Origin of the Moroccan Touissit-Bou Beker and Jbel Bou Dahar Supergene Non-Sulfide Biomineralization and Its Relevance to Microbiological Activity, Late Miocene Uplift and Climate Changes

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Abstract: Through integration of Pb-Zn ± Cu non-sulfide mineralogy, texture, and stable isotope (C, O, S) geochemistry, the world-class Touissit-Bou Beker and Jbel Bou Dahar Mississippi Valley-type districts of the Moroccan Atlasic system have been investigated in order to gain insights into the origin and processes that contributed to the formation of the base metal non-sulfide mineralization. In both districts, direct replacement (“red calamine”) and wallrock replacement (“white calamine”) ores are observed. Based on the mineral assemblages, ore textures, and crosscutting relations, three distinct mineralizing stages are recognized. The earliest, pre-non-sulfide gossanous stage was a prerequisite for the following supergene stages and constituted the driving force that ultimately promoted the leaching of most base metals such as Zn and Cu and alkalis from their rock sources. The following two stages, referred to as the main supergene “red calamine” and late “white calamine” ore stages, generated the bulk of mineable “calamine” ores in the Touissit-Bou Beker and Jbel Bou Dahar districts. Stable isotope compositions (δ13C V-PDB, δ18O V-SMOW, δ34S CDT) support a three-stage model whereby metals were released by supergene acidic fluids and then precipitated by bacteria and archaea-mediated metal-rich meteoric fluids due to a decrease in temperature and/or increase of fCO2. Oxygen isotope thermometry indicates decreasing precipitation temperatures with advancing paragenetic sequence from 33° to 18°C, with wet to semi-arid to arid climatic conditions. The close spatial relationships between coexisting sulfide and non-sulfide mineralization along with stable isotope constraints suggest that the oxidation of sulfides occurred concurrently after the main stage of the Alpine orogeny between 15 Ma and the present. More importantly, the data show for the first time the involvement of biologically controlled activity as the major driving process that triggered both oxidation and deposition of supergene mineralization at Jbel Bou Dahar and Touissit-Bou Beker districts. Conclusions drawn from this study therefore have implications for supergene Mississippi Valley-type (MVT) -derived non-sulfide deposits worldwide and account for the prominent role of biological processes in the origin of these ore deposits.

Keywords: supergene environment; calamine ores; biomineralization; C; O; S isotope geochemistry; High Atlas Morocco
1. Introduction

Long ignored by the mining industry owing to problems with the mineral processing and recovery of the ore, supergene non-sulfide Zn-Pb ± Cu deposits, formed from low-temperature alteration of exhumed sulfide-bearing protore, are becoming attractive exploration targets worldwide due to the new developments in hydrometallurgy [1–3]. In addition to these advances, the renewed interest in supergene non-sulfide ore at the expense of conventional Zn-Pb ± Cu deposits resides in their higher Zn-Pb ± Cu grades averaging up to 20 wt.% Zn [4] and 18.1 wt.% Pb [5] and less environmental impact than sulfide ore (e.g., no sulfur emissions or waste, little to no acid mine drainage). Equally important, Zn non-sulfide ore can greatly improve the smelting of sulfide-rich Zn and Pb ore. In addition to base metals, supergene non-sulfide deposits are considered as a potential resource for strategic metals such as Ge, Ga and In [6–8] and have the potential to become future alternative sources complementary to alkaline rocks and associated carbonatites. The high demand for these critical metals, which are also recovered as a by-product from non-sulfide Zn-Pb ± Cu deposits, has spawned worldwide exploration projects focusing on this new, emerging class of supergene ore deposits. Notable examples include Cristal Zn prospect of Bongará district in northern Peru, Kabwe Zn-Pb deposit (formerly known as “Broken Hill”) in central Zambia, Tsumeb in Namibia, and Apex deposit in the United States ([6,8–10] and references therein).

Since non-sulfide minerals are recovered through different hydrometallurgical processes such as pyrometallurgy, electrowinning, AmmLeach®, and acid leaching ([3] and references therein), the economic value of Zn-Pb ± Cu non-sulfide ores is dependent not only on the geologic setting of each deposit but also on the initial mineralogy, including the nature of the gangue minerals, the composition and flow rates of groundwater, and the permeability of the ores and host carbonates ([11–13] and the present study). The current understanding of the geological setting, mineralogy, geochemistry, and climatic evolution of the supergene non-sulfide deposits indicates that these deposits formed in a near-surface environment by low-temperature oxidation of a sulfide-rich protolith. The ore-forming process involved a series of reactions between downward-percolating meteoric waters, metal solubility, and reactive host rocks contemporaneously with tectonic uplift and/or water table depression [3,14,15]. The released metals could be trapped locally, directly above the sulfide protolith, or they may be transported and precipitated far away from the sulfide protore [3,16–19]. More importantly, recent investigations demonstrated the ability of bacteria and archaea as well to influence metal solubilization, mobilization, and subsequent formation of carbonate-hosted Pb-Zn ± Cu ± Au deposits and their supergene counterparts (e.g., [20–25]).

The Tethyan intra-continental Atlas system of Northwest Africa [26] hosts more than 1000 sulfide Pb-Zn ± Cu ± F ± Ba orebodies and deposits that range from small uneconomic showings to major districts of great size and productivity (Figure 1). Major and most productive deposits occur in Morocco within unmetamorphosed, flat-lying Lower and Middle Jurassic carbonates that experienced multiple overlapping episodes of dolomitization, dissolution (i.e., karstification), and uplift during the Mesozoic breakup of the Pangean continental shelf of Northwest Africa. Featured deposits and districts, which are introduced in this contribution, include the world-class and historically important Touissit-Bou Beker (>100 Mt at 4 wt.% Pb, 3.5 wt.% Zn, <1 wt.% Cu, 120 g/t Ag; [5]) and the Jbel Bou Dahar (>30 Mt at 4 wt.% Pb, 4 wt.% Zn) districts (Figure 1). Both sulfide and non-sulfide mineralization coexist in these two major districts, with the supergene Pb-Zn ± Cu-bearing occurrences being superimposed on the surviving oxidized sulfide protore. Earlier studies indicate that the primary sulfide mineralization is of Mississippi Valley-type (MVT) affiliation [5,26–28]. Major advances in understanding the genesis of MVT mineralization have been accomplished over the last two decades, principally as a result of field observations, mineralogical assemblages, and textural characterization, along with fluid inclusions and stable and radiogenic isotope geochemistry [5,26–29].
However, the mineralogy and origin of the non-sulfide ores have been neglected and have only recently become the subject of ongoing investigations [30–32].

In this contribution, we report the first comprehensive carbon, oxygen, and sulfur isotope dataset from the abovementioned two representative and the largest MVT districts of the Moroccan Atlas system (Figure 1). Although the selected districts share many similarities in terms of their geological environments and age of mineralization, they show, however, contrasting Zn/Pb ratios that make them distinct from each other (i.e., Pb-dominant vs. Zn-dominant deposits), providing an ideal methodological approach to investigate the role of initial sulfide mineralogy in controlling the distribution of the supergene non-sulfide ore. The aim was to fill the gap of knowledge on non-sulfide Pb-Zn ± Cu deposits and to better constrain the nature of the fluids involved in the mineralizing process within the framework of geodynamics and paleoclimatic evolution of the Atlas, assess the role of microbial processes in their formation, a topic that has started to attract the interest of economic geologists only recently, and draw a distinction between “white” and “red” calamine ores. The specific objectives were fourfold: (1) update the geologic attributes, petrography, mineralogy, and geochemistry of the non-sulfide Pb-Zn ± Cu orebodies; (2) refine the paragenetic relationship between sulfides, non-sulfides, and their associated gangue minerals; (3) discuss the close temporal links between non-sulfide ore formation, tectonic uplift, regional climatic evolution, and biogenic processes; and (4) combine the newly acquired isotopic results with published isotopic data from other non-sulfide deposits around the world to constrain the genesis of supergene mineralization.

In this study, we provide macrotextural and carbon and sulfur isotope evidence supporting the hypothesis that the microorganisms played a major role in the formation of the major Moroccan non-sulfide Pb-Zn ± Cu deposits in the Atlas system. This will thus contribute to the identification of new targets throughout this vast metallogenic province, which contains some of the best prospects for undiscovered non-sulfide Pb-Zn ± Cu deposits.

1.1. Regional Geological Setting and Mineralization

The regional geologic setting of the two investigated districts has been the subject of several pioneering studies [5,26,28,33–36]. A summary of these results is presented here, with an emphasis on the Mesozoic to Cenozoic stratigraphic and tectonic events that the MVT mineralization and the superimposed non-sulfide Pb-Zn ± Cu ores are related to.
Figure 1. Geologic map of northwestern Africa and southern Iberia illustrating the main lithostructural domains and spatial distribution of major Maghrebian (i.e., Moroccan, Algerian, and Tunisian) Mississippi Valley-type (MVT) districts within the Atlas system (adapted from [26]).
Overall, the Mesozoic and Cenozoic stratigraphy of the Atlas system, to which all of the investigated districts belong, consists of a succession of rift-related Triassic-Early Jurassic red-bed formations and intercalated carbonate rocks. These lithologies are overlain by a series of post-rift sub-horizontal and unmetamorphosed Middle-Late Jurassic to Early Cretaceous carbonate and marl strata that were deposited into a succession of alternating horst and graben structures related to the opening of the Tethyan and Atlantic oceans [37,38]. The sedimentary succession was folded and thrust in response to the Late Cretaceous-Cenozoic Atlasic orogeny, which developed in response to the convergence between the African and Eurasian plates. The regional stress regime inverted from extensional to compressional tectonics since the Late Cretaceous, with the fault network inherited from the rifting episodes playing a first-order control on the inversion [39–44]. Lower to Middle Jurassic carbonate strata are exceptionally well endowed in Pb-Zn ± Cu deposits of MVT affiliation [26], some of which are of world-class significance. In all cases, mineralization is epigenetic (i.e., ore genesis postdates deposition and diagenetic lithification of carbonate host rocks) and occurs either as replacement of the carbonate host rock or as open-space filling of karsts, veins, vugs, and brecciated fault zones. Recent contributions have established close temporal links between supergene ore formation and regional geotectonic and paleoclimatic evolution [26,30].

1.2. Touissit-Bou Beker District Geology

The Touissit-Bou Beker district of northeastern Morocco and adjacent western Algeria (Figures 1 and 2A) is one of the top ten MVT districts in the world, with a total production, over 75 years of mining activity, that exceeded 5 Mt of Pb and Zn [5,28].

