A review of major non-sulfide zinc deposits in Iran

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ABSTRACT

The numerous non-sulfide zinc ore deposits were the historical basis for the development of zinc mining in Iran. They include the Mehdiabad, Irankouh and Angouran world-class deposits, as well as the Zarigan and Haft-bar deposits. These deposits were formed by supergene oxidation of primary sulfide minerals during the complex interplay of tectonic uplift, karst development, changes in the level of the water table, and weathering. Zn (Pb) carbonates, Zn-hydrosilicates and associated hydrated phases directly replace the primary ore bodies or fill cavities along fractures related to uplift tectonics. Direct replacement of primary sulfides is accompanied by distal precipitation of zinc non-sulfide minerals in cavities or internal sediments filling. The mineralogy of the non-sulfide mineralization in all six deposits is generally complex and consists of smithsonite, hydrozincite, and hemimorphite as the main economic minerals, accompanied by iron and manganese oxy-hydroxides and residual clays. Commonly, non-sulfide minerals in these deposits consist of two types of ore: red zinc ore (RZO), rich in Zn, Fe, Pb-(As) and white zinc ore (WZO), typically with very high zinc grades but low concentrations of iron and lead. Typical minerals of the RZO are Fe-oxyhydroxides, goethite, hematite, hemimorphite, smithsonite and/or hydrozincite and cerussite. Common minerals of the WZO are smithsonite or hydrozincite and only minor amounts of Fe-oxyhydroxides and hemimorphite.

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1. Introduction

"Non-sulfide" is a term, which comprises a series of oxidized Zn-(Pb) ore minerals, mainly considered as related to weathering of Zn (Pb) sulfide concentrations. These deposits may result from the supergene evolution of primary sulfide deposits such as Mississippi Valley-type (MVT), carbonate replacement, sedimentary exhalative (SEDEX), Irish type, veins and less commonly skarns (Heyl and Bozion, 1962; Hitzman et al., 2003). They can also be due to hypogene processes: (1) structurally controlled replacement deposits (Appold and Monteiro, 2009; Borg, 2009; Slezk et al., 2014), (2) strataform lenses in highly metamorphosed terrains (Brugger et al., 2003; Peck et al., 2009) or (3) hydrothermal overprints of sulfide deposits (Boni et al., 2007; Boni, 2014). In the case of supergene processes, subtypes have been distinguished, including direct replacement deposits and wall-rock replacement deposits (Heyl and Bozion, 1962; Hitzman et al., 2003; Borg, 2015).

Direct replacement non-sulfide deposits, are characterized by local trapping of base metals liberated by sulfide dissolution with a very limited transport from the original protore zone. They usually display a ‘red ore’, that commonly consist of iron (hydr-) oxides (mainly goethite and hematite) associated with hemimorphite, smithsonite, hydrozincite and cerussite. Direct replacement non-sulfide deposits typically contain >20% Zn, >7% Fe and display high Pb ± As contents and commonly preserved sulfide minerals like galena, sphalerite, or pyrite (Reichert and Borg, 2008; Reichert, 2009).

Wall-rock replacement mineralization, also known as ‘white ore’ is characterized by the leaching and the transport of metals by percolating waters due to the groundwater/geochemical gradient (primary ore) (Heyl and Bozion, 1962; Hitzman et al., 2003). Wall-rock replacement deposits consist of smithsonite, hydrozincite and minor iron oxides, and contain <40% Zn, <7% Fe and very low concentrations of Pb (Boni et al., 2007; Reichert and Borg, 2008; Reichert, 2009).
Reichert, 2009; Boni and Mondillo, 2015). They may occur at the vicinity of the protore or up to several hundred metres away (Heyl and Bozion, 1962; Hitzman et al., 2003; Reichert and Borg, 2008; Reichert, 2009). As they are commonly richer in Zn and poorer in Pb and Fe (Reichert, 2009).

Despite the widespread occurrence of surficial zinc oxides in gossans and soils (e.g., Jacquat et al., 2009), economic non-sulfide deposits are far less common than sulfide deposits. Non-sulfide deposits have experienced a significant revival over the recent years, as a consequence of new developments in hydrometallurgical acid leaching, solvent extraction, and electrowinning techniques (Choulet et al., 2013). The non-sulfide Zn ore, called “calamine” by the miners includes a mixture of smithsonite, hemimorphite, hydrozincite locally associated with Fe-oxhydroxides and clays.

The economic value of zinc non-sulfide ores is strictly dependent on the nature of the ore and gangue minerals. Host-rock composition may have significantly influenced the mineralogy (and by extension processing) of ores in these deposits. Since the differences in dissolution rates of the zinc minerals may have strong implications for the production strategies and metallurgical requirements, a detailed mineralogical and petrographic study, in the early phases of an exploration process, is required for this kind of ores, even more than in the case of sulfide deposits.

Scientific research, resulting from the economic interest shown by many mining companies in this style of mineralization, has focused on several potentially economic “Non-sulfide” deposits (e.g., Skorpion, Namibia: Borg and Kärner, 2001; Borg et al., 2003; Shaimerden, Kazakhstan: Boland et al., 2003; Vazante, Brazil: Monteiro et al., 1999; Beltana, Australia: Muller, 1972; Brugger et al., 2001; Hitzman, 2001; Groves and Carman, 2003; Hitzman et al., 2003; Angouran, Iran: Daliran, 2003; Daliran and Borg, 2004; Reichert, 2007; Reichert and Borg, 2008; Daliran et al., 2009).

Iran possesses a large range of Zn—Pb deposits including SEDEX, Irish-type and MVT that occur in carbonate and siliciclastic rocks (Rajabi et al., 2012). There are more than 350 Zn—Pb deposits and occurrences in Iran, including world-class deposits such as Angouran, Mehdiabad and Iran (Borg, 2005; Rajabi et al., 2012) (Fig. 1). Only a few of them, however, have actually been explored.

![Figure 1. Distribution map of sediment hosted Zn—Pb deposits according to the type of ore deposits in the main tectonic elements of Iran. Al, Alborz zone; CIGS, Central Iranian geological and structural gradual zone; E, East Iran ranges; K, Kopet-Dagh; KR, Kermanshah Radiolarites subzone; KT, Khazar-Talesh-Ziveh structural zone; L, Lut block; M, Makran zone; Oph, ophiolite belts; PB, Posht-e-Badam block; SSZ, Sanandaj-Sirjan zone; T, Tabas block; TM, tertiary magmatic rocks; UDMA, Urumieh-Dokhtar magmatic arc; Y, Yazd block; Z, Zabol area; Za, Zagros ranges. Tectonic and structural map of Iran modified after Alavi (1996) and Aghanabati (1998, 2004).]
and exploited. The largest of these deposits is the Mehdiabad deposit. This study focuses on six ore deposits throughout the Iran (Table 1). These deposits include Angouran, Irankouh, Mehdiabad, Kuh-e-Surmeh, Zarigan and Haft-har (Fig. 1 and Table 1).

