Morphological and chemical analysis of tremolite related to natural asbestos in the road paving

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Tremolite group minerals could be present in the aggregate used as filler in the mix asphalt floor in the road surfaces and are subject to wear for vehicular traffic and materials ageing. Dust produced by the degradation of asphalt surfaces can disperse tremolite group minerals present in road paving causing exposure by asbestos fiber to an increasing number of people and animals. In recent years, attention has been focused on naturally occurring asbestos (NOA) as the greatest frequency of lung cancer has been observed in environments characterized by rocks and soils rich in Tremolite that characterize asbestos minerals. Asphalt samples object of this study were collected in a motorway segment of the Lombardia Region (north of Italy) for an exposure assessment research on road paving workers. The evaluation was performed using a metrological approach based on petrographic and mineralogical analyses in order to reduce the epidemiologic risks and to define the best treatment strategies on waste management with sustainable costs in fulfillment of Council Directive 67/548/EEC (1967) and EC Regulation 1272/2008 (2008) (Paglietti et al., 2016). The fibers are generally not broken down to other compounds in the environment, but asbestos hazard could be related to the occurrence of tremolite fiber in road paving arises when natural weathering processes (e.g., erosion) or human activities (e.g., continuous passage of motorized vehicles) separate and break down the fibers, which could be lost into the environment as airborne and easily penetrate into the human respiratory tract (Punturo et al., 2019). Despite this, this work is not an epidemiological risk analysis but obtained data could be useful to other specialists to define the exposure risks.

Introduction

Asphalt is a natural mixture of solid and semifluid hydrocarbons, mostly consisting of bitumen; solid and dark in color, it is found as an impregnating agent for calcareous rocks (Greene, 1992).

Asphalt concrete (AC) is a composite material consisting of aggregate, asphalt binder (mastic), and air void. AC has been primarily used as a material in constructing road and airport pavements. (Xu, 2010; Romera et al., 2016) and in some cases it is mixed to fibers fillers of reinforcement (Xiong et al., 2015; Topini et al., 2018). The effectiveness of bituminous concrete is strictly influenced by the combination of base binder, additive type and dosage (Moreno-Navarro et al., 2017). However, under the effects of repeated vehicle loading at high temperature, moisture cycling, and low-temperature contraction, AC mixture is susceptible to distresses of rutting (permanent deformation), stripping (separation of asphalts from aggregates), and cracking, etc. Loss of performance of road paving is also due to climatic factors, i.e. temperature range (Bonica et al., 2016), moisture (Androjić et al., 2017). However, under the effects of repeated vehicle loading at high temperature, moisture cycling, and low-temperature contraction, AC mixture is susceptible to distresses of rutting (permanent deformation), stripping (separation of asphalts from aggregates), and cracking, etc. Loss of performance of road paving is also due to climatic factors, i.e. temperature range (Bonica et al., 2016), moisture (Androjić et al., 2017), solar radiation, frequency and intensity of the rain and snow, frost, salt, etc.; and to anthropogenic impact as traffic and number of heavy vehicles traveling on the road, loads, speed, etc. (Qian et al., 2019; Assogba et al., 2020).

To guarantee safe circulation of road users, maintenance of road pavements with new asphalt overlays is required. Unfortunately, different studies to measure the potential environmental impacts of the asphalt mixtures life cycle rarely take into account the presence of asbestos for their reuse in the reconstruction of the road paving (Chen et al., 2009; Xu et al., 2010). From the 1950s for about thirty years, some road paving in areas subject to severe deterioration (Du et al., 2019) and wear have been built with hot mix asphalt (HMA) (Landi et al., 2020) reinforced to asbestos in order to improve resistance and to reduce action of thermoclastism and cryoclastism (Slebi-Acevedo et al., 2019). It is therefore necessary to identify the presence of asbestos to avoid the risk of dispersion into the atmosphere of dangerous solid particles which can be inhaled by human and especially by road maintenance personnel in order to prevent exposure of the population to lung cancer caused by asbestos (Noonan and Pfau, 2010; Bourgault et al., 2014; Pfau and Noonan, 2019). The presence of asbestos in air and water has become a subject of health and environment concern (Mokhtari et al., 2019; Sąkol and Muszyńska-Graca, 2019; Taghizadeh et al., 2019). While the hazards of prolonged asbestos inhalation have been well established (Cox Jr, 2019; Nishimura et al., 2020), there does not yet appear to be
clear answer concerning the toxicity of ingested asbestos (Fitzgerald and Rhodes, 2019; Zheng et al., 2019).

