique for analyzing the full three-dimensional density profiles of planar crystal–fluid interfaces in terms of density modes. These density modes can also be related to crystallinity order parameter profiles which are used in coarse-grained, phase field type models of the statics and dynamics of crystal–fluid interfaces and are an alternative to crystallinity order parameters extracted from simulations using local crystallinity criteria. We illustrate our results for the hard sphere system using finely resolved, three-dimensional density profiles from a density functional theory of fundamental measure type.

1. Introduction
The study of interfaces between thermodynamically stable bulk phases continues to be of great practical and theoretical interest. One main motivation is simply the fact that every finite sample of material in some environment contains interfaces. If the bulk sample is sufficiently small, the physical properties of these interfaces may even govern the behavior of the whole sample. Additionally, surface-specific processes are widely investigated in materials science.

The liquid–vapor interface appears to be the simplest of these interfaces. For a simple liquid, there is just one scalar order parameter, the one-particle density, which continuously changes when going from the vapor to the liquid. The basic statistical mechanical theory of the liquid–vapor interface was developed a while ago [1] but, of course, there are still open problems such as the question of the correct formulation of effective interface Hamiltonians and the behavior of the wavevector-dependent surface tension [2, 3].

The solid–liquid and solid–vapor interfaces appear to be more difficult to describe. Upon solidifying, the continuous translational symmetry of the liquid (vapor) is broken to a discrete symmetry describing the crystal. Connected to this, besides density a number of additional order parameters are needed which completely characterize the bulk crystal and which should vanish when going over to the bulk liquid (vapor) phase. Phenomenological attempts to capture the liquid–solid transition from the point of view of statistical mechanics have restricted the choice of order parameters to the density and one crystallinity parameter (models of phase field type [4]) and employed ad hoc free energies of Landau–Ginzburg type to incorporate the transition. A more fundamental approach is classical density functional theory (DFT) [1] in which all equilibrium properties are governed by a unique free energy functional of the one-particle density. In this approach, the crystalline order parameters can be related to the Fourier modes of the inhomogeneous density profile. Depending on the level of approximation used, only a few modes are needed (as in phase field crystal models [5]) or many (as in Ramakrishnan–Yussouff models, weighted density functionals, . . . [6]). In practical calculations, however, the density mode expansion is almost always severely truncated.

From the DFT point of view, the full three-dimensional density profile of a solid–liquid interface contains all necessary information about the crystalline order parameter profiles. As remarked, with very few exceptions existing DFT calculations have used restricted parameterizations and none of them have attempted a systematic analysis of the three-dimensional density profile in terms of modes. In this
Consider the density distribution and some conclusions. A discussion on the meaning and measurability of the modes and as well as the free energy modes. Section 4 contains a introduction to the fundamental measure functionals, which contains results of the mode analysis of our DFT data. Here introduce the mode analysis in general terms. Section 3 thereof.

is the number of next neighbors of a particle or a multiple variables which are, however, difficult to capture in theory to analyze simulation data involving solid–liquid interfaces crystal–liquid interface. It appears also to be interesting (using fundamental measure functionals) for the hard sphere paper, we present such a mode analysis of the full density profile which is general and the resulting modes are also quantities which are (in principle) observable in scattering experiments or real-space experiments (confocal microscopy). We exemplify the technique using very accurate DFT data experiments or real-space experiments (confocal microscopy).

quantities which are (in principle) observable in scattering papers. The reciprocal basis is given (in Cartesian coordinates where the axes span the cubic unit cell in reciprocal space) by \( \mathbf{B}_1 = (1, 1, 1) \), \( \mathbf{B}_2 = (1, -1, 1) \) and \( \mathbf{B}_3 = (-1, 1, 1) \). An arbitrary RLV is a linear combination of the \( \mathbf{B}_i \). The shells are characterized by a triple \((m, n, k)\) of natural numbers and the \( \mathbf{K}_j \) belonging to this shell have Cartesian components \( b(\pm m, \pm n, \pm k) \) and permutations thereof. Thus, if \( m, n, k \) are mutually distinct, there is a maximum of 48 RLVs in one shell. The shells with lowest modulus are given by \((1, 1, 1)\), \((2, 0, 0)\) and \((2, 2, 0)\). A listing of the RLV triples up to shell 15 is given in [7] (table 1).

