

ARTICLE TYPE

Cite this: DOI: 00.0000/xxxxxxxxxx

The Lennard-Jones potential: when (not) to use it.

Xipeng Wang, a,b Simón Ramírez-Hinestrosa, b Jure Dobnikar, a,b,c‡ and Daan Frenkela*

Received Date Accepted Date

DOI: 00.0000/xxxxxxxxxx

The Lennard-Jones 12-6 potential (LJ) is arguably the most widely used pair potential in Molecular Simulations. In fact, it is so popular that the question is rarely asked whether it is fit for purpose. In this paper, we argue that, whilst the LJ potential was designed for noble gases such as argon, it is often used for systems where it is not expected to be particularly realistic. Under those circumstances, the disadvantages of the LJ potential become relevant: most important among these is that in simulations the LJ potential is always modified such that it has a finite range. More seriously, there is by now a whole family of different potentials that are all called *Lennard-Jones 12-6*, and that are all different - and that may have very different macroscopic properties. In this paper, we consider alternatives to the LJ 12-6 potential that could be employed under conditions where the LJ potential is only used as a typical short-ranged potential with attraction. We construct a class of potentials that are, in many respects LJ-like but that are by construction finite ranged, vanishing quadratically at the cut-off distance, and that are designed to be computationally cheap. Below, we present this potential and report numerical data for its thermodynamic and transport properties, for the most important cases (cut-off distance r_c =2 σ ("LJ-like") and r_c =1.2 σ (a typical "colloidal" potential).

1 Background

One of the most widely used intermolecular potentials in classical many-body simulations, is the so-called Lennard-Jones 12-6 potential

$$v_{LJ}(r) = 4\varepsilon \left(\left[\frac{\sigma}{r} \right]^{12} - \left[\frac{\sigma}{r} \right]^{6} \right)$$
 (1)

where ε denotes the depth of the attractive well, and σ the interparticle distance where the potential changes sign. Lennard-Jones-type $r^{-\nu}$ - $r^{-\mu}$ pair potentials were proposed in 1925 by Jones (later Lennard-Jones) to describe the cohesive energy of crystals of noble gases, such as Argon. The now conventional LJ 12-6 form was proposed by Lennard-Jones in 1931 after London had derived that the dispersion interaction between atoms decays as r^{-6} (at least, in the non-retarded regime) . It later turned out the LJ 12-6 potential is not a particularly good pair potential for Argon 4,5. However, when the first Monte Carlo (MC) simulations of argon were carried out by Wood and Parker and

subsequently, the first Molecular Dynamics (MD) simulations by Rahman ⁷, there was an unexpectedly good agreement between the simulation results and the experimental data for liquid argon. The reason, as was argued in ⁸ was due to a fortuitous cancellation of errors. However, towards the end of the 20-th century, the LJ 12-6 potential had already achieved an almost unassailable status in classical simulations: it was (and is) used to describe atoms, molecules, coarse-grained models of proteins, and in some cases even larger particles such as nano-colloids. However, for these systems, there is no evidence at all that the LJ 12-6 potential is better than other possible choices. Yet, whenever new simulation techniques are tested, the LJ 12-6 potential is the first model to try.

However, even if the true Lennard-Jones 12-6 potential would have been a satisfactory all-purpose potential, there is a practical problem: the Lennard-Jones potential has an infinite range, which would make it less suited for efficient numerical simulations (note that the infinite range was an advantage for (Lennard-)Jones's analytical calculations). This problem is normally resolved in practice by truncating the potential at a finite distance r_c , e.g. r_c =2.5 σ . Unfortunately, not all authors use the same truncation procedure, and in recent years this confusion has become worse, as the cost of using a larger (but still finite) cut-off distance has become less prohibitive. In addition, in MD simulations, one should truncate the force, rather than the potential. So actually, the potential is truncated and shifted. Yet even such a

^a Institute of Physics, Chinese Academy of Sciences, 8 Third South Street, Zhongguancun, Beijing 100190, China,

 $^{^{\}it b}$ Department of Chemistry, University of Cambridge, Lensfield Road, CB21EW Cambridge, UK

^c Songshan Lake Materials Laboratory, Dongguan, Guangdong 523808, China

 $[\]dagger$ Electronic Supplementary Information (ESI) available: [details of any supplementary information available should be included here]. See DOI: 10.1039/cXCP00000x/

[±] id489@cam.ac.uk

^{*} Corresponding author. df246@cam.ac.uk

force truncation is often not good enough, because a discontinuity in the truncated force creates numerical problems in solving the equations of motion. Therefore, many simulators use a version of the LJ 12-6 potential that is truncated and shifted, and the force is modified such that it goes to zero continuously at r_c . And even that procedure is not unique. So, in the end, there are possibly dozens of different so-called LJ 12-6 potentials in use that are all different, mostly untested for noble gases, and not expected to be particularly good for other molecules either. All these models have different thermodynamic properties: their critical temperatures may differ by more than 30% and the liquid-vapour surface tension of these models may differ by much more than that 9. Finally, this confusion also makes it harder to compare algorithms, if different authors test their favorite algorithm on a different LJ flavor. This problem is well recognized by the community and, in a recent paper, Hafskjold et al 10 have shown that the properties of one particular LJ 12-6 variant (the LJ spline model with r_c = 1.74σ) are quite different from those of the full LJ 12-6 potential.

In summary: the LJ 12-6 potential is anything but a well-defined standard and, in particular for proteins and nano-colloids it is not a good model at all because the range of attraction is too large compared to the effective diameter. There are, of course, LJ n-m style potentials that have been designed to mimic the phase behavior of colloids, but there the range of choices that have been proposed is even larger than for the LJ 12-6 potential.

However, once we give up on the long-ranged dispersion interaction, it is not obvious why a truncated LJ 12-6 potential should have any special merits that outweigh its disadvantages. We argue that no such advantage exists. In fact, even for the quantitative prediction of the properties of noble gases and similarly simple substances, the LJ 12-6 potential is unlikely to remain the model of choice, as force-fields based on machine learning applied to ab-initio simulations are increasingly taking over.

What we need for computer experiments (rather than for simulations) is a simple, short-ranged potential that is at least as simple as LJ 12-6, has none of its draw backs and is defined unambiguously.

