Bulk composition of a zoned rare-earth minerals-bearing pegmatite in the Pikes Peak granite batholith near Wellington Lake, central Colorado, U.S.A.

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ABSTRACT

A previously undescribed small lenticular (~5 × 5 × 5 m) pegmatite, located near Wellington Lake in the NW part of the 1.08 Ga ‘A-type’ (anorogenic) ferroan Pikes Peak granite batholith, ~15 km SW of the South Platte pegmatite district in central Colorado, is concentrically zoned around a mostly monomineralic quartz core with interconnected miarolitic cavities. Major constituents of the Wellington Lake pegmatite are quartz, perthitic microcline, albite (variety cleavelandite), hematite, and biotite. Accessory minerals include fluorocerite, bastnäsite, columbite, zircon (var. ‘cyrtolite’), thorite, and secondary U phases. Fluorite is conspicuously absent, although it is a common phase in the South Platte district NYF-type pegmatites, which are rich in niobium (Nb), yttrium (Y), fluoride (F), and heavy rare-earth elements (HREE). Notable for the Wellington Lake pegmatite are a small quantity of well-developed tabular crystals of fluorocerite that reach up to 4 cm in diameter, with sub-mm epitaxial bastnäsite overgrowths, suggesting formation from F- and CO32- bearing solutions rich in light rare-earth elements (LREE), with decreasing a(F)/a(CO32-) during the last crystallization phase. An Nd-isotope value of εNd1.08Ga = -1.6 for the fluorocerite is within the range of εNd1.08Ga = -0.2 to -2.7 of the host coarse-grained, pink K-series Pikes Peak Granite (PPG), indicating that REE and other pegmatite constituents derived from the parental PPG magma. A calculation of total pegmatite composition based on whole-rock chemistry and volume estimates of the different pegmatite zones reveals an overall composition similar to the PPG with respect to Si, Al, Na, and K. Yet the pegmatite is depleted in Fe, Mg, Ca, Ti, Mn, and P; the high-field-strength elements (HFSE; Zr, Hf, Nb, Y, Th); and, most significantly, total REE compared to the PPG. Despite containing the LREE minerals fluorocerite and bastnäsite, the lack of a net overall REE enrichment of the pegmatite compared to the PPG reflects the large amount of REE-poor silicate minerals forming the wall, intermediate, and core zones of the pegmatite. The calculated total pegmatite composition suggests that the pegmatite formed by the separation from the PPG magma of an F-poor H2O-saturated silicate melt depleted in REE and HFSE compared to the F-rich melts, which formed the NYF-type HREE-rich (LaN/YbN < 1) pegmatites in the South Platte district. Homogenization temperatures of < 500°C for possibly primary fluid inclusions in large quartz crystals from the core of the Wellington Lake pegmatite are consistent with recent models of pegmatite petrogenesis leading to nucleation controlled mega-crystal growth resulting from supercooling.

KEY WORDS: Colorado, fluorocerite, Pikes Peak granite batholith, rare-earth elements, South Platte pegmatite district, Wellington Lake pegmatite.

INTRODUCTION

The petrogenesis of granitic pegmatites has been a subject of debate to consistently explain their textural and mineralogical features, notably textural and mineralogical heterogeneity, inward coarsening and oriented crystal growth, distinct zonation of mineral assemblages, and their mineralogical and geochemical diversity (Swanson, 1977; Simmonds and Webber, 2008; London,
2008, 2009, 2014, 2015, 2016, 2018; Thomas et al., 2012; Thomas and Davidson, 2015; Sirbescu et al., 2017). Although pegmatites are generally established to be of igneous origin, the role of late-stage hydrothermal processes is often hard to either discount or discern.

Few studies have systematically addressed the bulk composition of pegmatites, because of the difficulties associated with exposure, internal zoning, and paragenetic heterogeneity (Gordienko, 1996; Zagorsky et al., 1997). Two recent notable exceptions are for the very large, complexly zoned, rare-earth element (REE) and Li-, Cs-, and Ta-rich (LCT-type) Tanco pegmatite at Bernic Lake in southeast Manitoba (Stilling et al., 2006), and the smaller Mount Mica LCT-type pegmatite dike in southwest Maine (Simmons et al., 2016). Whereas the Mount Mica study suggests a derivation by the anatexis of the host rocks for this pegmatite, the Tanco pegmatite work supports an origin from a parental granitic magma.

