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Enthalpy of Mixing in Al–Tb Liquid

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Abstract
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Keywords
Al–Tb, mixing, enthalpy, electromagnetic levitation, solution calorimetry

Disciplines
Materials Science and Engineering

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Article

Enthalpy of Mixing in Al–Tb Liquid

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Abstract: The liquid-phase enthalpy of mixing for Al–Tb alloys is measured for 3, 5, 8, 10, and 20 at% Tb at selected temperatures in the range from 1364 to 1439 K. Methods include isothermal solution calorimetry and isoperibolic electromagnetic levitation drop calorimetry. Mixing enthalpy is determined relative to the unmixed pure (Al and Tb) components. The required formation enthalpy for the Al 3 Tb phase is computed from first-principles calculations. Based on our measurements, three different semi-empirical solution models are offered for the excess free energy of the liquid, including regular, subregular, and associate model formulations. These models are also compared with the Miedema model prediction of mixing enthalpy.

Keywords: Al–Tb; mixing; enthalpy; electromagnetic levitation; solution calorimetry

1. Introduction

Phase selection pathways in glass-forming metallic liquids are influenced by the development of short- and medium-range order [1–10], which contribute to local energetics and dynamics. Even in binary glass-forming systems, clear evidence of the influence of ordering in the liquid state has been reported [3–5,11–17]. For example, the marginal glass-forming ability in the Al–RE (aluminum rare-earth) systems has been associated with ordering in the liquid phase [18–29]. In addition, due to their relative simplicity compared to conventional many-component glass-forming alloys, Al–RE and Al–RE–TM (TM denotes transition metal) alloys are excellent model systems for the investigation of phase stability [30–33], metallic glass formation [34–37], liquid properties and crystallization [38–40], and glassy alloy structure/properties [25,41,42]. As with all glass-forming liquids, however, properties may become increasingly temperature dependent near and below the liquidus [43–45], highlighting the need for reliable thermodynamic descriptions of the liquid state. Numerous theoretical/computational studies, thermodynamic assessments, and associated CALPHAD models for the Al–RE systems are available [32,33,46–50], but reported experimental measurements of excess mixing quantities for Al–RE liquids are very limited, as listed for the Al–lanthanides in Table 1.

Table 1. A listing of thermodynamic mixing property measurements, calculations, and phase diagram assessments reported for the Al–RE (lanthanide) binary systems.

<table>
<thead>
<tr>
<th>System</th>
<th>CALPHAD Assessments</th>
<th>FP Calculations of Crystal Phase Energies</th>
<th>Experimental Measurement of Liquid Mixing Properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al–La</td>
<td>[51,52]</td>
<td>[53,54]</td>
<td>[55–57]</td>
</tr>
<tr>
<td>Al–Ce</td>
<td>[52,58,59]</td>
<td>[53]</td>
<td>[60–62]</td>
</tr>
<tr>
<td>Al–Pr</td>
<td>[52,63]</td>
<td>[53]</td>
<td>[63,64]</td>
</tr>
<tr>
<td>Al–Nd</td>
<td>[52,58]</td>
<td>[53]</td>
<td>[64]</td>
</tr>
<tr>
<td>Al–Pm</td>
<td></td>
<td>[53]</td>
<td>*</td>
</tr>
<tr>
<td>Al–Sm</td>
<td>[52,65]</td>
<td>[53,66]</td>
<td>[67]</td>
</tr>
</tbody>
</table>
In the current work, we focus on the Al–Tb binary system (see Figure 1) and make critical measurements of the enthalpy of mixing, \( \Delta H_{\text{mix}} \), for the liquid phase. Specifically, we perform solution calorimetry and electromagnetic levitation drop calorimetry to measure \( \Delta H_{\text{mix}} \) in dilute Al–Tb alloys with Tb content up to \( x_{\text{Tb}} = 0.2 \) (mole fraction). In addition, we compare three different semi-empirical solution models for the excess free energy of the liquid, incorporating regular, subregular, and associate [75] formulations for the mixing enthalpy. These are compared with the Miedema model [76–78] for this system.

