

## EPITHERMAL MANGANESE MINERALIZATION, KIMOLOS ISLAND, SOUTH AEGEAN VOLCANIC ARC, GREECE

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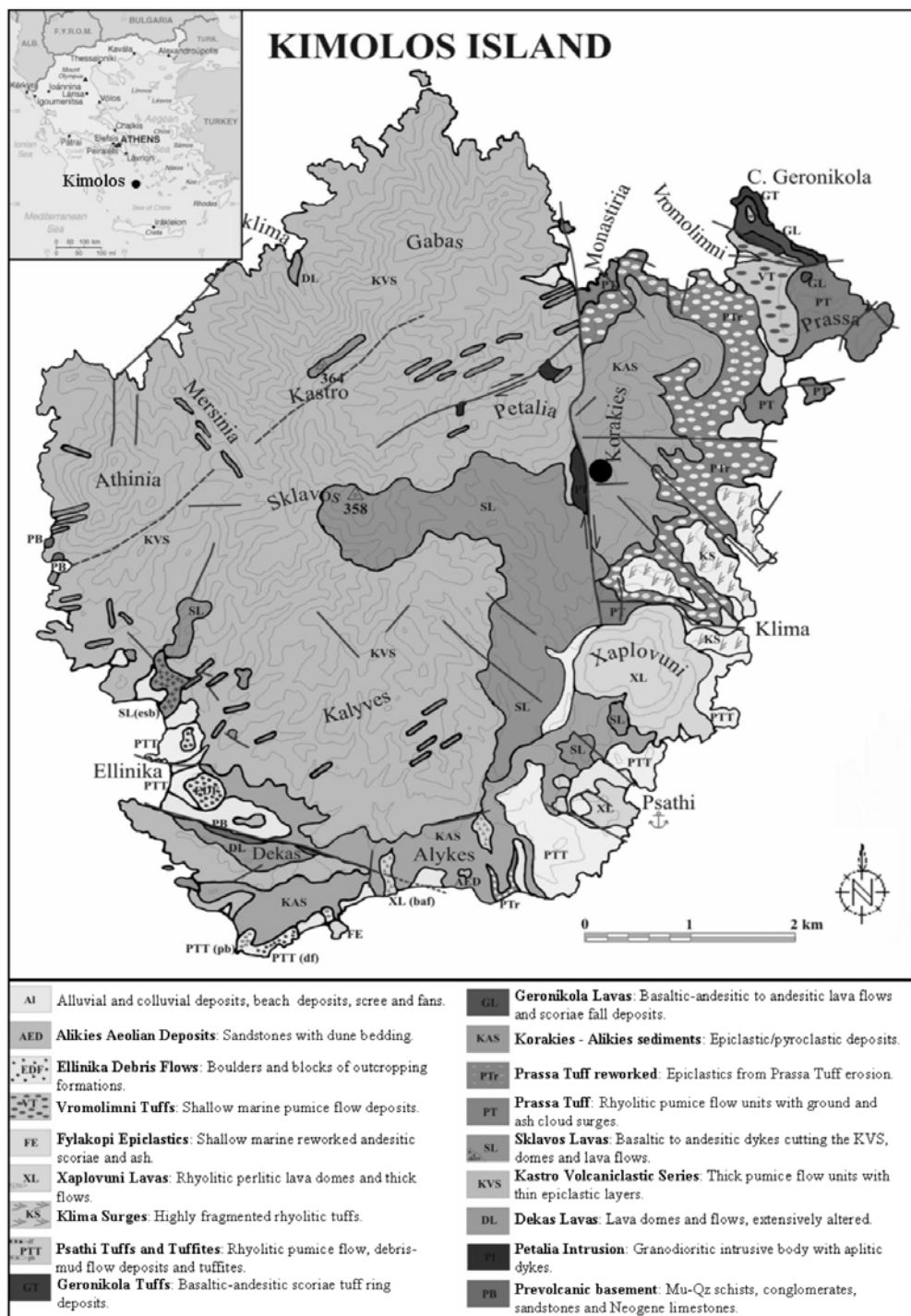
### Abstract