2 Constructing a computationally attractive alternative

If we accept that in most cases of practical interest, the choice of the LJ potential is neither justified by theory nor by experiment, it is logical to ask what requirements should be met by a computationally cheap, generic model pair potential for simple atomistic or molecular systems. Below, we formulate our wish list.

- 1. The potential should be repulsive at short distances $(r < \sigma)$ and attractive up to a cut-off distance r_c that should not be much larger than σ . As we explain below, we opt for $r_c = 2\sigma$ for atomic systems (which yields a rather LJ-like phase diagram), but for colloidal systems, a smaller value of r_c is preferable (we explore $r_c = 1.2\sigma$).
- 2. A potential to represent simple liquids should have liquid, vapor and crystalline phases, with a ratio between the critical temperatures and the triple point between ~ 1.7 (neopentane) and ~ 2.1 (methane). (LJ 12-6 has a ratio ~ 1.91 -

1.95, and argon \sim 1.8). In contrast, the "colloidal" version of the potential should not exhibit a transition between two thermodynamically stable fluid phases ("liquid" and "vapor").

- 3. At the cut-off, the potential should vanish (at least) quadratically, such that the pair force vanishes continuously at r_c .
- 4. Evaluating the potential should require only few arithmetic operations, and those should be cheap.

In addition, it is clear that a new pair potential is not very attractive, unless we know its most important thermodynamic and transport properties. In what follows, we will report the dependence of the pressure, energy and free energy on number density $\rho \equiv (N\sigma^3/V)$ and temperature $T \equiv k_BT/\varepsilon$. We give the predicted phase diagram, and the liquid-vapor surface tension. And finally, we report the relevant transport properties (diffusivity, viscosity, thermal conductivity) again as a function of ρ and T. For the solid phase, we only report the thermal conductivity. All data have been fitted to multi-variate polynomials. The raw data are accessible in the Supplementary Information.

3 The model

We wish to construct a simple pair potential that changes from repulsive to attractive at σ and that vanishes quadratically at r_c . In fact, we also give a more general form that vanishes as a higher power of $r-r_c$. However, whilst such generalization may have some applications in testing MD algorithms, we do not recommend them. The general form of the potential that we propose is

$$\phi(r) = \varepsilon \alpha \left(\left[\frac{\sigma}{r} \right]^{2\mu} - 1 \right) \left(\left[\frac{r_c}{r} \right]^{2\mu} - 1 \right)^{2\nu} , \qquad (2)$$

where α is a coefficient that ensures that the depth of the attractive well is $-\varepsilon$. μ and v are positive integers. We can obtain a simple analytical expression for α and the value of r where the potential reaches its minimum (see Appendix A). We shall consider the simplest case ($v = \mu = 1$), for which

$$\phi(r) \equiv \epsilon \alpha(1, 1; r_c) \left[\left(\frac{\sigma}{r} \right)^2 - 1 \right] \left[\left(\frac{r_c}{r} \right)^2 - 1 \right]^2 \text{ for } r \leq r_c$$

$$= 0 \qquad \text{for } r > r_c \qquad (3)$$

with

$$\alpha(1,1;r_c) = 2\left(\frac{r_c}{\sigma}\right)^2 \left(\frac{3}{2(\left(\frac{r_c}{\sigma}\right)^2 - 1)}\right)^3 \tag{4}$$

and

$$r_{min}(1,1;r_c) = r_c \left(\frac{3}{1+2\left(\frac{r_c}{\sigma}\right)^2}\right)^{1/2}$$
 (5)

3.1 The recommended model

In what follows, we use σ as our unit of length and ε as our unit of energy. A particularly simple expression for the pair potential results if r_c =2, because $\alpha(1,1;2)$ =1. This is our preferred model: it has a minimum at $r_{min}\approx 1.155$, compared to the LJ 12-6 minimum at $r_{min}(1,1;2)\approx 1.1225$. The (1,1:2)-potential has

a relatively short range and is therefore computationally cheap. Moreover, it approaches zero quadratically at r_c . However, for colloidal models, a smaller value of r_c is recommended (we will show results for $r_c = 1.2$ with $r_{min} \approx 1.055$).

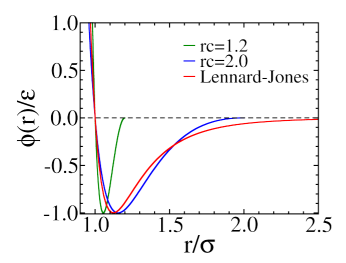


Fig. 1 Comparison of the untruncated Lennard-Jones 12-6 potential (red curve), and the short-ranged potentials described in the text: the "standard" form with $\mu=\nu=1$ and r_c =2.0 is shown as a blue curve. The green curve applies to the the "colloidal" form with $\mu=\nu=1$ and r_c =1.2.

Figure 1 shows a comparison of (1,1,2) and (1,1,1.2)-versions of simplified potential with the original LJ 12-6 potential. Of course, there is a wide choice for values of r_c , v and μ , but we argue that, as the advantage of this potential is its simplicity and not its realism for any specific system, there is usually little to be gained by choosing other values of μ and ν . In principle, increasing ν would make higher derivatives of the potential continuous at r_c , but this comes at a computational cost. The best way to vary the range of the potential is to vary r_c . Higher values of n might be interesting when comparing different Molecular Dynamics algorithms.

Of course, one could also construct a purely repulsive version of these potentials by shifting the potential by $+\varepsilon$ and truncating it beyond r_{min} . However, there is less need for such a model, as the repulsive version of the Lennard-Jones 12-6 potential 11 does not suffer from the ambiguities of the attractive LJ potential.

4 Properties of the v = 1, $\mu = 1$ model

One reason why the LJ 12-6 potential is widely used is not its realism, but simply the fact that by now many observable properties of this model have been computed (see, for instance: equation of state of the fluid $^{12-14}$ and solid 13,15 , transport properties $^{16-19}$ and phase diagram $^{20-22}$). The potential given in Eqn. 3 would therefore be of little practical use, if we did not present fairly complete and concise information about its most important equilibrium and transport properties.

