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# Comments on "Kinetic Study on the Hexacelsian-Celsian Phase Transformation"

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#### Comments on

### "Kinetic Study on the Hexacelsian-Celsian Phase Transformation" by D. Bahat, J. Mater. Sc., <u>5</u> (1970) 805-810

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Bahat [1] studied the hexacelsian-celsian phase transformation in crystallized grains of amorphous  $BaAl_2Si_2O_8$ . For the first stage, the transformation was reported to be controlled by the crystal growth rate. The activation energy for this transformation was evaluated to be 20.1  $\pm$  4 kcal/mole. It was suggested that this transformation does not involve Si-O and Al-O bond openings, since this would require an activation energy of 60 kcal/mole or higher for a bond breaking mechanism.

In recent investigations, the present authors [2,3] have reported a value of 125 kcal/mole for the activation energy of hexacelsian-to-celsian transformation in the Sr- analogue, SrAl<sub>2</sub>Si<sub>2</sub>O<sub>8</sub>. Furthermore, since the transformation involves a structural change, from a double-layered sandwich structure of

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hexacelsian to a three-dimensional framework structure of celsian, Si-O and Al-O bond breaking would be necessary for such a transformation. There appears to be a significant discrepancy between the results of the study by Bahat [1] and the requirements of the structural transformation from hexacelsian to celsian.

In an effort to understand the cause of this discrepancy, the experimental results of Bahat [1] have been analyzed using the Avrami equation [4]:

$$x = 1 - \exp[-(kt)^{n}]$$
 (1)

where x is the volume fraction transformed after time t, k is the reaction rate constant, and n is the Avrami exponent which determines the geometry for interface and diffusion controlled mechanisms. Rearranging eq. (1) and taking logarithm gives

$$\ln[-\ln(1-x)] = n \ln k + n \ln t$$
 (2)

Data from Figure 4 of Bahat's paper [1] were read and analyzed using eq. (2) in the form of plots of ln [-ln(1-x)] vs. ln t (Figure 1) at six different temperatures ranging from 1050 to 1300 °C and times from 0.5 - 8 hours. The values of n and k at various temperatures were evaluated from linear regression analysis of the data and are listed in Table I. An average value of n close to 0.5 suggests [5] the possibility of a diffusion controlled growth with a plate geometry.

Temperature dependence of k is expressed by the Arrhenius equation:

$$k = \nu \exp[-E/RT]$$
(3)

where  $\nu$  is the frequency factor, E the activation energy, R the gas constant, and T the isothermal reaction temperature in Kelvin. Arrhenius plots of *ln k vs. 1/T* for the data of Table I are shown in Figure 2. A linear regression analysis of the data (the 1050 and 1300 °C data excluded) gave a value of 89 ± 14<sup>1</sup> kcal/mole for E and  $\nu = 5 \times 10^8$ /s, with correlation coefficient of -0.977. Least-squares linear regression analysis of five data points (the 1050 °C data excluded) resulted in  $65 \pm 17^1$  kcal/mole for E,  $\nu = 1.05 \times 10^5$ /s, with correlation coefficient of -0.913 indicating a poorer fit. When all the six data points are used in the statistical analysis, a lower value of activation energy is obtained, but the 1050 °C data lies outside the 95% confidence interval.

Plots of hexacelsian (weight %) vs. time, when extrapolated to t = 0, should result in zero per cent transformation since the starting material was 100% hexacelsian in all cases. But the data in Figure 4 of Bahat's paper do not extrapolate to 100% hexacelsian at zero time of heat treatments. Furthermore, these extrapolated values at t = 0, were taken as the values of reaction rate constants k at various temperatures by Bahat which is not correct.

<sup>&</sup>lt;sup>1</sup>one standard deviation

The activation energy of hexacelsian-to-celsian transformation in  $BaAl_2Si_2O_8$ , obtained by reanalysis of Bahat's data using the Avrami equation, is significantly greater than the value of 20.1  $\pm$ 4 kcal/mole reported by Bahat. The recalculated values of E appear to be more reasonable and are consistent with a mechanism which involves breaking of the Al-O and/or Si-O bonds. The breaking of these bonds would be necessary for transformation of a layered hexacelsian structure into a three-dimensional feldspar structure of celsian. These values are also in better agreement with a value of 60 kcal/mole, the energy required for the ordering of albite [6] which involves Si-O and Al-O bond openings.

A further study is needed to obtain a more reliable value of the activation energy. We are presently investigating the kinetics of hexacelsian-celsian phase transformation using a quantitative Xray diffraction technique and the results of this study would be reported in the near future.

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Temperature (°C)	k x 10 <sup>6</sup> (s <sup>-1</sup> )	n
1050	20.9	1.4
1100	3.7	0.37
1150	7.2	0.38
1200	41.8	0.62
1250	73.3	0.79
1300	48.9	0.54

TABLE I. Values' of Avrami Parameter, n, and Reaction Rate Constant, k, for Hexacelsian-to-Celsian Transformation in BaAl<sub>2</sub>Si<sub>2</sub>O<sub>8</sub> at Various Temperatures

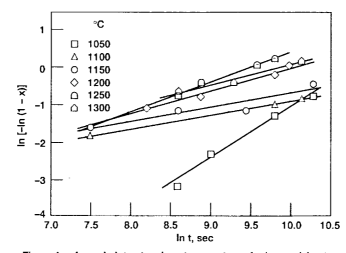
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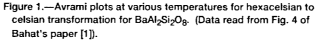
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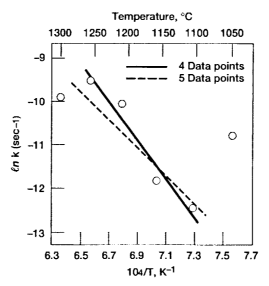
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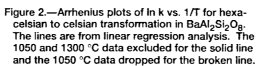
'Calculated from data of Bahat [1]

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