### FRIENDS OF FREZCHEM (Version 16)

Attached is a "beta" version of the FREZCHEM model that includes chloride, bromide, perchlorate, nitrate, sulfate, and bicarbonate-carbonate salts, strong acid chemistry, ferrous and ferric iron chemistry, aluminum and silicon chemistries, ammonia and ammonium chemistries, and gas hydrate chemistry. This version includes both temperature and pressure dependencies. This folder includes a FORTRAN program listing (which you can download directly), four input files, a list of chemical species in the model (Table 1), instructions for model input (Tables 2-5, 8, and 10), and four examples of model outputs (Tables 6-7, 9 and 11).

This model is very much a work in progress. I will be mainly adding new chemistries to the model in the next few years. I have not spent much time debugging the model or making it user-friendly. In addition, there are convergence problems, at times, with the model. My version of the model was created with Absoft's ProFortran for the Macintosh. Porting this code to another FORTRAN compiler is always problematic. Once you have a FREZCHEM model working, verify correctness from an example from published data (see examples below). If you have additional problems, contact me via e-mail (giles.marion@dri.edu). Indicate the FREZCHEM version you are using (e.g., FREZCHEM16) and your model input.

The model is an equilibrium chemical thermodynamic model, meaning it will always select the most stable minerals. There are a few minerals (e.g., aragonite and vaterite) that are always metastable with respect to other minerals (e.g., calcite). To explicitly include a metastable mineral in your calculations necessitates removing the stable mineral from the mineral database. This is most simply done by assigning the stable mineral an arbitrary high  $K_{sp}$  through SOLIDPHASE.txt. The # of the  $K_{sp}$  for a specific mineral in the FORTRAN program is the same as the solid phase # in Table 1 (e.g.,  $K_{52}$  is the solubility product for calcite). If you are using the model to calculate pH, then you should make sure that the initial solution is charge-balanced. Otherwise, the model will force a charge balance by changing the bicarbonate-carbonate or acid concentrations, which could lead to a serious error in calculated pH if the solution is badly charge-balanced. If necessary, force a charge-balance in the initial solution by changing a major constituent that minimizes the effect on pH (e.g., Na or Cl). If you input Fe, Al, Si, or alkalinity, then you will have four options on how to deal with pH; adding NH<sub>3</sub>-NH<sub>4</sub> adds a fifth option (see Table 2).

The validation of this model is discussed in 16 publications: (1) Spencer et al. (1990) The prediction of mineral solubilities in natural waters: A chemical equilibrium model for the Na-K-Ca-Mg-Cl-SO<sub>4</sub>-H<sub>2</sub>O system. Geochim. Cosmochim. Acta, 54:575-590; (2) Marion and Farren (1999) Mineral solubilities in the Na-K-Mg-Ca-Cl-SO<sub>4</sub>-H<sub>2</sub>O system: A re-evaluation of the sulfate chemistry in the Spencer-Møller-Weare model. Geochim. Cosmochim. Acta, 63:1305-1318; (3) Marion (2001) Carbonate mineral solubility at low temperatures in the Na-K-Mg-Ca-H-Cl-SO<sub>4</sub>-OH-HCO<sub>3</sub>-CO<sub>3</sub>-CO<sub>2</sub>-H<sub>2</sub>O system. Geochim. Cosmochim. Acta, 65:1883-1896; (4) Marion (2002) A molal-based model for strong acid chemistry at low temperatures (<200 to 298 K). Geochim. Cosmochim. Acta, 66:2499-2516; (5) Marion et al. (2003) Modeling aqueous ferrous iron chemistry at low temperatures with application to Mars. Geochim. Cosmochim. Acta, 67:4251-4266; (6) Marion et al. (2005) Effects of pressure on aqueous chemical equilibria at subzero temperatures with applications to Europa. Geochim. Cosmochim. Acta, 69:259-274; (7) Marion et al. (2006) Modeling gas hydrate equilibria in

electrolyte solutions. CALPHAD, 30:248-259; (8) Marion (2007) Adapting molar data (without density) for molal models. Computers & Geosciences, 33:829-834. (9) Marion and Kargel (2008) Cold Aqueous Planetary Geochemistry with FREZCHEM: From Modeling to the Search for Life at the Limits. Springer; (10) Marion et al. (2008) Modeling ferrous-ferric iron chemistry with application to Martian surface geochemistry. Geochim. Cosmochim. Acta, 72:242-266; (11) Marion et al., (2009) Br/Cl partitioning in chloride minerals in the Burns formation on Mars. Icarus, 200:436-445; (12) Marion et al. (2009). Modeling aluminum-silicon chemistries and application to Australian playa lakes as analogues for Mars. Geochim. Cosmochim. Acta. 73:3493-3511; (13) Marion et al. (2011). Modeling hot spring chemistries with applications to Martian silica formation. Icarus. 212:629-642; (14) Marion et al. (2012). Modeling ammonia-ammonium aqueous chemistries in the Solar System's icy bodies. Icarus. 220:932-946; (15) Marion et al. (2013). Sulfite-sulfide-sulfate-carbonate equilibria with applications to Mars. Icarus. 225:342-351; and (16) Marion et al (2014). Modeling nitrogen-gas, -liquid, -solid chemistries at low temperatures (173-298 K) with applications to Titan. Icarus. 236:1-8.

Compared to earlier versions, this Version 16 of the FREZCHEM model contains new parameterizations dealing with sulfite, sulfide, and nitrogen species. Also, Versions 15 and 16 have lowered the temperature level to 173 K (see Tables 8-11).

A fundamental change was made in FREZCHEM 13 on how to input data into the model. Earlier versions required inputs via the computer screen. Versions 13-16 require inputs via data files; this approach simplifies and speeds up model inputting. There are four input files that must be built to run FREZCHEM. Table 2 describes the main model inputs; Table 3 presents the main Input.txt file; and Table 4 describes in more detail how to handle gases for these inputs. There are three minor input files that are lumped together in Table 5. Table 2 is just a verbal description of the material that is in the Input.txt file of Table 3. Similarly, Table 4 describes in more detail how gases need to be dealt with in Input.txt (Table 3). In the Input.txt file, note that inputs are all placed to the left of the ",". Also, do not remove the ",". That comma separates model input from descriptive words. The three minor files in Table 5 include: (A) SOLIDPHASE.txt, (B) SOLIDMASS.txt, and (C) NUANCES.txt. SOLIDPHASE.txt allows the user to remove all solid phases from equilibrium calculations or some specific minerals. That option allows for a pure solution phase calculation without any minerals precipitating. In the molar to molal conversion example in Table 7, all solid phases had to be removed. Note the assigned exceptionally high equilibrium constants in Table 7, which is what keeps the solid phases from precipitating. In a previous Phoenix site calculation (Marion et al., 2010), we removed magnesite (MgCO<sub>3</sub>) and dolomite  $[CaMg(CO_3)_2]$  from model calculations (Table 5A), which led to calcite  $(CaCO_3)$  and  $hydromagnesite~[3Mg(CO_3)_2 \bullet MgSO_4 \bullet 3H_2O]~precipitating.~Generally, model calculations~start~$ with aqueous/gas phases, without initial solid phases; but if you want a particular solid phase to control the solution phase chemical composition, then you can specify the solid phase and its mole mass (Table 5); the mole mass is an arbitrary amount that must not completely dissolve in 1.0 kg H<sub>2</sub>O. For example, we assumed that Earth seawater would have been saturated with dolomite during Snowball Earth (Marion and Kargel, 2008). Changing the first line of SOLIDMASS.txt from No(0) to YES(1) and specifying dolomite would allow saturation with dolomite. The NUANCES.txt file allows for temperature, water content, or pressure changes to be adjusted during a specific run. For example, if you want to know the eutectic temperature of a salt assemblage, and you know that this will occur slightly below 259 K, you could change the MT term from 5 K between 298 and 263 K (as assigned by Input.txt) to 1 K between 263 and

259 K, and 0.1 K below 259 K (Table 5). This scenario would allow for a more accurate estimate of the eutectic temperature than using either a 5 K or 1 K term for the T decrement. With respect to NUANCES.txt, always retain two steps for temperature, water content, and pressure changes, even if you need to duplicate two steps (e.g., 263.15 0.1, 263.15 0.1). Table 1. A listing of chemical species in the FREZCHEM model (Version 16.1).

# Species			#	A. Solution and Atmospheric S Species	pecies #	Species	#
<u>Species</u>	#	Species	#	Species	#	<u>Species</u>	<u>#</u>
1 Na <sup>+</sup> (aq)			16	Cl <sup>-</sup> (aq)	201	CO <sub>2</sub> (aq)	216
$\text{Fe(OH)}_{2}^{0}(\text{aq})$	231	HF <sup>0</sup> (aq)					
2 K <sup>+</sup> (aq)			17	SO <sub>4</sub> <sup>2-</sup> (aq)	202	FeCO <sub>3</sub> °(aq)	217
Fe(OH) <sub>3</sub> (aq)	232	Sr <sup>2+</sup> (aq)					
$3  \text{Ca}^{2+}(\text{aq})$			18	OH (aq)	203	HCl(g)	218
FeOH <sup>2+</sup> (aq)		233 N	$H_3^{0}$	g)			
$4  Mg^{2+}(aq)$			19	HCO <sub>3</sub> (aq)	204	CaCO <sub>3</sub> °(aq)	219
$Fe(OH)_2^+(aq)$	234	$NH_3^{0}$ (aq)					
5 H <sup>+</sup> (aq)			20	$CO_3^{2}$ -(aq)	205	MgCO <sub>3</sub> °(aq)	220
$Fe(OH)_3^{0}(aq)$	235	$SO_2^{0}(g)$		-			
6 MgOH <sup>+</sup> (aq)			21	HSO <sub>4</sub> (aq)	206	HNO <sub>3</sub> (g)	221
Fe(OH) <sub>4</sub> (aq)	236	$SO_2^{0}(aq)$					
7 Fe <sup>2+</sup> (aq)			22	NO <sub>3</sub> (aq)	207	$H_2SO_4(g)$	222
$Al(OH)^{2+}(aq)$	237	$\mathrm{H}_2\mathrm{S}^0(\mathrm{g})$					
8 FeOH <sup>+</sup> (aq)			23	Br (aq)	208	$H_2O(g)$	223
$Al(OH)_2^+(aq)$	238	$\mathrm{H_2S}^0(\mathrm{aq})$					
9 Fe <sup>3+</sup> (aq)			24	ClO <sub>4</sub> (aq)	209	CO <sub>2</sub> (g)	224
$Al(OH)_3^{0}(aq)$	239	$N_2(g)$					
10 Al <sup>3+</sup> (aq)			25	$B(OH)_4$ (aq)	210	$H_2O(1)$	225
Al(OH) <sub>4</sub> (aq)	240	N <sub>2</sub> (aq)					
11 $NH_4^+$ (aq)			26	HSO <sub>3</sub> (aq)	211	$O_2(g)$	226
$Si(OH)_4^0$ (aq)							
·			27	SO <sub>3</sub> <sup>2-</sup> (aq)	212	O <sub>2</sub> (aq)	227
SiO(OH) <sub>3</sub> (aq)							
			28	HS (aq)	213	H <sub>2</sub> (g)	228
SrCO <sub>3</sub> (aq)				-		2 '	
			29	$S^{2}$ -(aq)	214	$CH_4(g)$	229

