Surface Science 603 (2009) 1145-1149

Contents lists available at ScienceDirect

Surface Science

journal homepage: www.elsevier.com/locate/susc



Realization of an atomic sieve: Silica on Mo(112)

Stefan Ulrich^a, Niklas Nilius^{a,*}, Hans-Joachim Freund^a, Umberto Martinez^b, Livia Giordano^b, Gianfranco Pacchioni^b

^a Fritz-Haber-Institut der MPG, Chemical Physics, Faradayweg 4-6, 14195 Berlin, Germany
^b Dipartimento di Scienza dei Materiali, Università di Milano-Bicocca, Via R. Cozzi 53 20125, Milano, Italy

ARTICLE INFO

Article history: Received 15 January 2009 Accepted for publication 26 February 2009 Available online 6 March 2009

Keywords: Adsorption Oxide surfaces Scanning tunneling microscopy Density functional theory

ABSTRACT

The adsorption of Pd, Ag and Au atoms on a porous silica film on Mo(112) is investigated by scanning tunneling microscopy and density functional theory. While Pd atoms are able to penetrate the holes in the silica top-layer with virtually no barrier, Ag atoms experience an intermediate barrier value and Au atoms are completely unable to pass the oxide surface. The penetration probability does not correlate with the effective size of the atoms, but depends on their electronic structure. Whereas Pd with an unoccupied valence s-orbital has a low penetration barrier, Ag and Au atoms with occupied s-states experience a substantial repulsion with the filled oxide states, leading to a higher barrier for penetration. In the case of Ag, the barrier height can be temporally lowered by promoting the Ag 5s-electron into the support. The Mo-supported silica film can thus be considered as a primitive form of an atomic sieve whose selectivity is controlled by the electronic structure of the adatoms.

© 2009 Elsevier B.V. All rights reserved.

1. Introduction

Zeolites and silicates have immense technological importance due to their unique structural properties, for instance as high-area catalysts, ion-exchangers and molecular sieves [1–3]. Such materials obey a universal building principle; an open –Al–Si–O– network encloses an array of nano-pores with rather homogenous diameter. These pores can accommodate single ions, molecules or atomic clusters that might conduct chemical reactions within the threedimensional (3D) matrix. The efficiency of such processes depends on the geometrical size of the species, which has to match the average pore size of the material. As pore diameters in zeolites and silicates are adjustable between 5 and 100 Å, size-selective adsorption and reaction schemes can be realized.

Recently, a 2D counterpart for 3D porous materials has been developed [4,5]. The system consists of an ultra-thin silicon-dioxide film that is prepared on a Mo(112) surface [6–8]. The fundamental building blocks of the oxide layer are six-membered – Si–O– rings, enclosing a hole of 3–4 Å diameter. These openings provide access to nano-pores located at the Mo–silica interface, which offer an attractive binding environment for metal atoms and small molecules [9,10]. The oxide surface, on the other hand, is chemically inert due to the high degree of bond saturation in the topmost silica layer [11]. The adsorption properties of SiO₂/ Mo(112) therefore depend on the ability of adsorbates to pass the openings in the oxide layer and to reach high-binding sites at

* Corresponding author. *E-mail address:* nilius@fhi-berlin.mpg.de (N. Nilius). the interface. So far, only the penetration of Pd atoms through the silica top-layer could be verified experimentally [10], although a similar behavior is predicted for Li and Na [12]. In contrast, Au atoms experience a substantial barrier for penetration and are in fact unable to bind to the defect-free oxide surface [9,10]. The relevant parameter that governs the energy barrier for passing the holes in the oxide film has not been identified yet for the different atoms. In a first, intuitive picture, it might be connected to the atom size with respect to the pore diameter. In this paper, we will show that this approach is too simple and metal atoms of comparable size have different penetration probabilities.