The number of workers exposed to asbestos since 1950 has greatly increased (U.S Department of Health and Human Services, 2001) caused to building construction and demolition and the deterioration and wearing of asbestos-containing materials. Worker exposure is a concern in the mining and milling of asbestos, during the manufacture of asbestos product (Heidrich et al., 2019; Kromhout et al., 2019).

This paper describes a method for the asphalt characterization. This study was requested by a private company from Ferrara city (North East of Italy) for the characterization of asphalts used in the road paving. The surfaces of each samples were analyzed using a portable Energy Dispersive X-Ray Fluorescence (ED-XRF). Fragment of each samples were prepared in order to perform X-ray diffractometric powder analysis (XRPD) and observation by scanning electron microscopy analysis (SEM-EDS). During XRPD and SEM-EDS analyzes, presence of asbestos fibers was unexpectedly observed in some samples.

Materials and methods

Sampling

Three samples were collected in the North of Italy in a motorway section in the Lombardia region, with a wet core drill of 5 cm cylindrical section (Fig. 1). The samples were stored individually in a double envelope to prevent contamination.

Petrographic study on the surfaces of the samples was conducted using a portable X-ray fluorescence (ED-XRF). No thin sections were made to avoid risks of exposure to hypothetical fiber particles. The samples were analyzed by ED-XRF as it is without any preparation or manipulation.

Fragments of degradation asphalt paving were crushed in water, sieved and collected on filter. Two different particle size fraction were obtained by sieving: samples pulverized granulate with a diameter between 1 mm and 0.063 mm; samples granulate crushed with a diameter less than 0.063 mm. All these samples were filtered and dried in stove in glass containers in order to perform X-Ray Diffractometric Powder analysis (XRPD) and observation by scanning electron microscopy analysis (SEM-EDS). For SEM-EDS analysis the samples were not metallized.

Chemical and Mineralogical Characterization

Chemical elements in asphalt samples were determined by ED-XRF, using a portable µ-XRF spectrometer ARTAX™ 200 (Bruker AXS Microanalysis GmbH, Germany) for the spectroscopic analysis. The instrument consisted of an X-ray tube with a Mo target and an SSD Peltier-cooled detector (10 mm² active area and resolution of <155 eV at 10 kcps). The system performs a simultaneous multi-ele-

Figure 1. Photo images that allow to realize the macroscopic characterization of the three asphalt samples: a) b) c) sample 1; d) e) f) sample 2; g) h) i) sample 3; a) d) g) section of samples; b) e) h) surface of samples; c) f) i) lateral part of each samples.
ment analysis in the element range from Na (11) to U (92). The maximum voltage and current of 50 kV and 1500 µA, respectively, were used to excite the secondary fluorescence X-rays. A collimator with a diameter of 1 mm was used to collect the emitted secondary X-rays from a surface area of about 0.79 mm² in air. The measurement is non-destructive. The major elements determined as oxides were SiO₂, TiO₂, Al₂O₃, Fe₂O₃, MnO, MgO, CaO, K₂O and the trace elements were Zn, Cr, Cu, Sr and S. The accuracy and precision of data were evaluated using certified international standards (JSd-1 and JSd-2 stream sediment powder, Geological Survey of Japan, Tsukuba, Japan). Accuracy was generally lower than 2% for major oxides and less than 5% for trace elements: the detection limits for major oxides were 0.03–0.05% and for trace elements 1-2 ppm. The qualitative data obtained were expressed as single element weight.

Moreover, mineralogical determinations of some aggregates, previously grinded using water and then dried, were carried out through X-ray diffractometric powder analysis (XRPD) using a Philips PW1860/00 diffractometer, with a graphite filtered and a Cu Kα radiation (1.54 Å) in a 2θ angular range 5-75°, with a 5 s/step (0.02° 2θ) (Marrocchino, et al., 2010).