# Species	# Species	B. Solid Phase Species	# Species	# Species
# Species #	# Species Species	# Species	# Species	<u>species</u>
31 H <sub>2</sub> O(cr,I)	51 Na <sub>2</sub> SO <sub>4</sub> •3K <sub>2</sub> SO <sub>4</sub> (cr)	71 H <sub>2</sub> SO <sub>4</sub> •4H <sub>2</sub> O(cr)	91 FeCl <sub>3</sub> •2KCl•H <sub>2</sub> O(cr)	111
$K_2$ Fe(II) <sub>5</sub> Fe(III) <sub>4</sub> (SO <sub>4</sub> ) <sub>1</sub>	2•18H <sub>2</sub> O(cr) 131 NaC	1O <sub>4</sub> •2H <sub>2</sub> O(cr)	_	
32 NaCl•2H <sub>2</sub> O(cr)	52 CaCO <sub>3</sub> (cr,calcite)	72 HCl•6H <sub>2</sub> O(cr)	92 Fe <sub>2</sub> (SO <sub>4</sub> ) <sub>3</sub> (cr)	112
AlCl <sub>3</sub> •6H <sub>2</sub> O(cr)	132 NH <sub>4</sub>	Cl(cr)	2	
33 NaCl(cr)	53 MgCO <sub>3</sub> (cr)	73 NaNO <sub>3</sub> •Na <sub>2</sub> SO <sub>4</sub> •2H <sub>2</sub> O(	cr) 93 Fe <sub>2</sub> (SO <sub>4</sub> ) <sub>3</sub> •2K <sub>2</sub> SO <sub>4</sub> •14H <sub>2</sub> O(c	r)
113 Al <sub>2</sub> (SO <sub>4</sub> ) <sub>3</sub> •17H <sub>2</sub> O(	cr) 133 (NH	$_4)_2$ SO <sub>4</sub> (cr)	2 .6 2 . 2	
		74 Na <sub>3</sub> H(SO <sub>4</sub> ) <sub>2</sub> (cr)	94 K <sub>2</sub> SO <sub>4</sub> •FeSO <sub>4</sub> •6H	O(cr)
	134 NH <sub>3</sub>		2 7 7	<u> </u>
	55 MgCO <sub>3</sub> •5H <sub>2</sub> O(cr)		95 Na <sub>2</sub> SO <sub>4</sub> •FeSO <sub>4</sub> •4H <sub>2</sub> O(cr)	115
MgBr <sub>2</sub>			2 4 4 2	
		76 $K_3H(SO_4)_2(cr)$	96 Fe <sub>2</sub> (SO <sub>4</sub> ) <sub>3</sub> •9H <sub>2</sub> O(cr)	116
	136 NH <sub>4</sub>		2 73 2	
	57 NaHCO <sub>3</sub> (cr)		97 Fe <sub>2</sub> (SO <sub>4</sub> ) <sub>3</sub> •H <sub>2</sub> SO <sub>4</sub>	•8H <sub>2</sub> O(cr)
	137 NH <sub>4</sub> ClO <sub>4</sub> (cr)	5 5 44	2 43 2 4	. 2
		78 $K_{\varrho}H_{6}(SO_{4})_{7} \cdot H_{2}O(cr)$	98 KFe <sub>3</sub> (SO <sub>4</sub> ) <sub>2</sub> (OH) <sub>6</sub> (cr)	118 S
	138 NH <sub>3</sub> •2H <sub>2</sub> O(cr)		3 42 0	
	59 NaHCO <sub>3</sub> •Na <sub>2</sub> CO <sub>3</sub> •2H <sub>2</sub> O		99 NaFe <sub>3</sub> (SO <sub>4</sub> ) <sub>2</sub> (OH) <sub>6</sub> (cr)	119
	139 K <sub>2</sub> S		3 42 0	
			) 100 H <sub>3</sub> OFe <sub>3</sub> (SO <sub>4</sub> ) <sub>2</sub> (OH) <sub>6</sub> (cr)	120
	140 Na <sub>2</sub>		3 3 42 0	
		81 FeSO <sub>4</sub> •7H <sub>2</sub> O(cr)	101 <b>▼</b> -Fe <sub>2</sub> O <sub>3</sub> (cr)	121
	141 CaSe		2 3	
	62 Na <sub>2</sub> CO <sub>3</sub> •7H <sub>2</sub> O(cr)		102 W-FeO(OH)(cr)	122
	142 MgS			
	63 KHCO <sub>3</sub> (cr)		103 W-FeO(OH)(cr)	123
	O(cr) 143 FeSO <sub>3</sub> •5H <sub>2</sub> O(c			
		84 FeCl <sub>2</sub> •4H <sub>2</sub> O(cr)	104 FeO(OH) <sub>0.75</sub> (SO <sub>4</sub> ) <sub>0.125</sub> (cr)	
	144 (NH <sub>2</sub>		0.75 4 0.125	
		85 FeCO <sub>3</sub> (cr)	105 FeSO <sub>4</sub> •4H <sub>2</sub> O(cr)	
	•22H <sub>2</sub> O(cr) 145 FeS <sub>2</sub>		4 2	
	O(cr) 66 HNO <sub>3</sub> •3H <sub>2</sub> O(cr)		106 Fe <sub>2</sub> (SO <sub>4</sub> ) <sub>3</sub> •7H <sub>2</sub> O(cr)	126
NaClO <sub>4</sub> •H <sub>2</sub> O(cr)	146 N <sub>2</sub> •6		2 43 2	
	2O(cr) 67 KNO <sub>3</sub> (cr)	=	107 Fe(II)Fe(III) <sub>4</sub> (SO <sub>4</sub> ) <sub>6</sub> (OH) <sub>2</sub> •2	20H <sub>2</sub> O(cr) 127
$Mg(ClO_4)_2 \bullet 8H_2O(cr)$	147 CaSO <sub>4</sub> •0.5H <sub>2</sub> O <sub>6</sub>		4.0	2
48 CaSO <sub>4</sub> •2H <sub>2</sub> O(cr)	68 NaNO <sub>3</sub> (cr)	88 CH <sub>4</sub> •6H <sub>2</sub> O(cr)	108 Fe <sub>5</sub> (SO <sub>4</sub> ) <sub>6</sub> O(OH) •20H <sub>2</sub> O(cr	)
128 Ca(ClO <sub>4</sub> ) <sub>2</sub> •6H <sub>2</sub> O(cr	-	+ 2	J 40 ' ' Z '	
49 CaSO <sub>4</sub> (cr)		Cl <sub>2</sub> •10H <sub>2</sub> O(cr) 109 Fe(	$(II)$ Fe $(III)_2$ (SO <sub>4</sub> ) <sub>4</sub> •22H <sub>2</sub> O(cr)	129 KClO₄(cr)
· ·		90 FeCl <sub>3</sub> •6H <sub>2</sub> O(cr)	110 Fe(II)Fe(III) <sub>2</sub> (SO <sub>4</sub> ) <sub>4</sub> •14H <sub>2</sub> O(	
$Mg(ClO_4)_2 \bullet 6H_2O(cr)$	2 42 ()	3 -2-()	2004/4	. ,

## Table 2. Description of Model Inputs (Version 16) (Compare with Table 3).

<u>Title</u>: Any alphanumeric character up to 50 characters.

<u>Freeze(1)</u> or <u>Evaporation(2)</u> or <u>Pressure (3)</u> <u>Pathway</u>: Enter 1, 2, or 3 depending on whether you want to simulate a temperature change (1), an evaporation (2), or a pressure change (3). For evaluating a single point, enter "1".

<u>Equilibrium(1)</u> or <u>Fractional(2)</u> <u>Crystallization</u>: In equilibrium crystallization (1), precipitated solids are allowed to re-equilibrate with the solution phase as environmental conditions change. In fractional crystallization (2), precipitated solids are removed and not allowed to re-equilibrate with the solution phase as environmental conditions change. <u>Seawater Salinity</u>: If you want seawater salinity to govern the calculations, enter 1 for Yes, and 0 for No.

<u>Practical Salinity</u>: If Yes in the above line, enter  $S_{\mathbf{p}}$ . If No, enter 0.0.

<u>Calcite Supersaturation in Seawater</u>: If you want this to be considered, enter 1 for Yes, or 0 for No.

Sodium (m/kg): Enter sodium molality (moles/kg(water)). Otherwise, enter 0.0.

<u>Potassium (m/kg)</u>: Enter potassium molality (moles/kg(water)). Otherwise, enter 0.0.

Calcium (m/kg): Enter calcium molality (moles/kg(water)). Otherwise, enter 0.0.

Magnesium (m/kg): Enter magnesium molality (moles/kg(water)). Otherwise, enter 0.0.

Strontium (m/kg): Enter strontium molality (moles/kg(water)). Otherwise, enter 0.0.

Ferrous Iron (m/kg): Enter ferrous iron molality (moles/kg(water)). Otherwise, enter 0.0.

Ferric Iron (m/kg): Enter ferric iron molality (moles/kg(water)). Otherwise, enter 0.0.

Aluminum (m/kg): Enter aluminum molality (moles/kg(water)). Otherwise, enter 0.0.

<u>Silicon (m/kg)</u>: Enter silicon molality (moles/kg(water)). Otherwise, enter 0.0.

Ammonium (m/kg): Enter ammonium molality (moles/kg(water)). Otherwise, enter 0.0.

<u>If Iron, Aluminum, Silicon, Alkalinity, or Ammonia-Ammonium are selected</u>, then choose an acidity option:

Acidity ignored(Option 1), enter 1.

Acidity fixed by pH(Option 2), enter 2.

Acidity fixed by H<sup>+</sup> ion concentration(Option 3), enter 3.

Acidity fixed by alkalinity(Option 4), enter 4.

Acidity fixed by  $NH_3(aq)$  and  $NH_4(aq)(Option 5)$ , enter 2.

<u>Initial pH</u>: Option 1, enter 0; Option 2, enter pH; Options 3 and 4, enter an approximate pH; Option 5, enter 10.

Chloride (m/kg): Enter chloride molality (moles/kg(water)). Otherwise, enter 0.0.

Bromide (m/kg): Enter bromide molality (moles/kg(water)). Otherwise, enter 0.0.

Perchlorate (m/kg): Enter perchlorate molality (moles/kg(water)). Otherwise, enter 0.0.

Sulfate (m/kg): Enter sulfate molality (moles/kg(water)). Otherwise, enter 0.0.

Nitrate (m/kg): Enter nitrate molality (moles/kg(water)). Otherwise, enter 0.0.

Carbonate Alkalinity: Enter as equivalents/kg(water). If alkalinity = 0.0, then you must

enter 0.0. The latter will cause the model to skip all bicarbonate-carbonate with pH chemistries in the model.

<u>Sulfite Alkalinity</u>: Enter as equivalents/kg(water). If alkalinity = 0.0, then you <u>must</u> enter 0.0.

<u>Sulfide Acidity</u>: Enter as equivalents/kg(water). If acidity = 0.0, then you <u>must</u> enter 0.0. <u>Acidity</u>: Enter as equivalents/kg(water). This is the total hydrogen concentration, if known initially. Generally this is only known for strong acid solutions. For example, for a 1 molal  $H_2SO_4$  solution, enter 2.00. Otherwise, enter 0.0. The equations used to calculate pH for the alkalinity and acidity cases are incompatible. So, a specification of either carbonate alkalinity or acidity requires that the other variable be assigned a value of 0.00. This will channel the calculations to the proper algorithm.