Another class of pegmatites of particular interest in Colorado are those associated with ‘A-type’ (anorogenic; Eby, 1990; Černy and Ercit, 2005; London, 2008) ferroan (Frost and Frost, 2011) granites, such as those in the South Platte pegmatite district at the northern end of the 1.08 Ga Pikes Peak granite batholith (Smith et al., 1999; our Fig. 1). This district has long been known for its abundant pegmatites, which locally are quite large (as large as ca. 50 m in width), and commonly contain unusual REE minerals and zoned mineral assemblages (Simmons and Heinrich, 1975, 1980; Simmons et al., 1987; Allaz et al., 2020). Several of the pegmatites in the central part of the South Platte district contain locally abundant HREE-rich minerals. They are well described examples of NYF-type (gadolinite subtype) pegmatites, which are characteristically rich in Nb, Y, F, and HREE, and commonly associated with ‘A-type’ ferroan granites (Eby, 1990; Černy and Ercit, 2005).

Early studies suggest that the South Platte pegmatites formed from the crystallization of a late-stage, volatile-rich magma of the Pikes Peak granite batholith (Haynes, 1965). However, aspects of the origin and formation of the South Platte pegmatites still remain unclear, specifically whether the REE-rich minerals in these pegmatites are the result of an already REE-enriched parental granitic magma (Simmons and Heinrich, 1980), result of extreme fractionation concentrating REE during pegmatite crystallization, or result from an influx of REE-rich fluids unrelated to the Pikes Peak Granite (PPG) (Gagnon et al., 2004). For the Oregon #3 pegmatite in the South Platte pegmatite district (Fig. 1), Simmons and Heinrich (1980) noted that although the PPG is enriched in total REE relative to many other granites, the total REE content of the pegmatite seems ~10× above the PPG value. However, although the PPG is enriched in LREE relative to HREE (La/YbN = 10), the Oregon #3 pegmatite appears to be enriched in HREE relative to LREE, with La/YbN < 1. This implies that processes involving fractionation of HREE relative to LREE, rather than simple overall enrichment in total REE relative to the granite, must be considered in the formation of the Oregon #3 and other South Platte NYF-type pegmatites.

In contrast to the large NYF-type pegmatites in the central part of the South Platte district, pegmatites in the southern and western part of the batholith are mostly smaller in size (<1 m to 15 m diameter) and often contain the LREE-rich minerals allanite and monazite. In this work we investigate an example of one of these smaller, internally zoned, LREE mineral-bearing pegmatites, located northwest of Wellington Lake (Fig. 1). This Wellington Lake pegmatite has not been previously described. To understand the chemical, textural, and mineralogical attributes of this pegmatite, we undertook mapping and transect sampling of the pegmatite, thin-section petrography, and whole-rock chemical analysis of the individual pegmatite zones via bulk sampling. Additionally, petrographic and thermometric analyses of fluid inclusions in quartz from the pegmatite core zone were performed, along with quantitative mineral analysis of the LREE-rich minerals in the pegmatite core. Finally, Nd-isotope data were obtained to determine if the pegmatite magma source was the magma that formed the PPG hosting the pegmatite.

GEOLOGIC SETTING

Figure 1 shows the geologic setting with the Wellington Lake pegmatite (lat. 39.322855°S, long. 105.399503°W) situated as an outlying body southwest of the central South Platte pegmatite district. The Wellington Lake pegmatite is located ~3 km northwest of Wellington Lake and ~2 km southeast of the McGuire pegmatite (Fig. 1), which is a large, irregularly zoned NYF-type pegmatite (Černy et al., 1999). Field work around the Wellington Lake pegmatite revealed several other pegmatites as isolated bodies or swarms. Most notably, ~500 m to the SW of the Wellington Lake pegmatite, a group of smaller, well-exposed granite-hosted pegmatites crop out. These contain fluorite in mioralitic cavities in their quartz cores, but lack apparent REE mineralization.

The Wellington Lake pegmatite is a steeply dipping, lenticular body hosted in medium- to coarse-grained, pink syenogranite corresponding to the major K-series phase of the Pikes Peak granite batholith. It is partially exposed in vertical cross section in an ~10-m-tall, steep west-facing outcrop, in sharp contact with minimally weathered pink syenogranite on at least three sides in its exposure (Fig. 2). The pegmatite body consists of a main, roughly prolate, ellipsoidal body ~4 m in diameter. It is exposed in a frontal plane with an upper zone over a vertical extent of ~3 m. Figure 2A shows a cross section of the upper mioralitic zone.
The pegmatite has recently been prospected for its large milky and smoky quartz crystals, which are abundant in the core zone of the pegmatite.

Wall Zone

As seen in cross sections B and C of Figure 2A, the host granite is in sharp contact with a narrow (~10-cm-wide) wall zone of graphic granite. The wall zone represents about 20 vol. % of the pegmatite (Fig. 2).

