






Article

Magnetite as a provenance and exploration tool for metamorphosed base-metal sulfide deposits in the Stollberg ore field, Bergslagen, Sweden

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Abstract

Magnetite is a common mineral in the Paleoproterozoic Stollberg Zn–Pb–Ag plus magnetite ore field (~6.6 Mt of production), which occurs in 1.9 Ga metamorphosed felsic and mafic rocks. Mineralisation at Stollberg consists of magnetite bodies and massive to semi-massive sphalerite–galena and pyrrhotite (with subordinate pyrite, chalcopyrite, arsenopyrite and magnetite) hosted by metavolcanic rocks and skarn. Magnetite occurs in sulfides, skarn, amphibolite and altered metamorphosed rhyolitic ash–siltstone that consists of garnet–biotite, quartz–garnet–pyroxene, gedrite–albite, and sericitic rocks. Magnetite probably formed from hydrothermal ore-bearing fluids (~250–400°C) that replaced limestone and rhyolitic ash–siltstone, and subsequently recrystallised during metamorphism. The composition of magnetite from these rock types was measured using electron microprobe analysis and LA–ICP–MS. Utilisation of discrimination plots (Ca+Al+Mn vs. Ti+V, Ni/(Cr+Mn) vs. Ti+V, and trace-element variation diagrams (median concentration of Mg, Al, Ti, V, Co, Mn, Zn and Ga) suggest that the composition of magnetite in sulfides from the Stollberg ore field more closely resembles that from skarns found elsewhere rather than previously published compositions of magnetite in metamorphosed volcanogenic massive sulfide deposits. Although the variation diagrams show that magnetite compositions from various rock types have similar patterns, principal component analyses and element–element variation diagrams indicate that its composition from the same rock type in different sulfide deposits can be distinguished. This suggests that bulk-rock composition also has a strong influence on magnetite composition. Principal component analyses also show that magnetite in sulfides has a distinctive compositional signature which allows it to be a prospective pathfinder mineral for sulfide deposits in the Stollberg ore field.

Keywords: magnetite, compositions, exploration, Stollberg deposit, Sweden

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Introduction

Trace-element studies of oxides, including gahnite (O'Brien *et al.* 2015a, 2015b, 2015c), hematite and maghemite (e.g. Schmidt Mumm *et al.*, 2012), chromite (e.g. Pagé and Barnes, 2009), ilmenite (e.g. Dare *et al.*, 2012), rutile (e.g. Clark and Williams-Jones, 2004) and magnetite (e.g. Dupuis and Beaudoin, 2011; Nadoll *et al.*, 2012a, 2014) have been used to determine the provenance of their host rocks and to explore for various types of ore deposits. Of these resistant oxides, magnetite is the most common and has been the subject of the largest number of trace-element studies as it occurs in a wide variety of geological settings including skarn, hydrothermal and magmatic ore deposits, glacial till and iron formations. The composition of magnetite is particularly useful for evaluating ore-forming

processes, as it is controlled by numerous factors, including: temperature; source rock/fluid composition; oxygen and sulfur fugacity; silicate and sulfide activity; host rock buffering; re-equilibration processes; and intrinsic crystallographic controls (i.e. ionic radius and charge balance; Nadoll *et al.*, 2014). Magnetite-bearing ore deposits include Ni–Cu–PGE, banded iron formation (BIF), iron oxide–Cu–Au (IOCG), skarn, porphyry Cu, Fe–REE–Nb, Cu–Au–Fe, iron oxide–apatite, and volcanogenic massive sulfide (VMS) deposits (e.g. Pagé and Barnes, 2009; Dupuis and Beaudoin, 2011; Nadoll *et al.*, 2012a; Dare *et al.*, 2012; Makvandi *et al.*, 2013; Acosta-Góngora *et al.*, 2014; Huang *et al.*, 2013, 2015; Chen *et al.*, 2015; Chung *et al.*, 2015; Pisiak *et al.*, 2015; Xu *et al.*, 2015; Zhao *et al.*, 2015; Qi *et al.*, 2021; Bédard *et al.*, 2022).

The objective of this study is to expand upon earlier trace-element studies of magnetite associated with metamorphosed massive sulfide and skarn deposits by evaluating the trace-element composition of magnetite in metamorphosed carbonate-associated massive sulfide deposits (Grängsgruvan, Tvistbo, Norrgruvan, Cedercreutz, Baklängen and Dammberget)

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in the Paleoproterozoic Stollberg Zn–Pb–Ag plus magnetite ore field, Bergslagen district, Sweden. We document the variability in the trace-element composition of magnetite in massive sulfides, different types of metamorphosed altered rocks (mainly altered rhyolitic ash–siltstone and limestone), mafic intrusions and massive magnetite bodies. The sulfide occurrences at Stollberg were considered by Jansson *et al.* (2013) and Frank *et al.* (2019) to be metamorphosed, replacement deposits overprinting limestone. The resulting deposits have geological features that resemble both skarn and metamorphosed VMS deposits. This study of magnetite in the Stollberg ore field, which is the first of its type in the Bergslagen district, specifically assesses: (1) the trace-element composition of magnetite in different rock types associated with metamorphosed sulfide occurrences; (2) whether or not the composition of magnetite can be used as a pathfinder to ore; and (3) the application of discrimination diagrams as aids in understanding the provenance of magnetite and ore deposit type.

Magnetite in metamorphosed massive sulfide deposits

Previous studies have been shown that the composition of magnetite can be used to fingerprint ore deposit types and provide insight into processes associated with the formation of VMS deposits (Singoyi *et al.*, 2006; Kamvong *et al.*, 2007; Dupuis and Beaudoin, 2011; Makvandi *et al.*, 2013; 2016a, 2016b; Bédard *et al.*, 2022). Using the composition of magnetite obtained from various VMS, skarn, iron oxide–Cu–Au, and Broken Hill-type Pb–Zn deposits, Singoyi *et al.* (2006) and Kamvong *et al.* (2007) suggested that these deposits can be distinguished on the basis of a plot of Sn/Ga vs. Al/Co. Dupuis and Beaudoin (2011) proposed that the average composition of magnetite from 17 VMS deposits and the Faro sedimentary exhalative deposit can be distinguished in Al/(Zn+Ca)–Cu/(Si+Ca) space from the average composition of magnetite from Fe–Ti, Cr, Cu vein, IOCG, Kiruna-type, banded iron formations, porphyry Cu, skarn, and magmatic Ni–Cu deposits. Included in the group of massive sulfide deposits that Dupuis and Beaudoin (2011) studied were Ducktown (Tennessee, USA) and Garpenberg (Sweden), which were metamorphosed to the amphibolite facies. In their study, Dupuis and Beaudoin (2011) only used samples from sulfide-bearing rocks that contained magnetite. Makvandi *et al.* (2016b) focused on both ore-bearing and unmineralised rocks from the Izok Lake Zn–Pb–Cu–Ag (Nunavut, Canada) and Halfmile Lake Zn–Pb–Cu (New Brunswick, Canada) VMS deposits, which were metamorphosed to the amphibolite and greenschist facies, respectively. They proposed that variability in trace-element composition of magnetite in samples of massive sulfides, altered rocks, less altered host rocks, and till from both locations was due to differences in geological setting and grades of metamorphism.

In addition to using scatter plots, multivariate statistical methods, including principal component analysis (PCA) and Random Forests, show promise in discriminating the trace-element signatures in different lithologies and ore deposit types (Bédard *et al.*, 2022). Makvandi *et al.* (2016b) showed, for example, that the trace-element composition of magnetite in massive sulfides, altered gahnite-rich dacite, gabbro and iron formation in the Izok Lake area could be distinguished using PCA. A further study by Makvandi *et al.* (2016a) applied PCA and partial least-squares discriminant analysis (PLS-DA) to magnetite from 18 VMS deposits in various types of rocks, including iron formations. Of these deposits, magnetite was analysed from some VMS

deposits metamorphosed to the greenschist (e.g. Brunswick #6 and #12, Austin Brook) and amphibolite facies (e.g. Izok Lake, Ducktown). The statistical approach of Makvandi *et al.* (2016a) allowed them to distinguish populations within large sets of trace-element compositions that would otherwise overlap in scatter plots, with PLS-DA being used by them to identify and classify different types of VMS deposits and BIF, as well as being able to discriminate magnetite in surficial sediments from those derived from other types of magmatic or hydrothermal ore deposits (e.g. iron oxide–Cu–Au, magmatic Ni–Cu and porphyry Cu).

Regional geology

The Paleoproterozoic Bergslagen mining district occurs in a supracrustal metavolcanic (predominantly calc-alkaline rhyolite)–metasedimentary province (1.91–1.75 Ga) belonging to the Fennoscandian Shield in south-central Sweden, together with pre- to post-tectonic intrusive rocks (1.90–1.87 Ga) (Allen *et al.* 1996; Stephens *et al.* 2009; Stephens and Jansson, 2020). It hosts a variety of ore deposit types including polymetallic Zn–Pb–Ag–(Cu)–(Au) sulfide and iron oxide deposits, BIF, skarn, and Mn oxide (Allen *et al.*, 2008; Stephens *et al.*, 2009). They are hosted typically by hydrothermally altered metavolcanic rocks and associated metacarbonates. Allen *et al.* (1996) described two types of polymetallic deposits in the Bergslagen district. They are bedded, stratiform Zn–Pb–Ag–(Cu)–(Ag)-rich mineralisation hosted in rhyolitic ash–siltstone (SAS-type) and stratabound volcanic-associated, limestone–skarn (SVALS-type). The district underwent amphibolite-grade metamorphism (with local areas of granulite and greenschist facies) during the Svecofennian orogeny (1.89–1.75 Ga), in association with four main phases of deformation that included rifting, isoclinal folding, shear-zone formation and widespread tight to isoclinal folding (Stephens *et al.*, 2009; Beunk and Kuipers, 2012). Extensive regional Na or K alteration is commonly early in the geological history and was associated with a diagenetic or hydrothermal event (Frietsch, 1982; Lagerblad and Gorbatschev, 1985). According to Allen *et al.* (1996), the Bergslagen district represents an extensional, continental margin, back-arc region developed on continental crust. Felsic pyroclastic and volcanoclastic material was deposited in a shallow marine basin by numerous caldera volcanoes, and stromatolite and microbial reef growth developed carbonate horizons during periods of relative volcanic inactivity (Allen *et al.*, 2003).

Base and precious metal deposits in the Bergslagen district, including the SVALS/SAS-type Falun (Sundblad, 1994; Kampmann *et al.* 2017), Garpenberg (Jansson and Allen, 2011, 2015), Sala (Jansson *et al.* 2021), Zinkgruvan (Hedström *et al.* 1989; Jansson *et al.* 2017) and Lovisa (Jansson *et al.* 2018) have been mined, however historically iron ore has been economically the most important. The iron deposits in the Bergslagen district have been classified traditionally on the basis of their phosphorous (or apatite) content and style of mineralisation (Geijer and Magnusson, 1944). Deposits with <0.2 wt.% P include hematite–magnetite BIF and skarn–limestone magnetite ores, and those with >0.2 wt.% P are classified as massive apatite-rich magnetite ores and disseminated apatite-bearing magnetite ores. Banded iron formation is subdivided typically into Mn-rich (>1 wt.%) and Mn-poor (<1 wt.%) types. Allen *et al.* (2008) divided the Fe oxide deposits hosted by skarn or carbonate rock into two types, the first of which is hosted by Mn-poor skarn or marble, and is up to 50 m wide. These occur mostly in the

supracrustal rocks in the central-western parts of the Bergslagen district and are the most common type of Fe oxide occurrence. The second type occurs in Mn-rich skarns, or, and in general, marble that is typically higher in the stratigraphic sequence in the Bergslagen district than the first type.

Local geology

A regional N–S trending, steeply dipping, F_2 syncline (herein referred to as the Stollberg syncline) is located ~4 km NE of the village of Ludvika, and is associated spatially with more than a dozen base-metal and magnetite deposits in the Stollberg ore field (Fig. 1). Sulfides are associated spatially with marble and skarn within bimodal felsic and mafic rocks and underwent amphibolite-grade metamorphism (560–600°C and 2–3.5 kbar; Beetsma, 1992; Björklund, 2011; Jansson *et al.*, 2013). Details of the local geology are described in detail by Ripa (1988, 1994, 2012), Jansson *et al.* (2013) and Frank *et al.* (2019) and are only summarised here.

Jansson *et al.* (2013) defined two stratigraphic successions (Stollberg and Staren) on the eastern side of the syncline. The lower part of the Staren succession consists of >200 m of massive metamorphosed quartz–feldspar rhyolitic sandstone that is overlain by the Staren limestone, which consists of coarse-grained locally pure marble that was, in places, altered to epidote-actinolite- and hedenbergite-bearing skarn. Interbeds of metamorphosed rhyolitic siltstone are also present. A mixture of rhyolitic and calcareous sandstones, breccias and conglomerates overlie the Staren limestone.

The Staren and Stollberg successions are separated by a rhyolitic intrusion that was affected by Na alteration. The base of the Stollberg succession consists of ~500 m of feldspar–phyric rhyolitic pumice breccia–sandstone that was altered hydrothermally and metamorphosed subsequently. This rock grades into the Stollberg limestone, which hosts most of the base-metal sulfide deposits (Table 1). A thin magnetite-bearing banded iron formation occurs a few tens of metres east of, and stratigraphically below, the Stollberg limestone. Near sulfides the limestone is altered to skarn, although where least altered, it consists of dolomitic and Mn-bearing marble. Over 700 m of metamorphosed planar-bedded rhyolitic ash silt–sandstone overlie the Stollberg limestone. The succession was altered hydrothermally to various intensities and metamorphosed subsequently to produce an array of metamorphosed altered rocks (e.g. Ripa, 1988, 1994, 2012; Jansson *et al.*, 2013). Amphibolite sills with a basaltic composition occur within the Stollberg succession (Björklund, 2011). Both the Stollberg and Staren successions were subsequently crosscut by diabase sills and dykes.

Mineral deposits in the Stollberg orefield consist of both Fe- and Mn-rich oxide deposits and stratabound carbonate replacement Zn–Pb–Ag–(Cu–Au) sulfide deposits (Jansson *et al.*, 2013) similar to deposits found elsewhere in the Bergslagen region (e.g. Geijer, 1917; Allen *et al.*, 1996; Stephens and Jansson, 2020). Metamorphosed altered rocks are associated spatially with base-metal sulfide deposits and include quartz–garnet–pyroxene rock, skarn, garnet–biotite, gedrite–albite, cordierite–biotite, siliceous, sericitic and quartz–fluorite rocks (Table 1). The term skarn here refers to stratabound or stratiform calc–silicate rocks dominated by clinopyroxene, garnet, and/or clinoamphibole without any necessary spatial or genetic link to igneous intrusion, as described by Jansson *et al.* (2013). Detailed descriptions of the metamorphosed altered rock types from the eastern limb of the

Stollberg syncline are given in Ripa (1988, 1994, 2012) and Jansson *et al.* (2013), whereas those associated spatially with the Grängsgruvan deposit, on the western side of the syncline, are summarised in Björklund (2011), Raat *et al.* (2013) and Frank *et al.* (2019).

Sampling and analytical methods

Magnetite-bearing samples were collected from drill core from the Grängsgruvan, Tvistbo, Norrgruvan, Cedercreutz, Baklängen and Dammerget deposits, and the Staren limestone, as well as from surface outcrops in BIF associated with the Grönkullan Fe deposit (Fig. 1). Polished thin sections were examined with an Olympus BX-60 dual reflected–transmitted light microscope. Electron microprobe analysis (EMPA) of magnetite was carried out at the University of Minnesota using a JEOL 8900 electron probe microanalyser at an accelerating voltage of 15 kV and a beam current of 20 nA. Trace-element concentrations of magnetite in 37 samples were measured at the U.S. Geological Survey in Denver, Colorado using laser ablation–inductively coupled plasma–mass spectrometry (LA-ICP-MS) with a New Wave Research UP-193 FX LA system (193 nm excimer) coupled to a PerkinElmer DRC-e ICP-MS. Magnetite was ablated at a beam diameter of 30, 65, or 135 μm depending on grain size and the presence or absence of mineral inclusions. Analyses were calibrated using synthetic glass reference standards GSD-1G and NKT-1G from the U.S. Geological Survey. Concentrations were determined using off-line calculations following the protocol of using GeoPro (CETAC Technologies) software (Longerich *et al.*, 1996). Iron (^{57}Fe) was used as the internal standard for concentration calculations. Values of FeO from EMPA were used for internal standardisation. Fifty-one elements were analysed. Al^{27} , Co^{59} , Cr^{52} , Ga^{71} , Mg^{24} , Mn^{55} , Sn^{118} , Ti^{47} , V^{51} and Zn^{66} were generally above detection limits, whereas Ag^{107} , As^{75} , Ba^{137} , Bi^{209} , Ca^{42} , Cd^{111} , Cu^{63} , Ge^{74} , K^{39} , Li^7 , Mo^{98} , Na^{23} , Ni^{60} , P^{31} , Pb^{208} , Rb^{85} , Sb^{121} , Sc^{45} , Si^{28} , Sr^{88} , Th^{232} , U^{238} , W^{184} , Y^{89} , Zr^{90} , and the rare earth elements (REEs) were generally near or below detection limits. Note that polyatomic interferences of Zr are not generally expected for magnetite (see Dare, 2012; 2014). This list includes Si, Ca and the REEs, which are useful for data screening and the identification of analyses that are affected by inclusions, some of which may be submicroscopic in size (e.g. Dare *et al.*, 2014; Nadoll *et al.*, 2014; Pisiak *et al.*, 2015). The detection limits are given in Supplementary table S1. GSD-1G was analysed three times during a 10 hour period to correct for instrument sensitivity drift.

Mineralogy of magnetite-bearing rocks in Stollberg ore field

Magnetite is present as layers or as disseminations in sulfides in the Staren limestone, skarn, banded iron formation, magnetite-rich conglomerate, and quartz–garnet–pyroxene, garnet–biotite, sericite–garnet–magnetite and gedrite–albite altered rocks. Magnetite-bearing assemblages identified here are given in Table 2.