2. Calorimetry Experiments

We employ here two different calorimetric techniques to determine \( \Delta H_{\text{mix}} \) for the Al–Tb liquid phase in the stable temperature range. Using an isothermal solution calorimetry (ISC) method, we measure the thermal transient associated with the dissolution of a small amount of Tb in a liquid Al solvent, permitting direct determination of \( \Delta H_{\text{mix}} \) for the liquid phase. Using an electromagnetic levitation drop calorimetry (EMLDC) method, we measure the heat evolved upon rapid cooling of the high temperature equilibrium alloy liquid to a low temperature multi-phase state, from which we determine the high temperature mixing enthalpy, \( \Delta H_{\text{mix}} \). All test specimens were prepared from the pure elemental components (0.9999 Al and 0.999 Tb, by weight (pure Al and Tb were supplied by Cerac, Inc., Milwaukee, WI, USA)). Alloy specimens of desired composition were prepared for EMLDC by arc melting the pure components five times in succession on a water-cooled copper hearth under a high-purity (ultra-high purity (99.999%) argon, supplied by Matheson Company, Des Moines, IA, USA) argon atmosphere. Test specimens were then cut to the desired size using a diamond saw.
The central components of the custom-built ISC are illustrated schematically in Figure 2. The entire assembly shown in the figure is contained within a vacuum chamber, evacuated to $10^{-5}$ torr, refilled to 760 torr with ultra-high purity argon prior to heating, and maintained at this pressure throughout the experiment. For the measurement, a mass ($m_{Tb}$) of pure Tb, initially at ambient temperature ($T_a$), is introduced into a bath of pure Al liquid of known mass ($m_{Al}$), with the system held isothermally at $T_0$ using an integrated thermocouple-based closed-loop control system. The resulting net thermal exchange between the calorimeter and the bath associated with isothermal control is given by

$$q_{cal} = q_{heat} + q_{melt} + q_{mix} + q_0 \quad (1)$$

where the first three terms on the right-hand-side (RHS) are exchange fluxes associated with heating, melting, and mixing of the added material, respectively. The fourth term is the heat flux required to maintain isothermal conditions at $T_0$ in the bath, with no addition of material. Upon the addition of the terbium mass, $m_{Tb}$, to the bath, the net exchange gives rise to a sensible thermal transient which is measured using a thermopile in contact with the melt crucible. An example thermal trace is shown in Figure 3. The measured temperature transient is related to the specimen-bath reaction enthalpy according to:

$$\delta q_{meas} = \int_0^\infty (q_{cal} - q_0)dt = \beta \int_0^\infty (T - T_0)dt, \quad (2)$$

and we compute the incremental heat of mixing for this transient as

$$\delta q_{mix} = \delta q_{meas} - \delta q_{heat} - \delta q_{melt}, \quad (3)$$

where

$$\delta q_{heat} = \frac{m_{Tb}}{M_{Tb}} \left( \int_{T_a}^{T_{melt}} c_p^{sol}dT + \int_{T_{melt}}^{T_0} c_p^{liq}dT \right) \quad (4)$$

and

$$\delta q_{melt} = \frac{m_{Tb}}{M_{Tb}} \left[ H_{Tb}^{sol}(T_{melt}) - H_{Tb}^{sol}(T_{melt}) \right]. \quad (5)$$

Here, $M_{Tb}$ and $m_{Tb}$ denote the molar and total mass of the Tb specimen, respectively, and $\beta$ is the ISC calorimeter constant. The temperature dependent molar heat capacities ($c_p$) were determined from SGTE free energy parameterizations [79].

The ISC calorimeter constant, $\beta$, was determined using pure Al solid specimens and a pure Al bath. In this case, $\delta q_{mix} = 0$, and the left-hand side (LHS) of Equation (2) can computed from the heat capacity and enthalpy of melting for Al, which are explicitly known. The right-hand side (RHS) of Equation (2) is determined from the measured temperature signal, permitting quantitative determination of $\beta$ for the specific temperature and mass of the bath, using Equations (3)–(5). Calibration measurements used to determine the calorimeter constant are summarized in Table 2, and the constant was determined as $-0.08336$ J/sK (The reported uncertainty of ±5% for $\beta$ is based on 1.5 standard deviations, with respect to pure Al measurements, as described in Appendix A).

