Effective equilibrium states in the colored-noise model for active matter II. A unified framework for phase equilibria, structure and mechanical properties

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Abstract. Active particles driven by colored noise can be approximately mapped onto a system that obeys detailed balance. The effective interactions which can be derived for such a system allow the description of the structure and phase behavior of the active fluid by means of an effective free energy. In this paper we explain why the related thermodynamic results for pressure and interfacial tension do not represent the results one would measure mechanically. We derive a dynamical density functional theory, which in the steady state simultaneously validates the use of effective interactions and provides access to mechanical quantities. Our calculations suggest that in the colored-noise model the mechanical pressure in the coexisting phases might be unequal and the interfacial tension can become negative.

Keywords: active matter, coarse-graining, Brownian motion, correlation functions
1. Introduction

Our understanding of equilibrium fluids tells us that all phases of a fluid at coexistence have the same pressure. Each phase boundary formed in the system increases its free energy, otherwise a spontaneous mixing occurs: the interfacial tension is positive. These familiar thermodynamic concepts need to be carefully reconsidered when the underlying time-reversal symmetry in equilibrium is broken, e.g. by a self-propelled motion of
active particles. The most striking observation is the phase separation of colloidal particles with purely repulsive interactions, only triggered by increasing their activity [1]. The corresponding phase diagram is similar in form to that of an equilibrium liquid-gas transition [2], However, a discontinuous drop of the virial pressure at the onset of phase separation [3] and a negative interfacial tension [4] measured in computer simulations by the virial theorem underline the exceptional role of active matter.

Several different routes have been explored [5–12] to obtain a thermodynamic description of the inherently non-equilibrium behavior of an active fluid in the steady state. The phenomenology of motility-induced phase separation is now well established in continuum models based on empirical arguments [13–17] or coarse-graining strategies [18, 19]. The driving force is a generic slow-down mechanism in the vicinity of other particles, e.g. due to collisions [1] or chemical signaling [20], resembling a passive system with attractive interactions. Two prominent models describing these experimental observations [16] are (interacting) active Brownian particles (ABPs) with a self-propulsion of a constant magnitude in the direction of the instantaneous orientation, and particles propelled by a velocity-dependent swim speed.

One of the most important challenges for active thermodynamics is to understand the role of pressure. Firstly, the notion of an active pressure is a matter of definition, even for a non-interacting fluid [21]. Secondly, there is no obvious link between the thermodynamic pressure derived from a governing free energy and the mechanical force-balance condition at the system boundaries. Finally, it depends on the particular model system whether the equality of mechanical pressure constitutes an appropriate criterion for phase coexistence [6, 9] or whether it is a state function at all [22]. Very recently, the mechanical contribution due to activity has been identified for ABPs [6, 23]. Much less is known about the chemical potential [24], i.e. the work necessary to insert a particle. Understanding this quantity would be an important step to develop the grand-canonical Monte-Carlo techniques required to observe interfacial phase behavior [11].

A model not considered in [9], but of particular theoretical appeal, consists of particles whose self-propulsion is mimicked by a fluctuating colored-noise variable. The resulting physics exhibit some intriguing similarities to equilibrium systems. For example, non-interacting particles can be described by introducing an effective temperature [12, 25] and at low activity there exists a regime where the principle of detailed balance still holds, even in the presence of interactions [7]. Going one step further, there exist convenient approximation schemes [26, 27] towards a system generally obeying detailed balance, the starting point of several effective-equilibrium studies on a microscopic level [10–12, 28–33]. Without further empirical input it is possible to calculate the $N$-body probability distribution [28] and an effective interaction potential [10, 12, 33] describing phase separation in a purely repulsive system [10] and related interfacial phase transition phenomena [11]. Considering a one-dimensional system at low activity, this effective-potential approximation (EPA) was shown to perfectly agree with simulation results for the full non-equilibrium colored-noise model in situations with [32] and without [29] a non-vanishing probability current. Recently, it has been demonstrated within the the effective equilibrium model that the mechanical and thermodynamical results for pressure and interfacial tension only coincide at lowest order in the activity [31].

In this paper we provide a new perspective on the EPA for the colored-noise model [10, 12, 33] and demonstrate that this microscopic approach is also capable of making

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predictions beyond the fluid structure. To this end we introduce the effective equilibrium approximation for the colored-noise model in section 2 and derive in section 3 an effective dynamical density functional theory (DDFT) \[34–37\] generalizing the original result of \[11\] by including an effective diffusion tensor. Applied in the steady state, this approach admits (i) an effective free energy \[11\] yielding coexisting densities, density profiles and correlation functions, (ii) a mechanical stability condition which we use in section 4 to modify the thermodynamic results for pressure and interfacial tension to obtain a definition consistent with the measurement in simulations and (iii) explicit calculations of these quantities without requiring further input. We compare in section 5 the different routes to make theoretical predictions and discuss that our framework does not, in principle, require the crude \[33, 39\] restriction to pairwise forces. In section 6, we conclude by comparing our findings with other theoretical frameworks.

2. Effective equilibrium approach for the colored-noise model

In the following we consider \(N\) active particles ‘propelled’ by the Ornstein–Uhlenbeck processes \(v_i(t)\), i.e. stochastic variables with zero mean and the non-Gaussian correlator \(\langle v_i(t)v_j(t') \rangle = (D_a/\tau_a)\delta_{ij} \exp(-|t - t'|/\tau_a)\). The active character enters via the finite orientational decorrelation time \(\tau_a\) and \(D_a\) is the active diffusion coefficient. The corresponding overdamped \(N\)-body Langevin equations read

\[
\dot{r}_i(t) = \gamma^{-1}F_i(r_1, \ldots, r_N) + v_i(t) \tag{1}
\]

where \(\gamma\) is the friction coefficient. We assume pairwise additive interaction forces

\[
F_i(r^N) = -\nabla_i U(r^N) \tag{2}
\]

arising from a many-body interaction potential \(U(r^N)\) due to one-body external fields \(\nu(r_i)\) and the interparticle potentials \(u(r_i, r_k) = u(|r_i - r_k|)\), such that

\[
F_i(r^N) = -\nabla_i \left( \nu(r_i) + \sum_{k \neq i} u(r_i, r_k) \right).
\]

For the reason of simplicity, we have neglected the contribution of translational Brownian diffusion in equation (1).

2.1. The multidimensional Fox approach

Although equation (1) does not resolve particle orientations, the non equilibrium nature of active particles becomes obvious in the impossibility of deriving an exact Smoluchowski equation describing the time evolution of the probability distribution \(f_N(r^N, t)\), as the dynamics are always non-Markovian when a colored-noise variable \(v_i(t)\) is involved. Following the multidimensional generalization \[10, 32, 39\] of the Fox approach \[27\], we obtain the following approximate Smoluchowski equation

\[
\frac{\partial f_N}{\partial t} = - \sum_{i=1}^{N} \nabla_i \cdot \sum_{k=1}^{N} D_{ik} \cdot (\beta F_{i\text{eff}}^k - \nabla_k) f_N \tag{3}
\]
with the inverse temperature $\beta = (k_B T)^{-1}$. This result gives rise to effective Markovian dynamics and thus allows an effective equilibrium description. In equation (3) we identify the two central quantities of our theory. Firstly, the effective forces

$$\beta \mathbf{F}_k^\text{eff}(\mathbf{r}^N) = \sum_j D_{jk}^{-1} \cdot \beta \mathbf{F}_j - \nabla_k \ln(\det D_{[N]}),$$

which for an interacting system are not anymore pairwise additive, and, secondly, the effective diffusion tensor $D_{[N]} = D_{[N]/(\beta \gamma)}$ with the components

$$D_{ij}^{-1}(r^N) = \frac{1}{D_a} \left( \delta_{ij} + \tilde{\tau} \nabla_i \nabla_j \mathcal{U}(r^N) \right)$$

of its inverse $D_{[N]}^{-1}$ where $\tilde{\tau} := \tau_a / \gamma$ and $D_a := \beta \gamma D_a$. This quantity comprises the total contribution of activity to the system, as it becomes trivial in the absence of activity ($\tilde{\tau} = 0$). Including the Brownian translational diffusion in equation (1), the form of the inverse diffusion tensor becomes more complicated [33], compare appendix A.

As an alternative to the Fox approach, applying the unified colored noise approximation (UCNA) [26] to equation (1) yields another evolution equation [12, 28, 33] which differs from equation (3) by a factor $D_{[N]}$. As will become clear later, it is rather instructive that we use the Fox picture here. In the (current-free) steady state with $f_N(\mathbf{r}^N, t) \to P_N(\mathbf{r}^N)$, however, both approximation schemes coherently yield [33]

$$0 = \sum_j D_{jk}^{-1} \cdot \beta \mathbf{F}_j P_N - \nabla_k P_N - P_N \nabla_k \ln(\det D_{[N]}) \simeq \beta \mathbf{F}_k^\text{eff} P_N - \nabla_k P_N,$$

$$0 = \beta \mathbf{F}_k^\text{eff} P_N - \sum_k \nabla_k \cdot (D_{ki} P_N) \simeq \sum_k D_{ki} \cdot (\beta \mathbf{F}_k^\text{eff} P_N - \nabla_k P_N),$$

where the second line is obtained after multiplying with $D_{ki}$ and summing over $k$.

