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Ionic charge distributions in silicon atomic surface wires†

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Using a non-contact atomic force microscope (nc-AFM), we examine continuous dangling bond (DB) wire structures patterned on the hydrogen terminated silicon (100)-2 × 1 surface. By probing the DB structures at varying energies, we identify the formation of previously unobserved ionic charge distributions which are correlated to the net charge of DB wires and their predicted degrees of freedom in lattice distortions. Performing spectroscopic analysis, we identify higher energy configurations corresponding to alternative lattice distortions as well as tip-induced charging effects. By varying the length and orientation of these DB structures, we further highlight key features in the formation of these ionic surface phases.

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Introduction

The development of novel atom-scale devices has led to the design and implementation of device components made of few to single atoms.^{1–8} Due to their decreased size, the charge distribution of such devices can no longer be treated as an ensemble average, instead requiring highly localized, individual assessment. The atomic force microscope (AFM) has been proven to be a useful tool capable of detecting discrete single-electron charge transitions in both molecular^{9–13} and atomic structures,^{14–16} including patternable silicon dangling bonds (DBs)^{6,17–19} on a hydrogen-terminated surface. A DB is patterned through atomically precise removal of a surface hydrogen atom through current injection from an atomically sharp tip^{20–25} leaving a single unsatisfied bond which extends into vacuum. Previous AFM studies have examined DBs, showing readout and manipulation of its quantized charge states through both electrostatic interactions with other local fixed charges (such as other DBs), or the probe tip.^{6,17–19} These previous studies, however, focused on DB structures that had intervening hydrogen-terminated lattice sites, *i.e.* non-continuous structures. Continuous structures, such as the DB wires examined in this work, are of particular interest as they have been proposed in prior studies to exhibit more exotic behav-

iors through enhanced DB–DB coupling, including spin or ionic ordering^{26–29} similar to that seen in low dimensional Mott insulators.^{30–32} Experimental exploration of such behaviours has so far been limited to scanning tunneling microscopy (STM) measurements at energies outside the band-gap of the material, where the expected ground state ordering of DB wire structures is convoluted with higher energy features associated with the perturbative nature of a necessary tunneling current.^{33–37}

In this work, we compliment these former studies with the use of the less perturbative AFM. We explore energies within the Si band gap revealing characteristics more easily associated to the charge state of DBs within these confined 1-D structures. Since the presence of either a spin or charge ordered configuration has significant implications on the predicted transport properties associated with such DB systems^{37–44} and their utility in device-focused applications, it is important to experimentally identify their preferred configurations.

Due to the anisotropy of the (2 × 1) reconstruction,^{45–48} these DB structures are either patterned along a single dimer (in the [011] direction), creating what is commonly referred to as a bare dimer,^{33,34,49,50} or along the same side of a dimer row (in the [01̄1] direction) creating a DB wire.^{35,37} DBs within a bare dimer are separated only by the shared dimer bond that forms a pi bond,^{26,51–53} while DBs within a DB wire are separated by a second layer back bonded Si atom. This difference in bonding leads to different constraints in regard to lattice relaxation, motivating differences in their atomic and electronic properties. These differing lattice geometries are emphasized in the ESI in Fig. S1.†

By using a charge sensitive Si-functionalized probe,^{25,54} we are able to clearly distinguish between the positive, neutral, and negative charge states of DBs within bare dimers and DB wires, yielding insights into the association of DB charging

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with lattice distortion. We identify the formation of an ionic charge distribution within bare dimers and DB wires, with the DB wires only exhibiting ionic character when accompanied by changes in their overall charge state. Through spectroscopic analysis, we identify the formation of both higher energy ionic configurations attributed to an alternative lattice distortion²⁷ as well as higher order charge distributions within DB wires, with differences observed for DB wires of varying lengths.

Results and discussion

The single dangling bond

DBs on the otherwise hydrogen terminated Silicon (H-Si) (100)- 2×1 surface have been shown to behave as quantum dots capable of holding 0, 1, or 2 electrons (rendering the DB in a positive, neutral, or negative charge state).^{45,46,55,56} The native charge state of a single DB can be modified by either varying the crystal doping level or the electrostatic environment surrounding the DB.^{6,17} Using a degenerately n-doped crystal (see Methods), we perform $\Delta f(V)$ spectroscopies which reveal the distinct charge transitions of the DB through discrete shifts in Δf associated with changes in the interaction between the tip apex atom and the DB, as well as a shift in the local contact potential difference.^{6,15,16,18,57} The transition bias between the neutral and negative ((0) to (-)) and the positive and neutral ((+) to (0)) charge states are routinely observed 0.2–0.4 V below the Fermi-level^{6,17,18} and at the onset of the valence band,^{46,58} respectively. Due to the location of the (+) to (0) charge transition, high tunneling current appears when probing Δf in the expected bias range, making it challenging to maintain tip integrity and distinguish the DB charging from higher bulk current contributions.⁵⁹ By raising the electrostatic potential surrounding the DB with local fixed charges,¹⁷ these charge transitions can be shifted in energy to a mid-gap region allowing for both charge transitions to be detected with $\Delta f(V)$ spectroscopies. Fig. 1(a) shows a spectroscopy of a DB positioned on the edge of a 14 DB cluster (shown in Fig. S2†) which caused its charge transitions to shift more mid-gap—the positive (purple), neutral (orange), and negative (blue) charge states are highlighted. The (+) to (0) charge transition is seen at $V_S = -0.8$ V and the (0) to (-) charge transition at $V_S = 0.7-0.8$ V, which is ~ 1.2 V higher than the (0) to (-) values previously reported^{6,17,18} (corresponding $I(V)$ spectroscopy and STM images can be seen in Fig. S3†). The slanted nature of the $\Delta f(V)$ charge step is not seen in other reported single electron charging systems^{11,57} and is thought to be a result of a current-induced averaging of the charge state based on competing emptying and filling rates of the DB.^{58,59}

