Fluid evolution in Tertiary magmatic-hydrothermal ore systems at the Rhodope metallogenic province, NE Greece. A review.

Vasilios Melfos1,2 and Panagiotis Voudouris2

1 Aristotle University of Thessaloniki, Department of Mineralogy, Petrology and Economic Geology, Thessaloniki, 54124, Greece
2 University of Athens, Department of Mineralogy-Petrology, Athens 15784, Greece

Keywords: Fluid inclusions, metallogeny, Rhodope, intrusion related systems, porphyry systems, epithermal mineralizations

ABSTRACT

Characterization of various fluid parameters in magmatic-hydrothermal ore mineralizations is potentially essential for interpretation of the conditions of formation and therefore for mineral exploration. Fluid inclusions can provide a useful and promising tool in the research of the ore forming processes in these systems. This review focuses on the nature, composition and origin of magmatic-hydrothermal ore forming fluids involved in the formation of representative Tertiary ore deposits at the Rhodope metallogenic province in NE Greece. These deposits are spatially related to Tertiary magmatism in NE Greece. Case studies are presented here and include an intrusion-hosted sheeted vein system (Kavala), a Au-rich carbonate replacement and quartz-vein mineralization (Asimotrypes), mineralized veins in Eptadendro-Rachi and Thasos island (Kapsalina and Panagia), porphyry Cu-Mo-Re-Au deposits in Pagoni Rachi and Maronia and epithermal Au-Ag mineralizations in Perama and Loutros. Hydrothermal fluids rich in CO2 together with elevated Au and Te content are common and occur at the Kavala intrusion hosted sheeted vein system, at the Asimotrypes Au-rich carbonate replacement mineralization and at the Panagia (Thasos) vein system. We classify all these ore mineralizations as intrusion-related gold systems (IRGS). Transport and precipitation of metals including Au and Te is favoured when CO2 is present. Precipitation of the ore mineralization takes place due to the immiscibility of the carbonic and the aqueous fluids which have a magmatic origin with the contribution of meteoric water. Cooling of magmatic hydrothermal fluids and dilution with meteoric water is a common cause for ore mineral formation in the vein mineralizations of Eptadendro/Rachi and Kapsalina Thasos. At the Pagoni Rachi and Maronia porphyry deposits, boiling and the high proportion of the vapour phase are the most essential fluid processes which affected ore formation. The epithermal veins overprinting the Pagoni Rachi and the Maronia porphyry systems and the HS-IS epithermal system in Perama Hill and the IS epithermal mineralization in Loutros are characterized by low to moderate temperatures and low to moderate salinities. Cooling and dilution of the ore fluids are the main process for gold precipitation. We conclude that the different fluid parameters and microthermometric data indicate a variety of fluid origin conditions and sources which can affect the strategy for exploration and prospecting for gold, rare and critical metals.

1. INTRODUCTION

Large-scale magmatic-hydrothermal fluid circulation in the Earth’s crust is derived from intermediate to felsic hydrous magmas mainly at convergent plate margins, providing endogenic heat and mass transfer for the formation of ore deposits including base, precious and rare metals (LEHMANN et al., 2000; BLUNDELL et al., 2005; SIMMONS & BROWN, 2006; BORTNIKOV, 2006; HEINRICH, 2007). These systems involve large amounts of volatiles including magmatic, meteoric or metamorphic water in aqueous or vapour form, CO2, NaCl, KCl, CaCl2, MgCl2, H2S, ±CH4, ±N2, as well as metals which originate from the magmas or the leaching processes from the rocks penetrated by the fluids through fluid-wall-rock interactions.

The detection of the various fluid characteristics is potentially important for interpretation of the formation of numerous ore deposit types and consequently for mineral exploration (BAKER & LANG, 2001; WILKINSON, 2001; HEINRICH, 2005; CHANG et al., 2011; KESLER et al., 2013). For example boiling has been a valuable guide in gold and copper exploration in porphyry systems. It is useful therefore to know how the volatile phases originate, how they dissolve and transport metals, which is the role of temperature and pressure and under what conditions the metallic minerals are deposited.

A useful fluid parameter which can be achieved by the fluid inclusions study, especially in the systems associated with Au and other sidirophile elements enrichment, is the CO2 content, because there is a strong correlation between CO2 content and gold mineralization, such as in the deep-seated intrusion-related systems (LOWENSTERN, 2001; BAKER 2002). In general, fluid inclusions with elevated homogenization temperatures and high salinities indicate magmatic productive fluids, while fluid inclusions with low ho-
mogenization temperature and low salinity show incorporation of meteoric waters (WILKINSON, 2001; KESLER et al., 2013). It is evident that homogenization temperatures, fluid salinity, density and pressure, can be useful in the determination of the fluid sources, the physical and chemical processes of transport and deposition of metals and also to explain the chemical environments of ore deposition and the possible zonation patterns of some ore systems.

This review paper summarizes the nature, composition and origin of magmatic-hydrothermal ore forming fluids which were involved in the formation of representative Tertiary ore deposits at the Rhodope metallogenic province in NE Greece (Fig. 1) which are developed within or around magmatic intrusive or extrusive rocks. We explore the evolution of high temperature and high salinity magmatic brines or CO$_2$-rich fluids, as well high temperature and low salinity magmatic vapours, to low temperature and low salinity meteoric fluids which were involved in the formation of these ore mineralizations.