We prepare a planar crystal–fluid interface with the interface normal in the \( z \)-direction and the density profile \( \rho(x, y, z) \). It is tempting to parameterize the density profile using the RLV as before, \[ \rho(x, y, z) = \sum_j \exp(i\mathbf{K}_j \cdot \mathbf{r}) p_j(z), \] but now with a \( z \)-dependent Fourier amplitude \( p_j(z) \) for the reciprocal modes. We call them (density) modes or order parameter profiles. One expects that upon crossing the interface coming from the crystal side, all \( p_j(z) \) go to zero (for nonzero \( \mathbf{K}_j \)). Only for \( \mathbf{K}_j \equiv 0 \) does the value for the associated mode cross from the average density of the crystal to the average density of the fluid. As discussed before, in the bulk crystal all RLVs within one shell with index \( m \) are degenerate. At the interface the degeneracy of the RLVs in one shell is lifted, and we introduce an index \( n \) which distinguishes the possible values of the \( z \)-component of the RLVs. Thus the decomposition becomes \[ \rho(x, y, z) = \sum_{mn} \sum_j p_{mn}(z) \exp(i\mathbf{K}_j_{mn} \cdot \mathbf{r}). \]

The sum over \( j \) is now only for those RLVs within shell \( m \) which have a common value of the \( z \)-component, as expressed by the index \( n \). In the literature, such a decomposition has been used to parameterize the full 3d density profile \( \rho(x, y, z) \) using only the leading mode [8–10] in order to facilitate a simplified order parameter description of the crystal–fluid

| \( \rho_0 \sigma^3 \) | \( \rho_b \sigma^3 \) | \( (\beta F/N)_{\text{ex}} \) | \( \beta\mu_{\text{ex}} \) | \( \beta p_{\text{ex}} \sigma^3 \) | \( n_{\text{vac}} \) |
|----------------|----------------|----------------|----------------|----------------|----------------|
| FMT            | 1.039          | 0.945          | 4.96           | 16.38          | 11.87          | \( 2 \times 10^{-5} \) |
| SIM            | 1.041\textsuperscript{a} | 0.940\textsuperscript{b} | 4.96\textsuperscript{b} | 16.09\textsuperscript{b} | 11.58\textsuperscript{a} | \( 3 \times 10^{-4d} \) |
| Error          | <0.001         | <0.001         | <0.01          | 0.01           |                |                |

\( \textsuperscript{a} \) From [19].

\( \textsuperscript{b} \) Free energy for \( \rho_0 \sigma^3 = 1.041 \) [19] using an improved fit in the form of the Speedy equation of state from [21].

\( \textsuperscript{c} \) Chemical potential for \( \rho_b \sigma^3 = 0.940 \) [19] from the Carnahan–Starling equation of state.

\( \textsuperscript{d} \) From [20].

\( \textsuperscript{e} \) From [24].

| \( \beta_1(100) \sigma^2 \) | \( \beta_1(110) \sigma^2 \) | \( \beta_1(111) \sigma^2 \) | \( \beta_1(211) \sigma^2 \) | \( \beta_1(210) \sigma^2 \) |
|----------------|----------------|----------------|----------------|----------------|
| FMT\textsuperscript{a} | 0.69 | 0.67 | 0.64 | 0.65 | 0.67 |
| SIM\textsuperscript{e} | 0.64 | 0.62 | 0.60 | 0.61 | 0.62 |
| Error | 0.01 | 0.01 | 0.01 | 0.01 | 0.01 |

2. Mode analysis

Consider the density distribution \( \rho_{\text{cr}}(x, y, z) \) in a perfect crystal. It can be Fourier expanded into a discrete sum,

\[ \rho_{\text{cr}}(x, y, z) = \sum_j P_j \exp(i\mathbf{K}_j \cdot \mathbf{r}), \]

where \( \mathbf{K}_j \) denotes the set of all reciprocal lattice vectors (RLVs) and \( P_j \) are the associated Fourier amplitudes. Owing to the symmetry of the crystal under consideration, not all \( P_j \) are independent of each other. It is convenient to group the \( \mathbf{K}_j \) in shells with index \( m \) where all \( \mathbf{K}_j \) belonging to one shell can be transformed into each other under the discrete symmetry group of the crystal under consideration (and thus all \( P_j \) associated with these \( \mathbf{K}_j \) are equal). As an example, for the face-centered cubic (fcc) crystal the reciprocal lattice is of body-centered cubic (bcc) symmetry. Let \( a \) be the side length of the cubic unit cell of fcc and correspondingly \( b = 2\pi/a \) the side length of the cubic unit cell of bcc in reciprocal space. The reciprocal basis is given (in Cartesian coordinates where the axes span the cubic unit cell in reciprocal space) by \( \mathbf{B}_1 = (1, 1, 1), \mathbf{B}_2 = (1, -1, 1) \) and \( \mathbf{B}_3 = (-1, 1, 1) \). An arbitrary RLV is a linear combination of the \( \mathbf{B}_i \). The shells with lowest modulus are given by \((1, 1, 1), (2, 0, 0)\) and \((2, 2, 0)\). A listing of the RLV triples up to shell 15 is given in [7] (table 1).

