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# An unusual assemblage of talc-phengite-chlorite-K-feldspar in quartz schists from the Nahavand area, Sanandaj-Sirjan zone, Iran

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# Abstract

For the first time, an unusual assemblage of talc-phengite-chlorite-K-feldspar was found in quartz schists from the Sanandaj-Sirjan zone in the Nahavand area in western Iran. The talc-bearing quartz schists occur as small bodies or lenses within pelitic schist layers and contain talc, phengite, chlorite, K-feldspar and quartz as major mineral constituents with subordinate amounts of calcite and graphite. Textural analysis revealed that talc, phengite, chlorite and K-feldspar are in sharp contact and no reaction rims between them were observed. Constructed petrogenetic gird in the K<sub>2</sub>O-FeO-MgO-Al<sub>2</sub>O<sub>3</sub>-SiO<sub>2</sub>-H<sub>2</sub>O (KFMASH) model system containing talc, phengite, chlorite, K-feldspar, phlogopite and kyanite with excess quartz and H<sub>2</sub>O shows that divariant assemblage of talc-phengite-chlorite-K-feldspar is stable over a wide P-T range defined by the following two univariant reactions: phengite + talc + quartz = chlorite + K-feldspar + kyanite + H<sub>2</sub>O and chlorite + phlogopite + quartz = talc + phengite + K-feldspar + H<sub>2</sub>O. Constructed Al<sub>2</sub>O<sub>3</sub>-KAlO<sub>2</sub>-MgO+FeO (AKM) compatibility diagrams predict that phengite (X<sub>Ph</sub> = 0.280, Y<sub>Ph</sub> = 0.860), chlorite (X<sub>Chl</sub> = 0.570, Y<sub>Chl</sub> = 0.640), talc (X<sub>Tlc</sub> = 0.160, Y<sub>Tlc</sub> = 0.02) and K-feldspar are stable at P = 11 kbar and T = 400°C. This relatively high-pressure assemblage could be formed during the subduction of the Neo-Tethys oceanic plate under Iranian microcontinent.

Keywords: High-pressure metamorphism; Nahavand; Sanandaj-Sirjan zone; Talc-phengite-K-felds parassemblage

## 1. Introduction

In natural rocks, talc-phengite assemblage was first found in the piemontite-quartz schist in Serbia (Abraham and Schreyer, 1976). Abraham and Schreyer (1976) suggested that talc-phengite assemblage is an indicator of high-pressure metamorphism based on theoretical grounds in the K<sub>2</sub>O-MgO-Al<sub>2</sub>O<sub>3</sub>-SiO<sub>2</sub>-H<sub>2</sub>O (KMASH) model system, although they also proposed an alternative hypothesis that this pair could be formed due to the presence of small amounts of additional non-KMASH components such as Fe<sub>2</sub>O<sub>3</sub>-MnO-CuO-NiO phyllosilicates under low-pressure in conditions. Massonne and Schreyer (1989) assessed the stability field of talc-phengite assemblage by the synthetic work in the KMASH model system containing kyanite, chlorite, talc, phlogopite and muscovite with excess quartz and H<sub>2</sub>O. They proposed that the following two reactions: chlorite+ phlogopite + quartz = talc + muscovite +  $H_2O$  and talc + muscovite = phlogopite + quartz +  $H_2O$ intersect at an invariant point at 10.8 kbar and 610°C, which supported hypothesis 2 of Abraham

\*Corresponding author Received: 4 February 2013 / Accepted: 7 May 2014 and Schreyer (1976) regarding the high-pressure for talc-phengite assemblages. origin This assemblage was subsequently recognized in the western Alps (Chopin, 1981) and thereafter in a number of areas around the world in high-pressure metamorphic terrains such as the Sanbagawa metamorphic belt, Japan (Izadyar et al., 2000; Hatanaka and Hirajima, 1999; Ubukawa et al., 2007). Talc-kyanite and talc-omphacite assemblages have also been identified from ultra high-pressure metamorphic rocks (Reinecke, 1991; Mattinson et al., 2004; Liou and Zhang, 1995). However, the natural parageneses of talc-phengite and talc-kyanite usually coexist with Fe-bearing mineral phases such as garnet, biotite, chloritoid and chlorite (Chopin, 1981; Frank et al., 1987) and do not belong to the constituents of the simple KMASH model system. Wei and Powell (2003) calculated the KFMASH model system for highpressure metapelites with mineral phases of garnet, chloritoid, chlorite, biotite, carpholite, talc, staurolite, kyanite with phengite, quartz/coesite and H<sub>2</sub>O in excess. They documented the typical parageneses of talc-phengite and talc-kyanite for a variety of bulk compositions from several highpressure terrains. Wei and Powell (2004) also calculated the stable mineral assemblages of Nabearing metapelites in high-pressure conditions as

occur widely in high-pressure thev also metamorphic terrains (Chopin, 1981; Izadyar et al., 2000; Meyre et al., 1999; Theye et al., 1992). By using Schreinemakers' analyses and thermodynamic calculation based on the dataset of Holland and Powell (1998), Izadyar et al. (2000) indicated that the talc-phengite-albite assemblage from piemontite-quartz schists of the Sanbagwa metamorphic belt is stable at 580-600°C and 11.6-12 kbar. However, Ubukawa et al. (2007) obtained temperatures of 565-580°C and pressures of 9.5-10 kbar for talc-phengite assemblage in NCKFe<sup>3+</sup> MASH model system in talc-Na pyroxene piemontite-quartz schist in the Sanbagawa metamorphic belt. None of these studies, either experimental or calculated, have addressed a talcphengite-K-feldspar assemblage. In fact, only Massonne and Schreyer (1989) proposed highpressure stability field for the talc-phengite-Kfeldspar assemblage purely on theoretical grounds but it is not reported in natural rocks from highpressure metamorphic terrains. During а petrological investigation in the Nahavand area of the Sanandaj-Sirjan zone, an unusual assemblage of talc-phengite-chlorite-K-feldspar was recognized in quartz schists. This paper is the first to describe this assemblage, to characterize its mineralogy and texture and to explain its equilibrium condition based on the petrogenetic grid in the KFMASH model system containing K-feldspar.