Magnetite in sulfide-bearing rocks

Magnetite is intergrown with sulfides in all base-metal occurrences except for Brusgruvan (Fig. 2a–d). Jansson *et al.* (2013) reported massive magnetite bodies grading into marble at

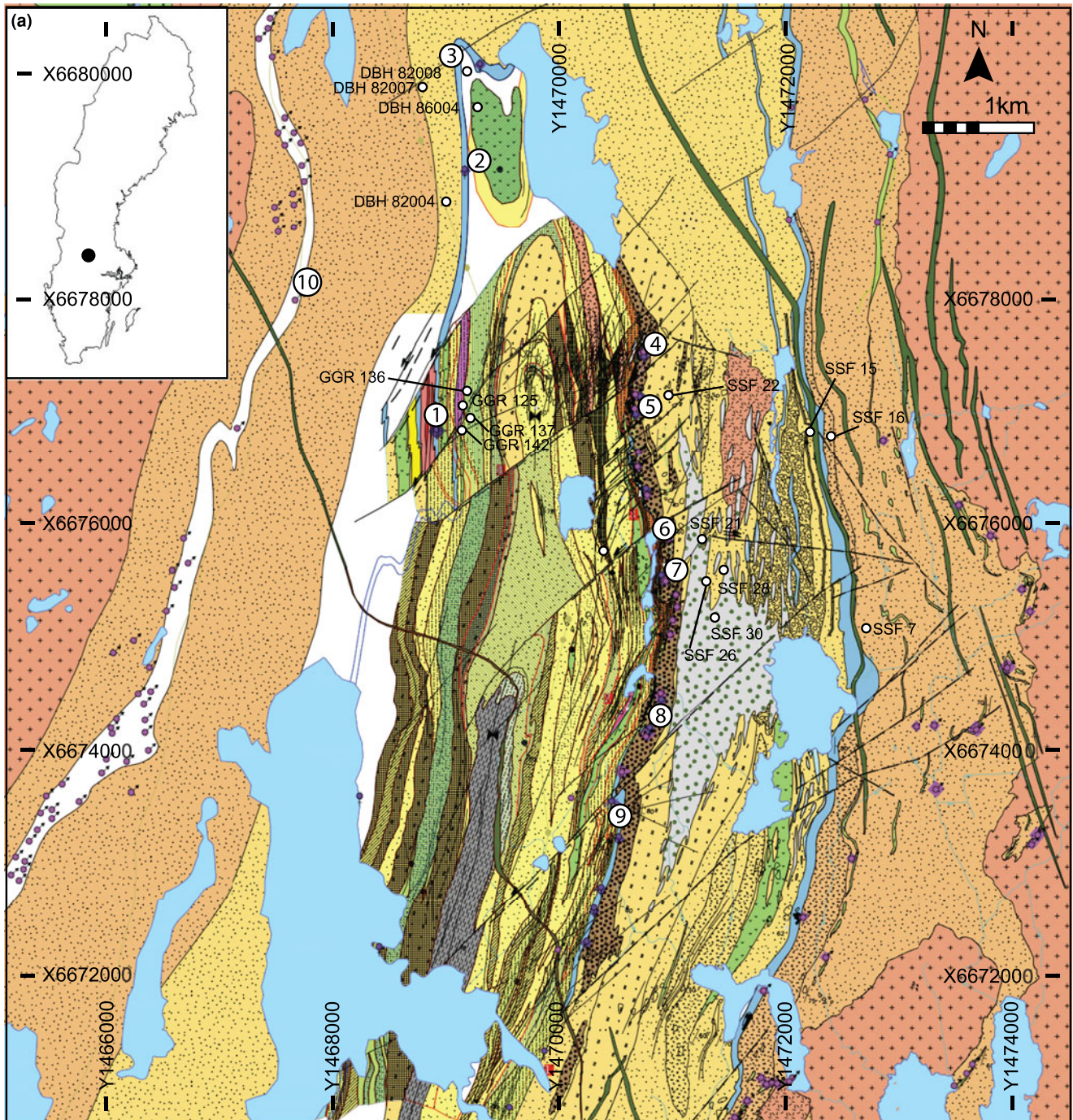


Fig. 1. (a) Geological map of the Stollberg area, showing the location of mines, mineral occurrences, and drill cores. 1 = Gränsgruvan, 2 = Norrgruvan, 3 = Tvistbo, 4 = Lustigkullagruvan, 5 = Cedercreutz, 6 = Baklängan, 7 = Dammerget, 8 = Stollmalmen, 9 = Brusgruvan and 10 = Grönkullan. Drill cores from which samples were taken are shown. Grid is Swedish National Grid RT90, and inset map shows the location of Stollberg in Sweden. (b) Key. Modified after Raat *et al.* (2013).

Stollmalmen, as well as veined magnetite in the marble. At Cedercreutz, massive magnetite contains porphyroblasts of knebelite (manganese variety of fayalite–tephroite series), whereas in the Marnäs deposit semi-massive magnetite is hosted by Al-rich skarn (Jansson *et al.*, 2013). Magnetite at Dammerget occurs primarily as disseminations in sulfide zones containing pyrrhotite and minor chalcopyrite (Fig. 2a, b). Just to the east of this deposit, the banded iron formation consists of layers of magnetite, quartz, garnet, biotite and cummingtonite–grunerite,

with lesser amounts of pyrite and chalcopyrite, and secondary chlorite and sericite in metamorphosed weakly altered rhyolitic ash–siltstone.

At Gränsgruvan, the main zone of sphalerite–galena–pyrite mineralisation lies stratigraphically between quartz–garnet–pyroxene and sericite altered rocks. Below these rocks, within sericite and silica altered rocks, is a pyrite–pyrrhotite zone that contains minor amounts of chalcopyrite and magnetite. Although quartz–garnet–pyroxene rocks at Gränsgruvan consist primarily

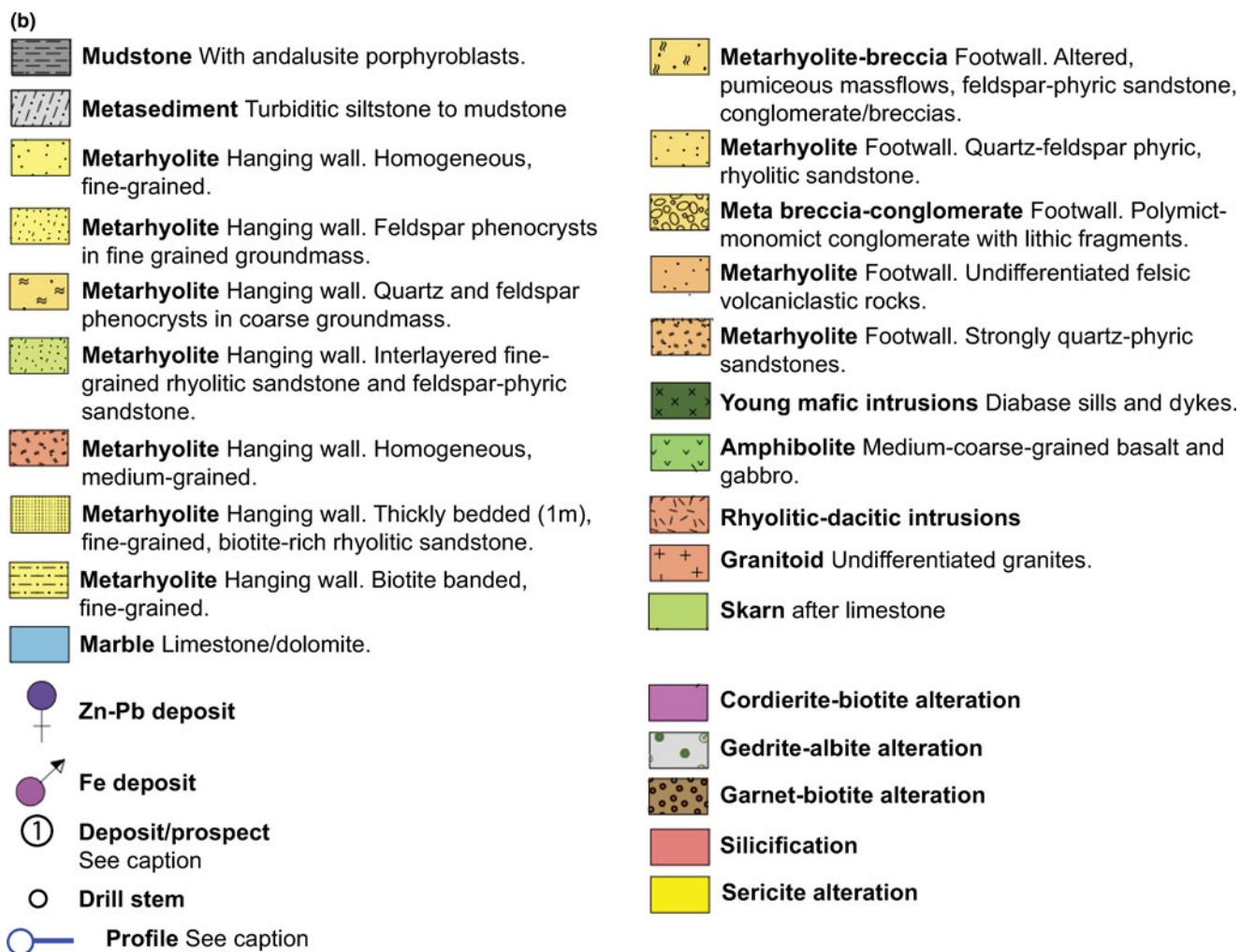


Fig. 1. Continued.

of quartz, garnet and diopside/hedenbergite with lesser amounts of biotite, microcline, clin amphibole and calcite, rare magnetite occurs with garnet and tremolite. Both magnetite and garnet are crosscut by tremolite.

Magnetite in sulfide mineralisation at Grönkullan, 2 km north-west of the Gränsgruvan deposit, is intergrown with galena, sphalerite, and minor pyrite and pyrrhotite. Sulfides are hosted by actinolite, garnet and clinopyroxene. Similar to other deposits, magnetite contains inclusions of sulfides. However, it should be noted here that samples from Grönkullan are skarn-like and from oxide-facies dominant banded iron formation that occurs stratigraphically below the Stollberg ore field.

Magnetite-bearing skarn

Skarn in the Stollberg ore field is a calc-silicate rock composed typically of clinopyroxene, calcite, garnet and clin amphibole, with lesser amounts of biotite, muscovite and microcline, and rare gahnite and fluorite. On the eastern side of the Stollberg syncline, skarn primarily hosts sulfide mineralisation, with altered magnetite-bearing marble present in the hanging wall and footwall of the sulfide zone. At Dammerberget, skarn contains disseminated, anhedral magnetite with calcite, diopside/hedenbergite and muscovite, whereas finer grained skarn at Cedercreutz contains

calcite, phlogopite and chlorite with disseminated grains of magnetite that contain inclusions of these three minerals (Fig. 2e, f). Magnetite-bearing skarn from Baklängen contains disseminated sulfides, primarily sphalerite, pyrrhotite and chalcopryrite, together with calcite, diopside/hedenbergite, magnetite, actinolite, hornblende, chlorite and garnet.

At Gränsgruvan, the skarn analysed occurs in the hanging wall and footwall of the ore zone but is devoid of sulfides. Instead, it contains magnetite as isolated grains in calcite, garnet, diopside/hedenbergite and actinolite. North of Gränsgruvan, a skarn-like rock at Norrgruvan is composed of calcite, quartz, microcline and anthophyllite, with locally abundant sericite, diopside/hedenbergite, garnet, hornblende and disseminated magnetite. Although rare at Stollberg, magnetite at the Norrgruvan prospect is altered locally to hematite (Fig. 2g). The Tvistbo prospect contains sphalerite, galena and magnetite with minor quantities of pyrrhotite, chalcopryrite and pyrite, hosted by skarn consisting of calcite, biotite, garnet, actinolite, diopside/hedenbergite and quartz. Magnetite contains inclusions of calcite.

On the eastern limb of the Stollberg syncline, in the Staren limestone, euhedral magnetite occurs in skarn-altered marble, which consists primarily of calcite, epidote, actinolite, Mg-hedenbergite, chlorite and sericite. In addition, Jansson *et al.* (2013) reported the presence of beds of magnetite-rich

Table 1. General geological characteristics of ore deposits in the Stollberg ore field.

Deposit	Grade and tonnage	Metallic minerals	Gangue minerals	Alteration Types	References
Brusgruvan	Zn 3.2 wt.%, Pb 15.6 wt.%, Ag 320 ppm, Fe <1.23 wt.%, Mn <0.10 wt.%, As 13 ppm	Gn, Sp, Py	Qz, Flr, Mcc, Ms, Bt, Grs, Czo	Pxn skarn, Sil	Ripa (1996), Jansson <i>et al.</i> (2013)
Stollgruvan	Zn 1–4 wt.%, Pb 0.5–1 wt.%, Ag 10 ppm, Fe 20–35 wt.%, Mn 7 wt.%, As <2786 ppm	Sp, Gn, Mag, Apy, Pyh, Py	Qz, Flr, Ms, Cal, Bt, Alm, Sps, Ghn, St, Crd, Hd, Ath, Tr, Ged, Chl, Mn-Ilm, Dsp	Ged-Ab, Grt-Bt, Pxn skarn	Ripa (1996), Jansson <i>et al.</i> (2013)
Dammsberget	Zn 3–5 wt.%, Pb 2–5 wt.%, Ag 20–60 ppm, Fe 10 wt.%	Sp, Gn, Mag, Apy, Pyh, Py	Qz, Flr, Ms, Bt, Cal, Phl, Chl, Hbl, Ath, Gru, Alm, Sps, Kne, Hd, Mn-Ilm, Tlc, Ghn	Ged-Ab, Pxn skarn, Grt-Bt	Ripa (1996)
Baklängen	Zn 2.6 wt.%, Pb 4.8 wt.%, Ag 43 ppm, Fe <26.20 wt.%, Mn <3.49 wt.%, As <450 ppm	Mag, Gn, Sp, Pyh, Apy	Qz, Flr, Ms, Cal, Bt, Alm, Sps, Cam, Kne, Hu-group, Ghn, Grs, Hd, Pxn, Chl, Mn-Ilm, Dsp, Alm	Ged-Ab, Qz-Bt, Grt-Amp-Ghn-Bt, Cam-Grt-Ghn-Flr skarn, Cam-Cpx skarn	Månsson (1979), Jansson <i>et al.</i> (2013)
Korrgruvan-Myggruvan	Zn 0.5 wt.%, Pb 0.5 wt.%, Ag 5 ppm, Fe 35 wt.%, Mn 11 wt.%, As <158 ppm	Mag, Sp, Gn, Pyh, Apy	Qz, Flr, Mcc, Ms, Cal, Bt, Alm, Sps, Cam, Hd, Ath, Chl, Mn-Ilm	Ged-Ab, skarn	Jansson <i>et al.</i> (2013)
Lustigkulla-Marnäs	Zn 1 wt.%, Pb 0.5 wt.%, Ag <2.8 ppm, Fe 30–40 wt.%, Mn 10–15 wt.%, As <3484 ppm	Sp, Gn, Mag, Apy, Lö, Pyh, Py	Qz, Flr, Ms, Cal, Bt, Alm, Sps, Cam, Mn-Gru, Hd, Chl, Mn-Ilm	Bt-Qz, Grt-Amp-Ghn-Bt, Cam-Grt-Kne-Flr, Cam-Cpx	Ripa (1996), Jansson (2011), Jansson <i>et al.</i> (2013)
Grängsgruvan	6.65 Mt, Zn 7.7 wt.%, Pb 2.6 wt.%, Ag 60 ppm	Sp, Gn, Ccp, Py, Pyh	Qz, Bt, Phl, Ms, Ser, Chl, Kfs, Mcc, Pl, Hbl, Act, Alm, Sps, Cal, Cpx, Ep	Crd-Bt, Qz-Grt-Pxn, Ser, Sil, Grt-Bt	Frank <i>et al.</i> (2019)
Tvistbo	575 kton, Zn 3.3 wt.%, Pb 2.6 wt.%, Ag 22 ppm	Sp, Gn, Ccp, Pyh, Mag	Qz, Bt, Phl, Ms, Ser, Chl, Kfs, Alm, Cpx, Cal, Ep	Grt-Bt, Pxn skarn, Qz-Grt-Pxn, Sil, Ser	Frank <i>et al.</i> (2019)
Norrgruvan	No data	Gn, Sp, Py, Pyh, Ccp, Au	Qz, Flr, Bt, Ms, Ser, Chl, Mcc, Pl, Grt, Cam	Pxn skarn, Qtz-Flr, Grt-Bt, Sil	Frank <i>et al.</i> (2019)

Abbreviations: Au – native gold; Ab – albite; Act – actinolite; Alm – almandine; Amp – amphibole; Apy – arsenopyrite; Ath – anthophyllite; Bt – biotite; Cal – calcite; Cam – clin amphibole; Chl – chlorite; Ccp – chalcopyrite; Crd – cordierite; Cpx – clinopyroxene; Czo – clinozoisite; Dsp – diaspore; Ep – epidote; Flr – fluorite; Gn – galena; Ged – gedrite; Ghn – gahnite; Grs – grossular; Grt – garnet; Gru – grunerite; Hbl – hornblende; Hd – hedenbergite; Hu-group – humite group; Kfs – K-feldspar; Kne – knebelite; Lö – löllingite; Mag – magnetite; Mcc – microcline; Mn-Ilm – Mn ilmenite; Mn-Gru – Mn – grunerite; Ms – muscovite; Phl – phlogopite; Pl – plagioclase; Pyh – pyrrhotite; Py – pyrite; Pxn – pyroxenoid; Qz – quartz; Ser – sericite; Sil – Sillimanite; Sp – sphalerite; Sps – spessartine; St – staurolite; Tlc – talc; Tr – tremolite. Most abbreviations from Warr (2021).

Table 2. Magnetite-bearing assemblages in the Stollberg ore field.