For this purpose, the binding behavior of Pd, Ag and Au atoms onto $SiO_2/Mo(112)$ is investigated by scanning tunneling microscopy (STM) and compared to results of density functional theory (DFT) calculations. The three species are chosen, because they either exhibit comparable van der Waals radii (Pd and Au) or similar electronic properties (Au and Ag). The derived binding characteristic demonstrates that the probability for an adatom to pass the holes in the silica top-layer is governed by its electronic structures, in particular by the spatial extension and electron filling of its valence orbitals.

2. Experiment and theory

The experiments are carried out in a custom-build, ultra-high vacuum STM operated at 10 K. Imaging is performed in the constant current mode, whereas the sample electronic properties are deduced from differential conductance (dI/dV) measurements applying lock-in technique. According to the Tersoff–Hamann

^{0039-6028/\$ -} see front matter @ 2009 Elsevier B.V. All rights reserved. doi:10.1016/j.susc.2009.02.030

theory for vacuum tunneling [13], the dI/dV signal provides a measure for the local density of electronic states (LDOS) in the sample.

The Mo support is cleaned by alternating cycles of Ar⁺ sputtering and annealing to 1800 K and afterwards exposed to 50 L O₂ to produce an oxygen-induced $p(2 \times 3)$ precursor structure [14]. Deposition of 1.2 ML Si in 1×10^{-7} mbar O₂ at 800 K and subsequent heating to 1200 K leads to the formation of a crystalline silica layer characterized by a sharp $c(2 \times 2)$ LEED pattern [4,8]. STM topographic images clearly reveal the porous structure of the silica film (Fig. 1A). Each hole is surrounded by a hexagonal –Si–O– ring of 5.5 Å diameter that is anchored to the Mo support via oxygen atoms below each Si (see model in Fig. 2B). The hole opens into a nano-pore located at the Mo-SiO₂ interface that offers a bridge and a top Mo site for binding adsorbates. The main structural defect in the oxide film are $[\overline{1} 1 0]$ -oriented antiphase-domain boundaries (APDB) that form between two domains shifted by a $[\overline{11}1]$ oriented Mo unit cell vector (arrows in Fig. 1). To bridge the displacement vector, the -Si-O- hexagons are replaced by four and eight-membered rings alternating along the APDB. The central hole in the octagonal rings is hereby two times larger than in the regular hexagons [8]. Single Pd, Ag and Au atoms are deposited from three different mini-evaporators consisting of the respective high-purity wires wrapped around a tungsten filament. Although atom deposition has been performed at 20 K sample temperature, the thermal energy of the incoming atoms ensures transient diffusion into their equilibrium adsorption sites.

Spin-polarized DFT calculations are performed in the VASP code, using the generalized gradient approximation, the PW91 exchange–correlation functional and a plane wave basis set (kinetic energy cut-off 400 eV) [15,16]. The electron–ion interactions are introduced by the projector augmented wave method [17]. The regular sites in the silica film are modeled with a (4×2) super-cell containing 7 Mo layers ($Mo_{56}Si_8O_{20}$), while a (5×2) cell with 4 Mo layers is used for the APDB ($Mo_{40}Si_{10}O_{25}$). The slabs are separated by 10 Å of vacuum. Penetration barriers are determined by moving the metal atom along the surface normal into the oxide pore and relaxing the metal/oxide complex for each vertical distance. The computed barriers do not correspond to a real transition state



Fig. 1. STM topographic images of (A) Pd ($U_s = 0.5 \text{ V}$), (B) Ag ($U_s = 1.5 \text{ V}$) and (C) Au ($U_s = 1.0 \text{ V}$) on a thin silica film grown on Mo(112) ($20 \times 20 \text{ nm}^2$). The nominal coverage was set to 5×10^{12} atoms per cm² in all cases. The arrows mark domain boundaries in the film. The Ag atom denoted by the square in (B) is bound to such a boundary.