*Manganese mineralization is hosted by a marine monomictic, lithic volcanoclastic breccia, possibly an andesitic in situ hyaloclastite, and shallow-marine or subaerial epiclastic conglomerates, in the Korakies area, NE Kimolos, active south Aegean volcanic arc. Old mine workings (in the form of rubble, adit and shaft), and abandoned rail and ship loading facilities, exist in the area. Mineralization occurs as a quartz/chalcedony vein system filling extensional NNE-SSW-trending faults and fractures, of Pliocene age. Maximum vein width reaches 5 m; length may extend to 250 m. The ore shares strong textural analogies with volcanic-hosted epithermal-style deposits, i.e. crustiform banding, vugs, hydrothermal breccias, cockade and comb textures. Vein wall rocks are hydrothermally altered to quartz-adularia±illite, chlorite and barite. Pyrolusite, hollandite, cryptomelane, and coronadite are the main ore minerals, with quartz, chalcedony, jasper and barite gangue. Ore samples contain up to 25.8 % MnO<sub>2</sub>, 14.7 % FeO<sub>TOT</sub>, 2860 ppm Zn, 1132 ppm Pb and 136 ppm Cu; Mn and Zn show mutual positive correlation ( $r^2=0.61$ ). Trace element enrichment (i.e. Zn, Pb, and Cu) may suggest a proximal base metal sulfide mineralization. Concentrations of 4.3 % Na, 0.09 % Mg and barite presence may suggest genetic involvement of sea water. The mineralization studied is similar to volcanic-hosted low-sulfidation epithermal ore deposits deposited from neutral pH fluids. This is a rare example of a vein-type epithermal-style hydrothermal manganese deposit formed in a marine environment.*

**Key words:** *epithermal manganese, Kimolos, South Aegean Volcanic Arc, hollandite, pyrolusite, cryptomelane, epithermal textures.*

### 1. Introduction

Kimolos volcanic island (36 km<sup>2</sup>) shares a common geological evolution with Milos, Polyegos and Antimilos the Milos Group of the Pliocene-modern South Aegean volcanic arc (SAVA) (Pe-Piper and Piper, 2002). To date, metallic mineralization has only been reported from Milos, a dormant volcano that documents the transition between the submarine and terrestrial volcanic environments (Fytikas et al., 1986, Stewart and McPhie, 2006). Milos is host to a large inventory of volcanic-hosted metallic deposits (e.g. Hauck, 1984; Plimer, 2000; Liakopoulos et al., 2001; Kiliias et al., 2001; Naden et al., 2005; Alfieris and Voudouris, 2007; Glasby et al., 2005) as well as industrial mineral deposits (Christidis, 2001, and references therein). Recent research on Milos has identified a new metallogenic environment—namely hybrid volcanic-hosted massive sulfide and continental magmato-hy-



**Fig. 1:** Geological map of Kimolos Island showing the main structural features and the location of the Mn ore deposit in the Korakies area (black dot) (modified from Francalanci et al., 2007).

drothermal mineralization (i.e. Profitis Ilias–Chondro Vouno Au–Ag–Cu deposit, Naden et al., 2005). In addition, epithermal and sea-floor exhalative textures have been documented in the Vani–Kondaros–Triades Mn–Ba–Pb–Zn–Ag hydrothermal field (Kilias et al., 2007, and unpublished data).

Published reports on the geology and hydrothermal mineralization on Kimolos are very scarce (Fytikas and Vougioukalakis, 1993, Pe-Piper and Piper, 2002; Francalanci et al., 2007); this is the first report on hydrothermal manganese mineralization that occurs in the Korakies area, NE Kimolos.

