I. INTRODUCTION

Dry sand trickles easily through chinks and crevices, as everyone knows well from the hour glass, or just personal experience. However, the addition of small amounts of liquid are sufficient to transform it into a plastic (or, more precisely, a viscoplastic) material. The same is true for all granular matter when a few volume percent of liquid are added, provided the latter wets the grains well and the grains are not too large. It is understood that this dramatic change, from a quasi-fluid to a solid behavior, is due to the formation of liquid bridges \[1\] and extends the effect to higher dimensional systems. Since the number of exact contacts is equal to \(2D\), in agreement with established simulations. When the particles are wetted, they interact by the formation and rupture of liquid capillary bridges. The same is true for all granular matter when a few volume percent of liquid are added, provided the latter wets the grains well and the grains are not too large. It is understood that this dramatic change, from a quasi-fluid to a solid behavior, is due to the formation of liquid bridges \[1\] and extends the effect to higher dimensional systems. Since the number of exact contacts is equal to \(2D\), in agreement with established simulations. When the particles are wetted, they interact by the formation and rupture of liquid capillary bridges.

A dry system of \(N\) hard spheres with diameter \(d\) confined to an area or volume \(V\) has no intrinsic energy scale, so that the equation of state is of the form \(P = T f(N/V)\) with the temperature \(T = \langle mv_i v_i\rangle\) and a nonlinear density dependence, \(f\). The defined size of hard particles is conveniently used to restate the density \(n = N/V\) as the

\[\phi\]
dimensionless occupied fraction \( \phi = \sigma_D n D^D / (2^D D) \) \( (\sigma_D \) the surface of a \( D \)-dimensional unit sphere), which is the area fraction \( \phi = 2^D n D^2 \) for two, and volume fraction \( \phi = 2^3 n D^3 \) for three dimensions.

The capillary interaction of wet granular matter has a well-defined binding energy \( E_{cb} \) \[10\], and it has been demonstrated experimentally \[14\] under realistic dynamical conditions with impact velocities typical for strongly fluidized wet granular matter that the hysteretic character of the interaction is essential: the dominant mechanism of dissipation is the hysteretic formation and rupture of capillary bridges, the energy \( E_{cb} \) of which is irreversibly taken from the kinetic energy of the granular motion whenever a liquid bridge ruptures \[11\]. The bridge energy has been quantified \[11, 14\] according to which \( E_{cb} \) is proportional \( d^\alpha \mathcal{V} \). Figure 1 illustrates this hysteresis in the Minimal Capillary model \[11\] applied here, which assumes a constant bridge force \( F_{cb} \). This may appear as an oversimplification at first glance, but there is increasing experimental evidence that the details of the force law are insignificant for the collective dynamics on which we focus here \[11, 14, 15\], as also confirmed from the point of view of dynamical systems theory \[10, 12\].

Obviously, an external energy current has to be continuously injected to drive the system into a nonequilibrium steady state. In the equilibrium limit, \( E_{cb} \to 0 \), we will have a pressure of the form \( P = T f(\phi) \). It is the objective of this article to derive the equation of state for the hysteretic liquid bridge interaction of wet granular matter in such a driven state. In view of the intrinsic energy scale \( E_{cb} \), this relation has to be of the form \( P = P(\phi, T/E_{cb}) \).

The equation of state is understood as an intrinsic property of homogeneous wet granular matter, kept in a stationary nonequilibrium state of granular temperature \( T \). With this given temperature we may subsume various ways in which the system can be externally driven to compensate for the dissipation by rupturing liquid bridges, so that this granular temperature \( T \) is maintained over many particle diameters.

We remark that in most experimental situations involving wet granular matter, the granular temperature is a nonlinear, even discontinuous, response depending on the details of the driving, such as boundary motion or air flow in air-fluidized beds. In this article we deliberately regard the granular temperature as the control parameter, so that the theoretical description of the boundary coupling is conveniently separated. Yet we emphasize that for the full description of an experimental situation one has to insert the equation of state into the equation for the external energy input, and then solve for the granular temperature as the nonlinear response to the external driving.

We aim at describing the steady nonequilibrium states of wet granular matter, which are so multifaceted that at first glance one might think that aside from density and granular temperature further physical parameters are necessary in order to describe such a state. Yet as simulations have shown, states of wet granular matter far from equilibrium \[1\] are very well described by a single granular temperature \( T \) assuming a Gaussian velocity distribution, neglecting higher cumulants. Furthermore, it is known that the self-organized velocity distribution of free cooling wet granular matter has a vanishing fourth cumulant \[18\]. We point out that the condition of a locally isotropic and homogenous
state used in this work implies that the temperature field may vary only slowly over many particle diameters so that there is no strong influence by a heat current, which would otherwise be considered as a third parameter of the local nonequilibrium state.

Throughout this study, we allow for a certain polydispersity, $0 \leq \Delta d / d < 0.1$. (For higher polydispersity, the dense system undergoes a kinetic glass transition [19, 20]. First of all, polydispersity is frequently used in simulations and experiments to prevent the monocrystalline state. Secondly, most systems of practical relevance exhibit some polydispersity. Another characteristic of ‘real’ granulates is that the surfaces of the grains are not ideal, bearing certain roughness. This does, however, only change the amount of liquid which must be added in order to achieve the capillary interaction: first some liquid is required to fill the crevices and tiny recesses in the grain surfaces, until the grains effectively have a smooth liquid coating, which is then completely wetted by all additional liquid. For glass beads, as those used in most of the experiments, this is typically the case above a volume fraction $W_{\text{min}} = 0.1\%$ of liquid with respect to the jammed granular sample volume. We also require an upper limit on the volume fraction of the wetting liquid, so that the maximal length $s_{\text{crit}}$ of liquid bridges is of the order or below the polydispersity $\Delta d$ of the spheres. This is to demand that $s_{\text{crit}} / d \approx \sqrt{W} / 3$ is smaller than $\Delta d / d < 0.1$, so that $W < W_{\text{max}} = 2.8\%$. This happens to closely coincide with the upper limit set on the liquid content to ensure that neighboring capillary bridge do not merge [21]. For this range of the liquid content the capillary interaction is a truly pairwise interaction with the capillary force acting radially between pairs of particles. Another implication of roughness is that there is a substantial tangential friction between adjacent grains. This means that in principle one has to include all rotational degrees of freedom in the kinetic considerations for any statistical physical treatment of our system. However, we are here focussing on the effects due to the liquid capillary bridges, which mediate central forces. These do not couple to the (tangential) rotational modes. We therefore expect that the rotational degrees of freedom play, in our system, the role of a spectator heat bath which follows the translational dynamics, but does not influence it greatly, aside from a quantitative increase of the granular specific heat. In fact, recent experiments and simulations of wet granular systems [15] show that this approach yields remarkable agreement with experimental data. In this work, we thus completely neglect all rotational degrees of freedom.

II. DRY SPHERES AS STARTING POINT

Before we add the wetting liquid to the hard sphere system, we investigate the dry case in this section and derive expressions for the pair correlation near contact, which will be extended to the wet case in the following section.

Due to their finite size, the positions of hard spheres are not distributed independently from each other, as it is the case for the point-particles of the ideal gas. The configuration space of $N$ spheres is not $V^N$, but restricted to a concave subset in which the systems moves chaotically as a high dimensional balliard. With the absence of an intrinsic energy scale, the dry system is athermal, which means that a change in temperature is equivalent to rescaling the time axis. The excluded volume gives rise to correlations in the particle positions, which are measured by the pair correlation function. Denoting by $n = N / V$ the mean macroscopic particle density and by $n_m(r) = \sum_n \delta(r - r_i)$ the microscopic density, the isotropic pair correlation $G(r)$ is defined as the probability

$$
\langle n_m(r) \rangle_{\text{particle at 0}} \ d \text{vol} = n \ G(|r|) \ d \text{vol} = n \ g(s) \ d \text{vol}
$$

(1)

to find the center of a particle in the shell $d \text{vol} = \sigma D \ r^{(D-1)} \ dr$ of radius $r = r_i + r_j + s$ and thickness $dr = ds$ centered around a reference particle. We have conveniently subtracted the particle radii $r_i, r_j$ in the last equality of (1), so that $s > 0$ is the surface separation. The function $g(s)$ is advantageous for polydispersity scattered around the mean diameter $d$, because of its defined contact point, $s = 0$, which is smeared out in the function $G(r)$. Furthermore it is the natural way to describe an interstitial liquid bridge between the considered pair of particles, with $s$ the length of the bridge. For a certain liquid volume per particle and contact angle of the wetting liquid, there is a well defined critical bridge length $s_{\text{crit}}$ at which the bridge becomes unstable and ruptures. The mean density $n$ is factored out in (1) so that the dimensionless $g$ would be equal to unity for all separations if there was no particle-particle correlation. Figure 2 shows the pair correlation of a fluidized state in which long range order is lost, so that $g(s)$, respectively $G(r)$, tends to unity for $r \gg d$.

The forces in wet granular matter, hard-core repulsion and liquid bridge attraction, are short-ranged and radial, acting between pairs of particles over a separation range $0 < s < s_{\text{crit}}$ with $s_{\text{crit}} \ll d$. We are therefore interested in the short-range behavior of the pair correlation $g(s)$ up to leading order in $s / d$. For such short particle separations the pair correlation $g(s)$ is (up to a normalization constant) just the probability to find next neighbors at a separation $s$. Put in equivalent words: decomposing the pair correlation function $g(s) = \sum \ g_k(s)$ in contributions $g_k$ of the $k$’s shell of Voronoi neighbors, we have $g(s) = g_1(s)$ in the range of interest, $0 < s < s_{\text{crit}} \ll d$. To shorten notation we suppress the subindex 1.
FIG. 2: The pair correlation of wet granular matter in a fluidized state resulting from a molecular dynamics-type simulation in $D = 2$ dimensions. The correlation function $G(r)$ vanishes in the range $(0, d)$ where the finite particle size leads to excluded volume. We use the function $g(s)$ with the surface separation $s$ of neighboring particles as it is convenient for wet granular matter where interstitial liquid bridges have the length $s$. Note that this is not exactly identical to the function $G(d+s)$ shifted by one particle diameter $d$, since a realistic granular system has some polydispersity $\Delta d$ around the mean diameter $d$. Aside from kinetic contributions, the pressure is due to the interaction forces which become dominant with increasing density. The internal forces in wet granular matter are short-ranged. Therefore our interest focuses on the sharp fall-off in the indicated range $0 < s < s_{\text{crit}}$ of capillary interaction. This highlighted region indicates the typical range of $s_{\text{crit}}$, and corresponds to the region highlighted in Fig. 4. Furthermore, we derive more detailed correlation functions, $g^u(s)$ and $g^b(s)$, for unbound and capillary connected pairs, respectively, in order to describe the hysteretic interaction in wet granular matter.

A. The Dense Limit

Figure 3 gives an overview of results by [20], [22] (Fig. 15 therein), and [23] for the phases of the two-dimensional system depending on density and polydispersity. For polydispersity below 0.1, there are two density regimes separated by the ordering transition at $\phi_o$ [64]. These transitions are a purely geometric property (i.e. excluded volume effect) of the configuration space and are therefore athermal. To compute the radial next-neighbor distribution at densities above the critical density, $\phi > \phi_o$, we consider the Voronoi tessellation of the system, which embeds each particle into a convex polygonal cell. The sizes $\{V_i\}$ of these Voronoi cells scales as $(d+s)^D$, with the particle separation $s$. The mean cell size $\sum_i V_i/N = V/N = 1/n$ is exactly the inverse density $n$. Hence,

$$\left\langle \left(1 + \frac{s}{d}\right)^D \right\rangle = \frac{n(s \to 0)}{n} = \frac{\phi(s \to 0)}{\phi},$$

where the triangle brackets denote averaging over next neighbors which are in contact $(s \to 0)$ with the center particle at the jamming density $\phi(s \to 0)$. We refer to those pairs of particles which come into contact at jamming as neighbors of type A, i.e the surface separation $s_A$ of A-neighbors is

$$s_A = 0 \text{ at } \phi = \phi_J.$$

In the monodisperse limit for $D = 2$, $\phi_J$ assumes the value of the triangular crystal, $\phi_{\text{max}} = \pi/(2\sqrt{3}) = 0.91$. Polydispersity decreases the (maximal random) jamming density $\phi_J$ and increases the critical density $\phi_o$ for the onset of triangular order as shown in Fig. 3.