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\[ \rho(x, y, z) = \sum_{mn} \sum_j p_{mn}(z) \exp(i\mathbf{K}_j_{mn} \cdot \mathbf{r}). \]

The sum over \( j \) is now only for those RLVs within shell \( m \) which have a common value of the \( z \)-component, as expressed by the index \( n \). In the literature, such a decomposition has been used to parameterize the full 3d density profile \( \rho(x, y, z) \) using only the leading mode [8–10] in order to facilitate a simplified order parameter description of the crystal–fluid
interface. Consistency with the bulk crystal solution $\rho_{ct}$ requires $\rho_{mn}(z \to -\infty) = P_m$, thus in a leading-mode description either the crystal description is also truncated [10] or higher-than-leading modes are parametrized proportional to the leading mode [9].

For a given 3d density profile $\rho(x, y, z)$ the extraction of the modes $\hat{\rho}_{mn}(z)$ does not appear to be straightforward since they cannot be projected out using a Fourier integral. Consider the Fourier transform of the 3d density profile

$$\hat{\rho}(k_x, k_y, k_z) = \int L_x L_y \sum_j \delta^2 (k_j + (K_{j,z})_{mn}) \hat{\rho}_{mn}(k_z + (K_z)_{mn}). \quad (4)$$

Here, the lateral dimensions of the system with interface are denoted by $L_x$ and $L_y$. The wavevector in the interface plane, $k_j = (k_x, k_y)$, is restricted to the discrete values allowed by the RLVs, $(K_{z})_{mn}$. The Fourier transforms of the modes, $\hat{\rho}_{mn}$, are shifted by the $z$-components of the associated RLVs, $(K_{z})_{mn}$, which are by our convention fully specified by the indices $m, n$. Hence we see that the $\hat{\rho}_{mn}(k_z)$ can be viewed as being centered around $-(K_z)_{mn}$ and are well separable from each other as long as their range in Fourier space does not extend further than the separation between two neighboring $(K_z)_{mn}$. This in turn is given by $2\pi/d_{[0]}$, with $d_{[0]}$ being the $z$-separation of lattice planes in the orientation [0]. This condition on the mode separability is a very physical one: it means that the density modes should cross across the interface much more smoothly than the actual density profile $\rho(x, y, z)$.

We illustrate the mode separability using explicit DFT results for the crystal–fluid interface in the hard sphere system; for more details see section 3. For the [100] orientation, figure 1(a) shows the laterally averaged density profile

$$\rho_{av}(z) = \frac{1}{L_x L_y} \int_0^{L_x} dx \int_0^{L_y} dy \ \rho(x, y, z), \quad (5)$$

and figure 1(b) its Fourier transform, corresponding to $\hat{\rho}(0, 0, k_z)$. This Fourier transform consists of localized ‘lumps’ around the values $i \pm \pi/a (i = 0, 1, 2, \ldots)$ which mark lattice planes separated by the distance $d = a/2$ (which is what we expect for the laterally averaged density of an fcc crystal). The inverse Fourier transform of the isolated ‘lumps’ (centered around 0) gives the associated modes in real space. As indicated graphically in figure 1(b), the Fourier transform of a particular mode can be isolated by cutting out the Fourier transform $\hat{\rho}(0, 0, k_z)$ symmetrically around $i \pi/a$ with a width $4\pi/a$. We formalize this a bit more generally as

$$\tilde{\rho}_{mn}(k_z) = \tilde{\rho}(K_x, K_y, k_z + (K_z)_{mn}) \ g(k_z), \quad (6)$$

where $(K_{z})_{mn} = (K_x, K_y)$ denotes the in-plane components of the RLV of interest for mode $(mn)$ (again, there is an additional degeneracy, expressed by $j$). The function $g(k_z)$ is a suitable cutoff function, varying monotonically from 1 at around $k_z = 0$ to zero near $k_z = 2\pi/d_{[0]}$ and with the additional properties

$$g(k_z) = 0 \quad (|k_z| > 2\pi/d_{[0]}),$$

$$g(k_z) - g(-k_z) = 0,$$

$$g(k_z) + g \left( \frac{2\pi}{d_{[0]}} - k_z \right) = 1 \quad (k_z \in \left[ 0, \frac{2\pi}{d_{[0]}} \right]). \quad (7)$$