For this reason, we have computed the equation of state, internal energy and free energy of the v=1, $\mu=1$, $r_c=2$ and the v=1, $\mu=1$, $r_c=1.2$ models between low temperatures and a

temperature of 1.4 in reduced units for r_c = 1.2 and 1.6 for r_c = 2.0, and between low densities and a density of 1.4 (in reduced units). In addition, we have computed the phase diagrams, the transport properties of the fluid phase (diffusivity, shear viscosity and thermal conductivity), and the liquid-vapor surface tension for the case of r_c = 2.0. Finally, we have also computed the thermal conductivity of the crystalline phase.

All data points can be found in the Supplementary Information (SI). Here we present the coefficients of a multi-variate polynomials fit the describes our simulation data to within the statistical error. To be precise: the fits of the free energy at high T and ρ have a deviation that is slightly larger than the statistical error. We did not try to improve this, as it would make the fit functions more complicated.

4.1 Results for $r_c = 2.0$

We first discuss the equation-of-state data. For the fluid, we computed the excess energy and excess pressure, *i.e.* the energy and pressure minus the corresponding quantities for an ideal gas at the same temperature and density.

For the solid (we considered face-centered cubic and hexagonal close packed) we also computed the excess energy and pressure, and the excess Helmholtz free energy.

We performed simulations over a grid of data points for $0 < \rho \le 1.14$ for the liquid, and between $\rho = 0.88$ and $\rho = 1.4$ for the solid, both over a temperature range between 0.52 and 1.4 (in reduced units). The resolution was 0.02 in both ρ and T, although some addition low density points ($\rho < 0.02$) where included for the vapor phase. Moreover, we left out a small number of densities and temperatures from the grid. As all fits turned out to be very smooth, we decided not to fill in these missing points later. All points that turned out to be located in a two-phase region (Liquid-Vapor or Solid-Liquid) where disregarded when performing a fit to the data. Hence the location of phase boundaries is only based on information about thermodynamically stable state points.

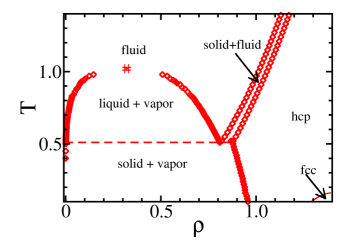


Fig. 2 Computed phase diagram of the potential given by Eqn. 3 for a cut-off distance r_c =2.0 ("Lennard-Jones-like"). This phase diagram was computed for a system size of N=1000 particles.

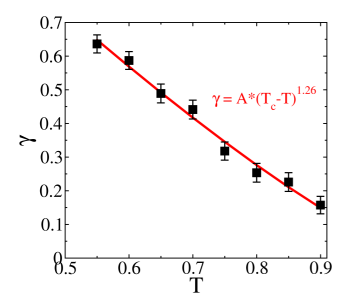


Fig. 3 Temperature dependence of the Liquid-Vapor surface tension of the $r_c=2.0$ model. The curve through the simulation data points was drawn assuming that the surface tension vanishes at the critical point $(T_c=1.04\pm0.02)$ with the 3D Ising critical exponent.

Figure 2 shows the phase diagram of the model potential given by Eqn. 3 for a cut-off distance r_c =2.0, for a system size of N=1000 particles. For this system size, our estimate of the critical temperature is $T_c \approx 1.04$ with a critical density $\rho_c \approx 0.32$. We have not tried to refine the estimate of the critical point using finite-size scaling 23 , as this level of detail is beyond the scope of the present paper.

We estimate the triple point to be located $T_{tr} \approx 0.505$, where the density of the solid is $\rho_S \approx 0.883$ and that of the coexisting liquid $\rho_L \approx 0.817$. We note that the ratio of the critical temperature to that of the triple-point temperature is approximately 2.06, which is higher than the corresponding ratio for argon but lower than that for methane. Figure 3 shows the temperature dependence of the Liquid-Vapor surface tension between $T=0.9T_c$ and the triple-point temperature. The simulation data agree well with the assumption that the surface tension approaches the value zero at the critical point ($T_c \approx 1.04$) with a power law with the 3D Ising critical exponent. Earlier simulations also found there to be no significant deviations from this scaling form, all the way down to the triple-point temperature. We note that the numerical value of the surface tension is comparable to the values found for various Lennard-Jones variants at the same reduced temperature T/T_c .

Note that the slope of the solid-liquid coexistence curves is lower than for the LJ 12-6 system. This is because the repulsive potential in the current model is less steep than in the LJ case $(r^{-6}$ rather than r^{-12}). If necessary, the slope could be increased by changing μ in the model from $\mu = 1$ to $\mu \geq 2$.

At the triple point, the density of the vapor is extremely low (but can be computed from the knowledge of the chemical potential, which can be computed from the multivariate fit using Eqn. 12).

For this Lennard-Jones-like model, we expect that the stable

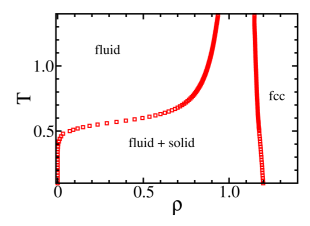


Fig. 4 Computed phase diagram of the potential given by Eqn. 3 for a cut-off distance r_c =1.2 ("colloid-like"). This phase diagram was computed for a system size of N=1000 particles.

solid phase is either face-centered cubic (fcc) or hexagonal close-packed (hcp). In fact, we find both phases. For the $r_c=2.0$ model, the hcp phase appears (very slightly) more stable than fcc, except for a small pocket at high densities and low temperatures. The numerical values of the free energy as a function of density and temperature are given in the SI. We have computed the transport properties and phase boundaries for the fcc crystal phase. Our free-energy calculations showed that, in fact, the hcp phase is slightly more stable for most densities. However, the observable properties of the two phases are so similar that we did not recompute them for the hcp solid.

4.2 Results for $r_c = 1.2$

For $r_c=1.2$, the phase diagram shown in Fig. 4 does not include a liquid-vapor transition in the stable region. Such behavior is typical for colloidal systems with a relatively short-ranged attractive interaction 24 . For the $r_c=1.2$ model, the fcc crystal phase is more stable than hcp, for all densities studied. Figure 4 shows the phase diagram of the model potential given by Eqn. 3 for a cut-off distance $r_c=1.2$, for a system size of N=1000 particles. This model corresponds to a typical "colloidal" system, which has no liquid-vapor phase transition in the thermodynamically stable region.