2. Methodology

The available information (as far as possible) on the Zn–Pb deposits of Iran has been incorporated into a sedimentary structural map of Iran and integrated into a comprehensive GIS database. Most of the data were derived from the inventories of the Geological Survey of Iran and from published and unpublished data of numerous source papers cited in the references. Because of the large number of nonsulfide Zn–Pb deposits and occurrences, and the lack of information about most of them, we tried to avoid unreliable occurrences and removed very small occurrences from the database. So, the authors have visited the six major nonsulfide Zn–Pb deposits and studied different geological characteristics of these deposits. Also we present field observations of macrostructures and try to decipher the relationships between the protore, the non-sulfide zone, and the host rocks.

3. Zinc–lead deposits in Iran

Most Zn–Pb deposits occur within continental blocks, which are key structures in the evolution of the Palaeo-Tethys and Neo-Tethys oceans. There is no information about the age of these deposits, but host rocks range from Devonian to Late Cretaceous. Iran is a part of the Tethyan Belt and, hence, the numerous deposits are related to the tectonic evolution of the Tethys Ocean. Although Iran is well known for porphyry and epithermal (Cu–Au) deposits (Aghazadeh et al., 2015), it also hosts a range of significant Zn–Pb deposits (Borg, 2005). They belong to the Tethyan Metallogenic Belt, similarly to those of China and Turkey (Reynolds and Large, 2010). Fig. 1 shows the distribution of the diversely aged deposits of Iran has been incorporated into a sedimentary structural map of Iran and integrated into a comprehensive GIS database. Because of the large number of nonsulfide Zn–Pb deposits and occurrences, and the lack of information about most of them, we tried to avoid unreliable occurrences and removed very small occurrences from the database. So, the authors have visited the six major nonsulfide Zn–Pb deposits and studied different geological characteristics of these deposits. Also we present field observations of macrostructures and try to decipher the relationships between the protore, the non-sulfide zone, and the host rocks.

3.1. Sedimentary exhalative massive sulfide or SEDEX deposits

The most important SEDEX Zn–Pb deposits of Iran occur in the NW, Central and SE parts of the Zarigan–Chahmir basin in the Poshteh-Badam and Yazd block (YB) and in the Malayer-Esfahan metallogenic belt (MEMB) (Fig. 1). The Koushk, Zarigan, Chahmir and Haft-har Zn–Pb deposits are the major ones (Rajabi et al., 2014). They commonly occur close to the intersection of syn-sedimentary faults (active throughout the sag-phase of the Zarigan–Chahmir basin evolution; Rajabi et al., 2012) and are hosted by sedimentary rocks deposited under anoxic conditions (Rajabi, 2012). Three major ore zone or facies are usually distinguished in the SEDEX Zn–Pb deposits in Iran (Rajabi, 2012; Rajabi et al., 2012): (1) massive sulfide ore or vent-complex, (2) stockwork or feeder zone, and (3) stratiform-bedded ore, which include exhalites and lenses of pyrite, sphalerite and galena.

3.2. Carbonate hosted (CH) Zn–Pb deposits

Based on the age of their host rocks, two major groups of CH Zn–Pb deposits can be distinguished and linked to different tectonic events that affected the Iranian Plate. The first group includes the Permian–Triassic-hosted MVT deposits mainly in the Central Alborz and Tabas-Posht e Badam metallogenic belts, and in the northeastern margin of the Sanandaj–Sirjan zone (SSZ) (Fig. 1). The Triassic carbonate rocks also host numerous F-rich deposits,
marked by stratabound mineralization and veins of fluorite, barite with minor galena and sphalerite. The second group includes the Zn–Pb deposits hosted in the Cretaceous series, located in the Malayer-Esfahan and Yazd-Anarak metallogenic belts (YAMB), and to a lesser extent, in the Central Iranian Geological and Structural (CIGS) transition zone and in the Central Alborz metallogenic belt (AMB) (Fig. 1).

4. Supergene non-sulfide zinc-lead deposits in Iran

Iran exposes numerous non-sulfide zinc deposits, and several centers for zinc smelting and refining were established near the mining sites. Furthermore, there are additional non-economic resources in mine dumps, flotation tailings, and smelter residues. Although these can be quite substantial in areas of ancient mining (e.g., Angouran and Mehdiabad), they will not be discussed further in this paper. Most of the Iranian examples of non-sulfide zinc mineralization belong to the group I (classification of Large, 2001) calamine deposits, hosted by carbonate rocks.

Non-sulfides deposits of Iran widespread in the different tectonic blocks of Iran. The most important metallogenic provinces for non-sulfide zinc mineralization are Central Iran, the Sanandaj-Sirjan zone, and the Alborz region (Fig. 1). The age of the host rocks ranges from Proterozoic to Tertiary. However, most of the host rocks are either Mesozoic or Cretaceous carbonates (Table 1). Only a few non-sulfide zinc deposits (Angouran, Irankouh, and Mehdiabad) are currently mined (Fig. 1). The giant Mehdiabad deposit is a special case, as both sulfide and non-sulfide ores are currently mined. In this paper we focus on the important non-sulfide deposits in Iran (Table 1).

4.1. Late Neoproterozoic–Early Paleozoic hosted non-sulfide deposits

4.1.1. The Angouran deposit

The Angouran Zn(–Pb–Ag) deposit is located in the western part of Zanjan Province, NW Iran (Fig. 1). This area belongs to the northwestern part of the Sanandaj-Sirjan Zone, a metamorphic belt related to the Zagros orogeny (Fig. 2).

The basement rocks at Angouran include imbricated thrust sheets of exhumed metamorphic rocks that consist of muscovite-, chlorite- and quartzite-schists with thin marble horizons and a discontinuous 50–100 m thick interval of meta-basic amphibolite schist (Daliran et al., 2013). Foliated pillow basalt and serpentinite bodies, which represent a mid-ocean ridge ophiolite assemblage are also present (Figs. 3 and 4) (Daliran et al., 2013). The footwall quartzite schist is equivalent to the Neoproterozoic Kahar Formation (Daliran et al., 2013). The U–Pb age of 511 Ma obtained by SHRIMP-dating on inherited cores of the zircon in pillow basalts indicates that the basalts probably formed during the Cambrian (Daliran et al., 2013). The 94 Ma U–Pb age for the rim of these zircon grains is interpreted as a metamorphic overgrowth during the Alpine orogeny (Daliran et al., 2009). The marble unit (Angouran Marble) was thrust onto the schists (Fig. 4) and contains discontinuous dolomitic horizons with supergene dissolution cavities, locally. The age of the marble unit is also Neoproterozoic to Early Cambrian based on the microfauna content (e.g. Siphogonuchites triangularis; Hamdi, 1995) (Fig. 4).

The host rocks at Angouran are part of a 70 km × 30 km block of stacked thrust sheets comprising marble, schists and ultrabasic basement rocks (Fig. 3) (Daliran et al., 2013). Footwall schists within the mine sequence are intensely deformed and comprise whithish to light green quartz-sericite schist and pale orange calcareous schist with intercalations of quartzite and subordinate muscovite-, chlorite-, epidote- and biotite-schist with thin marble bands (Fig. 4) (Daliran et al., 2009, 2013). Toward the north, the schist sequence is underlain by a discontinuous interval of 50–100 m of metamorphosed basic and ultrabasic rocks including pillow basalt, amphibolite, metabasalt and serpentinite. The hanging-wall consists of a 50–300 m thick marble unit (Angouran Marble) (Fig. 4) (Daliran et al., 2013). The marble is thick to medium bedded, light to medium gray with thin, discontinuous dark laminae (carbonaceous material) and subordinate dolomitic intercalations.

