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Nomenclature of Pyroxenes

Subcommitee on Pyroxenes Commission on New Minerals and Mineral Names (CNMMN) International Mineral Association (IMA)

by N. Morimoto, Chairman¹

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Abstract

This is the final report on the nomenclature of pyroxenes by the Subcommittee on Pyroxenes established by the Commission on New Minerals and Mineral Names of the International Mineralogical Association. The recommendations of the Subcommittee as put forward in this report have been formally accepted by the Commission. Accepted and widely used names have been chemically defined, by combining new and conventional methods, to agree as far as possible with the consensus of present use. Twenty names are formally accepted, among which thirteen are used to represent the end members of definite chemical compositions. In common binary solid-solution series, species names are given to the two end members by the "50% rule". Adjectival modifiers for pyroxene mineral names are defined to indicate unusual amounts of chemical constituents. This report includes a list of 105 previously used pyroxene names that have been formally discarded by the Commission.

Keywords: Pyroxenes, nomenclature, chemical composition, solid-solution series.

1. Introduction

The subcommittee on pyroxenes has, after a thorough evaluation of the group of pyroxene minerals, presented its recommendations for a new classification and nomenclature to the Commission on New Minerals and Mineral Names (hereafter abbreviated as CNMMN). These recommendations have been approved by the commission by a formal vote (20th May, 1987).

The classification and nomenclature of the pyroxenes have been largely based on their

crystal chemistry. In practice the chemical content of the pyroxene formula unit calculated to six oxygens, or to four cations (VIETEN and HAMM, 1978), is essential for the classification. This formula unit corresponds to one quarter of the unit cell for the monoclinic pyroxenes and to one eighth of the unit cell for the orthorhombic pyroxenes. The basic principle adopted for amphibole nomenclature (LEAKE and WINCHELL, 1978) is to denote principal stoichiometries by generally well-established names, with adjectival modifiers to indicate the presence of substantial substitutions that are

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not essential constituents of the end-members; this has been followed as far as possible in the pyroxene nomenclature.

No new names have been introduced in the proposed nomenclature. Accepted and widely used names have been chemically defined by combining new and conventional methods to agree as far as possible with the consensus of present use. Two kinds of adjectival modifiers are used: one to specify a part of the compositional range shown by a mineral that forms a wide solid solution (e.g. magnesium-rich augite and iron-rich augite); the other to specify elemental substitutions that are not essential constituents (e.g. titanian augite). 105 previously used pyroxene names, mostly synonyms, obsolete or almost unused, recommended for rejection, have formally been discredited by the CNMMN.

General publications dealing with the pyroxene group include Rock-Forming Minerals (DEER et al., 1978) (herafter DHZ), the Special Papers (ed. Papike, 1969) and Reviews in Mineralogy (ed. Prewitt, 1980) of the Mineralogical Society of America, which provide references to the voluminous literature.

2. Crystal chemistry of the pyroxenes

Pyroxenes are silicates that, in their simplest form, contain single SiO₃ chains of linked SiO₄ tetrahedra. Generally, small amounts of Si are replaced by Al and other small cations. The repeat along the chain (c axis) comprises two tetrahedra and is approximately 0.52 nm in length. The general chemical formula (formula unit) for all pyroxenes¹⁾ is M2M1T₂O₆, where M2 refers to cations in a generally distorted octahedral coordination, M1 to cations in a regular octahedral coordination, and T to tetrahedrally coordinated cations.

Any pyroxene belongs to either the orthorhombic or the monoclinic crystal system. There are two orthorhombic pyroxene types: orthopyroxene (Pbca) and orthopyroxene (Pbcn)²). Only the former has ben found in nature. Monoclinic pyroxenes are called clinopyroxenes. Their space groups are C2/c, $P2_1/c$

and P2/n, depending on their chemical composition and genetic history.

Throughout this report, the standard pyroxene formula is used with superscripted arabic numerals (e.g. Fe²⁺) referring to charges, and subscripted numerals (e.g. Mg₂) to numbers of atoms.

In order to derive a pyroxene formula from a chemical analysis, the calculation should be based on six oxygen atoms, when Fe²⁺ and Fe³⁺ are both determined. In microprobe analyses, only total Fe is determined and the option of calculating to four cations should at least be permitted if not actually preferred. VIETEN and HAMM (1978) show that calculation to four cations will be more reliable for microprobe analyses of the majority of pyroxenes. Therefore, for microprobe analyses it is recommended that the components be totalled to six oxygens and four cations by adjusting the ratios Fe²⁺/Fe³⁺, Ti⁴⁺/Ti³⁺, etc.

The standard pyroxene formula M2M1T₂O₆ contains two tetrahedral sites. In the allocation of the cations to obtain a pyroxene formula, the following procedure is recommended:

- 1) Sum T to 2.000 using Si^{4+} , then Al^{3+} , then Fe^{3+} .
- 2) Sum M1 to 1.000 using all Al³⁺ and Fe³⁺ in excess of that used to fill the T sites. If there is insufficient Al³⁺ and Fe³⁺ to sum to 1.000, then add Ti⁴⁺, Cr³⁺, V³⁺, Ti³⁺, Zr⁴⁺, Sc³⁺, Zn²⁺, Mg²⁺, Fe²⁺ and finally Mn²⁺ until the sum is 1.000.
- 3) Sum M2 using all Mg²⁺, Fe²⁺ and Mn²⁺ in excess of that used to fill the M1 sites. Then add Li⁺, Ca²⁺ and Na⁺ so that the sum becomes 1.000 or close to it. If the sum is far from 1.000, one must be suspicious about the results of the analysis.

A flow chart (Fig. 1) gives a diagrammatic representation of the site allocation of the principal cations in pyroxenes. However, because the distribution of cations among the M1, M2 and T sites in a given pyroxene is partly a function of temperature, the accurate site occupancy must be determined by structure determination. The site occupancy given in

¹⁾ In omphacite-P2/n, the M1 and M2 sites are further divided into M1a and M1b (for M1) and M2a and M2b (for M2).

²⁾ Orthopyroxene (*Pbcn*) is stable only at elevated temperatures for a limited composition near MgSiO₃.

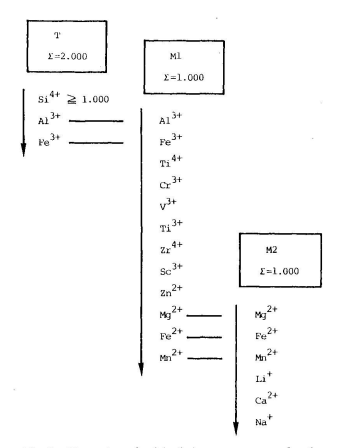


Fig. 1 Flow chart for ideal site occupancy of cations between the T, M1 and M2 sites of pyroxenes. Only representative cations are included. Arrows indicate order of filling of sites. Real site occupancy is usually slightly different from the ideal site occupancy.

