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# Free electron to electride transition in dense liquid potassium

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At high pressures, simple metals such as potassium have a rich phase diagram including an insulating electride phase in which electrons have a localised, anionic character. Measurements in the liquid phase have shown a transition between two states but experimental challenges have prevented detailed thermodynamic measurements. Using potassium as an example, we present numerical evidence that the liquid-liquid transition is a continuous transformation from free-electron to electride behaviour. We show that the transformation is manifest in anomalous diffusivity, thermal expansion, sound speed, coordination number, reflectivity and heat capacity across a wide range of pressure. The abnormalities stem from a significant change in local electronic and ionic structure. Although primarily a pressure-induced phenomenon, there is also a thermal expansion anomaly. By establishing the electride nature of the high-pressure liquid phase, we resolve the long-standing mystery of how

a liquid can be denser than a close-packed solid. Our work is relevant for high-pressure thermodynamic properties of all alkali metal liquids.

Phase transitions are defined by discontinuous changes in properties of the material in question. In crystalline structures, they are typically associated with a change in symmetry, and a discontinuity in the lattice parameters and density. In magnetic transitions, the spontaneous magnetization drops to zero, and by definition cannot become negative. The meaning of "secondorder" phase transition has evolved since Ehrenfest's original definition of a "discontinuity in second derivative of the free energy". Nowadays, any divergent thermodynamic property is regarded as an indicator of a second-order transition. A final category of "continuous" or "infinite-order" transitions exists which break no symmetries. In a liquid-liquid transition there is no symmetry change, and a first-order transition is possible via a volume discontinuity. Such a transformation is typically (but not always) driven by a change in bonding character. An example is phosphorus, where a molecular liquid comprising tetrahedral P<sub>4</sub> molecules changes to a polymeric form<sup>1</sup>. Others include the molecular-atomic transition in hydrogen<sup>2-5</sup>, superfluid helium<sup>6</sup>, and the transition in water where the coordination rises from 4 to 8, mimicking the transition from the open ice-Ih structure to the denser ice-VII<sup>7-12</sup>. Examples for electronic changes with less obvious effects on bonding are the f-electron delocalization in liquid cerium<sup>13</sup> and the spin collapse in iron-rich silicate melts<sup>14,15</sup>. Often, these transitions are described as a two-state system: a first-order transition sees a finite fraction of particles switch between states, while in a continuous transition the fraction of particles in each of two states varies continuously.

The electronic change underlying the liquid-liquid transition in the alkali metals is not im-

mediately obvious: these elements are the best examples of free electron metals, with simple bcc crystal structures. Under pressure, however, they have been observed to have series of complex crystal structures, associated with a drop in the melting point, reduced conductivity and even insulating phases <sup>16–19</sup>, including the discovered elemental electride (electrides have localised electrons at non-nuclear positions)<sup>17</sup>. This is related to a pressure-driven transformation from free-electron to electride behaviour<sup>20</sup>. This is an electronic change but, with two distinct types of bonding, it offers the possibility of a first-order transformation between two types of liquid. Alternatively, a continuous transition is possible, where the electride sites' proportion rises steadily from zero. The locations and signatures of any such transitions (if they exist) are not known. Electrides also exist at ambient pressure, often formed involving alkali metals in both crystalline and non-crystalline ionic salts <sup>21,22</sup>. Some exhibit exotic superconductivity or Mott insulator transitions <sup>23,24</sup>, and due to the localised electron have applications in catalysis<sup>25,26</sup>. Recent computational machinery has been developed for systematic prediction of stable electride structures <sup>27</sup>.

Although the electronic and structural properties of the crystalline alkali metals at different pressures have been studied in great detail, their liquid phases under high pressure are much less investigated 19,28,29 and not without controversy, e.g. on the liquid-liquid transition in Cs<sup>30–33</sup>. Experimental measurements of the macroscopic thermodynamic properties of the alkali metal melts have proved an insurmountable challenge due to their high reactivity. A few studies have suggested structural transformations affecting the short-range order, resembling those reported in solids 19,33. Ab initio molecular dynamics (AIMD) simulations were applied in high-pressure studies of structural features of liquid alkali metals, reporting tetrahedral clustering of nearest neighbours in liquid lithium at 150 GPa pressure 34, a possible metal-nonmetal transition in liquid sodium 35, a Widom

line crossing in rubidium<sup>36</sup>, and abrupt coordination changes in cesium<sup>37</sup>. AIMD simulations on liquid potassium reveal a drop in coordination number from 14 to 8-10 under pressure<sup>38</sup>, an effect also seen in other alkalis<sup>37</sup>. A major caveat in studying liquids with AIMD is the small system sizes: a liquid has some short-ranged structure, typically out to 3rd or 4th neighbour shells, and when looking for a structural phase transformation these must be correctly described, unaffected by periodic boundary conditions.