<u>HCl(bars)</u>: If the HCl atmospheric concentration is known, then specify here. Otherwise, enter 0.0. If you specify 0.0, then the model will calculate HCl(bars). Note that if you specify HCl(bars) or the other acids below, then these properties override the total acidity specification (see above). That is, the solution is equilibrated with the atmospheric concentration. Note, you can, if desired, specify atmospheric concentrations for some acids (e.g., HCl and HNO<sub>3</sub>) and leave other acid partial pressure unspecified (e.g.,  $H_2SO_4 = 0.0$ ).

 $\underline{\text{HNO}_3(\text{bars})}$ : If the  $\text{HNO}_3$  atmospheric concentration is known, then specify here. Otherwise, enter 0.0.

 $\underline{\text{H}_2\text{SO}_4(\text{bars})}$ : If the  $\underline{\text{H}_2\text{SO}_4}$  atmospheric concentration is known, then specify here. Otherwise, enter 0.0.

<u>Boron (m/kg)</u>: Enter boron molality (moles/kg(water)). Otherwise, enter 0.0. <u>Fluoride (m/kg)</u>: Enter fluoride molality (moles/kg(water)). Otherwise, enter 0.0. <u>Initial Total Pressure (bars)</u>: Enter the initial total pressure of the system. <u>Initial CO<sub>2</sub>(bars)</u>: If alkalinity > 0.0 or CO<sub>2</sub> hydrates are simulated, then specify the initial concentration of CO<sub>2</sub>(g) in bars.

Mole Fraction of  $CO_2$ : Enter the mole fraction of  $CO_2(g)$  for the system (mole fraction =  $P_{CO2}$ /total pressure). For pure  $CO_2$ , enter 1.0. If 0.0, then  $CO_2(g)$  is fixed and independent of total pressure.

 $\underline{O_2(bars)}$ : If the atmospheric concentration of oxygen is known, then specify here.

Otherwise, enter 0.0. If you are interested in ferrous iron chemistry, then you may want to assign  $O_2$  a value of 0.0. Otherwise, it is likely that the insolubility of ferric minerals in the presence of  $O_2$  will cause all the iron to precipitate as a ferric mineral [see discussions in Marion et al., (2003a) iron paper].

<u>Initial CH<sub>4</sub>(bars)</u>: If CH<sub>4</sub> is simulated, then specify the initial concentration of CH<sub>4</sub>(g) in bars.

Mole Fraction of  $CH_4$ : Enter the mole fraction of  $CH_4(g)$  for the system (mole fraction =  $P_{CH4}$ /total pressure). For pure  $CH_4$ , enter 1.0. If 0.0, then  $CH_4(g)$  is fixed and independent of total pressure.

Mixed CH<sub>4</sub>-CO<sub>2</sub> Gas Hydrate?: If both CH<sub>4</sub>(g) and CO<sub>2</sub>(g) are specified as inputs, then

you can use this data to estimate the stability of a mixed  $CH_4$ - $CO_2$  gas hydrate (YES = 1) or treat the two gases as independent gas hydrates (NO = 0).

<u>Initial NH<sub>3</sub>(bars)</u>: If NH<sub>3</sub>(g) are inputs, then specify the initial concentration of NH<sub>3</sub>(g) in bars. Do not enter positive values (> 0) for both NH<sub>3</sub>(bars) and NH<sub>3</sub>(aq).

 $\underline{\text{Initial NH}_3(\text{aq})}: \text{ If NH}_3(\text{aq}) \text{ are inputs, then enter NH}_3(\text{aq}) \text{ molality (moles/kg(water))}.$ 

Do not enter positive values (> 0) for both  $NH_3$ (bars) and  $NH_3$ (aq).

<u>Initial N<sub>2</sub>(bars)</u>: If the atmospheric concentration of nitrogen is known, then specify here. Otherwise, enter 0.0.

Mole Fraction of  $N_2$ : Enter the mole fraction of  $N_2(g)$  for the system (mole fraction =  $P_{N2}$ /total pressure). For pure  $N_2$ , enter 1.0. If 0.0, then  $N_2(g)$  is fixed and independent of total pressure.

<u>Mixed N<sub>2</sub>-CH<sub>4</sub> Gas Hydrate?</u>: If both N<sub>2</sub>(g) and CH<sub>4</sub>(g) are specified as inputs, then you can use this data to estimate the stability of a mixed N<sub>2</sub>-CH<sub>4</sub> gas hydrate (YES = 1) or treat the two gases as independent gas hydrates (NO = 0).

<u>Mixed N<sub>2</sub>-CO<sub>2</sub> Gas Hydrate?</u>: If both N<sub>2</sub>(g) and CO<sub>2</sub>(g) are specified as inputs, then you can use this data to estimate the stability of a mixed N<sub>2</sub>-CO<sub>2</sub> gas hydrate (YES = 1) or treat the two gases as independent gas hydrates (NO = 0).

Molar to Molal Conversion?: If you want to convert molar data into molal concentrations, then enter (YES = 1) or (NO = 0).

<u>Salinity/liter</u>: If yes above, then you must enter the total aqueous salinity (g salt/liter), which can be calculated from molar data [g salt/liter = (moles/liter) x (g salt/mole)]. In the case depicted in Table 7, the SL value is 316.57 g salt/liter (5.417 x 58.44). <u>Initial Temperature(K)</u>: Enter the temperature in absolute degrees (K) for start of simulation (e.g., 273.15).

For Temperature Change Pathway(1):

Final Temperature(K): Enter final temperature of simulation (e.g., 263.15).

Temperature Decrement(K): The temperature interval between simulations (e.g. 1). For the above temperature designations, the model would calculate equilibrium starting at 273.15 K and ending at 263.15 K at 1 K intervals. If you want to change the decrement in a run (e.g., to reduce the step size near an equilibrium), see File "NUANCES.txt."

For Evaporation Pathway(2):

<u>Initial Water (g)</u>: Normally enter "1000" at this point. The standard weight basis of the model is 1000 g water plus associated salts. In you enter 100, instead of 1000, the initial ion concentrations, specified above, will be multiplied by 10.0 (1000/100) as the starting compositions for calculations. This feature of the model is useful in precisely locating where minerals start to precipitate during the evaporation process without having to calculate every small change between 1000 g and 1 g.

<u>Final Water (g)</u>: Enter the final amount of water that you want to remain in the system (e.g., 100).

Water Decrement (g): Enter the water decrement for simulations (e.g., 50 g). Specifying initial = 1000, final = 100, and decrement = 50 would result in calculations at

1000g, 950g, ....100g. If you want to change the decrement in a run (e.g., to reduce the step size near an equilibrium), see Files "NUANCES.txt." For Pressure Pathway(3):

<u>Final Pressure(bars)</u>: Enter the final pressure of the simulation [e.g., 101.01325 bars (100 atm)].

<u>Pressure Increment(bars)</u>: Enter the pressure increment. For example, if initial pressure is 1.01 bars, final pressure is 101.01 bars, and pressure increment is 1.0 bars, then the simulation would calculate at 1.01, 2.01, 3.01, ....101.01325 bars. If you want to change the increment in a run, see File "NUANCES."

 $\textbf{Input.txt.} \quad \textbf{This is the main input for applications of FREZCHEM.}$ In this particular case, the model simulates SO2-H2S chemistry from 298.15 to 263.15 K (Table 6) (See Marion et al., 2013). TITLE: SO2-H2S model parameters 1, FREEZE(1) OR EVAPORATION(2) OR PRESSURE(3) SCENARIO ? CALLED PATH BELOW. 2, EQUILIBRIUM(1) OR FRACTIONAL(2) CRYSTALLIZATION? 0, WANT SEAWATER SALINITY(Sp) TO GOVERN THE CALCULATIONS, Y=1, N=0. 0.0, IF YES ABOVE, ENTER SEAWATER PRACTICAL SALINITY(Sp) OR 0.0. 0, WANT SEAWATER CARBONATE SUPERSATURATION TO BE CONSIDERED? Y=1, N=0. 0.00 SODIUM(M/KG). 0.00, POTASSIUM(M/KG). 0.10, CALCIUM(M/KG 1.00, MAGNESIUM(M/KG) 0.00, STRONTIUM(M/KG). 0.00, FERROUS IRON(M/KG). 0.00, FERRIC IRON (M/KG). 0.00, ALUMINUM(M/KG). 0.00, SILICA (M/KG). 0.00, AMMONIUM(M/KG). FOR FE, AL, SI, AND ALKALINE CHEMISTRIES, DO YOU WANT ACIDITY IGNORED(1), OR FIXED BY PH(2), OR ACIDITY(3), OR ALKALINITY(4)? IF YES, THEN ENTER NUM. NOTE THAT OPTIONS 3 AND 4 REQUIRE A FURTHER SPECIFICATION BELOW. OPTIONS 2-4 WILL ADJUST SOLUTION-PHASE CHARGE BALANCE AS AL, FE, OR SI REACTIONS PRODUCE ACIDITY BY ASSUMING H+ REACTS WITH ROCKS TO RELEASE NA, K, CA, MG, OR FE(II) IONS. SOME OF THE LATTER IONS MUST BE PRESENT AS INPUT TO SERVE AS AN ION SINK/SOURCE. FOR NH3(AQ) + NH4(AQ) CASE, SET OPTION = 2, WITH PH = 10.0. 3. SPECIFY ABOVE ACIDITY OPTION. 7.5, SPECIFY INITIAL PH. 1.00, CHLORIDE(M/KG). 0.00, BROMIDE(M/KG). 0.00, PERCHLORATE (M/KG). 0.40, SULFATE(M/KG). 0.00, NITRATE(M/KG). 0.00, CARBON ALKALINITY (EQUIVALENTS/KG). 0.40, SULFITE ALKALINITY (EQUIVALENTS/KG). 0.00, SULFIDE ACIDITY(EQUIVALENTS/KG). 0.00, ACIDITY(EQUIVALENTS/KG). 0.00, IF YOU WANT TO SPECIFY HCL(BARS), ENTER VALUE HERE. 0.00, IF YOU WANT TO SPECIFY HNO3(BARS), ENTER VALUE HERE. 0.00, IF YOU WANT TO SPECIFY H2SO4(BARS), ENTER VALUE HERE. 0.00, BORON (M/KG). 0.00, FLUORIDE(M/KG). 1.00, INITIAL TOTAL PRESSURE(BARS). 0.00, INITIAL CO2(BARS).

```
0.00, ENTER MOLE FRACTION OF CO2, 0=FIXED CO2, 1=PURE CO2.
0.00, INITIAL O2(BARS).
0.00, INITIAL CH4(BARS).
0.00, ENTER MOLES OF CH4, 0=FIXED CH4, 1=PURE CH4.
0.00, CONSIDER A MIXED CO2-CH4 GAS HYDRATE(YES=1, NO=0)?
0.00, INITIAL NH3(G)(BARS), DO NOT INCLUDE BOTH NH3 INPUTS.
0.00, INITIAL NH3(AQ)(M/KG), DO NOT INCLUDE BOTH NH3 INPUTS.
0.78, INITIAL N2(BARS)
0.00, ENTER MOLE FRACTION OF N2, 0=FIXED N2, 1=PURE N2.
0, CONSIDER A MIXED N2-CH4 GAS HYDRATE(YES=1, NO=0)?
0, CONSIDER A MIXED N2-C02 GAS HYDRATE(YES=1, NO=0)?
0, MOLAR TO MOLAL CONVERSION? YES=1, NO=0.
0.00, IF YES ABOVE, ENTER SALINITY(G)/LITER.
298.15, INITIAL TEMPERATURE(K).
263.15, FINAL TEMPERATURE(K), IF PATH =1, OTHERWISE, SET = 0.
5.00, TEMPERATURE DECREMENT(K), IF PATH = 1, OTHERWISE, SET = 0.
1000, INITIAL WATER(G), IF PATH = 2, OTHERWISE, SET = 1000.
0.00, FINAL WATER (G), IF PATH = 2, OTHERWISE, SET = 0.
0.00, WATER DECREMENT(K), IF PATH = 2, OTHERWISE, SET = 0.
1.00, FINAL PRESSURE(BARS), IF PATH = 3, OTHERWISE, SET = 0.
0.00, PRESSURE INCREMENT(BARS), IF PATH = 3, OTHERWISE, SET = 0.
```

