Review

Characterization of apatite resources in Norway and their REE potential — A review

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A B S T R A C T

Apatite is a necessity for the production of phosphorus fertilizers and presents a potential raw material for the extraction of REE and Y. A wide spectrum of apatite deposits is found in Norway including sedimentary, igneous, and vein type deposits. The igneous deposits which appear to have the greatest potential for exploitation occur associated with alkaline complexes, massif-type anorthosite complexes and monzonitic complexes. One of the most promising is found in the monzonoritic Bjerkreim–Sokndal Layered Intrusion of the early Neoproterozoic Rogaland Anorthosite Province in southwest Norway. The intrusion hosts three cumulate units with high-grade ore zones. The most promising resource is confined to MCU IV which is 50–170 m thick and nearly 10 km long with average normative contents of 10.2% apatite, 12.4% ilmenite and 7.3% vanadium-rich magnetite. The late Neoproterozoic–Cambrian carbonatite-bearing alkaline complexes are generally of low grade to represent potential resources of apatite as the sole commodity. However, apatite may represent a byproduct of potential Nb and REE + Y mineralisation in the Fen Complex in southern Norway. The late Ordovician–Silurian Misværdal complex comprising multiple alkali clinopyroxenite intrusions in the Uppermost Allochthon of the Caledonides in northern Norway contains 1–1.5 km long and 100–200 m wide ultrapotassic clinopyroxenite dykes with average contents of 7–10 wt.% apatite and with intermediate levels of TREY (~0.5 wt.%). However, high levels of Th in the apatite make it less suitable as a raw material for fertilizer production. Apatite–Fe–Ti oxide ores being characteristic for the monzonitic complexes are especially well developed in the Permian Oslo Igneous Province where apatite-rich magnetite clinopyroxenite cumulates are found in the Larvik Plutonic Complex. The Kodal body has an inferred open-pit ore reserve calculated to 70 Mt with approximately 11.6 wt.% apatite, 3.0 wt.% ilmenite and 26.5 wt.% ilmenomagnetite. The apatite contains about 1 wt.% REE. Comparable types of deposits of Palaeoproterozoic age have recently been recognized in the alkalic-calcic mangeritic to syenitic intrusions in the Lofoten–Vesteralen Mangerite Complex in northern Norway, whereas complexes with variable proportions of anorthosites (s.s.), jotunites and mangerites occurring in the Middle Allochthon of the Caledonides in South Norway also are known to host apatite-bearing Fe–Ti oxide deposits, some high in apatite. These complexes represent potential areas for green-field exploration. The TREY (TREE + Y) contents of the apatite in the igneous deposits are discussed and comprise very low levels in the Bjerkreim–Sokndal Layered Intrusion, intermediate levels in the carbonatites and pyroxenites of the alkaline complexes as well as nelsonite dykes and mangerite-associated deposits to high levels in the monzonite-associated deposits.

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Contents

1. Introduction .............................................................. 127
2. Geological setting ............................................................ 127
3. Sedimentary deposits .......................................................... 129
3.1. Stratiform Fe-oxide–apatite deposits ................................................ 129
3.2. Phosphorite occurrences ..................................................... 130
4. Igneous deposits ............................................................ 130
4.1. Deposits in alkaline complexes ................................................ 130
4.1.1. The Fen Complex .................................................... 130
4.1.2. Alkaline complexes of the Seiland Igneous Province ................. 132
4.1.3. The Misværdal Complex ............................................. 135

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1. Introduction

Apatite \([\text{Ca}_5(\text{PO}_4, \text{CO}_3)_{3}(\text{OH}, \text{F}, \text{Cl})]\) is the most abundant crystalline phosphate mineral which is found as an accessory mineral in most rock types on the earth surface. However, exploitable deposits of apatite are mainly found in sedimentary and igneous rocks. The former comprises stratiform phosphorite deposits in shelf-type shale-carbonate sequences that contain high-phosphorous ores of microcrystalline \(\text{CO}_2\)-rich fluorapatite (francolite) and cryptocrystalline collophane. The igneous deposits comprise fluorapatite ores which are mostly hosted by carbonatites and other types of alkaline intrusions. The magmatic ores are generally of lower grade, but give higher quality beneficiated products with low contents of unwanted contaminants (Cd, As, U, Th, Mg, and Al).

Beneficiation products of apatite ores as a commodity are traded as phosphate rock. It is the only significant global resource of phosphorus used dominantly in the manufacturing of nitrogen–phosphorus–potassium (NPK) fertilizers for food-crop nutrition and the production of animal feed supplements. Only 10–15% of the world production of phosphate rock has other applications (e.g. pharmaceuticals, ceramics, textiles and explosives). Sedimentary apatite rock makes the largest contribution to the world production of about 210 Mt (2012) whereas igneous apatite rock accounts for only 10–15% of this total (Jasinski, 2011, 2013). Although present resources of phosphate rock are large, the general high demand has over the years lead to a progressive depletion of high-grade ores and ores of good quality with few contaminants. In general, the phosphate content of economic ore appears to have decreased by about 0.45 wt.% \(\text{P}_2\text{O}_5\) per decade. The economic grade of a sizeable igneous apatite deposit is now 4–5 wt.% \(\text{P}_2\text{O}_5\).

Presently, there is no production of phosphate rock in Norway and it has in the past been of minor importance. Today, Yara International ASA which operates two NPK-fertilizer plants in Norway imports large volumes of phosphate rock. In spite of a long standing production of NPK-fertilizers by the company and its forerunner Norsk Hydro few efforts have been made to explore potential apatite provinces in Norway. However, renewed interest for potential apatite resources in Norway occurred when China, the dominant producer of REE in the world, implemented export restrictions for REE products in 2008. In the rest of the world, this led to intensified search for alternative sources of REE, including deposits of LREE-enriched apatite \([\text{Ca}_5\text{REE}_2(\text{PO}_4, \text{SiO}_4)_3(\text{F})]\) normally found in igneous deposits. Mammag apatite contains commonly more than 0.35 wt.% \(\text{REE}\) or 0.4 wt.% \(\text{REE}\) (see Tables 3 and 4). According to the mineral production statistics given by Brown et al. (2013) and Jasinski (2013) the world production of apatite from igneous deposits in 2011 was 20–30 Mt. If 25 Mt is estimated to contain apatite with an average content of 0.5 wt.% \(\text{REE}\), the total amount of \(\text{REE}\) residing in the world production of igneous apatite would be approximately 125,000 tonnes with predominance of LREE. This is roughly equal to the world production of \(\text{REE}\) in 2011 which suggests that extraction of \(\text{REE}\) from apatite as a byproduct of NPK-fertilizer production represents an important alternative source of \(\text{REE}\).

In Norway a number of geological provinces with a wide variety of apatite deposits are currently investigated by Norges geologiske undersøkelse (NGU; Geological Survey of Norway) including the \(\text{REE}\) contents of apatite. The present paper is based on unpublished results of on-going NGU projects and from data given in published papers, NGU reports, Norsk Hydro internal prospecting reports archived at NGU, NGU Ore Database and NGU Industrial Mineral Database. Most of these data are stored in NGU archives and reports are given in Norwegian like most of previously published accounts on the occurrence of apatite in Norway, e.g., Neumann (1985). Thus, the aim of the present paper is to present for the first time an up to date overview in English of phosphate deposits in Norway and their resources.

2. Geological setting

Norway is situated on the western margin of the Fennoscandian Shield which comprises an Archaean cratonic nucleus with Paleoproterozoic greenstone belts in the northwest being fringed by successively younger mobile belts towards the southwest (Fig. 1). These include the Paleoproterozoic Svecofennian mobile belt in the central part and the Late Mesoproterozoic–Early Neoproterozoic Sveconorwegian mobile belt (SMB) in the southwest. These two mobile belts are separated by the Transscandinavian Igneous Belt (TIB) which developed episodically over the period 1860–1650 Ma in a convergent continental-margin setting (Andersson et al., 2004, and references therein). The Archaean and Neoproterozoic mega-units are to a large part covered by nappes complexes of the Norwegian Caledonides and younger rocks of the Late Carboniferous–Early Triassic Oslo Paleorift including the Oslo Igneous Province (OIP) along the centre of the rift grabens.

The Fennoscandian Shield contains a large diversity of ore deposits and represents a major producer of metals and industrial-mineral raw materials in Europe. In 2013, the three sole producers in Europe of high-grade igneous phosphate rock are found at Siillinjärvi in Finland (Fig. 1) and at Kirovsk and Kolvod in Russia where an Archean carbonatite and Devonian alkaline complexes with nepheline syenite and phoscorite are being worked, respectively (Ilyin, 1989; Puustinen and Kauppinen, 1989). The apatite deposits in the Norwegian part of the Shield comprise sedimentary, igneous and vein type deposits (Table 1). The first two are largely syngenetic whereas the latter is epi- genetic. Potential apatite resources are only found associated with the igneous deposits which occur mainly in the SMB, in the plutonic rocks of the OIP, in the TIB intrusions, and in the Caledonides in order of potential economic importance.

The TIB comprises dominantly batholithic massifs of monzonitic to granitic composition intruding comagmatic volcanites. The majority of the plutonic rocks in the belt have sub-alkaline and alkal-calcic chemistry with I-type signature that in some areas may comprise A-type and rarely S-type varieties (Andersson et al., 2004). The batholiths of alkal-calcic mangerites (opx. monzonites) and monzonites (Malm and Ormaasen, 1978) of the Lofoten–Vesteralen Mangerite Complex (Fig. 1, LVMC) in the northern extension of the TIB contain abundant
Fig. 1. Simplified geological map of Norway showing the distribution of apatite deposits and important provinces with abbreviations used in the text where the numbered deposits are mentioned. Key map depicts the main mega-tectonic units of the Fennoscandian shield and the continent Baltica.
occurrences of Fe–Ti (nos. 4–6, Fig. 1). It was regarded by Gorbatschev (2004) to be part of the TIB plutons, though noting their association with abundant gabbroic and anorthositic intrusions and the absence of coeval volcanics uncommon in other parts of the belt. The rocks within the SMB were formed during the Mesoproterozoic (~1600 Ma) as a consequence of southwestward accretion of arc sequences truncated by calc-alkaline granitoid intrusions (Andersen et al., 2004; Slagstad et al., 2013). Locally these volcanosedimentary sequences contain stratiform Fe–Ti deposits in conjunction with jotunite intrusions (Roberts, 1988). The Lower Allochthon is characterized by enrichment of manganese and phosphorus and were deposited during the end of the Neoproterozoic. These deposits are stratiform Fe–Ti ores comparable to those in the Uppermost Allochthon (no. 3).

