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Trace elements in hazardous mineral fibres

(Article begins on next page)

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TG/DSC study of the thermal behaviour of hazardous mineral fibres

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ABSTRACT

For the first time, this paper reports a systematic and comparative study of the thermal behaviour of fibres of social, health, economic and industrial relevance using thermogravimetric and differential scanning calorimetry (TG/DSC). The mineral fibres selected for the study are: three chrysotiles samples, crocidolite, tremolite asbestos, amosite, anthophyllite asbestos and fibrous erionite. Powder X-ray diffraction (PXRD) and scanning electron microscopy combined with energy dispersive spectrometry (SEM/EDS) were used for the characterization of the mineral fibres before and after heating at 1000 or 1100 °C to identify the products of the thermal decomposition at a microscopic and structural scale and characterize their thermal behaviour.

TG/DSC data allowed the determination of the structural water content and temperature stability. Furthermore thermal analysis provided a sensitive and reliable technique for the detection of small quantities of different mineral phases occurring as impurities. After thermal treatment, fibrous samples were completely transformed into various iron oxide, cristobalite and other silicate phases which preserved the original overall fibrous morphology (as pseudomorphosis). Only crocidolite at 1100 °C was partially melted and an amorphous surface was observed.

Keywords: Mineral fibres, Asbestos, Thermal behaviour, Dehydroxylation, Pseudomorphosis.

Introduction

Among the minerals which are considered particularly hazardous for human health, the most notorious display a fibrous-asbestiform crystal habit [1]. It is now widely accepted to include in the family of the so-called asbestos mineral the following species: serpentine chrysotile, five amphibole species named tremolite and actinolite asbestos, crocidolite, amosite, anthophyllite asbestos and the fibrous zeolite erionite [2].

Chrysotile is a member of the serpentine group, a 1:1 layer silicate (constituted by a tetrahedral and a trioctahedral sheet) whose layers are rolled so to assume a characteristic fibrous habit [3, 4]. Amphiboles are double-chain silicates which may display a fibrous habit being structurally elongated in one preferred crystal direction. Finally, erionite is a common fibrous/acicular zeolite with an hexagonal, cage-like structure composed of a framework of linked tetrahedral [5, 6]. Despite their outstanding technological properties (e. g. low thermal conductivity, high mechanical strength, workability, among the others) which prompt their widespread industrial applications, asbestos minerals are considered hazardous. In general, all asbestos fibres if inhaled are thought to induce malignant mesothelioma, lung cancer (in combination with other factors), and other lung diseases [7, 8]. According to the existing regulations, amphibole asbestos fibres are banned worldwide whereas chrysotile is banned in only 28% of the countries worldwide. In the other countries, safe use of chrysotile is admitted. Asbestos erionite unfortunately is not regulated but listed by International Agency for Research of Cancer (IARC) as substance carcinogenic to humans. As a matter of fact, *in vivo* studies unequivocally proved that asbestiform erionite is more tumorigenic than chrysotile and crocidolite asbestos [9].

Since the advent of industrial age, asbestos fibres have been extensively used in an endless number of industrial applications and especially to manufacture various types of artefacts (asbestos cement, disc brake pads, pipes, reinforcing agents, fire retardants etc). In those countries where all asbestos minerals are banned, and remediation policies are fostered, many attempts were made to detoxify asbestos minerals by using different techniques [10-12]. In this regard, many projects and patents have dealt with the possible disposal and re-use of asbestos-containing materials (ACM) via the crystal-chemical transformation induced by thermal treatment [13-18]. However, it must be considered that the thermal transformations sequence of asbestos-containing materials (ACM), composed of a variety of different crystalline and amorphous phases, is totally different with respect to the transformations sequence of pure asbestos minerals.