Fig. 1 is called ideal site occupancy to distinguish it from real occupancy. A method for classifying pyroxenes by their ideal site occupancies has been proposed by Boku and Ginzburg (1985). In the present classification of pyroxenes, the M1 and M2 sites are considered together as a single M site in order to avoid the difference between the real and ideal site occupancies.

Starting from the most common pyroxene formula, $M2(R^{2+})M1(R^{2+})T_2(2R^{4+})O_6$, four coupled substitutions are possible if one assumes more than one R^{4+} in the T site. They are listed in Table 1 (p. 100), where the elements in parentheses are coupled substitutions.

Substitution (1) encompasses the endmembers jadeite (NaAlSi₂O₆), aegirine³⁾ (NaFe³⁺Si₂O₆), kosmochlor⁴⁾ (NaCr³⁺Si₂O₆), and jervisite (NaScSi₂O₆). Substitution (2) results in components such as NaFe²⁺ $_{0.5}$ Ti⁴⁺ $_{0.5}$ Si₂O₆, but is less important than the other substitutions.

In substitution (3) the Al-Al couple is often referred to as "Tschermak's component"; CaAlAlSiO₆, in particular, is called "calcium Tschermak's component". Substitution in esseneite5), CaFe3+AlSiO6, is obtained by this type of substitution. This substitution is also important in "fassaite" 6). Substitution resulting in CaTi3+AlSiO6 was reported by Dowty and Clark (1973) and Mason (1974) in pyroxenes from the Allende meteorite (Table 3, No. 4). In substitution (4) the component CaMg_{0.5}Ti⁴⁺_{0.5}AlSiO₆ is found in some pyroxenes. There are a few instances of the component of substitution (2) or (4) amounting to nearly 50%, as described later (Table 3). However, no particular names are given for the end member components of substitutions (2) and **(4)**.

3. Mineral names of the pyroxenes

3.1. TWENTY (20) MINERAL NAMES AND THEIR GROUPING

The pyroxenes form extensive solid solutions by various types of ionic substitutions, some of which are described above. To cope with the problem of pyroxene nomenclature, it is necessary to subdivide the solid-solution series into ranges with specified compositions and names. Whenever there is a complete solid-solution series between two end members, it is customary in mineral nomenclature to use only two names, and the division between them

³⁾ Aegirine is used in preference to "acmite" in this report. It is in common usage in the literature, and is consistent with the almost universal use of aegi-

rine-augite for minerals of intermediate compositions, though "acmite" has priority by 14 years (Dana, 1892). Common practice in experimental petrology has been to use the abbreviation Ac for NaFe³⁺Si₂O₆; Ae should now be used instead.

⁴⁾ The CNMMN, IMA, has recently voted in favour of the name kosmochlor instead of "ureyite" for the pyroxene of generalized composition NaCrSi₂O₆.

⁵⁾ Esseneite is a new pyroxene with the composition CaFe³⁺AlSiO₆ (Table 2, No. 13).

^{6) &}quot;Fassaite" has the general formula Ca(Mg,Fe³⁺,Al) (Si,Al)₂O₆. This name has been rejected as a formal name in this report.

	<u> </u>	and the second second		
Substitution site	М2	м1	T .	examples
standard	_R 2+	_R 2+	2R ⁴⁺	
substitution(1)	(R ⁺)	(R ³⁺)	2R ⁴⁺	Na-Al Na-Fe3+ Na-Cr3+ Na-Sc3+
substitution(2)	(R ⁺)	$R_{0.5}^{2+}(R_{0.5}^{4+})$	2R ⁴⁺	Na-(Ti ⁴⁺ /2)
substitution(3)	_R 2+	(R ³⁺)	(R ³⁺)R ⁴⁺	A1-A1 Fe ³⁺ -A1 Cr ³⁺ -A1
Substitution(4)	R ²⁺	$R_{0.5}^{2+}(R_{0.5}^{4+})$	(R ³⁺)R ⁴⁺	(Ti ⁴⁺ /2)-Al

Tab. 1 Four coupled substitutions of pyroxenes in the standard chemical formula $R^{2+}R^{2+}R^{4+}O_6$.

should be at A₅₀B₅₀ (the "50% rule"). However, this "50% rule" cannot be applied rigorously to the large groups of pyroxenes which show wide ranges of coupled substitutions. This is particularly so when the minerals concerned are abundant and widespread, and have a historically-established nomenclature in mineralogical and petrological circles. Taking this situation into consideration, 20 accepted and widely used names have been adopted as mineral species names of the pyroxenes (Table 2).

The definition of the pyroxene species has been based on thirteen end-members, or chemical components, listed in Table 2, and the component Ca₂Si₂O₆ (Wo)⁷⁾. These end-members are given the names of the minerals whose compositions they most closely approximate. The 20 pyroxene species are grouped into six chemical subdivisions on the basis of the cation occupancy of the M2 sites and crystal chemical similarity. This classification is a slight modification of the widely used scheme proposed by DHZ (1978).

For the precise classification of the pyroxenes into 20 mineral species, however, the following characteristics of the pyroxenes must be considered. First of all, the Mg-Fe pyroxenes and some of the Ca pyroxenes are the most common rock-forming pyroxenes and form wide solid solutions which cover the pyroxene quadrilateral of the ternary Ca₂Si₂O₆ (Wo) -Mg₂Si₂O₆ (En) - Fe₂Si₂O₆ (Fs) system. Therefore, these pyroxenes are better treated together as the Ca-Mg-Fe or "quadrilateral" pyroxenes. Secondly, Na pyroxenes form continuous solid-solution series with the Ca-Mg-Fe pyroxenes, forming the Na-Ca pyroxenes. Thirdly, donpeacorite and kanoite in the Mn-Mg pyroxenes, johannsenite, petedunnite and esseneite in the Ca pyroxenes, and spodumene are rare in occurrence and unique in chemistry. For simplicity they are treated together as 'other' pyroxenes⁸).

All the pyroxenes are thus divided into four chemical groups for the purpose of broad classification: Ca-Mg-Fe pyroxenes (Quad, 8), Ca-Na pyroxenes (Ca-Na, 2), Na pyroxenes (Na, 2) and 'other' pyroxenes (Others, 8). The abbreviations of the groups and the numbers of the accepted species are given between paren-

⁷⁾ $\text{Ca}_2\text{Si}_2\text{O}_6$ exists as wollastonite in nature, which belongs not to the pyroxenes but to the pyroxenoids. To represent the compositions of the Ca-Mg-Fe pyroxenes, the ternary $\text{Ca}_2\text{Si}_2\text{O}_6(\text{Wo})$ – $\text{Mg}_2\text{Si}_2\text{O}_6(\text{En})$ – Fe_2 $\text{Si}_2\text{O}_6(\text{Fs})$ system has been used, e.g. $\text{En}_{20}\text{Fs}_{38}\text{Wo}_{42}$.

