Apatite Composition and U-Pb Geochronology from the Ptarmigan and Tom Deposits, NWT Canada; Implications for Petrogenesis

Mark W. Richardson1*, Christopher R.M. McFarlane1, David R. Lentz1, Hendrik Falck2

1Department of Earth Sciences, University of New Brunswick, P.O. Box 4400, Fredericton, NB E3B 5A3, Canada
2Department of Industry, Tourism and Investment, Government of the Northwest Territories, Scotia Centre 4th Floor, 5102 50th Avenue, Yellowknife NT X1A 2L9 Canada

*Correspondence: m5b11@unb.ca (M.W.R)
dlentz@unb.ca (D.R.L.); crmm@unb.ca (C.R.M.M.); Hendrik_Falck@gov.nt.ca (H.F)

Abstract:
The Ptarmigan and Tom mesothermal gold deposits are located 10 km to the northeast of the city of Yellowknife, Northwest Territories in northern Canada. Both gold deposits comprise a series of en echelon veins that are hosted within upper greenschist to lower amphibolite facies ~2630 Ma (peak) rocks. Supracrustal units across the craton are intruded by the ca. 2610–2605-Ma granodiorite, tonalite, monzodiorite, quartz diorite, and affiliated rocks of the Concession Suite. Hydrothermal apatite is a common accessory mineral in both mineralized and non-mineralized quartz veins in the metasedimentary host rocks that constitute the Ptarmigan and Tom deposits. This study characterizes and compares turbidite-hosted hydrothermal apatite from the Ptarmigan and Tom deposits, non-mineralized veins adjacent to the ore body, and magmatic apatite from proximal LCT-pegmatites. Using electron probe microanalyses (EPMA), laser ablation inductively coupled plasma-mass spectrometry (LA-ICP-MS), micro-XRF, and cathodoluminescence (CL), the major, minor, and trace element abundances have been quantified and mapped. In addition to utilizing this data to determine if the chemistry of apatite can be used to constrain the source of hydrothermal fluids, the apparent age of the apatite is also evaluated utilizing in situ U-Pb dating.

The distribution and abundance of major, minor, and trace elements from in situ recovered apatite were studied to characterize the nature of mineralizing fluids. Most apatite from mineralized and non-mineralized veins show different Mn, Sr, and Pb contents, as well as chondrite-normalized rare-earth element (REE) and Y abundance patterns. REEs display five unique chondrite-normalized patterns: (1) negative sloped pattern with slight negative Eu anomaly, (2) a flat pattern with a positive Eu anomaly, (3) a positive slope with a negative Eu anomaly, (iv) light rare earth element (LREE) depleted pattern with positive Eu anomaly, and (v) bell-shaped pattern with a negative Eu anomaly. The REE patterns reflect both the source of the auriferous hydrothermal fluids and, perhaps, co-precipitating mineral phases. Apatite from the Ptarmigan vein occurs with both: (1) a flat pattern with a positive Eu anomaly and (2) bell-shaped pattern with a negative Eu anomaly. The bell-shaped and flat patterns typify orogenic gold deposits. Vein-hosted apatite commonly displays compositional zoning with a characteristic yellow cathodoluminescence (CL) emission spectra with darker cores and brighter rims. The cores have lower REE, whereas the rims are notably higher in REE. It is thought that the darker cores in CL images reflect a transition from an early low REE hydrothermal fluid to one enriched in REE. The hydrothermal apatite age of 2585 ± 15 Ma is consistent with the intrusions of the 2605 and 2590 Ma two-mica granites of the Prosperous Suite and associated LCT pegmatites.

The near-concordant 206Pb-corrected data of the LCT pegmatite hosted apatite reveals two clusters of ages. An older population with a Concordia age (n = 4) of 2581 ± 15 Ma, and a younger population with an age (n = 3) of 2519 ± 12 Ma. Furthermore, plotting a regression through all near -
concordant data for the pegmatite hosted apatite hints that metamorphic resetting occurred around ~2200 Ma.

**Keywords:** apatite; rare earth elements; hydrothermal fluids; Ptarmigan Mine;

1. Introduction

The Yellowknife Greenstone Belt (hereafter referred to as YGB) in the Northwest Territories, Canada (Figure 1) hosts a variety of gold occurrences including the world-class Con and Giant mines. The lesser-known Ptarmigan and Tom deposits occur 10 km NW of the city of Yellowknife. The Ptarmigan and Tom gold occurrences consist of en-echelon gold-bearing quartz veins hosted by metasediments and metasiltstones of the Burwash Formation. Supracrustal units across the craton are intruded by the ca. 2610–2605-Ma granodiorite, tonalite, monzodiorite, quartz diorite, and affiliated rocks of the Concession Suite [1]. The Concession Suite plutons have significantly enriched LREEs, high K2O, and overall geochemical affinities that are most like those derived from a subduction-enriched lithosphere that began melting because of post collision delamination [1]. Following this, 2600–2580-Ma granite sheets and plutons derived from crustal melts intruded all the Neoarchean rocks, except the late-orogenic conglomerates (e.g., [1-2]). Prosperous granite-associated LCT-pegmatites occur throughout the study area [3].

Both the margins of the quartz veins and the pegmatites observed at the Ptarmigan and Tom deposits typically have tourmaline-rich rims. Zoning in the pegmatites is simple with a quartz core, an intermediate zone of plagioclase, muscovite, tourmaline, and an outer finer grained zone (cf. [5]). Additionally, M. Seigel [6] found that tourmaline needles from the Prosperous Granite, Ptarmigan Mine, and the Con Mine are likely related. During mapping of Ross Lake area (25 km east of Ptarmigan Mine) a continuum between pegmatites and quartz veins was observed [7].