1. Contribution to the Contact Correlation: The A-Neighbors

Since the Voronoi cells exchange their free volume, $V - V_{\text{min}} \propto \left(1 + \frac{s}{d}\right)^D - 1$, and the total volume is conserved we assume an exponential distribution of the free volume, which is well confirmed by experiments with dry granulates.
FIG. 3: This plot reviews [20], [22] (Fig. 15 therein), and [23]. The athermal transitions shown are properties of the configuration space of hard discs. Since the wet granular dynamics takes place in this configuration space, results for hard discs form the starting point for a theory of wet granular matter in two dimensions. At low (gas) and moderate (fluid) densities $\phi$, the configuration space is probed ergodically and the system has low shear viscosity. As density is increased, the system gets trapped in a disordered state (glass for polydispersity above 0.1), or in a state with local triangular order. Both transitions, the glass transition (vertical line at $\phi \approx 0.80$) and the ordering transition (curved line ending at $\phi_c = 0.71$) can be detected by the rapid increase of the shear viscosity $\eta$ (cf. [24] for the ordering transition). While there is an athermal first order transition in three dimensions, it is at present discussed in the literature whether the transition region (double lines ending at $\phi_o$) represents a fluid/solid coexistence (corresponding to a weak first order transition with a small jump of the entropy per particle) or if there is an intermediate hexatic phase (according to the Kosterlitz-Thouless-Halperin-Nelson-Young scenario) [12, 13]. As the packing fraction $\phi$ is increased further, the islands to which the system is confined in the configuration space shrink to points. This jamming limit can be detected by the divergence of the pressure $p$ (at fixed granular temperature) under compression (for example using the particle expansion of the Lubachevsky-Stillinger algorithm). The maximal random jammed state (MRJ, for strict jamming as defined by Torquato, Truskett, and Debenedetti [25]) is the vertical curve at the right. Densities higher than MRJ are deep in the glassy regime. From the thermodynamic point of view, the system would cluster in phases separated according to the particle size, but this eutectic freezing-transition is kinetically suppressed [22] and unreachable.

The conditions (2) and (3) determine the A-neighbor distribution uniquely:

$$P_A(s) \, d\text{vol}(s) = \frac{D/(\sigma_D d^D)}{\phi_J/\phi - 1} \exp \left( - \frac{(1 + \frac{s}{d})^D - 1}{\phi_J/\phi - 1} \right) \, d\text{vol}(s).$$

The volume element for $D = 2$ is

$$d\text{vol}(s) = \sigma_D \, r^{(D-1)} \, dr = \pi d \, (1 + s/d) \, ds.$$  \hspace{1cm} (5)

The contribution $g_A$ which A-neighbors give to the pair correlation is equal to the A-neighbor distribution $P_A$ up to a prefactor, so that

$$g_A(s) = g_c^{at} \exp \left( - \frac{(1 + \frac{s}{d})^D - 1}{\phi_J/\phi - 1} \right)$$

is determined as soon as we know the athermal contact value, $g_c^{at} = g_A(0)$. This contact value follows from the classical free volume theory [27] (which was based on [28]),

$$P = nT = \frac{D}{\phi_J/\phi - 1} + O(1),$$

in conjunction with the general relation between the particle-wall correlation $g_{wall}^{at}$ and the pair correlation $g_c^{at}$,

$$P = g_{wall}^{at} = 1 + 2^{D-1} \phi \, g_c^{at}.$$  \hspace{1cm} (8)
As a consequence, we obtain
\[
\frac{2^{D-1}}{D} \phi g_{c}^{at} = \frac{1}{\phi / \phi_{1} - 1}
\] (9)

close to jamming. Expression (9) is exact for \( D = 1 \), and has been confirmed as the asymptotic behavior of the diverging pressure close to jamming for \( D = 2 \) \[24, 29\] in event-driven simulation with accuracy \( 10^{-4} \). We remark that this expression is not limited to weak polydispersity and has been confirmed for polydispersity far above 0.1 in the glass state \[30, 31\].

Inserting (9) in (6), we have as our first central result a closed expression for the near-contact pair correlation of neighbors which form exact contacts in the jamming limit (so-called A-neighbors):
\[
g_{A}(s) = g_{c}^{at} \exp \left( -\frac{2^{D-1}}{D} \phi g_{c}^{at} \left[ \left( 1 + \frac{s}{d} \right)^{D} - 1 \right]\right). \tag{10}
\]

Eq. (10) implies for the derivative at contact,
\[
dg_{A}(0) = -2^{D-1} \phi g_{c}^{2}(0), \tag{11}
\]
a quadratic dependence on the contact value \( g_{c}^{at} \). Eq. (11) can be viewed as a consequence of normalization: the height of the contact peak is \( g_{c}^{at} \) and so the width is of the order \( 1/g_{c}^{at} \), which means that the negative slope is of the order \( g_{c}^{2} \). In fact, writing the A-neighbor correlation function \( g_{A}(s) \) in terms of the contact value \( g_{c}^{at} \), as we did in (10), is the natural form to express the density dependence of \( g_{A} \) because this manifests that the coordination number of A-neighbors is density independent:
\[
K_{A} = n \int g_{A} \, d\text{vol} = \frac{2^{D}}{d} \phi g_{c}^{at} \int_{s=0}^{\infty} \exp \left( -\frac{2^{D-1}}{D} \phi g_{c}^{at} \left[ \left( 1 + \frac{s}{d} \right)^{D} - 1 \right]\right) \left( 1 + \frac{s}{d} \right)^{D-1} \, ds
\] (12)

More significantly, \( K_{A} \) equals exactly the isostatic contact value \( 2D \), which is obviously correct for particles on a line \( (D = 1) \) and is the accepted value for ideal discs and spheres in \( D = 2 \) and \( D = 3 \) dimensions respectively \[22, 32, 33\]. The finding (12) is an essential confirmation of consistency of our approach, since it is independent from conventional arguments based on the rank of the rigidity matrix (which accounts for global constraints on the degrees of freedom) \[32\].

As the contact value \( g_{c}^{at} \) grows to infinity in the jamming limit, \( \phi \to \phi_{1} \), the constant integral (12) implies that \( n g_{A}(s) \) becomes a delta distribution with \'weight\' \( 2D \) at contact, \( s = 0 \).

2. The Background Contribution: The B-Neighbors

The configuration space is spanned by all particle positions \( \{ \mathbf{r}_{i} \} \). Consequently, a jammed configuration is – aside from a small fraction of rattlers \[65\] – an isolated configuration point, and the set of jammed configuration is a set of discrete points. When the density is slightly relaxed, a finite system remains confined to a finite environment around the jamming point (cf. \[34\], p. 35). As density is lowered further, these environments are no longer isolated so that the system is able to migrate between these \'islands of jamming\'.

The stability analysis of contact networks \[32, 33, 36, 37, 38\] has put forth the result that frictionless spheres (except for the singular limiting case of a monodisperse crystal) jam strictly in an isostatic packing with \( 2D \) contacts per particle on average, as confirmed numerically \[32\] for \( D = 3 \), be the state random (glass regime in Fig. 3) or locally ordered. Therefore we can identify within an island of jamming on average four neighboring particles in \( D = 2 \) dimensions which are close to the reference particle, and which will be in contact with the reference particle, \( s_{A} \to 0 \), in the jamming limit, \( \phi \to \phi_{1} \). These are the A-neighbors with the contribution \( g_{A} \) to the pair correlation derived in (10). Furthermore, it is a mathematical fact that any discrete set of points in flat two-dimensional space has on average six Delaunay/Voronoi neighbors \[39\], two of which have no contact to the reference particle, \( g_{B}(0) = 0 \). Hence, on the mean field level the following picture arises: Beside the four A-neighbors there are two B-neighbors which are sterically hindered by other particles from further approach to the reference particle. Summing up the contributions of A- and B-neighbors,
\[
g_{\text{dense}}(s) = g_{A}(s) + g_{B}(s), \tag{13}
\]
FIG. 4: The pair correlation near contact resulting from free-volume considerations. Close to jamming we distinguish between neighbors which form exact contacts in the jamming limit (contribution $g_A$, Eq. (10)) and those that are blocked at positive separation $s$ (curve $g_B$, Eq. (14)). The near-contact correlation is the sum of both contributions. For this plot the density is chosen to $\phi = 0.8$. The dashed curve sketches a typical second shell consisting of the second Voronoï neighbors. They are out of the interaction range, $0 < s < s_{\text{crit}}$, which is indicated by the highlighted stripe.

gives us the pair correlation function near contact. The pair correlation near contact which arises from these blocked states, $g_B$, is discussed in detail in appendix A. The essential result is that the configuration space of blocked states tends quadratically to zero in $s_B$, so that to leading order the normalization of two B-neighbors for $D = 2$ determines the B-contribution in (13):

$$g_B(s_B) = N \frac{P_B(s_B)}{\left(\frac{s_B}{d}\right)^2} e^{-\left((1+s_B)\right)}/c_B \left[1 + O\left(\frac{s_B}{d}\right)\right]$$

(14)

with $c_B = \phi_{\text{max}}/\phi - 1$. In Fig. 4 the resulting near-contact pair correlation for the dense regime is shown as the sum of $g_A$ and $g_B$.

B. The Dilute and Moderately Dense Regime

In this part we turn to the free rheological regime, $0 < \phi < \phi_c$. When two spheres are closer than one diameter, $s < d$, they shield each other from certain collisions events. If one was to neglect three-particle correlations, the isotropic bombardment by third particles gives rise to the well-known attractive depletion force first proposed by S. Asakura and F. Oosawa [40, 41]. As is evident from Fig. 5 summing up equal contributions over the accessible cross section is equivalent to the pressure exerted onto the submanifold indicated by the solid line in Fig. 5C and denoted by $\Sigma$.

This depletion force, as well as the liquid bridge force which we will take into account in the next section, will affect the pair correlation function. A systematic way to study this effect has been worked out by Hansen et al. [42], resulting in a Fokker-Planck equation for the two-particle distribution function. After integrating out the momenta and the center of mass coordinates, one finds that the depletion force as well as other non-entropic pair forces (such as the liquid bridge force), give rise to a Boltzmann factor,

$$g(s) \propto \exp\left(-\frac{V(s)}{T}\right)$$

(15)

For the depletion force

$$F_{\text{depl}} = -V'_{\text{depl}} = T \ n \ g_{e}^{\text{at}} \ \Sigma$$

(16)
FIG. 5: Origin of the depletion force attracting neighboring particles that are separated by less than a particle diameter. One may either think of this as an entropic force, due to the decrease of excluded volume when the shells of excluded volume overlap. Equivalently one may view this as the net force due to isotropic bombardment. Obviously, the integration over the solid arc in A is up to a sign equivalent to the integration in plot B. In B the integration is over the outer solid arc, which is the configuration space of the third particle’s coordinate at impact. Since the integration in B is projected by a cos-factor to give the axial symmetric force component, we can equivalently drop the cos-factor and integrate over the submanifold indicated by the solid line $\Sigma$ in C.

where

$$n \ g_c^{at} \ \Sigma \ ds = \frac{dV_{conf}}{V_{conf}} = -d \ln g(s)$$

is the infinitesimal logarithmic change of the excluded area (or the configuration space per particle, $V_{conf}$), when the particles are separated by $s < d$, and $\Sigma$ denotes the size of the corresponding section (line or area) in Fig. 5C. At contact, $s = 0$, the size of the integration section $\Sigma$ is

$$\Sigma = \frac{\sigma D-1}{D-1} \left(\frac{\sqrt{3}}{2}d\right)^{D-1},$$

which yields $V_{depl}$ to leading order in $s$. The depletion effect with the potential

$$\frac{V_{depl}}{T} = \frac{9}{2} \phi \ g_c^{at} \ s \left(1 - \frac{s}{3d} - \left(\frac{s}{3d}\right)^2\right)$$

for $D = 3$ has been confirmed in [43] by computer simulations. Polydispersity is known to have a minor effect on the depletion attraction [44]. For $D = 1$ (17) gives the Poisson distribution $V_{depl}/T = \phi \ g_c^{at} \ s/d$ which is exact only for $D = 1$.