Fig. 2. (A) Experimental ($5 \times 5 \text{ nm}^2$) and (B) simulated STM images (left and center: $1.8 \times 1.6 \text{ nm}^2$ and right: $2.1 \times 1.4 \text{ nm}^2$) of a Pd atom (imaging bias: 0.5 V), an Ag (0.3 V) and an Au atom (0.8 V) on SiO₂/Mo(112). While Pd and Ag atoms are incorporated into regular six-membered rings, the Au is bound to a Si–O octagon located at a domain boundary. The ball model in (B) shows a single –Si–O– hexagon (Si: large blue, O: small red spheres). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

and slightly overestimate the actual barrier height. Topographic STM images are simulated using the Tersoff–Hamann approach [13].

3. Results and discussion

Fig. 1 shows STM topographic images of the silica film after deposition of single Pd, Ag and Au atoms (from left to right) for a nominal coverage of 5×10^{12} atoms per cm². The three kinds of ad-species appear with rather different shapes and spatial distributions on the surface, although similar preparation and imaging conditions have been applied in all cases. The Pd-related features are imaged with star-like shapes in a bias window between 0.5 and 1.5 V. According to earlier studies, this distinct contrast provides evidence for a penetration of Pd atoms through the silica holes and their attachment to a Mo bridge site at the interface [10]. In this binding geometry, the Pd 5s orbital hybridizes with the 2p states of the six O atoms in the hosting -Si-O- ring, thereby increasing the unoccupied LDOS at the oxygen positions. The availability of such Pd-induced states for tunneling is responsible for the star-like appearance of embedded Pd species in the STM images (Fig. 2). Only at higher imaging bias (above 2.0 V), the Pd 5s orbital becomes directly accessible for electrons from the tip and the contrast localizes at the Pd position in the ring center (not shown) [10]. The embedded Pd atoms are randomly distributed on the silica surface and show no preference for binding to APDB (Fig. 1A). From analyzing several STM images, an equal probability is deduced to find Pd in regular -Si-O- hexagons and octagonal rings along the APDB, indicating a similar penetration barrier in both cases.

After Ag deposition, the same hexangular stars appear on the silica surface as for Pd, although in a slightly different bias regime (Fig. 2). The features are visible at small negative and positive voltages, but transform into round protrusions already above 1.5 V (Fig. 1B). The presence of hexangular stars provides a first indication that also Ag atoms are able to penetrate the holes in the – Si–O– hexagons. In contrast to the Pd species, Ag shows a certain affinity to interact with the APDB. In Fig. 1B, one out of 18 atoms occupy a –Si–O– octagon, as marked by the square. The overall fraction of Ag atoms bound to APDB is determined to 5%, which compares to 1–2% of all available sites being located along the domain boundaries. Consequently, the probability to find an Ag in the –Si–O– octagon is 3–5 times higher than in a hexagon, suggesting a slightly lower penetration barrier at the APDB.

A completely different behavior is revealed for the Au atoms. No adsorbates are found on defect-free silica patches after gold deposition, suggesting that Au atoms are unable to pass the holes in the -Si-O- hexagons and do not bind atop the surface either. Apparently, the penetration barrier for regular -Si-O- rings is too large to be overcome by the thermal Au atoms, which arrive with \sim 90 meV according to the gold evaporation temperature. Instead, small Au aggregates line up along the APDB, indicating that the adatoms are able to interact with the eight-membered rings. Fig. 2 shows a close-up STM image of a single Au atom that has penetrated such an octagonal -Si-O- ring. It is identified by two bright spots emerging at the long sides of the ring opening at low imaging bias. As deduced from DFT calculations, this distinct contrast manifests a structural distortion of the -Si-O- octagon to allocate more space for the Au atom [10]. A single atom inserted in an eight-membered ring already forms the critical nucleus for Au aggregation and initiates the growth of small particles, as seen in Fig. 1C [18]. In summary, Pd, Ag and Au atoms interact with the silica surface in different ways. While a similar penetration probability of regular -Si-O- hexagons and octagons is found for Pd, the Ag shows a 3-5 times larger probability to pass the eight-membered rings at APDB. In contrast, Au is completely unable to penetrate the regular silica holes and exclusively binds to boundary sites of the surface.