# 2. Geological setting

The Sanandaj-Sirjan zone, which is 1500km long and up to 200km wide, stretches from the northwest to the southwest of Iran and is located between the central Iran block and Zagros Fold-thrust belt (Stoclin, 1968; Berberian and King, 1981; Alavi, 1994) (Fig. 1A). The Sanandaj-Sirjan zone consists of mafic-ultramafic rocks, metamorphic complexes and Paleozoic-Mesozoic sequences that underwent amphibolite/greenshist-facies metamorphism during the Cretaceous-Tertiary continental collision between the Afro-Arabian continent and the Iranian microcontinent (Sengör and Natalin, 1996; Mohajjel and Fergusson, 2000; Mohajjel et al., 2003). From the southwest to the northeast it consists of several elongated sub-zones (Fig. 1B): (1) a radiolarite subzone consisting of Triassic-Cretaceous shallow marine limestone and dominant deep-marine radiolarite; (2) a Bisotun sub-zone that includes Late Triassic to Late Cretaceous limestone; (3) an ophiolite sub-zone that is the location of the Zagros suture zone; (4) a marginal sub-zone that contains abundant Late Jurassic-Early Cretaceous volcanic rocks in addition to shallow marine strata in the Cretaceous unit and (5) a complexly deformed sub-zone (Mohajjel et al., 2003). The last sub-zone is distinguished from the other subzones by abundant metamorphic rocks. The main rock 244

types of the complexly deformed sub-zone are schist, marble, amphibolite, quartzite, dolomite-marble and metasandstone. All sub-zones are imbricated in a complicated thrust system with out-of-sequnce thrusts that locally juxtaposes the younger units over the older ones (Mohajjel et al., 2003). The Nahavand area belongs to the complexly deformed sub-zone and is located near the main Zagros reverse fault, which is proposed as a suture zone between the Arabian plate and Eurasia (Sengör, 1984; Agard et al., 2006; Ghasemi and Talbot, 2006) (Fig. 1B). Apart from a thin ophiolite unit, the lithologic units in the Nahavand region can be generally subdivided into two groups of metamorphosed and non-metamorphosed rocks (Alavi and Mahdavi, 1994) (Fig. 2). The metamorphic rocks include five different successions that are in fault contact with both the overlying and underlying rocks. The oldest one (PTrm) is a sequence of massive marbles of Permian-Triassic age that are strongly sheared and mylonitized near the contact with the underlying rocks. The next succession (TrJmv) consists of an assemblage of Middle to Upper Triassic-Lowermost Jurassic metamorphosed basaltic and andesite-basaltic lava flows (with tholeitic affinities). This assemblage contains locally occurring pillow structures interlayered with well-foliated marbles that have thin, upward increasing slate layers. Small plutons of gabbro and diorite (now metamorphosed) have locally intruded this assemblage (Alavi and Mahdavi, 1994) (Fig. 2). The third unit (TrJm) consists entirely of a sequence of fossiliferous marbles of Late Triassic to Early Jurassic. The fourth succession (Jsv) is a sequence of Lower Jurassic slates, phyllites and metagreywakes that have intercalations of marbles and metamorphosed volcanic rocks in the lower parts. The fifth lithological unit (Jh) consists of an Upper Liassic succession of cordieritebearing hornfels which is the product of contact metamorphism due to the intrusion of Late Cretaceous granitoid plutons (Alavi and Mahdavi, 1994) (Fig. 2).



Fig. 1. (A) Simplified structural map of Iran showing the location of the Sanandaj-Sirjan zone. (B) Tectonic map of southwestern Iran showing the divisions of the Sanandaj-Sirjan zone (modified after Mohajjel et al., 2003)



Fig. 2. Geological map of the Nahavand area (modified after Alavi and Mahdavi, 1994)

# 3. Petrography

Quartz schists were collected from the Jsv unit that is a Lower Jurassic sequence mostly of pelite and greywake origins (Fig. 3). This sequence usually occurs in the northwest of the Nahavand region. The quartz schists form 30-50 m thick layers interbedded with 10-30 m thick marbles and 100-150 m thick pelitic schists. They also occur as small bodies or lenses from one centimeter up to a maximum of three meters within pelitic schist layers. An unusual but stable mineral assemblage of talc-phengite-chlorite-K-feldspar could only be found in the quartz schist occurring as small bodies or lenses within pelitic schists. Therefore, based on the field and mineral assemblages, two types of quartz schists are recognized in the studied area which are named talc-free and talc-bearing quartz schists (Fig. 3). The mineral parageneses of these two quartz schists are tabulated in Table 1. The talc-bearing quartz schist contains coarser grains than talc-free quartz schist and commonly shows compositional banding of quartz-rich and mica-rich bands. The width of the quartz-rich bands ranges from 1 to 2 mm, and they are mainly composed of quartz, K-feldspar and subordinate amounts of phengite and chlorite. The mica-rich bands mostly contain phengite, talc and chlorite with minor amounts of quartz and K-feldspar. The width of these bands ranges from 1 to 2 mm. The textural relationship clearly shows that talc and phengite are in sharp contact, and no reaction rims between them

