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## ON STANDARDISED MOMENTS OF FORCE DISTRIBUTION IN SIMPLE LIQUIDS

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## ABSTRACT

The force distribution of a tagged atom in a Lennard-Jones fluid in the canonical ensemble is studied with a focus on its dependence on inherent physical parameters: number density (n) and temperature (T). Utilising structural information from molecular dynamics simulations of the Lennard-Jones fluid, explicit analytical expressions for the dependence of standardised force moments on n and T are derived. Leading order behaviour of standardised moments of the force distribution are obtained in the limiting cases of small density (n  $\rightarrow$  0) and low temperature (T  $\rightarrow$  0), while the variations in the standardised moments are probed for general n and T using molecular dynamics simulations. Clustering effects are seen in molecular dynamics simulations and their effect on these standardised moments is discussed.

## **I INTRODUCTION:**

Understanding the moments and measures of a distribution for a fully atomistic molecular dynamics (MD) simulation allow us to better fit coarser models that reproduce these 1–3. It is often the case in model coarse graining that we wish to directly reconcile the energy landscape of the fully atomistic system to a more basic representation that allows us to maintain as many physical properties of the system of interest, with as little computational cost as possible4. Though, it is also natural to match forces between the high and low resolution systems in an effort to reproduce the force distribution which will inherently give rise to the energy landscape5-10. Let F = [F1,F2,F3] denote a force on a tagged atom in a liquid. Depending on the relative positions of other atoms, force F can vary over a range of values and a detailed information on F can be obtained by calculating properties of its equilibrium distribution, which we will call force distribution in this manuscript.

Considering an isotropic system, the equilibrium distribution of each force coordinate is the same. We define the standardised moment of the force distribution by averaging over the k-th power of its first coordinate as

$$\alpha_k = \frac{\left\langle F_1^k \right\rangle}{\left\langle F_1^2 \right\rangle^{k/2}},\tag{1}$$

where F k 1 is the k-th moment of the force distribution and ak standardises the k-th moment by scaling it with the k-th power of the standard deviation of the force distribution. simple In а homogeneous radially fluid with symmetric interactions between particles, force distribution will exhibit the symmetry around the origin and thus all odd standardised moments vanish, i.e. 0 = $\alpha 1 = \alpha 3 = \alpha 5 = \dots$  As  $\alpha 2 \equiv 1$  by definition (1), the first non-trivial standardised moment is kurtosis, denoted  $\alpha 4$ , which provides a measure of spread that details how tailed the force distribution is relative to a normal distribution11. In this paper, we study how the force distribution



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depends on the number density of a homogeneous many-body system, and the temperature of the same system in a canonical ensemble. We will do this by studying the behaviour of the second moment of the force distribution F 2 1 and standardised even moments  $\alpha 4$ ,  $\alpha 6$ ,  $\alpha 8$ , ..... If the force distribution was Gaussian, then the even standardised moments would be

$$\alpha_k = (k-1)!! = \prod_{i=1}^{k/2} (2i-1), \text{ for } k = 2, 4, 6, 8, 10, \dots, (2)$$

and the second moment F 2 1 would be sufficient to parametrize the force distribution. However, the force distributions in simple liquids have been reported to deviate from Gaussian distribution12–14. In particular, by comparing the results of our analysis with Gaussian moments in equation (2), we can also quantify how non-Gaussian the real force distribution is.

Much work has been done in the area of force distributions of many-body systems: with seminal work from Chandrasekhar15 that employed Markov's theory of random flights to give an expression for the force distribution of a many-body system interacting through a 1/r gravitational potential. More recent work has been done with the help of MD by Gabrielli et al16, who derived an expression for the kurtosis of the force distribution for a lattice system of atoms interacting through the gravitational potential. Further, using the classical density functional theory, an expression for the probability distribution of force for a system interacting through an arbitrary weakly repulsive potential was derived by Rickayzen et al17,18.

In this paper, we study the number density and temperature dependence of the force distribution for a many-body system interacting through a Lennard-Jones 12-6 potential19,20, which is ubiquitously used and has been shown to model homogeneous systems of interacting (Argon) atoms well21–23.

In Section III, an in depth investigation is given to the simple two-body system in one spatial dimension, which provides the ideal platform to illustrate the underlying methods while retaining interesting dynamical behaviour. From first principles we derive first-order partial differential equations (PDEs) describing the dependence of the standardised moments of the force distribution has on parameters. In doing so we further derive an analytic expression for the partition function of a two-body system that depends solely on the standardised moments of the force distribution whereupon the expression is exact in an asymptotic limit of the density going to zero (n  $\rightarrow$  0). Similarly, an expression is derived relating the average energy of the system to standardised moments of force from the temperature dependent PDE. In parameter regimes where long-range forces between atoms dominate, we use a truncated Taylor series expansion to derive the leading order behaviour of the kurtosis of the force distribution in the limit  $n \rightarrow 0$ . Finally, we utilise a Laplace integral approximation to ascertain the leading order behaviour of the standardised moments of force at low temperatures  $(T \rightarrow 0)$ . Results from simple MD simulation are presented to provide evidence for the efficacy of these methods and underlying assumptions. This is followed by Section IV, where the natural idea that long range force calculations dictate asymptotic behaviour is extended from the 1D model to many-body systems of arbitrary size in three spatial dimensions. These systems exhibit the physical properties of standard MD simulations: i.e. cubic geometry with periodic boundary conditions that employ minimum image convention. In the



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particular, we can analyze the system by performing calculations on a central cubic cell. In Section IV B, MD results are displayed for many-body systems. We present the dependence of the standardised force moments on density, n, and temperature, T, and discuss the parameters and integrator schemes utilised in producing the results of MD simulations.

