## AX program - general

AX is an activity-composition calculation program for rock-forming minerals. It was never meant to be for public release, and so is not guaranteed to be always robust - however it is freeware and we hope that it will be as useful to you as it has been to us. The program performs by first recalculating the analysis to a mineral formula and then determining the activities of mineral endmembers. The uncertainties stemming from typical probe error (0.05wt% minimum + 1.5% relative on each oxide) are propagated to the calculated activities. These are therefore minimum errors - they do not take into account activity model uncertainties.

The models used in AX are kept deliberately simple, for two reasons: 1) Natural minerals are more complex than experimentally investigated equivalents, and simpler models probably extrapolate better than elaborate ones; 2) The main use of AX is to supply activities for thermometry and barometry, not to make the most precise phase diagram calculations. The errors involved, stemming from probe analysis, inhomogeneity or incomplete equilibrium, are often as large as any errors arising from simplifying the activity models.

The program is simple and should be relatively obvious even for the new user. The main steps involved are:

1. Create an input file; this may be done either with your favourite editor, or from within AX itself. The data file consists of a line of 11 oxides (these and ONLY these 11 are accepted currently by AX, although they can be placed in any order on the first line of the data file - see the example below).

Each analysis is entered as a pair of lines, the first of which gives the mineral code (g, cpx etc) and a brief title. (A list of the mineral codes can be found from the help menu in the program, and is also given below). The second line gives the oxide wt% values in the SAME order as the oxide names in the list at the top of the file. Data may be tab-, comma-, or space- delimited, and the file is termnated with an asterisk.

An example follows:

SiO2 TiO2 Al203 Cr203 Fe203 Fe0 Mn0 Mg0 Ca0 Na20 K20 g gt
39.70 0.00 23.20 0.00 0.00 23.60 0.50 7.60 7.00 0.00 0.00 cpx omph
57.20 0.00 12.50 0.00 0.00 4.70 0.00 7.00 10.20 8.00 0.00

- 2. Run the program. Double click the AX icon.
- 3. Examining the output. 2 kinds of output file are generated:
  - 1. AX output: what you see on screen.
  - 2. A file suitable for editing and submission to THERMOCALC as input. AX enters quartz automatically, and H2O (if hydrous silicates are processed) and/or CO2 (if carbonates are used).

N.B. Eliminate all doubtful endmember data before running THER-MOCALC (either because of disequilibrium character - e.g. retrograde phases, or because the end-member is the dilute endmember on a solvus limb - see below.

#### Some further notes on activities:

• A note on tiny values for activity:

Some minerals will have very small activities simply because there are many sites on which mixing occurs. To decide whether a calculated activity is too tiny to be reliable, use the following rule of thumb: for a mineral where mixing is dominated by mixing on n sites, raise the activity to the power of 1/n and check that the result lies in the range 0.1–1.0. This is effectively normalising to an equivalent one-site solution where we would be suspicious of activities for mole fractions less than 0.1 unless good Henry's law constants are available. As an example, although for garnet (n=3) an activity of 0.008 might seem at first sight to be far too small to be reliable, this would be equivalent to 0.2 on a one-site basis and would probably be acceptable. AX does not print activities which are far smaller than their uncertainties.

#### • Dilute limbs of a solvus.:

Avoid endmembers on the "dilute" limb of a solvus - ie do not use the paragonite activity in a K-rich white mica, or the muscovite or celadonite endmembers in a paragonite. The uncertainties on wronglimb endmembers are prohibitively large.

#### • Published thermobarometer calibrations

Many published thermometers or barometers rely on very specific recipes for activities for successful use. Do not use AX activities in such formulations, particularly in sensitive cation exchange equilibria (e.g. garnet-clinopyroxene thermometers), or solvus thermometers (e.g. two pyroxene thermometers).