Turbidite-hosted gold deposits occur world-wide (e.g., [8,9]). However, the tectonic setting, timing, and nature of veining and source reservoirs has led to a variety of genetic models (i.e., magmatic versus metamorphic versus meteoric (e.g., [9,10]). Rare earth elements and yttrium (REY)
compositions of accessory hydrothermal minerals have been shown to record the chemical make-up of ore-bearing hydrothermal fluids (i.e., apatite, [11-13]). The geochemical characteristics of hydrothermal fluids may help elucidate key differences between the proposed models.

Apatite Ca₅(PO₄)₃(F,Cl,OH) is a common calcium phosphate accessory mineral in igneous, sedimentary, and metamorphic rocks, and can also precipitate from hydrothermal fluids [14,15]. Apatite lattice can accommodate several cation substitutions through its ninefold M₁ and sevenfold M₂ polyhedral sites (also called the Ca₁ and Ca₂ sites). These substitutions include many divalent and trivalent cations, such as Na, Mg, Mn, Fe, Sr, Y, Ba, lanthanides (rare earth elements hereafter referred to as REE), Pb, Th, and U [16,17]. The variety of substitution possibilities has made apatite highly susceptible to various fluid induced (metasomatic) chemical and textural changes over a wide range of pressures and temperatures [18].

In hydrothermal fluids the apatite four-component system (Ca, P, O, and F) can form various aqueous complexes. The phosphate ion has a stepwise association with H⁺ to form the species H₄PO₄, H₃PO₄, and H₂PO₄. Calcium can also form complexes, such as CaCl₂, CaOH⁺, CaPO₄, CaH₂PO₄, and CaHPO₄ and lastly, F⁻ may complex with H⁺ to form HF. For example, it has been demonstrated that pressure, temperature, and dissolved silicates have limited influence on the solubility of apatite, whereas lower pH increases the solubility of apatite [19]. The occurrence of large apatite crystals in hydrothermal veins reflect the increased concentrations of mobile Ca, P, O, and F.

Since rare-earth elements (REE) are hard cations, Pearson’s rules [20] implies that they will bond preferentially with hard anions, such as F⁻, CO₃²⁻, O₂⁻, P₂O₇⁻, and SO₄²⁻. Therefore, fluids saturated in hard ligands will facilitate greater REE mobility in a hydrothermal solution. Thus, REE abundance in hydrothermal apatite is correlated to saturation of apatite in the vein and the presence of F⁻, OH⁻, CO₃²⁻, P₂O₇⁻ complexes available. Chloride complexes are typically most effective at mobilization of REE [21]. Additionally, they demonstrated that the interaction of saline hydrothermal fluid with primary magmatic accessory minerals enriched in REE will preferentially mobilize LREEs [21].

Trace element substitutions in natural apatites can be high; for example, Sr and Mn can substitute up to 8.96 and 7.59 wt.%, respectively [22]. The abundance of trace element substitution is dependent on many factors. For example, Mn²⁺ was observed to substitute into apatite more readily than Mn³⁺, so it was concluded that redox conditions have significant influence on the partitioning of Mn into apatite [23].

Apatite geochemistry (i.e., trace element abundances) has also been used to discriminate between different types of ore deposits [11] and to identify any influence that the host-rock in an ore-forming environment has on the composition of apatite [24]. It was also found that REE composition of apatite [16] can be used to provide information regarding the characteristics of fluids responsible for metasomatism as they record the nature of melts and fluids from which apatite saturates. Cathodoluminescence (CL) imaging is a particularly effective tool at observing compositional zonation in apatite. CL utilizes an electron beam that initiates inelastic scattering of primary electrons in the crystal lattice. This results in an emission of secondary electrons that lead to a cascade of scattering of more primary electrons. These secondary electrons may excite the valence electrons from the conduction band which recombine with the valence band and creates a photon. Mn²⁺ and REE ions are particularly effective at generating secondary electrons during interaction with an electron beam [25]. The compositional zoning can be further verified via elemental-distribution maps, e.g., micro-XRF elemental maps. Trace element zonation observed in hydrothermal apatite may be used to track fluid composition [26] and possibly aid mineral exploration [11].

In addition to metasomatic susceptibility, apatite is also a powerful geochronometer. For example, as new growth of apatite occurs in response to metasomatic processes, the U-Pb geochronometer will record the time at which slowing Pb diffusion leads to quantitative accumulation of radiogenic Pb (Pb⁴⁺). If metasomatism induces partial to complete loss of radiogenic-Pb (and (or) U) along with changes in trace-element composition, subsequent U-Pb dating will record the time of metasomatism. Th⁴⁺ and U⁴⁺ are the primary tetravalent cations (other than Si) found in apatite (few ppm to thousands of ppm) [11,16,22] and high initial concentrations facilitates apatite U-Pb geochronology, e.g., [27-30]. However, the precision of U–Pb apatite geochronology is lower.
compared to other minerals with high initial U/Pb, because apatite incorporates a substantial amount of common-Pb compared to U when it crystallizes. This leads to moderate to low ratios of $^{238}$U/$^{204}$Pb [27]. The high ratio of common to radiogenic Pb (Pb/ Pb*) presents challenges for apatite U-Pb geochronology and calculated dates (cf. [30]). Although precision on individual analyses can be low, the final age of a regression or concordant cluster can be very good. In natural apatite, the Pb-diffusion closure temperature estimates range between 375° to 600°C [26-29], therefore is useful for mid-range cooling ages typical of magmatic-hydrothermal processes.

This study characterizes and compares turbidite-hosted hydrothermal apatite from the Ptarmigan deposit, non-mineralized veins adjacent to the ore body, and magmatic apatite from proximal LCT-pegmatites. Using electron probe microanalysis (EPMA) and laser ablation inductively coupled plasma-mass spectrometry (LA-ICP-MS), micro-XRF, and cathodoluminescence (CL), the major, minor, and trace element abundances have been quantified and mapped. In addition to utilizing this data to determine if the chemistry of apatite can be used to constrain the source of hydrothermal fluids, the apparent age of the apatite is also evaluated utilizing U-Pb dating.

