



Violations of Löwenstein's rule in zeolites†

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Zeolites, microporous aluminosilicates, are amongst the most widely used catalysts in the petrochemical industry. Zeolite catalytic functionality is influenced by the location of tetrahedral alumina and associated counter-cations in the aluminosilicate framework, yet little is definitively known about the factors that govern the framework aluminium arrangement. It is generally accepted that all zeolites obey Löwenstein's rule of "aluminium avoidance", and that $-Al-O-Al-$ bond formation is forbidden. Here, we describe an unprecedented screening of aluminium distribution in catalytically active zeolite SSZ-13 (CHA) in both its protonated and sodium-containing forms, H-SSZ-13 and Na-SSZ-13, using density functional theory (DFT). We predict violations of Löwenstein's rule in high and low silica H-SSZ-13 and other protonated frameworks considered in this investigation: H-LTA, H-RHO, H-ABW and H-MOR. The synthetic realisation of these zeolites could spur the development of new catalytic routes and materials, and the optimisation of existing zeolite catalysts.

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Introduction

The use of zeolite catalysts in petrochemistry entirely revolutionised the industry over half a century ago. Since then, zeolites have become the workhorses of petrochemical processing, and are used extensively throughout the petrochemical industry.¹ Now, at a time of fast depleting traditional fuel sources and increasing toxic gas emission, zeolite catalysts are at the forefront of the development of 'green' alternatives to long-established petrochemical processes.² Green processes must operate at optimum efficiency³ and for catalytic processes this requires the structural elucidation of existing catalytic materials. Unequivocally resolving a material's structure can expedite the identification of structure–activity relationships, which in turn, can accelerate the development of material specific design rules that are necessary for the rational design of new, more sustainable and efficient catalysts.

It is well understood that zeolite catalytic functionality originates from negatively charged tetrahedral units of AlO_4 distributed throughout the aluminosilicate framework, and their associated charge-compensating cations located in nearby pores. Yet, despite major recent advances in experimental techniques,⁴ at present it is not possible to determine the absolute position of framework aluminium or accompanying counter-cations exactly. Furthermore, there are currently no established design rules that can be applied to infer framework aluminium's preferred and precise position. However, Löwenstein's rule⁵ of "aluminium avoidance" is commonly assumed; this states that on forming

the aluminosilicate zeolite framework there is a disinclination for tetrahedral units of alumina to exist adjacent to one another, forbidding formation of $-Al-O-Al-$ linkages, and restricting the minimum Si/Al ratio of any zeolite to unity.

Löwenstein's rule was conceptualised in 1954 and since then there have been few suggestions that violations of the rule are possible.^{6–13} Indeed, the scientific literature reports that "aluminium avoidance" is observed in zeolites almost without exception.^{5,14,15} Löwenstein's rule has hence become a fundamental law of zeolite science, and the possibility of non-Löwensteinian ordered zeolites is often not considered. This is true of most theoretical studies where the omission of non-Löwensteinian frameworks is considered a simple way to reduce unnecessary computational expense by shrinking the number of potential configurations.^{16–23} However, recent advances in supercomputing services and the development of increasingly efficient codes, mean it is now tractable to evaluate both Löwensteinian and non-Löwensteinian frameworks accurately through quantum mechanical approaches.

Characterisation techniques, such as X-ray diffraction, are currently unable to distinguish between framework silicon and catalytically important aluminium distributed throughout the zeolite, except in rare cases where there is strict ordering, such as in Goosecreekite.²⁴ On the other hand, atomistic modelling techniques can be used as a tool to provide insight into the most probable location of framework aluminium in real zeolites. Using framework crystallographic data for a particular zeolite, quantum mechanical methods can unambiguously resolve the most energetically favourable distribution of both Si and Al. However, there is a further complication. It is well documented that the framework aluminium distribution of a given zeolite is highly dependent on the synthesis details.²⁵ The Si/Al ratio of the initial synthesis gel, synthesis temperature, reaction times,

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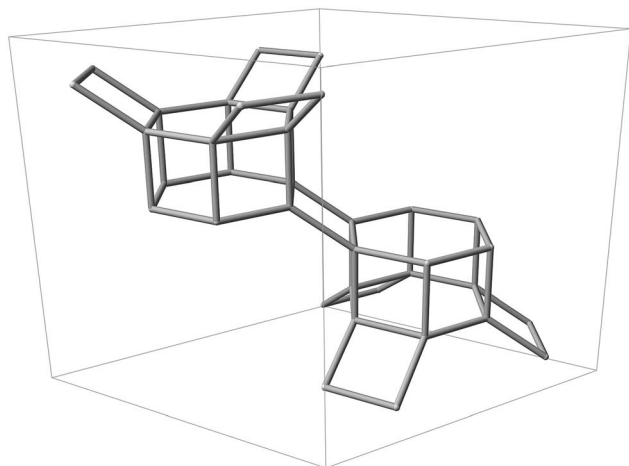


Fig. 1 Silicon backbone of the CHA framework contained within a single hexagonal unit cell, displaying two D6R units and adjoining tilted 4-rings.

counter-cation identity and kinetic factors may all cause differences in the final structural chemistry, and hence catalytic activity of the resultant framework. Furthermore, Perea *et al.* and Schmidt *et al.* recently showed that for a zeolite sample at a given Si/Al ratio, aluminium is inhomogeneously distributed throughout the framework,^{4,26} supporting earlier work by von Ballmoos and Meier, that reported the presence of Al zoning in single crystals.²⁷ Commercial zeolites are typically synthesised using alkali metal cations as the charge compensating species, and facile ion-exchange techniques may be used post-synthesis to replace the metal cation with a proton, hence generating Brønsted acid O–H sites proximal to the location of aluminium. A key open question, which we partially address here, is whether the nature of the counter-cation affects the positioning of aluminium. However, the broader question is whether there is a clear thermodynamic incentive to form ordered or partially ordered frameworks and whether the position of aluminium can be predicted. Here, we present results obtained for the active small-pore zeolite catalyst SSZ-13, which displays a CHA-type framework topology.²⁸ The CHA framework (Fig. 1) is made up of layers of hexagonally arranged double 6-ring (D6R) units connected by tilted 4-rings, giving rise to a characteristic ‘cha’ cavity accessible through an 8-ring pore system.²⁸ Using periodic density functional theory (DFT) implemented in the program CP2K^{29–31} we investigate all possible arrangements of framework aluminium, including non-Löwensteinian distributions, surveying the aluminium distribution of SSZ-13 at Si/Al ratios of 17, 11 and 8, in both Brønsted acidic H-SSZ-13 and the as-synthesised Na-SSZ-13. To our knowledge this is the only exhaustive study of zeolite framework aluminium distribution with different Si/Al ratios at this fully periodic quantum mechanical level of theory.