⁸⁾ Definition of the 'Other' pyroxenes is different from that given by CAMERON and PAPIKE (1981).

theses. Quad represents "quadrilateral" for the Ca-Mg-Fe pyroxenes. The four chemical groups are further divided into 20 mineral species by using 12 components (using the Wo component for Di and Hd components). The composition ranges for the accepted names will be given later.

The pyroxene names may be qualified by one or more adjectival modifiers according to definite rules described later to specify important (though relatively minor) departures from the composition ranges. When the composition range of the mineral species is large, as in augite, one or more adjectival modifiers are used to specify the composition more clearly (e.g. subcalcic augite, Fe-rich augite).

3.2. APPLICATION OF 50% RULE

The 50% rule has been applied to complete solid-solution series between two end members as far as possible. They are the Mg-Fe pyroxene series (enstatite-ferrosilite and clinoenstatiteclinoferrosilite series), Ca pyroxene series (diopside-hedenbergite series) and Na pyroxene series (jadeite-aegirine series). Subdivision names of the intermediate solid solution ranges, such as bronzite, hypersthene and eulite of the enstatite-ferrosilite series and salite and ferrosalite of the diopside-hedenbergite series, have been discarded. However, the 50% rule was not applied rigorously to the Ca-Mg-Fe pyroxenes and Na-Ca pyroxenes. The widely accepted terms such as augite, pigeonite, omphacite and aegirine-augite9) have been retained.

3.3. GEM NAMES OF SPODUMENE

Two names, "hiddenite" and "kunzite", are often used respectively for (pale) emerald-green and lilac colored spodumene of gem quality. They are not accepted as formal pyroxene names, but can be used as varietal gem names.

3.4. RELATIONSHIPS WITH THE PYROXENOIDS

Pyroxenoids are closely related to pyroxenes in that they have a similar type of chemical composition and a structure that also consists of SiO₃ single chains. However, the repeat of the chains, which is two SiO₄ tetrahedra in the pyroxenes, is three or more SiO₄ tetrahedra in the pyroxenoids. Though the tetrahedral sites are mostly occupied by Si ions, the large cations are mostly Ca, Mn and Fe²⁺ ions in the pyroxenoids. The classification and nomenclature of the pyroxenoids are beyond the scope of this report. However, the following two points may be noted. Firstly, there is a polymorphic relationship with some pyroxenes such as ferrosolite, hedenbergite and johannsenite. These show pyroxenoid structures at high temperatures or pressures. Secondly, the wollastonite chemical component (Ca₂Si₂O₆) is used to express the composition of the Ca-Mg-Fe pyroxenes, though wollastonite belongs to the pyroxenoid structural group.

4. Classification and nomenclature of the pyroxenes

4.1. PRELIMINARY CLASSIFICATIONS - CONSTRUCTION OF THE Q-J DIAGRAM AND APPLICATION OF PYROXENE DATA -

Before classifying the pyroxenes into the 20 mineral species listed in Table 2, the following procedure is recommended to divide them into four chemical groups: Ca-Mg-Fe pyroxenes (Quad), Na-Ca pyroxenes (Na-Ca), Na pyroxenes (Na), and other pyroxenes (Others) (Morimoto and Kitamura, 1983).

In this procedure the pyroxenes are classified by using the total numbers of specified cations at the M (M1 and M2) sites on the basis of six oxygens. The M1 and M2 sites are considered together as M sites, without considering the site preference of atoms between the two sites.

The numbers of Ca, Mg, Fe²⁺ and Na cations in the M sites are plotted in the Q-J diagram (Fig. 2) as Q=Ca+Mg+Fe²⁺ and J=2Na. The lines representing the following equations are used to subdivide the Q-J diagram:

- (1) Q + J = 2.0
- (2) Q + J = 1.5
- (3) J/(Q + J) = 0.2
- (4) J/(Q + J) = 0.8

The areas corresponding to the Ca-Mg-Fe pyroxenes, Ca-Na pyroxenes, Na pyroxenes

⁹⁾ The name aegirine-augite appears to be in more common usage than "aegirineaugite", and "acmite-augite".

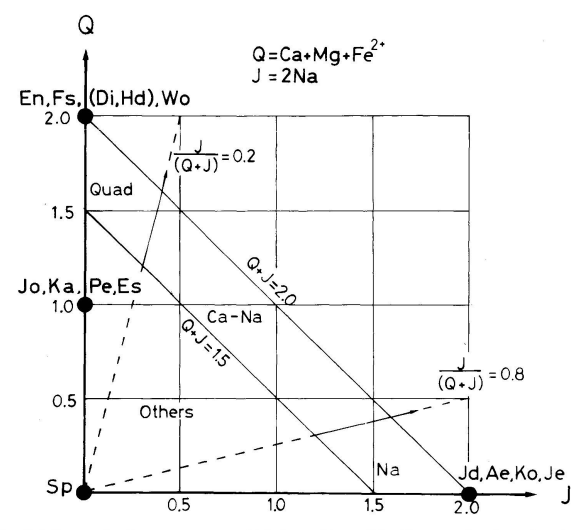


Fig. 2: Q-J diagram for the pyroxenes, on which the positions of the 13 accepted end-members have been indicated. Abbreviations and compositions of the end-members are listed in Table 2.

and other pyroxenes are labeled (Fig. 2) Quad, Ca-Na, Na, and Others, respectively.

In this diagram, J is meant to include the total number of Na and R³⁺, usually Al, Fe³⁺, Cr3+ and Sc3+, that couple with Na in substitution (1) mentioned in Table 1. When the coupling substitution in the pyroxene is not of type (1), but of type (2) or (3), the J value apparently does not represent the real numbers of Na and R^{3+} at the M sites. However, substitution (3) (e.g. Al-Al) works to move the J and Q values closer to the origin of the Q-J diagram, and substitution (2) (e.g. Na-Ti⁴⁺) to move the J value farther away from the O axis of ordinates. Therefore, the effects of substitutions (2) and (3) tend to cancel each other out in and near the Na pyroxenes area. Thus the J(=2Na) values in the Na-rich pyroxenes represent, to a good approximation, the total number of Na and $R^{3+}(Al, Fe^{3+}, Cr^{3+} \text{ and } Sc^{3+})$ at the M sites.

The boundary Q+J=2.0 represents the upper limit of Q+J at the M sites. The boundary Q+J=1.5 represents the limit below which more than half of the M1 or M2 sites may be occupied by ions other than Q and J ions. In this case, the pyroxenes are considered as belonging to 'Others', which include the Mn-Mg and Li pyroxenes, johannsenite, petedunnite and esseneite. The third and fourth equations represent the lines dividing the area limited by the two above-mentioned Q+J lines into Ca+Mg+Fe (Quad), Ca-Na and Na pyroxenes. The boundaries defined by J/(Q+J)=0.2 and 0.8 are used by DHZ (1978) and CAMERON and PAPIKE (1981).