Immediately following the calibration measurement, $\Delta H_{mix}$ measurements were performed at $T_0 = 1364$ K by introducing solid Tb ($T_a = 298$ K) to the melt in incremental amounts of 1.8528 g, 3.4288 g, and 4.861 g, yielding alloys with mole fractions of Tb ($x_{Tb}$) of 0.030, 0.054, and 0.075, respectively. The results are summarized in Table 3. Incremental mixing enthalpy (relative to the previous state) is denoted as $\delta q_{mix}$ while the mixing enthalpy relative to the unmixed pure Al and Tb components is denoted as $\Delta H_{mix}$. Attempts to perform ISC measurements with higher Tb content resulted in non-negligible reaction with the ZrO$_2$ calorimeter crucible, which interfered with the thermal transient signal. Measured quantities are shown in Table 3 for these cases, but no enthalpy of mixing is reported.
Table 2. ISC calibration data.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T_a$</td>
<td>298 K</td>
</tr>
<tr>
<td>$T_0$</td>
<td>1364 K</td>
</tr>
<tr>
<td>$m_{\text{crucible}}$</td>
<td>9.4869 g</td>
</tr>
<tr>
<td>$m_{\text{Al}}$</td>
<td>0.7009 g</td>
</tr>
<tr>
<td>$M_{\text{Al}}$</td>
<td>26.98 g/mol</td>
</tr>
<tr>
<td>$M_{\text{Tb}}$</td>
<td>158.93 g/mol</td>
</tr>
<tr>
<td>$\delta q_{\text{heat}}$</td>
<td>826.05 J</td>
</tr>
<tr>
<td>$\delta q_{\text{melt}}$</td>
<td>278.26 J</td>
</tr>
<tr>
<td>$\delta q_{\text{meas}}/\beta$</td>
<td>$-13,247.6$ sK</td>
</tr>
<tr>
<td>$\beta$</td>
<td>$-0.08336 \pm 5%$ J/sK</td>
</tr>
</tbody>
</table>

Figure 2. A schematic representation of the ISC, showing the essential calorimetry and specimen handling features (primary heating and vacuum systems not shown).

Figure 3. A typical ISC trace showing the thermal transient upon Tb specimen injection.

Table 3. Incremental ISC mixing enthalpy measurements for $T_0 = 1364$ K.

<table>
<thead>
<tr>
<th>$m_{\text{Al}}$ (g)</th>
<th>$\delta m_{\text{Tb}}$ (g)</th>
<th>$m_{\text{Tb}}$ (g)</th>
<th>$x_{\text{Tb}}$</th>
<th>$\delta q_{\text{mix}}$ (J/mol)</th>
<th>$\Delta H_{\text{mix}}$ (J/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10.1878</td>
<td>1.8528</td>
<td>1.8528</td>
<td>0.03</td>
<td>$-4955.05$</td>
<td>$-4955.05$</td>
</tr>
<tr>
<td>-</td>
<td>1.5676</td>
<td>3.4288</td>
<td>0.05</td>
<td>$-3994.04$</td>
<td>$-8949.09$</td>
</tr>
<tr>
<td>-</td>
<td>1.4322</td>
<td>4.8610</td>
<td>0.08</td>
<td>$-2044.05$</td>
<td>$-10,993.1$</td>
</tr>
<tr>
<td>10.85</td>
<td>6.4362</td>
<td>6.4362</td>
<td>0.09</td>
<td>$-1074.16$</td>
<td>-</td>
</tr>
<tr>
<td>-</td>
<td>1.4510</td>
<td>7.8872</td>
<td>0.11</td>
<td>$-943.902$</td>
<td>-</td>
</tr>
</tbody>
</table>
For higher solute content, we turn to a container-less approach and employ electromagnetic levitation coupled with drop-calorimetry (EMLDC). The basic setup is shown in Figure 4. In this method, a stable levitation condition is established with a specimen of known mass under vacuum \((10^{-6} \text{ Torr})\) at a fixed temperature \(T_{\text{drop}}\). Temperature is controlled by induction heating with forced gas (ultra-high purity (99.999\%) He + 5\% H_2 gas, supplied by Matheson Company) (He + 5\% H_2) and measured using a non-contact infrared pyrometer. After a stable condition is reached at \(T_{\text{drop}}\), the coil is de-energized and the specimen falls by gravity into the calorimeter vessel. The calorimeter vessel is simply a solid copper cup with a conical bottom and a total mass of 167.0 g. The specimen temperature at the time of contact with the calorimeter \(T_{\text{con}}\) is computed by accounting for heat loss during the fall period (see Appendix B). The temperature transient in the calorimeter is measured using a thermocouple positioned just below the contact surface. A typical thermal trace is illustrated in Figure 5. The measured signal reflects rapid heat transfer from the specimen to the calorimeter upon contact and the evolution of heat during solidification and other phase changes. After the initial local heating, the measured signal reveals a slow cooling effect associated with heat redistribution within the calorimeter and heat loss from the calorimeter to its surroundings as it returns to temperature, \(T_0\).

