Comparative study of morphology and kinetic-thermodynamic parameters of gel-grown lead iron mixed levo-tartrate crystals

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Abstract

Pure and mixed crystals of metal tartrate compounds have varieties of applications in the different fields of science and technology. Lead tartrate finds application as additive in gasoline to prevent knocking in motors and iron tartrate finds application as one of the prominent species in apple juice. In the present study, lead-iron mixed levo-tartrate crystals were grown by single diffusion gel growth technique. Silica hydro-gel was the growth medium. Dendrite crystals were obtained at the gel-liquid interface. The crystals were characterized by Thermo gravimetric analysis and kinetic as well as thermodynamic parameters were evaluated for the different stage of decomposition.

Keywords: Lead-iron mixed levo-tartrate crystals, gel growth, thermo gravimetric analysis, kinetic parameters, thermodynamic parameters.

1. Introduction

The metal tartrate compounds find various applications in the different fields of science and technology, for example, tanning action of iron tartrate to tan the skin [1], ferroelectric applications of sodium-pottasium tartrate [2], addition of lead tartrate in gasoline to prevent knocking in motors [3] and ferroelectric-ferroelastic applications of sodium-ammonium tartrate [4]. The gel growth technique is found to be suitable to grow tartrate compound crystals, which is elaborately described by Henisch [5]. The growth of several tartrate compound crystals by the gel technique and their characterization were reported, for instance, cadmium tartrate [6-11], iron tartrate [12] and lead tartrate [13, 14]. In the present study, the present author has attempted to grow the crystals of lead-iron mixed levo-tartrate and characterized them by TGA and evaluated the kinetic as well as thermodynamic parameters for the dehydration and decomposition stage.

2. Experimental

In the present study, the single diffusion method [14-17] was employed for the growth of lead-iron mixed levo-tartrate crystals. The silica hydro gel was used as a growth medium. To prepare the gel, a solution of sodium metasilicate of 1.05 specific gravity and 1 M solution of levo tartaric acid were mixed in such a manner that the pH of the mixture was set at 4.5. The gel solution was poured in to glass test tubes of 15 cm length and 2.5 cm diameter and allowed to set in the gel form. The supematant solution containing the mixture of 1 M, 6 ml lead nitrate solution and 1 M, 4 ml iron nitrate nonahydrate solution was poured on the set gel carefully without damaging the gel.

All the chemicals were of AR grade and obtained from Sigma Aldrich. The following reaction is expected to take place in the formation of lead-iron mixed levo-tartrate crystals.

\[
(1 - X)\text{Pb(NO}_3\text{)}_2 + X\text{Fe(NO}_3\text{)}_2 \cdot 9\text{H}_2\text{O} + \text{H}_2\text{C}_4\text{H}_4\text{O}_6 + n\text{H}_2\text{O} \rightarrow \text{Fe}\text{Pb}(1 - X)\text{C}_4\text{H}_4\text{O}_6 \cdot n\text{H}_2\text{O} + 4\text{HNO}_3 + 8\text{H}_2\text{O} + 1/2\text{O}_2
\]

The amount of HNO3 produced is very less in comparison to the nutrients being supplied to the growing crystals and hence no major limitation is imposed [5, 14-17].
Photograph of the grown crystals is shown in the figure 1.

Fig 1: Dendrite crystals of lead-iron mixed levo tartrate

Earlier, the present author has grown and reported [18] the lead-iron mixed levo tartrate crystals by taking the composition of the supernatant solutions as 1 M, 8 ml lead nitrate + 1 M, 2 ml iron nitrate nonahydrate. The crystals observed were reddish, long, dense and dendrite in nature.

In the present case, as the volume concentration of lead nitrate was decreased from 8 ml to 6 ml and iron nitrate nonahydrate was increased from 2 ml to 4 ml, the slight change in the morphology of the grown crystals was observed. Though the crystals were reddish and dendrite in nature, but the length and density of the crystals were altered. Both the length as well as density was found to decrease. This may be due to the reduced volume concentration of lead nitrate solution.