The district stratigraphy consists of a Paleozoic basement unconformably overlain by a series of tabular and unmetamorphosed Meso-Cenozoic to Quaternary terranes (Figure 2B). The Aalenian-Bajocian (Middle Jurassic) dolomitic sequence, which hosts nearly all economic deposits of the Touissit-Bou Beker district, rests either directly on the Paleozoic basement or conformably overlies the Early Jurassic unit. The mineralogy of the deposits is very simple, comprising sphalerite and galena, with minor pyrite-marcasite and sparse chalcopyrite ± sulfosalts (i.e., tetrahedrite), accompanied by different generations of hydrothermal dolomite. Supergene mineral assemblages are extensively developed, with the oxidized ore zones extending as deep as 120 m and accounting for nearly 60% of the extracted ore [5]. The geometry of the orebodies that parallel the Atlasic ENE–WSW- and E–W-trending faults [45–47] coupled with Pb-Pb geochronological data [5] suggest that the MVT mineralizing event occurred during the Late Miocene at <15 Ma [5,26,28], consistent with the time span determined for most of the north African MVTs [29,48].
Figure 2. (A). Geologic map of Touissit-Bou Beker district showing regional geology, major faults, distribution of ore-bearing dolostone, and associated MVT deposits (projecting to the surface). Regional geology simplified from [45,49–51]. The red rectangle shows the location of the Touissit-Bou Beker district. (B). Generalized stratigraphic column for the Touissit-Bou Beker district showing main lithostratigraphic units.
1.3. Jbel Bou Dahar District Geology

1.3.1. Mining History

Production of lead, zinc, and barite in the Jbel Bou Dahar district and the surrounding areas is carried out exclusively through artisanal mining under the auspices of the CADETAF “Purchasing and Development Cooperative for the Tafilalet and Figuig Mining Region”, a public institution created following the Royal Decree of December 1st, 1960. The CADETAF claims, covering an area that extends over 60,000 km² in the central and eastern High Atlas, together with the mineral exploration licenses currently owned by the mining company ManagemGroup, define the largest and most productive “calamine” province in North Africa. Compiled data provided by both the CADETAF and ManagemGroup indicate that >1 Mt of Zn ore, with a grade exceeding 30% ZnO, has been produced to date. For instance, from 2012 to 2019, ManagemGroup produced a total of 559,403 tons grading ~16% Zn.

1.3.2. Geologic Setting

The Jbel Bou Dahar plateau occurs as an elongated, ENE-trending, 35 to 40 km long and 4 to 15 km wide flat-topped mountain with all of the current and past mine workings being distributed along the margins of the Lower Jurassic carbonate platform (Figure 3). The core of the carbonate platform consists of a succession of greenschist-facies Cambrian-Ordovician to Silurian metasedimentary and volcanoclastic rocks made up dominantly of quartzite and schist. Unconformably overlying the metasedimentary Paleozoic package is a sequence of about 400–500 m thickness of Triassic sedimentary rocks consisting of red-bed siltstone and gypsiferous to salt-bearing argillite locally interbedded with CAMP tholeiitic basalts [52]. The reef deposits consist of unmetamorphosed, flat-lying, massive, thickly bedded coral-, algal-, and bivalve-bearing Pliensbachian platform carbonates. Platform growth was initiated during the Sinemurian and ended during the Domerian (Upper Pliensbachian)–Middle Toarcian [53–56]. The Jurassic to Cretaceous strata were dissected by an array of faults, some larger and more continuous, trending mainly E–W, with some trending ENE–WSW and a subsidiary set trending NNW–SSE (Figure 3). Sulfide Pb-Zn-Ba ± Cu orebodies are confined to the brecciated, coarse sand to gravel, bioclastic bivalve-coral-algal packstones of Pliensbachian age, and/or to pinchouts of the Liassic strata against Paleozoic basement high (Figure 4). The sulfide mineralogy is fairly simple and consists of variable amounts of sphalerite, galena, pyrite-marcasite, and chalcopyrite, accompanied by barite, celestite, and calcite as the main gangue minerals.
Figure 3. Geological map of Jbel Bou Dahar early Mesozoic annular reefal complex emphasizing regional geology, main tectonic structures, and stratigraphic distribution of major MVT deposits. Line A–B indicates cross section shown in Figure 4. Modified after [57].
Figure 4. N–S geological section along strike of Jbel Bou Dahar early Mesozoic carbonate platform showing the spatial distribution of main lithofacies associations and MVT deposits; see Figure 3 for location of section. Modified after [57].
2. Materials and Methods

The sample suite analyzed in this contribution for mineralogical and geochemical studies includes a total 59 mineral separates collected from surface outcrops, high-grade open pits, and underground workings to obtain a representative suite of the principal minerals deposited throughout the two main supergene stages recognized throughout the Touissit-Bou Beker and Jbel Bou Dahar districts. In all cases, and in order to avoid analysis of impurities, Pb-Zn ± Cu-bearing carbonates and gypsum separates were extracted from crushed and washed fractions. Only pure separates of each mineral phase were analyzed.

The selected samples were mounted on Epoxy resin for polished block preparation to perform microscopic observations. The entire surface of each polished block was imaged by an optical microscope (Zeiss AxioImager M2m, Carl Zeiss AG, Oberkochen, Germany) with an XYZ motorized stage to obtain tiled images that allowed the sample to be navigated under the Scanning Electron Microscope (SEM) with micrometric precision owing to correlative microscopy. Detailed mineral stoichiometry was addressed using a Zeiss (Sigma 300) field emission gun SEM equipped with two Bruker xFlash 6|30 X-ray energy-dispersive spectrometers (EDS) at the laboratory of Minerals Engineering, Materials and Environment (GeMMe) of University of Liège (Belgium). The SEM-EDS analyses were carried out using a probe current of 2.3 nA with an accelerating voltage of 20 kV at a working distance of 8.5 mm. Manual spot analysis was performed to identify the carbonate Pb-Zn ± Cu-bearing minerals and surrounding phases using backscattered electron (BSE) and cathodoluminescence (CL) imaging modes. Additional samples were observed on a Zeiss Axiophot reflection microscope equipped with a Jeol JSM-7500F SEM and with a Zeiss Sigma 300 FEG SEM (Carl Zeiss AG, Oberkochen, Germany) equipped with two Bruker EDS detectors (Bruker Corporation, Massachusetts, USA) at the University of Namur (Belgium).

Carbon and oxygen isotope analyses were carried out at the Radiometric Dating and Stable Isotope Research laboratory at the University of Erlangen-Nuremberg (Germany) and California State University on hand-picked carbonate minerals including cerussite, smithsonite, malachite, azurite, and anglesite as well as on calcite. Carbonate powders were reacted with 100% phosphoric acid at 70 °C using a Gasbench II connected to a ThermoFisher Delta V Plus mass spectrometer. Reproducibility and accuracy were monitored by replicate analysis of laboratory standards calibrated by assigning a δ^{13}C of + 1.95‰ to NBS19 and 47.3‰ to IAEACO9 and a δ^{18}O of 2.20‰ to NBS19 and 23.2‰ to NBS18. Analytical precision (1σ) was calculated on the basis of multiple measurements of well-characterized carbonate standards between sample runs and was better than 0.01‰ for δ^{13}C and 0.02‰ for δ^{18}O, after corrections for scale compression.

Isotopic data are reported here relative to the Standard Mean Ocean Water (V-SMOW) for oxygen and to the PeeDee Belemnite (V-PDB) for carbon [58]. Oxygen isotope data have been converted into temperature estimates using the isotope thermometer of [59] for cerussite, [60] for smithsonite, [61] for malachite, [62] for azurite, and [63] for calcite. For these calculations, groundwater involved in forming the carbonates was assumed to have an oxygen isotope value of −7.2‰. This value is based on both modern groundwater and fluid inclusion values, consistent with those reported in [62]. Reproducibility was checked by replicate analysis of laboratory standards and was better than ±0.07‰ (1σ) for both carbon and oxygen isotope analyses.

Isotope geothermometry was used in conjunction with the results of oxygen isotope analyses for the Pb-Zn ± Cu-bearing carbonates to produce estimates of the temperatures at which these divalent metal cation carbonates formed. In this respect, it must be noted that these are thermometry estimates, as they represent an estimation of the thermal conditions under which the minerals are likely to have formed. Critical to the accuracy of this technique is an understanding of the oxygen isotope value of the waters from which these minerals formed. It has been shown by many studies that modern meteoric water values often serve as a good proxy for the waters of formation, due to the relatively recent
formation of secondary carbonates [9,61,62,64,65]. Furthermore, the use of the hydrogen isotope values of water extracted from fluid inclusions may be used to extract a proxy for oxygen isotope values for the water of formation, using the meteoric water relationship [65]. However, the large quantity of pure mineral required for such analysis on a crushing stage precludes multiple analyses. It is also known that clay hydrogen isotope values may be used as a proxy for meteoric waters from as far back as tertiary weathering environments, with attendant implications for the waters of formation of metal carbonate minerals (e.g., [64,65]).

As with other studies [9,61,62,64,65], it is assumed that there may be as much as 1% variation in oxygen isotope values using these methods, based upon known issues such as seasonal variability in precipitation sourcing. Such a variation would yield a maximum thermometry error of less than 6°C; far less than the range of variation seen in the samples from this present study.

The sulfur isotope compositions were determined at Actlabs (Canada) using the online elemental analyzer (Carlo Erba 1108) continuous flow–isotope ratio mass spectrometer (Finnigan Mat Delta S). Data are reported as per mil (%) deviations relative to the Canyon Diablo troilite (CDT) standard. Analytical uncertainty (2σ) was ±0.2‰. The experimental procedure involved the liberation of SO2 gas by rapid combustion of the samples with vanadium pentoxide. In total, 16 mineral separates, including 11 sulfates (gypsum) and 5 sulfides (late-stage pyrite) were analyzed for sulfur isotope ratios (Table 1). Finally, it should be noticed that while carbon and sulfur cycling are usually associated in the oxidation zone, they are fundamentally independent cycles.

Table 1. Representative carbon, oxygen, and sulfur isotope compositions, expressed as ‰ δ18O, δ13C, δ34S, respectively for supergene carbonate (cerussite, smithsonite, calcite) and sulfate (gypsum) minerals along with late-stage pyrite from various mineralizing stages from the Touissit-Boû Bekar (TB) and Jbel Bou Dahar (JBD) districts.

