

PMW structure (Baldinozzi, Sciau, Buffat & Stadelmann, 1994).

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Structure of Na₃YSi₆O₁₅ – a Unique Silicate Based on Discrete Si₆O₁₅ Units, and a Possible Fast-Ion Conductor

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Abstract

Hydrothermal investigations in the high silica region of the Na₂O–Y₂O₃–Si₆O₁₅ system, carried out in a search for novel fast-ion conductors (FIC's), yielded the new compound trisodium yttrium hexasilicate, Na₃YSi₆O₁₅. Single-crystal X-ray methods revealed that Na₃YSi₆O₁₅ crystallizes in space group *Ibmm*, has lattice constants $a = 10.468$ (2), $b = 15.2467$ (13) and $c = 8.3855$ (6) Å, $Z = 4$, and 11 atoms in the asymmetric unit. Refinement was carried out to a weighted residual of 3.53% using anisotropic temperature factors for all atoms. The structure is unique in that the silica tetrahedra form isolated Si₆O₁₅⁶⁻ double dreier-rings, rather than layers as might be expected from the Si to O ratio of 0.4. No isomorphs to Na₃YSi₆O₁₅ have been reported.

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Introduction

High ionic conductivity has been observed in a number of alkali silicates. These include Na₅YSi₄O₁₂ (Shannon, Taylor, Gier, Chen & Berzins, 1978), Na₂ZnSiO₄ (Frostäng, Grins & Nygren, 1988), and the solid solution series Na_{1+x}Zr₂Si_xP_{3-x}O₁₂ [NASICON (Goodenough, Hong & Kafalas, 1976)]. In such materials, the alkali ion typically migrates *via* channels present in the silicate framework. The transport properties of alkali silicates are amenable to crystal-chemical tailoring because both the tunnel geometry and the size of the mobile ion can be altered *via* substitutions on the non-Si cation sites. Additionally, as with other ionic materials, the concentration of ionic defects can be modified *via* aliovalent substitutions, with the prospect of increasing the ionic conductivity. We were especially interested in Na₃YSi₆O₁₅ for a number of reasons. Firstly, we previously measured the conductivity of K₃NdSi₆O₁₅ (Haile, Wuensch, Siegrist & Laudise, 1992) and found

it to have a reasonably high conductivity for a material in which K⁺ is the mobile species. Secondly, Watanabe & Wuensch (1993) reported a rather high conductivity for Na₃YSi₆O₁₅ glass. These two observations suggested that the crystalline sodium analog to K₃NdSi₆O₁₅, having a smaller and typically more mobile species as the charge carrier, might exhibit fast-ion conduction. In addition, the anisotropy we observed in the properties of K₃NdSi₆O₁₅ suggested this class of compounds to be ideal for correlating structural features to observed conductivities.

Experimental

Crystal growth

Crystals of Na₃YSi₆O₁₅ were grown by the hydrothermal method, as described elsewhere (Haile, Wuensch & Laudise, 1993). This technique, which employs high temperatures and pressures to dissolve a normally insoluble precursor and precipitate the desired crystalline phase, has been utilized because of its effectiveness in inducing crystallization in glass-forming systems (Laudise, 1987). Typically, 0.05 g of a glass nutrient of composition 4Na₂O–Y₂O₃–17SiO₂ was placed in sealed cylindrical platinum capsules, 0.40 cm in diameter and 2.3–2.7 cm in length, with a total volume of 0.200–0.250 cm³ (after accounting for the loss in volume to the sealing process). These were, in turn, placed in Tuttle autoclaves (Tuttle, 1949), the design of which allowed the pressure to be controlled explicitly by the use of an intensifier, gauge and valve system. The pressure was held at either 825 or 1400 × 10⁵ Pa, the temperature at 773, 823 or 873 K, and the solvent was either pure water or a dilute solution of a sodium base (0.1 M NaOH, Na₂CO₃ or NaHF₂). After 7–12 d experiments yielded a crystalline powder of Na₃YSi₆O₁₅ with a crystallite size of several microns.

Composition determination

The composition was determined both from electron microprobe measurements and by inductively coupled plasma (ICP) spectroscopy. Microprobe measurements were performed on small crystals that had been mounted in an epoxy resin, polished and then sputter-coated using a gold/palladium electrode. Data were collected on a Jeol Superprobe 733 equipped with a wavelength dispersive detector. The intensities of the characteristic X-radiation peaks were converted to stoichiometric quantities using the ZAF data reduction program (Schamber, Wodke & McCarthy, 1981). The following materials were employed as standards: SiO₂ quartz glass, a single crystal of yttrium aluminum garnet, and sodalite or amelia albite glass for Na. These measurements yielded mole percentages of Na, Y and Si of 12.1(3), 4.0(3) and 24.6(8), respectively, values which are within experi-

mental error of those for ideal Na₃YSi₆O₁₅: 12, 4 and 24, respectively. Upon conversion of the experimental values to the weight percentages of the respective oxides, the sum of 101(2) wt% was found, indicating that no water of crystallization was present.

