Germanium, gallium and indium are in high demand due to their growing usage in high-tech and green-tech applications. However, the mineralogy and the mechanisms of concentration of these critical elements in different types of hydrothermal ore deposits remain poorly constrained. We investigated the mineralogical distribution of Ge, Ga and In at the Mt Carlton high-sulfidation epithermal deposit in NE Australia, using electron probe microanalysis and laser ablation inductively-coupled plasma mass spectrometry. Parageneses from which selected minerals were analyzed include: Stage 1 acid sulfate alteration (alunite), Stage 2A high-sulfidation enargite mineralization (enargite, argyrodite, sphalerite, pyrite, barite), Stage 2B intermediate-sulfidation sphalerite mineralization (sphalerite, pyrite, galena) and Stage 3 hydrothermal void fill (dickite). Moderate to locally high concentrations of Ga were measured in Stage 1 alunite (up to 339 ppm) and in Stage 3 dickite (up to 150 ppm). The Stage 2A ores show enrichment in Ge, which is primarily associated with argyrodite (up to 6.95 wt % Ge) and Ge-bearing enargite (up to 2189 ppm Ge). Co-existing sphalerite has comparatively low Ge content (up to 143 ppm), while Ga (up to 1181 ppm) and In (up to 571 ppm) are higher. Sphalerite in Stage 2B contains up to 611 ppm Ge, 2829 ppm Ga and 2169 ppm In, and locally exhibits fine colloform bands of an uncharacterized Zn-In mineral with compositions close to CuZn$_2$(In,Ga)S$_4$. Barite, pyrite and galena which occur in association with Stage 2 mineralization were found to play negligible roles as carriers of Ge, Ga and In at Mt Carlton. Analyzed reference samples of enargite from seven similar deposits worldwide have average Ge concentrations ranging from 12 to 717 ppm (maximum 2679 ppm). The deposits from which samples showed high enrichment in critical elements in this study are all hosted in stratigraphic sequences that locally contain carbonaceous sedimentary rocks. In addition to magmatic-hydrothermal processes, such rocks could potentially be important for the concentration of critical elements in high-sulfidation epithermal deposits.

Keywords: Mt Carlton; high-sulfidation; epithermal; germanium; gallium; indium; critical elements; LA-ICP-MS; enargite; sphalerite
optics (Ge) and solar cells (Ge, Ga, In) [1–5]. Due to their increasing economic importance combined with current supply security risks, Ge, Ga and In are commonly classified within the group of critical raw materials [6]. These elements can be concentrated in a variety of hydrothermal ore deposits, including volcanic hosted massive sulfide ( VHMS) deposits, Mississippi Valley-type (MVT) deposits, carbonate-hosted polymetallic deposits of Tsumeb-type (exemplified by Tsumeb in Namibia, Kipushi in D.R. Congo and the Apex Mine in Utah; e.g., [7]), syngenetic or early diagenetic sedimentary exhalative (SEDEX) deposits, epithermal deposits, polymetallic (±Sn) vein or stockwork deposits linked to granites, and skarn deposits [1,7,8]. Although Ge, Ga and In may form their own independent minerals, they are also incorporated as trace components into various minerals [1,7,9,10]. There are no hydrothermal ore deposits that are mined primarily for these elements. Instead, recovery of Ge, Ga and In occur as by-products from operations for other commodities, chiefly from the processing of sphalerite ores [1,2,11–14]. Enrichment in Ge and Ga in sphalerite is generally favored in low-temperature deposits that are not necessarily related to magmatic fluid (e.g., MVT deposits; [2,10,13,14]). Indium on the other hand is strongly linked to relatively high-temperature magmatic-hydrothermal systems, often related to felsic intrusions [10,15,16].

High-sulfidation epithermal (HS) Au ± Ag-Cu deposits occur in calc-alkaline island and continental arcs related to subduction zones [17,18]. These deposits are believed to form in a magmatic-hydrothermal environment equivalent to that beneath modern volcanic fumaroles and crater lakes [19]. Mineralization styles include hydrothermal breccias, veins, stockworks, and disseminations or replacements [20]. The high-grade ore typically occurs within cores of residual silica alteration, which reflect the leaching effects of highly acidic magmatic vapor condensates. The silicic cores are enveloped by a laterally extensive zone of advanced argillic alteration, and together they form a subhorizontal blanket, or lithocap, above the mineralizing intrusion [21–24]. A distinguishing feature of HS deposits is a high-sulfidation state ore mineral assemblage dominated by enargite and pyrite [20,25,26]. Most HS deposits also contain a later-stage ore mineral assemblage that is Au-rich and comprises intermediate-sulfidation state minerals (e.g., chalcopyrite, pyrite, tennantite-tetrahedrite, sphalerite, galena, electrum, and tellurides) [20,26,27]. High-sulfidation epithermal deposits and their associated lithocaps have a spatial and genetic link to porphyry Cu deposits (occurring in and around magmatic apophyses from the causative intrusion), intermediate-sulfidation epithermal (IS) deposits (commonly occurring on the flanks of the lithocap) and skarn deposits (occurring in carbonate-bearing wall rocks, if present) [23,24,28,29]. Mineralization in IS deposits shows close similarities to that in HS deposits except that the enargite-bearing assemblage is lacking, the Ag: Au ratio is typically higher, and advanced argillic assemblages form a smaller proportion of the wall rock alteration facies [26]. Despite major advances in the understanding of porphyry-epithermal mineral systems in recent decades, relatively little work has been done on the presence and distribution of critical elements such as Ge, Ga and In in these deposits.

The northern Bowen Basin is a metallogenic province related to Permocarboniferous magmatism in NE Queensland, Australia. Here, a variety of porphyry and epithermal mineral occurrences are hosted within an Early Permian backarc rift sequence (Figure 1; [30]). The most significant deposit in the northern Bowen Basin, and the only one currently in production, is the Mt Carlton HS deposit. Mt Carlton was discovered in 2006, and as of December 2016 it was estimated to contain a resource of 10.42 Mt averaging 2.92 g/t Au, 30.60 g/t Ag and 0.47% Cu [30]. An active open pit operation includes the Au-rich V2 pit in the NE, and the Ag-rich A39 pit in the SW (Figure 2). Mt Carlton represents one of the world’s few economic pre-Neogene HS deposits, and it formed during a stage of backarc rifting as the laterally extensive Bowen Basin was opening in the Early Permian [30].

In this study, we use electron probe microanalysis (EPMA) and laser ablation inductively-coupled plasma mass spectrometry (LA-ICP-MS) to determine the concentrations and distribution of Ge, Ga and In and related elements in key minerals formed throughout the paragenetic sequence at Mt Carlton. Samples from a selection of global HS and porphyry deposits are analyzed for reference, and the results are compared with available case studies from the literature. The main aim of the study is to improve...
the understanding of the mineralogical distribution of Ge, Ga and In in HS deposits. This information is a pre-requisite for the evaluation of HS deposits as potential sources for these elements, and for the development of efficient ore processing methods during recovery.

Figure 1. (A) Location of the northern Bowen Basin (red rectangle) in NE Queensland, Australia. (B) Geologic map of the northern Bowen Basin (center of map) and the corresponding Connors Arc (right-hand side of map), showing the location of the known porphyry, epithermal and mesothermal mineral occurrences. Grid—UTM zone 55; datum—GDA94. From Sahlström et al. [30].

2. Geologic Setting of the Mt Carlton Deposit

2.1. Regional Geology

The Bowen Basin is a NW-trending elongate sedimentary basin located in eastern Queensland, Australia. It formed as part of the New England Orogen during the Early Permian to Middle Triassic, and shows evidence for a complex and polyphase tectonic evolution ([31], and references therein). Backarc rifting in the Bowen Basin was initiated in the Early Permian, and resulted in a series of isolated graben and half graben basins infilled with volcanic and sedimentary rocks [32–34]. The Lizzie Creek Volcanic Group formed during this stage, and is the principal host of mineralization in the northern Bowen Basin. Thermal relaxation took over as the dominant mechanism of subsidence in the Middle Permian. This led to widespread transgression and deposition of marine, coastal plain and carbonate sedimentary units (e.g., the Back Creek Group and Tiverton Formation; [33,35–37]). The thermal relaxation stage was abruptly terminated in the Late Permian, due to the progression of a series of inland-stepping orogenic fronts during the ~265–235 Ma Hunter-Bowen Orogeny ([31], and references therein). This caused foreland loading, tectonic inversion and the development of a foreland basin. The foreland basin stage involved voluminous deposition of terrestrial sediments (e.g., the Blackwater Group) and coal measures (e.g., the Moranbah-German Creek CM and Rangal CM) until the Bowen Basin closed in the Middle Triassic [33,38–40].
2.2. Stratigraphy and Deformation Sequence at Mt Carlton

The basement in the northern Bowen Basin consists of fine- to medium-grained I-type granitoids belonging to the Urannah Batholith. At Mt Carlton, the granite basement (Unit 1, 297–294 Ma; [41]) is overlain by a volcano-sedimentary sequence belonging to the Early Permian Lizzie Creek Volcanic Group (288–275 Ma; [41]). The Lizzie Creek sequence consists of (from bottom to top): fine-grained porphyritic andesite (Unit 2), massive and locally flowbanded porphyritic rhyodacite (Unit 3), well-bedded fragmental rhyodacite lapilli tuffs with interbedded carbonaceous lacustrine sediments (Unit 4A), massively bedded dacite tuffs (Unit 4B), fragmental to agglomeritic dacite (Unit 5), fragmental to agglomeritic andesite and minor andesite lava (Unit 6), water-laid sediments with local coal seams (Unit 7), and a flowbanded rhyolite unit (Unit 8) forming the top of the sequence [30]. Mineralization at Mt Carlton is hosted partly in the massive rhyodacite porphyry (Unit 3) and partly in the overlying volcano-lacustrine sediments (Unit 4A; Figure 2).