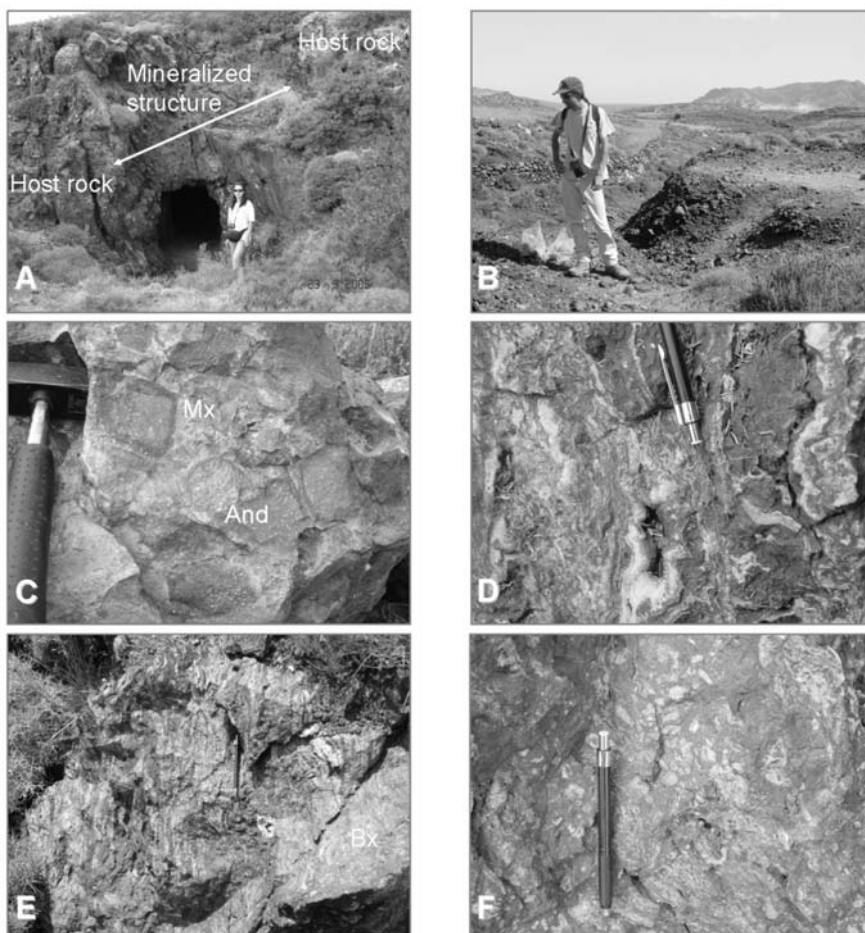
## 2. Geological setting

Volcanic centres of the Milos Group are positioned along NE-SW trending regional tectonic lineaments (Francalanci et al., 2007; Pe-Piper and Piper, 2002). Both Kimolos and Milos consist of marine and terrestrial lithologies with composition belonging to the calc-alkaline and high-K calc-alkaline series (Pe-Piper and Piper, 2002; Francalanci et al., 2007, and references therein). Volcanic activity occurred in two periods separated by a long serenity phase. Basaltic to rhyolitic lavas and voluminous pyroclastic rocks were erupted during the first period (3.5–2.0 Ma) in association with NE-SW-trending lineaments, whereas rhyolitic pyroclastic deposits and perlitic lava domes tied to NW-SE structural features, characterize the second period of activity (2.0–0.9 Ma) (Fig. 1). Structurally controlled NE-SW trending, active low-T (~55 °C) geothermal activity, NW Kimolos is recharged mainly by seawater (Pe-Piper and Piper, 2002, Fytikas and Vougioukalakis, 1993).

The host rocks of the manganese mineralization form part of the Korakies-Alikes pyroclastic-epiclastic formation (Fig. 1). This formation consists of a mixture of shallow marine and subaerial pyroclastic and epiclastic lithologies, such as pyroclastic breccias and debris flows, sandstones and siltstones, and epiclastic conglomerates (Fytikas and Vougioukalakis, 1993). The mineralized structures crosscut an ENE-WSW trending monomict, andesitic volcanoclastic lithic breccia, and locally unconformably overlying epiclastic conglomerates. The volcanic rocks show a conspicuous flow texture with individual 2-3 m thick flow-beds trending ENE-WSW. Wallrocks constitute: (1) a clast-supported lithic breccia which contains angular to subangular andesite lava fragments with an average diameter of 20-30 cm, and a matrix of fine grained lava fragments of the same composition (Fig. 2C); this formation is topographically controlled and shows maximum thickness of 30 m best seen along paleo-valleys (Fytikas and Vougioukalakis, 1993). These rocks represent possibly submarine, lithic-rich, volcanoclastic mass-flow deposits, or andesitic in situ hyaloclastites; (2) shallow-marine or subaerial epiclastic sediments which have clasts consisting of basaltic to andesitic Sklavos lavas (see Fig. 1) and Kastro volcanoclastics (see Fig. 1) cemented by hydrothermal silica. Host rocks come to a faulted N-S trending contact with the Kastro volcanoclastics to the east (Fig. 1).