ICP spectroscopy was performed after dissolution in a mixture of concentrated HCl and concentrated HF at 423 K. Measurements made on a sample of approximately 30 mg yielded Na, Y, Si and O weight percentages of 11.9, 13.9, 23.7 and 41.3 wt%, which, when normalized to 13.1, 15.3, 26.1 and 45.5 wt%, are comparable to the ideal values of 12.2, 15.7, 29.7 and 42.4 wt%, respectively. We attribute the deviation from 100 wt% to the small sample size.

Structure determination

A crystal measuring 0.02 × 0.06 × 0.1 mm³ was selected from the product of an experiment conducted at a temperature of 823 K, a pressure of 825 × 10⁵ Pa, in a 0.1 M NaOH solution for 12 d. Single-crystal intensity data were collected using a Siemens diffractometer and MoK α radiation ($\lambda = 0.71073$ Å). Refinement of the lattice parameters using data for the positions of 82 peaks revealed Na₃YSi₆O₁₅ to be orthorhombic with lattice constants $a = 10.468$ (2), $b = 15.2467$ (13) and $c = 8.3855$ (6) Å. With four formula units per unit cell, Na₃YSi₆O₁₅ has a calculated density of 2.811 g cm⁻³. Intensity data were collected using the ω - 2θ scan method, over 2θ values from 3 to 80°. The measured intensities were corrected for absorption; transmission factors ranged from 0.0334 to 0.0841. Of 2257 independent intensities, 1418 were considered significant (net intensity greater than three standard deviations). An analysis of the systematic absences showed the diffraction symbol to be mmm $I(bc)^{**}$, allowing space groups $Ibmm$ and $Ic2m$ (the latter in two possible orientations). The value of $E^2 - 1$, 0.863, gave slight preference to the centrosymmetric group, and thus $Ibmm$ was employed for the initial calculations and later proven to be correct, as discussed below. The structure was solved by first locating the Y atoms from a Patterson map, and the remaining atoms from successive Fourier-difference maps. All calculations were performed using the *SHELXS* package of programs (Sheldrick, 1985). The final weighted residuals, after employing anisotropic thermal parameters for all atoms, were 3.57 and 4.11%, for the limited and complete data sets, respectively, for a weighting scheme defined by $w = 3.2117\sigma(F)^{-2}$. Nonweighted residuals were 8.24 and 12.13%. A list of F_{obs} and F_{calc} is provided. After the last cycle of refinement, the deepest hole in the Fourier-difference map was 2.407 e Å⁻³, and the highest peak 3.910 e Å⁻³. No correlation terms in the least-squares matrix were greater than 0.5.

The atomic coordinates and thermal parameters for the 11 atoms in the asymmetric unit are reported in Table 1. The second Na atom, Na(2), has an occupancy

Table 1. Atomic coordinates and thermal parameters of Na₃YSi₆O₁₅ (e.s.d.'s given in parentheses)

The anisotropic temperature factors are of the form $\exp[-2\pi^2(h^2u_{22}^2 + \dots + hku_{12}ab^* + \dots)]$ and are given in units of 10^{-2} \AA^2 . The compound crystallizes in space group *Ibmm*, and has lattice constants $a = 10.468$ (2), $b = 15.2467$ (13) and $c = 8.3855$ (6) \AA.

	Position	<i>x</i>	<i>y</i>	<i>z</i>	<i>U</i> ₁₁	<i>U</i> ₂₂	<i>U</i> ₃₃	<i>U</i> ₁₂	<i>U</i> ₁₃	<i>U</i> ₂₃
Y	4c <i>.2/m</i>	1/4	1/4	1/4	1.147 (18)	1.068 (17)	1.068 (19)	1.126 (19)	0	0
Si(1)	8i <i>.m</i>	-0.03353 (8)	0.40244 (6)	1/4	1.18 (4)	1.19 (4)	1.34 (4)	0.10 (3)	0	0
Si(2)	16j 1	-0.28438 (6)	0.40016 (5)	0.42866 (8)	1.25 (3)	1.29 (3)	1.22 (3)	0.02 (2)	0.04 (2)	0.15 (2)
Na(1)	8f <i>.2</i>	1/2	0.24005 (9)	0	2.15 (7)	1.85 (9)	2.16 (8)	0	0.62 (6)	0
Na(2)	8i <i>.m</i>	0.4227 (4)	0.4606 (3)	1/4	3.57 (20)	5.20 (25)	2.46 (19)	0.17 (18)	0	0
O(1)	8i <i>.m</i>	0.3481 (2)	0.38768 (18)	3/4	1.30 (11)	2.28 (14)	1.46 (12)	-0.02 (10)	0	0
O(2)	8i <i>.m</i>	0.4338 (2)	0.17522 (16)	1/4	1.37 (11)	1.41 (11)	1.62 (12)	0.23 (9)	0	0
O(3)	16j 1	0.62972 (15)	0.10678 (12)	0.0952 (2)	1.27 (7)	1.96 (8)	1.45 (8)	0.10 (7)	0.09 (5)	0.08 (7)
O(4)	8h <i>.m</i>	0.3210 (2)	1/2	-0.0135 (3)	1.82 (13)	1.46 (12)	1.98 (13)	0	-0.40 (10)	0
O(5)	16j 1	0.16320 (17)	0.16573 (13)	0.0573 (2)	1.70 (8)	2.15 (9)	1.76 (9)	0.28 (7)	-0.15 (7)	-0.77 (7)
O(6)	4e <i>2mm</i>	0.4715 (3)	0	1/4	1.54 (16)	1.12 (15)	2.13 (19)	0	0	0