Concerning pure chrysotile, it has been demonstrated that its structure collapse at around 650 °C with early recrystallization at about 800 °C into anhydrous silicates such as forsterite and amorphous silica [19]. With respect to chrysotile, papers dealing with the thermal decomposition of pure amphibole asbestos and erionite are rare. As a matter of fact, only few studies were devoted to the thermal decomposition of pure amphibole asbestos up to 1100 °C and rare data report on the new phases appeared after thermal treatment. Thermal decomposition of fibrous amphibole minerals have been generally limited to asbestos minerals of commercial value or health concern such as crocidolite [20-25] and amosite [23, 26, 27]. Moreover, in some papers regarding the thermal analysis of amphiboles asbestos, a full picture of their thermal behaviour is not given [28]. Regarding erionite, the thermal behaviour of a sample from Jersey Nevada (USA) is described in only one paper [29] which showed that the main endothermic event has occurred at about 140 °C but did not report the TG curve neither the temperature of the structural collapse. In this scenario, the aim of this study was to systematically investigate and compare the thermal behaviour (TG/DSC) as well as the phase transformations of the most relevant mineral fibres during heating up to 1000 or 1100 °C. The study was performed on eight selected fibrous minerals including chrysotile, crocidolite (asbestiform riebeckite), tremolite asbestos, anthophyllite asbestos, amosite (grunerite asbestos) and fibrous erionite object of many important biomedical studies [30, 31]. Four fibrous species (chrysotile, crocidolite, amosite, and anthophyllite asbestos) were

distributed by the International Union Against Cancer (UICC).

The deep knowledge of thermal behaviour of these asbestos minerals may deliver data relevant for the understanding of the crystal-chemical transformations of asbestos through thermal treatment and for the identification of asbestos mineral fibres in bulk natural samples using TG/DSC.

Furthermore, since both asbestos samples and asbestiform erionite are used for *in vitro* studies to test their cyto-toxicity [30, 31], the determination of the presence of impurities may be crucial as such contaminants may adversely affect the experimental results.

Table 1 reports the nature and chemical formula (after Pollastri et al., 2015 [32]) of the eight investigated fibres: UICC chrysotile from Canada; chrysotile from Balangero (Italy); chrysotile from Val Malenco (Italy); UICC crocidolite from Koegas Mine, Northern Cape (S. Africa); tremolite asbestos from Val d'Ala (Italy); UICC amosite from Penge mine, Northern Province (S. Africa); UICC anthophyllite asbestos from Paakkila (Finland) and asbestiform erionite from Jersey Nevada (USA). The major focus of the work was to systematically characterize the mineral fibres by thermal analysis specifically thermogravimetric (TG), derivative thermogravimetric (DTG), differential scanning calorimetry (DSC), derivative differential scanning calorimetry (DDSC). TG and DSC were performed in an alumina crucible under a constant nitrogen flow of 30 cm³min⁻¹ with a Netzsch STA 449 C Jupiter in a $25 - 1000$ and $25 - 1100$ °C temperature range, with a heating rate of 10 °C/min. Instrumental precision was checked by six repeated collections on a kaolinite reference sample revealing good reproducibility (instrumental theoretical T precision of \pm 1.2 °C) and theoretical weight sensitivity of 0.10 µg, DSC detection limit < 1 µW. Samples were powdered by dry-grinding in an agate mortar; about 40 mg of each sample were used in all collections. Owing to the remarkable length of the chrysotile fibres from Val Malenco, before grinding they were cut with scissors.

A qualitative phase analysis, both of natural and heated samples, was performed according to the powder X-ray diffraction method (PXRD) using a Bruker D8 Advance X-ray diffractometer at 40 kV and 40 mA. The instrument is equipped with a copper tube and curved graphite monochromator. Scans were recorded in the range of 3–66 °2θ, with a step interval of 0.02 °2θ and a step-counting time of 3 s/step. EVA software (DIFFRACplus EVA) was used to identify the mineral phases and experimental peaks being compared with the 2005 PDF2 reference patterns. The morphology of the samples before and after thermal analysis was investigated by scanning electron microscopy (SEM)

using an Environmental Scanning Electron Microscope FEI QUANTA 200 equipped with an EDAX Genesis 4000 energy dispersive X-ray spectrometer (EDS), and a FEI Nova NanoSEM 450 equipped with an X-EDS Bruker QUATAX-200 system for the microanalysis.