![Figure 4](image1.png)

**Figure 4.** The levitation, heating, and calorimetry components of the EMLDC equipment.

![Figure 5](image2.png)

**Figure 5.** An example thermal trace, showing the characteristic temperatures (as listed in Table 4) for the EMLDC method.
The measured thermal exchange in the calorimeter is generally given as

$$\dot{q}_{\text{cal}} = \dot{q}_{\text{cool}} + \dot{q}_{\text{fr}} + \dot{q}_{\text{trans}} + \dot{q}_0,$$

where the first three terms on the right-hand-side give contributions from the dropped specimen, including cooling, freezing, and any other solid-state phase transitions that may occur, respectively. The last term is associated with thermal equilibration of the calorimeter, involving heat transfer within the calorimeter itself and thermal exchange with the surroundings. The thermal transient measured in the calorimeter includes all of these contributions, so the specimen contribution is given by

$$\delta q_{\text{spec}} = \delta q_{\text{cal}} - \delta q_0 = \phi \int_0^\infty (T - T_0) dt.$$  (7)

The enthalpy of mixing in the liquid can be determined as

$$q_{\text{mix}} = \delta q_{\text{spec}} - n \left( \sum f_p \Delta H_f - \sum x_i \delta q_i^T \right),$$  (8)

where \(n\) is the total number of moles (of atoms) in the specimen, \(\Delta H_f\) is the molar enthalpy of formation of a given phase at 298 K, \(f_p\) is the molar phase fraction of each phase, and the first term in parentheses is summed over all phases present in the quenched droplet. The second term in parentheses reflects the total molar enthalpy change for each elemental component, between the 298 K ground state and the liquid at \(T_{\text{con}}\), and it is summed over all components (Al and Tb in the present case), where \(x_i\) is the mole fraction of each component. Explicitly:

$$\delta q_{\text{Al}}^T = \int_{298}^{T_m} c_p^{\text{Al}, \text{fcc}} dT + \Delta H_m + \int_{T_m}^{T_{\text{con}}} c_p^{\text{Al}, \text{liq}} dT,$$

and

$$\delta q_{\text{Tb}}^T = \int_{298}^{T_m} c_p^{\text{Tb}, \eta} dT + \Delta H_m + \int_{T_m}^{T_{\text{con}}} c_p^{\text{Tb}, \text{liq}} dT.$$  (9)

Note that the expression in brackets in Equation (8) gives the total molar enthalpy difference between the quenched droplet at ambient temperature and the unmixed liquid at \(T = T_{\text{con}}\). The required formation enthalpy (for \(\text{Al}_3\text{Tb}\)) is computed using a DFT approach (see Appendix C).

For our experiments, the time scale for the measured local calorimeter heating upon specimen contact is much shorter than the time scale for cooling back to ambient conditions. Accordingly, we model the heating as instantaneous and locally adiabatic followed by Newtonian cooling with a characteristic exponential decay. We note that conditions are not adiabatic, but we define a local adiabatic equivalent temperature, \(T_A\), as the maximum temperature that would be reached in the limit of instantaneous adiabatic heating of a small local (i.e., effective) mass of the calorimeter upon specimen contact. The thermal transient measured upon contact with the non-adiabatic copper calorimeter is modeled as

$$T_0 + (T_A - T_0) \exp \left( -\frac{t}{\tau} \right),$$  (11)

where \(\tau\) is a calorimeter relaxation time, and \(T_A\) is a local adiabatic equivalent temperature corresponding to the limit of rapid heat extraction from the specimen. This temperature is treated as a fit parameter, along with \(\tau\). Equation (11) describes the measured behavior very closely, as shown by the root-mean-square-error (RMSE) values listed in Table 4, determined over the temperature range \(T < T_A - 0.25\) K. Once the values of \(\tau\) and \(T_A\) have been determined, the appropriate heat balance can be applied. Moreover, since \(q_{\text{cal}} = q_{\text{spec}}\) for this adiabatic heating, Equations (7)–(10) yield

$$q_{\text{spec}} = -\phi \int_{T_0}^{T_A} c_p^{\text{Cu}(s)} dT,$$  (12)
where the calorimeter constant, $\phi$, includes the effective calorimeter mass associated with non-uniform rapid heating upon contact, so that it includes the effects of specimen size and calorimeter size and shape.