of 0.5, and four Na atoms per unit cell are distributed over the eight equivalent sites of *8i*. This leads to a rather small Na–Na separation of 1.201 \AA. The proximity of these sites to the special position, *4e* ($x, \frac{1}{2}, \frac{1}{4}$) with $x = 0.4227$ (4), suggested that Na(2) might fully occupy a single site on one or the other side of the high symmetry position, thereby lowering the space group symmetry to *Ic2m*. A Fourier difference map in the space group of lower symmetry, however, showed two equally significant peaks at the positions related by pseudosymmetry. Attempts to refine the structure in *Ic2m* reduced the residual only nominally, while causing some bond lengths to assume unreasonable values. Additionally, a number of elements in the least-squares matrix became strongly correlated as a result of the pseudosymmetry. We are thus confident *Ibmm* is the true space group.

Bond distances in the cation polyhedra are given in Table 2. Bond lengths to oxygen, oxygen–oxygen separations, and oxygen–cation–oxygen bond angles are tabulated for the Y and Si tetrahedra. O atoms bonded to two Si atoms are designated as ‘bridging’ (br), whereas ‘terminating’ (term) indicates those bonded to only one. Silicon–oxygen–silicon angles for the bridging O atoms are also provided in Table 2. Only oxygen–cation distances are given for the Na polyhedra. In addition to the bond distances from the two Na sites that have been refined from X-ray data, the distances from what is considered to be the center of a ten-membered cage in which two Na(2) sites are located are also included. The nature of this cage is discussed in greater detail below.*

Discussion of the structure

Nature of the silicate unit

As can be seen from Table 2, three of the four oxygen atoms on both silica tetrahedra are bridging, while the

* A list of structure factors has been deposited with the IUCr (Reference: DU0394). Copies may be obtained through The Managing Editor, International Union of Crystallography, 5 Abbey Square, Chester CH1 2HU, England.

Table 2. Cation coordination polyhedra in Na₃YSi₆O₁₅ with bond distances and angles (e.s.d.'s in parentheses)

Number in parentheses after atom identifies the atom in terms of Table 2. 2×, 4× etc. indicates multiplicity of the bond. Oxygen–oxygen distances and angles are given for the yttrium octahedron and the silicon tetrahedra. Si–O–Si angles are also provided.

Central atom	Bond distance (\AA)	Oxygen separation (\AA)	O—M—O angle (°)
Y	O(2) 2× 2.237 (2)	O(2)—O(5) 4× 3.265 (3)	93.22 (6)
	O(5) 4× 2.256 (2)	O(2)—O(5) 4× 3.086 (3)	86.78 (6)
		O(2)—O(5) 4× 3.147 (2)	88.5 (1)
		O(5)—O(5) 2× 3.232 (4)	91.5 (1)
		O(5)—O(5) 4× 4.511 (4)	180
		O(2)—O(2) 4.473 (5)	180
Si(1)	O(2) _{term} 1.578 (3)	O(2)—O(6) 2.700 (2)	115.0 (2)
	O(6) _{br} 1.623 (2)	O(2)—O(3) 2× 2.642 (3)	109.87 (9)
	O(3) _{br} 2× 1.648 (2)	O(6)—O(3) 2× 2.661 (3)	108.8 (1)
		O(3)—O(3) 2.595 (3)	103.8 (1)
Si(2)	O(5) _{term} 1.573 (2)	O(5)—O(3) 2.672 (2)	112.8 (1)
	O(3) _{br} 1.635 (2)	O(5)—O(4) 2.601 (2)	108.0 (1)
	O(4) _{br} 1.643 (1)	O(5)—O(1) 2.705 (2)	114.1 (1)
	O(1) _{br} 1.651 (1)	O(3)—O(4) 2.670 (3)	109.1 (1)
		O(3)—O(1) 2.630 (3)	106.3 (1)
		O(4)—O(1) 2.635 (3)	106.3 (1)
Bridging oxygen bond-angles, Si—O—Si (°)			
	Si(2)—O(1)—Si(2) 130.3 (2)	Si(2)—O(4)—Si(2) 135.8 (2)	
	Si(2)—O(3)—Si(1) 134.1 (1)	Si(2)—O(6)—Si(2) 132.8 (2)	
Na(1) distances (\AA)		Na(2) distances (\AA)	
O(5) 2× 2.283 (2)	Na(2') 1.201 (8)		
O(2) 2× 2.419 (1)	O(4) 2× 2.525 (3)	O(4') 2× 3.390 (4)	
O(3) 2× 2.571 (2)	O(1) 2.644 (5)	O(1') 3.333 (5)	
	O(5) 2× 2.670 (4)	O(5) 2× 3.634 (4)	
From center of ten-membered cage at <i>4e</i> <i>2mm</i> (0.4596, $\frac{1}{2}, \frac{1}{4}$)			
	O(4) 2× 2.643	O(4') 2× 3.035	
	O(1) 2× 2.643	O(5) 4× 3.263	

remaining anion is terminating. This bonding scheme is the simplest way in which the silicon-to-oxygen ratio of 0.4 in Na₃YSi₆O₁₅ can be accommodated. Table 2 also reveals that the average Si—O_{term} distance, 1.576 (3) \AA, is smaller than the average Si—O_{br} distance, 1.641 (11) \AA, by 0.065 \AA. This difference, which is significantly larger than the standard deviation, results from the ‘lop-sided’ potential experienced by the terminal oxygen and agrees well with values encountered in other silicates (Liebau, 1985).

