On the rigidity of a hard sphere glass near random close packing

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Abstract. – We study theoretically and numerically the microscopic cause of the rigidity of hard sphere glasses near their maximum packing. We show that, after coarse-graining over time, the hard sphere interaction can be described by an effective potential which is exactly logarithmic at the random close packing $\phi_c$. This allows to define normal modes, and to apply recent results valid for elastic networks: rigidity is a non-local property of the packing geometry, and is characterized by some length scale $l^*$ which diverges at $\phi_c$ [1, 2]. We compute the scaling of the bulk and shear moduli near $\phi_c$, and speculate on the possible implications of these results for the glass transition.

Hard spheres present a glass phase between $\phi_0$, where the glass transition occurs and structural relaxation becomes unobservable, and $\phi_c$ where the pressure $p$ diverges. In this region this system is solid and resists to shear on any measurable time scales. Although a large amount of works focused on the super-cooled liquid, the glass itself received less attention. In particular, there is no undoubted microscopic theory to explain its mechanical properties and its rigidity. In the cage-escape picture [3], the cage formed by the neighboring particles tighten as $\phi$ increases, and the typical time for a particle to escape its cage grows and eventually diverges. Nevertheless, Maxwell showed that the stability against collective motions of particles is more demanding than against individual particle displacements: in particular $z = d + 1$ inter-particle contacts are sufficient to pin one particle in $d$ dimensions, whereas $z_c = 2d$ contacts in average are required to guarantee mechanical stability [4]. Thus considering a priori rigidity as a local property may be inappropriate.

Recently several works [1, 2, 5, 6, 8, 7] studied the mechanical properties of weakly-connected elastic networks with an average contacts number —the coordination number— $z$ close to the critical value $z_c = 2d$, such as those encountered for athermal repulsive short-range particles above $\phi_c$ [5, 6]. In particular it was shown that (i) these systems present an
excess of vibrational modes at low-frequency in comparison with normal solids [9]. These anomalous modes are characterized by some length \( l^* \sim \delta z^{-1} \) [11, 2], where \( \delta z \equiv z - z_c \), (ii) rigidity can occur only if \( \delta z \geq C_0 \sqrt{p/B} \) on any subsystems of size \( l \geq l^* \), where \( C_0 \) is a constant and \( B \) is the bulk modulus [2]. Thus rigidity is a non-local property of the packing geometry, (iii) the shear modulus \( G \) satisfies \( G/B \sim \delta z \), as observed numerically [6] and confirmed theoretically [3] for repulsive systems.

Can these results apply to hard sphere glasses? On the one hand, hard spheres are weakly-connected at high packing fraction, being exactly \( z = z_c \) at \( \phi_c \) as was shown theoretically [2] [10] [13] [14]. On the other hand, all the results established for elastic networks require a smooth potential to expand the energy and define normal modes. It is in principle problematic in hard sphere systems where the potential is discontinuous. In this Letter we show that, once a coarse-graining in time is made, hard spheres interact with a continuous effective potential, which becomes exactly logarithmic as \( \phi \rightarrow \phi_c \). This allows to define normal modes and to derive new results on the rigidity and the mechanical responses of the glass near \( \phi_c \).

We consider a hard sphere glass, where particles collide elastically, at high packing fractions \( \phi \) close to \( \phi_c \) where structural relaxation is frozen. The particle diameter defines the unit length. Since temperature only rescales the time unit we fix \( \beta = 1 \). Following [13] [15] [16] it is possible to define a contact force network. We introduce an arbitrary time \( t_1 \) much larger than the collision time \( \tau_c \). Two particles are said to be in contact if they collide with each other during a time interval of length \( t_1 \). This allows to define a coordination number \( z \equiv 2N_c/N \), where \( N_c \) is the total number of contacts and \( N \) is the particles number. Then, the contact force \( \vec{f}_{ij} \) between two particles \( i \) and \( j \) is defined as the total momentum they exchange per unit time:

\[
\vec{f}_{ij} = \frac{1}{t_1} \sum_{n=1}^{n_{\text{col}}[t_1]} \Delta \vec{P}_n,
\]

