

5-8-2007

Modeling of Thermodynamic Properties and Phase Equilibria for the Cu-Mg Binary System

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Abstract

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Keywords

Ames Laboratory, Materials and Engineering Physics

Disciplines

Metallurgy

Comments

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(Submitted November 2, 2006)

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Keywords thermodynamics, phase diagram, phase equilibria

1. Introduction

Given the promising mechanical, chemical, and magnetic properties of various metallic glasses and the potential widespread use of the so-called “bulk metallic glasses”,^[1–4] much recent attention has been given to the development of our general understanding of the thermodynamics and kinetics for glass formation and glass forming ability.^[5] While alloys exhibiting promising glass formation tendency are frequently ternary, quaternary, or higher order systems,^[1–4] several binary glass-forming metallic alloys have been identified. Of these, the Al-based alloys have been investigated most thoroughly.^[6] In addition, glass formation has been identified in several other binary systems,^[7–12] most notably including Cu-Mg,^[7] Pd-Si,^[8,9] and Cu-Zr.^[8] By virtue of their simplicity compared with their many-component counterparts, these binary alloys present us with an opportunity to investigate the detailed relationship between the thermodynamic descriptions of relevant phase

equilibria and observed glass-forming behavior. It is in this vein that the present thermodynamic analysis of the Cu-Mg binary system is motivated. In a subsequent paper, we will model the thermodynamic properties of the undercooled liquid and examine more closely the implications regarding glass formation tendency, with respect to the competing crystalline phases.

Thermodynamic models for the binary Cu-Mg system have been offered by Coughanowr et al.^[13] and by Zou and Chang.^[14] While these treatments are well posed and have resulted in phase diagrams that agree well with experimental reports, there are three specific features of the modeling approaches that limit their potential applicability to more general phase stability problems. First, in each of these treatments, the liquid phase is modeled as a regular solution, a model that cannot describe the chemical ordering observed in this system. For example, X-ray diffraction experiments reported by Lukens and Wagner^[15] indicate the existence of chemical short range order in the liquid phase with Cu_2Mg and CuMg_2 stoichiometries. This discrepancy is clearly observed in the inability of these models to accurately describe the heat capacity of the undercooled liquid, as shown in Fig. 1. Second, these previous models treat the $\text{Mg}_2\text{Cu-C}_b$ as a stoichiometric compound, precluding any description of nonequilibrium compositions for these phases. Finally, the models constrain the temperature dependence of the Gibbs free energy of this phase to be linear, implying that there is no contribution to the heat capacity from chemical mixing. Consequently, while the equilibrium phase diagram produced from these models may be useful, more fundamental thermodynamic quantities associated with these models, such as the heat capacity, do not agree well with experimental measurements, such as those reported by Feufel and Sommer.^[16]

In the work presented here, we address each of these limitations within the framework of a general CAL-

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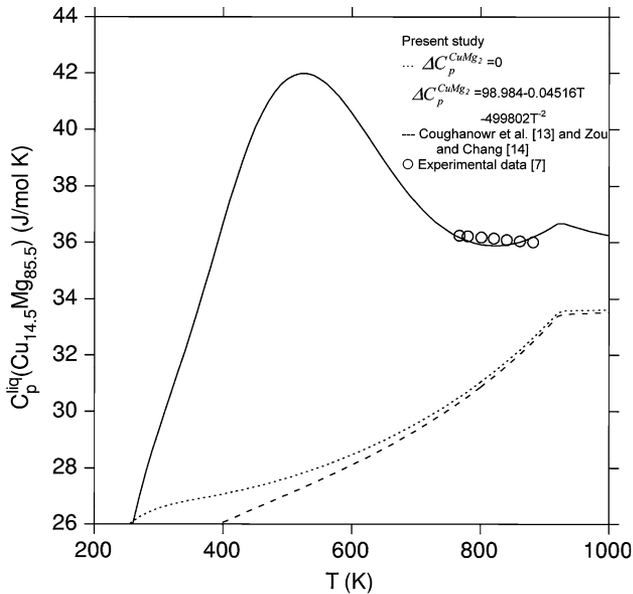


Fig. 1 The heat capacity of the liquid $\text{Cu}_{14.5}\text{Mg}_{85.5}$ alloy calculated using the parameters in Tables 2-4

PHAD^[17,18] formulation by (i) employing an association model capable of describing the nonlinear temperature dependence of chemical short range order in the liquid phase and (ii) treating the intermetallic phases as substitutional solid solutions on two sublattices, permitting quantification of the Gibbs free energies of the intermetallic phases at non-stoichiometric compositions. In addition, we incorporate first-principles calculations to compute the zero-Kelvin energies of end-member phases in unstable structures, since experimental data are not available for these phases. Remaining model parameters, describing the Gibbs free energies associated with the formation of compounds or solution phases, are determined through a systematic semi-empirical optimization, employing available experimental data from calorimetry, X-ray diffraction, electron-probe microchemical analysis, and optical micrography.^[7,16,19-40] The resulting thermodynamic properties and the associated equilibrium phase diagram are compared with the prior thermodynamic modeling reported by Coughanowr et al.^[13] and by Zou and Chang.^[14]

2. Thermodynamic Models

We describe the phase equilibria in the Cu-Mg binary system by modeling the Gibbs free energy for each relevant phase over the appropriate range of composition at constant pressure (1 atm). In any case where a temperature dependent parameter (P) is required, we use the general form,

$$P(T) = a + bT + cT \ln T + dT^2 + eT^{-1} + fT^3 + gT^7 + hT^{-9} \quad (\text{Eq 1})$$

The thermodynamic properties of pure Cu and Mg in various structures are computed using the parameters from the SGTE database.^[41] The liquid phase is described with an association model.^[42,43] The terminal fcc and hcp solid solutions are treated with a single lattice while intermediate phases are described using a two-sublattice model,^[44] where each sublattice is treated as a regular solution. In the following sections, the thermodynamic treatment of each phase is described in detail.