In two dimensions, the depletion potential is

$$\frac{V_{depl}}{T} = \frac{2}{\pi} \phi \ g_c^{at} \ \left(4 \arctan \frac{\gamma_{vol}(s)}{W} + \gamma_{vol}W - C\right)$$

$$= 2g_c^{at} \phi \left(\frac{s}{\phi_{max} d}\right) + O\left(\frac{s}{d}\right)^2$$

for $D = 2$ with $\gamma_{vol}(s) = 1 + s/d$, the square root $W(s) = \sqrt{(1-s/d)(3+s/d)}$ and the constant $C = 2\pi/3 + \sqrt{3}$ to have $V_{depl} = 0$ at $s = 0$. The first line (20) is valid for $0 \leq s \leq d$, and the second line (21) suffices for the region of interest, $0 \leq s \leq s_{crit} \ll d$. For the application of results on the near-contact decay of the pair correlation function, such as (21), we prefer the exponential notation (used before in the dense case (10)) because it is most elegant to perform volume integration:

$$g^{dilute}_{AO}(s) = g_c^{at} \exp\left(-\frac{\phi}{\phi_{max}} \ g_c^{at} \left(\left(1 + \frac{s}{d}\right)^2 - 1\right)\right) \left[1 + O\left(\frac{s}{d}\right)^2\right]$$

for $D = 2$. In this notation the dilute and dense behavior of the pair correlation are conveniently compared, showing that the result (22) for the gaseous/fluid regime differs by the factor $1/\phi_{max} = 1.10$ in the exponent from the dense result (10) close to jamming, so that according to (22) the depletion force falls-off slower than the configuration density $\phi g_c^{at}$. We will now show that this is due to an over-estimation of the depletion force, caused by neglecting correlated three-particle events: when the plane of incidence of the third particle closely coincides with the symmetry plane $\Sigma$. 
the incoming particle will hit in short sequence the pair of particles considered, which increases very effectively the exchange of momentum, i.e. the depletion attraction is reduced.

To determine analytically and numerically the effect of correlated collisions which correct the Asakura-Oosawa result [22] we define the dimensionless measure

$$Z = \frac{4}{\pi n T d g_e^3} = -\frac{d g_e^3}{\phi g_e^2}, \quad (23)$$

for which the Asakura-Oosawa approach [10] and [18] gives $Z_{AO} = \frac{4}{\pi} \sqrt{3} \approx 2.205$ (Line A in Fig. 7). When we take correlated three-particle events into account, there are three contributions. Firstly, an attractive contribution $Z_1 > 0$ due to collisions on the front side of the pair, indicated by '1' in Fig. 6 which fall in the range $-\pi/2 < \phi < \pi/2$. The corresponding value $Z_1$ is easily integrated. Isotropy of the state demands that the angle $\phi$ between the symmetry axis of the pair $P''P$ and the incoming momentum $p_i$ is uniformly distributed, as well as the impact parameter $b$ (cf. Fig. 6). These collision parameters are related by $(\alpha, b) = (\varphi + \theta, d \sin \theta)$ to the position $\varphi$ on P and the angle of incidence $\theta$ with respect to the normal of P, which implies that $\varphi$ is uniformly distributed and $\theta$ is weighted by the cosine-factor $\cos \theta$. Integration over $-\pi/2 < \varphi < \pi/2$ yields the axial force contribution

$$F_1 = 2 T n g_e^3 d,$$  

so that $Z_1 = 8/\pi \approx 2.546$.

Secondly, the attraction is weakened by collisions hitting P in the remaining range $\pi/2 < |\varphi| < \varphi_{max}(s)$ (which we refer to as the 'broad side') giving rise to $Z_2 < 0$. At contact $\varphi_{max}(s = 0)$ is $2\pi/3$. For these collisions the incidence is shadowed by the partner particle $P''$ so that the angle of incidence $\theta$ is restricted to $-\pi/2 < \theta < \theta_{max}(\varphi)$. (Confer the collision event '2' in Fig. 6). Some trigonometry determines $\theta_{max}(s, \varphi)$ by the relation

$$(1 + \gamma_{vol}(s) \cos \varphi) \sin \theta_{max} = 1 - \gamma_{vol}(s) \cos \theta_{max} \sin \varphi,$$  

which allows for an explicit function of $\varphi$ at $s = 0$:

$$2 \cos \theta_{max}(\varphi) = \tan \frac{\varphi}{2} - \sqrt{1 + 2 \cos \varphi}. \quad (26)$$

After integrating over the impact momenta $p_i$ in the rest frame of P, the axial force imposed on P is

$$F_2(s) = \frac{4}{\pi} T n g_e^3 d \int_{\pi/2}^{\varphi_{max}(s)} d\varphi \cos \varphi \int_{-\pi/2}^{\theta_{max}(\varphi)} d\theta \cos^2 \theta,$$  

where the $\cos \varphi$ projects the force on the symmetry axis of the pair PP'. The $\cos \theta$ factor appears quadratically in the integrand [22] because of the cosine-distribution (or equivalent, because the Enskog collision frequency is proportional to the radial velocity $(p_i/m) \cos \theta$), and the transferred momentum which is $p_i \sin \theta$. Symmetry allows us to integrate over the upper half, $\pi/2 < \varphi < \varphi_{max}(s)$ in [27] and multiply by 2 with the general result

$$Z_2(s) = \frac{8}{\pi} \int_{\pi/2}^{\varphi_{max}(s)} d\varphi \cos \varphi \left[ \frac{1}{2} + \frac{\theta_{max}(s, \varphi)}{\pi} + \frac{\sin 2\theta_{max}(s, \varphi)}{2\pi} \right], \quad (28)$$

and the numerical value $Z_2(0) = -0.32813(9)$.

Thirdly, the most obvious and important correction on the three-particle level comes from double collisions denoted by 3 in Fig. 6. The third particle hits first $P'$ (gray arrow in Fig. 6) from the broad side at $\varphi' \in (\pi/2, \theta_{max})$. The radial component $p_i \cos \theta'$ of its incoming momentum $p_i$ is transferred to $P'$, which is why the third particle moves on tangentially to the circular cross section of $P'$ with momentum $p_i \sin \theta'$ to collide shortly afterwards with particle P. Here the momentum transferred is the radial component with respect to P, $p_i \sin \theta' \cos \theta$, so that

$$F_3(s) = \frac{4}{\pi} T n g_e^3 d \int_{\pi/2}^{\varphi_{max}(s)} d\varphi' \cos \varphi(\varphi') \cos \theta(s, \varphi') \int_0^{\theta_{max}(s, \varphi')} d\theta' \cos \theta' \sin \theta'. \quad (29)$$

The collision point on $P'$ described by $\varphi'(\varphi)$ is related to $\varphi$ (the subsequent collision point on P) by $\cos(\varphi' - \varphi) = 1 + \gamma_{vol}(s) \cos \varphi$. The incident angle $\theta$ on P is independent of $\theta'$ and given by $\sin \theta(s, \varphi') = 1 + \gamma_{vol}(s) \cos \varphi'$. After the elementary $\theta'$-integration we find

$$Z_3(s) = \frac{8}{\pi^2} \int_{\pi/2}^{\varphi_{max}(s)} d\varphi' \cos \varphi(\varphi') \cos \theta(s, \varphi') \sin^2 \theta_{max}(s, \varphi'),$$  

(30)
and $Z_3(0) = -0.091593(7)$. Summing up the three contributions gives $Z_{\text{corr}} = \sum_{i=1}^{3} Z_i \approx 2.127$ which is shown as the line B in Fig. 7.

Based on our numerical data shown in Fig. 7 we shall in the sequel assume the value

$$Z_{\text{sim}} = 2.$$ \hspace{1cm} (31)

By virtue of good statistics the simulation at $\phi = 0.097$ gave $Z_{\text{sim}} = 2.0009 \pm 0.0050$, and Fig. 7 suggest this result to hold with few percent limits very well over the entire density regime $0 < \phi < \phi_o$ considered in this subsection. The value $Z = 2$ determines the near-contact pair correlation uniquely to be

$$g_{\text{dilute}}(s) = g_{\text{dilute}}^{\text{at}} \exp \left( -\phi g_{\text{dilute}}^{\text{at}} \left[ \left( 1 + \frac{s}{d} \right)^2 - 1 \right] \right) \left[ 1 + O \left( \frac{\phi}{d} \right)^2 \right]$$ \hspace{1cm} (32)

for $D = 2$. Satisfactorily, this result (32) has exactly the same functional dependence on the configuration density $\phi$ $g_{\text{dilute}}^{\text{at}}$ as the formula put forward for the dense case (10) in the previous subsection. While three-particle collisions are obviously important since they shift $Z$ in the right direction, no analytic explanation for this coincidence corresponding to the value $Z = 2$ is provided at present. Yet we shall see in the next section that any value other than $Z = 2$ would lead to inconsistencies when we introduce the liquid bridge interaction.

We finally remark that the result (32) strongly differs from the ‘Poissonian fluid’ [45], for which the contact correlation $g_{\text{dilute}}^{\text{at}} - 1 > 0$ is ignored. Even at the lowest density ($\phi = 0.1$) considered in Fig. 7 the Poisson fluid would give $Z_{\text{Poisson}}(\phi = 0.1) = 1.7$ which is 15% below the simulation value, and the deviation from $Z = 2$ grows with density $\phi$.

III. THE PAIR CORRELATION UNDER THE HYSTERETIC INTERACTION

In this section we dress up the pair correlation function in order to describe the status of the liquid-bonds which are created and ruptured hysteretically in wet granular matter. We will proceed in two steps: first, we introduce in part III A the liquid bridges as hysteretic but forceless objects which follow the unperturbed particle dynamics. As a result, a direct relation of the dynamical system and the limiting case of isostatic granular packings [33, 38] at rest is found. In III B we turn on the liquid bridge force to its physical value, so that the bridges unfold their back-reaction on the granular dynamics. In the limit of low granular temperatures, $T \ll E_{\text{cb}}$, the particles stick together. For this frozen state of wet granular matter the bridge coordination $K$ is computed analytically as a function of the rupture length $s_{\text{crit}}$, and we find very good agreement with simulations.
FIG. 7: Functional test of the near-contact pair correlation. The vertical axis is proportional to the depletion force, \( F_{\text{depl}} \propto -d \ln g/\,ds \) at contact, divided by the configuration density \( \phi \, g^\infty(\phi) \). This fraction \( Z \) is predicted to be density independent by (32) and to assume the value \( Z = 2 \) (line C). Line A corresponds to the classical Asakura-Oosawa result, which is only valid for large beads immersed in a bath of small beads. In line B the corrections due to temporally correlated collisions events (derived in the text) have been taken into account. These events occur when a third particle of equal size strikes a pair of particles with a given separation \( s \ll d \) as sketched in Fig. 6. We proceed using the value \( Z = 2 \) (line C) because it agrees best with the simulation. Furthermore, \( Z = 2 \) corresponds to a near contact correlation function which is of exactly the same form as the function \( g_A \) we use in the dense case, when expressed in terms of the configuration density \( \phi \, g^\infty(\phi) \).