Herein, we present an atomistic simulation of liquid potassium using a machine learning potential approach based on AIMD across a wide range of pressures and temperatures. This method allows our simulations to extend well beyond the short-range order in the liquids, eliminating finite size effects (Supplementary Fig. 1). We detect a range of thermodynamic and dynamic anomalies in the region of pressures between potassium's melting curve maximum and the appearance of electride solids<sup>20</sup> (10-20 GPa), which are mostly independent of temperature. Going back to the AIMD simulations, we relate these anomalies to changes in the electronic structure that suggest a continuous transition from a free electron-like metal to an electride liquid.

Interatomic potential The potential for potassium used in this work was derived by fitting to *ab initio* configurations of a range of high pressure phases<sup>20,39,40</sup>. It has been shown to reproduce the entire high pressure phase diagram with remarkable fidelity (see Fig. 1), including both free electron and electride solids. Comparison of the properties calculated on small cells using AIMD and using the potential shows a good agreement (see Methods and Supplementary Note 1), which allows us to use it on larger cells and longer time simulations.

Theory of partial electride liquid Our central idea in understanding these results is that the liquid structure can be modeled a mixture of two distinct electronic states, the low-pressure "atomic" state and a high-pressure "electride" state where electrons are localized in interstitial pockets. Mixing models with different complexity<sup>41–43</sup> are discussed in the Supplementary Note 3. Different models give slightly different results, but all the mixing models display anomalies in heat capacity, thermal expansion and compressibility. Crucially, peaks associated with these anomalies do not fall in the same place. Specifically, the heat capacity maximum is at the lowest pressure, the compressibility maximum at the highest pressure, and the thermal expansion dip at an intermediate pressure. Curiously, this is the opposite ordering to that observed in the supercritical Lennard-Jones system<sup>44</sup>.

**Thermodynamic anomalies** We have calculated a wide range of thermodynamic and electronic properties for potassium as a function of pressure. In all of these, there is evidence of anomalous behaviour in the same pressure regime from 10-20 GPa, over a wide temperature range, see Fig. 1. We can interpret them in terms of a crossover from an atomic to an electride liquid (as outlined in the above), and corroborate this interpretation with analyses of the electronic properties (see further below).

Figure 2 shows the thermal expansivity  $\alpha$ , heat capacity  $C_p$ , sound speed  $v_s$ , and volume equation of state, all as a function of pressure. The expansivity  $\alpha$  shows a strong dip in the crossover region, because crossing the transition line allows for more (compact) electride states to be occupied. The normal thermal expansion is still large enough to ensure positive  $\alpha$ . Above 20 GPa the electride states occupation saturates, so this volume-reduction mechanism is no longer

available. The heat capacity  $C_p$ , while dominated by the Dulong-Petit 3R limit, has an anomalous peak as a function of pressure around 10 GPa. The electronic heat capacity exhibits a similar anomaly but with reduced magnitude (see Supplementary Note 7). For the sound speed  $v_s$ , the change in the compressibility (see Supplementary Note 3 for details) manifests here as a dip around 18 GPa. The two-state thermodynamic model (see Supplementary Note 3) was fitted to MLMD data at 900 K to show the smooth oscillation in  $v_s$  with pressure. The p(V) equation of state shows the strong compressibility at low pressures typical for the alkali metals, while the inset reveals no measurable density discontinuity anywhere, just a subtle inflection at around 14 GPa.

**Non-equilibrium property anomalies.** Transport properties are non-equilibrium phenomena and therefore not directly associated with the Ehrenfest phase transition definitions. Nevertheless, when there are different structures, one can expect differences in the non-equilibrium properties.