## **II. NOTATION**

We consider a system of N identical atoms interacting via the Lennard-Jones 12-6 potential19. This is a ubiquitous interatomic pairwise potential; here the potential between atoms labelled i, j =1,2,...,N positioned at qi ,qj  $\in$  R 3 is given (in reduced units24) by the expression

$$U_{ij}(r_{ij}) = 4\left(\frac{1}{r_{ij}^{12}} - \frac{1}{r_{ij}^6}\right),\tag{3}$$

where ri j = qi -qj is the distance between atoms. The Lennard-Jones potential (3) between two atoms has a unique minima obtained at  $r_{ij} = r_* = 2^{1/6}$ .

We employ the framework of statistical mechanics for this closed many-body system and describe atom i = 1,2,...,N by phase space coordinates {qi ,pi}  $\in \mathbb{R}$  6 , were pi denotes the momentum of the i-th atom. We work in the canonical ensemble with temperature T; the partition function therefore becomes

$$\mathscr{Z}_{N}(T,V) = \frac{1}{h^{3N}N!} \iint_{\Omega_{\mathbf{q}} \times \Omega_{\mathbf{p}}} \exp[-\beta H(\mathbf{q},\mathbf{p})] \,\mathrm{d}^{3}\mathbf{q} \,\mathrm{d}^{3}\mathbf{p} \,,$$

where V is the volume of our closed system, and q = (q1,q2,...,qN) T and p = (p1,p2,...,pN) T are vectors containing the positions and momenta of all atoms in the system. Our integration domain is given by  $\Omega q \times \Omega p \subset R \ 3N \times R \ 3N$ . This denotes the phase space of our system. For systems of interest  $\Omega p \equiv R$  3N. The underlying geometry of the system (and principle simulation cell) is a cubic box of size L > 0, therefore  $\Omega q \equiv (-L/2, L/2] \times \cdots \times$ (-L/2, L/2]. The phase space volume elements in equation (4) are denoted by

$$d^{3}\mathbf{q} = \prod_{i=1}^{N} d^{3}\mathbf{q}_{i} \quad \text{and} \quad d^{3}\mathbf{p} = \prod_{i=1}^{N} d^{3}\mathbf{p}_{i}. \quad (5)$$

Throughout this work we make use of reduced units24, utilising Argon parameters25. In particular, all instances of T in this work can be translated back to SI units with the transformation  $T \rightarrow kBT$ where kB is the Boltzmann factor. Therefore, in the partition function (4), we have  $\beta = 1/T$  and h is the Planck constant ( $\approx 0.186$  in reduced units). Finally, H(q,p) is the classical Hamiltonian H(q,p) = K(p)+U(q) with kinetic energy K(p) = |p| 2/2(where the usual factor of mass is unity under reduced units) and a general potential U(q). The statistical average of a quantity X for this N-body system is given by

$$\langle X \rangle = \frac{1}{\mathscr{Z}_N h^{3N} N!} \iint_{\Omega_{\mathbf{q}} \times \Omega_{\mathbf{p}}} X \exp[-\beta H(\mathbf{q}, \mathbf{p})] \, \mathrm{d}^3 \mathbf{q} \, \mathrm{d}^3 \mathbf{p}, \quad (6)$$

where the Boltzmann factor acts as a statistical weighting for a configuration  $\{q,p\} \in \mathbb{R}$  6N, normalised such that h1i = 1. We label atoms so that the first one is the tagged atom. Denoting the force on the tagged atom produced from the j-th atom by Fj = [Fj,1,Fj,2,Fj,3]  $\in \mathbb{R}$  3, for j = 2,3,...,N, the total force F = [F1,F2,F3] on the tagged atom is

$$\mathbf{F} = \sum_{j=2}^{N} \mathbf{F}_j.$$

We define



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$$f_k = \int_{\Omega_{\mathbf{q}}} \left( \sum_{j=2}^N F_{j,1} \right)^k \exp[-\beta U(\mathbf{q})] \, \mathrm{d}^3 \mathbf{q}$$
(7)

for  $k = 0, 1, 2, \dots$  Then we have

$$\frac{f_k}{f_0} = \left\langle \left(\sum_{j=2}^N F_{j,1}\right)^k \right\rangle = \langle F_1^k \rangle.$$

Then the k-th standardised moment (1) is given by

$$\alpha_k = \frac{f_0^{k/2-1} f_k}{f_2^{k/2}},\tag{8}$$

where we are interested in cases k = 4,6,8,...

In order to study how the force distribution depends on the physical parameters of interest it is useful to identify how changes these parameters will manifest in themselves in the system. Indeed, we choose to work in the canonical ensemble with a target temperature of T: this is accomplished with the use of a thermostat which is discussed further in Section IV B and Appendix B. It is more illuminating to see that if we have a system with a fixed number of free interacting atoms N in a cubic box of side L; the (reduced) number density is given by n = N/L 3. Therefore the approach we employ in this paper to ascertain how values of standardised moments depend on number density, will be to keep the number of atoms fixed but vary the box width L - this will manifest as a change in density n. Similarly one could keep the volume of the cubic box the same and vary the number of atoms though this is a point of discussion in Section IV B.

For the remainder of the paper we will study systems with different spatial dimensions. The size of the system varies by changing the number of particles N; we will use equation (8) as a crucial initial point in each calculation. We will naturally proceed by investigating systems of increasing complexity; starting from a cartoon one-dimensional model and culminating to a general many-body system of arbitrary size in three spatial dimensions.