### 2.2. Interpretation of the two versions of the steady-state condition

Both versions, equations (6) and (7), of the steady-state condition are equivalent in the sense that they result in the steady-state probability distribution [28]

$$P_N(\mathbf{r}^N) \propto e^{-\beta \mathcal{H}_{[N]}(\mathbf{r}^N)},$$

where $\mathcal{H}_{[N]}(\mathbf{r}^N)$ is defined from equation (4) through $\mathbf{F}_k^\text{eff} = -\nabla_k \mathcal{H}_{[N]}$. The EPA [10, 12, 33] amounts to setting

$$\mathbf{F}_i^\text{eff} \approx -\nabla_i \mathcal{U}^\text{eff} = -\nabla_i \left( \nu^\text{eff}(\mathbf{r}_i) + \sum_{k \neq i} u^\text{eff}(\mathbf{r}_i, \mathbf{r}_k) \right)$$

using effective pairwise interactions defined in appendix A as $u^\text{eff}(\mathbf{r}_1, \mathbf{r}_2) = \mathcal{H}_{[2]}(\mathbf{r}_1, \mathbf{r}_2)$ (assuming $\nu = 0$) and $\nu^\text{eff}(\mathbf{r}) = \mathcal{H}_{[1]}(\mathbf{r})$.

The difference in form between equations (6) and (7) suggests an intriguing new interpretation of our theory. Equation (6) reminds of a thermodynamic condition involving a standard ideal-gas contribution $\nabla_k P_N$ and excess terms describing (activity-mediated) interactions. We thus consider an effective free energy functional [11, 12, 29].
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\[ \mathcal{F}^{\text{eff}}[\rho] = \mathcal{F}_{\text{id}}[\rho] + \mathcal{F}^{\text{eff}}_{\text{ex}}[\rho] + \int \text{d}\mathbf{r} \, \nu^{\text{eff}}(\mathbf{r}) \, \rho(\mathbf{r}), \]  

(10)

where the excess free energy \( \mathcal{F}^{\text{eff}}_{\text{ex}}[\rho] \) follows from \( u^{\text{eff}}(r) \) using standard methods (details in appendix A) and \( \beta \mathcal{F}_{\text{id}}[\rho] = \int \text{d}\mathbf{r} \, \rho(\mathbf{r})(\ln(\Lambda^3 \rho(\mathbf{r})) - 1) \) is the ideal-gas term with thermal wavelength \( \Lambda \) arising from \( \nabla_k P_N \). Knowing \( \mathcal{F}^{\text{eff}}_{\text{ex}} \), we can define in appendix A a hierarchy of direct correlation functions \(^34\) to characterize the fluid structure. In equilibrium density functional theory (DFT), the density \( \omega^{\text{eff}} \) of the grand potential is defined by

\[ \int \text{d}\mathbf{r} \, \omega^{\text{eff}}(\mathbf{r}) = \mathcal{F}^{\text{eff}}[\rho] - \mu N, \]

where \( \mu \) denotes an effective chemical potential (introduced to fix the average number of particles). In our approximate treatment of the active fluid we can define the thermodynamic bulk pressure

\[ \beta p^{\text{eff}} = -\omega^{\text{eff}}, \]

(11)

where \( \omega^{\text{eff}} \) is the grand potential density of the uniform system and the interfacial tension

\[ \beta \gamma^{\text{eff}} = \int \text{d}z \left( \beta p^{\text{eff}} + \omega^{\text{eff}}(z) \right) \]

(12)

at a planar interface. However, these formulas do not reproduce simulation results even qualitatively \(^4, 11, 39\).

Another point of view is to consider the first equality in equation (7) and separate external forces \( \mathbf{F}^{\text{ext}}_i = -\nabla_i \nu \) (acting on the boundary of the system) from internal ones due to both interactions and activity. Such a force balance allows us to identify the active pressure \( p = -\mathcal{V} (\{\mathbf{F}^{\text{ext}}_i\}) / (\partial V) \) by equating the virial \(^40\)

\[ \mathcal{V} (\{\mathbf{F}_i\}) := \int \text{d}\mathbf{r}_1 \ldots \int \text{d}\mathbf{r}_N \sum_{i=1}^{N} (\mathbf{F}_i \cdot \mathbf{r}_i) P_N \]

(13)

of the external forces with that of the internal contributions \(^31\). However, such a mechanical expression for the pressure does not necessarily allow us to study phase coexistence \(^6, 9\).

In summary, it appears that we need to consider a different condition depending on which property of the active fluid we are interested in. The second expression in equation (7), however, contains all the desired information. Upon substituting the exact effective force \(^4\), it becomes obvious how to properly calculate the virial of forces \(^13\). This trivial conversion will now serve to illustrate how the thermodynamical and mechanical aspects represented by equations (6) and (7), respectively, can both simultaneously be cast within the framework of DDFT.

3. Unified local force balance and free energy

For the sake of generality, we now return to the dynamical problem, equation (3), derived by (i) enforcing the Markovian property. We will further consider (ii) only pairwise interactions (EPA) instead of the full many-body effective potential, which is necessary to define the effective free energy, equation (10), and (iii) a diagonal diffusion tensor

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$D_{ij} \approx \delta_{ij} D^p$ with a pairwise additive expression for $D^p$ (compare appendix B). Based on approximations (i-iii) we will now derive local force balance condition which contains as an ingredient the effective free energy functional, equation (10), derived in the EPA.

### 3.1. Dynamical density functional theory for effective interactions

Recognizing the analogy of the form of equation (3) with that of a Smoluchowski equation describing hydrodynamic interactions in a passive system [37, 38], we integrate equation (3) over $N-1$ positions. Making the adiabatic assumption of DDFT [35–37] that the correlations in the dynamic system instantaneously follow from those of an equilibrium system at given configuration, we derive in appendix B the main result of this paper, the following equation of motion

$$\beta \gamma \frac{\partial \rho(r,t)}{\partial t} = \nabla \cdot \mathbf{D}(r,t) \cdot \left( \rho(r,t) \nabla \frac{\delta \beta \mathcal{F}^{\text{eff}}[\rho]}{\delta \rho(r,t)} \right)$$

(14)

for the time-dependent one-body density $\rho(r,t)$, where $\mathbf{D}(r,t)$ is a dimensionless ensemble-averaged diffusion tensor. In the following, we are interested only in the steady state and omit the time dependence. For reasons which will become clear at the end of this section, we identify [33]

$$\mathbf{D}^{-1}(r) \simeq \mathbf{D}_I^{-1}(r) = \frac{1}{D_a}(1 + \tilde{\tau}(\nabla \nabla \mathcal{U}))$$

(15)

with the ensemble average of equation (5), where we define $\langle \mathcal{U} \rangle := \mathcal{D} \nu(r) + \int d r' (\rho^{(2)}(r,r')/\rho(r)) \mathcal{D} \nu(r,r')$ for any (nontrivial) differential operator $\mathcal{D}_i$ acting on $\mathbf{r}_i$.

Our central time-evolution equation (14) is equal in form to a DDFT [37] for a passive colloidal system with hydrodynamic interactions, which do not affect the structure and the (osmotic) pressure in equilibrium. To check this analogy for an active system represented in terms of effective forces, we inspect the zero-flux condition

$$0 = \mathbf{D}(r) \cdot \left( \nabla \rho(r) + \rho(r) \langle \nabla \mathcal{U}^{\text{eff}} \rangle \right) \approx \mathbf{D}(r) \cdot \left( \rho(r) \nabla \frac{\delta \beta \mathcal{F}^{\text{eff}}[\rho]}{\delta \rho(r)} \right),$$

(16)

where the first equality is found before representing in appendix B the interaction term with an approximate excess free energy.

Firstly, it is apparent from equation (16) that the fluid structure in a steady state can be characterized alone using equilibrium DFT with effective potentials [11], as the underlying variational principle [34] $\delta \mathcal{F}^{\text{eff}}/\delta \rho = \mu$ always satisfies the equality. Secondly, this force-balance condition suggests that the (effective) thermodynamic pressure $p^{\text{eff}}$ from equation (11) does not represent the proper active pressure $p$ exerted on the system boundaries, in accordance with the discussion in section 2.2. To illustrate this finding we consider the wall theorem [40]

$$\beta p = -\int_0^\infty dz \frac{\partial \beta \nu(z)}{\partial z} \rho(z),$$

(17)

relating the bulk pressure to the density profile at a planar wall at $z=0$ with the bare potential $\nu(z)$. This condition is always fulfilled by the results of a (non-local) equilibrium DFT [34]. In the present case, however, we employ effective potentials

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and consistently obtain the pressure $p^{\text{eff}}$ exerted on the effective wall characterized by $\nu^{\text{eff}}(z)$. It is now obvious from equation (17) that $p \neq p^{\text{eff}}$. The general condition in equation (16) consolidates the DFT approach, as it also describes the balance of mechanical forces. As conjectured in [11], the averaged diffusivity $D(r)$ will play a key role in defining the pressure from bulk properties alone.