Figure 3a shows anomalous diffusion behaviour, with a distinct plateau in the ionic diffusivity across the 10-20 GPa transition region. We can understand this as follows: in the electride, electrons move to interstitial locations such that some potassiums can be thought of as positively charged cations. Cations are smaller than neutral atoms, and can therefore diffuse more easily. The increasing fraction of potassium-cations across the transition region leads to enhanced diffusion, largely cancelling out the normal decrease due to reduced free volume. The diffusivity at various temperatures can be collapsed onto a unique curve, suggesting that diffusion is thermally activated, but the liquid structure is not strongly affected by temperature. No hysteresis nor discontinuity in density was observed, see Fig. 3b, further suggesting that there is no first-order liquid liquid transition.

The diffusivity can be related to the viscosity  $\eta$  via the Einstein-Stokes relation  $D = \frac{k_{\rm B}T}{6\pi\eta r}$ , however that relation requires knowing the particle radius, r, which is uncertain, since the ions are converting from neutral to cations. However, the MLMD enables us to calculate the viscosity from the Green-Kubo relations and the fluctuations in the stress tensor, as shown in Fig. 3c. We then use this independent calculation of the viscosity to invert the Einstein-Stokes relation and obtain an effective particle size (Fig. 3d), which drops sharply in the transition region. This effective size should not be taken as the average size, because it is weighted by its contribution to the diffusion. A sharp drop occurs at about 10 GPa, suggesting that a small number of small ions can strongly influence the diffusion behaviour.

**Structural Properties.** The radial distribution function (RDF) is measurable by diffraction, as the Fourier transform of the liquid structure factor. Figures 4a and 4b show that below 12 GPa the RDF changes only by a scale factor. Above that pressure, the liquid has a very different structure. The first peak narrows and moves to a smaller radius - small even when the overall reduction in volume is factored out, which can be interpreted as the creation of smaller ions. In the normalised RDF, we see that the second neighbours are also much closer. These two effects drive the densification, despite the drop in near-neighbour coordination. Figure 4c and 4d clearly show the transformation in terms of the distributions of the coordination number.

Electronic Properties. Another potentially measurable quantity is the reflectivity. This cannot be obtained from the classical potential, but we can calculate it from the electronic structure of the *ab initio* data. In Fig. 5a we show the calculated reflectivity of liquid potassium at  $\lambda = 532$  nm and several temperatures, compared to the solid phases. The crossover in reflectivity across the liquid

transformation region is clear and follows the same trend as in the solid phases, though the dip is not as pronounced, and is virtually temperature-independent. A similar reduction of conductivity was observed in liquid lithium under pressure in experiments and simulations<sup>45,46</sup>.

We use Bader analysis<sup>47</sup> to determine the electride nature of the system (Fig. 5b) by decomposing the charge density. Specifically, we study the distribution of non-nuclear Bader charges,  $q_{NN}$ , which is a measure for the electride fraction x discussed above, and strength of the electride sites. The number of  $q_{NN}$ 's, the integral of the distribution, rises strongly to about x=0.3 electride sites/atom around 12 GPa. The peaks of the distributions rise most dramatically between 12 and 20 GPa where  $q_{NN}$ 's saturates around 0.4 e, close to the values of  $q_{NN}$ 's in phase-IV (oP8) at 40 GPa and the guest electride channels in phase-IIIa at 20 GPa. Counting charge within basins of the Electron Localization Function (ELF)<sup>48</sup> rather than charge basins for solid IIIa<sup>20</sup> found  $q_{NN}$  to be higher by a factor of 3, and this is also likely true for the liquid  $q_{NN}$  distribution.

Figures 5c-e use the ELF which provides a clearer picture of the pressure induced free electron-electride transitions. At 2 GPa, the ELF maxima are all nearly spherical, showing them to be associated with the potassium ions. In the transformation region there are also non-nuclear ELF maxima (non-circular red blobs), associated with the electride pseudo-anions. By 26 GPa, these non-nuclear maxima are even more prevalent. Figure 5f and Supplementary Movie 1 relates the Bader and ELF analyses, showing that the  $q_{NN}$ 's are associated with regions of localised electrons, i.e. high values of ELF.