## 3. Manganese ore: occurrence and textures

Abandoned mine workings (in the form of rubble, adit and shaft) and ship loading facilities, attest to past exploitation of the Mn deposit at Korakies. The deposit occurs in veins hosted exclusively by the Korakies pyroclastic/epiclastic formation. The exploited vein fills NNE-striking normal faults and related extensional fractures within a structural corridor of the regional Pliocene NE trend (Pe-Piper and Piper, 2002). It has maximum width of 5-6 m, a length of at least 250 m and, dips 70° to 90° to the SW; the depth of the vein is unknown (Fig. 2A). The vein displays complex and multi-episodic filling with epithermal-style textures characteristic of open-space precipitation such as crustiform banding, vugs, hydrothermal breccias, and cockade and comb textures (Figs. 2D&E&F, Fig. 3A); breccia mainly occurs on the vein margins (Hedenquist et al., 2000) (Fig. 2E). These textures suggest that vein opening and filling was episodic, with several episodes of fault movement related



**Fig. 2:** A. Photo of the Korakies vein in which the banded texture, the width of the mineralized structure and an old mining gallery can be seen. B. Photo of abandoned mine workings close to the vein, where a number of samples were collected. C. Photo of the host rock of Mn mineralization which is a monomict volcanoclastic lithic breccia. Blocky hornblende— and feldspar—phyric andesite clasts (And) are separated by small amounts of millimeter to submillimeter-sized granular matrix (Mx) of the same composition. The matrix includes splinters of andesite and scattered crystal fragments. Group of clasts in the matrix display jigsaw-fit texture. Lithic fragments are surrounded by pinkish jasper. D. Epithermal vein texture displaying: crustiform banded quartz-chalcedony-jasper intergrown with bands of manganese mineralization, and, cockade texture. E. Transitional relationship between banded quartz-Mn ore texture and hydrothermal breccia (Bx) F. Enlargement of (Bx) in previous photo. Hydrothermal fault-breccia structure consists of angular vein material and matrix with Mn-ore. (And: andesite; Mx: matrix; Bx: breccia).

to brecciation and mineralization. Primary Mn ore forms part of these textures. Banding may be related to cyclic pulses of fluid caused by a feeder mechanism in which fluid overpressure caused faulting events during low effective normal stress conditions. A supergene oxidation zone exists of Fe-Mn-oxy-hydroxides, affecting at least 5 m of the upper and exposed parts of the vein.



## 4. Methods

Thirty two mineralized, host rock and vein wallrock samples were collected both in-situ and from old nearby mine rubble (Fig. 2B). The samples were examined under transmitted and reflected light microscopes, and analysed by SEM-EDS, AAS and XRD methods in the laboratories of the Dept. of Economic Geology and Geochemistry, University of Athens. We used an SEM-EDS Jeol Jsm-5600 (Oxford ISIS 300 microanalysis system) with 20 kV voltage, current of 0.5 nA, analysis time of 50 sec and bundle diameter <2  $\mu$ . For the XRD studies, a Siemens D5005 CuK $\alpha$  radiation diffractometer was used, with 1.54 Å wavelength and 40 kV voltage. The concentrations of K and Na were measured with a Jenway PFP7 flamephotometer.