Note that these conditions are required to have (6) consistent with (4). A practical choice (which violates the first condition a tiny bit) is

$$g(k_z) > 0 = \frac{1}{2} \left( 1 - \text{erf} \left( \alpha \left[ \frac{k_z d_{[0]}}{\pi} - 1 \right] \right) \right). \quad (8)$$

where $\alpha$ tunes the width of the error function kink. Note that for $\alpha \to \infty$ the cutoff function becomes $g(k_z > 0) = \theta(\pi/d_{[0]} - k_z)$, literally corresponding to the graphical procedure indicated in figure 1(b). The disadvantage hereby is that through a hard cutoff in Fourier space unwanted oscillations arise in real space which simply might be due to insufficient resolution of the density profile in real space and are not due to physical oscillations of the modes. We found $\alpha > 3$ to be a convenient choice.
We had introduced the modes $p_{mn}(z)$ as belonging to the RLV shell $m$ but differing in the $z$-component of the associated RLV, as indicated by $n$. If that $z$-component is zero, the mode will be purely real. If for a particular $n$ that $z$-component is nonzero, the mode will be in general complex since from equation (6) it cannot be expected that the Fourier transform of the mode is symmetric in $k_z$. The reality of $\rho(x, y, z)$ requires that in the same shell $m$ there is a mode $p_{mn}(z)$ for which the $z$-component of the associated RLV is minus the previous one and which is the complex conjugate of $p_{mn}(z)$. Hence can define real-valued modes by separating the real and imaginary parts,

$$p_{mn}(z) = \frac{p_{mn}(z) + p_{mn}(z)}{2}, \quad p_{mn}^-(z) = \frac{p_{mn}(z) - p_{mn}(z)}{2i},$$

and the $p_{mn}^-(z)$ have the obvious interpretation of phase shifts of the associated density mode oscillations across the interface.

3. Results for the hard sphere crystal–fluid interface

3.1. Fundamental measure DFT

We apply the mode expansion technique introduced in the previous section to DFT results for the full crystal–fluid interface density profile. DFT is built on the existence of a unique free energy functional of the one-particle density field $\rho(r)$,

$$F[\rho] = F^\text{id}[\rho] + F^\text{ex}[\rho],$$

$$\beta F^\text{id}[\rho] = \int d^3r \rho(r) (\ln(\rho(r)) \Lambda^3 - 1),$$

which can be split into the exactly known ideal gas part $F^\text{id}$ (\& is the de-Broglie wavelength, $\beta = 1/k_B T$ is the inverse temperature) and a generally unknown excess part $F^\text{ex}$. The equilibrium density $\rho_\text{eq}(r)$ in the presence of an external (one-particle) potential $V^\text{ext}(\rho)$ is then given by

$$\delta F[\rho] \left|_{\rho = \rho_\text{eq}} \right. = \mu - V^\text{ext}(\rho),$$

where $\mu$ is the imposed chemical potential (e.g. by requiring a certain bulk density far away from the region where the external potential acts). For the description of the equilibrium density profile between crystal and fluid, $V^\text{ext} = 0$. Thus the bulk crystal far away from the interface on one side appears as a self-sustained, periodically inhomogeneous fluid at the coexistence chemical potential $\mu_\text{coex}$.

For the hard sphere system, fundamental measure theory (FMT) allows the construction of very precise functionals [11–13]. Essentially, FMT postulates an excess free energy with a local free energy density in a set of weighted densities $n_\alpha$,

$$F^\text{ex}[\rho] = \beta^{-1} \int d^3r \Phi(n_\alpha(r)).$$

The weighted densities are constructed as convolutions of the density with weight functions, $n_\alpha(r) = \rho * w^\alpha(r)$. The weight functions reflect the geometric properties of the hard spheres. For one species, the weight functions include four scalar functions $w^3, w^1, w^2$, two vector functions $w^0, w^1$ and a tensor function $w^4$ defined as

$$w^3 = \delta(r - |r|), \quad w^2 = \delta(r - |r|), \quad w^0 = \frac{w^2}{4\pi R^2}, \quad w^1 = \frac{2}{4\pi R^2}, \quad w^4 = \frac{\vec{r} \cdot \vec{r}}{R^2} \delta(r - |r|).$$

Here, $R = \sigma / 2$ is the hard sphere radius. Using these weight functions, the corresponding scalar weighted densities $n_0 - n_3$, vector weighted densities $n_1, n_2$ and one tensor weighted density $n_4$ are defined. In constructing the free energy density $\Phi$, arguments concerning the correlations in the bulk fluid, certain geometric consistencies and arguments pertaining to strongly inhomogeneous systems are used [13]:

$$\Phi([n_\alpha(r)]) = -n_0 \ln(1 - n_3)$$

$$+ \varphi_1(n_3) \frac{n_1 n_2 - n_1 \cdot n_2}{1 - n_3} + \varphi_2(n_3) \times \frac{3(-n_2 n_2 - n_2 n_3 \rho n_k_j)}{16\pi(1 - n_3)^2}$$

$$= \frac{3(-n_2 n_2 - n_2 n_3 \rho n_k_j)}{16\pi(1 - n_3)^2}.$$ (14)

Here, $\varphi_1(n_3)$ and $\varphi_2(n_3)$ are functions of the local packing density $n_3(r)$. The most accurate functional to date is the tensor version of the recently introduced White Bear II (WBII) functional [14] and corresponds to the choice

$$\varphi_1 = 1 + \frac{2n_1 - n_2 + 2(1 - n_3) \ln(1 - n_3)}{3n_3},$$

$$\varphi_2 = 1 - \frac{2n_3 - 3n_2 + 2(1 - n_3)^2 \ln(1 - n_3)}{3n_3^2}.$$ (15)

This functional has proved to be very reliable for strongly inhomogeneous fluids [15, 16] and also wall–fluid surface tension data at densities close to freezing compare excellently with recent simulation data [17].