### A. The Hysteretic Coupling

Due to the hysteretic interaction, the pair correlation \( g \) is no longer a function of the particle separation \( s \). In order to include the knowledge about the collision history the configuration space has to be enlarged in two respects: Obviously we distinguish between pairs with and without liquid bridges, which we denote by superscript indices, \( g^b(s) \) and \( g^u(s) \) respectively, for ‘bridged’ and ‘unbridged’ neighbors (cf. Fig. 8). In addition, time reversal-symmetry is broken by the formation of the capillary bridge at contact. Hence we distinguish approaching pairs (with a negative relative velocity) which might collide and form a liquid bridge in the future, and those that move apart so that they can rupture the liquid-bond in the future. This relative velocity is denoted by a subscript arrow.

As we discuss the radial pair distribution, contact and rupture become the important points on the \( s \)-axis of the pair correlation function. At theses points the functions \( g^b \) and \( g^u \) are coupled according to the hysteretic transition of the bond status. We use an intuitive notation to refer to these points:

\[
\begin{align*}
\mathbf{g}^{\text{u}}_{c\rightarrow\odot} &= \{ \text{The probability for a pair at rupture distance approaching without bridge.} \\
& \text{at contact moving away with bridge.} \\
\mathbf{g}^{\text{b}}_{\odot\rightarrow r} &= \{ \text{The probability for a pair} \\
& \text{etc.}
\end{align*}
\]

The contact point \( s = 0 \) (or to be more precise: the right-sided limit \( s = 0^+ \)) is denoted by a circled \( \odot \), and the rupture at \( s = s_{\text{crit}} \) by the circled \( \odot \). Infinitesimally close to contact, there are four detailed correlation values: the bridge-connected and the unconnected states, either particularized by the sign of the relative velocity. The same is true for the left-sided limit \( s = s_{\text{crit}}^- \) of the rupture point. An infinitesimal distance beyond this point, at \( s = s_{\text{crit}}^+ \), there is only the unbound state possible with the two signs for incoming and outgoing velocities. This gives us in total ten detailed pair correlation coefficients. These are determined by the following ten equations describing the
hysteric flow of probability, as it we can be read off from Fig. 8

Conditions on the contact shell

\[ g^u_{\text{c-\rightarrow r}} = 0 \]  
\[ g^b_{\text{c-\rightarrow r}} = g^u_{\text{c-\rightarrow r}} + g^b_{\text{c-\rightarrow r}} \]  

(33)  
(34)

The Eq. (33) expresses that no particles rebound without a liquid-bond, but rather that all return with a bridge as stated by (34). This implies that the particle number is conserved in collisions (in contrast to the absorbent dynamics modelled in [18] for \( D = 1 \)).

Domain of capillary interaction

\[ g^b_{\text{c-\rightarrow r}} = \gamma^b(s_{\text{crit}}) g^b_{\text{c-\rightarrow r}} \]  
\[ g^u_{\text{c-\rightarrow r}} = \gamma^u(s_{\text{crit}}) g^u_{\text{c-\rightarrow r}} \]  
\[ g^u_{\text{c-\rightarrow r}} = \gamma_{\text{pass}} g^u_{\text{c-\rightarrow r}} \]  

(35)  
(36)  
(37)

The functions \( \gamma^u(s) \) and \( \gamma^b(s) \) take into account the near-contact decay of the pair correlation without and with liquid bond, respectively. The last Eq. (37) describes spectator grains, i.e. grains which pass through the domain of possible capillary interaction without bridge formation. The fraction of these passing particles, \( \gamma_{\text{pass}} = 1 - (1 + s_{\text{crit}}/d)^{1-D} \) (= \( 1/(1 + d/s_{\text{crit}}) \) for \( D = 2 \)) equals the gap between the considered cross section \( (2d + 2s_{\text{crit}})^{D-1} \) of the capillary interaction and the hard-core cross section \( (2d)^D \).

Conditions on the rupture shell

\[ g^b_{\text{c-\rightarrow r}} = 0 \]  
\[ g^u_{\text{c-\rightarrow r}} = g^u_{\text{c-\rightarrow r}} \]  
\[ g^b_{\text{c-\rightarrow r}} + g^u_{\text{c-\rightarrow r}} = g^u_{\text{c-\rightarrow r}} \]  

(38)  
(39)  
(40)

The Eqs. (38, 39) state that only unbound particles enter the domain of capillary interaction, and (40) describes the rupture of a capillary bridge when the pair escapes from the domain.

The hysteretic capillary dynamics is coupled to the hard particle dynamics by the source term of new unbound pairs of particles entering the capillary interaction range:

Source Term

\[ g^u_{\text{c-\rightarrow r}} + g^u_{\text{c-\rightarrow r}}/\gamma^u(s_{\text{crit}}) = (1 - K/K_{\text{sites}}) g^b_{\text{c-\rightarrow r}} \]  

(41)

The left-hand side is the current of approaching unbound neighbors (measured at contact). If all neighbors were unconnected, \( K = 0 \), this current would equal the dry value \( g_{\text{c-\rightarrow r}}^{\text{at}} \). But since there are \( K \) neighbors with bonds out of the \( K_{\text{sites}} \) ‘docking sites’ which are sterically accessible for liquid bonds, the remaining unconnected fraction is \( 1 - K/K_{\text{sites}} \).

The final tenth equation is the stationary state condition, which demands that the rupture frequency equals the binding frequency:

Stationary state condition

\[ f_{\text{bind}} = f_{\text{rupt}} . \]  

(42)

These frequencies follow from the probability to have a particle on the collision or rupture shell, respectively, multiplied by the radial component of the relative velocity under the condition that the particle moves in the appropriate direction for the event to occur. This is analogous to the case \( D = 1 \) [1], with the only difference that here we have to integrate over shells:

\[ f_{\text{bind}} = 2^{D+1}D \sqrt{\frac{T}{\pi}} \frac{\phi}{d} g^u_{\text{c-\rightarrow r}}, \text{ and} \]  
\[ f_{\text{rupt}} = 2^{D+1}D \sqrt{\frac{T}{\pi}} \frac{\phi}{d} g^b_{\text{c-\rightarrow r}} \gamma_{\text{vol}}(s_{\text{crit}}) . \]  

(43)  
(44)

The volume factor \( \gamma_{\text{vol}}(s) = (1 + s/d)^{(D-1)} \) takes the increased size of the outer rupture shell as compared to the inner binding shell into account.
Eliminating those correlation coefficients that are identically zero, we can arrange the coupling equations for the domain of capillary interaction as a $6 \times 6$ matrix system:

\[
\begin{bmatrix}
1 & 1 & -1 & 0 & 0 & 0 \\
0 & \gamma^b & \gamma^b & 0 & -1 & 0 \\
\gamma^u & 0 & 0 & -1 & 0 & -1 \\
-1 & 0 & 0 & 0 & \gamma_{\text{vol}} & 0 \\
0 & 0 & 0 & -1 & 0 & \gamma_{\text{pass}} \\
1 & 0 & 0 & 0 & 0 & 1/\gamma^u
\end{bmatrix} \times \begin{bmatrix}
g^{\text{un}}_{00} \\
g^{\text{un}}_{00} \\
g^{\text{un}}_{00} \\
g^{\text{vol}}_0 \\
g^{\text{pass}}_0 \\
g^{\text{at}}_\circ
\end{bmatrix} = (1 - K/K_{\text{sites}}) \begin{bmatrix}
g^{\text{at}}_{00} \\
g^{\text{at}}_{00} \\
g^{\text{at}}_{00} \\
g^{\text{vol}}_0 \\
g^{\text{pass}}_0 \\
g^{\text{at}}_\circ
\end{bmatrix} \tag{45}
\]

The $\gamma$-functions in the matrix are to be evaluated at $s = s_{\text{crit}}$. As it has to be on physical grounds, this system is non-singular with determinant $(1 + s_{\text{crit}}/d)^{D-1} (2 + \gamma_{\text{pass}}) \gamma^b(s_{\text{crit}}) > 0$.

The last row of the system describes the creation of new liquid bridges as discussed before in the context of the equivalent Eq. (41). We remark that here we used that the correlation $g^{\text{at}}_\circ$ of the dry system at contact has equal contributions from positive and negative relative velocities, immediately before and after the collision, which is still true for the wetted elastic particles we consider. This symmetry between positive and negative radial relative velocities is broken if one wishes to introduce a restitution coefficient $0 < \epsilon < 1$ to model inelastic collisions: the contact correlation of positive velocities is then increased by a factor $1/\epsilon$ as compared to the negatives.

One should see clearly the very different meaning of $K$ and $K_{\text{sites}}$. The dynamical quantity $K$ is the number of instantaneously existing capillary bonds:

\[
K = 2^D D \phi \frac{g^{\text{un}}_{00}}{d} \int_0^{s_{\text{crit}}} \gamma^b(s) \gamma_{\text{vol}}(s) \, ds . \tag{46}
\]

$K$ rapidly decays close to zero in dilute systems. As $K$ comes closer to the value of $K_{\text{sites}}$ in a very dense system, the binding frequency $\gamma_{\text{bind}} \propto g^{\text{un}}_{00} \propto K - K_{\text{sites}}$ goes to zero because steric hindrance prohibits the formation of further capillary contacts: $K - K_{\text{sites}}$ gives the number of vacant sites for capillary bonds. Therefore $K_{\text{sites}}$ is the maximum number of docking sites for capillary bonds. It is a pure geometric property and grows with $s_{\text{crit}}$, because $s_{\text{crit}} > 0$ still allows for a slight rearrangement of particles in the formation of new capillary bridges without breaking existing ones. In the limit $s_{\text{crit}} \to 0$, $K_{\text{sites}}$ is the number of contact sites. We therefore expect $K_{\text{sites}}$ to equal the number of exact contacts, $2D = 4$. So let us compute $K_{\text{sites}}(\phi, s_{\text{crit}})$ in the following paragraph.

The maximum number of possible bonds, $K_{\text{sites}}$, is an athermal function of density $\phi$ and the critical liquid bridge length $s_{\text{crit}}$. We determine $K_{\text{sites}}$ from the obvious fact, that the granular dynamics is unaffected by the introduction of forceless bridges: for $\gamma^b = \gamma^u$ we recover the dry contact correlation $g^{\text{un}}_\circ + g^{\text{un}}_\circ = g^{\text{at}}_\circ$. This is the athermal limit, or high temperature limit of wet granular matter.

\[
\text{High temperature limit: } \left\{ \begin{array}{c}
\gamma^b = \gamma^u \\
g^{\text{un}}_\circ + g^{\text{un}}_\circ = g^{\text{at}}_\circ
\end{array} \right. \quad (47)
\]
FIG. 9: The capillary bridge coordination $K$ in the low temperature limit, $T \ll E_{cb}$. As proven in the text, $K$ converges to the athermal function $K_{\text{sites}}(\phi, s_{\text{crit}})$ in this low temperature limit. The solid line is $K_{\text{sites}}(\phi_0, s_{\text{crit}})$ over a wide range of maximal bridge lengths $s_{\text{crit}}$. Points represent final states of free cooling simulations with 1000 particles of uniformly distributed polydispersity $\Delta d = 0.06d$. The open symbols are clusters with winding number one (cylindrical topology), connected over one periodic boundary on a rectangular domain. Such structures have internal tensile strength which necessitates a slightly increased coordination, visible as a small shift compared to the closed symbols which represent localized clusters (as the two examples drawn in the plot). As predicted by Eq. (51) of the presented theory, the structures emerging with exact contacts, $s_{\text{crit}} \to 0$, are found to be precisely isostatic, $K_{\text{sites}} = 4$. The line is the the analytic result (50), for which very good agreement is found with the simulations over the entire range of the capillary bridge regime, $0 < s < 0.2r$ (with $r$ the particle radius), which is indicated in the figure. Beyond this regime, the theory does not hold because in the derivation we limited ourselves to the leading order in $s_{\text{crit}}/d$. More importantly, the rupture length $s_{\text{crit}}$ cannot be further increased beyond the capillary regime by simply increasing the liquid content in the granular sample. As mentioned in the introduction, liquid bridges residing on the same sphere would rather merge [21] into more complicated objects.