are observed (Figs. 4A, 4B, 4D and 4E). Chlorite is one of the major constituent minerals in both talcbearing and talc-free quartz schists. It occurs in quartz-rich and mica-rich bands in sharp contact with talc, phengite and K-feldspar in talc-bearing assemblage (Figs. 4C and 4F) and with muscovite in talc-free assemblage. It is hard to distinguish talc from phengite under the microscope, because talc has similar birefringence to that of phengite. However, talc can be distinguished from phengite by its lower interference colour under microscope, because tale's lower Mohs hardness (H = 1) should enhance the thinning during thin-section making rather than surrounding harder minerals, such as quartz (H = 7) and phengite (H = 2.5-3)(Figs 4A and 4B). Talc occurs only in talc-bearing quartz schists in sharp contact with phengite, chlorite and K-feldspar (Figs. 4A, 4B, 4C, 4D, 4E and 4F). Kfeldspar occurs in both talc-bearing and talc-free quartz schists and also in both mica-rich and quartzrich bands (Fig. 4A and 4C). Usually it shows an elongated form but sometimes in the talc-free samples a sigmoidal form is seen with asymmetrical pressure shadows. Calcite and graphite are not common, but are present in the matrix of both talc-bearing and talc-free quartz schists.



Fig. 3. Lithological map of the studied area and sample localities

 
 Table 1. Mineral assemblages of talc-bearing and talcfree quartz schists from the Nahavand area

Mineral Na.	Chl	Ms	Ph	Tlc	Kfs	Qz	Cal	Gr
C18	+	+	+	+	+	+	+	-
C19	+	+	-	-	-	+	+	+
C20	+	-	+	+	+	+	-	+
C21	+	+	+	+	+	+	-	-
C22	+	+	+	+	+	+	+	-
C44	+	-	+	-	+	+	+	+
C32	+	+	-	-	-	+	+	+
C53	+	+	-	-	+	+	-	+
C63	+	+	+	+	+	+	+	-

The mineral abbreviations are from Whitney and Evans (2010). Symboles (+) and (-) indicate present and absent minerals respectively



**Fig. 4.** Photomicrographs of talc-bearing quartz schists showing (A) coexistence of talc, phengite and K-feldspar; (B) intergrowth of talc and phengite; (C) talc, chlorite, phengite and K-feldspar coexistence; (D)-(E) BSE images of coexisting talc and phengite; (F) BSE image of intergrowth of chlorite and talc. Crossed polars. Abbreviations are from Whitney and Evans (2010)

#### 4. Mineral chemistry

Mineral chemistry was acquired by a Cameca SX50 electron microprobe equipped with three wavelength dispersive spectrometers at Toronto University, Canada and a Cameca SX100 at Iran Mineral Processing Research Center (IMPRC). The accelerating voltage and the beam current were maintained at 15 kv and 10 nA, respectively. Corrections were made using either the ZAF or Phi-Rho-Z method following Armstrong (1988). We did not increase the spot size to attempt to minimize the alkali loss, because this would have obscured evidence of any zoning in mica. However, the risk

of using mica analyses with a significant contamination or alkali loss was reduced using the method proposed by Vidal and Parra (2000). Unfortunately, in the absence of direct measurement, it is impossible to estimate precisely the Fe<sup>3+</sup>-content from EPMA. However, it is generally accepted that  $Fe^{3+}$  in chlorite group minerals is controlled by crystallochemical constraints rather than fo<sub>2</sub> conditions and is never abundant (Deer et al., 1992; Zane and Sassi, 1998). Therefore, chlorite octahedral contents were calculated from EPMA analyses with the assumption that  $Fe^{2+} = Fe^{total}$  could be a real feature. In contrast, Guidotti et al. (1994c) showed that proportion of Fe<sup>3+</sup> substituting for Al in mica is controlled by  $fo_2$  in which  $Fe^{3^+}$  is minimal in rocks containing carbonate and graphite. In order to limit the possibility for white micas to contain Fe<sup>3+</sup>, we preferentially used graphite or at least carbonatebearing samples. Under such low  $fo_2$  condition the assumption of  $Fe^{2+} = Fe^{total}$  was also considered for

#### 4.1. Chlorite

talc.