Deposit	Sample	Rock Type	Assemblage
Grängsgruvan	GGR 125 476.1	Sulfide mineralisation	Qz-Py-Bt-Pyh-Ms-Kfs-Mag-Chl-Ccp-Sp*
Grängsgruvan	GGR 125 519.0	Sulfide mineralisation	Qz-Chl-Bt-Act-Py-Ep-Mag-Mcc-Sil*-Ccp*-Pyh*
Grängsgruvan	GGR 136 180.3	Amphibolite	Tr-Grt-Mag-Sp-Chl*
Grängsgruvan	GGR 136 189.6	Skarn	Cal-Cpx-Grt-Mag-Tr-Ghn*
Grängsgruvan	GGR 142 408.4	Garnet-biotite rock	Qz-Bt-Grt-Cpx-Mcc-Mag-Py*-Pyh*-Sp*
Norrgruvan	DBH 82004 147.6	Magnetite-bearing quartz vein	Qz-Mag-Ser-Hem-Grt-Chl*-Act*
Norrgruvan	DBH 86004 48.9	Skarn	Cal-Tr-Phl-Mag*
Twistbo	DBH 82007 197.1	Skarn	Cpx-Grt-Tr-Mag-Chl-Pyh-Cal-Sp*-Ccp*-Py*
Twistbo	DBH 82007 198.9	Sulfide mineralisation	Cpx-Gar-Mag-Sp-Gn-Cal-Phl-Tlc-Ser-Pyh*-Cpy*-Py*
Twistbo	DBH 82007 201.0	Sulfide mineralisation	Grt-Phl-Cal-Mag-Cpx-Tlc-Gn-Ser-Tr*-Sp*
Twistbo	DBH 82007 203.5	Skarn	Cpx-Cal-Grt-Mag-Tr-Gn*-Sp*-Pyh*
Cedercreutz	SSF 22 844.8	Skarn	Cal-Tr-Mag-Pyh*-Chl*
Baklängen	SSF 21 416.1	Skarn	Cal-Grt-Chl-Mag-Tr-Pyh-Cam-Ccp*
Baklängen	SSF 21 519.5	Skarn	Cal-Tr-Sp-Mag-Pyh-Cpx-Ccp*
Dammsberget	SSF 26 832.5	Sulfide mineralisation	Hst-Mag-Grt-Phl-Pyh-Ccp*-Sp*-Cal*-Qz*
Dammsberget	SSF 26 841.0	Garnet-biotite rock	Phl-Grt-Mag-Ghn-Pyh-Ccp*-Qz*
Dammsberget	SSF 26 844.2	Sulfide mineralisation	Chl-Pyh-Sp-Ghn-Hst-Apy-Mag-Qz*
Dammsberget	SSF 26 854.3	Sulfide mineralisation	Pyh-Mag-Gru-Act-Grt-Ccp*-Py*-Qz*
Dammsberget	SSF 26 861.0	Amphibolite	Act-Pyh-Mag-Qz-Ccp*
Dammsberget	SSF 26 925.9	Garnet-biotite rock	Grt-Bt-Mag-Qz-Ser-Chl-Pyh*-Flr-Ccp*
Dammsberget	SSF 28 356.1	Gedrite-albite rock	Qz-Pl-Ged-Grt-Bt-Mag*-Py*
Dammsberget	SSF 28 883.4	Skarn	Cal-Cpx-Tr-Sp-Mag-Gn
Dammsberget	SSF 28 889.8	Skarn	Cal-Cpx-Tr-Mag
Dammsberget	SSF 30 553.8	Gedrite-albite rock	Qtz-Ged-Grt-Chl-Mag-Py-Pyh-Ccp*-Sp*
Dammsberget	SSF 30 556.0	Garnet-biotite rock	Qtz-Mag-Grt-Bt-Ged-Chl*-Ccp*-Pyh*
Dammsberget	SSF 30 556.6	Garnet-biotite rock	Qtz-Mag-Bt-Grt-Pyh-Chl-Ccp*
Dammsberget	SSF 30 561.1	Garnet-biotite rock	Mag-Bt-Qz-Grt-Cam-Chl-Py*-Ccp*
Dammsberget	SSF 30 562.2	Garnet-biotite rock	Qtz-Bt-Mag-Ser-Grt-Py
Dammsberget	SSF 30 564.3	Garnet-biotite rock	Qtz-Mag-Hst-Ser-Phl-Py*-Pyh*
Staren	SSF 15 354.0	Quartz-magnetite rock	Qtz-Mag-Bt-Kfs-Ep
Staren	SSF 16 137.7	Skarn	Cal-Chl-Mag*-Ccp*
Staren	SSF 16 195.9	Skarn	Cal-Tr-Mag*-Ser*-Ccp*
Staren	SSF 7 386.8	Skarn	Cal-Act-Mag-Pyh*-Ccp*
Staren	SSF 7 47.0	Skarn	Cal-Tr-Mag-Pyh*-Ccp*
Staren	SSF 7 85.8	Skarn	Cal-Mag-Act-Ccp*-Cpx*
Grönkullan BIF	Stol 13	Skarn	Cal-Tr-Pyh-Mag*-Cpx*
Grönkullan BIF	Stol 14b	Sulfide mineralization	Act-Grt-Cpx-Mag-Gn-Sp-Py*-Pyh*-Chl*

Notes: Minerals are listed in approximate order of abundance; * indicates <1% present; Most mineral abbreviations from Warr (2021); BIF = banded iron formation.

mafic conglomerate in the Staren limestone, which also contain actinolite-, epidote- and magnetite-bearing skarn, with clasts of bimodal mafic and felsic volcanic material. A sample collected here from the Staren limestone (SSF15 354.0) contains fine-grained magnetite, microcline and biotite with minor epidote. Skarn in banded iron formation at Grönkullan contains calcite and clusters of tremolite with minor pyrrhotite, magnetite, diopside/hedenbergite and chalcopyrite.

Other magnetite-bearing rocks

Garnet-biotite rocks are almost as widespread as skarn, and are also associated typically with magnetite-bearing sulfides. At Dammsberget, these rocks contains garnet, biotite, quartz, fibrolite and muscovite, with lesser amounts of fluorite, grunerite, gahnite, pyrrhotite, chalcopyrite and subhedral to anhedral magnetite. Magnetite-rich lenses composed of magnetite, quartz, and garnet also occur on the eastern limb of the Stollberg syncline (Fig. 2h).

Gedrite-albite rocks occur on the eastern limb of the Stollberg syncline in a zone that stretches from just north of the Cedercreutz deposit to more than 1 km south of the Brusgruvan deposit. They occur stratigraphically beneath these deposits and are composed of fine-grained quartz, acicular to radiating ferro-

gedrite, and plagioclase, with rare cordierite, epidote and fine-grained euhedral to subhedral magnetite.

Magnetite is also present in mafic sills either as small masses or as disseminations. A mafic sill at Dammsberget consists primarily of ferro-actinolite, with disseminated magnetite, pyrrhotite and chalcopyrite, and minor plagioclase.

Magnetite compositions

Trace-elements

Magnetite compositions ($n = 442$) were obtained from 37 samples of various types of metamorphosed altered rock. More compositional data were collected from magnetite in skarn (274 analyses from 15 samples) than from any other rock type due to the common association between magnetite, sulfides and skarn. Samples from altered Staren limestone were collected from two drill holes (SSF16 and SSF7), ~2 km apart, which are not associated spatially with sulfide mineralisation. The median concentrations of the ten most abundant trace elements (Al, Co, Cr, Ga, Mg, Mn, Sn, Ti, V and Zn) are shown for each sample in Table 3 and are discussed here. All data are given in Supplementary table S2. Element-element bivariate plots are shown for Mg vs. Al, V vs. Mn, Al vs. Ti, and Zn vs. Ga for magnetite in skarn (Fig. 3a-d) and for Al vs. Ti, V vs. Mn, and Zn vs. Ga for magnetite in sulfide mineralisation and garnet-biotite rocks (Fig. 4a-f).

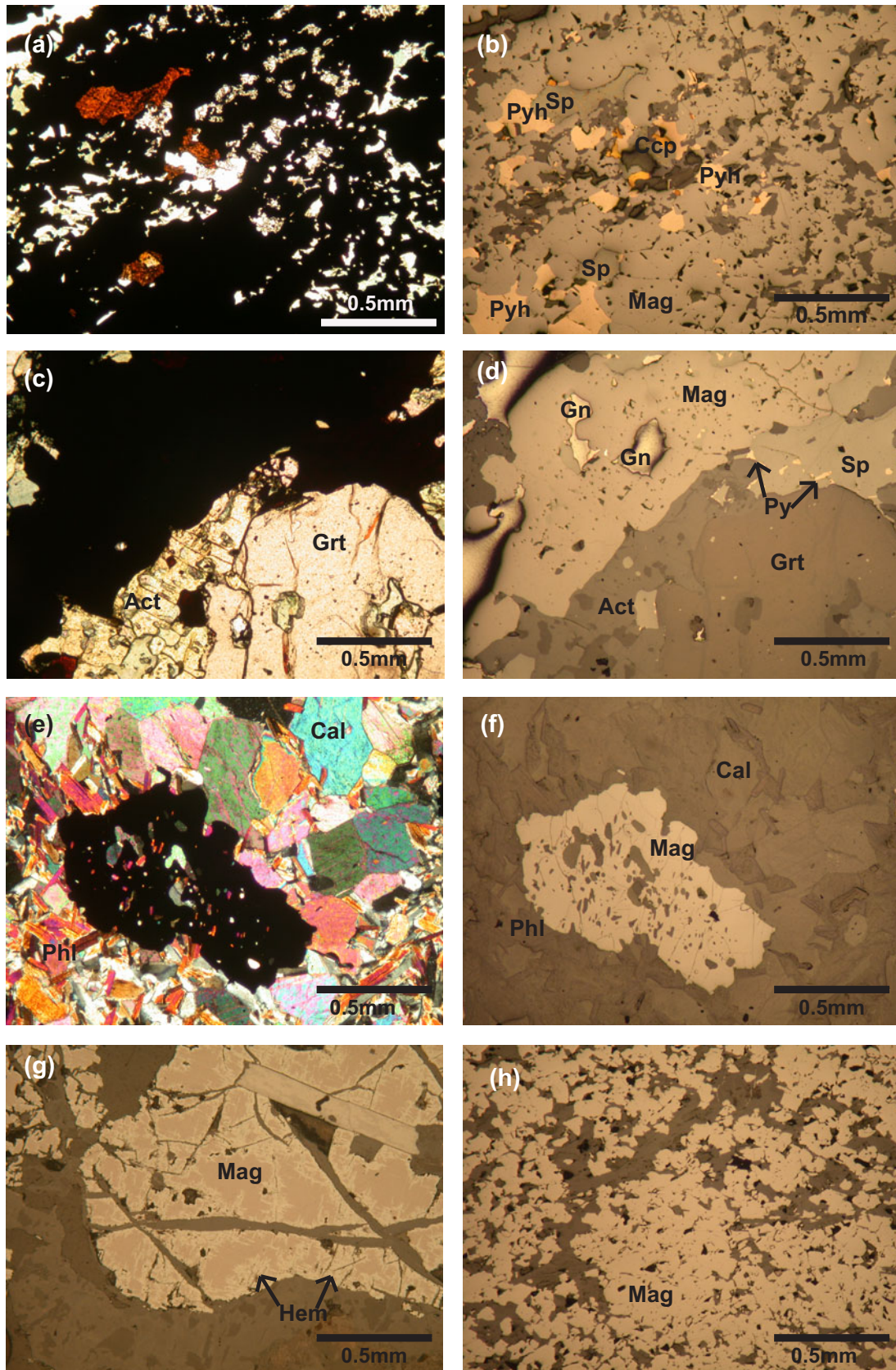


Fig. 2. Photomicrographs of magnetite in thin section from altered rocks in the Stollberg syncline. (a) Magnetite (Mag) with inclusions of pyrrhotite (Pyh), sphalerite (Sp) and chalcopyrite (Ccp) in massive sulfide mineralisation (Dammerget, sample SSF26 832.5), in transmitted light and (b) reflected light. Quartz is the colourless mineral in (a). (c) Magnetite with inclusions of galena (Gn) and sphalerite in massive sulfide mineralisation, with garnet (Grt) and actinolite (Act) (Grönkullan, sample 14b), in transmitted light and (d), reflected light. (e) Isolated grains of magnetite with inclusions of calcite (Cal) and phlogopite (Phl) in weakly altered skarn (Cedercreutz, sample SSF22 844.8), in transmitted light, and (f) reflected light. (g) Magnetite (Mag) in sericite-garnet-magnetite rock exhibiting alteration to hematite (Hem) along mineral grain margins and fractures (Norrgruvan, sample DBH 82004 147.6) in reflected light. (h) Massive magnetite in garnet-biotite rock (Dammerget, sample SSF 30 561.1), reflected light.

Table 3. Median trace compositions of selected elements in magnetite from the Stollberg district (ppm).

Sample	Deposit and rock type*	N	Mg	Al	Ti	V	Cr	Mn	Co	Zn	Ga	Sn
GGR 125 476.1	Grängsgruvan ¹	7	79.1	1159	90.6	5.71	17.1	684.0	0.95	198.7	27.0	1.26
GGR 125 519.0	Grängsgruvan ¹	16	148.9	489.6	213.5	6.55	9.51	929.7	1.01	160.1	32.9	0.78
GGR 136 180.3	Grängsgruvan ²	6	309.8	1557	9337	23.8	8.71	6680	6.76	3368	67.8	14.2
GGR 136 189.6	Grängsgruvan ³	12	990.5	4546	5943	35.0	5.15	9914	3.99	1936	138.5	19.3
GGR 142 408.4	Grängsgruvan ⁴	14	39.6	612.5	45.2	2.82	8.30	879.1	0.59	80.3	19.1	0.93
DBH 82004 147.6	Norrgruvan ⁵	16	13.6	1623	637.5	241.6	10.2	556.3	6.58	733.1	175.4	6.09
DBH 86004 48.9	Norrgruvan ³	18	80.2	274.2	428.5	52.3	7.12	1631	2.03	131.9	18.6	4.18
DBH 82007 197.1	Tvistbo ³	13	74.8	620	2503	5.09	8.77	4252	1.28	258.4	77.2	52.6
DBH 82007 198.9	Tvistbo ¹	12	353.8	3240	1557	2.22	2.19	10269	0.39	359.0	54.4	22.0
DBH 82007 201.0	Tvistbo ¹	19	302.3	1176	2839	4.32	12.6	8304	1.46	436.8	37.6	30.6
DBH 82007 203.5	Tvistbo ³	9	241.9	963	4137	20.3	14.6	9892	1.96	1093	38.4	20.9
SSF 22 844.8	Cederkreutz ³	9	176.1	1231	1651	288.7	16.1	2876	2.78	222.4	33.6	56.6
SSF 21 416.1	Baklängen ³	13	309.1	6305	2620	151.7	14.0	851	1.00	91.7	75.0	24.7
SSF 21 519.5	Baklängen ³	12	678.6	978	1347	96.9	11.6	5040	0.67	528.5	30.2	36.8
SSF 26 832.5	Dammsberget ¹	11	237.1	2857	1524	13.9	3.05	1317	0.21	339.2	38.8	57.6
SSF 26 841.0	Dammsberget ⁴	11	44.5	3282	2444	36.9	9.6	153.4	0.67	1040	134.0	73.4
SSF 26 844.2	Dammsberget ¹	10	130.8	3734	2564	88.8	2.60	255.4	0.60	1027	185.1	61.9
SSF 26 854.3	Dammsberget ¹	12	36.9	2112	1143	25.0	4.04	183.2	0.94	253.0	25.7	53.8
SSF 26 861.0	Dammsberget ²	12	30.0	1085	838.2	56.3	21.9	421.9	1.01	441.1	52.8	19.7
SSF 26 925.9	Dammsberget ⁴	13	48.2	3621	1708	21.2	3.11	56.5	1.73	484.3	560.8	21.8
SSF 28 356.1	Dammsberget ⁶	12	29.5	4263	2090	100.0	17.9	104.5	11.5	159.3	372.5	15.9
SSF 28 883.4	Dammsberget ³	1	794.2	2696	4286	241.1	-	7301	0.74	2195	225.1	95.6
SSF 28 889.8	Dammsberget ³	12	548.8	1513	285.8	49.6	11.5	6132	0.65	3270	20.8	89.4
SSF 30 553.8	Dammsberget ⁶	13	82.6	3860	1308	9.19	8.46	173.9	0.96	128.5	50.6	35.7
SSF 30 556.0	Dammsberget ⁴	15	60.4	3158	1081	7.83	7.61	443.0	0.82	139.5	22.9	25.4
SSF 30 556.6	Dammsberget ⁴	15	87.8	2910	743.0	28.7	15.5	457.2	1.17	214.6	27.8	34.4
SSF 30 561.1	Dammsberget ⁴	12	156.6	2635	268.0	13.1	2.95	677.6	1.41	116.1	22.6	2.84
SSF 30 562.2	Dammsberget ⁴	14	57.5	1858	829.3	27.1	9.29	456.9	1.39	240.1	47.8	2.16
SSF 30 564.3	Dammsberget ⁴	17	45.3	2500	594.3	24.6	15.2	706.2	1.67	188.8	19.2	9.13
SSF 15 354.6	Staren ⁷	9	28.7	129.7	76.6	1550	65.9	405.8	28.4	38.1	5.80	1.98
SSF 16 137.7	Staren ³	12	56.2	771.7	497.9	156.2	9.6	1336	4.07	233.8	5.40	1.61
SSF 16 195.9	Staren ³	11	123.5	514.4	254.2	290.0	25.6	409.5	15.4	336.6	25.0	1.47
SSF 7 386.8	Staren ³	11	219.6	444.9	194.1	20.1	12.5	400.0	5.87	61.7	23.0	11.8
SSF 7 47.0	Staren ³	15	253.9	652.5	608.7	138.7	11.1	611.7	4.09	212.9	28.4	1.44
SSF 7 85.8	Staren ³	12	120.9	2739	328.5	13.7	9.37	403.7	5.20	50.0	22.0	4.61
Stol 13	Grönkullan ³	7	435.8	66.8	83.3	160.3	6.30	3686	0.48	824.9	9.69	74.3
Stol 14b	Grönkullan ²	12	160.0	1843	2368	25.4	3.40	2560	0.46	601.9	36.3	85.8

Notes: N = number of analyses, Co contents <1 are rounded to 2 decimal places

*Key to deposit and rock type: 1 sulfide mineralization; 2 amphibolite; 3 skarn; 4 garnet-biotite altered rock; 5 magnetite-bearing quartz vein; 6 gedrite albite rock; 7 quartz-magnetite rock

These elements were chosen because most data were above the limits of detection.