The Mt Carlton deposit has experienced five stages of extensional deformation [30]. Hydrothermal alteration and subsequent epithermal mineralization occurred during a stage of rifting with associated high-angle normal faulting in response to both E-W and N-S extension (D1). Mineralization was controlled by steep normal faults and fracture networks related to the rifting, and occurred partly contemporaneously with volcanism and deposition of volcanic sediments into local graben and half graben basins. Continued E-W extension after mineralization provided rapid burial of the Mt Carlton deposit beneath a volcano-sedimentary cover. It also caused the development of a series of low-angle (and locally layer-parallel) detachment faults which have rotated and displaced the stratigraphic sequence and the ore zones at Mt Carlton (D2). High-angle normal faulting in response to N-S extension and emplacement of E-W trending basaltic dykes into the normal faults (D3) was followed by dominantly dextral strike-slip faulting along the dyke margins (D4). A final stage involving emplacement of WNW-trending basaltic dykes (D5) completes the deformation sequence at Mt Carlton.

Figure 2. NE-SW section along the Western ore zone at Mt Carlton. The section shows the distribution of stratigraphic units, major D2 structures, alteration assemblages and high-grade mineralization. Abbreviations: alu—alunite; anh—anhydrite; APS—aluminium-phosphate-sulphate minerals; brt—barite; chl—chlorite; dck—dickite; gyp—gypsum; hbx—hydrothermal breccia; hem—heamatite; ill—illite; kln—kaolinite; mnt—montmorillonite; prl—pyrophyllite; py—pyrite; qtz—quartz; rt—rutile. Modified from Sahlström et al. [30].
3. Mineral Paragenesis

3.1. Hydrothermal Alteration

The volcano-sedimentary host rocks at Mt Carlton are extensively hydrothermally altered. Stage 1A of the paragenetic sequence (Figure 3) comprises a suite of magmatic-hydrothermal alteration assemblages that are zoned outward, away from fluid conduits as follows: silicic (massive + vuggy + hydrothermal breccia) → quartz-alunite (Figure 4A) → quartz-dickite-kaolinite. This alteration halo envelops mineralization in Units 3 and 4A (Figure 2). The silicic cores locally contain veins and void fill made up of magmatic-steam alunite (Stage 1B; Figures 3 and 4B). Stage 1A alteration in the granite basement comprises chlorite and illite occurring in veinlets and as replacements of pre-existing phenocrysts of feldspar, biotite and hornblende.

![Figure 3. Paragenetic sequence of minerals at Mt Carlton. The minerals investigated in this study are highlighted in green. Modified from Sahlström et al. [30].](image)

3.2. Hydrothermal Mineralization and Void Fill

Proximal Au-Cu mineralization in the V2 pit occurs in networks of steep veins, fractures and hydrothermal breccia bodies in the rhyodacite porphyry (Unit 3). Three major ore zones (Eastern, Western and Link) are aligned in an en echelon fashion along an E-W trending corridor. The Eastern and Link ore zones terminate in the V2 pit, whereas the Western ore zone continues along ~600 m
strike length to the SW into the A39 pit. From proximal and deep to distal and shallow (at the time of formation), the Western ore zone shows a metal zonation of: Au-Cu → Cu-Zn-Pb-Ag → Ag-Pb-(Cu) → Ag (Figure 2). Distal Ag mineralization in the A39 pit occurs in the overlying volcano-lacustrine sediments (Unit 4A), in a stratiform position parallel to primary sedimentary layering [30].

Epithermal mineralization at Mt Carlton developed in three stages (Stages 2A–C; Figure 3). The initial Stage 2A mineralization is the most voluminous event, and comprises a high-sulfidation state mineral assemblage dominated by enargite. The assemblage also contains variable amounts of pyrite, luzonite, fahlores of tennantite-tetrahedrite-goldfieldite compositions, electrum, chalcopyrite, bornite, sphalerite, galena, argyrodite, pearceite-polybasite group minerals, an uncharacterized analogue to aguilarite and cervelleite [ideally Ag₄TeSe], and barite (Figure 3, Figure 4B–F and Figure 5C,D). Stage 2A mineralization is associated with silicic alteration (quartz ± alunite) of the wall rocks (Figure 4C). Whole rock analyses of Stage 2A ore samples from the V2 pit showed average concentrations of 61 ppm Ge (up to 246 ppm), 34 ppm Ga (up to 66 ppm) and 56 ppm In (up to 151 ppm In), while samples from the A39 pit yielded average concentrations of 109 ppm Ge (up to 342 ppm), 24 ppm Ga (up to 45 ppm) and 11 ppm In (up to 19 ppm) [30].

Stage 2A mineralization is crosscut by a younger intermediate-sulfidation state mineral assemblage dominated by sphalerite (Stage 2B). The Stage 2B assemblage also contains variable amounts of galena, pyrite, electrum, fahlores of tennantite-tetrahedrite-goldfieldite compositions, chalcopyrite, bornite and barite (Figures 3, 4D and 5E). Sahlström et al. [42] recently documented the presence of a mineral with compositions close to CuZn₂(In,Ga)S₄, which occurs as fine (~1 µm thick) colloform bands inside the Stage 2B sphalerite (Figure 5F, Table 1). This mineral corresponds to a Ga-rich variety of the uncharacterized Zn-In mineral [ideally CuZn₂InS₄] first described from the Toyoha deposit in Japan [43], and later from the Mt Pleasant deposit in Canada [44]. Stage 2B ore samples from the V2 pit showed average whole rock concentrations of 167 ppm Ge (up to 265 ppm), 284 ppm Ga (up to 396 ppm) and 255 ppm In (up to 408 ppm), while average concentrations of 7 ppm Ge (up to 13 ppm), 108 ppm Ga (up to 151 ppm) and 201 ppm In (up to 468 ppm) were observed for samples from the A39 pit [30].

### Table 1. List of critical element-bearing minerals identified at the Mt Carlton deposit.

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Class</th>
<th>Crystal System</th>
<th>Ideal Formula</th>
<th>Paragenesis</th>
<th>Contained Elements</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alunite</td>
<td>Sulfate</td>
<td>Trigonal</td>
<td>KAl₃( SO₄ )₂(OH)₈</td>
<td>1A and 1B</td>
<td>Ga</td>
<td>this study</td>
</tr>
<tr>
<td>Argyrodite</td>
<td>Sulfide</td>
<td>Orthorhombic</td>
<td>Ag₄GeS₆</td>
<td>2A</td>
<td>Ge</td>
<td>this study</td>
</tr>
<tr>
<td>Enargite</td>
<td>Sulfosalts</td>
<td>Orthorhombic</td>
<td>Cu₃AsS₄</td>
<td>2A</td>
<td>In, Ga</td>
<td>this study</td>
</tr>
<tr>
<td>Sphalerite</td>
<td>Sulfide</td>
<td>Isometric</td>
<td>ZnS</td>
<td>2A and 2B</td>
<td>In, Ga, Ge</td>
<td>this study</td>
</tr>
<tr>
<td>Zn-In mineral</td>
<td>Sulfide</td>
<td>Uncharacterized</td>
<td>CuZn₂InS₄</td>
<td>2B</td>
<td>In, Ga</td>
<td>[42]</td>
</tr>
<tr>
<td>Dickite</td>
<td>Silicate</td>
<td>Monoclinic</td>
<td>Al₂Si₂O₅(OH)₄</td>
<td>3</td>
<td>Ga</td>
<td>this study</td>
</tr>
</tbody>
</table>

Stage 2C is a minor event that overprinted Stage 2B mineralization, and it has only been observed at the microscopic scale. The mineral assemblage is of intermediate-sulfidation state and is dominated by tennantite. The tennantite forms anhedral masses that contain variable amounts of luzonite, chalcopyrite, galena, electrum and aggregates of intergrown hessite and petzite (Figure 3). A late stage of dickite and pyrite (Stage 3; Figures 3 and 4F) is prevalent at Mt Carlton and occurs in veins and as massive void fill overprinting Stage 2 mineralization.

3.3. **Syn-Tectonic Alteration and Supergene Oxidation**

Pervasive illite-montmorillonite ± red hematite alteration with local syn-tectonic gypsum veins developed regionally during the D₂ deformation event (Stage 4; Figure 3). In the area around the open pits, this alteration affects Units 2, 5 and 6, and locally Unit 4 (Figure 2). The final paragenetic stage at Mt Carlton involves minor degrees of supergene oxidation that affects the upper ~50 m of the deposit. This has caused local development of secondary covellite, chalcocite and malachite on Cu-bearing minerals such as enargite, luzonite and bornite, particularly along fractures within the minerals (Stage 5; Figures 3 and 5E).
Figure 4. Photographs of relevant alteration and mineralization assemblages from Mt Carlton. (A) Quartz-alunite-pyrite alteration (Stage 1A) developed in a quartz-feldspar phyric rhyodacite. Magmatic-hydrothermal alunite occurs as disseminations and as replacements of pre-existing feldspars. (B) Plumose alunite vein of magmatic-steam origin (Stage 1B). After deposition of alunite, the vein was re-opened along the central line to form an enargite vein (Stage 2A). The composite vein was then overprinted by dickite (Stage 3). (C) Silicic hydrothermal breccia cemented by Stage 2A enargite-pyrite ore. (D) Silicic altered rhyodacite crosscut by a fine enargite-cemented hydrothermal breccia (Stage 2A). The enargite breccia is crosscut by a sphalerite-pyrite-galena vein containing specules of electrum (Stage 2B). The sphalerite vein is in turn locally overprinted by tennantite (Stage 2C). (E) Stage 2A enargite veinlet with a halo of barite. (F) Disseminated Stage 2A enargite mineralization crosscut by a Stage 3 dickite-pyrite vein. Mineral abbreviations: Alu—alunite; Brt—barite; Dck—dickite; Elc—electrum; Eng—enargite; Gn—galena; Py—pyrite; Qtz—quartz; Sp—sphalerite; Tn—tennantite.
Figure 5. Associations of minerals investigated in this study. (A) Magmatic-hydrothermal alunite (Stage 1A) replacing a pre-existing feldspar crystal in a quartz-feldspar phryic rhyodacite (transmitted light, cross polarized). (B) Plumose texture in a magmatic-steam alunite vein (Stage 1B). Alunite is locally overprinted by Stage 3 dickite (transmitted light, cross polarized). (C) Argyrodite intergrown with barite and enargite in Stage 2A high-sulfidation ore (BSE image). (D) Stage 2A high-sulfidation assemblage comprising enargite, luzonite, sphalerite and galena (reflected light, plane polarized). (E) Stage 2B assemblage comprising sphalerite, pyrite, galena, bornite and barite. The galena contains abundant segregations of tennantite. Stage 5 covellite has locally overprinted the assemblage (reflected light, plane polarized). (F) Colloform texture in Stage 2B sphalerite. The fine BSE-bright band corresponds to the Zn-In mineral [CuZn$_2$(In,Ga)S$_4$] (BSE image). Mineral abbreviations: Alu—alunite; Arg—argyrodite; Bn—bornite; Brt—barite; Cv—covellite; Dck—dickite; Eng—enargite; Gn—galena; Luz—luzonite; Qtz—quartz; Sp—sphalerite; Tn—tennantite.
4. Methods