2.1 The Liquid Phase

We employ a four-species association model,^[42,43] where Cu_2Mg and CuMg_2 are chosen as the relevant intermediate chemical associates.^[15] Thus, the Gibbs free energy of the liquid phase is given by

$$G_m^{\text{liq}} = \sum_i x_i \circ G_i^{\text{liq}} + RT \sum_i x_i \ln x_i + \sum_i \sum_{j>i} x_i x_j \circ L_{ij}^{\text{liq}}, \quad (i, j = \text{Cu}, \text{Cu}_2\text{Mg}, \text{CuMg}_2, \text{Mg}) \quad (\text{Eq 2})$$

where x_i and $\circ G_i^{\text{liq}}$ denote the mole fraction and molar Gibbs free energy for the i th species in the liquid phase, respectively, and where we have considered only pair-wise interaction between species. The Gibbs free energy of the intermediate associates, Cu_mMg_n , are given as

$$\circ G_{\text{Cu}_m\text{Mg}_n}^{\text{liq}} = m \circ G_{\text{Cu}}^{\text{liq}} + n \circ G_{\text{Mg}}^{\text{liq}} + \Delta G_{\text{Cu}_m\text{Mg}_n}^{\text{liq}}, \quad (\text{Eq 3})$$

where $\Delta G_{\text{Cu}_m\text{Mg}_n}^{\text{liq}}$ represents the Gibbs free energy of formation. Substituting Eq 3 into Eq 2, the Gibbs free energy of the liquid phase is written as

$$\begin{aligned} G_m^{\text{liq}} = & (x_{\text{Cu}} + 2x_{\text{Cu}_2\text{Mg}} + x_{\text{CuMg}_2}) \circ G_{\text{Cu}}^{\text{liq}} \\ & + (x_{\text{Mg}} + x_{\text{Cu}_2\text{Mg}} + 2x_{\text{CuMg}_2}) \circ G_{\text{Mg}}^{\text{liq}} \\ & + x_{\text{Cu}_2\text{Mg}} \Delta G_{\text{Cu}_2\text{Mg}}^{\text{liq}} + x_{\text{CuMg}_2} \Delta G_{\text{CuMg}_2}^{\text{liq}} \\ & + RT(x_{\text{Cu}} \ln x_{\text{Cu}} + x_{\text{Cu}_2\text{Mg}} \ln x_{\text{Cu}_2\text{Mg}} + x_{\text{CuMg}_2} \ln x_{\text{CuMg}_2} \\ & + x_{\text{Mg}} \ln x_{\text{Mg}}) + x_{\text{Cu}} x_{\text{Cu}_2\text{Mg}} \circ L_{\text{Cu}, \text{Cu}_2\text{Mg}}^{\text{liq}} + x_{\text{Cu}} x_{\text{CuMg}_2} \circ L_{\text{Cu}, \text{CuMg}_2}^{\text{liq}} \\ & + x_{\text{Cu}} x_{\text{Mg}} \circ L_{\text{Cu}, \text{Mg}}^{\text{liq}} + x_{\text{CuMg}_2} x_{\text{Cu}_2\text{Mg}} \circ L_{\text{CuMg}_2, \text{Cu}_2\text{Mg}}^{\text{liq}} \\ & + x_{\text{Mg}} x_{\text{Cu}_2\text{Mg}} \circ L_{\text{Cu}_2\text{Mg}, \text{Mg}}^{\text{liq}} + x_{\text{Mg}} x_{\text{CuMg}_2} \circ L_{\text{CuMg}_2, \text{Mg}}^{\text{liq}} \end{aligned} \quad (\text{Eq 4})$$

The interaction energies ($\circ L_{ij}^{\text{liq}}$) are described here as constants, while the Gibbs free energies of formation ($\Delta G_{\text{Cu}_m\text{Mg}_n}^{\text{liq}}$) are described as functions of temperature, using the form given in Eq 1. Where sufficient heat capacity data are available, the parameters a – e are used. Otherwise, we employ only parameters a and b . This will be discussed in further detail in a subsequent section.

2.2 Terminal fcc and hcp Phases

The fcc and hcp phases are treated as simple binary substitutional solutions with Gibbs free energies expressed as

$$G_m^{\Phi} = \sum_{i=\text{Cu}, \text{Mg}} x_i \circ G_i^{\Phi} + RT \sum_{i=\text{Cu}, \text{Mg}} x_i \ln x_i + x_s G_m^{\Phi} \quad (\text{Eq 5})$$

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where x_i denotes mole fraction of element i , and ${}^\circ G_i^\Phi$ ($i = \text{Cu, Mg}$) denotes the molar Gibbs free energy of the pure element i with the structure Φ ($\Phi = \text{fcc or hcp}$). The excess Gibbs free energy ${}^{xs}G_m^\Phi$ is expressed as,

$${}^{xs}G_m^\Phi = x_{\text{Cu}}x_{\text{Mg}} \sum_{j=0}^n {}^jL_{\text{Cu,Mg}}^\Phi (x_{\text{Cu}} - x_{\text{Mg}})^j \quad (\text{Eq 6})$$

where the ${}^jL_{\text{Cu,Mg}}^\Phi$ coefficients are left as interaction parameters to be evaluated with experimental data. Here, we consider only the $j = 0$ term and assume that ${}^\circ L_{\text{Cu,Mg}}^\Phi$ is constant (i.e., a regular solution).