**Discussion and conclusions** We have simulated the behaviour of high pressure liquid potassium, using molecular dynamics with interactions derived from both DFT calculations and a machine-

learned potential. The simulations show anomalous behaviours in a wide range of properties between 10 and 20 GPa. However, we do not see any hysteresis or density discontinuity. These observations led us to propose that liquid potassium exhibits a gradual increase in electride content. The electride/ionic potassium ions generally have higher energy than the atomic potassium, but on mixing can occupy smaller volume. This transformation is primarily driven by pressure, so the lines tracing the peaks of anomalous behaviours are nearly vertical on the pressure-temperature phase diagram, see Fig. 1.

This crossover behaviour can be modelled as a two-state system, in which the "electride" state has higher energy than the atomic state. The simple two-state models considered predict that anomalies in  $C_p$ ,  $\alpha$  and  $v_s$  will occur in that sequence with pressure - just as observed in simulations here. In these models, the excess heat capacity comes from the extra degree of freedom offered by "ionisation", the reduced thermal expansion is due to thermal excitation into the (denser) electride state, and the excess compressibility comes from the additional compression mechanism offered by creating electride.

From the anomalies in the sound velocity and thermal expansion, we obtain in the two-state thermodynamic model  $\Delta V/V=0.05$  and  $\Delta S=-0.2$  R between the atomic and the electride states, which gives a negative, almost vertical slope of -500 K/GPa for the boundary between the two unmixed states where  $G_A=G_B$ , in agreement with the vertical slope marking the observed anomalies (Fig. 1). Notice that a negative slope is required by the observation that the denser electride state is obtained by heating the liquid at constant pressure.

The diffusion plateau can be understood by smaller "ions" moving more quickly. The dis-

tinctive change in the local structure can be seen in the radial distribution and coordination. The conductivity/reflectivity shows a crossover with pressure that mirrors the transition from nearly free electron metal to electride phases in the solid.

The low pressure liquid is extremely compressible with no characteristic ionic radius, as the nearest neighbour distance scales with the volume. At high pressure the interionic spacing approaches twice the ionic radius of 1.3 Å. The coordination number drop (Fig. 4c) is consistent with the transition from close-packed solid ( $C_n = 12$ ) to the incommensurate electride phase of host-guest structure ( $C_n = 9$ ). The electride-containing phase is denser, despite the lower coordination, because the near neighbours are closer together and the second neighbour shell is much closer. The liquid becomes denser than the solid after the melting maxima around 10 GPa, implying that the dense liquid takes on properties of the phases succeeding fcc. Solid phases of host-guest structures and phase-IV (oP8) were observed as moderate electrides<sup>20,49</sup>.

In conclusion, we have shown that potassium becomes a type of two-component liquid in which atomic and ionic/electride potassium coexists. The phase diagram for potassium is similar to the other alkali metals, and we can expect that this type of two component liquid will be a general feature. Our work resolves the mystery of the negative Clapeyron slope between phase II (fcc) and liquid: the negative slope means that the liquid is denser than the solid. This cannot be due to more efficient packing, because fcc is the most efficient possible for spheres. Herein, the cause is revealed as the larger electride fraction in the liquid, where the ions are smaller than the atoms, and electrons can adopt any shape, to fit into the interstitial region where they are located.

#### Methods

**Ab initio Molecular Dynamics.** DFT calculations utilized the CASTEP code<sup>50</sup>. We used the generalized-gradient approximated exchange-correlation<sup>51</sup> and a 9-electron ultrasoft pseudopotential with 1.7 Bohr inner-core radius and 400 eV plane wave cutoff with k-points sampled at the  $\Gamma$ -point only in AIMD, and with a grid density of 0.02 Å<sup>-1</sup> in structure optimisation. Forces are calculated using the Hellmann-Feynman theorem.

Simulations ran at fixed density (NVT) with a Nosé-Hoover thermostat, using a 0.75 fs timestep, a Gaussian smearing of 0.1 eV, and up to 25 ps sampling. Additional simulations were performed at fixed pressure (NPT) and all observables of the electride transition are consistent. The initial conditions for the liquid state for the 650 K istotherm were attained by quenching a 1000 K melt of 128 BCC potassium ions. For the 1500 K isotherm, the simulations ranged from 108 to 250 potassium ions from melting the stable solid phases of bcc, fcc, and IIIa at various pressures.