Results and discussion

Chrysotile

The thermal analysis for chrysotile UICC (Fig. 1) showed four endothermic peaks at 226, 401, 520 and 633 °C. The first peak at 226 °C may be due to the dehydroxylation of pyroaurite [33]; the peak at 401 °C to the dehydroxylation of brucite and de-oxygenation of pyroaurite [33, 34]; the very weak peak at 520 °C is thought to be generated by the decarbonation of siderite [35], likely present as very minor impurity; the wide peak at 633 °C to the chrysotile dehydroxylation [22, 36]. The weak endothermic event at 901 °C on DTG (Fig. 1) curve was ascribed to talc dehydroxilation [37, 38]. It is possible that the wide endothermic event at 633 °C hides minor endothermic events due to the decarbonation of calcite, dolomite, and/or dehydroxylation of clinochlore [34, 36, 37]. The TG curve showed a weight loss of 0.91 % below 110 °C due to adsorbed water while the main weight loss of 12.22 % was due to chrysotile dehydroxylation (Table 2). Our findings are in line with previous literature data on chrysotile dehydration mechanisms and

high-*T* crystallization [39-42]. The product of the dehydroxylation of chrysotile recrystallized to forsterite [19, 43, 44] caused a sharp exothermic peak at 823 °C (Fig. 1, Table 3). Indeed, the corresponding PXRD pattern (Fig. 2) of the chrysotile UICC after thermal treatment confirmed the presence of the forsterite.

The DSC curve of the chrysotile from Balangero exhibited one major effect (Fig. 3) at 660 °C related to the chrysotile dehydroxylation with a weight loss of 11.80 %. As already described above, the weak endothermic effect (Fig. 3, Table 3) at 402 °C is due to brucite [34] breakdown. In the DTG curve the peak at 869 °C was related to the decarbonation of dolomite [45].The DSC weak

shoulder effect at 938 °C which was clearly recorded on DDSC curve was related to the talc dehydroxilation [38]. Again, the wide endothermic event at 660 °C may hide minor endothermic events due to the I decarbonation of dolomite, calcite, and clinochlore [34, 36, 37]. The weak endothermic effect at 717 °C visible on the DTG curve (Fig. 3) is due to the antigorite dehydroxylation [36, 46]. The effect on TG curve (- 0.57 wt%) below 110 °C was ascribed to the release of humidity adsorbed at the sample surface while the total weight loss at 1000 °C was of 14.9 % (Table 2). On the DSC curve the exothermic peak at 822 °C (Fig. 3) was related to the crystallization of forsterite as determined by PXRD after thermal treatment at 1000 °C (Fig. 2). The TG curve for the chrysotile from Val Malenco (Fig. 4) showed a continuous weight loss mainly due to the decomposition of chrysotile in correspondence with the major endothermic event at 652 °C (see the DSC curve in Fig. 4). Dehydroxylation of chrysotile causes a weight loss of 12.01 %. DTG weak effect at 760 °C (Fig. 4) is the diagnostic signal [47] of the presence of antigorite in the sample which was also detected by TEM analysis on the same sample by Cattaneo et al. [48]. Broad DTG and DSC signals in the 25–110 \degree C range were due to adsorbed water (weight loss of 0.54 %) while the total weight loss at 1000 °C was 13.35 % (Table 2). A sharp exothermic peak at 820 °C indicates the crystallization of forsterite [19, 43, 44] as confirmed by PXRD data (Fig. 2). As showed in Fig. 5 the curves of the three chrysotile samples have similar trend; no significant variation in the exothermal peaks (range 820-823 °C) was observed, while a slight difference in the main endothermic effect occurred. The endothermic peak observed for the chrysotile UICC shows a shift versus lower temperature (633 °C) with respect to both chrysotile from Val Malenco (652 °C) and chrysotile from Balangero (660 °C). The different temperature of chrysotile decomposition (range 633-660 °C) of the three chrysotile samples may be due to the different fibre size (width and length) distribution and the presence of different amount of Fe [43, 49] and Ni [44] substitute for Mg in octahedral sheet [32]. The relationships between distribution size, Ni, Fe content and change of temperature stability of chrysotile are complex. In fact, as recently demonstrated, the thermal