2. Materials and Methods

Polished (~30 μm) thin sections were cut from samples collected from 50 quartz veins in surface outcrops. These samples were selected from mineralized veins, including the wall rocks immediately adjacent to the veins. Petrographic observations were also made on a 35 x 15 cm cut slab that spanned the entire width of the Ptarmigan vein in a trench located ~350 m NW of the Ptarmigan Mine shaft (Figure 2). Of the 60 samples taken, 16 contained enough apatite for use in this study.

Mineral identification, textural analysis, and imaging were carried out using a petrographic microscope in both transmitted and reflected light on polished thin (~30 μm) sections. Apatite grains were imaged with cathodoluminescence (CL) using a Nuclide Corporation Luminoscope cold-cathode luminescence stage microscope. Micro-XRF EDS maps were completed at the University of New Brunswick utilizing a Bruker M4 Tornado with Rh X-ray tube with polycapillary focusing to 20 μm, dual SDD detectors (130-140 eV resolution). In situ apatite geochronology was completed at the University of New Brunswick via laser ablation-inductively coupled plasma-mass spectrometry (LA-ICP-MS) using a Resonetics M-50-LR Ar-F 193 nm Excimer laser ablation system coupled to an Agilent 7700 quadrupole ICP-MS. The trace element and U-Pb data was collected simultaneously using laser fluence of 5 J/cm², 3 Hz (pulses per second), 60 second ablation time and analyzed with a beam diameter of 60 μm. Since both U-Pb and trace element data were collected simultaneously three standard reference materials standards were used. The MAD apatite standard (REF) was used as a calibration standard with accuracy checked using an in-house apatite from Phalaborwa, South African (assumed age of 2060 Ma) NIST610 glass was used as a primary standard for trace element analysis with an assumed stoichiometric value of 39 wt% Ca as internal standard. Non-apatite mineral phases were filtered during offline data reduction based on anomalous Ca, trace-element intensities in the ablation time-series, and silicate contamination. Before analysis, targeted apatite grains were visually inspected to be free of mineral inclusions. After measurements were obtained, signal versus time intensity graphs of the raw data were scrutinized for possible contamination by inclusions. Of note, apatite grains from several of the samples were beam sensitive, such that some
grains were extensively damaged (pitted out) or destroyed by the LA-ICP-MS beam. As a result, a complete dataset is not available for all the investigated apatite grains. Due to their large size, multiple ablations were done on a single apatite grain. Ablations sites varied from the centre to the rim of the apatite grains. Once the data was obtained, it was reduced offline using Iolite v.3.7 and the UcomPbine DRS of [30].

3. Results

Apatite occurs in most of the variably mineralized quartz veins in the study area. In the studied samples, apatites are commonly fractured, complexly zoned, and form euhedral to anhedral crystals between 30 – 100 μm (Figure 3-4). Larger (>200 μm) apatite grains were observed in the Ptarmigan vein. In many cases, apatite displays a mottled texture due to the presence of sub-micron to nanometre-scale inclusions of pyrite, as well as monazite and tourmaline. Previous researchers have interpreted this as an effect of fluid percolation (e.g., [31]). Cathodoluminescence imaging revealed distinctive changes in the CL-colour from core to rim. Although the host rocks varied from black – siltstones to greywacke, there was no correlation between host rock the measured REY abundance. Apatite commonly is found at the vein margins and accounts for up to 1-5 % of vein material.

3.1. Cathodoluminescence and Micro-XRF Element Variation

The apatite examined in this study exhibits near homogeneous cathodoluminescence (Figure 3-4), with brighter patches in some samples. The apatite taken from the Ptarmigan vein (sample P-14) consist of at least two generations: (1) a darker first-generation apatite that forms much of the grain, and (2) a brighter CL apatite that forms the rim and fills in fractures in early apatite. Variations in CL were confirmed utilizing micro-XRF (µ-XRF) trace-element maps. The micro-XRF maps of apatite from sample P-14 exhibit relative Mn-rich cores and Mn-poor rims. Spot analyses were preformed via LA-ICP-MS on the two generations. The early, darker CL apatite has higher Sr, Mn, Zn, Fe, Pb, Th, and U, and low REE and Y contents. The second-generation apatite is brighter in CL and is associated with low Sr, Mn, Zn, Fe, Pb, and higher REE and Y contents.
Figure 3: Top Left: P-Mn element map of sample P-14 taken from the Ptarmigan vein. The P-Mn element map reveals compositional zonation. Top Right: Cathodoluminescence emission spectra plots intensity vs. wavelength of sample P-14 (dark holes are from LA-ICP-MS spot) Bottom: Reflected light image of apatite grain inclusion in pyrite (sample P-14).

3.2. LA-ICP-MS – trace-element concentrations

3.2.1. REE and yttrium

Rare earth element (REE) patterns for quartz vein-hosted apatite are shown in Figure 4 and apatite from LCT pegmatites are shown in Figure 5. Characteristic chondrite-normalized (CN) patterns of REY, the vein-hosted apatite can be grouped into (5) distinct sub types, based on the values from [32]: type – i apatite exhibits negatively sloped patterns with no Eu anomalies; type – ii apatite exhibits relatively flat patterns with positive Eu anomalies; type – iii apatite have positive slopes with weakly negative Eu anomalies; type – iv apatite exhibit concave up REE patterns (i.e., relatively lower MREEs than LREEs and HREEs) with positive Eu anomalies; and (type – v) apatite exhibit weak concave down patterns (relatively higher MREEs than LREEs and HREEs) and weak negative Eu anomalies. The trace-element data for each sample are presented (see Appendix A). LCT-pegmatite-hosted apatite contains contrasting REE patterns. Sample MR-002 is relatively enriched in MREE producing a strong concave down pattern. Sample MR-003 is characterized by a broadly concave down pattern with a strong negative Eu anomaly. Sample MR-005 has a negatively sloped REE pattern resembling (type – i) apatite. The trace-element data for the pegmatite-hosted apatite are presented (see Appendix B). REY (REE + Y) substitute into apatite via coupled substitution processes. The REE + Y are correlated well with Na (Figure 6; a), which suggests that REE + Y were mainly incorporated via the following (Equation 1) coupled substitution reaction (e.g., [33-34]):

\[(\text{REE + Y})^{3+} + \text{Na}^+ = 2 \text{Ca}^{2+}, \tag{1}\]