2.3 Intermediate $\text{Cu}_2\text{Mg-C15}$ and $\text{Mg}_2\text{Cu-C}_b$ Phases

Using a two-sublattice model, we describe each intermediate compound as a solid solution of the form $(\text{Cu,Mg})_2(\text{Cu,Mg})_1$. The Gibbs free energy is given as

$$G_m^\theta = \sum_{i=\text{Cu,Mg}} y_i^I \sum_{j=\text{Cu,Mg}} y_j^{II} {}^\circ G_{ij}^\theta + RT \sum_{i=\text{Cu,Mg}} (2y_i^I \ln y_i^I + y_i^I \ln y_i^{II}) + {}^{xs}G_m^\theta \quad (\text{Eq 7})$$

where the colon separates the components on different sublattices, and y^I and y^{II} are the sublattice site occupancy fractions. The superscript, θ , represents the $\text{Cu}_2\text{Mg-C15}$ or $\text{Mg}_2\text{Cu-C}_b$ structure.

With the sublattice description we have adopted, each of the two structures (C15 and C_b) can assume four different stoichiometries (Cu_2Cu , Mg_2Cu , Cu_2Mg , and Mg_2Mg), with only one being that of the stable compound. These are $\text{Cu}_2\text{Mg-C15}$ and $\text{Mg}_2\text{Cu-C}_b$, and, for these compounds, we express ${}^\circ G_{\text{Cu:Mg}}^{\text{C15}}$ and ${}^\circ G_{\text{Mg:Cu}}^{\text{C}_b}$ using the form given in Eq 1 and evaluate the related coefficients using available enthalpy of formation, melting temperature, and heat capacity data, as discussed in a subsequent section. For the unstable compounds (i.e., $\text{Cu}_2\text{Cu-C15}$, $\text{Mg}_2\text{Cu-C15}$, $\text{Mg}_2\text{Mg-C15}$, $\text{Cu}_2\text{Cu-C}_b$, $\text{Cu}_2\text{Mg-C}_b$, and $\text{Mg}_2\text{Mg-C}_b$), there are no available heat capacity data with which to evaluate the coefficients c-g. Therefore, we express the Gibbs free energy as

$${}^\circ G_{ij}^\theta = 2 {}^\circ G_i^{\text{ref}} + {}^\circ G_j^{\text{ref}} + \Delta G_{ij}^\theta, \quad (i, j = \text{Cu, Mg}) \quad (\text{Eq 8})$$

where ${}^\circ G_i^{\text{ref}}$ and ${}^\circ G_j^{\text{ref}}$ are the molar Gibbs free energy of either fcc-Cu or hcp-Mg, and ΔG_{ij}^θ is the Gibbs free energy of formation for the compound $(i)_2(j)_1$ from the $2(i) + (j)$ mixture, which we treat as a constant. The excess Gibbs free energy term in Eq 7 is modeled as,

$${}^{xs}G_m^\theta = y_{\text{Cu}}^I y_{\text{Mg}}^I \sum_{i=\text{Cu,Mg}} y_i^{II} \sum_{k=0}^k {}^kL_{\text{Cu,Mg};i}^\theta (y_{\text{Cu}}^I - y_{\text{Mg}}^I)^k + y_{\text{Cu}}^{II} y_{\text{Mg}}^{II} \sum_{i=\text{Cu,Mg}} y_i^I \sum_{k=0}^k {}^kL_{i;\text{Cu,Mg}}^\theta (y_{\text{Cu}}^{II} - y_{\text{Mg}}^{II})^k \quad (\text{Eq 9})$$

Once again, we consider only the $k = 0$ term in each inner sum, and assume that ${}^\circ L^\theta$ is a constant.

2.4 Calculation of Zero-Kelvin Enthalpies from First Principles

To facilitate the determination of the Gibbs free energy of formation for the intermediate compound phases, we compute the enthalpy of formation for the end-members at zero Kelvin. For these calculations, we employ the VASP^[45] implementation of the plane wave method using the Vanderbilt ultrasoft pseudopotential^[46] with a generalized gradient approximation (GGA).^[47] The Monkhost $15 \times 15 \times 15$ k points are employed for high precision calculations. The $3s3p$ and $3d4s4p$ shells are treated as valence states with core radii of 2.88 a.u. and 2.48 a.u. for the hcp-Mg and fcc-Cu, respectively. To ensure that the unit cell corresponds to a stable structure, we fully relax the cell shape and the internal atomic coordinates of the stable end-members of the compounds and relax only the cell volumes of those end-members which are unstable.

The enthalpy of formation, ΔH_f^ϕ , of a compound ϕ is calculated as the difference between the energy of the compound and the linear combination of the energies of the pure elements in their reference states,

$$\Delta H_f^\phi = E^\phi - x_{\text{Cu}}^\phi E_{\text{Cu}}^{\text{fcc}} - x_{\text{Mg}}^\phi E_{\text{Mg}}^{\text{hcp}}, \quad (\text{Eq 10})$$

where x_i^ϕ is the mole fraction of component i in the ϕ structure. The values of E^ϕ , $E_{\text{Cu}}^{\text{fcc}}$, and $E_{\text{Mg}}^{\text{hcp}}$ are the computed zero-Kelvin energies of the indicated phases, each considered here to be stoichiometric. The calculated results are listed in Table 1 and compared with experimental data. Again, we note here that for the stable phases ($\text{Cu}_2\text{Mg-C15}$ and $\text{Mg}_2\text{Cu-C}_b$), both experimental and first-principles data are considered in our evaluation of thermodynamic parameters. For the unstable intermetallic phases ($\Delta H_f > 0$), however, experimental data are not attainable and the zero-Kelvin energies become essential in assessing the relative stability of these compounds. The determination of model parameters is discussed in the next section.