The time-dependent heat transfer will be influenced by specimen size for a given calorimeter configuration (size, shape, insulation, thermocouple placement, etc.). Accordingly, determination of the calorimeter constant requires a standard specimen of the approximate size/mass of the specimen for which the enthalpy measurement is to be performed. In this work, two sets of calibration measurements were performed using pure Al specimens, with masses of 0.25 g and 0.5 g, nominally. Calibration measurements are listed along with applicable alloy test measurements in Table 4. Underlined values of the calorimeter constant ($\phi$) indicate an average of measured values determined from applicable calibration experiments listed. The stated specimen temperatures correspond to those shown in Figure 5. Pure Al specimens were used to determine $\phi$ for the two different nominal specimen sizes, where:

$$M_{\mathrm{Al}} \frac{q_{\text{meas}}}{m} = \int_{T_h}^{T_m} c_p \text{Al}(L) \, dT + \Delta H_m + \int_{T_m}^{T_{\infty}} c_p \text{Al}(s) \, dT \quad (13)$$

where $m$ is the specimen mass. We note that Equation (13) can be recovered from Equations (7)–(9) for a pure Al specimen, since $q_{\text{mix}}$ and $\Delta H_f$ both are equal to zero. Combining Equations (12) and (13), the calorimeter constant was determined using the pure Al samples, and the enthalpies of the alloy specimens were measured using the mean calorimeter constant determined from the calibration runs, as listed (type “cal”) in Table 4 (standard relative uncertainty of 1.3%; see Appendix A).

<table>
<thead>
<tr>
<th>#</th>
<th>Type</th>
<th>$x_{\text{TB}}$</th>
<th>Mass (g)</th>
<th>$T_{\text{drop}}$ (K)</th>
<th>$T_{\text{con}}$ (K)</th>
<th>$T_g$ (K)</th>
<th>$T_A$ (K)</th>
<th>$T_{\infty}$ (K)</th>
<th>RMSE (K)</th>
<th>$q_{\text{cal}}$ (J/mol)</th>
<th>$\phi$</th>
<th>$\Delta H_{\text{mix}}$ (kJ/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>cal</td>
<td>0</td>
<td>0.5206</td>
<td>1245.0</td>
<td>1241.8</td>
<td>292.3</td>
<td>304.4</td>
<td>294.9</td>
<td>0.013</td>
<td>744.4</td>
<td>2.692</td>
<td>-</td>
</tr>
<tr>
<td>2</td>
<td>cal</td>
<td>0</td>
<td>0.5190</td>
<td>1161.8</td>
<td>1159.1</td>
<td>298.0</td>
<td>309.0</td>
<td>294.7</td>
<td>0.004</td>
<td>684.9</td>
<td>2.710</td>
<td>-</td>
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<tr>
<td>3</td>
<td>cal</td>
<td>0</td>
<td>0.5190</td>
<td>1176.3</td>
<td>1173.5</td>
<td>298.2</td>
<td>309.7</td>
<td>295.2</td>
<td>0.031</td>
<td>697.6</td>
<td>2.657</td>
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<tr>
<td>4</td>
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<td>1078.9</td>
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<td>1297.7</td>
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<td>0.016</td>
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<tr>
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<td>310.1</td>
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<td>747.4</td>
<td>2.718</td>
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<tr>
<td>7</td>
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<td>293.7</td>
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<td>1426.7</td>
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<td>298.8</td>
<td>295.2</td>
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<td>-26.37</td>
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</tbody>
</table>

