



Crystal structure data

Space group: Pnam (62)

S (goodness of fit) = 1.76

Lattice parameters: a=9.1230(4) Å

b=10.7469(4) Å

c=2.9562(1) Å

_=10.81% R

R_B=3.86%

$Na_{0.76}Fe_{0.79}Ti_{1.21}O_4$ – the new CaFe₂O₄ structure type compound obtained from natural ilmenite sand under hydrothermal conditions

Lidija Mancic¹, Bojan Marinkovic², Paula Jardim², Fernanado Rizzo², Katarina Marinkovic¹, Olivera Milosevic¹

¹Institute of Technical Sciences of Serbian Academy of Sciences and Arts, Belgrade, Serbia ²Departamento de Ciencia dos Materiais e Metalurgia, Pontificia Universidade Catolica, Rio de Janeiro, Brazil

 $CaFe_{2}O_{4}$ structure type, named after the prototype compound, is a less known crystal structure from the $A_{2}BX_{4}$ stoichiometry. Its basic motif is a double rutile chain composed of octahedras connected by edges. A double chain itself is connected to another four double chains through corners, forming in that way tunnels oriented along the shortest unit-cell axis. In order to compensate the negative charge of framework, the tunnels are occupied most frequently by Na⁺ or Ca²⁺ and more rarely by Sr²⁺ or Ba²⁺. CaFe₂O₄ structure type compounds are generally obtained through a high-temperature synthesis route. Here, we report a simple low-temperature hydrothermal synthesis of Na, Fe, Ti, 2, O₄ phase with CaFe₂O₄ type structure, by reacting natural ilmenite sand (Millenium Inorg Chem., Brazil) with 10M NaOH solution at temperatures bellow 200°C. The typical product is sodium-deficient Na, Fe, Ti_{2,4}O₄ phase obtained in content of 92wt%. Up to now there was just one report in literature on hydrothermal synthesis of a CaFe₃O₄ type compound related to Na₃Mn₄Te₂O₁₂ compound that adopts a superstructure intimately related to the CaFe₂O₄ structure type [1].

Table 1. Rietveld refinement data for $Na_{0.76}Fe_{0.79}Ti_{1.21}O_4$

0.75 0.26

0.75

0.75

0.75

0.47

Another possible indication from N_i for the Fe1 and Fe2 is that the Fe3+ might not be

randomly distributed over these two sites, so far as conventional XRD can determine. This is also indicative from determined bond lengths: Ti1,Fe1-O is 1.95 Å, while Ti2,Fe2-O is 1.98 Å. Having in mind that the mean Ti-O bond length within octahedral polyhedron for many simple and complex oxides is around 1.96 Å, while the mean Fe-O distance assumed from the CaFe₂O₄ is 2.02 Å, obtained value of for Ti2,Fe2-O bond length is expected, while for Til,Fel-O is somewhat smaller then it would be expected. Therefore, the indication of

TEM image of typical $Na_{0.76}Fe_{0.79}Ti_{1.21}O_4$ crystal and corresponding SEAD pattern

TEM image shows submicron to micron Na_{0.76}Fe_{0.79}Ti_{1.21}O₄ crystals with well defined crystal facets and growth direction. Zone axis is determined from SEAD pattern to be [100] while the direction of growth is [001] assuming Pnam space

group. This means that the tunnels are oriented along the growth direction of the

crystals. The crystals are thinnest in [100] direction. The crystallographic direction of [010] is perpendicular to the growth direction. The crystal facets are

composed of {100}, {010} and {031} crystal planes, being {100} the most

0.24444(99) 0.34783(82) 0.75

0.11390(35) 0.75 0.75 0.75 0.53 0.74

0.64888(90) 0.75

Fe3+ ordering still has to be verified by neutron diffraction.

0.11390(35)

0.60468(31)

0.60468(31)

0.99072(95)

0.21033(84)

0.92440(72)

0.05986(35)

0.05986(35)

0.08608(37)

0.08608(37)

0.31080(10)

0.39026(82) 0.46880(12)

0.05710(11)

4c

4c

4c 4c 4c

4c 4c

40



Magnetite Fe_3O_4 is the minority phase, which weight percentage is not higher than 7wt%. Residual un-reacted ilmenite can be remove by increasing synthesis time.





From the X-ray spectrum
measured on a TEM, through
Cliff-Lorimer ratio technique
it was possible to calculate
Ti/Fe ratio [2]:
N _{Ti} /N _{Fe} =k _{TiFe} I _{Ti} /I _{Fe}
to be 1.41 (estimating its
relative error due to counting
statistic of the peaks Ti-Ka
and Fe-K α , as lower than %).



This Ti/Fe ratio deviated ~ 8% from the one calculated from X-ray diffraction pattern (1.53), which suggests that the true Ti/Fe is most likely around 1.5, as indicated independently by two different approaches, Rietveld method and Cliff-Lorimer ratio technique

Intensity [counts

Acknowledgement:

This research is supported through COST 539 action. The author L.M. would like to express his gratitude for post-doc grant to the Ministry of Science and Environment Protection Republic of Serbia and CNPq Brazil.

Literature:

1. C.R. Feger and J.W. Kolis, Acta Cryst., C54 (1998) 1055. 2. Transmission Electron Microscopy IV, Spectrometry, D.B. Williams and C.B. Carter, Plenum Press NY, 1996, p. 600.

extended one.