MINERAL CHARACTERIZATION OF SOIL TYPE RANKER FORMED ON SERPENTINES OCCURRING IN SOUTHERN BELGRADE ENVIRONS BUBANJ POTOK

by

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The paper addresses the issue of health risk associated with the presence of chrysotile in the soil type ranker formed on massive serpentines occurring in the area of Bubanj Potok, a settlement located in the southern Belgrade environs, Serbia. Characterization of the ranker soil was conducted by scanning electron microscopy, X-ray diffraction, micro-Raman spectroscopy and transmission ⁵⁷Fe Mössbauer spectroscopy. Scanning electron microscopy figures showed regular shaped smectite (montmorillonite) particles, aggregates of chlorite, and elongated sheets of serpentines minerals antigorite. X-ray diffraction analysis confirmed the presence of detrital mineral quartz polymorph as well as minor amounts of other mineral species. Micro-Raman spectroscopy identified the presence of dominant minerals, such as montmorillonite, kaolinite, muscovite, gypsum, calcite, albite, amphiboles (hornblende/kaersutite) and orthoclase. Important polymorph silica modifications of quartz, olivine (forsterite), pyroxene (enstatite/ferrosilite, diopside/hedenbergite), and serpentine (antigorite/lizardite/chrysotile) were identified.

Key words: ranker soil, serpentine, chrysotile, Mössbauer spectroscopy, micro-Raman spectroscopy

INTRODUCTION

One of the most widely distributed groups of siliceous rocks in the hilly and mountainous areas of Serbia consists of serpentinised peridotite rocks. The qualitative and quantitative determination of various serpentine minerals present within massive serpentinites is rather difficult because of their similar structures and chemical compositions [1]. Besides the geodynamical and petrological significance of the precise identification of these minerals, there is a health risk issue associated with the presence of chrysotile or heavy metals. In order to identify the three principal mineral polymorphs of the serpentine group, i. e., chrysotile, lizardite and antigorite, micro-Raman and 57Fe Mössbauer spectroscopy were applied in the analysis of the massive serpentines occuring in the Bubanj Potok area, located near Belgrade, Serbia.

Micro-Raman spectroscopy is a non-destructive, rapid technique with no sample preparation, and has the advantage of high spectral resolution and easy data collection in identification of minerals and other species [2, 3]. ⁵⁷Fe Mössbauer spectroscopy has proved to be a valuable technique in providing information about the characterization of iron minerals in soil and clays [4]. ⁵⁷Fe Mössbauer characterization results in the detection of the iron site geometry, valence state, and co-ordination number, and also provides a quantitative estimate of the iron content of unknown species in a sample. Measured quantities such as the isomer shift (δ), the electric quadrupole splitting (QS) and the hyperfine magnetic field (B), can be used to distinguish between oxidation states of iron in various structural sites and to detect iron oxides of different particle sizes in the investigated sample. The relative amount of each structural state or phase can be obtained with this technique [5, 6].

MATERIALS AND METHODS

The studied site is located in the vicinity of Belgrade in Serbia (20°33'36" Eastern longitude, 44°43'30" Northern latitude, figs. 1 and 2). The soil type ranker formed on serpentinite was investigated. Soil samples were collected from nine small-depth profiles, and a single composite sample of A_h soil horizon was formed.

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Figure 1. The Bubanj Potok location chart



Figure 2. Terrain model – Topographic map with the locations of nine soil profiles of the studied Bubanj Potok soil, ranker type

Scanning electron microscopy (SEM) was done by JEOL Model JSM–6390LV at an accelerated voltage of 15 kV and 20 kV. The specimens were coated with a thin layer of gold (Au) using a Bal-Tec SCD005 Sputter Coater.