Figure 2. View of the Angouran open pit; the brown color correspond to the non-sulfide ore. Host rock of the non-sulfide ore is marble.
Recent mining (Fig. 3) began after the rediscovery of the Zn ore in cerussite-bearing gossans, in surface outcrops and dissolution cavities in the hanging-wall marble in 1922. The ore reserve in 2001 was 4 Mt sulfide ore (40.4% Zn, 1.9% Pb) and 18.23 Mt carbonate ore (28.1% Zn, 4.4% Pb, and 110 g/t Ag; Pride and Salehi, 2003).

The orebody exhibits a vertical zonation with (1) an underlying Zn-rich, hypogene, sulfide ore, and a mixed sulfide—carbonate ore at the bottom and, (2) a carbonate ore at the top. The deposit formed at a lithological boundary, which comprises a thrust contact between the footwall schists and the hanging-wall marbles (Figs. 3 and 4). The ore zone at Angouran displays a complex geometry. The hypogene sulfide deposit, which has been oxidized to form a significant non-sulfide orebody, mainly in the form of smithsonite, is located at the crest of an open anticline at the contact between Neoproterozoic to Cambrian footwall micaschists and hanging wall marbles (Boni et al., 2007). The dimensions of the mineralized zone are ca. 600 m in length (along the N–S axis) and 200 to 400 m in width. The orebodies are delimited by two

![Schematic regional geological map of the Angouran area, NW Iran (after Babakhani and Ghalamghash, 1990).](image-url)
major NNW–SSE and NW–SE trending faults and a third NE–SW fault trending (Fig. 3). The thickness of the zinc carbonate ore zone may reach up to 200 m; this ore zone occurs in the hanging-wall marbles and overlies a tabular sulfide orebody (Boni et al., 2007; Daliran et al., 2013; Boni and Mondillo, 2015). Smaller bodies of mixed sulfide and carbonate ore occur at the contact between sulfide and non-sulfide zones, as well as within the non-sulfide ores. Both ore types also fill a wide variety of breccias, especially along the three main fault zones that are laterally controlling the deposit.

Previous genetic models proposed for the Angouran sulfide deposit include a wide range of models (SEDEX and VMS) and ages (Proterozoic to Mesozoic) (e.g., Gazanfari, 1991; Ghorbani, 1999; Annels, 2005). These essentially syngenetic models were challenged by more recent studies that document an epigenetic model of formation for the sulfide ore (e.g., Daliran, 2003; Daliran and Borg, 2004; Gilg et al., 2006; Boni et al., 2007; Daliran et al., 2009). Hirayama (1968), however, recognized the unmetamorphosed nature of the sphalerite ore and concluded that the mineralization was post-metamorphic.

The mineralogy of the supergene and hypogene parts of the deposit mainly consists of smithsonite (Fig. 5), arsenopyrite, galena, sphalerite, pyrite, greenschist, Co-arsenopyrite, cerussite, pyromorphite, Zn-arsenates, Co–Ni-arsenates, Pb–Mn-(hydr) oxides, hemimorphite, litharge, several clay minerals, and apatite (Fig. 5) (Boni et al., 2007). The genesis of the Angouran Zn carbonate ores is

Figure 4. Generalized schematic columnar section of the Angouran area, with the main ore-bearing strata (modified after Daliran et al., 2013).
distinct from that of other “classical” non-sulfide Zn deposits that formed entirely by supergene processes, as demonstrated also by carbon and oxygen isotope geochemistry (Daliran, 2003; Daliran and Borg, 2004; Daliran et al., 2009; Boni and Mondillo, 2015). Therefore, non-sulfide mineralization at Angouran is interpreted to be the result of during two successive stages, following an early generation of hypogene sulfide minerals. The first, main stage of Zn carbonates (stage I carbonate ore) is associated with both preexisting and subordinate newly formed sulfides, whereas the second stage is characterized by supergene carbonates (Zn and minor Pb), associated with oxides and hydroxides (stage II carbonate ore) (Boni et al., 2007; Daliran et al., 2013; Boni and Mondillo, 2015).

### 4.1.2. Zarigan SEDEX-type deposit

The central lithotectonic domain of the Posht-e-Badam Block, within the Central Iranian microcontinent, contains the oldest (Late Neoproterozoic) basement of Iran (Rajabi et al., 2012). The Zarigan–Chahmir basin is located in the southern section of this lithotectonic domain (Fig. 1) and hosts abundant mineral deposits, especially iron oxide–apatite (IOA) deposits, Fe–Mn exhalative deposits, and sedimentary exhalative (SEDEX) Zn–Pb deposits (Rajabi et al., 2012). The economically most important SEDEX Zn–Pb deposits of this basin include the Koushk, the Chahmir and the Zarigan deposits (Rajabi et al., 2012).

The Zarigan Zn–Pb deposit is hosted within the upper part of the Early Cambrian Volcano-Sedimentary Sequence (ECVSS), in the northern part of the Zarigan–Chahmir basin (Fig. 6). Two sequences can be distinguished (Fig. 6): (I) a lower volcano-sedimentary (syn-rift) sequence well exposed in the Narian, Esfordi, Lakeh-Siyah and Zarigan areas, which comprises gray basic coarse-grained clastic rocks, various pyroclastic rocks and bimodal volcanic rocks; and (II) the upper sedimentary sequence (corresponding to the post-rift sag-phase), including calcareous shales, siltstones and carbonates with minor tuffaceous rocks. The intercalated calcareous black siltstone of the post-rift sequence hosts the Zarigan SEDEX Zn–Pb mineralization, typically in its middle part (Fig. 6A).

The Zarigan deposit differs from other SEDEX deposits in the basin by its intensive weathering and the development of an oxidized Pb(Zn) ore zone (Fig. 6). Ore minerals include cerussite, anglesite associated with abundant Fe-(hydr) oxides and minor pyrite, sphalerite and galena (Fig. 7 and Table 1). Therefore, red zinc ore (RZO) is dominant in the Zarigan SEDEX type deposit. Exploration of the sulfide ore is in early stages. Based on the previous mining activities (from 1957 to 1977), the primary sulfide ore reserve is estimated to be about 18–20 Mt, with 10 Mt of oxidized ore (Simiran Mining Co.). The host rocks at Zarigan include black siltstone and minor silty limestone (Fig. 6). In addition to some relict bands of sulfide minerals within the oxidized ore zone, laminated-disseminated sulfide minerals occur within the host rock (Fig. 7). Similar to the Lady Loretta deposit in Australia (Hancock and Purvis, 1990), zinc has been completely removed from the gossan cap and the oxidized zone (Figs. 6 and 7). The Zarigan deposit occurs as a single, wedge-shaped ore body within the host rock. The northern part of the deposit displays a strata-bound geometry, but toward the south, as thickness decreases, the mineralization becomes stratiform and concordant with the host sedimentary rocks. Based on crosscutting relationships, mineralogy and texture of sulfide and non-sulfide mineralization, three major ore facies types can be distinguished, from proximal to distal relative to the syn-sedimentary Zarigan Fault (Rajabi, 2012): (1) a complex stratiform oxide ore, formed by replacement, open space filling and vein and veinlets of cerussite, anglesite, goethite, hematite, pyrite and galena; (2) a bedded stratiform oxide ore, including laminated cerussite and hematite (relict textures of primary laminated sulfides) with disseminated pyrite and galena (Fig. 7A and B); and (3) a bedded stratiform sulfide ore, consisting of a banded body of...