where the sum is made on the total number of collisions \( n_{\text{col}}[t_1] \) between \( i \) and \( j \) that took place in the time interval \( t_1 \), and \( \Delta \vec{P}_n \) is the momentum exchanged at the \( n \)th shock. Fig. 1 shows a two-dimensional example of the contact force network obtained with a polydisperse configuration (1) at packing fraction \( \phi \) close to \( \phi_c \). To obtain high packing fractions numerically we used the 2-dimensional jammed configurations of [12] with packing fraction \( \phi_c = 0.83 \). At \( \phi_c \) the particles are in contact. We reduce the particles diameters by a relative amount \( \epsilon \). This leads to configuration of packing fraction \( \phi = \phi_c (1 - \epsilon)^2 \). Then, we assign a random velocity to every particle and launch an event-driven simulation. Such system is not at thermal equilibrium and displays aging [13]: “earthquakes” can occur which suddenly relax the system and decrease the pressure (2). In between these rare events, there are very long quiet periods where no structural relaxation is observed. All our measures are done during these periods. Note that coordination and contact forces could a priori depend on the arbitrary parameter \( t_1 \).

In the vicinity of \( \phi_c \) no significant dependence of these quantities with \( t_1 \) were observed as long as (i) \( \tau_c \ll t_1 \) and (ii) no earthquake occurs in the time interval \( t_1 \).

As we shall see below, the force networks of dense hard sphere glasses are weakly-connected. To build a correspondence between hard spheres at \( \phi < \phi_c \) and elastic spheres at \( \phi > \phi_c \), we change variables: instead of considering the instantaneous particles positions we consider their time-average position \( \bar{\vec{R}}_i^{(n)} \) over some time scale \( t_1 \gg \tau_c \). To define an effective potential we must relate the contact force \( f_{ij} \) to the average distance between particles \( i \) and \( j \). We

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(1) Half of the particles have a diameter unity, the other half as a diameter 1.4.

(2) Similar earthquakes have been observed in other aging systems such as colloidal pastes, laponite or Lennard-Jones simulations [19]. They correspond to a sudden collective rearrangement of a large number of particles.
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Fig. 1 – Contact forces for $N = 256$, $\epsilon = 10^{-4}$ and $t_1 = 10^5$ collisions. Particles centers are represented by red points and the contact forces by the segments whose width is proportional to the force amplitude. Note that the forces are balanced within our data precision on every particle, as it must be the case on time scales where the structure is stable. For similar force networks see [16].

start by considering the simple example of a line of hard spheres equilibrated at some pressure $p$. The isothermal isobaric partition function is easily computed by introducing the spacings $h_i$ between particles, defined as $h_i = r_{i,i+1} - r_i - r_{i+1}$, where $r_{i,i+1}$ is the distance between particles $i$ and $i + 1$, and $r_i$ is the radius of particle $i$. One obtains:

$$Z \sim \prod_i \int_{h_i=0}^{h_i=\infty} dh_i e^{-\beta p h_i}, \quad (2)$$

where the terms containing the kinetic energy of the particles have been integrated out. If an external force dipole $p_i = -p_{i+1} \equiv p_1$ is now applied on $i$ and $i + 1$, the work required to open the contact $i$ of an amount $h_i$ is now $\left(p + p_1\right) h_i$. Thus we obtain:

$$Z \sim \prod_{j \neq i} \int_{h_j=0}^{h_j=\infty} dh_j e^{-\beta p h_j} \int_{h_i=0}^{h_i=\infty} dh_i e^{-\beta(p+p_1) h_i}. \quad (3)$$

It is then straightforward to compute the average spacing $\langle h_i \rangle = 1/\beta(p+p_1)$. Since the contact force $f_i$ in the contact $i, i + 1$ is $f_i = p_1 + p$, one finds that all contacts satisfy the relation:

$$f_j = \frac{1}{\beta \langle h_j \rangle} \quad \text{for all } j \quad (4)$$

which is thus true whether external forces are present or not.

We now demonstrate that Eq. (4) can be extended to hard sphere glasses at $\phi = \phi_c$ for any spatial dimension. As discussed above, at $\phi_c$, if the “rattlers” are removed (3), the system is marginally connected, or isostatic: $z = z_c$. Isostatic states have the particularity to display as many degrees of freedom of displacements as number of contacts. Hence, (i) the configuration of the system can be defined by the set of distances between particles in contact and (ii) these