The silica tetrahedra in Na₃YSi₆O₁₅ are linked together to form double dreier-rings having point group

2mm. This discrete species is best described as a short tubular unit, with a circumference consisting of three silicate tetrahedra and a length of two tetrahedra. Alternatively, it can be considered as the unit resulting from the condensation of a pair of three-membered rings such that previously terminal O atoms are shared at a common interfacial plane. One of these isolated Si₆O₁₅⁶⁻ units is depicted in Fig. 1. The vast majority of silicates with a silicon-to-oxygen ratio of 0.4 form layered structures, while others take on various chain and single-ring configurations. Reports of double-ring compounds are few. Only one other phase that contains the double dreier-ring is known (Smolin, 1970). Indeed, the chemical analogs to Na₃YSi₆O₁₅ – Na₃NdSi₆O₁₅.xH₂O (Haile, 1992) and K₃NdSi₆O₁₅ (Haile, Siegrist, Laudise & Wuensch, 1991; Pushcharovskii, Karpov, Pobedinskaya & Belov, 1977) – are both based on single (Si₂O₅) layers. Other analogs, such as Na₂LiYSi₆O₁₅, take on a double-chain configuration (Gunawardane, Howie & Glasser, 1982). The structural differences between the Na–Y compound and the two Nd compounds are particularly striking as the three phases crystallize under a similar range of hydrothermal conditions (Haile, Wuensch, & Laudise, 1993; Haile, Wuensch, Siegrist & Laudise, 1993), with the exception of the slightly higher temperatures required for the synthesis of the Na–Y compound relative to the Na–Nd compound.

The rarity of the double dreier-ring results from the small separation between neighboring silicate tetrahedra encountered in this configuration relative to those typically observed in layered structures. A useful measure of the intratetrahedral separation is the distance between neighboring Si⁴⁺ ions. This latter quantity, $d(\text{Si}—\text{Si})$, is maximized when the angle defined by O—Si—O takes on a value of 180°. However, because of the partially covalent nature of the Si—O bond, this angle typically has a significantly smaller value of approximately 140°. We see from Table 2 that in order to both accommodate the small Si—O—Si angles within the double dreier-ring unit and maintain as large a Si—Si separation as possible, the Si—O bond lengths in Na₃YSi₆O₁₅ are unusually long: 1.641 (11) Å. The Si—O—Si angles, Si—Si distances, Si—O_{br} distances and cation properties of the three M₃RSi₆O₁₅ synthesized in our laboratory are compared in Table 3. For further comparison, the values for a linear and a bent Si₂O₇ double tetrahedra, with ideal bond angles (O—Si—O = 109°) and ideal bond lengths (Si—O = 1.62 Å), are also included. We notice immediately that smaller Si—O—Si angles do, in fact, correspond to smaller Si—Si separations. In the layered compounds, the distances take on a greater range of values, but are, on average, larger than in Na₃YSi₆O₁₅. We also note that, despite the sizable difference in the radii of K and Na, both neodymium compounds form layered structures. We would thus not expect the difference in ionic radii between Nd and Y to explain the

great differences between the two sodium compounds. Indeed, we can certainly imagine a layered compound in which the smaller ionic radius of Y is accommodated, even if it is not isostructural with either of the Nd phases.

For a plausible explanation of the structural differences, we look to the remaining significant chemical difference between Y and Nd: the electronegativity. Liebau (1985) has suggested that the electronegativity of a guest cation in a silicate can greatly influence the topology that is assumed by a structure by determining