# III. ONE ATOM IN A POTENTIAL WELL

We now go on to illustrate three approaches to obtain the dependence of the force distribution on parameters n and T. It is useful to note that, as we are now working in one spatial dimension, density n is proportional to 1/L, i.e. we have n  $\propto$ 1/L. We will consider a simple system in one spatial dimension consisting of two atoms interacting through the Lennard-Jones potential (3) in interval [0,L] with periodic boundary conditions. One of the atoms is considered to be fixed at position  $q0 = L/2 \in [0,L]$  and the other atom is free to move, therefore, we have N = 1 free atom. Its position is denoted  $x \in [0,L]$ . Therefore, the inter-atomic distance is r =|x - q0|. Using our simplified onedimensional set up, F1 = F and  $\Omega q = (0,L)$ , equation (7) reduces to

$$f_k(L) = \int_0^L F^k(|x - q_0|) \exp[-\beta U(|x - q_0|)] \,\mathrm{d}x, \qquad (9)$$

which is the marginalised expected value of the k-th moment of force F(x) = -dU/dx, where we have dropped subscripts in the Lennard-Jones potential (3) and we write it as U(z) = 4(z - 12 - z - 6). Utilising the symmetry of the potential (and therefore the force) we are left with

$$f_k(L) = 2 \int_{0}^{L/2} F^k(r) \exp[-\beta U(r)] \,\mathrm{d}r.$$
 (10)



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In what follows, we will assume that we are in a regime where the box width L satisfies  $L \gg r^*$ , where  $r^* = 2 \frac{1}{6}$  minimizes the Lennard-Jones potential U.

A. Differential equation for standardised moments

We consider a perturbation of the form L  $\rightarrow$  L+ $\delta$ L. Using equation (10) and considering terms to the order O( $\delta$ L), we obtain

$$f_k(L+\delta L) = f_k(L) + f'_k(L) \,\delta L + O(\delta L^2)$$
  
=  $f_k(L) + F^k(L/2) \exp[-\beta U(L/2)] \,\delta L + O(\delta L^2).$ 

Using equation (8), we approximate  $\alpha k(L+\delta L)$ 

$$\alpha_k(L) + \alpha_k(L) \upsilon_k(L) \exp[-\beta U(L/2)] \delta L + O(\delta L^2),$$

where our notation  $\alpha k(L)$  highlights the dependence of the standardised moments of force,  $\alpha k$ , on L, and function  $\nu k(L)$  is given

$$\upsilon_k(L) = \frac{k-2}{2f_0(L)} + \frac{F^k(L/2)}{f_k(L)} - \frac{kF^2(L/2)}{2f_2(L)}.$$
 (11)

Taking the limit  $\delta L \rightarrow 0$ , we obtain the derivative of the k-th standardised moment of force, with respect to L,

$$\frac{\partial \alpha_k}{\partial L}(L) = v_k(L) \exp[-\beta U(L/2)] \alpha_k(L), \qquad (12)$$

where vk(L) are expressed in terms of integrals (10) as given by equation (11).

## B. Far-field integral approximation

To further analyze integrals (10), we introduce a cutoff c, which satisfies that r\* < c < L/2, where r\* = 2 1/6 is a unique maximum of exp[- $\beta$  U(z)], which can be Taylor expanded as  $\beta(1+4z -6 +4z -12)$ -16/3z -18 +8z -24 ...). Considering sufficiently large L, we can choose the cutoff c, so that

$$\left| f_0(L) - 2 \left( \int_0^c \exp\left[-\beta U(r)\right] \mathrm{d}r + \beta \int_c^{L/2} 1 + \frac{4}{r^6} \mathrm{d}r \right) \right| \le \varepsilon, \quad (13)$$

where tolerance  $\varepsilon$  is chosen to be 10–4 in our illustrative computations. This splitting allows us to numerically calculate the bulk of the integral (10) as a constant independent of L and then use the second term to give an analytic expression for  $\alpha$ k with dependence on L, and ultimately on n.

The range of values of T that are of typical use are chosen in order to maintain the liquid state of Argon during simulation. These are approximately temperatures in the interval 0.70 < T < 0.73 under ambient conditions26. Therefore, as volume is varied we are in a regime where  $\beta = O(1)$ , for convenience we set  $\beta = 1$ . Though given that the density of our system changes between each simulation some systems will be in a liquid phase and others in a gaseous phase, this is a point of discussion in Section IV B.

Splitting the integration domain [0,L/2] of integral (10) into [0, c] and [c,L/2], we use the exact form of the integrand in [0, c] to obtain a 'near-field' contribution. Utilising an approximate form for the integrand given by the truncated Taylor expansion f(z) in the domain [c,L/2] gives rise to a density dependent 'far-field' contribution. Combining these we arrive at the approximate form for f0(L). Using cutoff c = 2, equation (13) is satisfied with  $\varepsilon =$ 10-4. Therefore, upon numerically calculating the bulk contribution for the integral with domain [0,2], we get

$$f_0(L) = 2 \int_0^{L/2} \exp[-\beta U(r)] dr = b_0 + L + O\left(L^{-6}\right) \quad (14)$$

with b0 = -0.71832, which depends on our choice of cutoff c = 2. Similarly, we can



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calculate far-field integral approximations of integrals (10) for general values of k = 2,4,6,8,10,12. The integrand F k (r) exp[ $-\beta$ U(r)] has maxima when r = r\* = 2 1/6 or when kU''(r) =  $\beta$  (U ' (r))2. This forms a cubic in r 6 that can be solved. For the values of k used in this work, this sometimes results in a global maximum, that always lies at a distance less than r < r\* from the origin. Therefore r\* = 2 1/6 is the furthest maximum of the integrand from the origin.