(3) At $\phi_c$ some particles ($\approx 5\%$) do not have any contact, and lie at a finite distance of their neighbors. These “rattlers” do not participate to the rigidity of the structure. Near $\phi_c$ these particles can be identified since the distance with their neighbors is much larger than the average. In all our measures we defined “rattlers” as the particles for which their second strongest contact force is smaller than 1% of the system-averaged contact force. We checked that our results, such as the scaling of the coordination and the pressure, are still valid when other definitions of rattlers are used (for example taking a threshold of 5% instead of 1% for the contact force).
degrees of freedom are independent. This implies that the isobaric partition function is a product of terms corresponding each to an individual contact. Consequently, if the system is at equilibrium in a meta-stable state where the contact forces field $\{f\} = \{f_{ij}\}$ is well-defined, the isobaric partition function can be written (4):

$$Z \sim \prod_{(ij)} \int_{h_{ij}=0}^{h_{ij}=\infty} dh_{ij} e^{-\beta f_{ij} h_{ij}}.$$  \hspace{1cm} (5)

Repeating the argument valid for $d = 1$, one obtains that $f_{ij} = \langle h_{ij} \rangle^{-1} \beta^{-1}$. Obviously, as is the case in one dimension, this result is valid with or without external forces. Note that since this derivation only invokes thermodynamic arguments, it also applies to Brownian particles. 

This relation $f/distance$ is checked numerically in Fig.(2-a) near $\phi_c$. The dependence of $f$ with $h$ is found to be in very good agreement with Eq.(4).

When $\phi$ is lowered from $\phi_c$, we shall see that the coordination $z$ increases. Then, the $h_{ij}$ are not independent variables anymore in Eq.(5) and Eq.(4) is invalid. Nevertheless the relative corrections to Eq.(4) are expected to be small, of order $\delta z$ [8]. We check this result in Fig.(2-b), where we compute numerically $C(\delta z) \equiv \langle f_{ij} \beta \langle h_{ij} \rangle \rangle_{ij}^{-1} - 1$, where $\langle \rangle_{ij}$ denotes the average over all contacts. In what follows, we are mainly interested in scaling relations near $\phi_c$, for which corrections of order $\delta z$ are not relevant. We shall neglect them.

![Fig. 2 – (a) – Log-log plot of the contact force amplitude vs. the spacing $h = r_i - r_i - r_j$ for various $\epsilon$ in systems of $N = 256$ particles in two dimensions. Each dot represents the pair of numbers $(f_{ij}, \langle h_{ij} \rangle)$ associated with the contact $ij$. Dots collapse on the dotted theoretical curve defined by Eq. (4). (b) – Average correction $C(\delta z)$ as defined in the text vs. excess coordination $\delta z$ for various $\phi$. The line is a linear fit consistent with the predictions of [8] at small $\delta z$. Corrections are small, of the order of 3 to 4 percent when $\delta z = 1$.](image)

In this approximation, it is straightforward to compute the thermodynamic potential from Eq.(5) (or by integrating Eq.(4)): $G = -\beta^{-1} \sum_{ij} \ln(\langle h_{ij} \rangle)$. A key remark is that this expression of $G$ corresponds precisely to the energy of an athermal assembly of particles of positions $\{\mathbf{R}_i\}$, interacting with a smooth potential $V_{ij}$ of the form:

$$V_{ij}(r) = \infty \quad \text{if} \quad r < r_i + r_j$$
$$V_{ij}(r) = -\beta^{-1} \ln(r - r_i - r_j) \quad \text{if} \quad i \text{ and } j \text{ are in "contact"}$$
$$V_{ij}(r) = 0 \quad \text{if} \quad i \text{ and } j \text{ are not in "contact"}$$  \hspace{1cm} (6)

(4)The upper limits of the integrals of Eq.(5) are not infinite, but bounded by some finite value $h_{max}$ which depends of the contact considered. Nevertheless, as in the one-dimensional case, $h_{max} \sim N/(f \beta)$ [13]. Since the integrals in Eq.(5) converge as soon as $h \gg 1/\beta f$, Eq.(5) becomes exact when $N \gg 1$ for any $\phi$ near $\phi_c$. 
where \( r \approx |\vec{R}_{av}^i - \vec{R}_{av}^j| \) is the average distance between \( i \) and \( j \). This analogy between hard spheres and systems with soft interactions implies that: (i) a configuration minimum of the thermodynamic potential is also a minimum of the energy in the corresponding elastic system, and must therefore satisfy the rigidity criterion evoked in introduction. (ii) when a hard sphere system is sheared (or compressed), the change of thermodynamic potential can be deduced from the shear (bulk) modulus of the elastic system. (iii) Since the effective potential of Eq. (6) is continuous the thermodynamic potential can be expanded around any configuration. This allows to compute the dynamical matrix \( \mathcal{M} \) and the normal modes defined as the eigenvectors of \( \mathcal{M} \).