Intermediate Zone

The wall zone then transitions into an ~95-cm-wide intermediate zone, which represents about 69 vol. % of the pegmatite. First is a coarser-grained ~20-cm-wide outer intermediate zone (OIZ) of graphically intergrown quartz and microcline-perthite with minor biotite laths. We estimate that the OIZ represents about 22 vol. % of the pegmatite (Fig. 2). Proceeding inward, the OIZ is in contact with an ~60-cm-wide middle intermediate zone (MIZ) composed of a still coarser graphic quartz and microcline-perthite without biotite. The MIZ represents ~41 vol % of the pegmatite. This zone is followed inward by an ~15-cm-wide inner intermediate zone (IIZ), which is characterized by microcline-perthite and small laths of albite, without quartz, along with irregular pods of 'cleavelandite' albite, biotite, and hematite. We estimate that the IIZ represents ~6 vol. % of the pegmatite.

Core Zone

The core zone consists of an ~2-m-diameter quartz core with open cavities lined with anhedral to euhedral, multi-generation milky and smoky quartz crystals as large as ca. 50 cm in size. Open spaces are partially filled with minor hematite, albite, and clay minerals. The core represents about 11% of the pegmatite volume. Mica-litic cavities within the core form void space in druses and between large quartz crystals. These constitute < 0.5 vol. % of the total pegmatite.

Accessory minerals in the pegmatite include fluorocerite and bastnäsite in the core, and columbite, 'crytoltite' zircon, thorite, and minor secondary U–Th species in the IIZ. Notable is the absence of fluorite or phosphate minerals (e.g., monazite, xenotime, or apatite) in any of the zones of the pegmatite body.

METHODS

Thin sections were examined, and bulk chemistry determined (Figs. 3 and 4) for samples across the transition from the granite into the core of the pegmatite (Fig. 2; Tables 1–3). Inductively coupled plasma-mass spectrometry (ICP-MS) trace-element and whole-rock major element compositions were obtained from Activation Laboratories Ltd. (Ontario, Canada).

Mineral chemistry was determined by electron microprobe (EMP) analysis using a JEOL JXA-8600 EMP at the University of Colorado Boulder. It is a four-spectrometer instrument equipped with argon X-ray detectors (P-10 mixture) on spectrometers 1 and 2 (PET and TAP crystals), and xenon X-ray detectors on spectrometers 3 and 4 (LiF crystals). Operating conditions using a W-cathode were 15 KV accelerating potential with a 20 nA probe current. Details on EMP analy-
sis and standards used for the REE minerals fluocerite and bastnäsite are described in Allaz et al. (2015, 2020).

Micro-Raman spectroscopy on polished thin sections was performed on an upright Raman microscope (Olympus BH51, with 632 nm HeNe laser excitation at 1 mW, a 50× objective of NA = 0.8, and a Princeton Instruments Acton SP500a imaging spectrograph, with a PIXIS 100 liquid nitrogen-cooled CCD camera, calibrated on multiple spectral lines using a He-lamp). Nd-isotopic data were obtained using established analytical procedures (Farmer et al., 1991). Measured $^{143}\text{Nd}/^{144}\text{Nd}$ was normalized to $^{146}\text{Nd}/^{144}\text{Nd} = 0.7219$. The Nd-isotopic compositions are reported as $\varepsilon_{\text{Nd}}$ values using a reference $^{143}\text{Nd}/^{144}\text{Nd}$ ratio of 0.512638.

Fluid-inclusion petrography and thermometry were performed on doubly polished thin sections of quartz from the top of the core of the pegmatite. Inclusions were analyzed thermometrically with an adapted U.S. Geological Survey gas-flow heating stage. Stage calibration was carried out at -56.6°C, 0.0°C, and 374°C, using synthetic fluid inclusion (SYN FLINC) standards. Uncertainties in the thermometric measurements are ± 0.2°C at 0.0°C, and ± 3°C at 374°C. Salinities were determined from thermometric measurements as described by Potter et al. (1977, 1978).

**PETROLOGY AND CHEMISTRY**

**Granite**

The coarse-grained, pink K-series Pikes Peak Granite (PPG) that hosts the pegmatite consists of quartz, perthitic microcline and orthoclase, and Fe-rich mica (annite), with minor plagioclase and accessory magnetite, zircon, fluorite, and trace monazite. Its mineralogy is typical for coarse-grained pink Pikes Peak granite, but lacks amphibole, which has been observed in small quantities in other K-series Pikes Peak granite samples elsewhere in the batholith (Simmons and Heinrich, 1980; Simmons et al., 1987; Smith et al., 1999). Three samples of granite, WL-19, WL8-1, and WL8-2 (Table 1), are similar in composition to previously published analyses (Smith et al., 1999). They have 600 to 800 ppm total REE, which is relatively high.
compared to other granites worldwide (Simmons and Heinrich, 1980; Simmons et al., 1987), La/Yb ratios of approximately 10, and significant negative Eu anomalies (Figs. 3A and 4). One sample, WL4-1 (Table 1), has somewhat lower FeO, Zr, Y, Nb, Hf, U, and HREE, but appears mineralogically similar to the other three. This sample was not included in the average composition of the granite used in our bulk pegmatite model described below.