### 3.2. Equivalence of structure and pressure from different routes for an ideal gas

To further illustrate the utility of equation (16), we consider non-interacting particles in an external field $\nu(r)$. This problem can be solved exactly within the present model, as $P_N(8)$ factorizes into one-body contributions [12]. Recognizing the equality of $D = D_1 = D_{[1]} = D_{1[1]} = D_a(1 + \hat{\tau}\nabla\nu(r))^{-1}$ in this single-particle ($N = 1$) limit [33], we have $\nabla \beta\nu^{\text{eff}}(r) = D^{-1}(r) \cdot (\nabla \beta\nu(r) + \nabla \cdot D(r))$ with the effective external potential $\nu^{\text{eff}}$ (compare appendix A). Now we plug the resulting effective free energy (with $\mathcal{F}_{\text{ex}} = 0$) into equation (16), which becomes

$$0 = D \cdot \rho \nabla \left( \ln \rho + \beta \nu^{\text{eff}} \right) = \nabla \cdot \left( D \rho \right) + \rho \nabla \beta \nu. \quad (18)$$

The variational principle of equilibrium DFT yields the steady-state profile $\rho(\mathbf{r}) = \rho_0 \exp(-\beta \nu^{\text{eff}}(\mathbf{r}))$ in agreement with the solution of equation (18), where $\rho_0 = \exp(\beta \mu / \Lambda^3)$ is the bulk density.

As discussed in [12] for a planar and spherical geometry and briefly recapitulated in appendix C, equation (18) allows the identification of the bulk pressure $\beta p = D_a \rho_0$ with constant bulk density $\rho_0$ or the different components of the pressure tensor at a curved surface [42], e.g. via the external virial (13) or the wall theorem (17). The effective pressure $\beta p^{\text{eff}} = \rho_0 = \beta p / D_a$ lacks a factor $D_a$, i.e. the bulk value of $D_{zz}$, which, in this special case, can be absorbed into an effective temperature. Alternatively, we can interpret the effective (ideal) pressure $p^{\text{eff}}$ as a local pressure [21], which, in contrast to the mechanical pressure $p$, does not depend on the activity. This close analogy could give a meaning to $p^{\text{eff}}$ also for interacting systems.

### 3.3. Effective free energy from local force balance

In the special case discussed above, the EPA is only one possibility to derive the exact free energy and pressure, see [12]. Moreover, equation (18) can be seen as the first member of an active equivalent of a Yvon–Born–Green (YBG) hierarchy [12, 31], a mechanical force-balance equation. In the presence of interparticle interactions such an equation can only be obtained approximately by considering the inverse of equation (5)

$$D_{ij}(\mathbf{r}^N) = D_a \left( \delta_{ij} - \hat{\tau} \nabla_i \nabla_j U(\mathbf{r}^N) \right) \quad (19)$$

to leading order in $\hat{\tau}$ and integrating equation (7) over $N - 1$ coordinates. The result [12, 33]

$$0 = \rho(\mathbf{r}) \langle \nabla \beta U \rangle + D_a \nabla \cdot (\rho(\mathbf{r})(1 - \hat{\tau} \langle \nabla \nabla U \rangle)) \quad (20)$$

has no obvious connection to thermodynamics: only in the special case of equation (18) it is possible to define a free energy, which allows the structure (density profile) of the fluid to be determined. In general, a free energy [12] reproducing upon functional
differentiation the (mechanical) condition in equation (20) differs from $F^{\text{eff}}$ identified in equation (16). Instead such a strategy amounts to a pseudo-thermodynamical picture obtained from equation (7) by expanding $D_{\delta_{i}}$ with $\delta_{i}$ being the leading order and constructing an excess free energy including all terms but $-\nabla_{i} P_{N}$. However, there is no effective pair potential representing this artificial case, as the solution (8) to equations (6) and (7) is unique [33].

To restore a unified mechanical and thermodynamic condition in the spirit of equation (16) from equation (20) we propose in appendix E an alternative (to the EPA) generalization of the first equality in equation (18) on a simple mean-field level. The other way round, we note that a YBG-like hierarchy can also be obtained from integrating the first equality in equation (6) over $N - 1$ coordinates [33] in a way that is similar to the calculation in appendix B (done in the more general context of DDFT). By doing so, the averaged inverse diffusion tensor $D_{\nabla}^{-1}(r)$, which we use in equation (15), emerges without any approximation on the pairwise inverse tensor from equation (5). In [33] this quantity is then used to establish the connection to equation (20) derived from equation (7), which motivates our choice to consider $D_{\nabla}^{-1}(r) \simeq D_{\nabla}^{-1}(r)$ in the closely related result (16) of the EPA. An alternative motivation of this choice stems from expanding $D_{\nabla}(r_{N})$ up to linear order in $\tau_{a}$ [12, 31].

Finally, we note that the presented calculations also allow us to understand further the nature of the Fox [27] and UCNA [26] approaches to derive an approximate Smoluchowski equation. Starting with the probability current of the UCNA instead of equation (3), the resulting effective diffusion tensor in our DDFT equation (14) would have a completely different form [33]. In the steady state, however, equation (16) equally applies to the UCNA result, as it can be directly derived from equation (7) without having to introduce the adiabatic DDFT argument. Therefore, the Fox approach presented here is more appealing for its self-consistency with the dynamical situation.

### 4. Mechanical properties of interacting active particles

In section 3 we discussed that, within the EPA, the active pressure can be measured indirectly by calculating the density profile at a planar wall, or, as demonstrated for an ideal gas, determined by rescaling the thermodynamic pressure obtained from an effective free energy $F^{\text{eff}}$. Our objective is to generalize the latter method for an interacting active fluid and relate the active pressure and interfacial tension to the thermodynamic results in equations (11) and (12).

#### 4.1. Rescaled effective pressure

The starting point to rescale the effective mechanical properties is the first equality in equation (16) to which we apply in appendix C the virial theorem [40]. In doing so, the effective diffusion tensor $D(r)$ from equation (15) allows us to recover the (averaged) external force $F^{\text{ext}}(r)$. Thus, we may, e.g. establish the connection to the mechanical pressure exerted on the system boundary via equation (13). Far away from such a boundary, we argue that all internal terms which depend on the external potential

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vanish. In $d = 3$ dimensions the diffusion tensor of interest for a fluid that is inhomogeneous at most in the $z$-direction only depends on particle interactions mediated by $u(r)$ and is diagonal. Its components read

$$D_{\alpha\alpha}(r) = D_\alpha \left(1 + \frac{\rho^{(2)}(r, r')}{\rho(r)} \left( \frac{\partial^2 u(r) - \partial_r u(r)}{r} \right) \left( \frac{(r_\alpha - r'_\alpha)^2}{r^2} + \frac{\partial_r u(r)}{r} \right) \right)^{-1},$$

with the Cartesian index $\alpha \in \{x, y, z\}$ and $\partial_r = \partial/\partial r$ denotes the partial derivative with respect to $r = |r - r'|$.

In a homogeneous bulk system, all components $D_{xx} = D_{yy} = D_{zz} = \text{Tr}[D]/3$ are equal and constant. The rescaled active pressure becomes

$$\beta p^{(R)} = \frac{\text{Tr}[D]}{3} \beta p_{\text{eff}} = \frac{\text{Tr}[D]}{3} \rho + \frac{\text{Tr}[D]}{3} \beta p_{\text{ex}}$$

with the effective pressure $p_{\text{eff}}$ given by equation (11). The second step illustrates the separation into ideal-gas and the excess contributions, where $p_{\text{ex}}$ is identified from the virial of the effective force. The derivation and a detailed discussion of this result can be found in appendix C. As it is obvious from equation (22), the effective contribution $\beta p_{\text{ex}}$ to the virial pressure is different from that $\beta p^{(R)} - \rho$ in an active system. As a consequence we can rationalize why the comparison of these two quantities made in [39] does result in a good agreement. These simulations do not reveal a failure of the EPA in general but rather serve to justify the route taken here.

Inspecting our result (21) for the active pressure, we make some interesting observations when it comes to the coexistence of two phases at low (g) and high density (l). In the EPA the calculation of the corresponding densities amounts to evaluating $p_{\text{eff}}(g) = p_{\text{eff}}(l)$, $\mu(g) = \mu(l)$ and $T(g) = T(l)$. Regarding equation (21) it appears that $\text{Tr}[D]$ is a monotonously decreasing function of the density and we thus expect a higher pressure $p^{(R)}(g) > p^{(R)}(l)$ in the dilute phase, as it was also observed in computer simulations for active Brownian particles with periodic boundary conditions [3]. However, this prediction has to be taken with care, as we discuss in section 4.2.

In order to quantify our observations, we perform some model calculations for the active pressure using the effective equilibrium DFT described in appendix A. We fix $\beta^{-1/\tilde{\gamma}}/d^2 = 0.065$ and introduce the Peclét number $\text{Pe} = \sqrt{3D_\alpha \beta d^2/\tilde{\gamma}}$ as a control parameter for activity. Note that the mean-field approximation chosen here differs from that in [11] and results in a more realistic location of the critical point at $\text{Pe} \approx 44.531$. In the homogeneous regime, we have $\rho^{(2)}(r, r') = \rho^2 g(r' = |r - r'|)$, where the radial distribution $g(r) = g_{\text{OZ}}(r)$ in general follows from solving the Ornstein–Zernicke equation in appendix A. This means that for a phase-separating system the active pressure is ill-defined at intermediate densities. We further use the simple approximation of $g_{\text{step}}(r) = \Theta(r/\sigma - 1)$ as a step function ($\sigma$ denotes the effective hard-sphere diameter), which allows for an analytic calculation.