Splitting integral (10) into a near-field and far-field contribution, using the general cutoff c = 2, we find

$$f_k(L) = b_k + O(L^{-7k})$$
 (15)

The near-field contributions, bk , generally increase vastly if we increase the value of k, for example

$$b_0 = -0.71832$$
,  $b_2 = 130.64$  and  $b_4 = 2.5727 \times 10^5$ ,

while the dependence on L decreases more rapidly for larger values of k. Therefore, the non-negligible density contributions to  $\alpha k(L)$  in the low density limit come exclusively from the normalisation f0(L) given by (14).

Substituting equations (14) and (15) in equation (8), we obtain an expression for the general k-th standardised moment of force

$$\alpha_k(L) = \frac{b_0^{k/2-1}b_k}{b_2^{k/2}} \left(1 + \frac{L}{b_0} + O\left(L^{-6}\right)\right)^{k/2-1}.$$
 (17)

Using the values of b0, b2 and b4 given by (16), we obtain the dependence of the kurtosis of the force distribution on the reduced number density n = 1/L in the dilute limit  $n \rightarrow 0$  as  $\alpha 4 = -10.828+15.074n -1 +O n 6$ . Figure 1 compares this result with the results obtained by MD simulation of the one

atom system. We observe that MD is in good agreement with the results obtained by formula (17).

C. Leading order behaviour for differential equation (12)

Since  $L/2 > r^*$ , the force F(L/2)monotonically decreases as a function of L. When looking at leading order approximations in the low density limit n  $\rightarrow 0$  (equivalent to limit  $L \rightarrow \infty$ ) to equation (12), we need to analyse vk(L). The second and third term in equation (11) converge to zero more rapidly than the first term as  $L \rightarrow \infty$ , therefore the leading order behaviour is given by the first term.

$$v_k(L) \sim \frac{k-2}{2f_0(L)}$$
 as  $L \to \infty$ . (18)

By utilising the far field integral approximation (14), we arrive at  $fO(L) \sim$ (b0 +L), where b0 = b0(c) is a constant term that depends on cutoff parameter c. With this, our leading order approximation of the k-th standardised moment,  $\alpha \ 0 \ k$ , obey

$$\frac{\partial \, \alpha_k^0}{\partial L}(L) = \frac{k-2}{2 \left( b_0 + L \right)} \, \alpha_k^0(L) \, .$$



FIG. 1. Plot of  $\alpha 4$  as a function of n = 1/L for the illustrative one-atom system. Results of MD simulations are compared with  $\alpha 4 = -10.828 + 15.074n - 1$  obtained by using equation (17) with b0, b2 and b4



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given by (16) (blue dashed line). MD simulation results for temperature T = 1 utilising Langevin dynamics27 described in equation (B1), with friction parameter  $\gamma$ = 0.1, are represented by red dots. The MD simulation length was a total of 1.1×108 time steps with the first 107 time steps used for initialisation.

Finally this gives us tha

$$\mathscr{Z}_1(T,V) = \frac{1}{h} \int_0^L \exp[-\beta U(q)] \, \mathrm{d}q \int_{-\infty}^\infty \exp\left[-\frac{\beta p^2}{2}\right] \, \mathrm{d}p \,, \quad (20)$$

where the Planck factor of 1/h arises instead of 1/h 3 due to the fact that we are in one-dimensional physical space. Using (10), we obtain

$$f_0(L) = h \mathscr{Z}_1(T, V) \sqrt{\frac{\beta}{2\pi}}.$$
(21)

Considering the low density limit  $n \rightarrow 0$ (i.e.  $L \rightarrow \infty$ ) in equation (12) and using (18) and (21), we obtain

$$\mathscr{Z}_{1}(T,V) \sim \frac{(k-2)\sqrt{2\pi}}{\sqrt{h^{2}\beta}} \left[ \alpha_{k}(L) \left( \frac{\partial \alpha_{k}}{\partial L}(L) \right)^{-1} \right],$$
 (22)

as  $L \rightarrow \infty$ . In particular, we can obtain the partition function (20) in the dilute (low density) limit by using information



FIG. 2. Approximation of the partition function Z1(T,V) obtained using the right hand side of equation (22) with k = 4 and values of kurtosis ( $\alpha$ 4) estimated from MD simulation (blue dashed line). The exact

values obtained by (20) are plotted as the red dots.

of force about the moments the distribution. The accuracy of equation (22) is illustrated in Figure 2, where we use k =4. We use MD simulations of a single atom, using a range of simulation box widths L. We estimate the values of kurtosis of the force distribution, its derivative with respect of L and use the right hand side of equation (22) to estimate the Z1(T,V). Considering  $L \ge 10$ , the result is within 5% error when compared with the exact result (20), while for larger values of box width L the error decreases to around 1%, confirming that the formula (22) is valid in the asymptotic limit  $L \rightarrow$ **%**.