Code Mineral groups white micas, including margarite mu bi biotites amphiboles amph fsp feldspars epidotes, zoisites ep garnets g срх clinopyroxenes orthopyroxenes opx chl chlorites talc tascapolites scap ol olivines chloritoid  $\operatorname{ctd}$  $\operatorname{cd}$ cordierite  $\operatorname{st}$ staurolites spinels spcarb carbonates ilmenites and hematites ilhem sapphirines spr osumilites osm

## Mineral end-member activity models

Activities of mineral endmembers for Average P, average T, and average PT calculations may be estimated with the help of the program AX which accepts raw microprobe data in the form of oxide weight percents and performs standard mineral recalculations, with attempts at ferric iron estimation. The program calculates activities for end-members which can

then be used for rock calculations in THERMOCALC . The assumptions used in deriving the activities and in estimation of ferric iron are listed briefly below. ( $R_{max}$  is the maximum allowed ratio of ferric to ferrous iron; HP90 is Holland & Powell 1990, J. Met. Geol. 8, 89–124. HP98 is Holland & Powell 1998, J. Met. Geol.16, 309–343.)

For more information on someof the mineral groups, see

- 1) HP90 p 100 & especially HP98 p 315-318
- 2) www pages at www.esc.cam.ac.uk/astaff/holland/

### • Clinopyroxene

Ferric from: Cation Sum = 4 for 6 oxygens,  $R_{max}$ =0.95. Wood & Banno model (M1–M2 mixing only) used. If Na < 0.3 then disordered C2/c *calcic cpx* is assumed with nonideal mixing. e.g.  $a_{di} = X_{Ca,M2}X_{Mg,M1}\gamma_{di}$ 

Nonideality is approximated by renormalising to the set of endmembers di-hed-cats-jd-en with symmetric formalism interaction energies (kJ):

| W             | hed | cats | $\mathrm{jd}$ | en |
|---------------|-----|------|---------------|----|
| di            | 3   | 7    | 24            | 30 |
| hed           |     | 4    | 24            | 20 |
| cats          |     |      | 20            | 24 |
| $\mathrm{jd}$ |     |      |               | 24 |

If Na > 0.7 then disordered C2/c **sodic pyroxene** is assumed with nonideality approximated by renormalising to the set of endmembers jd-acm-di-hed with symmetric formalism interaction energies (kJ):

| W   | acm | di | hed |
|-----|-----|----|-----|
| jd  | 0   | 24 | 24  |
| acm |     | 24 | 24  |
| di  |     |    | 3   |

Otherwise, (if 0.3 > Na < 0.7), P2/n **omphacite** is assumed. Ideal coupled mixing is assumed as an approximation (jd-di-hed-acm).

if Mg,M1 > Ca,M2 then 
$$a_{\rm di}=X_{\rm Ca,M2}$$
 else  $a_{\rm di}=X_{\rm Mg,M1}$  if Fe,M1 > Ca,M2 then  $a_{\rm hed}=X_{\rm Ca,M2}$  else  $a_{\rm hed}=X_{\rm Fe,M1}$  if Al,M1 > Na,M2 then  $a_{\rm jd}=X_{\rm Na,M2}$  else  $a_{\rm jd}=X_{\rm Al,M1}$   $a_{\rm acm}=X_{\rm Fe^{3+},M1}$ 

### Orthopyroxene

Ferric from: Cation Sum = 4 for 6 oxygens,  $R_{\text{max}}=0.2$ . The Wood & Banno model (M1–M2 mixing only) is used. e.g.  $a_{\text{en}} = X_{\text{Mg,M2}} X_{\text{Mg,M1}} \gamma_{\text{en}}$  Nonideality is approximated by renormalising to the set of endmembers en-fs-mgts-di with symmetric formalism interaction energies (kJ):

| W    | fs | mgts | di |
|------|----|------|----|
| en   | 1  | 0    | 30 |
| fs   |    | 1    | 28 |
| mgts |    |      | 30 |

#### • Olivine

Ferric from: Cation Sum = 3 for 4 oxygens,  $R_{max}$ =0.1

Mixing on sites used (M1-M2). e.g.  $a_{\rm fo} = X_{\rm Mg,M2} X_{\rm Mg,M1} \gamma_{\rm fo}$ 