We now precise (i) to derive a microscopic criterion for the rigidity, or meta-stability, of dense hard sphere glasses. Any meta-stable state must contain at least one configuration corresponding to a minimum of thermodynamic potential. Using (i), this implies that for this configuration \( \delta z > C_0 \sqrt{p/B} \). Anticipating on what follows, we have for hard spheres \( B \sim p^2 \) and \( p \sim (\phi_c - \phi)^{-1} \), therefore there is a constant \( C_1 \) such that:

\[
\delta z \geq C_1 p^{-1/2} \sim (\phi_c - \phi)^{1/2}.
\]

This is our main result, which relates rigidity and microscopic structure.

To test this prediction we study three different systems: the two-dimensional hard sphere glass introduced above, the mono-disperse crystal and the mono-disperse square lattice. We consider all these systems at their maximum packing fraction where particles are in permanent contact, then we reduce the particles diameter by a relative amount \( \epsilon \), and we launch a simulation. In the crystal case, the coordination is 6, therefore \( \delta z = 2 \gg p^{-1/2} \sim \epsilon^{1/2} \) for small \( \epsilon \): condition (7) is satisfied and the system is stable. On the other hand, the square configuration has \( z = 4 \), \( \delta z = 0 \), and the system cannot satisfy (7) without large structural rearrangements for any \( \epsilon \). These predictions are verified numerically. For small \( \epsilon \), the crystal is stable and displays not structural changes, whereas the square lattice collapses rapidly [9, 16], see Fig. 3(a)).

![Fig. 3](image)

Fig. 3 – (a) Collapse of a square lattice. (b) \( D(\omega) \) vs. \( \omega \) for \( \epsilon = 10^{-4} \) in a poly-disperse glass. All frequencies are rescaled by \( \epsilon^{-1} \). The particle positions were averaged over a time \( t_1 = 4 \times 10^5 \) to obtain \( \langle \vec{R}_{av} \rangle \). Eq. (6) was used to compute the dynamical matrix, from which the normal modes frequencies were inferred.

As discussed above, the poly-disperse glass we obtain near \( \phi_c \) presents long periods of stability, where no relaxation occurs. To check Eq. (7) we computed numerically both the coordination and the pressure for various packing fractions, and for various stable periods that appear along the aging regime. As shown in Fig. 4, the data are consistent with an equality of the inequality (7). This suggests that a hard sphere glass lies close to marginal stability, as is the case for soft spheres slowly decompressed toward \( \phi_c \).
In Fig. (3-b) we also furnish an example of density of states \( D(\omega) \) for \( \epsilon = 10^{-4} \) computed during such a plateau, when the “rattlers” are removed. Note that \( D(\omega) \) does not vanish as \( \omega \to 0 \), as in any isostatic system [1]. It is the case here, since \( \phi \) is very close to \( \phi_c \), and therefore \( \delta z \approx 0 \). Interestingly no unstable modes are observed at this packing fraction. This is not obvious: in a meta-stable state, the system could lie alternatively in several minima of the thermodynamic potential, since the temperature is non-zero. Condition (7) would then be satisfied, as is the case for each minimum. Nevertheless, after averaging, the position may lie in between several minima, near a saddle-point where unstable modes are present. This situation may well occur at lower \( \phi \). We leave this question and its possible relation with some observed structural relaxation processes [3] for future investigations.

![Log-log plot of \( \delta z \) versus the average contact force \( \langle f \rangle \sim p \). The data were obtained for different \( \epsilon \) and different time periods. The black line corresponds to the equality of the inequality (7).](image)

To compute the scaling of the elastic moduli of the glass near \( \phi_c \) we can use the results valid for repulsive weakly-connected elastic networks. For the bulk modulus one obtains [6, 8]:

\[
B \sim \frac{\delta p}{\delta \phi} \sim (\phi_c - \phi)^{-2}.
\]

as found previously [13, 14]. The same scaling holds for the crystal [20]. As discussed in introduction, in repulsive weakly-connected network the shear modulus does not scale as \( B \), but rather as \( G \sim B \delta z \). Making the assumption that in the glass phase, the system does not depart much from marginal stability, Eq. (7) is an equality, and one obtains the new result:

\[
G \sim p^{3/2} \sim (\phi_c - \phi)^{-3/2}.
\]

To conclude, we showed that an analogy can be made between a hard sphere glass and an elastic system once a coarse graining in time is made, and that the effective potential that describe particle interactions becomes exactly logarithmic at \( \phi_c \), where the pressure diverges. This allows to define normal modes, and to compute the scaling of the elastic moduli near \( \phi_c \). This implies that the rigidity that characterizes the glass phase near maximum packing is related to a non-local microscopic property of the system geometry: the coordination number \( z \) must be bounded below on any subsystems larger than some length \( l^* \sim \delta z^{-1} \) which diverges at \( \phi_c \). Finally, our numeric data suggest that the glass phase is only marginally stable, at least in the vicinity of \( \phi_c \), implying the presence of anomalous modes near zero-frequency.