![Figure 5. Images of supergene non-sulfide zinc minerals from the Angouran deposit. (A) Colloform smithsonite; (B) hemimorphite (Hem) crystals around a core of colloform smithsonite (Sm); (C) concretionary smithsonite; (D) crystalline hemimorphite; (E) botryoidal smithsonite filling vug; (F) white ore sample made of hydrozincite \([\text{Zn}_5(\text{CO}_3)_2(\text{OH})_6]\); (G) aggregate of goethite, hematite and smithsonite.](image-url)
cerussite, anglesite, smithsonite, hemimorphite, pyrite and sphalerite with disseminated galena. In addition, banded chert facies, consisting of chert beds with barite and disseminated pyrite, overlies these ore facies types. These ore facies types and deposit geometry, together with the presence of a syn-sedimentary fault adjacent to the deposit, have led us to classify the Zarigan deposit as a vent-proximal (Selwyn-type) SEDEX deposit, equivalent to the Koushk and the Chahmir deposits (Rajabi, 2012).

4.1.3. Haft-har SEDEX-type deposit

The Haft-har Zn–Pb deposit, located about 260 km to the NW of Mehdiahab deposit, is hosted within the middle part of the Cambrian series of the Yazd block in Central Iran (Fig. 8). The footwall rocks are gray shales with interbedded sandstone (litharenite) and sandy volcaniclastic rocks. The ore-bearing zone occurs within the Haft-har unit, which consists of carbonaceous, fine-grained black siltstone and shale with intercalated volcaniclastic sandstone (Fig. 8). Limestone and massive cherty dolomite of the
Figure 7. Images of supergene non-sulfide zinc samples in Zarigan deposit (Rajabi et al., 2012). (A and B) Hand sample of bedded non-sulfide ore, which have been completely oxidized to goethite, hematite and cerussite; (C) coarsely crystalline white-tan cerussite filling vugs in hematitic siltstone; (D and E) crystalline cerussite covering iron oxides.

Figure 8. Simplified geological map of the Haft-har area in the Yazd block, showing the location of Haft-har SEDEX deposits within this map.
Mila Formation with thrust fault cover the ore-bearing shale and siltstone in the Haft-har area (Fig. 9A).

The sulfide lens of the Haft-har deposit does not crop out at the surface; therefore, the geological setting of the deposit has been reconstructed from drill core logs and adits (Fig. 9). Sulfide mineralization occurs as a single wedge within the organic matter-rich siltstone and shale. The ore-bearing host rock is usually folded, whereas the more competent dolomite typically exhibits a brittle deformation resulting in extensive breccia zones cut by late calcite veins. The limestone of the Mila Formation in the vicinity of the ore body has developed a large karst system probably caused by sulfide oxidation in the vicinity. Evidence of remobilization includes the precipitation of secondary minerals along joints and fractures of the host rock (Fig. 9C). The potentially economic resource of Haft-har deposit mainly consists of non-sulfide Zn minerals, associated with abundant Fe-(hydr) oxides; this non-sulfide mineralogy results from weathering of presumed primary sulfide minerals (Fig. 9). The primary orebody is likely to have been deformed and oxidized during post-Cambrian episodes. Preliminary mineralogical studies have revealed that the main sulfide phases are sphalerite, pyrite and galena, and are accompanied by calcite, barite and quartz (Movahednia, 2014).

Non-sulfide mineralization at Haft-har mainly consists of Fe-(hydr) oxides, smithsonite and hemimorphite, with subordinate hydrozincite. Cerussite and anglesite also occur, generally associated with nodules and lenses of residual or supergene galena (Fig. 9). A complex association of Fe and Mn oxy-hydroxides, with a characteristic red-brown color (goethite, lepidocrocite, hematite), and residual clay minerals hosts the non-sulfide ore (Fig. 9). The ore grade of the non-sulfide zone is highly variable throughout the mining district, ranging from 8% to more than 30% in the areas where the alteration profile displays a total replacement of sulfide minerals by secondary carbonates. As the dominant sulfide ore is rich in pyrite, a red zinc ore was formed in this deposit. The non-sulfide ore is rich in Zn in the lower levels of the oxidation profile, whereas, near the surface (in the leaching zone), the ore grade is generally not economic. The mineralization is considered to be the result of in situ oxidation of the primary sulfide ore by meteoric fluids, with increased acidity owing to the dissolution of substantial amounts of pyrite. The dissolution was followed by a more or less total replacement of the sulfide minerals, as well as of parts of the host rocks, by secondary minerals. Subsequent remobilization and further deposition of secondary minerals within dissolution vugs and karst cavities locally represent the last stages of the deposit evolution.

4.1.4. Kuh-e-Surmeh deposit

The Kuh-e-Surmeh Zn–Pb deposit is the only mineralization occurring within the southern part of the Zagros ranges (Fig. 1) in the Firoozabad region. The total Zn:Pb ratio of the deposit is 3:1, with average ore grades around 19% Zn and 7% Pb; the orebody contains approximately 990,000 tons of pre-mining reserves (Liaghat et al., 2000). The stratigraphic sequence of the mine area (Fig. 10) begins with more than 1000 m of Ordovician sediments of the Zard-Kuh Formation, which are silty, micaceous, olive-green shales and associated fine-grained dark-brown sandstone

Figure 9. (A) General view of the Haft-har deposit. (B) Widespread non-sulfide ore (white) and Fe-oxide ore (red) forming the wall of underground tunnel. (C) Shale and siltstone host rocks overlain by iron oxide and a non-sulfide mineralization. (D) Crystalline white cerussite and anglesite coating the wall of tunnels. (E) Cerussite and associated to an iron oxidized facies.
(Ghavidel, 1973; Liaghat et al., 2000). These strata are overlain by about 85 m of Devonian and Early Permian Faraghan sediments composed mainly of purple and dark-gray shale with sandstone intercalations (Fig. 10). The Faraghan Formation is overlain in turn by about 120 m of Upper Permian Dalan Formation consisting of laminated limestone, dolostone, carbonaceous brown-red sandstone and, in some cases, gypsum and marl. The Dalan Formation is divided into two parts, the Early and Upper Dalan, by thin gypsum layers of the Nar Member. The Early Dalan Formation hosts the Kuh-e-Surmeh deposit (Fig. 10). The non-mineralized Upper Dalan is overlain by thin layers of shale, carbonate and anhydrite of the Triassic Kangan Formation (Solymani, 1996). The overlying Triassic Dashtak sediments are composed mainly of evaporitic dolomite and anhydrite (or gypsum) measuring about 850 m in thickness at the Kuh-e-Surmeh deposit (Fig. 10). The Jurassic sediments consist predominantly of shale, gypsum, dolostone and limestone of the Neyriz and Surmeh formations (Liaghat et al., 2000).