The Carboniferous to Triassic OIP comprises saturated to undersaturated alkaline to subalkaline basaltic, latitic, trachytic and ignimbritic volcanics formed in conjunction with fissure eruptions and subsequent formation of central volcanoes with associated calderas. The volcanics are truncated by gabbroic, monzonitic, syenitic and granitic plutons. The early Permian monzonite plutons and associated monzodiorites containapatite-rich ultrapotassic pyroxenite intrusions (Misvaerdal, no. 7).

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rocks suggests that the iron ores originated as chemical sediments de-
posited from Fe–Mn-rich hydrothermal plumes generated in conjunc-
tion with sub-aqueous volcanism.

The iron ores can be subdivided into two major types on the basis of
their average \( P_2O_5 \) contents including high-\( P (>0.7 \text{ wt.}\% \ P_2O_5) \) and low-
\( P (<0.7 \text{ wt.}\% \ P_2O_5) \) ores. These two types of ore are often found at differ-
ent levels in the tectonostratigraphy (Bugge, 1978). The low-\( P \) ores
previously being mined in the Rana district (no. 8 in Fig. 1) are frequently
bordered by Mn-silicate-banded (garnet-rich) units. The high-\( P \) ores are
characterized by low contents of manganese (<0.2 wt.\% MnO). The
largest of the explored high-\( P \) deposits is the Andørja deposit (no. 3 in
Fig. 1) in Troms comprising 8 closely spaced and 7.5–30 m thick ore ho-
argins hosted by biotite amphibolites over a distance of about 2 km
(Geis, 1967; Lindahl and Priesemann, 1999). Estimated reserves based on
125 drill holes are 91 Mt with 11.1–22.1 Fe\(_{\text{mag}}\) and 1.9–2.7 wt.\% \( P_2O_5 \). The fluorocarbonate apatite with 0.23 wt.\% CI is unsuitable for fer-
tilizer production (<0.2 wt.\% CI) and contains only 572 ppm total REE
(TREE) and 206 ppm Y (Lindahl and Priesemann, 1999) that are compara-
table with the contents in the apatite of the Rana district (Bugge, 1978).

3.2. Phosphorite occurrences

The earliest of several major episodes of phosphogenesis in the earth
history, i.e. in the latest Neoproterozoic and Cambrian (Cook and McIlhinney,
1979) has left its finger print in the form of thin beds of phosphatic glauconite sandstones and phosphoritic conglomerates representing littoral to shallow marine deposits. These occur mainly in the Neoproterozoic sedimentary sequences of intercalated shales, feldspathic sandstones and arkoses (sparagmites) in the Lake Mjøsa District and in the Cambro-Ordovician shale-carbonate sequences along the margin of the OIP (Bjørlykke, 1974; Bjørlykke et al., 1976; Neumann, 1985). The thickest of these phosphatic units is part the Neoproterozoic sub-marine fan at Biskopåsen (no. 9 in Fig. 1) a
10–20 m thick basal sequence of grain-supported conglomerates con-
tains abundant immature angular clasts and poorly rounded pebbles of
limestone and phosphorite (1–10 cm in diameter; Bjørlykke et al.,
1976, and references therein).

The Cambrian succession carries phosphate-enriched beds in the
basal shale–sandstone sequences covering the weathered Proterozoic
gneiss complexes along the sub-Cambrian penepane/unconformity. The
sandstone beds rarely exceeding a few metres in thickness contain phosphorite nodule and phosphatic fossil detritus as well as several
interlayered conglomeratic beds (cm to dm thick) with clasts of phosphatic sandstones and phosphorites (Neumann, 1985, and references therein). The thickest sequences of Early Cambrian age are found in
the autochthonous Dividalen Group in Reisadalen of northern Norway
(no. 2) where a 100–150 m thick sequence of sandstones and siltstones comprises phosphate-cemented units with scattered phosphorite
goils, concretions and thin beds of phosphorite conglomerates (Vogt,
1967; Vrålstad, 1976). Grab sampling of these sandstones by Norsk
Hydro at Aveaggave gave a maximum of 2.0 wt.\% \( P_2O_5 \) (Vrålstad, 1976).

Comparable low-grade phosphatic units (<1 m thick) are also found
along unconformities at the base of the Middle Cambrian, the Middle Ordo-
vician and of the Upper Ordovician (Neumann, 1985). The Middle Cambri-
an sandstone bed and overlying phosphorite-pebble conglomerate at
Steinsviken, Lake Mjøsa, (no. 10) contain 2.1 wt.% and 6.69 wt.% \( P_2O_5 \), re-
spectively (Strand, 1929, and references therein). Although REE analyses have not yet been conducted, all of the known occurrences have too
small dimensions and phosphorite grades to be of economic interest.

4. Igneous deposits

4.1. Deposits in alkaline complexes

The alkaline complexes in Norway range in age from the
Palaeoproterozoic to the Silurian and include those at Fen, Seiland
Igneous Province (SIP) and Misvær at the treated below (nos. 12, 1 and 7,
respectively). These complexes show different ratios of carbonatites
versus mafic igneous rocks that are highest in the former complex
and lowest in the latter. All the complexes are enriched in apatite, but
differ in contents of high field strength elements (e.g. Y, REE, Th, Nb)
which occur strongly enriched in the Fen carbonatites, but are compara-
ble low in the Lillevold (SIP) and Misvær at the complexes.

4.1.1. The Fen Complex

The Fen Complex has become famous by the pioneering work by
Brogger (1921) who recognized carbonatites as a specific group of mag-
matic rocks and established it among the type examples of its kind.

4.1.1.1. Geological outline. The Fen Complex is situated in Mesoproterozoic
granitic orthogneisses about 17 km west of the southwestern margin of
the OIP (Figs. 1 and 2). It is surrounded by a wide halo (1500 km\(^2\) of
small satellite intrusions and dykes of similar composition as the rocks of
the central complex (Bergstøl and Svindal, 1960; Dahlgren, 1987,
2004). Meert et al. (1998) advocated on the basis of their own \( ^{40}\)Ar/\( ^{39}\)Ar
dating of phlogopites in ultramafic lamprophyre (damtjernite) and
phonolite satellites that the best estimate for the emplacement of
the complex is 583 ± 10 Ma or late Neoproterozoic (Ediacaran).
The emplacement occurred at shallow level in the crust and probably at
a 1–3 km depth below the sub-Cambrian peneplane (Sæther, 1957).
Gravity data indicates that the central complex is underlain by a pipe-
shaped body of dense material, probably mafic–ultramafic silicate
rocks extending from a few hundred metres below the present surface
to at least 15 km in depth (Ramberg, 1973).

The central complex comprises an about 9 km\(^2\) sub-circular com-
posite intrusion of dominantly peralkaline to sub-alkaline carbonatites
and intrusive silicate rocks. The latter rocks represent, with the excep-
tion of the ultramafic lamprophyres, early formed members of the com-
plex (Andersen, 1987a, 1988), Fluids emanating from the early ijolitic
intrusions caused pervasive alkali metasomatism or fenitisation of the
granitic gneisses in the western half of the complex (Verschure and
Maier, 2005 and references therein). Hydrothermal veinng and meta-
somatism overprinting the early formed fentines and their gneissic
protoliths are developed in conjunction with subsequent carbonatite
intrusions. It includes widespread plagiogossitisation in association with
dykes of apatite-rich calcite-carbonate (soevite) in the
central and northern parts of the complex (Andersen, 1983, 1984, 1987b,
1989; Verschure and Maier, 2005). The eastern half of the complex is
according to Andersen and Qvale (1986) composed mainly of chlorite-
ankerite ferrocarbonatites possibly representing heterogeneous pyro-
clastic intrusions emplaced coevally with dykes and plugs of ultramafic
lamprophyres. The ferrocarbonatite is altered to a fine-grained hema-
tite–calcite– Dolomite carbonatite (redberg) caused by oxidation of an-
k erite and precipitation of microscopic grains of hematite in neofomed
calcite and dolomite. The formation of redberg or carbothermal carbonatite according to the classificiation by Mitchell (2005) was trig-
gered by influx of groundwater-derived hydrothermal fluids during the
final stage of the ferrocarbonatite magmatism (Andersen, 1984;
Andersen and Qvale, 1986). The late chlorite-bearing ferrocarbonatites,
carbothermal carbonatite and carbonated gneisses in their immediate
surroundings contain abundant hydrothermal veins of magnetite and
hematite which were mined in the period 1652–1957 yielding a total of
1 Mt of ore (Andersen, 1983; Bugge, 1978).

4.1.1.2. Mineralisation of Nb, Y-REE and Th. The Søve mine produced a
total of about 1 Mt of crude ore with 0.31–0.57 wt.% Nb\(_2O_5\) during the
period 1953–1965 (Dahlgren, 2005). Most of the ores were mined from a body of dominantly calcite-carbonatite that was followed under-
ground from the Cappelen quarry (CQ in Fig. 2) to a depth of about
150 m (Bjørlykke and Svindal, 1960; Sæther, 1957). Subordinate
amounts of ores were also excavated from a calcite-carbonatite dyke in
the Hydro quarry (HQ) and from dolomitic carbonatites in the Tufte
quarry and its underground workings (Fig. 2). The pyrochlore in the ores contains up to 2 wt.% ThO₂ + REO and 2 wt.% Ta₂O₅ and is locally uranium-rich (Bjørlykke and Svinndal, 1960).

Presently, several companies are investigating the Fen Complex for potential resources of Y, REE and Th. The contents of these elements appear according to Bugge (1978) to successively increase from calcite-carbonatite through ankeritic ferrocarbonatite to carbothermal carbonatites, whereas the Nb content shows the opposite trend. Analyses show that the ankeritic ferrocarbonatites and carbothermal carbonatites and associated iron ores are highly enriched in REE with contents normally exceeding 0.3 wt.% TREE and with frequent analyses yielding values of 1–2 wt.% TREE (Svinndal, 1970, and references therein; Møller et al., 1980; Andersen, 1987a,b). The REE enrichment is caused by fine-grained dissemination of monazite, bastnäsite, parisite and/or synchysite (Andersen, 1986a; Lie and Østergaard, 2011a). Thorium, mainly residing in thorite, is especially enriched in the carbothermal carbonatites and associated hematite veins. They yield normally analytical values in the range 0.1–1.0 wt.% Th in contrast to 0.05–0.12 wt.% Th in the ferrocarbonatites (Lie and Østergaard, 2011a, 2011b).