## **Table 4. Inputting Gases into the Model**

SYSTEM INPUTS	NO GASES	FIXED GAS CONCS.	VARIABLE GAS CONCS.
1. OPEN CARBON SYSTEM	1	1	1
2. INITIAL TOTAL PRESSURE	USER SPECIFICATION	USER SPECIFICATION	USER SPECIFICATION
3. INITIAL CO2	0	USER SPECIFICATION	USER SPECIFICATION
4. MOLE FRACTION OF CO2	0	0	USER SPECIFICATION
5. INITIAL O2	0	USER SPECIFICATION	0
6. INITIAL CH4	0	USER SPECIFICATION	USER SPECIFICATION
7. MOLE FRACTION OF CH4	0	0	USER SPECIFICATION
8. MIXED CH4-CO2 GAS HYDRATE	0	0	1
9. INITIAL NH3	0	USER SPECIFICATION	0
	CLOSED CARBON		
1. OPEN CARBON SYSTEM	2		
2. INITIAL TOTAL PRESSURE	USER SPECIFICATION		
3. INITIAL CO2	USER SPECIFICATION		
4. MOLE FRACTION OF CO2	USER SPECIFICATION		
5. INITIAL O2	0		
6. INITIAL CH4	USER SPECIFICATION		
7. MOLE FRACTION OF CH4	USER SPECIFICATION		
8. MIXED CH4-CO2 GAS HYDRATE	1		
9. INITIAL NH3	0		
NO GASES means that you do no	t want any gases considered in these	chemical equilibrium calculations.	
	you want the user specified gas conc	entrations to remain fixed	
as T, P, and H2O content change. 3.80e-4 bars.	For example, atm. CO2 is equal to		
VARIABLE GAS CONCS, were s	pecifically designed for CO2 and CF	[4 gas hydrate equilibrium which all	ows gas
pressures to increase as total pressurerincreases.			·
CLOSED CARBON was specifica	lly designed for CO2 and CH4 gas h	ydrate equilibrium in small volumes	(e.g, ice pockets).

The aqueous component contains 1.0 kg of water. The air component contains 0.1 liter.					
To change the aqueous/air ratio, change the 0.1 liter multiplier in FC13.3 lines 300 and 316.					

Table 5. Minor input files for FREZCHEM 13.3

#### (A). SOLIDPHASE.txt

0, WANT ALL SOLID PHASES TO BE REMOVED, YES=1, NO=0. 2, WANT SPECIFIC SOLID PHASES REMOVED, SPECIFY # OF CASES. 53, SPECIFY SAMPLE # TO BE REMOVED. 61, SPECIFY SAMPLE # TO BE REMOVED.

#### (B). SOLIDMASS.txt

0, WANT STARTING MINERALS TO CONTAIN MASS, YES=1, NO=0. 1, SPECIFY NUMBER OF MINERALS TO HAVE INITIAL MASS. 61 10.000, SPECIFY MINERAL NUMBER AND MASS(MOLES).

### (C). NUANCES.txt

0, WANT TO REDUCE TEMPERATURE DECREMENT AT LOWER T? YES=1, NO=0.

263.15 1.0, TEMPERATURE AND ☑T, WHERE CHANGE SHOULD BE MADE.

259.15 0.1, TEMPERATURE AND ☑T, WHERE CHANGE SHOULD BE MADE.

0, WANT TO REDUCE WATER DECREMENT AT LOWER WATER? YES=1, NO=0.

100 1, WATER CONTENT AND ☑H2O, WHERE CHANGE SHOULD BE MADE.

10 0.1, WATER CONTENT AND ☑H2O, WHERE CHANGE SHOULD BE MADE.

0, WANT TO INCREASE(OR DECREASE) PRESSURE INCREMENT AT HIGHER P? YES = 1, N=0.

17 1, PRESSURE AND ☑P, WHERE CHANGE SHOULD BE MADE.

50 10, PRESSURE AND ☑P, WHERE CHANGE SHOULD BE MADE.

## **Model Outputs.**

"Ion.Str." is the ionic strength of the equilibrium solution (see Table 6). "RHO" is the density of the solution. "Phi" is the osmotic coefficient of the equilibrium solution. "H2O(g)" is the amount of water remaining as liquid. "Ice" is the amount of water that formed ice between 268.15 K and 263.15 K. The total ice that formed between 273 and 263 is 519.24 g (28.822 x 18.0153). The mass basis for calculation in the model is 1.0 kg of water (except for evaporation); therefore, the water in liquid water + ice + hydrated salts should always sum to 1.0 kg. The data under "Initial Conc." are the input concentrations at 298 K (Table 3). "Final Conc." are the equilibrium concentrations at 263 K. Act. coef. (activity coefficient) and activity are selfexplanatory. Moles are the # of moles in the solution phase. For the major constituents, the "Mass Balance" column should generally agree with the input column ("Initial Conc."); this is the best check on the internal consistency of the calculations. The "Accum Moles" in the solids section are the net # of moles of that solid that have precipitated. For equilibrium crystallization, accum. moles = moles(solid). For fractional crystallization accum. moles ≥ moles (solids); in this "ice" case, moles represent the solids that have precipitated in the last interval (e.g., between 268 K and 263 K), while accum. moles represent the total precipitate of ice (e.g., between 273 K and 263 K).

Table 6 is a case where we examined calcium/magnesium/chloride/sulfate/sulfate, and nitrogen gas systems for Mars (Marion et al., 2013) (see the Input.txt file in Table 3). During the freezing process from 273 K to 263 K, 51.9 % of the original water at 298 K precipitated as ice. Also, the ionic strength changed from 3.32 m at 298 K to 5.36 m at 263 K (Table 6). The pH of

the system that began at 8.12 at 298 K changed to 7.30 at 263 K. During this 35 K drop in temperature, ice, MgSO<sub>4</sub>•11H<sub>2</sub>O, CaSO<sub>3</sub>•0.5H<sub>2</sub>O, and MgSO<sub>3</sub>•6H<sub>2</sub>O precipitated (Table 6).

Table 7 is a case where we converted molar into molal concentrations. In the upper table there are three columns labeled rho, SA, and CF that represent model calculated estimates of density (kg(soln.)/liter or g/cm $^3$ ), absolute salinity [g salt/kg(soln.)], and the conversion factor [liters/kg(H $_2$ O)]. The iterations quickly converted molar concentrations ( 5.4170 moles/liter under Initial Conc.) into molal concentrations [6.1458 moles/kg(H $_2$ O) under Final Conc.]. In addition to inputs of molar concentrations, this algorithm also requires salinity on a liter basis(SL)(see Table 3). In this case, the SL value is 316.57 g/liter [= 5.417 x 58.44 (molecular weight of NaCl)]. So even with no prior knowledge about solution density (the model arbitrarily assigns initial density = 1.00 g/cm $^3$ ), we were able to quickly calculate density and convert molar to molal concentrations. In turn, molal concentrations could be directly imported into FREZCHEM to explore geochemical processes. Note that all the potential solid phases were assigned high solubility products to prevent their precipitation (Table 7). FORTRAN model inputs to accomplish this negation of solid phases are in the SOLIDPHASE.txt file in Table 5 and must be implemented by the user. See the previously cited Marion (2007) paper for a fuller discussion of the techniques used in this algorithm.

Table 8 is an input file for a Titan simulation from 273 K to 173 K that is dominated with NH<sub>3</sub>(aq), NH<sub>4</sub>(aq), and CH<sub>4</sub>(g). Table 9 is the output of this simulation after the initial temperature at 273 K dropped to 173 K. At 173 K, ice had formed and methane hydrate, NH<sub>4</sub>Cl, and (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> had precipitated (Table 9). NH<sub>3</sub>(aq), which started at 10.0 molal, has risen to 36.27 molal that is approaching the eutectic where NH<sub>3</sub>•2H<sub>2</sub>O would precipitate. In this case, the pH started at 10.0 (Table 8) and rose to 18.03 (Table 9). The latter may not be accurate.

Table 10 is an input file for a Titan simulation from 273 to 173 K that is dominated with  $NH_3(aq)$ ,  $NH_4(aq)$ ,  $N_2(bars)$ , and  $CH_4(bars)$ . Table 11 is the output of this simulation after the initial temperature at 273 K dropped to 173 K. At 173 K, ice had formed,  $NH_4Cl$  and  $(NH_4)_2SO_4$  had precipitated, and  $N_2 \cdot 6H_2O \cdot CH_4 \cdot 6H_2O$  had formed.  $NH_3(aq)$ , which started at 10.0 molal, has risen to 30.34 molal. In this case, the pH started at 10.0 (Table 10) and rose to 17.89. These Tables 10-11 are very similar to Tables 8-9, except that a mixed gas hydrate formed in this case and methane hydrate formed in the previous case.

TABLE 6. TITLE. This is a SO2-H2S model test case (Version 16).