This paper is partly motivated by microthermometric measurements of fluid inclusions which have been published in previous works of several researchers (VAVELIDIS et al., 1995; ELIOPOULOS, 2000; MELFOS et al., 2002; VAVELIDIS & MELFOS, 2004; MELFOS et al., 2008, 2010; FORNADEL et al., 2011; ELIOPOULOS & KILIAS, 2011; VAVELIDIS & MELFOS, 2014; MELFOS et al., 2010; FORNADEL et al., 2011; ELIOPOULOS & KILIAS, 2011; VOUDOURIS et al., 2011, 2013b) and include a reduced intrusion-related system (Kavala), a Au-rich carbonate replacement mineralization (Asimotyres), mineralized veins in Eptadendro-Rachi and Thasos island (Kapsalina and Panagia), porphyry Cu±Au±Mo deposits in Pagoni Rachi and Maronia and epithermal mineralizations in Perama and Loutros (Fig. 1). The purpose of this work is to show that detailed exploration for the discovery of new ore deposits requires a thorough fluid inclusion study which will interpret fluid phase relations and evaluate fluid characteristics in view of the ore forming conditions.

2. GEOLOGICAL SETTING OF TERTIARY ORE DEPOSITS

Greece is traditionally a mining country with numerous ore deposits which include base (Cr, Ni, Mn, Al, Fe, Pb, Zn, Cu), precious (Au, Ag, Platinum Group Elements-PGE), rare (Bi, Te, Mo, Re, Ti) & critical (Mg, Sb, W, Co, Ga, Ge, In, REE) metals (MELFOS & VOUDOURIS, 2012). Mining dates back to prehistoric times focusing initially on gold, copper and iron mineralizations. Later, in the Classical and Hellenistic times, ore exploitation became more intense especially for Au, Ag, Cu and Fe. Famous ancient mining centres operated in Milos and Sifnos Islands (Aegean Sea), in Lavrio (Attica), in Chalkidiki, Palea Kavala, Pangaeon Mt and Thasos Island (N. Greece), where a great number of underground galleries and metallurgical furnaces are still preserved (PERNICKA et al., 1981; GEORGAKOPOULOU et al., 2011). Several metals, such as Ag, Au, Fe, Cu, were also produced during the Byzantine and Ottoman periods (NERANTZIS, 2009). Since the beginning of the 20th century a great number of ore deposits have been targeted for exploration and exploitation of Cr, Ni, Al, Mn, Fe, Pb, Zn, Cu, Ag and Au. Greece is the most important producing country in the EU for Ni-Fe (from laterites) and Al.

![Simplified geological map of the Rhodope metallogenic province, NE Greece, showing the studied mineralizations. Kavala (Kv) intrusion-related system, Asimotyres (As) Au-rich carbonate replacement mineralization, Eptadendro-Rachi (ER) mineralized veins, Kapsalina (Kp) and Panagia (Pn) Cu-Bi-Ag±Te vein systems in Thasos island, Pagoni Rachi (PR) and Maronia (Mr) Cu-Mo-Re-Au porphyry deposits, Perama Hill (PH) and Loutros (Lt) epithermal mineralizations.](image-url)
The geological setting of the Hellenides, part of the Alpine-Himalayan orogeny, comprises numerous geotectonic terrains, which resulted as a consequence of the complicated geodynamic events in the region of the Palaeoore and Neotethys Oceans, including ocean rifting, spreading, subduction, collision, post-collision and extensive subduction and post-subduction magmatism (PE-PIPER & PIPER, 2002). After closure of the Neotethys Ocean, syn-orogenic exhumation of HP-LT rocks occurred during the Late Cretaceous–Eocene, before an acceleration of slab retreat changed the subduction regime and caused collapse of the Hellenic mountain chain and thinning of the Aegean Sea from Oligocene to the present (JOLIVET & BRUN, 2010). During this post-orogenic episode large scale detachments formed which exhumed metamorphic core complexes in a back-arc setting. Tertiary to Quaternary calc-alkaline to alkaline magmatism in the Aegean region occurred mostly in a post-collisional setting behind the active Hellenic subduction zone (PE-PIPER & PIPER, 2001). Lithospheric delamination, roll-back and/or break-off of the subducting slab were the principal mechanisms resulting in magma generation, which was triggered by asthenospheric mantle upwelling (JO-LIVET & BRUN, 2010).

In this geotectonic regime, large ore deposits of economic importance were formed during the Tertiary, in relation to magmatism and the evolution of extensive magmatic-hydrothermal systems. The ore types include reduced intrusion-related, skarn, porphyry, high-, intermediate- and low-sulfidation epithermal, carbonate-hosted replacement Pb-Zn and polymetallic vein deposits (MELFOS et al., 2002; VOUDOURIS & ALFIERIS, 2005; VOUDOURIS, 2006; BONSALL et al., 2011; MELFOS & VOUDOURIS, 2012; VOUDOURIS et al., 2013a). They are concentrated into four regional metallogenic provinces: 1) the Rhodope metallogenic province (RMP), 2) the Serbomacedonian metallogenic province (SMMP), 3) the Atticocycladic metallogenic province (ACMP) and 4) the Eastern Aegean metallogenic province (EAMP). These regions are the most promising targets for the near future exploration and exploitation of precious, rare and critical metals in Greece.

Tertiary basins and magmatic settings in the Rhodope massif are associated with significant skarn, polymetallic vein, porphyry and epithermal ore deposits, which are occasionally rich in Au and critical metals (MELFOS et al., 2002; VOUDOURIS, 2006; VOUDOURIS et al., 2009, 2011, 2013b), which constitute a part of the Rhodope metallogenic province. The magmatic-hydrothermal deposits of the Rhodope metallogenic province were formed during the final stage of the Tertiary orogenic collapse, and are related to the emplacement of hot asthenosphere into shallow crustal levels above a detached lithospheric plate which generated widespread magmatism (DE BOORDER et al., 1998; BLUNDELL et al., 2005).