3. Determination of the Thermodynamic Model Parameters

Expressions for the standard Gibbs free energy (${}^\circ G_i^\phi$) of each pure component in the relevant phases are taken from the SGTE database^[41] as listed in Table 2. In addition, based on the models described in the preceding sections, we compute a standard Gibbs free energy for each of the two intermediate compounds and evaluate the parameters, as listed in Table 3. For the relevant excess Gibbs free energies, we employ a total of eight Gibbs free energy of formation terms and 12 interaction parameters (four of which we assume to be equal to zero). These are listed in Table 4, along with the results from our parameter evaluation. In the present section, the methodology used for determination of the parameters listed in Tables 3 and 4 is discussed.

Table 1 A summary of the results from the first-principles calculations

Phase	Formula	ΔH , kJ/mol of atoms		
		First principles	Modeling	Experiment
		0 K	298 K	
Cu-fcc	Cu	0
Mg-hcp	Mg	0
Cu ₂ Mg-C ₁₅	Cu ₂ Cu	15.5	15.5	...
	Cu ₂ Mg	-15.72	-11.4	-11.3 ^[36] -8.04 ^[37]
	CuMg ₂	34.72	34.72	...
	Mg ₂ Mg	7.00	7.00	...
CuMg ₂ -C _b	Cu ₂ Cu	20.39	20.39	...
	Cu ₂ Mg	38.94	38.94	...
	CuMg ₂	-13.20	-9.6	-9.55 ^[36]
	Mg ₂ Mg	12.92	12.92	...

Table 2 The thermodynamic parameters for pure Cu and Mg^[41]

Cu phases		${}^\circ G_{Cu}^{liq}$		${}^\circ G_{Cu}^{fcc}$	${}^\circ G_{Cu}^{hcp}$
T_{min} , K	298	1357.77	298	1357.77	298
T_{max} , K	1357.77	3200	1357.77	3200	3200
${}^\circ G^{ref}$	${}^\circ G_{Cu}^{fcc}$	0	0	0	${}^\circ G_{Cu}^{fcc}$
a	12964.735	-46.545	-7770.458	-13542.026	600
b	-9.511904	173.881484	130.485235	183.803828	0.2
c	...	-31.38	-24.112392	- 31.38	...
d	-2.65684×10^{-3}
e	52478
f	1.29223×10^{-7}
g	-5.8489×10^{-21}
h	3.64167×10^{29}	...
Mg phases		${}^\circ G_{Mg}^{liq}$	${}^\circ G_{Mg}^{fcc}$		${}^\circ G_{Mg}^{hcp}$
T_{min}	298	923	298	298	923
T_{max}	923	3000	3000	923	3000
${}^\circ G^{ref}$	${}^\circ G_{Mg}^{hcp}$	0	${}^\circ G_{Mg}^{hcp}$	0	0
a	8202.243	-5439.869	2600	-8367.34	-14130.185
b	-8.83693	195.324057	-0.9	143.675547	204.716215
c	...	-34.3088	...	-26.1849782	-34.3088
d	4.858×10^{-4}	...
e	78950	...
f	-1.393669×10^{-6}	...
g	-8.0176×10^{-20}
h	1.038192×10^{28}

Note: ${}^\circ G_i^l = {}^\circ G^{ref} + a + bT + cT \ln T + dT^2 + eT^{-1} + fT^3 + gT^7 + hT^{-9}$ (J/mol)

For the liquid phase, we evaluate the parameters $\Delta G_{Cu_mMg_n}^{liq}$ and ${}^\circ L_{ij}^{liq}$, from Eq 2-3, using reported values of activity,^[33] chemical potential,^[30-35] and enthalpy of mixing.^[29,30] Initially we model $\Delta G_{Cu_mMg_n}^{liq}$ using only the first two terms in Eq 1 (i.e., $a + bT$). This treatment, shown in Fig. 1, yields a heat capacity for Cu_{1.5}Mg_{84.5} liquid that is slightly higher than the models of Coughanowr et al.^[13] and Zou and Chang,^[14] but one that remains much lower than the reported experimental data.^[7] We assert that this

apparent excess heat capacity may be due to chemical (and perhaps structural) ordering in the liquid phase over this range of temperatures. Accordingly, we describe the associated nonlinear temperature dependence by fitting the parameters c-e in Eq 1 for $\Delta G_{CuMg_2}^{liq}$. The evaluated results are listed in Table 4 and the corresponding heat capacity is also plotted in Fig. 1, where a maximum in the $C_p(T)$ curve is exhibited. We note that such maxima in $C_p^{liq}(T)$ have been observed in a number of systems (e.g., toluene, Au-Si,

Table 3 Ground state standard Gibbs free energy parameters for the intermetallic phases $\text{Cu}_2\text{Mg-C15}$ and $\text{Mg}_2\text{Cu-C}_b$

$^{\circ}G_{ij}^0$	$^{\circ}G_{\text{Cu:Mg}}^{\text{C15}}, \text{J/mol}$		$^{\circ}G_{\text{Mg:Cu}}^{\text{Cb}}, \text{J/mol}$	
T_{min}, K	298	1070	298	850
T_{max}, K	1070	...	850	...
$^{\circ}G^{\text{ref}}$	0	0	0	0
a	-58201	-64662.18	-53491	-58610.6334
b	409.642	490.7029614	425.428	481.9243846
c	-76.1	-87.17102875	-77.9913484	-85.33353573
d	-9.9×10^{-4}	...	2.31×10^{-3}	...
e	183906	...	190378	...
f	-1.35×10^{-6}	...	-2.72115×10^{-6}	...
g
h