4.1. Sampling

The samples used in this study were collected from drill cores and in situ from outcrops in the open pits at Mt Carlton, and subsequently prepared as polished thin and thick sections. The present study focuses on key minerals formed throughout the paragenetic sequence at Mt Carlton, including minerals formed during Stage 1 acid sulfate alteration (alunite), Stage 2A high-sulfidation mineralization (enargite, argyrodite, sphalerite, pyrite and barite), Stage 2B intermediate-sulfidation mineralization (sphalerite, pyrite and galena) and Stage 3 hydrothermal void fill (dickite) (Figure 3 and Table 1). Analytical measurements were generally conducted on multiple samples to obtain a range of representative results, and when possible, samples from both the V2 pit (proximal) and the A39 pit (distal) were included.

Despite enargite being the principal ore mineral in HS deposits there is a lack of published trace element data for this mineral, in particular with regards to its content of Ge, Ga and In. Therefore we have also included reference samples of enargite from seven representative HS and porphyry deposits around the world. These samples were collected during previous field visits to the deposits, and include Lepanto (Philippines), Laurani (Bolivia), La Mejicana (Argentina), Motomboto (Indonesia), Chelopech (Bulgaria), Cerro Quema (Panama) and El Salvador (Chile) (Table 2).

4.2. WDS Spot Analyses

The major element composition of argyrodite was determined by wavelength dispersive spectroscopy (WDS) using a Jeol JXA8200 “Superprobe” (Jeol, Tokyo, Japan) housed at the Advanced Analytical Centre (AAC) at James Cook University. Analyses were done utilizing a focused beam (<1 μm) with an accelerating voltage of 20 kV and a beam current of 20 nA. Analytical conditions were [analyzed element (standard, line, crystal)]: Cu (chalcopyrite, Kα, LIF), S (chalcopyrite, Kα, PET), Pb (galena, Mα, PET), Ag (argentite, Lα, PET), Bi (bismuthinite, Mα, PET), Se (bismuthinite, Lα, TAP), As (gallium arsenide, Lα, TAP), Sb (metal Sb, Lα, PET), Te (metal Te, Lα, PET), Au (metal Au, Mα, PET) and Ge (metal Ge, Kα, LIF).

4.3. WDS Element Mapping

WDS element mapping of enargite was performed on the same electron microprobe, utilizing a 2 μm electron beam with an accelerating voltage of 20 kV and a beam current of 200 nA. Mapping was done with a step size of 2 μm and a 100 ms dwell time. GeLα was measured on a TAP crystal, ZnKα on a LIFH crystal, FeKα on a LIF crystal, SnLα on a PETJ crystal and SbLα on a PETJ crystal. Element maps were processed offline using the ImageJ software.

4.4. LA-ICP-MS Spot Analyses

Trace element analyses of alunite, enargite, pyrite, galena, sphalerite, barite and dickite were done in separate runs using a Varian 820 ICP-MS instrument (Varian, Melbourne, Australia) interfaced to a GeoLas 200M 193 nm ArF excimer laser ablation system (Coherent, Göttingen, Germany), housed at the AAC. Ablation was conducted in a custom built large volume cell using high-purity He as the carrier gas, which was subsequently mixed with Ar prior to introduction into the ICP-MS. For spot analyses, the laser fluence was generally 5–6 J/cm² at the sample site with the laser repetition rate set to 10 Hz and the beam diameter set to between 24 and 60 μm.
### Table 2. List of deposits from which reference samples of enargite were analyzed.

<table>
<thead>
<tr>
<th>Deposit</th>
<th>Type</th>
<th>Age</th>
<th>Tectonic Setting</th>
<th>Stratigraphic Column (excluding post-mineralization units)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lepanto (Philippines)</td>
<td>HS</td>
<td>Quarternary</td>
<td>Island arc</td>
<td>Andesite-dacite pyroclastic rocks and porphyry intrusions (Imbanguila Dacite)</td>
<td>[29]</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Andesite volcanioclastic rocks (Balili Volcaniclastics), intruded by a quartz-diorite porphyry complex</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Basement of metamorphosed submarine basalt-andesite volcanic rocks (Lepanto metavolcanic rocks)</td>
<td></td>
</tr>
<tr>
<td>Laurani (Bolivia)</td>
<td>HS</td>
<td>Neogene</td>
<td>Continental arc (foreland basin)</td>
<td>Andesite-rhyolite lava and pyroclastic rocks (Laurani volcanic complex)</td>
<td>[45]</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Red bed conglomerate, sandstone and siltstone (Chuquichambi Formation)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Basement comprising shale, siltstone and sandstone (Llalagua, Uncia and Catavi Formations)</td>
<td></td>
</tr>
<tr>
<td>La Mejicana (Argentina)</td>
<td>HS</td>
<td>Neogene</td>
<td>Continental arc</td>
<td>Dacite-rhyodacite porphyritic rocks (Mogote Formation)</td>
<td>[46]</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Continental sandstone and conglomerate (Agua Colorada and Patquía Formations)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Basement of metamorphosed shale, sandstone, siltstone (Negro Peinado Formation), intruded by Nuñarco granites</td>
<td></td>
</tr>
<tr>
<td>Motomboto (Indonesia)</td>
<td>HS</td>
<td>Quarternary</td>
<td>Island arc</td>
<td>Subaerial andesite-rhyolite lava and pyroclastic rocks (Motomboto Volcanics)</td>
<td>[47]</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Submarine basalt-andesite lava (Bilungala Volcanics)</td>
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</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Unknown basement</td>
<td></td>
</tr>
<tr>
<td>Chelopech (Bulgaria)</td>
<td>HS</td>
<td>Cretaceous</td>
<td>Island arc</td>
<td>Argillaceous limestone, calcarenite and sandstone interlayered with volcanic rocks (Chelopech Formation)</td>
<td>[48]</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Coal-bearing conglomerate and sandstone, intruded by subvolcanic andesite intrusions</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Metamorphic basement comprising migmatitic gneiss, amphibolite, phyllite</td>
<td></td>
</tr>
<tr>
<td>Cerro Quema (Panama)</td>
<td>HS</td>
<td>Paleogene</td>
<td>Island arc (forearc basin)</td>
<td>Andesite-dacite volcanic rocks, biomicritic hemipelagic limestone, turbidite, shale (Rio Quema Formation)</td>
<td>[49]</td>
</tr>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>Basement comprising basalt and pillow basalt (Azuero Igneous Basement)</td>
<td></td>
</tr>
<tr>
<td>El Salvador (Chile)</td>
<td>Porphyry</td>
<td>Paleogene</td>
<td>Continental arc</td>
<td>Rhyolite dome complex (Indio Muerto Rhyolite), intruded by rhyolite and granodiorite porphyry complexes</td>
<td>[50]</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Basement comprising andesite lava and volcaniclastic rocks</td>
<td></td>
</tr>
</tbody>
</table>
The primary purpose of the LA-ICP-MS analyses was to determine whether the investigated minerals have low (e.g., <10 ppm), moderate (e.g., 10–100 ppm) or high (e.g., >100 ppm) concentrations of Ge, Ga and In. The minerals differ significantly in their structure and major element composition, and thus, in which trace elements they can be expected to incorporate. We used two separate analytical protocols for sulfates/silicates and sulfides/sulfosalts, respectively. For alunite, barite and dickite, the collected analytes were $^{27}\text{Al}$, $^{31}\text{P}$, $^{44}\text{Ca}$, $^{53}\text{Cr}$, $^{55}\text{Mn}$, $^{57}\text{Fe}$, $^{65}\text{Cu}$, $^{66}\text{Zn}$, $^{71}\text{Ga}$, $^{72}\text{Ge}$, $^{75}\text{As}$, $^{77}\text{Se}$, $^{85}\text{Rb}$, $^{88}\text{Sr}$, $^{98}\text{Mo}$, $^{107}\text{Ag}$, $^{115}\text{In}$, $^{118}\text{Sn}$, $^{121}\text{Sb}$, $^{137}\text{Ba}$, $^{140}\text{Ce}$, $^{197}\text{Au}$, $^{208}\text{Pb}$, $^{209}\text{Bi}$, $^{232}\text{Th}$ and $^{238}\text{U}$. The internal standard isotopes used for data reduction were $^{27}\text{Al}$ (alunite and dickite) and $^{137}\text{Ba}$ (barite), based on contents confirmed previously by WDS. NIST-SRM-610 [51] was used as a bracketing external standard, and GSE-1G and GSD-1G [52] were analyzed as secondary standards to evaluate accuracy. For enargite, sphalerite, pyrite and galena, the collected analytes were $^{51}\text{V}$, $^{52}\text{Cr}$, $^{55}\text{Mn}$, $^{57}\text{Fe}$, $^{59}\text{Co}$, $^{60}\text{Ni}$, $^{65}\text{Cu}$, $^{66}\text{Zn}$, $^{71}\text{Ga}$, $^{72}\text{Ge}$, $^{75}\text{As}$, $^{77}\text{Se}$, $^{95}\text{Mo}$, $^{107}\text{Ag}$, $^{114}\text{Cd}$, $^{115}\text{In}$, $^{118}\text{Sn}$, $^{121}\text{Sb}$, $^{125}\text{Te}$, $^{197}\text{Au}$, $^{202}\text{Hg}$, $^{205}\text{Tl}$, $^{208}\text{Pb}$ and $^{209}\text{Bi}$. USGS Mass-1 [53] was used as a bracketing external standard, and NIST-SRM-610, GSE-1G and GSD-1G were used as secondary standards. $^{65}\text{Cu}$ (enargite), $^{66}\text{Zn}$ (sphalerite), $^{57}\text{Fe}$ (pyrite) and $^{208}\text{Pb}$ (galena) were used as the internal standard isotopes, based on contents confirmed previously by WDS.