The Oligocene-Miocene magmas were intruded to shallow depths along deep seated detachment faults, forming mafic to felsic mantle-derived plutonic, subvolcanic and volcanic rocks of calc-alkaline, high-K calc-alkaline and shoshonitic to ultrapotassic (lamprophyric) affinities (CHRISTOFIDES, 1996; MARCHEV et al., 2005).

3. FLUID INCLUSION CASE STUDIES
3.1. KAVALA INTRUSION-RELATED GOLD SYSTEM
The Kavala intrusion-related gold system (IRGS) is part of the Palea Kavala ore system (Fig. 1) which includes ~150 ore mineralizations occurring either as primary hypogene quartz-calcite-sulfide veins or as supergene oxidized ore bodies, associated with the ~21–22 Ma granodioritic Kavala pluton of I-type (FORNADEL et al., 2011). This pluton intruded marbles, schists, amphibolites and gneisses of the Palaeozoic Rhodope metamorphic core complex, along the regional E–W trending Kavala-Xanthi-Komotini fault (Fig. 1). The mineralization is considered to be part of an intrusion-related ore system in overlapping zones centred on the pluton. The marble hosts distal oxidized Fe-Mn (Pb±Zn±Ag) and Fe-Mn-Au ore bodies, whereas the gneiss and pluton contain proximal Fe-Cu-Au, Fe-Au-Au and Bi-Te-Au vein mineralization. These ore mineralizations were exploited in ancient times for gold and silver with extensive underground and surface mining operations (VAVELIDIS et al., 1996).

![Figure 2. Fluid inclusion types from Tertiary ore mineralizations in the Rhodope metallogenic province, NE Greece. a. Type L two phase aqueous liquid-vapour fluid inclusion coexisting with type C three phase inclusion which contain H2O-CO2 in Kavala sheeted vein system in the pluton; b. Type L two phase, liquid and vapour, fluid inclusion in a quartz vein at Eptadendro/Rachi; c. Type V vapour-rich two-phase aqueous inclusion coexisting with type L-S inclusion which contain three phases: a liquid, a vapour and a halite crystal in Pagoni Rachi porphyry system, indicating that they originate from a deep near-critical single phase fluid by phase separation; d. Type V two-phase vapour-rich inclusion with an opaque daughter crystal possibly chalcopyrite in Pagoni Rachi porphyry system; e. Type V vapour-rich two-phase aqueous inclusion coexisting with type L-S inclusion which contain three phases: a liquid, a vapour and a halite crystal in the Kavala Kranidi batholith; f. Type L aqueous liquid rich fluid inclusions which contain mainly NaCl in quartz at Perama Hill epithermal deposit.](image-url)
The SE-NW trending Bi-Te±Pb±Sb±Au quartz sheeted veins cut through the pluton and the subjacent metamorphic rocks in the Chalkero area, and contain pyrite, tetradymite, bismuthinite, cosalite and Sb-lillianite (FORNADEL et al., 2011). The veins consist of quartz, K-feldspar, albite and muscovite, altered to sericite and kaolinite. Fluid inclusions in these veins (Tab. 1) are rounded in shape, consistently distributed throughout the samples with a length of up to 60 μm (with the majority in the 25 to 35 μm range). They are classified into three types which coexist in each sample (MELFOS et al., 2008; FORNADEL et al., 2011) and therefore belong to the same primary fluid inclusion assemblage based on the criteria of GOLDSTEIN & REYNOLDS (1994).

Type L contains two phase aqueous liquid-vapour fluid inclusions (Fig. 2a) with NaCl and CaCl₂, and 20-30 volume % vapour. Homogenization temperatures range between 216° and 420°C, with a maximum at 380°C, and the salinities vary from 15.9 to 22.6 wt% NaCl equiv. Type C inclusions (Fig. 2a) are three phase: aqueous liquid, CO₂-liquid, CO₂-vapour approximately 50-70 volume %, with minor CH₄. They homogenize at 255°-414°C, with a peak at 320°C, demonstrating salinities from 5.5 to 11.2 wt% NaCl equiv. The third type V consists of two phase aqueous vapour-liquid fluid inclusions with a bubble of 90-95%, which homogenized to the vapour phase at Th between 210° and 323°C (max= 280°C).

The three fluid inclusion types are considered to represent coexisting immiscible carbonic and aqueous ore forming fluids (Fig. 3a). Isochores for fluid inclusions within the Kavala veins revealed a depth of formation, between 3 and 5 km (MELFOS et al., 2008; FORNADEL et al., 2011). Carbon dioxide played a significant role in the ore mineral precipitation in the Kavala veins.

3.2. ASIMOTRYPS CARBONATE REPLACEMENT AND VEIN TYPE Au DEPOSIT AT PANGEON MOUNTAIN

The Pangeon mountain includes numerous carbonate replacement, vein and shear-hosted Au-Ag ore mineralizations, which were mined intensively during ancient times for gold and silver (VAXEVANOPoulos et al., in press). They are hosted in the marbles and gneisses of the Rhodope massif as well as in the Pangeon granitoids.

