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Mineralogical and Geochemical Characterization of Asbestiform Todorokite, Birnessite, and Ranciëite, and Their host Mn-Rich Deposits from Serra D’Aiello (Southern Italy)

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Abstract: Manganese ores, especially the oxyhydroxides in their different forms, are the dominant Mn-bearing minerals that occur in marine and terrestrial environments, where they are typically found as poorly crystalline and intermixed phases. Mn oxyhydroxides have a huge range of industrial applications and are able to exert a strong control on the mobility of trace metals. This paper reports the results of a detailed study on the Mn oxyhydroxides occurring in the manganiferous deposit outcropping in the Messinian sediments from Serra D’Aiello (Southern Italy). Nine Mn samples were characterized in detail using X-ray powder diffraction (XRPD), differential scanning calorimetry (DSC), thermogravimetry (TG), transmission electron microscopy combined with energy dispersive spectrometry (TEM/EDS), and X-ray fluorescence (XRF). The results indicated that the Mn deposit included the oxyhydroxide mineral species birnessite, todorokite, and ranciëite. The size, morphology, and chemical composition of Mn oxyhydroxide samples were investigated in order to define their impact on the environment and human health. Todorokite displayed asbestiform shapes and could disperse fibers of breathable size in the air. Furthermore, since in-depth characterization of minerals within Mn deposits may be the first step toward understanding the genetic processes of manganese deposits, hypotheses about the genesis of the Mn oxyhydroxide deposits were discussed.

Keywords: Mn oxyhydroxides; todorokite; ranciëite; birnessite; asbestiform; human health

1. Introduction

Manganese (Mn) deposits occur in both marine and terrestrial environments as a mix of Mn oxyhydroxides. It is within the marine environment, independent of deep and oxygenated conditions, that the most important Mn deposits are formed. Considering their formation and composition, Mn oxyhydroxide deposits can be classified as hydrogenous, diagenetic, and/or hydrothermal [1]. In both hydrothermal and diagenetic occurrences, Mn oxyhydroxides may exhibit different crystal habits (lamellar, needle-like, fibrous, and asbestiform) [2–8].

Generally, hydrogenous Mn deposits are slowly precipitated from seawater, diagenetic Mn oxyhydroxides are formed by the diagenesis of Mn-rich protolith, and hydrothermal Mn oxyhydroxides are directly precipitated from geothermal solutions around hot springs [9].

Mn oxyhydroxide minerals are a potential economic resource [10]. In particular, todorokite [(Na, Ca, K, Ba, Sr)-(Mn, Mg, Al)-O]-, (Mn, Mg, Al)-O]-, 3-4H₂O] has many potential industrial applications, such as
sorbents, sensors, heterogeneous catalysts, and rechargeable batteries [11]. Regarding birnessite 
\([\text{Na,Ca}]_{0.5}(\text{Mn}_2\text{O}_4\cdot 1.5\text{H}_2\text{O})\), it has been used in previous works as a catalyst in soot combustion 
processes and in methylene blue degradation, showing appreciable catalytic activity compared to 
traditional catalysts [12]. Ranciéite \([\text{Ca,Mn}^{2+}]\text{Mn}^{4+}\cdot 4\text{O} \cdot 3\text{H}_2\text{O}\) is known to be an efficient adsorbent of 
toxic metals, such as plutonium [13]. Despite their economic use, Mn oxyhydroxide deposits could 
pose an environment problem, as they could contaminate the air, soils, sediments, surface water, and 
groundwater [14]. Some studies have reported associations between Mn exposure and adverse health 
effects [15–18], while other studies suggest that Mn-rich water can be neurotoxic for both adults and 
for children [19–22].

Another aspect to consider is that some Mn oxyhydroxides, such as todorokite and manjiroite, 
can crystallize, exhibiting an asbestiform morphology [5–7]. It is worth mentioning that asbestiform 
minerals (e.g., erionite, ferrierite, and fluor-edenite) are sometimes more dangerous than the six 
regulated asbestos fibers (chrysotile, amosite, crocidolite, anthophyllite, tremolite, and actinolite) 
[23,24]. Indeed, non-regulated fibers, such as erionite, ferrierite, and fluor-edenite, are considered to 
be positive carcinogen minerals [25–27]. As such, the US National Institute for Occupational Safety 
and Health (NIOSH) [28] has recently proposed to extend the definition of asbestos to all elongated 
mineral particles (EMPs) [29].