Effort has been made to only report data reflecting elements hosted in solid solution within the studied minerals. Samples were inspected under the polarized light microscope and in backscatter electron images prior to LA-ICP-MS analyses, so that only surfaces without visible mineral inclusions (and in the case of barite, fluid inclusions) were ablated. Segments in downhole spectra showing evidence for inclusions beneath the sample surface (i.e., sharp peaks in specific elements) were excluded during data processing, which was performed using the software GLITTER [54].

5. Occurrence and Composition of Key Minerals at Mt Carlton

5.1. Alunite

Alunite [trigonal $\text{KAl}_3(\text{SO}_4)_2(\text{OH})_6$] occurs within the silicic and quartz-alunite alteration zones at Mt Carlton. Stage 1A magmatic-hydrothermal alunite [55,56] comprises up to ~300 µm long euhedral, platy crystals occurring as disseminations, as infill in vugs, and as replacements of pre-existing feldspar crystals (Figures 4A and 5A). Stage 1B magmatic-steam alunite [55,56] occurs as coarse-grained, plumose and banded alunite in veins (Figures 4B and 5B). Previous WDS analyses of both types of alunite showed that they belong to the alunite-natroalunite solid solution series [$\text{KAl}_3(\text{SO}_4)_2(\text{OH})_6$–$\text{NaAl}_3(\text{SO}_4)_2(\text{OH})_6$] [30].

Trace element data was obtained for magmatic-hydrothermal alunite (n = 44, 11 samples) and magmatic-steam alunite (n = 21, 3 samples) from the V2 pit at Mt Carlton. Both types of alunite exhibit moderate to locally high concentrations of Ga. The average Ga concentrations are 32 ppm (up to 109 ppm) in magmatic-hydrothermal alunite and 72 ppm (up to 339 ppm) in magmatic-steam alunite (Table 3, Supplementary Table S1). Gallium shows smooth downhole signals in alunite, suggesting it is hosted in solid solution. Furthermore, the signals of Ga closely match the signals of Al. Indeed, incorporation of Ga into minerals of the alunite supergroup is generally attributed to direct substitution of $\text{Ga}^{3+}$ into the G site normally occupied by $\text{Al}^{3+}$ or $\text{Fe}^{3+}$ [57–60]. The concentrations of Ge and In in alunite were consistently below 10 ppm and 2 ppm, respectively. Overall, magmatic-hydrothermal and magmatic-steam alunite display similar trace element signatures. Elements with average concentrations in excess of 100 ppm include P, Ca, Sr, Ba and Pb (both types), Fe (magmatic-hydrothermal alunite) and Ce (magmatic-steam alunite) (Table 3, Supplementary Table S1).
Table 3. Summary table of LA-ICP-MS trace element compositions of minerals from Mt Carlton and similar deposits worldwide.

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Location</th>
<th>Mineral Assemblage</th>
<th>n (Samples)</th>
<th>Ge (ppm) Mean (max)</th>
<th>Ga (ppm) Mean (max)</th>
<th>In (ppm) Mean (max)</th>
<th>Au (ppm) Mean (max)</th>
<th>Other Elements Mean &gt;1000 ppm</th>
<th>Other Elements Mean &gt;100 ppm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alunite</td>
<td>V2 pit</td>
<td>Stage 1A</td>
<td>44 (11)</td>
<td>1.2 (9.1)</td>
<td>32 (109)</td>
<td>0.1 (1.3)</td>
<td>0.0 (0.2)</td>
<td>Ba</td>
<td>P, Ca, Fe, Sr, Pb</td>
</tr>
<tr>
<td>Alunite</td>
<td>V2 pit</td>
<td>Stage 1B</td>
<td>21 (3)</td>
<td>0.9 (3.7)</td>
<td>72 (339)</td>
<td>0.0 (0.2)</td>
<td>0.0 (0.2)</td>
<td>P, Sr</td>
<td>Ca, Ba, Ce, Pb</td>
</tr>
<tr>
<td>Enargite</td>
<td>V2 pit</td>
<td>Stage 2A</td>
<td>14 (2)</td>
<td>86 (330)</td>
<td>6.0 (37)</td>
<td>14 (155)</td>
<td>12 (62)</td>
<td>Sb, Bi</td>
<td>Fe, Zn, Se, Ag, Sn, Te</td>
</tr>
<tr>
<td>Enargite</td>
<td>A39 pit</td>
<td>Stage 2A</td>
<td>20 (2)</td>
<td>612 (2189)</td>
<td>5.2 (35)</td>
<td>7.7 (76)</td>
<td>0.2 (1.4)</td>
<td>Sb</td>
<td>Fe, Zn, Se, Ag, Te, Pb, Bi</td>
</tr>
<tr>
<td>Barite</td>
<td>A39 pit</td>
<td>Stage 2A</td>
<td>53 (4)</td>
<td>0.3 (1.8)</td>
<td>0.4 (1.4)</td>
<td>0.0 (0.0)</td>
<td>0.0 (0.0)</td>
<td>Pb, Sr</td>
<td>Ca, La, Ce</td>
</tr>
<tr>
<td>Pyrite</td>
<td>V2 pit</td>
<td>Stage 2A</td>
<td>14 (2)</td>
<td>3.5 (4.7)</td>
<td>0.2 (0.4)</td>
<td>0.1 (0.2)</td>
<td>2.4 (11)</td>
<td>-</td>
<td>Co, Ni, Cu, Zn, As, Ag</td>
</tr>
<tr>
<td>Pyrite</td>
<td>V2 pit</td>
<td>Stage 2B</td>
<td>14 (2)</td>
<td>3.7 (4.2)</td>
<td>0.1 (0.3)</td>
<td>0.1 (0.8)</td>
<td>56 (367)</td>
<td>Au</td>
<td>Cu, Pb</td>
</tr>
<tr>
<td>Sphalerite</td>
<td>V2 pit</td>
<td>Stage 2A</td>
<td>5 (1)</td>
<td>77 (143)</td>
<td>677 (1181)</td>
<td>469 (571)</td>
<td>10 (21)</td>
<td>Cu, Cd, Pb</td>
<td>As, Se, Ag, Sb, Te</td>
</tr>
<tr>
<td>Sphalerite</td>
<td>V2 pit</td>
<td>Stage 2B</td>
<td>44 (4)</td>
<td>206 (611)</td>
<td>444 (2829)</td>
<td>360 (2169)</td>
<td>13 (75)</td>
<td>Cu, Cd, Pb</td>
<td>As, Ag, Sb, Hg</td>
</tr>
<tr>
<td>Sphalerite</td>
<td>A39 pit</td>
<td>Stage 2B</td>
<td>10 (1)</td>
<td>5.1 (8.5)</td>
<td>260 (476)</td>
<td>20 (40)</td>
<td>0.5 (2.1)</td>
<td>Cd, Pb</td>
<td>Fe, Cu, As, Sb</td>
</tr>
<tr>
<td>Galena</td>
<td>V2 pit</td>
<td>Stage 2B</td>
<td>10 (1)</td>
<td>0.6 (0.9)</td>
<td>0.0 (0.1)</td>
<td>2.4 (12)</td>
<td>1.1 (2.1)</td>
<td>Cu, Sb</td>
<td>Zn, Cd</td>
</tr>
<tr>
<td>Galena</td>
<td>A39 pit</td>
<td>Stage 2B</td>
<td>7 (1)</td>
<td>0.3 (0.8)</td>
<td>bd</td>
<td>0.7 (1.2)</td>
<td>0.4 (0.9)</td>
<td>Ag, Sb</td>
<td>-</td>
</tr>
<tr>
<td>Dickite</td>
<td>V2 pit</td>
<td>Stage 3</td>
<td>10 (1)</td>
<td>3.1 (4.1)</td>
<td>100 (109)</td>
<td>0.1 (0.5)</td>
<td>0.0 (0.0)</td>
<td>P, Sr</td>
<td>Ca, Fe, Ba, Pb</td>
</tr>
<tr>
<td>Dickite</td>
<td>A39 pit</td>
<td>Stage 3</td>
<td>30 (3)</td>
<td>5.2 (9.1)</td>
<td>86 (150)</td>
<td>0.0 (0.2)</td>
<td>0.0 (0.0)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td><strong>Samples from other high-sulfidation epithermal and porphyry deposits</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Enargite</td>
<td>Lepanto</td>
<td>Eng, Py, Luz, Tn-Td, Cv</td>
<td>15 (1)</td>
<td>38 (57)</td>
<td>0.2 (1.3)</td>
<td>3.7 (8.0)</td>
<td>1.3 (7.6)</td>
<td>Sb, Te</td>
<td>Fe, Se, Sn</td>
</tr>
<tr>
<td>Enargite</td>
<td>Laurani</td>
<td>Eng, Py, Luz, Tn-Td</td>
<td>15 (1)</td>
<td>12 (15)</td>
<td>0.0 (0.3)</td>
<td>9.1 (17)</td>
<td>0.2 (1.4)</td>
<td>Sb, Sn</td>
<td>Zn, Cd, Te, Hg</td>
</tr>
<tr>
<td>Enargite</td>
<td>La Mejicana</td>
<td>Eng, Py, Luz</td>
<td>16 (1)</td>
<td>717 (2679)</td>
<td>4.2 (15)</td>
<td>5.2 (18)</td>
<td>1.4 (10)</td>
<td>Sb</td>
<td>Fe, Zn, Sn, Bi</td>
</tr>
<tr>
<td>Enargite</td>
<td>Motombo</td>
<td>Eng, Py</td>
<td>15 (1)</td>
<td>21 (50)</td>
<td>0.4 (3.3)</td>
<td>1.3 (1.8)</td>
<td>0.2 (0.8)</td>
<td>Sb, Bi</td>
<td>Fe, Se</td>
</tr>
<tr>
<td>Enargite</td>
<td>Chelopech</td>
<td>Eng, Py, Luz, Cp, Tn-Td</td>
<td>15 (1)</td>
<td>101 (186)</td>
<td>0.2 (0.6)</td>
<td>1.1 (1.3)</td>
<td>0.1 (0.4)</td>
<td>Sb</td>
<td>Fe, Bi</td>
</tr>
<tr>
<td>Enargite</td>
<td>Cerro Quema</td>
<td>Eng, Py, Luz, Cp, Cv</td>
<td>10 (1)</td>
<td>562 (1202)</td>
<td>0.1 (0.6)</td>
<td>1.4 (2.0)</td>
<td>0.0 (0.3)</td>
<td>Sb, Fe</td>
<td>Hg</td>
</tr>
<tr>
<td>Enargite</td>
<td>El Salvador</td>
<td>Eng, Py, Tn-Td, Cp</td>
<td>5 (1)</td>
<td>12 (24)</td>
<td>0.0 (0.1)</td>
<td>1.3 (1.6)</td>
<td>15 (72)</td>
<td>Sb</td>
<td>Sn, Te</td>
</tr>
</tbody>
</table>