One of the most important ore mineralizations occurs in Asimotrypes (Fig. 1) which is hosted in the Nikisiani pluton, part of the Pangeon granitoids, and in the marbles of the Rhodope massif (ELIOPOULOS, 2000; ELIOPOULOS & KIL-
Three types of primary fluid inclusions (Tab. 1) were distinguished by ELIOPoulos (2000) and coexist in each sample being in equilibrium with the sulfide minerals. Type L contains two phase aqueous liquid-vapour fluid inclusions with NaCl. The bubble occupies 10-20 volume % and the homogenization temperatures range from 135° to 256°C with a peak at 190°C and salinities from 5.62 to 9.05 wt% NaCl equiv. Type C inclusions are three phase and contain liquid, vapour, and a pour bubble (approximately 10 volume %). The fluid is dominated by NaCl with a minor contribution of KCl demonstrating a pour phase which occupies 20-30 % by volume. Two fluid inclusion assemblages at room temperature were identified MELFOS et al. (2010). Fluid inclusions are irregular to euhedral in shape with smooth boundaries reaching in size up to 20 μm.

Three types of fluid inclusions (Tab. 1) were distinguished by ELIOPoulos (2000) and coexist in each sample being in equilibrium with the sulfide minerals. Type L contains two phase aqueous liquid-vapour fluid inclusions with NaCl. The bubble occupies 10-20 volume % and the homogenization temperatures range from 135° to 256°C with a peak at 190°C and salinities from 5.62 to 9.05 wt% NaCl equiv. Type C inclusions are three phase and contain liquid, vapour, and a pour bubble (approximately 10 volume %). The fluid is dominated by NaCl with a minor contribution of KCl demonstrating a pour phase which occupies 20-30 % by volume. Two fluid inclusion assemblages at room temperature were identified MELFOS et al. (2010). Fluid inclusions are irregular to euhedral in shape with smooth boundaries reaching in size up to 20 μm.

The gold mineralization was formed by coexisting immiscible carbonic and aqueous fluids at pressures of 1.8 to 2.0 kbars under lithostatic conditions corresponding to depths of 6 to 9 km with the ore fluids at ~270°C (ELIOPoulos & KILIAS, 2011).

3.3. EPTADENDRO-RACHI VEIN TYPE Ag-Au ORE MINERALIZATION

The Cu-rich sulfide mineralization at the Eptadendro-Rachi ore district in Thrace, NE Greece, occurs at the contacts between granitoid intrusions (trodhjemites and pegmatites) of Upper Cretaceous to Early Tertiary age, and the metultramafic to slightly peraluminous post-collisional rocks and have a hornblende ⁴⁰Ar/³⁹Ar age of 21 to 22 Ma (ELEFTHERIADIS & KORONEOS, 2003).

The mineralization is associated with structurally controlled quartz veins, pods and lenses, and in gold-bearing replacement bodies, and is partly oxidized. It consists mainly of pyrite and arsenopyrite, with minor galena, sphalerite, chalcopyrite, pyrrhotite, tetrahedrite-tennantite, marcasite and gold (ELIOPOULOS, 2000; VAXEVANOPOULOS et al., in press). Fluid inclusions are irregular to euhedral in shape with smooth boundaries reaching in size up to 20 μm.

Microthermometric measurements were carried out on fluid inclusions of the mineralized quartz (VAVELIDIS et al., 1995). Based on their phase proportions at room temperature, three types of fluid inclusions were distinguished which belong to the same primary fluid inclusion assemblage based on the criteria of GOLDSTEIN & REYNOLDS (1994). Fluid inclusions of type L (up to 18 μm in length) consist of liquid rich two-
phase inclusions which contained mainly NaCl and the vapour ranges from 10 to 20 % in volume. The homogenization temperatures vary between 229° and 305°C, with a maximum at 390°C and the salinities are low, from 3.12 to 3.28 wt% NaCl equiv.

The type C fluid inclusions, with a length up to 20 μm, contain three phases dominated by aqueous liquid, CO₂-liquid and CO₂-vapour approximately 60-70 % of the total volume. A total fluid salinity of 3.7 to 6.5 wt% NaCl equiv was estimated. Final homogenization temperatures of this type of inclusions vary from 313 and 360°C, with a maximum at 330°C. A third type of fluid inclusion is made up by large irregularly shaped cavities, which contain pure CO₂. The vein type mineralizations in Panagia were formed by two coexisting immiscible carbonic and aqueous fluids and CO₂ played a significant role in the ore precipitation.

3.5. PAGONI RACHI PORPHYRY Cu-Mo-Re-Au DEPOSIT

The Pagoni Rachi Cu-Mo-Re-Au mineralization (Fig. 1) belongs to the Kirki mining district and is a telescoped porphyry-epithermal system which is spatially related to calc-alkaline granodiorite-tonalite porphyry stock of Oligocene age (VOUDOURIS et al., 2013b). Alteration types and vein relationships suggest that mineralization can be divided into four paragenetic stages, which are, from early to late: (1) sodic/potassic-calcic alteration with quartz- and magnetite-bearing veins (A- and M-type) and distal propylitic alteration, (2) sodic/potassic alteration with quartz-pyrite-chalcopyrite-molybdenite veins (B-type), (3) sericitic alteration with “transitional” porphyry to epithermal pyrite-chalcopyrite-molybdenite veins (D-type), and (4) argillic alteration with quartz-calcite base metal and precious metal rich veins (E-type) with epithermal affinity. The mineralization contains molybdenite with up to 4.7 wt% Re and the rare mineral rheniite (VOUDOURIS et al., 2009, 2013b).