| Sample ID | Mineral | Brief Description/Paragenesis | δ13C | δ18O | δ34S | T (°C) |
|-----------|---------|--------------------------------|------|------|------|--------|
| TB1       | Cerussite | Transparent cerussite lining vug (Cer-2) | -12.95 | 12.48 | -2   |       |
| TB2       | Cerussite | Cerussite post saddle dolomite crowning on massive galena (Cer-1) | -15.09 | 12.59 | -35  |       |
| TB3       | Cerussite | Transparent cerussite encrusted on massive galena (Cer-1) | -12.76 | 10.36 | -50  |       |
| TB4.1     | Cerussite | Transparent cerussite encrusted on galena (Cer-1) | -14.21 | 20.23 | -4   |       |
| TB5       | Azurite  | Crystalline azurite associated with late fibrous malachite (Az-2) | -7.68  | 26.21 | -29  |       |
| TB6       | Azurite  | Crystalline azurite (Az-1) | -2.77  | 26.80 | -26  |       |
| TB7       | Azurite  | Crystalline azurite with subordinate malachite (Az-1) | -7.40  | 26.17 | -30  |       |
| US-TB8    | Azurite  | Azurite | -7.51  | 26.02 | -30  |       |
| US-TB9    | Azurite  | Azurite | -7.37  | 26.53 | -28  |       |
| US-TB10   | Azurite  | Azurite | -7.72  | 26.21 | -29  |       |
| US-TB11-1 | Azurite  | Azurite | -7.73  | 26.19 | -30  |       |
| TB12      | Malachite | Bothroidal malachite (Mal-2) | -4.06  | 25.70 | -24  |       |
| TB13      | Malachite | Bothroidal malachite | -5.19  | 24.85 | -25  |       |
| US-TB11-2 | Malachite | Bothroidal malachite | -4.67  | 25.24 | -23  |       |
| TB15      | Calcite  | Late stage calcite calcite encrusted on dolomite | -8.77  | 20.65 | -25  |       |
| TB16      | Calcite  | Late-stage rhombohedral calcite | -8.69  | 21.70 | -20  |       |
| TB17      | Calcite  | Late-stage rhombohedral calcite | -1.17  | 21.60 | -20  |       |
| TB18      | Calcite  | Late-stage calcite rhombohedral calcite | -3.09  | 19.20 | -32  |       |
| TB19      | Calcite  | Late-stage calcite rhombohedral calcite | -8.51  | 23.30 | -13  |       |
| TB20      | Calcite  | Late-stage calcite rhombohedral calcite | -8.51  | 21.80 | -25  |       |
| US-TB11-3 | Calcite  | Late-stage euhedral calcite | -2.89  | 22.75 | -15  |       |
| TB22      | Gypsum   | Transparent gypsum encrusted on galena (Gyp-1) | - | 0.60  | 6.70 |       |
| TB23      | Gypsum   | Fibrous gypsum encrusted on dolomite (Gyp-2) | - | 2.40  | -15.80 |       |
| TB24      | Gypsum   | Fibrous gypsum encrusted on dolomite (Gyp-2) | - | 3.70  | -11.30 |       |
| TB25      | Gypsum   | Fibrous secondary gypsum (Gyp-2) | - | - | -6.18 |       |
| TB26      | Gypsum   | Fibrous secondary gypsum (Gyp-2) | - | - | -7.25 |       |
| TB27      | Gypsum   | Fibrous secondary gypsum (Gyp-2) | - | - | -13.46 |       |
| TB28      | Gypsum   | Fibrous secondary gypsum (Gyp-2) | - | - | -14.11 |       |
| TB29      | Gypsum   | Fibrous secondary gypsum (Gyp-2) | - | - | -2.28 |       |
| TB30      | Pyrite   | Euhedral late-stage pyrite encrusted on saddle dolomite | - | - | -12.60 |       |
| TB31      | Pyrite   | Euhedral late-stage pyrite encrusted on saddle dolomite | - | - | -16.80 |       |
| TB32      | Pyrite   | Euhedral late-stage pyrite encrusted on saddle dolomite | - | - | -16.80 |       |
| TB33      | Pyrite   | Euhedral late-stage pyrite encrusted on saddle dolomite | - | - | -16.20 |       |
| TB34      | Pyrite   | Euhedral late-stage pyrite encrusted on saddle dolomite | - | - | -16.60 |       |
| JBD1      | Smithsonite | Layered smithsonite (Sm-1) | -6.44  | 24.22 | -18  |       |
| JBD2      | Smithsonite | Bothroidal smithsonite from high-grade ore (Sm-1) | -5.43  | 21.93 | -28  |       |
| JBD3      | Smithsonite | Smithsonite from mixed Pb-Zn ore (Sm-1) | -6.15  | 24.52 | -17  |       |
| JBD4      | Smithsonite | Layered red ore smithsonite (Sm-1) | -1.31  | 23.61 | -21  |       |
| JBD5      | Smithsonite | Bothroidal smithsonite (Sm-1) | -5.64  | 25.05 | -14  |       |
Table 1. Cont.

| Sample ID | Mineral   | Brief Description/Paragenesis                                      | δ13C_V_PDB | δ18O_V_SMOW | δ34S | T (°C) |
|-----------|-----------|------------------------------------------------------------------|------------|-------------|------|--------|
| JBD6      | Smithsonite | Smithsonite from mixed ore (Sm-2)                            | −5.01      | 25.70       | -    | 12     |
| JBD8      | Smithsonite | Bothroidal Smithsonite (Sm-2)                                  | −4.83      | 25.88       | -    | 11     |
| JBD9      | Calcite   | Late stage “en dents de cochon” calcite                      | −3.07      | 18.65       | -    | 34     |
| JBD10     | Calcite   | Late stage columnar calcite                                   | −8.92      | 21.55       | -    | 20     |
| JBD11     | Calcite   | Late stage stalactitic calcite encrusted on dolomitized limestone | −7.74      | 24.37       | -    | 9      |
| JBD12     | Gypsum    | Transparent gypsum associated with sphalerite                | -          | -           | 5.1  | -      |
| JBD13     | Gypsum    | Transparent gypsum associated with sphalerite                | -          | -           | 9.6  | -      |
| JBD14     | Gypsum    | Transparent gypsum associated with sphalerite                | -          | -           | 6.1  | -      |

3. Results

3.1. Introduction

In the two studied districts, partial to complete oxidation of Zn-Pb ± Cu sulfides occurs at depths ranging from a few meters from the surface down to hundreds of meters below the present water table. It is noteworthy that oxidation of the primary sulfides reveals typical vertical alteration profiles within individual deposits, with the shallow oxides of Fe, Mn, and Cu predominating in the upper part of the deposits, grading downward through the mixed non-sulfide and sulfide Zn-Pb ± Cu mineralization to the unaltered buried sulfide protore. Moreover, the supergene mineralization shows intimate spatial relationships with major faults as most of the enriched orebodies occur in the footwall of, or adjacent to, major faults, thus allowing incursion of meteoric water and oxidation of the primary ore.

3.2. Mode of Occurrence

In the Touissit-Bou Beker district, approximately 70% of the total Pb-Zn ± Cu production is derived from non-sulfide counterparts, including Pb-carbonates and Pb-sulfates (i.e., “cerussitic” ore, [5]). Some orebodies are oxidized to the extent that the only surviving sulfide is galena (Figure 5A). The most impressive example is represented by the so-called “collapse zone”, which hosts some of the higher-grade orebodies of the district, approaching 20% Pb. Of these, the clay-hosted Hassi Msidira orebody is the most intriguing one, owing to the clayey, rather than carbonaceous, nature of the host rocks and to the higher grade of the ore, averaging 18% Pb. The mineralization consists of massive blocks of galena embedded in a residual reddish-brown to yellow-brown matrix made up of finely to strongly laminated Zn-bearing clays (i.e., illite and kaolinite ± smectite) (Figure 5B). The clays contain Fe (hydr)oxide pseudomorphs after pyrite as well as non-sulfide Zn-rich carbonate minerals grading locally 4.5% Zn, of which 1.9% is represented by ZnO. Within the same horizon, the boundary between the clay-hosted ore zone is unweathered Aalenian-Bajocian gray dolostone consisting of a distinctive reddish, hematitic Zn-bearing “sandy” dolomite and remnants of original dolostone (Figure 5C).
Figure 5. Selected underground exposures and hand samples from the Touissit-Bou Beker and Jbel Bou Dahar district deposits, illustrating representative styles of non-sulfide mineralization and textural variations of different ore types. (A) Surviving cubic galena (Gn) crystal rising up in a matrix of white calamine surrounded by aggregates of botryoidal smithsonite (Sm). (B) Underground exposure at the Hassi Msidira orebody, illustrating an example of clay-hosted galena. Notice that the normally 25 m thick ore-bearing dolostone (D2) has been completely dissolved, leaving an approximately 2 m thick layer of insoluble residue made of variously colored, laminated clays, accompanied locally by massive galena clasts (blue dots). TJ = “Toit jaune”. (C) Close up of an underground exposure showing a dissolution cavity filled with clastic internal sediments (i.e., speleothems). Note that the galena fragments and dispersed in a silt- to sand-sized iron-rich dolomite referred to as “sandy dolomite” recognizable by its oxidation ochre-yellow color. Note also that the base of the cavity is lined by a fine film of black earthy cerussite “cerussite pierreuse”. (D) Hand sample of high-grade copper rich-ore showing brecciated chalcopyrite (Cpy), pyrite and hydrothermal dolomite with malachite (Mal) and hematite (Hem) sealing the fractures. Note the later development of white clays (Cl) and azurite (Az) crystals lining the remaining vugs. (E) Vuggy to cellular sponge texture derived from dissolution-replacement of coarse-grained sphalerite (Sp) and its subsequent filling. The resulting boxwork cavities are characteristically filled by late-stage calcite (CC), concretionary smithsonite and later iron oxides. (F) Close up of a cm-wide dissolution cavity filled with botryoidal smithsonite.
This gradational contact led the authors of [5] to interpret the collapse zone as internal sediment “trash zone” formed by complete meteoric dissolution of the host carbonate rocks and leaving behind clays (illite and kaolinite ± smectite), quartz, and organics as insoluble residue, along with oxidized Pb-Zn occurrences of Pb-Zn carbonates (mainly cerussite and smithsonite) and sulfates (anglesite).

Rich azurite and malachite mineralization occurs in the central part of the Beddiane mine within the upper part of the Aalenian–Bajocian dolomitic strata, close to the Late Bajocian–Lower Bathonian sandstone to carbonate hanging-wall rocks (i.e., “Toit Jaune”; [5]). Copper mineralization may also extend into the overlying Callovian-Oxfordian rocks and the “Toit Jaune”, itself for 10 m or more [5]. It is noteworthy that the carbonate-dominated copper zone is exclusively constrained to the hanging wall of the ENE–trending North and South Beddiane faults [36]. From the fault geometry and siting of the MVT orebodies, it is inferred that the South and North Beddiane fault zones constituted favorable conditions for lateral transport and deposition of copper in the faults during supergene mineralization. Supergene copper mineralization consists of surficial, locally highly silicified, Fe- and Cu-cemented breccia of malachite, azurite, chrysocolla, cuprite, native copper, atacamite, connellite, and abundant iron oxide minerals (goethite and hematite) ([5] and this present study; Figure 5D). The presence of sharp-edged azurite crystals lining within the gouge zone indicates deposition after movement along the fault. Pods of remnants’ massive chalcocpyrite and tetrahedrite have been encountered at the Beddiane mine at a depth of 180 m [36].

Unlike the Touissit-Bou Beker district, where Pb dominates largely over Zn in a mass ratio of 5:1, the Jbel Bou Dahar district contains the largest non-sulfide zinc orebodies, which constitute the historical basis for the development of calamine mining in Morocco. The Jbel Bou Dahar district is also a special case, as both Pb-Zn ± Cu sulfide and non-sulfide ores ± barite are currently mined from the numerous mine workings scattered along the flanks of the reefal structure (Figure 3). Calamine grade is highly variable throughout the mining district, ranging from a few percent of combined Zn-Pb to more than 30% ZnO in the most enriched ore. In this respect, it should be emphasized that the minimum calamine ore grade required to be calcined by the CADETAF facilities is 18% ZnO, which ultimately allows enrichment up to 45% ZnO through roasting. From the mined open pits, digs, and geologic observations, it can be inferred that the calamine ore extends downward as deep as >200 m, but only calamine orebodies located above the present water table have been or are being exploited by artisanal miners.