FIG. 10: (A) The maximum number of 'docking sites' for capillary bonds on a particle, $K_{\text{sites}}$, which is possible in two dimensions given the pinch-off length $s_{\text{crit}}$ of capillary bridges and the density $\phi$. $K_{\text{sites}}$ is independent of the temperature because it is a pure geometric quantity: the number of possible neighboring sites. Here $K_{\text{sites}}$ is shown as a function of $\phi$ for different $s_{\text{crit}}$ ranging from $s_{\text{crit}}/d = 0.07$ (solid curve), 0.04, 0.02, to 0.01 (short dashes). As is shown in the text, $K_{\text{sites}}$ is the coordination number in the zero temperature limit. While the mean coordination number $K$ rapidly goes to zero with density for finite temperature, the zero-temperature limit is $\geq 4$ for all densities, because the system clusters. The dotted curve represents the limit $s_{\text{crit}} \to 0$. The longer $s_{\text{crit}}$ the closer $K_{\text{sites}}$ comes to six, the number of next neighbors in two dimensions. In the limit $s_{\text{crit}} \to 0$ the coordination $K_{\text{sites}}$ converges to the number of exact contacts which is precisely four. (B) The capillary bridge coordination $K$ drops down in the vicinity of the critical temperature as clustered structures break up. For this plot the mean density is chosen to be $\phi = 0.75$. 
From the hysteretic bridge system (45) follows in this forceless or high granular temperature limit (47):

\[
(1 - \frac{\gamma^u \gamma_{\text{vol}}}{1 + \gamma_{\text{pass}}}) K_{\text{sites}} = (1 + \gamma^u \gamma_{\text{vol}}) K \quad \text{and}
\]

\[
g_{\text{pass}}^b = \frac{g_{\text{pass}}^{\text{at}}}{1 + \gamma^u \gamma_{\text{vol}}}
\]

The \(\gamma\)-functions with the argument \(s\) suppressed are understood to be evaluated at \(s = s_{\text{crit}}\). Inserting the Eqs. (48) and (49) in (46) yields

\[
K_{\text{sites}} = 2^D D \phi \int_0^{s_{\text{crit}}} g_{\text{pass}}^{\text{at}} \frac{\gamma^u(s_{\text{crit}}) \gamma_{\text{vol}}(s_{\text{crit}})}{1 - \gamma^u \gamma_{\text{vol}}/(1 + \gamma_{\text{pass}})} \, ds/d
\]

\[
= 4 \frac{1 - \gamma^u + \mathcal{O}(s_{\text{crit}}^3)}{1 - \gamma^u + \mathcal{O}(s_{\text{crit}}^2)} = 4 + \mathcal{O}(s_{\text{crit}}) .
\]

In the last line we have set \(D = 2\), so that we could use (10) and (14). The result (51) is the second important consistency test. Finding the number of exact contacts in the jamming limit to equal four in (12) showed the consistency of the free-volume argument applied there. Here in (51) we find for any density that the different function \(K_{\text{sites}}\) for the number of possible bridges sites equals four as well when \(s_{\text{crit}} = 0\). This is as intuitively expected and a confirmation of the consistency between the hysteretic system (45) and the near-contact pair correlation. In view of the numerical finding \(Z_{\text{sim}} = 2\) for the derivative at contact of the pair correlation (as defined in (23)) we remark that the entirely analytic description by the hysteretic system gives in general \(K_{\text{sites}} = 8/Z + \mathcal{O}(s_{\text{crit}})\), which is why the consistency is non-trivial and the finding \(Z_{\text{sim}} = 2\) fits favorably into the entire picture.

Thus the hysteretic system (45) provides a direct connection between the static granular properties captured in \(K_{\text{sites}}\) and the granular system in motion at positive granular temperature which we are treating in general. We remind that \(K_{\text{sites}}\) is determined by the steric self-hindrance and therefore a pure geometric property independent of the granular temperature. When inspecting a snapshot of a close granular packing we can find local cases of contact coordination \((s_{\text{crit}} = 0)\) higher than four. These are fluctuations within the granular ensemble, while \(K_{\text{sites}}\) and \(K\) are mean-field quantities. Of course, for a finite bridge length \(s_{\text{crit}} > 0\), a mean bridge coordination \(K \in (0, K_{\text{sites}})\) with \(K_{\text{sites}}\) higher than four is possible due to elongated bridges, as described by (40). Before we evaluated the expression (50) of \(K_{\text{sites}}\) for positive \(s_{\text{crit}}\) (plotted in Fig. 9), it is enlightening to switch on the capillary forces in the following section because this allows us to apply \(K_{\text{sites}}\) to ‘frozen’ wet granular matter.

**B. Switching On the Force of Capillary Bridges**

Under the attraction of a liquid bridge, the pair correlation \(g^b(s)\) of connected neighbors falls off faster than \(g^u(s)\) for unbound particles, depending on the granular temperature \(T\)/\(E_{\text{ch}}\) compared to the bridge energy. The logarithmic derivative of the radial pair correlation is to be interpreted as the effective radial force \(\beta F = \partial_s \ln g(s)\), as discussed before in section 11B. This exponential dependence can be justified as the solution of the Fokker-Planck equation derived in [12]. Moreover, in the context of the hysteretic interaction of wet granular matter this exponential factor has been successfully applied in the case \(D = 1\) (cf. Eq. (3) in [1]). Therefore we proceed by switching on the liquid bridge force to the physical value of the Minimal Capillary model [11], \(F_b = E_{\text{ch}}/s_{\text{crit}}\), including this exponential in the short-range dependence of the pair correlation function for bridges neighbors:

\[
\gamma^b(s, T) = \gamma^u(s) \exp \left( -\frac{E_{\text{ch}}}{T} \frac{s}{s_{\text{crit}}} \right) .
\]

At low granular temperatures this exponential gives rise to shorter average bridge lengths, and describes the reduced probability that a bridge reaches its critical length \(s_{\text{crit}}\). Therefore the hysteretic system (45) describes the sticking of particles and the onset of clustering.

We have discussed in the previous section 11.1A that steric effects in the dynamical system limit the mean number of bonds to a maximum of \(K_{\text{sites}}\), and we derived that \(K_{\text{sites}}\) converges to the number of isostatic contacts in the limit \(s_{\text{crit}} \to 0\). Here this connection is put on firm grounds with a clear physical interpretation attributed to \(K_{\text{sites}}\): \(K_{\text{sites}}\) is the bridge coordination \(K\) of solid wet granular matter.
Proof of $K \to K_{\text{sites}}$ in the low temperature limit. Solving (15) for $K/K_{\text{sites}}$, we obtain

$$
\frac{K(T, s_{\text{crit}}, \phi)}{K_{\text{sites}}(s_{\text{crit}}, \phi)} = \frac{1}{1 + X(T)/Y(T)},
$$

(53)

with

$$
X(T) = \gamma^b(s_{\text{crit}}, T) K_{\text{sites}} (\gamma_{\text{pass}} + 2) \gamma_{\text{vol}} \quad \text{and}
$$

$$
Y(T) = 8 I(T) \phi g_{\text{at}}^b (\gamma_{\text{pass}} + 1),
$$

(54)

(55)

where $I(T)$ stands for the integral over bond states,

$$
I(T) = \int s_{\text{crit}}^{s_{\text{crit}}} \gamma^b \gamma_{\text{vol}} \frac{ds}{d} = \int s_{\text{crit}}^{s_{\text{crit}}} e^{-\frac{E_{\text{ch}}}{T}} \gamma^n \gamma_{\text{vol}} \frac{ds}{d} = \frac{T s_{\text{crit}}}{E_{\text{ch}} d} + O(T^2),
$$

(56)

which goes linearly to zero, while $\gamma^b(T) \propto e^{-E_{\text{ch}}/T}$ vanishes for $T \to 0$ faster than any power of $T$. Hence $X/Y \to 0$ so that Eq. (53) implies

$$
\lim_{T \to 0} K = K_{\text{sites}}
$$

(57)

as conjectured.

This low temperature limit ($T \ll E_{\text{ch}}$) is of general interest since it represents a sticky gas of ideal spheres, which serves as a model for aggregation in various areas of physics [48] and astrophysics [49]: once two particles had contact, the remaining degree of freedom is tangential motion. The analytic prediction of formula (51) is $K_{\text{sites}} = 4$ in the limit of exact contacts, $s_{\text{crit}} = 0$. In order to evaluate (50) for positive $s_{\text{crit}}$ we insert the near-contact decay $\gamma^n$ given by the general results [10], [14], and [22], setting $D = 2$. The explicit expression for $\gamma^n$ which we use throughout this article for results without free parameters is given in the appendix B. Here we take into account known formulas for the contact value $g_{\text{at}}^b$ at low densities, as well as higher corrections to the free volume theory. Inserting this expression in (50) results in the curve shown in Fig. 3. We have performed simulations in this low-temperature limit. The wet granular matter was initially prepared in a gas state with $T = 50 E_{\text{ch}}$ and cooled by the formation and rupture of bonds. The insets in Fig. 3 show final states when the granular temperature $T$ is more than one order of magnitude below $E_{\text{ch}}$ and no further change in the configuration was observed on exponential time scales. The symbols in Fig. 3 have been measured in this final state. In perfect agreement with the prediction of Eq. (51), we find in the contact limit, $s_{\text{crit}} \to 0$, the coordination to be exactly 4. Moreover, the increase in the number of bonds per particle with the increase of the maximal bridge length $s_{\text{crit}}$ is found to be in very good agreement with the simulations.

Further analytic results for high densities are shown in Fig. 10 (A). As is intuitively clear and shown by the family of curves in Fig. 10 (A), the convergence of the limit $s_{\text{crit}} \to 0$ is not uniform with respect to density, since $K_{\text{sites}}$ is pinned to the kissing number 6 of the monodisperse crystal density at $\phi_{\text{max}}$.

IV. THE EQUATION OF STATE

We are now in the position to derive the equation of state, $P = P(T, \phi)$, for wet granular matter with capillary bonds tensile up to the rupture length $s_{\text{crit}}$. The cohesion of capillary bridges will reduce the pressure as compared to a dry hard-sphere system of equal temperature. By virtue of Eq. (53), we have the bridge coordination number $K$ as a function of density $\phi$ and granular temperature $T$. Since in the Minimal Capillary model [11] the bridge force is assumed to be independent of the bridge length $s$, the knowledge of the mean number of bridges $K$ will allow us to evaluate the reduction of the pressure due to cohesion. Furthermore, the particle-particle collisions are enhanced by the bridge attraction, increasing the contact correlation. The contact correlation $g_{\text{wet}}^b$ for wet granular matter derives from the Eqs. (45) and (10):

$$
g_{\text{wet}}^b = \frac{(1 + \gamma_{\text{pass}})(1 + \gamma^b \gamma_{\text{vol}}) I^u}{(1 + \gamma_{\text{pass}} - \gamma^u \gamma_{\text{vol}}) I^b + (2 + \gamma_{\text{pass}}) \gamma^b \gamma_{\text{vol}} I^u}
$$

(58)

with the integrals

$$
I^{u/b} = \int_0^{s_{\text{crit}}} \gamma^{u/b}(s) \gamma_{\text{vol}}(s) \, ds.
$$

(59)
FIG. 11: A local configuration of two-dimensional wet granular matter at moderate density. Since there is one particle in each Voronoi cell, the mean area equals the inverse density. In section II A we have used the Voronoi tessellation to compute the derivative of the pair correlation at contact for a dry and dense system. For such a dense system, the Voronoi cell resembles a hexagon with a size proportional to \((d + s)^2\), where \(s\) is the particle separation. The cell borders are at half surface separation for polydisperse diameters (not half center distance), so that each cell contains one particle completely.