3.2. Thermodynamic properties of the crystal and the crystal–fluid interface

In equilibrium, hard spheres in their crystalline state form an fcc lattice, although the free energy difference to hcp is very small, about $10^{-3} k_B T$ per particle [18]. Bulk coexistence properties (densities, chemical potential, pressure, free energy) have been widely investigated by simulations and there seems to be consensus on the respective values with an error margin below 1%. The bulk crystal has been investigated in [21] using full minimization of the FMT functional discussed above and very good agreement has been found with simulations. The coexistence density on the liquid side is overestimated by 0.5%; this results in an overestimation of the coexistence chemical potential and pressure of about 2%. The relative vacancy concentration $n_{\text{vac}} = 2 \times 10^{-5}$ at coexistence is too small by a factor
of 10 compared to simulations; however, an assessment of possible systematic errors in the simulations is not available. The FMT results for $n_{\text{vac}}$ improve significantly on results from Ramakrishnan–Yussouff and weighted density functional theories where $n_{\text{vac}} \sim 0$. Also, in contrast to these earlier approaches, the FMT crystal density profiles agree with simulations with respect to the width of the crystal peak and the anisotropy in different lattice directions [21]. See table 1 for a summary.

The determination of crystal–fluid surface tensions in simulations has not yet resulted in consistent data across the literature. As an example, for the surface tension in the [110] direction integration methods give $\beta \gamma_{[110]} \sigma^2 = 0.58$ (cleaving, [22]) and 0.64 (umbrella sampling, [23]). Through the capillary wave method, interfacial stiffnesses are accessible but the conversion to tensions is not without difficulties owing to the finite cutoff in the stiffness and tension expansion in cubic harmonics [24]. Here, Laird et al find $\beta \gamma_{[110]} \sigma^2 = 0.58$ (expansion up to order two, [25]) and the newest results give 0.64 (expansion up to order three, [24]). The FMT results (also from [24]) are larger by about 7% and the anisotropies compare well with the simulation results, see table 1. It is perhaps not so surprising that the surface tension in FMT is higher than in simulations since the FMT functional does not capture the very long-ranged fluctuations present in capillary waves. Note that previous theories based on (i) severe approximations to the free energy functional and (ii) very restricted minimizations in density space have produced similar results [26, 27], consequently the agreement in numbers has to be regarded as fortuitous. See also below for the discussion of the density modes which puts simple density parameterizations in perspective.

### 3.3. Density modes

We analyze the density profiles of crystal–fluid interfaces in five different orientations of the interface normal: [100], [110], [111], [211] and [210]. The density is given for a cuboid with edge lengths $L_x$, $L_y$ and $L_z$ which contains the fluid in the middle ($z \sim L_z/2$) and the crystal phase at the boundaries ($z \sim 0$ and $z \sim L_z$). $L_{x,y}$ are given by the edge lengths in the $x,y$-direction, $a_{x,y}$, of the smallest cuboid unit cell of the crystal which has the desired orientation in the $z$-direction. Typically we chose $L_z = 32a_z$. The crystal cuboid unit cells are depicted in figure 2 for the [100], [110], [111] and [211]
orientations and in figure 3 for the [210] orientation. While they are still comparatively easy to imagine for the first four orientations, it is least trivial for the [210] orientation.

As explained in section 2, the density modes $p_{mn}(z)$ are labeled by their RLV shell index $m$ and by $n$ for the possibly different $z$-components of the RLV. The specific RLV assignment within one shell for the index $n$ then depends on the chosen interface orientation. As an example, we show the assignment for the [100] interface in table 2. We call the mode $p_{11}(z)$ the leading crystallinity mode since the associated RLV is in the lowest shell and in the bulk crystal the associated Fourier amplitude $P_1$ is largest. The meaning of $p_{11}(z)$ is linked to the strength of the density oscillations of the close-packed planes across the interface into the bulk liquid. For the [100] interface, the normal to the close-packed planes is not in the $z$-direction. Another mode of significance is the mode $p_{22}(z)$ which is linked to the strength of the density oscillations of square arrays of particles which are in the $x$-$y$-plane. This is the leading Fourier component of the lateral density average (often shown in simulation works) when the average density mode $p_{00}(z)$ has been subtracted.