shift in chrysotile decomposition can be also related to the presence of trace elements in its structure (i.e. Li, As, Sb) [50] which may control the range of temperature over that the structure collapses. The TG curves of the three specimens of chrysotile showed a weight loss of about 12 % (Table 2) due to their decomposition [19, 51]. These data match the theoretical and experimental values of mass loss observed in natural and synthetic chrysotile fibres reported in literature [19, 43, 47, 51]. A representative set of secondary electron SEM images showing the morphology of chrysotile before and after thermal analysis at 1000 °C is reported in Fig. 6. The unheated fibres of chrysotile samples appear arranged in bundles (Fig. 6a, 6d) and curved with their typical wavy appearance (Fig. 6g). The splitting in fibrils starting from the fibre bundle is shown in Figures 6a and 6d . After thermal treatment at 1000 °C, chrysotile fibres recrystallize in forsterite apparently retaining the original fibrous crystal habit (pseudomorphosis phenomenon) [52, 53] and appearing curved and still flexible (Fig. 6b, 6e, 6h). However, at higher magnification (HM) the apparent fibres turn out to be a continuous sequence of sub-cylindrical particles with basis both sharp and perpendicular to the original fibre axis [16], approximately 100 nm in length (Fig. 6c), Sometimes, the new silicate is constituted by sub-spherical particles disposed not very tidily along the axis (Fig. 6f). The original cleavage parallel to the fibre axis is lost (Fig. 6c). Therefore, the eventual fracture of the transformed pseudo-morphic fibres occurs at the particle boundaries and not along the fibre axis. Moreover, as observed in Fig 6i, the typical smooth surface of chrysotile fibres is completely lost becoming very rough.

Amphiboles

Crocidolite

The DTG curve of crocidolite (Fig. 7) shows a major endothermic event at 648 °C related to iron oxidation accompanied by dehydrogenation and/or dehydroxylation. The structure does not show collapse which occurs at higher temperature [24] with a corresponding weight loss of 2.18 % in the range 110-680 °C of the TG curve. No weight loss of crocidolite was observed above 700 °C in agreement with Fujishige et al. [54]. As it can be observed in the DTG curve (Fig. 7), the partial oxidation of ferrous iron content in the crocidolite takes place in the range 200-580 °C [55], as confirmed by the three weight gain peaks at 205, 360, and 570 °C (Fig. 7). Although the detailed discussion of the processes is not the object of this paper, it is necessary to point out that the oxidation of crocidolite is essentially a dehydrogenation, as long as dehydroxylation occurs [21] so that the mechanism of oxidation was considered to be dependent on migration of protons and electrons through the crystal. Indeed, when hydrated silicates containing ferrous iron are heated the constitutional hydroxyl decomposes and iron may change its valence state. Oxidation may result from either the incorporation of oxygen into the material (oxygenation) or a dehydrogenation with the following mechanism: $Fe^{2+} + OH^- \rightarrow Fe^{3+} + O^{2-} + H$ [56].

The DSC curve up to 700 °C showed two exothermic and two endothermic effect: 320 °C, 431 °C. °C and 649 °C respectively (Table 3). The first exothermic effect at 320 °C was related to the crocidolite dehydrogenation as confirmed by the weight gain peak at 360 °C in the DTG curve due to consequent Fe²⁺ oxidation. The endothermic effect at 354 °C should be related to the dehydroxylation of hydroxyl water of the crocidolite with a consequent formation of oxy-crocidolite (crocidolite partially dehydrogenated) [23]. The exothermic peak at 431 °C (Fig. 7) was related to the second dehydrogenation as confirmed by the DTG weight gain at 570 °C which is a consequence of Fe^{2+} oxidation. The weak endothermic effect at 649 °C represents the total dehydroxylation of crocidolite in good agreement whit data reported by Hodgson et al. [20] in which the total dehydroxylation of crocidolite was observed in the range of 570-700 °C. The broad exothermic effect at 850 °C (Fig. 7) was related to the structure collapse and formation of magnetite, cristobalite, and a pyroxene phase (acmite, $NaFeSi₂O₆$) [20].