4.3 Simulation details

All simulations were carried out using LAMMPS 25 , with a Hamiltonian thermostat, except in the computation of the thermal conductivity, where we used the Nosé-Hoover thermostat, and compared the results with constant NVE and with the results for a Hamiltonian thermostat: to within the statistical error, we found no difference between the results obtained using these three methods. The free-energy calculations for the crystalline solids were carried out in a system of 768 particles, with a periodic box shape that was compatible with both fcc and hcp packing: 12 close-packed planes and 8×8 primitive cells in the close-packed planes $(L_x : L_y : L_z = 1 : \sqrt{3/4} : \sqrt{3/2})$.

For the direct liquid-vapor coexistence calculations, we simulated a system consisting of N=1000 particles in an elongated box with dimensions $L_x:L_y:L_z=1:1:4$. For the thermal conductivity (only for the fcc solid), we used a box $10\times10\times9$ primitive cells (900 particles). All other simulations were carried out for cubic boxes containing 1000 particles. In all cases, periodic boundary conditions were employed. All pressures reported were computed using the virial expression for the stress.

4.4 Transport properties

We used Green-Kubo expressions to compute the diffusivity, viscosity and thermal conductivity. We did not attempt to correct for finite-size effects in the transport properties, even though these may be significant ²⁶. The most important finite-size effect is presumably in the diffusion constant. Dünweg and Kremer ²⁷ and Yeh and Hummer ²⁸, have proposed an expression to correct for this finite size effect for the computation of the diffusivity (at least for a periodically repeated cubic box, as used in this work):

$$D_{\infty} = D_{\text{observed}} + \frac{k_B T \xi}{6\pi n L} \tag{6}$$

where D_{∞} is the diffusion constant in an infinite (non-periodic) medium and D_{observed} is the diffusion coefficient observed in a periodic system with box diameter $L=(N/\rho)^{1/3}$. ξ is a numerical constant withe the value $\xi \approx 2.837297$, and η denotes the shear viscosity of the fluid. The finite-size correction described by Eqn. 6 is valid if the fluid behaves as a hydrodynamic continuum on length scales comparable to the system size. However, for highly viscous fluids, reaching this hydrodynamic regime may require very large system sizes. To our knowledge, there is no numerical evidence for a systematic system-size dependence of the shear viscosity η^{28} . In contrast, finite-size corrections to the thermal conductivity κ may be large, in particular for solids (see Chantenne and Barrat who studied finite-size effects in nonequilibrium MD simulations²⁹). However, there is little information about finite-size effects in the thermal conductivity obtained from Green-Kubo integrals, nor is there much information about such finite-size effect in liquids.

4.5 Free-energy calculations

For the free-energy calculations, we used the Einstein-crystal method 30 . The maximum spring constant in the Einstein crystal integration was chosen such that the mean-squared amplitude of the displacement of particles from their lattice sites was the same for the Einstein crystal as for the original crystal. To avoid possible problems with the diverging integrand of the thermodynamic integration in the Einstein limit (see *e.g.* 31), we did not start the thermodynamic integration at the Einstein limit, but at a point nearby where the integrand is well behaved. To be precise, writing the potential-energy function that interpolates between Einstein crystal and real crystal as

$$U(\lambda) = \lambda U_{\text{crystal}} + (1 - \lambda) U_{\text{Einstein}} , \qquad (7)$$

we separated the calculation of the free energy difference into two parts:

$$\beta \Delta F = -\ln \langle \exp(-\beta \Delta U_{\lambda_{min}}) \rangle + \int_{\lambda_{min}}^{1} d\lambda \langle \beta \Delta U(\lambda) \rangle,$$
 (8)

where

$$\Delta U(\lambda) \equiv \langle U_{\text{crystal}} - U_{\text{Einstein}} \rangle_{\lambda} . \tag{9}$$

For the system size used in the free-energy calculation, we used λ_{min} =0.0001. That value could have been further optimised, but optimisation had no noticeable effect on the value or accuracy of the result. The integration in Eqn. 8 was carried out using a 10-point Gauss-Legendre quadrature.

5 Multi-variate polynomial fits

All numerical data, can be found in the SI. Here we just present the results of the multi-variate polynomial fits to our numerical data. For convenience and ease of use, we have chosen polynomials as our basic fitting functions. We do not suggest that our choice of the fitting functions is optimal.

5.1 Thermodynamic properties

As the data for temperature, pressure and free-energy are related (see below) we used *a single* multi-variate fit for excess pressure and energy of the fluid, and a single fit of excess energy, pressure and Helmholtz free energy for the solid. The central quantity is the excess Helmholtz free energy $A_{exc}(N,V,T)$, or more precisely $\beta A_{exc}/V = \beta \rho a_{exc}$, where $\beta \equiv 1/k_B T$, and $a_{exc} \equiv A_{exc}/N$. We will make use of the following thermodynamic relations:

$$\beta P_{exc} = -\left(\beta \rho a_{exc} - \rho \left(\frac{\partial \beta \rho a_{exc}}{\partial \rho}\right)_T\right), \qquad (10)$$

and

$$\rho e_{exc} = \left(\frac{\partial \beta \rho a_{exc}}{\partial \beta}\right)_{0}.$$
 (11)

To compute the phase diagram, we will also use

$$\beta \mu_{exc} = \left(\frac{\partial \beta \rho a_{exc}}{\partial \rho}\right)_T . \tag{12}$$

Finally, ignoring irrelevant constants, the following relations hold for the ideal gas: $\beta P_{id} = \rho$ and $\beta \mu^{id} = \ln \rho$. Hence, if we have an expression for $\beta \rho a_{exc}$, we also have expressions for P, e and μ .