The orebearing Permian carbonate sequence is folded and the mineralization occurs on both flanks of an anticline. The main sulfide ore minerals are sphalerite, galena, and pyrite, while the main non-sulfide ore minerals are hydrozincite, smithsonite, and hemimorphite. Supergene weathering of the sulfide ore could have taken place at several intervals between the Paleozoic and the present day (Liaghat et al., 2000). Gossans representing the oxidation products of Zn, Fe and Pb sulfides exploited in the Kuh-e-Surmeh mineralized district, occurred not only on top of the primary ores, but they also represent the infill of deeper karstic cavities in Paleozoic carbonates. Non-sulfide minerals from Kuh-e-Surmeh have been investigated previously by several authors (Reichert and Borg, 2008). The shape of the non-sulfide orebodies ranged from bulk replacement bodies of entire blocks of the host carbonates and/or sulfides, to concretionary infilling of isolated druses and cavities with idiomorphic smithsonite and hemimorphite crystals at different levels in the upper part of the oxidation zone. Hydrozincite is also present, as well as remnants of primary sulfides. Calcite was commonly observed, either as a gangue mineral that precipitated with the primary sulfides, and as a newly formed phase associated with smithsonite.
4.2. Nonsulfide Zn–Pb deposits in Iran within Mesozoic host rocks

Cretaceous carbonates are the most common host rocks of Iranian nonsulfide Zn–Pb deposits, which are largely concentrated in (1) the MEMB (Momenzadeh, 1976) in the SSZ, and (2) the YAMB in the Yazd block of the CIGS (Fig. 1 and Table 1). Approximately 66% of the Zn–Pb deposits of Iran occur in these two belts (metallogenic provinces) and include world-class examples such as Mehdiabad (in the YAMB) and Irankouh (in the MEMB) (Fig. 1).

4.2.1. Irankouh deposit

The Irankouh district comprises several Zn–Pb deposits (Figs. 11 and 12). They are located in the Irankouh Mountain Range, 20 km south of Esfahan in West-Central Iran. The Irankouh district comprises several Zn–Pb deposits, including the Goushfi pit (Borg, 2009), the Kolahdarvazeh pit, and the small Tapeh-Sorkh pit (Figs. 11 and 12). The main difference between the Goushfi and the Kolahdarvazeh pits is the nature of the ore. The Goushfi pit is characterized by large amounts of sulfide ore and a minor nonsulfide zinc ore zone. The Kolahdarvazeh mine is an open pit mine, which produces a concentrate of nonsulfide zinc ore and only shows a small amount of sulfide minerals (Fig. 13). Estimated ore reserves are more than 20 million metric tons, grading 7.4 wt.% of Zn and 2.4 wt.% of Pb. The Zn/(Zn + Pb) ratio in the Irankouh district is usually high, averaging about 0.76 (Reichert, 2007; Borg, 2009).

Zn–Pb–Ba mineralizations in the Irankouh (and in the Isfahan) district were first reported by Zahedi (1976), who described mineralized veins occurring along faulted and fractured zones in the lower part of the hydrothermally dolomitized Barremian-Aptian limestone. Rastad (1981) suggested a diagenetic origin for such stratabound Zn–Pb–Ba–(Cu) mineralization. A very comprehensive isotopic (O, C, S and Sr isotopes) study by Ghazban et al. (1994) concluded that the deposit was an epigenetic Zn–Pb–Ba MVT-type. Hosseini-Dinani and Aftabi (2015) also proposed an epigenetic stratabound Zn–Pb–Ba mineralization related to MVT on the basis of the low Cu and Fe contents and geochemical features. Boviri and Rastad (2016) and Boveiri et al. (2017) proposed a SEDEX type mineralization for the Gushfi and Tapeh-Sorkh parts in the Irankouh district.

The regional stratigraphic sequence of the Irankouh area (Fig. 12) starts with Early Jurassic shale at the bottom, followed by Barremian–Aptian sandstone and limestone, Upper Aptian shale, and, on the top of this sequence, Tertiary sediments. Two major stratigraphic gaps can be found within this stratigraphic sequence (Fig. 12). The first one separates the Lower Jurassic strata from the Early Cretaceous strata. The second gap is located between the Cretaceous strata and the Oligocene strata (Rastad, 1981)(Fig. 12). The Early Cretaceous sediments unconformably overlie the Jurassic shales to form a continuous sequence beginning from the Barremian up to the Upper Aptian sediments (Fig. 12). The total thickness of this sequence is approximately 800 m. These Cretaceous strata comprise mainly dolomite, limestone, and minor amounts of volcanic rocks, shale and marble (Fig. 12). The Barremian and Lower Aptian sedimentary rocks host the sulfide and non-sulfide mineralization, which forms two discrete ore-bearing horizons, namely the Goushfi-Kolahdarvazeh horizon and the Goushfi-Gowdezandane horizon (Rastad, 1981; Boveiri et al., 2015)(Fig. 12).

At the Kolahdarvazeh mine, both sulfide and non-sulfide ore zones are found within the Cretaceous dolomite and limestone (Fig. 13). Sulfide mineralization occurs in several places at the contact (or close to) with shale. In most cases, the sulfide ore is partly oxidized and only local relics of sulfide minerals (mainly galena) are found. The primary sulfide mineralization consists of
sphalerite, galena, with minor amounts of pyrite, and traces of marcasite, and chalcopyrite (Reichert, 2007). Associated minerals are barite and calcite mainly as veins and fracture fillings. A partial oxidation is frequently observed. In an advanced state of weathering most of the sulfide minerals have been removed and only galena is present.

Gossans representing the oxidation part of the Zn, Fe and Pb sulfide ore zones exploited in the Kolahdarvazeh mineralized districts (Fig. 13), occurred not only on the top of the primary ores, but they also represent infilling within deep karstic cavities in the Cretaceous carbonates (Reichert, 2007). Oxidation of sulfide minerals generally extends from the surface to an average of 40 to 70 m in depth. However, in several parts, Zn carbonates as well as druses filled by cerussite were also irregularly enclosed in the primary sulfide bodies down to 130 m in depth. Pb and Fe secondary minerals (cerussite, anglesite, goethite, etc.) were always associated to the Zn ores. The mineralization is considered to be the result of in situ oxidation of the primary sulfide ores by meteoric fluids, with increased acidity owing to the dissolution of substantial amounts of pyrite, which circulated through the carbonates (Reichert, 2007) (Fig. 13). The dissolution was followed by the partial to total replacement by secondary minerals of the sulfide minerals. A distinct oscillatory zoning (zinc-enriched and zinc-poor bands) in the Kolahdarvazeh deposit is considered to have been caused by variations of the water table, cyclically altering the regular deepening grade of the oxidation profiles (Reichert, 2007). Subsequent

Figure 12. Regional stratigraphic sequence of the Irankouh deposit. The ore deposits are hosted within the Early Cretaceous Formations (modified after Rastad, 1981).