Constant height Δf maps of the DB in a fixed positive (+), neutral (0), and negative (-) charge state relative to the H-Si surface can be seen in Fig. 1(b), (c), and (d), respectively. Matching the relative frequency shifts seen in the $\Delta f(V)$ spectroscopy, the (+) DB appears “brighter” than the surrounding H-Si atoms, the (0) DB shows similar contrast to the surrounding H-Si atoms, and the (-) DB appears “darker” than the sur-

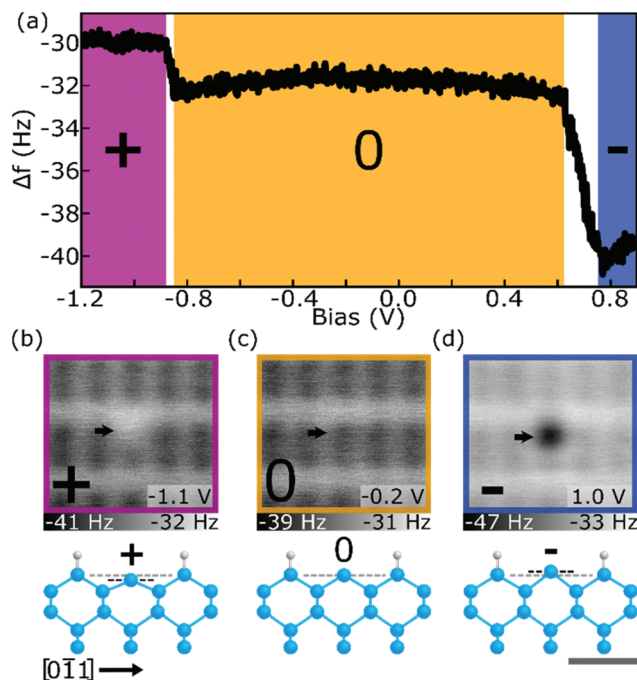


Fig. 1 Three charge states of the Si DB. (a) $\Delta f(V)$ spectroscopy over a Si DB showing the (+) to (0) and (0) to (-) charge transitions (occurring at ~ -0.9 V and $\sim 0.7-0.8$ V, respectively). (b–d) Constant height Δf images of the DB in a fixed positive (+), neutral (0), and negative (-) charge state, respectively, with corresponding qualitative ball and stick models of the Si DB in each charge state. Models are in a projected side view along the $[0\bar{1}1]$ direction (see Fig. S1†). The black dashed line in the models highlights the vertical shift in height of the charged host Si atom relative to the neutral state (grey dashed line), with dimensions listed in the text. Each image is 2.1×2.1 nm² with bias values indicated in the lower right of the panels. The grey scale bar is 0.5 nm relative to the ball and stick models. All measurements are performed at -300 pm from the height set point (see Methods).

rounding H-Si atoms. In addition to the change in charge state, there is also a predicted corresponding lattice distortion of the host Si atom.^{18,55,60} A (-) DB raises a host Si atom up (u) from the surface by roughly 30 pm as it adapts more sp^3 character while a (+) DB lowers the position of the host Si atom down (d) towards the surface by roughly 40 pm with it adapting more sp^2 character.⁵⁵ This lattice distortion is qualitatively shown in the ball-and-stick models in Fig. 1(b)–(d) with the horizontal black line showing the relative shift from the (0) Si atom in grey. It is difficult to extract the contribution of such lattice features from AFM data as the measured Δf signal is also convoluted with the changing covalent and charge induced dipole interaction of the surface DB with the neutral DB of the Si terminated tip.^{61–65} It is thus expected that the measured Δf signal will be a combination of the height of the host silicon atom relative to its surroundings, its covalent interaction with the tip (more sp^2 vs. more sp^3 like bonding interactions), and an ion-induced dipole component for charged DBs. Numerical determination of the weighting of each of these three components on the measured Δf awaits detailed theoretical modelling outside of the scope of this manuscript.



The bare dimer

With an understanding of the behaviour of single DB charging using a Si terminated probe, we now extend these measurements to a bare dimer and DB wires. Consistent with observations of the unterminated 2×1 surface,^{66–72} bare dimers are predicted to undergo a lattice distortion putting one atom in a raised (u) sp^3 electron-rich (–) state, and the other in a lowered (d) sp^2 electron-deficient (+) state.^{26,42,73} Fig. 2(a) shows the $\Delta f(V)$ spectroscopy of each DB in a bare dimer (labelled left and right) revealing six distinct regions within the previously unexplored Si band gap. $\Delta f(V)$ line scan maps along the bare dimer were also taken, with 50 line scans over the dimer acquired at incremented bias intervals of 0.02 V (corresponding $I(V)$ spectroscopy and $I(V)$ line scan maps are shown in Fig. S4†).