The EMLDC method described above requires determination of the thermodynamic state of the quenched specimen. This requires knowledge of the phases present in the quenched droplet, their relative amounts, and their formation energies. Based on the equilibrium phase diagram, we expect the quenched droplets to be comprised of the Al (fcc) and Al$_3$Tb ($\delta$) phases and estimate the expected molar fraction of the $\delta$ phase as 0.4 and 0.8 for alloy compositions ($x_{\text{TB}}$) of 0.1 and 0.2, respectively. We verified these estimates experimentally using X-ray diffraction (XRD) and scanning electron microscopy, as shown in Figure 6 (SEM analysis was done using a JEOL JSM-5910LV instrument. XRD was performed with a PANalytical X-Pert Pro diffractometer). Based on the XRD patterns shown in Figure 6a, the phases present in the quenched droplets were identified as Al (fcc) and Al$_3$Tb ($\delta$) for both compositions ($x_{\text{TB}}= 0.1, 0.2$). The corresponding microstructures are shown in Figure 6b,c, revealing primary $\delta$ phase (light) and a two-phase (fcc + $\delta$) eutectic constituent (dark). To measure the fraction of primary $\delta$, quantitative image analysis was performed using two images (1280 × 960 pixels) for each composition, with pixel sizes of 0.1 and 0.2 mm. Measurements yielding average $\delta$ fractions of 0.31 and 0.77 for $x_{\text{TB}}= 0.1$ and 0.2, respectively. Based on the eutectic composition
of $x_{Tb} = 0.03$ (see Figure 1), the two-phase eutectic structure is expected to include 12% δ and 88% Al (fcc). Accordingly, the total molar fraction of the δ phase was determined to be 0.396 and 0.797% for $x_{Tb} = 0.1$ and 0.2, respectively, in very good agreement with our original estimates of 0.4 and 0.8. Assuming that the formation energy at 0 K is equivalent to the formation enthalpy at 298 K, we computed the required enthalpy of formation for $\text{Al}_2\text{Tb}$ (δ) from first principles, obtaining a value of $-43.2 \text{ kJ/mol}$ (see Appendix C), relative to the pure Al and Tb reference states at 0 K [53,80].

$$
\Delta H_f^{\text{mix}} = \frac{1}{1 + \frac{2}{y_{\text{Al}_2\text{Tb}}} \left( \sum_i \left( \sum_j y_j \sum_{k=0}^n k_i y_k^{\delta} \right) + y_{\text{Al}_2\text{Tb}} \Delta H_f^{\text{Al}_2\text{Tb}} \right) \right)}
$$

where $x_i$ are the elemental mole fractions, and $y_i$ are mole fractions of associate species Al, Tb, and $\text{Al}_2\text{Tb}$. The model parameters determined on the basis of our measurements are listed in Table 5. The resulting model descriptions of the mixing enthalpy are plotted in Figure 7 for $T = 1364$ K and compared with measured values. Corresponding model descriptions of chemical activity and partial molar mixing enthalpy ($\Delta H_{\text{mix}}^{\text{mix}}$) are also plotted. In addition, the Miedema model prediction [76–78] is included in each figure for comparison.

It is significant to point out that the experimental data show very good agreement with the Miedema [76–78] prediction, which involves no adjustable parameters and serves as a baseline model. These figures also illustrate clearly that all three of the CALPHAD formulations (with the parameters listed in Table 5) provide similar descriptions of $\Delta H_{\text{mix}}^{\text{mix}}$ for this alloy over the dilute regime. Comparison of the modeled chemical activities suggests relatively small differences in the non-ideality for Al-rich compositions. This behavior is isolated more clearly in the partial molar mixing enthalpy, plotted in Figure 7c, which shows the different values of ($\Delta H_{\text{mix}}^{\text{mix}}$) in the limit as $x_{Tb} \rightarrow 0$. Generally, Figure 7 shows that multiple model formulations can provide reasonable descriptions of the measured behavior. Further discrimination will require experimental measurements in intermediate and Tb-rich compositional regimes. Considering the challenges related to crucible reactions, this calls for additional container-less experiments.

![Figure 6](image_url)

**Figure 6.** (a) XRD patterns indicating the fcc + δ constitution of the quenched alloys, with $f_δ = 0.4$ and 0.8 for $x_{Tb} = 0.1$ and 0.2, respectively (XRD specimens: sectioned and diamond-polished to 0.6 μm finish); (b,c) SEM images of EMLDC specimens showing the δ phase (light) and a two-phase eutectic constituent (dark) for alloy compositions of $x_{Tb} = 0.1$ and 0.2 (backscattered electron contrast, 20 kV accelerating voltage, 10 mm working distance).

### 3. Analysis and Discussion

To establish additional thermodynamic context for our enthalpy of mixing measurements for the Al–Tb liquid phase, we now consider appropriate models and specifically examine the application of three different semi-empirical treatments, all arising from the general model

There are no tables or figures mentioned in the raw text. The complete page content is provided for further analysis.
Table 5. Parameters for thermodynamic models of $\Delta H_{mix}^{Liq}$.