The issue is even more complicated as minerals with fibrous and lamellar habits are often found 
together within the same deposits and the same rocks [29–32], and these minerals must be 
discriminated correctly from a morphological point of view. In this regard, NIOSH highlights the 
difficulty in ascertaining the source of exposure in the case of mixed exposures (i.e., asbestos, 
asbestiform minerals, lamellar minerals) for some mining operations. A large amount of Mn 
 oxyhydroxides with fibrous-asbestiform habits are observed in deposits that can eventually be mined 
and used for industrial applications [5,7]. Disturbance (i.e., erosion, excavation, road construction, 
agricultural activities) of these outcrops can generate airborne fibers with a similar size to asbestos 
fibers, thus increasing the hazard to people who live near these deposits.

As a matter of fact, only sporadic studies were devoted to Mn deposit outcroppings in the 
Messinian sediments from the Serra D’Aiello area [33–35]. In this context, we have collected and 
studied in detail nine Mn-rich samples occurring in the manganiferous deposit outcropping in the 
Messinian sediments from Serra D’Aiello (Southern Italy) by combining X-ray powder diffraction 
(XRPD), differential scanning calorimetry (DSC), thermogravimetry (TG), transmission electron 
microscopy combined with energy dispersive spectrometry (TEM/EDS), and X-ray fluorescence 
(XRF) in order to: (i) inspect their possible mineralogical hazardous nature, (ii) evaluate their 
potential for releasing contaminant elements into soils or waters, and (iii) identify their 
environmental formation.

2. Geological Setting

The studied Mn deposit outcrop in the Serra D’Aiello area in the southern boundary of the 
Amantea Basin sedimentary sequence in Southern Italy is shown in Figure 1.
The basin is characterized by two main sedimentation cycles: a Tortonian-Messinian cycle (TMC) and a Messinian-Lower Pliocene cycle (MLPC), which are divided by a compressive tectonic intramessinian event [37–39]. The TMC formed during a transgressive phase that progressively affected the entire basin and consists of conglomerates and sandy-silty terrains, indicating a transition from a continental (coastal alluvial plain) to a marine environment [40], as recognized using the following terrains in the TMC: (i) red conglomerates, (ii) sandstone, (iii) cinerites, (iv) fossiliferous conglomerates, (v) arenaceous calcarenites, (vi) clays, and (vii) diatomites at the top of the sequence (deep sea conditions). The Mn deposit outcrops in correspondence to the cyclical arenaceous-clayey alternations, whose total thickness is just under 30 meters, overlies the gray fossiliferous clays (Figure 2) and is laterally observed (in the outcrop) for about 150 m, providing a detailed lithostratigraphic description of the Mn outcrop and highlighting the presence of mineralization made up by autogenous minerals, such as Mn hydrated oxides.
Figure 2. Distant view of the alternation of gray silty layers and manganese (black) layers from Serra D’Aiello (Southern Italy).

The first cycle is interrupted by the Messinian tectonic phase, which has a compressive character in the Amantea basin, as evidenced by the shortening of the original basin. After this tectonic phase, sedimentation resumed (cycle II—MLPC) and the terrains were deposited in discordance over the previous cycle, and in some places, directly on the crystalline-metamorphic basement. The recognized succession is constituted by: (i) polygenic conglomerates; (ii) evaporitic limestones and limestones; and (iii) sandstones, clay, and silty clay [40].

3. Materials and Methods

Nine Mn-rich samples were collected and studied using different analytical techniques to obtain a detailed mineralogical and geochemical characterization [41,42]. The collected samples were all black in color and showed a massive appearance. The samples were preliminarily inspected under reflected light using a Zeiss Axioskop 40 microscope (Zeiss, Jena, Germany) in order to choose representative single nodules (i.e., free from alterations) to be studied using X-ray powder diffraction analysis (XRPD), differential scanning calorimetry (DSC), and thermogravimetry (TG).