In all the studied deposits, non-sulfide mineralization occurs as (1) strata-bound tabular blankets and pods of variable thickness, width, and strike extent and disseminations of millimeter- to centimeter-scale patches resulting from the replacement of both the primary sulfides and the Jurassic host carbonates, (2) oxide boxworks (after sphalerite, Figure 5E), and (3) botryoidal concretions (Figure 5F) to columnar aggregate infills (Figure 6A) of interconnected karst cavities, vugs, veins, and fractures. The dominant ore zone geometry is strata-bound, and boundaries of mineralization correspond with the weathering front within fresh dolomitic limestone. The higher-grade ore zones are deeply oxidized with karstic cavities or fissures filled with variously colored clays, organic debris, lithic fragments derived from the overlying weathered carbonates, and speleothems (Figure 5A). The supergene ore shows different stages of in situ oxidation and replacement of the primary Pb-Zn-bearing sulfide mineralization and therefore could be assigned to direct replacement type of supergene mineralization [3]. In comparison with replacement ore, for most of the higher-grade orebodies, open-space filling mineralization accounts for a significant proportion of the recovered supergene Pb-Zn ore.
Figure 6. Selected representative rock slab photographs illustrating mineralogical and paragenetic attributes, styles of open-space filling mineralization, and textural variations of the non-sulfide ores from the Jbel Bou Dahar district. (A) Close up of a columnar “palissadic” late-stage calcite. (B) Bedded crustiform calamine ore showing crosscutting relationships. Note the stratiform texture shown by early smithsonite 1 which consists of a succession of subhorizontal white to grayish translucent smithsonite layers which are crosscutted by veinlet of later hydrozincite (Hyz). (C) Stromatolitic-like textured calamine crosscutted by thin veinlets of galena, with this latter being rimmed by dark cerussite (Cer). (D) Massive red calamine ore showing the coexistence of black smithsonite and iron hydr(oxydes) (Gth, Hem). (E) White calamine ore showing stromatolitic-like texture overprinted by the development of numerous mm-sized vacuoles interpreted as “bird-eyes” resulting from CO₂-degassing. (F) Close up of a crustiform stromatolitic-like textured white calamine ore showing laminations of smithsonite and hydrozincite.
3.3. Mineralogy and Geochemistry

Regardless of the considered deposit and/or district, the oxidized mineral paragenesis consists predominantly of variable proportions of cerussite (PbCO$_3$) and smithsonite (ZnCO$_3$), whose relative abundances are primarily controlled by the Pb:Zn mass ratios of the primary sulfide ore (i.e., Zn-dominant vs. Pb-dominant deposits). Subordinate minerals include anglesite (PbSO$_4$), hydrozincite (Zn$_5$(CO$_3$)$_2$(OH)$_3$), hemimorphite (Zn$_4$Si$_2$O$_7$(OH)$_2$·H$_2$O), willemitc (Zn$_2$SiO$_4$), malachite, azurite, and Mn-(hydr)oxides. Calcite, aragonite, gypsum, barite, celestine, and quartz are the most common gangue minerals. These mineral assemblages are typical of those characterizing non-sulfide occurrences derived from MVT deposits (3). Exotic mineral species include vanadinite and wulfenite, and, to lesser extent, pyromorphite, manganese oxides (coronadite, cryptomelane, and hollandite), motttramite (PbCu(VO$_4$)(OH)), paralaurionite (PbCl(OH)), phosgenite (Pb$_2$CO$_3$Cl$_2$), and sauconite ((Na$_{0.3}$Zn$_3$(Si, Al)$_4$O$_{10}$(OH)$_2$·4(H$_2$O)) [26,30]. In partially oxidized orebodies, remnants of minor sphalerite, chalcopyrite, and less commonly pyrite and abundant galena are locally preserved within the non-sulfide orebodies being enclosed within their unaltered carbonate gangue (calcite and dolomite).

The predominant non-sulfide zinc mineral of economic significance is smithsonite, which has been observed in various proportions in every deposit though the higher abundances are reported in the Jbel Bou Dahar mines. In addition to smithsonite, the complete sequence of Zn-bearing phases that collectively define the “calamine” group (Figure 6B,C) include hydrozincite, willemitc, hemimorphite, and sauconite, with variable amounts of Fe (hydr)oxides (goethite and hematite), cerussite, and anglesite. Owing to their mineralogy, the non-sulfide zinc deposits in the Jbel Bou Dahar district and by inference over the entire central and eastern High Atlas are assigned to the carbonate-hosted “calamine” class of [1], in which smithsonite, hydrozincite, willemite, and hemimorphite are the principal zinc-bearing minerals.

Based on the mode of occurrence of mineralization (i.e., replacement vs. open-space-filling), ore textures, and Zn-bearing mineral assemblages, two distinct types of calamine referred to as “red” and “gray to white” calamine ores are distinguished (Figure 6D,E). The red calamine occurrences are widespread throughout the studied districts and are easily recognizable in surface outcrops owing to their distinctive detrital, “gossanous”, and “earthy” appearance, exhibiting locally spongy textures. Their reddish to yellowish color is imputed to the widespread development of superficial blankets of Fe oxyhydroxides (hematite, goethite), which derived from oxidation of preexistent pyrite and/or marcasite. Conversely, the gray to white calamine occurrences are recognizable by their milky chalky appearance although locally transparent to translucent. Textural relationships indicate that the emplacement of the white calamine occurred after the development of the red calamine, in agreement with all the established chronologic reconstitutions for calamine deposits worldwide [6,19,30,64,65]. Structural constraints indicate that the emplacement of calamine ore is structurally controlled as most of the orebodies are distributed along the major ENE-trending faults.

Smithsonite is present in all deposits, occurring as fine disseminations variably replacing the sphalerite grains and the carbonate groundmass, as open space infills, and as concretionary crust on dolomitic carbonates (Figure 5F). Direct evidence for the replacive nature of smithsonite is provided by the vertical zonation of the Pb-Zn ore, which shows an underlying buried Pb-Zn sulfide protore grading upward to an overlying cap of carbonate, iron oxides, and oxyhydroxides ore through a mixed zone of sulfide-carbonate ore. Two generations of smithsonite are distinguished. Smithsonite-1 (Sm-1), which is not very common, is paragenetically related to the “red calamine ore”, being intimately associated with Fe oxyhydroxides. It consists of botryoidal aggregates of brownish-red color crystals with a rugged appearance and is chemically impure with FeO and CdO contents below 1 wt% [5]. Sm-1 shows evidence of in situ oxidation and progressive replacement of both the primary sulfides and the carbonate groundmass. In contrast to Sm-1, the second generation of smithsonite (Sm-2) occurs as transparent rice-shaped infills [65] of microkarsts,
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vein, and fracture networks and as grayish to whitish concretionary aggregates lining walls of the cavities (Figure 5F) and the remnant porosity. Geochemically, Sm-2 displays a quite pure chemical composition approaching the idealized formula ZnCO$_3$.

Primary sphalerite, from which the two generations of smithsonite derived, is characterized by low concentrations of Fe (122–6271 ppm, average = 1349 ± 1775 ppm), Cd (298–1462 ppm, average = 814 ± 38 ppm), As (0.1–84.1 ppm, average = 9.5 ± 25.0 ppm), Sb (0.0–56.6 ppm, average = 9.7 ± 16.9 ppm), and Pb (0.1–321.8 ppm; average = 70.9 ± 96.5 ppm). Ge (0.28 ± 0.04 ppm) and Ga (5.84 ± 8.46 ppm) (Table 2) show much lower concentrations than those abundances reported by [31] for the calamine ore (willemite), which approach 1000 ppm for Ge.

**Table 2.** Representative trace element compositions of sphalerite from Jbel Bou Dahar (JBD) district.

| Sample No. | Mn  | Fe      | Cd      | Sb  | Cu  | Pb   | As  | Ag  | Co  | In  | Ga  | Ge  | Se  | TI  | Bi  |
|------------|-----|---------|---------|-----|-----|------|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| JBD-Sp1    | 2.9 | 1285.9  | 298.1   | 1.2 | 32.1| 77.8 | 3.3 | 0.03| 11.9| 0.00| 1.59| 0.02| 0.00 | 0.00 | 0.00 |
| JBD-Sp2    | 1.8 | 586.4   | 333.2   | 7.7 | 39.1| 154.4| 2.3 | 0.81| 19.4| 0.00| 0.03| 0.01| 0.001| 0.001| 0.00 |
| JBD-Sp3    | 0.1 | 121.6   | 471.9   | 0.2 | 33.7| 0.1  | 0.1 | 0.03| 12.6| 0.01| 22.08| 0.00| 0.031| 0.000| 0.000|
| JBD-Sp4    | 0.1 | 131.9   | 551.4   | 0.3 | 33.8| 0.1  | 0.1 | 0.02| 13.6| 0.02| 22.18| 0.00| 0.030| 0.000| 0.000|
| JBD-Sp5    | 0.9 | 298.9   | 1265.5  | 0.1 | 1.6 | 10.3 | 0.1 | 0.04| 22.5| 0.01| 2.01 | 0.01| 0.047| 0.000| 0.000|
| JBD-Sp6    | 6.1 | 1696.9  | 877.1   | 5.8 | 31.7| 30.6 | 1.2 | 0.05| 23.2| 0.00| 0.46 | 0.02| 0.002| 0.001| 0.000|
| JBD-Sp7    | 0.9 | 372.6   | 799.4   | 4.8 | 17.6| 15.8 | 0.2 | 0.05| 44.3| 0.00| 8.36 | 0.01| 0.001| 0.000| 0.000|
| JBD-Sp8    | 1.4 | 390.1   | 926.1   | 0.8 | 6.6 | 102.4| 1.1 | 0.02| 37.1| 0.00| 5.86 | 0.01| 0.001| 0.000| 0.000|
| JBD-Sp9    | 5.2 | 1570.1  | 854.1   | 5.5 | 25.2| 321.8| 1.1 | 0.08| 24.3| 0.00| 0.98 | 0.03| 0.008| 0.000| 0.001|
| JBD-Sp10   | 6.8 | 2110.6  | 1462.2  | 56.6| 98.8| 107  | 11.4| 3.82| 39.3| 0.01| 0.38 | 0.03| 0.002| 0.011| 0.000|
| JBD-Sp11   | 5.6 | 6271.3  | 1112.1  | 23.5| 183.2| 55.9 | 84.1| 0.21| 78.9| 0.01| 0.35 | 0.14| 0.002| 0.004| 0.001|

Hydrozincite is widespread in all deposits of Jbel Bou Dahar district and to a lesser extent in the Touissit-Bou Beker district. Commonly associated with smithsonite, hydrozincite occurs as needle-shaped crystals locally replacing smithsonite and sphalerite directly and also as whitish millimeter- to decimeter-thick alternating laminae (banded white calamine) defining a “stromatolitic-like” texture (Figure 6F), accompanied locally by botryoidal aggregates filling the remaining cavities and voids on smithsonite.

