## NOMENCLATURE OF AMPHIBOLES

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for

## SUBCOMMITTEE ON AMPHIBOLES. I.M.A.

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Final report by the subcommittee on the Amphibole Group as approved by the International Mineralogical Association Commission on New Minerals and Mineral names. The amphibole subcommittee was composed of H. Winchell, Chairman (U.S.A.), R. A. Binns (Australia), M. Fleischer (U.S.A.) later replaced by A Kato (Japan), C. Guillemin (France) later replaced by G. Gottardi (Italy) and M. Fonteilles (France), E. Hilmy (Egypt), B. E. Leake (U.K.), K. J. Neuvonen (Finland), and L. van der Plas (Netherlands), later replaced by H. J. Kisch (Israel). All the reports were compiled by B. E. Leake.

This report is the fifth draft and could not have been compiled without the previous extensive work by R. Felix, L. van der Plas (The Netherlands), E. J. W. Whittaker (U.K.), R. A. Binns (Australia), K. J. Neuvonen (Finland), M. Ross, P. Robinson and H. Winchell (U.S.A.), together with many non-members of the sub-committee on amphiboles including E. K. Lazarenko (U.S.S.R.), I. V. Ginsburg (U.S.S.R.), V. A. Frank-Kamenetskii (U.S.S.R.), I. Kostov (Bulgaria), E. H. Nickel (Australia), M. Hey (U.K.), H. Micheelson (Denmark) and E. Wenk (Switzerland).

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#### 1. General classification of the amphiboles

It is proposed that the classification of the amphiboles should be largely based on crystal chemistry, as the optical and other physical determinative properties such as X-ray powder diffraction cannot differentiate unambiguously between different members of the group. Of course the traditional and important distinction between orthorhombic and monoclinic members has been retained. When it is necessary to distinguish different polytypes or polymorphs further (e.g. with cummingtonite) this may be done by adding the space group symbol as a suffix.

The proposed nomenclature has successfully avoided introducing new names by the use of adjectival modifiers (e.g. titanian) and prefixes (e.g. ferro-) which cover specified elemental ranges and which, for simplicity, are hereafter both called prefixes. Accepted and widely used names have been chemically codified to agree, as far as is possible, with the consensus of present use. About 200 previously used amphibole names, mostly synonyms or obsolete or almost unused names, are recommended for formal extinction.

The classification is based on the chemical contents of a standard amphibole calculated to 24(0,0H,F,Cl), but it is recognised that where there is no determination of H2O+ (e.g. electron microprobe analyses), or there is reason to suppose that the reported  $H_2O+$  is erroneous, or where it is probable that unreported F or C1 may be substantial, then the basis of 23(0) should be used to calculate the cation contents of the standard formula. This formula unit contains eight tetrahedral sites and corresponds to the half unit cell for monoclinic amphiboles and to one quarter of the unit cell for orthorhombic amphiboles.

Throughout this report the standard amphibole formula is used with superscript arabic numerals (e.g.  $Fe^2$ ) referring to charges; roman numerals (e.g.  $Mg_3$ ) to co-ordination numbers and subscript numerals to numbers of atoms (e.g.  $Mg_3$ ). General works dealing with the amphibole group include Deer et al. (1963), Ernst (1968) and the special papers of the Mineralogical Society of America (1969) and Great Britain (1968) which together provide a key to the voluminous literature.

The standard amphibole formula is taken to contain 8 tetrahedral sites and the general form of the standard formula is:

In the calculation of the standard amphibole formula the following procedure is recommended:

- (1) If the water and halogen contents are well established, or if there is physical evidence that the amphibole is an oxy-amphibole, the formula should be calculated to 24(0,0H,F,Cl).
- (2) If the water plus halogen content is uncertain the formula should be calculated on a water-free (and halogen free) basis to 23(0) and 2(0H,F,Cl) assumed, unless this leads to an impossibility of satisfying any of the following criteria, in which case appropriate change in the assumed number of (0H+F+Cl) should be made.
- (3) Sum T to 8.00 using Si, then Al, then Cr3, then Fe3, then Ti4.
- (4) Sum C to 5.00 using excess Al, Cr, Ti, Fe<sup>3</sup> from (3), then Mg, then Fe<sup>2</sup>, and then Mn.
- (5) Sum B to 2.00 using excess Fe<sup>2</sup>, Mn, Mg from (2), then Ca, then Na.
- (6) Excess Na from (5) is assigned to A, then all K. Total A should be between 0.00 and 1.00, inclusive.

These assignments normally correspond to the occupancies of the tetrahedral sites (T), the M1 + M2 + M3 sites (C), the M4 sites (B) and the A sites (A). Present knowledge of the distribution of ions is not sufficient to warrant making separate formal allocation to the three distinct sites that in total constitute the C position, nor does the available evidence suggest that calculation to a fixed number of cations is desirable.

When a standard amphibole formula has been determined in this way it is classified first into one of four principal amphibole groups on the basis of the numbers of atoms of  $(Ca + Na)_B$  and  $Na_B$ . Within each of these groups it can then be named by reference to the appropriate two-dimensional diagram (Figs. 2-5) using the number of Si atoms and the ratio  $Mg/(Mg + Fe^2)$ . The name so found is the name of the defined end-member to which the formula most closely approximates. This name may be qualified by one or more prefixes according to definite rules to specify important (but relatively minor) departures from the end-member formula. The four principal amphibole groups are defined as:

- (a) When (Ca+Na)<sub>B</sub> < 1.34, then the amphibole is a member of the <u>iron-magnesium-manganese amphibole group</u>.
- (b) When  $(Ca+Na)_B \ge 1.34$  and  $Na_B < 0.67$ , then the amphibole is a member of the <u>calcic amphibole group</u>. Nearly all such natural amphiboles have  $Ca_B > 1.34$ .

- (c) When  $(Ca+Na)_B \ge 1.34$  and  $0.67 \le Na_B < 1.34$ , then the amphibole is a member of the sodic-calcic amphibole group. Such natural amphiboles usually contain  $0.67 < Ca_B < 1.34$ .
- (d) When  $(Na)_B \ge 1.34$ , then the amphibole is a member of the <u>alkali amphibole</u> group.

The principal reference axes chosen for the calcic, sodio-calcic and alkali amphibole groups are Na<sub>B</sub>; (Na+K)<sub>A</sub>; and (8-Si), as shown in Fig. 1 based on Smith's (1959) proposals. Other choices of axes are of course possible, and have been considered, but for various excellent reasons the present choice is recommended.

In general the scheme seeks to avoid primary divisions at integral contents of the standard formula so that analyses near to formalised end-, or integral members, whose names are defined, are grouped together, rather than split apart.

The form of the Mg to Fe ratio usually used is  $Mg/(Fe^2+Mg)$ . An increasing number of amphibole analyses are being obtained by microprobe analysis (over 85% of those reported in 1976) and these analyses usually do not report Fe<sub>2</sub>O<sub>3</sub>. There are various different possible procedures to partially alleviate the problems raised by such partial analyses but no one procedure is recommended though calculation on the basis of 23(0) and then adjustment of the total cations, excluding (Ca+Na+K), to 5+8=13, by varying the Fe<sup>2</sup>/Fe<sup>3</sup>, has much to recommend it.

Provision is made to denote by prefixes the presence of substantial substitution by elements that are not essential constituents of the end-members. Prefixes that are generally applicable are:-

chlor	when	Cl	≥ 1.00 (about 4% Cl)
chromium	when	Cr	≥ 1.00 (about 9% Cr <sub>2</sub> 0 <sub>3</sub> )
chromian	when	Cr_	= $0.25-0.99$ (about $2.3-9\%$ $Cr_20_3$ )
ferri	when		≥ 1.00 (about 9% Fe <sub>2</sub> 0 <sub>3</sub> ) except in alkali amphiboles and hastingsite
ferrian	when	Fe <sup>3</sup>	= 0.75-0.99 (about 6.8-9% Fe <sub>2</sub> 0 <sub>3</sub> ) except in alkali amphiboles and hastingsite
fluor	when	F	≥ 1.00 (about 2% F)
hydro	when	OH	≥ 3.00 (about 3% H <sub>2</sub> 0)
lithian	when	Гī	≥ 0.25 (about 0.4% Li20) except in the alkali amphiboles when Li ≥ 0.50. Not used with holmquistite and clinoholmquistite.
manganese	when	Min	≥ 1.00 (about 10% MnO) except in end-members containing Mn
manganoan	when	Mn	= 0.25-0.99 (about 2.5-10% MnO) except in end- members containing Mn
oxy	when	OH+F+Cl	1.00. As many poor analyses have low recorded water and no F or Cl values, this prefix should be used with discretion.
plumbian	when	Pb	≥ 0.08 (about 1.1% Pb0)
potassium	when	K	≥ 0.50 (about 2.7% K <sub>2</sub> 0)
potassian	when	K	= 0.25-0.49 (about 1.3-2.7% K20) except in the alkali amphiboles
subsilicio	when	Si	< 5.75

CALCIC

ALKAL! AMPHIBOLES

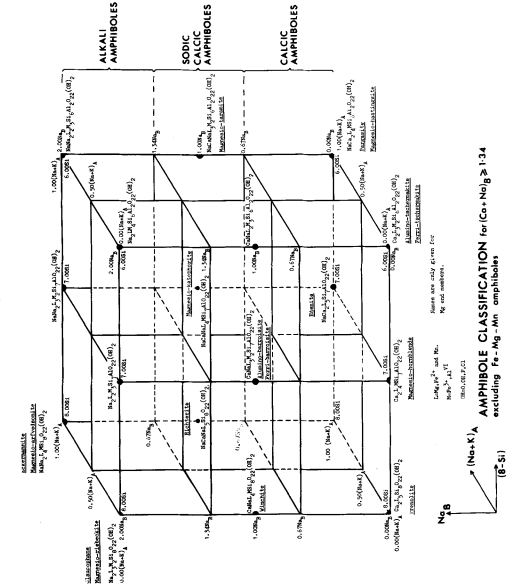


Fig. 1

titanium	when Ti	≥ 1.00 (about 10% TiO <sub>2</sub> ) except in kaersutite
titanian	when Ti	= 0.25-0.99 (about 2.5-10% TiO <sub>2</sub> ) except in kaer-
zino	when Zn	≥ 1.00 (about 5% ZnO) sutite
zincian	when Zn	= 0.25-0.99 (about 1.2-5% ZnO)

A few prefixes (alumino, calcian, subcalcic, and sodian) have to be defined differently in the different principal amphibole groups, and their definitions are given in the appropriate places.