The bulk geochemical composition of major (SiO$_2$, TiO$_2$, Al$_2$O$_3$, Fe$_2$O$_3$, MnO, MgO, CaO, Na$_2$O, K$_2$O, P$_2$O$_5$) and trace (Ni, Cr, V, Co, Ba, Sr, Cu, Zn, Pb) elements were obtained using X-ray fluorescence (XRF). Chemical elements were analyzed with a rhodium tube with a 40 kW intensity using a Bruker S8 Tiger (Bruker, Karlsruhe, Germany) X-ray fluorescence device (Tables 3 and 4). Chemical data were processed according to the Aitchison model in order to calculate the centered-log-ratio (clr) transformations.

XRPD patterns were obtained on a Bruker D8 Advance X-ray diffractometer (Bruker, Karlsruhe, Germany) with CuKα radiation, monochromated with a graphite sample monochromator at 40 kV and 40 mA. Scans were collected in the range of 3–66° 2θ, with a step interval of 0.02° 2θ and step-counting time of 3 seconds. EVA software (DIFFRACplus EVA version 11.0. rev. 0) was used to identify the mineral phases in each X-ray powder pattern with experimental peaks being compared with 2005 PDF2 reference patterns. Differential scanning calorimetry (DSC) and thermogravimetry (TG) were performed in an alumina crucible under a constant nitrogen flow of 30 cm$^3$ min$^{-1}$ with a Netzsch STA 449 C Jupiter (Netzsch-Gerätebau GmbH, Selb, Germany) in a 25–1000 °C temperature range.
range and a heating rate of 10 °C/min. Derivative thermogravimetry (DTG), derivative differential scanning calorimetry (DDSC) and endothermic peaks were obtained using Netzsch Proteus thermal analysis software, version 4.7.0. The instrumental precision was checked via repeated collections on a kaolinite reference sample (six collections), revealing a good reproducibility (instrumental theoretical T precision of ±1.2 °C) and a theoretical weight sensitivity of 0.10 μg. For DSC/TG and XRPD investigations, samples were ground in an agate mortar. The size, crystallinity, structural features, and chemical composition of single particles were determined using a Jeol JEM 1400 Plus (120 kV) transmission electron microscope (TEM; Jeol, Tokyo, Japan) equipped with a double tilt holder to obtain structural data using selected area electron diffraction (SAED) and with a Jeol large-area silicon drift detector SDD-EDS (Jeol Tokyo, Japan) for microanalyses. For TEM investigations, the sample was put in isopropyl alcohol and then sonicated. Two drops of the obtained suspension were deposited on a Formvar carbon-coated copper grid. In order to describe the size (length and diameter) of the fibers, several dozen TEM micrographs were recorded and six single fibers of todorokite for each sample were measured.

4. Results and Discussion

4.1. Field Observation

Serra D’Aiello’s Mn deposits were localized in correspondence with the cyclical arenaceous-clayey alternations (Figure 2). The deposits were stratified showing layers with thicknesses varying from a few mm up to 30 cm and were also oriented parallel to the direction of the silty layers (Figure 3a-d). Within the layers, the Mn oxyhydroxides were deposited in laminae or nodules of variable dimensions. The latter were constituted by a shell of hydroxides of Mn that enclosed a core of silt (Figure 3e). Their shape was ellipsoidal with a larger diameter ranging from 2–3 cm up to 22 cm and were oriented with the largest diameter parallel to the direction of the layers.

The larger nodules contained smaller nodules and had a thin shell, whereas those that were smaller in size did not contain nodules and had a thicker shell. Within the silty layers, there were also some levels (at least three) with iron oxides (Figure 3f). These were sometimes constituted by nodules and dimensions of less than 3 cm, whereas in others, they formed laminae with a thickness of about 1 cm. Both the layers and the iron oxide nodules were oriented parallel to the direction of the silty layers. The mineralized layers had been affected by a series of faults, bringing them to different heights and constructing a stepped structure.
Figure 3. Typical morphology of the Mn oxyhydroxide deposits at the mesoscale, (a–d) alternation of gray silty layers and black manganese layers, (e) ellipsoidal black manganese nodule, and (f) iron oxide nodules inside the black manganese layers.