Fluid inclusions were studied in the various vein types (A, B, and E) and are in equilibrium with the sulfide minerals and therefore represent the ore forming fluids (VOUDOURIS et al., 2013b). Their maximum size is 30 μm (mainly between 7 and 15 μm). They are isolated or arranged along quartz and calcite growth zones and are considered to have a primary origin based on the criteria of BODNAR (2003).

Three types of primary fluid inclusions were identified (VOUDOURIS et al., 2013b). Type L consists of two-phase aqueous inclusions with a vapour bubble occupying 10 to 20 volume %, which homogenize to a liquid phase. Type L-S inclusions contain three phases: a liquid, a vapour (10 to 20 volume %) and a halite crystal (Fig. 2c). Type V inclusions are two-phase vapour-rich (~80 volume %) aqueous inclusions (Fig. 2c,d). They homogenize to the vapour phase. They commonly contain an accidental-trapped opaque crystal (Fig. 2d) possibly chalcopyrite (VOUDOURIS et al., 2013b). Three distinct primary fluid inclusion assemblages (FIAs) were identified (VOUDOURIS et al., 2009, 2013b). The first two assemblages contain coexisting L, L-S and V inclusion types in A- and B-veins. The third assemblage includes late type L inclusions in the epithermal E-veins.

A- and B-veins were deposited at temperatures ranging between 360 and 510 °C and at pressures up to 690 bars (<2 km depth) for A-veins and up to 360 bars (<1.5 km depth) for B-veins, from boiling hydrothermal fluids. This process produced coexisting low to moderately saline (1.7-10.7 wt% NaCl equiv) and highly saline (36~74 wt% NaCl equiv) fluids (Fig. 3c). The fluid inclusions in the A- and B-type veins indicate that vapour was the predominant ore-forming fluid. Subsequent dilution of the moderately saline fluid resulted in a lower temperature (210–340 °C), less saline (1.4–2.9 wt% NaCl equiv) mineralizing fluid in late epithermal E-veins (Fig. 3c, VOUDOURIS et al., 2013b).

3.6. MARONIA PORPHYRY Cu-Mo-Re-Au MINERALIZATION

The Maronia Cu-Mo-Re-Au mineralization (Fig. 1) is hosted by a porphyritic microgranite (MELFOS et al., 2002). Three hydrothermal alteration zones have been recognized: an argillic, a sericitic and a propylitic zone. Additionally, three highly silicified zones crop out at the microgranite. Chalcopyrite-pyrite-molybdenite mineralization is located mainly in the silicified zones and is associated with areas of sericitic and propylitic alteration. The molybdenite revealed unusually high and variable rhenium concentrations (Re from 0.12 to 4.21 wt%).

Fluid inclusions which were studied in the ore-related quartz (MELFOS et al., 2002) have rounded to elongated or negative-crystal shapes and are isolated or arranged along quartz demonstrating a primary origin. Their size reaches up to 60 μm. Four types of fluid inclusions were identified (MELFOS et al., 2002). Type L contains 2-phase aqueous liquid-rich inclusions (vapour 20-35 volume %) and type L-S consists of 3-phases, an aqueous liquid phase, a vapour (20-40 volume %) and a halite (NaCl) crystal (Fig. 2e). Type V inclusions are two-phase vapour-rich (70-90 volume %), whereas type V-S are 3-phase aqueous vapour-rich (60-80 volume %) inclusions with a daughter mineral (halite). Fluid inclusions, mainly types L and L-S contain an accidentally-trapped opaque phase (pyrite and/or chalcopyrite?), which suggests that they represent the mineralizing fluid at the Maronia deposit.

Microthermometric results from the porphyry system showed that propylitic alteration and associated magnetite-pyrite-quartz veins are characterized by temperatures between 320 to 410°C, with a mean at 400°C. Trapping temperatures in the quartz-pyrite-chalcopyrite veins from sericitic alteration zone range from 360 to 420°C, with a peak at 380°C, and record the main temperature range of copper deposition under boiling conditions. Pyrite-molybdenite veins in sericitic zone were formed at slightly lower temperatures, from 250 to 360°C, with a maximum at 340°C. The ore-bearing fluids were dominated by NaCl, with salinities ranging from 7.0 to 16.4 wt% NaCl equiv in type L inclusions, from 28 to 55 wt% NaCl equiv in L-S fluid inclusions, from 5.5 to 14.0 wt% NaCl equiv in type V inclusions and from 38 to 51 wt% NaCl equiv in V-S inclusions (Fig. 3d). Trapping pressures of the ore-forming fluids were estimated to vary from 150 to 510 bar and boiling along with cooling of the fluids are
3.7. PERAMA HILL HS-IS EPITHERMAL Au-Ag DEPOSIT

An epithermal mineralization rich in Au, Ag, Te and Se occurs in Perama Hill at the Petrotata graben, NE Greece (Fig. 1). The ore mineralization is hosted in andesitic rocks and the overlying conglomerates and sandstones and exhibits characteristics of a high sulfidation system which was followed by deposition of an intermediate sulfidation mineralization (VOUDOURIS et al., 2011). It forms a sulfide rich vein-type in the andesite and a Au-rich stratabound oxidized type in the overlying sedimentary rocks. The ore mineral paragenesis includes a complex Pb-Zn-Bi-Se-Te-Sn-Au-Ag hypogene ore assemblage with native gold, native tellurium and electrum.