Abbreviations: bd—below detection; Cp—chalcopyrite; Cv—covellite; Eng—enargite; Luz—luzonite; Py—pyrite; Td—tetrahedrite; Tn—tennantite.
5.2. Argyrodite

Argyrodite [orthorhombic Ag₈GeS₆], the mineral from which Ge was first detected [61], occurs in the Stage 2A ores as up to ~100 μm big, anhedral crystals intergrown with barite and enargite (Figure 5C). The presence of argyrodite appears to be restricted to the distal, Ag-rich parts of the deposit (A39 pit). The analysed compositional range of Mt Carlton argyrodite is (Ag₅.⁷⁻⁷.₈₀Cu₀.₁₁⁻⁰.₆₇Pb₀.₀₀⁻₀.₀₁) (Ge₀.₈₆⁻¹.⁰₆As₀.₀₀⁻₀.₀₆Sb₀.₀₀⁻₀.₀₈Bi₀.₀₀⁻₀.₀₁) (S₄.₃₆⁻₅.₆₂Se₀.₆₁⁻¹.₄₄Te₀.₀₃⁻₀.₄₆), based on 23 spot analyses on two different grains from separate samples (Table 4, Supplementary Table S2). One grain showed a composition close to argyrodite *sensu stricto*, while the other grain corresponds to a Se-rich analogue of argyrodite [ideally Ag₈GeS₂S₄] (Figure 6). The presence of Cu in argyrodite suggests minor solid solution towards putzite [Cu₄.⁷Ag₃.₃GeS₆] [62]. The mineral also locally contains minor amounts of As, Sb and Bi, presumably substituting for Ge. The average Ge concentration in argyrodite is 5.76 wt % (up to 6.95 wt %) (Table 4, Supplementary Table S2).

<table>
<thead>
<tr>
<th>n = 23</th>
<th>Se</th>
<th>Ge</th>
<th>Ag</th>
<th>Te</th>
<th>S</th>
<th>As</th>
<th>Cu</th>
<th>Au</th>
<th>Sb</th>
<th>Bi</th>
<th>Pb</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean (wt %)</td>
<td>8.06</td>
<td>5.76</td>
<td>69.65</td>
<td>3.26</td>
<td>12.84</td>
<td>0.22</td>
<td>1.26</td>
<td>0.01</td>
<td>0.05</td>
<td>0.10</td>
<td>0.03</td>
<td>101.25</td>
</tr>
<tr>
<td>Minimum</td>
<td>4.34</td>
<td>5.19</td>
<td>68.56</td>
<td>0.38</td>
<td>11.66</td>
<td>0.03</td>
<td>0.59</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>0.00</td>
<td>99.94</td>
</tr>
<tr>
<td>Maximum</td>
<td>9.52</td>
<td>6.95</td>
<td>71.47</td>
<td>4.90</td>
<td>16.32</td>
<td>0.50</td>
<td>3.78</td>
<td>0.03</td>
<td>0.57</td>
<td>0.26</td>
<td>0.14</td>
<td>101.77</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>n = 23</th>
<th>Se</th>
<th>Ge</th>
<th>Ag</th>
<th>Te</th>
<th>S</th>
<th>As</th>
<th>Cu</th>
<th>Au</th>
<th>Sb</th>
<th>Bi</th>
<th>Pb</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean (apfu)</td>
<td>1.21</td>
<td>0.93</td>
<td>7.59</td>
<td>0.30</td>
<td>4.69</td>
<td>0.03</td>
<td>0.23</td>
<td>0.00</td>
<td>0.00</td>
<td>0.01</td>
<td>0.00</td>
<td></td>
</tr>
<tr>
<td>Minimum</td>
<td>0.61</td>
<td>0.86</td>
<td>7.03</td>
<td>0.03</td>
<td>4.36</td>
<td>0.00</td>
<td>0.11</td>
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</tr>
<tr>
<td>Maximum</td>
<td>1.44</td>
<td>1.06</td>
<td>7.80</td>
<td>0.46</td>
<td>5.62</td>
<td>0.08</td>
<td>0.67</td>
<td>0.00</td>
<td>0.05</td>
<td>0.01</td>
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</tr>
</tbody>
</table>

Table 4. Electron microprobe composition of argyrodite from Mt Carlton.

Figure 6. Compositional plot of argyrodite from Mt Carlton, based on WDS spot data (in *apfu*). The two clusters represent analyses of two individual grains.

5.3. Enargite

Enargite [orthorhombic Cu₃AsS₄] is the principal mineral in the Stage 2A ores at Mt Carlton. Enargite occurs as masses of anhedral to subhedral crystals that host most of the subordinate minerals (Figures 4C and 5D). Trace element data was obtained for enargite from the V2 pit (n = 14, 2 samples) and the A39 pit (n = 20, 2 samples) (Table 3, Supplementary Table S3). Mt Carlton enargite shows moderate to high enrichment in Ge, with average concentrations of 86 ppm in V2 samples (up to...
and 612 ppm in A39 samples (up to 2189 ppm). Both In (average 14 ppm in V2, 7.7 ppm in A39) and Ga (average 6.0 ppm in V2, 5.2 ppm in A39) are generally low, but significant concentrations are present locally (e.g., up to 155 ppm In and 37 ppm Ga in V2).

Comparing trace elements with average concentrations above 100 ppm in Mt Carlton enargites, the V2 samples are relatively enriched in Sb, Bi and Sn (Supplementary Table S3). Both Bi and Sn are elements indicative of higher temperature conditions [63] and both are commonly present in HS deposits (e.g., Pierina in Peru and Rodalquilar in Spain; [20]). Samples from A39 are in turn relatively enriched in Zn, Ag, Ge and Pb. The concentrations of Te, Se and Fe are similar in samples from V2 and A39. Gold has average concentrations of 12 ppm in V2 (up to 62 ppm) and 0.2 ppm in A39 (up to 1.4 ppm) (Supplementary Table S3). Previous studies of the Far Southeast-Lepanto porphyry-epithermal system (Mankayan, Philippines) have revealed a spatial variability in the trace element composition of enargite [64,65]. Enargite collected close to the Far Southeast porphyry deposit was enriched in Au-Te, enargite collected from the center of the Lepanto HS deposit was enriched in Ag-Fe-Pb, and enargite collected from the most distal ores was enriched in Zn (±Cd) [65]. From proximal to distal, the Sn content in enargite was progressively decreasing [64]. The trace element signatures for enargite from Mt Carlton in part corroborate the trends observed at Far Southeast-Lepanto, suggesting that this method may have some potential as a vectoring tool for exploration in the porphyry-epithermal environment. Furthermore, at the Rodalquilar HS deposit, the presence of anomalous Sn concentrations (high enough to form Sn-bearing minerals such as cassiterite (wood tin), colusite or rodalquilarite; [66]) was considered to be a prospective guide to high Au grade mineralized structures [67].

5.4. Barite

Barite [orthorhombic BaSO\(_4\)] is present in Stage 2A ores as up to ~0.5 cm large anhedral to subhedral crystals intergrown with the enargite assemblage (Figure 5C). More rarely it occurs as halos to enargite veinlets (Figure 4f). Barite is most abundant in the distal and shallow parts of the Mt Carlton deposit (A39 pit) [30]. The LA-ICP-MS analyses of barite from the A39 pit (n = 53, 4 samples) yielded concentrations of Ge, Ga and In consistently below 2 ppm (Table 3, Supplementary Table S1). Barite has high concentrations of Pb (average 1.3 wt %) and Sr (average 2406 ppm). This reflects significant solid solution toward hokutolite [(Ba,Pb,Sr)SO\(_4\)], a plumbian variety of barite reported from some acidic hot springs (e.g., the Tamagawa Hot Spring in Japan and the Peitou Hot Spring in Taiwan). The identification of Pb-rich barite at Mt Carlton has some significance for exploration, as this mineral potentially is responsible for the distal Pb geochemical anomaly observed at the deposit [30]. Other elements enriched in barite include Ca (average 333 ppm), La (average 202 ppm) and Ce (average 155 ppm), and to a lesser degree Al, P, Ti, Cu and Zn (all average 10–100 ppm) (Table 3, Supplementary Table S1).