4.2. XRPD and DSC/TG Characterization

The minerals detected in the studied samples are listed in Table 1. The evaluation of the XRPD patterns showed that Mn oxyhydroxides were the main phases identified in all samples. The diffraction peaks were found to match with the standard, provided by the International Centre for Diffraction Data (ICDD) [43], of birnessite (ICDD 23-1046), todorokite (ICDD 19-83 and ICDD 21-553) and ranciéite (ICDD 22-718). Quartz, calcite, plagioclase, K-feldspar, rhodochrosite, and muscovite were also detected in varying amounts but not in all samples.

The XRPD patterns of birnessite and ranciéite generally showed broad reflections (Supplementary Materials Figure S1), suggesting that poorly crystalline phases were present in the samples. On the other hand, the XRPD pattern of todorokite was characterized by less broad peaks, indicating a moderate level of crystallinity. The mixed Mn oxyhydroxides (i.e., todorokite, birnessite, and ranciéite) were also proven using a thermal analysis collapse temperature in agreement with the literature data [44–46]. In particularly, the DSC curves showed an endothermic effect at around 200 °C (Figure 4; Table 2) due to the breakdown of birnessite [44]. The DSC curve of ranciéite exhibited one endothermic effect in the range 640–680 °C (Figure 4; Table 2) [45]. Finally, the DSC curve of todorokite showed one shoulder effect at about 460 °C (Figure 4; Table 2), which was clearly recorded on the DSC curve, related to its structural collapse [46]. The very weak endothermic effect in the range 575–597 °C (Figure 4; Table 2) was due to the structural α → β transition of quartz [47], while the endothermic peak at 806 °C (sample 3S) corresponded to the breakdown of muscovite in agreement with Földvári [47]. The endothermic peaks on the DSC curves below 110 °C (Figure 4) were attributed to the release of water adsorbed on the samples’ surfaces.
Table 1. Location of each collected sample (geo-referenced using Universal Transverse of Mercator (UTM) coordinates system and World Geodetic System 1984 (WGS84) as ellipsoid), and their semi-quantitative mineralogical assemblage in order of decreasing abundance as detected using XRPD, DSC/TG, and TEM/EDS. Ranc = ranciéite, Birn = birnessite, Todor = todorokite, Qtz = quartz, Plag = plagioclase, Ms = muscovite, K-feld = K-feldspar, Cal = calcite, Rod = rhodochrosite.

| Samples | X (UTM) | Y (UTM) | Phases       |
|---------|---------|---------|-------------|
| 1S      | 597057.462 | 4327544.556 | Ranc>Birn>Todor>K-feld |
| 2S      | 597169.537 | 4327610.483 | Ranc>Birn>Todor>Qtz>Plag>K-feld |
| 3S      | 597288.205 | 4327528.075 | Birn>Ranc>Cal>Todor>Qtz>Plag>K-feld>Ms |
| 4S      | 597166.241 | 4327330.295 | Birn>Ranc>Todor>Cal>Qtz>Plag>K-feld |
| 5S      | 597301.39  | 4327294.035 | Ranc>Birn>Todor>Qtz>Plag>K-feld |
| 6S      | 597413.465 | 4327244.59  | Ranc>Birn>Qtz |
| 7S      | 597113.499 | 4327162.182 | Todor>Ranc>Rod>Qtz |
| 8S      | 597228.871 | 4327076.478 | Birn>Ranc>Qtz>Cal>Plag |
| 9S      | 597331.057 | 4327063.293 | Ranc>Birn>Qtz |

Table 2. Peak temperatures (°C) in DSC curves; w—weak, s—strong, sh—shoulder, endo—endothermic. Exothermic peaks have not been reported.