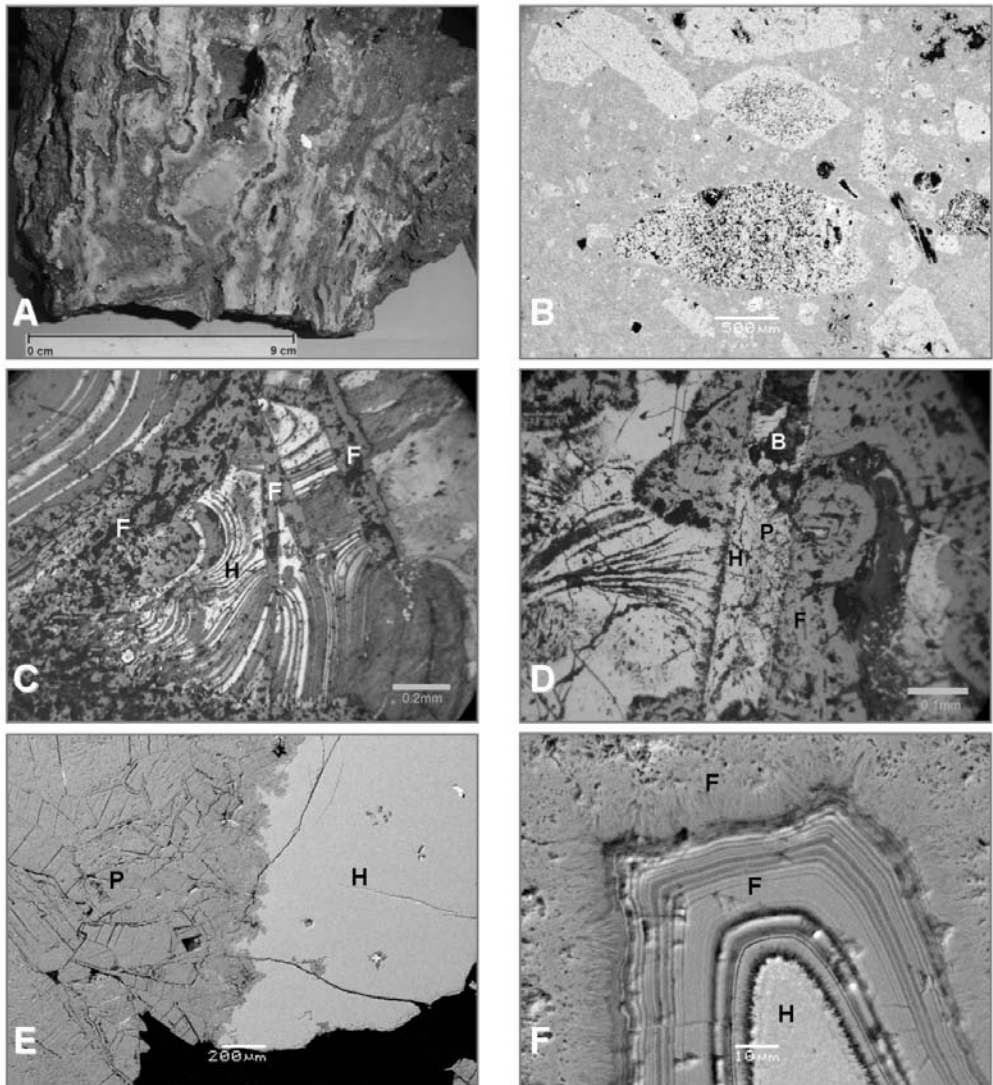
## 5. Mineralogy and mineral chemistry

**Hydrothermal alteration:** Due to overprint of supergene oxidation, hydrothermal alteration of the host rocks is not easily discernible. It seems that it is limited to the vicinity of the main vein, to an unknown extent. The most common minerals present in altered wall rock are quartz, adularia, Ba-adularia, illite (sericite), chlorite, and Fe-oxides. The main alteration type is quartz-adularia+chlorite $\pm$ illite. Quartz is mainly present as a pervasive replacement of the volcanic host rock matrix, whereas adularia, illite and chlorite grow over feldspar and hornblende clasts and crystal fragments of the groundmass (Fig. 3B). Traces of Mn-ilmenite were also found.

**Manganese minerals:** Primary manganese ore essentially consists of pyrolusite and intermixed hollandite-group Mn-oxide minerals (hollandite-cryptomelane-coronadite). Gangue minerals are quartz, chalcedony, jasper, barite, minor adularia and illite. Supergene minerals are iron oxides and hydroxides, and Mn-Fe wad. Formation of manganese, and iron, minerals may have been controlled by Eh changes, under either hypogene and/or supergene oxidizing conditions, or both (i.e. ascending and/or descending fluids) (see Figure 3C&D); temperature and Eh (oxidized sulfur) changes, may have caused gangue precipitation, and barite, respectively (e.g. Leal et al., 2008). Pyrolusite appears as masses of well-formed rhombohedral to prismatic and needle-like crystals that may reach a few millimeters in size. Hollandite-group minerals occur mainly as repetitive colloform microbands with variable grain size and porosity that commonly follow the same succession of precipitation (Fig. 3C), or vug filling wad (Fig. 3D&E). Goethite is the most common iron oxide; it is present as a botryoidal aggregate and usually occurs with hematite. Electron microprobe analyses of Mn minerals are shown in Table 1.

## 6. Geochemistry

Whole rock chemical analyses of manganese ores are shown in Table 2. The ore samples may contain up to 25.83% Mn, 14.7% Fe, 2860 ppm Zn, 1132 ppm Pb, 136 ppm Cu and 3780 ppm Ba, while concentrations in Ni and Co are very low. In addition, concentrations of 4.3 % Na, 0.75 % K, 0.77% Ca, and 0.09 % Mg have been measured. According to the correlations between all elements, they can be subdivided into different groups; Mn and Zn show mutual positive correlation (correlation coefficient  $r^2=0.61$ ) indicating that Zn is associated with the manganese oxide phase; Fe-Pb-Cu-Zn and Cr show mutual positive correlation ( $r^2=0.74-0.99$ ), suggesting that some of these elements may be adsorbed onto supergene goethite and hematite, however, association with Mn minerals cannot be excluded. Na, K, Al and Ca show positive mutual correlations but do not have a statistically significant correlation with Fe, Mn, or Si-Al. These elements are probably partitioned among all those phases, but predominantly with an aluminosilicate phase. The high Na content may suggest involvement of seawater (Hein et al., 2008); Mg does not correlate with any other element and may also originate from seawater. The Ca content probably reflects a carbonate sediment source.