In the following, the experimental adsorption behavior of the three atomic species is compared to DFT calculations. Pd atoms are found to pass the regular -Si-O- hexagons with virtually no barrier (below 50 meV) and bind with 3.3 eV to interfacial Mo bridge sites (Fig. 3). According to a Bader analysis, the Pd remains neutral during penetration (4d¹⁰5s⁰ configuration), but becomes partly negatively charged (-0.5 e) in its final binding configuration. The penetration and binding energies are similar for the six- and eight-membered rings, explaining the equal distribution of Pd atoms on regular and defect sites of the silica surface. The most pronounced change in the oxide electronic structure upon Pd incorporation is a hybridization between the Pd 5s and O 2p orbitals of the surface oxygen atoms. The resulting O2p-Pd5s hybrid state is calculated to be at 2.0 eV above the Fermi level and shows up also in the experimental dI/dV spectra of Pd atoms as a shoulder at 2.3 V (Fig. 4). The good agreement between theory and experiment in terms of imaging contrast (Fig. 2) and spectroscopic response (Fig. 4) verifies the incorporation mechanism of Pd atoms into the silica film.

The Ag atoms experience a much higher barrier of 0.3 eV for penetrating the six-membered -Si-O- rings (Fig. 3). Assuming an Arrhenius behavior, thermal Ag atoms have a 30 times lower penetration probability than Pd atoms, which is however still high enough to yield measurable penetration rates.¹ The Ag atoms are able to overcome this barrier only at the moment of impact. After thermalization, the atoms are trapped in a weak physisorption potential of roughly 0.1 eV and diffuse to adjacent APDB, where penetration of the octagonal rings remains possible. This diffusive transport might be responsible for the higher adsorption probability of Ag atoms at APDB with respect to regular oxide sites. After passing the nanohole, Ag binds to the Mo-SiO₂ interface with 1.6 eV, remaining in its neutral 4d¹⁰5s¹ configuration as above the surface (Fig. 3). The binding energy is substantially lower than for Pd due to the chemically inactive nature of the low-lying Ag 4d states. Similar to the Pd case, the largest change in the oxide electronic structure is the formation of an O2p-Ag5s hybrid state that peaks at +1.6 eV and exhibits a long tail crossing the Fermi level (Fig. 4A). Tunneling into this tail produces the typical star-like images of inserted Ag atoms, reflecting from the enhanced LDOS of the six O atoms of the hosting ring (Fig. 2). Above 1.5 V, the O2p-Ag5s hybrid state becomes directly available for electron transport and the imaging contrast localizes at the Ag atoms (Fig. 1B). Also the resonance state of inserted Ag shows up experimentally as a dI/dV peak at 2.05 V, supporting the proposed binding mechanism.

The calculated penetration barrier of the -Si-O- hexagons increases to 0.9 eV for Au atoms (Fig. 3). The probability to reach a high-binding site at the interface is therefore four orders of magnitude lower than for Pd, explaining the absence of Au species on defect-free silica patches. The penetration of regular SiO₂ rings is, however, not only kinetically hindered. Even at higher temperature, desorption and not penetration of Au atoms would be the dominant process, as the Au binding energy to the silica surface is with 0.1 eV considerably lower than the penetration barrier. Gold is able to pass the oxide surface exclusively at the APDB, where the pore sizes are larger and high-binding interfacial sites are accessible without substantial barrier. Whereas Au atoms are neutral above the silica film ($5d^{10}6s^1$ configuration), they become partly negatively charged (-0.4 e) at the interface due to an electron transfer from the Mo (Fig. 3). Also from an electronic point

¹ Quantitative values for the penetration rate cannot be given as the pre-factor for the Arrhenius process is unknown.