The proposals often do not involve uniform divisions at elegant and invariable mathematical points such as would clearly be proposed if usage could be ignored. On the contrary, the four separate amphibole-group schemes each endeavour to fit present usage and codify it. Consequently there are sometimes rather untidy aspects but this is preferable to schemes which cut across traditional and present usage. As there are already over 8000 published amphibole analyses it is important to provide for nearly every probable variation so as to avoid irregular proliferation of names and this is best prevented by providing ample scope for fairly detailed compositional indications.

Adjectival prefixes have been employed to keep the number of fundamental amphibole names to a minimum and to indicate specifically defined ranges of composition which seek to (1) avoid present and future haphazard and irregular naming, (2) enable between 15 and 20 variables to be conveyed in the name either explicitly or, more usually, implicitly (i.e., by the absence of a prefix), (3) give a non-specialist mineralogist or petrologist a name which in itself is meaningful (e.g. manganoan) even if the defined specific element-ranges covered by the adjectival prefixes are unknown. The absence of a prefix means that the element concerned is below, or occasionally above (e.g. with subsilicic and subcalcic), the limits prescribed for the use of the prefix, which in all instances has been defined after considering what is common and what is unusual and the limits defined endeavour to mark out the unusual from the common. Schaller's (1930) adjectives are used to indicate moderate enrichment of substituting elements.

The names proposed usually take into account and convey information about the following variables in the standard formula:-

$$\text{Si,Al}^{\text{IV}}, (\text{Ca+Na})_{\text{R}}, (\text{Na+K})_{\text{A}}, \text{Ca, Al}^{\text{VI}}, \text{Fe}^{\text{3}}, \text{Ti,F,Cl,K,Mm,Cr,Zn,Li,Fb,OH,O,and Mg/(Mg+Fe}^{\text{2}})$$

Prefixes magnesio-, ferro-, alumino-, and ferri- are often used with names that refer to part of a series. Alternate names are so widely used for the end or ends of some series that the alternative is sometimes preferable, such as tremolite instead of magnesio-actinolite and tschermakite as a synonym of alumino-tschermakite, particularly where two or more prefixes are otherwise required. If it is especially required to distinguish between pure theoretical end-members and natural compositions that will always only approach the theoretical end-member composition, then the prefix pure may (i.e. it is not obligatory) be used for the theoretical integral formula e.g. pure tremolite for Ca2 Mg5 Si8 O22 (OH)2.

For amphiboles whose general nature only is known, (for instance, from optical properties without a chemical analysis) it may not be possible to allocate a precise name. It is then recommended that the assigned amphibole name be made into an adjective to be followed by the word amphibole. Thus, anthophyllitic amphibole, tremolitic amphibole, pargasitic amphibole, richteritic amphibole and glaucophanic amphibole. The familiar word hornblende can still be used where appropriate for calcic amphiboles, because hornblende is never used without an adjective in the precise nomenclature. The adoption of these recommendations will not only avoid confusion between precisely and loosely named amphiboles but will not inhibit the giving of loose names that is obviously often inevitable when only paragenesis and optical properties are available.

Several names have been used for various asbestiform amphiboles. In mineralogy, as distinct from commercial use, the precise mineral name according to this report should be used, followed by -asbestos; e.g. anthophyllite-asbestos, actinolite-asbestos. Where the nature of the mineral is uncertain or unknown, asbestos alone may be appropriate. Where the approximate nature of the mineral is known but not its precise composition, the recommendations made above should be followed but amphibole should be replaced by asbestos, e.g. anthophyllitic asbestos, actinolitic asbestos. For this purpose crocidolite may also be retained to cover alkali amphibole asbestos as a general name whereas, e.g. riebeckite-, or magnesio-riebeckite-asbestos should be used when the precise composition is known.

Finally, it has been much in mind that the amphiboles constitute an extremely complex group: while even more detailed subdivisions are possible, the proposals attempt to be as simple as is reasonable so that ordinary mineralogists and petrologists will be able to rapidly, uniquely and unambiguously name most amphibole analyses.

Each of the four principal amphibole groups is dealt with separately below. The above section was approved by 12 votes for and 1 against.

## 2. Fe-Mg-Mn Amphiboles

The group is defined so as to include possessing (Ca+Na) $_{\rm B}$  < 1.34 in the standard formula. The detailed classification is based on Fig. 2.

# ORTHORHOMBIC FORMS

(1) Anthophyllite  $Na_x(Mg,Mn,Fe^2)_{7-y}Al_y(Al_{x+y}Si_{8-x-y})0_{22}(OH,F,Cl)_2$  where x+y<1.00, otherwise the mineral is gedrite.

End Members

Magnesic-anthophyllite  ${\rm Mg}_7{\rm Si}_8{\rm O}_{22}{\rm (OH)}_2$ Ferro-anthophyllite  ${\rm Fe}^2{}_7{\rm Si}_8{\rm O}_{22}{\rm (OH)}_2$ 

Sodium anthophyllite Na(Mg,Fe<sup>2</sup>)7AlSi7022(OH)2

Limits for use of end member names

Magnesio-anthophyllite  $Mg/(Fe^2+Mg) \ge 0.90$ Ferro-anthophyllite  $Fe^2/(Mg+Fe^2) \ge 0.90$ 

Sodium anthophyllite Na≥0.50

Prefix for particular substitution (see also below)

Alumino- when Al<sup>VI</sup>≥0.50

Na<sub>x</sub>(Mg,Mn,Fe<sup>2</sup>)<sub>7-y</sub>Al<sub>y</sub>(Al<sub>x+y</sub>Si<sub>8-x-y</sub>)0<sub>22</sub>(OH,F,Cl)<sub>2</sub>
when x+y≥1.00, the distinction from anthophyllite
being based on the total Al<sup>IV</sup>, which exceeds 0.99
in gedrite.

End Members

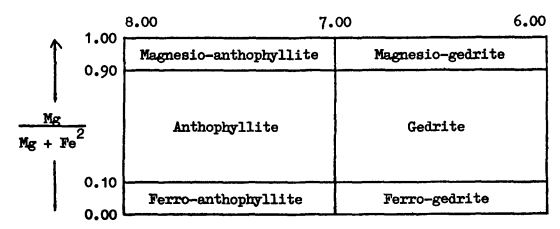
Magnesio-gedrite  $Mg_5Al_2Si_6Al_2O_{22}(OH)_2$ Ferro-gedrite  $Fe_5^2Al_2Si_6Al_2O_{22}(OH)_2$ 

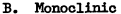
Sodium gedrite Na(Mg,Fe) AlSi Al 2022 (OH)

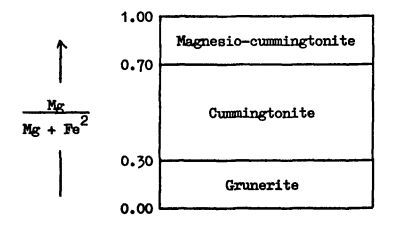
# Fig. 2. IRON-MAGNESIUM-MANGANESE AMPHIBOLES Li < 1.00; (Ca+Na)<sub>R</sub> < 1.34

A. Orthorhombic

← Si in the standard cell →







Limits for use of end member names

 $Mg/(Fe^2+Mg) \ge 0.90$ Magnesio-gedrite  $Fe^2/(Mg+Fe^2) \ge 0.90$ Ferro-gedrite

Prefix for particular substitution

when Na≥0.75 Sodium

 $\text{Li}_{2}(\text{Mg,Fe}^{2})_{3}(\text{Fe}^{3},\text{Al})_{2}\text{Si}_{8}\text{O}_{22}(\text{OH,F,Cl})_{2}$ (3) Holmquistite It is critical that Li ≥ 1.00 in structural formula (about 1.7%Li<sub>2</sub>0). End Members

Magnesio-holmquistite IT 5 WE 2 T 5 18 0 55 (OH) 5 LigFezAlgSigOgg(OH) Ferro-holmquistite

Limits of use of end member names

 $M_{\rm g}/({\rm Fe}^2 + M_{\rm g}) \ge 0.90$  ${\rm Fe}^2/(M_{\rm g} + {\rm Fe}^2) \ge 0.90$ Magnesio-holmquistite Ferro-holmquistite

MONOCLINIC FORMS

(Mg, Fe<sup>2</sup>, Mn)<sub>7</sub> Si<sub>2</sub>O<sub>22</sub>(OH)<sub>2</sub> (1) Cummingtonite Series

End Members

Magnesio-cummingtonite Mg7Si8022(OH)2 Fe<sub>7</sub>Si<sub>8</sub>O<sub>22</sub>(OH)<sub>2</sub> Grunerite Mn<sub>2</sub>Mg<sub>5</sub>Si<sub>8</sub>O<sub>22</sub>(OH)<sub>2</sub> Mn<sub>2</sub>Fe<sub>5</sub>Si<sub>8</sub>O<sub>22</sub>(OH)<sub>2</sub> Tirodite Dannemorite

Limits of use of end member names

Mg/(Fe<sup>2</sup>+Mg) ≥ 0.70 Magnesio-cummingtonite  $Fe^2/(Mg+Fe^2) \ge 0.70$ Grunerite

Mn/(Mn+Mg+Fe)≥0.10 and Mg≥Fe Tirodite Mn/(Mn+Fe+Mg)≥0.10 and Mg<Fe Dannemorite

Prefix for particular substitution (see also below)

when Na≥ 0.25 Sodian

 $\text{Li}_{2}(\text{Mg}, \text{Fe}^{2}, \text{Mn})_{3}(\text{Fe}^{3}, \text{Al})_{2}\text{Si}_{8}\text{O}_{22}(\text{OH}, \text{F}, \text{Cl})_{2}$ (2) Clinoholmquistite It is critical that Li ≥ 1.00 (i.e. about 1.7% Li\_0)

End Members

Magnesio-clinoholmquistite Li2Mg3Al2Si8022(OH)2 Li<sub>2</sub>Fe<sub>3</sub>Al<sub>2</sub>Si<sub>8</sub>O<sub>22</sub>(OH)<sub>2</sub> Ferro-clinoholmquistite

Limits for use of end member names

| Mg/(Fe<sup>2</sup>+Mg) \ge 0.90  $Fe^2/(Mg+Fe^2) \ge 0.90$ Ferro-clinoholmquistite

Special prefix for the whole Fe-Mg-Mn group of amphiboles

when Ca≥0.50 (about 3.5% CaO) Calcian

Nomenclature is given by reference to Fig. 2 or if  $\text{Li} \ge 1.00$  to the above text, combined with the prefixes given for the whole amphibole group and those special to the Fe-Mg-Mn amphiboles.