Results

A decisive variable in optimising catalytic activity is the Si/Al ratio as this dictates the density of charge compensating

species, such as acidic sites. We compare three quite distinct Si/Al ratios to probe how the Si/Al ratio affects aluminium ordering.

High silica SSZ-13 Si/Al = 17

A single hexagonal CHA unit cell contains 36 symmetry equivalent T-sites; in order to methodically explore all of the possible configurations of 2 Al per unit cell, corresponding to a Si/Al ratio of 17, a single aluminium atom, Al1, was substituted into an arbitrary T-site. Maintaining Al1's position, a second aluminium, Al2, was sequentially introduced into the remaining 35 T-sites. To maintain charge-neutrality, each individual aluminium substitution requires charge compensation by a cationic moiety. For H-SSZ-13, each cationic proton may reside at one of four oxygen sites at the apices of the alumina tetrahedra, yielding four potential topologically inequivalent Brønsted acid O–H sites per Al substitution, and hence a total of 560 unique combinations of 2 Al per unit cell. We used the periodic DFT method (at the PBE level)³² to fully optimize each individual framework model to equilibrium density; the resulting data is shown in Fig. 2, where the relative energy per unit cell (U.C.) (with respect to the average total energy) is given as a function of framework aluminium separation.

Assuming Löwenstein's rule⁵ is valid, and the principle of aluminium avoidance is adhered to, we would expect the highest energy SSZ-13 structures to be those containing aluminium atoms at separations equivalent to that of a “forbidden” –Al–O–Al– linkage, and structures with larger aluminium separations to become increasingly more stable, in accordance with Dempsey's rule³³ (a less sophisticated rule which states, on the basis of electrostatics, that negatively charged alumina units are inclined to be positioned as far away from one another as possible).

As predicted from Löwenstein's rule, the highest energy configurations for both H-SSZ-13 and Na-SSZ-13 are those containing adjacent aluminium atoms, with a separation of approximately 3 Å (Fig. 2). However, beyond this distance the relative energy landscapes for the two forms of the zeolite become dramatically different.

In accordance with Löwenstein's rule, and what has already been widely observed in sodium-containing zeolites, the Na-SSZ-13 global minimum (Fig. 3a) contains aluminium pairs as next-next-nearest neighbours (NNNN),¹⁶ and there is a +44.3 kJ mol^{−1} per U.C. energy penalty for forming the most favourable non-Löwensteinian (NL) structure. Ignoring barriers, the penalty to form a NL structure is at least 10kT, where k is the Boltzmann constant (assuming a typical synthesis temperature of ~450 K), which suggests –Al–O–Al– linkages are very unfavourable in Na-SSZ-13. In the global minimum structure, the aluminium ions are separated by a distance of 6.18 Å, and their associated Na⁺ cations reside at the parameters of the 8-ring apertures of the ‘cha’ cavity. However, the H-SSZ-13 global minimum structure (Fig. 3c) is remarkably different, containing adjacent aluminium ions along the edge of the 6-ring at a separation of 3.28 Å, violating Löwenstein's rule. In this structure the two associated protons, H1 and H2, are separated



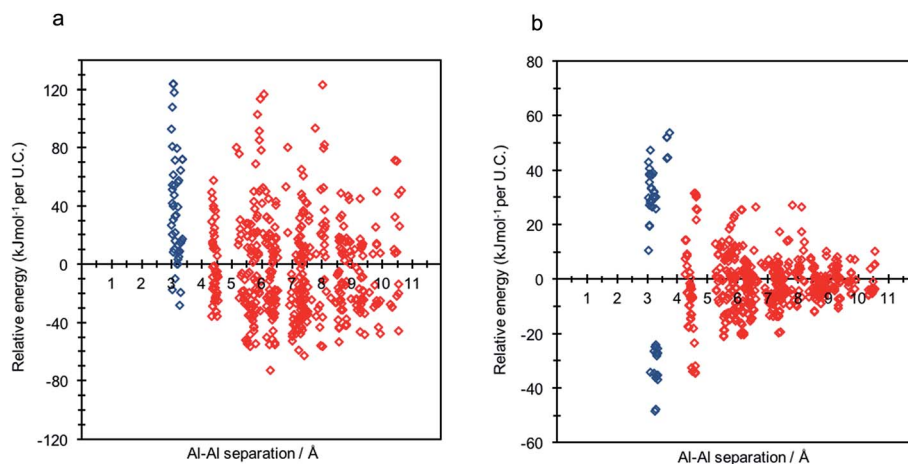


Fig. 2 Relative energy distribution (kJ mol^{-1} per U.C.) against framework aluminium separation for (a) Na-SSZ-13 and (b) H-SSZ-13. Frameworks possessing non-Löwensteinian (NL) ordered aluminium atoms ($-\text{Al}-\text{O}-\text{Al}-$) are shown in blue.