<table>
<thead>
<tr>
<th>Age</th>
<th>Symbol</th>
<th>Lithology</th>
<th>Ore-horizons</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oli. Mio.</td>
<td>O.M</td>
<td>Light-coloured limestone with basal conglomerate</td>
<td></td>
</tr>
<tr>
<td>U. Aqptan</td>
<td>K_{gl}</td>
<td>Marly bedded orbitolina limestone</td>
<td></td>
</tr>
<tr>
<td></td>
<td>K_{gsh}</td>
<td>Shale and marl with intercalated limestone</td>
<td></td>
</tr>
<tr>
<td>K. I</td>
<td></td>
<td>Thick-bedded to massive limestone</td>
<td></td>
</tr>
<tr>
<td>Barrem-Aptan</td>
<td>K_{d2}</td>
<td>Well-bedded limestone with dolomite lenses</td>
<td></td>
</tr>
<tr>
<td></td>
<td>K_{d1}</td>
<td>Dolomite and calcareous dolomite (K_{d2})</td>
<td></td>
</tr>
<tr>
<td></td>
<td>K_{d1}</td>
<td>Medium-bedded rudist limestone with dolomite lenses (K_{d1})</td>
<td></td>
</tr>
<tr>
<td></td>
<td>K_{d2}</td>
<td>Red-brown medium-bedded dolomite (K_{d2})</td>
<td></td>
</tr>
<tr>
<td></td>
<td>K_{d2}</td>
<td>Orbitolina limestone (K_{d2})</td>
<td></td>
</tr>
<tr>
<td></td>
<td>K_{d1}</td>
<td>Red sandstone and conglomerate with sandy dolomite at top (K_{d1})</td>
<td></td>
</tr>
<tr>
<td></td>
<td>K_{d1}</td>
<td>Shale with intercalation of siltstone and sandstone</td>
<td></td>
</tr>
</tbody>
</table>

Figure 12. Regional stratigraphic sequence of the Irankouh deposit. The ore deposits are hosted within the Early Cretaceous Formations (modified after Rastad, 1981).
remobilization and redeposition within dissolution vugs and karst cavities of the newly formed oxidation minerals locally followed the early replacement process (Fig. 13B and C). Carbon and oxygen isotope measurements were applied to hydrozincite of the Kolahdarvazeh mine (Reichert, 2007). The hydrozincite samples show a range of $\delta^{18}O$ (SMOW) values between 21.9‰ and 22.9‰. The $\delta^{13}C$ values range between $-3.8‰$ and $-7.1‰$ (Reichert, 2007).

4.2.2. The Mehdiabad non-sulfide Zn–Pb deposit

The Mehdiabad deposit is situated in the Yazd Province, the central part of Iran approximately 550 km southeast of Tehran (Fig. 1). Several occurrences, namely the Calamine mine (CM), Black-Hill Ore (BHO), East Ridge (ER) and Central Valley Ore body (CVOB), have been explored and CM has been mined in historical times (Maghfouri et al., 2015; Maghfouri, 2017). Exploration during the last two decades revealed huge economic reserves, with an oxide reserve of 45.2 Mt at 7.15% Zn and 2.47% Pb and a sulfide reserve of 116.5 Mt at 7.3% Zn and 2.3% Pb. The total geological resource of this deposit includes 218 Mt at 7.20% Zn, 2.30% Pb, and 51 g/t Ag (BRGM, 1994; Reichert, 2007). In spite of its size and importance, modern data on the Mehdiabad deposit are scarce. Ghasemi (2006) investigated the geology and mineralogy of the deposit, and Reichert (2007) provided some geochemical data of the oxide ores and weathering conditions. Maghfouri (2017) also proposed a syn-sedimentary Zn–Pb–Ba–(Cu) mineralization related to SEDEX for the Mehdiabad deposit on the basis of the clastic-carbonate dominate host rocks, relation between ore minerals and rock forming minerals, structures and textures, different ore facies and geological and geochemical features.

In the Mehdiabad deposit, a thick Early Cretaceous sedimentary sequence (ECSS) is exposed, unconformable lying on top of the Jurassic Shir-Kuh granite (Fig. 14). In the following, the Early Cretaceous formations of the Yazd area (Nabavi, 1972a,b; Majidifard, 1996; Maghfouri, 2017) will be briefly described from base to top.
Early Cretaceous terrestrial sediments of the Sangestan Formation cover the metasediments of the Shemshak Group and the Shir-Kuh granite that initially formed a major palaeo-relief (Fig. 14). The lower part of this Formation is exclusively siliciclastic (reddish conglomerate, sandstone and siltstone) reflecting semi-arid terrestrial conditions. Toward the top of sequence, the marine influence increases and is indicated by evaporites and carbonates, locally containing calcareous algae and agglutinating foraminifera. Although the carbonates from the upper part of the Sangestan Formation have been assigned to Hauterivian (Bucur et al., 2003), the Sangestan Formation is generally considered to be of Barremian age. The Sangestan Formation is conformably overlain by thick-bedded carbonates of the (Barremian?) Aptian Taft Formation (Fig. 13), consisting of thick-bedded to massive rudist-, algae- and orbitolinid-bearing shallow-water limestone, organic matter rich shale, silty shale, silty limestone and siltstone (Fig. 14) (e.g., Bucur et al., 2012; Schlagintweit et al., 2013a,b). The Taft Formation may reach up to 400 m in thickness (section at west BHO; Fig. 15) and is part of a large-scale carbonate platform system that characterized the Yazd Block during the Early Cretaceous (Wilmsen et al., 2013). The Zn–Pb–Ba mineralization of the Mehdiaabad deposit occurs within the Taft Formation (Fig. 14) (Maghfouri et al., 2015; Maghfouri, 2017). The upper part of the Taft Formation is covered by characteristic cherty limestone of the Abkuh Formation. Above the cherty limestone, dolomitic shale with calcarenite and chert-bearing limestone, are generally identical to those seen in the drill holes. 1–10 meters-sized lenses and olistoliths of para-reef limestone become frequent in the upper part of the unit. The upper part of this unit (60–80 m) consists of limy shale and laminar limestone. The CM orebody is located in the upper part of the Abkuh Formation (Fig. 14).

Zn–Pb–Ba (Cu) mineralization of the Mehdiaabad deposit occurs along two horizons that extend over a length of >4 km (Fig. 13). The sulfide and non-sulfide ores of the ore horizon I (OHI) are hosted by organic matter-rich shale, silty limestone, dolomite and silty shale.
of the Taft Formation, whereas non-sulfide ores of the ore horizon II (OHI) are enclosed by limy shale and thin-bedded limestone of the Abkuh Formation (Figs. 14e17) (Maghfouri et al., 2015; Maghfouri, 2017). The OHI is composed of several ore bodies, namely Black Hill Ore (BHO), Central Valley Ore Body (CVOB) and East Ridge (ER), located in a depression surrounded by hills and mountains (Figs. 15, 16 and 19D). The calamine mine (CM) exposes the OH II, which represents the highest part of the oxide ore mineralization (Figs. 17 and 19F). At sites of BHO, ER and CM, OHI is completely oxidized (Figs. 15e17), whereas the CVOB display a large proportion of sulfide minerals (Fig. 19E).

Detailed geological mapping of host rocks and structures in the Mehdiabad deposit shows that Zn-Pb-Ba-(Cu) mineralization (OHI and OHII) occurs to the east of the Black Hill Fault, a major and well-exposed fault within the Early Cretaceous sequence (Figs. 15 and 19D).