Table 4 Excess Gibbs free energy parameters (all temperatures)

Phase	Parameters	Value, J/mol
Liquid	$\Delta G_{\text{Cu}_2\text{Mg}}^{\text{liq}}$	$-28312 + 9.595 T$
	$\Delta G_{\text{CuMg}_2}^{\text{liq}}$	$-108077 + 748.301 T - 98.984 T \ln(T) + 0.02258 T^2 + 2499901 T^{-1}$
	$^{\circ}L_{\text{Cu,Cu}_2\text{Mg}}^{\text{liq}}$	-20012
	$^{\circ}L_{\text{Cu,CuMg}_2}^{\text{liq}}$	-24230
	$^{\circ}L_{\text{Cu,Mg}}^{\text{liq}}$	-22611
	$^{\circ}L_{\text{Cu}_2\text{Mg,CuMg}_2}^{\text{liq}}$	0
	$^{\circ}L_{\text{Cu}_2\text{Mg,Mg}}^{\text{liq}}$	-25845
hcp	$^{\circ}L_{\text{CuMg}_2,\text{Mg}}^{\text{hcp}}$	0
	$^{\circ}L_{\text{Cu,Mg:Va}}^{\text{hcp}}$	39230
fcc	$^{\circ}L_{\text{Cu,Mg:Va}}^{\text{fcc}}$	-19345
$\text{CuMg}_2\text{-C15}$	$\Delta G_{\text{Cu:Cu}}^{\text{C15}}$	46500
	$\Delta G_{\text{Mg:Cu}}^{\text{C15}}$	104160
	$\Delta G_{\text{Mg:Mg}}^{\text{C15}}$	21000
	$^{\circ}L_{\text{Cu,Cu,Mg}}^{\text{C15}}$	-27868
	$^{\circ}L_{\text{Cu,Mg:Mg}}^{\text{C15}}$	3521
$\text{CuMg}_2\text{-C}_b$	$\Delta G_{\text{Cu:Cu}}^{\text{Cb}}$	61170
	$\Delta G_{\text{Cu:Mg}}^{\text{Cb}}$	116820
	$\Delta G_{\text{Mg:Mg}}^{\text{Cb}}$	38760
	$^{\circ}L_{\text{Cu,Cu,Mg}}^{\text{Cb}}$	0
	$^{\circ}L_{\text{Cu,Mg:Mg}}^{\text{Cb}}$	0
	$^{\circ}L_{\text{Cu,Mg:Mg}}^{\text{Cb}}$	0

As_2Se_3 ,^[48] $\text{La}_{20}\text{Mg}_{50}\text{Ni}_{30}$, $\text{Al}_{30}\text{La}_{50}\text{Ni}_{20}$, $\text{Al}_{7.5}\text{Cu}_{17.5}\text{Ni}_{10}\text{Zr}_{65}$,^[49] $\text{Al}_{7.5}\text{Cu}_{27.5}\text{Zr}_{65}$,^[50]) again suggesting some type of clustering/ordering reaction in the liquid phase.

The interaction parameter for each of the terminal solid solution phases is determined from experimental estimates of the respective solidus and solvus boundaries, where the solubility of Cu in the hcp-Mg phase has been reported to be 0.15, 0.23, and 0.23 at.% by Hansen,^[25] Stepanov and Kornilov,^[26] and Yue and Pierre,^[27] respectively. The resulting interaction parameter for the hcp solution phase is high and positive, as listed in Table 4. For the fcc solid solution, the interaction parameter listed in Table 4 is determined from the fcc phase boundary data reported by Jones^[22] and Rogelberg.^[23]

Regarding the intermediate compounds and the evaluation of coefficients in the expressions for $^{\circ}G_{\text{Cu:Mg}}^{\text{C15}}$ and $^{\circ}G_{\text{Mg:Cu}}^{\text{Cb}}$, as in Eq 1 and 7, we require experimental or theoretical values for heat capacity, enthalpy of formation, and melting temperature. The parameters that contribute to the heat capacity (i.e., coefficients c-f in Eq 1 are evaluated using the experimental results of Feufel and Sommer,^[16] recognizing that, above their melting temperatures, the heat capacities of the $\text{Cu}_2\text{Mg-C15}$ and $\text{Mg}_2\text{Cu-C}_b$ phases should approach that of the liquid phase, as given by the SGTE model.^[41] The available enthalpy of formation data include only the stable $\text{Cu}_2\text{Mg-C15}$ and $\text{Mg}_2\text{Cu-C}_b$ phases. They are listed in Table 1 along with our first principles calculations, which include these phases as well as the unstable compounds for which experimental data are not available. In addition, we consider the electromotive force (EMF) measurements of activity and chemical potential, reported by Arita et al.^[38] and Eremenko et al.^[39] for the Cu-Mg binary system. These are shown in Fig. 2 and compared with the values of chemical potential associated with our first principles calculations and with experimental measurements of ΔH_f .^[36,37] In this figure, the fcc/ $\text{Cu}_2\text{Mg-C15}$ coexistence curve suggests that the experimental values reported by King and Kleppa^[36] may be the most accurate, since they exhibit excellent agreement with the EMF data of Eremenko et al.^[39] Based on this selection and the heat capacity considerations discussed above, the coefficients listed in Table 3 for $^{\circ}G_{\text{Cu:Mg}}^{\text{C15}}$ and $^{\circ}G_{\text{Mg:Cu}}^{\text{Cb}}$ are evaluated with the corresponding experimental data.^[16,21,22,24,36,38,39] Finally, the two interaction parameters for the $\text{Cu}_2\text{Mg-C15}$ phase ($^{\circ}L_{\text{Cu:Cu,Mg}}^{\text{C15}}$ and $^{\circ}L_{\text{Cu,Mg:Mg}}^{\text{C15}}$) are evaluated using available phase equilibrium data.^[20-24] The Gibbs free