At biases lower than -0.98 V (purple), the bare dimer appears to be in a symmetric configuration with both sides of the bare dimer appearing neutral also shown in the constant height Δf image of Fig. 2(c). Increasing the sample bias to between -0.98 V and -0.6 V (orange), the bare dimer reorders to the expected ionic (+–, du) configuration, with the left DB

appearing positive (less negative Δf) and the right DB appearing negative (more negative Δf), as supported in Fig. 2(d). As discussed with the single DB case, the measured difference in the Δf signal between DBs within the bare dimer is a convolution of the difference in height of the host Si atoms relative to the tip-apex, a difference in the covalent character of each surface DB, and differences in their ionic interaction with the tip, which are all correlated to their assigned charge state.

Between -0.6 V to -0.5 V (green), the tip becomes resonant with the pi state^{26,51–53} of the bare dimer, possibly allowing charge to quickly transfer between sides of the bare dimer as indicated by the fluctuations in signal in the $\Delta f(V)$ spectroscopy of Fig. 2(a), as well as the streakiness seen in the line scans of Fig. 2(b). Between -0.5 V and 0.2 V (yellow), the $\Delta f(V)$ spectroscopies show that both DBs appear negative, however looking at the line scans in Fig. 2(b) and the Δf map of Fig. 2(e), it is clear that the bare dimer is still in an ionic configuration, except now it readily switches between its (+–, ud) and (+–, du) configurations (geometry shown in Fig. S1(c)†), likely facilitated by the newly accessible pi state. The reason both DBs in the $\Delta f(V)$ spectroscopies appear negative is due to a localized attractive interaction between the DB and the tip