D. Temperature dependence of standardised moments

One can perform a similar analysis as in Section III A, viewing the moments  $\alpha k = \alpha k(T)$  as a function of temperature  $T = 1/\beta$ . To do that, we consider the moment definition (10) as a function of temperature T, namely, we define

$$f_k(T) = 2 \int_{0}^{L/2} F^k(r) \exp\left[-\frac{U(r)}{T}\right] dr.$$
 (23)

Considering small perturbations of these functions with respect to  $T \rightarrow T + \delta T$ , while fixing the domain length L, and collecting terms up to first order in  $\delta T$ , we obtain

$$\frac{\partial \alpha_k}{\partial T}(T) = \nu_k(T) \,\alpha_k(T), \qquad (24)$$

Where

$$\mathbf{v}_k(T) = \left(\frac{k}{2} - 1\right) \frac{f_0'(T)}{f_0(T)} + \frac{f_k'(T)}{f_k(T)} - \left(\frac{k}{2}\right) \frac{f_2'(T)}{f_2(T)}.$$
 (25)

Combining equations (24) and (25) with equation (21) where  $\beta = 1/T$ , we obtain



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$$\frac{\partial}{\partial T} \ln\left(\frac{\alpha_k^2 f_2^k}{f_k^2}\right) = (k-2) \left(\frac{\partial}{\partial T} \ln(\mathscr{Z}_1) - \frac{1}{2T}\right).$$

Since  $-\partial/\partial\beta(\ln Z1)$  is equal to the average energy of the system, hEi, we have

$$\langle E \rangle = \frac{T}{2} + \frac{T^2}{k - 2} \frac{\partial}{\partial T} \ln \left( \frac{\alpha_k^2 f_2^k}{f_k^2} \right), \tag{26}$$

where the first term on the right hand side of equation (26) is the average kinetic energy of our one-atom system. Substituting equation (8) into the second term on the right hand side, it can be rewritten as T  $2\partial$  (ln f0)/ $\partial$ T. Thus, using equation (6), we confirm that the second term on the right hand side of equation (26) is the average potential energy.

#### E. Low temperature limit

Next, we consider the behaviour of the kth standardised moment of force,  $\alpha k(T)$ , given by equation (8), in the low temperature limit,  $T \rightarrow 0$ , which is equivalent to the limit  $\beta \rightarrow \infty$ . Since the inter-atomic potential U(r) has a global minimum at r = r\* in interval [0,L/2], integrals of the form (10) and (23) can be approximated by Laplace's method in the limit  $\beta \to \infty$  and T  $\to 0$ , respectively. A general discussion of Laplace's method is given in Chapter 6 of the book by Bender and Orszag28. We calculate the asymptotic expansion of f0(T) by applying Laplace's method to integral (23) for k = 0. We approximate the integration limits of integral (23) to lie within the domain  $r \in$  $(r* - \varepsilon, r* + \varepsilon)$ , where  $\varepsilon \ll 1$ , and we Taylor expand U(r) at r = r\*. Using U ' (r\*) = 0, we have

$$\begin{split} U(r) &\approx U(r_*) + (r-r_*)^2 U''(r_*)/2 \\ &+ (r-r_*)^3 U^{(3)}(r_*)/6 + (r-r_*)^4 U^{(4)}(r_*)/24 \end{split}$$

where we denote the m th derivative of U as U (m) for  $m \ge 3$ . Substituting into

integral (23), we arrive at the asymptotic expansion

$$f_0(T) \sim \frac{\sqrt{\pi T} \exp[-U(r_*)/T]}{\sqrt{2U''(r_*)}} \Big[ 1 + B_0 T + O(T^2) \Big], \quad (27)$$

as  $T \rightarrow 0$ , where constant B0 is given by 28

$$B_0 = \frac{5 \left( U^{(3)}(r_*) \right)^2}{24 \left( U''(r_*) \right)^3} - \frac{U^{(4)}(r_*)}{8 \left( U''(r_*) \right)^2}.$$
 (28)

To apply Laplace's method to integral (23) for k = 2,4,6,..., we note that F k (r) = (U ' (r))k for even values of k. Using the truncated Taylor expansion around r = r\* and noting that U ' (r\*) = 0, we have

$$F^{k}(r) \approx (r - r_{*})^{k} \left( \left( U''(r_{*}) \right)^{k} + (r - r_{*}) C_{k,1} + (r - r_{*})^{2} C_{k,2} \right),$$
(29)

where Ck,1 and Ck,2 are constants, which can be expressed in terms of the derivatives of potential U(r) at r = r\* (see equations (A1) and (A2) in Appendix A). This gives the asymptotic expansion

$$f_k(T) \sim \frac{\sqrt{\pi T} \exp[-U(r_*)/T]}{\sqrt{2U''(r_*)}} \times A_k \left[ T^{k/2} + B_k T^{k/2+1} + O\left(T^{k/2+2}\right) \right].$$
(30)

as  $T \rightarrow 0$ , where constants Ak and Bk are given by

$$A_k = \left( U''(r_*) \right)^{k/2} (k-1)!!$$

And

$$B_k = \frac{(4k-15)(k^2-1)(U^{(3)}(r_*))^2}{72(U''(r_*))^3} + \frac{(k^2-1)U^{(4)}(r_*)}{8(U''(r_*))^2},$$

where the last formula reduces to equation (28) for k = 0. Substituting (27) and (30) into (8) gives the following expression in the limit  $T \rightarrow 0$ :

$$\alpha_k \sim (k-1)!! \left(1 + \frac{(k-2)B_0 + 2B_k - kB_2}{2}T + O(T^2)\right).$$

In particular, we have  $\alpha 2 \sim 1+OT2$  and



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$$\alpha_4 \sim 3 + 3 \left( \frac{(U^{(3)}(r_*))^2}{(U''(r_*))^3} + \frac{U^{(4)}(r_*)}{(U''(r_*))^2} \right) T + O(T^2)$$
  
=  $3 + \frac{203}{6}T + O(T^2).$  (31)

31) Therefore, Laplace's method predicts that the standardised moments of the force distribution,  $\alpha k(T)$ , tend to the values given in equation (2) for Gaussian moments in the low temperature limit. This limiting behaviour is to be expected as during the Laplace approximation we use a Gaussian distribution to approximate the Boltzmann factor. We can interpret this approach as approximating the force distribution as Gaussian and perturbations of the system around small temperatures give rise to non-Gaussian contributions to the standardised moments. Results from MD simulation are illustrated in Figure 3 over the range of values of temperature T. We see that the behaviour of kurtosis,  $\alpha 4$ , is well approximated by the linear approximation 3+203 T/6 given in equation (31) for the temperature values satisfying  $T \leq 0.1$ , though this agreement diverges as temperature T increases and higher order terms, O T 2 in equation (31), become significant. In Figure 3, we fix the box width as L = 10. Increasing the box width much further would take us to a regime where the particle is essentially free and the approximation calculated by the Laplace method around the potential minimum would lose validity.