The grown crystals were characterized by TGA. The TGA was recorder on Perkin Elmer make instrument, model Pyris-1 DSC, Pyris-1 TGA, DTA-7. The data were recorded from room temperature to 700 °C at a heating rate of 10 °C min⁻¹ in atmosphere of air.

3. Result and Discussion

There are reports available in literature on thermal studies on pure and mixed metal tartrate systems, for example, cadmium tartrate [19], mercuric iodate crystals [20], iron-manganese levo-tartrate tartrate compound [21], ternary iron-manganese-cobalt tartrate compound [15], ternary iron-manganese-nickel tartrate compound [16] and lead-cadmium mixed levo-tartrate crystals [17].

The TGA curve for lead-iron mixed levo-tartrate crystals is shown in figure 2.

There are three stages of decomposition starting from room temperature of 45 °C. There is no decomposition of sample upto temperature 160 °C. The first stage of decomposition is observed in the temperature interval of 160 to 200 °C with the weight loss is about 2.44%. The second stage of decomposition is in the temperature interval of 200 to 270 °C with the weight loss is about 26.06%. The third stage of decomposition is observed in the temperature interval of 270 to 400 °C with the weight loss is about 38.25%. The number of water molecules associated with the crystals was calculated and found to be 0.5.

Many researchers have used thermo gravimetric data to calculate the kinetic parameters of solid state reaction including mass [22-26]. By using kinetic parameters such as order of reaction, frequency factor and energy of activation, the shape of curve is determined. With the help of Coats and Redfern [24] relation the kinetic parameters were calculated. In the present study, the kinetic and thermodynamic parameters have been calculated for dehydration as well as for decomposition of crystals.
The thermodynamic parameters have been evaluated for the dehydration as well as decomposition stages by using the standard relations. The standard enthalpy of activation \( \Delta^\circ H^o \) was calculated by using the relation 3.

\[
\Delta^\circ H^o = E - 2RT
\]

The standard entropy of activation \( \Delta^\circ S^o \) was calculated by using relation 4.

\[
\Delta^\circ S^o = 2.303 \times R \times \log_{10} \left[ \frac{A h}{k T} \right]
\]

Here, \( k \) is the Boltzmann constant, \( h \) the Planck constant, \( T \) the temperature and \( A \) is the frequency factor.

The values of standard enthalpy, standard entropy and standard Gibbs free energy for the dehydration as well as decomposition stage are listed in the table 3.

**Table 3: Thermodynamic parameters for lead-iron mixed levo-tartrate crystals**

<table>
<thead>
<tr>
<th>Stage</th>
<th>Standard enthalpy ( \Delta^\circ H^o ) (kJ mol(^{-1}))</th>
<th>Standard entropy ( \Delta^\circ S^o ) (kJ mol(^{-1}) K(^{-1}))</th>
<th>Standard Gibbs free energy ( \Delta^\circ G^o ) (kJ mol(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dehydration</td>
<td>163.37</td>
<td>0.323</td>
<td>163.05</td>
</tr>
<tr>
<td>Decomposition</td>
<td>45.94</td>
<td>0.0458</td>
<td>45.86</td>
</tr>
</tbody>
</table>

Positive values of the standard enthalpy and the standard entropy of activation suggest that the process is spontaneous at high temperatures and the positive values of standard Gibbs free energy suggest that the process is thermodynamically unstable. Comparing with the earlier reported work, considerable change in the values of kinetic and thermodynamic parameters, clearly indicate the effect of variation of volume concentration of the supernatant solutions on the grown crystals.

**4. Conclusion**

Pure lead-iron mixed levo-tartrate crystals were grown in silica gel by using the mixture of 1 M, 6 ml lead nitrate and 1 M, 4 ml iron nitrate nonahydrate solution as supernatant solution. A change in the morphology of the crystals as well as kinetic-thermodynamic parameters was observed with the change in the volume concentration of the supernatant solutions. From TG curves, it was found that the crystals were thermally unstable. Upon heating, they become anhydrous and decomposed into metal oxide through a single stage of carbonate. The kinetic parameters were evaluated for both the stages by using Coats and Redfern relation. The thermodynamic parameters were evaluated for both the stages by using the standard relations. The presence of water molecules was detected and calculated.

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**6. References**