Because the Mn-Mg pyroxenes and johannsenite (Table 2) have Mn ions occupying more than half of the M2 and M1 sites, respectively, they appear along the Q axis between 1.0 and 1.5 of the Q value in the Q-J diagram. Similarly, petedunnite and esseneite appear along the Q axis with its Q value between 1.0 and 1.5. Spodumene concentrates at the origin of the Q-J diagram because both Q and J are zero. Thus, the thirteen end-members (Table 2) and Wo are located in the Q-J diagram (Fig. 2).

Application of this classification procedure to 406 pyroxene analyses presented in DHZ has shown that most of the analyses, except those of johannsenite and spodumene, are included in the area between the lines Q+J=2.0 and 1.5. The 103 DHZ pyroxenes selected by CAMERON and PAPIKE (1981), for which the Q-

values are less than 1.90 and Mn is less than 0.08 atoms per formula unit, are plotted in the Q-J diagram of Fig. 3. The 'CaMgTAL' pyroxene (CAMERON and PAPIKE, 1981) is included in the *Quad* area as described later (Table 3, No. 1). Only twenty analyses among 406 plot slightly over the line Q+J=2.0, and most of these show unusual total numbers of cations. The results of the classification of the pyroxenes into the four chemical groups by this procedure are in almost complete agreement with the results obtained by DHZ (1978) and by CAMERON and PAPIKE (1981). A few unusual

Tab. 2 Accepted pyroxene mineral names and their chemical subdivisions. Name, abbreviation and composition are given for any pyroxene that is used as an end-member of a pyroxene solid solution; such end-members are numbered between parentheses from 1 to 13. Main compositions are given for solid solutions. Space groups are also given.

mineral names	composition as end-member	main composition as solid solution	space group
I. Mg-Fe pyroxenes			
1. enstatite (En)(1)	Mg ₂ Si ₂ O ₆	(Mg,Fe) ₂ Si ₂ O ₆	Pbca
<pre>2. ferrosilite(Fs)(2)</pre>	Fe2+Si2O6	(119710) 251206	7000
3. clinoenstatite	,	(Mg,Fe) ₂ Si ₂ O ₆	$P2_1/c$
4. clinoferrosilite		(Mg, re/251206	121/ 0
5. pigeonite		$(Mg,Fe,Ca)_2Si_2O_6$	$P2_1/c$
II. Mn-Mg pyroxenes			
6. donpeacorite		(Mn,Mg)MgSi ₂ O ₆	Pbca
7. kanoite (Ka)(3)	MnMgSi ₂ O ₆	(Mn,Mg)MgSi ₂ O ₆	$P2_1/c$
III. Ca pyroxenes			
8. diopside (Di)(4)	CaMgSi ₂ O ₆	Gr (No. Br)Gi O	~
<pre>9. hedenbergite(Hd)(5)</pre>	CaFe ²⁺ Si ₂ O ₆	Ca(Mg,Fe)Si ₂ O ₆	C2/c
10. augite		(Ca,Mg,Fe) ₂ Si ₂ O ₆	C2/c
11. johannsenite(Jo)(6)	CaMnSi ₂ 0 ₆	*	C2/c
12. petedunnite(Pe)(7)*1	CaZnSi ₂ O ₆		C2/c
13. esseneite(Es)(8)*2	CaFe ³⁺ AlSiO ₆		C2/c

Tab. 2 Continued

<pre>IV. Ca-Na pyroxenes 14. omphacite 15. aegirine-augite</pre>	(Ca, Na)(R ²⁺ , Al)Si ₂ O ₆ (Ca, Na)(R ²⁺ , Fe ³⁺)Si ₂ O ₆	C2/c,P2/n C2/c
V. Na pyroxenes		
16. jadeite (Jd)(9)	NaAlSi ₂ 0 ₆ \ Na/Al Fe ³⁺ \Si ₂ 0 ₆	C2/c
17. aegirine (Ae)(10)	$ \begin{array}{c} \text{NaAlSi}_{2^{0}6} \\ \text{NaFe}^{3+\text{Si}_{2^{0}6}} \end{array} $	C27 C
18. kosmochlor(Ko)(11)	NaCr ³⁺ Si ₂ O ₆	C2/c
19. jervisite (Je)(12)*3	NaSc ³⁺ Si ₂ O ₆	C2/c
VI. Li pyroxene		
20. spodumene (Sp)(13)	LiAlSi ₂ 0 ₆	C2/c

- *1 Petedunnite has been determined by ESSENE and PEACOR (1987) to have the composition $(Ca_{0.92}Na_{0.06}Mn_{0.02})$ $(Zn_{0.37}Mn_{0.19}Fe_{0.19}^{2+}Fe_{0.12}^{3+}Mg_{0.14})$ $(Si_{1.94}Al_{0.06})O_6$ by means of an electron microprobe. This mineral was approved as a valid species by the CNMMN, IMA, in 1983.
- *2 Esseneite has been determined by Cosca and Peacor (1987) to have the composition (Ca_{1.01}Na_{0.01}) (Fe³⁺_{0.72}Mg_{0.16}Al_{0.04}Ti_{0.03}Fe²⁺_{0.02}) (Si_{1.19}Al_{0.81})O_{6.00} by means of an electron microprobe. This mineral was approved as a valid species by the CNMMN, IMA, in 1985.
- *3 Jervisite has been determined by M. Mellini et al. (1982) to have the composition (Na_{0.43}Ca_{0.31}Fe²⁺_{0.14-0.12}) (Sc_{0.66}Fe²⁺_{0.15}Mg_{0.19})Si₂O₆ by means of an electron microprobe. This mineral was approved as a valid species by the CNMMN, IMA, in 1982.

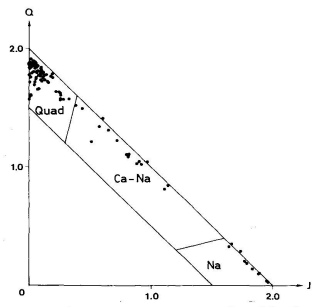


Fig. 3: The 103 DHZ pyroxenes selected by CAMERON and PAPIKE (1981) plotted on the Q-J diagram. For these pyroxenes the Q values are less than 1.90, and Mn is less than 0.08 atoms per formula unit.

pyroxenes with Mn less than 0.08 atoms for the chemical formula unit have been found to lie outside the area between Q+J=2.0 and Q+J=1.5 lines in the Q-J diagram. The classification of these unusual pyroxenes will be discussed later.