As shown in figure 1(a) the expected drop of the rescaled pressure at the phase transition (for $\text{Pe} = 50$) is apparent for both choices of $g(r)$. Interestingly, our model calculations suggest at intermediate activity ($\text{Pe} = 40$), that the active pressure exhibits a loop that does not admit phase separation. This is because the thermodynamic result
p_{\text{eff}} does not show such a loop in this case and, therefore, we do not find a divergence in the structure factor. If this was the case, the active pressure calculated with $g_{\text{OZ}}(r)$ would fluctuate in this region, as for $P_e=50$. For even smaller activity, the active pressure increases monotonously with increasing density, as expected for a non-phase-separating system. Quite generically, we see that the increase of the active pressure at small densities becomes steeper at higher activity, which can be expected by inspection of the leading term $\beta p_{\text{act}}(R) \propto D_a \rho$ in equation (22). Thus, by showing the activity-dependence of the pressure in figure 1(b), we clarify that our modification of the purely thermodynamical result $p_{\text{eff}}$ conveniently explains the contradiction pointed out in [39] with the expected behavior. The renormalized pressure increases with increasing activity, which is now consistent with simulations.

4.2. Comparison to the direct calculation of the virial pressure

The strange behavior of $p^{(R)}$ at higher densities, i.e. the loop in the absence of phase separation and the sharp decrease when $\mathbf{D}$ approaches zero, as well as, the slightly non-monotonic behavior as a function of $P_e$ appears to be an artifact of our derivation being based on the simplified condition in equation (16) instead of that in equation (7). Therefore it becomes instructive to compare the rescaled result from equation (22) to the virial pressure [31]

$$\beta p^{(V)} = \frac{\text{Tr}[\mathbf{D}]}{3} \rho + \beta p_0 \quad (23)$$

Figure 1. DFT results for the effective thermodynamic $p_{\text{eff}}$ and different versions of the active bulk pressure for $\beta^{-1}\tau/d^2 = 0.065$. (a) Dependence on the bulk density $\rho$ for different Péclet numbers $P_e$. As labeled in the subscripts, we consider our rescaled result $p^{(R)}$ from equation (22) with the theoretical radial distribution $g_{\text{OZ}}(r)$ and the approximation $g_{\text{step}}(r)$ as a step function (see text), as well as the common first term $p^{\text{act}} := \text{Tr}[\mathbf{D}]\rho/3$ (for $g_{\text{OZ}}(r)$ of equations (22) and (23). The dots denote the coexisting densities at $P_e = 50$ and the thin dotted lines denote the unstable region (the results for $g_{\text{OZ}}(r)$ are unphysical in this case). (b) Dependence on $P_e$ for different values of $\rho$. Here we show the virial pressure $p^{(V)}$ from equation (23), calculated with $g_{\text{step}}(r)$, instead of only the first term $p^{\text{act}}$. At small $P_e$, the assumption to neglect the translational Brownian motion is no longer justified. This is corrected here by adding the unit matrix to the averaged diffusion tensor in equation (21), in the spirit of the Fox approach [33] (compare appendix A), such that for $P_e = 0$ we have $p^{(R)} = p_{\text{eff}}$, i.e. the passive pressure.
directly obtained from the first equality in equation (7) via the condition in equation (20). The second contribution \( p_0 \) equals the formula for the virial pressure [40] in a passive system, only depending on the bare interactions. It increases monotonically with increasing density \( \rho \) and is independent of the activity. In appendix C we show how to recover this result from an empirical modification of our more approximative approach.

The first term in equation (23) was argued [31] to be closely related to the swim pressure [6] in an active Brownian system and is the same as that arising in equation (22) from the effective ideal-gas pressure. Indeed, we observe in figure 1(a) that this term shows the expected parabolic behavior of a swim pressure [5, 6, 23]. This is most apparent for \( g(r) = g_{OZ}(r) \), as in this case the diffusion tensor vanishes more rapidly. Approximating once again \( g(r) = g_{\text{step}}(r) \) as a step function, we obtain a simple quadratic expression \( p_0 \propto \rho^2 \) for the virial pressure in equation (23). In contrast, the second term in equation (22) always depends on the density in a more general form, even when a simple radial distribution is assumed to calculate the effective diffusion tensor.

As expected, the behavior of \( p^{(R)}(r) \) and \( p^{(V)}(r) \) at small densities is observed in figure 1(b) to be quite similar over a large range of activities. At very high densities, we recognize in figure 1(a) the limitation of rescaling the effective pressure with the effective diffusivity, since both quantities eventually vanish. We illustrate in appendix C that the terms resulting in the attractive part of the effective interaction potential, which are necessary to describe the phase transition, should cancel in the mechanical picture and thus not significantly contribute to the pressure. This is not the case for the rescaled pressure in equation (22), as only the low-density limit of the diffusion tensor enters the effective potential. A more detailed discussion of this point can be found in appendix D.

We thus conclude that the direct calculation of the virial pressure, resulting in equation (23), is the method of choice when we are not interested in the fluid structure. Since the contribution \( p_0 \) to the pressure due to bare interactions is a monotonously increasing function of the density, this expression is also useful for high densities. Another striking difference is that \( p_{(g)}^{(V)} < p_{(l)}^{(V)} \) appears to jump to a higher value at coexistence, which we already observe in figure 1(a) for the common first term of \( p^{(R)}(r) \) and \( p^{(V)}(r) \). On the downside, as we require the radial distribution function as an input to calculate \( p^{(V)}(r) \), the results from this approach are not self-consistent. It is thus not possible to make a definite statement of how active pressure behaves in our model without going beyond present approximations. Another scenario is that of an equal pressure at coexistence, which is suggested by calculating the pressure on a planar wall following equation (17), when taking into account the wetting transition predicted in [11]. This behavior agrees with the expectation for ABPs [6, 9].

### 4.3. Rescaled and virial formulas for the interfacial tension

Considering now a planar geometry, we define \( \mathcal{D}_{\alpha\alpha} := \iint dxdy \mathcal{D}_{\alpha\alpha}(r) \) with \( \mathcal{D}_{xx} = \mathcal{D}_{yy} \neq \mathcal{D}_{zz} \) and derive in appendix C the rescaled active interfacial tension

\[
\beta \gamma^{(R)} = \int dz \left( \mathcal{D}_{zz}(z) \beta \rho^{\text{eff}} + \mathcal{D}_{xx}(z) \omega^{\text{eff}}(z) + \rho(z) z \partial_z \mathcal{D}_{zz}(z) \right)
\]
at the free interface, where we identify the negative grand potential density $\omega^{\text{eff}}(z)$ with the effective tangential pressure. With unequal bulk pressures at phase coexistence, it is obvious that in our model also the normal pressure, i.e. the terms depending on $D_{zz}$, is not a constant along the interface. However, as expected, there is no contribution to the active interfacial tension $\gamma(R)$ in equation (24) at $|z| \gg 0$ sufficiently far away from the interface, as in the bulk all components of the effective diffusion tensor are equal and constant.

Again, it is instructive to compare equation (24) to the result [31]

$$\beta \gamma^{(V)} = \int dz \left( D_{zz}(z) - D_{xx}(z) \right) \rho(z) + \beta \gamma_0$$

(25)

derived from equation (20), where $\gamma_0 > 0$ is a standard virial expression by Kirkwood and Buff [31, 45] (see appendix C), which is independent of the activity. As discussed in appendix C, the first two terms in equation (24) imply a similar separation into an ideal-gas-like term and an effective virial contribution, which after switching to the DFT picture is not explicit any more. As a major difference to equation (25), we identify in the last term in equation (24) an additional contribution to the normal pressure, which vanishes in either bulk state. The presence of this term is explained in appendix C to counteract a similar contribution arising from the effective interaction potential in an approximate way.

In figure 2 we show how the integrand of the interfacial tension is modified by increasing activity. As for the pressure, we employ the DFT from appendix A with the averaged diffusion tensor in equation (21) and use $g_{\text{step}}(r) = \Theta(r/\sigma - 1)$. In contrast to the effective thermodynamic interfacial tension, with the integrand shown in figure 2(a), the contribution of the activity-modified ideal-gas term in equation (25) is highly negative and outweighs the smaller positive virial contribution $\gamma_0$ (not shown). Therefore, the overall interfacial tension $\gamma_{\text{YBG}}$ becomes negative [4] and figure 2(b) suggests a decrease with increasing activity. However, the behavior of this modified ideal-gas contribution does not become manifest implicitly in equation (24). In contrast, both contributions to the integrand of $\gamma$, shown in figure 2(c) and figure 2(d), are positive and increase with increasing activity.

Figure 2. Different contributions to the integrand of the interfacial tension calculated with $g_{\text{step}}(r)$ for different Peclét numbers increased from Pe = 46 to Pe = 55 in steps of 1. (a) Effective thermodynamic integrand of $\gamma^{\text{eff}}$ without rescaling. (b) Ideal contribution to $\gamma^{(V)}$, equation (25). (c) Rescaled thermodynamic integrand in equation (24). (d) Correction for $\gamma^{(R)}$ by the last term in equation (24).
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Table 1. Overview of different approaches to make explicit calculations within the effective equilibrium approximation for the colored-noise model, i.e. equation (3). The arrows with double line denote exact operations (or established methods) and those with a single line denote approximate operations. Each approximation is also necessary for the subsequent steps.