Temp(K)	Ion.Str.	RHO	Phi	H2O(g)	Ice(g)	Press.(bars)
263.15	5.3606	1.138740	1.2185	432.15	325.63	1.0000
Solution	Initial	Final				Mass
SPECIES	Conc.	Conc.	Act.Coef.	Activity	Moles	Balance
CA	0.10000	0.31816E-02	0.76184E-01	0.24239E-03	0.13749E-02	2 0.10000
MG	1.0000	1.6262	0.14589	0.23726	0.70278	0.98776
H	0.00000	0.47007E-07	1.0699	0.50291E-07	0.20314E-0	7
CL	1.0000	2.3140	1.4452	3.3441	1.0000	1.0000
SO4	0.40000	0.46419	0.49115E-01	0.22799E-01	0.20060	0.38838
HSO4	0.00000	0.22046E-07	1.9580	0.43165E-07	0.95271E-08	3
HSO3	0.40000	0.44125E-04	2.5684	0.11333E-03	0.19069E-0	1 0.39876

SO3	0.00000	0.8190	2E-02	0.54583E	-01	0.44704E-03	0.35394E-02	
SO2	0.00000	0.7618	2E-10	2.3544		0.17937E-09	0.32922E-10	
SO2(BAR)	0.00000	0.1253	3E-10	1.0000		0.12533E-10		
N2	0.00000	0.8978		2.1884		0.19649E-04	0.38802E-05	
N2(BAR)	0.78000	0.7800	0	0.98983		0.77206	0.00000	
H2O(BAR)	0.00000					0.25954E-02		
H2O(L)	55.508					0.90762	23.988	55.508
Solid			Equ	<b>i</b> 1	7.	ccum.		
SPECIES		Moles	_	stant		oles		
ICE		18.075	0.90			.822		
NACL.2H2O		0.00000	12.			0000		
NACL		0.00000	27.			0000		
KCL		0.00000	2.5			0000		
CACL2.6H2O		0.00000	125	2.1	0.0	0000		
MGCL2.6H2O		0.00000	554	13.	0.0	0000		
MGCL2.8H2O		0.00000	468	2.9	0.0	0000		
MGCL2.12H2O	1	0.00000	199	.88	0.0	0000		
KMGCL3.6H2O		0.00000	452	6.5	0.0	0000		
CACL2.2MGCL	2.12H2O	0.00000	0.11	069E+20	0.0	0000		
NA2SO4.10H2	0	0.00000	0.90	535E-03	0.0	0000		
NA2SO4		0.00000	0.48			0000		
MGSO4.6H2O		0.00000		596E-01		0000		
MGSO4.7H2O		0.00000		651E-02		0000		
K2SO4		0.00000		944E-02		0000		
MGSO4.K2SO4		0.00000		354E-05		0000		
NA2SO4.MGSO CASO4.2H2O	4.4HZO	0.00000		859E-02		0000		
CASO4		0.00000		490E-04 047E-04		0000 0000		
MGSO4.11H2O		0.18778		624E-02		8778		
NA2SO4.3K2S		0.00000		926E-09		0000		
CACO3(CALCI		0.00000		032E-08		0000		
MGCO3	,	0.00000		260E-07		0000		
MGC03.3H20		0.00000	0.15	228E-04	0.0	0000		
MGCO3.5H2O		0.00000	0.96	136E-05	0.0	0000		
CACO3.6H2O		0.00000	0.32	915E-07	0.0	0000		
NAHCO3		0.00000	0.14	566	0.0	0000		
NA2CO3.10H2		0.00000		292E-02		0000		
NAHCO3.NA2C		0.00000		154E-01		0000		
3MGCO3.MG(O	н)2.3н20	0.00000		729E-34		0000		
CAMG(CO3)2		0.00000		737E-16		0000		
NA2CO3.7H2O	1	0.00000		253E-01		0000		
KHCO3 CACO3 (ARAGO	NITUE	0.00000	0.50	021 984E-08		0000 0000		
CACO3 (VATER	•	0.00000		193E-07		0000		
HNO3.3H2O	1111)	0.00000	347			0000		
KNO3		0.00000		268E-01		0000		
NANO3		0.00000	2.3			0000		
HCL.3H2O		0.00000	124			0000		
H2SO4.6.5H2	0	0.00000	22.	485	0.0	0000		
H2SO4.4H2O		0.00000	999	.88	0.0	0000		
HCL.6H2O		0.00000	100	0.0	0.0	0000		
NANO3.NA2SO	4.2H2O	0.00000	0.84	020E-01	0.0	0000		
NA3H(SO4)2		0.00000	0.14	086	0.0	0000		
NAHSO4.H2O		0.00000	30.			0000		
K3H(SO4)2		0.00000		417E-04		0000		
K5H3(SO4)4		0.00000		412E-07		0000		
K8H6(SO4)7.	H2U	0.00000		716E-12		0000		
KHSO4		0.00000	0.97			0000		
MGSO4.H2O FESO4.7H2O		0.00000	11.	328 913E-02		0000 0000		
FESO4.7H20		0.00000	0.13			0000		
FECL2.6H2O		0.00000	398			0000		
FECL2.4H2O		0.00000	177			0000		
FECO3		0.00000		199E-10		0000		
FE(OH)3		0.00000		277E+06		0000		
•								

```
0.00000
CO2.6H2O
                                   3.8247
                                               0.00000
                     0.00000
CH4.6H2O
                                   9.8303
                                               0.00000
FECL3.10H20
                     0.00000
                                  0.67444E-02 0.00000
                     0.00000
                                  0.81058E-01 0.00000
FECL3.6H2O
FECL3.2KCL.H20
                     0.00000
                                  0.34623
                                               0.00000
FE2(SO4)3
                     0.00000
                                   17465.
                                               0.00000
                                  0.14131E-13 0.00000
FE2(SO4)3.2K2SO4.14H 0.00000
                                  0.26654E-05 0.00000
K2SO4.FESO4.6H2O
                     0.00000
                                  0.12040E-02 0.00000
NA2SO4.FESO4.4H2O
                     0.00000
FE2(SO4)3.9H2O
                     0.00000
                                  1.2561
                                               0.00000
                                  1130.8
FE2(SO4)3.H2SO4.8H2O 0.00000
                                               0.00000
                                  0.62064E-09 0.00000
KFE3(SO4)2(OH)6
                     0.00000
NAFE3(SO4)2(OH)6
                     0.00000
                                  0.16518E-02 0.00000
H3OFE3(SO4)2(OH)6
                     0.00000
                                  0.17876
                                               0.00000
a-FE203
                     0.00000
                                  1273.5
                                               0.00000
a-FEO(OH)
                     0.00000
                                  43.379
                                               0.00000
g-FEO(OH)
                     0.00000
                                  3733.3
                                               0.00000
FEO(OH)3/4(SO4)1/8
                     0.00000
                                   735.54
                                               0.00000
                                  0.20471E-01 0.00000
FESO4.4H2O
                     0.00000
FE2(SO4)3.7H2O
                     0.00000
                                   9.9750
                                               0.00000
FE(II)FE(III)4(SO4)6 0.00000
                                  0.26295E-18 0.00000
                     0.00000
                                  0.36333E-18 0.00000
FE(III)5(SO4)6O(OH).
FE(II)FE(III)2(SO4)4 0.00000
                                  0.21801E-16 0.00000
FE(II)FE(III)2(SO4)4 0.00000
                                  0.34750E-09 0.00000
                                  0.30995E-30 0.00000
K2FE(II)5FE(III)4(SO 0.00000
ALCL3.6H2O
                     0.00000
                                   12997.
                                               0.00000
                                  0.10093E-05 0.00000
AL2(SO4)3.17H2O
                     0.00000
NABR
                     0.00000
                                  0.00000
                                               0.00000
MGBR2
                     0.00000
                                  0.00000
                                               0.00000
AL(OH)3
                     0.00000
                                  0.24779E+11 0.00000
                                  0.27437E-04 0.00000
SIO2(QUARTZ)
                     0.00000
SIO2 (AMORPHOUS)
                     0.00000
                                  0.90848E-03 0.00000
                     0.00000
                                  0.53769E+06 0.00000
KAL3(SO4)2(OH)6
NAAL3(SO4)2(OH)6
                     0.00000
                                  0.13263E+10 0.00000
                                  0.41764E-07 0.00000
                     0.00000
KAL(SO4)2.12H2O
NAAL(SO4)2.12H2O
                     0.00000
                                  0.22928E-04 0.00000
FESO4.AL2(SO4)3.22H2 0.00000
                                  0.27651E-08 0.00000
AL2SI2O5(OH)4
                     0.00000
                                  0.46063E+10 0.00000
MGSO4.AL2(SO4).22H2O
                     0.00000
                                  0.96694E-08 0.00000
NACLO4.H2O
                     0.00000
                                   62.357
                                  0.24262E+06 0.00000
MG(CLO4)2.8H2O
                     0.00000
CA(CLO4)2.6H2O
                     0.00000
                                  0.84562E+06 0.00000
                     0.00000
                                  0.91970E-03 0.00000
MG(CLO4)2.6H2O
                     0.00000
                                  0.10000E+31 0.00000
                     0.00000
NACLO4.2H2O
                                  35.934
                                               0.00000
NH4CL
                     0.00000
                                   5.7644
                                               0.00000
NH42SO4
                     0.00000
                                  0.45087
                                               0.00000
NH3.H20
                     0.00000
                                  0.39362E+09 0.00000
                     0.00000
NH4NO3
                                  2.9136
                                               0.00000
                                  0.29822
NH4HCO3
                     0.00000
                                               0.00000
NH4CLO4
                     0.00000
                                  0.11862
                                               0.00000
NH3.2H2O
                     0.00000
                                  370.58
                                               0.00000
K2SO3
                     0.00000
                                   42.371
                                               0.00000
NA2SO3.7H2O
                     0.00000
                                  0.28386E-01
                                               0.00000
CASO3.0.5H2O
                     0.00000
                                  0.35972E-06 0.98625E-01
MGSO3.6H2O
                     0.84464E-02 0.59290E-04 0.97205E-01
FESO3.5H2O
                     0.00000
                                  0.15372E-05 0.00000
(NH4)2SO3.H2O
                     0.00000
                                  0.37862
                                               0.00000
                     0.00000
                                  0.39798E-16 0.00000
FES2
N2.6H2O
                     0.00000
                                   59.688
                                               0.00000
CASO4.0.5H2O
                     0.00000
                                  0.55003E-03 0.00000
```

pH= 7.29851 pHF= 7.32784 pHT= 7.16082 pH(SWS)= 7.16082 pHMacinnis = 7.82664 Temp. = 263.150

CONVERGENCE CRITERION = 0.100000. Iterations = 10

Table 7. Conversion of a molar NaCl solution into a molal concentration.

1.20181 20 1.19726 20 1.19812 20 1.19796 20 1.19799 20 1.19798 20 1.19798 20	53.411 1.1 54.411 1.1 54.223 1.1 54.259 1.1 64.252 1.1 64.253 1.1 64.253 1.1	6086 2964 3547 3437 3458 3454 3455 3454 3455				
Temp(K) 298.15	Ion.Str. 6.1458	RHO 1.19798	Phi 1.2846	H2O(g) 1000.0	Ice(g) 0.00000	Press.(bars) 1.0132
Solution SPECIES NA CL H2O(BAR)	Initial Conc. 5.4170 5.4170 0.24948E-01 55.508	Final Conc. 6.1458 6.1458	Act.Coe 1.0109 1.0109	f. Activity 6.2128 6.2128 .23840E-01 .75242	Moles 6.1458 6.1458	Mass Balance 6.1458 6.1458
Solid SPECIES ICE NACL.2H2O NACL KCL CACL2.6H2O MGCL2.6H2O MGCL2.8H2O MGCL2.12H2O KMGCL3.6H2O CACL2.2MGCL2 NA2SO4.10H2O NA2SO4 MGSO4.6H2O MGSO4.7H2O K2SO4 MGSO4.8E3O4 CASO4 MGSO4.2H2O CASO4 MGSO4.3H2O CACO3(CALCITI MGCO3 MGCO3.3H2O MGCO3.5H2O CACO3.6H2O NAHCO3 NA2CO3.10H2O NAHCO3 NACO3.7H2O KACO3 CACO3(CATENT MGCO3 MGCO3.7H2O KACO3 CACO3(CATENT MGCO3 NA2CO3.7H2O KHCO3 CACO3(VATERIT HNO3.3H2O KNO3	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	Moles .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000 .00000	Equil. Constant 0.10000E+31 0.1000E+31 0.10000E+31	Accum. Moles 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000 0.00000		