<table>
<thead>
<tr>
<th></th>
<th>Regular</th>
<th>Subregular</th>
<th>Associate</th>
</tr>
</thead>
<tbody>
<tr>
<td>$i$</td>
<td>$j$</td>
<td>$i$</td>
<td>$j$</td>
</tr>
<tr>
<td>$L_{ij}^0$ (J/mol)</td>
<td>$-160,000$</td>
<td>$-128,000 + 29.991T$</td>
<td>$-75,252$</td>
</tr>
<tr>
<td>$L_{ij}^1$ (J/mol)</td>
<td>$-$</td>
<td>$-80,455 + 30.998T$</td>
<td>$-$</td>
</tr>
<tr>
<td>$L_{ij}^2$ (J/mol)</td>
<td>$-$</td>
<td>$30,342$</td>
<td>$-$</td>
</tr>
<tr>
<td>$\Delta G_{Al_{2}Tb}$ (J/mol)</td>
<td>$-$</td>
<td>$-$</td>
<td>$-113,233 + 18.904T$</td>
</tr>
</tbody>
</table>

Notes

$\begin{align*}
  y_i &= x_i \\
  y_i &= \frac{x_i}{y_{Al_{2}Tb} = 0} \\
  \Delta G_{Al_{2}Tb} &= -RT\ln\left(\frac{x_{Al_{2}Tb}}{x_{Al_{2}Tb}}\right)
\end{align*}$

Figure 7. The liquid models: (a) mixing enthalpy with measured values; (b) chemical activity; and (c) partial molar enthalpy of mixing, calculated for $T = 1364$ K (all referenced to pure Al and Tb liquid states).

4. Conclusions

We report here measurements of the enthalpy of mixing in the liquid phase for Al–Tb alloys, for Tb mole fractions up to $x_{Tb} = 0.2$. Specifically, we utilize isothermal solution calorimetry and isoperibolic electromagnetic levitation drop calorimetry to measure the enthalpy of mixing, employing
X-ray diffraction, scanning electron microscopy, and first principles calculations to provide necessary quantification of the quenched states and component reference states. Based on our measurements, we offer three different assessed semi-empirical solution models for the excess free energy of the liquid, including regular, subregular, and associate model formulations for the enthalpy of mixing. All three of these model formulations provide very good representation of the measured behavior over the limited range examined. Our results highlight the need for additional measurements targeting the composition range of $0.5 < x_{Tb} < 0.7$, which would enable further refinement of the general model.

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Author Contributions: Shihuai Zhou designed the ISC experiments. Ralph Napolitano and Carl Tackes designed the EMLDC experiments. Shihuai Zhou and Carl Tackes performed the experiments; Shihuai Zhou and Ralph Napolitano analyzed the data; Shihuai Zhou and Ralph Napolitano wrote the paper. All authors have read and approved the final manuscript.

Conflicts of Interest: The authors declare no conflict of interest.

Appendix A

The uncertainties for the calibration constants, $\beta$ and $\phi$ for ISC and EMLDC, respectively, were determined through a series of measurements using pure aluminum (0.9999) standards. For ISC, the measurements were performed with the specimen under argon at atmospheric pressure and a constant temperature of 1411 K. By successive incremental additions of aluminum (at ambient temperature, 298 K) in small amounts (~0.5 g) to a liquid Al bath held in an Al$_2$O$_3$ crucible, ISC measurements of the thermal transient yielded a series of independent measurements of $\beta$, with a relative standard deviation of 3.4%, shown in Table A1. Similarly, a set of six measurements were performed to determine the EMLDC calibration constant ($\phi$), using pure aluminum specimens, nominally 0.5 g each (also shown in Table 5). The relative standard deviation was determined to be 0.85%, as shown in Table A1. Based on these measurements and taking the standard uncertainties as 1.5 standard deviations as measured in the calibration experiments, we estimate the standard uncertainties for the ISC and EMLDC methods as 5% and 1.3%, respectively.

<table>
<thead>
<tr>
<th>Incremental Mass, Al (g)</th>
<th>Cumulative Mass, Al (g)</th>
<th>Calibration Const. ($\beta$)</th>
<th>Total Mass, Al (g)</th>
<th>Calibration Const. ($\phi$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.5206</td>
<td>2.692</td>
</tr>
<tr>
<td>0.06331</td>
<td>0.10152</td>
<td>-0.33261</td>
<td>0.5190</td>
<td>2.657</td>
</tr>
<tr>
<td>0.04129</td>
<td>0.16483</td>
<td>-0.32971</td>
<td>0.5133</td>
<td>2.693</td>
</tr>
<tr>
<td>0.04251</td>
<td>0.20612</td>
<td>-0.31686</td>
<td>0.5173</td>
<td>2.717</td>
</tr>
<tr>
<td>0.06162</td>
<td>0.31025</td>
<td>-0.32572</td>
<td>0.5168</td>
<td>2.718</td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.5168</td>
<td>2.718</td>
</tr>
</tbody>
</table>

Table A1. Summary of experiments for determination of the standard uncertainty for ISC and EMLDC.