As for smithsonite and hydrozincite, cerussite has been observed in all deposits, particularly in the Touissit-Bou Beker district, where this mineral constitutes the principal Pb-carbonate phase of economic significance. Where observed, cerussite, together with anglesite, occurs as (i) dark gray massive to partial replacement of the dolomitic host rocks and the associated galena ore, giving rise to earthy or stony “cerussitic” ore (i.e., “cerussite pierreuse” of [5], Cer-1) (Figures 5B and 6C), or (ii) vugs, fractures, and remnant porosity infills, mostly as euhedral crystals of variable dimension (up to several cm) (Cer-2) filling irregularly distributed druses within the massive galena orebodies down to 150 m in depth (Figure 7A). Cerussite and anglesite are also observed together within fractures in galena and along grain boundaries. The earliest replacive stony or earthy cerussite (Cer-1) is not chemically pure and contains oxide concentrations that can locally reach 8–10 wt.% MgO, up to ~3 wt.% MnO, up to 5–6 wt.% FeO, and up to 6 wt.% CaO. The late cerussite (Cer-2), however, is chemically pure, showing little variation in chemistry and approaches the idealized formula CaCO$_3$. 
Figure 7. Representative photographs of hand specimens (A–C, E–H) and reflected light microscopy (D) displaying representative aspects of non-sulfide mineralogy, textural variations, and paragenetic attributes from the Touissit-Bou Beker and Jbel Bou Dahar districts. (A). Close up of a cm-wide dissolution cavity filled with euhedral transparent cerussite (Cer, Dol = Dolomite). (B) Botryoidal malachite. (C) Cauliflower-textured azurite. (D) Reflected plane-polarized light photomicrograph of brecciated and altered chalcopyrite (Cpy) with covellite (Cv) and the fractures and grain boundaries. Note also the later development of hematite (Hem) needles within the remaining vugs. (E) Late-stage calcite (Cc-2) encrusted on galena (Gn). (F) Close up of radiating aragonite (Arg) cementing cm-sized fragments of galena partially oxidized to cerussite. (G) Close up of transparent tabular gypsum (Gyp-1) intimately associated with sphalerite (Sp). (H) Late-stage “miscellaneous” to “wired gypsum” (Gyp-2).
Malachite and azurite (Figure 7B,C), along with covellite (Figure 7D) and chalcocite, occur as partial to complete replacement of Cu-bearing minerals, especially of chalcopyrite and Ag-rich tetrahedrite [5] and are generally associated with cerussite and anglesite. A complex association of iron and manganese oxy-hydroxides (e.g., goethite, hematite, lepidocrocite) and residual Zn-rich clays complete the list of the recognized supergene mineral species. Subordinate mineral species include a wide panel of supergene phases including carbonates silicates (hemimorphite, chrysocolla), oxides (psilomelane, wulfenite, and cuprite), rare phosphates (pyromorphite, scholzite), vanadates (vanadinite, desclozite, and mottramite), arsenates (beudantite, mimetite), and native copper [5]. The main gangue minerals include late-stage calcite (Figure 7E), aragonite (Figure 7F), quartz, gypsum (Figure 7G,H), and celestine. Gypsum, which is widespread in the Touissit-Bou Beker district but less common in Jbel Bou Dahar deposits, occurs either as transparent to translucent euhedral crystals intimately associated with primary sulfides (sphalerite and galena, Figure 7G) (Gypsum 1, Gyp-1) or as replacement crust and stalactitic encrustations lining cave walls (Gypsum 2, Gyp-2) formed at the contact of sulfuric acid and carbonate rocks (Figure 7H). Similarly, late-stage calcite shows different habits, varying from columnar-like radiating aggregates (Figure 6A) through botryoidal precipitates to rhombohedral/scalenohedral crystals (Figure 7E) filling cavities, veins, and fractures.

3.4. Paragenetic Sequence

Although no consistent paragenetic sequence can be defined, as the supergene mineral assemblages appear to be the product of multiple overlapping pulses, an idealized paragenetic sequence of mineral deposition within the non-sulfide Pb-Zn ± Cu mineralization for all deposits is proposed (Figure 8). In summary, the sequence of mineral deposition shows the existence of two main stages of supergene mineralization referred to as “red calamine” and “white calamine” ore stages (Figure 8). A further earlier event characterized by weathering-related processes is also recognized. The proposed succession is consistent with those paragenetic sequences drawn for non-sulfide Pb-Zn ± Cu deposits worldwide [19,30,66–68].

The earliest stage, referred to as pre-non-sulfide immature “gossanous” stage I, corresponds to the oxidation of pyrite and subsequent formation of yellowish to reddish massive to pulverulent mixture of Fe oxi-hydroxides (hematite, goethite) with a variable proportion of subordinate silicates. The second stage, which constitutes the main stage of sulfide oxidation, accounts for a significant proportion of the calamine ore recovered from all the deposits of the High Atlas calamine province. The initiation of large-scale supergene zone-refining reactions leads to a progressive in situ replacement of all the preexistent sulfides with a very limited transport of the released base metals by sulfide dissolution from the original protore zone. The resulting mineral paragenesis consists predominantly of smithsonite-1, cerussite-1 and anglesite-1, accompanied by various amounts of willemite-1 and Fe oxy-hydroxides (goethite and hematite principally). Sulfide remnants (minor sphalerite and abundant galena) are commonly observed within the non-sulfide protore, together with the unaltered carbonate gangue (calcite and dolomite).

The latest stage of sulfide oxidation is economically the most attractive as it accounts for most of the higher-grade Zn-rich orebodies in all of the investigated deposits of the calamine province of the Central High Atlas. The mineralization occurs as late-phase fillings of cavities, fractures and cementing breccias as well as replacements of earlier phases. The mineralogy is dominated by smithsonite-2, cerussite-2, anglesite-2, and, more importantly, hydrozincite, which characterizes this stage. Owing to their high mobility in comparison to lead, zinc ions were transported farther away from the sulfide precursor and then precipitated distally in cavities or as internal sediment infillings. The resulting wall-rock replacement mineralization (i.e., white calamine ore) is Zn-rich and correlatively Pb- and Fe-poor, in agreement with previous works of [3,16].
3.5. Carbon and Oxygen Isotope Geochemistry

The carbon and oxygen isotope datasets comprise a total of 39 samples, including 4 cerussite, 8 smithsonite, 10 calcite, 8 azurite, and 3 malachite. A further set of four gypsum specimens and two single crystals of anglesite and vanadinite were also analyzed for their oxygen isotopic compositions. The recorded $\delta^{13}$C and $\delta^{18}$O values are indicated in Table 1 and plotted in Figures 9 and 10.
Figure 9. $\delta^{13}C_{\text{VPDB}}$ vs. $\delta^{18}O_{\text{V-SMOW}}$ data for supergene carbonate minerals (cerussite, smithsonite, calcite) from various mineralizing stages from the Touissit-Bou Beker and Jbel Bou Dahar districts. The compositional fields of smithsonite (dark grey), calcite (light grey), and cerussite (pink colored) from supergene Zn-Pb deposits [59] are indicated for comparison. Composition of Middle Jurassic marine limestone is from [69,70]. Arrows show the generalized burial, and meteoric diagenetic trends for carbonates are from [71].

Figure 10. $\delta^{13}C_{\text{VPDB}}$ vs. $\delta^{18}O_{\text{V-SMOW}}$ data for natural azurite and malachite from various mineralizing stages from the Touissit-Bou Beker district. The compositional fields of caliche and speleothem are from [62].
The δ\(^{13}\)C\(_{\text{PDB}}\) and δ\(^{18}\)O\(_{\text{SMOW}}\) values of cerussite range from −15.1 to −12.8‰ (average = −13.7 ± 1.1‰, n = 4) and 10.4 to 20.2‰ (average = 13.9 ± 4.3‰, n = 4), respectively. The δ\(^{13}\)C\(_{\text{PDB}}\) and δ\(^{18}\)O\(_{\text{SMOW}}\) values for smithsonite range from −6.4 to −1.4‰ (average = −5.0 ± 1.7‰, n = 7) and 21.9 to 25.9‰ (average = 25.2 ± 2.2‰, n = 7), respectively (Table 1). Overall, the isotopic compositions of smithsonite fall within the compositional field of the typical supergene smithsonite (Figure 9), although some of the isotopic compositions plot outside this field, toward lower δ\(^{18}\)O\(_{\text{SMOW}}\) values. Conversely, none of the δ\(^{18}\)O\(_{\text{SMOW}}\) values of cerussite fit within the range of typical isotopic compositions that characterize the compositional field of supergene cerussite (Figure 9). Instead, most of the oxygen isotope signatures of cerussite are shifted toward lower oxygen isotope ratios characteristic of a hydrothermal overprint, but δ\(^{13}\)C\(_{\text{PDB}}\) ratios typical of supergene cerussite (Table 1; Figure 9). Such isotopic patterns have also been documented for cerussite from the Tunisian Bou Jaber deposit [72].

The azurite crystals from the Touissit-Bou Beker district yield δ\(^{18}\)O\(_{\text{SMOW}}\) values of 26.2 to 26.8‰ (average = 26.3 ± 0.2‰, n = 7), whereas their corresponding δ\(^{13}\)C ratios range from −7.7 to −2.8‰ (average = −6.9 ± 1.8‰, n = 7) (Table 1). The three malachite specimens show δ\(^{13}\)C and δ\(^{18}\)O\(_{\text{SMOW}}\) values of −5.2 to −4.1‰ (average = −4.6 ± 0.6‰, n = 3) and 25.2 to 25.7‰ (average = 25.3 ± 0.4‰, n = 7), respectively. Overall, these isotopic compositions are similar to those determined for natural azurite and malachite worldwide and plot within the caliche field, as defined by [62] (Figure 10). The single anglesite and vanadinite samples display, however, contrasting δ\(^{18}\)O\(_{\text{SMOW}}\) values of 12.5 and 33.3‰, respectively (Table 1).

The late fracture- and cavity-filling calcite from both the Touissit-Bou Beker and Jbel Bou Dahar districts display overlapping δ\(^{13}\)C\(_{\text{PDB}}\) values between −8.9 and −12.2‰ (average = −6.1 ± 3.1‰, n = 10) and δ\(^{18}\)O\(_{\text{SMOW}}\) ratios between 18.6 and 24.4‰ (average = 21.5 ± 1.7‰, n = 10) (Figure 9). Overall, these isotopic compositions plot within the compositional range that characterizes supergene late-stage calcite elsewhere. In comparison to the Jurassic host carbonates and associated hydrothermal dolomite and calcite [5,29], the isotopic compositions of late-stage calcite show substantially lower and distinctive negative δ\(^{13}\)C values (Figure 9). Of particular interest is the temporal trend displayed by the distribution of δ\(^{13}\)C and δ\(^{18}\)O ratios of smithsonite and cerussite, which shows a concomitant increase in δ\(^{13}\)C and δ\(^{18}\)O values with advancing paragenetic sequence (i.e., from Sm-1 and Cer-1 to Sm-2 and Cer-2) (Figure 9). A similar pattern is also displayed by the distribution of δ\(^{13}\)C and δ\(^{18}\)O values of coexisting azurite, malachite, and late-stage calcite, which shows an overall increase in δ\(^{13}\)C and δ\(^{18}\)O values from older azurite to younger calcite through intermediate malachite (Table 1).