4.1.1.3. Distribution of apatite. Presently, there exist no systematic investigations of apatite-bearing rocks in the Fen Complex. The different lithologies in the complex contain according to published major element analyses highly variable concentrations of P₂O₅ or apatite (Table 2). The granitic country rocks and their fenitised equivalents carry only accessory amounts of apatite. This is also the case for most of the nepheline syenites with the exception of the more melanocratic varieties which occasionally exceed 2 wt.% P₂O₅. A similar concentration of P₂O₅ is also commonly encountered in the ultramafic melteigite which according to Verschure and Maijer (2005) contains locally up to 14% apatite. The melteigite possibly representing cumulates of an ijolitic magma (Andersen, 1988) is somewhat higher in P₂O₅ compared with the more nepheline-rich ijolite–urrite intrusive phases and later intrusions of

Table 2

<table>
<thead>
<tr>
<th>Rock type</th>
<th>Sample Numbers</th>
<th>P₂O₅ Average</th>
<th>St dev</th>
<th>Maximum</th>
<th>Minimum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Country rock gneisses</td>
<td>15</td>
<td>0.08</td>
<td>0.06</td>
<td>0.25</td>
<td>0.02</td>
</tr>
<tr>
<td>Alkali syenites</td>
<td>9</td>
<td>0.71</td>
<td>0.76</td>
<td>2.34</td>
<td>0.03</td>
</tr>
<tr>
<td>Ijolite–urrite</td>
<td>4</td>
<td>0.98</td>
<td>0.77</td>
<td>2.12</td>
<td>0.44</td>
</tr>
<tr>
<td>Nepheline pyroxenite (melteigite)</td>
<td>7</td>
<td>1.86</td>
<td>0.61</td>
<td>2.58</td>
<td>0.64</td>
</tr>
<tr>
<td>Aegirine calcite-carbonatite</td>
<td>2</td>
<td>2.15</td>
<td>1.05</td>
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<td>1.40</td>
</tr>
<tr>
<td>Biotite-amphibole pyroxenite</td>
<td>2</td>
<td>3.69</td>
<td>0.25</td>
<td>3.87</td>
<td>3.51</td>
</tr>
<tr>
<td>Biotite–calcite–carbonatite</td>
<td>6</td>
<td>3.17</td>
<td>2.41</td>
<td>6.92</td>
<td>0.21</td>
</tr>
<tr>
<td>Dolomite-carbonatite</td>
<td>2</td>
<td>2.02</td>
<td>1.15</td>
<td>2.83</td>
<td>1.21</td>
</tr>
<tr>
<td>Ankeritic ferrocarbonatite</td>
<td>4</td>
<td>1.18</td>
<td>1.18</td>
<td>2.80</td>
<td>0.24</td>
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<td>1.01</td>
<td>0.37</td>
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<td>0.52</td>
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<td>1.76</td>
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<td>1.25</td>
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<td>Hematite ore</td>
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<td>1.47</td>
<td>0.90</td>
<td>2.10</td>
<td>0.83</td>
</tr>
</tbody>
</table>

Fig. 2. Simplified geological map of the Fen Complex with locality names. Modified from Dahlgren (2004). Abbreviations: CQ = Cappelen quarry at Søve mine, HQ = Hydro quarry and TQ = Tufte quarry north of Tufte underground mine.
ankeritic ferrocarbonatites and associated ultramafic lamprophyres. Higher P₂O₅ grades are detected in the biotite–amphibole pyroxenite intrusion at Vipeto. This intrusion contains 3–4 wt.% P₂O₅ and characterized by up to cm long prisms of apatite and finer grained aggregates which often occur associated with abundant segregations, veinlets and narrow cross-cutting dykes and lenses of calcite- and dolomite-carbonatites. Comparable apatite-rich dykes and lenses are also found in fenites and granitic gneisses along the southern margin of the Fen Complex (Brogger, 1921).

Most of the carbonatites in the complex are enriched in apatite which may become segregated and yield analyses exceeding 10 wt.% P₂O₅ (Tore Vrålstad, pers. com. 2013). Although there is a considerable spread in the analytical values for the carbonatites (Table 2), it appears from the average values that the calcite-carbonatites (3.2 wt.% P₂O₅) are the main carrier of apatite in the complex. The calcite-carbonatite dykes and associated dolomite-carbonatites in the Tufte–Cappelen mining area yield analytical values of up to 6.9 wt.% P₂O₅ (Hornig-Kjarsgaard, 1998). Ore reserve calculations by Bjørllykke (1953) prior to the opening of the Seve mine gave an estimate of 1.4 Mt with 0.24 wt.% Nb₂O₅ and 2% pyrochlore and 62% calcite (Hornig-Kjarsgaard, 1998).

In the Tufte area there occur a number of apatite-rich lens-shaped bodies varying in composition from almost carbonate-free apatite–phlogopite–amphibole–magnetite rocks to apatite–rich calcite- and dolomite-carbonatites (Andersen, 1986b). On the basis of fluid inclusion studies and intergrowth textures Andersen (1986b, 1987a, 1990) advocated that the apatite represent an early liquidus mineral which largely crystallised prior to the final emplacement of the carbonatite magma. The carbonate-poor and apatite-rich rocks probably formed as largely crystallised prior to the final emplacement of the carbonatite magma. The bulk of the various intrusions of the SIP was emplaced in the Northern Norway (Roberts and Gee, 1985). The intrusions constituting the Kalak Nappe Complex in the Middle Allochthon of the Caledonides are more than 5000 km². They were emplaced into arenaceous meta-sedimentary rocks containing pelitic to semi-pelitic turbidites and carbonatites (Andersen, 1986a).

The ankeritic ferrocarbonatite is lower in apatite than the other carbonatite types (Table 2) and carries normally 1–2% modal apatite, locally rising to 5–10% (Sæther, 1957). The carbothermal carbonatites show a similar spread in analytical values as the calcite-carbonatites, but with a lower average (2.4 wt.% P₂O₅). The hydrothermal hematite ores worked by the Fen iron mines contained according to Bugge (1978) an average of 1.05 wt.% P₂O₅. Three apatite concentrates from the carbonatite-carbonatites yielded analyses exceeding 10 wt.% P₂O₅ and characteristic low contents of REE and high field strength elements (Hornig-Kjarsgaard, 1998).

A proper evaluation of the apatite resources in the Fen Complex has to await more systematic sampling. Presently it appears that apatite in the carbonatites may represent a potential by-product if exploitable Nb and REE + Y deposits are defined in the future.

4.1.2. Alkaline complexes of the Seiland Igneous Province

4.1.2.1. Geological outline. The Seiland Igneous Province (SIP) is part of the Kalak Nappe Complex in the Middle allochthon of the Caledonides in Northern Norway (Roberts and Gee, 1985). The intrusions constituting the SIP cover an area of more than 5000 km². They were emplaced into arenaceous meta-sedimentary rocks containing pelitic to semi-pelitic units with some interlayered marbles (Roberts, 1973). Most of the plutonic rocks and their wall rocks are affected by two tectonothermal events (D1 and D2) which mainly occurred under amphibolite to granulite facies conditions (Elvevold et al., 1994; Roberts, 1996, and references therein). The bulk of the various intrusions of the SIP was emplaced in the Ediacaran (570–560 Ma) (Pedersen et al., 1989; Roberts et al., 2010). The SIP comprises a wide spectrum of magmas. The early stage intrusions according to Roberts (1996) comprise sub-alkaline gabbros of calc-alkaline affinity that are intruded by late diorites, monzonites and granodiorites in their interior (Fig. 3). The intermediate stage is the most voluminous. It consists of layered masses of gabbro-norites and norites (some of them showing tholeiitic trends) that are followed by emplacement of syenogabbros with magma compositions transitional to alkali basalts. Subsequent intrusions are represented by layered olivine gabbros often carrying associated pegmatitic gabbros (Tappendal) and ultramafic masses derived from more alkaline magmas (alkali olivine basaltic or alkali picritic). The late stage intrusions comprise mainly strongly alkaline rocks (syenites, nepheline syenites and carbonatites) derived from nephelinitic magmas (Robins, 1996). The rocks constituting the phosphoritic alkaline complexes at Lillebukt, Breivikbotn, and Pollen include alkaline nepheline syenites, hornblende clinopyroxenites and carbonatites with associated envelopes of mafic-ultramafic fenites. The miasitic chemistry of the alkaline rocks that was identified by Barth (1927) may have an indirect influence on the characteristics low contents of REE and high field strength elements in these rocks (Heier, 1961). The mafic–ultramafic fenites which represent metasomatic alteration of the gabbroic wall rocks are very heterogeneous in composition and contain frequently nebulitic remnants of the protolith. It is a coarse-grained ultramafic rock composed of variable proportions of amphibole, aegirine–augite, biotite, apatite and calcite. The dykes spilling into early Cambrian (530–520 Ma) (Pedersen et al., 1989; Roberts et al., 2010). The SIP comprises a wide spectrum of magmas. The early stage intrusions according to Roberts (1996) comprise sub-alkaline gabbros of calc-alkaline affinity that are intruded by late diorites, monzonites and granodiorites in their interior (Fig. 3). The intermediate stage is the most voluminous. It consists of layered masses of gabbro-norites and norites (some of them showing tholeiitic trends) that are followed by emplacement of syenogabbros with magma composition transitional to alkali basalts. Subsequent intrusions are represented by layered olivine gabbros often carrying associated pegmatitic gabbros (Tappendal) and ultramafic masses derived from more alkaline magmas (alkali olivine basaltic or alkali picritic). The late stage intrusions comprise mainly strongly alkaline rocks (syenites, nepheline syenites and carbonatites) derived from nephelinitic magmas (Robins, 1996). The rocks constituting the phosphoritic alkaline complexes at Lillebukt, Breivikbotn, and Pollen include alkaline nepheline syenites, hornblende clinopyroxenites and carbonatites with associated envelopes of mafic-ultramafic fenites. The miasitic chemistry of the alkaline rocks that was identified by Barth (1927) may have an indirect influence on the characteristics low contents of REE and high field strength elements in these rocks (Heier, 1961). The mafic–ultramafic fenites which represent metasomatic alteration of the gabbroic wall rocks are very heterogeneous in composition and contain frequently nebulitic remnants of the protolith. It is a coarse-grained ultramafic rock composed of variable proportions of amphibole, aegirine–augite, biotite, apatite and calcite. The
fenites are enriched in P, C, Zr, Sr, Nb, Ba, LREE, Ti, Fe, Ca, K and Mn and depleted in Si, Al and Na relative to the composition of the gabbroic protolith (Robins and Tysseland, 1983). The overall rock association, and geochemistry indicate formation of SIP in a continental rift during crustal extension (Roberts et al., 2010).

4.1.2.2. Apatite distribution in selected intrusions. Apatite represents an accessory mineral in most of the intrusions, but is especially enriched in those derived from alkali basaltic and nephelinitic magmas, including the Tappeluft, Breivikbotn, Pollen and Lillebukt intrusions which have been explored in some detail by a number of companies. The Tappeluft pegmatitic gabbro (Fig. 3; Mørk and Stabel, 1990) which was sampled by Norsk Hydro contains an average of 1.88 wt.% P₂O₅ and a maximum of 4.29 wt.% P₂O₅ (Aggerholm and Parr, 1984). The Breivikbotn alkaline complex yields modal apatite contents in the following ranges for the different rocks of the complex: nepheline syenites: 0.1–0.5%; aegerine-augite pyroxenites: 0.4–7.7%; melanocratic syenites: 0.5–8.7%; syenites: 0.2–2.7% and carbonatites: 0.3–6.4% (Sturt and Ramsay, 1965).