## **IV. MANY-BODY S**

TEMS In this section we employ the far field approximation approach introduced in Section III B and we will vary the number density of the system by changing the size L of the integration domain, which will be given as the three-dimensional



20 FIG. 3. Kurtosis,  $\alpha 4$ , as a function of temperature, T, for T  $\leq$  0.3. The linear behaviour is estimated as  $\alpha 4(T) \sim$  2.9388+37.002T for T  $\in$  (0.01,0.10) (using the MD computed data, with density n = 0.1, visualized as red dots). We compare this to the theoretical linear result 3+203T/6 predicted by equation (31) (illustrated by the blue dashed line.

cube [0,L] 3. Using notation introduced in Section II, the distance between atoms labelled i, j = 1,2,...,N positioned at qi ,qj  $\in$ R 3 is denoted by ri j = qi -qj. Taking into ccount the periodic boundary conditions, the distance qi -qj is the minimum image inter-atomic distance given by

$$|\mathbf{q}_i - \mathbf{q}_j| = \left(\overline{(q_i^x - q_j^x)}^2 + \overline{(q_i^y - q_j^y)}^2 + \overline{(q_i^z - q_j^z)}^2\right)^{1/2}, \quad (32)$$

(32) where the overline denotes  $\zeta = \zeta$ -L[ $\zeta$ /L] for  $\zeta \in R$  and [.] rounds a real number to the nearest integer. For an interacting N-body system the dimensionality of the integral given by equation (7) is 3N. We first present an illustrative calculation with N = 2 interacting atoms in Section IV A and then we study systems with larger values of N in Section IV B.

A. Dependence of  $\alpha k$  on density for N = 2 interacting atoms

In Section III, we have considered two atoms in the onedimensional spatial



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domain, where one atom was fixed at position q0, i.e. we have effectively studied a single atom in a one-dimensional potential well. Here, we will consider N = interacting atoms 2 in the threedimensional cubic domain [0,L] 3 with periodic boundary conditions. We calculate the k-th standardised moment of force according to equation (8). To do so, we consider equation (7), where we have d 3q = d 3q1 d 3q2, U(q) = U(r12), F1(q) =F1(r12) and we integrate over the domain  $\Omega = [0,L] \ 3 \times [0,L] \ 3$  to get

$$f_k = \int_{\Omega} F_1^k(r_{12}) \exp[-\beta U(r_{12})] \,\mathrm{d}^3 \mathbf{q}_1 \,\mathrm{d}^3 \mathbf{q}_2. \tag{33}$$

(33) It is useful to introduce a change of coordinates  $\xi \ell = q \ell 1 - q \ell 2$  and  $\eta \ell = q \ell 1 + q \ell 2$  for  $\ell = x$ , y,z. We note that r12 is only dependent on the  $\xi \ell$  variables, therefore one can trivially integrate (33) through the  $\eta \ell$  variables as the integrand has no dependence on these to obtain

$$f_k = \frac{L^3}{8} \int_{-L-L-L}^{L} \int_{-L}^{L} F_1^k(r_{12}) \exp[-\beta U(r_{12})] \,\mathrm{d}\xi^x \,\mathrm{d}\xi^y \,\mathrm{d}\xi^z,$$

where r12 is the minimum image interatomic distance (32). This integral can be written in terms of standard Euclidean distance r 2 =  $(\xi x) 2 + (\xi y) 2 + (\xi z) 2$ as

$$f_k = 8L^3 \int_0^{L/2} \int_0^{L/2} \int_0^{L/2} F_1^k(r) \exp[-\beta U(r)] \,\mathrm{d}\boldsymbol{\xi}\,, \qquad (34)$$

where  $d\xi = d\xi \ x \ d\xi \ y \ d\xi \ z$ . In order to analyse fk further by implementing a far field approximation, we need to make sure we are in a regime where the integrand is small - we do this by introducing a cutoff  $\gamma$ , which will divide the cube [0,L/2] 3 into 8 cuboid subdomains, including

$$egin{aligned} \Omega_1 &= [0,\gamma]^3, & \Omega_2 &= [0,\gamma]^2 imes [\gamma,L/2], \ \Omega_3 &= [0,\gamma] imes [\gamma,L/2]^2, & \Omega_4 &= [\gamma,L/2]^3. \end{aligned}$$

Utilising the symmetry of the problem, we can rewrite integral (34) as

$$f_k = 8L^3 \left( \int_{\Omega_1} + 3\int_{\Omega_2} + 3\int_{\Omega_3} + \int_{\Omega_4} \right) F_1^k(r) \exp[-\beta U(r)] \,\mathrm{d}\boldsymbol{\xi} \,.$$
(35)

Considering (35) for k = 0, the integral over  $\Omega 1$  is independent of L and provides a bulk contribution to f0 that will depend on  $\gamma$ . The remaining three terms have integration domains that allow the integrand to be accurately described by a Taylor expansion giving the leading order contribution in the asymptotic limit  $L \rightarrow \infty$ as f0  $\propto$  L 6, which can be rewritten in terms of the density, n, in the form

$$f_0 \propto n^{-2}$$
 as  $n \to 0$ . (36)