Nonideality is approximated by renormalising to the set of endmembers fo-fs with symmetric formalism interaction energy 4.0 kJ per site  $(W_{\text{fo,fa}} = 8 \text{ kJ})$ 

#### • Talc

Ferric from: Cation Sum = 7 for 11 oxygens,  $R_{\text{max}}$ =0.1

Ideal mixing on sites model of Holland & Powell 1998 is used. e.g.  $a_{\rm ta}=X_{\rm Mg,M1}X_{\rm Mg,M2}^2X_{\rm Si,T1}^2$ 

### • Garnet

Ferric from: Cation Sum = 8 for 12 oxygens,  $R_{max}$ =0.99

2 site mixing model for ideal mixing part e.g.  $a_{py} = X_{Mg,M2}^3 X_{Al,M1}^2 \gamma_{py}$ Nonideality is approximated by renormalising to the set of endmembers py-gr-alm-spss-andr with symmetric formalism interaction energies (kJ):

| W    | gr | alm | $\operatorname{spss}$ | andr |
|------|----|-----|-----------------------|------|
| py   | 33 | 2.5 | 0                     | 73   |
| gr   |    | 0   | 0                     | 0    |
| alm  |    |     | 0                     | 60   |
| spss |    |     |                       | 60   |

## • Epidote

Ferric from: Si + Al + Fe<sup>3+</sup> = 6 for 12.5 oxygens,  $R_{max}$ =0.99

2 site mixing and ordering model of Holland & Powell 1998, involving endmembers clinozoisite (cz AlAl), epidote (ep AlFe) and ferricepidote (fep FeFe). e.g.

$$a_{\rm cz} = X_{\rm Ca,M2} X_{\rm Al,M1} X_{\rm Al,M3} \gamma_{\rm cz}$$

$$a_{\rm ep} = X_{\rm Ca,M2} X_{\rm Al,M1} X_{\rm Fe^{3+},M3} \gamma_{\rm ep}$$

$$a_{\text{fep}} = X_{\text{Ca,M2}} X_{\text{Fe}^{3+},\text{M1}} X_{\text{Fe}^{3+},\text{M3}} \gamma_{\text{fep}}$$

Nonideality is approximated by renormalising to the set of endmembers cz-ep-fep. The proportion of fep acts as an order parameter and is determined at the temperature selected in the program. The symmetric formalism interaction energies (kJ):

 $\Delta H = -26.10 \text{ kJ}$  for the reaction cz + fep = 2 ep

## • Feldspar

Ferric: all iron taken as ferric.

If Ca < 0.05, then **plagioclase feldspar** is assumed, with the ordered model of Holland & Powell 1992 (model 1)

If Ca > 0.05, then **alkali feldspar** is assumed. The subregular solution model of Waldbaum & Thompson 1969 is used. Tetrahedral mixing terms are ignored.

## • Scapolite

Ferric: all iron is taken as ferrous.

Ideal mixing on large cations site, ignoring tetrahedral terms. e.g.

$$a_{\text{me}} = X_{\text{Ca,A}}^4$$
$$a_{\text{miz}} = 9.48X_{\text{Ca,A}}^3 X_{\text{Na,A}}$$

#### Chloritoid

Ferric from: Cation Sum = 4 for 6 oxygens.  $R_{max} = 0.2$ 

For the half formula size used, there is one M2 site (Fe, Mg, Mn) and half a M1 site (Al, Fe<sup>3+</sup>) and so:  $a_{\text{mctd}} = X_{\text{Mg,M2}} X_{\text{Al,M1}}^{0.5} \gamma_{\text{mctd}}$ 

Nonideality is approximated by renormalising to the set of endmembers mctd-fctd-mnctd with symmetric formalism interaction energies (kJ)

| W    | fctd | mnctd |
|------|------|-------|
| mctd | 1.5  | 1.5   |
| fctd |      | 1.5   |

### Amphibole

Ferric from: The method of Holland & Blundy 1993.