The morphological and chemical characterization was carried by a Scanning Electron Microscope (SEM) Zeiss EVO MA15 (Carl Zeiss AG, Oberkochen, Germany) equipped with an Energy Dispersive X-ray Spectrometer (EDS) INCA 300 (Oxford Instruments, Abingdon, UK) for X-ray microanalysis. The SEM-EDS high magnification images of the sample surfaces were performed using the SmartSEM software (Zeiss) (Telloli, et al., 2018). For SEM examinations, a piece of each sample was fixed on an SEM stub utilizing double-sided conductive adhesive tape.

Results

Macroscopic Characterization

All samples can be classified as bituminous conglomerates characterized by good quality aggregate of limestone breccia with good physical-mechanical resistance incorporated in a bituminous matrix (Fig. 1).

In all analyzed samples, the aggregate was constituted of limestone granules characterized by sharp-edged fragments with jagged contours and it was well graded, with a prevalent fraction with a diameter major than 4 mm. These features led to hypothesize that the aggregates used was probably obtained by crushing and sieving limestone rocks, or they could come from gravel quarries.

The aggregates were predominantly composed by microcrystalline rocks with carbonate composition and were submerged in an asphalt matrix mixed with an asbestos filler (not exceed 5%, observed using stereomicroscope). In the samples 1 and 2 grains orientation was almost absent, the aggregates were well graded and well mixed with matrix. Sample 3, instead, was less well graded than samples 1 and 2, with fractions with different dimensions. In all samples, the grains were well submerged in the matrix; this implies that granules do not have contact each other, being evenly coated with bitumen. They were characterized by a medium degree of thickening: low macro-porosity in samples 2 and 3, while in sample 1 several macro vacuoles were present in the bituminous fraction. No ravelling phenomena or other forms of degradation were observed, testifying optimal aggregate bitumen bonds. The surface of the bituminous layer was rough in sample 1, while sample 2 had a surface without roughness and intermediate roughness characterized sample 3. All samples seem to have a positive correlation between macro porosity and surface roughness. Sample 3 was characterized by a layer of dust due to accumulation phenomena, conversely in sample 2 was not evident any accumulation of dust, but a biological attack with abundant development of fungal hyphae.

ED-XRF Data Analysis

Clasts and bituminous fraction had been analyzed for each samples. Sample 1 was characterized by different particle size, as shown in figure 2a. The nature of the clasts was mono-compositional consisting exclusively of calcium carbonate. Iron and strontium were also present (Fig. 2a). In other analyzed clasts, Zn, Ti, V, Mn and S were also reported. The bituminous fraction (Fig. 2b) was characterized not only by Ca element, that could be related to the presence of carbonate mineral fragments, but also high peaks of Fe and Cr and alum potassium silicates, represented by high peaks of Si and K. Finally, presence of small peaks of Ni, Ti, Zn and Sr elements were also detected, as shown in the graph of figure 2b. These elements could be related to vehicular traffic pollution that could be adsorbed by the fracture in the asphalt.

Figure 2c shown the ED-XRF spectrum of clasts in sample 2, that were characterized by calcium carbonate. Small peaks of Fe, Mg, Si and K elements were also present, as shown in the graph of figure 2c. High peak of Ca reflected the low thicknesses of the bituminous layer therefore the presence of the carbonate aggregate emerged compositionally. Abundant in the bituminous fraction were S and Fe; Ni, Cr, Cu, Zn and Sr elements were also detected (Fig. 2d) probably related to vehicular traffic pollution that could be adsorbed by the fracture in the asphalt as in the sample 1.

Sample 3 was characterized by high concentration of calcium in the clasts analyzed, probably due to the presence of calcium carbonate in the majority fraction of the sample (Fig. 2e). The bituminous fraction appeared wrinkled and thin in correspondence with the clasts this could cause a partially transparent to X-rays from which an important peak of calcium related to the presence of carbonate clast. Compared to sample 1 and 2, sample 3 shown the bituminous fraction with a lower peak of transition metals, especially for iron and sulphur. Low concentrations of Sr and K were also detected (Fig. 2f).

XRPD Data Analysis

Samples pulverized granulate using water and then dried with a diameter between 1 mm and 0.063 mm and samples granulate crushed with a diameter less than 0.063 mm were analyzed by XRPD. The semi-quantitative data obtained were processed and the average of the results were inserted in a single graph shown in figure 3.