Following the method proposed by Vidal and Parra (2000), only chlorite compositions that could be expressed as a linear combination of amesite. clinochlore, daphnite and sudoite end-members were retained. The distribution of cations on the different sites (Table 2) follows Holland et al. (1998), whereby octahedral Al in clinochlore, daphnite and amesite is restricted to the octahedral M1 and M4 sites with a strong preference for M4, and tetrahedral Al is restricted to T2 on which it substitutes randomly for Si. In addition, we assume that vacancies in sudoite are restricted to the M1 site and that Fe, Mg and Al mix randomly over the (M1+M2) (equal Mg/Fe proportions in these sites). Comparing cation site distributions between chlorite from two assemblages indicates that the chlorite from talc-free assemblage contains higher amounts of  $Si_{(T1+T2)}$ ,  $Fe_{(M1)}$ ,  $Al_{(M1)}$ ,  $\Box_{(M1)}$  and lesser amounts of  $Al_{(T2)}$ ,  $Mg_{(M2+M3)}$ ,  $Fe_{(M2+M3)}$ ,  $Al_{(M2+M3)}$ while  $Al_{(M4)}$ ,  $Mg_{(M1)}$  are almost similar (average composition of chlorite from talc-bearing assemblage:  $Si_{(T1+T2)} = 2.7$ ,  $Fe_{(M1)} = 0.25$ ,  $Al_{(M1)} = 0.44$ ,  $\Box_{(M1)} = 0.2$ ,  $Al_{(T2)} = 1.3$ ,  $Mg_{(M2+M3)} = 1.1$ ,  $Fe_{(M2+M3)} = 2.5$ ,  $Al_{(M2+M3)} = 0.27$ ,  $Al_{(M4)} = 1$ ,  $Mg_{(M1)}$ = 0.11; average composition of chlorite from talcfree assemblage:  $Si_{(T1+T2)} = 3.01$ ,  $Fe_{(M1)} = 0.32$ ,  $\begin{array}{l} \text{Al}_{(M1)} = 0.70, \ \Box_{(M1)} = 0.43, \ \text{Al}_{(T2)} = 0.98, \ \text{Mg}_{(M2+M3)} \\ = 0.87, \ \text{Fe}_{(M2+M3)} = 2.25, \ \text{Al}_{(M2+M3)} = 0.14, \ \text{Al}_{(M4)} = 1, \ \text{Mg}_{(M1)} = 0.12) \quad (\text{Table } 2 \ ). \ \text{The chlorite} \end{array}$ compositions of both talc-bearing and talc-free assemblage deviate from the ideal clinochlore composition by Tschermak substitution (MgSi = Al.  $_{1}Al_{1}$ ) towards amesite, and they also deviate from

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Fe<sup>2+</sup>=Mg the clinochlore composition by substitution towards daphnite (Fig. 5A). Considering Vidal and Parra (2000) diagrams for major substitution, chlorite of the talc-free assemblage contains more amesite and sudoite endmembers. The deviation towards sudoite is related to di/trioctahedral substitution (Mg,  $Fe^{2+}$ )<sub>3</sub> =  $\Box Al^{-2}$ ) (Fig. 5A).

 
 Table. 2. Representative analyses of chlorites from talcbearing and talc-free quartz schists

Mineral Asset	Tlc-bearing	Tlc-bearing	Tlc-free	Tlc-free
Point No.	21	23	9	18
SiO <sub>2</sub>	24.51	23.18	28.44	28.50
TiO <sub>2</sub>	0.06	0.09	0.07	0.02
$Al_2O_3$	22.61	22.37	22.63	22.71
FeO	28.55	29.40	29.61	28.15
MnO	0.53	0.43	0.38	0.32
MgO	7.17	7.27	6.54	6.07
CaO	0.19	0.09	0.20	0.18
Na <sub>2</sub> O	0.18	0.13	0.10	0.06
$K_2O$	0.57	0.37	1.40	1.62
Total	84.37	88.33	89.37	87.63
Atom site (O=14)				
Si <sub>(T1+T2)</sub>	2.75	2.65	2.99	3.04
Al <sub>(T2)</sub>	1.25	1.35	1.01	0.96
Al <sub>(M4)</sub>	1.00	1.00	1.00	1.00
Mg(M1)	0.11	0.11	0.13	0.11
Fe <sub>(M1)</sub>	0.25	0.26	0.33	0.30
Al <sub>(M1)</sub>	0.40	0.47	0.55	0.86
□ <sub>(M1)</sub>	0.24	0.16	0.40	0.46
Mg(M2+M3)	1.08	1.13	0.90	0.85
Fe <sub>(M2+M3)</sub>	2.43	2.55	2.28	2.22
Al(M2+M3)	0.34	0.20	0.25	0.03

Cation site distributions follow Holland et al. (1998). All Fe was considered as FeO following Zane and Sassi (1998) and Guidotti et al. (1994c)



**Fig. 5.** Relevant substitutions responsible for the compositional variability of chlorites (A) and white micas (B) represented in SMFA and AKMF ternary diagrams. The mineral abbreviations are from Whitney and Evans (2010). TK: Tschermak substitution; DT: di/trioctahedral substitution; P: pyrophyllite substitution; △ : talc-bearing assemblage; □: talc-free assemblage

## 4.2. White mica

Following the method proposed by Vidal and Parra (2000), only white mica compositions that could be expressed as a linear combination of the celadonite, muscovite, paragonite and pyrophyllite end-members were used. Structural formulae of white micas were calculated from the remaining analyses on a 11-oxygen basis. Six distinct sites can be recognized within the crystal structure of white