The above section was approved by 11 votes for and 2 against.

## 3. Calcic Amphiboles

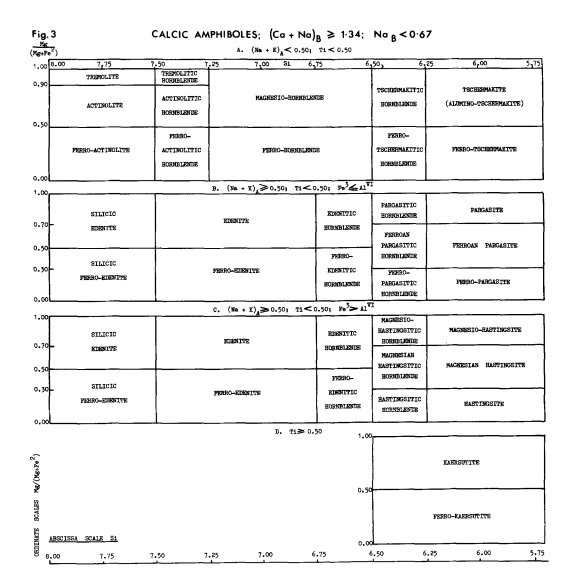
The group is defined as monoclinic amphiboles in which  $(Ca+Na)_B \ge 1.34$  and  $Na_R < 0.67$ . Generally  $Ca_R > 1.34$ .

## End Members

Tremolite	$^{\mathrm{Ca_2Mg_5Si_80_{22}(OH)_2}}$
Ferro-actinolite	$^{\mathrm{Ca}_{2}\mathrm{Fe}_{5}^{2}\mathrm{Si}_{8}\mathrm{O}_{22}\mathrm{(OH)}_{2}}$
Edenite	${\tt NaCa_2Mg_5Si_7Alo_{22}(OH)_2}$
Ferro-edenite	$\text{NaCa}_2\text{Fe}_5^2\text{Si}_7\text{AlO}_{22}(\text{OH})_2$
Pargasite	$NaCa_2Mg_4AlSi_6Al_2O_{22}(OH)_2$
Ferro-pargasite	$NaCa_2Fe_4^2AlSi_6Al_2O_{22}(OH)_2$
Hastingsite	$NaCa_2Fe_4^2Fe_5^3Si_6Al_2O_{22}(OH)_2$
Magnesio-hastingsite	$NaCa_2Mg_4Fe^3Si_6Al_2O_{22}(OH)_2$
Alumino-tschermakite	$^{\mathrm{Ca_2Mg_3Al_2Si_6Al_2O_{22}(OH)_2}}$
Ferro-alumino-tschermakite	$\text{Ca}_{2}\text{Fe}_{3}^{2}\text{Al}_{2}\text{Si}_{6}^{2}\text{Al}_{2}^{0}\text{C}_{22}^{0}$
Ferri-tschermakite	$^{\mathrm{Ca_2Mg_3Fe_2^3Si_6Al_2O_{22}(OH)_2}}$
Ferro-ferri-tschermakite	$^{\text{Ca}_{2}\text{Fe}_{3}^{2}\text{Fe}_{2}^{3}\text{Si}_{6}^{\text{Al}_{2}^{0}}\text{O}_{22}^{\text{OH}})_{2}}$
Alumino-magnesio-hornblende	$Ca_2Mg_4AlSi_7AlO_{22}(OH)_2$
Alumino-ferro-hornblende	$\text{Ca}_{2}\text{Fe}_{4}^{2}\text{AlSi}_{7}\text{AlO}_{22}\text{(OH)}_{2}$
Kaersutite	NaCa2Mg4TiSi6Al2(O+OH)24
Ferro-kaersutite	NaCa <sub>2</sub> Fe <sub>4</sub> TiSi <sub>6</sub> Al <sub>2</sub> (O+OH) <sub>24</sub>
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## Limits for use of end member names and nomenclature of the group

The nomenclature of the group is tabulated in Fig. 3. Assignment of the name is as follows: If  $Ti \ge 0.50$  go to Fig. 3D; If Ti < 0.50 and  $(Na+K)_A < 0.50_{VI}$  go to Fig. 3A; If Ti < 0.50 and  $(Na+K)_A \ge 0.50$ , then go to Fig. 3B if  $Fe^{3+} < Al^{VI}$  and to Fig. 3C if  $Fe^{3} \ge Al^{VI}$ . Further subdivisions depend upon Si and  $Mg/(Fe^2 + Mg)$ . These give the fundamental name of the particular amphibole. The final step is to scan the range of the elements dealt with by prefixes to finally obtain a name which implicitly or explicitly conveys an indication of the composition with respect to no less than 19 variables — Si,  $Al^{VI}$ ,  $Al^{VI}$ ,  $Fe^3$ ,  $(Na+K)_A$ ,  $Na_B$ , Ca, Ti, F, Cl, K, Na, Mn, Zn, Cr, Fb, CH, O and  $Mg/(Fe^2 + Mg)$ . Although it would appear that very long and cumbersome names would be common, the reverse is true because the



prefixes are only used for unusual compositions and so over 80% of the available analyses in this group give names containing two or fewer adjectives, including adjectives which form part of the fundamental name.

## Special prefixes for the calcic amphibole group

Alumino when Al<sup>VI</sup> ≥ 1.00

Sodian when Na 2 1.00 (about 3.5% Na<sub>2</sub>0) Subcalcic when Ca < 1.50 (about 9.5% CaO)

The compositions of the two tschermakite end-members, one with Al<sup>VI</sup> and the other with Fe<sup>2</sup>, can be clearly indicated and the prefixes ferri- or alumino- are in practice dropped for most, but not all, natural tschermakites because neither Fe<sup>2</sup> nor Al<sup>VI</sup> reach or exceed 1.00. With tschermakite, tschermakitic hornblende, ferro-tschermakite and ferro-tschermakitic hornblende, alumino- and ferri-immediately precede the word tschermakite, e.g. ferro-alumino-tschermakite. Otherwise the order in which prefixes are used is not fixed. Neither ferri- nor ferrian should be used with hastingsite because hastingsite implies high Fe<sup>2+</sup>.

The problem of what to call amphiboles that have Si and, or,  $(Na+K)_A$  in excess of that contained in compositions between tremolite and edenite has not been satisfactorily resolved. Such amphiboles plot near the back left-hand bottom corner of Fig. 1 and have compositions that fall outside the theoretical range of possible substitutions. However, as some such compositions exist it is suggested that they be prefixed, silicic if Si exceeds 7.25 when  $(Na+K)_A \ge 0.50$  but for the compositions involved in which  $(Na+K)_A < 0.50$  no special name is proposed as these compositions are quite close to the names given in Fig. 3A.

This section was approved by 13 votes for and 0 against.

## 4. Sodic-calcic Amphiboles

This group is defined as monoclinic amphiboles in which  $(Ca+Na)_B \ge 1.34$  and  $0.67 \le Na_B \le 1.34$ . Generally  $0.67 \le Ca_B \le 1.34$ .

## End Members

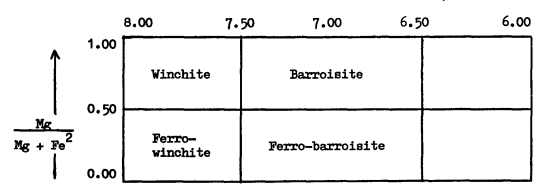
Richterite	${\tt NaCaNaMg}_5{\tt Si}_8{\tt O}_{22}{\tt (OH)}_2$
Ferro-richterite	$\text{NaCaNaFe}_{5}^{2} \text{Si}_{8} \text{O}_{22} \text{(OH)}_{2}$
Ferri-winchite	CanaMg <sub>4</sub> Fe <sup>3</sup> Si <sub>8</sub> O <sub>22</sub> (OH) <sub>2</sub>
Alumino-winchite	$Canamg_4Alsi_6O_{22}(OH)_2$
Ferro-alumino-winchite	$\text{Canafe}_{4}^{2} \text{Alsi}_{8} \text{O}_{22} \text{(OH)}_{2}$
Ferro-ferri-winchite	$\text{Canafe}_{4}^{2}\text{Fe}^{3}\text{Si}_{8}\text{O}_{22}\text{(OH)}_{2}$
Alumino-barroisite	$Canamg_3Al_2Si_7Alo_{22}(OH)_2$
Ferro-alumino-barroisite	$CaNaFe_{\overline{3}}^{2}Al_{2}Si_{\overline{7}}Alo_{22}(OH)_{2}$
Ferri-barroisite	$\operatorname{Canamg}_{3}\operatorname{Fe}_{2}^{3}\operatorname{Si}_{7}\operatorname{AlO}_{22}(\operatorname{OH})_{2}$
Ferro-ferri-barroisite	$CaNaFe_{2}^{2}Fe_{2}^{3}Si_{7}Alo_{22}(OH)_{2}$
Magnesio-ferri-katophorite	$NaCaNaMg_4Fe^3Si_7AlO_{22}(OH)_2$
Magnesio-alumino-katophorite	NaCaNaMg_AlSi_AlO22(OH)2

# Fig. 4. SODIC - CALCIC AMPHIBOLES

 $(Ca+Na)_B \ge 1.34$ ;  $Na_B$  between 0.67 and 1.34

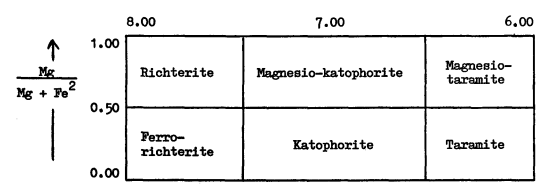
A.  $(Na+K)_A < 0.50$ 

← Si in the standard cell →



B.  $(Na+K)_A \ge 0.50$ 

← Si in the standard cell →



Ferri-katophorite	$NaCaNaFe_4^2 Fe^3 Si_7 Alo_{22} (OH)_2$
Alumino-katophorite	NaCaNaFe4AlSi7AlO22(OH)2
Ferri-taramite	$NaCaNaFe_{3}^{2}Fe_{2}^{3}Si_{6}Al_{2}O_{22}(OH)_{2}$
Magnesio-ferri-taramite	$NaCaNaMg_3Fe_2^5Si_6Al_2O_{22}(OH)_2$
Alumino-taramite	${\tt NaCaNaFe}_3^2{\tt Al}_2{\tt Si}_6{\tt Al}_2{\tt O}_{22}{\tt (OH)}_2$
Magnesio-alumino-taramite	NaCaNaMe3Al2Si6Al2O22(OH)2

## Limits for use of end member names and nomenclatures of the group

The nomenclature of the group is tabulated in Fig. 4. Assignment of the name is as follows: if  $(Na+K)_A < 0.50$  go to Fig. 4A otherwise to Fig. 4B. Si, then the ratio  $Mg/(Fe^2+Mg)$ , and then the  $Al^{VI}$  and  $Fe^2$  values decide the fundamental name of the amphibole. Analyses with  $Al^{VI} \ge 1.00$  or  $Fe^2 \ge 1.00$  have in the name alumino— or ferri— respectively. The final step is dealt with by considering the prefixes already given plus that given below which then gives a name which implicitly or explicitly conveys an indication of the composition with respect to 15 variables.