Brighter in CL in sample P-14 (associated with low Sr, Mn, Zn, Fe, Pb, and higher REE and Y contents) was found to have relatively flat CN-REE patterns with greater concentrations of REEs + Y and rims with negatively sloped CN patterns and less REEs.

Binary plots (Figure 6) allow for easy comparison of LREE-, MREE-, and HREE-enrichment patterns for both LCT- and vein-hosted apatite. (Gd/Yb)n versus (La/Yb)n biplot (Figure 6b) of all apatite shows a distinct positive correlation. The (La/Sm)n versus (Eu/Eu*) (Equation 2) biplot (Figure 6c) shows a clear negative correlation between REE composition and the magnitude of the Eu anomaly. In detail there are two groups of data that plot along this trendline. In the first group are the type – iii, type – iv, and type – v apatites and the second group contains the LCT pegmatites.
and type – I, and type – ii apatite. Biplots of (La/Yb)$_N$ versus (Eu/Eu$^*$) also show that apatite cluster into two groups (Figure 6d). Apatite from type – iii, type – iv, and type – v plot with lower (La/Yb)$_N$ ratios with respect to type – i, type – ii, and LCT pegmatite-hosted apatite. Biplot of (La/Sm)$_N$ versus (Sm/Yb)$_N$ also cluster the REE apatite data into two separate groups (Figure 6 e). The first group type – iii, type – iv, and type – v apatite. The second grouping contains the LCT-pegmatites, type – i, and type – ii apatite.

**Figure 3:** (A,C,E,G) Cathodoluminescence images of apatite (*type – ii apatite see figure 4) next to (B,D,F,H) Plane polarized light image of apatite with assemblages in veins.
A negative or positive correlation between Ce and Eu relates to the oxidation state [33]. In the studied apatite the δCe anomalies (Equation 2, [35]) are mostly between 0.85 and 1.2 in the studied apatite. Additionally, these Ce anomalies are negatively correlated to δEu anomalies (Equation 3, [35]) (Figure 6f)

\[
\frac{\text{Ce}}{\text{Ce}^*} = \frac{\text{Ce}_{\text{CN}}}{{\sqrt{\text{La}_{\text{CN}} \times \text{Pr}_{\text{CN}}}}},
\]

(2)

\[
\frac{\text{Eu}}{\text{Eu}^*} = \frac{\text{Eu}_{\text{CN}}}{{\sqrt{\text{Sm}_{\text{CN}} \times \text{Gd}_{\text{CN}}}}},
\]

(3)

**Figure 4:** Chondrite-normalized REE and Y diagrams for LA-ICP-MS spot analyses of apatite in this study. The patterns are colour coded based on type (see Results and Discussion section). Chondritic values from [36].
Manganese, Lead, Yttrium, Iron, Strontium

Manganese contents of the examined apatite are between 147 and 7551 ppm. Apatite collected from the Ptarmigan vein have the greatest Mn values (1000 – 7551 ppm) with the exception type-i apatite (collected from a zone of intense quartz hydrofracturing proximal to the mine shaft and the mineralized vein). The Mn concentrations are positively correlated with $^{204}$Pb concentration (Figure 7a) and Sr concentrations (Figure 7b). The concentration of $^{204}$Pb is independent of the U and Th concentration, therefore $^{204}$Pb concentrations are likely to be a better discriminator of fluid source. Additionally, the biplots separate the non-LCT pegmatite apatite and vein-hosted apatite into two distinct clusters. $^{204}$Pb concentrations are greatest in type – i apatite followed by apatite from type – ii mineralized quartz veins. Non-mineralized or barren quartz veins are depleted in both Pb and Sr with respect to type – i and type – ii apatite. Sr concentrations in the LCT pegmatites are lowest in sample MR-003 and highest in sample MR-002. Yttrium concentrations are negatively correlated to the Mn concentrations, except for high Mn apatite (sample P-14) (Figure 7c). Highest Y concentrations are associated with the non-mineralized vein-hosted apatite, whereas the LCT pegmatites contain considerably lower concentrations. Iron concentrations plotted against Mn also cluster into two groups with two distinct trendlines (Figure 7d). The first group includes type – iii, type – iv, and type – v apatite. This group has a strong positive correlation between Fe and Mn concentrations. The second group includes the LCT-pegmatites, type – i, and type – ii apatite; however, the type – i apatite plot slightly above the main cluster (i.e., enriched in Fe). The biplot of Mn versus Eu/Eu* (Figure 7e) shows that apatite from the LCT-pegmatites, type – i, and type - ii have greater concentrations of Mn than type – iii, type – iv, and type – v apatite. Furthermore, Mn concentrations show no correlation with respect to the magnitude of the Eu anomaly. Strontium concentrations in apatite range from 83 to 7300 ppm, with a mean of 1244 ppm and a median of 653
ppm. Apatite from mineralized veins contain the greatest concentrations of Sr (1000 – 3569 ppm), whereas the unmineralized veins contain 100 – 500 ppm Sr. Biplots of Sr versus Eu/Eu* show that Sr concentrations are positively correlated to the magnitude of the Eu anomaly (Figure 7f).