NANO3	0.00000	0.10000E+31	0.00000
HCL.3H2O	0.00000	0.10000E+31	0.00000
H2SO4.6.5H2O	0.00000	0.10000E+31	0.00000
H2SO4.4H2O	0.00000	0.10000E+31	0.00000
HCL.6H2O	0.00000	0.10000E+31	0.00000
NANO3.NA2SO4.2H2O	0.00000	0.10000E+31	0.00000
NA3H(SO4)2	0.00000	0.10000E+31	0.00000
NAHSO4.H2O	0.00000	0.10000E+31	0.00000
K3H(SO4)2	0.00000	0.10000E+31	0.00000
K5H3(SO4)4	0.00000	0.10000E+31	0.00000
K8H6(SO4)7.H2O	0.00000	0.10000E+31	0.00000
KHS04	0.00000	0.10000E+31	0.00000
MGSO4.H2O	0.00000	0.10000E+31	0.00000
FESO4.7H2O	0.00000	0.10000E+31	0.00000
FESO4.71120	0.00000	0.10000E+31	0.00000
FECL2.6H2O	0.00000	0.10000E+31	0.00000
FECL2.4H2O	0.00000	0.10000E+31	0.00000
FECO3	0.00000	0.10000E+31	0.00000
FE(OH)3	0.00000	0.10000E+31	0.00000
CO2.6H2O	0.00000	0.10000E+31	0.00000
CH4.6H2O	0.00000	0.10000E+31	0.00000
FECL3.10H20	0.00000	0.10000E+31	0.00000
FECL3.6H2O	0.00000	0.10000E+31	0.00000
FECL3.2KCL.H20	0.00000	0.10000E+31	0.00000
FE2(SO4)3	0.00000	0.10000E+31	0.00000
FE2(SO4)3.2K2SO4.14H	0.00000	0.10000E+31	0.00000
K2SO4.FESO4.6H2O	0.00000	0.10000E+31	0.00000
NA2SO4.FESO4.4H2O	0.00000	0.10000E+31	0.00000
FE2(SO4)3.9H2O	0.00000	0.10000E+31	0.00000
FE2(SO4)3.H2SO4.8H2O	0.00000	0.10000E+31	0.00000
KFE3(SO4)2(OH)6	0.00000	0.10000E+31	0.00000
NAFE3(SO4)2(OH)6	0.00000	0.10000E+31	0.00000
H3OFE3(SO4)2(OH)6	0.00000	0.10000E+31	0.00000
a-FE2O3	0.00000	0.10000E+31	0.00000
a-FEO(OH)	0.00000	0.10000E+31	0.00000
g-FEO(OH)	0.00000	0.10000E+31	0.00000
FEO(OH)3/4(SO4)1/8	0.00000	0.10000E+31	0.00000
FESO4.4H2O	0.00000	0.10000E+31	0.00000
FE2(SO4)3.7H2O	0.00000	0.10000E+31	0.00000
FE(II)FE(III)4(SO4)6	0.00000	0.10000E+31	0.00000
FE(III)5(SO4)6O(OH).	0.00000	0.10000E+31	0.00000
FE(II)FE(III)2(SO4)4	0.00000	0.10000E+31	0.00000
FE(II)FE(III)2(SO4)4	0.00000	0.10000E+31	0.00000
K2FE(II)5FE(III)4(SO	0.00000	0.10000E+31	0.00000
ALCL3.6H2O	0.00000	0.10000E+31	0.00000
AL2(SO4)3.17H2O	0.00000	0.10000E+31	0.00000
NABR	0.00000	0.10000E+31	0.00000
MGBR2	0.00000	0.10000E+31	0.00000
AL(OH)3	0.00000	0.10000E+31	0.00000
SIO2(QUARTZ)	0.00000	0.10000E+31	0.00000
SIO2(AMORPHOUS)	0.00000	0.10000E+31	0.00000
KAL3(SO4)2(OH)6	0.00000	0.10000E+31	0.00000
NAAL3(SO4)2(OH)6	0.00000	0.10000E+31	0.00000
KAL(SO4)2.12H2O	0.00000	0.10000E+31	0.00000
NAAL(SO4)2.12H2O	0.00000	0.10000E+31	0.00000
FESO4.AL2(SO4)3.22H2	0.00000	0.10000E+31	0.00000
AL2SI2O5(OH)4	0.00000	0.10000E+31	0.00000
MGSO4.AL2(SO4).22H2O	0.00000	0.10000E+31	0.00000
` '			

Iterations = 9

Table 8. A Titan simulation. This is the main input file for applications of

# FREZCHEM. In this particular case, the model simulates a NH3-NH4 chemistry from 273.15 K to 173.15 K (Table 9)(Icarus, 2012, 220:932-946) (Version 15).

```
TITLE: This is a Titan NH3-NH4 case.
1, FREEZE(1) OR EVAPORATION(2) OR PRESSURE(3) SCENARIO? CALLED PATH BELOW.
2, EQUILIBRIUM(1) OR FRACTIONAL(2) CRYSTALLIZATION?
1, OPEN(1) OR CLOSED(2) CARBON SYSTEM?
0, WANT SEAWATER SALINITY(Sp) TO GOVERN THE CALCULATIONS,Y=1,N=0.
0.0, IF YES ABOVE, ENTER SEAWATER PRACTICAL SALINITY(Sp) OR 0.0.
0, WANT SEAWATER CARBONATE SUPERSATURATION TO BE CONSIDERED? Y=1,N=0.
0.00, SODIUM(M/KG).
0.00, POTASSIUM(M/KG).
0.00, CALCIUM(M/KG).
0.00, MAGNESIUM(M/KG).
0.00, STRONTIUM(M/KG).
0.00, FERROUS IRON(M/KG).
0.00, FERRIC IRON (M/KG).
0.00, ALUMINUM(M/KG).
0.00, SILICA (M/KG).
3.00, AMMONIUM(M/KG).
FOR FE,AL,SI, AND ALKALINE CHEMISTRIES,DO YOU WANT ACIDITY
IGNORED(1), OR FIXED BY PH(2), OR ACIDITY(3), OR
ALKALINITY(4)? IF YES, THEN ENTER NUM. NOTE THAT OPTIONS 3
AND 4 REQUIRE A FURTHER SPECIFICATION BELOW. OPTIONS 2-4
WILL ADJUST SOLUTION-PHASE CHARGE BALANCE AS AL, FE, OR SI
REACTIONS PRODUCE ACIDITY BY ASSUMING H+ REACTS WITH ROCKS
TO RELEASE NA,K,CA,MG,OR FE(II) IONS. SOME OF THE LATTER
IONS MUST BE PRESENT AS INPUT TO SERVE AS AN ION SINK/SOURCE.
FOR NH3(AQ) + NH4(AQ) CASE, SET OPTION = 2, WITH PH = 10.0.
2, SPECIFY ABOVE ACIDITY OPTION.
10.00, SPECIFY INITIAL PH.
1.00, CHLORIDE(M/KG).
0.00, BROMIDE(M/KG).
0.00, PERCHLORATE(M/KG).
1.00, SULFATE(M/KG).
0.00, NITRATE(M/KG).
0.00, ALKALINITY(EQUIVALENTS/KG).
0.00, ACIDITY(EQUIVALENTS/KG).
0.00, IF YOU WANT TO SPECIFY HCL(BARS), ENTER VALUE HERE.
0.00, IF YOU WANT TO SPECIFY HNO3(BARS), ENTER VALUE HERE.
0.00, IF YOU WANT TO SPECIFY H2SO4(BARS), ENTER VALUE HERE.
0.00, BORON (M/KG).
0.00, FLUORIDE(M/KG).
10.0, INITIAL TOTAL PRESSURE(BARS).
0.00, INITIAL CO2(BARS).
0.0, ENTER MOLE FRACTION OF CO2, 0=FIXED CO2, 1=PURE CO2.
0.00, INITIAL O2(BARS).
5.00, INITIAL CH4(BARS).
0.0, ENTER MOLE FRACTION OF CH4, 0=FIXED CHR, 1=PURE CH4.
0, CONSIDER A MIXED CO2-CH4 GAS HYDRATE(YES=1, NO=0)?
0.00, INITIAL NH3(G)(BARS), DO NOT INCLUDE BOTH NH3 INPUTS.
10.00, INITIAL NH3(AQ)(M/KG), DO NOT INCLUDE BOTH NH3 INPUTS.
0, MOLAR TO MOLAL CONVERSION? YES=1,NO=0.
0.0, IF YES ABOVE, ENTER SALINITY(G)/LITER.
273.15, INITIAL TEMPERATURE(K).
173.15, FINAL TEMPERATURE(K), IF PATH =1, OTHERWISE, SET = 0.
5.00, TEMPERATURE DECREMENT(K), IF PATH = 1, OTHERWISE, SET = 0.
1000, INITIAL WATER(G), IF PATH = 2, OTHERWISE, SET = 1000.
0.00, FINAL WATER (G), IF PATH = 2, OTHERWISE, SET = 0.
```

0, WATER DECREMENT(K), IF PATH = 2, OTHERWISE, SET = 0. 0, FINAL PRESSURE(BARS), IF PATH = 3, OTHERWISE, SET = 0. 0, PRESSURE INCREMENT(BARS), IF PATH = 3, OTHERWISE, SET = 0.

Table 9. A Titan simulation (Icarus, 2012. 220:932-946).

TITLE: This is a Titan NH3-NH4 case.

