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# Thermodynamics of the formation of surface PtO<sub>2</sub> stripes on Pt(111) in the absence of subsurface oxygen†

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This paper examines the thermodynamics of PtO<sub>2</sub> stripes formed as intermediates of Pt(111) surface oxidation as a function of the degree of dilation parallel to the stripes, using density functional theory and atomistic thermodynamics. Internal energy calculations predict 7/8 and 8/9 stripe structures to dominate at standard temperature and pressure, which contain 7 or 8 elevated PtO<sub>2</sub> units per 8 or 9 supporting surface Pt atoms, respectively. Moreover, we found a thermodynamic optimum with respect to mean in-stripe Pt–Pt spacing close to that of  $\alpha$ -PtO<sub>2</sub>. Vibrational zero point energies, including bulk layer contributions, make a small but significant contribution to the stripe free energies, leading to the 6/7 stripe being most stable, although the 7/8 structure is still close in free energy. These findings correspond closely to experimental observations, providing insight into the driving force for oxide stripe formation and structure as the initial intermediate of platinum surface oxidation, and aiding our understanding of platinum catalysts and surface roughening under oxidative conditions.

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## 1. Introduction

Surface oxidation, which happens both in gas-phase catalysis and in electrochemical environments, impacts profoundly the selectivity, activity and stability of (electro)catalysts.<sup>1,2</sup> A commonly-used electrocatalyst material, platinum, has long been known to undergo surface oxidation during catalyst operation.<sup>1–4</sup> Whether as an electrocatalyst or as a gas-phase heterogeneous catalyst, the metal surface is exposed to oxygen species, which may lead to surface reconstruction, or in the case of aqueous media, dissolution.<sup>5</sup> Dissolution,<sup>6–8</sup> and reduction of surface oxides,<sup>9</sup> drive further reconstruction and (irreversible) roughening of the surface.<sup>4,8,10,11</sup> In order to mitigate any deleterious effects and improve our understanding of oxygen-induced surface roughening, we need to know the adsorption patterns of oxygen on – and in – platinum.

There is ample evidence for more complicated oxygenation of Pt(111) than simple on-surface adsorption. At low oxygen coverages generated under ultrahigh vacuum (UHV) conditions, Devarajan *et al.* observed this on-surface adsorption of oxygen in fcc hollow sites using temperature programmed desorption and scanning-tunneling microscopy (STM) experiments.<sup>12</sup> Yet for oxygen coverages greater than 0.40 ML, some Pt atoms are elevated, forming PtO<sub>x</sub> stripe segments and (above 0.71 ML) honeycombs. Density functional theory (DFT) calculations by Hawkins *et al.* and force field simulations by Farkas *et al.* corroborated part of these findings by finding stripe segments at coverages equivalent to 0.75 ML and stable PtO<sub>2</sub> stripes for 1 ML coverage; all these structures were supported by subsurface oxygen.<sup>13,14</sup> Holby *et al.*, on the other hand, indicated that it is not necessary to incorporate subsurface oxygen: they found striped structures alternating between flat and buckled stripes.<sup>15</sup>

In practice, platinum catalysts are usually operated under standard or elevated pressures and close to standard temperature in the presence of O<sub>2</sub>, as opposed to UHV and high-temperature conditions using more strongly oxygenating reagents. Luo *et al.* found that for platinum films, exposure to either pure O<sub>2</sub> or mixtures of O<sub>2</sub> and water vapor led to near-monolayer oxygen coverage.<sup>16</sup> In an *in situ* STM study of oxygenation patterns, Van Spronsen *et al.* exposed Pt(111) to O<sub>2</sub> at pressures ranging from 1–5 bar and at temperatures between 300–538 K.<sup>17</sup> During O<sub>2</sub> exposure at 1 bar, a *spokewheel structure* is observed, above 2.2 bar accompanied by *stripes* as illustrated in Fig. 1.<sup>17</sup> The linear segments, or spokes of the spokewheel, are PtO<sub>2</sub> stripe

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† Electronic supplementary information (ESI) available: Thermodynamics of stripe formation and free molecules, computational details of DFT calculations and stripe geometries, DFT formation energies and corrections for free molecules, surface and stripe vibrational analysis including bulk modes, formation energies of PtO<sub>2</sub> structures under standard conditions, and stripe formation energies with respect to Pt binding energies. All calculations are performed at 300 K, and simulations were performed using the PBE exchange–correlation functional through VASP. All structural images were rendered using VESTA.<sup>33</sup> See DOI: 10.1039/c9cp05107d