Skarn is host to ore along the eastern side of the syncline and at Tvistbo, where magnetite is a common accessory mineral. When compared to the trace-element compositions of magnetite from other metamorphosed altered rock types in the Stollberg ore field, magnetite in skarn generally has among the highest Mg (up to 1867 ppm; Grängsgruvan), Mn (up to 19,423 ppm; Grängsgruvan), Ni (up to 142 ppm, Norrgruvan) and Cd (up to 792 ppm, Norrgruvan) contents (See Supplementary table S2). Clockwise around the Stollberg syncline, starting from Norrgruvan in the north to deposits on the eastern limb of the Stollberg syncline (i.e. Norrgruvan, Tvistbo, Cedercreutz and then Dammsberget), magnetite in skarn generally shows a steady increase in the median content of Mg (Fig. 3a). For the same sequence of deposits (excluding Dammsberget), the concentration of Al also increases for magnetite in skarn, with the highest values occurring in the Baklängen deposit, which then decreases at Dammsberget (Fig. 3a, c). Elevated concentrations of Sn in magnetite occur in sulfide-bearing skarn from Tvistbo (up to 68 ppm), Cedercreutz (up to 83 ppm), Baklängen (up to 52 ppm) and Dammsberget (up to 101 ppm). Magnetite contains elevated values of Ga in skarn from the Grängsgruvan (up to 178 ppm), Tvistbo (93 ppm) and Baklängen (98 ppm) deposits,

with one anomalous value from Dammsberget of 225 ppm Ga (Fig. 3d), whereas Zn is enriched in magnetite from the Grängsgruvan (up to 3963 ppm) and Dammsberget (up to 4049 ppm) (Fig. 3d). The composition of magnetite in skarn at Grängsgruvan, which is not associated spatially with sulfides, has the highest concentrations of Mg (1868 ppm) and Ga (178 ppm), as well as elevated levels of Al (up to 6274 ppm) (Fig. 3a, c, d). On average, magnetite in marble/skarn from the Staren limestone contains the highest concentration of Co (up to 22 ppm) and among the lowest concentrations of Mn compared to magnetite from other marbles/skarns (Fig. 3b).

Magnetite in sulfide mineralisation from the Dammsberget, Grängsgruvan, Tvistbo and Grönkullan deposits is generally enriched in Mn (particularly at Tvistbo, ~9000 ppm), Sb (up to 9.79 ppm, 13.5 ppm and 7.54 ppm at Tvistbo, Dammsberget and Grönkullan, respectively), and Mo (up to 9.63 ppm, Tvistbo), relative to magnetite in all other rock types, except for skarn which can also contain elevated concentrations of Mo (up to 8.97 ppm). Where elevated in Mo, samples of magnetite are also generally enriched in Zn (up to 6236 ppm, Dammsberget). Relative to magnetite compositions in sulfide mineralisation from other deposits in the Stollberg district, those from Grängsgruvan are depleted in Ti (Fig. 4a) and Zn (Fig. 4b). Similarly to those from Tvistbo, magnetite compositions are depleted in V compared to those from Grönkullan and Dammsberget (Fig. 4c).

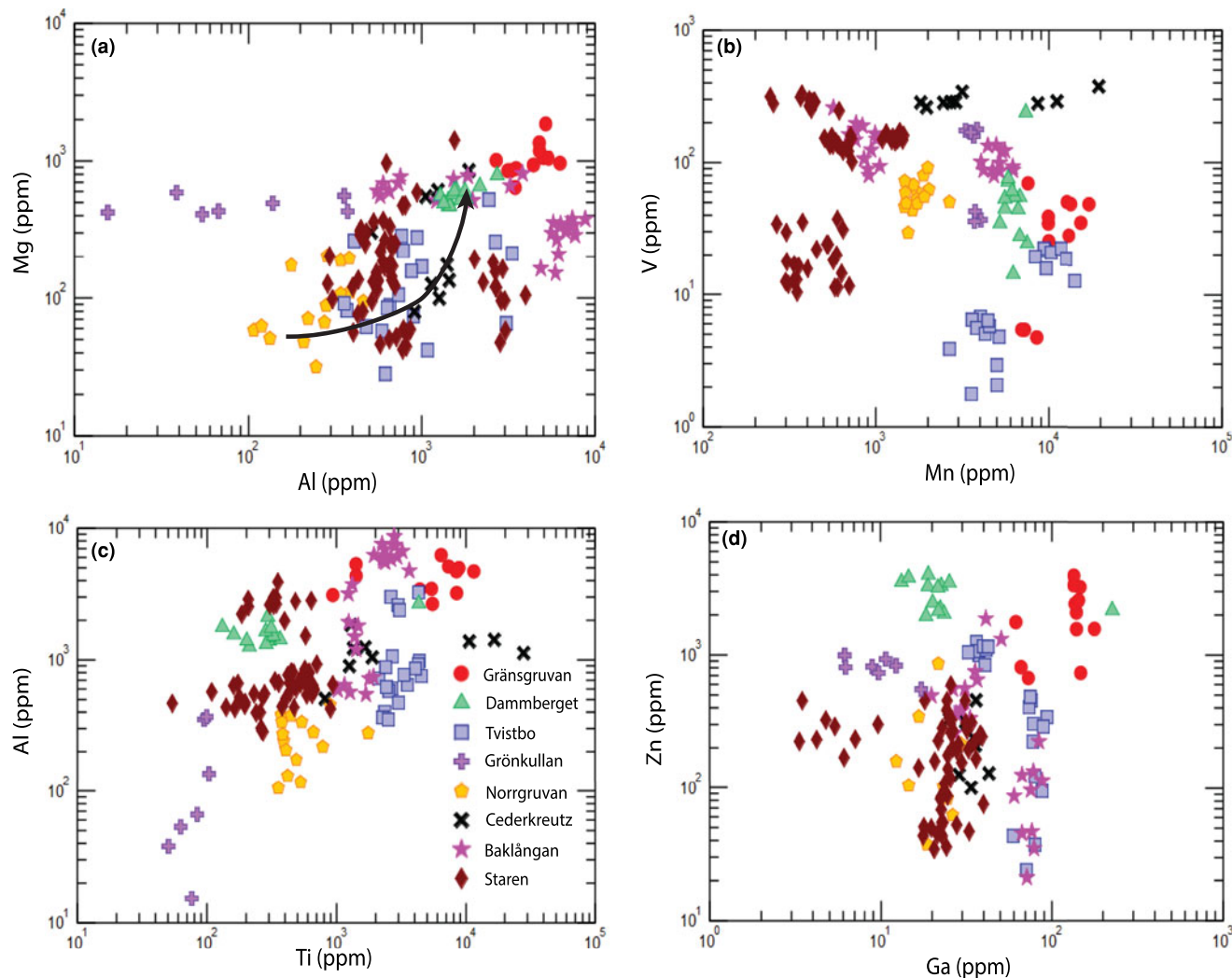


Fig. 3. Element–element plots of magnetite composition (ppm) in skarn from the Stollberg ore district. (a) Mg vs. Al; (b) V vs. Mn; (c) Al vs. Ti; and (d) Zn vs. Ga. The arrow in Fig. 3a shows the increase in Al and Mg content going from Norrgruvan through to Tvistbo, Cederkreutz and Dammerget.

Those from the last deposit are generally more enriched in Ga and depleted in Mn (Fig. 4b, c).

Magnetite in garnet–biotite rock from Grängsruvan and Dammerget is generally depleted in trace elements relative to other metamorphosed altered rocks, particularly in sample GGR142 408.4 from the Grängsruvan deposit. Magnetite in samples from the Dammerget deposit is relatively depleted in Ti, V, Mn, Co, Zn, Ni, Cu, Mo, Cd and Pb, except for magnetite in sample SSF26 841.0, which occurs in close proximity to sulfide mineralisation and is enriched in Al, Ti, Zn, Ga and Sn. Compared to magnetite in garnet–biotite rocks from Grängsruvan those from Dammerget are enriched in Al, Ti, Zn, Ga and V, though more depleted in Mn (Fig. 4d–f).

Quartz–garnet–pyroxene rock (GGR136 180.3) from the Grängsruvan deposit contains magnetite with elevated concentrations of Ti, Mn, Zn, Co and W, and low amounts of Cr, Na, Cu, Sb and Ba. Magnetite in a sample of matrix-supported mafic polymict conglomerate (SSF15 354.6) associated spatially with the Staren limestone contains the highest V (1550 ppm), Co (28.4 ppm) and Cr (65.9 ppm), and the lowest Zn (38.1 ppm) of any rock analysed here. This magnetite is also among the poorest in terms of Mg (29.7 ppm), Al (129.7 ppm), Ga (5.80 ppm) and Sn (1.98 ppm).

Magnetite in a sericitically altered garnet–magnetite rock from Norrgruvan (DBH 82004 147.6) contains the lowest Mg content (13.6 ppm) and among the highest concentrations of V (241.6 ppm), Ga (175.4 ppm) and Co (6.58 ppm). Magnetite in amphibolite (SSF26 861.0) near the ore zone at Dammerget contains low amounts of trace elements but the highest concentration of Ge (27.2 ppm) and is enriched in Ga and Cr, though depleted in Mg, Al, Co, Li, Na, Ca, Ni, Cu, Zr, Cd and Pb.

Magnetite in a gedrite–biotite–albite rock (sample SSF28 356.1) in the footwall of the Dammerget deposit contains among the highest median values of Al (4263 ppm), Co (11.5 ppm) and Ga (372.5 ppm) and lowest Mg (29.5 ppm) and Mn (104.5 ppm), whereas sample SSF30 553.8 contains similar median concentrations of Al (3860 ppm) and Mn (173.9 ppm), but lower Ga (50.6 ppm) and Co (1.35 ppm) contents.

Principal component analysis

Although the element–element variation diagrams shown in Figs 3 and 4 have been used to demonstrate which two elements are elevated or depleted in a given rock type within and between the various ore deposits, they do not show trends for multiple

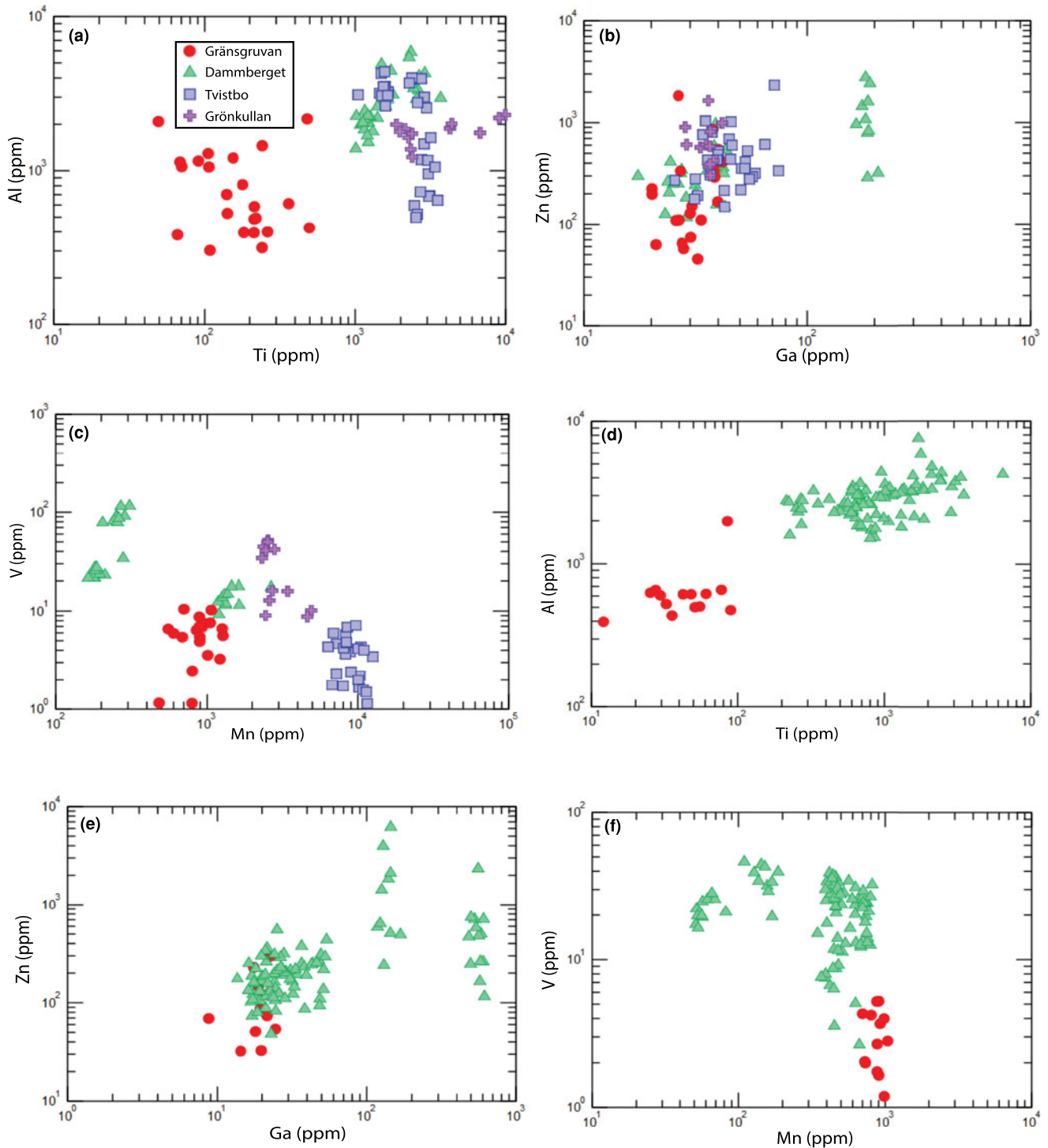


Fig. 4. Element–element plots of magnetite composition (ppm) in sulfide mineralization from the Stollberg ore district. (a) Al vs. Ti; (b) Zn vs. Ga; and (c) V vs. Mn; and in garnet–biotite rocks (d) Al vs. Ti; (e) Zn vs. Ga; and (f) V vs. Mn.

elements. Multivariate statistical methods, including, for example, principal component analysis (PCA) and Random Forests, can be used instead to identify underlying patterns in datasets with many variables (e.g. Jolliffe and Cadima, 2016). A PCA allows for the reduction of multivariate data sets into a two-dimensional representation of variance, which enables factors that contribute the greatest variability within the data set to be identified. Here we apply a PCA to various magnetite-bearing rocks in the Stollberg ore field.

Makvandi *et al.* (2016b) suggested that the most important discriminator elements in magnetite from VMS deposits are Al, Ca, Co, Cr, Ga, Mn, Mg, Ni, Si, Ti, Zn and Zr. We have focused principally on these elements here. Data gathered in the present study via LA-ICP-MS were pre-treated to avoid problems associated with the constant-sum constraint and to correct for heteroscedasticity using a centred log-ratio transformation in *CoDaPack* version 2.01.8 (Comas-Cufí and Thió-Henestrosa,

2011). Pre-treated data were then subjected to PCA in *JMP Pro* version 11.0.

Magnetite analysed here typically contain concentrations of Al, Co, Cr, Ga, Mg, Mn, Sn, Ti, V and Zn above detection limits, with Ca, Si and Zr generally below detection limits. For the present study, we included censored geochemical data, which contains values below detection limits for some elements. The censored data were substituted with values equal to the detection limit/ $\sqrt{2}$ for a given element, which according to Croghan and Egeghy (2003) and Verbovsek (2011) produces a mean value of the population that is close to the statistical mean of the censored data. On the basis of the statistical studies of Hron *et al.* (2010), populations with up to 40% censored data can be evaluated. We conducted a PCA on the ten most abundant elements listed above for 435 of 442 LA-ICP-MS analyses obtained here (Fig. 5a). The seven compositions that were excluded from the PCA were of mineral inclusions in magnetite.

Principal component 1 accounts for 33.7% of variance with V, Cr and Co, correlating negatively with Al, Ti, Zn, Ga and Sn (Fig. 5b). Principal component 2 represents 25.3% of variance and shows that Mn and Mg correlate negatively with Al, Co, Ga, Ti and V. Principal components 3 to 9 range from 10.6% to 1.6% (Fig. 5c). Component 1 and 2 scores for each element are shown in Figs 5c and d, respectively. Magnetite from Tvistbo and Grängsgruvan have similar scores, regardless of rock type, whereas magnetite scores from the Dammerberget and Norrgruvan deposits are more scattered. The composition of magnetite in garnet–biotite and gedrite–albite rocks from Dammerberget, which have high component 2 scores, is dictated by the elevated concentrations of Ga and Al. Magnetite in magnetite-rich conglomerate from Staren has component 1 scores <0 and component 2 scores near 0 that can be distinguished from magnetite from other locations.

Because of the high number of magnetite compositions obtained here, scores from various rock types in Fig. 5a overlap, so the data have been separated to distinguish the data more clearly in Figs 6a–c. Figure 6 shows PCAs for magnetite compositions in garnet–biotite rocks from Dammerberget and Grängsgruvan (Fig. 6a, $N=107$), sulfide mineralisation from Grängsgruvan, Tvistbo, Dammerberget and Grönkullan (Fig. 6b, $N=95$), and skarn from the Staren limestone and the Grängsgruvan, Norrgruvan, Tvistbo, Baklångan, Cedercreutz and Dammerberget deposits (Fig. 6c, $N=274$). Magnetite in garnet–biotite rocks from Dammerberget cluster near zero for components 1 and 2 with some data showing positive values. Although some data from Dammerberget overlap with those from Grängsgruvan, magnetite in garnet–biotite rocks in the latter have component 2 scores between –1 and 0 (Fig. 6a). It should be pointed out that magnetite in these rocks from Dammerberget contains higher amounts of Al, Ti, Ga and V as shown in Fig. 4d–f, relative to those in garnet–biotite rocks from Grängsgruvan. This is reflected in the PCA (Fig. 6a) where Ga, Al and Ti show positive values of components 1 and 2.

Principal component analysis scores for magnetite in sulfide mineralisation from Tvistbo, Dammerberget, Grängsgruvan and Grönkullan are somewhat clustered with magnetite from Grängsgruvan being characterised by component 1 and 2 showing scores <0, whereas component 1 and 2 scores for magnetite from Dammerberget are primarily >0 and controlled principally by the presence of Al, Ga, Sn and Ti. Principal component analysis scores for magnetite in sulfides from Tvistbo, Grönkullan, and some data from Dammerberget exhibit component 1 scores >1

and component 2 scores mostly between –1 and 0 (Fig. 6b). Deposits in the eastern side of the Stollberg syncline are associated spatially with a prominent gedrite–albite alteration, which is not observed at Grängsgruvan on the western side of the syncline. The differences in the physicochemical conditions of the ore-forming fluid undoubtedly accounts for the differences in PCA scores in Fig. 6b and the element–element plots in Figs 4a–c.