Temp(K)	Ion.Str	. RHO		Phi	н20	D(g)	Ice(g)	Press.(bars)
173.15	1.3862	0.8759973		1.4671		5.69	0.00000	10.000
Solution	Initial	Final						Mass
SPECIES	Conc.	Conc.		Act.Coe	f. Act	tivity	Moles	Balance
Н	0.00000	0.77718E	-18	1.1999	0.93	3250E-18	0.21426E-18	
NH4	3.0000	1.3861		0.40767	0.50	5509	0.38215	3.0000
CL	1.0000	1.3861		0.40768	0.50	5507	0.38213	1.0000
SO4	1.0000	0.41533E	-04	909.86		7789E-01	0.11450E-04	1.0000
ОН	0.00000	0.72284E	80-	884.69	0.63	3948E-05	0.19928E-08	
HSO4	0.00000	0.38913E	-16	0.56845	0.22	2120E-16	0.10728E-16	
NH3	10.000	36.272		3.5713	129	9.54	10.000	10.000
NH3(BAR)	0.00000	0.11087E	-03	1.0000	0.1	1087E-03	0.00000	
CH4	0.00000	9.1645		1.0000	9.1	1645	0.00000	
CH4(BAR)	0.00000	5.0000		0.92416	4.0	6208	0.00000	
H2O(BAR)	0.00000				. 92	2816E-08		
H2O(L)	55.508				.2	7966	15.303	55.509
` ,								
Solid			Equ	ıil.	Accum			
SPECIES		Moles	Con	nstant	Moles			
ICE		0.00000	0.45	5268	18.629			
NACL.2H2O		0.00000	81.	415	0.00000			
NACL		0.00000	0.84	1048E-02	0.00000			
KCL		0.00000	0.65	5448E-02	0.00000			
CACL2.6H2O		0.00000	0.21	L833E-05	0.00000			
MGCL2.6H2O		0.00000	30.	.019	0.00000			
MGCL2.8H2O		0.00000	0.54	1878E-03	0.00000			
MGCL2.12H20	)	0.00000	0.10	0669E-33	0.00000			
KMGCL3.6H2C	)	0.00000	0.39	9141E-01	0.00000			
CACL2.2MGCI	L2.12H2O	0.00000	0.34	1336E+25	0.00000			
NA2SO4.10H2	20	0.00000	0.65	083E-08	0.00000			
NA2SO4		0.00000	0.44	1571	0.00000			
MGSO4.6H2O		0.00000		0811E-01	0.00000			
MGSO4.7H2O		0.00000		5567E-04	0.00000			
K2SO4		0.00000		7467E-05	0.00000			
MGSO4.K2SO4		0.00000		2453E-11	0.00000			
NA2SO4 MGSC	04.4H2O	0.00000		3068E-02	0.00000			
CASO4.2H2O CASO4		0.00000		351E-07	0.00000			
MGSO4.11H20	<b>.</b>	0.00000		1270E-05 3729E-06	0.00000			
NA2SO4.11H2C		0.00000		7521E-15	0.00000			
CACO3(CALCI		0.00000		9496E-08	0.00000			
MGCO3	111)	0.00000		3880E-04	0.00000			
MGC03.3H20		0.00000		9706	0.00000			
MGC03.5H20		0.00000		3181E+22	0.00000			
CACO3.6H2O		0.00000		7849E-11	0.00000			
NAHCO3		0.00000		470	0.00000			
NA2CO3.10H2	20	0.00000		854E-04	0.00000			
NAHCO3.NA20	СО3.2Н2О	0.00000	0.45	061E-02	0.00000			
3MGCO3.MG(C	ОН)2.3H2O	0.00000	0.86	5235E-23	0.00000			
CAMG(CO3)2		0.00000	0.98	3928E-12	0.00000			
NA2CO3.7H2C	)	0.00000	0.31	1124E-03	0.00000			
KHCO3		0.00000	0.42	2063E-01	0.00000			
CACO3 (ARAGO	ONITE)	0.00000	0.38	3711E-08	0.00000			
CACO3 (VATER	RITE)	0.00000	0.26	699E-07	0.00000			
HNO3.3H2O		0.00000	0.62	2103	0.00000			
KNO3		0.00000	0.51	L379E-05	0.00000			
NANO3		0.00000	0.15	5269E+11	0.00000			
HCL.3H2O		0.00000		94.2	0.00000			
H2SO4.6.5H2	20	0.00000	0.42	2082E-02	0.00000			

H2SO4.4H2O	0.00000	0.25389	0.00000
HCL.6H2O	0.00000	146.93	0.00000
NANO3.NA2SO4.2H2O	0.00000	0.58559E+06	0.00000
NA3H(SO4)2	0.00000	0.93785E-49	0.00000
NAHSO4.H2O	0.00000	0.60938E+18	0.00000
K3H(SO4)2	0.00000	0.15135E-01	0.00000
K5H3(SO4)4	0.00000	0.41467E-10	0.00000
K8H6(SO4)7.H2O	0.00000	0.43211E-06	0.00000
KHSO4	0.00000	0.31505E-01	0.00000
MGSO4.H2O	0.00000	0.90352E+11	0.00000
FESO4.7H2O	0.00000	0.84993E-02	0.00000
FESO4.H2O	0.00000	4.8511	0.00000
FECL2.6H2O	0.00000	81.034	0.00000
FECL2.4H2O	0.00000	0.10524E+07	0.00000
FECO3	0.00000	0.19735E-09	0.00000
FE(OH)3	0.00000	0.36265E+14	0.00000
CO2.6H2O	0.00000	0.40474E-03	0.00000
CH4.6H2O	0.48527E-01	0.22105E-02	3.5960
FECL3.10H20	0.00000	0.43891E-12	
			0.00000
FECL3.6H2O	0.00000	0.16793	0.00000
FECL3.2KCL.H20	0.00000	0.81402E-14	0.00000
FE2(SO4)3	0.00000	0.22676E+30	0.00000
FE2(SO4)3.2K2SO4.14H	0.00000	0.19306E-13	0.00000
K2SO4.FESO4.6H2O	0.00000	0.96795E-09	0.00000
NA2SO4.FESO4.4H2O	0.00000	0.14419E-02	0.00000
FE2(SO4)3.9H2O	0.00000	0.13795E+12	0.00000
FE2(SO4)3.H2SO4.8H2O	0.00000	0.37146E+20	0.00000
KFE3(SO4)2(OH)6	0.00000	56.527	0.00000
NAFE3(SO4)2(OH)6	0.00000	0.53531E+12	0.00000
H3OFE3(SO4)2(OH)6	0.00000	0.58130E+23	0.00000
a-FE2O3	0.00000	0.42230E+17	0.00000
a-FEO(OH)	0.00000	0.10846E+09	0.00000
g-FEO(OH)	0.00000	0.10294E+12	0.00000
FEO(OH)3/4(SO4)1/8	0.00000	0.26660E+11	0.00000
FESO4.4H2O	0.00000	0.92593E-02	0.00000
FE2(SO4)3.7H2O	0.00000	0.13315E+16	0.00000
FE(II)FE(III)4(SO4)6	0.00000	1869.1	0.00000
FE(III)5(SO4)6O(OH).	0.00000	0.10855E+06	0.00000
FE(II)FE(III)2(SO4)4	0.00000	0.79320E-17	0.00000
FE(II)FE(III)2(SO4)4	0.00000	7.9464	0.00000
K2FE(II)5FE(III)4(SO	0.00000	0.34573E+07	0.00000
ALCL3.6H2O	0.00000	0.29269E-07	0.00000
	0.00000	0.69490E-08	0.00000
AL2(SO4)3.17H2O	0.00000		
NABR		0.00000	0.00000
MGBR2	0.00000	0.00000	0.00000
AL(OH)3	0.00000	0.18472E+22	0.00000
SIO2(QUARTZ)	0.00000	0.75777E-07	0.00000
SIO2(AMORPHOUS)	0.00000	0.32652E-04	0.00000
KAL3(SO4)2(OH)6	0.00000	0.45047E+30	0.00000
NAAL3(SO4)2(OH)6	0.00000	0.54098E+36	0.00000
KAL(SO4)2.12H2O	0.00000	0.19265E-10	0.00000
NAAL(SO4)2.12H2O	0.00000	0.25714E-05	0.00000
FESO4.AL2(SO4)3.22H2	0.00000	0.37090E-08	0.00000
AL2SI2O5(OH)4	0.00000	0.42725E+27	0.00000
MGSO4.AL2(SO4).22H2O	0.00000	0.12873E-07	0.00000
NACLO4.H2O	0.00000	0.22603	0.00000
MG(CLO4)2.8H2O	0.00000	0.54775E+07	0.00000
CA(CLO4)2.6H2O	0.00000	9140.1	0.00000
KCLO4	0.00000	0.23292E-04	0.00000
MG(CLO4)2.6H2O	0.00000	0.10053E+31	0.00000
NACLO4.2H2O	0.00000	0.42354	0.00000
NH4CL	0.42506E-01	0.31928	0.61787
NH42SO4	0.10009E-04	0.12067E-01	0.99999
NH3.H2O	0.00000	40.353	0.00000
NH4NO3	0.00000	0.30043E-01	0.00000
NH4HCO3	0.00000	0.33153E-03	0.00000
111000	0.0000	0.33133E-03	3.00000

```
NH4CLO4
                     0.00000
                                 0.49215E-04 0.00000
                     0.00000
NH3.2H2O
                                 15,150
                                             0.00000
pH= 18.0303 pHF= 18.1095 pHT= 16.4013 pH(SWS)= 16.4013
pHMacinnis = 18.2358 Temp. = 173.150
CONVERGENCE CRITERION = 0.100000 %
Iterations = 263
```

Table 10. A Titan simulation. This is the main input for applications of FREZCHEM. In this particular case, the model simulations a N2-CH4 chemistry from

```
273.15 to 173.15 K(Table 11)(Icarus, 2014, 226:1-8)(Version 16).
TITLE: SO2-H2S model parameters
1, FREEZE(1) OR EVAPORATION(2) OR PRESSURE(3) SCENARIO ? CALLED PATH BELOW.
2, EQUILIBRIUM(1) OR FRACTIONAL(2) CRYSTALLIZATION?
0, WANT SEAWATER SALINITY(Sp) TO GOVERN THE CALCULATIONS, Y=1, N=0.
0.0, IF YES ABOVE, ENTER SEAWATER PRACTICAL SALINITY(Sp) OR 0.0.
0, WANT SEAWATER CARBONATE SUPERSATURATION TO BE CONSIDERED? Y=1, N=0.
0.00 SODIUM(M/KG).
0.00, POTASSIUM(M/KG).
0.00, CALCIUM(M/KG
0.00, MAGNESIUM(M/KG)
0.00, STRONTIUM(M/KG).
0.00, FERROUS IRON(M/KG).
0.00, FERRIC IRON (M/KG).
0.00, ALUMINUM(M/KG).
0.00, SILICA (M/KG).
3.00, AMMONIUM(M/KG).
FOR FE, AL, SI, AND ALKALINE CHEMISTRIES, DO YOU WANT ACIDITY
IGNORED(1),OR FIXED BY PH(2), OR ACIDITY(3), OR
ALKALINITY(4)? IF YES, THEN ENTER NUM. NOTE THAT OPTIONS 3
AND 4 REQUIRE A FURTHER SPECIFICATION BELOW. OPTIONS 2-4
WILL ADJUST SOLUTION-PHASE CHARGE BALANCE AS AL, FE, OR SI
REACTIONS PRODUCE ACIDITY BY ASSUMING H+ REACTS WITH ROCKS
TO RELEASE NA, K, CA, MG, OR FE(II) IONS. SOME OF THE LATTER
IONS MUST BE PRESENT AS INPUT TO SERVE AS AN ION SINK/SOURCE.
2, SPECIFY ABOVE ACIDITY OPTION.
10.0, SPECIFY INITIAL PH.
1.00, CHLORIDE(M/KG).
0.00, BROMIDE(M/KG).
0.00, PERCHLORATE(M/KG).
1.00, SULFATE(M/KG).
0.00, NITRATE(M/KG).
0.00, CARBON ALKALINITY (EQUIVALENTS/KG).
0.00, SULFITE ALKALINITY (EQUIVALENTS/KG).
0.00, SULFIDE ACIDITY(EQUIVALENTS/KG).
0.00, ACIDITY(EQUIVALENTS/KG).
0.00, IF YOU WANT TO SPECIFY HCL(BARS), ENTER VALUE HERE.
0.00, IF YOU WANT TO SPECIFY HNO3(BARS), ENTER VALUE HERE.
0.00, IF YOU WANT TO SPECIFY H2SO4(BARS), ENTER VALUE HERE.
0.00, BORON (M/KG).
0.00, FLUORIDE(M/KG).
1.467, INITIAL TOTAL PRESSURE(BARS).
```

0.00, INITIAL CO2(BARS).

0.00, INITIAL O2(BARS).