Analyzing non-nuclear charge was performed using the Bader partitioning scheme<sup>47</sup> for the charge density. For a particular pressure 100 snapshots were selected for charge density and ELF calculations. Most Bader charges were found close to potassium ions (0.4 Å), while we deem non-nuclear charges,  $q_{NN}$ , as those that were found far away (2.0 Å) from potassium ions. These distances are consistent with the interstitial regions of the ion-ion g(r), and the ion- $q_{NN}$  g(r) shows clear peaks in the minimum regions of the ion-ion g(r). The majority of  $q_{NN}$ 's found with either Bader and ELF were found in similar positions, with some differences as expected, and may underestimate the total amount of non-nuclear localized electrons.

**Reflectivity.** The interband dielectric function  $\varepsilon(\omega)$  was determined within the random phase approximation using CASTEP and its OptaDOS extension<sup>52</sup>, with dense k-point grids for all solid phases and (8,8,8) grids for AIMD snapshots of 108 atoms. An intraband contribution  $\varepsilon^{intra}(\omega)$  was added using a Drude term with 100 THz broadening. Four snapshots per AIMD trajectory were sufficient to converge the spectral features of the dielectric function. The reflectivity was then determined using the usual expressions<sup>52</sup>.

Machine Learning the inter-atomic potential. We used a domain knowledge based machine learning approach to match the true potential energy surface of potassium<sup>39,40</sup>. Here, the machine learning interatomic potential (MLIP) is directly learned from an accurate reference database of first principle calculations<sup>53</sup> using VASP<sup>54</sup>. In order to capture the different phases in the temperature-pressure phase diagram of potassium, more than 20000 initial reference configurations were accumulated from the ab initio molecular dynamic (AIMD) simulations of different stable phases and their melts. The local chemical environment of these reference configurations were then fingerprinted by some atom-centered symmetry function descriptors (or fingerprints). These descriptors are mapped into the corresponding atomic energy/forces with the kernel ridge regression (KRR) method. The MLIP was further optimized by an iterative scheme, in which the potential is fitted to the existing reference database, and then simulations with the potential are run to create a more diverse set of structures. This way, the MLIP "steers" itself into regions of configuration space that need to be further explored. Additional information and benchmarks about the MLIP, can be found in Supplementary Note 1 as well as our recent work<sup>39</sup>.

The AIMD database includes Hellmann-Feynman forces which are calculated in VASP<sup>54</sup> as

the derivatives of the generalized free energy. Consequently the electronic entropy is included in both AIMD and MLMD forces; this is expected to be a small effect<sup>55,56</sup>. We used a fixed Gaussian smearing of 0.2 eV in all AIMD runs. In the Supplementary Note 8 we show that resulting forces deviate from those obtained with temperature-adjusted Fermi-Dirac smearing by only around 10 meV/Å. We note that this value is less than the uncertainty of the MLIP. However, electronic entropy can add a significant contribution to the heat capacity<sup>57</sup> which we calculate separately from the electronic DOS using the Sommerfeld approximation (see Supplementary Fig. 18).

Classic molecular dynamic simulations were then performed using a time step of 1 fs, and periodic boundary conditions were applied along all three dimensions. The Nosé-Hoover thermostat and the Parrinello-Rahman barostat<sup>58,59</sup> were used for controlling temperature and pressure, respectively. All simulations were carried out using the LAMMPS package and the atomic configurations were visualized with the AtomEye package. Typical models of potassium liquid containing 3456 atoms were obtained by heating a single crystal at 2000 K and specific pressures, and these liquid structures are used to study the corresponding properties of liquid (such as diffusivity, heat capacity and sound speed) at selected temperatures and pressures.

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**Author Contributions** H.Z., V.N.R., A.H. and G.J.A. conceived the research, H.Z., V.N.R., A.H. and L.Z. conducted the simulation; S.S. and G.J.A. created the two-state liquid model; All authors participated in the

analysis and interpretation of the results as well as the writing of the manuscript.

**Competing Interests** The authors declare no competing interests.

**Data availability** All The data represented in Figs. 1–5 are available as Source Data. All other data that

support the plots within this paper and other findings of this study are available from the corresponding

author upon reasonable request.