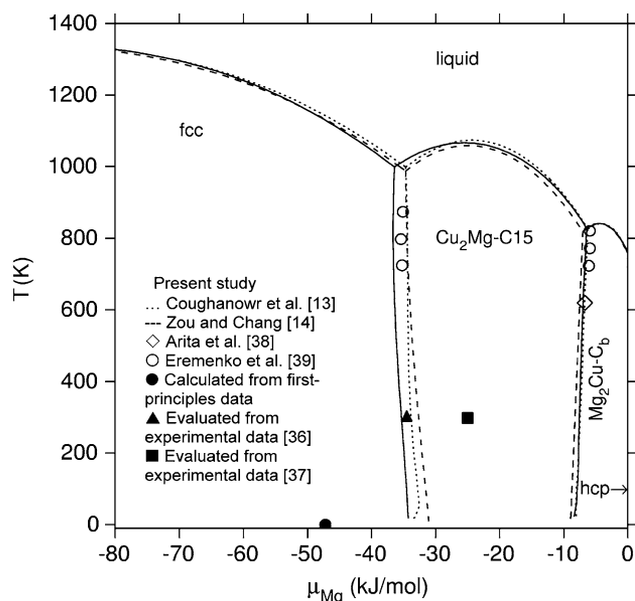


Fig. 2 The chemical potential of Mg associated with the indicated two-phase equilibrium, plotted over the relevant temperature range for the Cu-Mg system. The solid curves are computed and compared with prior modeling and experiment. The reference states are fcc-Cu and hcp-Mg

energy of formation, ΔG_{ij}^{θ} , for each unstable end-member in Eq 8, is treated as a constant and determined using the first-principles data in Table 1. The parameter assessment

practice is iterative and culminates with an optimized fit involving all selected data and calculation results. The final parameters from the overall optimization are listed in Tables 3 and 4.

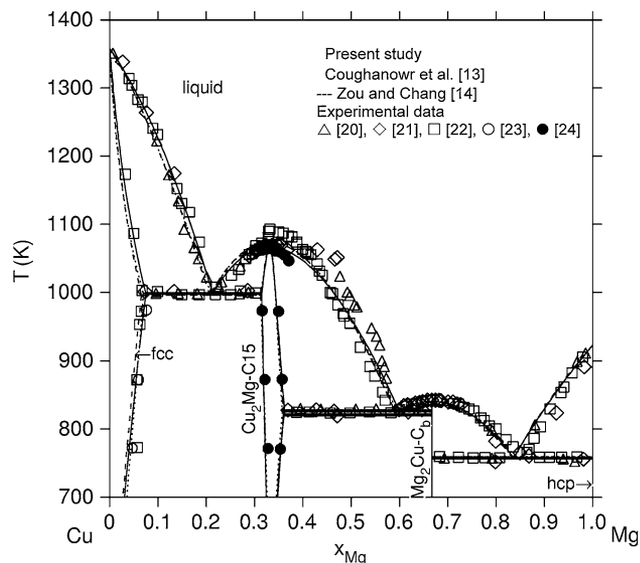


Fig. 3 The Cu-Mg phase diagram calculated using the parameters listed in Tables 2-4, shown with the relevant experimental data

4. Phase Equilibrium Results

The phase diagram yielded by our modeling effort is shown in Fig. 3 along with previously proposed phase diagrams and relevant experimental data. In Table 5, the invariant reactions are compared with experimental reports^[20-22] and with the models of Coughanowr^[13] and Zou and Chang.^[14] We note that the equilibrium phase boundaries produced by our model do not differ significantly from prior reports, except that our results may exhibit slightly better agreement with experimental liquidus and solidus data for the terminal solution phases. However, there are several important features of our thermodynamic description of this binary system that do differ substantially from previously reported models but are not dramatically evident in the equilibrium phase diagram itself. These differences primarily arise from (i) the use of chemical associates in our treatment of the liquid, (ii) our two-sublattice solution description of the intermediate phases, and (iii) inclusion of non-linear temperature dependent free energies of formation, all of which offer improvements over prior treatments