The Lillebukt Complex (no. 1) is the only one that has been mapped in detail by MSc students from the University of Bergen and explored by Elkem, Yara International and NGU. It is situated on the southern central part of the island Stjernøy (Fig. 3) and covers an area of about 13 km² (Gautneb and Ihlen, 2009; Skogen, 1980b). It is the largest of the alkaline complexes within the SIP and is hosted by layered metagabbros belonging to the intermediate stage of plutonic activity in the province. The development of the complex started with the emplacement of hornblende clinopyroxenite dykes followed by intrusions of monzonites, syenites, nepheline syenites, carbonatites and late nepheline syenite pegmatites (Fig. 4). Mafic fenites derived from the layered gabbros and felsic fenites formed by metasomatism of early syenites and monzonites in the south constitute a 500–1000 m wide zone around the central carbonatite intrusion.
The main apatite-bearing rock types are hornblende clinopyroxenite dykes and calcite-carbonatites. The former dykes occur abundantly in the mafic fenites and gabbros along the northern and eastern margins of the complex where they form a 50–600 m wide and 11 km long belt of sub-parallel dykes separated by screens of ultramafic fenites constituting about 40% of the rocks. Individual dykes are generally 50–100 cm wide with a maximum width of 10 m. They consist of aluminous titanian salite, magnesian hastingsite and apatite together with commonly sub-ordinate ilmenite and ilmenomagnetite, as well as minor hercynitic spinel and calcite (Robins, 1985). These steeply-dipping coarse-grained to pegmatitic dykes commonly show an inward directed growth of up to 50 cm long crystals of clinopyroxene and hornblende forming comb structures with interstitial aggregates of coarse-grained apatite which is locally developed in up to 10 cm large skeletal crystals. Some of the dykes are unusually apatite-rich, often with cores containing up to 50 vol.% apatite and Fe–Ti oxides (Robins, 1985).

The mineralogy and petrochemistry of the carbonatites were investigated by Strand (1981) who revealed that the carbonatites of the complex represent calcitic silicocarbonatite with about 40% of strontian calcite. The main minerals comprise calcite, biotite, alkali hastingsite, apatite, ilmenite and titanomagnetite and occur together with minor alkali feldspar and nepheline. Apatite occurs as evenly distributed mm-sized prisms (Fig. 5a). Fe–Ti oxides are common locally. The carbonatite has developed a crude banding defined by variable proportions and alignments of mafic minerals that are the result of mechanical redistribution and recrystallization during post-magmatic deformation and metamorphism (Skogen, 1980a; Strand, 1981).

A total of 330 grab and percussion-drilling samples have been collected from the carbonatites and hornblende-clinopyroxenite dykes. The latter includes probably also samples of the thin screens of ultramafic fenites which generally are low in apatite. The carbonatites (218 analyses) show an average of 2.33 wt.% P₂O₅ with a maximum of 6.45 wt.% P₂O₅. 23 of these samples contain more than 6 wt.% P₂O₅. The hornblende clinopyroxenites (56 analyses) yield an average of 2.35 wt.% P₂O₅ which is similar to the average for the carbonatites. 4 of these analyses gave values in excess of 8.00 wt.% P₂O₅ with a maximum of 13.49 wt.% P₂O₅. The average content of apatite in all samples is 5.6 wt.% apatite. In well-exposed and densely sampled areas of

Fig. 5. Macroscopic features of the different types of apatite ores. a) Apatite-bearing biotite calcite-carbonatite of the Lillebukt Complex (length of match is 44 mm), b) ultrapotassic coarse-grained biotite pyroxenite in the Mivardal Complex with interstitial white alkali feldspar aggregates and light grey acicular apatite (diameter of coin is 26 mm), c) layered apatite–Fe–Ti oxide ore of MCU IV in the Bjerkreim–Sokndal Layered Intrusion of the RAP (length of hammer is 0.7 m), d) close-up of nelsonite (black) breccia-dyke at Nordre Falstad in the LVMC containing fragments of the wall rocks comprising charnockite and mangerite dykes in fine-grained felsic granulites (length of hammer is 0.5 m), and e) patchy dark apatite–Fe–Ti oxide ore in mangerite at Grindvik in the LVMC (length of magnet pen is 13 cm).
about 300 m × 300 m in the carbonatite and clinopyroxenite dykes yield averages of 3.00 wt. % P2O5 and 2.93 wt. % P2O5, respectively. Analyses of hand-picked coarse-grained crystals show that the apatite from the carbonatite contains 5601 ppm TREE and 288 ppm Y that are higher than apatite from a hornblende clinopyroxenite dyke with 3562 ppm TREE and 335 ppm Y (Table 3; Gautneb, 2010). However, one of the hand-picked apatite crystals from the carbonatite contained 11,084 ppm TREE and 5045 ppm Ce which may suggest intergrowth with monazite.

The average grades of 3.0 wt. % P2O5 in both carbonatites and clinopyroxenites suggest that the Lillebukt complex is too low grade to represent any important resource of apatite. In addition, the low to moderate contents of TREE + Y in the apatite comparable to those in the Fen alkaline complex give probably no extra value to the deposit. This resource estimate appears also to be the case for the Tappeluft, Breivikbotn and Pollen deposits.

4.1.3. The Misværدل Complex

The Misværodelist Complex (MC) comprises multiple intrusions of feldspar and apatite. Four types of clinopyroxenites can be distinguished (Fig. 6). Major element analyses reveal ultrapotassic bulk compositions (i.e. molar ratio of K2O/Na2O > 3) for most types due to high contents of biotite and alkali feldspar. The most voluminous of these is the fine- to medium-grained biotite clinopyroxenites (termed fine-grained pyroxenites in the text below) which even on an outcrop scale may grade into a number of sub-types. One of these are characterized by dense dissemination and irregular segregations, lenses and veinlets of coarse-grained, often monomineralic aggregates of biotite. This glimmeritic type is frequently rich in apatite especially along the gradational contacts with coarse-grained to pegmatic segregation which are composed of acicular clinopyroxene crystals (1–10 cm long) in a matrix of up to 80% meso- and phyllosilicate alkali feldspar together with apatite and biotite. These up to several tens of metres sized segregations, possibly representing residual liquids, are similar in composition and appearance to thin dykes and major intrusive bodies of coarse-grained apatite–biotite–alkali feldspar clinopyroxenites (Fig. 5b; termed as coarse-grained pyroxenites in the text below) cutting the fine-grained pyroxenites in different directions (Fig. 6). The coarse-grained pyroxenites are commonly only weakly deformed and retrogradated and have largely retained their igneous texture and mineralogy. In some of them alkali feldspar is missing and this sub-type is generally rich in biotite (20–60 vol.%), but low in apatite. The biotite-rich rocks are probably of metasomatic origin assumed to be caused by volatiles expelled from crystalising magmas of coarse-grained pyroxenites or from carbonatites at depth. All of the clinopyroxenites are cross-cut by abundant thin dykes of coarse-grained to pegmatic alkali feldspar syenites and monzonites, as well as some small dykes of calcite-carbonatites. The fine-grained pyroxenites often form globular mingling structures with the monzonites that are commonly encountered in the southern satellite intrusion where a number of intermediate rock types are developed.

The mineral assemblage of the clinopyroxenites (modal and normative) is characterized by the absence of foids, indicating that the magmas were silica saturated. Subordinate biotite-free clinopyroxenites have a sub-alkaline bulk composition whereas those containing biotite and/or alkali feldspar are alkaline and ultrapotassic. The composition of the rocks fits none of the common classification schemes for mafic–ultramafic igneous rocks given by Le Maitre (2002). The silica-saturation, the frequent mingling structures with felsic intrusions and low contents of TiO2 = 0.99 ± 0.28 (normally titanite, rarely ilmenomagnetite) suggest a link to the temporally associated calc-alkaline and alkali-calcic granitic plutons. One possibility is that the rocks formed by differentiation of a magma generated by mixing of granitic, and phosphorus-rich lamproitic or shoshonitic magmas.

### 4.1.3.1. Geological outline

The MC comprises multiple intrusions of clinopyroxenites forming two bodies measuring 6 km2 and 2 km2, respectively: Breivikbotn and Pollen deposits. This resource estimate appears also to be the case for the Tappeluft, Breivikbotn and Pollen deposits.

**Table 3**

| DEPOSIT   | HOST ROCK               | La   | Ce   | Pr   | Nd   | Sm   | Eu   | Gd   | Tb   | Dy   | Ho   | Er   | Tm   | Yb   | Lu   | TREE | Y   | Th   |
|-----------|-------------------------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|
| Lillebukt | Bio. calcite-carb.      | 1052.7 | 2501.2 | 295.3 | 1195.9 | 194.9 | 57.8 | 158.1 | 18.0 | 78.0 | 11.1 | 23.2 | 2.4  | 11.4 | 1.3  | 5601.2 | 287.8 | 0.5  |
|           | Bio. calcite-carb.      | 2258.3 | 5045.5 | 578.6 | 2255.3 | 310.9 | 160 | 258.3 | 29.2 | 128.3 | 18.6 | 38.1 | 4.0  | 20.8 | 2.3  | 11,064.1 | 485.6 | 0.2  |
|           | Hbl. pyroxenite         | 842.5 | 1648.5 | 196.7 | 676.8 | 82.4 | 21.5 | 62.3 | 6.7  | 32.5 | 4.8  | 6.7  | 2.7  | 0.9  | 1.1  | 3561.7 | 131.8 | 50.2 |
| Misværدليل | Fgr. pyroxenite         | 486.5 | 1506.2 | 248.5 | 1277.1 | 269.6 | 59.8 | 107.4 | 20.7 | 129.5 | 29.9 | 71.8 | 8.6  | 47.4 | 6.1  | 6364.0 | 818.4 | 9.5  |
|           | Cgr. pyroxenite         | 638.6 | 1576.9 | 233.4 | 1074.9 | 208.7 | 45.8 | 160.7 | 16.1 | 67.6 | 8.7  | 16.9 | 2.1  | 12.0 | 1.4  | 4074.7 | 253.0 | 62.8 |
|           | Cgr. pyroxenite         | 456.3 | 1457.7 | 226.6 | 1112.6 | 208.7 | 50.2 | 183.1 | 20.0 | 85.0 | 12.1 | 26.1 | 3.1  | 16.7 | 2.3  | 3882.9 | 334.8 | 60.1 |
| N. Fjeldåد | Nelsonite              | 651.6 | 1248.9 | 236.9 | 1224.9 | 286.5 | 10.4 | 304.7 | 45.4 | 279.2 | 52.9 | 138.8 | 18.9 | 103.3 | 13.2 | 4615.6 | 1394.0 | 51.1 |
| Grindvikåد | Pyroxenite ore         | 946.3 | 2411.3 | 1688.8 | 3130.0 | 2741.1 | 333.7 | 1703.0 | 299.7 | 71.8 | 8.6  | 47.4 | 6.1  | 63640.0 | 818.4 | 9.5  |
|           | Pyroxenite ore         | 1554.6 | 3381.8 | 422.4 | 1729.1 | 263.2 | 52.3 | 228.7 | 26.4 | 129.5 | 23.0 | 57.0 | 6.9  | 38.1 | 5.1  | 7958.1 | 630.1 | 44.1 |
| Kodal     | Pyroxenite ore, h.w.   | 2084.2 | 4366.3 | 524.6 | 2595.3 | 314.5 | 45.0 | 267.8 | 30.4 | 152.2 | 26.6 | 64.4 | 6.1  | 46.6 | 6.0  | 9597.8 | 737.4 | 44.2 |
| Rossavika | Carbonate lens         | 744.4 | 16119.2 | 219.7 | 905.6 | 123.0 | 28.7 | 90.2 | 8.7  | 39.5 | 6.0  | 13.9 | 1.8  | 10.9 | 1.4  | 3805.6 | 183.7 | 22.7 |

