J.C.S. Chem. Comm., 1981

## A Reassessment of Zeolite A: Evidence that the Structure is Rhombohedral with Unexpected Ordering in the Aluminosilicate Framework

By John M. Thomas,\* Leslie A. Bursill, and Elizabeth A. Lodge (Department of Physical Chemistry, University of Cambridge, Lensfield Road, Cambridge CB2 1EP)

## ANTHONY K. CHEETHAM

(Chemical Crystallography Laboratory, University of Oxford, 9 Parks Road, Oxford OX1 3PD)

and Colin A. Fyfe

(Guelph-Waterloo Centre for Graduate Work in Chemistry, University of Guelph, Ontario, Canada N1G 2W1)

Summary A new structural model for zeolite A (rhombohedral  $R\overline{3}$ ) differs from the one currently accepted (cubic Fm3c) in that each tetrahedrally co-ordinated Si<sup>4+</sup> ion is surrounded, via oxygen bridges, by three (not four) Al<sup>3+</sup> ions and one Si<sup>4+</sup> ion, and each Al<sup>3+</sup> ion by three (not four) Si<sup>4+</sup> and one Al<sup>3+</sup>.

RECENTLY<sup>1</sup> we raised the question as to whether the structure of zeolite A (idealized formula M+12Al12Si12-O48.27H2O) had been correctly identified. This discussion arose initially because our electron microscopic and, in particular, our electron diffraction studies of dehydrated Na-A yielded results which implied that the currently accepted picture [see Figure (a)], in which each Si4+ ion is surrounded, via oxygen bridges, by four Al3+ ions and each Al<sup>3+</sup> likewise by four Si<sup>4+</sup> ions, space group Fm3c, is wrong. Further, independent evidence that the accepted structure<sup>2,3</sup> merits re-investigation came from the high-resolution n.m.r. studies of Lippmaa et al.,4 who reported that each Si<sup>4+</sup> is surrounded, via oxygens, by three Al<sup>3+</sup> and one Si<sup>4+</sup>, and that, conversely, each central Al3+ also displays 3:1 co-ordination (i.e. there are three Si3+ and one Al3+ ions in the first tetrahedral shell).

On the tacit assumption that the zeolite A structure has a cubic unit cell, we tentatively suggested that, when the Si/Al ratio is unity, the space-group is Pm3 and that, in general (Si/Al  $\neq$  1), the space group is Fm3.

We have now completed neutron diffraction studies (Rietveld powder analysis  $^{5,6}$ ) at 5 and 298 K, as well as high-resolution (magic-angle spinning), solid-state  $^{28}$ Si n.m.r. measurements  $^{7,8}$  and further electron microscopic analysis on a specially prepared sample of dehydrated Na-A possessing a Si/Al ratio of  $1\cdot00\pm0\cdot02$ . Our results indicate the following.

- (a) There is a doubled unit cell, *i.e.* two joined cuboctahedra ( $\beta$ -cages) constitute the repeat unit (from electron diffraction). Due allowance has been made for the complications that arise from double or multiple diffractions, and we conclude that the structure has ca. 24.6 rather than 12.3 Å repeats.
- (b) The co-ordination is unmistakably 3:1 (from solid-state n.m.r. studies).
- (c) The space group is  $R\overline{3}$  (from neutron diffraction). The diffraction peaks cannot be indexed with, and the structure does not refine in, Fm3c. Moreover, the <sup>29</sup>Si n.m.r. resonance at ca. -84 p.p.m. characteristic of 4:0 co-ordination, in sodalite for example, is absent, the peak instead appearing at ca. -89 p.p.m., which signifies 3:1 co-ordination.

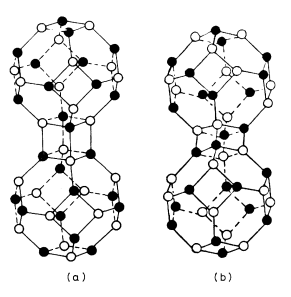


FIGURE. (a) In the currently accepted model for the structure of zeolite-A, two cuboctahedra (i.e.  $\beta$ -cages) are joined at the so-called 'double-four' rings. At the vertices of the cubocta-hedra  $Si^{4+}$  ( ) and  $Al^{3+}$  ions ( ) alternate such that 4:0 co-ordination results, *i.e.* each  $Si^{4+}$  is surrounded tetrahedrally by four Al3+ ions, via oxygen bridges, which, for simplicity are not This double unit of cuboctahedra repeats itself threedimensionally resulting in a cubic (Fm3c) space group. changeable cations not shown). (b) In the new model the cuboctahedra ( $\beta$ -cages) are slightly distorted (not shown) and again repeat themselves three-dimensionally, yielding a rhombohedral space-group  $R\overline{\bf 3}$ . The ordering scheme within the aluminosilicate framework is such that each  ${\rm Si}^{4+}$  ion is surrounded by three Al<sup>3+</sup> and one Si<sup>4+</sup>, and each Al<sup>3+</sup> by three Si<sup>4+</sup> and one Al<sup>3+</sup> (3:1 co-ordination). Al<sup>3+</sup>-O-Al<sup>3+</sup> bridges, hitherto discounted (see text), are an integral feature of the structure. In the new model there are centres of inversion at the midpoints of the double-four rings. Note that two kinds of sixrings, differing in Si, Al sequence, occur in the new model, and there are also two types of four-ring. (Exchangeable cations not shown).