| Phases         | 1S     | 2S     | 3S     | 4S     | 5S     | 6S     | 7S     | 8S     | 9S     |
|----------------|--------|--------|--------|--------|--------|--------|--------|--------|--------|
| Water/humidity | 92 endo s | 80 endo s | 80 endo s | 93 endo s | 80 endo s | 92 endo s | 70 endo s | 80 endo s | 108 endo s |
| Birnessite     | 214 endo s | 190 endo w | 197 endo s | 216 endo s | 210 endo s | 206 endo s | -       | 204 endo | 208 endo |
| Todorokite     | 472 sh   | 460 sh  | 423 sh  | 443 sh  | 480 sh  | -       | 474 endo w | -       | -       |
| Quartz         | -       | 575 endo | 575 endo | 576 endo | 576 endo | 597 endo | -       | 577 endo | 598 endo |
| Rhodochrosite  | -       | -       | -       | -       | -       | -       | 570 endo | -       | -       |
| Ranciéite      | 661 endo s | 650 endo s | 670 sh  | 681 endo | 663 endo s | 677 endo s | 643 endo s | 678 endo | 678 endo s |
| Calcite        | 743 endo s | 751 endo s | 798 endo | 743 endo s | 751 endo s | 798 endo | 743 endo s | 751 endo s | 798 endo |
4.4. TEM/EDS Characterization

The identification of mineral components and the characterization of the morphologies using scanning electron microscopy (SEM) were not undertaken due to the colloidal texture and nanometer-scale sizes of the Mn oxyhydroxides. Indeed, both secondary and backscattered electron images did not show any detailed features of the Mn oxyhydroxides. In light of this, as well as the characteristics and regarding the nanometric dimensions of the Mn oxyhydroxides layers, TEM was used due to its high magnification power.

A representative set of TEM micrographs showing the morphology of Mn oxyhydroxides observed in the various samples are given in Figures 5,6. In accordance with literature data [48], TEM observations of birnessite showed crumpled sheet morphologies (Figure 5a), all characterized by rather poor crystallinity (Figure 5a). The surface of the birnessite seemed to peel off in many plate-like layers typical of phyllomanganate Mn oxyhydroxides [48]. These layers were partially wrapped, and in some places, it was possible to observe the interlayer of its structure (Figure 5b). Rancièite were found to be elongated platelets (Figure 5c) from fractions of a few nanometers to micrometers in length (Figure 5d). The particles of rancièite in all samples were found to be irregularly shaped platelets similar to those of birnessite under TEM (Figure 5). Under TEM, todorokite showed a strain-shaped morphology with parallel sides and regular termination, and occasionally, the longitudinal splitting of such fibers into thinner fibrils was observed (Figure 6). In some cases, todorokite fibers showed tight extremities assuming a conical morphology (Figure 6b). The fibrous todorokite was found in all samples, and always in conjunction with lamellar birnessite and/or rancièite. It is worth remembering that todorokite fibers grow via the crack-and-seal mechanism [7]. Similar crack-and-seal structures are characteristic for the occurrences of chrysotile (asbestos) and asbestiform amphiboles in weakly metamorphosed formations [30–32]. Regarding the particle sizes detected using TEM, the todorokite fiber lengths in all samples ranged from a few nanometers to 10 μm, and from about 10 nm to 500 nm in width. In all the samples, most of the fibers measured were generally longer than 5 μm, with a width of < 3 μm and with aspect ratios > 3:1, and were therefore classified as asbestiform fibers [49,50].

The semi-quantitative chemical composition of several particles, as determined using TEM/EDS analyses, was in line with the general chemical composition of birnessite (Na,Ca)0.5(Mn)2O4·1.5H2O, todorokite (Na,Ca,K,Ba,Sr)1-x(Mn,Mg,Al)6O12·3-4H2O, and rancièite (Ca,Mn)0.2(Mn)O2·0.6H2O. The variation of the calcium percentage in the rancièite and birnessite particles was used in order to discriminate between the two minerals, in agreement with Chukhrov et al. [51].

