Defect mediated manipulation of nanoclusters on an insulator

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With modern scanning probe microscopes, it is possible to manipulate surface structures even at the atomic level. However, manipulation of nanoscale objects such as clusters is often more relevant and also more challenging due to the complicated interactions between the surface, cluster and apparatus. We demonstrate the manipulation of nanometer scale gold clusters on the NaCl(001) surface with a non-contact atomic force microscope, and show that the movement of clusters is in certain cases constrained to specific crystallographic directions. First principles calculations explain this kinetic anisotropy as the result of the cluster attaching to surface defects: cation vacancies allow the clusters to bond in such a way that they only move in one direction. Constraining the movement of clusters could be exploited in the construction of nanostructures or nanomechanical devices, and the manipulation signatures may also be used for identifying cluster-defect complexes.

Controlling structures at the atomic level is the ultimate challenge in nanofabrication – absolute control over the positions of individual atoms, molecules, or clusters on substrates allows for nearly complete freedom in material composition or in the fabrication of nanodevices. Although arranging structures atom-by-atom is attractive from the perspective of fundamental interest in surface physics and chemistry, for many applications manipulation of larger units is more relevant. With respect to metallic clusters, interest stems from catalysis where clusters promote many catalytic reactions, but also from nanotechnology where clusters can be used as building blocks for nanoelectronic devices. In tribology, clusters can even serve as units for probing the friction at the nanoscale. In all these domains, manipulating clusters as single objects enables the systematic study of the impact of the specific and voluntarily chosen adsorption site on the physical, mechanical, electronic and chemical properties of a single cluster. While manipulation of atoms was achieved already in 1990 with scanning tunneling microscope (STM), atomic manipulation with non-contact atomic force microscopy (nc-AFM) has been a much more recent development. In particular, manipulation on insulating surfaces remains particularly challenging, yet insulators are a crucial class of materials in nanoelectronics and nanocatalysis.

Similar to STM, two methods are used to manipulate clusters by nc-AFM: the first method consists of moving the cluster by pushing during a spectroscopy step (Fig. 1 (a)). Here, the AFM tip is brought down vertically near the cluster until at some point the repulsive interaction between the tip and the cluster is strong enough to make the cluster move. In the second method the tip is scanned above the cluster (Fig. 1 (b)), preferentially in the constant height AFM mode. At a specific distance, as the tip approaches the cluster laterally, a force between both moves the cluster. For simplicity, we will call the former method kicking and the latter sliding.

This work investigates methods for directing the manipulation of a nanocluster beyond the constraint of the tip’s scanning direction. In particular we demonstrate the possibility of using intrinsic defects on the surface to mediate the diffusive behaviour of the clusters. We have chosen the NaCl(001) surface as a model insulator substrate firstly because it can be easily prepared and examined by nc-AFM even at the atomic scale, but also because the characteristic defects at its surface are well-established. Furthermore, NaCl and other alkali halide surfaces have been key in breakthrough nc-AFM studies in atomic and charge identification, as well as providing prototype substrates for the study of adsorbed molecules, self-assembly, friction, and local electronic structure.

Results

Manipulation on terraces. When single neutral gold atoms are deposited on perfect NaCl(001) surfaces, the gold atoms diffuse on the terraces due to their high mobility at room temperature until they reach a step-edge, where they aggregate into clusters. At room temperature and low gold coverage, the steps are decorated by gold...
clusters whereas on the flat terraces very few clusters can be found because of the lack of surface defects that would act as nucleation sites (see further discussion in the Supplementary information). The image in Fig. 2 (a) exemplifies a typical decoration of steps (along (100) equivalent directions) by gold clusters. The clusters have a mean diameter of 5 nm (~2500 atoms), which was estimated from the flux of incoming gold during the deposition.