# Special prefix for the sodic-calcic amphibole group

Alumino when Al<sup>VI</sup> ≥ 1.00

The words alumino- and ferri- immediately precede the fundamental amphibole name (i.e. the noum) otherwise the order in which the prefixes are used is not fixed.

This section was approved by 10 votes for, 2 against and 1 abstention.

## 5. Alkali Amphiboles

This group is defined as monoclinic amphiboles in which  $Na_{R} \ge 1.34$ .

## End Members

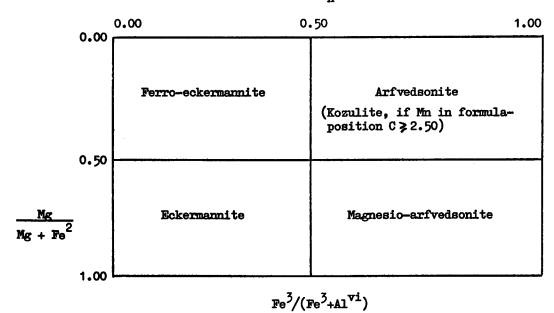
Glaucophane	Na <sub>2</sub> Mg <sub>3</sub> Al <sub>2</sub> Si <sub>8</sub> O <sub>22</sub> (OH) <sub>2</sub>
Ferro-glaucophane	$Na_2Fe_3^2Al_2Si_8O_{22}(OH)_2$
Magnesio-riebeckite	$Na_2Mg_3Fe_2^3Si_8O_{22}(OH)_2$
Riebeckite	$Na_2Fe_3^2Fe_2^3Si_8O_{22}(OH)_2$
Eckermanni te	NaNa2Mg4AlSi8022(OH)2
Ferro-eckermannite	$NaNa_2Fe_4^2AlSi_8O_{22}(OH)_2$
Magnesio-arfvedsonite	$\operatorname{NaNa_2Mg_4Fe^3Si_8O_{22}(OH)_2}$
Arfvedsonite	$NaNa_2Fe_4^2Fe_5^3Si_8O_{22}(OH)_2$
Kozulite	NaNa <sub>2</sub> Mn <sub>A</sub> (Fe <sup>3</sup> ,Al)Si <sub>8</sub> 0 <sub>22</sub> (OH) <sub>2</sub>

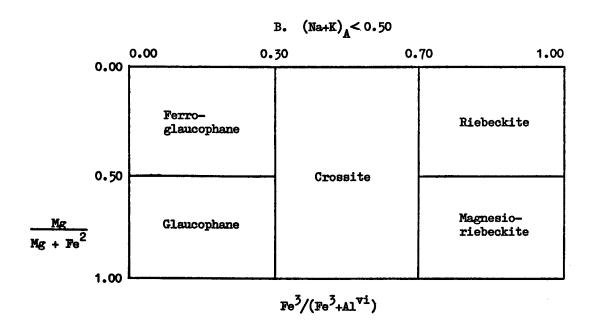
## Limits for use of end member names

The nomenclature of the group is tabulated in Fig. 5. Three factors decide which fundamental name applies; the (Na+K) values (Fig. 5A or 5B) then the ratio

Fig. 5. <u>ALKALI AMPHIBOLES</u>
Na<sub>B</sub>≫ 1.34

A. (Na+K)<sub>A</sub>≥0.50





 ${\rm Fe}^3/({\rm Fe}^3+{\rm Al}^{\rm VI})$  and thirdly the ratio Mg/(Fe $^2+{\rm Mg}$ ). The final step is dealt with by the prefixes already given together with those given below and 16 variables are implicitly or explicitly conveyed by the name — Si, Ca, Ti, F, Cl, K, Ii, Mn, Zn, Cr, OH, O, Fe $^3/({\rm Fe}^3+{\rm Al}^{\rm VI})$ , Pb, Fe $^2/({\rm Fe}^2+{\rm Mg})$  and (Na+K)A. Kozulite is newly described (Nambu et al., 1969).

## Special prefixes for the alkali amphibole group

Calcian when Ca ≥ 0.50 (about 3% Ca0)

Lithian when Li  $\geq$  0.50 (about 1.0% Li<sub>2</sub>0)

The optical variations in this group are so complex and so irregularly related to composition that no formal recommendations regarding them are made at this time. The optical orientations may be indicated conveniently and precisely following Borg's (1967) method by prefixing the symbol G, C, O or R for the four different orientations if it is required to emphasize this aspect.

This section was approved by 12 votes for, 0 against and 1 abstention.

## References

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DEER, W. A., HOWIE, R. A. and ZUSSMAN, J. 1963. Rock-forming minerals. Vol. 2, Longmans, Green, London.

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NAMBU, M., TANIDA, K. and KITAMURA, T. 1969. Kozulite, a new alkali amphibole from Tanohata Mine, Iwate Prefecture, Japan. J. Jap. Ass. Mineral., Petrol., Econ. Geol. 62, 311-328. Abst., Amer. Mineral. 55, p.1815.

## 6. Formal Resolutions adopting the Proposed Amphibole Nomenclature

Throughout, roman superscripts refer to co-ordination numbers and arabic superscripts to charges.

 For the purposes of the following resolutions the standard amphibole formula is taken to contain 8 tetrahedral sites and the general form of the standard formula is:

$$A_{0-1} B_2 C_5^{VI} T_8^{IV} O_{22}(OH,F,C1)_2$$

In the calculation of the standard amphibole formula the following procedure is recommended:

(1) If the water and halogen contents are well established, or if there is physical evidence that the amphibole is an oxy-amphibole, the formula should be calculated to 24(0,0H,F,Cl)

- (2) If the water plus halogen content is uncertain the formula should be calculated on a water-free (and halogen free) basis to 23(0) and 2(OH,F.Cl) assumed.
- (3) Sum T to 8.00 using Si, then Al, then Cr<sup>3</sup>, then Fe<sup>3</sup>, then Ti<sup>4</sup>.
- (4) Sum C to 5.00 using excess Al, Cr, Ti, Fe<sup>3</sup> from (2), then Mg, then Fe<sup>2</sup>, and then Mn.
- (5) Sum B to 2.00 using excess Fe<sup>2</sup>, Mn, Mg from (3), then Ca, then Na.
- (6) Excess Na from (5) is assigned to A, then all K. Total A should be between 0.00 and 1.00, inclusive.
- The iron-magnesium-manganese amphiboles are amphiboles defined by possessing (Ca+Na)<sub>B</sub> < 1.34 in the standard formula.</p>
- The formalised end-member formulae for the orthorhombic members are as follows.

3.1	Magnesio-anthophyllite	${\rm Mg_7Si_80_{22}(OH)_2}$
3.2	Ferro-anthophyllite	$\text{Fe}_7^2 \text{Si}_8 \text{O}_{22} (\text{OH})_2$
3.3	Sodium anthophyllite	$Na(Mg, Fe^2)_7Si_7Al(OH)_7$
3.4	Magnesio-gedrite	${\rm Mg_5^{Al}_2Si_6^{Al}_2O_{22}(OH)_2}$
3.5	Ferro-gedrite	$\text{Fe}_{5}^{2}\text{Al}_{2}\text{Si}_{6}^{2}\text{Al}_{2}^{0}\text{OH})_{2}$
3.6	Sodium gedrite	$Na(Mg, Fe^2)_6 AlSi_6 Al_2 O_{22} (OH)_2$
3.7	Magnesio-holmquistite	$\text{Li}_{2}\text{Mg}_{3}\text{Al}_{2}\text{Si}_{8}\text{O}_{22}\text{(OH)}_{2}$
3.8	Ferro-holmquistite	$\text{Li}_{2}\text{Fe}_{3}^{2}\text{Al}_{2}\text{Si}_{8}^{0}\text{C}_{22}^{2}\text{OH}\text{O}_{2}^{2}$

4.1 <u>Magnesic-anthophyllite</u> is to be used for orthorhombic amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B < 1.34$$
; Li < 1.00; Si  $\ge 7.00$ ; Mg/(Mg+Fe<sup>2</sup>)  $\ge 0.90$ .

4.2 <u>Anthophyllite</u> is to be used for orthorhombic amphiboles chemically defined with respect to the standard formula as follows:

 $(Ca+Na)_B < 1.34$ ; Li < 1.00; Si  $\ge 7.00$ ; Mg/(Mg+Fe<sup>2</sup>) between 0.10 and 0.89 inclusive.

4.3 <u>Ferro-anthophyllite</u> is to be used for orthorhombic amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_{B} < 1.34$$
; Li < 1.00; Si  $\geq 7.00$ ; Mg/(Mg+Fe<sup>2</sup>) < 0.10.

4.4 <u>Magnesio-gedrite</u> is to be used for orthorhombic amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_{B} < 1.34$$
; Li < 1.00; Si < 7.00; Mg/(Mg+Fe<sup>2</sup>)  $\geq$  0.90.

4.5 Gedrite is to be used for orthorhombic amphiboles chemically defined with respect to the standard formula as follows:

 $(Ca+Na)_B < 1.34$ ; Li < 1.00; Si < 7.00; Mg/(Mg+Fe<sup>2</sup>) between 0.10 and 0.89 inclusive.