**Code availability** Code is available at https://github.com/zhaolongxjtu/KMLP.

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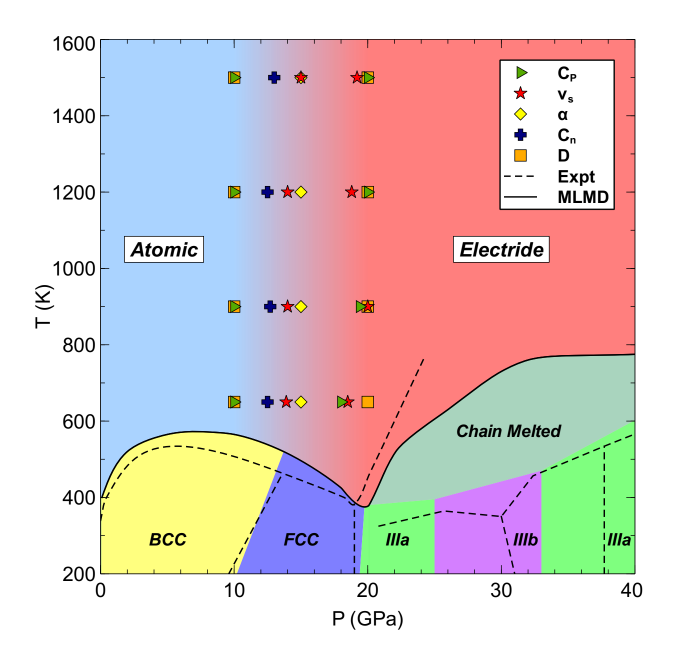


Figure 1 | Phase diagram of potassium from MLMD simulations. The colours denote stability regions of phases as labelled. Dashed lines are experimental phase boundaries (solid-solid, solid-chain melt and solid-liquid) taken from Ref. <sup>60</sup>. Points in the liquid region show maxima and minima in heat capacity  $C_p$  and sound speed  $v_s$ , minima in the thermal expansion  $\alpha$ , maxima in high-coordination environments ( $C_n > 15$ , see the Supplementary Figs 10-11), and boundaries of the plateau of diffusivity D.

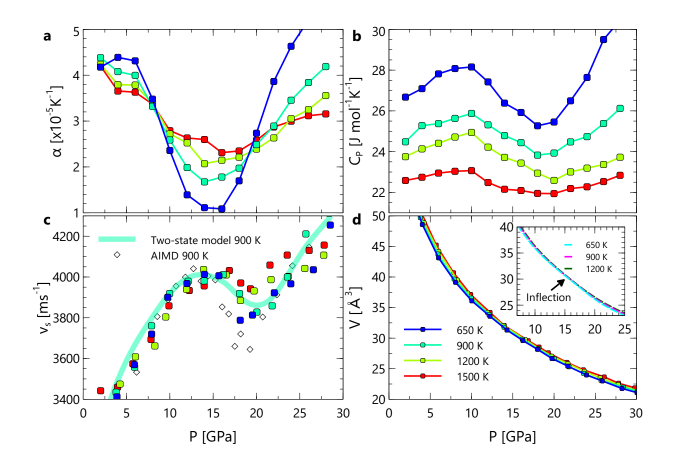


Figure 2 | Thermodynamic quantities of liquid potassium, shown as a function of pressure for selected temperatures. a, Thermal expansivity  $\alpha$ . b, Heat capacity  $C_p$  (excluding electronic contribution - see Fig S17). c, Sound speed  $v_s$  (open diamonds show AIMD at 900 K; cyan line shows the thermodynamic two-state model (see Supplementary Note 3) fitted to MLMD data at 900 K). d, Equation of state p(V) (inset shows AIMD results).