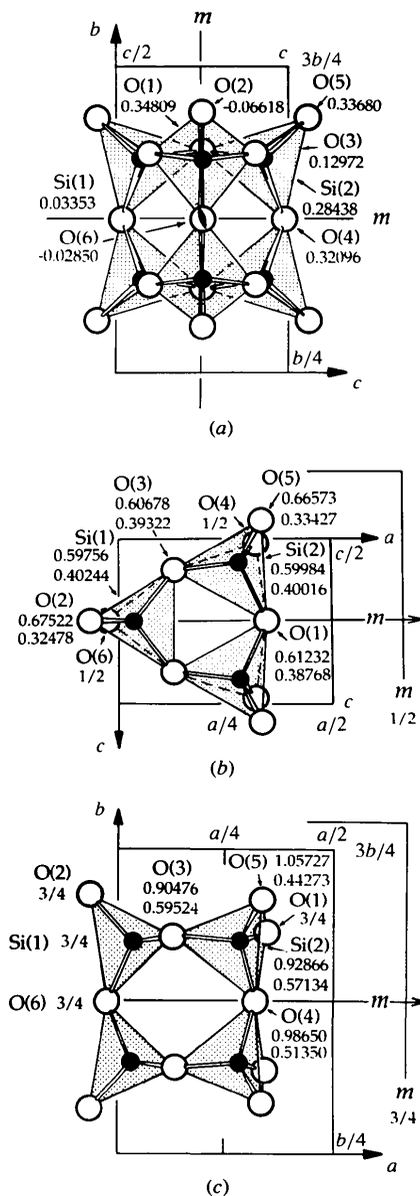


Fig. 1. The isolated Si₆O₁₅⁶⁻ double dreier-ring with point group 2mm, which is found in Na₃YSi₆O₁₅. The unit that is depicted is on the symmetry line ($x, \frac{1}{2}, \frac{3}{4}$) with the dreier rings centered at $x \sim 0.200$. (a) Projection along b; (b) projection along a; (c) projection along c.

Table 3. Comparison of structural parameters in three $A_3\text{MeSi}_6\text{O}_{15}$ compounds and an ideal Si_2O_7 double tetrahedra in two configurations (ionic radii listed are those of Shannon & Prewitt, 1969)

	'Ideal' Si_2O_7		$\text{Na}_3\text{YSi}_6\text{O}_{15}$	$\text{Na}_3\text{NdSi}_6\text{O}_{15} \cdot x\text{H}_2\text{O}$	$\text{K}_3\text{NdSi}_6\text{O}_{15} \cdot x\text{H}_2\text{O}$
Structure	Linear	Bent	Double ring	Layered	Layered
Si—O—Si ($^\circ$)	180	140	133 (2)	142 (6)	145 (8)
d (Si—Si) (\AA)	3.24	3.08	3.02 (2)	3.08 (5)	3.08 (7)
d (Si—O _{br}) (\AA)	1.62	1.62	1.641 (11)	1.628 (14)	1.620 (10)
3+ cation	—	—	Y	Nd	Nd
Coordination number	—	—	6	7	6
Ionic radius (\AA)	—	—	0.900	~ 0.99	0.983
Electronegativity	—	—	1.22	1.14	1.14
1+ cation	—	—	Na	Na	K
Coordination number	—	—	5, 6	4, 5	6, 7
Ionic radius (\AA)	—	—	1.00, 1.02	0.99, 1.00	1.38, 1.46
Electronegativity	—	—	0.93	0.93	0.82

the effective charge of the silicate anionic framework. In the limit of purely ionic bonding between the silicate anion and the guest cation, the effective charge per silicate tetrahedron in an ideal phyllosilicate (three bridging O atoms and one terminating per Si) is -1 . As the electronegativity of the incorporated cation(s) increases, however, the effective charge per silicate tetrahedra decreases. Consequently, maintenance of a large spatial separation between $\text{SiO}_4^{-(1-\delta)}$ tetrahedra becomes less important. Liebau has further suggested that when iso-valent cations are incorporated into a silicate, the electronegativity of the cation has, in fact, a greater impact on the structure of the silicate anion than the ionic size of the cation. It is thus plausible that the greater electronegativity of Y (1.22) over Nd (1.14) facilitates the formation of the double dreier-ring by decreasing the effective charge per silicate tetrahedra, thereby reducing repulsive coulombic interactions and permitting covalent bonding considerations to dominate. It is of interest to note that the single heretofore reported compound

that contains a double dreier-ring is an organosilicate: $[\text{Ni}(\text{H}_2\text{NCH}_2\text{CH}_2\text{NH}_2)_3]_3[\text{Si}_6\text{O}_{15}] \cdot \text{H}_2\text{O}$ (Smolin, 1970), a compound in which we would expect the 'cation' not to be particularly electropositive.

Turning from the origin of the structure to its topology, a view of the structure along *c*, Fig. 2, shows that the (YO_6) octahedra link the isolated Si_6O_{15} units together to form a three-dimensional framework. Each apex of a particular yttrium octahedron is connected to a different silica unit. These octahedra are rather regular, with the three transpolyhedral angles precisely equal to 180° (consistent with the $..2/m$ site symmetry for Y). The remaining 12 bond angles do not differ from orthogonality by more than 3.5° . Additionally, all O atoms participating in yttrium octahedra are bonded to only one Si atom (*i.e.* are terminating), as one might expect *a priori* for charge-balance reasons.

Sodium polyhedra: configuration, linkage and possible conduction paths

One of several criteria necessary for high ionic conductivity in a compound is the presence of pathways through which the mobile species can readily migrate. A second criteria for high ionic conduction is that the pathways for ion migration connect energetically equivalent sites that are present in numbers greater than the number of cations available to fill them. Fig. 2 reveals that the silica–yttria framework defines a set of large tunnels that extend along *c*. These are ellipsoidal in cross-section, measuring approximately 3.75 \AA along *a*, and extending 11.99 \AA in the direction of *b*.