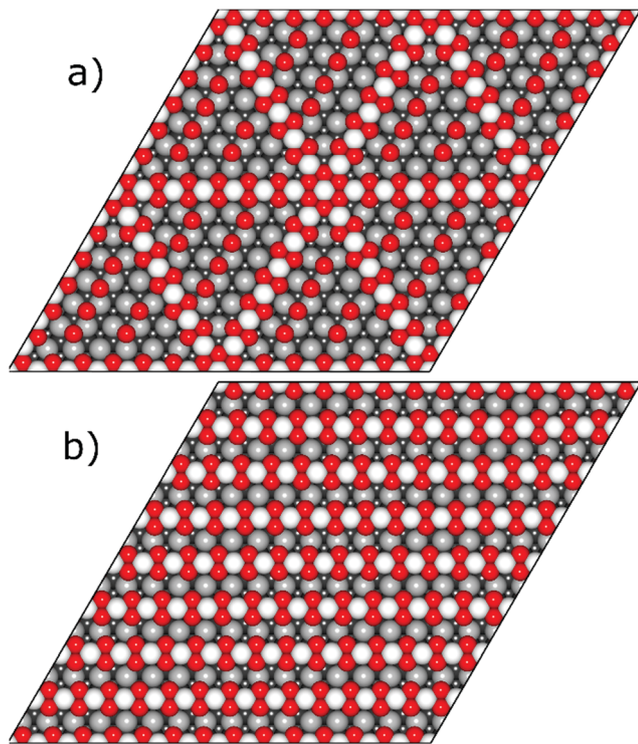


Fig. 1 Artist's impression of spokewheel (a) and stripe (b) models adapted from Van Spronsen *et al.*<sup>17</sup>

structures which are platinum deficient relative to the underlying surface, causing a  $\sim 14\%$  lattice mismatch of  $\text{PtO}_2$  structures relative to the pristine surface lattice vectors. In other terms, these structures contain  $7/8$   $\text{PtO}_2$  units per Pt atom partaking in the reaction.

This lattice mismatch appears to be mirrored in sub-monolayer  $\text{PtO}_x$  structures studied by Hawkins *et al.*, who found non-contiguous  $\text{PtO}_x$  clusters supported by subsurface oxygen to be dilated relative to the underlying platinum bulk lattice parameter.<sup>13</sup> The presence of subsurface oxygen supporting Pt atoms corresponds to the concept of place exchange, which was first proposed by Reddy *et al.* for electrochemical oxidation of platinum,<sup>18</sup> and later supported by a kinetic study by Conway *et al.* and X-ray measurements by You *et al.*<sup>19–21</sup> The traditional definition of place-exchange, as mentioned by Drnec *et al.*, involves switching one platinum atom in the surface by one oxygen atom, lifting the former from the surface and integrating the latter into it. Yet, platinum atoms in Van Spronsen *et al.*'s study appear to be elevated in continuous stretches, and the oxygen coverage is twice the fraction of surface platinum atoms in the stripes, which are lifted through oxidation. This suggests that contrary to Hawkins *et al.*'s  $\text{PtO}_x$  structures, Van Spronsen's structures on average consist of  $\text{PtO}_2$  units even below 1 ML oxygen coverage. All these lifted atoms are found to be adjacent in spokewheel structures, which one would not expect if buckled stripes play a major role as in the model proposed by Holby *et al.*<sup>15</sup> XPS measurements on stripe structures by Van Spronsen show 0.88 ML O coverage, indicating regular  $7/8$  dilation, yet could not be elucidated

using STM imaging. Additionally, platinum atoms are expelled from the pristine surface domain on which these stripes are formed, an effect not taken into account by Hawkins *et al.*<sup>13</sup> This may in turn aid the formation of serpentine islands otherwise attributed to adsorbate-induced strain by Van Spronsen *et al.* Most strikingly, Van Spronsen *et al.*'s results indicate that subsurface oxygen is not necessary to stabilize  $\text{PtO}_2$  surface structures under (near-)standard oxidizing conditions.

Apart from their role in gas-phase Pt oxidation,  $\text{PtO}_2$  stripes may also be first intermediates in the electrochemical oxidation of Pt electrodes. Furuya *et al.* detected oxidized platinum atoms exposed on-surface under electrochemical conditions at  $\sim 1.05$ – $1.20$  V, suggesting the formation of surface  $\text{PtO}_x$  in a so-called quasi-3D structure as in gas-phase Pt oxidation.<sup>5</sup> *In situ* electrochemical STM measurements by Jacobse *et al.* show that cycling of Pt(111) through relatively mild oxidative potentials of 1.35 V initially causes island formation, and subsequently irreversibly roughens the surface during multiple oxidation–reduction cycles,<sup>11</sup> in agreement with X-ray spectroscopy measurements by Ruge *et al.*<sup>22</sup> Lifting of Pt atoms from the surface has been observed through *in situ* X-ray observations of electrochemical Pt oxidation and assumed to involve place exchange;<sup>4,23</sup> Drnec *et al.*, however, noted that their measurements do not show either an ordered overlayer with a different lattice parameter, nor an amorphous oxide structure, but they cannot exclude the possibility of (long-range) oxide chain formation.<sup>4</sup> Together, these findings suggest a role for a  $\text{PtO}_x$  based intermediate not supported by subsurface oxygen under electrochemical conditions, both regarding reactivity of Pt electrodes at high potentials and the surface transformation as a result of oxidation–reduction cycles.