The chemical composition of the respective Mn oxyhydroxides hosted many major, minor, and trace elements. For example, todorokite had a variable composition where elements such as Zn, Mg, Ba, Sr, Ca, Na, K, Cu, Pb, and Ni could be detected in variable amounts. Calcium and alkali contents in birnessite fluctuated, and other elements, such as Mg, Ba, Zn, Cu, Co, Li, Al, and Fe, were also detected. These elements were often embedded in a sandwich–like manner as hydroxides in the form of disordered interlayers between well-ordered Mn oxyhydroxides or as Mn substitutes in the octahedral sites [45,48]. In rancièite, variable amounts of Ca and Fe, and minor amounts of Mg, Al, Na, K, Si, P, and Cu were detected. As a matter of fact, variations in the chemical composition are linked with complex structure of Mn oxyhydroxides and can be explained easily by the manifold substitutions of the cations [45]. In this regard, Mn oxyhydroxides’ high reactive surface area particles play innumerable roles in the transformation and cycling of major and trace elements in hydrogenetic, diagenetic, and hydrothermal environments [52,53].
Figure 5. Representative TEM micrograph: (a) aggregates of sheet-like birnessite in sample 4S (the SAED pattern is shown in the inset); (b) detail of (a) where the interlayer of the birnessite structure is indicated by the black arrow; (c) rancièite occurrence as elongated platelets (sample 5S), and (d) detail of (c) where the interlayer spacing of rancièite structure of ≈ 8 Å is indicated by the white arrow.
Figure 6. Representative TEM micrograph of the samples examined: (a) fibrous todorokite (sample 1S); (b) thin fibers of todorokite (sample 3S); (c) fibrous todorokite (note the curvature flexibility; sample 7S); (d) fibrous todorokite (cleavage fragment, note the irregular sides; sample 4S); (e) fibrous todorokite (sample 9S); and (f) bundle of todorokite fibers (sample 2S).

4.5. XRF Characterization

Through the X-ray fluorescence analytical technique, the following major and trace elements were found in the samples of manganese mineralization from Serra D’Aielo: MgO, Al2O3, SiO2, P2O5, K2O, CaO, TiO2, MnO, Fe2O3, Na2O, Rb, Sr, Cr, V, Ba, Ni, Co, Zn, Pb, and Cu. Tables 3 and 4 report the content of the oxides (in wt%) and trace elements (in ppm) for each sample.

As expected, MnO was the main component in all samples with mean values of 53.7 ± 8 wt% followed by SiO2 and CaO (13 and 6 wt%, respectively).

The relatively high values of SiO2, Al2O3, and K2O were explained by the occurrence of minor amounts of silicate minerals, such as quartz, K-feldspar, plagioclase, and muscovite (Table 1).

Generally, a strong correlation did not exist between MnO and any other major elements (Table 3).

It is noteworthy to mention, however, that the mineralized beds contained some iron oxide abundances, which were separated by the manganiferous layers, suggesting that the solution from which the Mn precipitated also contained a fair amount of iron. It is good to note that in submarine hydrothermal Mn deposits, a common geochemical feature is the fractionation of Fe from Mn.

The initial precipitation of Fe-bearing phases is due to their higher precipitation rates with respect to Mn ones [54–58].

Concerning trace elements (Table 4), samples showed mutually comparable Sr, Cr, V, Zn, and Cu concentrations, whereas different distributions have been recognized for the remaining trace
constituents (Ba, Ni, Co, and Pb). The high barium value in samples 5S and 9S was probably linked to the high content of rancéite detected, in which often the calcium present in its structure is replaced by barium [56].

In detail, samples 1S, 2S, 3S, 4S, and 5S showed anomalous Ni concentrations, close to or slightly below 500 ppm, which was different to what was highlighted in the remaining samples (below 150 ppm). Furthermore, samples 3S and 9S, and 4S and 6S, were characterized by high concentrations of Pb and Co, respectively (Table 3). High concentrations in these trace elements made them potentially available for subsequent accumulations in soils, water, and air with consequent negative effects on human health.

Table 3. Major element concentrations (in oxides wt%) obtained using XRF analysis. (LOI.: loss on ignition). Fe$_2$O$_3$ stands for total Fe.