Finally, gypsum from both the Touissit-Bou Beker and Jbel Bou Dahar districts has δ\(^{18}\)O values of 21.9 to 0.6‰ (mean = 7.0 ± 9.6‰, n = 4). Similar to the evolutionary trends displayed by the δ\(^{13}\)C and δ\(^{18}\)O values of smithsonite and cerussite, as well as azurite, malachite, and late-stage calcite, the distribution of the oxygen isotopic compositions of gypsum show distinct temporal variation. This variation is expressed by an overall increase in δ\(^{18}\)O\(_{\text{SMOW}}\) values with advancing paragenetic sequence toward younger sulfate generation. In this respect, it is noteworthy that early gypsum (Gyp-1) derived from in situ oxidation of sphalerite displays δ\(^{18}\)O\(_{\text{SMOW}}\) value of 0.6‰ whereas late-stage wall-rock replacement gypsum (Gyp-2) yields lower δ\(^{18}\)O\(_{\text{SMOW}}\) ratios of 2.4 and 3.7‰ (Table 1).

3.6. Sulfur Isotope Geochemistry

A total of 11 gypsum specimens along with five late-stage pyrite separates from the different non-sulfide orebodies of the Touissit-Bou Beker and Jbel Bou Dahar districts were analyzed for their sulfur isotope compositions (Table 1).

The gypsum samples display a wide range of δ\(^{34}\)S values from −15.8 to 9.6‰ (average = −3.5 ± 9.6‰, n = 11), with the highest ratios characterizing the early gypsum (i.e., Gyp-1) from the red calamine ore stage and the lowest values from the late wall-rock replacement gypsum (i.e., Gyp-2) (Table 1). In this respect, it is noteworthy that all of the δ\(^{34}\)S values of
Gyp-1 are positive, ranging from 2.3 to 9.6‰ (average = 6.0 ± 2.6‰, n = 5), whereas values from Gyp-2 are all negative and vary between −15.8 and −6.2‰ (average = −11.3 ± 3.9‰, n = 6). The δ³⁴S values of late-stage pyrite whose precipitation is related to the collapse of the hydrothermal MVT system and the initial episode of exhumation and weathering of sulfides are all negative, ranging from −16.8 to −12.6‰ (average = −15.8 ± 1.8‰, n = 5). Overall, δ³⁴S values of gypsum and pyrite are distinctly below the range of sulfur isotopic compositions for Jurassic seawater sulfate (i.e., 13 to 20‰; [73,74]).

4. Discussion

Owing to their strong buffering capacity, carbonate host rocks constitute ideal protoliths for the formation of supergene non-sulfide Zn-Pb ores. The distribution of the resulting mineral assemblages is controlled primarily by the initial mineralogy; the composition and flow rates of groundwater; the tectonic evolution, which triggers uplift; uncapping and oxidation of sulfide-rich protore; and climate conditions prevailing at the time of sulfide alteration [1,3,13]. The presence of appreciable amounts of pyrite in the alteration zones seems to constitute the best conditions for supergene enrichment as the oxidation of pyrite makes the percolating water more acidic and consequently the replacement of sulfides by supergene minerals more effective. Moreover, fluctuation in the depth of the water table is also an important factor that triggers supergene alteration as the rise and fall of the water table introduces oxygenated water to the primary sulfide body, driving oxidation reactions. Similarly, a climate with alternating wet and dry seasons could promote the formation of weathering and oxidizing solutions, which ultimately would lead to the development of karst networks and replacement of the sulfides [3]. Equally important, uplift and long-wavelength vertical movements [15,32,75–78] may lead to enhanced fracturing of the host rock, creating preferential pathways for the downward and lateral flow of metal-bearing groundwaters and consequently migration of the supergene alteration front.

4.1. Origin of the Supergene Mineralizing Fluid(s)

Field observations along with mineralogic and petrographic examinations show that the primary Pb-Zn ± Cu ores are intensely weathered and strongly oxidized. Textural evidences for oxidation and replacement of primary sulfides by low-temperature and low-pressure non-sulfide mineral assemblages include development of reaction fronts, the stalactitic, columnar, and concretionary structures shown by late-stage calcite, smithsonite, and hydrozincite. Further evidence for the involvement of widespread and extensive recent meteoric alteration as the trigger for the oxidation of primary sulfide mineralization includes the development of vertical karsts displaying morphological features associated with normal meteoric speleogenesis (soil carbonates and speleothems). Additionally, the intense and widespread reddening of surface outcrops and internal sediment infills, the ubiquitous occurrence of sandy dolomite, and more importantly the extremely light carbon and oxygen isotope compositions shown by late-stage calcite, which indicate meteoric oxidation of organic matter as shown below, reflect meteoric alteration. Altogether, these textural and mineralogic characteristics indicate that the non-sulfide mineralization from the Touissit-Bou Beker and Jbel Bou Dahar districts precipitated from downward percolating groundwaters of meteoric origin in either a supergene or a low-temperature environment.

In this respect, carbon and oxygen isotope compositions of the Pb-Zn ± Cu-bearing carbonates provide useful information on the fluid characteristics responsible for precipitation of non-sulfide minerals and indirectly constrain the climatic environment that prevailed at the time of deposition (e.g., [19] and references therein). The obtained δ¹³C and δ¹⁸O values of smithsonite, cerussite, azurite, malachite, and late-stage calcite from the two investigated districts form distinct clusters whose overall isotopic signatures plot in the supergene compositional fields of cerussite, smithsonite, azurite-malachite, and late-stage calcite as defined by [59,62] (Figures 9 and 10). The carbon and oxygen isotope compositions of the Jurassic carbonate host rocks are, however, consistent with
altered carbonates that underwent isotopic re-equilibration during deep burial fluid–rock interaction [5] (Figure 9).

Assuming that the coexisting cerussite and smithsonite precipitated under equilibrium exchange conditions, the oxygen isotope composition and related temperature of the supergene ore-forming fluid ($\delta^{18}O_{V-SMOW}$) can be calculated using the cerussite-water (1000 ln $\alpha_{\text{cerussite-water}} = 2.29(10^6/T^2) - 3.56$) fractionation equation of [59] and the smithsonite-water (1000 ln $\alpha_{\text{smithsonite-water}} = 3.21(10^6/T^2) - 3.63$) equation of [60] (T given in Kelvin). By doing so, the calculated $\delta^{18}O_{V-SMOW}$ and related temperature from the paired samples TB4.1 and TB4.2 (Table 1) are $-8.6\%$ and 25 °C, respectively. In this respect, it is noteworthy that the calculated $\delta^{18}O_{V-SMOW}$ and ambient temperature values match perfectly, within error, the mean isotopic value of $-8\%$, which characterizes the present-day $\delta^{18}O$ value of meteoric waters in the Atlas system [79–81]. More importantly, the oxygen isotope thermometry of azurite, malachite and late-stage calcite, listed in order of their appearance from oldest to youngest and assuming a local meteoric water value of $-8\%$, indicates decreasing temperature of formation over time from 34 to 19 °C (Table 1). Overall, these calculated temperatures are similar to those determined for smithsonite, cerussite, and azurite-malachite in other base metal supergene deposits worldwide (11–35 °C, avg 20 °C; [13,59,68]) and are in agreement with the continental environment in a temperate climate [68,82]. Cerussite oxygen isotope thermometry, suggesting temperatures above ambient air temperatures (>30 °C) (Table 1), reflects minor heat production by exothermic oxidation of sulfides.

4.2. Source(s) of Carbon and Sulfur: Evidence for Microbial Activity

The broad range of $\delta^{13}C$ values of smithsonite, cerussite, azurite, malachite, and late-stage calcite, ranging from $-1.2\%$ to as low as $-15.1\%$ (Table 1), indicates contributions from various carbon sources. In this regard, it is noteworthy that the obtained $\delta^{13}C$ ratios are intermediate between the isotopic compositions of Jurassic marine limestones (i.e., 0–2‰; [69,70]), organic matter ($\delta^{13}C = -24\%$; [5]), and atmospheric carbon ($\delta^{13}C = -7\%$; [83]) (Figure 9). The negative values recorded for $\delta^{13}C$ (Table 1) suggest that most of the carbon present during supergene oxidation and subsequent Pb-Zn ± Cu carbonate formation was derived from different degrees of mixing along the fluid flow path between a $^{13}C$-depleted carbon source derived from the decomposition of C3 ($\delta^{13}C = -32.0$ to $-25.0\%$; [84]) and/or C4 surface vegetation ($\delta^{13}C = -14.0$ to $-10.0\%$; [85]), atmospheric carbon ($\delta^{13}C = -7\%$; [83]), and dissolution of the Jurassic carbonate host rocks. While the dissolution of the local carbonate rocks and atmospheric carbon undoubtedly makes a contribution, there is a clear relationship between oxygen and carbon isotope values that indicate a diagenetic or lithification overprint (Figure 9). Irrespective of the exact mechanism for such trends, the leading contenders are bacteria and archaea metabolism and deep basin fluid mixing, e.g., [64,65].

In this respect, it is well documented that colonies of chemolithotrophic bacteria and archaea (i) greatly accelerate the alteration of primary sulfide minerals (e.g., [9,86,87]), and (ii) are associated with the formation of speleothems in meteoric environment [23,24]. As the precipitation of azurite requires partial pressures of CO$_2$ that greatly exceed present atmospheric levels [86–88], this precludes a carbon source in equilibrium with the atmosphere for at least a portion of the supergene period. Indeed, the lighter $\delta^{13}C$ compositions displayed by all the analyzed carbonates from the studied districts (Table 1) are suggestive of the involvement of significant biological carbon input in the mineralizing process. Further support for the involvement of a biological source in the formation of the supergene mineralization at the Touissit-Bou Beker and Jbel Bou Dahar districts is provided by the light sulfur isotopic compositions shown by Gyp-2 (Table 1). In fact, the highly negative $\delta^{34}S_{\text{Gyp-2}}$ values of $-15.8\%$ to $-6.2\%$ are interpreted to have resulted from isotopic fractionation during multi-stage microbial oxidation of primary sulfide, at different rates of oxygen availability at or near the water table (open vs. closed reservoirs with respect to oxygen), given that most bacteria and archaea associated with supergene processes are
found near the water table and are thermophilic, flourishing deep in the oxidation zone at temperatures above ambient (e.g., [89]). Accordingly, Gyp-2 is interpreted to have been produced by microbially mediated oxidation of sulfides as a result of the reaction between sulfuric acid (H$_2$SO$_4$) and the carbonate host rocks. In this regard, it is well documented that phototrophic sulfide oxidation can cause $^{34}$S depletions of up to $\sim$4‰, or even higher, for $\delta^{34}$S [90], consistent with the sulfur isotope data reported in this study.