4.6 Ferro-gedrite is to be used for orthorhombic amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_{R} < 1.34$$
; Li < 1.00; Si < 7.00; Mg/(Mg+Fe<sup>2</sup>) < 0.10.

4.7 <u>Magnesic-holmquistite</u> is to be used for orthorhombic amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_R < 1.34$$
; Li  $\ge 1.00$ ; Mg/(Mg+Fe<sup>2</sup>)  $\ge 0.90$ .

4.8 <u>Ferro-holmquistite</u> is to be used for orthorhombic amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_{R} < 1.34$$
; Li  $\ge 1.00$ ; Mg/(Mg+Fe<sup>2</sup>)  $< 0.10$ .

4.9 <u>Holmquistite</u> is to be used for orthorhombic amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B < 1.34$$
; Li  $\ge 1.00$ ; Mg/(Mg+Fe<sup>2</sup>) between 0.10 and 0.89 inclusive.

- 5.1 The prefix sodium is to be used within the orthorhombic amphibole group for amphiboles with Na≥ 0.50 in the standard formula.
- 5.2 The prefix alumino— is to be used within the anthophyllite subgroup for amphiboles with Al $^{VI} \ge 0.50$  in the standard formula.
- 6. The formalised end-member formulae for the monoclinic members are as follows:

Dannemorite Mn<sub>2</sub>Fe<sub>5</sub>Si<sub>8</sub>O<sub>22</sub>(OH)<sub>2</sub>
 Magnesio-cummingtonite is to be used for monoclinic amphiboles chemically

defined with respect to the standard formula as follows:

$$(Ca+Na)_{B} < 1.34$$
; Li < 1.00; Mn < 0.50; Mg/(Mg+Fe<sup>2</sup>)  $\geq$  0.70.

7.2 <u>Cummingtonite</u> is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B < 1.34$$
; Li < 1.00; Mn < 0.50; Mg/(Mg+Fe<sup>2</sup>) between 0.30 and 0.69 inclusive.

7.3 Grunerite is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B < 1.34$$
; Li < 1.00; Mn < 0.50; Mg/(Mg+Fe<sup>2</sup>) < 0.30.

7.4 <u>Magnesio-clinoholmquistite</u> is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_{B} < 1.34$$
; Li  $\geq 1.00$ ; Mg/(Mg+Fe<sup>2</sup>)  $\geq 0.90$ .

7.5 Ferro-clinoholmquistite is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_{R} < 1.34$$
; Li  $\ge 1.00$ ; Mg/(Mg+Fe<sup>2</sup>)  $\le 0.10$ .

7.6 Clino-holmquistite is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_R < 1.34$$
; Li  $\ge 1.00$ ; Mg/(Mg+Fe<sup>2</sup>) between 0.10 and 0.89 inclusive.

7.7 <u>Tirodite</u> is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B < 1.34$$
; Li < 1.00; Mn  $\ge 0.50$ ; Mg/(Mg+Fe<sup>2</sup>) < 0.50.

7.8 <u>Dannemorite</u> is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B < 1.34$$
; Li < 1.00; Mn  $\ge 0.50$ ; Mg/(Mg+Fe<sup>2</sup>)  $\ge 0.50$ .

- 8.1 The prefix sodian is to be used within the monoclinic iron-magnesium-manganese amphiboles when Na≥0.25 in the standard formula.
- 8.2 The prefix calcian is to be used within the iron-magnesium-manganese amphiboles when Ca≥ 0.50 in the standard formula.
- 9. The calcic amphiboles are monoclinic amphiboles in which the standard formula contains  $(Ca+Na)_B \ge 1.34$  and  $Na_B < 0.67$ . Usually  $Ca_B \ge 1.34$ .
- 10. The formalised end-member formulae are as follows:

10.1	Tremolite	Ca <sub>2</sub> Mg <sub>5</sub> Si <sub>8</sub> O <sub>22</sub> (OH) <sub>2</sub>
10.2	Ferro-actinolite	${^{ m Ca}_2Fe}_5^2{^{ m Si}_80}_{22}{^{ m (OH)}_2}$
10.3	Edenite	${\tt NaCa_2Mg_5Si_7AlO_{22}(OH)_2}$
10.4	Ferro-edenite	$NaCa_2Fe_5^2Si_7AlO_{22}(OH)_2$
10.5	Pargasite	$NaCa_2Mg_4AlSi_6Al_2O_{22}(OH)_2$
10.6	Ferro-pargasite	$NaCa_2Fe_4^2AlSi_6Al_2O_{22}(OH)_2$
10.7	Hastingsite	$NaCa_2Fe_4^2Fe_5^3Si_6Al_2O_{22}(OH)_2$
10.8	Magnesio-hastingsite	$\text{NaCa}_2\text{Mg}_4\text{Fe}^3\text{Si}_6\text{Al}_2\text{O}_{22}\text{(OH)}_2$
10.9	Tschermakite (Alumino-tschermakite)	Ca2Mg3Al2Si6Al2O22(OH)2
10.10	Ferro-alumino-tschermakite	$^{\mathrm{Ca_{2}Fe_{3}^{2}Al_{2}Si_{6}Al_{2}O_{22}(\mathrm{OH})_{2}}$
10.11	Ferri-tschermakite	$^{\text{Ca}_{2}\text{Mg}_{3}^{\text{Fe}_{2}^{3}\text{Si}_{6}^{\text{Al}_{2}^{0}}\text{O}_{22}^{}(\text{OH})_{2}}$
10.12	Ferro-ferri-tschermakite	$^{\text{Ca}_{2}\text{Fe}_{3}^{2}\text{Fe}_{2}^{3}\text{Si}_{6}^{\text{Al}_{2}^{0}}\text{O}_{22}^{}\text{(OH)}_{2}}$
10.13	Magnesio-hornblende	Ca <sub>2</sub> Mg <sub>4</sub> AlSi <sub>7</sub> AlO <sub>22</sub> (OH) <sub>2</sub>
10.14	Ferro-hornblende	$\text{Ca}_{2}\text{Fe}_{4}^{2}\text{AlSi}_{7}\text{AlO}_{22}\text{(OH)}_{2}$

10.15 Kaersutite

$$NaCa_2Mg_4TiSi_6Al_2(O+OH)_{24}$$

10.16 Ferro-kaersutite

$$\mathtt{NaCa_2Fe_4TiSi_6Al_2(0+OH)_{24}}$$

11.1 <u>Tremolite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_{B} \ge 1.34$$
;  $Na_{B} < 0.67$ ;  $(Na+K)_{A} < 0.50$ ;  $Si \ge 7.50$ ;  $Mg/(Mg+Fe^{2}) \ge 0$ .

11.2 <u>Actinolite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A < 0.50$ ;  $Si \ge 7.50$ ;  $Mg/(Mg+Fe^2)$  between 0.50 and 0.89 inclusive.

11.3 <u>Ferro-actinolite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A < 0.50$ ;  $Si \ge 7.50$ ;  $Mg/(Mg+Fe^2) < 0$ .

11.4 <u>Tremolitic hornblende</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B^{\geq} 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A < 0.50$ ;  $Mg/(Mg+Fe^2) \geq 0.90$ ; Si between 7.25 and 7.49 inclusive.

11.5 <u>Actinolitic hornblende</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
; Na<sub>B</sub> < 0.67; (Na+K)<sub>A</sub> < 0.50; Mg/(Mg+Fe<sup>2</sup>) between 0.50 and 0.89 inclusive and Si between 7.25 and 7.49 inclusive.

11.6 <u>Ferro-actinolitic hornblende</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A < 0.50$ ;  $Mg/(Mg+Fe^2) < 0.50$ ; Si between 7.25 and 7.49 inclusive.

11.7 <u>Magnesic-hornblende</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A < 0.50$ ;  $Mg/(Mg+Fe^2) \ge 0.50$ ; Si between 6.50 and 7.24 inclusive.

11.8 Ferro-hornblende is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34; \quad Na_B < 0.67; \quad (Na+K)_A < 0.50; \quad Mg/(Mg+Fe^2) < 0.50;$$
 Si between 6.50 and 7.24 inclusive.

11.9 <u>Tschermakitic hornblende</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_{B} \stackrel{>}{>} 1.34$$
; Na  $< 0.67$ ;  $(Na+K)_{A} < 0.50$ ; Mg/(Mg+Fe<sup>2</sup>)  $\ge 0.50$ ; Si between 6.25 and 6.49 inclusive; Ti < 0.50.

11.10 <u>Ferro-tschermakitic hornblende</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A < 0.50$ ;  $Mg/(Mg+Fe^2) < 0.50$ ; Si between 6.25 and 6.49 inclusive;  $Ti < 0.50$ .

11.11 <u>Tschermakite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A < 0.50$ ;  $Mg/(Mg+Fe^2) \ge 0.50$ ;  $Si < 6.25$ ;  $Ti < 0.50$ .

11.12 Ferro-tschermakite is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_{B} \ge 1.34$$
;  $Na_{B} < 0.67$ ;  $(Na+K)_{A} < 0.50$ ;  $Mg/(Mg+Fe^{2}) < 0.50$ ;  $Si < 6.25$ ;  $Ti < 0.50$ .

11.13 Edenite is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A \ge 0.50$ ;  $Mg/(Mg+Fe^2) \ge 0.50$ ; Si between 6.75 and 7.25 inclusive.

11.14 Ferro-edenite is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A \ge 0.50$ ;  $Mg/(Mg+Fe^2) < 0.50$ ; Si between 6.75 and 7.25 inclusive.

11.15 Edenitic hornblende is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_{B} \ge 1.34$$
;  $Na_{B} < 0.67$ ;  $(Na+K)_{A} \ge 0.50$ ;  $Mg/(Mg+Fe^{2}) \ge 0.50$ ; Si between 6.50 and 6.74 inclusive.

11.16 Ferro-edenitic hornblende is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A \ge 0.50$ ;  $Mg/(Mg+Fe^2) < 0.50$ ; Si between 6.50 and 6.74 inclusive.

11.17 Pargasitic hornblende is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A \ge 0.50$ ;  $Mg/(Mg+Fe^2) \ge 0.70$ ; Si between 6.25 and 6.49 inclusive;  $Ti < 0.50$ ;  $Fe^{5+} \le A1^{VI}$ .