Figure 3a shows the analyses of the fraction with diameter less than 0.063 mm. Sample 1 and sample 3 had similar pattern characterized by a high mineral abundance of carbonate clasts represented by high peak of calcium and magnesium. The presence of magnesium was probably due to the presence of dolomite clasts.

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In addition, small peaks of tremolite species and the absence of chrysotile. On the other side, sample 2 shown the same abundant of carbonate clasts represented by high peak of calcium and magnesium as in the other two samples, peaks of tremolite species but also a small abundant of chrysotile, that could be related with asbestos.

For this reason, sample 2 has been much more wet grinded and the fraction with diameter between 1 and 0.063 mm was analyzed. Figure 3b shows the results: confirming the presence of chrysotile and tremolite species.

**SEM-EDS Data Analysis**

Morphological and microstructural characterization were performed using SEM analysis in order to detect the morphology of the particles, recognize the presence of fiber phases (Tremolite asbestos), identify the fibers dimension (diameter and length) and any degree of aggregation (presence of growths, micro fractures, etc.). The morphological analysis was conducted by microanalysis aimed to identify the chemical and mineralogical composition of the observed granules.

Two different particle size fraction were obtained by sieving and analyzed by SEM-EDS: samples pulverized granulate with a diameter between 1 mm and 0.063 mm and samples granulate crushed with a diameter less than 0.063 mm.

**Size fraction < 0.063 mm**

Table 1 shows the average of the data obtained by SEM-EDS analysis of the three samples collected and expressed in oxide weight (%).
For each samples, superficial layer called S and internal layer called I were analyzed.

Both in the internal and external layers of sample 1 the results show an absence of asbestos particles. The matrix was granular with rounded particles (as shown in Fig. 4a and 4b) and the chemical composition not highlight high Mg concentration that could be typical of the asbestosis phases. Mg was low abundant and compatible with the clay composition of the matrix or associated with the presence of dolomite clast. For this reason, it was not possible to define if Mg concentration was due to the presence of dolomite or to asbestos particles.

Figure 4c shows particles in sample 2S, that were higher than particles in sample 1S, so the sample 2S was less powdery. Also in sample 2 the particles were generally rounded. High Ca and Mg concentrations could be related to the presence of calcite and dolomite; the percentage of silicates was low being the SiO₂ very low. On the opposite, sample 2I results (Table 1) show the presence of asbestos particles: Si-Mg ratio was consistent with the presence of chrysotile (Gronow, J., 1987; Kimizuka et al., 1987; Falini et al., 2002).

The particles in the sample 3S were mainly massive (Fig. 4e). A few slightly fiber phases are observed, but due to the presence of
dolomite there is a high concentration of Mg which does not allow to identify its nature. Table 1 and figure 4f show the total absence of asbestos particles in sample 3I.

**Size fraction between 1 mm and 0.063 mm**

Only in sample 2, SEM-EDS analyzes show the presence of asbestos particles in the form of chrysotile (Table 2 and Fig. 5).

Table 2 shows the high concentration of Si and the low concentration of Ca respect to Table 1, that could confirm the presence of the fiber.

SEM observations highlighted that tremolite asbestos and actinolite appear straight and show a slender needle-like crystal habit with a length longer than 10 μm. It is worth remembering that fibers are composed of many fibrils, which tend to split up along the fiber elongation axis (Belluso et al., 2017).

**Discussion and Conclusion**

Morphological and chemical characterization of asphalt samples
for road paving have shown the presence of asbestos fibers especially in the size fraction with diameter between 1 mm and 0.063 mm. The results of the analysis in the fine fraction with diameter less than 0.063 mm show the total absence of asbestos, probably too small and