Considering fk for k 6=0, the integral over  $\Omega 1$  in equation (35) is again independent of L. However in the far field expansion the integrals over  $\Omega 2$ ,  $\Omega 3$  and  $\Omega 4$  all decay with L due to the force factor. As the integration domain has essentially been transformed into that of inter-atomic distances about the three coordinates, when we increase the domain length, the interatomic force necessarily decays to 0. Therefore in the limit  $L \rightarrow \infty$  the dominant term arises from integrating over  $\Omega 1$ , and we see that, for k = 2,4,6,8,...,

$$f_k \propto n^{-1}$$
 as  $n \to 0$ . (37)

This leaves us with the final result that in the low density limit  $n \rightarrow 0$ , combining equation (8) with asymptotic expressions (36) and (37),

$$\alpha_k \propto n^{1-k/2}$$
 as  $n \to 0$ . (38)

While this result has been calculated for N = 2 interacting atoms, it is also confirmed for larger values of N by estimating the k-



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th standardised moments using MD simulations, as it is shown in the next section.

N	t <sub>sim</sub>	$L_0$	<i>n</i> <sub>0</sub>
2	10 <sup>9</sup>	5	0.016
8	107	3	1/64
64	10 <sup>6</sup>	5	1/64
512	$10^{4}$	10	1/64

TABLE I. The length of MD simulation, tsim, the (smallest) box width, L0, used for simulations with N atoms and density n0 for MD simulations with varying temperatures.

B.MD simulations with N interacting atoms

In this section we present the results from MD simulations of many-body systems in three spatial dimensions using different values of N, including the case N = 2(analyzed in Section IV A). Atoms are subject to pairwise interactions governed by a Lennard-Jones potential, given in equation (3). For each system we use a velocity-Verlet23 integrator and maintain the system in the canonical ensemble by incorporating Nosé-Hoover а thermostat37, see Appendix B. We perform two types of MD simulation studies: those that are used for studying how the number density, n, of a system affects standardised moments, and those that aim to probe temperature dependency. In all cases we utilise a time step  $\Delta t = 0.01$ . In the case of the simulation with N = 2 atoms, we initialise the positions of atoms by setting q1 = 0 and q2 = (L/2, L/2, L/2), whereas for the N = 8, 64, 512 atom systems, we choose to initialise these on a uniform cubic lattice.

The MD simulation parameters are summarised in Table I, where tsim is the total simulation time used for calculating the required statistics, which is preceded by the initial simulation of length tsim/10 used for equilibrating the system. When investigating the number densitv dependence, we perform 20 simulations each with a box width of  $L = L0 \times (6/5)$ i-1, where  $i = 1, 2, \dots, 20$  labels the simulation number and L0 is the smallest cubic box width. We simulate the N =8,64,512- atom systems with L0 = 3, 5, 10,This enables respectively. direct comparison because we can identify triplets of simulated systems corresponding to systems of the same number densities. The two-atom system however is simulated in a sparser regime with L0 = 5. We calculate statistics on the fly for every time step, for every atom and for each coordinate - therefore we average the computed results over the number of time steps (tsim/ $\Delta$ t) and atom coordinates (3N). In particular, the statistics are calculated over 3N tsim/ $\Delta t$  data points. This is equal to  $6 \times 1011$  (resp.  $1.536 \times 109$ ) data points in the simulation with N = 2(resp. N = 512) atoms.

Calculating the number density in three spatial dimensions by n = N/L 3, we can study the behaviour of kurtosis  $\alpha 4$  as n varies. The results are presented in Figure 4. We see general agreement between behaviour of each of the four systems. We see when n is equal, the values of kurtosis are larger for N = 2 than for the manybody systems with N = 8,64,512, which agree well amongst themselves.

The results in Figure 4 enable us to test the asymptotic expression (38) for k = 4 derived in the limit  $n \rightarrow 0$ . Util-





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FIG. 4. Dependence of kurtosis  $\alpha 4$  on density n. Each of the larger atomic systems (N = 8,64,512) is simulated over the same domain of number densities, while the N = 2 system is simulated in a sparser domain, though all are simulated in three spatial dimensions. We truncate the results of the N = 2 simulation in the plot, however the additional data points are used to calculate the results displayed in Figure 5.



FIG. 5. Comparison of the results of MD simulations for a range of values of the number of atoms, N. After long time simulation, we compute the asymptotic behaviour  $\alpha k \propto n - \kappa$  and compare the leading order power scalings for each system. We compare this with the theoretical result (38) (denoted as a blue dashed line) that in the limit  $n \rightarrow 0$  we expect the universal behaviour  $\kappa = k/2 - 1$ , where k = 2,4,6,... denotes which standardised moment of force we are looking at.

ising similar log-log plots for MD data, we estimate the power law behaviour of each standardised moment,  $\alpha k$ , for k =4,6,8,10,12. Figure 5 illustrates the results. All systems agree well with the predicted asymptotic behaviour (38), in particular the N = 512 atom system. There is a slight deviation between the results due to the fact that the smaller atom systems require a larger tsim in order to converge fully to the predicted value. This discrepancy is amplified when looking at higher standardised moments due to the fact that we are calculating statistics resulting from F 12 1 (i.e. for  $\alpha$ 12) compared to F 4 1 (i.e. for  $\alpha$ 4), for example.



FIG. 6. Dependence of kurtosis  $\alpha 4$  on temperature T . Each atomic system is simulated at approximately the same density n = n0 given in Table I.