11.18 Ferroan pargasitic hornblende is to be used for amphiboles chemically defined with respect to the standard formula as follows:

11.19 Pargasite is to be used for amphiboles chemically defined with respect to the standard formula as follows:

11.20 <u>Ferroan Pargasite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A \ge 0.50$ ;  $Ma/(Ma+Fe^2)$  between 0.30 and 0.69 inclusive;  $Si < 6.25$ ;  $Ti < 0.50$ ;  $Fe^3 \le Al^{VI}$ .

11.21 <u>Ferro-pargasite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

11.22 <u>Magnesic-hastingsitic hormblende</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_{B} \ge 1.34$$
;  $Na_{B} < 0.67$ ;  $(Na+K)_{A} \ge 0.50$ ;  $Mg/(Mg+Fe^{2}) \ge 0.70$ ; Si between 6.25 and 6.49 inclusive;  $Ti < 0.50$ ;  $Fe^{5} \ge A1^{VI}$ .

11.23 <u>Magnesian hastingsitic hornblende</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A \ge 0.50$ ;  $Mg/(Mg+Fe^2)$  between 0.30 and 0.69 inclusive; Si between 6.25 and 6.49 inclusive;  $Ti < 0.50$ ;  $Fe^2 > Al^{VI}$ .

11.24 <u>Hastingsitic hornblende</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_{B} \ge 1.34$$
;  $Na_{B} < 0.67$ ;  $(Na+K)_{A} \ge 0.50$ ;  $Mg/(Mg+Fe^{2}) < 0.30$ ; Si between 6.25 and 6.49 inclusive;  $Ti < 0.50$ ;  $Fe^{3} \ge AI^{VI}$ .

11.25 <u>Magnesio-hastingsite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A \ge 0.50$ ;  $Mg/(Mg+Fe^2) \ge 0.70$ ;  $Si < 6.25$ ;  $Ti < 0.50$ ;  $Fe^2 > Al^{VI}$ .

11.26 <u>Magnesian hastingsite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A \ge 0.50$ ;  $Ma/(MarFe^2)$  between 0.30 and 0.69 inclusive;  $Si < 6.25$ ;  $Ti < 0.50$ ;  $Fe^2 \ge A1^{VI}$ .

11.27 <u>Hastingsite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $(Na+K)_A \ge 0.50$ ;  $Mg/(Mg+Fe^2) < 0.30$ ;  $Si < 6.25$ ;  $Ti < 0.50$ ;  $Fe^{-1} > Al^{VI}$ .

11.28 <u>Kaersutite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_B \ge 1.34$$
;  $Na_B < 0.67$ ;  $Mg/(Mg+Fe^2) \ge 0.50$ ;  $Si < 6.50$ ;  $Ti \ge 0.50$ .

11.29 <u>Ferro-kaersutite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$(Ca+Na)_{R} \ge 1.34$$
;  $Na_{R} < 0.67$ ;  $Mg/(Mg+Fe^{2}) < 0.50$ ;  $Si < 6.50$ ;  $Ti \ge 0.50$ .

- 12.1 The prefix subcalcic is to be used within the calcic amphibole group for amphiboles with Ca<1.50 in the standard formula.
- 12.2 The prefix alumino- is to be used within the calcic amphibole group for amphiboles with Al in six fold co-ordination ≥1.00 in the standard formula.

- 12.3 The prefix sodian is to be used within the calcic amphibole group for amphiboles with Na≥1.00 in the standard formula.
- 12.4 The prefix silicic is to be used within the calcic amphibole group for amphiboles with Si > 7.25 when (Na+K) $_{\Lambda} \ge 0.50$ .
- 13. The sodic-calcic amphiboles are monoclinic amphiboles in which  $(Ca+Na)_B \ge 1.34$  and  $Na_B$  is between 0.67 and 1.33 inclusive.
- 14. The formalised end member formulae are as follows:

14.1	Alumino-winchite	Canamg Alsi 8022 (OH) 2
14.2	Ferro-alumino-winchite	Canafe <sup>2</sup> AlSi <sub>8</sub> O <sub>22</sub> (OH) <sub>2</sub>
14.3	Ferri-winchite	$\operatorname{Canamg}_{4}\operatorname{Fe}^{3}\operatorname{Si}_{8}\operatorname{O}_{22}(\operatorname{OH})_{2}$
14.4	Ferro-ferri-winchite	$\operatorname{CaNaFe}_{4}^{2}\operatorname{Fe}^{3}\operatorname{Si}_{8}\operatorname{O}_{22}(\operatorname{OH})_{2}$
14.5	Alumino-barroisite	$^{\mathtt{CaNaMg}_{3}\mathtt{Al}_{2}\mathtt{Si}_{7}\mathtt{Alo}_{22}\mathtt{(OH)}_{2}}$
14.6	Ferro-alumino-barroisite	$CaNaFe_3^2Al_2Si_7AlO_{22}(OH)_2$
14.7	Ferri-barroisite	$Caname_{3}Fe_{2}^{3}Si_{7}Alo_{22}(OH)_{2}$
14.8	Ferro-ferri-barroisite	$CaNaFe_{3}^{2}Fe_{2}^{3}Si_{7}Alo_{22}(OH)_{2}$
14.9	Richterite	${\tt NaCaNaMg}_5{\tt Si}_8{\tt O}_{22}{\tt (OH)}_2$
14.10	Ferro-richterite	${\tt NaCaNaFe}_5^2{\tt Si}_8{\tt O}_{22}{\tt (OH)}_2$
14.11	Magnesio-ferri-katophorite	NaCanamg <sub>4</sub> Fe <sup>3</sup> Si <sub>7</sub> AlO <sub>22</sub> (OH) <sub>2</sub>
14.12	Magnesio-alumino-katophorite	${\tt NaCaNa^4Mg_4^AlSi_7^AlO_{22}(OH)_2}$
14.13	Alumino-katophorite	$NaCaNaFe_4^2AlSi_7AlO_{22}(OH)_2$
14.14	Ferri-katophorite	$NaCaNaFe_4^2 Fe_5^3 Si_7^{AlO}_{22}(OH)_2$
14.15	Ferri-taramite	$NaCaNaFe_{2}^{2}Fe_{2}^{3}Si_{6}^{Al}_{2}O_{22}^{OH}_{2}$
14.16	Magnesio-ferri-taramite	NaCaNaMg_Fe2Si6Al2O22(OH)2
14.17	Alumino-taramite	$NaCaNaFe_{3}^{2}Al_{2}Si_{6}^{Al}_{2}O_{22}(OH)_{2}$
14.18	Magnesio-alumino-taramite	NaCaNaMg_Al_Si6Al_O22(OH)2

15.1 <u>Winchite</u> is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

 $(Ca+Na)_B \ge 1.34$ ; Na<sub>B2</sub> between 0.67 and 1.33 inclusive;  $(Na+K)_A < 0.50$ ; Si  $\ge 7.50$ ; Mg/(Mg+Fe<sup>2</sup>)  $\ge 0.50$ .

15.2 <u>Ferro-winchite</u> is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

(Ca+Na)<sub>B</sub>  $\geq$  1.34; Na<sub>B2</sub> between 0.67 and 1.33 inclusive; (Na+K)<sub>A</sub> < 0.50; Si  $\geq$  7.50; Mg/(Mg+Fe<sup>2</sup>) < 0.50.

15.3 <u>Barroisite</u> is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

 $(Ca+Na)_B \ge 1.34$ ; Na<sub>B2</sub> between 0.67 and 1.33 inclusive;  $(Na+K)_A < 0.50$ ; Si < 7.50; Mg/(Mg+Fe<sup>2</sup>)  $\ge 0.50$ .

15.4 <u>Ferro-barroisite</u> is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

 $(Ca+Na)_B \ge 1.34$ ; Na<sub>B2</sub> between 0.67 and 1.33 inclusive;  $(Na+K)_A < 0.50$ ; Si < 7.50; Mg/(Mg+Fe<sup>2</sup>) < 0.50.

15.5 Richterite is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

(Ca+Na)<sub>B</sub> $\geq$ 1.34; Na<sub>B</sub>between 0.67 and 1.33 inclusive; (Na+K)<sub>A</sub> $\geq$ 0.50; Si $\geq$ 7.50; Mg/(Mg+Fe<sup>2</sup>) $\geq$ 0.50.

15.6 <u>Ferro-richterite</u> is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

(Ca+Na)<sub>B</sub>  $\geq$  1.34; Na<sub>B2</sub> between 0.67 and 1.33 inclusive; (Na+K)<sub>A</sub>  $\geq$  0.50; Si  $\geq$  7.50; Mg/(Mg+Fe<sup>2</sup>) < 0.50.

15.7 <u>Magnesio-katophorite</u> is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

 $(Ca+Na)_B \ge 1.34$ ; Na<sub>B</sub> between 0.67 and 1.33 inclusive;  $(Na+K)_A \ge 0.50$ ; Si between 6.50 and 7.49 inclusive;  $Mg/(Mg+Fe^2) \ge 0.50$ .

15.8 <u>Katophorite</u> is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

 $(Ca+Na)_B \ge 1.34$ ; Na\_B between 0.67 and 1.33 inclusive;  $(Na+K)_A \ge 0.50$ ; Si between 6.50 and 7.49 inclusive; Mg/(Mg+Fe<sup>2</sup>) < 0.50.

15.9 <u>Magnesio-taramite</u> is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

(Ca+Na)<sub>B</sub>  $\geq$  1.34; Na<sub>B2</sub> between 0.67 and 1.33 inclusive; (Na+K)<sub>A</sub>  $\geq$  0.50; Si < 6.50; Mg/(Mg+Fe<sup>2</sup>)  $\geq$  0.50.

15.10 Taramite is to be used for monoclinic amphiboles chemically defined with respect to the standard formula as follows:

 $(Ca+Na)_B \ge 1.34$ ; Na<sub>B</sub> between 0.67 and 1.33 inclusive;  $(Na+K)_A \ge 0.50$ ; Si < 6.50; Mg/(Mg+Fe<sup>2</sup>) < 0.50.