In this study, we investigate the formation energies of subsurface oxygen-free  $\text{PtO}_2$  stripes as proposed by Van Spronsen, and attempt to reproduce their structures, using DFT methods. In the process, we compare various stripe  $\text{PtO}_2/\text{Pt}(\text{displaced})$  ratios, calculate their energies as a function of oxygen chemical potential and electrochemical potential, and discuss their similarity to the ordering of the spokewheel segments, and how these energies are related to  $\text{PtO}_2$  geometries and bulk platinum properties.

## 2. Density functional theory methods and initial structure generation

All calculations on Pt(111) and oxidized Pt(111) were performed using density functional theory (DFT) in the Vienna Ab Initio Simulation Package,<sup>24</sup> with the spin-restricted Perdew–Burke–Ernzerhof (PBE) exchange correlation functional.<sup>25</sup> All plane-wave cutoffs are equal to 450 eV, while ionic cores are described using plane augmented wave (PAW) pseudopotentials for PBE supplied in the VASP 5.2 distribution.<sup>24</sup> Orbital occupancies in slab systems and bulk solids are subject to 0.20 eV first order Methfessel–Paxton smearing,<sup>26</sup> whereas gas-phase molecules are subject to 0.0001 eV Gaussian smearing.<sup>27</sup> The bulk Pt



lattice parameter  $a$  was found to be equal to 3.9754 Å through minimization of internal energy of a single atom fcc bulk cell under variation of lattice vector lengths. Computational details pertaining to gas phase molecules ( $\text{H}_2(\text{g})$ ,  $\text{H}_2\text{O}(\text{g})$ ) are listed in Section 3 of the ESI.†

We defined cells for  $(111) M \times N$  surface slabs for each stripe coverage ratio; similarly, we created bulk cells by stacking six  $(111)$  layers of  $M \times N$  platinum atoms. Cell dimensions and parameters used for these slabs and bulk cells are listed in Table S.1 in the ESI.† Specifically, each unit cell is defined as having a  $2 \times N-60^\circ$  translational surface symmetry, and lattice vectors (including the additional lattice vector in the bulk) run parallel to shortest-distance atom pair vectors. The  $k$  point samples along each lattice vector are obtained from gamma-centered  $(24/M) \times (24/N) \times 1$  grids for surface slabs, and gamma-centered  $(24/M) \times (24/N) \times 4$  grids for bulk crystal, respectively; 24 points were chosen per reciprocal atom for that density provides a sufficient balance between computational efficiency and convergence of the formation energy.

Within the constraints posed by the surface slab cells, we created one  $M \times N$  surface for each slab cell defined in Table S.1 (ESI†), which we subsequently relaxed below a maximum atomic force of  $0.01 \text{ eV } \text{Å}^{-1}$ . We generated simulation cells containing on-surface adsorbed  $\text{O}^*$  configurations for 0.25 ML, 0.33 ML, and 1.0 ML coverages, using  $4 \times 4$ ,  $\sqrt[3]{3} \times \sqrt[3]{3}$  and  $4 \times 4$  unit cells respectively. For these on-surface  $\text{O}^*$  configurations, all oxygen atoms are placed strictly above the top layer of uncorrugated Pt(111) surfaces, and distributed as equidistantly as possible. In these cells, no Pt atoms were lifted from the top layer of the Pt(111) surface.

We generated initial  $\text{PtO}_2$  stripe supercells (1 ML) from the  $2 \times 4$  slab cell by lifting one surface row or a grid consisting of eight atom-wide triangles, respectively, and suspending them using oxygen atoms following the suggestions by Van Spronsen *et al.*<sup>17</sup> From these structures we created cells containing dilated stripes (stripe/surface ratios: 2/3, 3/4, 4/5, 5/6, 6/7, 5/8, 7/8, 8/9, 9/10) by removing one  $\text{PtO}_2$  unit from each stripe supercell, and scaling the remaining  $\text{PtO}_2$  segments along the long surface vector to close the gap left behind. For the stripe cells, Pt (in the oxide)-to-Pt (in the surface) ratios vary from 2/3 to 9/10. To more broadly sample low-energy stripe structures, platinum and oxygen coordinates in initial  $\text{PtO}_2$  stripes were either shifted systematically relative to the surface (by either half or a quarter of bulk interatomic spacing) or randomly displaced, and oxygen atoms were selectively moved to obtain geometries containing low-coordinated Pt sites in specific stripe segments. This process is illustrated in Fig. 2.