A powder X-ray diffraction (XRD) pattern was obtained on the Philips PWW1710 powder diffractometer with monochromatic (graphite monochromator) CuK $\alpha_{1,2}$ radiation (= 1.54178 Å*). The working voltage and current in the course of data collection were 40 kV and 30 mA, inside 4-70° 2θ range in steps 0.03°, and scanning time of 2s per step. The identification of minerals in total soil and clay fraction of soil was made using the computer program package DRX Win 1.4c. The Raman scattering measurements were performed using TriVista 557 Raman system in backscattering micro-Raman configuration. The system was in subtractive mode with 900 g/mm, 900 g/mm, and 2400 g/mm gratings and 200 m slits. The 514.5 nm line of an Ar^+/Kr^+ mixed gas laser was used as an excitation source. The laser beam focusing was achieved by a long distance microscope objective (magnification 50 times). Laser power on the sample was about 1 mW.

⁵⁷Fe Mössbauer absorption spectra were obtained in a standard transmission geometry using a source of ⁵⁷Co in Rh (920 MBq) at room temperature. The spectrometer was operated with a triangular velocity waveform in constant acceleration mode. The measurements were made on a powder sample contained in a plexiglas holder; the surface density of the absorber was 16.7 mg/cm². The data were stored in 1024 multichannel analyzer. The laser spectrum was recorded and fitted in order to recalculate channels in mm/s. The sample thickness corrections were carried out by transmission integral. All quoted isomer shifts (δ) are referred to natural alpha iron at ambient temperature. The spectra were fitted using the WinNormos program created by Brand (2008), using a least squares method [7].

RESULTS AND DISCUSSION

SEM investigation of the Bubanj Potok soil clearly showed the regular-shaped smectite (montmorillonite) particles and aggregates of chlorite, well developed, with different grain sizes (figs. 3 and 4). These aggregates have small deformations on the borders of grains and do not have regular hexagon surfaces. Elongated sheets of lizardite mixed with antigorite were also observed (fig. 4).

Results of semiquantitative XRD analysis of the investigated ranker soil provided limited information on the presence of the following crystal phases: the domi-



Figure 3. SEM micrograph at magnification 700 of the Bubanj Potok soil sample. Larger single particles indicate smectite phase. Smaller aggregates of particles represent chlorite phase. Marker 20 μ m



Figure 4. SEM micrograph at magnification 1800 of the Bubanj Potok soil sample. Clay fraction with elongated sheets of lizardite mixed with antigorite. Marker 10 μ m

nant detrital mineral quartz polymorph 41%, was identified, along with numerous fractions of minerals in the soil and five oxides up to 59% (fig. 5). The following minor fractions of minerals were present in the soil: interstratifications of chlorite-vermikulite-montmorillonite, chlorite-vermikulite, amphibole (hornblende), talc, muskovite, serpentine-lizardite, quartz polymorph, orthoclase, kaolinite, pyroxenes (enstatite, ferrosilite, diopside and hedenbergite), albite, haloysite and montmo-



Figure 5. XRD pattern of the Bubanj Potok ranker soil. Present mechanical fractions of minerals in the soil: interstratifications of chlorite-vermikulite-montmorillonite (Ch-V-M), chlorite-vermikulite (Ch-V), amphibole (hornblende) (Af), talc (Tc), muskovite (Mus), serpentine-lizardite (Liz), quartz polymorph (Q), orthoclase (Or), and kaolinite (K); pyroxene (enstatite, ferrosilite, diopside, and hedenbergite) (Es), albite (Ab), haloysite (Ha), and montmorillonite (M)

rillonite. To assess the productive capacity of the studied ranker soil, the percentage of the five oxides containing Si, Al, Fe, Ca, and Mg were determined by atomic absorption spectrometer (AAS-4000 Perkin-Elmer). Active acidity of the soil, pH in H_2O (1:2.5), was obtained potentiometrically with a pH meter (tab. 1).

Raman spectrum of the Bubanj Potok soil is presented in fig. 6. Raman measurements were done in the range 200-1060 cm⁻¹. Extraction of the data has been performed using a Lorentzian lineshape. The most intense peaks are centered at 465 cm⁻¹ (P8), 602 cm⁻¹ (P10), and 875 cm⁻¹ (P16), whereas groups of less intense peaks are found in the ranges 208-440 cm⁻¹, 520-680 cm⁻¹, and 760-970 cm⁻¹.