**Figure 6.**

a) ΣREY vs Na ppm. b) (Gb/Yb)_N vs (La/Yb)_N bivariate plots illustrate the compositional differences that reflect the evolution of hydrothermal fluids from one enriched in MREE and depleted in HREE to ones depleted in MREE and enriched in HREE. c) (La/Sm)_N vs. (Eu/Eu*)_cn bivariate plot illustrating the co-evolution of two hydrothermal fluids. Increases in MREE is also observed to correlate with an increase in the Eu anomaly. d) (La/Yb)_N vs. Eu/Eu*cn shows that the LREE/HREE elements plot into four separate clusters. Apatite from samples P-14 (high and low Mn) and P-20 are...
partitioned with the cluster that includes two samples taken from the LCT-pegmatite dikes. d) Biplot of \((\text{La/Yb})_N\) versus \(\text{Eu/Eu}^*\) illustrating the trends of the two clusters of apatite. e) \((\text{La/Sm})_N\) versus \((\text{Sm/Yb})_N\) biplot show how hydrothermal fluids may have evolved along two independent trajectories. f) \((\text{Ce/Ce}^*)\) versus \(\text{Eu/Eu}^*\) indicated that a negative correlation exists.

**Figure 7.** a) Bivariate plot of Mn versus Pb illustrating the positive correlation between Mn and Pb concentrations. Group 1 contains apatite from mineralized and non-mineralized quartz veins, whereas Group 2 contains apatite from the LCT pegmatites. b) Biplot of Mn versus Sr illustrating the positive correlation between Mn and Sr concentrations. Group 1 contains apatite from non-
mineralized quartz veins, whereas Group 2 contains apatite from the LCT pegmatites and mineralized quartz veins. c) Biplot of Y versus Mn illustrating the negative correlation between Y and Mn. Additionally, this plot shows that Mn-rich apatite taken from the Ptarmigan vein plot as a separate cluster likely indicating that Y was sequestered in xenotime. d) Biplot of Mn versus Fe illustrating the positive correlation between Mn and Fe concentrations. Group 1 contains apatite from non-mineralized quartz veins, whereas Group 2 contains apatite from the LCT pegmatites and mineralized quartz veins. e) Biplot of Mn versus Eu/Eu* illustrating that apatite from the LCT-pegmatites, type – i, and type - ii have greater concentrations of Mn than type – iii, type – iv, and type – v apatite. f) Biplots of Sr versus Eu/Eu* show that Sr concentrations are positively correlated to the magnitude of the Eu anomaly.

3.2.3. LA-ICP-MS – Uranium, Thorium, and Lead

U, Th, and Pb can replace Ca in apatite. Although Pb²⁺ can directly substitute for Ca²⁺, the incorporation of Th⁴⁺ and U⁴⁺ cations require a vacancy to maintain charge balance [33]. In the quartz vein-hosted apatite, the U and Th contents are low 0.5 – 10 ppm in comparison to the LCT pegmatites that have concentrations that range from 9 to 180 ppm. It is noted that U and Th contents of apatite taken from the Ptarmigan vein are on average greater than the non-mineralized veins, except for sample X-16. The U and Th concentrations do not show any correlation with ΣREY concentrations (Figure 8). Furthermore, U and Th are not correlated with Y, HREE, or Eu anomaly.

![Figure 8](image_url)

**Figure 8.** Bivariate plot of a) U + Th (ppm) versus Σ REY

3.3 U-Pb in situ Hydrothermal Apatite Geochronology

Uncorrected common-Pb data from the hydrothermal apatite was plotted on a Tera–Wasserburg concordia plot (Figure 9), where ²³⁸U/²⁰⁶Pb is plotted along the x-axis and ²⁰⁷Pb/²⁰⁶Pb is plotted along the y-axis. Tera–Wasserburg concordia plots are used to calculate concordia-intercept ages. The analysis assumes that samples (uncorrected for common Pb) are cogenetic and diverge from concordia because of variable contamination with common-Pb with a restricted ²⁰⁷Pb/²⁰⁶Pb composition. Therefore, a linear regression through the series of discordant points will intercept the concordia at the age of initial crystallization. Theoretically this is true; however, the data is likely to have non-target phases (e.g., inclusions or neighboring grains), or record differential Pb loss that disperses data towards lower ²³⁸U/²⁰⁶Pb values. Therefore, it was necessary to refine the linear
regression to obtain the lowest Mean Square of the Weighted Deviates (MSWD), while using the greatest number of data points possible. After omitting data via weighed residuals and data with large errors, a linear regression yielded an age of 2590 ± 13 Ma (MSWD = 0.94) (n = 88/284) from 10 different samples (P20, P-15, P-14, P-09, S3063, S3052, S3052b, X-16 and B8a/b).

Figure 9. U-Pb apatite isochron for hydrothermal quartz veins. Anchored regression lower intercept at 2585 ± 15 Ma; upper intercept anchored terrestrial common-Pb value at 2600 Ma (1.066) [29]. Error ellipses are 2σ.