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*[P.M. Ihlen et al. / Ore Geology Reviews 58 (2014) 126–147]*
contain apatite as 1–30 mm long acicular crystals which frequently have recrystallised to needle-shaped granular aggregates of fine-grained apatite. The apatite represents an early liquidus mineral which locally segregates to form cumulates with interstitial biotite and alkali feldspar. The cumulus-forming process may explain the very uneven distribution of apatite between outcrops and within individual outcrops (Ihlen and Furuhaug, 2012). The total average for 283 clinopyroxenite samples is 2.38 wt.% P$_2$O$_5$. The highest values are found associated with the coarse-grained pyroxenite segregations and their intrusive counterparts which both contain averages of 2.5 wt.% P$_2$O$_5$ or about 6 wt.% apatite. In spite of these rather low values, presently of no economic interest, it has been possible so far to define a zone, 200 m wide and more than 650 m long, in the western part of the complex where sampled outcrops of coarse-grained pyroxenite yield an average of 4.1 wt.% P$_2$O$_5$ or about 10 wt.% apatite.

Laser ablation ICP-MS analyses of single grains of apatite in polished sections of coarse-grained pyroxenites yield TREE in the range of 1243–11,180 ppm with averages in the range of 4150–6353 ppm TREE for the individual sections (Table 4). The large variations in TREE may be caused by the retrograde formation of allanite frequently forming micro-networks in the apatite aggregates and possibly extracting variable amounts of REE from the apatite. ICP-MS analyses of apatite concentrates from a fine-grained pyroxenite gave 4270 ppm TREE, whereas concentrates from coarse-grained pyroxenites yield values in the range of 3883–4075 ppm TREE (Table 3). The latter range is considerably lower than the averages of the LA-ICP-MS analyses from the same samples, i.e. 3883 ppm TREE (ICP-MS) versus 5581 ppm TREE (LA-ICP-MS). Such interlaboratory and intermethod differences are not uncommon. Thus, a direct comparison of analytical values for apatite should be made with caution as long as the analytical method is not known. However, both methods yield high average contents of Th reaching 100 ppm in some samples which is about the double of the amount accepted for raw materials to NPK-fertilizers. This is the main obstacle regarding the utilization of apatite from the MC.

Fig. 6. Simplified geological map of the Misværdal Complex based on mapping by the first author. Black areas and dots represent outcrops. Numbers on the frame are kilometre units for UTM coordinates (EUREF 89) in grid zone 33 W. 50 m altitude contours. Lake Skardsvatnet is situated at 162 m above sea level. Roads shown as red lines.
et al. (2011) was emplaced during two short lived plutonic events orogeny and according to Schärer et al. (1996) and Vander Auwera et al. (1999; Schiellerup et al., 2000; Vander Auwera et al., 2011). The last plutonic event terminating the magmatic activity in the RAP comprises charnockitic to mangeritic intrusions and dykes of jotunitic to noritic compositions — including the world class Tellnes ilmenite deposit (Schärer et al., 1996; Vander Auwera et al., 2011). Genetic models tend to involve primitive jotunitic to high alumina basaltic parental melts subjected to a polybaric differentiation history to account for all the various units in the province (Charlier et al., 2010; Duchesne and Hertogen, 1988; Longhi et al., 1999; Robins et al., 1997). A lower crustal source for these magmas has been advocated by a number of authors. (Duchesne et al., 1999; Longhi, 2005; Longhi et al., 1999; Schiellerup et al., 2000; Vander Auwera et al., 2011).

Apatite is present in most intermediary and evolved lithologies such as titaniferous jotunites and gabbronorites. Jotunite dykes of variable size and form transect both the anorthosite masses and other major intrusions, such as the BKSK. They represent the youngest intrusive event in the province. The Lomland and Varberg dykes intersecting the Egersund–Ogna anorthosite massif are reported to contain 2.99 wt.% P₂O₅ and 4.05 wt.% P₂O₅, respectively (Duchesne and Schiellerup, 2001).

Jotunite enriched in phosphorus also occurs in a more primitive form as marginal chills to both the BKSK and the Hidra massif anorthosite, as well as in marginal rocks of the Tellnes intrusion. Apatite-bearing gabbronitites found higher in the stratigraphy of the BKSK are invariably more evolved cumulates saturated in multiple minerals. The RAP also contains a large variety and number of Fe–Ti deposits and occurrences (Schiellerup et al., 2003) and some of these are also enriched in apatite (e.g. Hestenes; Fig. 7). The apatite-bearing types include narrow nelsonite dykes and lenses intersecting the northern sheared margin of the Håland–Helleren anorthosite massif (Fig. 7). The nelsonite dykes are almost devoid of silicates and consist of fairly equal amounts of apatite and Fe–Ti oxides (Duchesne and Schiellerup, 2001).

4.2.1. Deposits in massif-type anorthosite complexes

4.2.1.1. Deposits in the Bjerkreim–Sokndal Layered Intrusion. The most prominent of the apatite deposits is hosted by the Bjerkreim–Sokndal Layered Intrusion (BKSK). This intrusion contains a more than 7000 m thick Layered Series consisting of anorthosite, leucorutile, leucotroctolite, norite and gabbronit (e.g., Wilson et al., 1996; Fig. 8). The Layered Series is overlain by a jotunitic Transition Zone and finally capped by increasingly monotonous mangerites, quartz-mangerites and charnockites where apatite mineralisation appears to be missing. Fine to medium grained jotunite rocks found along the margins of the intrusion have

The RAP is dominated volumetrically by a number of massif-type anorthosite plutons consisting of more or less monotonous anorthosites or leucorutiles. The somewhat younger Bjerkreim–Sokndal Layered Intrusion (BKSK) measuring 230 km², is considered to be the largest layered intrusion in Western Europe, and comprises cumulates fractionated from a jotunitic magma (Duchesne and Hertogen, 1988; Robins et al., 1997). The last plutonic event terminating the magmatic activity in the RAP comprises charnockitic to mangeritic intrusions and dykes of jotunitic to noritic compositions — including the world class Tellnes ilmenite deposit (Schärer et al., 1996; Vander Auwera et al., 2011). Genetic models tend to involve primitive jotunitic to high alumina basaltic parental melts subjected to a polybaric differentiation history to account for all the various units in the province (Charlier et al., 2010; Duchesne and Hertogen, 1988; Longhi et al., 1999; Robins et al., 1997). A lower crustal source for these magmas has been advocated by a number of authors. (Duchesne et al., 1999; Longhi, 2005; Longhi et al., 1999; Schiellerup et al., 2000; Vander Auwera et al., 2011).

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been interpreted as marginal chills and suggest a Fe– and Ti-enriched, relatively Ca-poor basaltic parental magma (Duchesne and Hertogen, 1988; Robins et al., 1997). The P₂O₅ content of the fine grained marginal rocks is between 0.6 and 1.1 wt. %.

The intrusion has been deformed into a roughly north–south elongated syncline or trough, at least partly due to gravitational subsidence of the core of the intrusion at a late- to post-magmatic stage (Bolle et al., 2002; Paludan et al., 1994). The cumulate sequence is thickest along the south plunging axis of the northern Bjerkreim-lobe.

The Layered Series are organised into 6 megacyclic units (MCU 0, IA, IB, II, III, and IV) each resulting from recharge of relatively primitive magma. Phase layering is distinct within the megacyclic units reflecting the general order of crystallisation: plagioclase (± olivine), ilmenite, orthopyroxene, magnetite, clinopyroxene and apatite. In addition magnetite is a cumulus phase in olivine-bearing cumulates in MCUs III and IV. The index minerals defining the phase layering are used to divide each MCU into a maximum of 6 cumulate zones (a–f) as outlined in Fig. 8.

Within each megacyclic unit phase layering is accompanied by cryptic layering of mineral compositions with stratigraphic height (Duchesne, 1972; Nielsen and Wilson, 1991; Wilson et al., 1996). The most primitive silicate compositions are found in the leucotroctolitic zones at the base of the MCUs. In the Layered Series the most evolved compositions are found in MCU IV containing antiperthitic plagioclase and inverted pigeonite. The Fe–Ti oxides also display rhythmic cryptic layering with magnetite exhibiting an increasing TiO₂-content and decreasing magnesium, vanadium, nickel and chromium contents with differentiation (Duchesne, 1972; Jensen et al., 1993). The highest-temperature ilmenites are characterized by high hematite contents of up to 19 vol.%, high MgO contents exceeding 4 wt.% as well as elevated chromium and nickel contents (Duchesne, 1972; Duchesne et al., 1999). Due to the strong partitioning of chromium and nickel into the oxides their concentration rapidly fall in both magma and cumulus oxides. MgO in ilmenite gradually decreases to a few tenths of percent in the most evolved cumulates.

Fig. 8. Generalised zone stratigraphy and phase layering of the Layered Series in the Bjerkreim–Sokndal Layered Intrusion in the RAP.
The BKSK represents a very large, but low-grade resource of apatite, ilmenite and magnetite, carrying significant amounts of vanadium. Detailed maps and knowledge of phase layering and mineral chemistry combined with a predictable stratigraphic control on the resources have made it possible to identify three well-constrained zones rich in apatite and oxides. All these zones are located in the northern part of the intrusion, i.e., in the Bjerkreim-lobe (Fig. 9).