mica: two tetrahedral  $(T1)_2$  and  $(T2)_2$ , three octahedral (M1), (M2) and (M3), and one interlayer A sites (Parra et al., 2002). The distribution of cations among the different sites is constrained by the following observations: tetrahedral ordering is favored in the 3T structure, for phengitic compositions, and for Si: Al<sup>IV</sup> ratios near 1:1 (Bailey, 1984). For the other polytypes, refinements did not lead to the detection of any substantial degree of tetrahedral ordering in micas (Bailey, 1984). However, Haselton et al. (1995) suggest that the distribution of Al and Si in tetrahedral sites of 2M1 muscovite is nearly ordered. This indicates short-range Al/Si ordering, at least to the extent of obeying the Al-avoidance role. In the following we assume a partial Si/Al ordering with T1 filled by Si only (Parra et al., 2002). Octahedral cation ordering occurs between cations of different sizes and charges. In particular,  $Al^{Vl}$  shows a strong preference for M2 and M3 sites, whereas the vacancies and the excess of Fe + Mg cations resulting from the di/trioctahedral substitution are primarily located in the larger M1 octahedrons. The cation site distribution model used here, which stems from the above constraints, is given in Table 3. This model does not differentiate the M2 and M3 sites of similar sizes (collectively referred to as the M2 site of multiplicity 2) (Parra et al., 2002). Comparing cation site distributions between white mica from two assemblages shows that white mica from talc-bearing assemblage contains higher amounts of  $Si_{(T1,T2)}$ ,  $Mg_{(M2+M3)}$  and  $K_{(A)}$  and lesser amounts of  $Al_{(T2)}$ ,  $Al_{(M2+M3)}$  and  $\Box_{(A)}$  than those of talc-free assemblage (average cation site distributions of white mica from talc-bearing assemblage:  $Si_{(T1+T2)} = 3.3$ ,  $Mg_{(M2+M3)} = 0.43$ ,  $K_{(A)} = 0.77$ ,  $Al_{(T2)} = 0.70$ ,  $Al_{(M2+M3)} = 1.54$ ,  $\Box_A = 0.16$ ; average cation site distributions of white mica from talc-free assemblage:  $Si_{(T1+T2)} = 2.98$ ,  $Mg_{(M2+M3)} =$  $0.03, K_{(A)} = 0.62, Al_{(T2)} = 1.01, Al_{(M2+M3)} = 1.96, \Box_A = 0.23$  (Table 3). By plotting white mica compositions on the Vidal and Parra (2000) diagram for major substitutions in white micas, it is evident that white mica composition from talcbearing assemblage deviates from ideal muscovite end-member by Tschermak substitution (MgSi = Al<sub>1</sub>Al<sub>1</sub>) towards celadonite and by Mg =  $Fe^{2+}$ substitution towards Fe-celadonite (Fig. 5B). But, the white mica from the talc-free assemblage is close to ideal muscovite end-member composition and only deviates towards pyrophyllite by pyrophyllite substitution (KAl = Si<sub>.1</sub>  $\Box$ ) (Fig. 5B).

### 4.3. Talc

Following Holland and Powell (1998, 1990) four distinct sites have been considered within the crystal structure of talc; two tetrahedral  $(T1)_2$  and

 $T2)_2$ , two octahedral (M1)<sub>2</sub> and (M3). The distribution of cations on the different sites (Table 4) follows Wei and Powell (2003), whereby octahedral Al is assumed to order onto the M3 site and tetrahedral Al is restricted to T1 on which it substitutes for Si. The two T2 sites are filled by Si only. Octahedral M3 site is filled by  $Al^{VI}$  and Mg + Fe, where as the excess of Mg + Fe and the possible resulting vacancies from di/trioctahedral substitution are located in two M1 octahedrons. The chemical composition of the talc deviates from the ideal talc composition by  $Mg = Fe^{2+}$  substitution towards Fe-talc and by  $MgSi = Al_{-1}Al_{-1}$  towards Tschermak-talc. A remarkable feature of talc analyses is the high percentage of iron especially in the talc not coexisting with phenigite or chlorite that leads to molar contents of minnesotatite  $(Fe_3Si_4O_{10}(OH_2))$ . Talc coexisting with phengite or chlorite contains higher amounts of Mg(M3) and  $Mg_{(M1)}$  and lesser amount of  $Fe_{(M1)}$  ( $Mg_{(M3)} = 0.98$ ,  $Mg_{(M1)} = 1.41$  and  $Fe_{(M1)} = 0.45$  for talc coexisting with phengite or chlorite and  $Mg_{(M3)} = 0.93$ ,  $Mg_{(M1)}$ = 0.40 and  $Fe_{(M1)}$  = 1.10 for talc not coexisting with phengite or chlorite) (Table 4). The relative distribution of Fe-Mg among coexisting phengite, chlorite and talc shows that relative to phengite and talc, chlorite is enriched in Fe-content while distribution coefficient for phengite-talc pair is close to one indicating relatively equal Fe-Mg partitioning between two minerals ( $K_D^{Ph-Chl} = 0.07$ ,  $K_D^{Ph-Tlc} = 0.93$ ,  $K_D^{Chl-Tlc} = 12.17$ ).

 Table 3. Representative analyses of white micas from talc-bearing and talc-free quartz schists

Mineral Asset	Tlc-bearing	Tlc-bearing	Tlc-free	Tlc-free
Point No.	17	12	16	25
SiO <sub>2</sub>	50.05	50.85	45.27	45.83
TiO <sub>2</sub>	0.26	0.14	0.16	0.18
$Al_2O_3$	29.35	28.92	37.86	39.33
FeO	1.48	1.50	1.29	1.65
MnO	0.27	0.28	0.05	0.07
MgO	4.36	5.12	0.33	0.64
CaO	0.05	0.04	0.04	0.13
Na <sub>2</sub> O	0.54	0.62	1.16	1.07
$K_2O$	9.34	9.21	8.10	6.82
Total	95.70	96.68	94.26	95.72
Atom site (O=11)				
Si(T1+T2)	3.29	3.30	3.10	2.97
Al <sub>(T2)</sub>	0.71	0.69	1.00	1.03
Al <sub>(M2+M3)</sub>	1.56	1.52	1.96	1.97
Mg(M2+M3)	0.41	0.45	0.05	0.02
Fe(M2+M3)	0.02	0.02	0.02	0.009
Mg <sub>(M1)</sub>	0.02	0.05	0.006	0.04
Fe <sub>(M1)</sub>	0.06	0.06	0.004	0.08
□(M1)	0.93	0.90	0.94	0.88
K <sub>(A)</sub>	0.78	0.76	0.69	0.56
Na <sub>(A)</sub>	0.07	0.08	0.15	0.13
□(A)	0.15	016	0.16	0.30