Fluid inclusions have negative-crystal or elongated isometric shapes with a maximum diameter of 30 μm, and were observed in quartz (Fig. 2f) and barite from the high sulfidation veins in andesite at depths of 88 m and 97 m (VOUDOURIS et al., 2011). They are aqueous liquid rich (Type L) with ~20 volume % vapour and contain mainly NaCl. They are isolated or arranged along the quartz or barite growth zones and consequently are defined as primary.

Homogenization temperatures in quartz vary from 211 to 331°C, with a maximum at 240°C, and salinities from 0.7 to 4.9 wt% NaCl equiv. In barite the inclusions display Th values of 222 to 256°C, with a maximum at 240°C, overlapping those of quartz, and very low salinities, between 0.8 and 2.3 wt% NaCl equiv. The mineralization was formed by cooling and dilution of the ore fluids which were possibly derived by condensation of magmatic vapour with limited mixing with meteoric water at 330 to 220°C (Fig. 3f).

KILIAS (2011) have shown that the main cause of ore precipitation is related gold system (IRGS). Panagia vein mineralization at Pangaiion mountain demonstrate the co-existence of two immiscible fluids, a low to moderate salinity (5.5 to 11.2 wt% NaCl equiv) CO₂-rich fluids are common and co-exist with moderate to high salinity fluids (salinities 15.9 to 22.6 wt% NaCl equiv). Methane is detectable in the CO₂-rich fluid inclusions. Kavala veins are characterized as a deep seated system (~3 to 5 km) which was formed by two co-existing high temperature (Th=300° and 400°C), moderate to high salinity, immiscible carbonic and aqueous fluids (FORNADEL et al., 2011).

Fluid inclusions from Asimotrypes carbonate replacement mineralization at Pangaiion mountain demonstrate the co-existence of two immiscible fluids, a low to moderate (2.61 to 7.31 wt% NaCl equiv) carbonic fluid and a moderate salinity (5.62 to 9.05 wt% NaCl equiv) aqueous fluid. ELIOPOULOS & KILIAS (2011) have shown that the main cause of ore precipitation should be attributed to unmixing of two fluids, a carbonic and an aqueous, at depths from 6 to 9 km at ~270°C. In the quartz veins at Panagia of Thasos the fluids are dominated by two immiscible fluids, an aqueous-carbonic and an aqueous fluid, which have low to moderate salinities (3.7 to 6.5 wt% NaCl) and trapping temperatures from 340° to 360°C at 500-700 kbar (VAVELIDIS et al., 1995). The presence of Te and the elevated temperatures of the Panagia vein system possibly show that it is located proximally to a magmatic rock, which however is not exposed on Thasos island but can only be assumed.

Fluid inclusions from Asimotrypes carbonate replacement mineralization at Pangaion mountain demonstrate the co-existence of two immiscible fluids, a low to moderate (2.61 to 7.31 wt% NaCl equiv) carbonic fluid and a moderate salinity (5.62 to 9.05 wt% NaCl equiv) aqueous fluid. ELIOPOULOS & KILIAS (2011) have shown that the main cause of ore precipitation should be attributed to unmixing of two fluids, a carbonic and an aqueous, at depths from 6 to 9 km at ~270°C. In the quartz veins at Panagia of Thasos the fluids are dominated by two immiscible fluids, an aqueous-carbonic and an aqueous fluid, which have low to moderate salinities (3.7 to 6.5 wt% NaCl) and trapping temperatures from 340° to 360°C at 500-700 kbar (VAVELIDIS et al., 1995). The presence of Te and the elevated temperatures of the Panagia vein system possibly show that it is located proximally to a magmatic rock, which however is not exposed on Thasos island but can only be assumed.