**Fig. 3:** A. Polished section showing Mn-ore forming part of epithermal-style crustiform banding, vugs and cockade structures. B. Back-scattered electron image of hydrothermally altered wallrock. Equant to cuneiform andesite clasts are packed in a matrix that contains scattered crystal fragments. The clasts consist of feldspar and hornblende altered to Ba-adularia, chlorite, illite, and quartz; groundmass comprises fine-grained silica and altered feldspar and hornblende. C. Microphotograph of colloform hollandite (light grey) cut by hematite and goethite. D. Microphotograph of hollandite wad and pyrolusite within an older elongate barite crystal; goethite-hematite are the supergene phase E. Back-scattered electron image illustrating primary assemblages of pyrolusite (P) and hollandite/coronadite (H). F. Back-scattered electron image of finely laminated bulbous and smooth undulating aggregates of Fe-rich (goethite-hematite) phases, of possible biogenic (microbial) origin (P:pyrolusite; H:hollandite/coronadite; B: barite; F: goethite-hematite)

**Table 1.** Electron microprobe analyses (wt%) of selected manganese oxides from Korakies. Samples 1-13: hollandite – cryptomelane, 14-17: hollandite – coronadite, 18-26: pyrolusite, 27-36: goethite – hematite (n.d= not detected).

	MgO	Al <sub>2</sub> O <sub>3</sub>	SiO <sub>2</sub>	K <sub>2</sub> O	CaO	MnO <sub>2</sub>	Fe <sub>2</sub> O <sub>3</sub>	ZnO	BaO	PbO	As <sub>2</sub> O <sub>3</sub>	TOTAL
1	0.5	n.d	n.d	4.5	n.d	80.4	1.7	n.d	5.8	n.d	n.d	93.3
2	1.0	n.d	0.3	1.5	0.2	81.1	n.d	0.7	6.2	n.d	n.d	91.1
3	n.d	n.d	0.4	2.8	n.d	76.9	2.2	n.d	7.2	0.6	n.d	90.5
4	3.2	n.d	n.d	2.4	0.3	81.9	n.d	n.d	7.2	n.d	n.d	95.3
5	1.0	n.d	n.d	2.3	n.d	79.0	n.d	n.d	7.2	n.d	n.d	90.4
6	3.3	n.d	n.d	2.5	n.d	83.1	n.d	0.6	7.3	n.d	n.d	97.3
7	n.d	n.d	n.d	2.0	0.2	80.3	n.d	n.d	7.4	n.d	n.d	90.8
8	1.1	n.d	n.d	1.3	n.d	79.7	n.d	0.8	7.5	0.1	n.d	90.5
9	3.2	n.d	n.d	2.5	0.4	82.5	n.d	n.d	7.5	n.d	n.d	96.7
10	n.d	n.d	0.8	0.7	0.6	65.2	3.6	0.8	10.8	n.d	n.d	82.6
11	n.d	n.d	0.3	1.7	0.3	75.9	1.9	n.d	9.9	n.d	n.d	91.2
12	n.d	n.d	n.d	1.0	n.d	77.6	n.d	0.8	12.5	n.d	n.d	92.8
13	n.d	n.d	n.d	1.2	n.d	76.8	0.9	n.d	12.7	n.d	n.d	92.5
14	n.d	n.d	0.8	0.3	n.d	68.3	n.d	0.8	1.5	28.0	n.d	99.7
15	n.d	n.d	0.7	0.2	n.d	68.0	n.d	0.7	1.5	28.1	n.d	99.2
16	n.d	n.d	0.8	n.d	n.d	66.1	n.d	0.9	1.3	28.5	n.d	97.8
17	n.d	n.d	0.8	n.d	n.d	65.8	n.d	0.8	1.1	28.7	n.d	97.2
18	n.d	n.d	0.7	0.2	0.3	87.0	n.d	0.7	2.2	n.d	n.d	91.7
19	n.d	n.d	1.2	0.2	n.d	87.2	0.6	n.d	1.7	n.d	n.d	91.7
20	0.4	n.d	0.6	n.d	n.d	89.2	n.d	0.6	n.d	n.d	n.d	91.4
21	n.d	n.d	0.5	n.d	n.d	89.3	n.d	n.d	n.d	n.d	n.d	90.3
22	n.d	n.d	0.7	n.d	n.d	89.4	n.d	1.1	n.d	n.d	n.d	91.6
23	n.d	n.d	n.d	n.d	n.d	89.9	n.d	n.d	n.d	n.d	n.d	91.1
24	1.1	n.d	0.8	n.d	n.d	90.0	n.d	n.d	n.d	n.d	n.d	92.7
25	n.d	n.d	0.7	0.0	n.d	90.6	n.d	0.7	n.d	n.d	n.d	92.7
26	n.d	n.d	1.1	n.d	n.d	90.7	n.d	0.9	n.d	n.d	n.d	92.3
27	n.d	n.d	3.0	n.d	n.d	3.5	87.3	n.d	n.d	n.d	1.0	95.2
28	n.d	n.d	5.7	n.d	0.2	21.1	58.2	3.0	0.5	n.d	1.7	90.8
29	n.d	0.4	6.5	n.d	0.2	18.2	61.3	2.9	0.5	n.d	2.2	91.8
30	n.d	n.d	2.8	n.d	0.8	4.1	73.7	1.1	n.d	n.d	4.8	87.7
31	1.4	n.d	6.5	n.d	n.d	1.7	74.4	2.0	n.d	0.2	n.d	86.5
32	n.d	n.d	7.5	n.d	0.2	2.5	74.7	1.2	n.d	n.d	n.d	85.7
33	n.d	n.d	5.2	n.d	n.d	n.d	73.8	1.2	n.d	n.d	n.d	83.0
34	1.2	n.d	6.5	n.d	n.d	1.4	74.7	1.4	n.d	n.d	n.d	85.6
35	n.d	n.d	6.7	n.d	0.4	2.9	67.1	1.5	n.d	n.d	n.d	79.7
36	n.d	0.5	4.2	n.d	n.d	0.7	79.5	1.3	n.d	n.d	n.d	86.1