The ideal mixing part follows mixing on sites, following Holland & Powell 1998 (which allows only half the configurational entropy contribution from tetrahedral sites). Thus

$$a_{\rm tr} = X_{\Box,A} X_{{\rm Ca,M4}}^2 X_{{\rm Mg,M13}}^3 X_{{\rm Mg,M2}}^2 X_{{\rm Si,T1}}^2 \gamma_{\rm tr}$$

For *calcic amphiboles*, taken when  $X_{\text{Ca,M4}} > 0.5$ , nonideality is approximated by renormalising to the set of endmembers tr-fact-ts-parg-gl-fits-kpa with symmetric formalism interaction energies (following Dale, et al., 2005) (kJ)

| W                  | fact | ts   | parg  | gl   | fits  | kpa |
|--------------------|------|------|-------|------|-------|-----|
| $\overline{ m tr}$ | 10   | 20   | 33    | 65   | 20    | 0   |
| fact               |      | 12.5 | -1.9  | 39.3 | 12.5  | 0   |
| ts                 |      |      | -38.5 | 25   | 0     | 0   |
| parg               |      |      |       | 50   | -38.5 | 0   |
| gl                 |      |      |       |      | 45.9  | 0   |
| fits               |      |      |       |      |       | 0   |

For **sodic amphiboles**, here taken when  $X_{\text{Na,M4}} > 0.5$ , nonideality is approximated by renormalising to the simple set of endmembers gl-fgl-rieb with symmetric formalism interaction energies (kJ)

For Fe-Mg amphiboles, taken when  $X_{\text{Ca,M4}} < 0.3$  and  $X_{\text{Na,M4}} < 0.3$ , nonideality is approximated by renormalising to the simple set of endmembers cum-grun with a symmetric formalism interaction energy  $W_{\text{cum,gr}} = 17.5$  (kJ). This approximates the behaviour modelled in Holland & Powell (1996) in the metamorphic temperature range.

#### • Chlorite

Ferric from: Cation Sum = 10 for 14 oxygens.  $R_{max} = 0.2$ 

Mixing is taken from Holland, Baker & Powell (1998), simplified for chlorites more aluminous than clinochlore (Al assumed ordered into the M4 site). e.g.

$$a_{\text{clin}} = 4X_{\text{Mg,M23}}^4 X_{\text{Mg,M1}} X_{\text{Al,M4}} X_{\text{Al,T2}} X_{\text{Si,T2}} \gamma_{\text{clin}}$$

Nonideality is approximated by renormalising to the set of endmembers clin-daph-ames with symmetric formalism interaction energies (kJ)

#### • White mica

Ferric from: Tet + Oct cation sum = 6.05 for 11 oxygens.  $R_{max} = 0.7$  Mixing is taken from Holland & Powell (1998) e.g.

$$a_{\mathrm{mu}} = 4X_{\mathrm{K,A}}X_{\square,M1}X_{\mathrm{Al,M2}}X_{\mathrm{Al,M3}}X_{\mathrm{Al,T1}}X_{\mathrm{Si,T1}}\gamma_{\mathrm{mu}}$$

For *margarites*, here taken when  $X_{\text{Ca,A}} > 0.5$ , nonideality is approximated by a regular solution of Ca-Na- $\square$  on the A site (i.e the small amount of K is ignored) with interaction energies (kJ)

For **paragonites**, here taken when  $X_{\text{Na,A}} > 0.5$ , nonideality is approximated as follows: for margarite activity the solid solution is treated as a ternary Ca-Na- $\square$  as above; for muscovite and paragonite activities the solid solution is treated as a ternary K-Na- $\square$  (ignoring the small Ca content) with a DQF increment to the muscovite free energy: (kJ)

$$DQF (mu) = -3.28 \text{ kJ}$$

For *muscovite-phengites*, here taken when  $X_{K,A} > 0.5$ , nonideality is approximated by renormalising to the set of endmembers mu-pacel-fcel with symmetric formalism interaction energies (kJ)

$$DQF (pa) = 1.42 + 0.4P kJ$$

#### • Biotite

Ferric from: Tet + Oct cation sum = 6.9 for 11 oxygens.  $R_{max} = 0.15$  Mixing is taken from the ordering model of Powell & Holland (1999) e.g.:

$$a_{\rm phl} = 4X_{\rm K,A}X_{\rm Mg,M1}X_{\rm Mg,M3}^2X_{\rm Al,T1}X_{\rm Si,T1}\gamma_{\rm phl}$$