Kavala, Asimotrypes and Panagia ore systems show many similarities regarding their fluid characteristics. According to the intrusion related model presented by HART (2005) the Kavala sheeted veins in the pluton and the adjacent metamorphic rocks represent the most proximal system, whereas Asimotrypes carbonate replacement mineralization occurs at some distance from the Nikisiani pluton, being part of an intrusion-related gold system (IRGS). Panagia vein mineralization at
<table>
<thead>
<tr>
<th>Ore district</th>
<th>Metals</th>
<th>Ore style</th>
<th>Host mineral</th>
<th>FI types</th>
<th>Th°C (max)</th>
<th>Salinity wt% NaCl equiv</th>
<th>Fluid composition</th>
<th>Ore formation conditions</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kavala</td>
<td>Pb, Sb, Bi, Te, Au</td>
<td>Intrusion-hosted, sheeted veins</td>
<td>Quartz</td>
<td>L</td>
<td>216-420 (380)</td>
<td>15.9 - 22.6</td>
<td>NaCl and CaCl</td>
<td>Coexisting immiscible carbonic and aqueous fluids at a depth of 3-5 km. CO₂ played a significant role in the ore precipitation</td>
<td>MELFOS et al. (2008); FORNADEL et al. (2011)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>C</td>
<td>255-414 (320)</td>
<td>5.5 - 11.2</td>
<td>H₂O-CO₂, minor CH₄</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>V</td>
<td>210-323 (280)</td>
<td>-</td>
<td>NaCl and CaCl</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Asimotrypes</td>
<td>Fe, As, Pb, Zn, Cu, Sb, Au</td>
<td>Carbonate replacement style, part of an IRGS</td>
<td>Quartz</td>
<td>L</td>
<td>135-256 (190)</td>
<td>5.62 - 9.05</td>
<td>NaCl</td>
<td>Coexisting immiscible carbonic and aqueous fluids at a depth of 6-9 km at ~270°C. CO₂ played a significant role in the ore precipitation</td>
<td>ELIOPOULOS (2000); ELIOPOULOS &amp; KILIAS (2011)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>C</td>
<td>275-340 (320)</td>
<td>2.61 - 7.31</td>
<td>H₂O-CO₂</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>V</td>
<td>-</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Thasos (Kapsalina)</td>
<td>Cu, Fe, As, Sb, Bi, Ni, Ag, Au</td>
<td>Vein style</td>
<td>Quartz</td>
<td>L</td>
<td>300-380 (330)</td>
<td>1.6 - 7.2</td>
<td>NaCl</td>
<td>Cooling and dilution of fluids with a magmatic origin mixed with meteoric water</td>
<td>MELFOS et al. (2010)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>L</td>
<td>210-260 (240)</td>
<td>3.2 - 6.3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Thasos (Panagia)</td>
<td>Cu, Fe, As, Sb, Bi, Ni, Te, Ag, Au</td>
<td>Vein style, part of an intrusion related system</td>
<td>Quartz</td>
<td>L</td>
<td>229-305 (290)</td>
<td>3.12 - 3.28</td>
<td>NaCl</td>
<td>Coexisting immiscible carbonic and aqueous fluids. CO₂ played a significant role in the ore precipitation</td>
<td>VAVELIDIS &amp; MELFOS (2004)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>C</td>
<td>313-360 (330)</td>
<td>3.7 - 6.5</td>
<td>H₂O-CO₂</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pagoni Rachi</td>
<td>Cu, Fe, Mo, As, Sb, Pb, Zn, Bi, Re, Te, Ag, Au</td>
<td>Vein style, early stage</td>
<td>Quartz</td>
<td>A</td>
<td>247-514 (450)</td>
<td>A: 2.4 – 10.7</td>
<td>NaCl</td>
<td>A- and B-type porphyry style veins were deposited at a depth &lt;2 km from a boiling hydrothermal fluid. Subsequent dilution of the moderately saline fluid resulted in the formation of a lower temperature and less saline fluid in the late epithermal E-veins</td>
<td>VOUDOURIS et al. (2013)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>B</td>
<td>210-513 (390)</td>
<td>B: 1.7 – 9.2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>L-S</td>
<td>204-600 (370)</td>
<td>A: 38.9 - &gt;74</td>
<td>NaCl</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>L-S</td>
<td>361-600 (390)</td>
<td>B: 36.4 - &gt;74</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>V</td>
<td>245-522 (390)</td>
<td>A: 4.9 – 10.4</td>
<td>NaCl</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>V</td>
<td>276-494 (400)</td>
<td>B: 4.6 – 9.2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maronia</td>
<td>Cu, Fe, Mo, As, Sb, Pb, Bi, Re, Au</td>
<td>Porphyry Cu-Mo-Re-Au style (A, B veins)</td>
<td>Quartz</td>
<td>L</td>
<td>200-580 (365)</td>
<td>7 – 16</td>
<td>NaCl</td>
<td>Boiling and cooling of the fluids</td>
<td>MELFOS et al. (2002)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>L-S</td>
<td>210-515 (331)</td>
<td>28 – 55</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>V</td>
<td>275-590 (393)</td>
<td>5 – 14</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>V-S</td>
<td>307-455 (381)</td>
<td>38 – 51</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Perama Hill</td>
<td>Pb, Zn, Bi, Se, Te, Sn, Ag, Au</td>
<td>HS-IS epithermal style</td>
<td>Quartz</td>
<td>L</td>
<td>211-331 (240)</td>
<td>0.7 - 4.9</td>
<td>NaCl</td>
<td>Cooling and dilution of fluids with a magmatic origin mixed with meteoric water</td>
<td>VOUDOURIS et al. (2011)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Barite</td>
<td>222-256 (240)</td>
<td>0.8 – 2.3</td>
<td>NaCl</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Loutros Hill</td>
<td>Fe, Pb, As, Ag</td>
<td>IS epithermal style</td>
<td>Barite</td>
<td>L</td>
<td>220-329 (310)</td>
<td>8.65 – 16.14</td>
<td>NaCl</td>
<td>Cooling of a magmatic ore fluid possibly mixed with meteoric water</td>
<td>This study</td>
</tr>
</tbody>
</table>
Thasos island also demonstrates features which are similar to an IGGS, although there is no plutonic rock occurrence on the island.

In these three systems aqueous-carbonic and aqueous fluids coexist in the same samples, demonstrating that carbon dioxide played a significant role in ore formation. It has been emphasized that $\text{CO}_2$ is the medium which can induce transport and precipitation of some metals, including gold, and aids the process of ore deposition by phase separation in magmatic-hydrothermal systems (LANG & BAKER, 2001; BAKER, 2002). This process is common in many intrusion related gold systems and has been cited as direct evidence for a magmatic fluid source (BAKER & LANG, 2001), although decarboxylation reactions in wall-rock and dissolution of $\text{CO}_2$ from crustal hydrothermal fluids may be also encountered (LOWENSTERN, 2001).

Vein type ore mineralizations in the Rhodope metallogenic province are fault-controlled vein structures. They show similar formation conditions (temperatures-salinities), but relatively different fluid chemical compositions. In the Eptadendro-Rachi vein system the mineralization was caused by mixing between ascending magmatic fluids and downward migrating dilute meteoric water during cooling at homogenization temperatures from 380° to 210°C. These fluids with low to moderate salinities (1.6-7.2 wt% NaCl equiv) probably show an association with a granitoid body at depth, of Tertiary age, which is not exposed at the surface. The mineralized veins in Kapasalina of Thasos island were formed under similar conditions by low to moderate temperatures and low density fluids (Th=120-280°C, salinities=1.0-3.3 wt% NaCl equiv) also by cooling of the hydrothermal fluids which were derived mainly by meteoric waters (VANELIDIS & MELFOS, 2004).