Resulting stripe structures and on-surface  $\text{O}^*$  configurations were relaxed, and vibrational analysis was performed on the lowest DFT energy configurations pertaining to individual stripe  $\text{PtO}_2/\text{Pt}(\text{displaced})$  ratios. These computations are detailed in the ESI.† Relaxed geometries of the 2/3, 3/4, 7/8, and 1/1 dilated  $\text{PtO}_2$  stripes are shown in Fig. 3. We calculated Helmholtz free energies for Pt(111) and oxidized Pt(111) surface slabs using the methods listed in Section 1 of the ESI.†

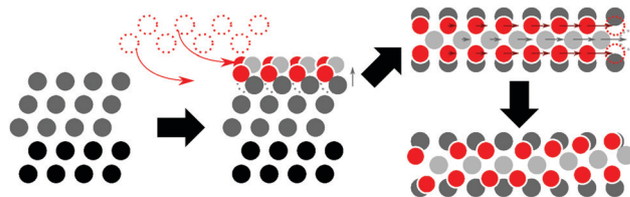


Fig. 2 Initial geometry production process, from left to right: (side view) surface optimization, formation of full coverage  $\text{PtO}_2$  stripe by lifting one surface row, (top view) omission of one  $\text{PtO}_2$  unit from an  $\text{PtO}_2$  covered surface supercell, and rearrangement of Pt and O atoms (illustrating final structure after both initial manual displacement/dilation and DFT geometry relaxation).

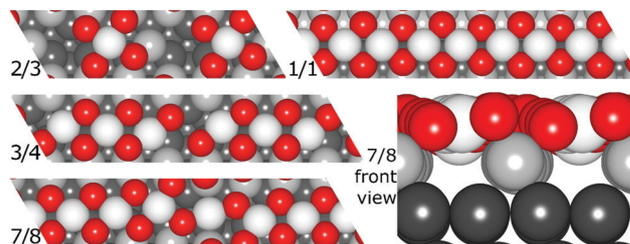


Fig. 3 Relaxed  $\text{PtO}_2$  stripe structures in top and front views. Platinum atoms are gray (darker gray is deeper into the slab), and oxygen atoms are red and relatively smaller.

## 3. Results and discussion

### 3.1 Results without vibrational energy

In all geometry optimizations (both in- and excluding vibrational energy), the most stable  $\text{PtO}_2$  stripes contain stretches of nearly rectangular planar 4-coordinated stripe Pt atoms with surrounding oxygen atoms slightly misaligned with respect to the surface vectors, alternating with single 2-coordinated stripe Pt atoms as shown for the 7/8 structure in Fig. 3. This holds for all structures ranging from 4/5 to 8/9. Fig. 4 plots stripe energies against individual and mean Pt–Pt spacing (excluding stripe vibrations), and shows that the formation energies are directly related to the degree of lattice mismatch. This assumption is reasonable given there are no apparent phase changes in either the  $\text{PtO}_2$  stripes or in respective decorated surfaces. Excluding vibrations, the optimum  $\text{PtO}_2/\text{Pt}(\text{displaced})$  ratio lies between 7/8 and 8/9, judging from a least-squares quadratic fit of the four lowest-energy points (see Fig. 4). In general, the shorter Pt–Pt distances in the four most stable stripes cluster between the Pt–Pt distance in bulk  $\alpha\text{-PtO}_2$  and that of the shorter Pt–Pt spacing in bulk  $\beta\text{-PtO}_2$ , while the longer Pt–Pt distances are close to that of longer Pt–Pt spacing in bulk  $\beta\text{-PtO}_2$ . The optimum Pt–Pt spacing appears to be 3.20 Å, while for 7/8 and for 8/9, they are 3.22 Å and 3.17 Å, respectively ( $R^2 = 0.930$ ). The average Pt–Pt distances in the 7/8 and 8/9 stripes are relatively close to those in bulk  $\alpha\text{-PtO}_2$ , which is the most stable bulk  $\text{PtO}_2$  phase. The relation between  $\text{PtO}_2/\text{Pt}(\text{displaced})$  ratios and stripe thermodynamics including vibrational energies will be discussed later in this paper.

The most stable stripe structures found in our calculations, *i.e.* the 6/7, 7/8, and 8/9 stripes, correspond closely to the