Fig. 3. Energy profiles for the insertion of Pd, Ag and Au atoms into regular six-membered rings of the silica film. Left and right values in each plot denote the atom binding energy on top and inside the film, respectively, while the central value gives the energy barrier for penetration. All energies are given with respect to a gas-phase atom. The Bader charges during penetration are reported in parenthesis. The binding configurations of the different atoms above and below the silica film are shown in the insets (Mo: large gray, Si: medium blue and O: small red spheres). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)



Fig. 4. (A) Total LDOS of the pristine silica film (gray) and plots of the hybrid states formed between s-orbitals of incorporated Pd, Ag and Au atoms and the O 2p states of the surface oxygen atoms. (B) Experimental dI/dV spectra of the pristine oxide film and inserted Pd and Ag atoms, showing similar resonance states as in (A).

of view, Au atoms bound to APDB behave differently than their Pd and Ag counterparts. The Au 6s orbital strongly couples to the Mo states, while hybridization with the oxide LDOS is negligible due to a large energy mismatch (Fig. 4A). In contrast to the Pd and Ag-induced states that have substantial density probability even above the oxide surface, the Au 6s-derived states are localized at the Mo-silica interface and therefore inaccessible for *dI/dV* spectroscopy (Fig. 4B). In fact, the imaging contrast of Au atoms bound to APDB results mostly from a structural distortion of the silica lattice upon Au incorporation [10].

The origin for the dramatically different penetration barriers for Pd, Ag and Au atoms shall be discussed in the last part of the paper. In a first intuitive picture, the barrier height might be connected to the size of the respective gas-phase atoms, as given by their van der Waals radius [19]. Following this model, smaller atoms should exhibit a reduced barrier with respect to larger ones, a trend that is indeed observed for Pd (van der Waals radius of 163 pm) versus Ag (172 pm). However, the penetration barrier for Au is substantially higher, although its van der Waals radius of 165 pm is relatively small due to the relativistic contraction of Au orbitals. The atomic radius seems therefore to be an inappropriate parameter to rationalize the observed trend in barrier height. In a more sophisticated model, the penetration barrier might be linked to the repulsion that is exerted by the oxide charge density, especially the occupied O 2p states, on the incoming atom. This interaction will be dominated by the spatially most expanded adatom orbitals, viz the s-like valence states. Due to its unfilled nature, the Pd 5s orbital produces only a small repulsion and the resulting penetration barrier is low. In contrast, the half-filled Ag 5s and Au 6s orbitals will strongly interact with the surface O2p states during penetration. The surprising fact that Ag has a three times lower energy barrier than Au is explained by a transient positive charging of Ag atoms when passing the ring plane. The Ag atom transfers charge density from its 5s orbital into the Mo states, thereby reducing the electron–electron interaction with the oxide states. This effect is clearly revealed from the computed Bader charge of +0.6 e for an Ag atom in the ring plane (Fig. 3).

A similar mechanism is not accessible for gold. An Au atom is a strongly electro-negative species and rather accumulates excess electrons in its 6s orbital instead of transferring charge density to adjacent atoms [20,21]. Consequently, the Au remains neutral when passing the hole in the silica top-layer (Fig. 3). Also an internal charge redistribution, e.g. by promoting the 6s electron into the 'leaner' $6p_z$ state having its electron density localized along the penetration axis, is inhibited by the high activation energy of 4.6 eV [22]. The Au atoms are therefore unable to lower the repulsive interaction with the oxide surface electrons and the penetration barrier remains high.

4. Conclusion

The nano-porous SiO_2 film grown on Mo(112) exhibits essential properties of an atomic sieve. Single metal adatoms can either penetrate the openings in the top-most silica layer, followed by a strong binding at the metal-oxide interface, or are repelled at the inert oxide surface. As demonstrated by the combined STM/ DFT study, the oxide penetration barrier varies considerably for different atomic species. Whereas Pd atoms experience virtually no barrier, the activation energy for penetration steeply increases when going to Ag and Au atoms. The actual barrier height is connected to a repulsive interaction between the valence electrons of the incoming atom and the electron density at the oxide surface, and therefore a purely electronic effect. The identification of this interaction mechanism enables a reliable prediction of penetration barriers for other atomic and molecular species, being a first step towards the construction of a true atomic sieve based on the SiO₂/Mo system.