**Pegmatite Wall Zone**

The 5- to 15-cm-thick wall zone consists of a relatively fine-grained graphic intergrowth of quartz and perthitic microcline with biotite and occasional zircon grains. Quartz content is greater and biotite content is lower than in the granite, and Na-plagioclase occurs only in the perthite and not as independent grains. Fe-oxides and fluorite are absent.

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Figure 3. Whole-rock REE contents of: A, host coarse pink Pikes Peak granite; B, pegmatite zones from the wall zone, through the outer, middle, and inner intermediate zones (OIZ, MIZ, and IIM); and C, samples from the “replacement” zone of albite, thorite, and columbite bearing rock, and the hematite breccia. Fluocerite ICP-MS analyses, as well as fluocerite and bastnäsite electron microprobe (EMP) analyses are also plotted in C. Fields in C have the same colors as the individual samples in A (red for the granite), and B (shades of brown for the wall and intermediate zones). Note the continuous inward REE depletion, yet increasing La/Yb, and the HREE enrichment in the hematite breccia and albite, thorite, and columbite bearing rocks of the “replacement zone,” in contrast to and compensated by the LREE-rich mineral phases.
Samples of the wall zone (Figs. 3B and 4; Table 2) have compositions similar to the granite host (Table 1), but with slightly lower total REE contents, a feature also noted for this zone in South Platte pegmatites (Simmons and Heinrich, 1980; Simmons et al., 1987). This depletion in total REE relative to the granite host possibly reflects the greater abundance of quartz, and lower abundance of feldspars compared to the granite. The Lanthanum/Ytterbium ratios of these samples vary from 10 to 5. The negative Eu anomaly for the wall zone is also less pronounced than for the granite, possibly due to the higher proportion of potassium feldspar, which relative to plagioclase concentrates Eu as it crystallizes. In this zone plagioclase only occurs in the microcline-perthite, not as a primary crystallization phase. This zone also has lower Zr, Y, Th, and U contents than the granite (Table 2), which is reflected mineralogically in the scarcity and small size of zircon crystals.

Pegmatite Intermediate Zone

In the outer intermediate zone (OIZ), the graphic intergrowth of quartz and perthitic microcline becomes coarser and can be observed macroscopically, biotite occurs in decreased abundance as scattered large plates, and zircon is absent. Further inward toward the core of the pegmatite and away from the contact with the granite, this ~20-cm-wide OIZ changes to a middle intermediate zone (MIZ) of even coarser microcline-perthite without any biotite. There is then an inner intermediate zone (IIZ) of coarse microcline-perthite without quartz and the appearance of small euhedral laths of albite. These appear to have crystallized directly from the pegmatite parental magma.

Four samples from the OIZ (Table 2) have lower Fe$_2$O$_3$ and TiO$_2$ than the granite, reflecting the lower concentration of biotite, and significantly lower REE contents, with lower Ln/Yb$_n$ ratios between 2 and 3. These samples lack a negative Eu anomaly (Figs. 3B and 4). The lack of a negative Eu anomaly in this zone may reflect the fact that potassium feldspar, which was the dominant feldspar crystallizing from the pegmatite parental magma prior to evolving into perthite during cooling, concentrates Eu relative to other REE. Ba, Sr, Zr, Nb, Th, Hf, and U contents are also lower than in either the granite or wall zone (Table 2). Four samples from the MIZ (Table 2) have Ln/Yb$_n$ ratios between 1 and 5. Three samples from the IIZ (Table 2) have significantly lower REE concentrations despite the absence of quartz. Two of these samples have Ln/Yb$_n$ of ~1.

Albite and Hematite-rich Pods

Local segregations between the perthitic microcline-rich intermediate zone and the quartz-rich core of the pegmatite are characterized by white-bladed albite (var. cleavelandite). These pods are more heterogeneous than the intermediate zone, as indicated by the chemical analysis of 10 samples from this zone (Table 3). They contain (1) areas of albite intergrown with quartz (samples WL5-1 and WL5-5); (2) areas of almost pure albite (WL5-4A, WL5-4B, and WL7-1); (3) albite with accessory thorite (WL5-2) and/or columbite (WL5-3), as well as pockets of albite and hematite (WL4-6A and WL4-6B); and (4) breccias dominated by hematite (WL6-1). Simmons and Heinrich (1980) and Simmons et al. (1987) suggested that similar associations observed in South
Together, these so-called “replacement zone” associations comprise only ~1 vol. % of the pegmatite. The albite-rich zones have La\textsubscript{o}/Yb\textsubscript{o} ranging from 1 to 10, while the hematite-rich samples have La\textsubscript{o}/Yb\textsubscript{o} < 1 (Figs. 3C and 4).