The pyroxenes that plot in the area between Q+J=2.0 and 1.5 have components other than Q and J ions at less than 25% of the M sites. Therefore, we can classify such pyroxenes on the basis of the normalized Q and J components, thereby neglecting the effects of the other components. The following procedures are adopted for further classification:

- 1) The pyroxenes in the *Quad* area are classified on the pyroxene quadrilateral Wo-En-Fs diagram with normalized Ca, Mg and Σ Fe = Fe²⁺ + Fe³⁺ + Mn atoms.
- 2) The pyroxenes in the Na area are jadeite,

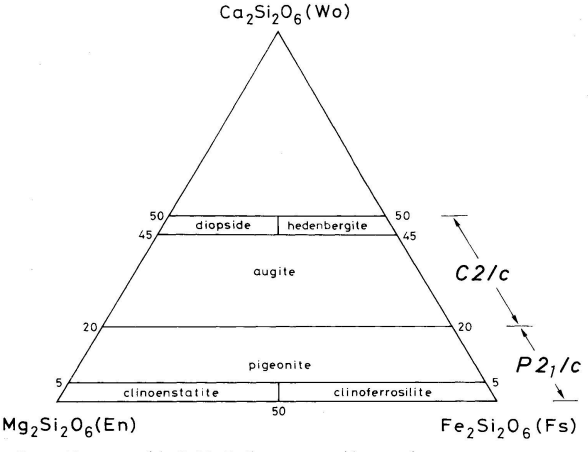


Fig. 4: Composition ranges of the Ca-Mg-Fe clinopyroxenes with accepted names.

aegirine, kosmochlor and jervisite. Because kosmochlor and jervisite show little or no solid solution towards other end-members, they play no role in the classification. Jadeite and aegirine are classified on the Quad-Jd-Ae diagram together with the Ca-Na pyroxenes, aegirine-augite and omphacite.

4.2. CLASSIFICATION OF THE Ca-Mg-Fe "QUADRILATERAL" PYROXENES

The common rock-forming pyroxenes form wide ranges of solid solution of the Ca-Mg-Fe pyroxenes and can be expressed by the pyroxene quadrilateral of the Mg₂Si₂O₆ (En) – Fe₂²⁺Si₂O₆(Fs) – CaMgSi₂O₆(Di) – CaFe²⁺Si₂O₆ (Hd) system. The Ca-Mg-Fe pyroxenes include varieties that have orthorhombic symmetry. They consist essentially of a simple chemical series (Mg,Fe)₂Si₂O₆, thus contrasting with the clinopyroxenes which have wide ranges of chemical composition in the Ca-Mg-Fe pyroxenes. Therefore, the

Ca-Mg-Fe pyroxenes are defined on the basis of symmetry and relative amounts of $\text{Ca}_2\text{Si}_2\text{O}_6(\text{Wo})$, $\text{Mg}_2\text{Si}_2\text{O}_6(\text{En})$ and $\text{Fe}_2^{2+}\text{Si}_2\text{O}_6(\text{Fs})$. The composition ranges of the clinopyroxenes and orthopyroxenes are indicated in Figures 4 and 5, respectively, where the composition is normalized to

Ca+Mg+
$$\Sigma$$
Fe=100 with
 Σ Fe=Fe²⁺+Fe³⁺+Mn²⁺¹⁰).

¹⁰⁾ For the nomenclature of the Ca-Mg-Fe pyroxenes, normalization must be made to Ca+Mg+ΣFe = 100, where ΣFe = Fe²⁺+Fe³⁺+Mn. Hereafter the mol percent of the end-member components is always used without remark and is represented simply by %. If the mole % quadrilateral components are calculated by the atomic % of Ca to the total cations at the M sites, no pyroxenes should contain more than 50% Ca₂Si₂O₆. However, if Ca, Mg and Fe are normalized, or, calculated as 100 Ca/(Ca+Mg+ΣFe), 100 Mg/(Ca+Mg+ΣFe) and 100Σ Fe/(Ca+Mg+ΣFe), respectively, then some augites will plot on a Wo-En-Fs triangular diagram above the 50% Ca₂Si₂O₆ line. Especially, when the plot in the Q-J diagram is very close to

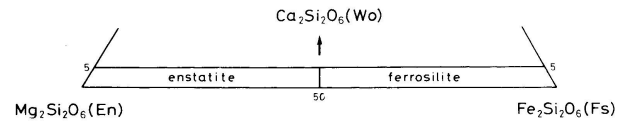


Fig. 5: Composition ranges of orthopyroxenes with accepted names.

The distinction between augite and pigeonite in the Ca-Mg-Fe pyroxenes is primarily structural, their space groups being C2/c and $P2_1/c$, respectively. There is a miscibility gap between augite and pigeonite, and many pyroxenes with 15-25% Wo have proved to be mixtures of the two. Augite with less than about 25% Wo is often called subcalcic augite. On heating, pigeonite undergoes a rapid displacive transformation to a C2/c structure which cannot be quenched. Augite does not show this type of transformation.

The most calcium-rich orthopyroxene contains approximately 5% Wo. The high-temperature form of enstatite has the space group *Pbcn* and can be expressed as 'enstatite-*Pbcn*''. This form is not quenchable and has not been found in nature. "Protoenstatite" has been used conventionally to describe this form, but this name is not adopted as a mineral name. The Wo value of "enstatite-*Pbcn*" does not exceed 2% and the En value commonly exceeds 90%. Thus the composition field of "enstatite-*Pbcn*" is different from that of enstatite-*Pbca*.

4.3. THE CLASSIFICATION OF THE Na AND Ca-Na PYROXENES

The Na pyroxenes, jadeite and aegirine, commonly contain more than 90% of the NaAlSi₂O₆ or NaFe³⁺Si₂O₆ component, respectively, but contain neither the Ko nor the Je component. Because kosmochlor is a rare accessory constituent of some iron meteorites and only one terrestrial locality is known for

each of kosmochlor and jervisite, these two species are separately treated in the classification of the Na pyroxenes. Both jadeite and aegirine, however, show extensive solid solution with the Ca-Mg-Fe pyroxenes, especially with the diopside-hedenbergite series and augite, leading to the Ca-Na pyroxenes. The Na and Ca-Na pyroxenes are classified on the Quad-Jd-Ae diagram (Fig. 6) with normalized Q (Wo+En+Fs), Jd and Ae components¹¹). The arbitrary divisions between the Ca-Mg-Fe pyroxenes, Na-Ca pyroxenes and pyroxenes are defined at 20 and 80% of Q = (Wo + En + Fs). Omphacite displays a $C2/c \Rightarrow P2/n$ polymorphic transition, and both high-temperature C2/c and low-temperature P2/n polymorphs appear in nature. Omphacite can thus be divided into two subspecies: omphacite-C2/c and omphacite-P2/n. Because omphacite-P2/n shows a unique crystal structure different from that of jadeite and augite, it is accepted as an independent pyroxene species. Aegirine-augite is also accepted as an independent species to keep balance with omphacite, though it is not known to occur with the P2/n structure. The classification of the Ca-Na pyroxenes by Essene and Fyre (1967) is not followed in this report.

4.4. THE CLASSIFICATION OF OTHER PYROXENES

Most naturally occurring pyroxenes in the 'Others' area are johannsenite (CaMnSi₂O₆),

or outside of the boundary Q+J=1.5, the effect of johannsenite and petedunnite components must be considered. If the effect is negligible, the pyroxene must be considered to have an unusual composition and must be referred to the section of unusual pyroxenes.