Acknowledgments

The work has been supported by the COST Action D41. Part of the computing time was provided by the Barcelona Supercomputing Center (BSC-CNS).

References

- D.W. Breck, Zeolite Molecular Sieves: Structure, Chemistry, and Use, Wiley, New York, 1974.
- [2] H. van Bekkum, E.M. Flanigen, J.C. Jansen (Eds.), Introduction to Zeolite Science and Practice, Elsevier, Amsterdam, 1991.
- [3] S. Bhatia, Zeolite Catalysis: Principles and Applications, CRC Press, Boca Raton, 1990.

- [4] T. Schroeder, M. Adelt, B. Richter, M. Naschitzki, M. Bäumer, H.-J. Freund, Surf. Rev. Lett. 7 (2000) 7.
- [5] D. Stacchiola, S. Kaya, J. Weissenrieder, H. Kuhlenbeck, S. Shaikhutdinov, H.-J. Freund, M. Sierka, T.K. Todorova, J. Sauer, Angew. Chem. Int. Ed. 118 (2006) 7798.
- [6] Y.D. Kim, T. Wei, D.W. Goodman, Langmuir 19 (2003) 354.
- [7] B.K. Min, W.T. Wallace, D.W. Goodman, J. Chem. Phys. B 108 (2004) 14609.
- [8] J. Weissenrieder, S. Kaya, J.L. Lu, H. Gao, S. Shaikhutdinov, H.-J. Freund, M. Sierka, T.K. Todorova, J. Sauer, J. Phys. Rev. Lett. 95 (2005) 076103.
- [9] L. Giordano, A. Del Vitto, G. Pacchioni, J. Chem. Phys. 124 (2006) 034701.
- [10] S. Ulrich, N. Nilius, H.-J. Freund, U. Martinez, L. Giordano, G. Pacchioni, ChemPhysChem 9 (2008) 1367.
- [11] J.B. Giorgi, T. Schröder, M. Bäumer, H.-J. Freund, Surf. Sci. Lett. 498 (2002) L71.
- [12] U. Martinez, L. Giordano, G. Pacchioni, J. Chem. Phys. 128 (2008) 164707.
- [13] J. Tersoff, D.R. Hamann, Phys. Rev. Lett. 50 (1983) 1998.
- [14] T. Schröder, J.B. Giorgi, A. Hammoudeh, N. Magg, M. Bäumer, H.-J. Freund, Phys. Rev. B 65 (2002) 115411.
- [15] J.P. Perdew, J.A. Chevary, S.H. Vosko, K.A. Jackson, M.R. Pederson, D.J. Singh, C. Fiolhais, Phys. Rev. B 46 (1992) 6671.
- [16] G. Kresse, J. Hafner, Phys. Rev. B 47 (1993) 558;
- G. Kresse, J. Furthmüller, Phys. Rev. B 54 (1996) 11169.
- [17] P.E. Blöchl, Phys. Rev. B 50 (1994) 17953.
- [18] K. Min, W.T. Wallace, A.K. Santra, D.W. Goodman, J. Chem. Phys. B 108 (2004) 16339.
- [19] D.R. Lide (Ed.), Handbook of Chemistry and Physics, CRC Press, Boca Raton, 2008.
- [20] G. Pacchioni, L. Giordano, M. Baistrocchi, Phys. Rev. Lett. 94 (2005) 226104.
- [21] N. Nilius, V. Ganduglia-Pirovano, V. Bradzova, M. Kulawik, J. Sauer, H.-J. Freund, Phys. Rev. Lett. 100 (2008) 096802.
- [22] C.E. Moore (Ed.), Atomic Energy Levels, US-Gov. Print Office, Washington, 1952.