In the BKSK apatite is stratigraphically constrained to the e- and f-zones (Fig. 8) which are found in the uppermost part of three of the megacyclic units; MCU IB, MCU III and MCU IV. In these zones, apatite coexists with plagioclase, Ca-poor and Ca-rich pyroxenes, ilmenite and magnetite. The P2O5 content of the Bjerkreim–Sokndal cumulates correlates with MgO, TiO2 and Fe2O3, implying that apatite is more abundant in oxide-rich mafic layers (Fig. 5c). The sequences richest in apatite and oxides have been constrained to the lower parts of the apatite-bearing zones and three sequences are evaluated as potential resources. In these potentially ore-grade zones, the coexisting magnetite typically contains around 0.9 wt.% V2O3 with no apparent stratigraphic variability. Whereas Cr remains low in ilmenite, the MgO-content varies strongly with stratigraphy and the modal composition of the individual layers. Generally the MgO-content decreases as the phase assemblage becomes more evolved, and the MgO-content fluctuates around an average of 1–2 wt.% in the three mineralised zones. EPMA analyses of Fe–Ti oxides in drill cores across the mineralised zone in MCU IV reveal an average of 1.74 wt.% MgO in ilmenite and 0.89 wt.% V2O3 in magnetite. Apatites are generally fluor-apatites with Cl content below 0.1 wt.%. Apatites from each of the zones have been analysed for REE-content by LA-ICP-MS showing a maximum TREE-content of 0.23 wt.% and averages for the individual sections are in the range of 1400–1484 ppm TREE. The apatites are low in thorium with averages in the range of 1–7 ppm Th (Table 4).

The largest apatite and oxide rich zone is found in MCU IV (Fig. 9). It is estimated to be laterally persistent for more than 10 km and is made up of a stratigraphic sequence 50 m to 170 m thick, but is partly covered by lakes. 38 samples from a number of different traverses across the zone give an average composition of 4.1 wt.% P2O5 (Max: 5.5%, Min: 2.4%) and 6.2 wt.% TiO2 (Max: 8.5%, Min: 3.8%). A normative recalcula-
tion yields an average composition of 10.2% apatite, 12.4% ilmenite and 7.3% magnetite giving a total of 32% of the three value minerals. The data have been confirmed by two drill holes through the MCU-IV mineralised zone at Mjåsund and Ollestad where continuous analyses of 30 m drill cores yield an average of 31% and 27% value minerals, respectively. At the same time, modal analyses of large samples for beneficiation testing, collected at various sites in the mineralised zones, corroborate the calculated normative compositions given here.

The apatite-rich zones of MCU IIIe consist of two isolated parts. The southern part has an estimated thickness of 130 m with a lateral extent of approximately 1500 m. The average normative composition from 9 samples in a profile of 120 m gives 13.4% ilmenite (Min 8.9; Max 15.7), 8.3% apatite (Min 4.7; Max 10.3) and 8% magnetite (Min 5.6; Max 11.5). The northern part of the zone (MCU IIIe) is represented by two profiles 2 km apart. The maximum thickness is 120 m, thinning towards the east to 90 m. The average normative composition from 18 samples in this part of the prospect is 11.4% ilmenite (Min 4.3; Max 17.5), 7.8% apatite (Min 3.8; Max 12.7) and 6.9% magnetite (Min 2.1; Max 11).

The apatite bearing part of MCU IBe forms a 3 km long zone with a maximum thickness of 60 m gradually thinning out at the flanks. The potential extent of the mineralised sequence is poorly documented due to extensive quaternary cover and a challenging topography. The sequence is, however, interesting in terms of the magmatic evolution because the entry of cumulus magnetite is delayed with respect to MCUs III and IV, and magnetite appears at the same time as apatite. Presumably as a result of this the Fe2O3 contents of the apatite-bearing cumulates are higher than for the equivalent sequences in MCUs III and IV. Based on XRF whole rock analyses from 10 samples in two profiles of...
40–50 m each, the average normative abundances of the three minerals have been calculated to 8.3% apatite, 15.2% ilmenite and 10.6% magnetite.

In the Bjerkreim–Sokndal Layered Intrusion the apatite-rich zone of MCU IV is particularly interesting with a considerable exposure of cumulates containing more than 10% normative apatite and 20% Fe–Ti oxides. Future development of the Layered Series as an apatite resource may, however, depend on parallel sales in the widely different market regimes of phosphate, titanium and Fe–V.

4.3. Deposits in monzonitic complexes

Enrichment of apatite in association with Fe–Ti oxides typical for the RAP is also a characteristic for members of voluminous plutonic complexes of monzonitic composition developed in both the lower and upper crusts. Thus, they can be subdivided into complexes characterized either by abundant mangerite intrusions or by monzonites. Monzonitic complexes crystallised under granulite facies conditions are found in the Palaeoproterozoic LVMC of northern Norway (Malm and Ormaasen, 1978) and in the late Palaeoproterozoic to early Neoproterozoic mangeritic and jotunitic rocks of the JVNC and MBA of the Caledonides in southern Norway (Fig. 1; Roffeis and Corfu, 2013, and references therein). The apatite occurrences in the two latter areas are poorly known, but appear similar to those in the LVMC and include patchy ultramafic ores and pyroxenitic ores (types given below) with 2–11 wt.% P₂O₅ (Neumann, 1985; NGU Ore Database). The LVMC was emplaced during two distinct events at 1870–1860 Ma and 1800–1790 Ma (Corfu, 2004). Most of the apatite deposits are temporally related to quartz normative mangerites of the second and most important magmatic event which represents according to Markl et al. (1998) and Corfu (2004) extraction of magma from the subcontinental mantle and consolidation at a depth of about 15 km (0.4 GPa). According to major element analyses given by Malm and Ormaasen (1978) the mangerites contain an average of about 0.4 wt.% P₂O₅. The mangerites of the second event are intruded by alkali basaltic dolerites containing 1.4–2.4 wt.% P₂O₅ (Misra and Griffin, 1972). Their relationship to the mangerite magmatism is, however, unclear.

The Oslo Igneous Province (OIP) comprises a number of intrusions with enhanced phosphorus contents, including olivine tholeiitic gabbros, monzodiorites, alkaline troctolites, monzonites and nepheline-rich syenomonzonites. These yield ages in the range 297–256 Ma (Dahlgren et al., 1998; Sundvoll et al., 1990). The plutons crystallised at a depth of 3–4 km below the surface of the former lava plateau, i.e. –0.1 GPa lithostatic pressure. The monzonitic plutons of the OIP are more enriched in phosphorus in comparison with the mangerites of the LVMC by yielding an average of 0.63 ± 0.25 wt.% P₂O₅ (Neumann, 1978). There is also a general enrichment of phosphorus in rocks associated with the monzonites which contain small rafts and larger bodies of monzodiorite and
alkaline troctolite (e.g. at Rånerød, Fig. 10), possibly representing cumulates with 1.6–4.7 wt.% P₂O₅ (Brogger, 1934; Barth, 1945; Ihlen, unpublished data). Late stage monzonite dykes in the Larvik Plutonic Complex (Fig. 10) commonly contain more than 1 wt.% P₂O₅ (Ihlen, unpublished data). This is the normal level for the nepheline-rich olivine monzonites and syenomonzonites (lardalites) intruding the monzonite massifs in the west where they locally may reach 2 wt.% P₂O₅ (Bergstøl, 1972; Ofstedal and Petersen, 1978).

4.3.1. Apatite–Fe–Ti ores in the Lofoten–Vesteralen Mangerite Complex

The description of the apatite accumulations in the LVMC is based on the results from an on-going NGU project. They are characterized by large quantities of Fe–Ti oxides comprised by ilmenomagnetite, titanomagnetite and subordinate amounts of ilmenite and hemoilmenite that together far exceeds the amount of apatite. The apatite–Fe–Ti ores are often diluted by silicate gangue comprised by feldspars, clinopyroxene, orthopyroxene, biotite and/or minor zircon and titanite. Three major types of ores can be distinguished which are termed 1) nelsonite, 2) patchy ultramafic, and 3) pyroxenitic. The latter two are also found in the JVNC and OIP.

Nelsonite according to the definition by Philpotts (1967) occurs in the LVMC as decimetre to several metre wide veins and breccia dykes intersecting the mangerites and their granulate facies supracrustal wall rocks. The cement of the intrusion breccia at Nordre Foslåt (no. 5, Figs. 1 and 5d) are composed of about 85 vol.% Fe–Ti oxides carrying inclusions and interstitial crystals of fine-grained apatite together with minor biotite and zircon. Monomineralic aggregates of biotite replace the wall-rock feldspars and form a thin veneer separating the Fe–Ti–Mg melt interacting with the feldspatic granulate wall rocks. Two of the nelsonite deposits in the LVMC yield analytical values in the range of 2.75–4.23 wt.% P₂O₅. The nelsonite ores are also enriched in Y and TREY with whole-rock values in the range of 120–229 ppm Y and 984–2329 ppm TREY. Apatite concentrates extracted from the Nordre Foslåt ores contain 4616 ppm TREY and 1394 ppm Y (Table 3). The high enrichment of Zr is a characteristic signature for the type of deposit reaching a maximum of 4345 ppm Zr. Bodies of very biotite-rich (50 vol.%) nelsonite ores are locally encountered containing up to 0.9 wt.% Zr, but only 1–2 wt.% P₂O₅.

Patchy ultramafic ore is the most widespread ore type in the mangerites. The mafic minerals filling the interstices between the cm-sized feldspar crystals in the mangerites frequently segregate into irregular ultramafic patches composed mainly of pyroxenes, Fe–Ti oxides, biotite and apatite. The metre-sized segregations of small irregular patches, stringers and schlieren of semi-massive Fe–Ti–P ore grade into the host-rock mangerites via thin zones with intergranular networks (Fig. 5e). These ultramafic segregations have an uneven patchy distribution in up to 100 m wide and more than kilometre long linear zones, frequently close to the border of the individual monzonite plutons. Although the individual semi-massive patches may reach 4.5 wt.% P₂O₅, most of the bulk ores are of low grade and rarely exceed 3 wt.% P₂O₅ due to the common dilution by porphyritic feldspar aggregates.