At 928 °C, the DSC curve shows an endothermic peak due to the conversion of magnetite to hematite. Indeed, in the DTG curve the peak at 928 °C stems from the oxidation of the ferrous iron present in magnetite ($Fe^{+2}Fe^{+3}O_4$) which involves a weight gain (Fig. 7). The DSC shoulder effect at 960 °C, which is clearly recorded on DDSC curve, was related to the incongruent melting point of the acmite with separation of hematite [57, 58]. The endothermic peak observed at 1064 °C in the DDSC curve indicated that cristobalite is being dissolved by the liquid [20]. Indeed, at 1100 °C the final minerals products detected by PXRD were mainly hematite (33-0664 JCPDS card.) while cristobalite (02-0278 JCPDS card) is evidently in a small amount because its reflections are close to the detection limit (Fig. 8).

The structural changes of crocidolite with increasing temperature can be summarized in the following steps: dehydrogenation and/or dehydroxylation accompanied by iron oxidation, structure collapse and crystallization of newly formed crystalline phases, early melting. Crocidolite, blue at room temperature, turned into dark red at 1100 °C, mostly due to hematite formation.

At SEM, the raw fibres of crocidolite appear as straight and rigid, looking like needles (Fig. 9a). After heating at 1100 °C, the original morphology is strongly altered (Fig. 9b). The single fibres, originally arranged in fibre bundles (Fig. 9a), now appear as thick sticks, confirming that partial melting occurred during heating. However, it is still possible to recognize some fibrous-like structure that was melt-bonded (Fig. 9b). At higher magnification (HM) crocidolite showed meltfragments composed of an aggregate of particles with totally different morphology with respect to the original morphology (Fig. 9c).

Tremolite

DSC curve of tremolite asbestos from Val d'Ala (Fig. 10) exhibits a number of both endothermic and exothermic peaks in the range 500-1100 °C that can be explained by the presence of impurities in the sample [59]. The endothermic events at 729 °C and 776 °C are due to the dehydroxylation of minor chlorite and antigorite, respectively [37, 36] (see Fig. 10 and Table 3). The exothermic peak at 842 °C was related to oxidation of $\rm Fe^{2+}$ [60] present in the chlorite, its presence being also proved by the weight gain in the DTG curve. The exothermic effect at 898 °C is interpreted as

recrystallization to forsterite and hematite. The shoulder at 955 °C on DSC curve was ascribed to talc breakdown [38], while the sharp endothermic peak at 1046 °C corresponds to breakdown of tremolite in agreement with Luckewicz [61]. The main TG weight loss of 2.02 % between 850 and 1050 °C due to the tremolite dehydroxylation (Fig. 10, Table 2) was in agreement with the theoretical tremolite water content [62]. Finally, the exothermic DSC signal at 1077 °C was related to the crystallization of diopside. The effect on TG curve (-0.17 wt%) below 110 °C was ascribed to the release of humidity adsorbed at the sample surface. The mineral products after heating to 1100 °C were diopside (JCPDS card 11-0654), forsterite and hematite deriving from tremolite and chlorite breakdown (Fig. 8).

Figure 9d depicts the typical morphology of tremolite fibres; they are stiff and exhibit a prismatic rod shaped morphology. After thermal treatment, the crystal habit is preserved but almost all the individuals result more brittle and fractured nearly perpendicular to the fibre axis (Fig. 9e). The HM Fig. 9f depicts the presence of newly-formed polyhedral crystals on the surface of the pristine tremolite asbestos some of which exhibit definite morphology with clear evidence of edges and faces. The growth occurs through the formation of recrystallizing-islands with heterogeneous lens shape. On the surface of the pristine tremolite asbestos the growth and coalescence of the recrystallizing-islands evolve from lens to faceted crystals. EDS/SEM investigations confirmed that the pseudomorphic process involves a complete recrystallization of the original tremolite asbestos into diopside.