The SiO₂ quartz polymorph is characterized by peaks at 354 cm⁻¹ (P6), 428 cm⁻¹ (P7), 465 cm⁻¹ (P8), 813 cm⁻¹ (P13), and 836 cm⁻¹ (P14), [8-10].

The smectite montmorillonite, was characterized with following peaks: 257 cm^{-1} (P2), 295 cm^{-1} (P3), 875 cm^{-1} (P16), and 916 cm⁻¹ (P18), [11]. Raman modes of the end-member Mg-Fe-Ca pyroxenes consist of enstatite: 543 cm^{-1} (P9) and 857 cm^{-1} (P15), ferrosilite: 654 cm^{-1} (P12), diopside: 255 cm^{-1} and hed-enbergite: 234 cm^{-1} , [12, 13]. Other minerals present are gypsum (CaSO₄ 2H₂O): 616 cm⁻¹ [14], calcite (CaCO₃): 890 cm⁻¹ (P17) [14], kaolinite: 335 cm^{-1} (P5), and 918 cm⁻¹ (P18) [15, 16], albite: 335 cm^{-1} (P5) and 815 cm^{-1} (P13) [14, 17], mixture of pyroxene and forsterite: 602 cm^{-1} (P10) [14], muscovite: 316 cm^{-1} (P4), [18] silicate amphibole (hornblende): 917 cm⁻¹ (P18) [19] and orthoclase: 813 cm^{-1} (P13) [17].



Figure 6. Raman spectrum of the Bubanj Potok soil ranker type. The labeled peaks are explained in the main text

Table 1. Average properties of investigated Bubanj Potok soil, type ranker: chemical composition [%], heat loss [%], and active acidity of the soil, pH in H_2O

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Horizon depth [cm]	SiO ₂ [%]	Al ₂ O ₃ [%]	Fe ₂ O ₃ [%]	CaO [%]	MgO [%]	Loss in heating [%]	pH in H ₂ O
$A_{\rm h} (0-20)$	46.00(4.33)	5.67(1.61)	20.60(4.39)	0.84(0.25)	7.80(1.61)	20.20(2.89)	6.27(0.06)
(Range)	(41.79-52.21)	(4.12-8.29)	(16.5-27.2)	(0.48-1.11)	(5.98-10.1)	(15.4-24.1)	(6.2-6.36)

The assignment of the Raman peak at 234 cm⁻¹ (P1) needs a more careful consideration. The three principal mineral polymorphs of the serpentine group are chrysotile, lizardite and antigorite [20]. Previous experimental results determining the Raman peak within the range of 230-236 cm⁻¹ (P1) suggest that these three serpentine minerals are present [2, 14, 21].

The presence of health-risk chrysotile was investigated by taking the ⁵⁷Fe Mössbauer absorption spectrum at 294 K of the Bubanj Potok soil (fig. 7). The spectrum consists of one Zeeman sextet and four paramagnetic doublets. The sextet corresponds to the γ -Fe₂O₃, maghemite, in which Fe³⁺ is in octahedral position.



Figure 7. ⁵⁷Fe Mössbauer absorption spectrum at 294 K of the Bubanj Potok soil ranker type

For the investigated sample, 57 Fe Mössbauer data indicate the presence of octahedral Fe²⁺, octahedral Fe³⁺ and tetrahedral Fe³⁺ (11.2%, 47%, and

14.9% of total iron, respectively) in the present mixed serpentine phases and the presence of octahedral Fe^{3+} (22.2%) in maghemite. The fourth unresolved doublet $-FWHM = 0.6(3) \text{ mms}^{-1} - \text{ is presented with } 4.7\%$. The relative contents of the iron containing components were derived from the intensity of the corresponding spectral components omitting the possible influence of the Lamb-Mössbauer factor.