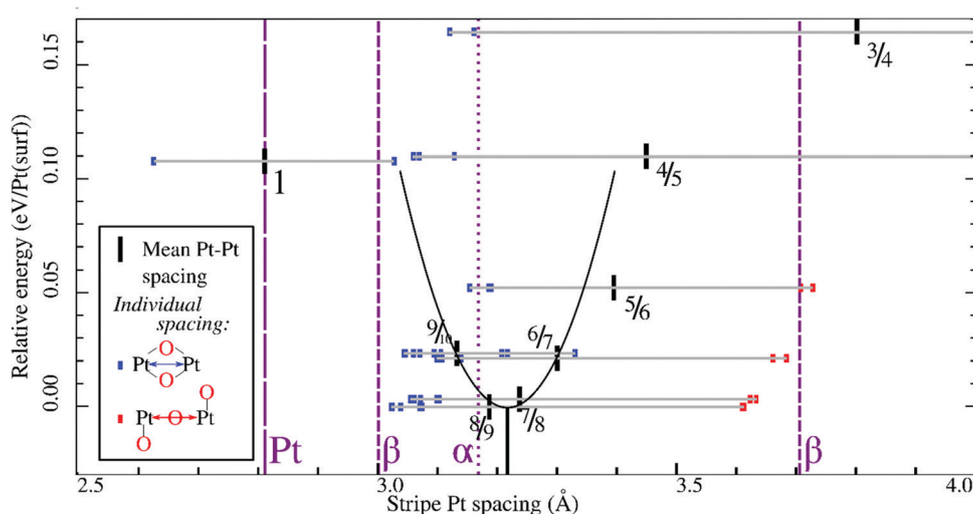


Fig. 4 Relative stripe formation energies (per surface platinum atom, without vibrational contribution to the energy) versus individual Pt–Pt distances (in the stripe) and mean spacings, including a quadratic fit of mean spacings near the energy minimum (marked by a descender from the apex of the parabola) and bulk Pt–Pt spacings for Pt,  $\alpha$ -PtO<sub>2</sub> and  $\beta$ -PtO<sub>2</sub>; legend in inset.

spokewheel structures observed by Van Spronsen *et al.* There is a close correlation between the Pt–Pt atom spacing in the stripe and the formation energy. The correspondence of most stable Pt–Pt spacing with those in most stable bulk structures indicates the physics underlying Pt–O bulk and stripe oxide interactions are remarkably similar; hence, such structures may aid bulk formation. Taken together, both observations suggest that the degree to which Pt–Pt or Pt–O bonds can relax dominates stripe thermodynamics, and suggests that removal of platinum from the stripe relative to the density of Pt in the underlying surface is favorable, allowing the stripe to dilate.

Energies of formation for each individual stripe – excluding vibrational free energies of the PtO<sub>2</sub> units and the underlying surfaces – are plotted against the O<sub>2</sub> chemical potential and the electrochemical potential relative to the computational hydrogen electrode ( $E_0 - E_{\text{SHE}}$ ) in Fig. 5. The plot includes on-surface O\* configurations, in which no platinum is lifted from the surface as described previously. For a Pt (in stripe)–Pt (in surface) ratio of 1, all original surface atoms lifted from the surface remain in the stripe. For other PtO<sub>2</sub>/Pt(displaced) ratios, some platinum atoms are instead expelled from the area in which a stripe is being formed, and migrate on top of the original surface. Hence, a plot of the free energy of formation against the chemical potential of the expelled Pt atom(s) is included in Figure S.2 in the ESI.†

We can infer from Fig. 5 that forming non-dilated stripes, *i.e.* surface oxidation without emitting Pt atoms from the surface, is stable only at high oxygen chemical potential or high electrochemical potential. In contrast, 5/6, 6/7, 7/8, 8/9 dilated stripes are significantly more stable than either the Pt(111) surface or stripes with other PtO<sub>2</sub>/Pt(displaced) ratios at 1 bar O<sub>2</sub> pressure and standard temperature. This means there is a clear (electronic) driving force towards lower PtO<sub>2</sub>/Pt(displaced) ratios in general. Stripe formation is also more favorable than high-coverage (1 ML) on-surface oxygen adsorption, as shown in Fig. 5.

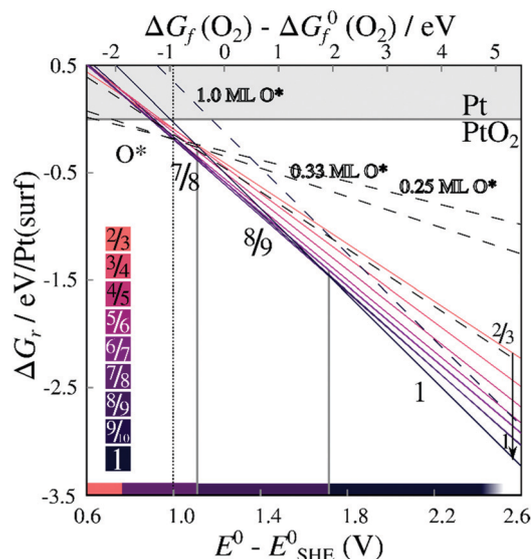


Fig. 5 Energies of formation, excluding the contribution of the vibrational modes, for oxygenated Pt surfaces with respect to the O<sub>2</sub> chemical potential and the electrochemical potentials. Lines are labeled by the corresponding stripe PtO<sub>2</sub>/Pt(displaced) ratios M/N or coverages  $\times$  ML O\*. Contiguous lines refer to PtO<sub>2</sub> stripes; colors correspond to PtO<sub>2</sub>/Pt(displaced) ratios as defined in the color bars (bottom-left inset). Dashed lines refer to on-surface O\*. Both figures are based on DFT binding energies for Pt bulk (see Table S.5 in the ESI).†

### 3.2 Results with vibrational energy

As the internal energy difference between 7/8 and 8/9 stripes is  $\sim 2$  meV/Pt(surf), and the difference between 6/7 or 9/10 and the 8/9 stripe is  $\sim 20$  meV/Pt(surf), vibrational energies might influence the trend as for which stripe is most stable.