Amosite

Characteristic DSC/TG thermogram of amosite is presented in Fig. 11. The first DSC weak peak at 288 °C is caused by the breakdown of goethite [63] present as impurity which transforms into hematite [64]. The second broad endothermic effect at 741 °C (Fig. 11) was related to the structural breakdown of amosite (Table 3), although amosite decomposition started at 600 °C with the formation of oxy-amosite [23, 26] and ended at about 850 °C in agreement with the literature data

[22, 23, 27]. Finally, the DSC curve shows a broad exothermic peak at 878 °C due to enstatite and hematite formation, as confirmed by the PXRD pattern (Fig. 12). The very weak endothermic effect in the range 530-580 °C (Fig. 11), evidenced in the DDSC curve, could be due to the structural $\alpha \rightarrow \beta$ transition of quartz [65] which is present in the sample as impurity (Fig. 12).

The TG curve of amosite showed a continuous weight loss of 1.94 % between 110 and 690 °C, due to dehydroxylation and dehydrogenation reactions [21, 26] which involve a weight gain of 0.24 % between 690 and 1000 °C, due to oxidation of ferrous iron (Fig. 11). In fact, hematite (JCPDS card 24-0072) was also found among the final mineral products (Fig. 12) in addition to enstatite (JCPDS card 07-0216) and quartz (JCPDS card 07-0346). The effect on TG curve (-0.33 wt%) below 110 °C was ascribed to the release of humidity adsorbed at the sample surface.

Raw amosite (Fig. 13a) shows fibres which look like flexible needle arranged in bundles. After heating at 1100 °C, the newly-formed silicate (enstatite) preserved the original fibrous morphology (pseudo-morphosis) but fibres appear more rigid and thicker (Fig. 13b). In some cases, single fibres seem to be fused together at forming prismatic crystal (Fig. 13b, 13c) and when observed at higher magnification, they appear partially covered by pseudo-spherical particles growing along the axial direction of the fibres (Fig. 13c).

Anthophyllite asbestos

Figure 14 presents the thermal behaviour of anthophyllite asbestos. DTG curve shows one main peak of maximum weight loss (2.30 %) at 868 °C due to anthophyllite dehydroxylation in correspondence with the shoulder at 824 °C on DSC curve. However, DDSC confirm the presence of an endothermic peak at 861 °C which corresponds to the structural breakdown of this phase [22, 27] followed by recrystallization of enstatite as showed by the exothermic peak at 915 °C (Fig. 14). According to Freeman [27] the decomposition temperatures of anthophyllite asbestos as well as tremolite asbestos were higher than those reported for amosite and crocidolite. The weak endothermic peaks at 246 and 509 °C (Fig. 14) should be due to the dehydration and

dehydroxylation of vermiculite [66]. The DTG peaks at 972 and 1004 °C (Fig. 14) are interpreted as the dehydroxylation of talc [38] and biotite [67] respectively, present as impurities. PXRD of anthophyllite asbestos after heating to 1100 °C (Fig. 12) exhibits the lines of enstatite (02-0520 JCPDS card.) and cristobalite (03-0267 JCPDS card).

Fibres of anthophyllite asbestos appear straight, poorly flexible and thin and exhibit a slender needle-like crystal habit both before and after heating treatment (Fig. 13d, 13e). SEM images collected at higher magnification (Fig. 13f) showed that the new phase formed after heating (enstatite) preserves the original fibre morphology (pseudomorphosis) but the surface becomes rough.

Fibrous erionite

The TG curve of Fig. 15 showed a continuous weight loss due to the dehydration of erionite $(H₂O)$ loss of 17.00 wt%), corresponding to the broad endothermic peak at 126 °C [29] and to the weak endothermic peak at 356 °C (Table 3) on the DSC curve. In the first endothermic effect, the water loss is 16.11 wt% while in the second endothermic the water loss is 0.89 wt%. The complete dehydration is attained at 450 °C [68] without loss of crystallinity which started at temperature above 700 °C (as verified by PXRD) followed by recrystallization of K-feldspars and plagioclase as evidenced by the DSC exothermic peak at 911 °C (Fig. 15). The curves are in agreement with those reported in Gottardi and Galli [69] and the estimated temperature of breakdown and recrystallization is comparable to that (840 °C) reported by Ballirano and Cametti [70]. Differences may be related not only to different experimental conditions but also to different Si/Al ratio, ionic potential and size of exchangeable cations [19], and crystallite size of the various samples [70]. The products of erionite recrystallization after heating to 1000 °C were K-feldspar, plagioclase and quartz (see the PXRD in Fig. 16), according to the reaction sequence $Na₅K₃Al₈Si₂₈O₇₂•28H₂O$ (approximated erionite formula) \rightarrow Na₅K₃Al₈Si₂₈O₇₂ \rightarrow 3KAlSi₃O₈ + 5NaAlSi₃O₈ + 4SiO₂.