5.5. Pyrite

Pyrite [isometric FeS\(_2\)] is present in Stage 2A ores as up to ~1 cm large subhedral crystals intergrown mainly with massive enargite and luzonite (Figure 4C). Stage 2B in turn contains up to ~0.5 cm large subhedral pyrite crystals mainly intergrown with sphalerite and galena (Figure 5E). Pyrite grains from Stage 2A (n = 14, 2 samples) and Stage 2B (n = 14, 2 samples) consistently show low concentrations of Ge, Ga and In (below 5 ppm; Table 3, Supplementary Table S3). Both stages of pyrite exhibit similar enrichment in Cu (average >700 ppm; Supplementary Table S3). Stage 2A pyrite shows a relative enrichment in Co, Ni, Zn and Ag (average 100–500 ppm), while Stage 2B pyrite is richer in As (average 2443 ppm, up to 1.2 wt %) and Pb (average 314 ppm). The high As concentrations in Stage 2B pyrite could be related to the low abundance of independent As minerals (e.g., tennantite) in this mineral assemblage. Furthermore, Stage 2B pyrite is significantly enriched in Au (average 56 ppm and up to 367 ppm) relative to Stage 2A pyrite (average 2.4 ppm and up to 11 ppm). This is consistent with previously made observations of high incorporation of Au in arsenian pyrite [68].
5.6. Sphalerite

Sphalerite [isometric ZnS] is present in small quantities in the Stage 2A ores. It occurs intergrown with enargite and luzonite as up to ~150 µm large, subhedral crystals that exhibit a dark brown to black color (Figure 5D). Sphalerite is the principal mineral in the Stage 2B ores, where it has a massive to colloform texture, with colors varying from almost transparent to dark brown (Figures 4D and 5E,F).

Trace element data was obtained for Stage 2A sphalerite from V2 (n = 5, 1 sample) and Stage 2B sphalerite from V2 (n = 44, 4 samples) and A39 (n = 10, 1 sample). Stage 2A sphalerite shows average concentrations of 77 ppm Ge (up to 143 ppm), 677 ppm Ga (up to 1181 ppm) and 469 ppm In (up to 571 ppm) (Table 3, Supplementary Table S3). Stage 2B sphalerite from V2 yielded average concentrations of 206 ppm Ge (up to 611 ppm), 444 ppm Ga (up to 2829 ppm) and 369 ppm In (up to 2169 ppm). Stage 2B sphalerite from A39 in turn shows significantly lower average concentrations of these elements, including 5.1 ppm Ge (up to 8.5 ppm), 260 ppm Ga (up to 476 ppm) and 20 ppm In (up to 40 ppm).

Both Stage 2A and Stage 2B sphalerite are highly enriched in Cu, Cd and Pb (average > 1000 ppm) (Table 3, Supplementary Table S3). The Cu content in Stage 2B sphalerite decreases from proximal (average 2013 ppm Cu in V2) to distal (average 486 ppm Cu in A39), which is consistent with the deposit scale metal zonation (Figure 2). Other elements with average concentrations in excess of 100 ppm in sphalerite include As, Ag and Sb (both stages), Se and Te (Stage 2A), and Fe and Hg (Stage 2B). Concentrations of Au are locally significant in sphalerite from both stages (up to 21 ppm in Stage 2A and up to 75 ppm in Stage 2B) (Table 3, Supplementary Table S3).

Germanium, Ga and In all show smooth downhole profiles in sphalerite, confirming that they are predominantly hosted in solid solution in this mineral [69]. Substitution of In and Ga into sphalerite can occur via coupled substitution with monovalent cations, and with Cu$^+$ in particular [1,15,16,42,69–73]. A positive correlation was observed in the Cu + Ag vs. In + Ga plot (Figure 7), corroborating the coupled substitution $2Zn^{2+} \leftrightarrow (Cu,Ag)^+ + (In,Ga)^{3+}$. In general there is an excess of Cu over Ag in sphalerite, suggesting that Cu is the principal cation involved in the substitution. No statistically significant correlation was observed between Ge and other trace elements in sphalerite, something that has also been noted in previous studies [69,74,75]. This could potentially suggest that Ge substituted for Zn with vacancies in the structure to allow for charge balance, i.e., $2Zn^{2+} \leftrightarrow Ge^{4+} + \square$ [74,75].

![Figure 7. Correlation plot of Cu + Ag vs. In + Ga in Mt Carlton sphalerite, based on LA-ICP-MS spot data.](image-url)
5.7. Galena

Galena [isometric PbS] in Stage 2B occurs as up to ~0.5 cm large, euhedral to subhedral crystals intergrown mainly with sphalerite and pyrite (Figure 5E). This galena contains abundant segregations of tennantite (Figure 5E). Trace element data for Stage 2B galena from V2 (n = 10, 1 sample) and A39 (n = 7, 1 sample) reveals concentrations of Ge, Ga and In that are consistently below 5 ppm (Table 3, Supplementary Table S3). Elements with average concentrations in excess of 100 ppm include Sb, Ag, Cu, Zn and Cd, which are all common trace components in galena [76]. The Ag content in galena is particularly high in samples from the A39 pit (average 1443 ppm, up to 3447 ppm). No independent Ag minerals have been identified in the Stage 2B assemblage, however, whole rock analyses of these ores have showed significant Ag grades (up to 2143 ppm Ag in samples from A39; [30]). The abundance of galena combined with the high Ag content in the mineral, therefore, suggest it is an important Ag carrier in the Stage 2B ores.

5.8. Dickite

Dickite [monoclinic Al$_2$Si$_2$O$_5$(OH)$_4$] within the Stage 3 void fill is generally massive, microcrystalline and exhibits a translucent to whitish or bluish color (Figure 4F). Trace element data was obtained for dickite from the V2 pit (n = 10, 1 sample) and the A39 pit (n = 30, 3 samples). Dickite exhibits moderate to locally high enrichment in Ga, with average concentrations of 100 ppm (up to 109 ppm) in V2 samples and 86 ppm (up to 150 ppm) in A39 samples (Table 3, Supplementary Table S1). Smooth downhole signals were observed for Ga in dickite, and the signals closely matched those of Al, again suggesting direct substitution of Ga$^{3+}$ for Al$^{3+}$. Germanium and In were always below 10 ppm and 1 ppm, respectively. Other trace elements present in significant concentrations in Mt Carlton dickite include P, Ca, Fe, Sr, Ba and Pb, and the V2 sample showed distinctly higher concentrations of these elements relative to the A39 samples (Table 3, Supplementary Table S1).

6. Composition of Enargite from Reference Deposits

Reference samples of enargite were analyzed from Lepanto (n = 15), Laurani (n = 15), La Mejicana (n = 16), Motomboko (n = 15), Chelopech (n = 15), Cerro Quema (n = 10) and El Salvador (n = 5) (Table 2). Only one polished section lacking detailed spatial information was analyzed from each locality. Given that a spatial variability can be expected for many trace elements in enargite ([64,65]; this study), these samples are not necessarily representative for the entire deposit from which they are derived. Nevertheless, the samples provide valuable information on the trace element content of enargite from HS deposits, which can be compared with the results from Mt Carlton.

Similar to enargite from Mt Carlton, moderate to high Ge concentrations were observed in all of the reference samples. This includes average Ge concentrations of 717 ppm for La Mejicana (up to 2679 ppm), 562 ppm for Cerro Quema (up to 1202 ppm), 101 ppm for Chelopech (up to 186 ppm), 38 ppm for Lepanto (up to 57 ppm), 21 ppm for Motomboko (up to 50 ppm), 12 ppm for El Salvador (up to 24 ppm) and 12 ppm for Laurani (up to 15 ppm) (Table 3, Supplementary Table S4). Downhole profiles of Ge in enargite (also including Mt Carlton samples) were smooth to undulating. This indicates that Ge occurs in solid solution, with a locally zoned distribution, something that is also visible in WDS element maps (Figure 8). Germanium shows a broad correlation with Zn in both WDS element maps (Figure 8) and in LA-ICP-MS spot data (Figure 9). Assuming that Cu is monovalent and As is pentavalent in the enargite lattice [77], this might suggest that Ge is in part incorporated via the coupled substitution As$^{5+}$ + Cu$^+$ $\leftrightarrow$ Ge$^{4+}$ + Zn$^{2+}$, as proposed by Bernstein (1985) [9]. Specifically, enargite from La Mejicana exhibits a strong correlation between Ge and Zn (Figures 8B and 9); however, the absolute concentrations of Zn are consistently lower than those of Ge (Figure 9). This suggests that additional substitution mechanisms must also be involved in the incorporation of Ge into enargite.
Both Ga and In are generally low in the reference enargite samples, however, moderate concentrations were observed locally in the samples from La Mejicana (up to 15 ppm Ga and 18 ppm In) and Laurani (up to 17 ppm In) (Table 3, Supplementary Table S4). Other trace elements with average concentrations in excess of 100 ppm in the reference enargite samples include Sb (all deposits), Fe (Cerro Quema, Motomboto, Chelopech, La Mejicana, Lepanto), Zn (La Mejicana, Laurani), Se (Lepanto, Motomboto), Cd (Laurani), Sn (Laurani, Lepanto, La Mejicana, El Salvador), and As (Chelopech, La Mejicana).

A particularly strong correlation between Zn and Ge, however, the concentrations of Zn are consistently lower than those of Ge (below the 1:1 correlation line).

Figure 8. BSE images (left) and X-ray intensity maps of selected elements in enargite. (A) Enargite crystal from Mt Carlton showing an oscillatory zoning primarily related to Sb content. Zinc and Ge are homogenously distributed in the crystal, while Sn shows a faint enrichment in Sb-rich zones. (B) Zoned enargite crystal from La Mejicana. Growth zones enriched in Ge are also enriched in Zn. A relatively homogenous distribution is observed for Sn, but locally it shows high enrichments in areas also marked by a high Sb content. Mineral abbreviations: Eng—enargite; Py—pyrite.