Moreover, the occurrence of the “stromatolitic” and “peloïdal” textures exhibited by the Jbel Bou Dahar “calamine” ore (Figures 6F and 11A), which are remarkably similar to the present-day-forming stromatolites [91], provide ample evidence for the involvement of a biologically controlled mechanism as the main driving process that triggered the supergene mineralization at Jbel Bou Dahar and Touissil-Bou Beker districts. Even though these fabrics resemble the collomorph textures commonly described in both hydrothermal and supergene deposits, their microbial-mediated origin has been demonstrated (e.g., [22] and references therein). In fact, the development of these laminated rhythmically banded macrostructures that are very similar to nanotextures observed in recent bacterial biofilms are interpreted as fossil microbial mats that resulted from in situ metabolic products of sulfate-reducing/oxidizing bacteria and archaea (e.g., [22–24]). Similarly, the aforementioned macro- and microstructures are comparable to those described in biologically controlled hydrozincite-bearing mineralization whose genesis has been linked to photosynthetic alkalization of the cyanobacterial cell surface during the seasonal alternation between spring and summer [92–94]. More interestingly, the stromatolitic- to peloïdal-like textured smithsonite, which consists of a layered succession of alternating mm- to cm-thick laminae of white and colorless hydrozincite and smithsonite-2 (Figure 11B,C), could reflect seasonal variations in temperature and fluctuations in pH. In this regard, EDS analyses show that compositional variations in Ca, O, and Zn contents (Figure 11D) between the alternating white and colorless microfabrics could be attributed to calcification of cyanobacterial sheaths related to photosynthetic activity, which, in turn, is related to seasonal variations. While debate remains open on the utility of texture as a biogenic indicator, and despite the fact that interpretation of the aforementioned textures may evolve over time, our observations should be considered for the benefit of future researchers and their interpretations.

In summary, we suggest that the combined widespread occurrence of the biogenic rhythmically banded macrotextures, along with the lighter carbon and sulfur isotope compositions shown by the analyzed carbonates, are consistent with the involvement of chemolithotrophic bacteria and archaea in the formation of the Atlasic supergene carbonate-hosted Pb-Zn mineralization.
Figure 11. Hand samples and cathodoluminescence images of the banded white calamine ore. (A) Hand sample of white calamine ore from the Jbel Bou Dahar district showing a well-developed “oolitic” to “oncolytic” texture similar to the columnar build-ups produced by modern shallow subtidal stromatolites [90]. (B) Cathodoluminescence (CL) microphotograph of white calamine ore from Jbel Bou Dahar district showing a stromatolitic-like texture materialized by an alternation of mm-thick crusts of smithsonite and hydrozincite. The laminations (i.e., crusts) are interpreted to represent calcified biofilms (Type 2 mats) as defined by [90]. (C) Closeup of the layered stromatolitic-like textured white calamine ore from the Jbel Bou Dahar district showing cycling between smithsonite and hydrozincite. (D) Distribution of Ca, O, and Zn abundances as a function of layering.

4.3. Proposed Evolution of the Atlasic Non-Sulfide Pb-Zn ± Cu Deposits

Field evidence, mineralogical observations, and stable isotope (C, O, S) geochemistry indicate that the supergene Pb-Zn ± Cu mineralization at Touissit-Bou Beker and Jbel Bou Dahar districts occurred during two main contrasting supergene mineralizing events (i.e., “red calamine” vs. “white calamine” stages). Both events were preceded by development of an earlier immature “gossanous” weathering-related stage. All three stages probably occurred synchronously with the last uplift stage related to the Atlasic orogeny, given that the non-sulfide mineralization shows no evidence of deformation and is therefore interpreted to postdate the Atlasic regional metamorphism and deformation of the Jurassic carbonate rocks.

The exhumation of the Jurassic carbonate platform and the associated sulfide orebodies from late Miocene to present facilitated the downward infiltration of meteoric fluids, oxidation of sulfides and subsequent non-sulfide formation. The resulting low-temperature, low-pressure mineral assemblages precipitated predominantly as open-space infills (pore space, veins, open fractures, and breccia zones) and also as replacement of carbonate precursors. Although the age(s) of supergene alteration in the studied districts remains poorly constrained, published radiometric ages (Ar-Ar on jarosite, [95] and refer-
ences therein [96,97]) along with thermochronological data from adjacent lithostructural domains [42–44,96–101] indicate that the emplacement of the non-sulfide mineralization likely occurred during the late Miocene (Messinian–Tortonian), Pliocene, and between the late Pleistocene and Holocene, under wet to semi-arid to arid climatic conditions. Arid conditions seem to have prevailed during the last tens of millions of years [102,103] in the Mediterranean area, although alternating wet and arid conditions have been reported [104,105]. Uplift and increasing aridity would progressively lower water tables, possibly with substantial seasonal fluctuations in a monsoonal climate, enhancing the effectiveness of the dissolution and the leaching of base metals, especially Zn and Cu, from the sulfide orebodies. As a result, downward moving surface water, charged with oxygen, percolates through a thicker rock column in the vadose zone between the surface and the water table. This setting provided more frequent replenishment of oxygen to the primary sulfide body, resulting in both a larger degree and a greater thickness of supergene alteration [106,107]. These favorable conditions would have promoted deep weathering of the sulfide orebodies and the production of highly acidic fluids, owing to widespread occurrence of pyrite and organic matter within the Jurassic carbonate host rocks.

The influx of the resulting acidic fluids into the carbonate host rocks triggered deep weathering and karst formation, allowing supergene mineralization to extend down dip in the carbonate rocks for at least several hundred meters and vertically for a similar distance in steep structural zones. Equally important, the highly acidic chemistry of the supergene fluids allowed these fluids to progressively leach and transport Zn and Cu and, to a lesser extent, Pb owing to its low mobility. Higher-grade mineralization could have been generated by a supergene “zone refining” process, whereby Zn, Cu, and, to a lesser extent, Pb are progressively dissolved and then reprecipitated within secondary trap sites. Galena seems to be more resistant to the oxidizing groundwater than sphalerite and chalcopyrite and can thus occur in association with smithsonite, hydrozincite, and hemimorphite.

The experimental work of [108,109] showed that temperature, pH, activity of CO$_2$ and SiO$_2$ within the ambient groundwater, and concentrations of Zn and Pb in solution constitute the principal factors in determining the paragenesis of Zn, Pb, and Cu non-sulfides and their spatial distribution. Based on the abovementioned studies and building upon recently published literature (e.g., [6]), we propose a three-stage model for the genesis of the Moroccan non-sulfide Pb-Zn ± Cu deposits, all of which are coincident with the D$_3$ event of [42] as the primary MVT mineralization (protore) has been dated <15 Ma [5,26,28]. In this respect, it should be emphasized that large-scale, vertical, positive movements were able to produce uncapping and weathering of the primary hypogene sulfides by creating a gradient for the percolation of meteoric fluids. In Morocco, weathering episodes have been related to four main Atlasic geodynamic events referred to as D$_0$ (Late Cretaceous), D$_1$ (Late Eocene), D$_2$ (Early Miocene), and Late Miocene to Recent, consistent with the conclusions of [30,42,73,110–112].

4.4. Early Pre-Non-Sulfide Gossanous Stage

The earlier phase of weathering, which resulted in the development of earthy yellowish to reddish goethite- and hematite-rich gossanous rocks, represents the transitional stage that encompasses the time span between the waning stage related to the emplacement of MVT mineralization dated at <15 Ma [5,28] and the initial episode of exhumation and weathering of sulfides in concert with the latest Atlasic phase of tectonic uplift in the late Miocene (~10 Ma, D$_3$ of [42]). During this early stage, the occurrence of pyrite has been critical in producing sufficient circulating acid fluids and favoring the high metal concentration in the aqueous solutions ([14]). The presence of pyrite is therefore considered a prerequisite for the development of the two following supergene mineralizing stages, the red calamine and the white calamine ore stages. The oxidation of pyrite will prepare the ground for future non-sulfide mineralization by creating additional secondary porosity and providing the required metals (i.e, Zn, Cu, and Pb) for the ore. Although sphalerite,
chalcopyrite, and galena are very susceptible to oxidation [113,114], they produce relatively small quantities of acid sulfate-bearing solutions compared to iron sulfides [88].

During this initial stage, oxidation of pyrite constituted the driving force that led to the generation of strongly oxidized and acidic fluids, which ultimately promoted the leaching of most base metals, such as Zn and Cu, and alkalis from their host rocks with the catalytic participation of sulfide-oxidizing bacteria and archaea. The released metals will migrate away from the sulfide precursor, guided by the high permeability of the host rocks, the ENE- and E-W-trending steeply dipping dilational faults, and the lower water table and seasonal flushing. The light sulfur isotopic compositions shown by late-stage pyrite (δ34S = −16.8 to −16.2‰, average = −15.8 ± 1.8‰, n = 5), whose precipitation is interpreted to have occurred during the collapse of the MVT hydrothermal system and its shift to supergene conditions [28], strongly support the involvement of low-temperature bacteriogenically mediated processes in the oxidation of pyrite. At this stage, the temperature of the shallow subsurface acidic fluids was probably higher than ambient temperatures, as indicated by the crystallographic constraints of the paragenetically related goethite, which indicate temperatures up to 50 °C [115].

Except for the radiochronological ages provided by [32], who used paleomagnetism to timely constrain the supergene oxidation episodes in the High Atlas, no other direct radiometric ages on the non-sulfide mineral phases from the Touissit-Bou Beker and Jbel Bou Dahar districts are yet available. By regional geological correlations [42] and comparison to equivalent dated supergene ore deposit types in the nearby Mediterranean basin [97], the first episode of oxidation described in this study could have occurred synchronously with the latest phase of tectonic uplift that affected the High Atlas system at ~10 Ma (Tortonian–Messinian time [42]).

4.5. Main Stage of Sulfide Oxidation: The “Red Calamine” Ore Stage

During this stage, which accounts for significant amounts of the extracted non-sulfide ore in the Touissit-Bou Beker and Jbel Bou Dahar districts, large-scale supergene zone-refining reactions led to a progressive replacement of all the preexistent sulfides, including sphalerite, chalcopyrite, and, to a lesser extent, galena, as Zn\(^{2+}\) and Cu\(^{2+}\) are more mobile than Pb\(^{2+}\) in the supergene environment [14,116]. Transport of Zn and Cu out of the sulfide orebody was facilitated by a falling water table caused by the uplift of the Atlassian system and probably by a change from wet tropical to monsoonal or semiarid climatic conditions.

The main chemical change during this stage resides in the complete release of sulfur from the pre-existent sulfides and the subsequent formation of Pb-Zn ± Cu carbonates and sulfates as a result of the neutralizing effect of the Jurassic host carbonate rocks. The resulting mineral paragenesis occurs mainly as vein and fracture infills, and also as in situ replacement of primary sulfide ore minerals. The development of this low-temperature paragenesis implies the complete oxidation of sulfide minerals, including oxidation of pyrite to oxyhydroxides, sphalerite to smithsonite-1, galena to anglesite-1 and cerussite-1, and chalcopyrite to malachite and azurite. The oxidation involves mainly a series of reactions similar to those described by [19,62,117–119]:

$$\text{PbS} + 2\text{O}_2 \rightarrow \text{PbSO}_4 \text{ under relatively low pH and high sulfate contents}$$

$$\text{PbSO}_4 + \text{H}_2\text{CO}_3 \rightarrow \text{PbCO}_3 + \text{SO}_4^{2-} + 2\text{H}^+, \text{ which is dependent on the activities of } \text{H}^+,$$

$$\text{H}_2\text{CO}_3(aq), \text{ and } \text{SO}_4^{2-}$$

$$\text{PbS} + \text{H}_2\text{O} + \text{CO}_2 + 2\text{O}_2 \rightarrow \text{PbCO}_3 + \text{H}_2\text{SO}_4$$

Smithsonite was generally produced during early oxidation of sphalerite, when Zn-bearing solutions reacted with the host carbonates or with carbonate-bearing fluids [17,108,120] at pH values between 6.2 and 8.1 and under high partial pressures of CO\(_2\) (P\(_{\text{CO}_2(g)}\)). Oxidation of galena occurred either as direct replacement of galena by anglesite, or as subsequent replacement of galena and anglesite by cerussite. The Pb-carbonate,
which is widespread throughout most orebodies of the Touissit-Bou Beker district as void infills and replacement of the host dolostones (i.e., “cerusite pierreuse”), precipitated when solutions carrying carbonate reacted with galena, whereas anglesite formed in the presence of carbonate-poor fluids, as documented in other Moroccan supergene Pb-Zn ± Cu deposits [75]. Altogether, these observations indicate that during the main stage of sulfide oxidation, metals were quickly neutralized by the carbonate host rocks and did not migrate over long distances. The stronger evidence for short-distance migration of the metal-bearing fluids is provided by the occurrence of gypsum, whose precipitation in supergene environments is triggered by limited fluid migration [17].