¹¹⁾ To normalize Q, Jd and Ae components, Ca+Mg+Fe²++2Na at the M sites must be made to total 100%. Then the normalized 2Na% must be divided to the ratio of Al/Fe³+ to give the ratio of Jd/Ae. Thus Q+Jd+Ae must always give 100%.
When the plot in the Q-J diagram is significantly outside the boundary Q+J=2.0, the effect of substitution (2) must be considered, as in the section of unusual pyroxenes.

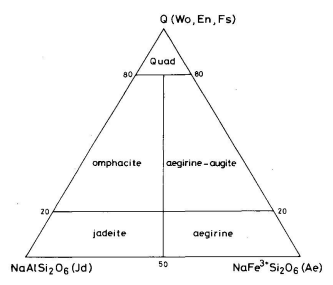


Fig. 6: Ca-Mg-Fe and Na pyroxenes with accepted names. Quad represents the Ca-Mg-Fe pyroxene area (see Fig. 4).

petedunnite (CaZnSi₂O₆) and spodumene (LiAlSi₂O₆) (Fig. 2). Recent investigations of natural manganese-bearing pyroxenes have yielded two new minerals, kanoite and its dimorph donpeacorite, (Mn,Mg)MgSi₂O₆, which seem to form a solid solution with En (Petersen et al., 1984). They too occur in the 'Others' area. These results suggest a possible Mn-Mg-Fe pyroxene quadrilateral. Esseneite (CaFe³⁺AlSiO₆) is the first pyroxene with the substitution (3) as described in Table 1.

4.5. THE CLASSIFICATION OF UNUSUAL PYROXENES

Several pyroxenes with unusual chemical compositions (Table 3) appear outside the area between the Q+J=2.0 and Q+J=1.5 lines in the Q-J diagram, though they do not belong to the 'other' pyroxenes mentioned above (Fig. 7). They contain large amounts of chemical constituents of substitutions (2), (3) and (4) mentioned in Table 1 in the standard chemical formulae.

These pyroxenes can be divided into two groups: firstly Ca-rich pyroxenes with the $S3(CaR^{3+}AlSiO_6)$ and $S4(CaR^{2+}_{0.5}Ti^{4+}_{0.5}AlSiO_6)$ components representing substitutions (3) and respectively, and Na-rich (4),secondly pyroxenes with the S2 component $(NaR_{0.5}^{2+}Ti_{0.5}^{4+}Si_2O_6)$ representing substitution (2). The former shows a significant deficiency

of Si atoms such as Si<1.60 in the standard formula resulting in the Q-value close to or less than 1.5 (point S4 in Fig. 7). The latter appears outside the line Q+J=2.0 approaching point S2 in Fig. 7. All these unusual pyroxenes are classified by using the accepted pyroxene names and the adjectival modifiers mentioned below, except the Allende pyroxene (Table 3, No. 4) which is called subsilicic titanoan aluminian pyroxene.

The Allende pyroxene (No. 4) contains 39% of the S3(Ti) component and can be considered as a new mineral. However, we have decided only to use the accepted names in this report and if a species has not yet been approved, we use *pyroxene* as for No. 4 in Table 3. The names used in literature for the unusual pyroxenes are listed in Table 3 in comparison with those in this report. The "CaMgTAL" pyroxene (No. 1) is diopside in this classification.

5. Adjectival modifiers

Adjectival modifiers for mineral names are used to indicate unusual amounts of chemical

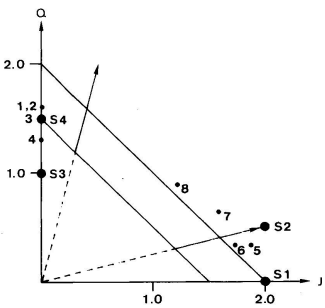


Fig. 7: Q-J diagram for eight unusual pyroxenes with Q-value less than 1.62 and Mn less than 0.08 atoms per formula unit (Table 3). The components formed by the substitutions (1) to (4), as indicated in Table 1, are plotted in the diagram. They represent the following compositions:

 $S1 = NaR^{3+}Si_2O_6$, $S2 = NaR^{2+}_{0.5}Ti_{0.5}^{4+}Si_2O_6$,

 $S3 = CaR^{3+}AlSiO_6$ and $S4 = CaR^{2+}_{0.5}Ti^{4+}_{0.5}AlSiO_6$.

Tab. 3 Chemical composition and classification of eight unusual pyroxenes. Numbers such as 320-8, etc. represent pages and analysis number in DHZ (1978). Other references are in text. With the exception of 320-8 (= 406-16), all the DHZ analyses in this table were not included in the 103 selected analyses of CAMERON and PAPIKE (1981). All pyroxenes in the table are shown with their numbers in the Q-J diagram (Fig. 7). S2, S3 and S4 represent the following components of substitutions (2), (3) and (4), respectively: S2=NaR $_{0.5}^{2+}$ Ti $_{0.5}^{4+}$ Si $_{0.5}$ Si $_{0.5}$ Oi, S3=CaR $_{0.5}^{3+}$ AlSiO $_{6}$, and S4=CaR $_{0.5}^{2+}$ Ti $_{0.5}^{4+}$ AlSiO $_{6}$. To indicate R ions explicitly in these components, the notation S(R), such as S2(Mg) and S3(Al), is used. S3(Fe) is a new pyroxene, esseneite (Es).

a) Ca-rich group related to S3 and S4								
No.	1. 320- (406-	-8	2. 403-		3. D an	d S*1	4. T an	nd R ^{*2}
Si	1.443	2 00	1.506	2.00	1.434	2.00	1.196	2.00
Al	.557	2.00	.494	2.00	.566	2.00	.804	2.00
Al	.091	10 Mar 2010	.171		.306		.186	
Ti4+	.165		.065		.022	Ti	4+ .111	
Fe ³⁺	.128		.159		.218	Ti	3+ .394	
Mg	.385		.570		.408		.289	
Fe ²⁺	.229	2.00	.063	2.02	.060	2.00		2.00
Mn	.005		.007		.005			
Ca	.992		.975	te.	.979		1.021	
Na	.006		.007		.002			
K	.000		.001		-			
Q	1.61	3	1.61		1.45		1.31	
J	0.01		0.01		0.00		0.00	
Mineral names	subsilic titanian ferrian diopside		subsilic aluminia ferrian diopside	ın	subsilic aluminia ferrian diopside	n	subsilic titanoar aluminia pyroxena	n an
Names in litera- ture	titanaug (320-8) titanium fassait (406-16 CaMgTAL (C and	e)	fassaite		fassaite		titanaug	gite

Component ratios;