| ID | LOI  | MgO | Al$_2$O$_3$ | SiO$_2$ | P$_2$O$_5$ | K$_2$O | CaO | TiO$_2$ | MnO | Fe$_2$O$_3$ | Na$_2$O | Tot  |
|----|------|-----|------------|---------|------------|-------|-----|---------|-----|------------|--------|------|
| 1S | 14.21| 3.50| 5.33       | 15.32   | 0.34       | 1.13  | 9.20| 0.34    | 47.58| 2.72       | 0.32   | 100  |
| 2S | 16.32| 2.56| 3.48       | 10.04   | 0.22       | 0.79  | 6.77| 0.18    | 57.53| 1.86       | 0.25   | 100  |
| 3S | 15.57| 3.34| 3.19       | 9.62    | 0.09       | 0.46  | 4.17| 0.13    | 61.76| 1.53       | 0.13   | 100  |
| 4S | 17.02| 3.54| 3.90       | 10.88   | 0.14       | 0.66  | 8.07| 0.20    | 53.34| 2.03       | 0.21   | 100  |
| 5S | 16.09| 2.18| 2.44       | 7.52    | 0.14       | 0.45  | 4.10| 0.25    | 65.38| 1.44       | 0.15   | 100  |
| 6S | 14.52| 2.16| 3.50       | 11.24   | 0.21       | 0.95  | 7.04| 0.11    | 57.65| 2.32       | 0.31   | 100  |
| 7S | 14.59| 2.52| 4.55       | 13.27   | 0.27       | 0.81  | 7.46| 0.20    | 53.64| 2.27       | 0.40   | 100  |
| 8S | 19.70| 4.09| 4.85       | 15.34   | 0.19       | 0.91  | 6.34| 0.22    | 45.67| 2.37       | 0.30   | 100  |
| 9S | 13.83| 6.40| 6.69       | 23.24   | 0.12       | 1.06  | 4.29| 0.28    | 40.92| 2.87       | 0.30   | 100  |

Table 4. Trace element concentrations (in ppm) obtained using XRF analysis.

| ID | Sr  | Cr  | V   | Ba  | Ni  | Co  | Zn  | Pb  | Cu  |
|----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
|    | ppm | ppm | ppm | ppm | ppm | ppm | ppm | ppm | ppm |
| 1S | 159 | 23  | 58  | 135 | 555 | 115 | 0   | 172 | 57  |
| 2S | 153 | 14  | 28  | 114 | 511 | 38  | 0   | 75  | 57  |
| 3S | 157 | 24  | 52  | 158 | 472 | 32  | 0   | 309 | 49  |
| 4S | 160 | 24  | 27  | 132 | 509 | 566 | 0   | 169 | 65  |
| 5S | 261 | 21  | 43  | 578 | 379 | 101 | 0   | 101 | 38  |
| 6S | 254 | 11  | 22  | 112 | 94  | 298 | 37  | 92  | 56  |
| 7S | 241 | 25  | 40  | 164 | 57  | 57  | 0   | 95  | 38  |
| 8S | 219 | 24  | 55  | 125 | 151 | 94  | 0   | 0   | 57  |
| 9S | 243 | 39  | 90  | 1086| 127 | 109 | 54  | 746 | 73  |

Indeed, nickel compounds can cause a variety of adverse effects on human health, such as a nickel allergy in the form of contact dermatitis, lung fibrosis, cardiovascular and kidney diseases, and cancer of the respiratory tract [59–61]. In the same way, an excessive amount of cobalt exposure could cause a complex clinical syndrome with a varying set of neurological, cardiovascular, and endocrine deficits that are directly related to the uptake of Co ions in the tissue and in blood circulation [62]. Moreover, lead is also an extremely harmful environmental pollutant. If it is adsorbed, it can cause several negative affects to the respiratory and digestive body systems [63]. The threshold values of potential hazardous elements in each environmental matrix (soil, water, and air) are reported in Table S1 (Supplementary Materials). As such, defining the concentration of these compounds (minerals,
elements) in these geological settings is the first step to limiting the exposure and consequently the problems that they can cause to those who live within the surrounding areas.

4.6. Origin of the Mn-rich Mineralizing Fluid

The geochemical characterization is an indispensable tool for understanding the genesis of Mn deposits. Using geochemical parameters, marine Mn deposits can be classified into three main types [1]: (i) hydrothermal, (ii) hydrogenetic, and (iii) diagenetic deposits, as well as through a combination of these processes [57].

A (Cu + Ni + Co) × 10 – Fe – Mn diagram is usually used to identify the possible origin of Mn ores [64–65]. Figure 7 shows that the studied samples fell within the hydrothermal field. The high Mn/Fe ratio recorded suggests a typical deposit of a distal environment [9,66], which was also reflected by the lack of Fe phases in the selected Mn deposits.