The porphyry Cu-Mo-Re-Au ore systems in Pagoni Rachi and Maronia demonstrate the contribution of the vapour phase and the highly saline fluids under boiling conditions for ore mineral precipitation. Boiling processes resulted in the generation of coexisting low to moderately saline and highly saline fluids at temperatures from 300 to 500°C and the formation of the different hydrothermal alteration and ore zones (MELFOS et al., 2002; VOUDOURIS et al., 2013b). Vapour and brine dominance in both porphyry systems indicate that they originate from a deep near-critical single phase fluid by decompression and subsequent phase separation. In this case a low-density magmatic vapour phase is the principal agent for metal transportation, according to the model proposed by HEINRICH (2005), WILLIAMS-JONES & HEINRICH (2005), HURTIG & WILLIAMS-JONES (2015).

The released magmatic derived vapour rich fluids can transport high concentrations of metals mainly Cu and Au. Rapid expansion of vapour at a shallow depth favours Cu, Fe, Mo, Re and Au saturation and precipitation of the related ore minerals. WILLIAMS-JONES & HEINRICH (2005) emphasize the role of vapour volatile and precipitation of the bulk of Cu, Au and Mo at temperatures between 320 and 425°C at a shallow depth of about 2 km. At this temperature in the presence of iron, $\text{SO}_2$ in the fluid phase is reduced to $\text{H}_2\text{S}$ and $\text{H}_2\text{SO}_3$, which favours the precipitation of sulfides, especially pyrite, chalcopyrite, bornite and molybdenite, and increases gold solubility. The same process decreases drastically the fluid pH and causes the generation of an increasingly acidic fluid, which is associated with the related hydrothermal alteration minerals, mainly sericite and after mixing with groundwater, kaolinite (HEINRICH, 2005).

The shallow epithermal veins in the Pagoni Rachi porphyry system and the epithermal overprinting identified in Maronia (MELFOS et al., 2002; VOUDOURIS et al., 2013b) were formed under lower temperature conditions due to cooling and dilution of the ore fluids, which were evolved from initial high temperature and low salinity fluids towards cooler and very low salinity hydrothermal fluids.

The Perama Hill HS-IS and the Loutros IS epithermal mineralizations show similar temperature conditions (220-330°C) but slightly different salinities. In Perama Hill the salinities range between 0.7 to 4.9 wt% NaCl equiv indicating condensation of a magmatic vapour in the hydrothermal fluid with limited mixing with a fluid that had a major meteoric water component. In Loutros the salinities are higher (8.65 to 16.14 wt% NaCl equiv) which show a significant magmatic contribution in the hydrothermal fluid with possibly a limited mixing with meteoric water. The epithermal mineralizations in Perama Hill and Loutros were formed by cooling and dilution of the ore fluids. These fluids may represent late stage magmatic fluids that probably mixed with meteoric or seawater fluids at shallow levels.

5. CONCLUSIONS

Fluid inclusion studies of Tertiary magmatic-hydrothermal ore mineralizations in the Rhodope metallogenic province, NE Greece, focus mainly on the chemical composition of the ore forming fluids and the formation conditions of the mineralizations, the role of the magmatic and meteoric fluids and their possible interaction for the metallogenesis, the immiscibility and boiling processes and the role of $\text{CO}_2$ in ore precipitation. The different fluid characteristics and microthermometric data indicate a variety of fluid origin conditions and sources which can affect the strategy for exploration and prospecting of ore deposits.

Carbon dioxide rich hydrothermal fluids and Au and Te mineralization is common and occurs at the Kavala intrusion hosted sheeted vein system, at the Asimotrypes Au-rich carbonate replacement mineralization and at the Panagia (Thasos) vein system. Transport and precipitation of metals like Au and Te is favoured when $\text{CO}_2$ is present. Precipitation of the ore mineralization takes place due to the immiscibility of the carbonic and the aqueous fluids resulting in their unnmixing. These fluids have a magmatic origin with the contribution of meteoric water and are derived from deep seated systems. Based on geological and mineralogical criteria and the fluid inclusion data, we classify all these three ore mineralizations in Kavala, Asimotrypes and Panagia of Thasos, as parts of IGGS.

Cooling of magmatic hydrothermal fluids and dilution with meteoric water is a common cause of ore mineral formation in vein mineralizations (Eptadendro/Rachi, Kapasalina Thasos). Boiling and the vapour phase predominance are the most important processes for metallogenesis in the porphyry deposits (Pagoni Rachi and Maronia). These features are common in
porphyry systems and the mining industry should focus on them during early stage exploration work.

The epithermal veins overprinting the Pagoni Rachi and the Maronia porphyry systems and the HS-IS epithermal system in Perama Hill and the IS epithermal mineralization in Loutros are characterized by low to moderate temperatures and low to moderate salinities. Cooling and dilution of the ore fluids are the main processes for gold precipitation.

Following these criteria of the fluid characteristics, exploration for gold, rare and critical metals could have encouraging results to promote the mining industry in the future by using fluid inclusion techniques.

Acknowledgment

The authors would like to thank the reviewer Ladislav PALINKAŠ and an Anonymous Reviewer for their helpful and constructive comments on the manuscript. Ms Alisa MARTEK is especially thanked for her editorial handling of the paper.

REFERENCES