Figure 9. Correlation plot of Zn vs. Ge in enargite from all deposits included in this study (red diamonds), based on LA-ICP-MS spot data. Enargite from La Mejicana (highlighted in green) show a particularly strong correlation between Zn and Ge, however, the concentrations of Zn are consistently lower than those of Ge (below the 1:1 correlation line).
Laurani), Se (Lepanto, Motomboto), Cd (Laurani), Sn (Laurani, Lepanto, La Mejicana, El Salvador), Te (El Salvador, Lepanto, Laurani), Hg (Laurani, Cerro Quema) and Bi (Motomboto, Chelopech, La Mejicana) (Table 3, Supplementary Table S4). The Au content in these enargites varied from below detection limit to tens of ppm (up to 72 ppm at El Salvador), confirming that enargite can be a significant Au carrier in HS and porphyry deposits [65].

7. Discussion

The results of the present study, combined with available evidence from case studies in the literature, suggest that there are three distinct mineral associations in HS deposits that may host critical element enrichment. These include: (1) Ga enrichment in Al-bearing hydrothermal sulfates and silicates, (2) Ge ± In-Ga enrichment in high-sulfidation enargite ores, and (3) In-Ga ± Ge enrichment in intermediate-sulfidation sphalerite ores.

7.1. Ga Enrichment in Al-Bearing Sulfates and Silicates

This study highlights how Ga may be present in moderate to high concentrations in Al-bearing hydrothermal sulfates and silicates such as alunite and dickite, due to substitution of Ga$^{3+}$ into the structural sites normally occupied by Al$^{3+}$ in these minerals. Equivalent substitutions (Ga$^{3+}$ ↔ Fe$^{3+}$) should be possible in Fe$^{3+}$-analogues to such minerals. Exceptionally high Ga contents in minerals of the alunite supergroup have previously been documented in the oxidized zones of Tsumeb-type polymetallic deposits. The Tsumeb deposit in Namibia locally contains supergene gallobudantite [PbGa$_3$(AsO$_4$)(SO$_4$)(OH)$_6$] [58] and galloplumbogummite [Pb(Ga,Al)$_3$(PO$_4$)$_2$(OH)$_6$] [78], while similar ores at the Apex Mine in Utah contain supergene jarosite [the Fe$^{3+}$-analogue to alunite; KFe$_3$(SO$_4$)$_2$(OH)$_6$] with up to 7000 ppm Ga [79,80]. At the Apex Mine, the world’s only deposit ever considered as a primary Ga and Ge producer, jarosite is one of the principal Ga carriers [79,80].

Compared to the above examples, the Ga concentrations in alunite and dickite at Mt Carlton are distinctly lower. However, they are comparable to those of worldwide bauxite deposits (average 57 ppm Ga; [81]), the main source of the global Ga supply [14]. Similar Ga enrichment to that at Mt Carlton has previously been documented in whole rock samples from the advanced argillic alteration zone at the Paradise Peak HS deposit in Nevada (up to 120 ppm Ga; [82]). The similarities between X-ray absorption spectra of these samples and those of gallium sulfate suggest that the Ga there is predominantly hosted in alunite [82]. Alunite is a primary ore mineral for Al, K and S in exceptionally alunite-rich HS deposits similar to those at Alunite Ridge in Utah [83] and Red Mountain in Colorado [84], several of which have been exploited historically. The added possibility of recovering Ga as a by-product could spark a renewed interest in such occurrences. Furthermore, there could be potential for recovery of Ga from alunite present in the gangue in HS deposits that are primarily mined for their precious and/or base metal contents. Methods for extracting Ga from alunite concentrate have been developed in Japan (e.g., [85]), and more recently in China (e.g., [86]).

7.2. Ge ± In-Ga Mineralization in High-Sulfidation Enargite Ores

The mineralogical distribution of Ge in hydrothermal ores is dependent on the activities of sulfur and oxygen. In relatively reduced fluids with a high sulfur activity, Ge preferentially forms thio germanate minerals and substitutes into sulfosalts rather than sphalerite (Figure 10; [7,9]). This style of Ge mineralization is best known from Tsumeb-type deposits [7,9,87,88], however, it appears to be common also in HS deposits.

The presence of argyrodite at Mt Carlton is not unique for HS deposits, and a variety of independent Ge sulfide minerals have previously been documented in HS deposits elsewhere. Deposits from the Cretaceous Panagyurishte district of Bulgaria (including the Chelopech, Radka, Krassen and Elshitsa HS deposits) locally contain germanite [Cu$_{13}$Fe$_2$Ge$_2$S$_{16}$], renierite [(Cu,Zn)$_{11}$Fe$_4$(Ge,As)$_2$S$_{16}$], germanocolusite [Cu$_{13}$V(Ge,As)$_2$S$_{16}$], briartite [Cu$_2$(Zn,Fe)GeS$_4$] and Ge-bearing arsenosulvanite [Cu$_3$(As,V,Ge)S$_4$] [89–94]. Neogene epithermal mineralization in the South Apuseni Mountains
in Romania (including the Bucium, București–Rovina, Băița Crăciunești and Părâul lui Avram occurrences) contain argyrodite and/or germanite associated with enargite ([95], and references therein). The Capillitas deposit (Catamarca province, Argentina) comprises a number of mainly intermediate-sulfidation epithermal veins, some of which show high-sulfidation affinities [96]. Oxidized enargite-bearing ores in the mine dumps from the La Rosario vein exhibit a supergene mineral assemblage that contains several original species of Ge sulfides, including putzite [Cu4.7Ag3.3GeS6], catamarcaite [Cu6GeWS8] and omariniite [Cu8Fe2ZnGe2S12], with zincobriartite also being present in trace amounts [62,96–100].

Sulfosalts such as enargite, luzonite and tennantite-tetrahedrite are known to incorporate significant concentrations of Ge in Tsumeb-type deposits (up to ~5000 ppm; [7,9,87,88]). On the basis of LA-ICP-MS analyses of enargite from seven HS deposits and one porphyry deposit conducted in this study, moderate Ge concentrations (10–100 ppm) seem to be common in enargite in these systems. Higher Ge concentrations in enargite (e.g., 100–3000 ppm) are potentially restricted to a smaller number of deposits (e.g., Mt Carlton, La Mejicana, Cerro Quema and Chelopech). Previously, Maydağan et al. [101] reported up to 41 ppm Ge in enargite from the Altar linked porphyry-HS deposit in Argentina. Dekov et al. [102] studied seafloor chimneys made up almost entirely of luzonite and enargite in the Manus backarc basin (offshore of Papua New Guinea). Enargite from these chimneys, which were interpreted as submarine analogues of terrestrial HS deposits, was found to contain up to 137 ppm Ge and 11 ppm Ga, while luzonite contained up to 120 ppm Ge and 106 ppm Ga [102]. Indium was consistently below 1 ppm in both enargite and luzonite. Contrary to the results presented in this study, the highest Ge concentrations in these chimneys were observed in pyrite and marcasite (up to 1287 ppm Ge; [102]).
Sphalerite is typically a volumetrically minor component of high-sulfidation enargite ores [20]. Based on Figure 10, the Ge concentrations in such sphalerite is expected to be comparatively low (e.g., one order of magnitude lower than those in enargite at Mt Carlton). Sphalerite is probably the preferred host for In and Ga in this assemblage. However, moderate enrichment of these elements may also be present in sulfosalt minerals.

At Mt Carlton, the presence of argyrodite and the high Ge content of enargite are predominantly confined to the distal and shallow, Ag-rich parts of the deposit. This may suggest that mixing of the ore fluid with cooler meteoric waters, and a high activity of Ag, were important controls on this type of Ge mineralization. High-sulfidation enargite ores are generally voluminous in most HS deposits and, thus, represent a potential source of Ge, in addition to their commonly exploited Au, Ag and Cu contents. A potential drawback is the high concentrations of “penalty elements” (e.g., As, Sb and Bi) commonly present in unoxidized ores of this type, which can make them challenging to process economically at lower grades [22,103]. This would probably be less of an issue in oxidized HS deposits, where the ores are significantly easier to process [22]. Indeed, the gossans of some HS deposits (e.g., the La Rosario vein), Tsumeb-type deposits (e.g., Tsumeb and Apex Mine) and polymetallic vein deposits (e.g., San Roque and Pingüino, Argentina; [104,105]) are known to be enriched in critical elements, which can be attributed to the general immobility of Ge, Ga and In during weathering [9,80,105,106]. In the weathering environment, these elements may form their own exotic minerals, or occur in high concentrations in various supergene minerals (e.g., sulfides, oxides, hydroxides, hydroxy-sulfates and arsenates; [9,80,98,104]). From both a mineralogical and economic standpoint it can, therefore, be of interest to further investigate potential mineral hosts of Ge, Ga and In in oxidized HS deposits.

### 7.3. In-Ga ± Ge Mineralization in Intermediate-Sulfidation Sphalerite Ores

Indium and Ga display a similar distribution in the intermediate-sulfidation sphalerite ores at Mt Carlton. Both elements are primarily hosted in sphalerite and in the “Zn-In mineral”. The highest concentrations of In and Ga in sphalerite occur in the proximal, Cu-rich parts of the deposit. Indeed, a high Cu activity in the ore fluid is a strong control on In enrichment and, to a lesser degree, Ga enrichment in sphalerite [2], as it allows for extensive incorporation of these elements in coupled substitution with Cu.

Germanium also appears to be hosted predominantly in sphalerite in the Stage 2B ores at Mt Carlton. Indeed, Ge is expected to primarily concentrate into sphalerite at low to moderate sulfur activities (Figure 10; [7,9]). However, a high activity of Cu or Ag may still promote the formation of Ge-bearing Cu or Ag sulfides at lower sulfur activities [9]. The precise nature of Ge incorporation into sphalerite is less clear, and potentially more complex, than that of In and Ga [10,69,74,107,108]. It is, therefore, difficult to assess possible physicochemical controls on Ge enrichment in these ores.