In figure 4 we show the modes up to $m = 4$ for the [100] interface. In figure 4(a), the real parts of the mode profiles are shown, including the rescaled density mode $p_{00}(z)$ (large dots). The density mode shows a depletion zone in front of the crystalline phase. In previous work on the crystal growth at walls [28], a more pronounced depletion zone has been found in instantaneous profiles and attributed to the finite velocity with which the interface moves. The absence of the depletion zone for equilibrium profiles as seen in [28] can be attributed to the averaging procedure: there the density was averaged between minima of the laterally averaged density profile. Due to the phase shift of the density oscillations across the interface, such an averaging procedure does not give our density mode $p_{00}(z)$; we verified that using the procedure of [28], the depletion zone also vanishes. Further inspection of figure 4(a) shows that the leading crystallinity mode $p_{11}(z)$ displays a smooth, monotonic behavior which corresponds to the expectations on a phase-change order parameter profile and which could be used to define an interface location. However, this obviously involves some arbitrariness: the kinks in higher modes $m \geq 2$ are clearly positioned deeper in the crystal phase and the mode $p_{22}(z)$ (leading crystallinity mode of the lateral density average) differs also in the qualitative shape by showing a local minimum (similar to the depletion zone in $p_{00}(z)$). Furthermore we note that the the interface locations as extracted from the leading crystallinity mode $p_{11}(z)$ and from the density mode $p_{00}(z)$ differ by about one unit cell length $a$. We discuss this further below. In figure 4(b) we show the imaginary parts of the mode profiles, corresponding to the phase shift of oscillations across the interface. Positive values indicate that the distance between maxima becomes larger in the interface region, see also

![Figure 3](image-url)

**Figure 3.** Cuboid unit cells for the [210] orientation of the interface normal (z-direction). Different layers with $z = $ const. are given in different colors. Layers $n = 7–11$ are omitted for clarity. They can be obtained from layer $k = 12 – n$ using the reflection of coordinates $x \rightarrow \sqrt{5}a - x$ for all particles in the layer. The $z$-separation between lattice planes, needed in equations (6) and (7), is given by $a_{[210]} = a/10 = a/(2\sqrt{5})$.

**Table 2.** The assignment of reciprocal lattice vectors to the main shell index $m = 0–3$ and $z$-component index $n$ for the [100] interface. Note that modes with indices $n$ and $\bar{n}$ are related by complex conjugation.

| $m$ | $n$ | $j$ | $K_x$ | $K_y$ | $K_z$ | Significance |
|-----|-----|-----|-------|-------|-------|--------------|
| 0   | 0   | 1   | 0     | 0     | 0     | Average density                      |
| 1   | 1   | 1−4 | ±1    | ±1    | ±1    | Leading crystallinity mode           |
| 2   | 1   | 1−2 | 0     | ±2    | 0     | Leading cryst. mode                  |
| 3   | 1   | 1−4 | ±2    | ±2    | 0     | Leading density average              |
| 0   | 2   | 1−2 | 0     | ±2    | ±2    |                                             |
| 3−4 | 1   | 0   | 0     | ±2    | 0     |                                             |
| 3−4 | 1   | 0   | 0     | ±2    | ±2    |                                             |
| 3−4 | 1   | 0   | ±2    | ±2    | 0     |                                             |

Note: The lateral density average (LDA) is given by $p_{ll}(z) = \bar{p}_{ll}(z)$, where $\bar{p}_{ll}(z)$ is the lateral density average of $p_{ll}(z)$ over all layers $n$ with $z = n$. This is linked to the strength of the density oscillations across the interface, see also [28].
the crystalline bulk. Line thickness decreases with increasing $m$ interface orientations in the $z$-direction. Within a shell $m = \text{const}$, the RLVs are sorted with increasing $|K_z|$, for simplicity $n$ and $\bar{n}$ have been identified.

| $m$ | $n$ | $K_x$ | $K_y$ | $K_z$ | $K_x$ | $K_y$ | $K_z$ | $K_x$ | $K_y$ | $K_z$ | $K_x$ | $K_y$ | $K_z$ |
|-----|-----|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|-------|
| 1   | 1   | ±1    | ±1    | 0     | ±1    | −1    | −3    | ±1    | −1    | ±3    | 0     | 0     |
| 1   | 0   | ±1    | ±1    | 0     | ±1    | −1    | 3     | ±1    | 1     | 0     | ±2    | ±2    |
| 1   | 2   | 0     | ±1    | ±1    | 0     | ±3    | 0     | 1     | ±1    | −3    | 1     | ±1    |
| 1   | 3   | 0     | ±1    | −3    | 0     | ±1    | −3    | 1     | ±1    | 1     | 0     | 1     |


$^a$ The monotonically varying leading modes shown in figure 6.

Figure 4. Density mode profiles $p_{mn}(z)$ in units of $\sigma^{-1}$ up to $m = 4$ for the [100] interface. Profiles with the same $m$ reach the same value in the crystalline bulk. Line thickness decreases with increasing $m$. (a) Real parts of the profiles, note that the density mode $p_{00}(z)$ has been rescaled and shifted, see the tick labels on the right $y$-axis. (b) Imaginary parts of the profiles.