We assume that $\beta \rho a_{exc}$ for phase α can be expanded in a multivariate polynomial. For the liquid we have:

$$\beta \rho a_{exc}^L = \sum_{n=n_{min}}^{n_{max}} \sum_{m=m_{min}}^{m_{max}} a_{n,m}^L \rho^n \beta^m . \tag{13}$$

However, to reproduce the correct (harmonic) low-temperature behavior of the solid, we write

$$\beta \rho a_{exc}^S = \frac{3}{2} \rho \ln \beta + \sum_{n=n_{min}}^{n_{max}} \sum_{m=m_{min}}^{m_{max}} a_{n,m}^S \rho^n \beta^m . \tag{14}$$

n	m=-3	m=-2	m=-1	m=0	m=1	m=2
2	0.76902	-3.15391	2.88640	5.17074	-10.4462	0.865638
3	3.26974	-77.7391	368.594	-723.031	646.553	-219.3018
4	-8.29352	377.872	-1929.64	3888.37	-3522.43	1219.511
5	-31.8858	-597.874	3874.72	-8352.09	7764.53	-2720.454
6	102.257	349.298	-3783.50	8937.04	-8548.36	3013.089
7	-97.1753	-4.10970	1783.38	-4738.69	4672.47	-1647.794
8	31.1298	-44.7404	-315.575	985.658	-1004.95	353.2964
2	19.1395	-92.8140	151.915	-99.3440	16.8835	
3	-21265.4	75376.4	-100238	59293.4	-13166.0	

Table 1 Fit coefficients for the expression for excess free-energy density of the liquid phase and the vapor phase for the model with r_c =2.0 (Eqn. 13). The upper part of the table gives the coefficients that correspond to a fit to the simulation data for all simulated fluid densities at temperatures above T_c and, for supercritical densities below T_c . The lower part of the table gives the coefficient for temperatures below T_c and densities below T_c . In the SI, we give the coefficients with full numerical accuracy.

We then have (for phase $\alpha = S$ or L):

$$\beta P_{exc}^{\alpha}(\rho,\beta) = \sum_{n=n_{\text{min}}}^{n_{\text{max}}} \sum_{m=m_{\text{min}}}^{m_{\text{max}}} (n-1) a_{n,m}^{\alpha} \rho^n \beta^m$$
 (15)

and

$$\rho e_{exc}^{S} = \frac{3}{2}\rho T + \sum_{n=n}^{n_{max}} \sum_{m=m}^{m_{max}} a_{n,m}^{S} m \rho^{n} \beta^{m-1}$$
 (16)

or

$$\rho e_{exc}^{L} = \sum_{n=n_{min}}^{n_{max}} \sum_{m=m_{min}}^{m_{max}} a_{n,m}^{L} m \rho^{n} \beta^{m-1}$$
 (17)

When fitting the excess pressure and excess energy-density of the fluid/liquid phase, we used n_{min} =2 (because at low densities, the excess pressure scales as ρ^2), n_{max} =8, m_{min} =-3 and m_{max} =2. In the case of r_c =2.0, this form was used for all densities of the fluid at temperature above T_c and for the liquid down to the triple point. Note that all these data points correspond to temperatures above T_{tr} .

For the vapor phase (r_c =2.0), we used a polynomial with n_{max} =6, m_{min} =2 and m_{max} =2 and m_{min} =-3.

Note that the coefficient of ρ^2 follows from the second virial coefficient, that we have computed separately. For the ease of use, it is however, more convenient to treat all fit coefficients on an equal footing (that is: the coefficient of ρ^2 was not derived from $B_2(T)$, but was fitted with all the other coefficients.

5.2 Transport properties

We used multi-variate polynomial fits to approximate the diffusivity, viscosity and thermal conductivity of the fluid, and the thermal conductivity of the solid. As the diffusivity D diverges as $1/\rho$ at low densities, we fitted ρD . We used the following functional forms: for the diffusivity

$$\rho D = \sum_{n=n_{\text{min}}}^{n_{\text{max}}} \sum_{m=m_{\text{min}}}^{m_{\text{max}}} \alpha_{n,m}^{(D)} \rho^n \beta^m . \tag{18}$$

with $n_{min} = 0, n_{max} = 3, m_{min} = -3$ and $m_{max} = 0$. For the viscosity:

$$\eta = \sum_{n=n_{min}}^{n_{max}} \sum_{m=m_{min}}^{m_{max}} \alpha_{n,m}^{(\eta)} \rho^n \beta^m.$$
 (19)

with n_{min} =0, n_{max} =6, m_{min} =0 and m_{max} =6. For the thermal conductivity of the fluid and the solid:

$$\kappa = \sum_{n=n,\dots,m}^{n_{max}} \sum_{m=m_{min}}^{m_{max}} \alpha_{n,m}^{(\eta)} \rho^n \beta^m . \tag{20}$$

with $n_{min}=0$, $n_{max}=6$, $m_{min}=0$ and $m_{max}=6$.

m=-2	m=-1	m=0	m=1
59.9947	-111.081	23.4348	19.8311
-254.708	458.412	-85.2926	-88.2034
427.727	-749.243	124.562	154.892
-356.028	607.871	-75.9448	-157.340
147.127	-245.199	21.1821	76.6613
-24.1761	39.3881	-1.96241	-12.3529
	59.9947 -254.708 427.727 -356.028 147.127	59.9947 -111.081 -254.708 458.412 427.727 -749.243 -356.028 607.871 147.127 -245.199	59.9947 -111.081 23.4348 -254.708 458.412 -85.2926 427.727 -749.243 124.562 -356.028 607.871 -75.9448 147.127 -245.199 21.1821

Table 2 Fit coefficients for the expression for excess free-energy density of the *fcc* crystal phase for the model with r_c =2.0 (Eqn. 14). In the SI, we give the coefficients with full numerical accuracy.

n	m=-3	m=-2	m=-1	m=-1 $m=0$		
0	0.69663	-2.56961	3.31891	-1.33620		
1	-3.40977	12.1701	-14.5135	5.95227		
2	5.10453	-17.8776	20.7046	-8.46850		
3	-2.41105	8.35025	-9.54006	3.85669		
0		-2.79549	5.85772	-2.94945		
1		48.2219	-97.7224	49.7001		
2		-175.126	355.337	-180.773		

Table 3 Fit coefficients for the expression for ρD of the liquid phase and vapor phase for the model with r_c =2.0 (Eqn. 18). The upper part of the table gives the coefficients that correspond to a fit to the simulation data for all simulated fluid densities at temperatures above T_c and, for supercritical densities below T_c . The lower part of the table gives the coefficient for temperatures below T_c and densities below P_c . In the SI, we give the coefficients with full numerical accuracy.