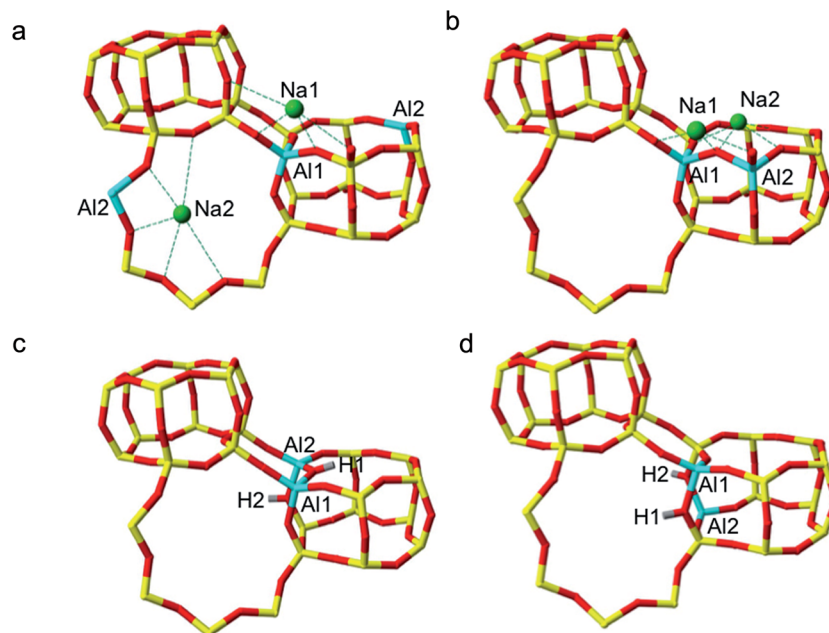


Fig. 3 Global minima L and NL 2 Al per U.C. SSZ-13 structures predicted by DFT. (a) Global minimum Na-SSZ-13 structure, with Löwensteinian ordered aluminium atoms at the NNNN position, (b) lowest energy NL Na-SSZ-13 structure, (c) Global minimum NL H-SSZ-13 structure, with protons oriented *trans* to one another (d) lowest energy Löwensteinian ordered structure with Al at NNN position. Where; silicon (yellow), oxygen (red), aluminium (blue), sodium (green), hydrogen (grey).

by 4.36 Å and arranged *trans* to one another; H1, which mediates the aluminium ions, is directed into the plane of the 6-ring, and H2, positioned at the connecting edge of the D6R, is oriented away from H1, and directed into the 8-ring window of the 'cha' cavity. The most stable Löwensteinian (L) structure (Fig. 3d) contains aluminium ions as next-nearest neighbours (NNN), at a 'non-Dempsey' separation of 4.60 Å, with both protons directed into different 8-ring windows of the 'cha' cavity. The energy penalty for forming the L structure rather than the NL structure is $+14.2 \text{ kJ mol}^{-1}$ per U.C., approximately one third of the energy difference between the global minimum NL/L structures for Na-SSZ-13. $\Delta E(\text{NL}_{\text{global minimum}} - \text{L}_{\text{global minimum}})$ for

Na-SSZ-13 is $+44.3 \text{ kJ mol}^{-1}$ per U.C., whilst $\Delta E(\text{NL}_{\text{global minimum}} - \text{L}_{\text{global minimum}})$ for H-SSZ-13 is $-14.2 \text{ kJ mol}^{-1}$ per U.C., indicating a strong enthalpic incentive for Löwensteinian configurations when Na^+ is the charge compensating cation and a modest enthalpic incentive to adopt non-Löwensteinian linkages when the charge compensator is a proton. Free energy calculations that include the vibrational entropy contributions to the energy show that the relative stability of the L and NL H-SSZ-13 configurations is maintained beyond typical synthesis temperatures (see ESI†), demonstrating a clear thermodynamic preference for adopting non-Löwensteinian structures for the proton compensated structure, a result that clearly conflicts with accepted wisdom.



Furthermore, the NL H-SSZ-13 global minimum is not unique and seven other NL ordered frameworks (excluding the global minimum structure), all of which contain proton arrangements similar to those displayed in the NL global minimum structure (Fig. 3c), are more stable than the global minimum L H-SSZ-13 structure.

To test our unexpected H-SSZ-13 result we further investigated the lowest energy structures using the higher level hybrid functional PBE0,^{34,35} and van der Waals corrected functionals, vDW-DF2 and PBE + D3. The relative energies calculated using these methods show good correlation with those calculated using the standard PBE functional, confirming the robustness of our predictions. The results for these calculations, presented as energy correlation plots, are included in the ESI† (S2 and S3).

Low silica SSZ-13 Si/Al < 17

Exploring SSZ-13 with lower Si/Al ratios becomes increasingly complicated with each introduction of additional aluminium. To avoid calculating the prohibitively large number of combinations of 3 Al per unit cell SSZ-13 (Si/Al = 11), we employed a method of stepwise aluminium incorporation. In this approach, the NL and L ordered 2 Al per unit cell global minima according to the prior DFT (Fig. 3c and d) were used as the initial structures. A single Al, Al₃, was sequentially introduced into each of the remaining silica T-sites of both structures, and the appropriate counter-cation positioned at one of the four apical oxygen sites. For each NL and L initial global minimum structure, a total of 136 distinct framework arrangements were created for Na-SSZ-13 and H-SSZ-13, respectively (544 calculations in total).

Each structure was optimised and the NL and L H-SSZ-13 initial configurations gave the same 3 Al per unit cell global minimum structure. The structure, Fig. 4a, contains a chain of three oxygen linked aluminium atoms, [O–Al–O]₃, with each charge-compensating proton located at a bridging oxygen and arranged *trans* to its neighbour(s). Once again, Na-SSZ-13 did not follow the same trend as H-SSZ-13, where each initial structure yielded different global minimum structures; the Löwensteinian structure favouring the third aluminium at the NNN position,¹⁹ and the non-Löwensteinian structure favouring

Al at the NNNN position. The corresponding figures for these structures can be found in the ESI (S6†).

Using the H-SSZ-13 (Si/Al = 11) global minimum as the new initial structure, we then proceeded to investigate 4 Al per unit cell, equivalent to a Si/Al ratio of 8. The global minimum structure, shown in Fig. 4b, contains a chain of four oxygen linked aluminium atoms arranged in a 4-ring, with protons arranged *trans* to one another. In the sodium form of this structure (S6†), the fourth Al resides in the NNN position, again in accordance with Löwenstein's rule, and what has already been documented for similar zeolites.¹⁹ All four sodium cations position themselves proximal to the aluminium ions, at the centre of both faces of the aluminium doped D6R unit and at the parameters of the proximal 8-rings. It appears that as the aluminium content of the zeolite is increased, the aluminium clusters into zones of concentrated –Al–O–Al–, this is contrary to the general belief that aluminium is reasonably well dispersed throughout the frameworks of real samples.^{4,26} It should be noted that these results cannot imply whether the minimum Si/Al ratio is 1.