Table 3. The assignment of reciprocal lattice vectors to the mode indices $m, n$ for the first two main shells and for the orientations used (except [100]). $K_{[n,z]}$ are given in units of $2\pi/a_{[n,z]}$, where $a_{[n,z]}$ are the side lengths of the cuboid unit cells of figure 2 with the desired interface orientations in the $z$-direction. Within a shell $m = \text{const}$, the RLVs are sorted with increasing $|K_z|$, for simplicity $n$ and $\bar{n}$ have been identified.

As is the case for the surface tensions, the modes are also not rotationally invariant. For a new orientation, one takes the rotation matrix which transforms the coordinate system of the [100] unit cell to the cuboid unit cell with the desired orientation (see figure 2) and applies it to the RLVs. The RLVs within a shell $m$ need to be regrouped with respect to their new $z$-components (see table 3). Whenever a $z$-degeneracy is lifted when going to another orientation (such as for the leading mode $m = 1$ which is completely degenerate in the [100] orientation), there will be new modes. As an example, in figure 5 we show the modes up to $m = 2$, both real and imaginary parts, for the [111] interface. The shell $m = 1$ is split into two modes where the associated RLVs in reduced units are given by $(0, 0, 3)$ (for $p_{11}(z)$) and $(0, 2, 1)$ (for $p_{12}(z)$). The reduced units are defined by normalization of the RLV component $x[y, z]$ to the value $2\pi/a_{[y,z]}$, where $a_{[y,z]}$ are the side lengths of the cuboid unit cell, see figure 2.) As can be seen in figure 5, mode $p_{11}(z)$ is similar in shape to the same mode for [100] and corresponds to the decay of the density oscillations of the close-packed $x$-$y$-planes. However, the decays of the density oscillations of the other close-packed planes which are not perpendicular to the interface normal are different, as the behavior of $p_{12}(z)$ shows.

We also found that for the other orientations there is always one leading mode $(m = 1)$, which shows the monotonic decay of the leading mode for the [100] orientation, for the associated RLVs see table 3. The shape is roughly similar for all orientations, and can be roughly fitted to a tanh-profile with width $a$ (see figure 6(a)). The average density mode also qualitatively has the same shape (especially...
the depletion zone) for all orientations, see figure 6(b). Therefore our previous finding holds regarding the clear separation between the interface position as determined from the leading crystallinity mode and the interface position as determined from the average density mode. This separation is approximately of size $a \approx 1.6 \sigma$ which is not small. So, coming from the liquid side, the hard sphere fluid first orders and then densifies, in contrast to the time sequence in homogeneous nucleation where nuclei form by first densifying and then ordering [29].

Since the interparticle interaction is short-ranged, one would expect an (oscillatory) exponential decay of the modes into the respective bulk phases. An analysis within Landau theory predicts an upper bound of the decay length of mode $p_{mn}(z)$ of $1/|K_{\perp, j}^{mn}|$ [30]. A more refined analysis taking into account the direct correlation function of the bulk liquid [8, 31] also finds that the decay lengths decrease with increasing modulus $1/|K_{\perp, j}^{mn}|$ but the magnitude is enhanced by a factor of 2–3. We defer a detailed analysis of the decay lengths to future work and merely remark that the aforementioned simple rules on the decay lengths do not seem to capture the qualitative importance of modes in the approach to the bulk phases, see figure 4. (For example, $p_{00}$ and $p_{22}$ should have the same decay length but the corresponding amplitudes are very different in the approach to the liquid phase. Furthermore, the different $z$-positions of the mode kinks are very important for the relative importance of the modes at a given location on the $z$-axis.)

In view of all the features we have seen in figures 4–6 and discussed above, it is clear that restricted density parameterizations cannot be very reliable. This has been shown already for the hard sphere crystal–liquid interface evaluated with a simpler functional of weighted density type. Full minimization [32] reduces the surface tension by a factor of 2 compared to a parameterization [26]. Also for FMT functionals quite elaborate parameterizations have been tested in [33]. The average surface tension is still larger by about 30% than the results given in table 1 and the ordering $\gamma_{[100]} > \gamma_{[110]} > \gamma_{[111]}$ is not reproduced.

### 3.4. Free energy modes

The primary focus of the analysis in the last section was on the density modes which are observable quantities (see also section 4). Nevertheless, the mode analysis can also be performed for the free energy density $f$ or the grand free energy density $\omega$ using the DFT results. The particular significance of the leading mode $\omega_{00}(z)$ of the grand free energy density is the connection to the surface tension,

$$\gamma = \int_{-\infty}^{\infty} dz \left( \omega_{00}(z) + p_{\text{coex}} \right).$$

In figure 7 we show the leading modes for both the free energy density (a) and the grand free energy density (b). The leading free energy modes for the five different orientations actually look very similar and almost monotonically connect from the higher free energy density of the solid to the

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**Figure 5.** Density mode profiles $p_{mn}(z)$ in units of $\sigma^{-3}$ up to $m = 2$ for the $[111]$ interface. Note that the density mode $p_{00}(z)$ has been rescaled and shifted, see the ticks on the right $y$-axis.