**Table 2.** Major element (wt%) and selected trace element (ppm) content with the resulting Mn/Fe ratios of the Mn ore samples.

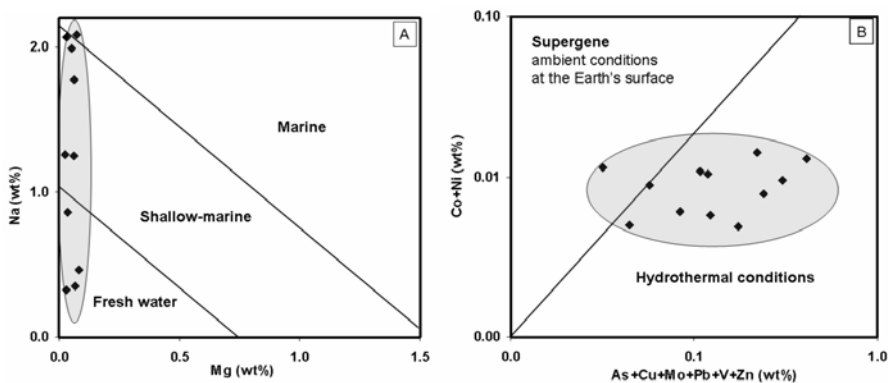
Sample	1	2	3	4	5	6	7	8	9	10	11	12
(wt %)												
<b>Mn</b>	4.88	9.57	4.49	3.76	17.08	25.75	18.02	1.18	17.91	25.83	10.20	10.97
<b>Fe</b>	2.68	3.06	5.33	1.57	14.70	4.84	4.79	1.64	1.93	3.97	1.82	7.77
<b>Na</b>	4.30	2.07	1.26	1.78	2.09	1.25	3.02	0.87	1.99	0.35	0.46	0.33
<b>K</b>	0.53	0.35	0.24	0.31	0.50	0.17	0.75	0.13	0.29	0.28	0.14	0.21
<b>Al</b>	2.29	0.91	0.56	0.79	0.85	0.50	1.13	0.23	0.22	0.24	0.23	0.12
(ppm)												
<b>Zn</b>	860	651	1045	336	2860	2835	1962	202	488	1047	657	1796
<b>Pb</b>	296	375	136	69	1132	174	189	87	60	626	121	447
<b>Ni</b>	57	79	39	30	70	48	47	77	50	20	30	49
<b>Co</b>	48	30	19	20	60	48	94	38	40	29	30	29
<b>Cu</b>	29	49	39	40	119	29	75	29	30	78	61	136
<b>Cr</b>	96	108	194	59	447	183	160	58	70	127	71	252
<b>Mg</b>	783	335	261	633	715	646	547	366	507	695	848	330
<b>Ca</b>	7727	740	823	742	1549	1080	3745	558	617	1321	646	553
<b>Ba</b>	2259	2395	1983	1294	3780	1995	1894	1629	1450	2458	1970	2109