The analytic expression (58) for the contact correlation of wet granular matter, \(g_{\text{wet}}^{c}\), is indeed strictly greater than the one of the dry system, \(g_{\text{at}}^{c}\), to which it converges in the high temperature limit when the capillary energy \(E_{cb}\) is small compared to the granular temperature \(T\). This limit follows obviously from (58) because the functions with superscript index 'b' turn into those with 'u' for \(T \gg E_{cb}\). In the low temperature limit, liquid bonds oscillate with an amplitude proportional to the kinetic energy which equals \(T\) on average, so that the probability to find the particles at contact, \(g_{\text{wet}}^{c}\), grows proportional to \(1/T\), as can be derived easily from (58) using the expansion (56).

### A. Frozen Degrees of Freedom

As the system starts to cluster at temperatures close to \(E_{cb}\), voids remain between the clusters with linear dimensions large compared to the particle diameter. Clearly, this growing length scale, which is set by the sizes of clusters and voids, is not captured by the short-range behavior of the pair correlation function. Here we advance the theory beyond the level of two-particle correlations to take correlation on large scales, such as the collective particle motion in a cluster, in an approximative fashion into account.

The collective motion of a cluster is due to stable capillary bonds which impose constraints, such that the internal degrees of freedom of clusters are frozen. Since \(K\) is the number of instantaneous capillary bridges of which the fraction \(\operatorname{erf}\left(\sqrt{E_{cb}/T}\right)\) with kinetic energies below \(E_{cb}\) forms stable bonds, we have

\[
K_{\text{frozen}} = K \operatorname{erf}\left(\sqrt{E_{cb}/T}\right)
\]

for the number of frozen degrees of freedom.

We are interested in the density of the remaining degrees of freedom. The idea is simple and powerful: As a general mathematical property of triangulations, there are on average precisely six Voronoi neighbors \([39]\), independent of density or ordering. In Fig. 11 we can observe that the Voronoi neighbors with stable bonds contribute less to the area \(1/n\) of the Voronoi cell. This picture suggest a two-fluid model with frozen and free neighborhoods as the two constituents. The fraction of frozen and free triangulation bonds is proportional to \(K_{\text{frozen}}\) and \(K_{\text{free}}\) respectively, and the area contributions associated to each bond sum up to the total size of the Voronoi cell:

\[
\begin{align*}
K_{\text{frozen}} + K_{\text{free}} &= 6 \\
\frac{K_{\text{frozen}}}{n_{\text{frozen}}} + \frac{K_{\text{free}}}{n_{\text{free}}} &= \frac{6}{n}.
\end{align*}
\]

Because of this reciprocal sum rule for the densities one may call this a reciprocal two-fluid model.

The density \(n_{\text{frozen}}\) of the stable bond component follows analogously to (2) when averaged with the additional
We consider for instance a compressed state of wet granular matter with has contributions only by the unfrozen pairs of particles with density $n_{\text{free}}(T, s_{\text{crit}}, \phi)$, i.e. $r/s/d$. Without affecting the leading order in $s/d$ one is free to replace the last $s$ in the exponent (65) by $s + s^2/(2d)$, so that the integral (63) is elementary resulting in

$$\frac{n_J}{n_{\text{frozen}}} - 1 = \left[\left(1 + \frac{s_{\text{crit}}}{d}\right)^D - 1\right] \left[\frac{1}{\alpha} - \frac{1}{e^\alpha - 1}\right]$$

(66)

with $\alpha = \left[\left(1 + \frac{s_{\text{crit}}}{d}\right)^D - 1\right] \left[\phi \frac{g^\text{at}}{T} + \frac{E_{\text{cb}}}{D s_{\text{crit}}}\right]$

(67)

We point out that Eq. (66) implies Eq. (4) in [1] for $D = 1$.

From the Eqs. (63), (65), and (66) follows the density of degrees of freedom which are not frozen out by capillary bonds, $n_{\text{free}}(T, s_{\text{crit}}, \phi)$. One may regard $n_{\text{free}}$ as the density of clusters.

We remark that the two-fluid model of neighborhoods is the only concept presented in this theory of wet granular matter which cannot be generalized in a straight forward manner to three dimensions, because for $D = 3$ the number of Voronoi neighbors (double counted per particle) is not a universal constant (such as 6 for $D = 2$ and 2 for $D = 1$), but depends on the granular order (reaching its minimum value 12 for close packing and its maximum of approximately 15.5 in the ideal gas limit) in D. The reason for this is that three-dimensional space cannot be filled with tetrahedrons, while flat space can be tiled by triangles. As a consequence, the number of constituents in the two-fluid model of neighborhoods would not be conserved for $D = 3$ and the numerator on the right-hand side of (62) is not a constant.

### B. The Pressure of Wet Granular Matter

Here we arrive at the pressure $P(T, \phi)$ using the density $n_{\text{free}}(T, \phi)$ of degrees of freedom, the coordination $K(T, \phi)$, and the contact correlation $g^\text{vol}(T, \phi)$. The pressure is the trace of the stress tensor

$$P = -\frac{1}{D} \text{tr} \sigma$$

(68)

The stress tensor $\sigma = \sigma^\text{kin} + \sigma^\text{force}$ describes the flow of momentum. The kinetic term has components $\sigma_{i,j}^\text{kin} = -\sum_k \langle mv_i(i) v_j(k) \delta(\text{r} - r(k)) \rangle$. With the granular temperature $T = \langle mv_i v_i \rangle$, its trace yields $nT$ for uncorrelated particle motion (as in an ideal gas). In general we have the kinetic contribution

$$P^\text{kin} = n_{\text{free}}T$$

(69)

wherein there frozen degrees of freedom have been taken out. For moderate densities, one may interpret (69) as the kinetic contribution to the pressure due to a gas of clusters.

The interparticle forces $F$ give rise to the Cauchy tensor $\sigma^\text{force}$, which is the tensor product of the center-to-center vector $\text{r}$ and the pair force $F$, $\sigma^\text{force} = \frac{n_{\text{free}}}{2} (F \otimes \text{r})$.

(70)

so that $\sigma^\text{force}$ is diagonal for radial forces. The factor $1/2$ assigns half of the momentum current to either of the interaction particles, i.e. $r/2$ may be seen as the transport vector within the Voronoi cell. The Cauchy tensor (70) has contributions only by the unfrozen pairs of particles with density $n_{\text{free}}$, because in frozen neighborhoods the repulsive momentum exchanged in collisions is exactly balanced by the bridge attraction under the time average on the right-hand side of (70).

A comment on the significance of the reciprocal two-fluid model as represented by Eq. (61) and (62) is in order here. We consider for instance a compressed state of wet granular matter with $K_{\text{frozen}}$ around five and $K_{\text{free}}$ around unity.
FIG. 12: The pressure $P$ of wet granular matter is shown as function of the granular temperature $T$. The dimensionality is $D = 2$ and the covered area fraction is $\phi = 0.1$, so that at high temperatures the system is a dilute gas. The maximum bridge length is $s_{\text{crit}} = 0.07d$. The behavior below the critical temperature $T_c = 0.274E_{\text{cb}}$ of wet granular can be understood in the following way: the system agglutinates to clusters. With these effective particles the pressure is reduced according to the reduced number density of effective particles. The breakup of clusters is reflected by the rising pressure around $T_c$. The straight line is the athermal pressure of hard discs, $P^\text{dry} = n_{g,\text{wall}} T$ which is reached asymptotically when the granular temperature is higher than the energy scale $E_{\text{cb}}$ set by the capillary interaction.

While $K_{\text{free}}$ is small, the prefactor $n_{\text{free}}$ in (70) is not necessarily small. From (61) and (62) follows that both, $n_{\text{free}}$ and $n_{\text{frozen}}$, converge to $n_1$ as the system gets jammed ($n \to n_1$), so that the repulsive dominated state is correctly described by the Cauchy tensor (70) which grows beyond all bounds as $n \to n_1$. If one had (in contradiction to the additivity of areas) summed up densities linearly instead of the reciprocal sum rule (62), the free density would vanish or could even become negative under such conditions.

It is finally easy to determine the time average on the right-hand side of (70) for the two different forces acting in wet granular matter, the delta-force in collisions of hard particles and the flat force $F_{\text{cb}} = E_{\text{cb}}/s_{\text{crit}}$ of the capillary bonds. In a collision at time $t_{\text{coll}}$ the radial momentum $\Delta p$ is transferred instantaneously:

$$\langle F_{\text{coll}} \otimes r \rangle = \langle \Delta p \otimes r \delta(t - t_{\text{coll}}) \rangle = I \langle \Delta p \otimes r \delta((r, -v) \cdot \delta(r - d)) \rangle = -I g_{\text{wet}} \sigma_D d^{D-1}. \quad (71)$$

In the last equality the $\delta$-function gives rise to the contact correlation $g_{\text{wet}}$ and the trivial integration of angles leaves $\sigma_D d^{D-1}$. $I$ is the unity matrix and $\theta$ is the Heaviside step function. Inserting (71) in (70) and taking the trace (68) yields

$$P_{\text{coll}} = 2^{D-1} n_{\text{free}} T \phi g_{\text{wet}}. \quad (72)$$

The cohesive virial due to capillary bridges is

$$\langle F_{\text{cb}} \otimes r \rangle = \left( K \frac{E_{\text{cb}}}{s_{\text{crit}}} \frac{r \otimes r}{r} \right) = \frac{1}{D} K \frac{E_{\text{cb}}}{s_{\text{crit}}} (d + s) \approx \frac{1}{D} K \frac{E_{\text{cb}}}{s_{\text{crit}}} d. \quad (73)$$

Hence the final result

$$P = n_{\text{free}} T \left( 1 + 2^{D-1} \phi g_{\text{wet}} \right) - n_{\text{free}} E_{\text{cb}} \frac{K d}{2D s_{\text{crit}}}, \quad (74)$$

where the last term is the bridge cohesion (63). Since $n_{\text{free}}$, the contact correlation $g_{\text{wet}}$ and $K$ have been derived explicitly in (61), (68) and (63) as functions of $\phi$ and $T$, we have the equation of state for wet granular matter, $P = P(\phi, T)$.

The Figs. 12 and 13 show the analytic result (74) as a function of the granular temperature $T$ and the density $\phi$. In the high temperature limit wet granular matter behaves as a hard-spheres system. Below the critical point granular
FIG. 13: Isotherms of wet granular matter for the realistic rupture length $s_{\text{crit}} = 0.07d$. In the high temperature limit the liquid bridges forfeit their influence on the dynamics, so that the equation of state reduces to the hard sphere pressure. This can be seen by the two black isotherms of wet granular matter, of which the higher is at $T = E_{\text{cb}}$ and converges to the green curve in the limit $T \gg E_{\text{cb}}$. The lower black isotherm is at $T = 0.2E_{\text{cb}}$ and exhibits an unstable branch. The critical point is at $T_c \approx 0.274 E_{\text{cb}}$ (cf. Fig. 14 for a close-up).