5. Summary

We developed in section 2.2 a new interpretation of the steady-state condition in the effective equilibrium approximation for the colored-noise model. With the goal of arriving at a predictive theory, it becomes necessary to integrate the steady-state conditions and thereby consider approximate pairwise additive quantities. There are different ways to do so, as reviewed in the first paper of this series [33] and resumed in section 3 with the focus on mechanical properties of the active fluid. The degree of accuracy, discussed in section 4, of the thermodynamic route (‘rescaled’ formulas based on section 3.1) and the mechanical route (‘virial’ formulas based on section 3.3) is reversed when it comes to characterizing the structure of the fluid. The interconnection of these two routes is explained in the following and illustrated in table 1.

5.1. Connection between mechanic and thermodynamic route

The form of equation (6) corresponds to a thermodynamic condition to determine the structure and phase behavior in the steady state. This enables the interpretation of an effective attraction as the driving force of phase separation, which on the two-particle
level can be cast in an effective pair potential [10, 33]. However, even if it was possible to represent the exact effective interaction force within a thermodynamic free energy, we explained that the thermodynamic results for pressure and interfacial tension cannot coincide with the respective definitions based on the balance of mechanical forces. This reflects the non-equilibrium nature of active systems, even in our effective equilibrium model, and is not a consequence of assuming only a pairwise effective force as argued in [39].

The proper way to switch to the mechanical picture provided by equation (7) in order to calculate pressure and interfacial tension is to multiply equation (6) with the effective diffusion tensor, ensuring that the bare interaction forces are recovered from the effective ones that are responsible for the fluid structure. This marks the fundamental difference to a passive system with hydrodynamic interactions (described by an evolution equation similar to equation (3)), where thermodynamical and mechanical pressure are equal and the diffusion tensor does not contribute to the pressure. We argued that the effective ‘thermodynamic’ attraction originating from the term $\nabla_k \ln(\det \mathcal{D}^{(N)})$ does not result in an equal decrease of the mechanical pressure. We stress that these theoretical insights are in general not restricted to the (highly criticized [39]) approximations involved in defining pairwise effective forces [33] to simplify the general condition in equation (7).

5.2. Inconsistency arising from pairwise approximations

Employing a pairwise approximation eventually results in an inconsistency between the thermodynamic route based on equation (6) and the mechanical route based on equation (7), which becomes manifest in either direction. On the one hand, the approach based on the EPA (first introducing an effective free energy to be later recovered in an approximate local force balance equation from equation (16)) seems to underestimate the virial pressure in equation (22). A perhaps more accurate formula (23) can be derived directly from a YBG-like force-balance equation in equation (20). On the other hand, as a prerequisite to do explicit calculations, we require $g(r)$ and therefore an implicit knowledge of the phase diagram, which can only be provided by an appropriate free energy. In this sense, the EPA is particularly appealing as it allows both defining equation (22) (or equation (23)) and providing the required input. In appendix D we discuss how a different choice of the effective diffusion tensor could improve the mechanical EPA results. Vice versa, we derive in appendix E a free energy by combining equation (20) with the effective diffusion tensor from equation (21), such that equations (22) and (23) become practically equivalent. However, due to its mean-field nature, such a free energy cannot be used to predict the fluid structure, in contrast the one derived from the EPA.

Another interesting difference between the mechanical and thermodynamical picture lies in the validity of low-activity expansions. In the thermodynamic picture, the expansion in the parameter $\tilde{\tau}$ is not justified [33] because of the finite radius of convergence of the logarithmic term in equation (6). However, it was shown [31, 33] that at first order of such an expansion the mechanical and the thermodynamical (without the EPA) route to define pressure and interfacial tension are consistent. Moreover, for the EPA we discuss in appendix D that at this level the spurious negative contribution in the formula for the effective virial pressure, which arises from the logarithmic term in
the effective force, can be compensated. As the expansion in $\tilde{\tau}$ converges for all terms present in equation (7), we understand that such an approximation is suitable to derive formulas for mechanical quantities. Hence, we see that it is a difficult task in future work to arrive at a higher level of self-consistency between the typical approximations to simplify equation (7) or equation (6), which cannot be achieved by including higher-order terms in $\tilde{\tau}$ (see the note in table 1).

5.3. Outlook

Despite its simplicity, our DDFT in equation (14) clearly demonstrates the importance of switching between a mechanical and thermodynamical interpretation of the steady-state condition obtained for our model. We stressed that the perhaps most important advantage of the thermodynamic route is the possibility to determine the fluid structure as input for the results obtained from the mechanical route. As an alternative to the resulting bulk formulas based on the radial distribution, the bulk pressure can be calculated from the profile of the one-body density at a wall, see, equation (17), which can be easily calculated with the help of the EPA. It is also interesting to investigate the force on a curved [42] or structured [46] wall along these lines.

Our results should further guide the way to perform more elaborate numerical calculations. For example, the ‘exact’ simulation of passive particles interacting with the effective potential could answer some open questions regarding the behavior of the pressure at coexistence. Moreover, one can determine pair [43] (and higher-order [44]) interaction potentials (which may depend on the density) from a many-body simulation of any suitable active system, or simply take the structural information to calculate the pressure and interfacial tension with the formulas derived from a YBG-like condition in [31] and recovered here. With such a numerical insight, we could also improve the purely theoretical approach by incorporating some semi-empirical corrections of the free energy to account for (many-body) effects ignored in the definition of the effective pair potential.

6. Conclusions

In this paper we have explicitly shown that there is no equal thermodynamical and mechanical interpretation of the effective steady-state condition derived for active particles propelled by colored noise. This main conclusion agrees with the statement of [9], established on a more coarse-grained level for an arbitrary model system. For the study of mechanical quantities of the (non-equilibrium) steady state, the appropriate starting point is equation (7), as the bare interaction forces are separated from all activity-dependent quantities. On the other hand, a thermodynamic theory requires that there is an ordinary passive ideal-gas contribution as in equation (6), as the ‘reference term’ to construct a free energy. In general, we established and quantified that the effective thermodynamical pressure is unequal, although related, to the pressure that an active fluid exerts on its boundary.
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Within the EPA, we approximately separated the governing equation (16) of the steady state including a purely thermodynamic condition and a mechanical ‘correction’ factor $D(r)$. This average diffusion tensor exhibits certain similarities with the concept of an effective temperature. However, as a result of its position dependence and tensorial nature, it is not constant over the whole system but remains irrelevant for calculating phase equilibria. According to the interpretation chosen here all Boltzmann factors just comprise the ordinary thermal weight $\beta$, as in equation (8), whereas all activity dependence relevant for the fluid structure is captured by effective interactions [33]. Only the effective thermodynamic results for pressure and interfacial tension are then to be rescaled according to (22) or (24). In this step it does not appear to be sufficient to use the constant factor $D_\alpha$, which is commonly used to define the effective temperature [12, 25, 28–31] and equals $\beta^{-1}D_{\alpha\alpha}(r)$ only in the absence of any interaction (or in the special case of a linear external potential [12]).

Our generalized DDFT (14) admits a similar form of the one-body current as found phenomenologically by taking into account a density-dependent swim speed [14, 15]. This approach does, however, not admit a direct connection to a force-balance equation in the steady state, as it has been established here. Moreover, in our derivation we obtained from first principles a tensorial diffusivity and a non-local free energy [11] without any empirical input. It would be interesting to apply our approach to study, for example, the coarsening dynamics in the early state of phase separation. In the steady state this analogy suggests that the approximate connection between the excess chemical potential and the swim pressure found in [11] is consistent but requires further investigation.

Since the effective equilibrium system constructed from the EPA respects detailed balance, we can further make contact to the discussion in [17]. The phase separation is driven by (effective) attractive interactions, whereas we showed that the effective diffusivity does not affect phase coexistence. This quantity thus plays a role similar to that of non-integrable gradient terms in the probability current. Coherent with our (approximate) observation from the EPA, the equivalence of thermodynamic pressure in such a situation close to equilibrium can be argued [17] to result in a discontinuous mechanical pressure. Despite this observation our equation (22) suggests that the rescaled pressure is a (wall-independent) state function. In order to clarify this seeming contradiction with some findings for different model systems [9, 22] and identify the role of a swim pressure [6, 23] and active chemical potential [24] in the EPA, it would be enlightening to integrate active particles propelled by colored noise, see, equation (14), into a generalized thermodynamical framework.

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Appendix A. Effective free energy for the full Fox approach

In the main text, we derived the probability distribution $P_N(r^N)$ (8). In the single-particle case $N = 1$, we can thus identify from $P_1(r) = \exp(-\beta \nu^\text{eff}(r))$ the effective external field

$$\nabla \beta \nu^\text{eff}(r) = D_1^{-1} \cdot (\nabla \beta \nu(r) + \nabla \cdot D_1)$$

(A.1)

where $\nu(r)$ denotes the bare interaction of a passive particle. For curved potentials [42, 46], the second term can be conveniently replaced by different approximate forms. This makes the EPA, i.e. the expression on the left-hand side of equation (18) of the main text more suitable for such situations.