Other zeolite framework types

To ascertain whether our unexpected findings manifest in other proton compensated zeolite frameworks, or are unique to CHA, we investigated a selection of framework-types by the same methods previously discussed. The selected frameworks, LTA, RHO and ABW, are shown in Fig. 5, and their corresponding densities are included in the ESI† (S7).²⁸

Each of the frameworks contain a single symmetry equivalent T-site, and exhibit contrasting densities and topologies to that of CHA. The least dense of the frameworks, LTA, has a highly controversial history regarding aluminium distribution at Si/Al ratios tending to 1, where previous work has suggested the existence of non-Löwensteinian (NL) linkages.^{6,7} Investigation of each framework (2 Al per unit cell) using DFT revealed that all three framework types possess NL global minimum structures in their protonated forms, and that the protons adopt the same '*trans*'-like orientation as seen for CHA. The data for each structure is shown in Fig. 6, and the corresponding global minimum framework structures for each framework type can be found in the ESI† (S8). The energy penalty for forming the L

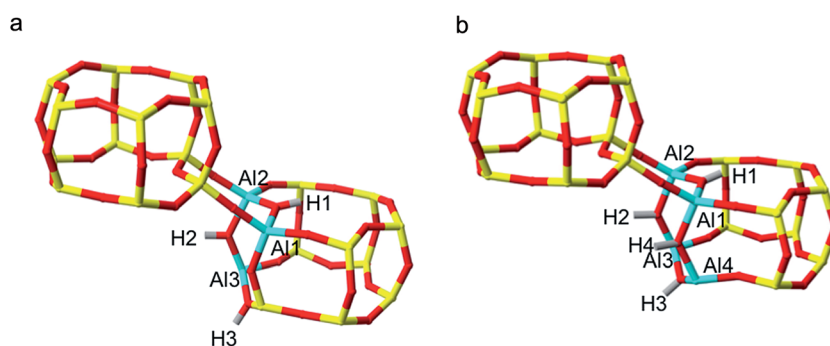


Fig. 4 (a) H-SSZ-13 3 Al per unit cell (Si/Al = 11) global minimum structure, containing a chain of –Al–O–Al–O–Al–. (b) H-SSZ-13 4 Al per unit cell (Si/Al = 8) global minimum structure, containing a chain of –Al–O–Al–O–Al–O–Al–.



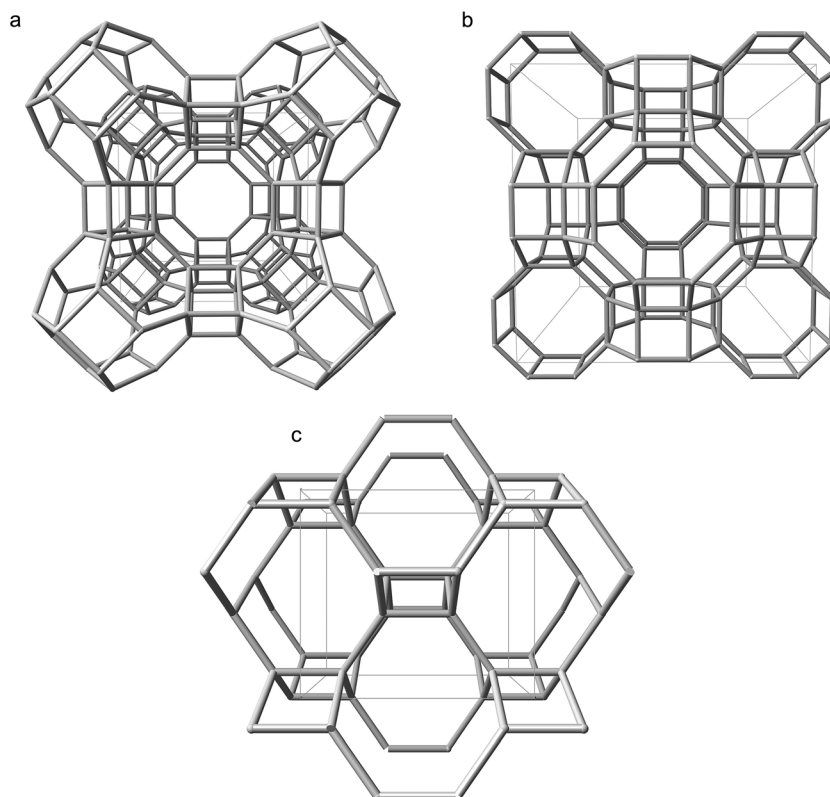


Fig. 5 Silicon backbones for the unit cells of additional zeolite frameworks considered in this work: (a) LTA viewed along [100] (b) RHO viewed along [100] (c) ABW viewed along [010].²⁸

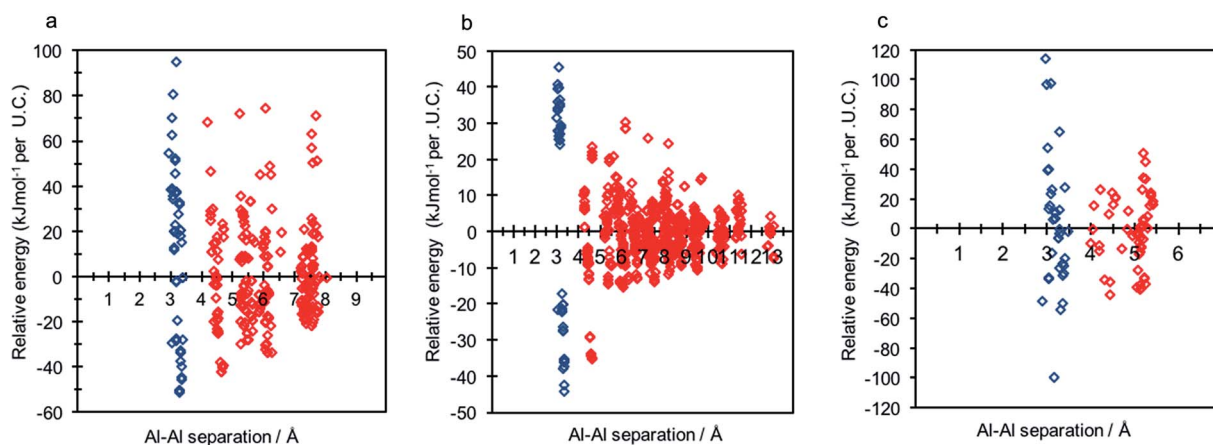


Fig. 6 Relative energy dispersion (kJ mol^{-1} per U.C.) against framework aluminium separation for (a) LTA (1 : 1 : 1) (b) RHO (1 : 1 : 1) (c) ABW (2 : 2 : 2). Frameworks possessing non-Löwensteinian (NL) ordered aluminium atoms ($-\text{Al}-\text{O}-\text{Al}-$) are shown in blue.