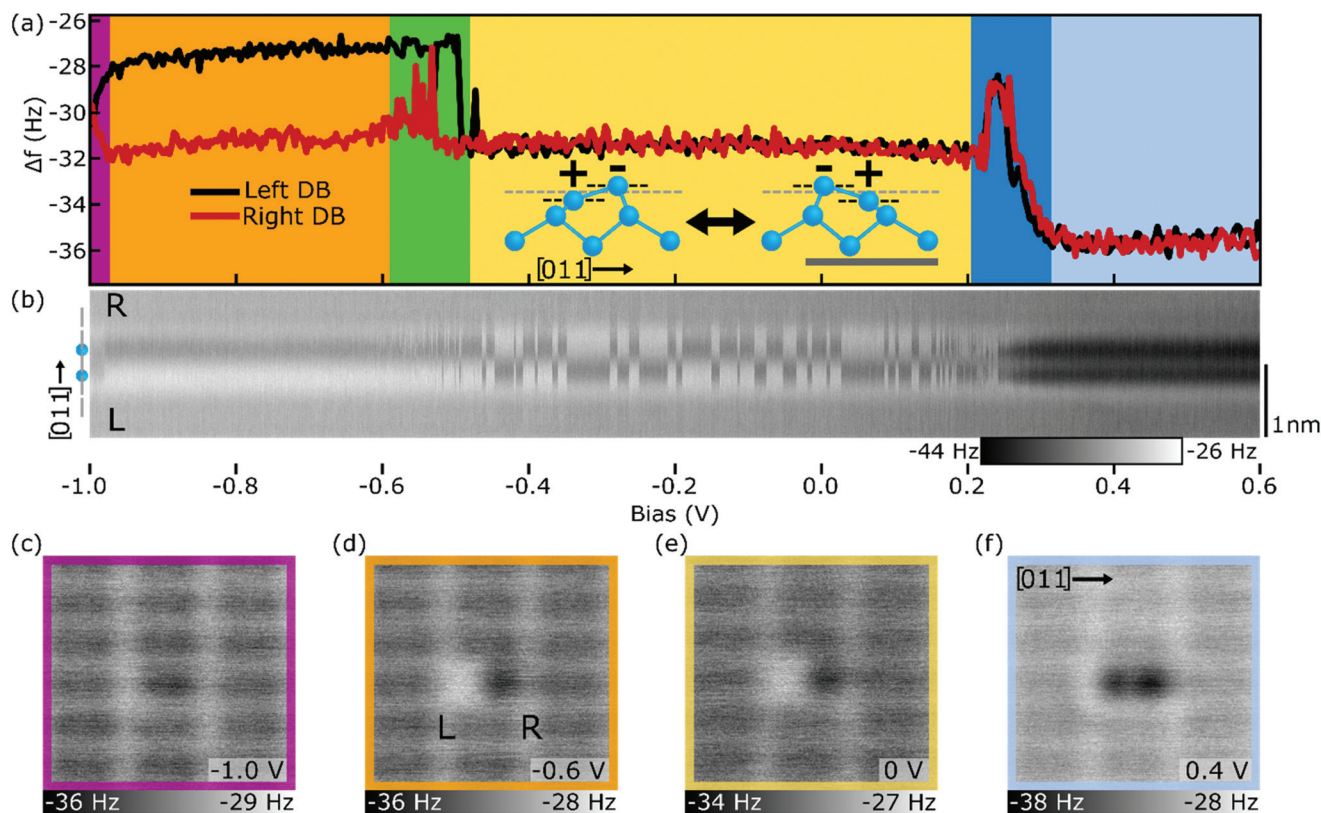


Fig. 2 Charge distributions of the bare dimer. (a) $\Delta f(V)$ spectroscopy over the right (red) and left (black) DB of the bare dimer. The inset shows a side-view along the [011] direction of the buckled geometry of the bare dimer in the two degenerate charge configurations (+–/–+, du/ud) (dimensions taken from ref. 74). Scale bar in grey corresponds to 0.5 nm. (b) $\Delta f(V)$ line scans over the bare dimer with the position of the bare dimer within the line scans highlighted by the model on the left of (b) with a black scale bar of 1 nm on the right. 50 line scans from a bias of -1.0 to 0.6 V are shown, in increments of 0.02 V. (c)–(f) Constant height Δf images of the bare dimer in four of the six bias regions of (a), as indicated by the coloured border and bias. All measurements were performed at a tip–sample distance of ~ 300 rpm from the height set point (see Methods). Corresponding band diagrams are shown in Fig. S8.†



causing the negative charge within the bare dimer to “follow” the tip.^{18,63,67} As detailed by Rashidi *et al.*,¹⁸ they found that a silicon atom directly underneath the tip experiences an attractive tip–sample interaction which mechanically pulls the host Si atom towards the tip, adopting more sp^3 character. This enables the surface DB to switch to a negative charge state if the tip is positioned over it for a sufficient time (\sim seconds). DBs which are geometrically coupled to the newly localized negative charge state and corresponding change in lattice geometry of the DB under the tip, undergo corresponding transitions to minimize the overall energy of the DB structure. In this case, it requires the adjacent DB within the bare dimer to transition to a positive (+) charge state ((d) lattice configuration), accounting for the ionic configuration observed in Fig. 2(e). The line scans of Fig. 2(b) for biases between -0.5 V and 0.2 V (yellow) show a switching between each degenerate ionic configuration due to the tip scanning across the surface. As the tip probes the (+) DB during the line scan, there is some probability that this localized attractive tip–sample interaction will cause a transition from the (+) to (–) charge state. The system can then reverse the configuration through a similar transition over the newly (+) DB within the bare dimer. The probability of a tip induced switching to occur in the bare dimer structure is shown to be strongly dependent on the tip sample separation as shown in Fig. S5 and S7.†

At higher bias values between 0.2 V and 0.3 V (dark blue), the bare dimer becomes resonant with the π^* state^{52,53} as indicated by the peaks seen in the $\Delta f(V)$ spectroscopy. Looking at the line scans in Fig. 2(b), it appears that the DBs experience this increase in Δf simultaneously suggesting that this peak is a measurement artifact, *i.e.*, electron dynamics are occurring at a faster rate than the AFM is capable of sampling. Corresponding features are also seen in the $I(V)$ plots in Fig. S4,† corresponding to resonant tunneling through the π^* state. Above bias values of 0.3 V (light blue), both DBs in the bare dimer appear to be in a negative charge state as supported by Fig. 2(f) and the more negative Δf signal seen for both DBs in Fig. 2(a). It is unclear if the bare dimer is in a $2e^-$ (–) charge configuration due to electron injection from the tip, or is switching between a degenerate $1e^-$ (0–/–0) charge configuration at rates faster than the sampling of the AFM. Since the origin of the characteristic butterfly shape^{33,34} of the bare dimer in STM images (Fig. S4†) has been attributed to a flipping between two degenerate states,³⁴ it is more likely that the bare dimer exists in a $1e^-$ (0–/–0) configuration. Assigning the same up and down lattice features with the higher charge structures is omitted since such geometries have yet to be analyzed with theoretical modelling. Further discussions regarding the structural and charge distributions of the bare dimer are given in the ESI.†

Dangling bond wires

DB wires are predicted to undergo a similar lattice distortion attributed to a Peierls effect,^{75–77} thus reducing the overall energy of the DB wire (as shown in Fig. S1(d)†). Features in STM measurements have been attributed to such reorderings through measured variations in the height of DBs within a

wire, as explained by Jahn–Teller distortions³⁵ in the pairing of second layer Si atoms. DBs which are raised through the pairing of second layer Si atoms exhibit more sp^3 qualities creating a more electron-rich state, while DBs which are lowered through the pairing of second layer Si atoms exhibit more sp^2 qualities and create a more electron-deficient state.^{35,76,78} This resulting lattice shift is predicted to cause an almost ionic charge redistribution throughout the wire *via* an sp^3 – sp^2 (ud) pairing of DBs within the wire. More recent theoretical studies suggest that spin ordering through electron–electron interactions is the more stable ground state configuration when compared to a charge reordering from electron–phonon interactions.^{26,27,29,42,79} Wires are predicted to align in an anti-ferromagnetic fashion at finite length, with the non-magnetic (ionic) ordering becoming more stable at longer wire lengths due the additional stability of longer ionic configurations.^{27,29}