Both the independent Na^+ ions are situated within these tunnels. Fig. 3 presents a projection of the contents of the channel along *a*, the direction of its shortest axis. As can be seen from Fig. 3, two Na(2) sites are located within a cage defined by ten oxygen ions. The ten-cornered cage is centered at $4e \ 2mm \ (x, \frac{1}{2}, \frac{1}{4})$, with $x \approx 0.4596$. Of the ten anions that form the cage, four oxygen ions are located at a distance of 2.643 \AA , two are at 3.035 \AA , and another four are 3.226 \AA from its center, Table 3. Tenfold coordination is rare for sodium and unfavorable here as well. Six of the interatomic distances from the center to the oxygen anions are much too large even for a ninefold-coordinated sodium ion with an ionic radius of 1.32 \AA [$R_{\text{O}^{2-}}^{[3]} = 1.36 \text{ \AA}$ (Shannon & Prewitt, 1969)]. The Na^+ accordingly pulls off to either side and slightly towards O(4), Fig. 4. There, it can acquire five neighbors at comparable distances of 2.525 – 2.670 \AA , while another five second-nearest neighbors are located at distances of 3.333 – 3.634 \AA . Taken as a single group, the 'coordination polyhedron' formed by the five nearest oxygen neighbors has a rather remarkable configuration. The Na(2) site is almost coplanar with the triangle defined by O(1) and a pair of O(4) ions and, in fact, barely within the polyhedron; the plane defined by Na(2) and the remaining pair of

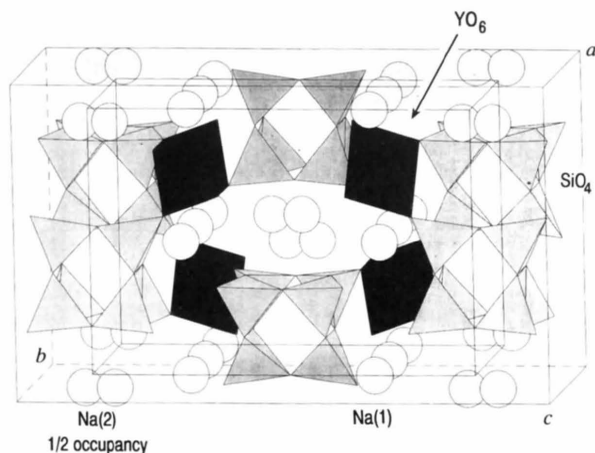


Fig. 2. Structure of $\text{Na}_3\text{YSi}_6\text{O}_{15}$ viewed along *c* using the ATOMS program from Shape Software. The SiO_4 and YO_6 units are represented as polyhedra. The contents of two cells along *c* are depicted.

nearest neighbors, O(5), is close to being normal to the preceding triangle of neighbors. The two Na(2) sites within the cage are separated only in the *b* direction and by only 1.2 Å. The proximity suggests they are too close to be simultaneously occupied. The coordination about the saddle-point at the center of the cage suggests that the ion hops rather easily between the sites.

The alkali ion designated Na(1) (also located in the tunnel) sits within a distorted octahedron formed by six neighboring oxygen ions at distances that range from 2.283 to 2.571 Å, Fig. 3. These polyhedra share O(2) corners to build chains extending along *c* and situated at either side of the tunnel. The Na(2) cages are also joined into chains along *c*, in this case, through sharing of the O(4)–O(4') edge, which is oriented approximately along *a*. These two chains of polyhedra are connected to each other *via* the O(5) atoms: the O(5)–O(5') edge of the Na(2) cage joins the corners of two Na(1) octahedra along *c*. Face-sharing is a mode of polyhedral linkage that would be more conducive to a pathway of high mobility than the corner- and edge-sharing exhibited, respectively, by the Na(1) and Na(2) polyhedra. However, the spaciousness of the channels depicted in Fig. 2 suggests that there are many local minima in which a cation migrating through the structure might temporarily reside, rather than jumping directly between normal sites. Two such potential sites have been identified in Fig. 3 and their nearest neighbors are provided in Table 4. In fact, the presence of such sites suggests that Na₃Si₆O₁₅ will be an *intrinsic* fast-ion conductor – a material in which the structure contains vacant sites easily accessible to a jumping ion without the introduction of aliovalent dopant ions.

Fig. 3 reveals that the first site, at *8i m* (0.54, 0.28, $\frac{1}{2}$), might be easily accessible to a cation jumping from one Na(1) site to another *via* the rather open sides of the distorted octahedra. The distance from the proposed interstitial site to the two closest O atoms,

O(1) and O(2), is 1.98 Å, a value not too much less than the expected interatomic distance between Na⁺ and O²⁻ of 2.35 Å, when the ions are four- and threefold coordinated, respectively (Shannon & Prewitt, 1969). In fact, the coordination about this site, by four more O atoms at distances of less than 3 Å, suggests that it should not be too much higher in energy than the normal sites.