BHO veins and brecciated mineralization occurs adjacent to the Black-Hill fault (Fig. 19D). The BHO also referred as the BHO gossan consists of irregular veins of barite, quartz, chalcopyrite, pyrite, sphalerite, galena, limonite, hematite, Zn-Pb carbonate and dolomite (Fig. 15). To the east, the BHO passes into CVOB (Fig. 19D). The organic matter-rich shale, dolomite and silty limestone of the Taft Formation, host the main part of the preserved sulfide ore of the CVOB (Fig. 19E), whereas ores are enclosed is shale and siltstone in the easternmost areas. The most abundant minerals are galena, sphalerite, barite and pyrite with traces of chalcopyrite (Maghfouri, 2017).

Further to the east, the ER orebody consists of an essentially strata-bound non-sulfide mineralization developed within the silty limestone of the Taft Formation, close to the uppermost part of the Sangestan Formation, which exposes sandstone and oolitic limestone (Figs. 16A and 19D). Mineralization consists of iron (hydr-) oxide (hematite, limonite), Mn oxide and Zn and Pb carbonates (Fig. 18). A deep preserved sulfide mineralization, including galena, sphalerite and pyrite was recognized by drill cores at 100 m (Reichert, 2007; Maghfouri, 2017).

The CM represents the highest part of the exposed mineralization at Mehdiabad (Fig. 17). The mineralization is predominantly hosted within limy shale with intercalated laminar limestone of the Abkuh Formation and particularly at the contact between limy shale and chert-bearing bedded limestone (Figs. 17 and 19F). Zn-Pb mineralization exclusively consists of non-sulfide minerals, such as smithsonite, hemimorphite, and hydrozincite (Fig. 18).

The non-sulfide zinc minerals in the Mehdiabad deposit belong to the carbonate-hosted “non-sulfide” category (Large, 2001), in which hemimorphite, hydrozincite, and smithsonite are the principal zinc-bearing minerals. Cerussite and anglesite also occur, generally associated with lenses of residual or supergene galena. A complex association of iron and manganese oxi-hydroxides (goethite, lepidocrocite, hematite), and residual clay minerals hosts the non-sulfide ore. The mineralogy of the ores is generally complex and, besides the most common Zn and Pb carbonates and silicates, it also comprises exotic species (Billows, 1941; Moore, 1972; Stará et al., 1996). The non-sulfide zinc ore of the Mehdiabad deposit can be subdivided into a red zinc ore (RZO) and a white zinc ore (WZO) (Fig. 18). The RZO is rich in Zn (up to 30%), iron (17%) and other metals such as Pb and As (Reichert, 2007). In contrast, the WZO typically shows high zinc grades (up to 40%) and low concentrations of iron (<7%) with minor Pb and As (Reichert, 2007). The RZO is dominant in the OHI, while the WZO is characteristic of the OHII. Common minerals of the RZO are Fe-oxhydroxides, goethite, hematite, hemimorphite, hydrozincite, smithsonite, and cerussite. Hemimorphite is one of the most important zinc minerals of this ore-type. Most common minerals of the WZO are hydrozincite, smithsonite, and hemimorphite. Iron-bearing minerals are rare compared with the RZO. Hemimorphite is less common compared to what observed in the RZO.
Carbon and oxygen isotope measurements have been made in ore samples of the Mehdiabad deposit (Reichert, 2007). The hydrozincite samples show a restricted range of $\delta^{18}O$ values between 20.4‰ and 21.9‰. The $\delta^{13}C$ values are more variable and range between $-0.4‰$ and $-3.8‰$ (Reichert, 2007). This signature is similar to that of smithsonite samples, which have been analyzed by Boni et al. (2003) and Gilg and Boni (2004). It indicates that at least two isotopically different sources of carbon contributed to the formation of zinc carbonates. The roughly constant $\delta^{18}O$ values and the variably low- to medium-depleted $\delta^{13}C$ values reflect the influence of meteoric water associated with subaerial exposure (Allen and Matthews, 1982; Lohmann, 1988) and point to the precipitation of the zinc carbonates within the vadose and/or phreatic zone (Allen and Matthews, 1982; Reichert, 2007; Borg, 2015). The isotopically light component of the carbon is probably the result of dissolved CO$_2$ derived from the decay of organic matter (Criss, 1995).

5. Discussion

5.1. Critical parameters on the formation of non-sulfide deposit

Based on studies of Borg (2005), Reichert (2007) and Reichert and Borg (2008), non-sulfide zinc ores in Iran are mostly secondary and was formed by the supergene oxidation of primary sulfide deposits (Figs. 19 and 20A), although there is some evidence for primary origin of non-sulfide zinc minerals in Irankouh (Reichert, 2007). Silicate-dominated non-sulfide Zn deposits, such as those in the Upper Proterozoic volcano-sedimentary successions in the Americas and Australia (e.g., Vazante, Brazil; Beltana, Australia), have not been identified in Iran. Non-sulfide zinc deposits in Iran are essentially zinc-rich gossans exposing the replacement of sphalerite by smithsonite, hemimorphite and hydrozincite. In addition, zinc sulfide deposits originally contained galena, copper and lead sulfides. As these minerals are oxidized, they may be replaced by copper and lead carbonates, oxides, and even secondary sulfides like chalcocite.

Direct-replacement non-sulfide deposits in Iran represent the supergene evolution of carbonate hosted-type (MVT-type), carbonate replacement-type and Sedex deposits (Fig. 20A). Deposits characterized by carbonate environment (Fig. 19A and C–F), such as Mehdiabad, Irankouh, Kuh-e-Surme and Angouran tend to be mineralogically simple and are dominated by smithsonite, hemimorphite, and hydrozincite. In addition, like at Mehdiabad and Irankouh deposits Zn-rich Mn minerals, copper-zinc carbonates and complex arsenic minerals are also present. The occurrence of primary iron sulfide minerals tends to produce sufficient acid
during weathering to completely leach zinc from the near-surface environment (Figs. 7, 9 and 16). For example, the OHI in Mehdiabad deposit is oxidized up to a depth of 100 m (Figs. 15 and 19D); despite high primary grade in the sulfide zone (approx. 12% zinc and 5% lead), the gossan cap and oxidized zone show complete non-sulfide minerals features (Fig. 15). The sulfide mineral content in this horizon is dominated by pyrite, which constitutes more than 30 vol.% of the ore. The \( \text{H}_2\text{SO}_4 \) generated by pyrite oxidation, together with \( \text{Fe}^{3+} \) as an oxidant (Reichert, 2007), allows for the complete removal of zinc, which is highly mobile in low pH conditions (Andrew-Jones, 1968). Such complete leaching results in the formation of a gossan with iron oxides, cerussite, hemimorphite, and copper carbonates in the BHO and ER.