Using $\Delta f(V)$ spectroscopies, it is now revealed that with a net neutral charge, DB wires do not exhibit any charge reordering. Only through the addition of a negative charge (or positive charge as shown in Fig. S9†) does the DB wire reorder into the theoretically predicted ionic distribution, likely facilitated by an additional polaronic effect,^{76,80–82} as seen in Fig. 3. Fig. 3(a) shows the $\Delta f(V)$ spectroscopy taken over the second DB in a five DB wire (indicated by the black arrow in Fig. 3(b)–(f)) revealing four distinct charge distributions, with a $\Delta f(V)$ line scan map shown in (b). Between -1.0 V and -0.5 V (orange), the DB wire exists in a net neutral charge state with each DB in the wire imaged in a neutral charge state, as shown in Fig. 3(b) and (c) (insights into the predicted spin ordering require a spin sensitive probe and are currently outside our abilities). Between -0.5 V and 0.25 V (dark blue), the DB wire enters a net $1e^-$ charge state, resulting in an ionic charge redistribution and corresponding lattice distortion (–+–+, ududu) as shown in Fig. 3(b) and (d).

As the bias is increased between 0.25 V and 0.45 V (light blue), the 5 DB wire is shown to periodically enter an alternative net $1e^-$ (0–+–0, dudud) charge distribution shown in the top half of Fig. 3(e) and in the line scan map between 0.05 V and 0.45 V in Fig. 3(b). Such a charge configuration is predicted to have a higher overall energy²⁷ than the (–+–+, ududu) configuration and is therefore less stable when imaged. Although the outer atoms in this charge configuration are expected to be in a more electron-depleted sp^2 configuration, the rigidity of the end H–Si atoms, which cannot similarly re-hybridize in combination with the crystal doping level, prevent the formation of positive DBs at the ends of the wire.^{27,29} Increasing the bias between 0.45 V and 0.8 V (red), the 5 DB wire enters a higher order charge state (multiple e^-). Looking at the Δf map of Fig. 3(f), the DB wire appears to be in a (–0–0–) charge state, however, Fig. 3(b) shows the DB wire appears to be in a (–000–) charge configuration. Since Fig. 3(b) was imaged with a 50 pm greater tip–sample separation, the central DB in Fig. 3(f) is thought to charge due to an increased attractive interaction between the tip and surface DB (Fig. S10† highlights the height dependence of charging within 5 DB line scan maps).

Extending these measurements to DB wires of varying length, the same charge reordering is seen for DB wires with a



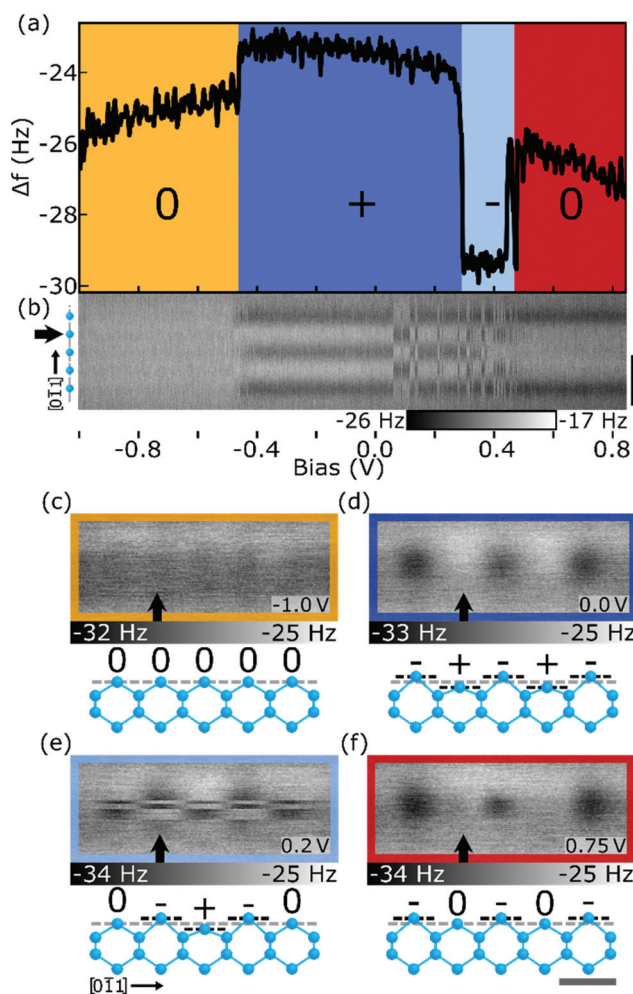


Fig. 3 Charge distributions of the 5 DB wire. (a) $\Delta f(V)$ spectroscopy over the second Si DB (marked with a black arrow in (b)–(f)) of the 5 DB wire showing four distinct charge regions ($\Delta f(V)$ spectroscopy of all 5 DBs within the wire is shown in Fig. S11†). (b) Stacked $\Delta f(V)$ line scans over the DB wire. 50 line scans at bias increments of 0.02 V from -1.0 to 0.82 V are shown. Black scale bar is 1 nm as indicated on the right of (b). (c)–(f) Constant height Δf images of the neutral, lower energy net $1e^-$, higher energy net $1e^-$, and higher net negative charge ($2e^-$ or $3e^-$) configuration, respectively. Ball and stick models provide a side view along the $[0\bar{1}1]$ direction (grey scale bar is 0.5 nm) which shows the expected trends in lattice distortion as calculated in ref. 27 and 78. Bias values of (c)–(e) are indicated in the lower right of each panel. The relative tip heights for (b) is -200 pm and (a), (c)–(f) is -250 pm (see Methods). All images are 1.0×2.5 nm 2 .