The magnetic phase is observed only in maghemite (tab. 2). The magnetic microstructure of maghemite (γ -Fe₂O₃) is characterized with isomer shift of 0.33(2) mms⁻¹, the magnitude of 50.1(2) T and negligible quadrupole shift that are in excellent agreement with previously reported data [22, 23]. The three named doublets have origine in serpentine phases [24, 20]. Their Mössbauer parameters are under the influence of presented pyroxenes and amphiboles which are much higher in Fe. The fourth unresolved doublet, characterised with $\delta = 0.8 \text{ mms}^{-1}$, and QS = 1.3 mms⁻¹, appeared as the consequence of undistinguished octahedral positions of Fe²⁺ and Fe³⁺ in pyroxene hedenbergite and amphibole kaersutite [25].

The results of the least squares fit of microcomposites are summarized in tab. 2; the values of isomer shift (δ), quadrupole splitting ($\Delta = eQV_{ZZ}/2$), hyperfine internal magnetic field (B_{hf}) and area of each component, are reported.

As we have previously shown, the analysis of Bubanj potok soil revealed five oxides with Si, Al, Fe, Ca, and Mg and 18 minerals: quartz, gypsum, calcite, agregates of chlorite, antigorite, lizardite, montmorillonite, muscovite, kaolinite, albite, forsterite, amphibole (hornblende), orthoclase, four modes of pyroxene (enstatite, ferrosilite, diopside, hedenbergite), including maghemite from analysis of Mössbauer data. The present study indicates that the Bubanj Potok soil is very enriched in silica, with a rough estimation of 41%. The presence of health risk related to chrysotile mineral in the Bubanj Potok soil ranker type, was successfuly excluded during the Mössbauer analysis.

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Table 2. Selected fit of the ⁵⁷Fe-Mössbauer parameters for naturally-occuring Fe²⁺ and Fe²⁺ oxides in the Bubanj Potok soil. Site occupancies are marked with (T) for tetrahedral site and (M) for octahedral site

Compound		[mms ⁻¹]	QS [mms ⁻¹]	B (T)	FWHM [mms ⁻¹]	A^*
Serpentines -Fe ₂ O ₃	M T M	1.134 0.30 0.02 0.355 0.007 0.33 0.02	2.68 0.01 0.27 0.03 0.66 0.02 0.06 0.05	50.1 0.2	$\begin{array}{cccc} 0.24 & 0.02 \\ 0.28 & 0.07 \\ 0.40 & 0.03 \\ 0.58 & 0.08 \end{array}$	0.112 0.013 0.149 0.052 0.470 0.076 0.222 0.046
Kaersutite/Hedenbergite	М	0.8 0.2	1.3 0.3		0.6 0.3	0.047 0.022

* Relative ration of areas M – octahedral site

T – tetrahedral site

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МИНЕРАЛНА КАРАКТЕРИЗАЦИЈА РАНКЕРА ФОРМИРАНОГ НА СЕРПЕНТИНИТУ ИЗ ОБЛАСТИ БУБАЊ ПОТОКА ЈУЖНО ОД БЕОГРАДА

Рад се бави питањем здравственог ризика повезаног са присуством хризотила у ранкеру (тип земљишта) формираном на масивном серпентиниту у области Бубањ Потока, насељу које се налази у јужној околини Београда. Карактеризација ранкера је извршена скенирајућом електронском микроскопијом, рендгенском дифракцијом праха, микро-Раман спектроскопијом и трансмисионом ⁵⁷Fe Месбауер спектроскопијом. Слике добијене скенирајућим електронским микроскопом показују честице смектита регуларног облика монтморионита, агрегата хлорита и издужене листове серпентинског минерала антигорита. Рендгенска дифракциона анализа потврдила је присуство детритал минерала кварцног полиморфа као и мање количине других минералних врста. Микро-Раман спектроскопија идентификовала је присуство доминантних минерала, као што су монтморионит и каолинит, мусковит, гипс, калцит, албит, амфибол (каерсутит/хорнбленда) и ортоклас. Важни полиморфи силика модификације кварца, оливина (форстерит), пироксена (енстатит, феросилит, хеденбергит, диопсид), и серпентина (антигорит, лизардит, хризотил) су били идентификовани.

Кључне речи: ранкер, серџеншиниш, хризошил, Месбауерова сџекшроскоџија, микро-Раман сџекшроскоџија