Magnetite in skarn from the Staren limestone and Norrgruvan, which shows no spatial association to sulfides, exhibit component 1 scores that overlap and are <0, whereas magnetite in skarn from Baklångan, Dammerberget, Tvistbo and Cedercreutz, that are associated with sulfides, as well as Grängsgruvan, which is also not associated with sulfides, plot near the centre of the score plot and generally have component 1 scores between 0 and 2 (Fig. 6c). Although there is an overlap in the component 1 and 2 scores of magnetite among the various deposits, most deposits are clustered and show little compositional variation, a feature that is also shown in the element–element plots in Fig. 3a–d.

Discussion

Origin of magnetite

Ripa (2012) and Jansson *et al.* (2013) proposed that sulfides in the Stollberg ore field formed as a result of sub-seafloor replacement of limestone to form stratabound replacement mineralisation. On the basis of the sulfide assemblage and positive Eu anomalies (Eu/Eu* up to 2.35) in the magnetite-bearing Stollberg limestone and skarn at the transition with proximal footwall altered rocks, together with the sulfide mineralisation (Jansson *et al.*, 2013), vent temperatures of >250°C were likely, with most of the ore-forming components being derived from felsic volcanoclastic rocks prior to metamorphism. The spatial association between magnetite and sulfides in massive sulfides and in metamorphosed altered rocks (e.g. garnet–biotite and gedrite–albite rocks) suggests that at least one generation of magnetite formed at the same time as the sulfides, and was subsequently recrystallised and metamorphosed (Fig. 2a–d). Further support for the pre-metamorphic origin of magnetite is provided by carbon and oxygen isotope studies of calcite in limestone at Stollberg (Billström *et al.*, 1985) and petrography of limestones and dolomites in the Bergslagen district, which suggested to Allen *et al.* (2003) and Jansson *et al.* (2013) that many of the carbonate horizons are marine stromatolitic limestone altered locally by hydrothermal fluids. Part of the pre-metamorphic alteration process added magnetite to both the Staren and Stollberg limestones and developed skarn-hosted sulfide mineralisation in the latter. Although, magnetite in skarn rocks is generally devoid of inclusions of sulfides (Fig. 2e, f). A second generation of magnetite was recognised by Ferrow and Ripa (1991) who proposed that some magnetite formed during retrograde metamorphism as gedrite was altered to chlorite and quartz. Such magnetite was not observed in the present study.

Conditions of formation of magnetite

Nadoll *et al.* (2014) and Makvandi *et al.* (2016b) proposed that the most important factors controlling the trace-element compositions of magnetite derived from “low-temperature hydrothermal fluids” (i.e. < 500°C) include f_{O_2} , f_{S_2} , temperature, pressure, cooling rate, silica activity, host rock and fluid phase compositions, and the degree of fluid–rock interaction. However, it should be

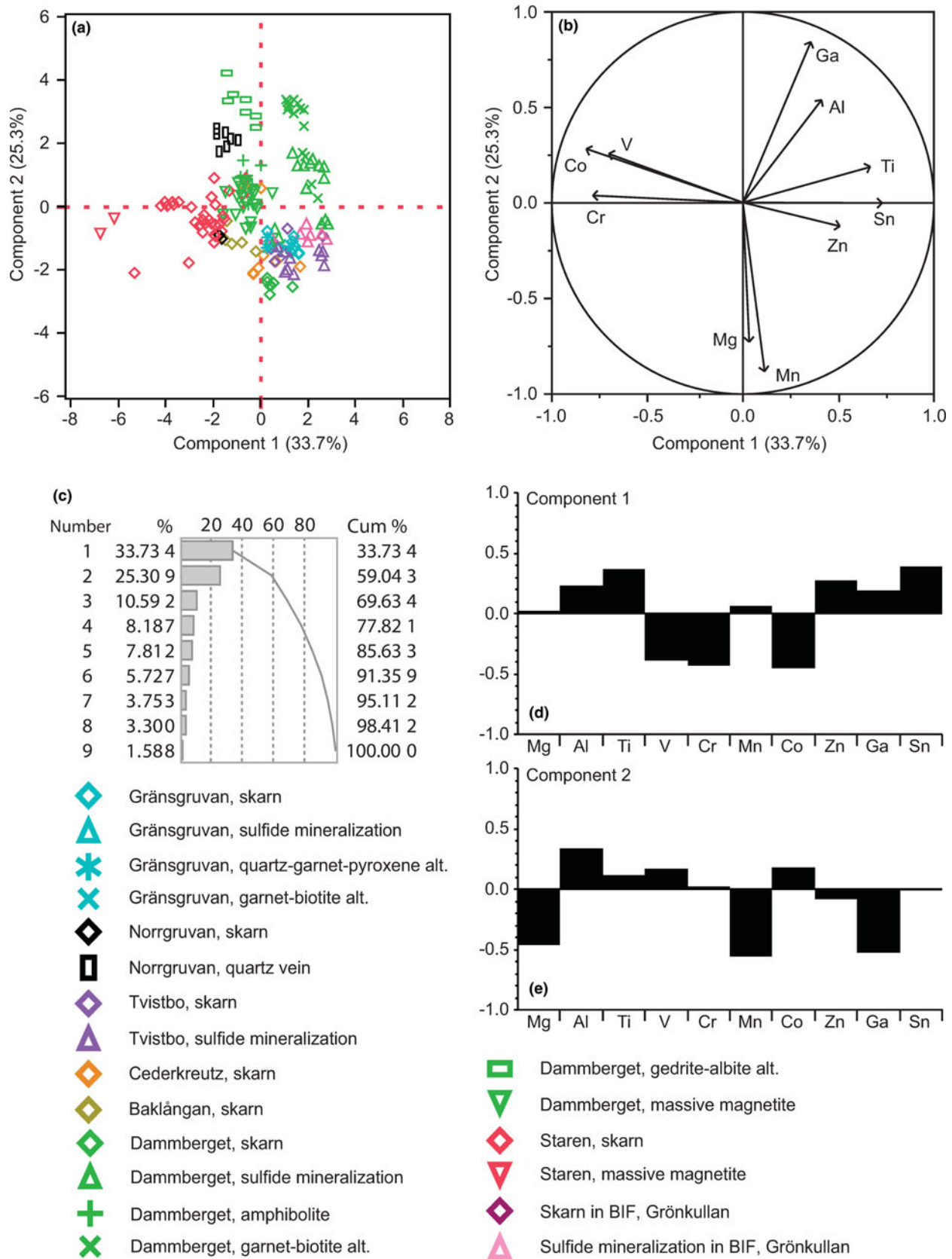
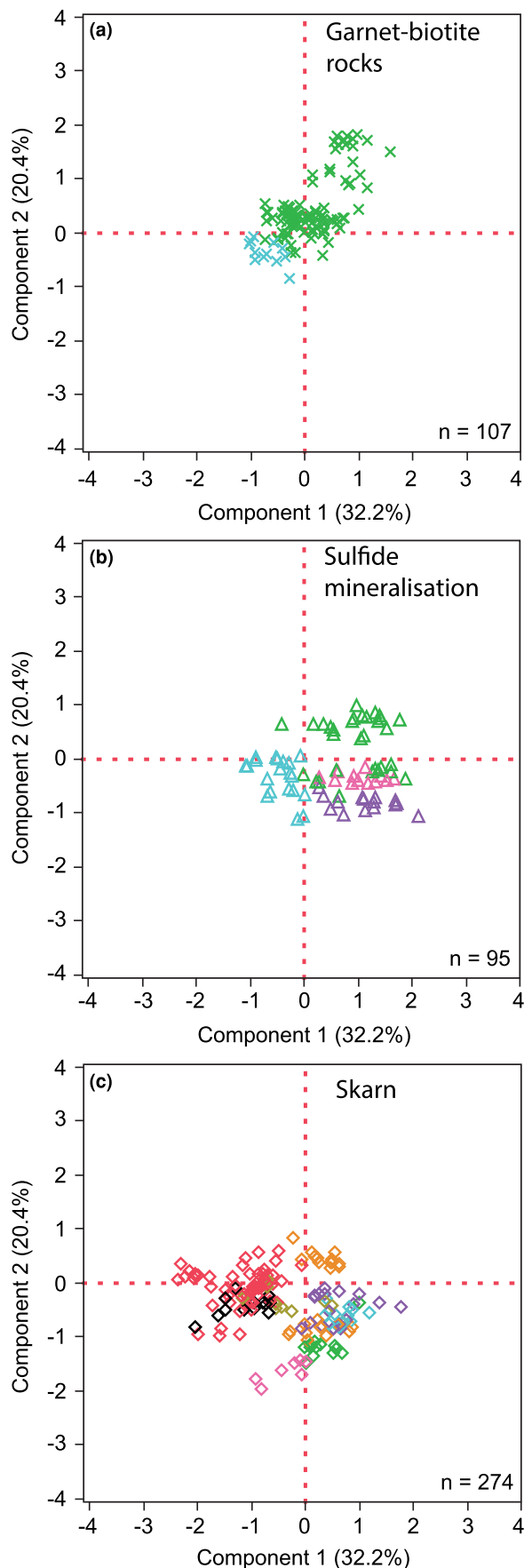


Fig. 5. Principal component analysis for 10 elements (Al, Co, Cr, Ga, Mg, Mn, Sn, Ti, V and Zn) in magnetite ($n = 435$) for all rocks studied here from the Stollberg syncline. (a) Score plot of the first two principal components, with the percentage of variance for each component noted in parentheses. (b) Loading plot showing the geometric representation of how data were projected onto the score plot with respect to each element. (c) Percentages of the first nine principal components. (d) Score contribution plot for principal component 1. (e) Score contribution plot for principal component 2.



noted that studies of magnetite in sulfide deposits derived from lower temperature hydrothermal fluids that were metamorphosed subsequently are few in number (e.g. Singoyi *et al.*, 2006; Kamvong *et al.*, 2007; Dupuis and Beaudoin, 2011; Makvandi *et al.*, 2013; 2016b), and the relative importance of these factors is still poorly understood. On the basis of their studies of magnetite in the metamorphosed Izok Lake bimodal-felsic Zn–Pb–Cu–Ag and the Halfmile Lake felsic siliclastic Zn–Pb–Cu VMS deposits, Makvandi *et al.* (2016b) suggested that the composition of metamorphic magnetite was essentially dependent on the bedrock composition, the grade of metamorphism, and oxygen fugacity.

High field strength and other immobile elements (Ti, Al, Zr, Hf, Nb, Ta and Sc) are commonly depleted in hydrothermal magnetite, as they are not present in, or transported by, hydrothermal fluid (Dare *et al.*, 2014). At Stollberg, Al and Zr are generally depleted, although Ta and Nb were not analysed for in the present study. Silicon and Ca are also locally enriched in hydrothermal magnetite, although both elements were mostly below detection limits in samples from the Stollberg ore field. In contrast, elements such as Mg, Mn and Zn that are transported readily in hydrothermal fluids can be enriched, especially compared to igneous magnetite (e.g. Nadoll *et al.*, 2014). Manganese and Zn are enriched in magnetite from Stollberg, whereas Mg is generally depleted.

Dare *et al.* (2014, fig. 1, p. 790) generated a trace-element variation diagram for magnetite that incorporates the 25 most common trace elements generally present in magnetite. They ordered these elements on the basis of their compatibility in magnetite in a silicate magma, so that the concentration of trace elements should increase from left to right. Any variation from this increase was due to other factors that control magnetite composition. The data of Dare *et al.* (2014) were normalised to bulk continental crust of Rudnick and Gao (2003), which was chosen for its compositional similarities to a magmatic/hydrothermal magnetite source. In evaluating the trace-element composition in different geological settings, Dare *et al.* (2014) suggested that compatible element concentrations (e.g. Ni, V, Co, Zn, Mn and Sn) in magnetite derived from high-temperature hydrothermal fluids (500–700°C) were higher than in magnetite derived from low-temperature fluids. The median composition of magnetite from high-temperature deposits (i.e. porphyry, IOCG, magnetite–apatite and Fe–Ti–V) derived from Dare *et al.* (2014) is shown in Fig. 7 together with median compositions of magnetite they analysed from low-temperature deposits (Fe skarn, Ag–Pb–Zn veins and BIF). In addition, the range of magnetite compositions analysed from sulfide mineralisation in the Dammerget, Tvistbo and Grängsgruvan deposits, and the Izok Lake Zn–Pb–Cu–Ag VMS (Makvandi *et al.*, 2013), Dongyuan and Tengtie Fe skarn (Zhao and Zhou, 2015) and Vegas Peledas retrograde Fe-skarn deposits (Dare *et al.*, 2014) are shown in Fig. 8. Izok Lake occurs in an Archean-age bimodal felsic and mafic terrane metamorphosed to the amphibolite facies, with magnetite occurring in glacial till and a variety of rock types including massive sulfides, gabbro and syenite, whereas magnetite in the Vegas Peledas deposit occurs as masses in calcic Fe-skarn associated with diorite to granite intrusions.

Fig. 6. Principal component analysis of magnetite in: (a) garnet-biotite rock from Dammerget and Grängsgruvan ($N=107$); (b) sulfide mineralisation from Dammerget, Grängsgruvan, Tvistbo, and Grönkullan ($N=95$); and (c) skarn from the Staren limestone, and the Grängsgruvan, Norrgruvan, Tvistbo, Baklångan, Cedercruz and Dammerget deposits ($N=274$). Data are derived and also shown in Fig. 3.

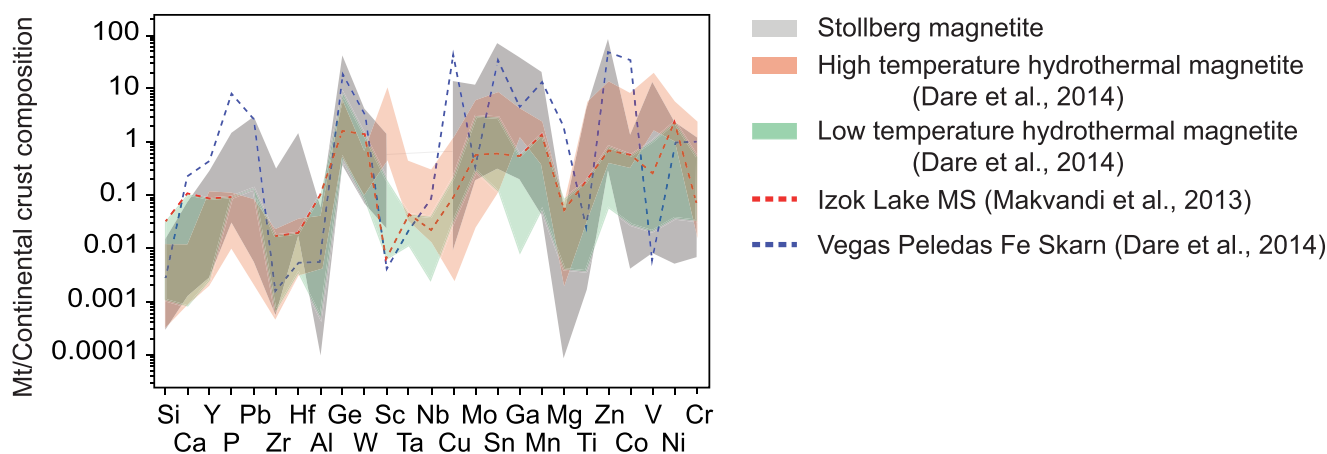


Fig. 7. Multi-element variation diagram of 25 elements for magnetite, comparing compositions of magnetite in the Stollberg ore field to high temperature (500–700°C) hydrothermal magnetite (Dare *et al.*, 2014), low temperature (<500°C) hydrothermal magnetite (Dare *et al.*, 2014), the Izok Lake MS (Makvandi *et al.*, 2013), and the Vegas Peledas Fe skarn (Dare *et al.*, 2014). Data normalised to bulk upper continental crust (Rudnick and Gao, 2003); diagram modified after Dare *et al.* (2014).

Dare *et al.* (2014) proposed that these compatible elements were depleted in magnetite derived from lower temperature fluids as compared to higher temperature fluids due to their low solubility at lower temperatures. Nevertheless, there are concerns with this suggestion because the trace-element pattern of magnetite from the Izok Lake deposit, which formed from lower temperature hydrothermal fluids (probably <400°C), have normalised concentrations that are greater for Mg, Mn, Co and Zn than magnetite derived from high-temperature fluids. The composition of magnetite from the Vegas Peledas Fe skarn overlaps that derived from low-temperature fluids, whereas magnetite in sulfides from the Stollberg district overlap both the low- and high-temperature fields for magnetite. Compared to trace-element concentrations of Mg, Al, Ti, V, Cu, Mn, Zn and Ga in magnetite from the Dongyuan and Tengtie Fe skarn deposits, China (Zhao and Zhou, 2014), as well as in magnetite reported by Nadoll *et al.* (2012b) and Makvandi *et al.* (2013) from unnamed skarn deposits and a porphyry Cu–Mo from the southwestern United States, the Couer d’Alene Ag–Pb–Zn deposit, and the Izok Lake Zn–Pb–Cu–Ag VMS deposit, the composition of magnetite from sulfides in

the Tvistbo, Dammerget, and Gränsgruvan deposits are more similar to those from the Mg skarn deposit reported by Nadoll *et al.* (2012b) and Makvandi *et al.* (2013) and Fe skarns from China. Compared to magnetite from the Izok Lake deposit, Mg, Zn, Co and Ga in magnetite from the Stollberg ore field are more enriched whereas Al, Ti and V are more depleted (Fig. 8). Similarly, compositions of magnetite in sulfide mineralisation from the Stollberg ore field that are normalised to continental crust compositions have a pattern more similar to that of magnetite in Mg and Fe skarns. Compared to Izok Lake, samples from Stollberg are more enriched in Hf, Sc and V, and depleted in Ca, Y, P, Cu, Mg and Co (Fig. 7). Compared to samples from the Stollberg ore field, a single sample of magnetite from the Vegas Peledas skarn deposit is within the range of Stollberg magnetite compositions except that Si and Ca are more enriched and Sn is more depleted in magnetite from Vegas Peledas.