The hitherto unsuspected rhombohedral distortion (or pseudo-cubic nature) of the structure was uncovered using neutrons of wavelength 2.96 Å. The splitting of the peaks, e.g. 8,8,0, at first suggested that the sample was biphasic, but the 12,0,0 peak is not split, signifying that the sample is monophasic. We have analysed crystallographically all the possible combinations of the two cuboctahedra linked by a cube that satisfy the known facts pertaining to the structure, taking cognizance of the n.m.r. results and the conclusion from our neutron scattering measurements that

the cuboctahedra are slightly distorted such that they no longer possess four, but only one 3-fold axis. The unique result is shown schematized, in part, in Figure (b), from which we note that the 'doubled' entity (pseudo-cubic unit cell repeat 24.6 Å) consists of an enantiomeric pair of (slightly distorted) cuboctahedra, i.e.  $C_3^1$ .  $C_3^2$  in the Schoenflies notation.

We have also discovered a possible new cubic model for dehydrated Na-A when the Si/Al ratio is not unity. It is made up of a combination of distorted  $(C_{3i})$  cuboctahedra, arranged such that their unique 3-fold axes are aligned along the four <111> directions. This cubic structure (space group Pn3n) possesses pseudo-cubic (i.e. face-centred cubic Fm3c) and pseudo Pm3 symmetry.

When Si/Al = 1.00 the unit cell dimensions in R3 are  $a = 17.401 \pm 0.001 \,\text{Å}$ ,  $\alpha = 59.53^{\circ} \, (\pm 0.01^{\circ})$ . When Si/Al = 1.10,  $a = 17.352 \pm 0.001$  Å,  $\alpha = 59.84 \pm 0.01$ °. Both sets of dimensions refer to 5 K. A full description of the new structural analysis, and the implications of the new model are given elsewhere.9 It is of interest that the

structure contains hitherto discounted Al-O-Al linkages and that, in the new model, two types of hexagonal rings occur, one symmetrical the other unsymmetrical. Both these features, as well as others, are likely to influence the chemical properties of zeolite-A.

We note also that this relatively unconventional combination of techniques: high-resolution, magic-anglespinning n.m.r. spectroscopy, neutron powder diffraction (profile analysis), and electron microscopy, constitutes a powerful and potentially widely applicable method of structural analysis.

We acknowledge support from the S.R.C., the Universities of Cambridge and Melbourne, and Unilever Ltd. (for a maintenance grant to E. A. L.). We are also grateful to Dr. A. C. Bishop and colleagues at the British Museum for the Si, Al analyses. L. A. B. is on leave from the School of Physics, University of Melbourne.

(Received, 15th January 1981; Com. 040.)

<sup>1</sup> E. A. Lodge, L. A. Bursill, and J. M. Thomas, J. Chem. Soc., Chem. Commun., 1980, 875; see also L. A. Bursill, E. A. Lodge, and J. M. Thomas, Nature, 1980, 286, 111

J. Pluth and J. V. Smith, J. Phys. Chem., 1979, 83, 741.
J. J. Pluth and J. V. Smith, J. Am. Chem. Soc., 1980, 102, 4704.
G. Engelhardt, D. Kunath, A. Samoson, M. Tarmak, and M. Mägi, Workshop in 'Adsorption of Hydrocarbons in Zeolites,' Berlin Aldershof, 19-20 Nov., 1979; E. Lippmaa, M. Mägi, A. Samoson, G. Engelhardt, and A. R. Grimmer, J. Am. Chem. Soc., 1980, 102, 4889.

M. Rietveld, J. Appl. Crystallogr., 1969, 2, 65.
A. K. Cheetham and J. C. Taylor, J. Solid State Chem., 1977, 21, 253; A. W. Hewat in 'Accuracy in Powder Diffraction' (eds. S. Bloch and C. R. Hubbard), National Bureau of Standards, Spec. Publ. 567, U.S. Dept. of Commerce, Washington, 1980.
J. A. Lyerla, C. A. Fyfe, and C. S. Yannoni, J. Am. Chem. Soc., 1979, 101, 1351.
C. A. Fyfe, J. Klinowski, J. M. Thomas, L. A. Bursill, et al., in preparation.
L. A. Bursill, E. A. Lodge, J. M. Thomas, and A. K. Cheetham, J. Phys. Chem., submitted for publication.