Our oxygen coverages are high, entire rows of Pt atoms are lifted out of the surface, there are strong adsorbate–surface interactions, and minute energy differences between



individual configurations. A traditional adsorbate-only vibrational analysis (where the surface vibrational modes are assumed un-perturbed by the adsorbate) may hence lead to incorrect approximations of vibrational enthalpies and entropies. Hence we attempt to subsume standard adsorbate-only vibrational analysis into a adsorbate-bulk vibrational analysis. This methodology is based on the assumption that the amplitudes of forces exerted between two atoms in a lattice decay quickly enough with increasing distance between them. This enables us to describe their vibrational modes using only short-range interactions where necessary, including between different simulation models.

Using this method, we may stack bulk layers under the surface of interest, aiming to converge to a vibrational energy which accurately represents both surface, bulk, and surface-bulk interactions. While selecting the atoms taking part in this vibrational analysis, one must take into account to which degree the vibrational data correctly represents the atoms affected by the chemical reaction under study. In the context of the stripe-oxygenated Pt surfaces we are studying, four categories of atoms can be defined: stripe atoms, the (remaining) top surface atoms, the first underlying layer and the underlying layers or bulk. For pristine surfaces, the former two correspond to the top surface layer, while the second corresponds to the first subsurface layer.

We decided to analyze 5/6, 6/7, 7/8, and 8/9 stripes as their internal energies are closest to the optimum and, hence, they are the structures of which the relative energies may be affected most by vibrational energy contributions (see Table S.6, ESI†). In calculating the vibrational contribution to the free energy, all vibrational modes were calculated by  $\pm 0.02 \text{ \AA}$  single atom displacements along Cartesian unit vectors, of which one is parallel to the short edge of the surface, and the other is perpendicular to the surface. The forces exerted by these stripe/surface/subsurface atoms on each other and on the first two underlying layers are compiled into a group of Hessian matrix elements – note that we only have information on stripe-to-underlying layer coupling rather than the other way around – which is combined with a large Hessian matrix describing the force gradients of a set of bulk layers with the two underlying layers mentioned above and (if applicable) two layers above and below the current bulk layer. This matrix is illustrated in Fig. S.1 in the ESI†; some further practical details are listed in ESI Section 4.†

When including only the first (corrugated) platinum layer and the oxygen atoms, vibrational contributions destabilize stable stripes by  $\sim 0.35 \text{ eV}$  per Pt(surf) relative to their respective pristine surfaces. Including three bulk layers, for which the vibrational energy converges while remaining computationally reliable as shown in ESI Section 4,† causes a further increase of vibrational free energy for less dilated structures, and a decrease in energy for the more dilated structures. Adding two additional bulk layers beyond the first layer results in vibrational free energy changes ranging from 10–40 meV per Pt(surf), depending on the stripe PtO<sub>2</sub>/Pt(displaced) ratio (see Table S.4 in the ESI†). The resulting vibrational energies depend monotonously on PtO<sub>2</sub>/Pt(displaced) ratios, indicating that stripes with PtO<sub>2</sub>/Pt(displaced) ratios between those

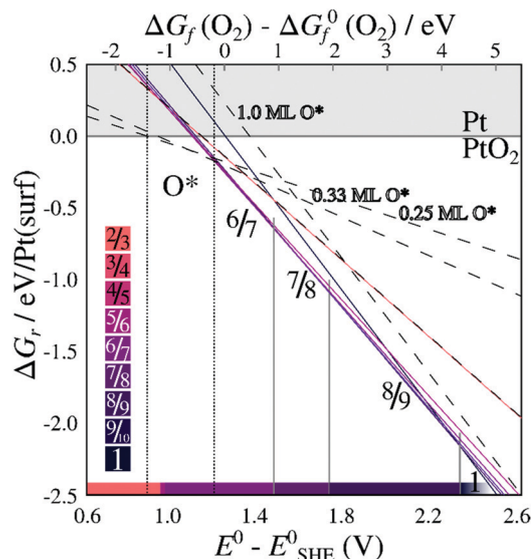


Fig. 6 Energies of formation, including the contribution of vibrational modes, for PtO<sub>2</sub> superstructures with respect to O<sub>2</sub> chemical potentials (a) and electrochemical potentials relative to CHE (b), labeled by PtO<sub>2</sub>/Pt(displaced) ratios M/N. Dashed lines refer to on-surface O\*. Shades correspond to PtO<sub>2</sub>/Pt(displaced) ratios as defined in the color bars. All energies are based on DFT binding energies for Pt bulk (see Table S.5, ESI†).