structure ($\Delta E(\text{NL}_{\text{global minimum}} - \text{L}_{\text{global minimum}})$) for high density H-ABW is $+55.7 \text{ kJ mol}^{-1}$ per U.C., $+14.2 \text{ kJ mol}^{-1}$ per U.C. for H-CHA, $+9.2 \text{ kJ mol}^{-1}$ per U.C. for H-RHO and $+8.3 \text{ kJ mol}^{-1}$ per U.C. for H-LTA, which correlates with their respective densities. These results suggest that NL linkages are more strongly preferred in denser zeolites, but even in LTA, which is one of the lowest density zeolites, the energy penalty for forming L structures is $\sim 2kT$ at typical synthesis temperatures ($\sim 450 \text{ K}$).

To discern whether the preference for NL ordering over L ordering can be extended to other ring systems we examined the protonated MOR framework. Due to the large number of symmetry inequivalent tetrahedral framework sites of the MOR framework, we augmented our approach to focus only on the thermodynamic stability of aluminium pairs as nearest neighbours compared to next-nearest neighbours for each of the four symmetry equivalent T-site present within the framework. Once again, the DFT data showed the NL ordering to be preferred over



the 'traditional' Löwensteinian ordering. For these structures $\Delta E(\text{NL}_{\text{global minimum}} - \text{L}_{\text{global minimum}}) = -16.1 \text{ kJ mol}^{-1}$, this value is consistent with the trend observed between density and preference for the formation of Al–O–Al. Further information about these calculations, results and the NL MOR global minimum structure are included in the ESI† (S9).

Discussion

This work provides evidence for non-Löwensteinian ordering in protonated zeolite frameworks, where there is a thermodynamic preference for Al³⁺ ions to exist adjacent to one another linked by a bridging hydroxyl moiety oriented 'trans' to its nearest neighbour proton. This prediction holds true across a range of different frameworks, and we have shown that in low silica frameworks there is a preference for the formation of discrete aluminium clusters. However, this is not the case for sodium-containing zeolites, where the global minimum structures are Löwensteinian ordered frameworks. In low silica sodium frameworks, next-nearest neighbour aluminium distribution is favoured, but next-next-nearest neighbour distributions are preferred with increasing aluminium content. Marked differences between the most thermodynamically stable aluminium distributions of protonated and sodium-containing zeolites have been discerned demonstrating the influence of counter-cation identity on framework aluminium location. In addition, Dempsey's rule³⁶ is violated in the global minimum structures of all investigated frameworks.

The literature contains several reports of violations of Dempsey's rule in zeolites,^{16,19,37} and it has been established that non-covalent interactions, present between framework oxygen and extra-framework cations, may distort aluminium distributions away from true Dempsey ordering.³⁷ On close inspection, violations of Dempsey's rule in Na-SSZ-13 can be rationalised by simple electrostatics. As shown in the global minimum structure for 2 Al per unit cell (Fig. 3a), there is a preference for Na⁺ cations to maximise their coordination with framework oxygen whilst minimising unfavourable cation–cation interactions – as illustrated by the collection of unusually high energy structures (with aluminium separations of 5.80–8.20 Å) in Fig. 2a, all of which contain Na⁺ cations at relatively unfavourable short separations, causing these structures to be destabilised compared with what would be expected from Dempsey's rule. The importance of Na⁺ cations in determining the distribution of framework aluminium throughout a zeolite is also reflected in the variation in the position of the third aluminium for the two Na-SSZ-13 structures with 3 Al per unit cell. In the Löwensteinian structure, the NNN Al position is favoured, and the associated Na⁺ cations occupy two 8-rings and one 6-ring, with minimal repulsions between the counter-cations. However, the NNNN Al position is favoured for the non-Löwensteinian structure, in which the Na⁺ cations occupy only one 8-ring, and the two 6-rings of the D6R. In this structure, a single 6-ring and 8-ring occupancy are filled by virtue of the initial non-Löwensteinian arrangement of the aluminium ions. NNN substitution would result in Na⁺ occupancy of a six-ring that is already filled. Despite the NNNN position traditionally being

thought of as more unfavourable, in the NL case, it is the only aluminium position which can satisfy the Na⁺ coordination requirements whilst minimising unfavourable Na–Na interactions.

Rationalising the non-Dempsey aluminium distribution in sodium-containing frameworks is straightforward, whilst untangling the thermodynamic preference for NL ordering in protonated frameworks is more complex. As demonstrated by sodium-containing frameworks, non-covalent interactions play a significant role in determining aluminium distribution, we hence speculated that hydrogen-bonding interactions could be the cause of the unanticipated stability of the NL ordering in protonated zeolite frameworks. Fujita *et al.* demonstrated that hydrogen bonding interactions cause aluminium atoms to reside in close proximity to one another in zeolite Beta.³⁷ The separation between framework oxygen and H1 and H2 in 2 Al per unit cell H-SSZ-13 indicates the existence of two hydrogen bonds (O–H...O < 2.5 Å) per aluminium in both the global minimum structure and the lowest energy Löwensteinian structure. We hence examined the robustness of the order of stability predicted in this work by using other density functionals. A representative subset of structures were selected and re-optimised with the revPBE³⁸ and BLYP functionals,^{39,40} which have been shown to underbind hydrogen bonding interactions in water and ice structures (whilst PBE overbinds).⁴¹ The results (S1†) show that decreasing the hydrogen bonding strength in this way has no qualitative effect on the results and little quantitative effect, indicating that whilst hydrogen bonding must play a part in stabilising the H-SSZ-13 structures, it is not the decisive factor that controls whether NL is favoured over L.