In this first series of manipulation experiments, we focus on one of the rare clusters on the terrace (highlighted in green in Fig. 2) and consider its manipulation characteristics. The cluster was first imaged with a high magnification in the constant height mode (Fig. 2 (b)). In the figures, the slow scanning direction of the AFM tip is always from the bottom to the top. The same cluster was imaged several times while gradually bringing the tip closer to the surface. At a specific tip-surface distance the cluster suddenly disappeared during imaging (arrow in Fig. 2 (e)) by escaping behind the tip to the region that had just been imaged to the position shown in Fig. 2 (c). This kind of "backheel" led to a 16 nm long movement of the cluster along the [110] direction. In a second manipulation step the same cluster was imaged again and the tip-surface distance was further reduced. Now, the cluster could be moved by sliding in front of the tip along the [110] direction (Fig. 2 (f)). Because the cluster is moved in front of the tip from line to line, it appears with an elongated contrast, reflecting its movement. The new position of the cluster, which is 11 nm further away from the old one, can be seen in Fig. 2 (d).

**Manipulation from step-edges.** In Fig. 3, similar clusters of about 2000 atoms decorate intersecting [110] and [100] steps on the NaCl(001) surface. One of the clusters is highlighted (I in Fig. 3 (a), green color) at the (110) step-edge. After recording a series of high resolution images at this site, the last image of this series shows that the cluster was moved 12 ± 1 nm from the step (to position II in Fig. 3 (b)). We continued to move the same cluster, now on the terrace, by sliding about 15 ± 1 nm along the [110] direction (Fig. 3 (c)), which has been verified in a following image with a larger scale (Fig. 3 (d)). We could move the cluster also by kicking with a vertically approaching tip in a spectroscopy step. Figure 3 (e) shows that in this case the cluster could also be moved by 16 ± 1 nm from its initial position. Altogether, the cluster was moved by the tip dozens of times during the whole manipulation process. Regardless of the method of manipulation, the events always resulted in cluster movement along equivalent (110) directions on the surface (see path of cluster in Fig. 3 (f)). This extremely anisotropic movement is in clear contrast to the cluster found on the terrace, which could easily be moved in different directions.

**Calculated manipulation kinetics.** In order to understand the movement of clusters during manipulation and pinpoint the origin of the experimentally observed differences, we sample the possible pathways for moving 20 and 100 atom gold clusters (Fig. 4 (a)) using density functional calculations. With the smaller Au20 cluster we perform extensive analysis on the adsorption configurations and diffusion pathways while the larger Au100 cluster is used in selected cases to assess scaling with respect to cluster size.

The Au20 cluster is found to adsorb on the perfect surface (ad sorption energy 0.5 eV) and the energy barriers for moving the cluster in any given direction are very low, about 0.2 eV (Fig. 4 (b)). At room temperature, such a diffusion barrier corresponds to a Boltzmann factor of ~10^-4, and thus such a small cluster should move spontaneously due to thermal energy. Furthermore, the barriers for moving the cluster in either (100) or (110) directions are practically the same and thus no direction should be preferred.

The calculated high mobility on the ideal terrace strongly implies the presence of a defect under the cluster in Fig. 2 — else the cluster would have diffused to a step during growth. Hence, we also consider the role of defects in cluster adsorption and diffusion. The calculations show that in the presence of surface vacancies, or F-centres, the cluster adsors even more strongly than on the perfect surface, as the adsorption energies are 2.1, 2.5, and 1.1 eV for adsorption at neutral Cl, Na and double vacancies, respectively. (Defect charge state influence is discussed in the Supplementary information.) Also the diffusion energy barriers are much higher when defects are present, as discussed below, meaning it is much more difficult to move the cluster, and this verifies that defects can act as anchors for the clusters.