5.3 Fits for $r_c = 2.0$

5.3.1 Thermodynamic Properties

Table 1 summarizes the fit coefficients for Eqn. 13 for the liquid phase, and separately for the vapor phase, for the model with

n	m=0	m=1	m=2	m=3	m=4	m=5	m=6
0	-1025.67	3674.72	13.3259	-17568.4	31676.6	-22985.9	6218.70
1	3356.81	-23366.1	61687.4	-74998.8	33867.7	8135.96	-8738.63
2	11489.7	-51202.6	71889.4	-2816.70	-65814.6	34706.1	2083.05
3	-10159.4	53558.7	-115496	90391.8	4125.24	-11895.1	-11487.9
4	-1577.01	50211.2	-145801	177803	-101049	-5929.13	27800.4
5	-43091.4	147984	-201402	147912	-87827.2	66770.3	-31457.5
6	42832.8	-191063	352507	-348827	203671	-75284.1	16505.3
0	-24.2194	130.612	-195.143	89.0932			
1	-875.221	1481.66	-235.286	-374.664			
2	-2435.26	5949.15	-4915.41	1419.74			
3	15240.2	-10443.2	-25016.1	20194.2			

Table 4 Fit coefficients for the expression for the shear viscosity η of the liquid phase and vapor phase for the model with r_c =2.0 (Eqn. 19). The upper part of the table gives the coefficients that correspond to a fit to the simulation data for all simulated fluid densities at temperatures above T_c and, for supercritical densities below T_c . The lower part of the table gives the coefficient for temperatures below T_c and densities below T_c . In the SI, we give the coefficients with full numerical accuracy.

n	m=0	m=1	m=2	m=3	m=4	m=5	m=6
0	149.596	-577.751	762.037	-269.597	-232.326	210.741	-42.3878
1	121.498	-1762.49	4762.55	-4701.22	1831.24	-467.709	215.772
2	1267.22	-1646.78	-1088.86	-743.600	3964.64	-897.456	-841.283
3	-2770.85	6173.25	-1969.91	-1254.57	-591.950	-883.117	1299.32
4	65.6882	-1829.26	1879.76	1899.70	-1720.49	-1707.09	1333.25
5	2825.56	-2287.38	-6347.60	6021.19	-379.015	3047.14	-2738.52
6	-1904.17	3644.62	-2652.01	5624.72	-7964.28	2738.41	444.527
0	-210.615	1042.19	-1402.49	570.548			
1	-54.5982	-3446.25	6110.59	-2600.56			
2	2675.78	-12716.3	23099.6	-13092.7			
3	55913.5	-128202	78195.6	-5837.50			

Table 5 Fit coefficients for the expression for κ of the liquid phase and vapor phase for the model with r_c =2.0 (Eqn. 20). The upper part of the table gives the coefficients that correspond to a fit to the simulation data for all simulated fluid densities at temperatures above T_c and, for supercritical densities below T_c . The lower part of the table gives the coefficient for temperatures below T_c and densities below T_c . In the SI, we give the coefficients with full numerical accuracy.

n	m=0	m=1	m=2	m=3	m=4	m=5	m=6
0	31380.5	-54228.7	17548.3	-1481.55	-3480.41	3635.92	-611.898
1	-41206.1	51105.3	22036.7	-2702.04	1864.24	-6366.19	510.533
2	93.1931	10395.4	-34834.1	-10094.2	3169.27	2109.39	1711.08
3	10319.3	-4986.17	-12617.4	3328.08	4016.16	2709.28	-2490.02
4	3854.17	-778.979	11250.5	8097.52	-67.0914	-1984.86	-429.411
5	-3438.01	-9826.25	2586.75	3108.49	-6384.14	-2068.63	2338.61
6	225.031	4188.17	-4.36.24	-3944.54	2129.02	1781.09	-1023.35

Table 6 Fit coefficients for the expression for κ of the *fcc* crystal phase for the model with r_c =2.0 (Eqn. 20). In the SI, we give the coefficients with full numerical accuracy.

 $r_c = 2.0.$

Table 2 summarizes the fit coefficients for Eqn. 14 for the crystalline *fcc* phase for the model with r_c = 2.0.

5.3.2 Transport Properties

In Tables 3-5 we summarize the fitting coefficients for the diffusivity, more precisely the product ρD (Eqn. 18); the shear viscosity (Eqn. 19) and the thermal conductivity (Eqn. 20). All of these transport coefficients were computed for the fluid phase and we have not attempted to include the data for the low-density gas (ρ < 0.1), as in this regime, the Green-Kubo integrals are rela-

tively noisy and the transport coefficients can better be calculated from the Boltzmann equation, using the approach of Chapman and Cowling $^{\rm 32}$

Table 6 summarizes the fit coefficients for Eqn. 20 for the thermal conductivity of the *fcc* solid phase for the model with r_c = 2.0.

5.4 Fits for $r_c = 1.2$

5.4.1 Thermodynamic Properties

Table 7 summarizes the fit coefficients for Eqn. 13 for the fluid phase, for the model with r_c =1.2.

n	m=-3	m=-2	m=-1	m=0	m=1	m=2
2	-1.10009	6.27642	-13.5539	15.5901	-7.05497	0.63096
3	4.79833	-31.7208	76.0381	-84.1657	46.4876	-10.4596
4	-0.47723	19.8875	-62.2031	78.4142	-55.7024	18.4785
5	-7.60729	39.2723	-131.249	205.539	-110.462	9.10103
6	-22.4105	65.7709	66.3596	-315.674	244.739	-43.4275
7	56.4536	-228.002	243.158	30.6737	-133.904	33.7645
8	-29.5309	127.279	-174.625	68.3272	18.5640	-9.79081

Table 7 Fit coefficients for the expression for excess free-energy density of the fluid phase for the model with r_c =1.2 (Eqn. 13). In the SI, we give the coefficients with full numerical accuracy.

n	m=-2	m=-1	m=0	m=1
0	1585.85	-2137.72	-1114.08	-0.10780
1	-6114.51	8239.19	4047.61	-195.525
2	9422.37	-12692.5	-5907.11	682.374
3	-7253.89	9769.46	4342.62	-807.773
4	2789.93	-3757.12	-1600.17	364.210
5	-428.864	577.560	236.283	-46.0042

Table 8 Fit coefficients for the expression for excess free-energy density of the *fcc* crystal phase for the model with r_c =1.2 (Eqn. 14). In the SI, we give the coefficients with full numerical accuracy.