FIG. 14: A close-up of the transition region in wet granular matter. The dashed line in the main panel is the spinodal of the homogeneously driven wet granular system in $D = 2$ dimensions. The solid black lines are wet granular isotherms around the critical point, which is located at $T_{\text{crit}} = 0.273(5)E_{\text{cb}}$ for $s_{\text{crit}} = 0.07d$. The change of the critical point with the amount of added liquid (represented by $s_{\text{crit}}$) is shown in Fig. 14. The curve in the upper left corner is the athermal pressure $P_{\text{dry}}$ of the hard disc system without liquid bridges, and the line at the bottom is the ideal gas pressure ($P_{\text{id}} = \phi$) has a defined slope). $P_{\text{dry}} = \phi^2$, $P_{\text{id}}$ is increased compared to the ideal gas by the Enskog factor $g_{\text{at}}$. The pressure of wet granular matter is reduced compared to the dry system $P_{\text{dry}}$ due to the capillary cohesion. The inset shows the spinodal in the temperature-density plane, where the critical temperature can be clearly determined.
clusters are predicted to segregate due to the mechanically unstable branch of the pressure as a function of density, which appears in Fig. 13 below the critical temperature. Figure 14 provides a close-up of the critical point of wet granular matter and its spinodal. The critical density of this transition is high, because the particles have to be close enough in order to form a dynamical capillary network. As we show in Fig. 15 the critical density is determined by the length scale of capillary bridges, such that the rupture length $s_{\text{crit}}$ scales with the mean particle separation $s = d \left( \sqrt{3} / \phi_c - 1 \right)$ scales with the rupture length $s_{\text{crit}}$. This shows that both intrinsic characteristics of the capillary interaction, the rupture length and the bridge energy $E_{\text{cb}}$, determine the critical point of wet granular matter.

V. CONCLUSION

Starting with the hard-sphere fluid, an expression (10) for the narrowing of the near contact pair correlation was derived, which describes in the jamming limit the delta-peak of $2D$ isostatic contacts per particle, in agreement with the expected value of simulations. In the gas and fluid regime the fall-off predicted by this expression for the pair

FIG. 15: The influence of the rupture length $s_{\text{crit}}$ on the position of the critical point in the phase diagram Fig. 14 of wet granular matter. The position of the critical point is described by the critical parameters ($\phi_c, T_c$), which are plotted on the left and right vertical axis respectively. Solid lines result from the full theory (74) by solving for the intersection of $\partial \phi P(\phi, T) = 0$ and $\partial^2 \phi P(\phi, T) = 0$. For the critical temperature we find a very mild variation with the rupture length, so that over the entire physically relevant range of capillary interaction we have $T_c \approx E_{\text{cb}}/4$. The influence of the rupture length $s_{\text{crit}}$ on the critical density $\phi_c$ can be understood very clearly with the help of the dashed line. The critical density is such that the mean particle separation $s = d \left( \sqrt{3} / \phi_c - 1 \right)$ scales with the rupture length $s_{\text{crit}}$. This shows that both intrinsic characteristics of the capillary interaction, the rupture length and the bridge energy $E_{\text{cb}}$, determine the critical point of wet granular matter.
correlation at contact was found to be well confirmed by simulations. We then addressed the nonequilibrium case of wet granular matter by the introduction of capillary bridges which are formed hysteretically. The description in terms of the pair-correlation function was extended with six different non-vanishing correlation coefficients which take the bridge status into account and allow for the hysteretic dissipative dynamics. The coordination number of bonds was computed analytically as a function of the rupture length of the capillary bridges, the granular temperature, and the density. The limiting case of strong bonds led to the sticky gas dynamics for which simulations have been performed which showed very good agreement with the analytic prediction of the coordination number. Based on the derived expressions for the contact correlation and the bridge coordination, we finally computed the pressure of wet granular matter analytically as a function of density and granular temperature. Here a method was put forward, which describes the effective degrees of freedoms in order to take the correlated motion of particles glued to clusters into account. The isotherms of wet granular matter were found to have an unstable branch which gives rise to the segregation of dense clusters. The critical temperature of this transition was derived to be approximately one quarter of the capillary bond energy. The critical density is directly related to the pinch-off distance of the capillary bridges. A close relation to the un-clustering effect reported in one dimension [1] was shown, for which reason this effect persists also in higher dimensions.

It will be interesting to probe the critical point of wet granular matter experimentally and by direct simulations. As we have shown, the position of the critical point is determined by the length and energy of the capillary bridges. These quantities can be controlled very accurately in an experiment of shaken wet granular matter. Therefore the measurement of the critical temperature will allow to discern between extensions such as the nonlinear coupling discussed in appendix.

Future analytic work includes the background contribution $g_B$ in the dense regime, since our numerics indicate that the pair correlation is flatter near the contact as predicted by $g_A$ alone. This task might be addressed in conjunction with the analogous background contribution in three dimensions, for which in the jamming limit an integrable power-law divergence, $g_B \propto 1/s^6$, has been reported in numerical studies (with $\delta = 0.5$ [51] or $\delta = 0.6$ [32]) and experiments [52], but is as well lacking a theoretical explanation at present.

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APPENDIX A: THE BACKGROUND CONTRIBUTION $g_B$

1. The Weighting Factors

With $g_A$ in [10] we considered the four (cf. Eq. [12]) A-neighbors, which form isostatic contacts at jamming, $s_A \to 0$ for $\phi \to \phi_J$ [8], and are separated by $s_A$ according to Eq. [1] before jamming. Analogously, the separation $s_B$ of the two B-neighbors is weighted by

$$P_B(s_B) \propto \exp \left( -\frac{(1 + \frac{mB}{\phi})^2 - 1}{\phi_{\text{max}}/\phi - 1} \right). \tag{A1}$$

While in Eq. [11] the denominator in the exponential is $c_A = \phi_J/\phi - 1$ so that $s_A \to 0$ at the jamming density, in Eq. [A1] the denominator is $c_B = \phi_{\text{max}}/\phi - 1$ since the blocked B is only forced to form a contact, $s_B \to 0$, for a perfect crystal with $\phi \to \phi_{\text{max}}$. Of course this limit is kinematically unreachable because the system comes to rest at the jamming density $\phi_J < \phi_{\text{max}}$. $\phi_{\text{max}}$ would be reached. We note that $c_B$ is a small dimensionless quantity: for $\phi > \phi_o = 0.71$ we have $0 < c_B < 0.2774$.

Close to jamming, the B-neighbors are fixed in space by particles other than the reference particle. Except for arch-like constructions which are rare for frictionless particles, and would include second Voronoi neighbors keeping B at a separation larger than our region of interest, $s_B > s_{\text{crit}}$, this hindrance is due to the A-neighbors. Therefore the probability $g_B(s_B)$ to find a B-neighbor at separation $s_B$ from the reference particle (sketched with hatching in Fig. [17]) is given by the integral over all configurations where four A-neighbors hinder two B-neighbors.

The configurations will be weighted by a phase space factor $C$ and the exponential factor $P_B$. We are above the ordering density $\phi_o$, so that the neighborhood has (by definition of the phase) hexagonal order as sketched in the inset of Fig. [16]. Projecting the configurations with the two B-neighbors blocked (gray subset in Fig. [16]) on a single $\theta$-axis, we find the configuration space factor

$$C(\theta) = \frac{3(5\pi - 6\theta)}{2\pi^2}. \tag{A2}$$
FIG. 16: The angular configuration space of four neighbors close to the reference particle. These we denote as A-neighbors. The faceted inner subset shown in gray is the subspace conditioned to the property that two further particles, the B-neighbors, are hinder by the A-particles in approaching the reference particle. The projection of this subset onto an $\theta$-axis (for the angle between a blocking A-pair, $\theta_1$ or $\theta_3$ in this example) gives rise to a linear configuration space factor $C(\theta)$. Obviously a B-neighbor acts like a wedge driven between two A-neighbors, and therefore increases $\theta$. This is taken into account by the weighting factor $P_B(s_B)$ which favors shorter separations $s_B$ between the particle B and the reference particle, depending on the density $\phi$.

In the sequel we abbreviate

$$\gamma_{\text{vol}}(s) \propto 1 + s/d .$$  \hfill (A3)

for the volume factor (A1) in $D = 2$. Wide gaps of length $s_B$ are exponentially suppressed by $P_B$.

2. The Configuration Space

Let us now address the configuration space plotted in Fig. 17. If the opening angle $\theta$ of the A-neighbors exceeded $\theta_T(s_A)$,

$$\cos \frac{\theta_T(s_A)}{2} = \sqrt{s_A(2d + s_A)} \frac{1}{d + s_A} ,$$  \hfill (A4)

the B-particle could slip through and turn into an A-neighbor, which is defined by having a free path towards the reference particle. This transition corresponds to the neck connecting different jamming island in the configuration space. Only along the line (PQ in Fig. 17) defined by $\theta = \theta_C(s_A)$,

$$\cos \frac{\theta_C(s_A)}{2} = \frac{s_A + d}{2d} ,$$  \hfill (A5)

the B-neighbor can touch the reference particle, so that $s_B = 0$. The Eqs. (A4) and (A5) define the upper boundary of the domain of integration for all $s_A$,

$$\theta_{\text{max}}(s_A) = \begin{cases} \theta_C(s_A) & s_A/d \leq \sqrt{2} - 1 \\ \theta_T(s_A) & s_A/d \geq \sqrt{2} - 1 \end{cases} ,$$  \hfill (A6)

which is continuously differentiable but not smooth at the point Q. The lower boundary is

$$\cos \frac{\theta_{S}(s_A, s_B)}{2} = \frac{(s_A + d)^2 + s_B^2 + 2ds_B}{2(s_A + d)(s_B + d)} ,$$  \hfill (A7)
FIG. 17: A section of the configuration space of neighboring particles. Within the gray domain the particle denoted by B is blocked: the two neighbors labelled A sterically hinder the particle B from approaching the reference particle (shaded). Only at the boundary $\theta_{\text{C}}(s_A)$ (curve PQ ranging from $[s_A, \theta]^P = [0, 2\pi/3]$ to $[s_A, \theta]^Q = [(\sqrt{2} - 1)d, \pi/2]$) the B-neighbor can touch the reference particle. The probability $g_B(s_B)$ to find a B-neighbor at a separation $s_B$ follows from integrating over the gray domain, which grows with increasing $s_B$. The lower bound, $\theta_{\text{S}}(s_A, s_B)$, is plotted for the values $s_B = 0.05$, $0.30$, and $0.60$. Large areas spanned by this neighborhood are exponentially rare the higher the mean density $\phi$, so that the probability distribution in this plot concentrates in the vicinity of the upper left corner P as we come closer to the jamming limit. At the line QR the B-neighbor slips through and turns into an A-neighbor, so that QR is the transit to another jamming island in configuration space. The corresponding transition rate is proportional to the probability density along QR and therefore vanishes in the jamming limit.

FIG. 18: The $s_A-\theta$-plot of Fig. 17 with the full $s_B$ dependence shown on the additional vertical axis.
where B hits A.

The simple lower bound on $\theta$,

$$\cos \frac{\theta_{\text{min}}(s_A)}{2} = \sqrt{1 - \left( \frac{d/2}{d + s_A} \right)^2}, \quad (A8)$$

which ensures that the A-neighbors do not overlap is without applicatory relevance, as it implies that the B-neighbor is pushed out to $s_B/d > \sqrt{3} - 1 \approx 0.73$. This is suppressed in the dense regime $\phi > \phi_0$ by the factor $F$ of Eq. (A1).

The configuration space ends to its right in a cusp where the lower and upper bound intersect at

$$s_A^{\text{cusp}}(s_B) = \sqrt{s_B^2 + 2d s_B + 2d^2 - d}. \quad (A9)$$

This cusp converges to the point Q for $s_B \to 0$.