net $1e^-$ charge. Odd DB wires, shown in Fig. 4(a), are observed in a lower energy configuration where the edge DBs are both in the $(-, u)$ orientation and higher energy configuration where the edge DBs are both in the $(0, d)$ configuration. Each of the line profiles in Fig. 4 is an averaging of multiple line scans in either configuration taken from full line scan maps in Fig. S12† where the local minima in Δf correspond to a negatively charged $(-, u)$ Si DB and the local maxima in Δf correspond to a positively charged $(+, d)$ Si DB. Like the 5 DB wire, the 3 and 7 DB wires show a lower energy $(-+-, udu)$ and

$(-+-+--+, udududu)$ charge reordering, respectively. In the higher energy configuration of the 3 and 7 DB wires, each host Si atom switches its orientation resulting in $(0-0, dud)$ and $(0-+-+0, dududud)$ configuration respectively. As predicted in ref. 27, the energy difference between the (udu) and (dud) states of the 3 DB wire is larger than the energy difference between these states in the 5 and 7 DB cases. As a result, the higher energy $(0-0, dud)$ configuration of the 3 DB wire is less stable and was never observed for a complete line scan. The line scan presented in Fig. 4(a) only showed a momentary excitation as the tip scans over the centre DB. As a result, the line profile appears much noisier (due to reduced averaging) with each DB appearing negative. The blue line profile was drawn to show the expected configuration if the structure was stable enough to be completely imaged by the tip. The transition of the middle DB in the 3 DB wire from the $(+)$ to $(-)$ charge state is also seen in the $\Delta f(V)$ spectroscopies shown in Fig. S11.†

Unlike the odd length DB wires, the even length DB wires demonstrate mirrored degenerate configurations when in the net $1e^-$ charge state. Fig. 4(b) shows line profiles of both degenerate distributions for DB wires of length 2, 4, and 6. Much like the odd wire case, DBs which are raised through second layer pairing (sp^3) are imaged as negative, while DBs which are lowered through second layer pairing (sp^2) are imaged as positive if they sit within the wire and neutral if they sit at the ends of the wire. Due to the even length geometry, edge DBs now sit in opposite configurations $(0...-/-...0, d...u/u...d)$ where the energy difference between either configuration is null. The ionic distributions in the 2, 4, and 6 DB wires are therefore observed in the $(-0/0-, -+-0/0-+-, -+--+0/0-+-+)$ charge distributions, respectively, as shown in Fig. 4(b) which corresponds to $(ud/du, udud/dudud, ududud/dududud)$ respectively. In the 2 DB wire case, the predicted barrier between degenerate configurations is much lower than that of the longer DB wires,^{35,83} so it toggles more easily between each configuration. As a result, the 2 DB wire configuration often falsely appeared in a $(--)$ configuration, as the negative DB easily “followed” the tip during scanning. The left configuration $(-0, ud)$ appeared for only a few line scans resulting in greater noise seen in the line profile, with the right configuration $(0-, du)$ never appearing in this dataset, possibly due to the unseen presence of a local electrostatic perturbation from subsurface charges making the negative charge favour a side.^{17,21} The blue curve has been drawn to highlight the expected configuration. Full $\Delta f(V)$ spectroscopies and $\Delta f(V)$ line scan maps are shown in Fig. S11 and S12† respectively.

As the DB wire length is increased, the bias at which each wire transitions from the net neutral to net negative ($1e^-$) charge state is found to occur at more negative biases, as predicted in ref. 27 and 29. Fig. 5 highlights this reduction in charge transition voltage for DB wires up to 15 DBs long (extracted from Δf line scan maps in Fig. S14†). In addition to this associated decrease in charge transition bias with increased length, a slight increase in the charge transition bias is observed between odd length and 1 DB longer even length wires (3 to 4, 5 to 6, etc.). Since these odd and even wires