**Pegmatite Core**

In the roof of the quartz core, in contact with the intermediate zone, an ~0.8 m × 0.5 m × 0.4 m cavity was discovered during the investigation. The cavity is formed by large intergrown microcline-perthite and quartz crystals, and
filled with elongated euhedral smoky quartz crystals up to 15 cm long, albite-hematite breccia fragments, yellow-green clay, and hematite dust. In addition, -30 euhedral 0.5 cm to 4 cm in size fluorite crystals, with bastnasite overgrowths, were found concentrated at the bottom of the cavity. The fluorite has La₆/Yb₆ ~ 1,000 (Fig. 3C).

REE Mineralogy

REE and HFSE minerals found in the pegmatite include columbite, zircon, thorite, fluorite, and bastnasite. Columbite is found both as well-formed crystals up to 6 cm in association with albite and microcline as well as subhedral masses associated with biotite. In the biotite, columbite is associated with secondary uranium minerals that occur as coating and thin fracture fillings. Nodules of zircon and thorite up to 5 cm in size are also associated with the biotite, locally with cassiterite inclusions of 10s of mm in size. The zircon-thorite (‘cyrtolite’) is also found locally in small pits on the surface of large euhedral quartz crystals.

Most notable for the pegmatite are the well-developed tabular crystals of fluorite-(Ce) [(REE)F₆], which are overgrown by bastnasite-(Ce) [(REE)(Ce)CO₃]F₂, as shown in Fig. 5A. Such fluorite-bastnasite crystals were first described from the Pikes Peak granite batholith (Cheyenne Canyon; Mount Rosa intrusive center) by Allen and Comstock (1880), who gave the fluorite cores a new name, ‘tysomite,’ after its discoverer, S. T. Tyson. Allen and Comstock were apparently not aware of the previous work of Berzelius (1818), who described a new mineral from granitic pegmatites at Finno and Broddbo near Falun, Sweden, as “Neutrals Fluss-spatsynat Cerium,” later named ‘fluorite’ by Haidinger (1845). Geijer (1921) restudied the type material from Sweden, recognizing that the Swedish fluorite is identical with tysomite, although the name tysomite can be found in the literature up to 1965 (Sverdrup et al., 1965).
Table 3. Compositions of different albite-rich pods occurring at the contact of the intermediate zone and the core zone of the pegmatite.

| Sample No. | OX Wt. % | W.L.5-1 | W.L.5-5 | W.L.5-A | W.L.5-B | W.L.7-1 | W.L.5-2 | W.L.5-3 | W.L.4-6B | W.L.4-6A | W.L.6-1 |
|------------|----------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| SiO₂       |          | 74.7    | 77.4    | 68.1    | 67.6    | 65.3    | 66.5    | 63.7    | 47.1    | 50.1    | 7.95    |
| TiO₂       | <0.01    | <0.01   | <0.01   | <0.01   | 0.03    | 0.01    | 0.06    | 0.27    | 0.03    | <0.01   |         |
| Al₂O₃      | 15.5     | 13.9    | 19.8    | 19.9    | 19.8    | 19.9    | 18.9    | 17.1    | 15.7    | 1.60    |         |
| Fe₂O₃      | 0.36     | 0.17    | 0.04    | 0.05    | 2.20    | 0.52    | 2.59    | 19.7    | 19.5    | 76.7    |         |
| MnO        | <0.01    | <0.01   | <0.01   | <0.01   | <0.01   | <0.01   | <0.01   | <0.01   | <0.01   | <0.01   | <0.01   |
| MgO        | <0.01    | <0.01   | <0.01   | <0.01   | 0.04    | 0.01    | 0.04    | 0.30    | 0.17    | <0.01   |         |
| CaO        | 0.28     | 0.24    | 0.14    | 0.53    | 0.33    | 0.56    | 0.44    | 1.61    | 2.38    | 0.65    |         |
| Na₂O       | 8.41     | 7.67    | 11.44   | 11.48   | 10.46   | 10.93   | 9.88    | 3.91    | 6.96    | 0.05    |         |
| K₂O        | 0.26     | 0.18    | 0.13    | 0.15    | 0.55    | 0.37    | 0.34    | 3.23    | 0.66    | 0.12    |         |
| P₂O₅       | <0.01    | <0.01   | <0.01   | <0.01   | <0.01   | <0.01   | 0.01    | <0.01   | <0.01   | <0.01   | <0.01   |
| LOI        | 0.4      | 0.4     | 0.5     | 0.4     | 1.2     | 0.7     | 1.8     | 6.2     | 4.6     | 13.4    |         |
| Sum        | 99.9     | 99.9    | 100.0   | 100.0   | 100.0   | 99.4    | 97.8    | 99.5    | 99.9    | 99.8    |         |