In the presence of a Cl vacancy, the barriers for moving the cluster are 0.8 – 1.1 eV and the anisotropy is relatively weak (Fig. 4 (c)). A cluster bonded to a Na vacancy, on the other hand, prefers a [110] direction where the barrier is calculated to be 0.7 eV. In comparison, the (100) barriers are 0.9 eV, and in the perpendicular [110] direction the barrier is 1.2 eV (Fig. 4 (d)). Similar calculations with a 100 atom gold cluster show that this is still the case even when the clusters grow. Figure 4 (e) shows the calculated barriers for the Au100 + Na vacancy complex. When the vacancy is located near the cluster edge, in the energetically favored position, the anisotropy is even stronger than for the small cluster (barriers 0.5 – 1.1 eV).
Discussion
The charged Na⁺ and Cl⁻ vacancies (latter are F⁻-centres) are expected to be mobile on the NaCl(001) surface and diffuse to steps during annealing. Recent nc-AFM and Kelvin Probe Force Microscopy (KPFM) studies have further shown that Na⁺ vacancies are the dominant defect at step-edges and kinks, which is due to the Debye-Frenkel layer formation on alkali halide (001) surfaces. This suggests that a cluster taken from a step, as shown in Fig. 3, is with a high probability anchored by a Na⁺ vacancy. The experimentally observed kinetic anisotropy of these clusters can then be explained as the characteristic behaviour of this particular cluster-defect complex, as revealed by the simulations. The third type of possible defect on NaCl(001) is the Cl⁻-center (Cl vacancy + electron, i.e., a Cl⁺ vacancy), and these defects are expected to be stable on the terrace (calculated diffusion barrier for the defect is 0.9 eV). Therefore, they are also the most probable defect on which clusters found on terraces are adsorbed (as in Fig. 2). From this consideration, the theoretically predicted small diffusion anisotropy for a cluster adsorbed on a Cl⁻ vacancy agrees with the experimentally found isotropic movement of the cluster on the terrace. The reason why the defects influence the movement of clusters differently can be understood by examining the calculated structures of the bonded cluster-vacancy pairs. In Fig. 5 (a) and (b), the differential charge densities are shown for clusters in the presence of Cl and Na vacancies, respectively. This density represents the change in electronic charge upon bringing the cluster and the surface together. In the presence of a Cl⁻ vacancy, the Au cluster bonds almost exclusively at one of its corners, attaching to a Cl⁻ ion next to the vacancy. In fact, the cluster tilts 10° as the bonded corner is attracted to the Cl⁻ ion. Because of this, the cluster prefers to move by pivoting around the bonded corner, as schematically shown in Fig. 5 (c) (although the preference is not strong, as shown by the almost isotropic barriers in Fig. 4 (c)). Simultaneously, the Cl⁻ vacancy jumps to a neighboring site, allowing the cluster to bond to another corner. Combining these rotations leads to movement in the [100] direction, but since the cluster is constantly reorienting itself, on average none of the [100] surface directions is preferred and the cluster may in fact move in any direction by taking a zig-zag path.

On the other hand, if the cluster attaches to a Na vacancy, it does not bond at a corner, but along an edge which orients itself with an entire row of Cl ions on the surface. This is seen in Fig. 5 (b). Again, the cluster is seen to slightly tilt (5.1°) towards the active edge to facilitate the bonding. For such a configuration, the energetically favored mode of movement of the cluster is sliding along the row of Cl ions with which it has bonded, so that the active edge always stays on the Cl row. This is represented in Fig. 5 (d). Since this movement follows a single row of Cl ions, the cluster moves in a [110] direction. Notably, since the cluster cannot move perpendicularly to the Cl row without detaching or rotating first, a much more difficult task than sliding in the preferred direction, this configuration results in truly anisotropic movement. Exactly the same kind of attaching to a Cl row occurs with a larger cluster as shown in Fig. 5 (f), demonstrating that this behaviour does not depend on the cluster size.

There is a discrepancy between the cluster sizes in the presented theory and experiments (100 and 1000 atom clusters, respectively). Although such a difference in the number of atoms only means a linear size difference by a factor of two, difference in size at this scale may lead to changes in the cluster morphology. On ionic surfaces like NaCl(001) or MgO(001), gold clusters expose either their (001) or (010) facets to form rectangular or triangular shapes, respectively, and the shape may depend on the size and growth conditions. However, as our simulations predict the anisotropic movement for the cluster–vacancy complex to be due to bonding of low-coordinated gold atoms at the cluster edge with Cl ions on the substrate near a Na vacancy, the anisotropy should not depend on the size and exact shape of the cluster. Instead, the behaviour is a characteristic feature of this kind of cluster–defect pair. We therefore expect the same characteristics also for the larger clusters, as seen in the experiments.