investigated have intermediate vibrational energies. After adding the vibrational contributions, Fig. 6 shows that the 6/7 stripe is more stable than the 7/8 stripe by 9 meV per Pt(surf), while the 7/8 stripe is more stable than the 8/9 stripe by 13 meV per Pt(surf). This suggests the most stable PtO<sub>2</sub>/Pt(displaced) ratio to be close to 6/7 or 7/8, which still corresponds closely to the experimentally observed data.

Similarly, Fig. 7 shows that the minimum of the quadratic fit of the stripe energies *versus* the Pt–Pt spacing when including vibrations, shifts from 3.20 Å at 3.28 Å, close to the 6/7 structure spacing, while maintaining a clear relation between energies and Pt–Pt spacing ( $R^2 = 0.997$ ). Although the 6/7 stripe is most stable in our calculations including vibrational energy corrections, the 7/8 structure is less than 0.5  $k_B T$ /Pt(surf) higher in energy, suggesting that even if there are no additional entropy terms favoring 7/8 stripes, significant populations of 7/8 stripe structure domains can be found on an oxidized Pt(111) surface. Note that the spoked wheel structures include nodes and non-reconstructed domains adjacent to stripe segments, which may affect the optimal Pt–Pt spacing. The total vibrational correction raises the predicted onset potentials for 0.25 ML oxygen adsorption to 0.9 V, while shifting the onset potential for stripe formation to  $\sim 1.2 \text{ V}$ . This onset potential lies within the expected range for initial oxide formation as discussed above.

The model defined above does not take into account repulsion between adjacent stripes. Leaving out every second stripe in a double-spaced unit cell stabilizes each individual stripe by 0.248 eV per Pt(surf) or less (see Table S.6, ESI†). Repulsion between stripes appears to be significant, but is smaller than the binding energy of adjacent stripes. This means the adjacent stripe model is representative of real stripe coverages.



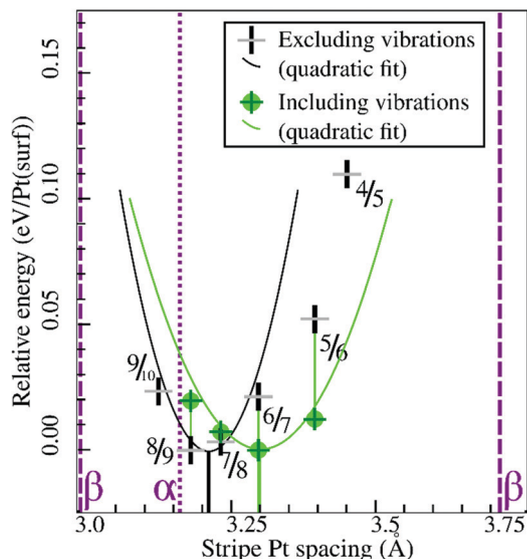


Fig. 7 Comparison of selected stripe energies with and without vibrational energy corrections and their respective quadratic fits; legend in inset.

The onset (electro)chemical potential for the formation of atomic oxygen and surface oxide stripes compares well with oxygen adsorption/oxide formation observed experimentally. Before the formation of the  $\text{PtO}_2$  film, and in the absence of specific adsorption of  $\text{Pt}(111)$  (*i.e.* in perchloric acid), there is a voltammetric peak at 1.10 V (with an onset at  $\sim 1.05$  V) typically attributed to surface atomic oxygen (but see studies by Huang *et al.*<sup>28,29</sup>).<sup>30</sup> Importantly, the formation of this surface species does not involve any major structural rearrangement of the platinum surface atoms. Using *in situ* X-ray spectroscopy, Liu *et al.* have shown that higher densities of on-surface  $\text{O}^*$  on a  $\text{Pt}(111)$  electrode, which they find near 0.8 to 1.0 V, lead to partial perpendicular displacement of Pt atoms from the top surface layer. This suggests that higher coverages may cause Pt atoms to be lifted from the surface, which is a first step towards reconstruction into  $\text{PtO}_2$  stripes. Further oxidation of  $\text{Pt}(111)$  leads to a new voltammetric feature with an onset at *ca.* 1.25 V. *In situ* Raman spectroscopic measurements of  $\text{Pt}(111)$  by Huang *et al.* have shown that this feature is accompanied by the formation of an  $\alpha$ - $\text{PtO}_2$  film.<sup>28</sup> Once this film is formed and subsequently reduced, the  $\text{Pt}(111)$  structure changes and develops defect sites,<sup>11</sup> in agreement with the idea that the formation of the oxide film (or oxide stripes) leads to an expulsion of Pt atoms, which do not return to their original position after oxide reduction. These observations are in excellent agreement with the predictions of Fig. 5 and 6. The relative stability of stripe structures also corresponds well with the findings by Van Spronsen *et al.* mentioned in the Introduction,<sup>17</sup> who observed 7/8 dilated spokewheel segments in SEM images. Van Spronsen's stripes were created at 2.2 bar and 535 K, which is equivalent to an oxygen potential shift of  $-0.488$  eV or an electrode potential at 300 K of 1.11 V. The  $\text{PtO}_2/\text{Pt}(\text{displaced})$  ratio of the stripes determines the number of Pt atoms expelled from the Pt surface during a single oxidation–reduction cycle (this number is larger with a smaller  $\text{PtO}_2/\text{Pt}(\text{displaced})$  ratio). The larger this