Nonideality is approximated by renormalising to the set of endmembers phl-ann-east-obi with symmetric formalism interaction energies (kJ)

| W    | ann | east | obi |
|------|-----|------|-----|
| phl  | 9   | 10   | 3   |
| ann  |     | -1   | 6   |
| east |     |      | 10  |

 $\Delta H = -32.3 \text{ kJ}$  for the reaction ann + phl = 2 obi

#### • Cordierite

Ferric from: Cation Sum = 11 for 18 oxygens,  $R_{max}$ =0.2

Mixing on M sites only is used. e.g.  $a_{\rm crd} = X_{\rm Mg,M}^2 \gamma_{\rm crd}$ 

Nonideality is approximated by renormalising to the set of endmembers crd-frcd-mncrd with symmetric formalism interaction energies (kJ)

#### • Staurolite

Ferric from: all ferrous assumed

Ideal mixing on 4 M sites only is used, e.g.  $a_{\text{fst}} = X_{\text{Fe,M}}^4$ 

### • Spinel

Ferric from: Cation Sum = 3 for 4 oxygens. Max Ratio = 0.9;

If an *aluminous spinel* (Al > 1.0) then a 3-site mixing model is used, e.g.

$$a_{\rm sp} = X_{\rm Mg,M}^2 X_{\rm Al,T} \gamma_{\rm sp}$$

Nonideality is approximated by renormalising to the set of endmembers sp-herc-mt with symmetric formalism interaction energies (kJ)

For magnetites (Al < 1.0) the model involves simple ideal mixing in inverse spinels:

$$a_{\rm mt} = X_{\rm Fe^{3+},M}^2 X_{\rm Fe,T}$$

### • Sapphirine

Ferric from: Cation Sum = 14 for 20 oxygens,  $R_{max}$ =0.7 Ideal mixing on sites model of Holland & Powell 1998

#### • Osumilite

Ferric from: Cation Sum = 18 for 30 oxygens,  $R_{max}$ =0.4

Ideal mixing on sites model of Holland & Powell 1998 e.g.:

$$a_{\text{osm1}} = X_{\text{K,A}} X_{\text{Mg,M1}}^2 X_{\text{Al,T1}}^3 X_{\text{Al,T2}}^2$$

#### • Carbonates

Ferric from: all ferrous

For **dolomite-ankerites**, a 2-site model is used e.g.  $a_{\rm dol} = X_{\rm Ca,M2} X_{\rm Mg,M1} \gamma_{\rm dol}$ Nonideality assumes  $W_{\rm FeMg,M1} = 3.0 \text{ kJ}$ 

For *calcite-magnesite-siderite-rhodachrosite* disordered carbonates a simple 1-site model is used e.g.  $a_{cc} = X_{Ca,M}\gamma_{cc}$ 

Nonideality is approximated by renormalising to the set of endmembers cc-mag-sid-rhc with symmetric formalism interaction energies (kJ)

| W                    | mag | $\operatorname{sid}$ | $\operatorname{rhc}$ |
|----------------------|-----|----------------------|----------------------|
| cc                   | 22  | 18                   | 0                    |
| mag                  |     | 4                    | 0                    |
| $\operatorname{sid}$ |     |                      | 0                    |

# • Ilmenite-hematite

Ferric from: Cation Sum = 2 for 3 oxygens.

Simple 2-site mixing in ordered ilm-hem is used.

$$a_{\text{ilm}} = X_{\text{Fe,M1}} X_{\text{Ti,M2}}$$

$$a_{\text{hem}} = X_{\text{Fe}^{3+},\text{M1}} X_{\text{Fe}^{3+},\text{M2}}$$