At the SEM observation, erionite displays bundles composed of many fibres resembling amphibole fibres morphology (stubby prismatic and acicular crystals) (Fig. 17a). After heating treatment the new forms have feldspar-like composition as detected by EDS/SEM analyses. At low magnification SEM, imaging show that the fibres are shorter but the original fibrous morphology is preserved (Fig. 17b) and the surface is smooth. However, at higher magnification (Fig. 17c) the surface of the fibrous crystals show irregularities and appear as rather rough (Fig. 17c).

Conclusions

Thermal decomposition of eight selected mineral fibres (chrysotile, four amphibole asbestos species and erionite) has been investigated using DSC/TG supplemented by PXRD and SEM/EDS. These investigations have shown that the decomposition of UICC chrysotile from Canada occurs at 633 °C while the structure breakdown for both chrysotile from Val Malenco and from Balangero shift versus higher temperatures taking place at 652 °C and 660 °C respectively. The decomposition of chrysotile is followed by the recrystallization into forsterite at about 822 °C. As it concerns amphiboles, the breakdown of crocidolite, amosite, anthophyllite asbestos and tremolite asbestos ranged from 850 to 1046 °C. Crocidolite asbestos UICC is decomposed at around 850 °C with the formation of magnetite, cristobalite and acmite. However, at 1100 °C the final minerals products were mainly hematite and partially melted cristobalite. Amosite UICC decomposes into enstatite and hematite at 878 °C. Anthophyllite asbestos UICC breakdown leads to the formation of enstatite and cristobalite at 861 °C. The breakdown of tremolite asbestos occurs at 1046 °C with the consequent formation of diopside.

While for tremolite and anthophyllite asbestos the complete dehydroxylation occurs in one main effect (1046 °C and 861 °C respectively) and leads to the breakdown of the mineral structure, the dehydroxylation of amosite and crocidolite occurs in various steps in the temperature range 570 – 850 °C and it does not correspond with the structural collapse. Fibrous erionite shows the same

behaviour as that reported for amosite and crocidolite: although the complete dehydroxylation took place between 25 and 425 °C, the loss of its crystallinity starts above 700 °C. The products of the recrystallization after heating to 1000 °C are K-feldspar, albite and quartz.

The thermogravimetric (TG) analysis allowed the calculation of the water content in the fibres which could be useful for the determination of their chemical formulae.

Despite the thermal treatment, all fibrous samples preserve the same external fibrous habit but the structure is completely changed at a molecular scale: this phenomenon called pseudomorphosis lead to the complete transformation of asbestos minerals into non-hazardous silicates such as forsterite and enstatite. However, potentially hazardous minor phases such as cristobalite and quartz were found in the new phases appeared after thermal treatment of anthophyllite asbestos, crocidolite and asbestiform erionite; these may hinder a safe reuse of the processed asbestos samples.

Moreover, it has been demonstrated that DSC and DTG analyses are very effective for the identification of minerals impurities both in chrysotile, amphibole asbestos and asbestiform erionite specimens. Indeed, DSC and DTG analysis clearly showed the presence of low impurities, not relievable by the semi-quantitative PXRD analysis, such as pyroaurite, talc, brucite, smectite, dolomite, siderite, goethite, and biotite. However, all identified minerals are non-fibrous, mostly having platy morphology.

The knowledge of the thermal behaviour of the eight fibrous samples studied in this work will allow interpret thermograms obtained from natural samples and asbestos treated by heating with more confidence.