The bulk-rock compositions of marble and skarns in the Stollberg ore field are significantly different from those of the spatially associated variably altered rhyolitic ash–siltstone (Jansson *et al.*, 2013). As a means to test the relative importance of the

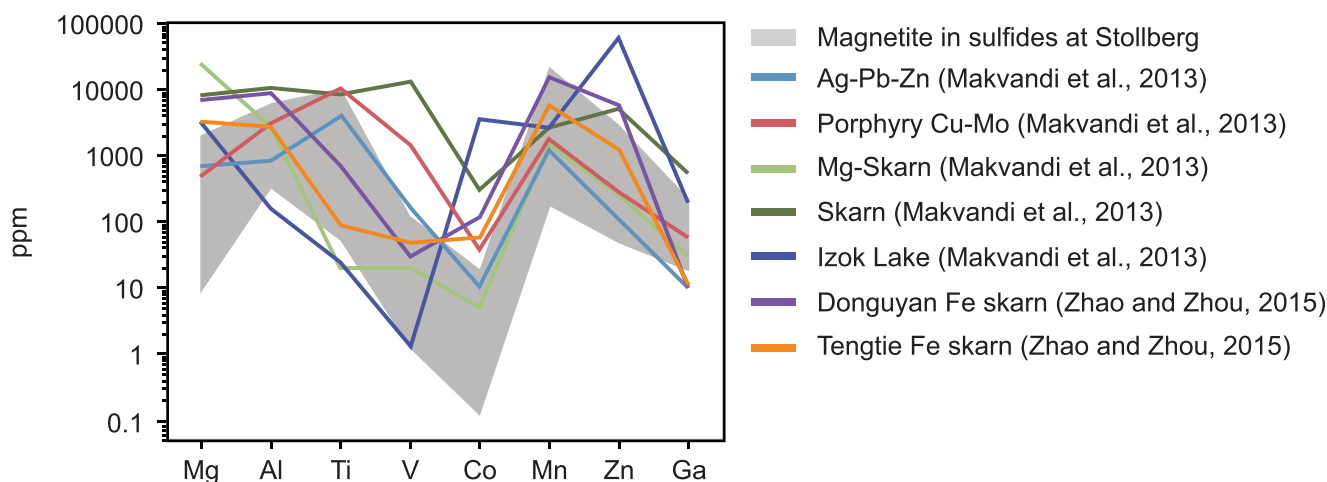


Fig. 8. Elemental concentrations of Mg, Al, Ti, V, Co, Mn, Zn and Ga, for magnetite in sulfide mineralisation from the Stollberg ore field relative to the composition of magnetite from the Dongyuan and Tengtie Fe skarns, China (Zhao and Zhou, 2015), as well as the Coeur D’Alene Ag–Pb–Zn deposit, unnamed porphyry Cu–Mo, Mg-skarn, and skarn deposits from southwestern United States (Nadoll *et al.*, 2012b; Makvandi *et al.*, 2013, and the Izok Lake Zn–Pb–Cu–Ag VMS deposit, Nunavut (Makvandi *et al.*, 2013, Fig. 7).

influence that bulk-rock composition and fluid–rock interactions have on the trace-element composition of magnetite at Stollberg, the trace-element compositions of magnetite in skarn, sulfide mineralisation, and garnet–biotite rocks are compared. The basis for comparison are the variation diagrams that show normalised magnetite composition from different rock types in each deposit (Fig. 8a–f). Overall, the variation diagrams have the same general shape, and it could be argued that regardless of differences in host-rock composition, the trace-element patterns, though similar, are not exactly the same. If this interpretation is correct, it would support the concept that there was a high fluid to rock ratio and that the interaction of the hydrothermal fluid with the host rock was controlled principally by the composition of magnetite, assuming that subsequent metamorphism and recrystallisation were not liberating trace elements to the fluid. However, in detail, the trace-element pattern of magnetite in sulfide mineralisation from the Gränsgruvan and Dammerget deposits resembles that of magnetite in garnet–biotite and quartz–garnet pyroxene rocks in the same deposit, however it is generally different from that for magnetite in skarn (Fig. 9a, d). Moreover, the patterns for magnetite in skarn versus magnetite in sulfides can be distinguished at Tvistbo (Fig. 9b) and Grönkullan (Fig. 9f), thus suggesting that bulk-rock composition is also an important factor controlling magnetite composition.

The relationship between magnetite composition and the composition of its host rock is further demonstrated in a downhole plot of ten trace elements (Mg, Ti, V, Cr, Co, Zn, Ga, Sn, Al and Mn) for magnetite in six samples (sulfide mineralisation, garnet–biotite rock and amphibolite) from 832.5 to 925.0 m in drill hole SSF26 from the Dammerget deposit (Fig. 10), element–element bivariate plots (Figs 3 and 4) and PCA (Fig. 6). Magnetite in amphibolite has low concentrations of many elements, with the exception of V and Cr, which are relatively enriched compared to the other rocks in drill hole SSF26. Box and whisker plots show that magnetite in marble at each location in the Stollberg area has different trace-element signatures (Fig. 11), notwithstanding the similar trace-element compositions of magnetite in samples of the Staren marble located up to 2 km apart from each other (see also Fig. 6c, Table 3). Magnetite in skarn at Gränsgruvan appears to be distinct from magnetite in skarn from other locations along the Stollberg ore trend in that it is more enriched in Mg, Ga, Mn and Ti. Magnetite in skarn from Gränsgruvan and Dammerget is also more enriched in Zn relative to magnetite in skarns elsewhere in the district. It should be noted here that samples of skarn collected from the Gränsgruvan deposit did not contain sulfides as the most recent drilling of this deposit did not target sulfide-bearing skarn, which is associated with other deposits in the Stollberg ore field. It is likely that skarn analysed here is probably skarn after dominantly felsic volcanoclastic rock. Whether this means that the distinct composition of magnetite from Gränsgruvan reflects differences in f_{O_2} and f_{S_2} conditions relative to samples of skarns at other locations, which are associated spatially with sulfides, remains uncertain. The composition of magnetite in each of the altered rock types also probably reflects the composition of the unaltered precursor rock type.

The application of discrimination diagrams to the composition of magnetite

Dupuis and Beaudoin (2011) proposed a series of discrimination diagrams that can be used to distinguish the average trace-element

compositions of magnetite between several deposit types (e.g. iron oxide–copper–gold (IOCG), skarn, banded iron formation, porphyry copper, magmatic nickel and magnetite apatite). They considered such diagrams as being useful as a means to identify the provenance of magnetite and to serve as a potential exploration guide to ore deposits, especially where magnetite is found in surficial sediments including glacial, soil, and alluvial sediments (e.g. Makvandi *et al.*, 2013, 2016b; Pisiak *et al.*, 2015). A plot of individual trace-element compositions of magnetite in sulfide mineralisation, skarn, amphibolite, magnetite-rich conglomerate, and garnet–biotite, gedrite–albite, sericite–garnet–magnetite and garnet–pyroxene rocks from the Stollberg ore field, as a function of $Ca+Al+Mn$ vs. $Ti+V$, show that the composition of magnetite broadly overlaps the skarn field of Dupuis and Beaudoin (2011, fig. 6, p. 328), although magnetite in some samples of sulfide mineralisation from Dammerget and BIF from Grönkullan plot in the porphyry field (Fig. 12a). When plotted in terms of $Ni/(Cr+Mn)$ vs. $Ti+V$ (Fig. 12b), magnetite compositions from the Stollberg ore field fall within the skarn field of Dupuis and Beaudoin (2011, fig. 5, p. 327). However, the $Ni/(Cr+Mn)$ values are less than those shown on fig. 5 of Dupuis and Beaudoin (2011). Huang *et al.* (2013) and Zhao and Zhou (2015) showed an almost identical range of magnetite compositions from the Cihai and Tengtie Fe skarns, respectively, to those reported here from Stollberg when $Ni/(Cr+Mn)$ is plotted as a function of $Ti+V$. The lower $Ni/(Cr+Mn)$ values may better represent the range for skarn deposits than proposed by Dupuis and Beaudoin (2011). Note that where magnetite from these same samples is plotted as a function of $Al/(Zn+Ca)$ vs. $Cu/(Si+Ca)$, which Dupuis and Beaudoin (2011; fig. 4, p. 327) used to discriminate magnetite in VMS deposits from other ore types, there is no overlap in the composition of magnetite from Stollberg with that from VMS deposits (Fig. 12c). This is because Cu, Si and Ca contents of magnetite from Stollberg commonly occur near or below detection limits. When Cu and Ca are above detection limits, Ca generally has a much higher concentration than Cu. The lack of overlap with VMS mineralisation is in keeping with a re-evaluation of the usefulness of this discrimination diagram by Bédard *et al.* (2022) who showed in a study of magnetite compositions from 33 VMS deposits that 82% of the magnetite compositions fall outside of the VMS field.

In a study of several porphyry Cu deposits in British Columbia, Pisiak *et al.* (2015) showed that the composition of magnetite in igneous rocks could be distinguished from magnetite in hydrothermal deposits on the basis of Sn versus Ti. Regardless of the host rock of magnetite in the Stollberg ore field, nearly all magnetite plots within the hydrothermal field of Pisiak *et al.* (2015), even though it is likely that the hydrothermal deposits they studied were derived from “higher temperature fluids”, using the terminology of Dare *et al.* (2014), whereas those in the Stollberg ore field were derived from lower temperature fluids (Fig. 13a). Pisiak *et al.* (2015) questioned the use of the magmatic versus hydrothermal magnetite discrimination diagram of Dare *et al.* (2015) that plots the composition of magnetite as a function of Ti vs. Ni/Cr . As was shown by Pisiak *et al.* (2015) and also shown here in Fig. 13b, the data fail to distinguish magmatic magnetite from hydrothermal magnetite, even though Dare *et al.* (2014) included the trace-element compositions of magnetite from high- and low-temperature hydrothermal deposits. Although the data set for magnetite from the Stollberg ore field includes compositions from all rock types analysed here, the discrimination diagrams clearly yield conflicting conclusions

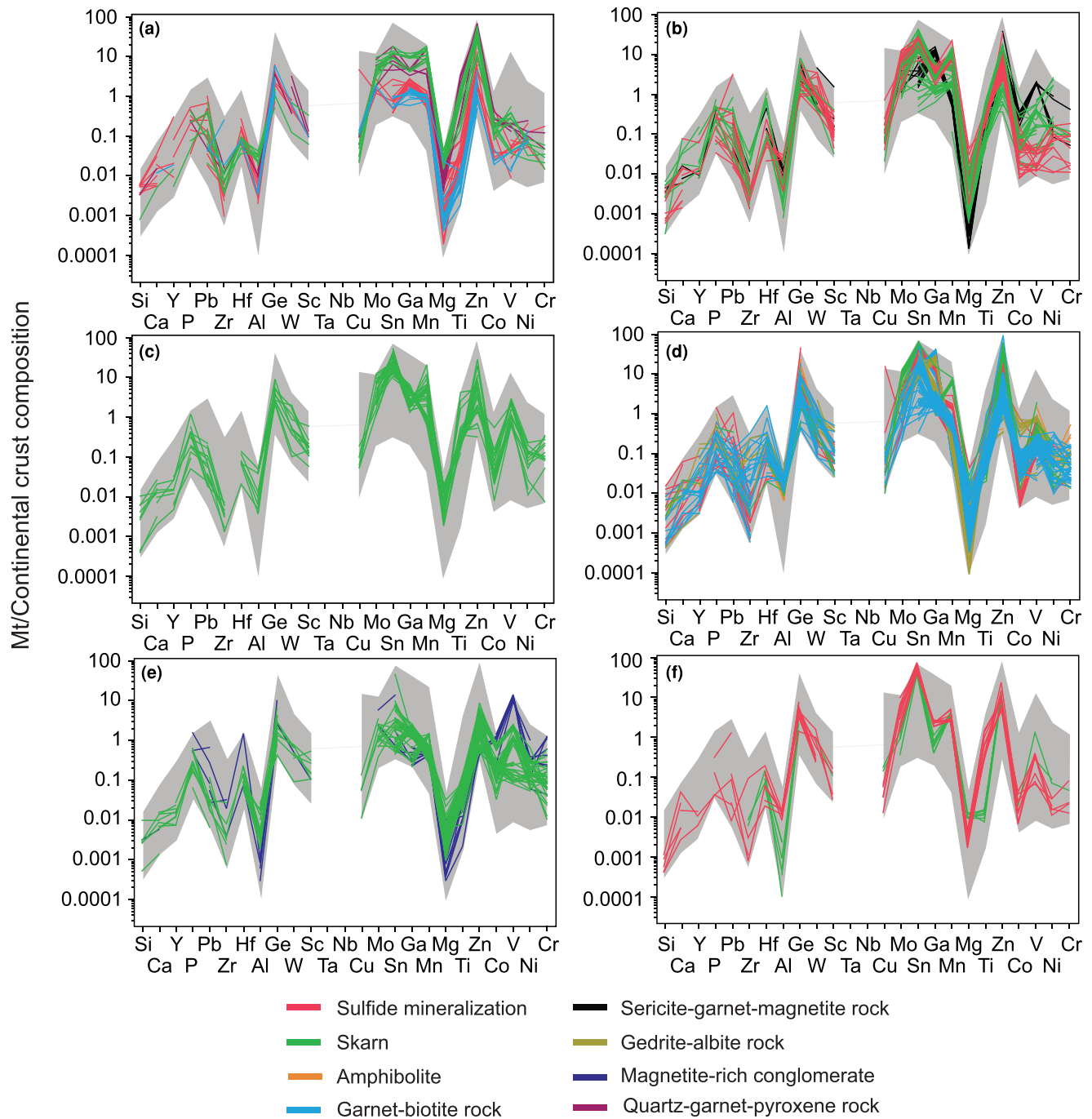


Fig. 9. Multi-element variation diagrams of 25 elements for magnetite compositions from (a) Gränsgruvan; (b) Tvistbo and Norrgruvan; (c) Cedercreutz and Baklängan; (d) Dammberget; (e) Stären; and (f) Grönkullan. Each line represents data from one analysis, and the grey shape represents the furthest extent of all data. The red lines are for magnetite in sulfide mineralisation, green lines for skarn, purple lines for quartz-garnet-pyroxene rocks, light blue lines for garnet-biotite rocks, black lines for magnetite-rich conglomerate, yellow lines for gedrite-albite rocks, dark blue lines for sericite-garnet-magnetite rocks, and orange lines for amphibolite. Data normalised to bulk upper continental crust (Rudnick and Gao, 2003), diagram modified after Dare *et al.* (2014).

regarding the contribution of hydrothermal versus magmatic fluid. This is in a large part due to the variety of rock types that contain magnetite at Stollberg and the variable physico-chemical conditions of the replacement deposits over the 5 km length of the Stollberg ore field. There is no evidence for a causative intrusion related to the formation of the magnetite-bearing rocks and it raises the question regarding the suitability of a plot involving Ti vs. Ni/Cr (Fig. 13a, b) as a discriminator between igneous and hydrothermal processes.

Implications for exploration

The presence of magnetite, distinctive PCA scores, and the elevated contents of some elements in magnetite from sulfides can be used potentially as a prospecting tool for sulfides in the Stollberg area. Previously, the trace-element geochemistry of magnetite has been used as a pathfinder to ore deposits in general by distinguishing the composition of magnetite in rocks associated with ore zones from those not associated spatially with ore (e.g.

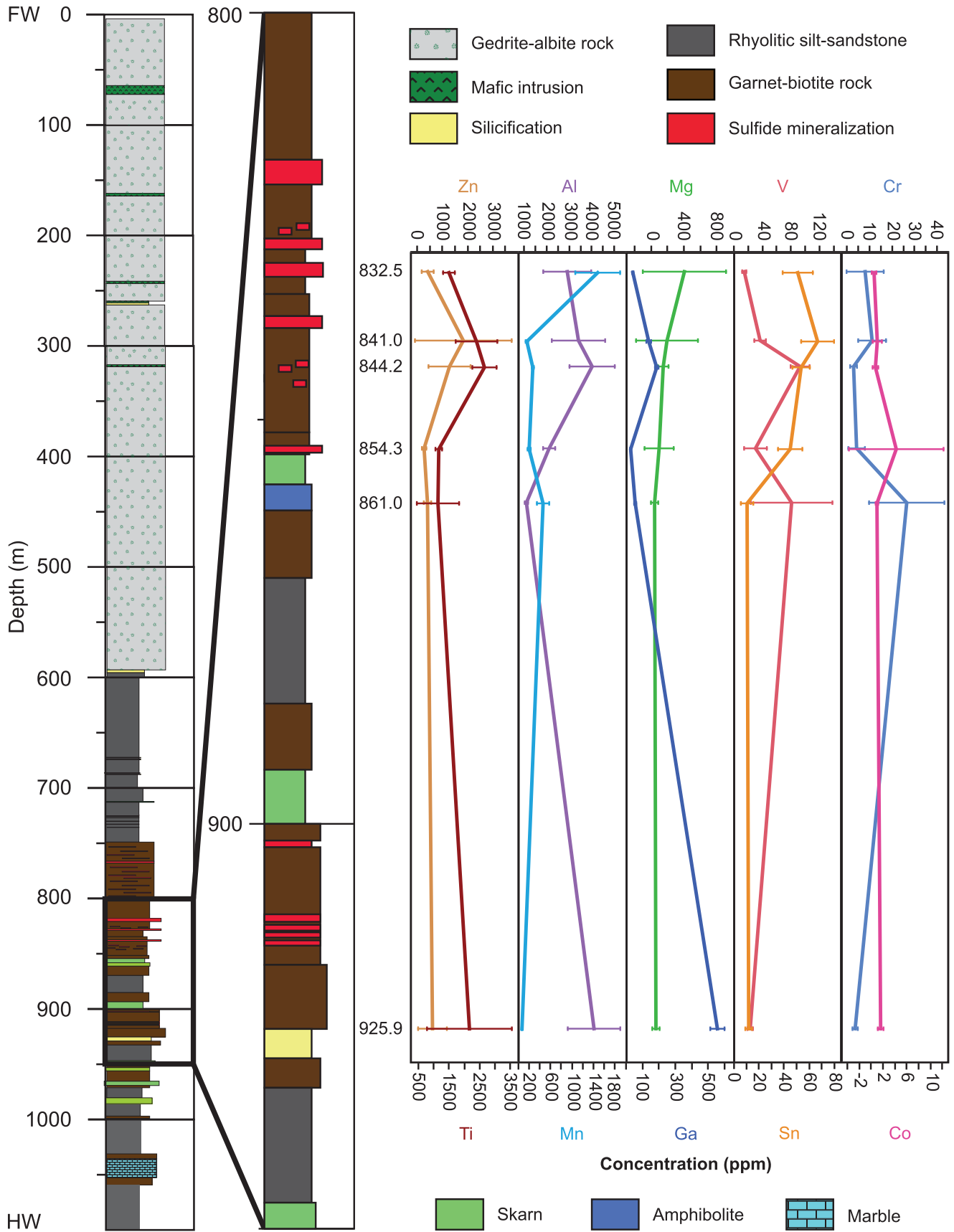


Fig. 10. Downhole variation of the mean value of Al, Co, Cr, Ga, Mg, Mn, Sn, Ti, V and Zn in magnetite from various rocks in drill core SSF26. Error bars represent 1 standard deviation from the mean.