3.4 U-Pb in situ LCT-Pegmatite Hosted Apatite Geochronology

The range of LCT-pegmatite apatite ages were plotted on a conventional concordia (Wetherill) diagram. The weighted mean 207Pb/206Pb ages were determined based on 204Pb-corrected isotope ratios. Common-Pb correction utilized the measured 204Pb assuming an initial Pb composition from [36] Pb evolution model. In this model, no assumptions of U/Pb* concordance are made, allowing for identification of concordance of the 204Pb-corrected data [30]. Apatite typically incorporates substantial amounts of common-Pb compared to U when it crystallizes, leading to a low 238U/204Pb ratio. Lead loss is generally regarded as the most important cause for age discordance [37]. Additionally, discordance (normal or reverse) may be attributed to mobility of Pb and U during dynamic recrystallization or dissolution-reprecipitation reactions [27]. Plotting of the LCT-Pegmatite data revealed that several analyses fall above and below concordia, reflecting minor matrix-mismatch and recent Pb loss. By examining the near-concordant 204Pb-corrected data (Figure 10; a) of the LCT pegmatite hosted apatite two clusters of ages emerge. An older population with an intercept age (n = 4) of 2581 ± 15 Ma (Figure 10; b), and a younger population with an intercept age (n = 3) of 2519 ± 12 Ma (Figure 10; c). Furthermore, plotting a regression through all near – concordant data for the pegmatite hosted apatite hints that metamorphic resetting occurred around ~2200 Ma (Figure 10; d). It is likely that detailed imaging using a scanning electron microscopy (SEM) of these apatite grains may reveal older domains.
Figure 10. a) Plots of near-concordant $^{206}\text{Pb}$-corrected data of the LCT pegmatite-hosted apatite illustrating two clusters of ages that emerge; b) LCT pegmatite-hosted apatite focused on the older cluster with a concordant age ($n = 4$) of $2581 \pm 15$ Ma; c) LCT pegmatite-hosted apatite focused on the younger cluster with a concordant age ($n = 3$) of $2519 \pm 12$ Ma; d) Error ellipses are $2\sigma$. Linear regression through all near-concordant data for the pegmatite-hosted apatite hints that metamorphic resetting occurs around $-2200$ Ma.

4. Discussion

Hydrothermal apatite forms large (up to 5 mm) subhedral to euhedral crystals and occurs within most quartz veins in the study area. Petrological observations suggest that apatite is precipitated early, as sulphide-bearing phases are observed to infill fractures and occasionally partially surround apatite grains. Additionally, in the gold-bearing Ptarmigan vein, tourmaline is found as inclusions in apatite.

The spectra observed via cathodoluminescence and micro-XRF mapping revealed compositional variations with respect to trace-elements in most of the examined apatite grains. This spectral variation is explained by compositional zoning in apatite. In order to determine whether the compositional zoning reflects primary elemental distributions and abundances, the REE abundances and zonation were contrasted with the $(\text{La/Sm})_N$ established conditions [38]. Additionally they also noted that since LREEs-MREEs are readily substituted into the Ca(2) site, homogenization of (La/Sm) ratios between compositional zones may elucidate whether intragrain diffusion of elements has occurred [38]. If post-crystallization, intra-grain diffusion of LREEs-MREEs in apatite has occurred, there will be an absence of zone-to-zone variation in La/Sm within a single grain. Most of the apatite grains were too small to perform detailed trace element spot analysis. However, the apatite in sample P-14 was large enough to use LA-ICP-MS spot analysis on the observed spectral zonation observed by cathodoluminescence and micro-XRF trace element maps. As noted above, reflected light
petrography and micro-XRF trace element maps of sample P-14 indicate the presence of tourmaline inclusions in the apatite grain. The inclusions of tourmaline in the apatite were small enough to avoid, with careful placement of the laser spot. Sample P-14 had two clearly identifiable zones (core and rim) in micro-XRF mapping. The core of the apatite grain appeared enriched in manganese and depleted in REY with respect to the rim (Figure 2). CL-imaging of the apatite grains indicated that yellow was the main colour of the CL emission among the studied apatite grains. The core of P-14 had La/Sm values ranging from 0.91 to 4.61, whereas the surrounding rims are found with lower La/Sm varying between 0.34 and 2.09. Therefore, based on the zone-to-zone variation of La/Sm in sample P-14, intra-grain diffusion did not occur, and the observed zoning pattern of apatite records the primary distribution of elements. This zonation pattern developed (as [39] suggest) from changes in the hydrothermal system, such as varying fluid composition, pressure, temperature, as well as changes in mineral growth rate and not as separate and distinct hydrothermal events.

It was found that the fluids responsible for deposition of intergrown quartz and tourmaline in gold-bearing vein samples from the Ptarmigan vein had a temperature of 560° to 600° ± 50°C [40]. Additionally, the arsenic concentrations in the Ptarmigan also indicated that the mineralizing fluid was 550° ± 50°C supporting the role of a high-temperature fluid in formation of these gold deposits. The character of wall-rock alteration adjacent to the Ptarmigan veins also supports that the vein formed at a high temperature. Furthermore, they [40] found that the “static” growth of cordierite porphyroblasts in wall rocks that host the Ptarmigan deposit indicates that the country rocks experienced temperatures of ≥500°C during amphibolite-grade metamorphism.

4.1. Trace element substitution

Elements can substitute into apatite either by direct (i.e., the substituting element has a charge and effective ionic radius compatible with that of the element it is replacing) or by heterovalent substitution (i.e., coupled substitution or the presence of a vacancy). For any given element to substitute into apatite, it must adhere to Goldschmidt’s Rules. Thus, elements with an ionic radius closest to the substituted ion will preferentially be incorporated into apatite. For example, substitution of REY in apatite follows two rules based on the availability of the two Ca positions in apatite: The LREE preferring the Ca2 site and the HREE the Ca1. The selective incorporation of REEs into apatite is driven by crystallographic features and by fluid composition and temperature [15].

The trace element composition of apatite is dependent on the hydrothermal fluid composition, host rock composition, temperature, pressure, pH, and fO2, as well as the partition coefficients between fluid, apatite, and co-precipitating minerals. It was noted that auriferous hydrothermal fluids in orogenic gold deposits are typified as forming from mixed aqueous-carbonic fluid with a low salinity with a minor difference in composition [9]. Additionally, these fluids have been shown to contain Na, minor K and B, and low levels of trace elements [41]. However, in the present study area, the concentrations of REEs show measured variation in apatite taken from different quartz veins. This contrasts with low variance in hydrothermal fluid compositions in orogenic gold deposits.