- 1. $(Wo_{22}En_{12}Fs_{10})_{44}S4(M\tilde{g})_{18}S4(Fe)_{16}Es_{13}S3(Al)_{9}$
- 2. $(Wo_{26}En_{25}Fs_2)_{53}S3(Al)_{17}Es_{16}S4(Mg)_{12}S4(Fe)_2$
- 3. $(Wo_{22}En_{20}Fs_2)_{44}S3(Al)_{31}Es_{21}S4(Mg)_4$
- 4. $S3(Ti)_{39}S4(Mg)_{22}(Wo_{11}Fs_9)_{20}S3(Al)_{19}$

b) Na-rich group related to S2

No.	5. 488-	9	6. 491-	14	7. 492- (C and	-19 1 G ^{*4})	8. C an	nd G*4
Si	1.994	2 00	2.024	2 02	2.026	2 02	2.009	2.01
Al	.032	2.00	.000	2.02	.000	2.03	.000	2.01
Al	.000		.021		.098		.348	
Ti4+	.265		.023		.227		.104	
Fe ³⁺	.458		.728		.192		.031	
Mg	.150		.070		.070		.168	
Fe ²⁺	.107	2.00	.113	2.00	.420	1.98	.356	2.00
Mn	.003		.006		.021	4	.011	
Na	.933		.872		.794		.610	
Ca	.083		.155		.152		.361	
К	20 <u></u>)		.009		-		.006	
Q	0.34		0.34	n	0.64		0.89	
J	1.87	A 3 000	1.74		1.59		1.22	
Mineral names	titanian magnesia ferroan aegirine	n	calcian ferroan aegirine		titanian aegirine augite		titaniar ferroan omphacit	
Names in litera- ture	titanian aegirine		aegirine augite	-	titanian aegirine augite (492-19 titanian aegirine (C and)) 	titanian ferro-om	

Component ratios;

- 5. $Ae_{46}S2(Mg)_{28}S2(Fe)_{20}^{\Delta_6}^{*5}$
- 6. $(Ae_{73}Jd_2Wo_8Fs_6En_3)_{92}S2_4\Delta_4$
- 7. $(Ae_{19}Jd_{10}Fs_{12}Wo_8En_2)_{51}S2(Fe)_{42}S2(Mg)_4\Delta_3$
- 8. $(Jd_{35}Ae_3Wo_{18}Fs_{15}En_7)_{78}S2(Fe)_{14}S2(Mg)_6\Delta_2$

^{*1} DEVINE and SIGURDSSON (1980), Table 1 for fassaite.

^{*2} Tracy and Robinson (1977), Table 3, Analysis I for pyroxenes from the Allende meteorite (Mason, 1974).

^{*3} CAMERON and PAPIKE (1981), Table A3, Analysis 320-8 and 406-16.

^{*4} Curtis and Gittins (1979), Table 2, Analysis 5 for No. 7 and Table 5, Analysis 5 for No. 8.

^{*5} \triangle represents minor components, some of which have unusual metal ratios for the pyroxene structure.

Tab. 4 Extreme chemical compositions of pyroxenes in DHZ (1978). Number of cations per formula unit, minimum values for Si and maximum values for other cations. Bold numbers are for the main constituent elements. Numbers in the parentheses such as 42-9, etc. indicate pages and analysis numbers in DHZ (1978). Other references are in text.

	Mg-Fe pyroxenes	Ca pyroxenes	Na pyroxenes
Si	1.76 (42-9)	1.44 (320-8)*1	1.94 (488-9)
A13+	0.24 (42-9)	0.56 (320-8)	0.07 (488-8)
Fe ³⁺	0.04 (49-8)	0.09 (320-11)	0.02 (488-9)
A1 ³⁺	0.15 (49-6)	0.35 (320-11)	0.98 (464-1)
Ti 4+	0.04 (40-30)	0.17 (320-8) ^{*2}	0.27 (488-9)*3
Fe ³⁺	0.12 (170-8)	0.37 (321-5)*4	0.97 (487-1)
Mg ²⁺	1.99 (41-1)	1.27 (208-4)	0.15 (488-9)
Fe ²⁺	1.72 (47-33)*5	1.09 (220-13)	0.11 (488-9)
_{Mn} 2+	0.27 (45-21)*6	0.36 (217-5) ^{*7}	0.03 (487-4)
Cr ²⁺	0.02 (36-9)	0.06 (207-11)	_ *8
Ni ²⁺	-	0.003 (317-1)	-
zn ²⁺	_	0.21 (216-11) ^{*9}	-
Ca ²⁺	0.26 (169-2)	1.03 (202-4)	0.16 (466-14)
Na ⁺	0.10 (169-2)	0.31 (323-7)	0.98 (464-1)

^{*1} Table 3, No. 1. Table 3, No. 4: Pyroxene from the Allende meteorite 1.20 (MASON, 1974; TRACY and ROBINSON, 1977).

constituents. In order to define the unusual amounts for the pyroxene mineral group quantitatively, extreme compositions of pyroxenes have been listed in Table 4, where the values for the main cations are shown as well as those for the accessory cations. DHZ (1978) and ROBINSON'S (1980) table were mainly used in constructing the table.

An element specified as a modifier should be present as a general rule in a quantity larger than 0.1 or 0.01 atoms in the standard chemical formula of 6 O or 4 metal atoms (Table 5) depending on the maximum content in Table 4.

The suffixes are those proposed by SCHALLER (1930) and adapted by CNMMN (NICKEL and MANDARINO, 1987). The suffix -ian is used for the higher valence state (e.g., ferrian) or for an element with a non-variable state (e.g., lithian). The suffix -oan implies the lower valence state (e.g., ferroan). It is recom-

^{*2} Probe analysis 0.252 and 0.282, half of $CaR_{0.5}^{2+}Ti_{0.5}^{4+}AlSiO_6$ (S4) (Tracy and Robinson, 1977; Robinson, 1980).

^{*3} Table 3, No. 5. Half of $NaR_{0.5}^{2+}Ti_{0.5}^{4+}Si_2O_6$ (S2).

^{*4 406-15 0.67,} omitted because of possible errors in chemical analysis.

^{*5} Probe analysis 1.880 (JAFFE et al., 1978).

^{*6} Probe analysis 0.301 (ROBINSON, 1980), Kanoite 1.04 (KOBAYASHI, 1977).

^{*7} Johannsenite 0.963 (417-2).

^{*8} Kosmochlor 0.90 (522-1).

^{*9} Petedunnite 0.37 (Table 2, remark *1).

mended that such modifiers never be used for main cations normally contained in the named mineral, for example, in terms like calcian augite, aluminian omphacite, and sodian aegirine-augite, in which the modifiers are obviously superfluous.

If there is less than the amount necessary for the assignment of the modifiers such as 'aluminian' in Table 5, or Al < 0.1, but if the increased content of the element must be stressed, a modifier 'aluminum-bearing' may be used. This second type of modifier should be used also (1) if only an incomplete analysis is available preventing the calculation of a full chemical formula or (2) for pyroxenes where the valence state of a cation is unknown. With regard to the Si content in pyroxenes, it is suggested that Si < 1.75 is a suitable limit for use of the term 'subsilicic', though one should bear in mind that the Si < 5.75 limit for 'subsilicic' in amphiboles corresponds to Si < 1.5 for pyroxenes.