Table 8 summarizes the fit coefficients for Eqn. 14 for the crystalline *fcc* phase for the model with r_c = 1.2.

n	m=-3	m=-2	m=-1	m=0
0	0.04176	-0.15423	0.30690	-0.03915
1	-0.09000	0.28129	-0.36580	0.25672
2	0.41788	-1.36771	1.48079	-0.90280
3	-0.47069	1.57829	-1.76868	0.80454

Table 9 Fit coefficients for the expression for ρD of the fluid phase for the model with r_c =1.2 (Eqn. 18). In the SI, we give the coefficients with full numerical accuracy.

5.4.2 Transport Properties

In Tables 9-11 we summarize the fitting coefficients for the diffusivity, more precisely the product ρD (Eqn. 18); the shear viscosity (Eqn. 19) and the thermal conductivity (Eqn. 20). All of these transport coefficients were computed for the fluid phase and we have not attempted to include the data for the low-density gas (ρ < 0.1), as in this regime, the Green-Kubo integrals are relatively noisy and the transport coefficients can better be calculated from the Boltzmann equation, using the approach of Chapman and Cowling 32

Table 12 summarizes the fit coefficients for Eqn. 20 for the thermal conductivity of the *fcc* solid phase for the model with $r_c = 1.2$.

6 Conclusions

In this paper we have presented a simple model pair-potential that can be used in numerical studies of systems of particles with a short-ranged attraction. The model potential that we use vanishes quadratically at a cut-off distance r_c . Our main reason for proposing this simple model is that, in practice, the widely used LJ 12-6 model is implemented differently (truncated, shifted, in-

terpolated etc.) by different authors. In contrast, the current model is uniquely defined once r_c is specified.

We report a fairly complete set of thermodynamic and transport properties for the cases r_c =2.0 ("atomic liquid") and r_c =1.2 ("colloidal suspension"). The raw simulation data are available online.

We stress that the models that we present are not designed to model any specific substance. Rather, they represent generic models. However, in many cases the ubiquitous LJ 12-6 potential is used in exactly the same way.

A very substantial amount of work has been invested in constructing force fields for realistic models, which are at least partially based on the LJ 12-6 potential. We neither expect nor suggest the reparametrization of all these force fields. In fact, as the current potential is softer than the LJ 12-6 potential, combining it with, for instance, attractive Coulomb interactions, should be done with great care, as the minimum of the resulting interacting could shift deep into the repulsive regime of the potential given by Eqn. 3. In such cases, a using larger value of \boldsymbol{v} may be advisable.

We note that the present model can be easily be extended to describe purely repulsive interactions. However, as the Weeks-Chandler-Andersen representation of the LJ 12-6 potential is not ambiguous, there is less need for a "repulsive" version of the current potential.

Finally, we note that the present model shows LJ-like behavior for a cut-off radius $r_c=2.0$, which should make it cheaper than most LJ 12-6 models, which tend to have larger cut-off radii and therefore have more interacting neighbors.

Comparing speeds is tricky, as the speed of a code depends on the details of the implementation (e.g. computing forces "on the fly" versus tabulated forces). To give an indication of the performance of the potential in Eqn. 3 with r_c =2.0, compared with LJ 12-6 (r_c =2.5), we have carried out simulations of equal length, for 768 particles at ρ =1.2 and T=1.0. We compute the forces on the fly. For the comparison, we chose a state point in the solid, because then there will be no difference in the rate of updating neighbor lists due to possible differences in the particle dynamics. We find that the duration of the simulation for the potential given by Eqn. 3 is about 60% of the duration of the corresponding LJ 12-6 potential. However, once again, we stress that speed is not our primary concern: it is just a bonus.

n	m=0	m=1	m=2	m=3	m=4	m=5	m=6
0	-21.0627	73.8838	-101.523	76.3315	-35.9647	9.97495	-1.10172
1	304.977	-949.825	1118.91	-742.064	358.002	-109.781	11.2826
2	-1514.92	4454.07	-5269.71	4512.62	-3153.32	1166.78	-126.363
3	2031.52	-3468.57	1668.04	-5065.03	7428.87	-3017.70	160.901
4	920.307	-9087.33	15730.5	-2095.78	-7314.55	1478.52	894.480
5	-897.404	2981.78	3051.34	-16181.4	10282.7	2798.33	-2560.19
6	24.0482	611.894	-1780.61	-1038.12	8783.96	-9202.61	2810.34

Table 10 Fit coefficients for the expression for η of the fluid phase for the model with r_c =1.2 (Eqn. 19). In the SI, we give the coefficients with full numerical accuracy.

n	m=0	m=1	m=2	m=3	m=4	m=5	m=6
0	-2.03719	114.606	-433.051	672.146	-528.419	210.761	-34.1161
1	-459.380	592.988	2059.10	-5723.11	5714.97	-2669.76	489.759
2	4792.68	-15213.3	14970.1	1976.83	-14492.9	10377.1	-2424.31
3	-8414.27	30113.4	-43241.1	21722.6	14714.3	-21369.3	6537.43
4	-12862.1	44845.5	-40351.8	2485.73	-9523.57	25508.3	-10224.1
5	31717.7	-98686.9	69562.6	46534.2	-58861.8	3308.63	6557.54
6	-16527.1	48302.2	-27515.2	-33077.9	34935.2	-2969.87	-3195.33

Table 11 Fit coefficients for the expression for κ of the fluid phase for the model with r_c =1.2 (Eqn. 20). In the SI, we give the coefficients with full numerical accuracy.

n	m=0	m=1	m=2	m=3	m=4	m=5	m=6
0	-84237.9	214416	-198388	39887.8	-1683.09	22660.5	-3314.31
1	126293	-179489	231537	-15974.1	-42312.7	-12879.4	3719.35
2	-79926.1	-103018	-50331.8	69792.9	-23547.0	4661.70	-8736.66
3	57608.1	25353.5	-70553.9	26570.2	-10978.8	4057.21	7647.01
4	1760.96	137339	-34058.8	-6416.11	33321.7	1897.50	2704.36
5	-62117.8	-16507.1	18451.1	-32836.2	-4996.54	-5931.71	-6129.40
6	32370.3	-42227.5	38845.9	-16073.6	11788.1	-2072.94	2478.03

Table 12 Fit coefficients for the expression for κ of the *fcc* crystal phase for the model with r_c =1.2 (Eqn. 20). In the SI, we give the coefficients with full numerical accuracy.