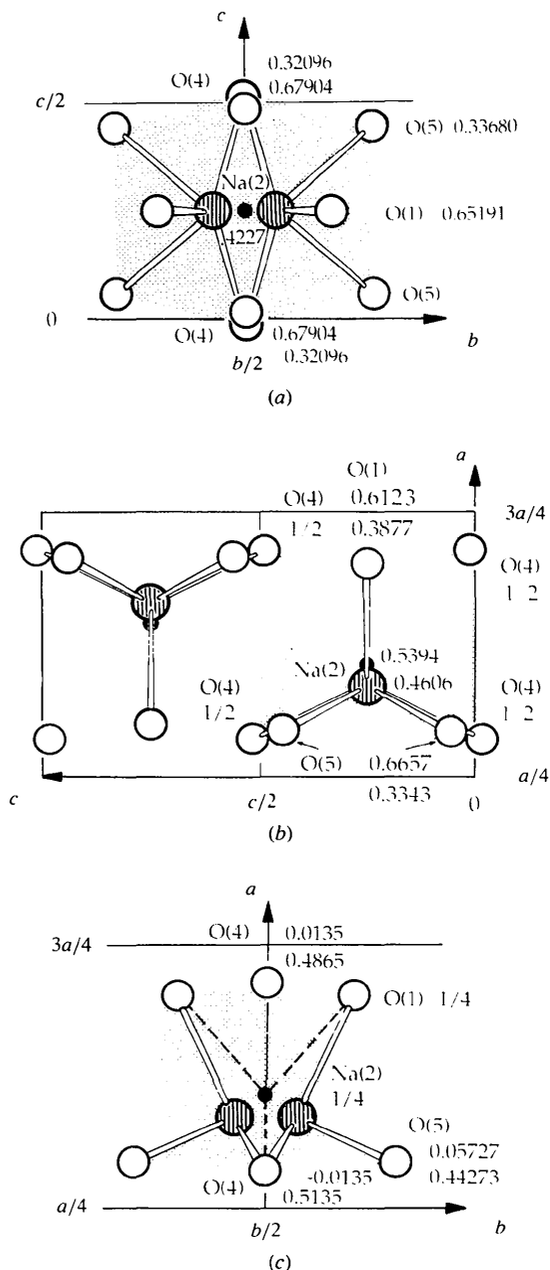


Fig. 4. Projections of the ten-membered cage surrounding Na(2) sites. Location of the proposed saddle point is indicated with a small dot. (a) Projection along *a*; (b) projection along *b*; (c) projection along *c*.

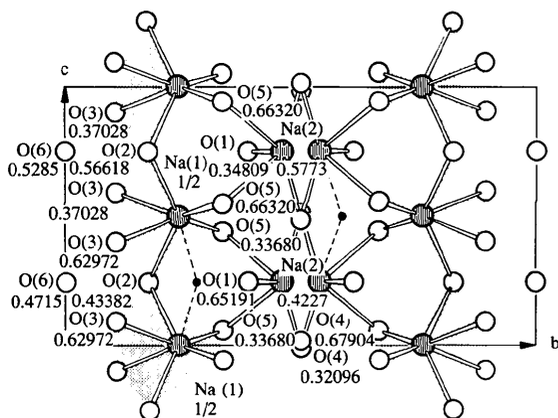


Fig. 3. Projection along *a*, showing the contents of the tunnel, Na polyhedra, connectivity and the possible interstitial site along the Na(1)–Na(1) jump.

Table 4. Locations and nearest neighbors of proposed interstitial sites

Numbers in parentheses identify atoms in terms of Table 1. 2× indicates multiplicity of the bond. Standard deviations are not given as sites are proposed and the positions have not been refined from X-ray data.

Site	Coordinates	Neighbor	Distance	Neighbor	Distance
8i .m	0.54, 0.28, $\frac{1}{2}$	O(1)	1.981	Si(1)	2.806
		O(2)	1.982	O(5) 2×	2.813
		Na(1) 2×	2.235	O(5') 2×	2.979
		O(5) 2×	2.119	Si(1) 2×	2.340
8f .2.	$\frac{1}{2}$, 0.41, $\frac{1}{2}$	O(4) 2×	2.324	Na(2) 2×	2.375
		O(1) 2×	2.654	Na(1)	2.594

The second interstitial site at 8f 2 ($\frac{1}{2}$, 0.41, $\frac{1}{2}$), Table 4, appears to be an even more favorable local minimum. All the four nearest neighbors to this site are O atoms, located at distances of 2.12 and 2.32 Å, almost ideal distances for a fourfold-coordinated sodium ion. In fact, only the presence of two Si atoms at distances of 2.34 Å appear to make the site unsuitable for normal occupation. In addition to its favorable coordination polyhedra, the 8f site is likely to be an integral part of a migration pathway, because of its accessibility from the normal sites. For example, migration from an Na(2) site might occur through the O(5)–O(4)–O(4') face of the ten-membered cage, or from an Na(1) site *via* the O(5)–O(5') edge of the distorted octahedron.

In addition to the two proposed interstitial sites, vacancies also appear to result from the one-half occupancy of the Na(2) sites. However, the proximity of neighboring Na(2) sites, as noted above, suggests that simultaneous occupation is unlikely. This, in turn, implies that unoccupied Na(2) sites cannot serve as truly accessible vacancies to Na⁺ ions moving from one cage to another. Alternatively, given the dimensions of the ten-cornered cage, it is quite likely that high concentrations of a divalent dopant such as Ca or Sr can be incorporated at this site to render the material an extrinsic conductor.