Fluocerite, while overall uncommon in Pikes Peak pegmatites, has been reported locally in abundance, notably in the Black Cloud, Little Patsy, and Mount Rosa area pegmatites, some with, some without bastnäsite association. Additional descriptions of fluocerites crystals overgrown by bastnäsite in Colorado were made by Hidden (1891) from the Crystal Peak area, Hillebrand (1899), and Glass and Smalley (1945). The exact structural relationship between the two minerals was only recently confirmed from crystallographic and high-resolution TEM analysis as an epitaxial growth along the crystallographic c-axis associated with the hexagonal symmetry, with close lattice match (Müller et al., 2011; our Fig. 5B). In addition to the epitaxial overgrowth relationship, bastnäsite replacement of fluocerite (typically on crystal or grain margins or along cleavage planes) has been noted by several authors (Lahti and Suominen, 1988; Beukes et al., 1991).

Polarization resolved micro-Raman spectroscopy, with vibrational modes characteristic for the two mineral species and their spatial invariance across the crystal (Fig. 5C), confirms the single crystal nature of both the fluocerite and bastnäsite in the Wellington Lake pegmatite. Both fluocerite and bastnäsite have space group P6₁/mcm and corresponding symmetry point group D₆h. Based on the Raman active modes and the respective Raman tensors (A₁g, E₁g, and E₂g), the crystallographic orientation was determined through choice of input and output polarization (H and V respective to the c-axis) (Kuzmanyak, 2009). This indicates the parallel orientation of the (0001) crystal planes of the fluocerite core and bastnäsite. Together with the bastnäsite primarily grown on the (0001) crystal plane of fluocerite, of even thickness and a straight phase boundary (see also Fig. 6A and B showing a region of the only major disturbance of
Table 4. Electron microprobe analyses of a euhedral fluocerite and the bastnäsite overgrow on this crystal, in comparison with a bulk ICP-MS analysis of another fluocerite crystal from the same mineralized pocket.

| Sample          | Bastnäsite | Fluorocerite |
|-----------------|------------|-------------|
|                 | Ox, Wt. %  | Low Gd     | High Gd | High Ce | average | average | ICP-MS |
| ThO₂            |            | 0.53       | 0.9     | 0.38    | 1.09    | 1.07    |        |
| Y₂O₃            |            | 0.58       | 0.2     | 0.29    | 0.46    | 0.95    |        |
| La₂O₃           | 14.87      | 11.88      | 17.22   | 15.52   | 21.35   | 19.82   |        |
| Ce₂O₃           | 35.86      | 32.01      | 36.33   | 34.72   | 41.9    | 36.08   |        |
| Pr₂O₃           | 3.83       | 3.83       | 3.22    | 3.51    | 3.86    | 3.77    |        |
| Nd₂O₃           | 14.3       | 16.24      | 10.84   | 13.33   | 14.21   | 11.66   |        |
| Sm₂O₃           | 2.1        | 4.09       | 2.02    | 2.48    | 1.87    | 1.68    |        |
| Eu₂O₃           |            |            |         |         | 0.03    |         |        |
| Gd₂O₃           | 0.6        | 2.14       | 1.04    | 1.19    | 0.79    | 0.67    |        |
| Tb₂O₃           |            | 0.14       | 0.12    | 0.12    | 0.07    | 0.07    |        |
| Dy₂O₃           |            | 0.31       |         | 0.16    | 0.17    | 0.24    |        |
| Yb₂O₃           |            |            |         |         | 0.02    |         |        |
| MgO             |            |            |         | 0         | 0.01    | 0.01    |        |
| CaO             |            |            | 0       | 0        | 0.25    | 0.86    |        |
| F               | 8.77       | 8.35       | 8.6     | 8.62    | 32.64   | 19.9    |        |
| O-F             | -3.69      | -3.52      | -3.62   | -3.63   | -13.74  | -       |        |
| Total           | 95.71      | 95.65      | 96.69   | 95.79   | 104.93  | 96.82   |        |

Table 4. Electron microprobe analyses of a euhedral fluocerite and the bastnäsite overgrow on this crystal, in comparison with a bulk ICP-MS analysis of another fluocerite crystal from the same mineralized pocket.

Neodymium Isotopes

Neodymium isotopic compositions (Table 5) were determined for a fluocerite crystal from the Wellington Lake pegmatite and for two bulk samples of REE mineral-rich pods from the White Cloud pegmatite, a classic NYF-type pegmatite located to the east in the central South Platte district (Fig. 1). The isotopic ratios of these samples plot close to a 1.08 Ga isochron (Fig. 7A), as do samples from the PPG (Smith et al., 1999). The initial εNd values at 1.08 Ga for the Wellington Lake pegmatite fluocerite is -1.6, and for the REE mineral-rich pods from the White Cloud pegmatite are -2.0 and -2.1 (Fig. 7B). These values fall within the range of -0.2 to -2.7 determined for the potassic series rocks from the PPG. They are consistent with the formation of the potassic series rocks of the PPG by either melting of tonalitic crustal rocks (Smith et al., 1999; Frost and Frost, 2011) or a more complex history of crystallization of man-
tle-derived basaltic magma combined with crustal assimilation (DePaolo, 1981; Guitreau et al., 2016).