Next we considered the charge distributions in the structures. On comparison of the Bader charges for 2 Al per unit cell Na and H-SSZ-13 we found the charge on the Na⁺ cation is +0.92, far greater than that of the proton in corresponding H-SSZ-13 NL structure, H⁺ = +0.66. Consequently, the charge on the framework oxygen atoms covalently bound to the protons is reduced in comparison to the corresponding oxygen atoms in the Na-SSZ-13 structure, O = –1.47 *versus* –1.60 (Mulliken = –0.71 and –1.08) in the H-SSZ-13 and Na-SSZ-13 structures, respectively. The reduced charge on the oxygen in the protonated case, results in the formation of long T–O(H) bonds (where T is Si or Al). According to the DFT data, the Al–O bond is more deformable than Si–O, in line with expectation as the absolute charge on Al is smaller than on Si and the radius of Al³⁺ is greater than Si⁴⁺. Hence, Si⁴⁺ forms shorter, stronger more ionic bonds in comparison to Al³⁺. For the H-SSZ-13 structure with 2 Al per U.C. Al–O(H) bonds are 11% longer compared to Al–O, stretched to a maximum of 1.90 Å, compared to an average Al–O bond length of approximately 1.71 Å. Si–O(H) bonds are only 7% longer than Si–O, stretched to a maximum bond length of 1.74 Å, compared to a framework average of approximately 1.63 Å. Each of the stable NL H-SSZ-13 structures contain a hydroxyl species mediating two aluminium ions, these structures therefore contain a total of three long Al–O(H) bonds, and a single long silanol Si–O(H) bond. In the high-energy NL structures and all L structures, there are two Al–O(H) bonds, and two comparatively unfavourable long Si–O(H) bonds. Adopting the



NL configuration minimises the number of long Si–O(H) bonds and maximises the number of the short, strong, more ionic Si–O bonds. In the sodium loaded zeolites, the interaction between Na⁺ and framework oxygen is primarily electrostatic and there is essentially complete charge transfer between Na⁺ and framework oxygen, as reflected by the computed Na⁺ charge and so the difference in ionicity/charge between a framework oxygen coordinated to Na⁺ and those not coordinated to Na⁺ is rather small. In H-SSZ-13, the electrons are smeared across the covalent O–H bond and the effective charge on the bridging oxygen is reduced and the alumina units favour adopting next-nearest neighbour structures. The clustering or islanding of aluminium has been noted in silicon–aluminium phosphate zeolites¹⁶ but not in aluminosilicate zeolites.

To check whether the qualitative result is sensitive to the extra-framework cation, we performed further calculations, substituting Na⁺ and H⁺ cations in the 2 Al per unit cell SSZ-13 model for intermediate sized Li⁺ cations and optimising all configurations to equilibrium density. The DFT results are included in the ESI† (S4 and S5), and are remarkably similar to that of Na-SSZ-13, showing a thermodynamic preference for ‘traditional’ Löwensteinian ordering over non-Löwensteinian with a ΔE ($N_{\text{Lglobal minimum}} - L_{\text{global minimum}}$) = +51.2 kJ mol⁻¹ per U.C. However, for Li-SSZ-13, the global minimum structure shows marked differences in alkali cation position, with Li⁺ ions capping the faces of individual D6R units, rather than located at the parameter of the 8-rings, as was the case for Na-SSZ-13 at this Si/Al ratio. Because Li⁺ cations are considerably smaller than Na⁺ cations, the Li⁺ ions are able to get closer to the D6R due to their higher charge density.

SSZ-13 is typically synthesised from a sodium solution with a nitrogenous structure-directing agent, yielding Na-SSZ-13, which is subsequently ion-exchanged post synthesis to give the protonated, Brønsted acid active form of the zeolite catalyst, H-SSZ-13.⁴² It is this form of the zeolite that is used to catalyse methanol-to-olefin conversion, a proposed lucrative, non-petroleum route for the production of short-chain organic compounds. At present, there is no viable way to synthesise H-SSZ-13 directly, most probably due to the role of counter-cations in directing the progress of zeolite formation during synthesis. As shown by our results, the location of framework aluminium is directly affected by counter-cation identity, and we can therefore assume that the distribution of aluminium in the global minimum Na-SSZ-13 framework is most representative of what would likely be seen in typical samples of SSZ-13, as it is this cation which determines the position of Al. However, our predictions suggest that a direct synthesis of H-SSZ-13 (H-CHA), H-LTA, H-RHO, H-ABW and H-MOR should favour NL aluminium ion ordering. Interestingly, high resolution mass spectrometry data concerning the incorporation of aluminium in prenucleating silicate species by Schaack *et al.*,⁴³ indicates that Löwenstein’s rule is not obeyed for all silicate species. The work provides evidence of 4-ring units containing –Al–O–Al–, but concludes that whilst these species may occur in solution, species that obey Löwenstein’s rule are preferentially formed. Nevertheless, the observation of pre-nucleating building units with –Al–O–Al– linkages hints that this motif may not be as

elusive as is generally believed, and these sequences may be found in crystals.

Because direct synthesis of proton compensated zeolites has not yet been achieved, direct validation of NL ordered frameworks in protonated zeolites cannot be assessed immediately. However, with regard to the synthesis of H-zeolite frameworks, we propose that the formation of –Al–O–Al– might be facilitated *via* two post-synthetic methods. The first is to use water, which has been shown to facilitate the making and breaking of –Si–O–Si– and –Al–O–Si–.⁴⁴ Long-term steeping of H-SSZ-13 in water could be expected to lead to the redistribution of Al in the framework, yielding –Al–O–Al– as the thermodynamically preferred arrangement. Potentially, very slightly acidic or basic water might enhance the rate of rearrangement without dealumination or desilication of the zeolite framework. A second potential approach is, in essence, reverse-dealumination; placing a zeolite crystal in a solution containing an excess of alumina units with the assumption that for high alumina zeolites, the aluminium content will rise, increasing the likelihood of alumina units situated adjacent to one another. Previously, this has been achieved in high-silica ZSM-5 *via* AlCl₃ vapour treatment, and in very low-silica zeolite Y using non-crystallisation inducing alkaline solutions (*e.g.* KOH) in the presence of large concentrations of extra-framework aluminium.^{45,46}