- 16. The prefix alumino— is to be used within the soda calcic amphibole group when Al in six fold co-ordination≥ 1.00 in the standard formula.
- The alkali amphiboles are monoclinic amphiboles in which Na<sub>B</sub>≥ 1.34.
- 18. The formalised end member formulae are as follows:

18.1 Glaucophane  $Na_2Mg_3Al_2Si_8O_{22}(OH)_2$ 

18.2 Ferro-glaucophane Na<sub>2</sub>Fe<sub>3</sub>Al<sub>2</sub>Si<sub>8</sub>O<sub>22</sub>(OH)<sub>2</sub>

18.3	Magnesio-riebeckite	$\text{Na}_{2}\text{Mg}_{3}\text{Fe}_{2}^{3}\text{Si}_{8}^{0}_{22}(\text{OH})_{2}^{2}$
18.4	Riebeckite	$Na_2Fe_3^2Fe_2^3Si_80_{22}(OH)_2$
18.5	Eckermannite	$NaNa_2Mg_4AlSi_8O_{22}(OH)_2$
18.6	Ferro-eckermannite	$\operatorname{NaNa_2Fe_4^2AlSi_8O_{22}(OH)_2}$
18.7	Magnesio-arfvedsonite	${\rm NaNa_2Mg_4Fe^3Si_8O_{22}(OH)_2}$
18.8	Arfvedsonite	${\rm NaNa_2Fe_4^2Fe^3Si_8O_{22}(OH)_2}$
18.9	Kozulite	$NaNa_2Mn_4Fe^3Si_8O_{22}(OH)_2$

19.1 Glaucophane is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$Na_B \ge 1.34$$
;  $(Na+K)_A < 0.50$ ;  $Fe^2/(Fe^2+Mg) < 0.50$ ;  $Fe^3/(Fe^3+Al^{VI}) < 0.30$ .

19.2 <u>Ferro-glaucophane</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$Na_B \ge 1.34$$
;  $(Na+K)_A < 0.50$ ;  $Fe^2/(Fe^2+Mg) \ge 0.50$ ;  $Fe^3/(Fe^3+Al^{VI}) < 0.30$ .

19.3 <u>Crossite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

Na<sub>B</sub> 
$$\geq$$
 1.34; (Na+K)<sub>A</sub> < 0.50; Fe<sup>3</sup>/(Fe<sup>3</sup>+Al<sup>VI</sup>) between 0.30 and 0.69 inclusive.

19.4 <u>Magnesio-riebeckite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$Na_{B} \ge 1.34$$
;  $(Na+K)_{A} < 0.50$ ;  $Fe^{2}/(Fe^{2}+Mg) < 0.50$ ;  $Fe^{3}/(Fe^{3}+Al^{VI}) \ge 0.70$ .

19.5 <u>Riebeckite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$Na_{R} \ge 1.34$$
;  $(Na+K)_{A} < 0.50$ ;  $Fe^{2}/(Fe^{2}+Mg) \ge 0.50$ ;  $Fe^{3}/(Fe^{3}+Al^{VI}) \ge 0.70$ .

19.6 <u>Eckermannite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$Na_B \ge 1.34$$
;  $(Na+K)_A \ge 0.50$ ;  $Fe^2/(Fe^2+Mg) < 0.50$ ;  $Fe^3/(Fe^3+Al^{VI}) < 0.50$ .

19.7 <u>Ferro-eckermannite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$Na_B \ge 1.34$$
;  $(Na+K)_A \ge 0.50$ ;  $Fe^2/(Fe^2+Mg) \ge 0.50$ ;  $Fe^3/(Fe^3+Al^{VI}) \le 0.50$ .

19.8 <u>Magnesio-arfvedsonite</u> is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$Na_{B} \ge 1.34$$
;  $(Na+K)_{A} \ge 0.50$ ;  $Fe^{2}/(Fe^{2}+Mg) < 0.50$ ;  $Fe^{3}/(Fe^{3}+Al^{VI}) \ge 0.50$ .

19.9 Arrivedsonite is to be used for amphiboles chemically defined with respect to the standard formula as follows:

s standard formula as follows:  

$$Na_B \ge 1.34$$
;  $(Na+K)_A \ge 0.50$ ;  $Fe^2/(Fe^2+Mg) \ge 0.50$ ;  $Fe^3/(Fe^3+Al^{VI}) \ge 0.50$ ;  
 $Mn_C < 2.50$ .

19.10 Kozulite is to be used for amphiboles chemically defined with respect to the standard formula as follows:

$$Na_{B} \ge 1.34$$
;  $(Na+K)_{A} \ge 0.50$ ;  $Mn^{2}/(Mg+Fe^{2}+Mn^{2}) \ge 0.33$ ;  $Fe^{3}/(Al^{VI}+Fe^{3}) \ge 0.50$ ;  $Mn_{C} \ge 2.50$ .

- 20.1 The prefix calcian is to be used within the alkali amphibole group for amphiboles with Ca≥0.50 in the standard formula.
- 20.2 The prefix lithian is to be used within the alkali amphibole group for amphiboles with Li≥0.50 in the standard formula.
- 21. The following are specified prefixes for the whole amphibole group in terms of contents in the standard formula.
- 21.1 when Cl ≥ 1.00 chlor when Cr ≥ 1.00 21.2 chromium when Cr = 0.25-0.9921.3 chromian when Fe<sup>2</sup> ≥ 1.00 except in alkali amphiboles and hastingsite 21.4 ferri when  $Fe^{3} = 0.75-0.99$  except in alkali amphiboles and hastingsite 21.5 ferrian 21.6 when F ≥ 1.00 fluor 21.7 hydro when OH ≥ 3.00 when Li ≥ 0.25 except in alkali amphiboles when lithian is 21.8 lithian used when Li≥ 0.50. Not used with holmquistite and clinoholmquistite. when Mn ≥ 1.00 except in end-members containing Mn 21.9 manganese 21.10 manganoan when Mn = 0.25-0.99 except in end-members containing Mn 21.11 oxy when (OH+F+Cl) is confirmed as < 1.00 21.12 plumbian when Pb ≥ 0.08 21.13 potassium when K ≥ 0.50 21.14 potassian when K = 0.25 - 0.4921.15 subsilicie when Si < 5.75when Ti ≥ 1.00 except in kaersutite 21.16 titanium 21.17 titanian when Ti = 0.25-0.99 except in kaersutite 21.18 zinc when  $Zn \ge 1.00$ 21.19 zincian when Zn = 0.25-0.99
- 22. Physically identified amphiboles should be named according to the nearest identifiable end-member which should be made into an adjective to be followed by the word amphibole.
- 22.1 <u>Hornblende</u> is to be used for calcic amphiboles identified solely or largely by their physical properties and not confidently identifiable as near to an end-member.

Each part of the above section 6 was voted on separately and received at least 10, and usually 12 or 13 affirmative votes (out of 13) except for sections 11 and 19 which received 9 for, 2 against and 2 abstentions.

## 7. Amphibole Names Recommended for Extinction

It is agreed that the following amphibole names be formally abandoned.

```
Abkhazite
                              = tremolite
Abriachanite
                              = riebeckite
Achromaite
                              - hornblende
                              = actinolite
Actynolin
Actynolite
                              = actinolite
Actinote
                             = actinolite
Aktinolitischer tschermakite = magnesio- or ferro- hornblende
Alkali-femaghastingsite = sodian potassian magnesian hastingsite
Alkali-ferrohastingsite
                             = sodian potassian hastingsite
Alkali-hastingsite
                             = sodian potassian (hastingsite to magnesic-
                                   hastingsite)
Amiant(h)
                              = asbestos
Amianthoide
                              = asbestos
Amianthinite
                              = asbestos
                              = asbestos
Amianthus
Amosite
                              = asbestiform grunerite or anthophyllite pre 1948
Amphibole-anthophyllite
                              = cummingtonite
Amphibolite
                              = hornblende
Anophorite
                              = titanian calcian magnesio-arfvedsonite
Anthogrammatite
                             = anthophyllite
Anthogrammi te
                             = anthophyllite
Antholite
                             = anthophyllite and cummingtonite
Antholith
                             = anthophyllite
Anthophylline
                              = anthophyllite
Anthophyllite rayonné
                             = anthophyllite
Antiglaucophane
                             = glaucophane or crossite
Arfwedsonite
                             = arfvedsonite
Asbeferrite
                             = asbestos
Asbestinite
                             = asbestos
Asbestoide
                             = asbestos
                             = asbestos
Asbestus
Astochite
                             = manganoan richterite
Astorit(e)
                             = richterite
Bababudanite
                             = magnesio-riebeckite
Barkevicite
                             = (sometimes sodian) ferroan or ferro-pargasitic
                                   hornblende, but has been used for other compositions and has never been chemically defined
Barkevikite
Basaltic hornblende
                             = an oxyhormblende, often ferri- or ferrian
                                   titanian (magnesio or magnesian hastingsite)
Basaltine
                              = oxyhormblende + augite
Bedenite
                              = ferrian actinolitic hornblende
Bergamaskite
                              = hastingsite
Bergamaschite
                              = hastingsite
Bergflachs
                              = asbestos
Bergfleisch
                              = asbestos
Berghaar
                              = asbestos
Berghaut
                              = asbestos
Bergholz
                              = asbestos
Berghork
                              = asbestos
                              = asbestos
Bergpapier
                              = asbestos
Bergwolle
Bidalotite
                                 gedrite
                                 sodium amphibole
Borgmiezite
Breadalbanite
                              = hornblende
Byssolite
                              = asbestos
Calamite
                              = tremolite
```

= hornblende, often pargasitic hornblende Carinthine

Carystine = asbestos Cataphorite = kataphorite = katophorite Catophorite = katophorite Cataforite = katophorite Chernyshevite = sodium amphibole

Chiklite = manganoan ferri-ferro-richterite

= tremolite or actinolite Chrome-tremolite = magnesio-cummingtonite Clino-anthophyllite Clinokupfferite = cummingtonite Crocidolite = asbestiform riebeckite Daschkesanit = chlor potassian hastingsite Dashke(s)sanite = chlor potassian hastingsite

Diastatite = hornblende Eckrite = winchite Eisenrichterite = ferro-richterite Fasciculite = hornblende Femaghastingsite = magnesian hastingsite Feranthophyllite = ferro-anthophyllite

= sodian manganoan magnesio-hastingsite Ferrian pargasite

Ferri- edenite = ferro-edenite = magnesio-riebeckite Ferriglaucophane Ferrihedrite = ferri-gedrite

Ferririchterite manganoan magnesio-arfvedsonite

Ferri-tremolite = ferri-ferro-actinolite Ferrohastingsite = hastingsite

Ferro-tremolite = ferro-actinolite Gamsigradite = manganoan (magnesio-hornblende or edenite)