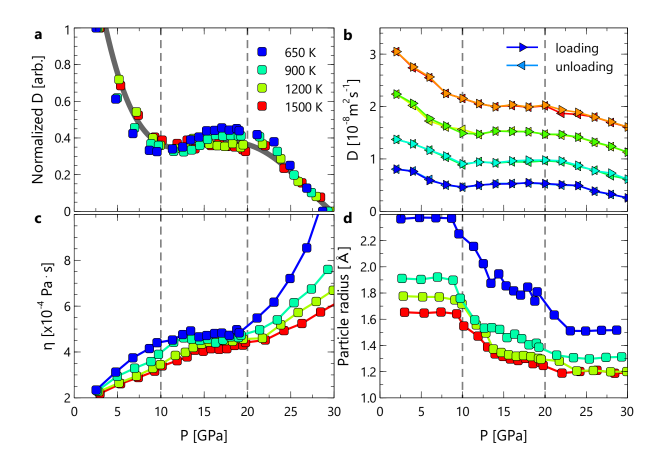


Figure 3 | Dynamical properties of liquid potassium from MLMD simulations. a, Normalised diffusion constant vs pressure for selected temperatures: line is a guide to the eye of the data collapse onto a single, T-independent curve once normalized as  $D_{norm} = \frac{D(P) - D_{min}}{D_{max} - D_{min}}$ . b, Absolute diffusion constants D(P,T) from a series of calculations showing an absence of hysteresis on increasing and decreasing pressure. c, Viscosity  $\mu$  calculated using the Green-Kubo stress fluctuation approach, d, Effective atomic radii R derived from  $\mu$  by inverting the Stokes-Einstein relation.

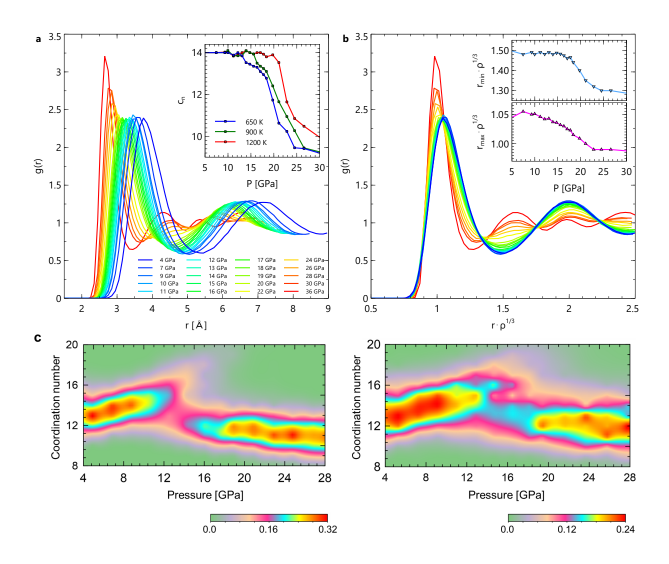
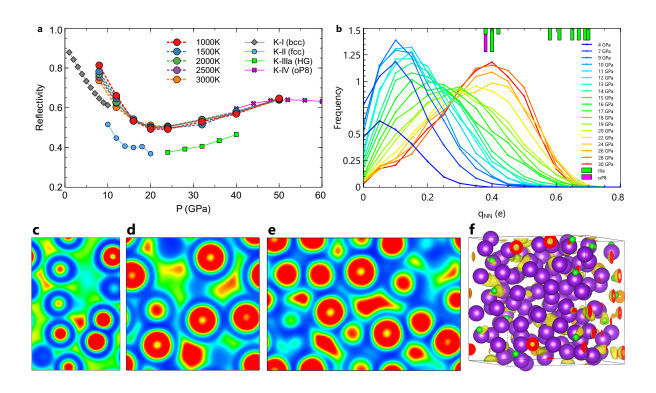


Figure 4 | Structural properties of liquid potassium from AIMD simulations. a, Radial distribution functions (RDFs) at selected pressures and 650 K, inset shows the average coordination number  $C_n$  defined as the cumulative distribution integrated to the first minimum  $r_{min}$ . b, RDFs with distances scaled by particle density  $\rho^{\frac{1}{3}}$ , insets show the positions of the first maximum and minimum with respect to pressure  $(r_{max} \text{ and } r_{min})$ . c, Heat map of distribution of coordination numbers  $C_n$  vs pressure, at 650 K (left) and 1200 K (right).



**Figure 5** | **Electronic properties of liquid potassium. a**, Reflectivity at 532 nm (green laser) based on DFT calculations of AIMD snapshots and optimized solid phases. **b**, Distributions of non-nuclear charges,  $q_{NN}$ , for liquid potassium as a function of pressure (lines) at 650 K, and  $q_{NN}$ 's for related solid electride phases are included above (bars): phase IIIa/oP8 at 20/40 GPa. **c-e**, Electron localization function (ELF) along typical (010) slices in simulation snapshots at 1200 K and (2, 16, 26) GPa. ELF values are shown in RGB scale from 0.0 (blue) to 0.7 (red), with 0.35 (green). **f**, AIMD snapshots of potassium (purple), sites of  $q_{NN}$ 's (green) and ELF = 0.7 isosurfaces (yellow).