The hydrothermal apatite in the study area exhibits five different REE patterns. Since the behavior of the REE in hydrothermal fluids is most affected by parameters, such as pH, temperature, salinity, redox conditions, and fluid composition, it follows that changes in these parameters are the most likely explanations for the variation in apatite REE patterns. In any hydrothermal system, changes in REE parameters are a result of the preferential incorporation of REEs into co-crystallizing phases; or due to changes in fluid composition in response to an evolution from metamorphically derived fluid to a magmatically derived fluid. The chondrite-normalized La/Yb, La/Sm, and Gd/Yb ratios (Figure 6) show distinct trends that track fluid that reflect the evolution of the fluid or the partitioning of REE among apatite and other co-crystallizing phases. It is apparent that two distinct fluid compositions (a magmatic and metamorphic fluid) are responsible for precipitation of the apatite observed in this study. This concept was previously suggested [40], as they found that the Ptarmigan deposit was likely formed by at least two mineralizing fluids. The first fluid, a ~380°C H2O-CO2 fluid with dissolved solids, gas contents, and a temperature akin to fluids involved in the formation of nearby pegmatite dykes, was responsible for low grade Au (500 ppb). The second fluid,
a 550°C, H2O-CH4 fluid of metamorphic origin, is thought to have deposited the high-grade gold in the core of the Ptarmigan vein.

Fluids derived from the adjacent S-type Prosperous plutons metasomatized the country rock during contact metamorphism, and that fluorine and phosphorus in the fluid complexed with U, Th, Y, and REE. Consequently, these intrusions likely were an important source of U, REE, Y, as well as P and F for the studied mineralization.

Examining the CN-REE patterns in detail, type – i apatite have LREE enriched profiles with only a minor positive Eu anomaly. This LREE enrichment likely is a result of dehydration reactions during intrusion of the Prosperous Granite. This event would have released LREE-enriched fluid [15], since LREEs are generally more mobile than HREEs, the REE pattern of type – i apatite is consistent with a metamorphically derived fluid. Additionally, magmatically driven hydrothermal fluids would inherit increased LREE along with U and Th. Over time, concentrations of trace elements would be reduced as an increased abundance of other phases competing for the same elements would deplete their contents in hydrothermal fluids. It follows that trace element concentrations in type – ii apatite were inherited from these LREE depleted hydrothermal fluids. The metamorphically derived hydrothermal fluids account for the CN-REE patterns observed in type – iii, type – iv, and type – v apatite. In type – iii and type – iv apatite, an increase in HREE in the CN-patterns are best explained by the early or co-crystallizing LREE sinks (e.g., epidote, titanite, and calcite) that are observed to occur in thin section with type – iv apatite. In several studies examining scheelite from orogenic gold deposits (which partitions REEs like that of apatite, i.e., [40], REE patterns have been documented to evolve from a bell-shaped pattern with negative Eu anomalies (e.g., type – v apatite) to flat REE patterns with positive Eu anomalies (e.g., type- ii apatite) [43-44]. Interestingly, a REE-enriched bell-shaped pattern (e.g., type – v apatite) was observed in samples from the Ptarmigan vein and a smaller gold-bearing vein on the Ptarmigan Mine property. Apatite from the Ptarmigan vein contains both type – ii and type – v REE patterns and reflect the change in the hydrothermal fluid from one that was metamorphically derived type – v apatite to one that is magmatically derived (type – i apatite).

Strontium is more mobile than the REE [45]; therefore, Sr concentrations in apatite formed at higher levels in the system will have inherited Sr concentrations. Comparatively, apatite from LCT pegmatites have much lower Sr than the vein-hosted apatites. This maybe a consequence of Sr scavenging in a closed hydrothermal system. In this scenario hydrothermal quartz veins are distal end members of the LCT-pegmatites found locally. It was observed that the intrusion-related gold at the Clarence Stream gold deposit in southwestern New Brunswick [46], which found that aplite and granophyric granitic laterally change into auriferous sulfide-bearing quartz veins, appears to support this possibility. This suggests that Sr content in apatite was likely inherited from Sr concentrations in the host rocks as the hydrothermal fluids ascended thru the crust. Additionally, shales commonly have high Pb, U, and Th compared to other rock types [47], which may also explain why Pb, U, and Th signature in the type – i apatite is elevated. This reasoning suggests that the host rock compositions have an influence on the trace element composition because of the reaction and exchange between hydrothermal fluids and host rocks e.g., [9,48].

Apatite is not observed to overprint any mineral phases and likely precipitated early. It is interpreted that apatite documented in this study are primary hydrothermal apatite that record the initial fluid compositions.

The source of U, REE, Y, as well as P and F for the studied mineralization was likely derived from melting of the Burwash sedimentary package during crustal anatexis. The evolution of a fluid phase from a melt is controlled by the solubility of H2O in the melt, which is in turn dependent on temperature, pressure, and composition. The Prosperous plutons are interpreted as 2 km thick sill-like intrusive sills emplaced at depths between 9 and 13 km (3.0-3.5 kbar (300-350 MPa) and 550° C; [49]). As sill-like intrusions cool, a boundary zone forms at the margins of the body. The last region to cool is the core. This cooling and crystallization of the magma causes saturation, and evolution of a H2O-rich volatiles. As crystallization continues, one of two events must occur, either the magma expands, or the internal pressure increases. Once internal pressures overcome the lithostatic load, fracturing occurs in the vertical plane [50]. This allows fluids to be channeled upwards. These fluids contain metals, S, Cl, and CO2 that have been separated from the cooling magma. Additionally, REEs
are shown to form stable complexes with Cl [51]. The high concentrations of Cl exsolved from a pluton may transport large amounts Ca as Ca-Cl and REE-Cl complexes, which may react with H3PO4 complexes leading to the crystallization of apatite. Furthermore, sericitization of host rock likely elevated Ca²⁺ activity in the hydrothermal fluid. As a result, the elevated Ca²⁺ content this would limit the precipitation of monazite, while promoting the saturation and growth of apatite. At temperatures greater than 350°C (such as in magmatic-hydrothermal systems) and low pH Au⁺ forms stable bonds with harder anions such as Cl [21]. Therefore, with decreasing temperature from magmatic to <350°C facilitated gold saturation.