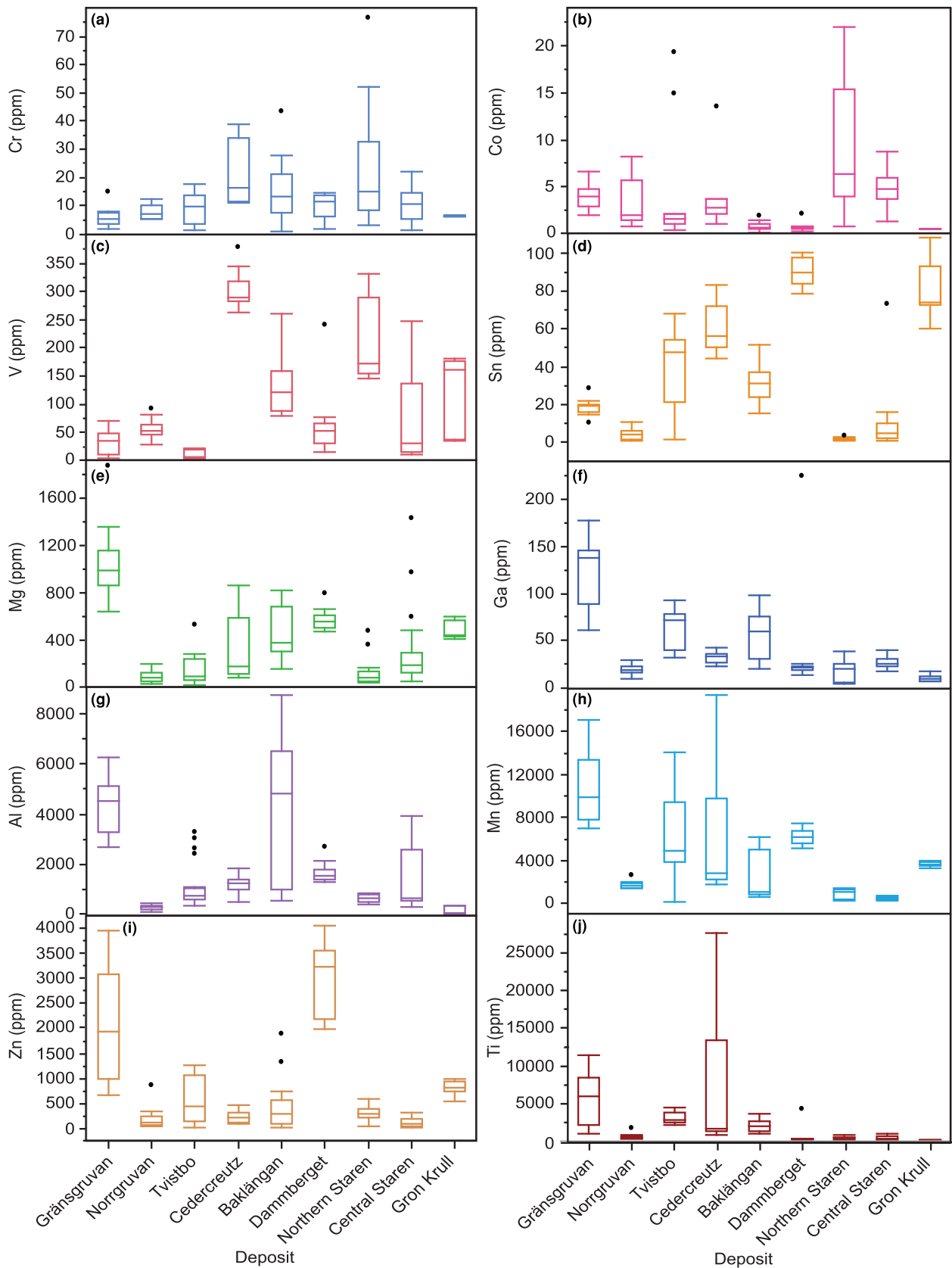


Fig. 11. Box and whisker plots comparing the concentrations (in ppm) of selected trace elements (a) Cr, (b) Co, (c) V, (d) Sn, (e) Mg, (f) Ga, (g) Al, (h) Mn, (i) Zn and (j) Ti in skarn-hosted magnetite from various locations in the Stollberg ore field. Northern Staren refers to drill core SSF16, and Central Staren refers to drill core SSF7. The edges of whiskers represent the 5th percentile (bottom) and 95th percentile values (top), the edges of the box represent the lower quartile and upper quartile (50th percentile), and the solid line across the box represents the median. Outliers are shown as disconnected points.

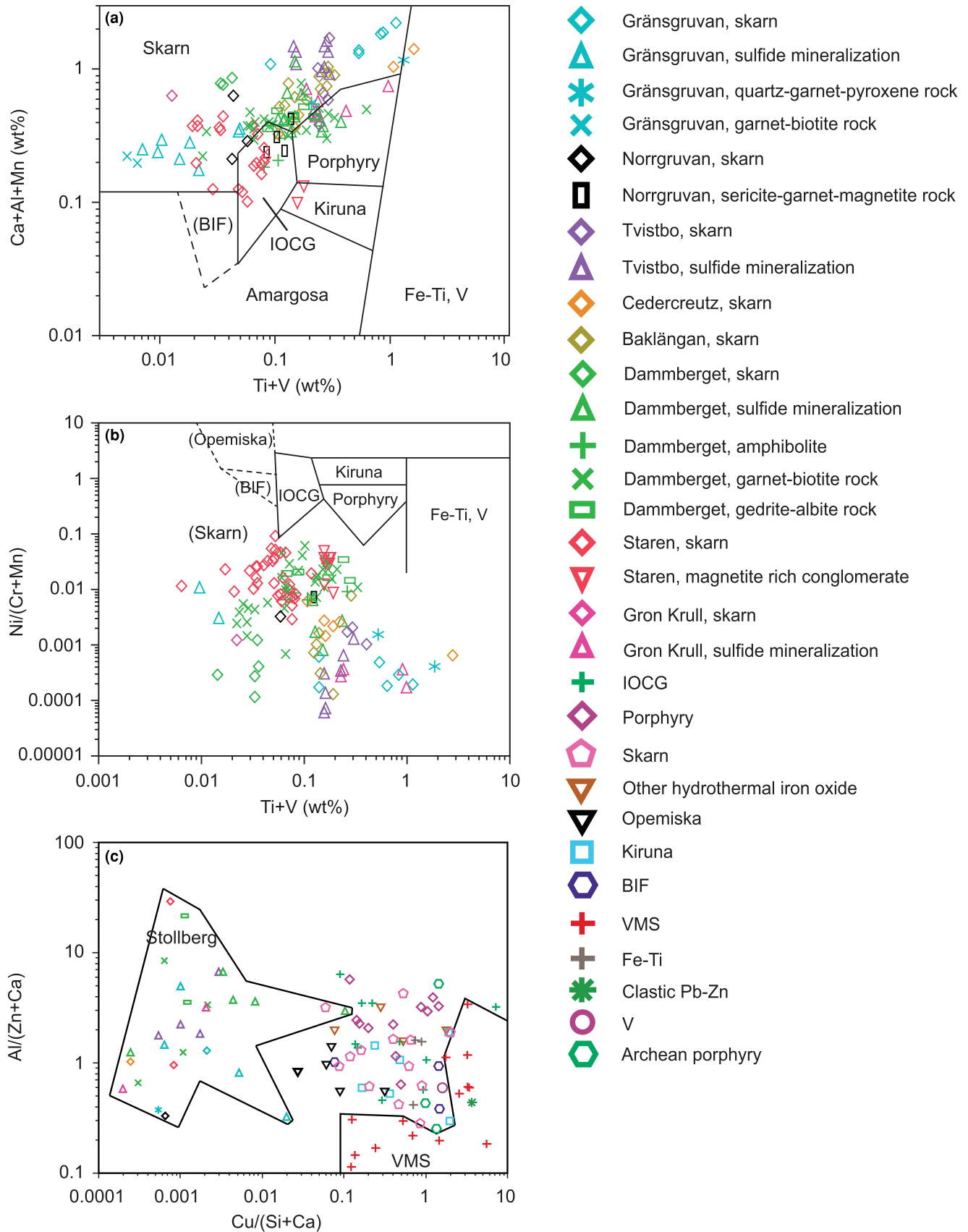


Fig. 12. Discrimination diagrams for magnetite from the Stollberg ore field in terms of (a) Ca+Al+Mn vs. Ti+V; (b) Ni/(Cr+Mn) vs. Ti+V; and (c) Al/(Zn+Ca) vs. Cu/(Si+Ca). Fields for various deposit types [skarn, porphyry, iron oxide-copper-gold (IOCG), banded iron formation (BIF), magmatic Fe-Ti oxide (Fe-Ti, V), Amargosa Fe oxide, Kiruna-type Fe, Opemiska Cu vein and volcanogenic massive sulfide (VMS)] are derived from Dupuis and Beaudoin (2011). Note that the compositions of magnetite from the Stollberg ore field in (c) are distinct from those for magnetite in VMS deposits.

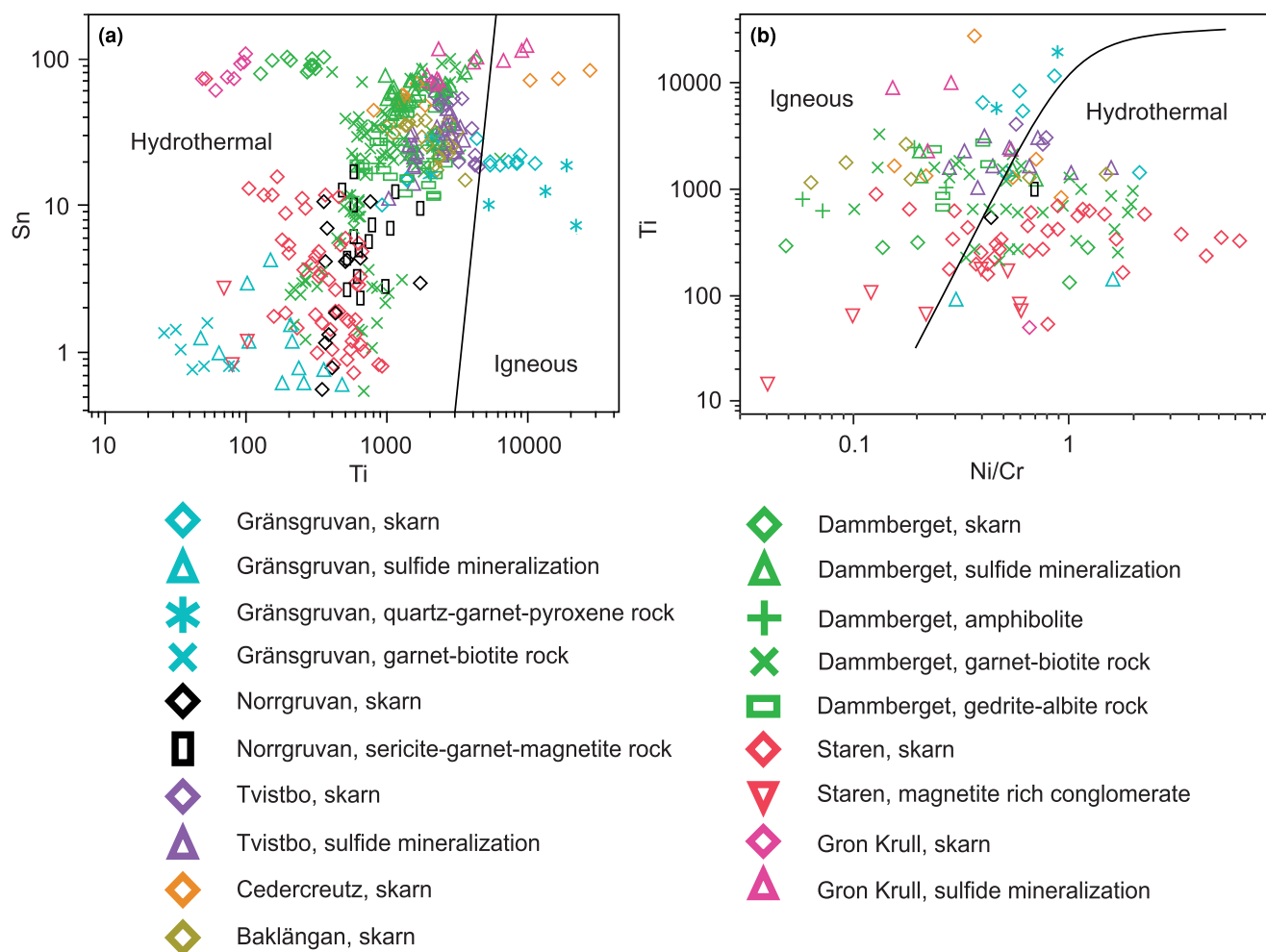


Fig. 13. Igneous vs. hydrothermal magnetite discrimination diagrams based on concentrations of (a) Sn vs. Ti, modified after Pisiak *et al.* (2015), and (b) Ti vs. Ni/Cr, modified after Dare *et al.* (2014). Whereas the composition of magnetite in the Stollberg ore field occurs predominantly in the hydrothermal field of (a) it drapes both igneous and hydrothermal fields in (b).

Dupuis and Beaudoin, 2011; Nadoll *et al.*, 2014; Dare *et al.*, 2015; Makvandi *et al.* 2016b; Ward *et al.*, 2018). In the Stollberg ore field, magnetite occurs in either banded iron formation generally unrelated to sulfides, in altered rocks associated spatially with sulfide mineralisation, and in the Staren limestone. In these settings, the very presence of magnetite is a good indication of hydrothermal activity, as magnetite has not been reported in any igneous rocks in the Stollberg area except for rare magnetite in amphibolite. On that basis alone, regardless of its composition, magnetite should be considered a potential prospective tool for the identification of sulfide mineralisation. Of note, is that the current study focused on the carbonate-hosted replacement plus magnetite deposits associated spatially with the Stollberg marble and not the banded iron formation found stratigraphically below the Staren limestone.

Magnetite in sulfide mineralisation from Dammlerget, Tvistbo and Grönkullan are characterised by PCA scores near the centre of the score plot with a slight distribution to the right (Fig. 6b), and by an overlap in the scatter plots for magnetite in sulfide and garnet-biotite rock from the Dammlerget deposit. In addition, the PCA scores for magnetite in sulfides and garnet-biotite rocks from the Grängsgruvan deposit are essentially identical. Although the concentration of individual trace elements for

magnetite in sulfides overlap with their concentrations in other rock types, magnetite in sulfides is characterised by being enriched in Mn, Sb and Mo relative to most other rocks. Economically important ore elements (Zn, Pb and Cu) are high in some analyses of magnetite in sulfides (the highest for Cu and Pb) in this investigation, however, in general, magnetite is no more enriched than in other rock types. The highest individual Zn concentration (up to 3270 ppm) is from magnetite in skarn from the Dammlerget deposit.

To evaluate whether magnetite that occurs in alluvial sediments or soils in the Stollberg area is derived from the various types of magnetite-bearing rocks would also require a detailed study of the composition of magnetite from the banded iron formation, which was not a focus of the present study, even though magnetite in two samples of iron formation from Grönkullan were analysed. These two samples, Stol 13 and Stol 14B, are from skarn in the iron formation and from a small mass of sulfides within the iron formation, respectively. Notably, magnetite in skarn from Grönkullan contains the lowest median values of Al (66.8 ppm), and among the lowest Ti (83.3 ppm), Cr (5.47 ppm), Co (0.48 ppm) and Ga (6.15 ppm) contents of any sample in the Stollberg area. However, the sample of magnetite in sulfides has trace-element concentrations that cannot be distinguished

from sulfide mineralisation associated with the Stollberg marble. Further trace-element studies of magnetite in iron formation need to be conducted.

Conclusions

Sulfide deposits in the Stollberg ore field are products of sub-seafloor replacement of carbonate and have physical characteristics that superficially resemble both skarn and metamorphosed VMS deposits. The compositions of magnetite from the ore field when plotted in terms of $\text{Ca}+\text{Al}+\text{Mn}$ vs. $\text{Ti}+\text{V}$, and $\text{Ni}/(\text{Cr}+\text{Mn})$ vs. $\text{Ti}+\text{V}$ more closely resemble those of magnetite associated with skarns elsewhere (particularly Fe skarns) rather than with VMS deposits. Moreover, normalised trace-element variation patterns of magnetite composition, using data from Dare *et al.* (2014), Makvandi *et al.* (2013), and Zhao and Zhou (2014), indicate that magnetite from Stollberg has patterns more similar to magnetite from a Mg skarn and the Vegas Peledas, Dongyuan and Tengtie Fe skarns than the metamorphosed Izok Lake VMS deposit.

Discrimination plots using the composition of magnetite to distinguish between a magmatic versus hydrothermal contribution to the ore fluid yield conflicting results, although a plot of Sn versus Ti is consistent with a hydrothermal ore-bearing fluid.

Data obtained here and from other skarn deposits in the literature suggest that the 'skarn field' in the plot of $\text{Ni}/(\text{Cr}+\text{Mn})$ vs. $\text{Ti}+\text{V}$ occurs at $\text{Ni}/(\text{Cr}+\text{Mn})$ between 0.0001 and 0.1 and over a wide range of $\text{Ti}+\text{V}$ concentrations (~0.01 to 5). These broad ranges of compositions reinforce the previously held view that there are a variety of physicochemical conditions that control the compositions of magnetite and that the use of such discrimination diagrams as a way to identify the type of deposit from which the magnetite occurs in must be treated with caution.

Trace-element variation diagrams show that magnetite in various rocks associated with the Stollberg ore field have similar patterns, supportive of magnetite formation from hydrothermal fluids that interacted with their host rocks under conditions of a high fluid to rock ratio. The relative importance of the effects of bulk rock versus fluid composition on the trace-element concentrations of magnetite are unknown though both parameters appear to be factors.