To summarize, we experimentally demonstrate nanomanipulation of gold clusters roughly of the size of 2000 atoms (5 nm diameter) as single units on the terraces of the insulating NaCl(001) surface using nc-AFM. From first principles calculations we show that stable clusters on the terrace are expected to be adsorbed at defects, which always accompany a cluster during its movement on the surface. Since the NaCl(001) surface only has vacancies as potential defect candidates at the surface, the manipulation properties are dominated by the type of vacancy: clusters on cation vacancies exhibit an anisotropic movement along the [110] surface directions whereas clusters on anion vacancies do not. Especially with respect to the cation vacancies, which are known to be located at the step edges, theory perfectly agrees with experiments since clusters that are detached from steps by the tip move indeed along ⟨110⟩ surface directions.

In general, this work demonstrates both experimentally and theoretically that even on smooth surfaces cluster-defect pairs can have extremely anisotropic movement patterns, and that this anisotropy can be strong enough to be the dominant factor controlling their manipulation. Such constrained movement can be exploited when building structures through nanomanipulation or even in the design.
of nanomechanical machines. Also conversely, observing specific anisotropy patterns could be used for distinguishing the types of defects on which clusters are adsorbed. Of course, this is not necessarily a one-to-one mapping: in a system where different kinds of defects exist, it is possible that different cluster-defect complexes have very similar manipulation patterns. Still, examining the cluster movement may allow one to distinguish the most likely defect types, especially if the intrinsic occurrence probabilities of defects are known.

Aside from providing an important characterization of defects, the ability to control and create defect-cluster complexes offers great potential for future studies in catalysis where, for instance, the cluster reactivity must be studied as a function of adsorption site. Such manipulation experiments can be combined with a control over the defects, which can be created by e.g. the AFM tip followed by imaging and spectroscopy identification, or by doping or irradiation of the substrate material such that known defects appear on the surface. When manipulation can be used for an overall characterization of the defect-cluster system, it will be a powerful tool for linking observed properties with the atomic and electronic structure of the defect-cluster complex.

**Methods**

**Sample preparation.** The undoped NaCl crystals are grown in an argon atmosphere by the Czochralski growth method in our laboratory at the CINaM. Before surface preparation, crystals and sample holders are cleaned by annealing (~250°C) in an oven located inside the main UHV chamber. After cooling, the (001) surfaces are prepared by cleavage of the crystals along the (001) plane at room temperature in the UHV chamber. The crystals are then annealed a second time in the same oven at ~150°C for a few hours to remove surface charges. The preparation of a clean NaCl(001) surface by cleavage and following annealing, the deposition of gold onto such a NaCl(001) surface, and following AFM experiments are done in one UHV chamber, which is equipped with a sample transfer system.
Gold coverages were estimated by measuring the flux of a beam of neutral gold atoms from a Knudsen cell using a quartz crystal microbalance. The pressure during the deposition is in the lower 10⁻¹¹ mbar range. The nc-ADF images were not used for coverage estimation due to the strong tip-cluster convolution, which is of importance when the clusters have a size of only a few nanometers. Especially for the small clusters studied in this work, their size (diameter ≈ 5 nm) is smaller than the tip apex, the diameter of which is typically 20 nm for a truncated pyramid, respectively, as shown in Fig. 4 (a). Much larger structures are not feasible at this level of accuracy. Gold clusters prefer to grow in (100) epitaxy but also in (111) epitaxy with a triangular or hexagonal shape. For consistency, we focus on small Au₃₀ and Au₄₄ clusters in (111) epitaxy in the calculations. For Au₄₄ our calculations show the (111) faceted truncated tetrahedron structure to be energetically preferred to the best (100) epitaxial geometry we could find by 2.5 eV.

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Acknowledgements
We acknowledge financial support through the FANAS project NOMCIS. CB and GC 
highly acknowledge stimulating discussions with C. R. Henry. TH and ASF acknowledge 
support by the Academy of Finland through its Centres of Excellence Program (project no. 
251748), as well as the computational resources offered by CSC, Finland.

Author contributions
All authors contributed to writing the article. CB and GC designed and performed 
experiments and analysis. TH and ASF designed the computational study. TH performed 
the calculations and analysis.

Additional information
Supplementary information accompanies this paper at http://www.nature.com/ 
scientificreports
Competing financial interests: The authors declare no competing financial interests. 
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How to cite this article: Hynninen, T., Cabalh, G., Foster, A.S. & Barth, C. Defect mediated 
motion of nanoclusters on an insulator. Sci. Rep. 3, 1270; DOI:10.1038/srep01270 
(2013).