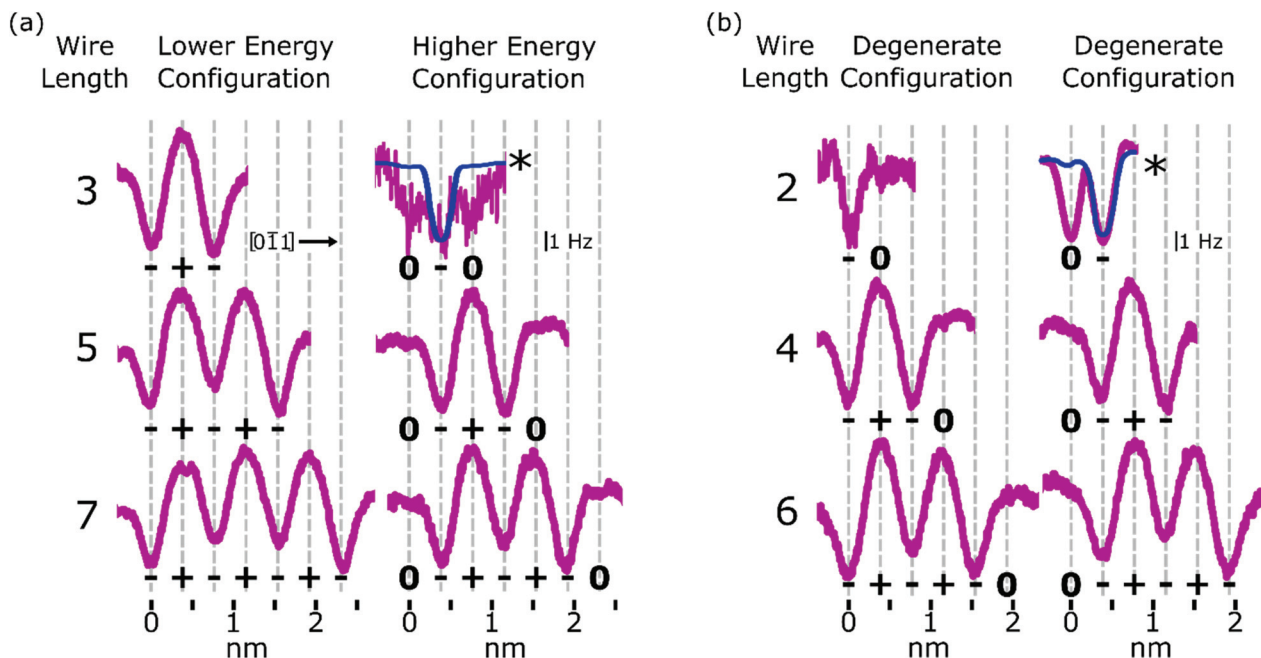


Fig. 4 DB wire charge distributions. (a) and (b) Line profiles of odd and even length DB wires from 2 to 7 DBs along the $[0\bar{1}1]$ direction in lower energy and higher energy charge configurations (displayed configurations are degenerate for even length DB wires). Each profile shown is an average of 20–50 individual line scans from Fig. S12,† except the higher energy configuration (0–0, dud) of the 3 DB wire and the degenerate (–0, ud) configuration of the 2 DB wire; these were only visible for 1 and 6 line scans respectively. The blue lines show the expected configuration for each as described in the text.

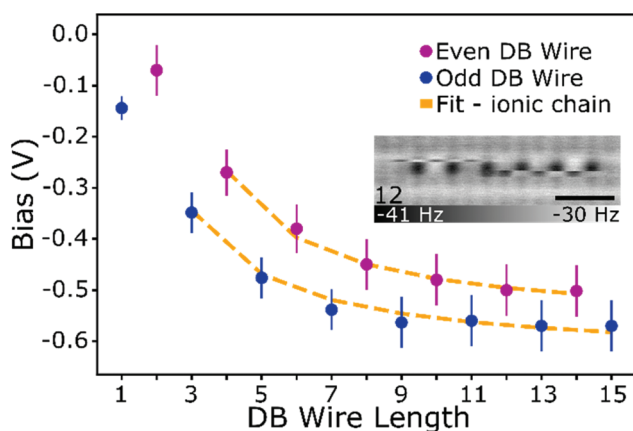


Fig. 5 Charging voltage of DB wires. Bias at which the DB wire transitions from a net neutral to net negative charge state ($1e^-$). Error bars account for variations in charging energy between DBs in the $\Delta f(V)$ spectroscopies (Fig. S11†) and line scan images (Fig. S12 and Fig. S14†) taken at varying tip heights. The fit correlates to an alternating ionic chain of varying length as described in the text. The inset shows a 12 DB wire imaged at 0 V and -300 pm to show an example of domain ordering for longer length wires. Scale bar in black is 1 nm. A full set is provided in Fig. S13.†

possess the same number of positive and negative DBs, we speculate that this shift in charge transition bias is due to an increase in energy from the additional lattice strain of the end DB without the energy lowering coulombic contribution from ionic pairing.^{27,29} Increasing the wire length further from an

even to odd length wire (4 to 5, *etc.*) creates an additional ionic (+–, du) pair which lowers the voltage at which the charge transition occurs. The opposite trend between odd and even length DB wires was predicted in ref. 27 and 29, which we attribute to their models assuming an ionic configuration in the net neutral charge state. At a DB wire length above 9 DBs, this charge transition voltage is found to reach a threshold value of roughly -0.55 V. This threshold in charging bias corresponds to a saturation in the coulombic energy gained per DB associated with a finite alternating ionic wire.⁸⁴ Such a physical process can be represented by relating the coulombic energy gained per DB ($\frac{U(N)}{N}$) as a function of the length of an alternating ionic wire (N for odd DBs, $N - 1$ for even DBs – due to the presence of the additional neutral DB). Developed by Ciftja *et al.*,⁸⁴ it is written as:

$$\frac{U(N)}{N} = \frac{k}{Na} \sum_{i=1}^{N-1} \sum_{j=i+1}^N \frac{(-1)^{j-i}}{(j-i)} + U_0 \quad (1)$$

where k is $\frac{e^2}{4\pi\epsilon_0\epsilon}$ (e is the elementary charge and ϵ is the effective dielectric constant), a is the DB lattice spacing (0.384 nm), and U_0 is some offset potential.