Table 5 Invariant reactions in the Cu-Mg system

Reaction		Calculated results			Experimental data	
		This work	Ref. ^[13]	Ref. ^[14]		
Liq → Cu ₂ Mg-C15	<i>T</i> , K	1066.3	1073.0	1066.2	1063 ^[20] 1070 ^[21] 1092 ^[22] 1066 ^[24]	
	<i>x</i> _(liquid, Mg)	0.333	0.333	0.333	0.333 ^[24]	
	<i>x</i> _(C15, Mg)	0.333	0.333	0.333	0.333 ^[24]	
	Liq → CuMg ₂ -C _b	<i>T</i> , K	840.5	841	841	841 ^[22,24]
		<i>x</i> _(liquid, Mg)	0.667	0.667	0.667	0.667 ^[24]
Liq → fcc + Cu ₂ Mg-C15	<i>x</i> _(Cb, Mg)	0.667	0.667	0.667	0.667 ^[24]	
	<i>T</i> , K	998.8	998	998.3	995 ^[22] 998 ^[20,24]	
	<i>x</i> _(liquid, Mg)	0.213	0.209	0.214	0.230 ^[20]	
Liq → CuMg ₂ -C _b + Cu ₂ Mg-C15	<i>x</i> _(fcc, Mg)	0.074	0.069	0.075	0.069 ^[24]	
	<i>x</i> _(C15, Mg)	0.314	0.313	0.316	0.310 ^[24]	
	<i>T</i> , K	826.5	825.5	825.2	828 ^[20,21] 825 ^[22,24]	
	<i>x</i> _(liquid, Mg)	0.594	0.595	0.588	0.585 ^[20]	
	<i>x</i> _(Cb, Mg)	0.667	0.667	0.667	0.667 ^[20]	
Liq → hcp + CuMg ₂ -C _b	<i>x</i> _(C15, Mg)	0.361	0.356	0.359	0.353 ^[20]	
	<i>T</i> , K	757.3	759.0	758.0	753 ^[20] 758 ^[21,22]	
	<i>x</i> _(liquid, Mg)	0.838	0.839	0.841	0.855 ^[21,22]	
	<i>x</i> _(hcp, Mg)	1.000	1.000	1.000	1 ^[22]	
	<i>x</i> _(Cb, Mg)	0.667	0.667	0.667	0.667 ^[22]	

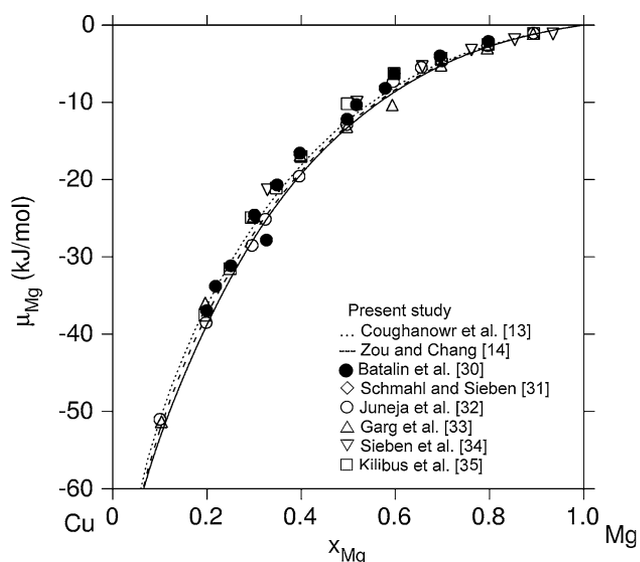


Fig. 4 The chemical potential of Mg as a function of composition for the Cu-Mg liquid phase calculated for 1,100 K and compared with prior modeling and experiment. The reference states are liq-Cu and liq-Mg

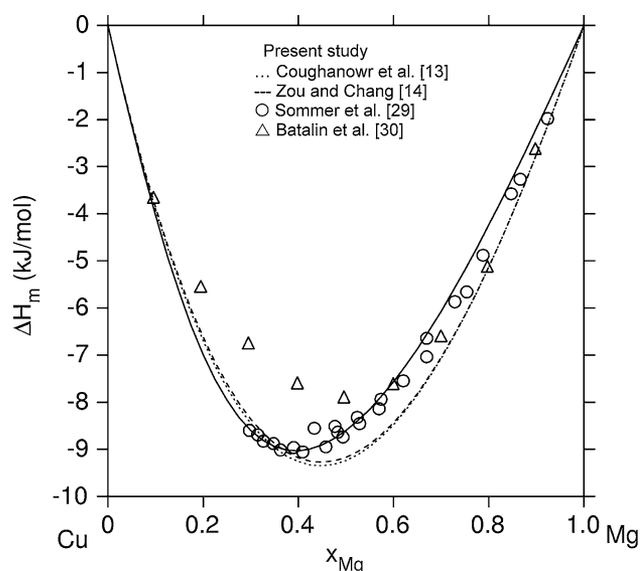


Fig. 6 The enthalpy of mixing of the liquid phase calculated for 1,120 K and compared with prior modeling and experiment. The reference states are liq-Cu and liq-Mg

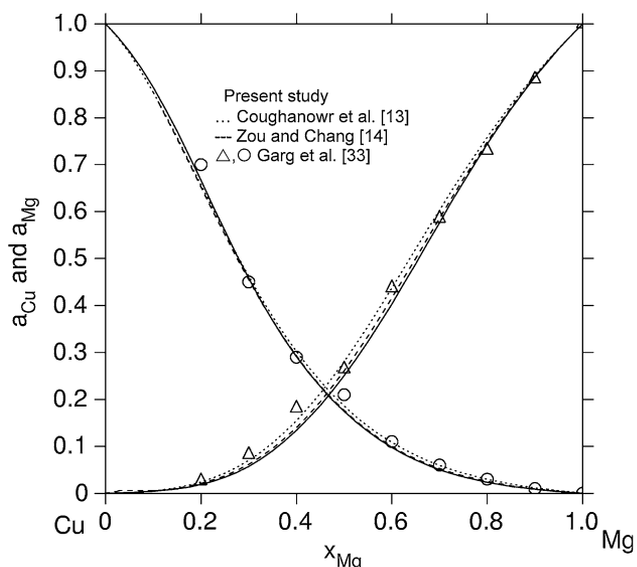


Fig. 5 The activity of Cu and Mg calculated for 1,200 K and compared with prior modeling and experiment. The reference states are liq-Cu and liq-Mg

and permit the quantitative description of thermodynamic parameters for both undercooled liquids and nonequilibrium compositions of the intermetallic phases.