Pyroxenitic ore is represented by pyroxene-dominated ultramafites (<40% SiO₂) forming bodies composed mainly of fine-grained apatite and Fe–Ti oxides intergrown with pyroxene, biotite and olivine, as well as variable amounts of porphyritic plagioclase and/or alkali feldspar aggregates. These ore bodies may represent cumulate rocks, although their irregular contacts with multiple branches extending into the mangerite host may suggest that they formed as separate intrusions. The ores at Grindvik and adjacent areas (no. 4, Fig. 1) define an en echelon array of ore zones, each of them consisting of up to 20 m wide and several hundred metres long zones composed either of a single dyke-like body or of densely spaced lenses, each several metres wide and tens of metres long. Reconnaissance grab-sampling of these ore zones yield analytical values in the range of 2.0–5.5 wt.% P₂O₅. Apatite concentrates from the Grindvik ore zone contain 6364 ppm TREY and 630 ppm Y (Table 3). The largest of the mangerite-hosted pyroxenite deposits occurs at Utaker (no. 6) where a 350 m by 700 m and crescent-shaped magnetite-olivine pyroxenite is exposed along the beach. However, only two grab-samples have been analysed so far yielding 3.3–3.4 wt.% P₂O₅ (Malm and Ormaasen, 1978; A. Korneliussen, pers. com., 2013).

The existence of potential resources of apatite in the mangerites of the LVMC, JVNC and MBA has just recently been recognized and our knowledge is presently rather immature. Thus it is difficult to make an assessment of their potential based on a restricted number of analysed samples.

4.3.2. Apatite–Fe–Ti ores in the Larvik Plutonic Complex

The monzonites of the Larvik Plutonic Complex consist of coarse-grained augite ± biotite ± olivine monzonites frequently containing larkivite zones where cryptoperthitic alkali feldspar with schiller effect occurs. The common alignment of the usually rhomb-shaped feldspars frequently generates a linear fabric in the monzonites that parallels the outer border of the individual plutons. The Larvik Plutonic Complex comprises a series of crescent-shaped monzonite intrusions formed in response to plutonic centres moving successively from the east towards the west (Fig. 10). Concurrently, the monzonite magmas changed composition from quartz normative in the east to nepheline normative in the western half of the complex.

All of the known occurrences of apatite–Fe–Ti ores in the monzonites of the Larvik Plutonic Complex occur inside nepheline-normative monzonites and syenomonzonites in contrast to quartz normative mangerites in the LVMC. The monzonite and olivine–nepheline monzonite (lardalite) intrusions both contain patchy ultramafic ores similar to those in the LVMC. These are invariably found close to the contact of the individual monzonite plutons. Phosphate grades of the patchy ores are similar to those in the LVMC, i.e. generally of low-grade, but with samples of the interior of the semi-massive patches yielding 4.50 wt.% P₂O₅ as in the Kjose zone (Fig. 10; Brogger, 1934). Tabular semimassive bodies of apatite-rich magnetite clinopyroxenites previously termed jacupirangite by Brogger (1934) and Bergstøl (1972) are encountered in several places e.g. at Kvelde where ores contain 4.85 wt.% P₂O₅ (Fig. 10). However, the most prominent of these are the magnetite pyroxenite bodies constituting the Kodal apatite–Fe–Ti oxide deposit (Fig. 11).

4.3.2.1. The Kodal deposit.

The Kodal deposit was investigated by Norsk Hydro due to its potential as an apatite raw material for the company's fertilizer plants. Between the years 1959 and 1984 the company conducted three exploration campaigns including 58 drill holes (18 mm), beneficiation tests and apatite resource estimations. Renewed interest for the deposit has recently been taken by Kodal Minerals Ltd. which presently is conducting systematic core drilling according to JORC/ PERC standards and advanced ore dressing tests.

The deposit is comprised of a 1900 m long ore zone of closely spaced lenses of massive pyroxenitic ores, 18–20 m thick, that have been drilled to a vertical depth of 150 m. The individual ore lenses show sharp contacts with the monzonites in the foot wall and gradational contacts with the mangerites in the hanging wall (Lindberg, 1985). The greyish monzonites, commonly altered to pink varieties along fracture zones, are composed of cm-sized perthitic alkali feldspar and oligo-clase, some antiperthitic, which carry interstitial aggregates of augite, hornblende, apatite, Fe–Ti oxides rimmed by biotite, and occasional grains of orthopyroxene (Starmer, 1974). The apatite represents an early liquidus mineral occurring both as scattered inclusions in the feldspar crystals and as densely spaced prisms enclosed by the minerals of the mafic aggregates where it often exceeds 10 vol.%. The ore zone terminates against a younger intrusion of alkali feldspar syenite in the west, whereas it tapers off in a series of small
en echelon lenses in the east. It is enveloped by a transitional zone up to 200 m wide of patchy ores (Fig. 11) composed of widely distributed lenses and more irregularly shaped pods of massive ore that have dimensions from cm to several decametre. The lenses increase in abundance when approaching the main ore zone (Lindberg, 1985). The ore zone shows small fault displacements and is cross-cut by dykes of alkali-feldspar syenite, trachyandesitic porphyry (RP) and camptonite (Fig. 11).

The massive pyroxenitic ores which contain small patches of monzonites are fine-grained with anhedral to subhedral clinopyroxene grains in a matrix of ilmenite, magnetite and apatite, the latter forming 0.1–0.5 mm long prisms and needles occurring interstitially to the Fe–Ti oxides (Bergstøl, 1972). Modal analyses of the ores show the following variations: 15–24% apatite (average 17%), 25–60% ilmenomagnetite and titanomagnetite, 5–15% high-Mg ilmenite, 20–40% diopside, and 3–10% phlogopite, magnesiohastingsitic hornblende, olivine, feldspar, nepheline and carbonates (Andersen and Seiersten, 1994; Bergstøl, 1972; Lindberg, 1985). Biotite is especially enriched along the contact zone and titanomagnetite, 5–10% phlogopite, magnesiohastingsitic hornblende, olivine, feldspar, nepheline and carbonates (Andersen and Seiersten, 1994; Bergstøl, 1972; Lindberg, 1985). Biotite is especially enriched along the contact zone and titanomagnetite, 5–10% phlogopite, magnesiohastingsitic hornblende, olivine, feldspar, nepheline and carbonates (Andersen and Seiersten, 1994; Bergstøl, 1972; Lindberg, 1985). Biotite is especially enriched along the contact zone and titanomagnetite, 5–10% phlogopite, magnesiohastingsitic hornblende, olivine, feldspar, nepheline and carbonates (Andersen and Seiersten, 1994; Bergstøl, 1972; Lindberg, 1985). Biotite is especially enriched along the contact zone and titanomagnetite, 5–10% phlogopite, magnesiohastingsitic hornblende, olivine, feldspar, nepheline and carbonates (Andersen and Seiersten, 1994; Bergstøl, 1972; Lindberg, 1985). Biotite is especially enriched along the contact zone and titanomagnetite, 5–10% phlogopite, magnesiohastingsitic hornblende, olivine, feldspar, nepheline and carbonates (Andersen and Seiersten, 1994; Bergstøl, 1972; Lindberg, 1985). Biotite is especially enriched along the contact zone and titanomagnetite, 5–10% phlogopite, magnesiohastingsitic hornblende, olivine, feldspar, nepheline and carbonates (Andersen and Seiersten, 1994; Bergstøl, 1972; Lindberg, 1985). Biotite is especially enriched along the contact zone and titanomagnetite, 5–10% phlogopite, magnesiohastingsitic hornblende, olivine, feldspar, nepheline and carbonates (Andersen and Seiersten, 1994; Bergstøl, 1972; Lindberg, 1985). Biotite is especially enriched along the contact zone and titanomagnetite, 5–10% phlogopite, magnesiohastingsitic hornblende, olivine, feldspar, nepheline and carbonates (Andersen and Seiersten, 1994; Bergstøl, 1972; Lindberg, 1985).

Bergstøl (1972) concluded that the deposit was formed by intrusion of an immiscible Fe–Ti–P melt containing cumulus crystals of clinopyroxene that was emplaced into a not yet fully consolidated monzonite, whereas Lindberg (1985) suggested formation by in situ fractional crystallisation of a monzonitic parent magma. The latest model was given by Andersen and Seiersten (1994) who envisage that the accumulation of apatite and ultramafic cumulates originally occurred in an ijolitic magma in the middle or lower crust. Later these cumulates were captured by ascending monzonite magma and brought to their present position where they re-settled to form semi-continuous layers of cumulate minerals. The present authors prefer the latter model although we have some problems on fully understanding how a large mass of high density cumulate minerals can be transported by a low-density magma from the lower or middle crust to near surface environments.

The Kodal deposit has according to the numbers given by Lindberg (1985) inferred open pit resources of about 70 Mt with an average content of 4.9 wt.% P2O5 (massive ore + transition zone) or alternatively underground resources of 35 Mt with an average of 6.8 wt.% P2O5 (massive ore). The apatite in the massive ores of the Kodal deposit shows Th levels acceptable for NKP fertilizer production (Tables 3 and 4), whereas the high TREY contents of about 1 wt.% may represent a potential extra value, if extracted.

5. Vein deposits

This type of apatite deposit is only found inside the Sveconorwegian Mobile Belt (SMB). Geological evidence indicates that they developed episodically during the evolution of the Sveconorwegian orogeny (1140–900 Ma) in response to structurally controlled fluid flow.

5.1. Kiruna-type apatite–Fe oxide deposits

Magnetite–hematite deposits occur in several places in the Nissedal area in Telemark (no. 14, Fig. 1) where they comprise massive oxide veins, some of them high in apatite. The host rocks consist largely of amphibolite facies biotite and/or hornblende gneisses representing felsic to intermediate meta-volcanites of assumed early Mesoproterozoic age. The largest of the apatite-bearing deposits at Søftestad was mined for 30 years in the last century (Bugge, 1978). The underground workings followed a linear ore zone composed of three en echelon, 2–5 m thick, with a total strike length of about 500 m. The frequently mylonitic ores are orientated at an angle to the regional structural trend of the country rocks. The ore zone comprises alternating massive bands of magnetite and hematite (ratio 3:2), up to 4 m thick, separated by cm to metre thick gneiss bands with disseminated Fe-oxides (Vogt, 1895). Disseminated and m-thick parallel stringers of fluorapatite are mainly found in the magnetite-dominated ores which contain up to 8 wt.% P2O5 (Bugge, 1978; Vogt, 1895). The ore mined in the past contained 47–50 wt.% Fe and 1.9–2.4 wt.% P2O5 (Aamo, 1957). Ore reserves left in the mine were estimated by Bugge (1978) to be in the order of 0.5 Mt with 55 wt.% Fe and 3.9 wt.% P2O5. Both Vogt (1895) and Bugge (1978) noticed the similarities with the much larger Kiruna-type Fe–P deposits (Gallivare) in northern Sweden.