The dependence of kurtosis  $\alpha 4$ on temperature T is presented in Figure 6, where we keep the density fixed at n = n0given in Table I. We observe that as temperature increases so does the kurtosis of the force distribution associated with each system. This can be explained in terms of the dynamics of the interacting atom system. If we maintain each system in the canonical ensemble, we expect on average that each atom will have a kinetic energy equivalent to 3T/2 (when in reduced units). As we increase this target temperature, the atoms become more energetic and thus are able to probe closer interatomic distances before a large repulsive force overcomes this inertial attraction. The range of forces on the tagged particle widens as temperature increases and therefore contributes to more outlier results in the distribution - leading to heavier tails and therefore distributions which become increasingly leptokurtic.



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In Figure 6, we observe that there is a qualitative difference between the results for N = 2 and larger atom systems. We see a bifurcation for the N = 64 and N = 512systems at some temperature  $T^* \in$ (0.6, 0.65), where a steady increase in kurtosis changes to a rapid increase. This bifurcation point in the phase plane lies on the coexistence boundary with (n,T) =(1/64,T\*) and is due to a clustering mechanism which has been seen in MD simulations of Lennard-Jones fluids29. From our results we see that the N = 2missed this system has behaviour completely. Snapshots of the N = 512atom system at some T = 0.6 < T\*, and T =0.66 > T\* are displayed in Figure 7. For T = 0.6, we see a large cluster has formed in the many-atom system. There would be far fewer outlier force results in this case due to the fact that the large majority of atoms are moving as a collective and effectively have fixed inter-atomic forces. Compared to the T = 0.66 snapshot, where we see that the atoms are too kinetically unstable to form these larger stable cluster structures, this results in more outlier forces felt between atoms due to the fact that the system is intrinsically more disordered. It is useful to note that this bifurcation point is located on the vapour-liquid coexistence boundary, the mechanisms of which have been studied on dilute Lennard-Jones fluids30; here we see that this results in a bifurcation on standardised moments of the force distribution.

To understand the underlying variations of kurtosis,  $\alpha 4$ , with respect to changes in temperature and density, we use  $12 \times 16$  MD simulations with N = 512 atoms and tsim =  $3 \times 106$ , varying simulation parameters (n,T), where n = 10-2 + (i-1)/10, for i = 1,2,...,12, and T = 10-1 + j/10, for j = 1,2,...,16. The sampled values of excess kurtosis ( $\alpha 4$  -3) are displayed in Figure 8. Here a bifurcation can be seen

when using the smallest density n = 0.01, as the change in colour is prominent in this vertical strip, indicating a large change of kurtosis. This occurs around T = 0.6, which is consistent with the result in Figure 6, where we saw the bifurcation similarly located, though the slight shift in temperature is accounted for by the shift in density parameters used in each simulation (namely n = 0.01 in Figure 8 and n = 1/64in Figure 6).

In general, this low density strip contains the largest values of kurtosis, and covers much of the purely gas phase of the Lennard-Jones fluid. This paper has so far probed the low density limit in an attempt to understand why the standardised moments of force are so large, though Figure 8 gives a good overview that in general, regardless of phase, a decrease in temperature, or an increase in density, systematically lead to a lower value of standardised moments. In this case as  $n \rightarrow$  $\infty$  or T  $\rightarrow$  0, we expect the  $\alpha 4 \rightarrow 3$  (excess kurtosis tends to zero). This limiting regime corresponds to the solid phase of a Lennard-Jones system, where the force variations are minimal and the distribution is Gaussian. There is not enough space, nor energy, that lead to (many) outlier forces experienced by any atom, so the force distribution becomes less and less skewed from Gaussian, the deeper we probe in these regions. This intuition was demonstrated analytically in Section III E when we showed this limiting behaviour on a 1D cartoon model with equation (31). It is interesting to note that these changes in values of  $\alpha 4$  appear smooth about changes in temperature and density (in absence of the bifurcation point for larger values of n), regardless of phase transitions.



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FIG. 7. Snapshots31 of the MD simulation are taken for the system with N = 512 atoms at time t =  $7.5 \times 105$  for: (a) T = 0.6 <T\*; and (b) T = 0.66 > T\*. Density is n = 1/64.

## **V. DISCUSSION AND CONCLUSIONS**

In Section III we have demonstrated use of a variety of methods to study the standardised moments of the force distribution in order to probe both their and number density temperature dependence. This gave way to a rich structure where we show that the partition function for a 1D system can be calculated entirely from these standardised moments. Extending the far field method introduced in Section III B to a system with N atoms in three-dimensional physical space. Section IV studies the dependence of  $\alpha k$ on number density n,



FIG. 8. The excess kurtosis,  $\alpha 4 -3$ , calculated as a function of density n and temperature T for  $n \le 1.11$  and  $T \le 1.7$ . The white dotted lines describe coexistence lines of different phases of a LennardJones fluid taken from the

literature32–35. The solid black dots indicate (from left to right), the critical point and vapour-liquid-solid triple points.

deriving the asymptotic expression (38). Our analytic results are contrasted with MD simulations of four systems of N =2,8,64,512 interacting Lennard-Jones atoms and these are compared. The results agree well with theoretical predictions though the results for systems with larger values of N are seen to converge more readily to the theoretically predicted results. In particular, rich dynamics such as clustering of Lennard-Jones fluids is completely missed by the systems with smaller values of N, but captured for systems with N as small as N = 64 atoms. In general, as temperature increases  $\alpha k$ increases due to energetic nature of atoms allowing them to push closer together and larger forces. experience Clustering exhibited at the vapour-liquid coexistence phase incurs a bifurcation point whereby a large increase is seen in the standardised moments of force in Figure 6, though a general increase in temperature, or decrease in number density, results in an in a4 regardless of increase the temperature/number density domain studied, as shown in Figure 8.

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