In-Ga ± Ge enrichment in intermediate sulfidation sphalerite ores similar to those at Mt Carlton have been documented elsewhere. The most prominent example is the world-class Toyoha IS deposit in Japan. Toyoha was the world’s largest In producer in the 1990s—early 2000s, containing 33,700 kt of ore with an average In grade of 138 ppm [11]. Similar to Mt Carlton, the main carriers of In at Toyoha are In-bearing sphalerite (up to 7 wt % In, 3.6 wt % Cu and 273 ppm Ga; [15,69]) and the “Zn-In mineral” [43]. Other In-bearing minerals at Toyoha include roquesite [CuInS2], stannite-kösterite [Cu2FeSnS4–Cu2ZnSnS4], sakuraiite [(Cu,Zn,Fe,In,Sn)S], laforetite [Ag(InS2)] and chalcopyrite [43]. The Ge content of Toyoha sphalerite is low (1–4 ppm; [69]). The Agios Philippos HS deposit in Thrace, Greece, contains a stage of Zn-Pb-rich intermediate-sulfidation ore rich in sphalerite and wurtzite (the hexagonal polymorph of sphalerite). Sphalerite and wurtzite at Agios Philippos exhibit high concentrations of In (up to 3.5 wt %), Ga (up to 1.6 wt %) and Ge (up to 800 ppm) [69]. The Roșia Montană IS deposit in the South Apuseni Mountains in Romania locally contains alburnite [Ag8GeTe2S4], the Te-rich analogue of argyrodite [95,112], as well as sphalerite with up to 73 ppm Ge, 366 ppm Ga, 38 ppm In and 190 ppm Ag [69]. The Nuevo Esperanza HS vein at the Capillitas deposit in Argentina is the type locality of the rare Ga-In mineral ishiharaite [(Cu,Ga,Fe,In,Zn)S]. The mineral
is associated with a late intermediate-sulfidation state assemblage containing In-bearing sphalerite and tennantite, as well as pyrite, chalcopyrite and galena [113]. A final example is the Palai-Islica IS deposit in Spain, where sphalerite is variably enriched in Ge (up to 0.42 wt %), Ga (up to 0.16 wt %), In (up to 0.43 wt %) and Cu (up to 1.72 wt %) [114].

There is, thus, a noteworthy potential for In-Ga ± Ge enrichment in (Cu-rich) intermediate-sulfidation sphalerite ores in the epithermal environment. This mineralization style may be present in both HS deposits (e.g., Mt Carlton, Agios Philippos, Nueva Esperanza vein) and IS deposits (e.g., Toyoha, Roșia Montană and Palai-Islica). When sphalerite in such deposits is volumetrically significant enough to be processed, the possibility of extracting critical elements as by-products could increase the value of these ores (e.g., Toyoha).

7.4. Petrogenetic Considerations

Studies of active volcanic fumaroles further highlight the potential for significant transport and enrichment of Ge, Ga and In in epithermal settings. Analyses of volcanic gases and precipitates from Merapi Volcano, Indonesia [115,116] and Kudryavy Volcano, Russia [117,118] show significant enrichment factors for these elements (1–10$^2$ for Ge and Ga, 10$^4$–10$^5$ for In) in the fumarolic discharge products compared to the underlying, degassing magma. For comparison, the hydrothermal system related to the submarine Brothers Volcano (Kermadec Arc, New Zealand) has a strong magmatic contribution, and mineralogical similarities to terrestrial HS deposits have been documented (e.g., enargite, alunite and anhydrite; [63,119]). Analyses of precipitates from several seafloor chimneys related to Brothers revealed locally high concentrations of Ga (up to 1870 ppm), and to a lesser degree Ge (up to 194 ppm) and In (up to 144 ppm) [63]. In most hydrothermal fluids, transport of Ge and Ga is expected to dominantly be in the form of hydroxide complexes. However, fluoride complexes or sulfur-bearing species may also be important in particularly F-rich or S-rich solutions, respectively ([106], and references therein). The behavior of In is expected to be more variable, and hydroxide, chloride, fluoride or bisulfide complexes may all contribute to its transport, depending on conditions ([106], and references therein).

The incompatible elements Ge, Ga and In may be enriched through fractional crystallization, remaining in the residual igneous liquids [1,7,9,120]. Due to their chalcophile properties, enrichment of these elements has been linked to sulfur-rich and oxidized magmatism [1,9,11,16,121], typical of a porphyry-epithermal environment [23,122]. Additionally, Ge, Ga and In may be mobilized into magmas and hydrothermal fluids from sedimentary country rocks, in particular those rich in organic matter. Both Ge and Ga form strong bonds with organic compounds in coals and related organic materials, and coal deposits represent a significant resource of Ge and Ga globally [7,9,13,14]. As an example, coal seams from the upper Devonian Chattanooga black shale in Tennessee were found to contain up to 760 ppm Ge (up to 4.5 wt % Ge and 50 ppm Ga in the coal ash), with the shale itself containing up to 18 ppm Ge [123]. Uptake of Ge and Ga from such carbonaceous sedimentary rocks by migrating hydrothermal fluids has been suggested as an important process for Ge-Ga mineralization in several MVT and Tsumeb-type deposits [7,9,88]. Indium also has a slight tendency to be enriched in reduced, carbonaceous sedimentary rocks [124]. In the Japanese island arc, assimilation of basement pelitic rocks is believed to have played an important role in supplying In and reducing agents to the oxidized magmas that formed Toyoha and other In-Sn-bearing polymetallic vein deposits [11,16,121]. The HS deposits from which samples showed high enrichment in critical elements in this study (Mt Carlton, La Mejicana, Chelopech, Cerro Quema) are all hosted in stratigraphic sequences that locally contain carbonaceous sedimentary rocks (Table 2, and references therein; [30]). For example, the Panagyurishte district that hosts Chelopech and several other Ge-(Ga-In) enriched HS deposits (Radka, Krassen and Elshitsa) contains an up to 400 m thick basal sequence of coal-bearing sandstones and conglomerates [48,94]. Such rocks appear to be largely lacking or ambiguous in the deposits from which samples showed more moderate critical element enrichment (Lepanto, El Salvador, Motomboto), with Laurani being the exception (Table 2, and references therein).
In addition to magmatic-hydrothermal processes, carbonaceous sedimentary rocks could potentially be important for the concentration of Ge, Ga and In in HS deposits. Although this link requires more rigorous testing, we suggest that the presence of carbonaceous sedimentary rocks in the stratigraphic sequence hosting a HS deposit could provide a first-order exploration indicator for potential critical element mineralization.

7.5. Concluding Remarks and Future Developments

In summary, the evidence from this study and from the literature highlight the potential for various styles of Ge, Ga and In mineralization in high-sulfidation epithermal deposits, as well as in the closely related intermediate-sulfidation epithermal deposits. This is something that is seldom considered during exploration and mining in these environments. In addition to their well-known resources of precious and base metals, the potential for critical elements could increase the prospectivity of these deposits.

Holistic mineralogical studies of similar deposits elsewhere, including more trace element data for common ore and alteration minerals, can help improve our understanding of the distribution of Ge, Ga and In in the epithermal environment. Data are particularly limited for deposits affected by supergene processes. To improve exploration for these commodities, a better understanding of the source(s) as well as the mechanisms of enrichment, transport and deposition of Ge, Ga and In in epithermal systems is also needed. We find the potential link between critical element enrichment and carbonaceous sedimentary rocks particularly interesting and deserving of more detailed investigation.

Supplementary Materials: The following are available online at www.mdpi.com/2075-163X/7/11/213/s1, Table S1: LA-ICP-MS data for sulfates and silicates from Mt Carlton, Table S2: WDS data for argyrodite from Mt Carlton, Table S3: LA-ICP-MS data for sulfides and sulfosalts from Mt Carlton, Table S4: LA-ICP-MS data for enargite from reference localities.

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References


19. Hedenquist, J.W.; Simmons, S.F.; Giggenbach, W.F.; Eldridge, C.S. White Island, New Zealand, volcanic-hydrothermal system represents the geochemical environment of high-sulfidation Cu and Au ore deposition. *Geology* 1993, 21, 731–734. [CrossRef]


27. Jannas, R.R.; Beane, R.E.; Ahler, B.A.; Brosnahan, D.R. Gold and copper mineralization at the El Indio deposit, Chile. *J. Geochem. Explor.* 1990, 36, 233–266. [CrossRef]


42. Sahlström, F.; Blake, K.; Corral, I.; Chang, Z. Hyerspectral cathodoluminescence study of indium-bearing sphalerite from the Mt Carlton high-sulphidation epithermal deposit, Queensland, Australia. Eur. J. Mineral. 2017, in press. [CrossRef]

43. Ohta, E. Occurrence and chemistry of indium-containing minerals from the Toyoha mine, Hokkaido, Japan. Min. Geol. 1989, 39, 351–357. [CrossRef]


47. Perelló, J. Geology, porphyry Cu-Au, and epithermal Cu-Au-Ag mineralization of the Tombulilato district, North Sulawesi, Indonesia. J. Geochem. Explor. 1994, 50, 221–256. [CrossRef]

49. Corral, I.; Cardellach, E.; Corbella, M.; Canals, À.; Gómez-Gras, D.; Grier, A.; Cosca, M.A. Cerro Quema (Azuro Peninsula, Panama): Geology, alteration, mineralization, and geochronology of a volcanic dome-hosted high-sulfidation Au-Cu deposit. Econ. Geol. 2016, 111, 287–310. [CrossRef]

50. Gustafson, L.B.; Hunt, J.P. The porphyry copper deposit at El Salvador, Chile. Econ. Geol. 1975, 70, 857–912. [CrossRef]


56. Rye, R.O.; Bethke, P.M.; Watson, M.D. The stable isotope geochemistry of acid sulfate alteration. Econ. Geol. 1992, 87, 225–262. [CrossRef]


100. Bindi, L.; Putz, H.; Paar, W.H.; Stanley, C. Omariniite, Cu$_8$Fe$_2$ZnGe$_2$S$_{12}$, the germanium-analogue of stannoidite, a new mineral species from Capillitas, Argentina. *Mineral. Mag.* 2017, 81, 1151–1159. [CrossRef]


103. Lane, D.J.; Cook, N.J.; Grano, S.R.; Ehrig, K. Selective leaching of penalty elements from copper concentrates: A review. *Miner. Eng.* 2016, 98, 110–121. [CrossRef]