0.00, ENTER MOLE FRACTION OF CO2, 0=FIXED CO2, 1=PURE CO2.

```
0.073, INITIAL CH4(BARS).
0.00, ENTER MOLES OF CH4, 0=FIXED CH4, 1=PURE CH4.
0.00, CONSIDER A MIXED CO2-CH4 GAS HYDRATE(YES=1, NO=0)?
0.00, INITIAL NH3(G)(BARS), DO NOT INCLUDE BOTH NH3 INPUTS.
10.00, INITIAL NH3(AQ)(M/KG), DO NOT INCLUDE BOTH NH3 INPUTS.
1.394, INITIAL N2(BARS)
0.00, ENTER MOLE FRACTION OF N2, 0=FIXED N2, 1=PURE N2.
1, CONSIDER A MIXED N2-CH4 GAS HYDRATE(YES=1, NO=0)?
0, CONSIDER A MIXED N2-C03 GAS HYDRATE(YES=1, NO=0)?
0, MOLAR TO MOLAL CONVERSION? YES=1, NO=0.
0.00, IF YES ABOVE, ENTER SALINITY(G)/LITER.
273.15, INITIAL TEMPERATURE(K).
173.15, FINAL TEMPERATURE(K), IF PATH =1, OTHERWISE, SET = 0.
5.00, TEMPERATURE DECREMENT(K), IF PATH = 1, OTHERWISE, SET = 0.
1000, INITIAL WATER(G), IF PATH = 2, OTHERWISE, SET = 1000.
0.00, FINAL WATER (G), IF PATH = 2, OTHERWISE, SET = 0.
0.00, WATER DECREMENT(K), IF PATH = 2, OTHERWISE, SET = 0.
1.467, FINAL PRESSURE(BARS), IF PATH = 3, OTHERWISE, SET = 0.
0.00, PRESSURE INCREMENT(BARS), IF PATH = 3, OTHERWISE, SET = 0.
```

#### Title 11. A Titan simulation (Icarus, 2014. 236:1-8)

TITLE: SO2-H2S model parameters.

```
Temp(K) Ion.Str. RHO Phi H2O(g) Ice(g) Press.(bars) 173.15 1.3866 0.9598179 1.4682 329.63 0.00000 1.4670
```

Solution Initial Final Mass SPECIES Conc. Conc. Act.Coef. Activity Moles Balance Н NH4 3.0000 1.3862 0.40642 0.56338 0.45693 2.9292 CL1.0000 0.38447E-03 94.720 0.36417E-01 0.12673E-03 0.96596 SO4 1.0000 OHHSO4 NH3 10.000 30.337 2.9001 87.979 10.000 10.000 NH3(BAR) 0.00000 0.36472E-02 1.0000 0.36472E-02 0.00000 0.00000 0.14507 1.0000 0.14507 0.47820E-01 CH4(BAR) 0.73000E-01 0.73000E-01 0.98875 0.72179E-01 0.00000 N2(BAR) 1.3940 1.3940 0.82061 1.1439 0.00000 H2O(BAR) 0.00000 0.13762E-07 55.508 H2O(L) 55.508 0.41465 18.297

Solid Equil. Accum. SPECIES Constant Moles Moles 0.00000 0.45274 34.184 ICE NACL.2H2O 0.00000 79.233 0.00000 NACL 0.00000 0.81395E-02 0.00000 0.00000 0.63477E-02 0.00000 KCL. CACL2.6H2O 0.00000 0.21027E-05 0.00000 MGCL2.6H2O 0.00000 28.814 0.00000 0.00000 0.52980E-03 0.00000 MGCL2.8H2O 0.00000 0.10419E-33 0.00000 MGCL2.12H2O KMGCL3.6H2O 0.00000 0.36320E-01 0.00000 CACL2.2MGCL2.12H2O 0.00000 0.29714E+25 0.00000 NA2SO4.10H2O 0.00000 0.60449E-08 0.00000 NA2SO4 0.00000 0.40677 0.00000 MGSO4.6H2O 0.00000 0.10076E-01 0.00000 MGSO4.7H2O K2SO4 MGSO4.K2SO4.6H2O 0.00000 0.27934E-11 0.00000 NA2SO4.MGSO4.4H2O 0.00000 0.15338E-02 0.00000

```
CASO4.2H2O
                 0.00000 0.14168E-07 0.00000
CASO4
              0.00000 0.10338E-05 0.00000
MGSO4.11H2O
                 0.00000 0.41319E-06 0.00000
NA2SO4.3K2SO4
                  0.00000 0.33860E-15 0.00000
                   0.00000
CACO3(CALCITE)
                           0.17744E-08 0.00000
MGCO3
              0.00000 0.17232E-04 0.00000
                 0.00000 1.8115 0.00000
MGCO3.3H2O
MGCO3.5H2O
                 0.00000 0.44639E+22 0.00000
CACO3.6H2O
                 0.00000 0.35212E-11 0.00000
NAHCO3
               0.00000
                        47.160 0.00000
NA2CO3.10H2O
                  0.00000 0.43490E-04 0.00000
NAHCO3.NA2CO3.2H2O 0.00000 0.39256E-02 0.00000
3MGCO3.MG(OH)2.3H2O 0.00000 0.58049E-23 0.00000
                 0.00000 0.82208E-12.0.00000
CAMG(CO3)2
NA2CO3.7H2O
                 0.00000 0.28653E-03 0.00000
KHCO3
              0.00000 0.40237E-01 0.00000
CACO3(ARAGONITE) 0.00000 0.35291E-08 0.00000
CACO3(VATERITE)
                   0.00000 0.24288E-07 0.00000
                0.00000
                         0.61102 0.00000
HNO3.3H2O
KNO3
             0.00000 0.49287E-05 0.00000
NANO3
              0.00000 0.14625E+11 0.00000
HCL.3H2O
               0.00000
                        2879.2 0.00000
H2SO4.6.5H2O
                 0.00000
                         0.40031E-02 0.00000
                0.00000
                         0.24035 0.00000
H2SO4.4H2O
HCL.6H2O
               0.00000
                         147.94 0.00000
NANO3.NA2SO4.2H2O 0.00000 0.51509E+06 0.00000
NA3H(SO4)2
                0.00000 0.93785E-49 0.00000
                 0.00000 0.58236E+18 0.00000
NAHSO4.H2O
K3H(SO4)2
               0.00000 0.15135E-01 0.00000
K5H3(SO4)4
                0.00000 0.41467E-10 0.00000
K8H6(SO4)7.H2O
                  0.00000 0.43211E-06 0.00000
KHSO4
              0.00000 0.30115E-01 0.00000
MGSO4.H2O
                0.00000
                         0.83112E+11 0.00000
FESO4.7H2O
                0.00000
                         0.79480E-02 0.00000
                0.00000
                         4.4608 0.00000
FESO4.H2O
                          77.594
FECL2.6H2O
                0.00000
                                 0.00000
FECL2.4H2O
                0.00000
                         0.10026E+07 0.00000
             0.00000 0.17986E-09 0.00000
FECO3
FE(OH)3
              0.00000
                       0.36265E+14 0.00000
CO2.6H2O
               0.00000 0.40848E-03 0.00000
               0.48650E-01 0.36688E-03 0.83384E-01
CH4.6H2O
FECL3.10H20
                0.00000 0.42606E-12 0.00000
                0.00000
                         0.16131 0.00000
FECL3.6H2O
FECL3.2KCL.H20
                  0.00000 0.72905E-14 0.00000
              0.00000 0.18464E+30 0.00000
FE2(SO4)3
FE2(SO4)3.2K2SO4.14H 0.00000 0.14357E-13 0.00000
K2SO4.FESO4.6H2O 0.00000 0.83703E-09 0.00000
NA2SO4.FESO4.4H2O 0.00000 0.12152E-02 0.00000
FE2(SO4)3.9H2O 0.00000 0.11488E+12 0.00000
FE2(SO4)3.H2SO4.8H2O 0.00000 0.29078E+20 0.00000
KFE3(SO4)2(OH)6
                 0.00000
                           49.109 0.00000
NAFE3(SO4)2(OH)6 0.00000
                            0.46291E+12 0.00000
H3OFE3(SO4)2(OH)6 0.00000
                            0.50881E+23 0.00000
a-FE2O3
              0.00000
                      0.41772E+17 0.00000
a-FEO(OH)
               0.00000
                      0.10815E+09 0.00000
               0.00000
                        0.10255E+12 0.00000
g-FEO(OH)
FEO(OH)3/4(SO4)1/8 0.00000
                            0.26329E+11 0.00000
                0.00000
                        0.86002E-02 0.00000
FESO4.4H2O
FE2(SO4)3.7H2O
                 0.00000 0.11091E+16 0.00000
FE(II)FE(III)4(SO4)6 0.00000
                           1274.6 0.00000
FE(III)5(SO4)6O(OH). 0.00000
                           75097. 0.00000
FE(II)FE(III)2(SO4)4 0.00000 0.62687E-17 0.00000
FE(II)FE(III)2(SO4)4 0.00000
                          6.1920 0.00000
K2FE(II)5FE(III)4(SO 0.00000
                           0.14736E+07 0.00000
ALCL3.6H2O
                0.00000 0.27362E-07 0.00000
AL2(SO4)3.17H2O
                 0.00000 0.56149E-08 0.00000
```

0.00000 0.00000 0.00000 NABR MGBR2 0.0000.0 00000.0 00000.0 AL(OH)3 0.00000 0.17975E+22 0.00000 SIO2(QUARTZ) 0.00000 0.70843E-07 0.00000 SIO2(AMORPHOUS) 0.00000 0.32652E-04 0.00000 KAL3(SO4)2(OH)6 0.00000 0.36156E+30 0.00000 NAAL3(SO4)2(OH)6 0.00000 0.43250E+36 0.00000 KAL(SO4)2.12H2O 0.00000 0.16824E-10 0.00000 NAAL(SO4)2.12H2O 0.00000 0.22345E-05 0.00000 FESO4.AL2(SO4)3.22H2 0.00000 0.28067E-08 0.00000 AL2SI2O5(OH)4 0.00000 0.35225E+27 0.00000 MGSO4.AL2(SO4).22H2O 0.00000 0.98113E-08 0.00000 0.00000 0.22234 0.00000 NACLO4.H2O MG(CLO4)2.8H2O 0.00000 0.55057E+07 0.00000 0.00000 9175.8 0.00000 CA(CLO4)2.6H2O KCLO4 0.00000 0.23020E-04 0.00000 0.00000 0.10003E+31 0.00000 MG(CLO4)2.6H2O NACLO4.2H2O 0.00000 0.41764 0.00000 NH4CL 0.89823E-01 0.31728 0.54059 NH42SO4 0.22109E-03 0.11558E-01 0.96584 NH3.H2O 0.00000 40.337 0.00000 0.00000 0.29481E-01 0.00000 NH4NO3 NH4HCO3 0.00000 0.33196E-03 0.00000 0.00000 0.49606E-04 0.00000 NH4CLO4 NH3.2H2O 0.00000 15.162 0.00000 K2SO3 0.00000 465.91 0.00000 NA2SO3.7H2O 0.00000 0.43191E-03 0.00000 0.00000 0.11518E-05 0.00000 CASO3.0.5H2O 0.00000 0.17424E-05 0.00000 MGSO3.6H2O FESO3.5H2O 0.00000 0.15407E-05 0.00000 (NH4)2SO3.H2O 0.00000 0.50045E-01 0.00000 0.00000 0.39899E-16 0.00000 FES2 N2.6H2O 0.24785 0.58146E-02 0.42116 0.00000 0.12851E-01 0.00000 CASO4.0.5H2O

pH= 17.8857 pHF= 17.5010 pHT= 9.54937 pH(SWS)= 9.54937 pHMacinnis = 17.6238. Temp. =173.15. CONVERGENCE CRITERION = 0.100000. Iterations = 5