One may question if these results apply in the vicinity of the glass transition. On the one hand, near \( \phi_0 \) the coordination is rather large, similar to the one of the crystal. One the other
hand, the distribution of contacts stiffness is certainly broader in the glass. This enhances the presence of anomalous modes at low-frequency, since softer contacts affect only weakly the vibrational spectrum \[8\]. Thus, anomalous modes could be an appropriate concept to study how rigidity appears when \(\phi\) increased toward \(\phi_0\). In particular, it has been proposed that the dramatic slow down near \(\phi_0\) corresponds to a transition in the topology of the free-energy landscape \[21, 22\]: at high \(\phi\), the system lies near free-energy minima, and the dynamics is activated. At lower \(\phi\), the system lives near saddle-points, and the dynamic consists in going down the unstable directions of the free-energy. Our work suggests the following hypothesis: these unstable directions correspond to anomalous modes. Since such modes are collective particles motions, they may cause the heterogeneous dynamics \[23\] observed near the glass transition.

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REFERENCES

[1] M. Wyart, S.R. Nagel, T.A. Witten, Euro. Phys. Letters, 72, 486-492, (2005)
[2] M. Wyart, L.E. Silbert, S.R. Nagel, T.A. Witten, Phys. Rev. E 72, 051306 (2005)
[3] Gotze W. and Sjorgen L., Rep. Prog. Phys., 241 (1992)
[4] Maxwell, J.C., Philos. Mag., 27, 294-299 (1864)
[5] C.S. O’Hern, S.A. Langer, A.J. Liu and S.R. Nagel, Phys. Rev. Lett. 88, 075507 (2002).
[6] C.S O’Hern, L.E Silbert, A. J. Liu and S.R. Nagel, Phys. Rev. E, 68, 011306 (2003)
[7] D.A. Head, Phys. rev. E, 72, 021303 (2005)
[8] M. Wyart, Ann. Phys. Fr., Vol. 30, N3 May-June 2005, pp. 1-96, or arXiv cond-mat/0512155
[9] S.Alexander, Phys. Rep., 296, 65 (1998)
[10] A.V. Tkachenko and T.A Witten, Phys. Rev. E 60, 687 (1999); A.V. Tkachenko and T.A Witten, Phys. Rev. E 62, 2510, (2000); D.A. Head, A.V. Tkachenko and T.A Witten, European Physical Journal E, 6 99-105 (2001)
[11] C.F. Moukarzel, Phys. Rev. Lett. 81, 1634 (1998)
[12] J-N Roux, Phys. Rev. E 61, 6802 (2000)
[13] A. Donev, S. Torquato, and F.H. Stillinger, Phys. Rev. E, 71, 011105 (2005)
[14] G. Parisi and F. Zamponi, cond-mat/0506445
[15] A. Ferguson, B. Fisher, B. Chakrabarty, Europhys. Lett., 66, 277 (2004)
[16] A. Donev, S. Torquato, F.H. Stillinger, and R. Connelly, J. Compt. Phys., 197, 139 (2004)
[17] Neil Ashcroft and N. David Mermin, Solid state physics, New York (1976).
[18] C. Brito and M. Wyart, in preparation
[19] W. Kob W, JL. Barrat, F. Sciortino., P. Tartaglia J., Phys. Condensed Matter 12 6385 (2000); A.Duri, P Ballesta, L. Cipelletti, H. Bissig and V. Trappe, Fluctuation and Noise Lett., 5, I-15, (2005); L Buisson, L Bellon and S Ciliberto, J. Phys.: Condens. Matter 15 S1163S1179 (2003)
[20] Daan Frenkel and Tony Ladd, Phys. Rev. Lett. 59, 1169 (1987)
[21] T.S. Grigera, A. Cavagna, I. Giardina, G.Parisi, Phys. Rev. Lett., 88, 055502 (2002)
[22] J. Kurchan and L. Laloux, J. Phys. A: Math Gen. A, 40, 1045, (1989)
[23] see e.g. M. D. Ediger, Ann. Rev. Phys. Chem. 51, 99 (2000); E. Weeks and D. A. Weitz, Chemical Physics 284 361-367