An intriguing question is whether the NL linkages that we have predicted are present in existing samples, and if so, what signatures could be used to unambiguously identify these –Al–O–Al– sequences. Recent atom tomography work^{4,26} has vividly demonstrated that the distribution of aluminium in a typical ZSM-5 zeolite sample is very heterogeneous. At present there is no available method that can accurately distinguish framework aluminium from framework silicon with Ångström resolution.⁴ A 2010 work by Shin *et al.*⁸ concerning possible non-Löwensteinian structures observed in gallosilicates, discusses the possibility of using of ¹⁷O magic angle spinning (MAS) NMR to detect non-Löwensteinian ordering, a method which has been successful in identifying –Al–O–Al– linkages in aluminosilicate glasses.⁴⁷ We have examined the global minimum H-SSZ-13 structures (Si/Al = 17) and predicted ²⁹Si, ²⁷Al solid-state, MAS NMR shifts and vibrational frequencies, in an attempt to discern whether spectroscopic signatures exist that would be indicative of the presence of non-Löwensteinian ordering (see ESI†). However, at a Si/Al ratio of 17, typical for SSZ-13, the predicted ²⁹Si NMR data shows that there is a slight decrease in the negativity of the chemical shift values for –Al–O–Al– containing frameworks. However, these shifts are well within the anticipated range for a zeolite at this Si/Al ratio, and far too similar to the chemical shifts of the surrounding Si atoms to be used practically as a characterisation method. Similarly, predicted vibrational frequencies indicate that characteristic stretches would not be detectable due to overlap of Al–O(H)–Al stretches with that of Si–O(H)–Si and Si–O(H)–Al, and –Al–O–Al– stretches with Si–O–Al. This data is included and discussed in the ESI† (S10 and S11).

If the proposed post-synthetic techniques or alternative synthetic strategies are successful in realising zeolites with NL framework aluminium distributions, as predicted by this work,



these materials would be potentially invaluable for the development of new zeolite catalysts. Despite the advantages of using zeolites in catalysis, for example, specificity and size exclusion properties, it is well documented that the catalytic efficiency of microporous materials is often limited by restricted access to active sites. Introducing ordered, controllable meso- and macroporosity to the framework provides a solution to mass transport limited diffusion through the porous zeolite network. The introduction of hierarchy has also been shown not only enhance catalytic activity, but also stability in a range of zeolite frameworks. A variety of both bottom-up and top-down strategies have proved successful for hierarchically ordered zeolite synthesis. The post-synthetic introduction of mesoporosity by the extraction of framework atoms is a particularly popular method, and can be achieved by acid, base or steam treatment of the zeolite material.^{48,49} One can imagine how techniques such as these could be used to dealuminate low-silica aluminium cluster-containing materials, similar to those predicted in this work. For example, removing all four alumina units in the 4 Al per unit cell H-SSZ-13 global minimum structure predicted by DFT would increase the 7 Å, 8-ring aperture cavity system, with a void-space of approximately 10 Å in diameter, to up to 17 Å, approaching mesoporosity. Crucially, the calculations indicate that not only is aluminium clustered, but it is also located in predictable, ordered positions, which suggests that introduced porosity *via* selective dealumination could be controllable in H-zeolites.

The reaction mechanisms and deactivation pathways of real catalytic zeolite materials is relatively a poorly understood area of zeolite science, not withstanding remarkable recent advances.^{50,51} In part this is due to a lack of molecular-level information concerning the location of framework alumina and associated counter-cations, which are thought to be integral to the catalytic reaction mechanism. Clustering of aluminium and the associated clustered acid sites, as predicted by the DFT results, is suggestive of new, potentially more reactive sites (due to the density of acid sites) and new reaction pathways which have not yet been considered. Perea *et al.* have shown that the Si/Al distribution can be inhomogeneously distributed throughout the zeolite framework,⁴ furthermore, silicon islanding (formation of silicon rich regions that must also give rise to aluminium rich areas) has been shown to be present in SSZ-13's silicoaluminophosphate counterpart SAPO-34.⁵² Hence, it is conceivable that aluminium cluster motifs, including non-Löwensteinian linkages already exist in real zeolite materials, and may impact reaction and deactivation pathways operating in current catalysts.

The realisation of zeolite materials with contradistinct aluminium distributions to those synthesised by traditional routes holds enormous potential for the future of zeolite catalysis. We hope this work stimulates experimental investigation into the direct or post-synthesis of non-Löwensteinian ordered zeolites and further characterisation of existing materials.

Methods

The majority of the periodic DFT calculations were performed using the CP2K code,^{29–31} and additional benchmark

calculations for energetics and solid-state NMR were performed using the CASTEP code.⁵³ Results were calculated using the PBE³² functional, although further calculations using revPBE³⁸ and BLYP^{39,40} were included to verify our initial 2 Al per unit cell SSZ-13 findings. These calculations, and methods, are discussed in detail in the ESI,[†] along with a full description of the single-point energy PBE0 (ref. 34 and 35) calculations mentioned in our Results and Discussion. All framework structures were obtained in their all-silica form from the database of zeolite structures,²⁸ and permutations of 2, 3 and 4 Al per unit cell models created by the methodology discussed in the main body of the text. Individual models were fully geometry optimized to equilibrium density, with variable lattice parameters in CP2K as 1 : 1 : 1 cells using the high quality TZV2P basis set and an energy cutoff of 650 Ry. Only the ABW framework was optimized as a 2 : 2 : 2 supercell, due to its small unit cell size. We also tested a selection of larger 2 : 2 : 2 supercells for each of the frameworks, although we saw no meaningful change in the relative energies using the larger cells. Additional computational details are included in the ESI.[†]

Conflicts of interest

There are no conflicts to declare.