The relatively high amount of some elements, such as Pb, Zn, Cu, and Ba, provides evidence of the genetic role of hydrothermal solutions (Nicholson, 1992). Some of the elements permit differentiation between manganese oxides formed in fresh water, shallow-marine and marine environment, whereas others distinguish between manganese oxides formed under surface conditions and those precipitated at deeper levels (Nicholson, 1992). For the manganese deposit described in this paper, the oxides must have precipitated under hydrothermal conditions (Fig. 4B), from fluids with a marine component, possibly in a sub-seafloor environment (Fig. 4A).

Hydrothermal enrichment of Zn in manganese deposits has been mentioned from the Galapagos area (Moore and Vogt, 1976, Cronan, 1986, Rogers, 1996), the Mariana-Bonin arc (Hein et al., 2008) and the Tonga-Kermadec Ridge and Lau Basin (Rogers et al., 2001). According to Cronan (1986), Hein et al. (2000), and Rogers et al. (2001), Zn enrichment in manganese deposits may represent the proximity of sulfide deposits that may be located near or below the Zn rich zones.

In terms of the geological, geochemical and mineralogical characteristics, Kimolos shares similarities with volcanic-hosted epithermal systems (Hedenquist et al., 2000). This is a rare example of a vein-type epithermal-style hydrothermal manganese deposit formed in a submarine environment (Roy, 1968, 1997; Canet et al., 2005; Leal et al., 2008).





**Fig. 4:** A. Geochemical signature of manganese oxides from Korakies. A. Marine signature. B. Hydrothermal signature (after Nicholson, 1992).

## 8. Conclusions

As a result of the present study the following conclusions can be drawn:

1. Base metal-rich manganese oxide mineralization is hosted by a possibly subaqueous monomictic andesitic volcanoclastic lithic breccia, possibly an andesitic in situ hyaloclastite, and shallow-marine or subaerial epiclastic conglomerates, in the Korakies area in NE Kimolos. The deposit has been exploited in the past. Exploitation was centered in a main vein that fills extensional NNE-SSW tectonic lineaments; it has maximum width of 5-6 m, extends for ~250 m, and has unknown depth. The distribution of the mineralized structures coincides with Pliocene NE-SW regional tectonic lineaments, along which the volcanic centres of the island were developed.
2. The vein displays complex and multiepisodic filling with textures characteristic of epithermal open-space precipitation such as crustiform banding, asymmetric banding, vugs, hydrothermal breccias, and cockade and comb textures. The Mn ore forms part of these textures. Ore formation occurred by hydrothermal solutions at shallow depths and at low temperatures.
3. Hydrothermal alteration associated with the mineralization is weak and confined to less than 2 m around the vein, and includes silicification (quartz-adularia and quartz-illite) and propylitization (chlorite-illite). Metallic ore minerals include basically primary pyrolusite and oxides of the isostructural series hollandite-cryptomelane-coronadite, and supergene goethite and hematite. Gangue minerals include quartz, chalcedony, jasper, barite and adularia.
4. The ore contains up to 25.83 % Mn, 14.7 % Fe, 2860 ppm Zn, 1132 ppm Pb, 136 ppm Cu, and 3780 ppm Ba. In addition, concentration of 4.3 % Na, 0.75 % K, 0.77% Ca and 0.09 % Mg has been measured.
5. Seawater together with subordinate contribution of meteoric sources could have been the source(s) for the mineralizing fluids, which formed the deposit in a subseafloor environment.
6. All geological, geochemical and mineralogical evidence compiled in this paper indicate that the Mn deposit of Korakies is similar to volcanic-hosted low-sulfidation epithermal ore deposits, deposited by neutral pH fluids. This is a rare example of a vein-type epithermal-style hydrothermal manganese deposit formed in a marine environment (Canet et al., 2005, Leal et al., 2008).

## 9. Acknowledgements

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