number, the faster the surface will roughen during multiple oxidation–reduction cycles (for a detailed model of this surface roughening, see Rost *et al.*<sup>31</sup>). Pt from the platinum oxide may also dissolve into the electrolyte during the reductive scan, leading to platinum corrosion; however, this Pt dissolution is expected to be more dominant when a thicker oxide is formed.<sup>6,32</sup>

When comparing the stripe model and our corresponding results to the electrochemical place-exchange studies, we find that oxide chains are plausible products of  $\text{Pt}(111)$  oxidation under electrochemical conditions, compatible with X-ray diffraction analysis by Drnec *et al.*<sup>4</sup> Although the structures we studied exhibit long-range order, which runs contrary to the conclusions of Drnec *et al.*, the observation of disordered and isolated spokewheels by Van Spronsen *et al.* in early  $\text{Pt}(111)$  oxidation is consistent with both our dilated stripe structures and findings by Drnec *et al.*<sup>4,17</sup> Pt atoms are suspended by more than one oxygen atom; in fact, two oxygen atoms per Pt atom are adsorbed above surface platinum atoms. This agrees with electron distributions found by Drnec *et al.*,<sup>4</sup> yet contradicts their assumed place exchange model in which one O atom displaces one Pt atom and is incorporated into the surface.<sup>4,22</sup> Additionally, even though they consider their data based on the traditional concept of place-exchange, their observation of two discrete zones of oxygen atoms between two layers of Pt atoms, rather than being embedded within a surface Pt layer,<sup>4</sup> suggests no oxygen is incorporated into the surface during place exchange, and that no oxygen is found above the Pt atoms. This contradicts the alternately buckled stripe model as proposed by Holby *et al.*<sup>15</sup> The stripes found through our relaxation had O atoms and lifted Pt atoms elevated roughly the same amount above the original  $\text{Pt}(111)$  surface, which seems to disagree with the  $\text{Pt}(\text{surf})\text{--O--Pt}$  layering found by Drnec *et al.*; however, we did not include dipole stabilization, which does happen in aqueous environments and may provide a driving force for lifted Pt atoms to pucker out of the stripe. Altogether, we find that non-buckled stripe – or spokewheel – formation is a more likely process of electrochemical oxidation of  $\text{Pt}(111)$  than either buckled stripes or traditional place exchange.

## 4. Conclusion

Understanding the driving forces for stripe structure formation allows us to better understand Pt catalyst structure evolution under oxidative and electrochemical conditions. We have found a clear correlation between increasingly oxidative potentials and the surface phase transition from adsorbed oxygen to the  $\text{PtO}_2$  stripes, and the thermodynamics of stripe formation suggests that the optimum average Pt–Pt spacing, and the spacing observed in spokewheel  $\text{PtO}_2$  structures by Van Spronsen *et al.*, is close to that of both bulk  $\text{PtO}_2$  species. This results in expulsion of platinum atoms relative to the Pt density in the surface layer, allowing the dilation of the stripe to occur. Formation of these  $\text{PtO}_2$  stripes is plausible at electrochemical potentials similar to those suggested in literature for oxygen adsorption and surface oxide formation. This indicates we have found a plausible early oxidation structure which not only closely corresponds to the stripe hypothesis by



van Spronsen *et al.* and their observations in gas phase, but also provide a good description of the electrochemical oxidation of the Pt(111) surface. The platinum oxide structures we have found are a key towards understanding the roughening observed during oxidation–reduction cycles, as well as to the formation of thicker oxide layers on Pt(111).

Our findings show that accurate computational modeling of electrochemical oxidation reactions on Pt(111) is possible, including active sites specific to PtO<sub>2</sub> stripes. This knowledge is equally applicable to Pt(111) surfaces under elevated O<sub>2</sub> pressure. Hence, future computational work may aid the design of more active real-life industrial platinum(-like) catalysts.

## Conflicts of interest

There are no conflicts to declare.

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