Cation distributions follow Parra et al. (2002). All Fe was considered as FeO following Guidotti et al. (1994c)

Table 4. Representative analyses of talc

Point No.	7	8	8	6
SiO <sub>2</sub>	61.12	61.20	62.32	60.21
TiO <sub>2</sub>	0.02	0.07	0.02	0.08
$Al_2O_3$	0.25	0.30	1.36	0.27
FeO	8.17	8.02	19.22	19.02
MnO	0.05	0.00	0.18	0.00
MgO	23.62	24.98	10.40	15.97
CaO	0.12	0.02	0.30	0.04
$Na_2O$	0.00	0.00	0.09	0.10
$K_2O$	0.00	0.00	0.30	0.32
Total	94.19	96.01	94.19	96.01
Atom site (O=11)				
$Si_{(T1+T2)}$	4.00	4.00	4.30	4.11
Al <sub>(M3)</sub>	0.02	0.02	0.11	0.02
$Mg_{(M3)}$	0.98	0.98	0.89	0.98
$Mg_{(M1)}$	1.41	1.46	0.40	0.64
Fe <sub>(M1)</sub>	0.45	0.44	1.11	1.08
□(M1)	0.14	0.10	0.75	0.28

Cation site distributions follow Wei and Powell (2003). All Fe was considered as FeO following Guidotti et al. (1994c)

#### 4.4. K-feldspar

EPMA analyses of K-feldspar show that its chemical composition is close to the ideal K-feldspar and is homogeneous (Table 5).

 Table 5. Representative analyses of K-feldspars from talc-bearing and talc-free quartz schists

Mineral Asset	Tlc-bearing	Tlc-bearing	Tlc-free	Tlc-free
Point No.	27	28	30	31
SiO <sub>2</sub>	65.21	64.71	64.02	65.43
TiO <sub>2</sub>	0.16	0.21	0.10	0.05
$Al_2O_3$	18.62	19.71	19.86	18.11
MgO	0.14	0.06	0.02	0.10
CaO	0.15	0.12	0.14	0.11
Na <sub>2</sub> O	0.67	0.98	0.84	0.76
K <sub>2</sub> O	14.96	14.13	14.44	14.85
Total	99.92	99.93	99.43	99.42
Atom site (O=8)				
Si	3.00	2.97	2.95	3.00
Ti	0.006	0.007	0.003	0.002
Al	1.00	1.06	1.08	0.98
Mg	0.01	0.004	0.001	0.007
Ca	0.007	0.006	0.007	0.005
Na	0.06	0.09	0.08	0.07
K	0.88	0.83	0.85	0.87
Total	4.96	4.96	4.97	4.96

#### 5. Discussion

Massonne and Schreyer (1989) purely on theoretical grounds in simplified KMASH model system, proposed a high-pressure condition for talcphengite-K-feldspar assemblage. They suggest that two reactions:

(1) chlorite + phlogopite + quartz = talc + phengite + H<sub>2</sub>O

(2) chlorite + K-feldspar = phengite + phlogopite + quartz +  $H_2O$  intersect at an invariant point about 11 kbar and around 350°C. The locations of both reactions have been determined experimentally by Massonne and Schreyer (1987, 1989). Thus, the resulting invariant point contains chlorite, K-feldspar, phengite, phlogopite, quartz, talc and  $H_2O$ .

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From this invariant point, three more reactions could theoretically be obtained. Reaction (3) represents the low-temperature limit of the talc-phengite assemblage:

(3) chlorite + K-feldsper + quartz = talc + phengite  $+ H_2O$ 

The other two are phlogopite + quartz breakdown reactions:

(4) phlogopite + quartz +  $H_2O$  = talc + K-feldspar + chlorite

(5) phlogopoite + quartz +  $H_2O$  = talc + phengite + K-feldspar

With this configuration, the talc-phengite-Kfeldspar assemblage could be stable under highpressure conditions, whereas the chlorite-Kfeldspar-phlogopite is a stable assemblage under low-pressure conditions. However, talc-bearing quartz schists in this study contain significant amount of Fe as well as phases with considerable solid-solutions such as phengite, chlorite and talc. Therefore, the petrogenetic gird proposed by Massonne and Schreyer (1989) cannot be directly applied for the mineral assemblage of the present samples. To consider the P-T stability of talcphengite-chlorite-K-feldspar assemblage in the studied rocks, a petrogenetic gird was constructed in the K<sub>2</sub>O-FeO-MgO-Al<sub>2</sub>O<sub>3</sub>-SiO<sub>2</sub>-H<sub>2</sub>O (KFMASH) model system with talc, chlorite, phengite, Kfeldspar, phlogopite and kyanite assuming excess quartz and H<sub>2</sub>O. We used the internally consistent thermodynamic dataset of Holland and Powell (1998) and the THERMOCALC 3.21 software (Powell et al., 1998). A quaternary symmetric mixing model was adopted for chlorite following Holland et al. (1998). Interaction parameters are W(afchl, clin) = 18 kjmol-1, W(afchl, daph) = 14.5kjmol-1, W(clin, ames) = 18 kjmol-1 and W(daph, ames) = 13.5kjmol-1. Following Parra et al. (2002) a non-ideal mixing model was used for white mica considering muscovite, celadonite and Feceladonite end-members. The interaction margules parameters ( $W_{Al-Mg}$  and  $W_{Al-Fe}$  at M2 site) were calculated similar to that adopted by Parra et al. (2002) and the intersite interactions of  $W_{Al-Si}$  at T2 and W<sub>Fe-Mg</sub> at M2+M1 were assumed to be zero. For talc, an ideal mixing model was used in which the Al is assumed to order onto the M3 site and to enter only the two T1 sites. Based on Gibbs phase rule, an invariant point for a 6 system component in a P-T field requires that 8 phases be in equilibrium, and a univariant reaction curve requires 7 phases. For the general case, one system with 8 phases (phengite, talc, chlorite, K-feldspar, kyanite, phlogopite, quartz, H<sub>2</sub>O) and 6 components (K<sub>2</sub>O, FeO, MgO, Al<sub>2</sub>O<sub>3</sub>, SiO<sub>2</sub>, H<sub>2</sub>O) contains one invariant point, 6 reaction curves and 15 divariant fields considering quartz and H<sub>2</sub>O as excess phases (Fig. 6) (Table 6). In the obtained petrogenetic gird,

the divariant field of talc-phengite-chlorite-K-feldspar, comparable with the observed assemblage in talc-bearing quartz schist, is stable over a wide range of P-T condition defined by the following two univariant reactions (Fig. 6).

phengite + talc + quartz = chlorite + K-feldspar + kyanite +  $H_2O$ 

chlorite + phlogopite + quartz = talc + phengite + K-feldspar +  $H_2O$ 



**Fig. 6.** Petrogenetic grid in the KMASH model system containing chlorite, phengite, K-feldspar, kyanite and phlogopite with excess  $SiO_2 + H_2O$ . The mineral abbreviations are from Whitney and Evans (2010)

**Table 6.** Univariant reactions (A) and divariant assemblages (B) in the KFMASH model system containing phengite, chlorite, talc, kyanite and phlogopite with excess quartz and  $H_2O$ 

A	
Absent phase	Univariant reactions
[Chl]	Ph + Phl + Qz = Tlc + Kfs + Ky + H2O
[Ph1]	Chl + Phl + Qz = Tlc + Kfs + Ky + H2O
[Tlc]	Ph + Phl + Qz = Chl + Kfs + Ky + H2O
[Kfs]	Chl + Ky + Phl + Qz = Ph + Tlc + H2O
[Ky]	Chl + Phl + Qz = Tlc + Ph + Kfs + H2O
[Phl]	Ph + Tlc + Qz = Chl + Kfs + Ky + H2O
B	
Absent phase	Univariant assemblages
[Chl + Ph]	Tlc + Kfs + Ky + Phl
[Chl + Tlc]	Ph + Kfs + Ky + Phl
[Chl + Kfs]	Ph + Tlc + Ky + Phl
[Chl + Ky]	Ph + Tlc + Kfs + Phl
[Chl + Phl]	Ph + Tlc + Kfs + Ky
[Ph + Tlc]	Chl + Kfs + Ky + Phl
[Ph + Kfs]	Chl + Tlc + Ky + Phl
[Ph + Ky]	Chl + Tlc + Kfs + Phl
[Ph + Phl]	Chl + Tlc + Kfs + Ky
[Tlc + Kfs]	Chl + Ph + Ky + Phl
[Tlc + Ky]	Chl + Ph + Kfs + Phl
[Tlc + Phl]	Chl + Ph + Kfs + Ky
[Kfs + Ky]	Chl + Ph + Tlc + Phl
[Kfs + Phl]	Chl + Ph + Tlc + Ky
$[K_{V} + Phl]$	Chl + Ph + Kfs + Tlc