**FLUID INCLUSIONS**

Fluid inclusions in pegmatite core-zone quartz of three different generations were analyzed by microthermometry. The quartz samples include one large crystal (W-26) from the center of the core, which had no clear terminations, a second large crystal (W-3) from along the contact between the core and microcline of the IIZ, and a third doubly terminated euhedral crystal (W-29) spatially associated with the fluocerite and bastnäsite mineralization.

Inclusions are abundant in all three samples (Fig. 8). They include both clearly secondary inclusions trapped along fractures, and possible primary inclusions randomly distributed or parallel to crystal growth planes defined by crystal faces. Four types of inclusions were identified: (1) type A two-phase (> 80% liquid, and < 20% vapor) liquid-rich inclusions (Fig. 8B); (2) type B two-phase (> 60% vapor, and < 40% liquid) vapor-rich inclusions; (3) rare type C inclusions containing transparent cubic crystals we interpret as halite (Fig. 8C); and (4) type D CO$_2$-bearing inclusions (Fig. 8D). Types A, B, and C had no visual CO$_2$ bubbles at room temperature. This suggests the presence of an aqueous H$_2$O-NaCl fluid with little or no CO$_2$, and a distinct H$_2$O-CO$_2$ fluid. Quartz crystals W-26 from the center of the core and W-3 from the edge of the core had all four types of inclusions, whereas sample W-29, associated with the REE mineralization, had only type A and type D inclusions.

The thermometric behavior of possible primary inclusions belonging to the three CO$_2$-free populations (types A, B, and C) suggests, by eutectic points very close to -21°C, that most of these inclusions trapped solutions of the H$_2$O-NaCl-system. A small group of these inclusions has slightly lower eutectic temperatures, between -22 and -24°C, indicative of other cations in solution, possibly K$^+$. The absence of any low eutectic point (-50°C) inclusions precludes the presence of CaCl$_2$ in the solutions. The majority of inclusions are liquid-rich type A of intermediate salinities between 10 and 16 wt. % NaCl equivalent, which homogenize between 119 and 196°C (Fig. 8A). In sample W-26, from the center of the core, type B vapor-rich inclusions, with > 60% vapor, homogenize at 340 to 360°C, and have salinities of 3 to 6 wt. % NaCl (Fig. 8A). In addition, a few of the relatively rare halite-bearing type C fluid inclusions, with salinities around 28 wt. % NaCl and homogenization temperatures of 320 to 380°C, were measured in sample W-26. Both types B and C also occurred in sample W-3 from the edge of the core, but could not be measured in this sample. A pressure correction of 1,500 bar, equivalent to a crystallization depth of 5 km as estimated by Simmons and Heinrich (1980) for the depth of solidification of the PPG, indicates a maximum entrapment temperature of -490°C for the halite-bearing type C inclusions in quartz crystal W-26.

**DISCUSSION**

**Pegmatite Bulk Chemical Composition**

The mineralogy and chemistry of the pegmatite appears to reflect a series of primary magmatic processes, as indicated by the absence of any evidence of major secondary hydrothermal or weathering processes in either the pegmatite or surrounding host granite. Thus, we see no indication of secondary exchange of elements with the larger volume of surrounding Pikes Peak granite. Minor hydrous sec-
ondary uranium mineralization on columbite, and host rock alteration around the cyrtolite, suggest some late-stage alteration possibly associated with metamictization. The pods of albite—in the form of euhedral crystal aggregates intergrown with quartz, hematite, columbite, zircon, and thorite—are similar to the associations in the “replacement” zones of the South Platte pegmatite district, which Simmons and Heinrich (1975, 1980) and Simmons et al. (1987) suggest were formed from volatile-rich pegmatite fluids developed during a latest stage of crystallization. Alternatively, Černý et al. (1999) suggested that hematite in the nearby McGuire pegmatite may have formed by oxidation of magnetite by subsolidus pegmatite fluids at high fO2. However, it remains unclear why the Wellington Lake and other South Platte pegmatites hosted in the magnetite-bearing ferroan PPG contain hematite.