Geochemical features of marine siliceous sediments, in particular the abundances of quartz and/or of its polymorphs common in deep-ocean floor sediments and in ophiolite sequences [67,68], have been studied to obtain information about their sources and depositional environments [69,70].

In response to this, the Al/(Al+Fe+Mn) and MnO/TiO₂ ratios were used to evaluate the relative contributions of the terrigenous and hydrothermal input of marine sediments, with Al and Ti being representative of a terrigenous source and Mn and Fe of a hydrothermal one, as reported by Kato et al. [71]. For the Serra D’Aiello samples, very low Al/(Al+Fe+Mn) values (lower than 0.04) and very high MnO/TiO₂ values (>218) were detected. These ratios allowed us to exclude the option of a terrigenous input, suggesting that silica in the studied mineralizations may be indicative of further evidence of a hydrothermal origin (Figure 8).

Based on obtained data, and considering the geological setting of the studied area, this could suggest that hydrothermal processes have driven the deposition of the Mn-bearing minerals in the Serra D’Aiello sedimentary basin. The obtained results agree with those shown by Sinisi et al. [9], who have studied Mn-rich mineralization in the metasedimentary succession from the Calabria region (Italy).

![Figure 7. Triangular plot for the genesis discrimination of the Mn deposits.](image-url)
Figure 8. (a) Al/(Al + Fe + Mn) and (b) MnO/TiO2 ratios for each sample.

5. Conclusions

A multidisciplinary approach based on mineralogical and geochemical analyses was used in order to characterize the Mn deposits from Serra D’Aiello (Southern Italy). The results obtained using XRPD, DSC/TG, TEM/EDS, and XRF showed that birnessite, todorokite, and ranciéite occurred as Mn oxyhydroxides phases in the analyzed samples. The Mn oxyhydroxides were poorly crystallized, and their morphology could only be recognized under TEM/EDS.

The release of Mn in the groundwater/soils, as well as other heavy metals from the Mn oxyhydroxide deposit, was regulated by a multitude of variables; therefore, the mineralogical and geochemical characterization of Mn oxyhydroxides minerals is important in order to define the potential environmental contamination in soils and groundwaters in areas where Mn oxyhydroxides occur. This investigation highlighted that ranciéite and birnessite with platy morphology occurred in all the analyzed samples, whereas the asbestiform todorokite was detected in seven samples.

Todorokite occurred in large amounts as microcrystalline fibrous–asbestiform phases in diagenetic rocks with fibers of breathable sizes (particles with length $\geq 5 \, \mu m$, width $\leq 3 \, \mu m$, length/width ratio $\geq 3:1$). It is important to note that, in addition to the minerals regulated as asbestos by law, asbestiform minerals, such as fibrous todorokite, could also potentially be dangerous if inhaled.

In the Amantea basin, a complete separation between Fe and Mn geochemistry during deposition and subsequent alteration was observed. This could be explained by an early segregation, with a release of a much greater abundance of Mn in comparison to Fe. From a geochemical point of view, some samples showed anomalous concentrations of Ni, Co, and Pb. The accumulations of these compounds in soils, water, and air can cause negative effects on human health, causing mild pathologies. Consequently, the widespread nature of both trace metals and asbestiform todorokite into the environment may be a cause of serious health problems for people who live near the sampling sites. For these reasons, an environmental monitoring plan in these potential contaminated sites is needed.

Finally, based on mineralogical and geochemical data, as well as considering the geological setting of the studied area, it was ascertained that hydrothermal processes have driven the Mn-bearing mineral deposition in the Serra D’Aiello sedimentary basin according to similar evidence presented in other metasedimentary successions outcropping in the Calabria region (Italy).

Supplementary Materials: The following are available online at www.mdpi.com/xxx/s1, Figure S1: XRPD pattern of the samples examined. Peaks were assigned according to the literature (International Centre for Diffraction Data (2005). ICDD Products/PDF-2, http://www.icdd.com/products/pdf2.htm). Table S1: Threshold
values of potential hazardous elements in soils and groundwaters (Italian Legislative Decree No. 152 of 03/04/2006) and air (NIOSH 2007).

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