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Table 1 Calculated chemical formulae (After Pollastri et al., 2015 [38]) and details of the investigated minerals fibres. ^aMixture of fiber from the firms Bells, Carey, Cassair, Flintkote, Johns-Manville, Lake, Normandie and National, proportioned roughly to represent Canadian production of asbestos products at that time. *UICC standard Chrysotile "B" Canadian NB #4173-111-1; §UICC standard Amosite from Penge mine South African; ⁺UICC standard Anthophyllite Finnish NB #4173-111-5; #UICC standard Crocidolite South African NB #4173-111-3

Table 2 Main TG data (weight loss %)

Table 3 Peak temperatures in DSC curves; $w = weak$, $s = strong$, $sh = shoulder$, $endo = endothermic$, $exo = exothermic$

Figure captions

Fig. 1 Thermal analysis of chrysotile UICC from Canada. Solid line: Solid line: DSC. Dashed line: DTG. Dotted line: TG

Fig. 2 PXRD patterns recorded before (b) and after (a) heating at 1000 °C for the three kinds of chrysotile; from the bottom to top chrysotile UICC (CCb and CCa); chrysotile from Balangero (CBb and CBa); chrysotile from Val Malenco (CVMb and CVMa)

Fig. 3 Thermal analysis of chrysotile from Balangero. Solid line: DSC. Dashed line: DTG. Dotted line: TG

Fig. 4 Thermal analysis of chrysotile from Val Malenco. Solid line: DSC. Dashed line: DTG. Dotted line: TG

Fig. 5 Comparison of DSC curves recorded for chrysotile from Balangero (CB), from Val Malenco (CVM) and UICC from Canada (CC)

Fig. 6 Secondary electron SEM images of the three kinds of chrysotile: UICC from Canada, first line; from Balangero, second line; from Val Malenco, third line. From left to right: first column (**a**, **d**, **g**), before heating treatment at 1000 °C; second and third column (**b**, **c**, **e**, **f**, **h**, **i**), after heating. Images **c**, **f**, **i** acquired with high resolution SEM, testifying the pseudomorphic transformation of the chrysotile fibres with temperature

Fig. 7 Thermal analysis of crocidolite UICC. Solid line: DSC. Dashed line: DTG. Dotted line: TG

Fig. 8 PXRD patterns recorded before **b** and after **a** heating at 1000 °C of crocidolite UICC (Crb and Cra); tremolite asbestos (Tb and Ta)

Fig. 9 Secondary electron SEM images of crocidolite UICC (first line) and tremolite asbestos (second line). From left to right: first column (**a**, **d**), before heating treatment at 1000 °C; second and third column (**b**, **e**, **c**, **f**), after heating. Images **c**, **i** acquired with high resolution SEM, showing a complete pseudo-morphic recrystallization occurred to fibres. Arrow in **b** indicates crocidolite fibres partially melted

Fig. 10 Thermal analysis of tremolite asbestos from Val d'Ala. Solid line: DSC. Dashed line: DTG. Dotted line: TG

Fig. 11 Thermal analysis of amosite. Solid line: DSC. Dashed line: DTG. Dotted line: TG

Fig. 12 PXRD patterns recorded before **b** and after **a** heating at 1000 °C of anthophyllite asbestos UICC (Anb and Ana)

Fig.13 Secondary electron SEM images of amosite (first line) and anthophyllite asbestos UICC (second line). From left to right: first column (**a**, **d**), before heating treatment at 1000 °C; second and third column (**b**, **e**, **c**, **f**), after heating. High resolution SEM images **c** and **i** show the fibrous recrystallized forms

Fig. 14 Thermal analysis of anthophyllite asbestos UICC. Solid line: DSC. Dashed line: DTG. Dotted line: TG

Fig. 15 Thermal analysis of asbestiform erionite. Solid line: DSC. Dashed line: DTG. Dotted line: TG

Fig. 16 PXRD patterns from asbestiform erionite before (Erb) and after (Era) heating at 1000 °C

Fig.17 Secondary electron SEM images of asbestiform erionite **a** before and **b**, **c** after; heating treatment at 1000 High resolution SEM Image **c** shows rough surface

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