Fitting the charge transition biases as a function of length in Fig. 5 (length 1 and 2 are omitted since there is no ionic pair present), we extract values of $\epsilon = 2.50 \pm 0.12$ and $U_0 = 0.40 \pm 0.04$ eV for the odd length wires, and $\epsilon = 2.37 \pm 0.11$ and $U_0 =$



0.52 ± 0.04 eV for the even length wires. Interestingly, the effective dielectric constant is slightly smaller than that calculated for DBs with intervening H atoms (4.1 ± 0.2)¹⁷ highlighting the potentially different properties associated with a continuous DB wire. Using the offset potential, we can also extract an energy difference of 0.12 ± 0.06 eV between the even and odd length wires due to the addition of some lattice strain associated with the neutral DB without the additional coulombic contribution from the charged DBs. At longer lengths, the $1/r$ dependence results in vanishingly small contributions to the charge transition bias with added DBs. The inset of Fig. 5 shows a manifestation of this effect, with a 12 DB wire in which a domain fracturing behaviour is seen through the decoupling of the two ends which are now able to switch between charge configurations independently. A full discussion of the longer lines is provided with Fig. S13 and S14.†

Conclusion

By using a Si terminated tip sensitive to the three different charge states of surface Si DBs, we were able to show the preferred configurations of DB structures at varying probe biases. We were able to confirm the charge redistribution and corresponding buckling of a net neutral bare dimer, as well as identify the pi and pi* states through $\Delta f(V)$ spectroscopy. Examining DB wires, we report that the theoretically predicted charge redistributions and lattice distortions are only favoured when the entire DB structure is negatively charged. The net neutral charge state of the DB wire reveals no such charge redistribution, allowing for a possible spin ordering to be present. Further measurements using spin sensitive microscopy is needed to confirm this. DB wires in the net negative ($1e^-$) charge state were observed to hold two different lattice configurations, corresponding to a lower (udu) and higher (dud) energy configuration in odd length DB wires and a degenerate (ud/du) configuration in even length DB wires, with the charging voltage dependent on the wire length. Higher order charge configurations were also revealed where additional charging breaks the ionic pairing. These higher order charge distributions likely contribute to the observed character in STM images. These results have allowed for the clear interpretation of charge distributions within 1 dimensional DB structures giving insights into their expected ground state phase formation and viability as charge carriers for atomic scale electronics.

Methods

Experiments were performed using an Omicron qPlus LT AFM^{85,86} and an Omicron LT STM system operating at 4.5 K and 3×10^{-11} Torr. AFM tips used a third generation Giessibl tuning fork with a focused ion beam (FIB) mounted Tungsten tip ($f_0 \sim 28$ kHz, Q -factor $\sim 16k$ – $22k$, Amplitude = 50 pm).⁸⁷ The tip was cleaned and sharpened using a combination of field-

evaporation and nitrogen etching in a field ion microscope (FIM).⁸⁸ Further *in situ* conditioning was done *via* controlled tip contacts with hydrogen desorbed patches of silicon until it returned characteristics corresponding to a Si terminated tip.^{25,54} DB structures were created *via* controlled bias pulses.^{20,23,89} The bias ranges for each set of measurements refer to the sample bias (V_s) and were chosen to probe as great a bias window as possible while trying to prevent any unwanted tip changes due to high tunneling current through the valence and conduction band of the crystal. The AFM measurements taken in this work were taken over several months, so while each tip shows the desired Si tip contrast of the surface, the exact reactivity of the tip varies between some data sets.

Samples were degenerately arsenic-doped ($1.5 \times 10^{19} \text{ cm}^{-3}$). They were prepared by first degassing the sample using resistive heating at 600 °C overnight followed by multiple annealing flashes at 1250 °C. The samples were subsequently hydrogen terminated by exposing the system to molecular hydrogen (1×10^{-6} Torr) for two minutes while holding the sample at 330 °C. A tungsten filament held at 1600 °C was used to crack the hydrogen.^{56,90}

Images and data were collected using a Nanonis SPM controller, with additional custom-built LabVIEW controllers. The height setpoint references in constant height measurements are given relative to a tip-sample separation distance initialized with a tunneling current feedback of 50 pA and sample bias of -1.8 V with the tip positioned over a H-Si atom. Negative values indicate a reduction in the separation between the tip and sample. While we use a height reference defined by these STM setpoints, previous work has estimated this to correspond to an absolute tip-sample distance of ~ 700 pm as established by Rashidi *et al.*¹⁸ where they defined $z = 0$ as a point where noticeable changes in tip-structure from surface contact occurred. It is also found that the absolute tip-sample distance can vary slightly (~ 100 pm) due to different apex functionalizations,⁵⁴ tip geometries, and surface dopant profiles. Thus, we stick with the easily accessible STM set-point to enable easy reproduction of our results. The fits in Fig. 5 were performed using `scipy.optimize`.

Author contributions

JC curated and analysed all experimental data except Fig. S9† which was curated and analysed by TH. The manuscript was written by JC with review and editing from all authors. TH, MR, JW, and EL further supported the development of this manuscript with additional investigations not shown. JP and RAW supervised the project.

Conflicts of interest

The authors declare competing financial interests. Some of the authors are affiliated with Quantum Silicon Inc. (QSi). QSi is seeking to commercialize atomic silicon quantum dot-based technologies.



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