Lastly, boron- and F-bearing volatiles likely escaped from this proximal granite intrusion into the adjacent metamorphic rocks. The margins of the quartz veins and the pegmatites observed at the Ptarmigan and Tom deposits typically have tourmaline-rich rims. [52] suggests that boron rich vapors were channelized along fractures that preceded pegmatite emplacement. An early boron-rich vapor likely deposited the 5-10 % concentrations of tourmaline observed in the Ptarmigan vein.

4.2. Eu Anomaly

Europium in apatite is controlled by coupled substitutions, so that Eu can substitute directly for Ca in apatite. Europium occurs as a divalent or trivalent ions. The multivalent nature of Eu allows for preferential substitution, based upon the ratio of Eu³⁺/Eu²⁺ in any given hydrothermal fluid. Since Eu³⁺ is more readily incorporated into apatite than Eu²⁺, a large Eu³⁺/Eu²⁺ ratio may manifest as a negative Eu anomaly, while small Eu³⁺/Eu²⁺ ratios may produce positive Eu anomalies on REE spider diagrams. This ratio of Eu³⁺/Eu²⁺ is controlled primarily by the initial Eu³⁺/Eu²⁺ ratio in the hydrothermal fluids, redox state, and temperature. In magmatic environments, the proportion of these two species is a function of oxygen fugacity. In contrast, the Eu³⁺/Eu²⁺ ratio in hydrothermal environments is primarily controlled by the pH, as Eu³⁺ is more dominant at high pH conditions (>7 at 300 °C [53]). In general, Eu anomalies in hydrothermal systems reflect the Eu anomalies of the fluids. During the breakdown of plagioclase, Eu²⁺ is released along with additional REEs that were incompatible with micas [54]. Therefore, the release of Eu³⁺ and REEs during sericite alteration of feldspar will be incorporated into coeval apatite. The REE patterns of type – i, type – ii, and, type – iv apatite have positive Eu anomalies. The type – i apatite has a weak Eu anomaly in comparison to type – ii and type- iv apatite. The primary difference between type – ii and type – iv apatite is the relative enrichment in HREEs and a depletion in LREE in type – iv apatite. Additionally, type – iii and type – v apatite have large negative Eu anomalies. Two potential explanations for these variations in the magnitude of the Eu anomaly is the dissolution of plagioclase or (2) an increased in pH or oxidation state of the mineralizing fluid. The crystallization or breakdown of feldspars alters the ratio of Eu³⁺/ Eu²⁺, thereby influencing the presence of an Eu anomaly. For example, crystallization of plagioclase prior to apatite mineralization will facilitate the uptake of Eu²⁺ and result in an Eu³⁺-depleted fluid. Furthermore, the breakdown of feldspar also releases additional Sr. Therefore, if the breakdown of plagioclase added Eu²⁺ to the fluid (i.e., type – ii and type – iv REE patterns), the Sr contents, when plotted against Eu anomalies should also be correlated. However, Sr vs Eu/ Eu⁺ in the apatite show a scattered distribution (Figure 8 f). Therefore, the enrichment in REEs from the dissolution of plagioclase was likely not responsible for the MREE depletion and positive Eu anomalies that characterize the type – iv patterns.

4.3. Apatite Geochronology

The apatite in this study likely formed coeval with early stages of sulphide precipitation. The age of the hydrothermal apatite (2585 ± 15 Ma) is consistent with the intrusions of the 2605 and 2590 Ma two-mica granites of the Prosperous Suite [1-2]. The near-concordant ⁸⁷Sr/⁸⁶Sr-corrected data of the LCT pegmatite-hosted apatite reveals two clusters of ages. An older near-concordant cluster with a mean concordia age (n = 4) of 2581 ± 15 Ma, and a younger population with a mean Concordia age (n = 3) of 2519 ± 12 Ma. Furthermore, plotting a regression through all near – concordant data for the pegmatite hosted apatite hints that metamorphic resetting occurred around ~2200 Ma.
5. Conclusions

The vein-hosted apatite documented in this study typically display a dark yellow core surrounded by a brighter yellow rim in cathodoluminescence. Apatite exhibits oscillatory zoning and breccia texture, which indicates that apatite precipitated from hydrothermal fluids. The apatite is petrographically indistinguishable from vein to vein. Apatite appears as an early phase that has been overprinted by later mineral phases. U-Pb dating reveals that the apatite has a crystallization age of 2590 ± 13 Ma.

Rare earth elements display five patterns in the hydrothermal vein-hosted apatite – (i) negative sloped pattern with minor Eu anomaly, (ii) a flat pattern with a positive Eu anomaly, (iii) a positive slope with a negative Eu anomaly, (iv) LREE depleted pattern with positive Eu anomaly, and (v) bell-shaped pattern with a negative Eu anomaly. The variations in REE accompanied by changes in the abundances of other trace elements could be related to separate and distinct pulses of hydrothermal fluids with slightly different compositions. Alternatively, trace element contents of apatite reflect the chemical evolution of a single fluid through mineral precipitation and fluid-rock interaction or reflecting in situ fractionation. It is interpreted that the source of U, REE, Y, as well as P and F for the studied mineralization was likely derived from melting of the Burwash sedimentary package during crustal anataxis, which then forms S-type granitic magmas that evolved to pegmatites and apatites during emplacement then volatile saturation contributing to gold mineralized quartz veins in the region.

Supplementary Materials: The following are available online at www.mdpi.com/xxx/s1, Table S1. Apatite isotopic data; Table S2. Trace element composition of apatite

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