Table 1. SEM-EDS average data analysis of the three samples analyzed expressed in oxide weight (%): “S” means superficial layer and “I” means internal layer

| Element | 1S   | 1I   | 2S   | 2I   | 3S   | 3I   |
|---------|------|------|------|------|------|------|
| C       | 11.58| 7.38 | 13.61| 17.41| 12.51| 8.37 |
| O       | 45.96| 51.54| 55.24| 55.24| 44.38| 50.72|
| Na      | 0.74 | 0.28 | 0.00 | 0.45 | 0.26 | 0.11 |
| Mg      | 2.27 | 6.11 | 7.64 | 5.45 | 2.24 | 6.13 |
| Al      | 6.26 | 2.43 | 0.45 | 1.22 | 7.18 | 3.58 |
| Si      | 12.27| 13.40| 4.32 | 2.53 | 13.98| 14.13|
| S       | 0.00 | 0.00 | 0.16 | 0.29 | 0.00 | 0.00 |
| Cl      | 0.00 | 0.00 | 0.00 | 0.32 | 0.00 | 0.00 |
| K       | 0.67 | 0.14 | 0.15 | 0.25 | 0.85 | 0.46 |
| Ca      | 18.60| 14.95| 13.81| 15.93| 17.02| 13.72|
| Ti      | 0.00 | 1.01 | 0.00 | 0.00 | 0.00 | 0.98 |
| Mn      | 0.00 | 0.00 | 0.40 | 0.00 | 0.00 | 0.00 |
| Fe      | 1.65 | 2.77 | 4.20 | 0.92 | 1.58 | 1.80 |
| Totals  | 100.00| 100.00| 100.00| 100.00| 100.00| 100.00|

Figure 3. XRPD spectrum: a) spectrum of the fraction with diameter less than 0.063 mm of all the three samples analyzed (sample 1 in blue, sample 2 in green and sample 3 in red); b) spectrum of the fraction with diameter between 1 and 0.063 mm of sample 2.
not detected by the instrument. Consequently, the fibers are concentrated in the coarse fraction (diameter between 1 mm - 0.063 mm). This is confirmed by XRPD and SEM-EDS analysis.

ED-XRF analysis show high presence of Fe and S and other metals as Ni, Cr, Cu and Zn in the bituminous fraction. These elements could be related to vehicular traffic pollution that could be adsorbed by the fracture in the asphalt. The bituminous composition could be related to a typical bitumen of Mexican Blend or California type (Riala, 1998). Instead of, the majority of the clasts are characterized by calcium carbonate and also dolomite. No asbestos had been observed.

XRPD results confirm that clasts in all the analyzed samples are constituted by the prevalence of calcite and dolomite, with also some quartz. Small amount of tremolite species are present in the three analyzed samples, while small peaks of chrysotile appears in sample 2.

SEM-EDS imaging and data confirm the presence of asbestos in sample 2, especially in the fraction with diameter between 1 mm and 0.063 mm.

As suggested from Militello et al. (2019), the observations under OM allowed us to investigate the textures and the spatial relations between the mineralogical phases and the identification of asbestos minerals could be easy under OM, but even at higher magnifications a fiber habit has the same geometric parameters of prismatic/acicular crystals. Under SEM it is better characterize the aspect ratios and the textural relations between the various phases-often the habit is detectable, but discrimination among the serpentine polymorphs is not simple and not always possible. For these reasons, could be plausible implement the analysis with TEM observations, that allow high-resolution discrimination for EMP and an effective distinction between regulated and non-regulated asbestos minerals. Moreover, for better characterize samples and mineralogical phases with more attention to fiber phases, synchrotron radiation X-ray micro-tomography could be a technique that offers the possibility to observe and image the 3D morphology and the spatial relationship between the mineral phases that constituted the investigated samples. Furthermore, it is a semi-destructive technique which allows the analysis of samples under high magnification without grinding/milling and or the loss of their morphology. Nevertheless, it provides qualitative and volumetric but not quantitative information of the mineralogical phases; therefore, it proved a complementary technique to other conventional ones.

In general, asbestos fibers in asbestos-containing roofing materials are encapsulated in bituminous matrices or resins, thereby binding the fibers and preventing their release (NRCA, 1994). During their work, many roofers may have been exposed to asbestos during the repair or tear-off of old roofing materials, including roofing cements and coatings. Very few studies have reported airborne concentrations of asbestos associated with the application, repair or removal of roof coatings or cements.
Mowat (2007) indicates a low probability that asbestos will become airborne during manipulation of coating- and cement-type roofing products such as those tested. The results indicate an exceedingly small asbestos exposure associated with the repair of these products, even during the unlikely 30-min application, hand scraping and sanding of these products from roof substrates.

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