Study OF binary SYSTEMS NdF3- MF (M = Li, Na, K): experimental, modeling and thermodynamic computation

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Abstract. The excess molar enthalpies HEm of the binary systems MF-NdF3 (M = Li, Na, K) were measured in the present work by high temperature calorimetry on a wide temperature (1220 K < T < 1400 K) and composition range. Some points of the equilibrium phase diagram have been also obtained by differential thermal analysis. Using the Hoch-Arpshofen model we represent the excess quantities of different systems. This will allow us thereafter to calculate the phase diagrams of binary systems NdF3-MF (M = Li, Na).

1 Introduction

The rare earth metals commonly called lanthanides are very reactive and form very stable compounds characterized by free energy of formation very high. Reduction with reducing agents (carbon, carbon monoxide, hydrogen) to obtain these metals is not feasible. The electrochemical method to extract the rare earth metals from their oxides or halides is widely used industrially. This method is also used to refine the crude metals obtained after the first electrolysis. Thousands of tons of rare earth metals are currently produced worldwide each year showing the industrial issue. The lanthanide fluorides are generally used as the electrolyte because of their low vapor pressure and their handling in the air is possible because much less hygroscopic than the corresponding chlorides. The electrolyte consists mainly of lanthanide fluoride, barium fluoride and lithium fluoride ... Besides the production processes of rare earth metals from ores by electrolysis of molten salts or by metallothermy [1], many researches are underway in the nuclear field such as: reprocessing of nuclear wastes [2], recycling of spent nuclear fuel [3], and in the lighting industry [4] (high pressure discharge halide lamps). All processes in development often encounter the lack of data on the compounds of lanthanides solid/liquid at high temperature, in particular, lanthanide halides and their mixtures with alkali metal halides.

In this work, we present the results of mixing enthalpy and differential thermal analysis obtained in our study of binary systems MF-NdF3 (M = Li, Na, K). The results will be compared with those of systems "neighbors" LnF3-MF (Ln = La, Y, Yb, Al). Using the Hoch-Arpshofen model we represent the excess quantities of different systems. The variation of the interaction parameter, derived from the thermodynamic model for many binary systems, against the ionic potential has been studied. The thermodynamic quantities represented using Hoch-Arpshofen model have allowed us then to calculate the phase diagrams of binary systems NdF3-MF (M = Li, Na).

2 Experimental

2.1 Quality of salts used

Salts used are products of very high purity: 99.99%, (Merck Suprapur for alkali fluorides and Johnson Matthey RE acton for NdF3). Before use, each salt was thermally treated to obtain complete dehydration and are then stored in a glove box under purified argon. These fluorides are very corrosive with high temperature, that required using a platinum crucible or a graphite crucible for these experiments.

2.2 Apparatus and experimental technique

Two calorimetric apparatus have been used for this study:
- A Calvet-type high-temperature microcalorimeter
- A Setaram high-temperature calorimeter operable up 1800 K. The "direct drop method" or the "indirect drop method" were used to synthesize in situ the mixtures and to measure the corresponding enthalpy of formation in the temperature range 1220-1400 K. The calorimeter was calibrated under the same experimental conditions with platinum.

2.3 Precision of results
The precision of our results of enthalpy of mixing was evaluated from the formula of "propagation of errors" described previously [6, 7]; It varies between 6 and 25% depending on the binary mixtures.

3 Experimental results and modeling

The excess enthalpies obtained for the studied systems NdF$_3$ - MF (M = Li, Na, K) are negative over the whole concentration range studied. As for systems "neighbors" LnF$_3$-MF (Ln = La, Y, Yb, Al), excess enthalpy becomes more negative when the radius of the alkali metal increases (Figs. 1-3).

The mixing enthalpy was calculated using the thermodynamic model of Hoch-Arpschofen [5] represented with a function of type:

$$\Delta_{mix}H = W n x [1 - (1 - y)^{(n-1)}]$$

where: $W$ is a binary interaction parameter and n integer $\geq 2$.

In this expression, $x$ represents the molar fraction of the atom to which the mixing enthalpy has an extremum for $x > 0.5$, $y$ is the concentration of the other atom (in the case of a binary mixture $y = 1 - x$). According to the authors, this atom must be less "binding".

This model, developed for metallic solutions has been applied to different types of systems: metals - salts, metals - oxides, silicates and ionic solutions [6, 8].

By optimization from the experimental values, the result of the calculation indicate that the value $n = 3$ is the best except for mixtures containing lithium where $n = 2$ allows a better representation of the data (Figs. 1-3). $W$ values were compared to those obtained in mixtures "neighbors" LnF$_3$-MF (Ln = La, Y, Yb, Al) and the variation of the interaction parameter has been studied as a function of the difference of potential ionic cations.

The difference of potential ionic cations, noted $\Delta IP$, is defined by:

$$\Delta IP = \frac{1}{r_{M^+}} - \frac{3}{r_{Nd^{3+}}}$$

$r_{Nd^{3+}}$ is the radius of the rare earth cation and $r_{M^+}$ is the radius of the alkali cation.

A linear representation is obtained for each "family" cation. The slope of the lines obtained with Na and K is substantially the same: therefore we wanted to represent all of these results by a single analytical form involving the size of the alkali cation. The results are satisfactory (Fig. 4). These results are not derived from a purely theoretical but nevertheless allow an estimate of the enthalpy of mixing systems containing rare earth metal for which this quantity has not been measured.

![Figure 1. NdF$_3$-LiF system](image1)

$n = 2$, $x = x$(LiF), $W = -8.3 \pm 1.2$ kJ.mol$^{-1}$.

![Figure 2. NdF$_3$-NaF system](image2)

$n = 3$, $x = x$(NaF), $W = -7.0 \pm 0.8$ kJ.mol$^{-1}$.

![Figure 3. NdF$_3$-KF system](image3)

$n = 3$, $x = x$(KF), $W = -15.4 \pm 1.3$ kJ.mol$^{-1}$.

![Figure 4. LnF$_3$-NaF et LnF$_3$-KF (Ln = La, Nd, Y, Yb, Al)](image4)

$W = 6.89 - 9.68 \Delta IP$ kJ.mol$^{-1}$. 

The precision of our results of enthalpy of mixing was evaluated from the formula of "propagation of errors" described previously [6, 7]; It varies between 6 and 25% depending on the binary mixtures.
The regression coefficient for the binary analysis is R = 0.90.

4 Calculation of phase diagrams

Some points of the phase diagram were obtained by DTA for NdF$_3$-MF (M = Li, Na) systems. However, the coordinates of the invariant points were taken from the literature: NdF$_3$-LiF system [9] and NdF$_3$-NaF system [10].

4.1 NdF$_3$-LiF system

In the case where the solution is regular $S^E = 0$, the calculated phase diagram is shown in Fig. 5. We note that the liquidus NdF$_3$ rich side is not in agreement with our points obtained by DTA. Taking into account the excess entropy, the diagram is calculated and presented in Fig. 5. The agreement between the experimental and calculated values is correct.

![Figure 5. NdF$_3$-LiF system.](image)

4.2 NdF$_3$-NaF system

The calculated phase diagram is presented in Fig. 6 with our experimental points. The agreement is very good except for the decomposition temperature of the compound 5NaF:9NdF$_3$.

![Figure 6. NdF$_3$-NaF system.](image)

5 Summary

The analysis of our experimental results shows that the enthalpy of formation of liquid mixtures is negative over the whole concentration range studied. Model Hoch-Arpshofen is well suited to represent correctly the excess enthalpy of these systems.

References

6. M. Berkani, Thèse de Doctorat, Université de Provence, Marseille (1994)