This assemblage is stable above 5 kbar pressure and between 400°C and 800°C. Therefore, to find the certain P-T condition of the observed assemblage one way is to find a relationship between compositions of talc, phengite and chlorite in different P-T conditions by construction compatibility diagrams. The compatibility diagrams show all phase relationships for all compositions in a defined model system at the specified P-T conditions. To illustrate the changes in the compositions of coexisting minerals with respect to P-T conditions, a series of calculated Al<sub>2</sub>O<sub>3</sub>-KAlO<sub>2</sub>-MgO+FeO (AKM) compatibility diagrams with phengite, chlorite and talc were drawn for a traverse from 5 kbar to 11 kbar (Fig. 7). For construction the compatibility diagrams at given P-T points (e.g. 5kbar-400°C, 5kbar- 600°C), the mode 1 (phase diagram calculations) of the THERMOCALC 3.21 software (Powell et al., 1998) and internally consistent thermodynamic dataset of Holland and Powell (1998) were used. In the obtained compatibility diagrams, the compositions of coexisting minerals (phengite, chlorite and talc) have been calculated by the following parameters:  $X_{Ph} = Fe/Fe+Mg$ ,  $Y_{Ph} = X(Al, M2A)$ ,  $X_{Chl} =$ Fe/Fe+Mg,  $Y_{Chl} = X(Al, T2)$ ,  $X_{Tlc} = Fe/Fe+Mg$  and  $Y_{Tlc} = X(Al, M3)$ . In the calculated compatibility diagrams, increasing pressure and temperature, with the exception of phengite composition at P = 11kbar and T = 400°C, leads to increasing of  $X_{Ph}$ ,  $X_{Chl}$ and  $X_{Tlc}$  (for example at P = 5 kbar and T = 400°C:  $X_{Ph}$  = 0.473,  $Y_{Ph}$  = 0.776,  $X_{Chl}$  = 0.163,  $Y_{Chl}$  = 0.517,  $X_{Tlc} = 0.064$ ,  $Y_{Tlc} = 0.066$  but at P = 11 kbar and T = 600°C:  $X_{Ph} = 0.944$ ,  $Y_{Ph} = 0.528$ ,  $X_{Chl} = 0.865$ ,  $Y_{Chl} = 0.544$ ,  $X_{Tlc} = 0.750$ ,  $Y_{Tlc} = 0.130$ ) (Fig. 7). Comparing the compositions of the observed minerals of phengite, chlorite and talc from talc-bearing quartz schists of the Nahavand area with those of calculated compatibility diagrams shows that the greatest similarity can be seen at P = 11 kbar and T = 400°C (compositions of phengite, chlorite and talc from talc-bearing quartz schits:  $X_{Ph} = 0.280$ ,  $Y_{Ph} = 0.860$ ,  $X_{Chl} = 0.570$ ,  $Y_{Chl} = 0.640$ ,  $X_{Tlc} = 0.160$ ,  $Y_{Tlc} = 0.02$ ; compositions of phengite, chlorite and talc shown by compatibility diagram;  $X_{Ph} = 0.244$ ,  $Y_{Ph} = 0.828$ ,  $X_{Chl} = 0.610$ ,  $Y_{Chl} = 0.644$ ,  $X_{Tlc} = 0.150$ ,  $Y_{Tlc} = 0.03$ ) (Fig. 7). Therefore, the observed coexisting minerals of phengite ( $X_{Ph} = 0.280$ ,  $Y_{Ph} = 0.860$ ), chlorite ( $X_{Chl}$  $= 0.570, Y_{Chl} = 0.640), talc (X_{Tlc} = 0.160, Y_{Tlc} =$ (0.020) and K-feldspar are stable at P = 11 kbar and  $T = 400^{\circ}C$ 

The other way to find the relationships between chemical compositions of minerals in equilibrium, is the construction of P-T diagram with the chemical isopleths of coexisting minerals. Because, the Si content in phengite is changing more markedly, it has been used to conclude the pressure of equilibrations at a defined temperature. P-T diagram showing the stability of divariant assemblage of talc-phengite-chlorite-K-feldspar is represented in Fig. 8. Univariant reaction curves restricting the assemblage to the low- and high-T sides are taken from Fig. 6. The Si content of phengite was calculated by the mode 1 of the THERMOCALC 3.21 software (Powell et al., 1998). Considering average Si content of the analysed phengite (Si=3.3) (Table 3), indicate formation pressure of 12 kbar at specified temperature of 580°C which is invariant temperature in Fig. 6.



Fig. 7. Calculated  $Al_2O_3$ -KAlO<sub>2</sub>-MgO+FeO (AKM) compatibility diagrams illustrating the changes in the compositions of coexisting minerals of phengite, chlorite and talc with excess quartz and  $H_2O$  with respect to P-T conditions. The mineral abbreviations are from Whitney and Evans (2010)



**Fig. 8.** Calculated P-T diagram with isopleths of Si content in phengite. Univariant curves restricting the assemblage of talc-phengite-chlorite-K-feldspar to the low- and high-T sides are taken from Fig. 6

The relatively high-pressure condition of talcphengite-chlorite-K-feldspar assemblage in talcbearing quartz schists from north of the Nahavand

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area indicates that this assemblage could be formed during subduction of the Neo-Tethys oceanic plate under the Iranian microcontinent. Such an interpretation is in agreement with Berberian and King (1981), Alavi (1994), Mohajjel et al. (2003), Ghasemi and Talbot (2006) and Dvoudian et al.'s (2007) conclusions.

### 6. Conclusion

1-For the first time, an unusual assemblage of talcphengite-chlorite-K-feldspar was found in quartz schists in the Nahavand area from the Sanandaj-Sirjan zone.

2-Based on the field occurrences and mineral assemblages, the quartz schists of the Nahavand region can be divided into talc-bearing and talc-free types.

3-Constructed petrogenetic grid in the KFMASH model system shows that talc-phengite-chlorite-K-feldspar assemblage is stable over a wide range of P-T condition. However the AKM compatibility diagram predicts that phengite ( $X_{Ph} = 0.280$ ,  $Y_{Ph} = 0.860$ ), chlorite ( $X_{Chl} = 0.570$ ,  $Y_{Chl} = 0.640$ ), talc ( $X_{Tlc} = 0.160$ ,  $Y_{Tlc} = 0.02$ ), K-feldspar with excess quartz and H<sub>2</sub>O could be stable at P = 11 kbar and T = 400°C.

4-The relatively high-pressure assemblage of talcphengite-chlorite-K-feldspar could be formed due to the Neo-Tethys oceanic plate subduction under Iranian microcontinent.

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