**Figure 6.** (a) Leading mode in units of $\sigma^{-3}$ for all five orientations and a fit of the form $p_{00}(z) = 0.45(1 + \tanh((z - z_0)/(wa)))$ with $z_0/a = 6.35$ and width $w = 1.0$. (b) Average density mode for all five orientations. For comparison the fitted leading mode, rescaled and shifted to match the asymptotic densities, is given.

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lower free energy density of the liquid (save for a small hump on the crystalline side). It becomes clearer in the plot for the grand free energy density that there is indeed a broadening of the interface (in free energy terms) in the sequence [100]–[211]–[110]–[111]–[210]. This is also in rough agreement with the widths of the leading crystallinity modes shown in figure 6(a).

Modes other than the leading one appear to be without much physical content. However, these may be used to gauge more phenomenological approaches such as phase field crystal models [5].

4. Summary and conclusion

In this paper we have presented an analysis of the full 3d density profile of a crystal–liquid interface in terms of density modes associated with the reciprocal lattice vectors of the bulk crystal. We have exemplified the extraction of the density modes on finely meshed density functional data obtained for the hard sphere system and found a number of interesting results:

- a separation of about $1.6\sigma$ between the interface location as determined by the average density and the interface location as determined by the leading crystallinity mode;
- a weak density depletion zone just in front of the bulk crystal which is essentially due to the increase of density oscillation wavelength in the interface region;
- strongly non-monotonic mode profiles also for next-to-leading modes.

The stronger density depletion found in [28] for the case of a moving interface (crystal growth at a hard wall) is of different origin: for a moving interface, the crystal accumulates material from the fluid phase which cannot be instantaneously provided from the fluid reservoir. Note also that average densities can be defined in a coarse grained way via averages between layers (maxima of laterally averaged density profiles). For our DFT results, such a discrete average density does not show a depletion zone. However, the prescription adopted here (average density defined as the lowest Fourier mode) naturally fits into the general mode expansion and allows for a continuous profile, in contrast to the definition via the average between layers. See also [34] for a simulation study on the crystal–fluid interfaces in the Lennard-Jones system where different density averages are discussed and compared; at low temperatures the average density looks similar to our results.

We suggest that simulation data on crystal–fluid interfaces should be analyzed for these modes. According to equation (6), the Fourier transform $\tilde{\rho}_{mn}(k_z)$ of the mode $(mn)$ is obtained by ‘cutting out’ the $k_z$-component of the Fourier transform of the 3d density profile $\tilde{\rho}(k_x, k_y, k_z)$ around the reciprocal lattice vector $\mathbf{k} = (K_{j})_{mn}$ (different $j$ for an index pair $(mn)$ signal mode degeneracy). Practically, this means just averaging $\langle \exp(i(K_{\perp})_{mn} \cdot \mathbf{r}_\perp) \rangle$ in the $x$-$y$-plane as a function of $z$. The average would guarantee that the statistics would be good enough for standard runs so that one would obtain a smooth Fourier transform and would be able to perform the extraction procedure. Alternatively, the Fourier transform in the $z$-direction can also be sampled directly. This procedure appears to be less prone to statistical uncertainty than sampling a full 3d profile.

In principle, the Fourier transforms of the modes $\tilde{\rho}_{mn}(k_z)$ are observable quantities in scattering experiments. In practice, one would need to irradiate a volume containing the crystal, the liquid and the interface, fix the lateral momentum transfer to a particular $(K_{\perp})_{mn}$ and finely scan the momentum transfer in the $z$-direction. In view of additional effects like the incoherent background and the finiteness of the volume with a well-defined interface orientation, this seems to be a very demanding task. Previous studies on surface melting using LEED [35], for instance, only obtained some integral information on the modes by recording a smaller Debye–Waller factor (enhanced decrease in peak intensity) for a certain reciprocal lattice vector due to the decay of the associated mode. The theoretical analysis made at that time was very rough and could be improved using the techniques presented here.

It is perhaps easier to use confocal microscopy for the observation of colloidal crystals [28, 36, 37] and to carry out
a planar averaging as described for the analysis of simulation data. In fact, confocal studies on crystals have previously frequently employed techniques which are borrowed from simulation such as bond-order analysis. As remarked in the introduction, observables connected to the bond order are difficult to handle theoretically, so the mode analysis of simulation and confocal experiments will be beneficial for development of the theory.

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