Tab. 5 List of adjectival modifiers to be used for pyroxene mineral names. The limit of the content is determined based on the values listed in Table 4.

cation	content*1	name
A13+	>0.10	aluminian
Ca ²⁺	>0.10	calcian
Cr ³⁺	>0.01	chromian
Fe ²⁺	>0.10	ferroan
}e³+	>0.10	ferrian
Li+	>0.01	lithian
Mg ²⁺	>0.10	magnesian
_{Mn} 2+	>0.10	manganoan
_{Mn} 3+	>0.01	manganian
Na+	>0.10	sodian
Ni ²⁺	>0.01	nickeloan
Si ⁴⁺	<1.75	subsilicic
Ti 3+	>0.01	titanoan
Ti 4+	>0.10	titanian
$2n^{2+}$	>0.01	zincian

^{*1} Number of cations per formula unit M2M1T₂O₆. If the mineral name itself implies the presence of certain cations, adjectival modifiers for these cations should not be used ('subsilicic' is an exception).

In certain cases, particularly for the augite series, it is convenient to use the following adjectival modifiers: iron-rich, magnesium-rich, and subcalcic. A prefix actually attached or hyphenated to a mineral name, however, is incorrect and should be avoided (NICKEL and MANDARINO, 1987), because it would cause the mineral to be indexed alphabetically under the prefix rather than the proper mineral name. This is why such terms as "ferropigeonite", "ferro-augite", etc., should not be used as mineral names.

It is often useful to give the space group of the mineral, particularly when it can occur in two or more forms. For example, we could distinguish between the two forms of omphacite by adding the space group symbol, i.e., omphacite-C2/c, omphacite-P2/n, or by adding the lattice-type symbol, i.e., omphacite-C, omphacite-C (BAILEY, 1977).

6. Obsolete pyroxene names

The names of 105 pyroxenes or altered pyroxenes listed in Appendix have formally been discarded by the CNMMN and are therefore obsolete. The preferred name is printed in *italics* in the same table.

Acknowledgements

We are thankful to Professor J.H.D. Donnay, McGill University, Montreal, who contributed greatly to the improvement of the report by his careful review. We also appreciate criticisms and comments by Dr. A. Kato, National Science Museum, Tokyo, Dr. M. Kitamura, Kyoto University, Kyoto, and the members of the Commission on New Minerals and Mineral Names, IMA.

Tab. 6 Obsolete pyroxene names (p. 110). The following pyroxene mineral names, or names which refer to altered pyroxenes, have been formally discarded by the CNMMN. The correct names are printed in *italics*. The original form of this table was compiled by Malcolm Ross using the following references: Dana (1892); Tschermak (1897); Chester (1886); Ford (1932); Winchell and Winchell (1951); Deer, Howie and Zussman (1963, 1978); Strunz (1970); and the unpublished Thesaurus of Mineralogical Terms of the International Mineralogical Association, which has been available since August 1974.

	1
acmite = aegirine	lavrovite = diopside
aegirite (aegyrite) = aegirine	lawrowite = diopside
aegerine-hedenbergite = augite	leucaugite = diopside
agalite = probably enstatite partly altered to talc	lime-bronzite = probably pigeonite or enstatite plus
aglaite = altered spodumene	augite, ("inverted" pigeonite)
alalite = diopside	loganite = diopside + actinolite + talc
alkali augite = aegirine-augite	lotalite = hedenbergite
amblystegite = enstatite	malacolite = diopside with good (001) parting, also
anthochroite = augite	diopside from Sala, Sweden
asteroite = iron-rich augite	mansjoite = augite or diopside or hedenbergite
baikalite = diopside	mayaite = omphacite
bastite = enstatite which has altered to serpentine,	mellcrite = orthopyroxene
talc, or perhaps anthophyllite	mondradite = probably an altered pyroxene
blanfordite = manganoan aegirine-augite	mussite = diopside
bronzite = enstatite	orthobronzite = enstatite
calc-clinobronzite = pigeonite	orthoenstatite = enstatite
calc-clinoenstatite = pigeonite	orthoeulite = ferrosilite
calc-clinohypersthene = pigeonite	orthoferrosilite = ferrosilite
calc-pigeonite = subcalcic <i>augite</i>	orthohypersthene = enstatite or ferrosilite
canaanite = diopside	paulite = enstatite
chladnite = enstatite	peckhamite = enstatite
chloromelanite = omphacite or aegirine-augite	phastine = altered enstatite
chrome-acmite = chromian aegirine	picrophyll = altered pyroxene?
chromejadeite = chromian jadeite	pigeonite-augite = probably subcalcic augite
clinohypersthene = clinoenstatite or clinoferrosilite	pitkarantite = pyroxene?
coccolite (kokkolith) = iron-rich augite	potash-aegirine = synthetic product, probably not
cymatolite = altered spodumene	properly characterized
diaclasite = altered enstatite	protheite = augite
diallage = diopside which has altered or which has	protobastite = enstatite
good (100) parting; also used for alteration	pyrallolite = altered pyroxene?, talc?
products of other pyroxenes	pyrgom = pyroxene
diopsidjadeite = omphacite	sahlite = diopside
endiopside = magnesium-rich augite	salite = diopside
enstatite-diopside = magnesium-rich augite	schefferite = manganoan diopside
eulite = ferrosilite	schillerspar (schillerspat) = enstatite which is altered
eulysite = ferrosilite	to serpentine, talc, or anthophyllite
fassaite = ferrian aluminian diopside or augite	shepardite = enstatite
fedorovite = diopside	soda-spodumene = sodian spodumene
ferroaugite = augite	strakonitzite = altered pyroxene, steatite?
ferrohedenbergite = augite	szaboite = partly altered enstatite
ferrohypersthene = ferrosilite	titanaugite = titanian augite
ferro-johannsenite = iron-rich johannsenite	titandiopside = titanian diopside
ferropigeonite = iron-rich pigeonite	titanpigeonite = titanian pigeonite
ferrosalite = hedenbergite	trachyaugite = augite
ficinite = enstatite	traversellite = diopside
funkite = hedenbergite	triphane = spodumene
germarite = altered enstatite	tuxtlite = omphacite
hiddenite = spodumene	uralite = pseudomorph of amphibole after
hudsonite= hedenbergite	pyroxenes
hyperstehne = enstatite or ferrosilite	urbanite = iron-rich augite or aegirine-augite
jadeite-aegirine (jadeite-aegirite) = jadeite or	ureyite = kosmochlor
aegirine	vanadinaugite = vanadium-bearing augite
jeffersonite = zincian manganoan diopside or augite	vanadinbronzite = vanadium-bearing enstatite
killinite = altered spodumene	vargasite = altered pyroxene?
korea-augite = augite	victorite = enstatite
kunzite = spodumene	violaite = augite
lavroffite = diopside	violan = magnesium-rich augite or diopside

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