Considering a system of $N = 2$ particles interacting with the bare potential $u(r)$ and neglecting the external forces, we find the effective pair potential

$$\nabla_i \beta u^\text{eff}(r) = D_i^{-1} \cdot (\nabla_i \beta u(r) + \nabla_i \cdot D_i)$$

(A.2)

in the diagonal approximation [33]. Another way to derive these effective potentials is to solve equation (6) for the respective number of particles [10, 33]. This strategy becomes necessary if we do not neglect the Brownian translational diffusivity in equation (1) and make the Fox approximation, which then does not admit a closed-form result for $P_N$.

In order to be consistent with our earlier calculations [10, 11], we will construct an excess free energy functional $\mathcal{F}_\text{ex}^\text{eff}[\rho]$ from the effective potential $u^\text{eff}(r)$, where we approximate the effective diffusion tensor in the Laplacian form [33] as

$$D_{ij} \approx \delta_{ij} \left( 1 + D_a \frac{\rho^2}{r^2 + \tilde{\gamma} \tau \partial_r (r^2 \partial_r u(r))} \right).$$

(A.3)

The first term stems from the translational diffusion and the second term constitutes an approximation for equation (5) of the main text. The effective potential is then separated into a repulsive and an attractive contribution [11, 48]. The first gives rise to an effective hard-sphere diameter [11, 47]

$$\sigma = \int_0^{r_0} dr \left( 1 - e^{-\beta u^\text{eff}(r)+\beta u^\text{eff}(r_0)} \right)$$

(A.4)

where $r_0$ is the position of the minimum of $u^\text{eff}(r)$. This term will be treated as the excess free energy $\mathcal{F}_\text{ex}^\text{has}[\hat{\rho}]$ of hard spheres at the rescaled density $\hat{\rho} = \rho \, d^d/\sigma^d$, where we employ Rosenfeld’s fundamental measure theory [41].

Finally, we obtain

$$\mathcal{F}_\text{ex}^\text{eff}[\rho] = \mathcal{F}_\text{ex}^\text{has}[\hat{\rho}] + \int \int dr_1 dr_2 \rho(r_1) \rho(r_2) \frac{u^\text{att}(r_{12})}{2}$$

(A.5)

in a generalized mean-field approximation, where $r_{12} = |r_1 - r_2|$ and $u^\text{att}(r) = 0$ if $r < \sigma$, $u^\text{att}(r) = \tilde{u}^\text{eff}(r)$ if $\sigma < r < r_0$ and $u^\text{att}(r) = u^\text{eff}(r)$ otherwise. By making this choice for the attractive part $u^\text{att}(r)$ of the effective potential, we implicitly assume the form of a step function for the radial distribution function of the reference fluid instead of setting it to unity as in [11]. Given any excess free energy functional, we calculate the direct correlation functions [34]
of order \(n\). When we are interested in the pair structure \((n = 2)\), we find the radial distribution function
\[
g(r_{12}) = 1 + c^{(2)}(r_{12}) \rho \int d\mathbf{r}_3 c^{(2)}(r_{13})(g(r_{23}) - 1).
\]
(A.7)

This result is required to calculate the averaged diffusion tensor in equation (21) of the main text, and, therefore, the active pressure and interfacial tension.

Appendix B. Effective dynamical density functional theory

In this appendix we demonstrate how the approximate DDFT in equation (14) can be deduced from equation (3). As stated in the main text this starting point represents an approximate Markovian time evolution of active particles driven by Gaussian colored noise. We further assume

\[
\mathcal{D}_{ij}(\mathbf{r}^{N}) \approx \delta_{ij} \sum_{l \neq i}^{N} \mathcal{D}^{p}(\mathbf{r}_i - \mathbf{r}_l)
\]

with the yet unspecified contributions \(\mathcal{D}^{p}\) used to construct an approximate pairwise additive representation for the diagonal components \(\mathcal{D}_{ii}\) of the effective diffusion tensor. Note that \(\mathcal{D}^{p}\) should also contain one-body terms, which can be treated in a much simpler way [33, 37] and thus will be conveniently omitted in the following presentation.

Upon substituting the effective force, equation (9), in the EPA we rewrite equation (3) as

\[
\beta \gamma \frac{\partial f_{N}^{\text{eq}}}{\partial t} = \sum_{i=1}^{N} \mathbf{\nabla}_i \cdot \sum_{l \neq i}^{N} \mathcal{D}^{p}(\mathbf{r}_i - \mathbf{r}_l) \cdot \left( \nabla_i f_{N}^{\text{eq}} + f_{N}^{\text{eq}} \nabla_i \nu^{\text{eff}}(\mathbf{r}_i) + f_{N}^{\text{eq}} \sum_{k \neq i}^{N} \nabla_i \beta u^{\text{eff}}(\mathbf{r}_i, \mathbf{r}_k) \right).
\]

(B.2)

Integration of this approximate time evolution for the probability density \(f_{N}(\mathbf{r}^{N}, t)\) over \(N - 1\) coordinates yields \(\beta \gamma \frac{\partial \rho(\mathbf{r}, t)}{\partial t} = \mathbf{\nabla} \cdot \mathbf{J}(\mathbf{r}, t)\) with the one-body probability current

\[
\mathbf{J} = \int d\mathbf{r}'' \mathcal{D}^{p}(\mathbf{r} - \mathbf{r}'') \cdot \left( \nabla \rho^{(2)}(\mathbf{r}, \mathbf{r}'', t) + \rho^{(2)}(\mathbf{r}, \mathbf{r}'', t) \nabla \left( \beta \nu^{\text{eff}}(\mathbf{r}) + \beta u^{\text{eff}}(\mathbf{r}, \mathbf{r}'') \right) \right)
\]

\[
+ \int d\mathbf{r}' \rho^{(3)}(\mathbf{r}, \mathbf{r}', \mathbf{r}'', t) \nabla \beta u^{\text{eff}}(\mathbf{r}, \mathbf{r}')
\]

(B.3)

Now we assume that the system evolves in time in such a way that the time-dependent correlations instantaneously follow those of an equilibrium system. This adiabatic assumption [35–37] enables us to employ exact equilibrium sum rules to rewrite the expression in brackets in terms of two-body densities [33, 37]. The resulting probability current reads

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\[ J = \mathcal{D}(r) \cdot \left( \nabla \rho(r, t) + \rho(r, t) \nabla \beta u^{\text{eff}}(r) + \int dr' \rho^{(2)}(r, r', t) \nabla \beta u^{\text{eff}}(r, r') \right), \]  

where we have defined the ensemble-averaged diffusion tensor

\[ \mathcal{D}(r) = \int dr'' \frac{D^p(r - r'') \rho^{(2)}(r, r'', t)}{\rho(r, t)}, \]

which we choose to be given by equation (15) of the main text, following the result of a calculation in [33] similar to the one presented here.

As a final step, we recognize that the term in brackets in equation (B.4) can be written as \( \rho(r, t) \delta \beta \mathcal{F}^{\text{eff}}[\rho] / \delta \rho(r, t) \) in terms of the functional derivative of a free energy functional. The exact excess free energy (corresponding to the term involving \( u^{\text{eff}} \)) is only known implicitly and we choose here an approximate representation with \( \mathcal{F}^{\text{eff}}[\rho] \) from equation (A.5). With this identification, the probability current in equation (B.4) directly leads to equation (14) and equation (16) of the main text.

**Appendix C. Effective route to the active pressure and interfacial tension**

Consider a fluid interacting with the pair potential \( u(r) \) confined by an external field \( \nu(r) \) and define according to equation (16) the average net force as

\[ X(r) := \mathcal{D}(r) \cdot \left( \nabla \rho(r) + \rho(r) \nabla \beta u^{\text{eff}}(r) \right) \]

and using the effective potentials defined in appendix A we obtain:

\[ X(r) = \mathcal{D}(r) \cdot \left( \nabla \rho(r) + \rho(r) \nabla \beta u^{\text{eff}}(r) + \int dr' \rho^{(2)}(r, r') \nabla \beta u^{\text{eff}}(|r - r'|) \right), \]  

where the diffusion tensor \( \mathcal{D}(r) \) is given by equation (15). Now, according to equation (16), \( X(r) = 0 \) is a consequence of the steady-state condition, equation (7), so that equation (C.1) constitutes an effectively mechanical force-balance condition. Our objective is to analyze the equality

\[ \frac{1}{\mathcal{V}} \int dr X(r) \cdot r = 0, \]  

which results from the virial (13) of forces \( X(r) \).

Recalling that, by the standard external virial, the mechanical pressure is defined as

\[ \beta p = \frac{1}{\mathcal{V}} \int dr \rho(r) \left( \nabla \beta \nu(r) \right) \cdot r \simeq \frac{1}{\mathcal{V}} \int dr \rho(r) r \cdot \mathcal{D}(r) \cdot \left( D_{[1]}^{-1} \cdot \nabla \beta \nu(r) \right). \]  

We make contact to equation (C.2) in the last step by inserting the first term of the effective external external potential \( \nu^{\text{eff}}(r) \) from equation (A.1). As discussed in the main text when deriving equation (18), this approximate conversion becomes exact when we consider a non-interacting fluid \( (u(r) = u^{\text{eff}}(r) = 0) \). In this case it is easy to show that equation (C.2) yields

\[ \mathcal{V} \beta p = - \int dr \left( \nabla \cdot \rho(r) \mathcal{D}(r) \right) \cdot r = - \int dr \nabla \cdot \left( \rho(r) \mathcal{D}(r) \cdot r \right) + \int dr \rho(r) \text{Tr}[\mathcal{D}(r)] \]

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after integration by parts, where the first term on the right-hand side is a boundary term and vanishes. In the bulk limit, where \( \nu(\mathbf{r}) \to 0 \) and \( \rho(\mathbf{r}) \to \rho \) is constant, we thus find the active pressure \( \beta P = D_a \rho \) of an ideal gas.