With the integration bounds $[A6], [A7], [A9]$, and the weighting factors $[A1], [A2]$ we have

$$g_B(s_B) = N P_B(s_B) \left[ \int_{0}^{s_B^{\text{cusp}}(s_B)} ds_A P_A(s_A) \gamma_{\text{vol}}(s_A) \int_{\theta_{0}(s_A,s_B)}^{\theta_{\text{max}}(s_A)} d\theta \ C(\theta) \right]^2 \quad (A10)$$

$$= N P_B(s_B) \left[ \frac{s_B}{d} I_1(n) + \left( \frac{s_B}{d} \right)^2 I_2(n) + O \left( \left( \frac{s_B}{d} \right)^3 \right) \right]^2. \quad (A11)$$

We emphasize that the configuration space $(s_A, \theta)$ describes the relative position of one A-neighbor sketched symmetrically in Fig. (17) Since there are two independent A-neighbors involved, their configuration is the direct product $(s_{A1}, \theta_1) \times (s_{A2}, \theta_2)$. On this account the configuration integral is squared in (A10), with the important consequence that the leading order in $g_B(s_B)$ is quadratical. The normalization constant $N$ is determined by the knowledge that there are two B-neighbors. While the exponential prefactor dominates the long range decay, we expand the near-contact increase in $s_B/d$. Substituting the dimensionless area $z_A = (1 + s_A/d)^2 - 1 / c_A$ for integration in favor of the particle separation $s_A$, the expressions $I_i$, $i = 1, 2$ are of the form

$$I_i = c_A \int_{0}^{1/c_A} e^{-z_A} f_i(c_A z_A) \; dz_A \quad (A12)$$

with

$$f_1(x) = \frac{3(x - 1)\alpha(x)}{2\pi^2 \sqrt{(3 - x)(x + 1)}} \quad (A13)$$

$$f_2(x) = \frac{2}{x - 3} - \frac{2}{x - 1} + \frac{6 \sqrt{x + 1}}{\alpha(x) \sqrt{3 - x}} - \frac{6 \sqrt{3 - x}}{\alpha(x) \sqrt{x + 1}} - 1 \quad (A14)$$

$$\alpha(x) = \pi - 12 \arcsin \frac{\sqrt{x + 1}}{2}$$

The integrals $I_i$ can be treated by expanding the functions $f_i = \sum_{\nu} f_i^{(\nu)} x^\nu$:

$$I_i = \sum_{\nu=0}^{\infty} f_i^{(\nu)} c_A^{\nu+1} \int_{0}^{1/c_A} e^{-z} z^{\nu} \; dz = \sum_{\nu=0}^{\infty} f_i^{(\nu)} c_A^{\nu+1} \int_{0}^{1/c_A} e^{-z} z^{\nu} \; dz. \quad (A15)$$

All incomplete Gamma functions can be eliminated by virtue of the recurrence relation (cf. (6.5.2) and (6.5.22) in [53])

$$\Gamma(\nu + 1, 1/c_A) = \nu \Gamma(\nu, 1/c_A) + (-1)^\nu c_A^{-\nu} e^{-1/c_A}. \quad (A16)$$

As is apparent from the recurrence relation, the result will be of the form

$$I_i = R_i(c_A) + e^{-1/c_A} S_i(c_A). \quad (A17)$$

The regular part, for instance in first order of $s_B/d$,

$$R_1(c_A) = c_A \frac{\sqrt{3}}{2\pi} + c_A^2 \frac{9 - 2\sqrt{3} \pi}{3\pi^2} + c_A^3 \frac{-27 + 2\sqrt{3} \pi}{3\pi^2} + \ldots, \quad (A18)$$
FIG. 19: The radii of convergence $r_k$ for expansions around the jamming point. The contribution blocked B-neighbors give to the pair correlation can be expanded in a series around the jamming point, $c_A = 0$. The radius of convergence is given by the Cauchy-Hadamard formula $r_j = 1/\sqrt[3]{K_j}$ for the term $K_j(s_B/d)^j$. (A) The asymptotic divergence of the $R$-series in (A18) poses no practical problem since few terms (less than 10) give sufficient accuracy. (B) The $S$-series in (A18) converges.

is a series expansion about the point of jamming, $c_A = 0$. It is asymptotically diverging due to the factorial which appears in the recurrence relation. Fortunately this does not restrain us from an excellent approximation, since for the relevant density, $\phi > \phi_o$, the quality of the expansion increases for more than 10 terms in the expansion (cf. panel (a) of the Fig. 19).

The second part in (A17), for which the first order of $s_B/d$ is given by

\[ S_1(c_A) = c_A \sqrt{\frac{3}{2\pi}} + c_A^2 \frac{9 - 2\sqrt{3\pi}}{3\pi^2} + c_A^3 \frac{-27 + 2\sqrt{3\pi}}{3\pi^2} + \ldots , \]  

(A19)

and has a positive radius of convergence (cf. panel (b) in Fig. 19). This part is over-exponentially suppressed by the prefactor $\exp(-1/c_A)$ close to jamming.

In the application to wet granular matter the sub-leading order $(s_B/d)^3 < 4 \cdot 10^{-4}$ is negligible for a realistic value of $s \leq s_{\text{crit}} \approx 0.07d$, whereby we have the concise result

\[ g_B(s_B) = \mathcal{N} e^{-z_B^2} + O(z_B^3) \]  

(A20)

with the abbreviation $z_B = ((1 + s_B/d)^2 - 1) / c_B$ and $1/c_B \approx \phi g_{\text{at}}$. The normalization $8\phi^{\frac{2}{3}} \int g_B \, dz_B = 2$ according the two B-neighbors determines $\mathcal{N}$ in (A20). Hence the result (14).

APPENDIX B: EXPPLICIT EXPRESSIONS FOR THE PAIR CORRELATION IN TWO DIMENSIONS WITHOUT FREE PARAMETERS

In our general derivation of the theory of wet granular matter we distinguished between the jamming density $\phi_J$ and the (highest possible) crystalline packing $\phi_{\text{max}} = \pi/(2\sqrt{3})$ achieved in monodisperse domains. The exact value of the jamming density $\phi_J$ depends on many details such as the distribution of polydispersity and the jamming protocol for the increase of density. When we want to give explicit results without free parameters on the bridge coordination $K(T, \phi, s_{\text{crit}})$ and the equation of state $P = P(T, \phi, s_{\text{crit}})$ we do this for weak polydispersity, where the difference between $\phi_J$ and $\phi_{\text{max}}$ is negligible and the limiting case of 'dry' discs has been studied extensively.

1. High Density

For monodisperse 'dry' discs, $\phi_J = \phi_{\text{max}}$, there are higher order corrections to the free volume result (9) available in the literature which are incorporated in the final results on the bridge coordination and the equation of state for wet granular matter. These corrections are expansions with respect to $x = \phi_J - \phi$ fitted to simulations:

\[ g_{\text{dense}} = \left( \frac{1}{x} + a_0 + a_2 x^2 + \ldots \right) \frac{\phi_J}{\phi} = \left( \frac{1}{x} + a_0 + a_2 x^2 + \ldots \right) \left( 1 + \frac{x}{\phi_J} + \ldots \right), \]  

(B1)
Equation (B1) holds in the dense regime, \( \phi_o < \phi < \phi_{\text{max}} \), above \( \phi_o = 0.71 \). The numerical coefficients are \( a_0 = -1.07 \) and \( a_2 = 5.89 \) [29], confirmed by our own simulations. Similar empirical expressions are also available for polydisperse discs in the glass state (Eq. (6) in [22]).

2. Low and Moderate Density

For the analytic treatment an explicit expression for the contact correlation \( g_c^{\text{at}} \) in Eq. (32) is needed (as the counterpart to the dense expression (B1)). Aside from the trivial one-dimensional case [66], exact expressions for the contact correlation of hard spheres are unknown for the dilute regime. Yet there are well-established approximations in the literature resulting from Scaled Particle theory [54, 55], from the virial expansions [56], as solutions of the Percus-Yevick closure [57], as well as heuristic expressions [58] such as the Carnahan-Starling formula with corrections to better fit simulation results (cf. Tab. I).

As in the dense regime [B1] we shall use the Henderson-Luding expression [29]

\[
\begin{align*}
g_c^{\text{dilute}}(0) &= g_c^{\text{dilute}}(s) = 1 - \frac{7\phi/16}{(1-\phi)^2} - \frac{\phi^3/128}{(1-\phi)^4} \\
g_c^{\text{dense}}(0) &= g_c^{\text{dense}}(s) = 1 - \frac{\phi^2/8}{(1-\phi)^2} - \frac{\phi^4/64}{(1-\phi)^4}
\end{align*}
\] (B2)

for the uncaged regime, \( 0 < \phi < \phi_o \), and the merging function \( m(\phi) = 1/(1 + \exp((\phi - \phi_o)/m_0)) \) with a cross-over width \( m_0 = 0.0111 \) to smoothly connect the dense (B1) and dilute (B2) expressions [29]:

\[
g(s) = m(\phi) g^{\text{dilute}}(s) + (1-m(\phi)) g^{\text{dense}}(s)
\] (B3)

with \( g^{\text{dilute}}(0) = g_c^{\text{dilute}} \) and \( g^{\text{dense}}(0) = g_c^{\text{dense}} \) as given by the Eqs. (B1) and (B2). The near-contact decay has been established in the Eqs. (10) and (14) for \( \phi > \phi_o \), and in Eq. (32) for \( 0 < \phi < \phi_o \):

\[
\begin{align*}
g_c^{\text{dilute}}(s) &= g_c^{\text{at}} \gamma_c^{\text{dilute}} \\
g_c^{\text{dense}}(s) &= g_c^{\text{at}} \gamma_c^{\text{dense}} = g_c^{\text{at}} \gamma_c^{\text{dilute}} \left[ 1 + \left( \phi g_c^{\text{at}} \frac{s}{d} \right)^2 \right]
\end{align*}
\] (B4, B5)

up to leading order in \( s_{\text{crit}} \) with

\[
\gamma_c^{\text{dilute}}(s) = \exp \left( -\phi g_c^{\text{at}} \left[ \left( 1 + \frac{s}{d} \right)^2 - 1 \right] \right)
\] (B6)

With the contact expressions (B1) (B2), as well as the short-range decay formulas (B4) (B5), we have sufficient information on the dry system over the entire density range. We may therefore proceed by introducing the hysteretic capillary bridges.
APPENDIX C: SELF-CONSISTENCY OF BRIDGE COORDINATION $K$

All results presented so far on the coordination $K(\phi, T)$ and pressure $P(\phi, T)$ allowed explicit analytic results. Here we want to demonstrate how to treat more complicated source terms of the hysteretic system (45) numerically. Such an extension of the theory could be motivated as follows. The current of free (unbound) approaching particles could be a function of the free density $n_{\text{free}}$ instead of the mean density, since some of the unconnected neighbors traverse the voids between clusters, so that Eq. (41) is changed to

$$\phi g_{c\rightarrow r}^n + \phi g_{c\rightarrow r}^n / \gamma (s_{\text{crit}}) = (1 - K/K_{\text{sites}}) \phi_{\text{free}} g_{\text{free}}^n (\phi_{\text{free}}). \quad (C1)$$

Obviously this approach is a lower estimate for the current of freely approaching particles, which is why (C1) is considered as a methodical example rather than a physical competitor to the theory presented above.

With the altered Eq. (C1) the hysteretic system (45) can still be solved analytically to find the correlation coefficients $g = \{g_{c\rightarrow r}^b, g_{c\rightarrow r}^b, g_{c\rightarrow r}^b, g_{c\rightarrow r}^b, g_{c\rightarrow r}^b, g_{c\rightarrow r}^b\}$. Unlike before, due to the coupling (C1) and the Eqs. (60)-(62), the correlations $g$ are a highly nonlinear function of $K$. Therefore Eq. (46) becomes a nonlinear self-consistent equation:

$$K(g(K, \phi, T), \phi, T) = K. \quad (C2)$$

The physical value $K(\phi, T)$ of the coordination is the solution $K$ of (C2). The numerical solution of (C2) is found to be very robust, as Fig. 20 indicates. Plugging the resulting self-consistent $K(\phi, T)$ back into the equation for the pressure (74) of wet granular matter, we find that the critical point is shifted from $T_c = 0.273(5)E_{cb}$ to $T_c = 0.216(5)E_{cb}$. This reduction of the critical temperature is intuitively clear since with less particles arriving to form bonds, the wet granular matter `evaporates' at lower granular temperatures.

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The configuration space of the one-dimensional gas is \( L - N_d \), so that the equation of state is \( P(L - N_d) = NT \). Comparison with the general expression \( P = g_{wall}^\text{at} nT \) yields \( g_{wall}^\text{at} = (1 - \phi)^{-1} \).