For the EMLDC method, specimen cooling during the drop period was computed to determine the specimen temperature upon contact with the calorimeter:

$$T_{con} = T_{drop} - \Delta T_{drop} = T_{drop} - \Delta T_C - \Delta T_R$$
where \( \Delta T_C \) and \( \Delta T_R \) describe convective and radiative cooling during freefall. Using the parameters defined in Table A2, these were computed, respectively, as

\[
\Delta T_C = B_0 h (T_{\text{drop}} - T_0) \quad \text{and} \\
\Delta T_R = B_0 \sigma \epsilon (T_A^4 - T_0^4),
\]

where for the spherical drop:

\[
B_0 = A_s t_f W / m c_p \\
A_S = (36 \pi V^2)^{1/3} \\
t_f = \sqrt{2z / g} \\
h = \frac{k}{D} \left[ 2 + \left( 0.4 \text{Re}^{1/2} + 0.6 \text{Re}^{2/3} \right) \text{Pr}^{2/5} \right],
\]

and the Prandtl and Reynolds numbers were taken, respectively, as

\[
Pr = \mu c_p / k \\
Re = \rho V_\infty D / \mu.
\]

Table A2. Parameters used in the calculation of \( T_{\text{con}} \).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \sigma )</td>
<td>Stephan-Boltzmann constant (taken here as ( 5.67 \times 10^{-8} ))</td>
<td>J/m²K⁴</td>
</tr>
<tr>
<td>( \epsilon )</td>
<td>Total hemispherical emissivity (taken here as 0.084)</td>
<td>-</td>
</tr>
<tr>
<td>( c_p )</td>
<td>Specimen heat capacity</td>
<td>J/mol</td>
</tr>
<tr>
<td>( m )</td>
<td>Mass of specimen</td>
<td>g</td>
</tr>
<tr>
<td>( W )</td>
<td>Specimen molar mass</td>
<td>g/mol</td>
</tr>
<tr>
<td>( A_S )</td>
<td>Surface area of specimen</td>
<td>m²</td>
</tr>
<tr>
<td>( V )</td>
<td>Volume of specimen</td>
<td>m³</td>
</tr>
<tr>
<td>( t_f )</td>
<td>Fall duration</td>
<td>s</td>
</tr>
<tr>
<td>( h )</td>
<td>Heat transfer coefficient</td>
<td>W/m²K</td>
</tr>
<tr>
<td>( \mu )</td>
<td>Dynamic viscosity (gas) (taken here as ( 1.90 \times 10^{-5} ))</td>
<td>Ns/m²</td>
</tr>
<tr>
<td>( k )</td>
<td>Thermal conductivity (gas) (taken here as ( 0.1513 ))</td>
<td>W/mK</td>
</tr>
<tr>
<td>( D )</td>
<td>Specimen diameter</td>
<td>m</td>
</tr>
<tr>
<td>( V_\infty )</td>
<td>Fluid velocity (taken as the average fall velocity)</td>
<td>m/s</td>
</tr>
<tr>
<td>( \rho )</td>
<td>Specimen density</td>
<td>Kg/m³</td>
</tr>
</tbody>
</table>

Appendix C

The total energy \( E^\theta \) is computed for each phase using the Vienna ab initio simulation package (VASP) [81–83] employing projector augmented wave method [83,84] and Perdew–Burke–Ernzerhof generalized gradient approximation (PBE-GGA) [85]. The “high precision” choice to ensure that the computed absolute energies were converged to a few meV. The spin-polarized electronic-structure calculations were enabled and the Monkhost 24 × 24 × 20 “k-points” were employed for high precision. For each compound, the cell was fully relaxed with respect to distortion and dilatation, and it was confirmed the crystal structure was maintained. The enthalpy of formation for the a given compound is calculated as the difference between the energy \( E^\theta \) of the compound and the linear combination of the pure element reference state energies, \( E_{Al}^{\text{fcc}} \) and \( E_{Tb}^{\text{hcp}} \), so that \( \Delta H_f^\theta \) can be calculated as

\[
\Delta H_f^\theta = E^\theta - x_{Al} E_{Al}^{\text{fcc}} - x_{Tb} E_{Tb}^{\text{hcp}}.
\]
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