Consider now the analogy for the bulk limit of an interacting active fluid. First we recognize that the second step in equation (C.3) is not exact since \( \mathbf{D}(\mathbf{r}) \) depends on the pair potential. However, it can be easily shown that at linear order in \( \tau_a \), all additional terms depend on both the external field and the pair potential [33]. Secondly, all terms in equation (C.1) depending on the external field (including these mixed terms) will not contribute to the internal bulk stress after taking the limit \( \nu(\mathbf{r}) \to 0 \) and \( \rho(\mathbf{r}) \to \rho \) as for equation (C.3). This further means that there exists a wall-independent equation of state, which is not the case for generic active systems [22]. Thus we argue that, in a good approximation, the internal contributions to equation (C.2) can be calculated with an effective diffusion tensor \( \mathbf{D}(\mathbf{r}) \) whose components

\[
D_a D_{a\beta}^{-1}(\mathbf{r}) = \delta_{a\beta} + \tilde{r} \int d\mathbf{r}' \frac{r^{(2)}(\mathbf{r}, \mathbf{r}')}{\rho(\mathbf{r})} \partial_\alpha \partial_\beta u(\mathbf{r})
\]

only depend on derivatives \( \partial_\alpha = \partial/\partial \alpha \) of the interparticle potential \( u(\mathbf{r} = |\mathbf{r} - \mathbf{r}'|) \), where the indices \( \alpha \) and \( \beta \) run over \( x, y \) and \( z \). Considering a system that is homogeneous in the \( x-y \) plane, the integrand in equation (C.5) is antisymmetric with respect to \( \alpha \) if \( \alpha \neq \beta \) (and \( \alpha \neq z \)) and equally for \( \beta \), so that \( \mathbf{D}^{-1} \) becomes diagonal and easily invertible. The result for \( \mathbf{D}_{\alpha\alpha} \) is stated as equation (21) of the main text.

Without loss of generality we consider in the following the case of \( d = 3 \) spatial dimensions. Separating the result of the external virial from other contributions, we define for each term of the sum \( \sum \mathbf{X}_a(\mathbf{r}) \mathbf{r}_a \) over the vector components in equation (C.2) the diagonal elements \( P_{\alpha\alpha}(\mathbf{r}) = P_{\alpha\alpha}^{id}(\mathbf{r}) + P_{\alpha\alpha}^{vir}(\mathbf{r}) \) of the pressure tensor \( P(\mathbf{r}) \), so that \( 3V_P = \int d\mathbf{r} \text{Tr}[P(\mathbf{r})] \). The first term in equation (C.1) results in the ideal part

\[
\beta P^{id}_{\alpha\alpha}(\mathbf{r}) = \rho(\mathbf{r}) \mathbf{D}_{\alpha\alpha}(\mathbf{r}) + \rho(\mathbf{r}) \mathbf{r}_a \partial_a \mathbf{D}_{\alpha\alpha}(\mathbf{r})
\]

after performing an integration by parts of \( \int d\mathbf{r} \beta P^{id}_{\alpha\alpha}(\mathbf{r}) := -\int d\mathbf{r} \mathbf{r}_a \mathbf{D}_{\alpha\alpha}(\mathbf{r}) \partial_a \rho(\mathbf{r}) \) as in equation (C.4) and subsequently taking the bulk limit. From the third expression in equation (C.1) we identify

\[
P^{vir}_{\alpha\alpha}(\mathbf{r}) = -\mathbf{r}_a \mathbf{D}_{\alpha\alpha}(\mathbf{r}) \int d\mathbf{r}' \rho^{(2)}(\mathbf{r}, \mathbf{r}') \partial_\alpha u^{eff}(\mathbf{r}) = -\mathbf{r}_a \mathbf{D}_{\alpha\alpha}(\mathbf{r}) \int d\mathbf{r}' \rho^{(2)}(\mathbf{r}, \mathbf{r}') \frac{r^{(2)}(\mathbf{r}, \mathbf{r}')}{r} \partial_\alpha u^{eff}(\mathbf{r}),
\]

which is identical to the standard virial contribution for a passive system, but here involving the effective interactions (and correlations).

To understand the role of the (uncommon) second term in the ideal pressure tensor (C.6), we substitute the effective pair potential (A.2) into equation (C.7), while assuming \( \mathbf{D}_{11}(\mathbf{r}, \mathbf{r}') \to \mathbf{D}(\mathbf{r}) \), which is only true for single-particle interactions. The result is \( \beta P^{vir}_{\alpha\alpha}(\mathbf{r}) := \beta P^{vir,0}_{\alpha\alpha}(\mathbf{r}) - \rho(\mathbf{r}) \mathbf{r}_a \partial_a \mathbf{D}_{\alpha\alpha}(\mathbf{r}) \), where the first term

\[
P^{vir,0}_{\alpha\alpha}(\mathbf{r}) := -\mathbf{r}_a \int d\mathbf{r}' \rho^{(2)}(\mathbf{r}, \mathbf{r}') \partial_\alpha u(\mathbf{r})
\]

equals the standard virial formula for a passive system interacting with the bare potential \( u(\mathbf{r}) \) and the second term is the negative of the expression in \( P^{id}_{\alpha\alpha}(\mathbf{r}) \) from
Equation (C.6). Defining $P^0 := P_{\text{id}} + P_{\text{vir}} = P_{\text{id},0} + P_{\text{vir},0}$ with $\beta P_{\text{id},0}(r) := \rho(r)D_{\text{id},0}(r)$, we recover the YBG results [31] for pressure and interfacial tension stated in equations (23) and (25) of the main text, respectively. In practice, the contributions to $P$ arising from the effective pair potential are only partially compensated by the second term in equation (C.6), such that $P$ does not actually reduce to $P^0$. This cannot be explained alone with the approximations of the EPA (discussed in appendix B of [33]), as we also ignore higher-order correlations in defining $D(r)$ from a pairwise quantity. To understand this we notice that we could also recover equation (C.8) from equation (C.7) when including $D(r)$ to the integrand in the form $D_{11}(r, r')$. This idea is further elaborated in appendix D. On the other hand, the effective free energy derived from $u_{\text{eff}}(r)$ in appendix A induces higher-order correlations, which are required to accurately describe the effective structure. Hence, the deviation in $P - P^0$ can be interpreted as both an undesired artifact of the EPA and an approximate compensation accounting for higher-order terms neglected in $P_{\text{id},0}$ as we require an approximate pairwise additive diffusion tensor to define $D(r)$.

For a homogeneous system, all components $D_{\alpha\alpha}$ are equal and independent of $r$. We thus recover equation (22) of the main text by calculating

$$\beta p = \frac{1}{3} \text{Tr}[P] = \frac{1}{3} \text{Tr}[D] (\rho + \beta p_{\text{vir}}) \simeq \frac{1}{3} \text{Tr}[D] \beta p_{\text{eff}}$$

(C.9)

and identifying the effective virial pressure $p_{\text{vir}} \simeq p_{\text{eff}}$ with the excess pressure $p_{\text{ex}}$ from DFT, which is equivalent to the compressibility result. In practice, this final step is subject to a slight thermodynamic inconsistency, which also occurs, e.g. between the solutions of the Percus-Yevick integral equation [40].

Proceeding analogously for the interfacial tension of the planar interface, the diagonal components $D_{\alpha\alpha}(z)$ in equation (21) will depend on the normal coordinate $z$ and thus we have $D_{11}(z) = D_{22}(z) \neq D_{33}(z)$, which allows us to differentiate between the tangential $P_T(z) := P_{xx}(z) = P_{yy}(z)$ and normal $P_N(z) := P_{zz}(z)$ components of the pressure tensor. The interfacial tension thus becomes

$$\beta \gamma = \frac{1}{A} \int dr \left[ P_N(z) - P_T(z) \right] = \int dz \left[ D_{zz}(z) \beta p_{zz}^{\text{eff}} + \rho(z) z \partial_z D_{zz}(z) - D_{xx}(z) \beta p_{xx}^{\text{eff}}(z) \right] \simeq \int dz \left[ D_{zz}(z) \beta p_{zz}^{\text{eff}} + D_{xx}(z) \omega(z) + \rho(z) z \partial_z D_{zz}(z) \right]$$

(C.10)

where, according to the definitions in a passive system, we identify from equation (C.7) and the first term in equation (C.6) the components $p_{\text{eff}}^{\text{vir}} = \rho(z) + P_{\text{vir},0}(z)/D_{\alpha\alpha}(z)$ of the effective pressure tensor. Note that in a passive system, only the virial stress (C.8) is relevant to calculate the interfacial tension $\beta \gamma_0 = \int dz \left( P_{N_{\text{vir},0}}^{\text{vir}}(z) - P_{T_{\text{vir},0}}^{\text{vir}}(z) \right)$. Quite in contrast, the linear term in $\rho(z)$ also contributes to the interfacial tension at non-zero activity, as $(D_{zz}(z) - D_{xx}(z))$ does not vanish. Translating equation (C.10) to DFT language in the last step, similar to equation (C.9), results in equation (24) of the main text. Here the (constant) effective normal pressure $p_{zz}^{\text{eff}} \simeq p_{\text{eff}}^{\text{vir}}$ corresponds to the bulk pressure and the effective tangential pressure $p_{xx}^{\text{eff}} \simeq -\omega(z)$ to the negative density of the grand potential along the interface. Note that $\omega(z)$ explicitly contains the ideal contribution $\rho(z)$, such as all components of the pressure tensor, whereas the DFT.
result for the bulk pressure corresponds to the value of the functional for either constant density at coexistence and thus cannot be related to the inhomogeneous density profile along the interface. Therefore, an explicit separation of the interfacial tension into ideal and excess contributions as in equation (C.9) is not possible.