Gastaldite = glaucophane

Girnari te = subsilicic titanian sodian magnesian hastingsite

Grammatite = tremolite Grammatit-strahlstein = tremolite = crocidolite Griqualandite = grunerite Grünerite Heikolite = crossite Heikkolite = crossite

Heterotype = amphibole and pyroxene

Hexabolit = oxyhormblende = manganoan tremolite Hexagonite Hillängeite = dannemorite Hoepfnerite = tremolite

Holzasbest = asbestos Hudsonite = hastingsite

Imerinite = magnesio-arfvedsonite Iron-anthophyllite = ferro-anthophyllite

= oxy-manganoan potassian ferrian ferro-hornblende Iron-hornblende

Iron-richterite = ferro-richterite Isabellite = richterite

Juddite = manganoan magnesio-arfvedsonite

Kalamite

= tremolite = titanian Kalio-magnesio-katophorite titanian potassian richterite

Karinthin = hornblende, often pargasitic hornblende

Kidney stone = actinolite Kievite = cummingtonite

Kirwanite = impure altered amphibole Kokscharowit = edenitic amphibole Kokscharovite = edenitic amphibole Krokidolite = crocidolite Krokvdolith = crocidolite

Kupfferite (Allen & Clement) = magnesio-anthophyllite

Kupfferite (Hermann) - chromian anthophyllite

Kupfferite (Koksharov) = chromian anthophyllitic amphibole

**Kymatine** = asbestos Labrador hornblende = orthopyroxene

oxyhornblende Lamprobolite

m ferroan or ferro-pargasitic hornblende Laneite

= ferri or ferrian oxy kaersutite
= holmquistite Linosite

Lithionglaukophan

= lithian amphibole, holmquistite and clino-Lithium-amphibole holmquistite

Maganthophyllite = magnesio-anthophyllite Magnesia-arfvedsonite = magnesio-anfvedsonite

= glaucophane Magnesian glaucophane

= titanian potassian richterite Magnophorite = magnesio-anthophyllite Magnesium anthophyllite = mangananoan actinolite Mangan-actinolite Mangan amphibole = rhodonite (not an amphibole) = manganoan riebeckite Mangan crocidolite

Mangan krokidolith = manganoan riebeckite

Mangano-anthophyllite = tirodite

Mangan-tremolite = manganoan tremolite

Mangamuralite = manganoan magnesio-arfvedsonite

Marmairolite = manganoan richterite Mboziite = potassian taramite Mountain wood = asbestos Montasite = asbestiform grunerite

= richterite Natrongrammatit

Natronrichterite = manganoan richterite Naurodite = alkali amphibole Nephrite = actinolite = ferro-hornblende Noralite Nordenskiöldite tremolite Orniblende hornblende Orthoriebeckite = riebeckite

= riebeckite Osannite Philipstadite = ferrian ferro-hornblende Picroamosite = ferrian anthophyllite = actinolite pseudomorph Pilite

Pseudoglaucophane = glaucophane or crossite Prismatic schillerspar = anthophyllite

Raphilite = tremolite

= magnesio-riebeckite or magnesio-arfvedsonite Rezhikite

= magnesio-riebeckite Rhodusite

Rimpylite hornblende Sebesite tremolite = Silbölite = actinolite Sillbölite = actinolite Silfbergite = dannemorite

= titanian potassian richterite Simpsonite Smaragdite = actinolite or hornblende

Smaragditic grammatite = tremolite

= tschermakite or tschermakitic hornblende Smaragditic techermakite

Soda asbestos = magnesio-arfvedsonite Soda hornblende = arfvedsonite = manganoan richterite Soda richterite Soda tremolite = richterite = magnesian hastingsite Soretite

Speziatite = hornblende Strahlstein = actinolite

Strelite = actinolite or anthophyllite Subglaucophane = crossite

Svidneite = oxy magnesio-riebeckite Syntagmatite (Troger 1952) = titanian hastingsite

Szechenyiite = richterite Szechonyit = richterite

Ternovskite = magnesio-riebeckite

Thalackerite = anthophyllite

Tibergite = manganoan sodian magnesio-hastingsite

Titanhornblende = aenigmatite
Tonerdehaltiger strahlstein = tremolite

Torendrikite = magnesio-riebeckite

Tremolite-glaucophane = richterite
Tschernischewit = sodium amphibole
Uralite = actinolite pseudomorph

Valleite = calcian manganoan anthophyllite

Waldheimite = richterite
Wallerian = hornblende

Weinschenkite = ferri-magnesio-hornblende or magnesio-hastingsite

Zillerite = actinolite
Zillerthite = actinolite
Zino-manganese-cummingtonite = zinc tirodite

M. H. Hey (1962 and appendix 1963), <u>Index to mineral species and varieties arranged chemically</u> should be consulted for further details of the above names.

This section was approved by 13 votes for, 0 against.

The compiler particularly draws the attention of mineralogists to the abandonment of barkevikite, basaltic hornblende, carinthine, ferrohastingsite, grammatite, karinthine, kataphorite and mboziite as these names are more commonly used than the remainder.

The compiler comments that the main practical difficulty in naming amphiboles by the agreed procedure is that the ratio Mg/(Mg+Fe²) cannot be accurately obtained from electron microprobe analysis. Agreement to use Mg/(Mg+Fe²+Fe³) could not be obtained and so it will be essential to examine critically the procedure adopted to calculate Fe² and Fe³ when only the total Fe has been determined. Different procedures could give different names to same chemical analysis. In addition, in view of the very large number of incorrectly calculated standard amphibole formulae in the literature, authors are urged to always calculate these carefully, never to avoid checking that the positive and negative charges balance and that the determined oxides have been precisely transcribed — a common error in computer-calculated results. The whole procedure including outputting the full name will be most conveniently dealt with by one computer programme.

Addendum to The nomenclature of amphiboles (by B. E. Leake and M. H. Hey)

The foregoing report is directed to the formulation of a set of species names with clearly defined limits, together with a system of prefixes and adjectival modifiers (see below). An appropriate name can be found for any amphibole for which a complete chemical analysis is available, including the  $Fe^{2+}/Fe^{3+}$  ratio; if this ratio has not been determined various procedures are possible, involving special assumptions: thus we may calculate on the basis of 13 cations exclusive of Ca, Na, and K, assuming that these cations are wholly in the A and B sites, and then adjust the  $Fe^{2+}/Fe^{3+}$  ratio to

bring O+OH to 24 (or, if H<sub>2</sub>O has not been determined or is unreliable, to bring O to 23) [this order of calculation is simpler and safer than that suggested on p. 3 of the report, which involves adjustment of all the cation ratios].

But we also need to be able to name amphiboles for which no chemical analysis is available, or only a partial one; the paragenesis and the physical properties, especially colour, refractive indices, extinction angle, and pleochroism, offer suffice to place the mineral, but only approximately. The problem is no new one; for example, anthophyllite is widely used both for the Al-free or Al-poor orthoamphibole, but also as a group name covering both anthophyllite sensu stricto and gedrite.

The Subcommittee recommend (p. 4, last paragraph; resolution 22) that where an unanalysed amphibole can be placed, on the strength of its physical properties and paragenesis as, for example, 'not far from richterite', it should be termed a 'richteritic amphibole', et sim. No group names are recommended.

Though some 51 'end-members' or species are recognized, multiplication of trivial names is avoided (they are kept to 23) by the use of prefixes, which indicate substantial amounts of substitution by various elements. These are an inseparable part of the name, and should be attached by a hyphen; they should not normally be translated, and should not be separated in indexing (thus, for example, Ca<sub>2</sub>Mg<sub>3</sub>Mn<sub>2</sub>Si<sub>8</sub>O<sub>22</sub>(OH)<sub>2</sub>, a manganese-tremolite, should be indexed under M, with perhaps a cross-reference under tremolite). [In languages such as German, where all nouns are capitalized, the form Mangan-tremolit or Mangantremolit might be preferred.]

These true *prefixes* are: alumino-, chlor-, chromium-, ferri-, ferro-, fluor-, hydro-, manganese-, magnesio-, oxy-, potassium-, titanium-, sodium-, and zinc-.

On the other hand there are a set of 'prefixes', more correctly termed adjectival modifiers, usually ending in -ian or -oan (according to the valency) denoting minor substitutions, and two denoting deficiencies, which are all simple adjectives. These are not an essential part of the name, and should be ignored in the first stage of indexing (e.g. manganoan tremolite should be indexed under T, as tremolite, manganoan. These adjectives may properly be translated by appropriate adjectives, calcian = calcifère (French) = kalkhaltig (German); with those that incorporate an indication of valency (ferrian, manganoan, et sim.) this may call for special treatment (e.g. ferroan = eisen(II)haltig).

These adjectival modifiers include: chromian, ferrian, ferroan, lithian, manganoan, plumbian, potassian, titanian, zincian, also subsilicic and subcalcic. Additional adjectives may be formed on this model as needed, e.g. nickeloan, cuprian.

## Index of names and prefixes

The following index refers to the list of formal definitions in the numbered resolutions of the Report (pp. 16-25). The two special adjectival modifiers subcalcic and subsilicic are included, as they too are formally defined. Limits are proposed in the Report (pp. 3-4) for the use of the adjectival

modifiers chromian, ferrian, lithian, manganoan, plumbian, potassian, titanian, and zincian.

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page 534, step (5) of the calculation of the standard formula should read: (5) Sum B to 2 00 using excess Fe<sup>2</sup>, Mn, Mg from (4), . . .. page 536, fig. 1, the point labelled 0·67 Na<sub>B</sub> along the left-hand back edge should read: 1·34 Na<sub>B</sub>. page 538, resolution 3.3, for (OH)<sub>7</sub> read (OH)<sub>2</sub>. page 549, resolution 6.4, for Fe<sub>3</sub> read Fe<sup>2</sup><sub>3</sub>. page 549, resolution 6.6, for Fe<sub>5</sub> read Fe<sup>2</sup><sub>5</sub>. page 551, resolution 11.1 should conclude: Mg/(Mg+Fe<sup>2</sup>) ≥ 0·90. page 551, resolution 11.3 should conclude: Mg/(Mg+Fe<sup>2</sup>) < 0·49. page 554, resolution 14.12, for NaCaNa<sup>4</sup> read NaCaNa.