Remarkably, the consistent increase of δ\(^{13}\)C and δ\(^{18}\)O values (Figure 9) along with the temporal shift from heavy (i.e., Gyp-1) to light sulfur (i.e., Gyp-2) sources with advancing paragenetic sequence could be explained in terms of microbial oxidation under decreasing \(\mathrm{H}_2\mathrm{S}/\mathrm{O}_2\) ratios and temperature conditions. This isotopic evolution could therefore reflect significant microbiologically sourced carbon input at relatively higher temperatures of formation during the paragenetically early sulfide oxidation stage as chemolithotrophs exothermically digested sulfides. Over time, as sulfide nutrients became scarcer in the oxidation zone, secondary carbonate minerals recorded progressively heavier δ\(^{13}\)C and lighter δ\(^{34}\)S values at decreasing temperatures. This reflects the diminishing role of both microbes in the oxidation process, and the associated heat production from their digestion of sulfides, similar to processes documented in Australia [65]. Such evolution is also reflected in deposit mineralogy, as the resulting drop in CO\(_2\) partial pressures inhibited azurite formation at the expense of malachite, and as pseudomorph formation of malachite after azurite. In the final stages of oxidation, temperatures are near ambient, and δ\(^{13}\)C values are dominated by dissolution of local carbonate rocks and atmospheric sources. In support of this conclusion, oxygen isotope thermometry from the coexisting azurite, malachite and late-stage calcite indicates decreasing temperatures from 34 °C to 19 °C, assuming a local meteoric water δ\(^{18}\)O value of −7.2‰.

Overall, the calculated temperatures are similar to those determined for smithsonite, cerussite and azurite-malachite in other base metal supergene deposits worldwide (11–35 °C, avg 20 °C; [13,59,68,72]) and are in agreement with a continental environment in a temperate climate [66,80]. Although climatic variations and complex weathering histories could have produced the higher calculated temperatures that prevailed during the ‘red calamine” stage, we suggest that the calculated higher temperatures of 32 ± 3°C could reflect the heat produced by the exothermic oxidation of sulfides. If so, this indicates that oxidation of primary MVT ores could produce shallow subsurface temperatures up to 30 °C higher than local mean air temperatures [62].

Given the predominance of smithsonite over hemimorphite and the presence of silica-rich mineral species such as willemite, we assume that the supergene weathering occurred under warm-dry climate conditions and low precipitation rates. In this regard, it is well documented that under humid-tropical climate, silica can be more efficiently leached from silicate rocks than Al or Fe [121,122]. Moreover, the negative δ\(^{13}\)C values, up to −15‰ (Table 1), shown by smithsonite, suggest the involvement of a light component of carbon dominated by surface vegetation and/or secondary carbonate dissolution caused by a rising water table.

Thermochronological data [42–44,94–99] along with structural constraints indicate the involvement of a major pulse of exhumation (i.e., D\(_3\) of [42]) that occurred during the last ~10 Ma, from middle/late Miocene to present, at an approximative shortening rate of 2.6 mm/year [44]. Additionally, geological correlations and comparison to similar supergene ore deposit types in the Mediterranean basin (e.g., [93]) and North Africa (e.g., [123]) indicate that the emplacement of the “red ore stage” could have occurred during the Pliocene-Pleistocene (8–7 Ma, [93]), coincident with an episode of sea-level rise in the study area at about 8 Ma [28,124]. Regionally, the authors of [123] show the occurrence of a main weathering event in Northern Tunisia during late Tortonian (8.6 ± 0.9
Ma, (U-Th)/He on supergene goethite) and a later episode during late Pleistocene (0.8 ± 0.2 Ma).

4.6. Late Stage of Sulfide Oxidation: The “White Calamine” Ore Stage

During this stage, the base metals released by the oxidation of sulfides were not trapped locally but transported by downward percolating meteoric fluids farther away from the sulfide protore, both vertically and laterally, and precipitated under more “oxidizing” conditions. The supergene fluids moved more than 150 m down dip and laterally through the Jurassic porous carbonate rocks and along the associated faulted corridors, where their acidity promoted further karstic dissolution. The dissolution was followed by partial to total replacement of the sulfide minerals by Pb-Zn ± Cu carbonates and sulfates and, to a lesser extent, silicates. The resulting supergene mineral assemblages consist of various proportions of cerussite-2, anglesite-2, smithsonite-2, malachite-azurite-2, and, more importantly, hydrozincite as the most diagnostic phase that characterizes this stage. Gangue minerals include gypsum-2, calcite, and quartz.

The higher δ¹³C and δ¹⁸O values of the Pb-Zn ± Cu-bearing carbonates and sulfates from this stage, which reflect isotopic interaction at lower ambient temperatures between meteoric fluids and carbonate wall rocks, indicate carbonate and sulfate precipitation under more oxidizing conditions as the partial pressure of CO₂(g) of the system reached equilibrium with the atmosphere and as pH increased from neutral to slightly alkaline values. The occurrence of hydrozincite constitutes further support that the supergene ore-bearing aqueous solutions were in equilibrium with the atmosphere (undersaturated zone) since hydrozincite is stable at pCO₂(g) levels occurring in near-surface settings and thus lower than those allowing smithsonite precipitation (e.g., [108]). Higher aeration would have provided higher O₂ concentration, allowing for a higher contribution of O₂ during oxidation from air and producing carbonates and sulfates with higher δ¹³C and δ¹⁸O values. In addition to an atmospheric carbon source, the formation of hydrozincite should have benefited from the incorporation of an isotopically light component of carbon as a result of dissolved CO₂ derived from the bacteria and archaea-mediated decay of organic matter (i.e., C₃-C₄ plants and soils) as discussed above. Paleoclimatic and/or seasonal variations, with lower δ¹⁸O values characterizing the winter season and higher δ¹⁸O values in summer, could also be invoked.

Given these relationships, hydrozincite precipitation must have occurred in relatively recent times, when the carbonate platform was already relatively uplifted. This observation suggests that the late stage of supergene oxidation occurred at higher altitudes, while the Atlas system continued to rise under the oblique convergence between Africa and Eurasia [125,126]. In this respect, structural and thermochronological data indicate relatively high exhumation rates of up to 0.3 km/Ma from 5 Ma to 0 Ma (e.g. [43,44]). If so, we document here a causal relationship between the fast exhumation rates and the timing of supergene mineralization in the Atlas system between 5 Ma and 0 Ma, consistent with Ar-Ar dates (2–1 Ma to present; [93]) in the Mediterranean area and (U-Th)/He age determinations in Tunisia [123]. It is noteworthy that the amount of shortening of the Central High Atlas, which hosts the most productive calamine deposits in North Africa, is more than double compared to that of the Western High Atlas (approximately 30–34 km; [38,41,127]) and thus records higher exhumation rates. This observation could explain why the most significant supergene deposits are concentrated within the Central High Atlas and not elsewhere.

5. Conclusions

The present study constitutes the first integrated contribution that combines field observations with thorough mineralogical and geochemical investigations to constrain the origin(s) and evolution of the supergene fluid(s) and depositional processes that led to the precipitation of the non-sulfide mineralization in the Jurassic carbonate-hosted Touissit-Bou Beker and Jbel Bou Dahar Mississippi Valley-type districts. Economic supergene
mineralization, which occurs as open-space fillings in intergranular voids, karsts, fractures, and breccias and as replacements of the host carbonates, consists of oxidized minerals resulting from weathering and alteration of hypogene sulfide mineralization. The secondary enrichment of the mineralized orebodies is assigned to both direct replacement and wall-rock replacement types. The resulting low-pressure, low-temperature non-sulfide ore assemblages consist predominantly of various proportions of smithsonite, hydrozincite, cerussite, anglesite, malachite, and azurite with subordinate amounts of willemite, pyromorphite, and sauconite. The main gangue minerals comprise calcite, aragonite, gypsum, and quartz. Three stages of supergene mineralization are recognized, the earliest of which constitutes a prerequisite for the development of the following two mineralized supergene stages. The first stage involves the oxidation of pyrite and subsequent development of gossanous manganese and iron hydroxide-rich blankets. The second stage of supergene mineralization, referred to as the main oxidation red calamine stage, accounts for significant amounts of the extracted non-sulfide ore and corresponds to the complete release of sulfur from the pre-existent sulfides and the subsequent formation of Pb-Zn ± Cu carbonates and sulfates as a result of the neutralizing effect of the Jurassic host carbonates. The third post-oxidation stage, referred to as the late oxidation white calamine stage, is typified by the ultimate precipitation of hydrozincite.

Combined carbon, oxygen, and sulfur isotope compositions of ores are consistent with precipitation of the zinc and lead ± copper non-sulfide mineralization in a supergene environment from in situ oxidation and replacement of the primary sulfide phases. The oxidation and replacement were due to meteoric fluids circulating in the carbonate rocks at ambient temperatures, which decreased with time from 32 to 18 °C. More importantly, the layered stromatolitic-like textures, along with the light carbon and sulfur isotope compositions, reflect the involvement of a bacteriogenically mediated origin for the carbon and sulfur under temperate to warm climatic conditions. The released acidic fluids were channeled by the Jurassic permeable carbonates and by steep fracture and fault zones. The transport of dissolved metals out of the precursor sulfide bodies was facilitated by a falling water table caused by the tectonic uplift related to the Atlasic orogeny and a change from wet tropical to monsoonal or semiarid climatic conditions. The close spatial relationships between sulfide and non-sulfide ore zones and coexisting sulfide-non-sulfide mineral assemblages and geological settings along with stable isotope data suggest that the deposition of the supergene non-sulfide mineralization of the Moroccan Atlasic system may have occurred during the latest stages of uplift at ~10 Ma, during intervals when the meteoric fluids were supplied by a relatively wet climate between the Late Miocene and Holocene. Periods of late Tortonian and late Pleistocene, recognized as weathering events in adjacent areas, are proposed as time spans for stages II and III of supergene mineralization. The process of oxidation may be ongoing today. Based on these findings, we recommend that for future regional to local-scale exploration campaigns for supergene lead-zinc ± copper deposits, explorationists should combine all aspects of climatic evolution, long-wavelength neotectonics movements, and hydrogeology for a successful operation.

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