Figures 4 and 5 show that there is little difference between prior models and ours with regard to quantification of chemical potential of the liquid phase over the full range of composition at 1100 K and 1200 K. However, Fig. 6 shows that our model provides much better agreement with

Table 6 The melting enthalpies of the compounds $\text{Cu}_2\text{Mg-C15}$ and $\text{Mg}_2\text{Cu-C}_b$

Compound	Calculated results, kJ/mol			Experimental data, kJ/mol ^[16]
	This work	Ref. ^[13]	Ref. ^[14]	
$\text{Cu}_2\text{Mg-C15}$	14.1	12.4	13.5	15.2 ± 1.5
$\text{Mg}_2\text{Cu-C}_b$	12.3	11.5	11.4	13.7 ± 1.5

the experimental measurements of enthalpy of mixing for the liquid phase, as reported by Sommer et al.^[29] This feature is essential if driving forces for various phase transitions involving both equilibrium and nonequilibrium compositions of the relevant phases are to be computed accurately and compared. In addition, we employ the temperature-dependent description given in Eq 1 for the Gibbs free energy of $\text{Cu}_2\text{Mg-C15}$ and $\text{Mg}_2\text{Cu-C}_b$ rather than the simpler description used in earlier reports.^[13,14] As a result, it is clear from Fig. 7 that our description reproduces the experimental heat capacity data^[16] far better than previously reported models.^[13,14] In addition, the modeled melting enthalpies for $\text{Cu}_2\text{Mg-C15}$ and $\text{Mg}_2\text{Cu-C}_b$ are listed in Table 6, where our results show better agreement with the experimental data for these phases^[16] than those reported by Coughanowr et al.^[13] and Zou and Chang.^[14] Finally, for the purpose of further direct comparison, we include in Fig. 1 our modeling results for the heat capacity of the $\text{Cu}_{14.5}\text{Mg}_{85.5}$ liquid, showing the dramatic improvement in low-temperature heat capacity compared to the models by Coughanowr et al.^[13] and Zou and Chang.^[14]

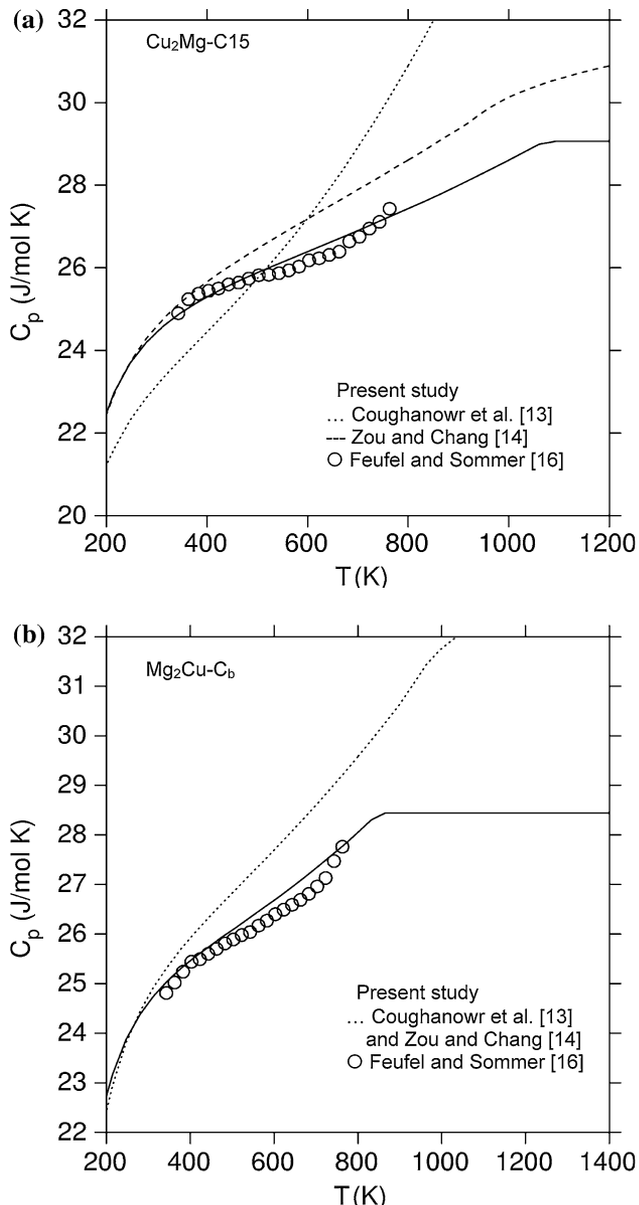


Fig. 7 The heat capacity (a) $\text{Cu}_2\text{Mg-C15}$ and (b) $\text{Mg}_2\text{Cu-C}_b$ computed using our model parameters in Tables 1-3 and compared with previously reported models and data

5. Summary and Conclusions

Employing a solution thermodynamics approach, the Gibbs free energy vs. composition curves for all relevant phases in the Cu-Mg binary system were estimated, and the associated binary phase diagram is reported here. While the resulting equilibrium phase diagram exhibits only modest deviation from previously reported diagrams, the current thermodynamic description exhibits several key differences from earlier thermodynamic models. These differences arise from (i) the two-sublattice solution treatment of the $\text{Mg}_2\text{Cu-}$

C_b phase, (ii) the use of first principles calculations to obtain the zero-Kelvin energies for the unstable compounds, and (iii) the use of an association model for the liquid phase. As a result, the current model provides a more realistic quantification of heat capacity for the relevant phases and a better description of the composition-dependent enthalpy of mixing for the liquid.

Acknowledgments

Work at the Ames Laboratory was supported by the U.S. Department of Energy, Basic Energy Sciences, under Contract No. DE-AC02-07CH11358.

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