A Minimum of generalized potential

Eqn 27 implies that

$$r_{min}(v,\mu;r_c) = \left(\frac{1}{u_{min}}\right)^{1/2\mu}$$

$$= r_c \left(\frac{1+2v}{1+2vr_c^{2\mu}}\right)^{1/2\mu}$$
(28)

and

$$\alpha(v,\mu;r_c) = 2vr_c^{2\mu} \left(\frac{1+2v}{2n(r_c^{2\mu}-1)}\right)^{2v+1}$$
 (29)

Here we derive the expressions for the location and depth of the minimum of the generalized v, μ, r_c potential. Note that in the main text, we use v = 1, $\mu = 1$ and $r_c = 2.0$ and 1.2.

Acknowledgements

DF acknowledges more than 40 years of stimulating discussions with Michiel Sprik, who does not take anything for granted. We gratefully acknowledge discussions with Jürgen Horbach.

Notes and references

- 1 J. Jones, Proc R Soc London A, 1924, 106, 441-462.
- 2 J. E. Lennard-Jones, Proc Phys Soc, 1931, 43, 461-482.
- 3 F. London, Zeitschrift für Physik, 1930, 63, 245-279.
- 4 J. Barker, R. Fisher and R. Watts, Mol. Phys., 1971, 21, 657-&.
- 5 G. Maitland and E. Smith, Mol. Phys., 1971, 22, 861-&.
- 6 W. Wood and F. Parker, J. Chem. Phys., 1957, 27, 720-733.
- 7 A. Rahman, Phys. Rev., 1964, 136, A405.
- 8 M. van der Hoef and P. Madden, *J. Chem. Phys.*, 1999, **111**, 1520–1526.
- 9 V. Baidakov, G. Chernykh and S. Protsenko, *Chem Phys Lett*, 2000, **321**, 315–320.
- B. Hafskjold, K. P. Travis, A. B. Hass, M. Hammer, A. Aasen and O. Wilhelmsen, *Mol Phys*, 2019, 0.1080/00268976.2019.1664780.
- 11 J. D. Weeks, D. Chandler and H. C. Andersen, *J. Chem. Phys.*, 1971, **54**, 5237.
- 12 M. Thol, G. Rutkai, A. Koester, R. Lustig, R. Span and J. Vrabec, *J Phys Chem Ref Data*, 2016, **45**, 023101.
- 13 A. J. Schultz and D. A. Kofke, J Chem Phys, 2018, 149, 204508.
- 14 J. Johnson, J. Zollweg and K. Gubbins, *Mol. Phys.*, 1993, **78**, 591–618.
- 15 M. van der Hoef, J Chem Phys, 2000, 113, 8142–8148.
- 16 D. Levesque, L. Verlet and J. Kürkijarvi, *Phys. Rev. A*, 1973, 7, 1690–1700.
- 17 K. Meier, A. Laesecke and S. Kabelac, *J Chem Phys*, 2005, **122**, 014513.
- 18 K. Meier, A. Laesecke and S. Kabelac, *J Chem Phys*, 2004, **121**, 9526–9535.
- 19 K. Meier, A. Laesecke and S. Kabelac, *J Chem Phys*, 2004, **121**, 3671–3687.
- 20 J.-P. Hansen and L. Verlet, Phys. Rev., 1969, 184, 151–161.
- 21 A. Travesset, J Chem Phys, 2014, 141, 164501.
- 22 M. van der Hoef, J Chem Phys, 2002, 117, 5092–5093.
- 23 K. Binder, Z. Phys. B-Condens. Mat., 1981, 43, 119–140.
- 24 M. Noro and D. Frenkel, J Chem Phys, 2000, 113, 2941-2944.

To locate the minimum of the potential of the type given in Eqn. 2, we define an auxiliary quantity u as

$$u \equiv \left[\frac{\sigma}{r}\right]^{2\mu} \tag{21}$$

and

$$u_c \equiv \left[\frac{\sigma}{r_c}\right]^{2\mu} \ . \tag{22}$$

As before, we use σ as our unit of length and ϵ as our unit of energy. Then

$$\phi(u) = \alpha (u - 1) \left(\frac{u}{u_c} - 1\right)^{2v}, \qquad (23)$$

The condition for the extremum (minimum) is:

$$0 = \left(\frac{\partial \phi(u)}{\partial u}\right) = \alpha \left[\left(\frac{u}{u_c} - 1\right)^{2\nu} + \frac{2\nu}{u_c} (u - 1) \left(\frac{u}{u_c} - 1\right)^{2\nu - 1} \right],$$
(24)

which implies that at u_{min}

$$\left(\frac{u_{min}}{u_c} - 1\right) = -\frac{2v}{u_c} \left(u_{min} - 1\right) , \qquad (25)$$

and hence

$$u_{min}(1+2v) = u_c + 2v \tag{26}$$

or

$$u_{min} = \frac{u_c + 2v}{1 + 2v} \ . \tag{27}$$

- 25 S. Plimpton, Journal of Computational Physics, 1995, 117, 1–19.
- 26 D. M. Heyes, M. J. Cass, J. G. Powles and W. A. B. Evans, *J Phys Chem B*, 2007, **111**, 1455–1464.
- 27 B. Dünweg and K. Kremer, J. Chem. Phys. , 1993, 99, 6983–6997.
- 28 I. Yeh and G. Hummer, *J Phys Chem B*, 2004, **108**, 15873–15879.
- 29 P. Chantrenne and J. Barrat, *J. Heat Transfer Transactions ASME*, 2004, **126**, 577–585.
- 30 D. Frenkel and A. J. C. Ladd, *J Chem Phys*, 1984, **81**, 3188–3193.
- 31 D. Frenkel and B. Smit, *Understanding Molecular Simulation*, Academic Press, London, 2nd edn, 2002.
- 32 S. Chapman and T. G. Cowling, *The Mathematical Theory of non-uniform gases*, Cambridge University Press, Cambridge, 1952.