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References

- 1 W. Vermeiren and J.-P. Gilson, *Top. Catal.*, 2009, **52**, 1131–1161.
- 2 M. Guisnet and J.-P. Gilson, *Zeolites for Cleaner Technologies*, Imperial College Press, 2002.
- 3 P. T. Anastas and J. C. Warner, *Green Chemistry: Theory and Practice*, Oxford University Press, 1998.
- 4 D. E. Perea, *et al.*, *Nat. Commun.*, 2015, **6**, 7589.
- 5 W. Loewenstein, *Am. Mineral.*, 1954, **39**, 92–96.
- 6 L. A. Bursill, E. A. Lodge, J. M. Thomas and A. K. Cheetham, *J. Phys. Chem.*, 1981, **85**, 2409–2421.
- 7 J. Klinowski, J. M. Thomas, C. A. Fyfe and J. S. Hartman, *J. Phys. Chem.*, 1981, **85**, 2590–2594.
- 8 J. Shin, *et al.*, *Dalton Trans.*, 2010, **39**, 2246–2253.
- 9 S. E. Tarling, P. Barnes and J. Klinowski, *Acta Crystallogr.*, 1980, **37**, 652–636.
- 10 A. Putnis, C. A. Fyfe and G. C. Gobbi, *Phys. Chem. Miner.*, 1985, **12**, 211–216.
- 11 C. A. Fyfe, G. C. Gobbi and A. Putnis, *J. Am. Chem. Soc.*, 1986, **108**, 3218–3223.
- 12 P. Florian, E. Veron, T. F. G. Green, J. R. Yates and D. Massiot, *Chem. Mater.*, 2012, **24**, 4068–4079.



- 13 S. E. Dann, P. J. Mead and M. T. Weller, *Angew. Chem., Int. Ed. Engl.*, 1995, **34**, 2414–2416.
- 14 R. G. Bell, R. A. Jackson and C. R. A. Catlow, *Zeolites*, 1992, **70**, 870–871.
- 15 C. R. A. Catlow, A. R. George and C. M. Freeman, *Chem. Commun.*, 1996, 1311–1312.
- 16 D. E. Akporiaye, I. M. Dahl, H. B. Mostad and R. Wendelbo, *J. Phys. Chem.*, 1996, **100**, 4148–4153.
- 17 M. Melchior, D. Vaughan and A. Jacobson, *J. Am. Chem. Soc.*, 1982, **104**, 4859–4864.
- 18 C. M. Soukoulis, *J. Phys. Chem.*, 1984, **88**, 4898–4901.
- 19 K. P. Schroeder and J. Sauer, *J. Phys. Chem.*, 1993, **97**, 6579–6581.
- 20 C.-S. Yang, J. M. Mora-Fonz and C. R. A. Catlow, *J. Phys. Chem. C*, 2011, **115**, 24102–24114.
- 21 C. Lo and B. Trout, *J. Catal.*, 2004, **227**, 77–89.
- 22 K. Muraoka, W. Chaikittisilp and T. Okubo, *J. Am. Chem. Soc.*, 2016, **138**, 6184–6193.
- 23 M. Zokaie, U. Olsbye, K. P. Lillerud and O. Swang, *Microporous Mesoporous Mater.*, 2012, **158**, 175–179.
- 24 R. C. Rouse and D. R. Peacor, *Am. Mineral.*, 1986, **71**, 1494–1501.
- 25 V. Gábová, *et al.*, *Chem. Commun.*, 2003, **11**, 1196–1197.
- 26 J. E. Schmidt, *et al.*, *Angew. Chem., Int. Ed.*, 2016, 11173–11177.
- 27 R. von Ballmoos and W. M. Meier, *Nature*, 1981, **289**, 782–783.
- 28 C. Baerlocher, L. McCusker and D. H. Olson, *Atlas of zeolite framework types*, Elsevier, 6th edn, 2007.
- 29 Available at: <https://www.cp2k.org/>.
- 30 J. VandeVondele, *et al.*, *Comput. Phys. Commun.*, 2005, **167**, 103–128.
- 31 J. VandeVondele and J. Hutter, *J. Chem. Phys.*, 2007, **127**, 114105.
- 32 J. P. Perdew, K. Burke and M. Ernzerhof, *Phys. Rev. Lett.*, 1997, **78**, 1396.
- 33 E. Dempsey, G. H. Kuehl and D. H. Olson, *J. Phys. Chem.*, 1969, **73**, 387–390.
- 34 C. Adamo and V. Barone, *J. Chem. Phys.*, 1999, **110**, 6158.
- 35 M. Guidon, J. Hutter and J. VandeVondele, *J. Chem. Theory Comput.*, 2010, **6**, 2348–2364.
- 36 E. Dempsey, *J. Catal.*, 1977, **49**, 115–119.
- 37 H. Fujita, T. Kanougi and T. Atoguchi, *Appl. Catal., A*, 2006, **313**, 160–166.
- 38 Y. Zhang and W. Yang, *Phys. Rev. Lett.*, 1998, **80**, 890.
- 39 A. D. Becke, *Phys. Rev. A*, 1988, **38**, 3098–3100.
- 40 C. Lee, W. Yang and R. G. Parr, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1988, **37**, 785–789.
- 41 M. J. Gillan, D. Alfè and A. Michaelides, *J. Chem. Phys.*, 2016, **144**, 130901.
- 42 S. I. Zones, *US Pat.* 4544538, 1985.
- 43 B. B. Schaack, The Solid State Formation of Zeolite Materials Studies by Mass Spectrometry, PhD thesis, Ruhr Universität Bochum, 2009.
- 44 R. Von Ballmoos and W. M. Meier, *J. Phys. Chem.*, 1982, **86**, 2698–2700.
- 45 X. Liu, J. Klinowski and J. M. Thomas, *J. Chem. Soc., Chem. Commun.*, 1986, 582–584.
- 46 R. D. Bezman, *J. Chem. Soc., Chem. Commun.*, 1987, **606**, 1562–1563.
- 47 J. F. Stebbins, S. K. Lee and J. V. Oglesby, *Am. Mineral.*, 1999, **84**, 983–986.
- 48 D. Verboekend and J. Pérez-Ramírez, *Catal. Sci. Technol.*, 2011, **1**, 879–890.
- 49 K. Möller and T. Bein, *Science*, 2011, **333**, 297–298.
- 50 E. de Smit, *et al.*, *Nature*, 2008, **456**, 222–225.
- 51 I. L. C. Buurmans and B. M. Weckhuysen, *Nat. Chem.*, 2012, **4**, 873–886.
- 52 G. Sastre, D. W. Lewis and C. R. A. Catlow, *J. Phys. Chem. B*, 1997, **101**, 5249–5262.
- 53 S. J. Clark, *et al.*, *Z. Kristallogr.*, 2005, **220**, 567–570.