Although potential interstitial/vacant sites have been identified, there is certainly no proof that these sites truly represent local minima that are, at least, temporarily occupied. If, in fact, these interstitial sites are not energetically favorable and thus have extremely low occupancies, alkali-ion migration directly between normal sites would still remain a possibility. For example, an alkali ion might jump between Na(2) cages sites *via* the shared O(4)–O(4') edge. As the midpoint of the edge is at $4a .2/m$. ($\frac{1}{2}$, $\frac{1}{2}$, $\frac{1}{2}$), a site of $\bar{1}$ symmetry, the Na⁺ ion would be required to squeeze directly through the midpoint, or be able to pass at two locations along the edge related by inversion symmetry. The length of the O(4)–O(4') edge is 3.755 Å, and, being somewhat smaller than twice the sum of the ionic radii of sodium and oxygen, suggests that a straight line path is the more probable of the two options. Following a path that leads around the O(4)–O(4') edge, on the other

hand, is essentially equivalent to utilizing the site earlier identified at the 8f position.

The nature of the structure of a fast-ion conductor is such that the energy barriers to motion of the mobile species are very small and thus the mobile ion resides in a shallow potential well. This is commonly reflected in large thermal vibration parameters for the mobile ion, particularly along the direction of the diffusion jump. Table 1 shows the anisotropic temperature factor coefficients for Na(2) are indeed large. In particular, the value of $5.20 \times 10^{-2} \text{Å}^2$ for U_{22} further supports the notion of jumps along **b** between adjacent sites within the ten-cornered cage. The value of $2.46 \times 10^{-2} \text{Å}^2$ for U_{33} , approximately in the direction of the shared edge between neighboring cages, is the smallest of the diagonal elements, but is still of appreciable magnitude. The temperature factor coefficients for Na(1) are moderate, consistent with the shorter Na–O bond distances, and indicate a smaller degree of anisotropy, consistent with the more even distribution of nearest-neighbor anions. The anisotropy, nevertheless, reflects the distorted nature of the octahedron in that the largest thermal displacement occurs within the *ac* plane and in a direction along which the arrangement of surrounding anions was noted to be more open.

To this point we have been unable to grow crystals large enough to perform conductivity measurements and thus confirm the presence of fast-ion conduction in the structure. Attempts to densify pressed pellets led to melting and the formation of a stable glass. It is hoped that the transport properties and their dependence on crystallographic direction can be determined from the synthesis of larger crystals, as done for phases encountered in the K₂O–Nd₂O₃–SiO₂ system (Haile, Wuensch, Siegrist & Laudise, 1992).

Concluding remarks

We have found that the structure of Na₃YSi₆O₁₅ contains a silicate unit based on unique double dreier-rings. We have proposed that such a configuration, rather than the layered structures observed in the Na₃NdSi₆O₁₅·*x*H₂O and K₃NdSi₆O₁₅ analogs, is obtained because of the greater electronegativity of Y relative to Nd, rather than the difference in size. In order to accommodate the small Si–O–Si angles which result from the double dreier-ring configuration, the Si–O bond lengths in Na₃YSi₆O₁₅ are unusually long. When taken together with the (YO₆) octahedra, the structure formed is a three-dimensional framework with channels along **c**. The placement of Na⁺ ions within these tunnels, the existence of potential interstitial sites and the nature of the polyhedral connectivity between these and the normal sites suggest easy ion migration and thus high intrinsic conductivity in the *c* direction. Very high thermal vibration amplitudes are indeed observed for Na(2), often indicative of high-ionic mobility. However, the

highest thermal displacements of this species occur along directions perpendicular to the proposed direction of motion, approximately towards the neighboring partially occupied site.

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Structure of Cs₃(HSO₄)₂(H₂PO₄) – a New Compound with a Superprotonic Phase Transition

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Abstract

Solid-solution studies in the CsHSO₄–CsH₂PO₄ system carried out in a search for low-temperature proton conducting materials yielded the new compound Cs₃(HSO₄)₂(H₂PO₄) [tricesium bis(hydrogensulfate) dihydrogenphosphate]. Single-crystal X-ray measurements (performed at room temperature) revealed that Cs₃(HSO₄)₂(H₂PO₄) crystallizes in space group *P2₁/n* and has lattice parameters *a* = 19.546 (3), *b* = 7.8798 (10), *c* = 9.1854 (17) Å and *β* = 100.536 (14)°. With four formula units in the unit cell, and a cell volume of 1390.7 (4) Å³, Cs₃(HSO₄)₂(H₂PO₄) has a calculated density of 3.295 Mg m⁻³. 18 non-H atoms were located in the asymmetric unit. Refinement using anisotropic temperature factors for all 18 non-H atoms

yielded weighted residuals based on *F*² and *F* values, respectively, of 9.32 and 4.56% for all observed reflections. Hydrogen sites were identified (but not refined) on the basis of geometric considerations. The structure contains zigzag chains of hydrogen-bonded anion tetrahedra that are, in turn, bonded to one another to form a three-dimensional structure. In the temperature range 381–398 K the compound transforms into a structure which is rhombohedral, pseudo-body-centered cubic, with lattice constants *a* = 6.95 (2) Å and *α* ≈ 90°.

Introduction

Many classes of solid acid sulfates and selenates are known to undergo superprotonic phase transitions