Like supergene copper “oxide” deposits (Chávez, 2000), non-sulfide zinc deposits in Iran formed by oxidation of a sulfide-bearing protolith (Fig. 20A) following a series of reactions between meteoric waters, metal sulfides, and reactive host rocks (Borg, 2005, 2015; Reichert and Borg, 2008; Choulet et al., 2013). Dissolution of sulfides by \( \text{O}_2 \) and \( \text{Fe}^{3+} \) provides low pH, sulfate-bearing solutions, which are able to transport metals. Although sphalerite and, to a much lesser extent, galena are very susceptible to oxidation (Bladh, 1982; Boyle, 1994), they produce relatively small quantities of acid sulfate-bearing solutions compared to iron sulfides (Williams, 1990). Sphalerite is also extremely susceptible to oxidation by moderately thermophilic (30–50 °C) bacteria in a highly oxidized, acidic, ferric sulfate-rich environment. It is normally consumed by oxidation before pyrrhotite, pyrite, galena, or chalcopyrite (Rose et al., 1979). Normally, acids produced by such oxidation of sulfide minerals would be expected to effectively remove zinc from the system (Sangameshwar and Barnes, 1983). However, given the buffering action of the carbonate host rocks, sulfuric acid generated by the oxidation of pyrite and other sulfide minerals will be neutralized, thereby ensuring a buffered, nearly
neutral pH environment. Under these conditions, zinc carbonate like smithsonite is the normal products of oxidation of sphalerite deposits through contact with meteoric waters (Reichert, 2007; Reichert and Borg, 2008; Choulet et al., 2013). The isotopic pattern of the hydrozincite samples of Kolahdarvazeh deposit show close ranged $\delta^{18}O$ values and a more variable range of $\delta^{13}C$ data (Reichert, 2007). The $\delta^{13}C$ values of the Kolahdarvazeh are up to 3.3‰ lower than those from hydrozincite of Mehdiabad (Reichert, 2007). The formation of the hydrozincite should be potentially influenced by an isotopically light component of carbon as a result of dissolved CO$_2$ derived from the decay of organic matter. However, the hydrozincite of the Kolahdarvazeh mine shows lower $\delta^{13}C$ values than its pendant from Mehdiabad deposit (Reichert, 2007). This suggests a more important role of organic and meteoric CO$_2$ than in Mehdiabad (Reichert, 2007). A possible reason is a variable contribution of carbonate carbon from the limestone and/or the dolomite host rock and reduced organic carbon derived from the overlying soils and is comparable to pedogenic carbonates (Cerling, 1984; Coniglio et al., 1996).

5.2. Genesis and controls on the formation of Iranian non-sulfide deposits

Some of the key controls on the formation of carbonate-hosted non-sulfide Zn–Pb deposits are the nature and availability of near-surface sulfide protore, lithology, subaerial exposure, tectonic uplift, climate and favorable hydrology (Large, 2001; Hitzman et al., 2003). In the Iranian non-sulfide deposits, tectonic uplift resulted in the near-surface exposure of the sulfide protore enabling their oxidation under a favorable climate (Fig. 20A and B) (Reichert, 2007; Reichert and Borg, 2008; Borg, 2015). Tectonic uplift and related faulting during post-sulfide mineralization deformation events played a role for the fracturing of hosting carbonates (Fig. 20B) (Aghanabati, 1998, 2004), facilitating the action of meteoric waters (Fig. 20C). Climate also strongly controlled the oxidation and therefore the transport and the redeposition of metals. The most favorable conditions for oxidation are achieved under arid (Reichert and Borg, 2008; Borg, 2015) to temperate (Hitzman et al., 2003; Santoro et al., 2013) climates,
Figure 19. Cross-sections and plan view of the ore sequence and extension of non-sulfide mineralization in Iranian deposits. (A) NE–SW cross section (800 N) of the Angouran pit through the ore body; the pipe-like geometry are shown together with the distribution of the various ore types and the drill hole intervals with Zn assays (modified after Daliran et al., 2013). (B) Geological cross section of the Haft-har deposit showing the small-scale folding of the sequence. (C) General section of the Irankouh mining district, including Kolahdarvazeh mine and Tapeh-Sorkh mine (Rastad, 1981). (D) Schematic cross section of the OHI in the Mehdiabad deposit, including the o–p profile along the BHO mineralization (Fig. 15) and the q–r profile along the ER mineralization (Fig. 16A). (E) N–S cross section of the ore sequence with extension of sulfide and non-sulfide ore mineralizations in the CVOB (Mehdiabad deposit). (F) Schematic NWW–SSE section of the folded and faulted strata at the Calamine Mine area in the Mehdiabad deposit (Location of the m–n section is shown in Fig. 17A). The non-sulfide-bearing strata are repeated due to folding and thrusting (modified after Nosratian, 1991).
in which a maximum amount of metals is available for transport by supergene solutions. As the biogenic activity in the soil is limited, more oxygen is available for sulfide oxidation, and consequently a lot of metals are liberated into the solutions (Reichert and Borg, 2008). It is important to note, that the climate as well as the local geology (fragmentation, karstification) of the carbonate host rocks influence the O$_2$ and CO$_2$ concentrations of the descendent fluids, and thus, the pH and the ability of the fluid to dissolve the carbonate host rock (Fig. 20C). Reichert and Borg (2008) showed that arid environments provided the best conditions for the oxygen-driven oxidation of sulfide ores. The oxidation by Fe (III) is not included here, since Fe (III) itself is provided by oxygen and affects the oxidation process only on a regional scale. In arid environments like in Iran, the dissolved O$_2$ reaches its maximum concentration, compared to other climates, and will not be consumed by biological activities within soils (Reichert, 2007; Reichert and Borg, 2008). The groundwater table in arid climates is commonly low. This will lead to an opening of the water-filled pores and joints after an individual rainfall event and will thus provide an inward flow of gasses (O$_2$, CO$_2$) to any available sulfide ore body (Reichert, 2007; Reichert and Borg, 2008; Choulet et al., 2013). This system will commonly change

Figure 19. (continued).
to a highly permeable system due to (karstic) dissolution processes of the carbonate host rock (Fig. 20C). Due to the limited availability of water in arid and hyper-arid climates, the fluids, which have been generated during the oxidation process, would be highly enriched in zinc and other metals (Fig. 20C). These high metal concentrations support an effective precipitation of non-sulfide base metal minerals (Fig. 20C). These high element concentrations support an effective precipitation process within the carbonate host rock. Thus, an arid or semi-arid climate of Iranian non-sulfide environments provides best conditions for the
6. Conclusion

The main economic minerals of the “non-sulfide” type deposits in Iran consist of Zn (hydroxides) and silicates (smithsonite, hydrozincite, and hemimorphite) associated with Fe and Mn oxides (hydroxides) and residual clays. The mineralization is considered to be the result of in situ oxidation and replacement (with limited transport, locally) of the primary sulfide phases, due to meteoric fluids circulating in the carbonate rocks. Consequently, non-sulfide Zn ore deposits that originate from stratiform sulfide lenses formed either by sulfide replacement or by infilling of karst conduits due to precipitation or internal sedimentation.

Although the six studied ore deposits present a similar first-order mineralogical evolution, we have observed small discrepancies between these sites owing to various local and external factors. Climate conditions were generally similar in Mehdiabad, Irankhoo, Kuh-e-Surme, Zarigan and Haft-haahr and different with Angouran, but studied zones have experienced differential deforestation, oxidation.

The association of carbonate-hosted non-sulfide zones with underlying massive sulfides in combination with the characteristics of the non-sulfide ore suggests that most of the known non-sulfide mineralized zones in Iran are the result of supergene, direct replacement process of CH and SEDEX deposits. Zinc and Pb non-sulfides can also be used as indirect indicator for exploration of MVT, SEDEX, Irish-type, and carbonate replacement and vein-type Zn–Pb deposits.

References