5.2. Metasomatic deposits

More than hundred small apatite and/or rutile veins are distributed in the Bamble–Lillesand Block (Fig. 1) where they mainly intersect meta-gabbroic rocks affected by regional scale metasomatism including

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**Fig. 11.** Simplified geological map of the Kodal ore zone with drill-hole section redrawn from Lindberg (1985). RP in the legend: Rhomb porphyry = porphyritic latite-trachyandesite.
scapolitisation, albitionisation and/or calc-silicate alteration (Brøgger, 1935; Engvik et al., 2011, and references therein). The apatite veins and lenses are decimetre to several metres thick and rarely exceed 100 m in length. They are composed of variable proportions of biotite, amphibole, clinopyroxene, scapolite, apatite, albite, quartz, rutile, calcite, magnetite, pyrrhotite and titanite (Brøgger and Reusch, 1875; Bugge, 1922). The apatite commonly occurs as coarsely crystalline aggregates and local crystals of up to 1 m in length (Brøgger and Reusch, 1875). Comparable veins are widespread in the Nissedal area where they also intersect the sheared oes of the Sømestad deposit and in the areas surrounding the Rossavika deposit in southwestern Norway (no. 16, Fig. 1).

These veins in the Bamble–Lillesand Block are the only ones that have been mined solely for phosphate in Norway. About 50 vein systems were worked from about 1850 to shortly after the first world war when a total of about 250.000 tonnes of apatite were produced, including 160,000 tonnes from the Ødegården deposit (Bugge, 1978). The Ødegården deposit (no. 13, Fig. 1) occurs in a deformed troctolitic leucogabbro dated to 1149 ± 7 Ma that has been scapolitised over a distance of about 1.5 km (Engvik et al., 2011). Metasomatism is centred on and occurs outwards from a system of parallel and up to metre thick apatite–enstatite–phlogopite veins containing about 30 wt.% apatite (Bugge, 1978). Detailed studies by Engvik et al. (2009, 2011) reveal a gradual decrease in apatite content in successive metasomatic zones away from the veins. The composition of apatite changed from fluor–chlorapatite with 1.9 wt.% Cl in the leucogabbro protolith to chlorapatite with up to 6.8 wt.% Cl in the scapolite metagabbros and apatite–phlogopite veins (Engvik et al., 2009). The chlorapatite shows pseudomorphic replacement by hydroxy-fluorapatite containing µm-sized inclusions of monazite, xenotime and allanite (Engvik et al., 2009; Lieftink et al., 1994). LA-ICP-MS analyses of the different types of apatite show that they all are enriched in Y (1481–1919 ppm) whereas the primary fluor-chlorapatite is somewhat higher in TReE (average 5820 ppm) than the others yielding averages of 5106 ppm TReE in the chlorapatite and 522 ppm TReE in the hydroxy-fluorapatite. The metamorphic overprint on the gabbro intrusion yields a Rb–Sr age of 1068 ± 7 Ma for phlogopite which indicates that the fluid migration at temperatures of ~600–700 °C represents a continuum of the second tectono-thermal event (amphibolite facies) in the Bamble–Lillesand Block (Engvik et al., 2011).

The high-grade paragneisses at Rossavika in SW Norway (no. 16, Fig. 1) host an enigmatic apatite-deposit represented by three tabular and dyke-like carbonate lenses which have been explored in the past by trial workings. The lenses which are oriented roughly parallel to the foliation and metamorphic banding of the wall rocks are 3–5 m thick and have a length of more than 50 m along strike and dip (Ihlen, unpublished data). The contact zone is characterized by abundant veinlets and dissemination of alkali feldspar and biotite in both the carbonate rocks and adjacent gneisses; a feature resembling fenitisation. The medium-grained carbonate lenses carry abundant parallel-oriented coarse crystals of apatite in a matrix of calcite and minor chlorite-altered biotite, alkali feldspar, epidote, quartz and titanite. Three drill samples of the lenses yield 7.63–21.34 wt.% P2O5. Although the calcite lenses look like a deformed carbonate dyke, they are very low in Th (<1 ppm), Sr (<1384 ppm) and Ba (<278 ppm). An apatite concentrate yields 3806 ppm TReE and 184 ppm Y (Table 3). Thus, the chemical signature of the calcite–apatite lenses does not fit the chemical characteristics of magmatic carbonatites. But stable isotope data (δ18O, δ13C) given by Bol (1990) plot within the fields of magmatic carbonatites (Bell, 2005) and outside the fields for high-grade meta-sedimentary marbles of the area. A possible metasomatic origin is supported by the widespread occurrences of small apatite-rich hornblende veins in the surrounding areas (Ihlen, unpublished data).

6. The REE chemistry of apatite

The chondrite normalized patterns for the Fen carbonatites are characterized by flat LREE distribution and comparatively high HREE values resulting in very low La/Yb 3 ratios in the range of 28–58 (Andersen, 1987a,b; Hornig-Kjarsgaard, 1998; Mitchell and Brunfelt, 1975; Möller et al., 1980). Apatite concentrates from the Fen carbonatites display a fairly linear steeply dipping REE pattern with strong enrichment in LREE. The resulting La/Yb 3 lies between 150 and 160 for both samples presented in Fig. 12a. The trends do show a slight concavity indicative of higher MREE over HREE enrichment compared to LREE over MREE. The two samples have parallel patterns and the apatite cumulate is systematically higher in all REEs than the calcite-carbonatite sample also presented. There are no single anomalies in the patterns and cerium is not obviously affected by coprecipitation of pyrochlore.

The patterns for apatite concentrates from the Lillebukt Complex in the SIP are very similar but even more linear. The fractionation among the REEs is a little less than in the Fen apatite and the pattern somewhat flatter, with La/Yb 3 ratios between 90 and 110. Generally the REE content of apatite in the Lillebukt Complex (5601 ppm TReE) is somewhat higher than the maximum value for the Fen Complex (5220 ppm TReE; Hornig-Kjarsgaard, 1998). The apatite concentrate from one of the carbonate lenses in the Rossavika deposit is also linear, but even flatter again with a La/Yb 3 ratio of 69. The apatite of Rossavika is similar to those of the Fen Carbonatites in terms of LREE but more enriched in HREE and there is a tendency towards less fractionation among the LREE. The Misværdal Complex is genetically poorly understood. The REE patterns in collected apatite separates are even flatter than the other alkaline and carbonatitic samples (Fig. 12a). Apatite from fine-grained pyroxenite shows a La/Yb 3 ratio of 22 with a flatter pattern than the apatite-rich coarse-grained pyroxenite (La/Yb 3, ~27–55). However, the most prominent difference in comparison with the alkaline and carbonatitic samples is the lack of, or reversed, fractionation among the LREE. We speculate that this pattern may be related to the secondary growth of strongly LREE-fractionated allanite in the Misværdal Complex. This trend becomes even more apparent when looking at LA-ICP-MS analyses of individual apatite grains (Table 4). Also here there is a generally linear trend with a conspicuous negative La-anomaly and the data now resolve a very poorly defined negative Eu-anomaly (Fig. 13a). The LA-ICP-MS data indicate a more strongly fractionated REE-pattern than the concentrate analyses with La/Yb 3 of 51–87, even though the data are overlapping.

Four samples from three different e-zones in the Bjerkreim–Sokndal Intrusion have also been analysed by LA-ICP-MS. The REE patterns of the cumulus apatites are virtually identical and thus not dependent on the megacyclic unit sampled (Fig. 13b). The apatite trends have very flat LREE with a slightly negative La-anomaly and a moderate Eu-anomaly, resulting from the prior fractionation of cumulus plagioclase. The convex pattern curves have previously been used to model a parental melt for the Bjerkreim–Sokndal Intrusion with a linear fractionation pattern across the complete REE-spectrum, La/Yb 3 of less than 10, no Eu-anomaly and characteristically low in TReE (Duchesne and Demaiffe, 1978). The pattern fits well with the documented patterns found in the fine grained marginal rocks of the intrusion (Duchesne and Hertogen, 1988; Robins et al., 1997).

Apatite concentrates of nelsonite samples from the LVMC and the RAP have very flat and poorly fractionated REE patterns with La/Yb 3 ratios of 6–11 (Fig. 12b). In particular the LREE pattern remains conspicuously flat. All samples have a profound negative Eu-anomaly reflecting the importance of plagioclase fractionation in the magmatic evolution of the anorthositic and mangeritic complexes. The same pattern is evident for the apatite concentrate from the pyroxenitic ore at Grindvik, LVMC, (Fig. 12c) showing a flat LREE pattern, distinct Eu-anomaly and modest fractionation across the REE range (La/Yb 3, 20). The Grindvik pattern is, however, slightly convex. Again the Eu-anomaly reflects prior plagioclase fractionation, and plagioclase is an early cumulus phase of the primary melts in this type of setting.

The concentrates from the apatite-rich pyroxenite of the Kodal deposit display a slightly convex REE-pattern with a distinct Eu-anomaly (Fig. 12c), but there is now a clear fractionation among the LREE. The
The total fractionation across the REE spectrum is higher (La/Yb<sub>n</sub> 40–44). The Eu-anomaly implies a foregoing plagioclase fractionation, which possibly links the Kodal pyroxenites to the formation of the hosting monzonites, or is inherited from a presumed ijolitic parental melt. However, LA-ICP-MS data for the apatite grains in the various zones of the Kodal deposit reveal a distinctly parallel signature for the ore zones and the foot wall and hanging wall monzonites (Table 4; Fig. 13c). The main difference is the TREE content of the apatites and the size of the negative Eu-anomaly. All patterns for both ore zones and host rock are slightly concave with poor fractionation among the HREE, and all have La/Yb<sub>n</sub> between 42 and 73. The apatites from both the massive ore and host monzonites are generally high in REE content and appear to be highest in the ore zone.

7. Conclusion

Norway exhibits a wide spectrum of different genetic types of apatite deposits ranging in age from the Palaeoproterozoic to the Permian.
The sedimentary deposits of phosphorites and stratiform volcanicogen Fe–P ores are generally narrow and of low-grade, and their potential resources are very small and comparable to estimated resources of the numerous vein-type deposits (Kurina-type Fe–P, metasomatic). Major resources of apatite are only found among the igneous deposits comprising alkaline complexes, massif-type anorthosite complexes and monzonitic complexes. The latter two representing apatite–Fe–Ti ores are the most interesting and occur associated with jutonic intrusions in the RAP, mangeritic intrusions in the LVMC, JVNC and MBA and monzonitic intrusions in the Larvik Plutonic Complex of the OIP. The apatite in the monzonite- and mangerite-associated deposits contains extractable levels of TREE (>6000 ppm), which are higher than the normal levels for TREE inapatite from alkaline complexes (3700–6000 ppm) and much higher than the levels for apatite in the BKSK of the RAP (1400–1500 ppm). The BKSK has resources exceeding 300 Mt with average normative contents of 8.3–10.5%apatite, 12.5–15.2% ilmenite and 21.7–23.5% V-magnetite. The Larvik Plutonic Complex hosting the Kodal deposit contains widespread small occurrences of high-REE apatite ores which are comparable to those in the LVMC, JVNC and MBA where detailed data are missing. The Kodal deposit has ore reserves calculated about 1 Mt TREY which may add extra value to the deposit.

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