Appendix D. Alternative rescaling of effective mechanical quantities

In order to derive equation (14) of the main text, we assumed in appendix B the effective diffusion tensor to be pairwise additive. We then argued that, regarding the equivalence of equations (6) and (7), it is convenient to define the averaged diffusion tensor $\mathcal{D}(r)$ as the inverse of the average of its inverse. As a consequence, the formulas we find in appendix C have the same structure as those derived in [31] from an expansion of the diffusion tensor $\mathcal{D}_{[N]}(r^N)$ defined in equation (5). Since we further employ the EPA, the present approach lacks of self-consistency.

In order to achieve a higher level of self-consistency, we reevaluate the argumentation line from appendix C considering the effective virial pressure tensor

$$\beta P^{\text{vir}}(r) = -r \cdot \mathcal{D}(r) \cdot \int dr' \rho^{(2)}(r, r') \mathcal{D}^{-1}_{[1]}(r, r') \cdot \nabla \cdot \left( \beta u(r) + \mathcal{D}_{[1]}(r, r') \right),$$

(D.1)

after substituting $u^{\text{eff}}(r)$ into equation (C.7), where

$$\mathcal{D}_{[1]}(r, r') = D_a (1 + \hat{\tau} \nabla \nabla u(r))^{-1}. \quad (D.2)$$

As a limitation of the EPA [33], we recognize artificial three-body correlations in equation (D.1). At leading order in the activity parameter $\hat{\tau}$, however, the term involving $\nabla \cdot \mathcal{D}_{[1]} \propto \hat{\tau}$ simplifies dramatically. This calculation suggests a redefinition of the correction term in the ideal pressure tensor from equation (C.6) as

$$\beta P^{\text{id}}_{\alpha\alpha}(r) = \rho(r) \mathcal{D}_{\alpha\alpha}(r) + r_a \int dr' \rho^{(2)}(r, r') \partial_{\alpha} \mathcal{D}_{[1]}(r, r'). \quad (D.3)$$

By doing so, we ensure that at low activity only the first terms in equations (D.1) and (D.3) contribute to the pressure tensor. In contrast to $P^{\text{id}}$, this new expression does not vanish in the bulk.

The spurious fact that the two integrations in equation (D.1) are carried out independently (mind that $\mathcal{D}(r)$ is an averaged quantity) stems from making the EPA [33]. Actually, the integrand of $\mathcal{D}(r)$ should cancel with $\mathcal{D}^{-1}_{[1]}$, such that the term involving $\nabla \cdot 1 \beta u(r)$ reduces to equation (C.8), which does not depend on the activity at all. This means that within the EPA we cannot recover the passive virial contribution to the pressure tensor even at leading order in $\hat{\tau}$. However, this consideration suggests that choosing the effective diffusion tensor $\mathcal{D}(r) \approx \mathcal{D}^{[2]}(r)$ with

$$\mathcal{D}^{[2]}(r) = \int dr' \rho^{(2)}(r, r') \mathcal{D}_{[1]}(r, r') \rho(r) \quad (D.4)$$

is more consistent with the EPA than the expression in equation (C.5). In a manner of speaking we thereby replace the approximate many-body average in equation (C.5) with an exact two-body average, equation (D.4).
Again, it is easy to verify that only the diagonal components
\[ (\mathcal{D}_{[11]})_{\alpha\alpha}(r, r') = D_{\alpha} \frac{1 + \tilde{\tau} \partial_{\alpha}^{2} u(r) - \tilde{\tau} \left( \frac{\partial_{\beta} u(r) - \delta_{\alpha\beta}}{r} \right) (x_{\alpha} - x_{\beta})^{2}}{(1 + \tilde{\tau} \partial_{\alpha}^{2} u(r)) \left( 1 + \tilde{\tau} \frac{\partial_{\alpha} u(r)}{r} \right)}. \] (D.5)
of \( \mathcal{D}^{[2]}(r) \) are important when the system is at most inhomogeneous in the \( z \) direction. In practice, however, the tensor \( \mathcal{D}_{[11]} \) may be ill-defined if the validity criteria of the underlying theory are violated [33]. Therefore, a similar correction as for defining the effective potentials would have to be employed in equation (D.5).

**Appendix E. Mean-field free energy in the mechanical picture**

In this appendix we derive a steady-state condition of the form (16) which gives rise to a different (mean-field) free energy functional. Our starting point is the YBG-like expression in equation (20) of the main text, which we write here as
\[ 0 = \rho(r) \langle \nabla \beta U \rangle + \nabla \cdot (\mathcal{D}(r) \rho(r)), \] (E.1)
assuming \( \mathcal{D} \simeq \mathcal{D}_{1} \) as in (15). This is exact at linear order in \( \tilde{\tau} \) [33], i.e. when we define \( \mathcal{D} \) from equation (19). Multiplication of equation (20) with \( I \equiv \mathcal{D} \cdot \mathcal{D}^{-1} \) does not alter its mechanical character, whereas we may rewrite it in the form
\[ 0 = \mathcal{D}(r) \cdot \rho(r) \left( \nabla \ln \rho(r) + \mathcal{D}^{-1}(r) \cdot \langle \nabla \beta U \rangle - \nabla \ln \det(\mathcal{D}^{-1}(r)) \right) \]
\[ \approx \mathcal{D}(r) \cdot \rho(r) \nabla \left( \frac{\delta \beta \mathcal{F}_{\text{id}}(\rho)}{\delta \rho(r)} + \frac{\delta \beta \mathcal{F}_{\text{ex}}^{(1)}(\rho)}{\delta \rho(r)} + \frac{\delta \beta \mathcal{F}_{\text{ex}}^{(2)}(\rho)}{\delta \rho(r)} \right) \] (E.2)
resembling equation (16) of the main text. In the last step we identify a free energy functional \( \mathcal{F} = \mathcal{F}_{\text{id}} + \mathcal{F}_{\text{ex}}^{(1)} + \mathcal{F}_{\text{ex}}^{(2)} \), such that the left-hand-side arises from its functional derivative.

The first term corresponds to the standard ideal-gas free energy \( \mathcal{F}_{\text{id}}(\rho) = \int dr \rho(r) \left( \ln(\Lambda^{3} \rho(r)) - 1 \right) \). The second and third terms contain the inverse of the ensemble-averaged diffusion tensor \( \mathcal{D}^{-1} \), which for an interacting system depends on the pair density. To obtain the free energy as a functional of the one-body density alone, we employ the simplest mean-field approximation \( \rho^{(2)}(r, r') \approx \rho(r) \rho(r') \) in (15) and identify
\[ \frac{\delta \beta \mathcal{F}_{\text{ex}}^{(1)}(\rho)}{\delta \rho(r)} = \frac{\beta}{D_{\alpha}} \left( \tilde{U}(r) + \frac{\tilde{\tau}}{2} \left( \nabla \tilde{U}(r) \right)^{2} \right), \] (E.3)
\[ \frac{\delta \beta \mathcal{F}_{\text{ex}}^{(2)}(\rho)}{\delta \rho(r)} = - \ln \det \left( 1 + \tilde{\tau} \nabla \nabla \beta \tilde{U}(r) \right), \] (E.4)
where we have defined \( \tilde{U}(r) = \nu(r) + \int dr' \rho(r') u(r, r') \). By construction of the free energy \( \mathcal{F} \) we recover the form of the mechanical condition in equation (E.1), but at the cost of neglecting all correlations. For a non-interacting system a functional integration of equations (E.3) and (E.4) becomes possible and \( \mathcal{F} \equiv \mathcal{F} \) becomes exact and thus equivalent to the free energy derived in appendix A within the EPA.

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There is no analytic expression for \( \tilde{F} \) in the general interacting case, which means that the EPA is more useful. Moreover, with the notion of an effective potential, it is easy to construct a closed theory beyond the simplest mean-field level, e.g. by treating the repulsive part employing an elaborate functional for hard spheres [11, 41]. In contrast, the structure predicted from the free energy \( \tilde{F} \) is highly inconsistent with the mean-field assumption \( g(r) \equiv 1 \). However, in the presence of both an external field and interparticle interactions, equations (E.3) and (E.4) account for the a coupling between the corresponding potentials, which is ignored in the EPA. Connecting to the discussion in appendix D, we recognize that at linear order in \( \tilde{r} \) the expression in equation (E.4) is also found in the EPA when we additionally make the same mean-field approximation.

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