The composition of magnetite has been used in the past as an indicator of provenance and as an exploration guide to various types of ore deposits. The presence of magnetite in the Stollberg ore field is a good indicator of hydrothermal activity. Principal component analysis indicates that the composition of magnetite from the iron formation at Grönkullan can be distinguished from the composition of magnetite in the Stollberg ore trend on the basis of Al, Ti, Cr, Co and Ga concentrations. The identification of PCA scores similar to those identified here for magnetite in sulfide mineralisation and garnet–biotite rock, together with elevated Mn, Sb, Mo, Cu, Pb and Zn, appear to have the greatest promise as indicators to base-metal mineralisation in the Stollberg ore field.

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References

- Acosta-Góngora P., Gleeson S.A., Samson I.M., Ootes L. and Corriveau L. (2014) Trace element geochemistry of magnetite and its relationship to Cu–Bi–Co–Au–Ag–U–W mineralization in the Great Bear Magmatic Zone, NWT, Canada. *Economic Geology*, **109**, 1901–1925.
- Allen R.L., Lundström I., Ripa M., Simeonov, A. and Christofferson H. (1996) Facies analysis of a 1.9 Ga, continental margin, back-arc, felsic caldera province with diverse Zn–Pb–Ag–(Cu–Au) sulfide and Fe oxide deposits, Bergslagen region, Sweden. *Economic Geology*, **91**, 979–1008.
- Allen R.L., Bull S., Ripa M. and Jonsson R. (2003) Regional stratigraphy, basin evolution, and the setting of stratabound Zn–Pb–Cu–Ag–Au deposits in Bergslagen, Sweden. *Geological Survey of Sweden (SGU) and Boliden Mineral AB, Final report for SGU/FoU project 03–1203/99*, 80 pp.
- Allen R.L., Ripa M. and Jansson N. (2008) Palaeoproterozoic volcanic- and limestone-hosted Zn–Pb–Ag–(Cu–Au) massive sulphide deposits and Fe oxide deposits in Bergslagen, Sweden. *Field guide to IGCP project 502 field workshop and 33 IGC excursion No 12, August 14–20, 2008*. 33rd International Geological Congress, Oslo, 84 pp.
- Bédard É., De Bronac de Vazelhes V. and Beaudoin G. (2022) Performance of predictive supervised classification models of trace elements in exploration. *Journal of Geochemical Exploration*, **236**, 106959.
- Beetsma J.J. (1992) Retrograde fluid evolution of the Stollberg Pb–Zn–Fe–Mn (Ag) ore deposit, central Bergslagen, Sweden. *Geologiska Föreningen i Stockholm Förhandlingar*, **114**, 279–290.
- Beunk F.F. and Kuipers G. (2012) The Bergslagen ore province, Sweden: Review and update of an accreted orocline, 1.9–1.8 Ga BP. *Precambrian Research*, **216–219**, 95–119.
- Billström K., Åberg G. and Nord A.G. (1985) Stable isotope data of Bergslagen carbonate and their potential use for sulphide ore prospecting. *Geologiska Föreningen i Stockholm Förhandlingar*, **107**, 169–173.
- Björklund E. (2011) *Mineralogy and lithochemical signature of a stratigraphic profile through the Gränsgruvan Zn-mineralization, Bergslagen, Sweden*. MS thesis, Uppsala University, Sweden.
- Chen W.T., Zhou M.-F., Li X., Gao J.-F. and Hou, K. (2015) In-situ LA-ICP-MS trace elemental analyses of magnetite: Cu–(Au, Fe) deposits in the Khetri copper belt in Rajasthan Province, NW India. *Ore Geology Reviews*, **65**, 929–939.
- Chung D., Zhou M.-F., Gao J.-F. and Chen W.T. (2015) In-situ LA-ICP-MS trace elemental analyses of magnetite: The late Palaeoproterozoic Sokoman Iron Formation in the Labrador Trough, Canada. *Ore Geology Reviews*, **65**, 917–928.
- Clark J.R. and Williams-Jones A.E. (2004) Rutile as a potential indicator mineral for metamorphosed metallic ore deposits. *Diversification de L'Exploration Minérale au Québec, Sous-projet SC2 Final Report*. DIVEX, Montreal, Canada, 17 pp.
- Comas-Cufí M. and Thió Henestrosa S. (2011) CoDaPack 2.0: A stand-alone, multi-platform compositional software. In: *CoDaWork'11: 4th International Workshop on Compositional Data Analysis* (J.J. Egozcue, R. Tolosana-Delgado and M.I. Ortego, editors). Sant Feliu de Guixols, Spain.
- Croghan C.W. and Egeghy P.P. (2003) Methods of dealing with values below the limit of detection using SAS. *Southeastern SAS User Group*. September 22–24, 2003, St. Petersburg, Florida, USA.
- Dare S.A.S., Barnes S.-J. and Beaudoin G. (2012) Variation in trace element content of magnetite crystallized from a fractionating sulfide liquid, Sudbury, Canada: Implications for provenance discrimination. *Geochimica et Cosmochimica Acta*, **88**, 27–50.
- Dare S.A.S., Barnes S.-J., Beaudoin G., Méric J., Boutroy E. and Potvin-Doucet C. (2014) Trace elements in magnetite as petrogenetic indicators. *Mineralium Deposita*, **49**, 785–796.
- Dupuis C. and Beaudoin G. (2011) Discriminant diagrams for iron oxide trace element fingerprinting of mineral deposit types. *Mineralium Deposita*, **46**, 319–335.
- Ferrow E.A. and Ripa M. (1991) Chemistry, reaction mechanisms, and microstructures during retrograde metamorphism of gedrite-biotite-plagioclase bearing rocks from Bergslagen, south-central Sweden. *Lithos*, **26**, 271–285.

- Frank K.S., Spry P.G., Raat H., Allen R.L., Jansson N.F. and Ripa M. (2019) Variability in the geological, mineralogical, and geochemical characteristics of base metal sulfide deposits in the Stollberg ore field, Bergslagen, Sweden. *Economic Geology*, **114**, 473–511.
- Frietsch R. (1982) Alkali metasomatism in the ore-bearing metavolcanics of central Sweden. *Swedish Geological Survey Series C*, **791**, 54 pp.
- Geijer P. (1917) Falutraktens berggrund och malmfyndigheter. *Sveriges Geologiska Undersökning*, **C275**. Geological Survey of Sweden, Stockholm.
- Geijer P. and Magnusson N.H. (1944) De mellansvenska järnmalmernas geologi. *Sveriges Geologiska Undersökning*, **Ca 35**. Geological Survey of Sweden, Stockholm.
- Hedström P., Simeonov F. and Malström L. (1989) The Zinkgruvan ore deposit, south central Sweden: a Proterozoic proximal Zn–Pb–Ag deposit in distal volcanic facies. *Economic Geology*, **84**, 1235–1261.
- Hron K., Templ M. and Filzmoser P. (2010) Imputation of missing values for compositional data using classical and robust methods. *Computational Statistics and Data Analysis*, **54**, 3095–3107.
- Huang X.-W., Zhou M.-F., Qi L., Gao J.-F. and Wan Y.-W. (2013) Re–Os isotopic ages of pyrite and chemical composition of magnetite from the Cihai magmatic-hydrothermal Fe deposit, China. *Mineralium Deposita*, **48**, 925–946.
- Huang X.-W., Zhou M.-F., Qiu Y.-Z. and Qi L. (2015) In-situ LA-ICP-MS trace elemental analyses of magnetite: The Bayan Obo Fe–REE–Nb deposit, North China. *Ore Geology Reviews*, **65**, 884–899.
- Jansson N.F. (2011) *The Relationship Between Metamorphosed, Syn-Volcanic Fe Oxide and Polymetallic Sulphide Ores in the Classic Bergslagen Mining District of Southern Sweden*. PhD thesis, University of Luleå, Sweden.
- Jansson N.F. and Allen R.L. (2011) The origin of skarn beds, Ryllshyttan Zn–Pb–Ag+ magnetite deposit, Bergslagen, Sweden. *Mineralogy and Petrology*, **103**, 49–78.
- Jansson N.F. and Allen R.L. (2015) Multistage ore formation at the Ryllshyttan marble and skarn-hosted Zn–Pb–Ag–(Cu) + magnetite deposit, Bergslagen, Sweden. *Ore Geology Reviews*, **69**, 217–242.
- Jansson N.F., Erismann F., Lundstam E. and Allen R.L. (2013) Evolution of the Paleoproterozoic volcanic-limestone-hydrothermal sediment succession and Zn–Pb–Ag and Fe-oxide deposits at Stollberg, Bergslagen, Sweden. *Economic Geology*, **108**, 309–335.
- Jansson N.F., Zetterqvist A., Malmström L. and Allen R.L. (2017) Genesis of the Zinkgruvan stratiform Zn–Pb–Ag deposit and associated dolomite-hosted Cu ore, Bergslagen, Sweden. *Ore Geology Reviews*, **82**, 285–308.
- Jansson N.F., Sädbom S., Allen R.L., Billström K. and Spry P.G. (2018) The Lovisa stratiform Zn–Pb deposit, Bergslagen, Sweden – Structure, stratigraphy and ore genesis. *Economic Geology*, **113**, 699–739.
- Jansson N.F., Allen R., Skogsmo G. and Turner T. (2021) Origin of Palaeoproterozoic, sub-seafloor Zn–Pb–Ag skarn deposits, Sala area, Bergslagen, Sweden. *Mineralium Deposita*, **57**, 455–480.
- Jolliffe I.T. and Cadima J. (2016) Principal component analysis: A review and recent developments. *Philosophical Transactions. Series A*, **374**, 20150202.
- Kampmann T.C., Jansson N.F., Stephens M.B., Majka J. and Lasskogen, J. (2017) Systematics of hydrothermal alteration at the Falun base metal sulfide deposit and implications for ore genesis and exploration, Bergslagen ore district, Fennoscandian Shield, Sweden. *Economic Geology*, **112**, 1111–1152.
- Kamvong T., Zaw K. and Siegel R. (2007) PIXE/PIGE microanalysis of trace elements in hydrothermal magnetite and exploration significance: a pilot study. In: *15th Australian Conference on Nuclear and Complementary Techniques of Analysis and 9th Vacuum Society of Australia Congress*. University of Melbourne, Australia [Abstract].
- Lagerblad B. and Gorbatshev R. (1985) Hydrothermal alteration as a control of regional geochemistry and ore formation in the central Baltic Shield. *Geologische Rundschau*, **74**, 33–49.
- Longerich H.P., Jackson S.E. and Gunther D. (1996) Laser ablation-inductively coupled plasma-mass spectrometric transient signal data acquisition and analyte concentration calculation. *Journal of Analytical Atomic Spectrometry*, **11**, 899–904.
- Makvandi S., Beaudoin G., Ghasemzadeh-Barvarz M.G. and McClenaghan B.M. (2013) Fingerprinting volcanogenic massive sulfide deposits using magnetite chemistry: Application to till from Izok Lake, Nunavut, Canada. In: *Mineral Deposit Research for a High-tech World*. Proceedings of the 12th Biennial Geology Applied to Mineral Deposits Meeting, 12–15 August, 2013, Uppsala, Sweden.
- Makvandi S., Ghasemzadeh-Barvarz M., Beaudoin G., Grunsky E.C., McClenaghan B.M. and Duchesne C. (2016a) Partial least squares-discriminant analysis of trace element compositions of magnetite from various VMS deposit subtypes: Application to mineral exploration. *Ore Geology Reviews*, **78**, 388–408.
- Makvandi S., Ghasemzadeh-Barvarz M., Beaudoin G., Grunsky E.C., McClenaghan B.M. and Duchesne C. (2016b) Principal component analysis of magnetite composition from volcanogenic massive sulfide deposits: Case studies from the Izok Lake (Nunavut, Canada) and Halfmile Lake (New Brunswick, Canada) deposits. *Ore Geology Reviews*, **72**, 60–85.
- Månsson S. (1979) Preliminär rapport. *Geologiska undersökningar Stollbergfältet*, Grb 103. Häksberg, Sweden, 8 pp.
- Nadoll P., Mauk J.L., Hayes T.S., Koenig A.E. and Box, S.E. (2012a) Geochemistry of magnetite from hydrothermal ore deposits and host rocks of the Mesoproterozoic Belt Supergroup, United States. *Economic Geology*, **107**, 1275–1292.
- Nadoll P., Mauk J.L., Leveille R., Fisher L. and Hough, R. (2012b) Magnetite – An indicator mineral for hydrothermal ore deposits. *Goldschmidt Conference Abstract*. 2012, Montreal, Canada.
- Nadoll P., Angerer T., Mauk J.L., French D. and Walshe, J. (2014) The chemistry of hydrothermal magnetite: A review. *Ore Geology Reviews*, **61**, 1–32.
- O'Brien J.J., Spry P.G., Nettleton D., Ruo X., Teale G.S., Jackson S.E. and Rogers, D. (2015a) Random forests as a statistical method for distinguishing gahnite compositions as an exploration guide to Broken Hill-type Pb–Zn–Ag deposits in the Broken Hill domain, Australia. *Journal of Geochemical Exploration*, **149**, 74–86.
- O'Brien J.J., Spry P.G., Teale G.S., Jackson S.E. and Koenig, A.E. (2015b) Gahnite composition as a means to fingerprint metamorphosed base metal deposits. *Journal of Geochemical Exploration*, **159**, 48–61.
- O'Brien J.J., Spry P.G., Teale G.S., Jackson S.E. and Rogers, D. (2015c) Major and trace element chemistry of gahnite as an exploration guide to Broken Hill-type Pb–Zn–Ag mineralization in the Broken Hill domain, New South Wales, Australia. *Economic Geology*, **110**, 1027–1057.
- Pagé P. and Barnes S.-J. (2009) Using trace elements in chromites to constrain the origin of podiform chromitites in the Thetford Mines Ophiolite, Québec, Canada. *Economic Geology*, **104**, 997–1018.
- Pisiak L.K., Canil D., Grondahl C., Plouffe A., Ferbey T. and Anderson R.G. (2015) Magnetite as a porphyry copper indicator mineral in till: A test using the Mount Polley porphyry copper-gold deposit, south-central British Columbia (NTS 093A). Pp. 141–150 in: *Geoscience BC Summary of Activities 2014, Report 2015-1*. British Columbia, Canada.
- Qi Y., Hu R., Gao J., Gao W. and Gong H. (2021) Trace element characteristics of magnetite: Constraints on the genesis of the Lengshuikeng Ag–Pb–Zn deposit, China. *Ore Geology Reviews*, **129**, 103943.
- Raat H., Jansson N.F. and Lundstam E. (2013) The Grängsgruvan Zn–Pb–Ag deposit, an outsider in the Stollberg ore field, Bergslagen, Sweden. In: *Mineral Deposit Research for a High-tech World*. Proceedings of the 12th Biennial Geology Applied to Mineral Deposits Meeting, 12–15 August, 2013, Uppsala, Sweden.
- Ripa M. (1988) Geochemistry of wall-rock alteration and of mixed volcanic-exhalative facies at the Proterozoic Stollberg Fe–Pb–Zn–Mn(–Ag)-deposit, Bergslagen, Sweden. *Geologie en Mijnbouw*, **67**, 443–457.
- Ripa M. (1994) The mineral chemistry of hydrothermally altered and metamorphosed wall-rocks at the Stollberg Fe–Pb–Zn–Mn(–Ag) deposit, Bergslagen, Sweden. *Mineralium Deposita*, **29**, 180–188.
- Ripa M. (1996) *The Stollberg Ore Field – Petrography, Lithochemistry, Mineral Chemistry, and Ore Formation*. PhD thesis, Lund University, Sweden.
- Ripa M. (2012) Metal zonation in alteration assemblages at the volcanogenic Stollberg Fe–Pb–Zn–Mn(–Ag) skarn deposit, Bergslagen, Sweden. *Geologiska Föreningen i Stockholm Förhandlingar*, **134**, 317–330.
- Rudnick R.L. and Gao S. (2003) The composition of the continental crust. Pp. 1–64 in: *The Crust, Treatise on Geochemistry* (R.L. Rudnick, editor). Elsevier-Pergamon, Oxford, UK.

- Schmidt Mumm A., Dart R.C. and Say P. (2012) Hematite/magnetite trace element geochemistry in base metal exploration. *Journal of Geochemical Exploration*, **118**, 1–13.
- Singoyi B., Danyushevsky L., Davidson G.J., Large R. and Zaw K. (2006) Determination of trace elements in magnetites from hydrothermal deposits using the LA ICP-MS technique. *Society of Economic Geologists Conference*, Keystone, Colorado, USA. CD-ROM.
- Stephens M.B. and Jansson N.F. (2020) Paleoproterozoic (1.9–1.8 Ga) syn-orogenic magmatism, sedimentation and mineralization in the Bergslagen lithotectonic unit, Svecokarelian orogen. Pp 155–206 in: *Sweden: Lithotectonic Framework, Tectonic Evolution and Mineral Resources* (M.B. Stephens, S. Bergman and J. Weihed, editors), Geological Society, London, Memoirs, Vol. 50.
- Stephens M.B., Ripa M., Lundström I., Persson L., Bergman T., Ahl M., Wahlgren C.H., Persson P.H. and Wickström, L. (2009) Synthesis of the bedrock geology in the Bergslagen region, Fennoscandian Shield, south-central Sweden. *Geologiska Föreningen i Stockholm Förhandlingar*, **Ba58**.
- Sundblad K. (1994) A genetic reinterpretation of the Falun and Ämmeberg ore types, Bergslagen, Sweden. *Mineralium Deposita*, **29**, 170–179.
- Verbovsek T. (2011) A comparison of parameters below the limit of detection in geochemical analyses by substitution methods. *Materials and Geoenvironment*, **58**, 393–404.
- Ward L.A., Holwell D.A., Barry T.L., Blanks D.E. and Graham S.D. (2018) The use of magnetite as a geochemical indicator in the exploration for magmatic Ni-Cu-PGE sulfide deposits: A case study from Munalu, Zambia. *Journal of Geochemical Exploration*, **188**, 172–184.
- Warr L.N. (2021) IMA-CNMNC approved mineral symbols. *Mineralogical Magazine*, **85**, 291–320.
- Xu L., Bi X., Hu R., Tang Y., Wang X. and Xu Y. (2015) LA-ICP-MS mineral chemistry of titanite and the geological implications for exploration of porphyry Cu deposits in the Jinshajiang – Red River alkaline igneous belt, SW China. *Mineralogy and Petrology*, **109**, 181–200.
- Zhao W.W. and Zhou M.-F. (2015) In-situ LA-ICP-MS trace elemental analyses of magnetite: the Mesozoic Tengtie skarn Fe deposit in the Nanling Range, South China. *Ore Geology Reviews*, **65**, 872–883.