We have estimated the bulk composition of the pegmatite, for comparison with the host granite. The whole-rock analyses of different pegmatite zones allow for an estimate of the average composition of each zone (Data Supplement 1 [DS 1]). From mapping the pegmatite to the best extent possible within the constraints of external and internal exposure (Fig. 2), and translating the spheroidal shape into a sphere, the volumes of the different zones were obtained and converted to masses based on measured densities of their constituent mineral phases (Data Supplement 2 [DS 2]). For the thorite, columbite, and hematite bearing zones, as well as the fluorocerite, total masses were estimated based on the field observations of their total modal abundances, which are small. Overall, with a 5% void space in the quartz core, the total pegmatite mass is just slightly lower than the mass of granite of equivalent volume. This is consistent with the pegmatite being derived from the magma that formed the granite, but enriched in water, which after crystallization escaped from the system.

### Table 5. Nd-isotope analysis of a fluorocerite crystal from the Wellington Lake pegmatite and REE-mineral rich segregations from the White Cloud pegmatite in the South Platte district.

| Sample          | Sm (ppm)1 | Nd (ppm)1 | 147Sm/144Nd | 143Nd/144Nd2 | εNd(0)3 | εNd(T)4 |
|-----------------|-----------|-----------|-------------|--------------|---------|---------|
| Wellington Lake fluorocerite | 2,842 ± 1 | 20,331 ± 2 | 0.0846 | 0.51176 ± 0.00001 | -17.2 | -1.6 |
| White Cloud WhC R3A | 7,242 ± 1 | 24,209 ± 4 | 0.1810 | 0.51242 ± 0.00001 | -4.3 | -2.1 |
| White Cloud WhC R4A | 7,605 ± 3 | 26,050 ± 4 | 0.1766 | 0.51239 ± 0.00001 | -4.8 | -2.0 |

Total procedural blanks averaged ~100 pg for Nd during study period. Analyses were dynamic mode, three-collector measurements. Thirty-two measurements of the La Jolla Nd Standard during the study period yielded a mean 143Nd/144Nd = 0.511840 ± 1 (2-σ mean).

1 εNd(0) are measured (present-day) values, initial εNd calculated at T = 1080 Ma.
2 Measured 143Nd/144Nd normalized to 143Nd/144Nd = 0.7219.
3 εNd values calculated using a present-day 143Nd/144Nd (CHUR) = 0.512638.

![Figure 6](image)

**Figure 6.** A, Backscattered electron image of fluorocerite-bastnäsite interface. B–E, Microprobe X-ray element maps for Ce, Gd, Sm, and Y (Figs. B–E, respectively) highlighting mostly spatially homogeneous fluorocerite composition, yet concentric zoning of the bastnäsite overgrowth. F, Integrated X-ray intensity (arbitrary units) transect of REE variation of the area indicated in Figure 6B, with anticorrelation of LREE (Ce) and HREE (Gd, Sm, and Y).

With the average compositions from DS 1 and estimated volumes for each zone from DS 2, the corresponding masses of the individual element oxides for each zone can be calculated. From their sum, the overall composition of the whole pegmatite can be obtained and compared to a granite...
body of the same size (DS 2). The results of the comparison of model pegmatite composition and granite and their fractional difference for the different elements are shown in Table 6, and in graphical representation in Figure 9. Constrained by exposure, coarse-grained texture, and heterogeneity of the mineralized zones, this approach of pegmatite bulk composition analysis has its limitations. However, given the good exposure of this pegmatite and extrapolating from the even distribution of mineralized zones observed along the margin of the quartz core, we estimate an uncertainty of <50% in the bulk pegmatite concentrations of the different trace elements, and much better for the estimation of the major elements. The lateral extent and top of the pegmatite are exposed and has been investigated well enough to conclude with good confidence that we have not missed any major element, REE- or HFSE-bearing phases. Although we were unable to probe the extent and mineralogy of the pegmatite at depth, the inward curvature of the pegmatite wall at the floor of the accessible cavities suggest a spheroidal shape. Further, from field observations of pegmatites in the South Platte district, it appears that the REE and HFSE mineral phases are generally concentrated in the upper regions of the pegmatites, making it unlikely that larger amounts of the associated elements have been missed.

The contents of Si, Al, K, and Na estimated for the Wellington Lake pegmatite are similar to that of Pikes Peak granite. Although the relative concentrations of these elements vary across the wall, intermediate, and core zones, the good agreement of the total content of these elements between the pegmatite and Pikes Peak granite indicate that the volume estimates, especially of the intermediate zone and quartz core, are approximately correct, with the implication

Figure 7. A, Measured Nd-isotopic ratios of fluorocerite from the Wellington Lake pegmatite and REE-rich aggregates from the White Cloud pegmatite in comparison with bulk samples of Pikes Peak granite from Smith et al. (1999) compared to a 1.08 Ga isochron. B, Corresponding $\epsilon_{\text{Nd}}$ versus age compared to the Proterozoic crustal rocks from Colorado, including the metamorphic basement, Boulder Creek granodiorites, Sherman and Silver Plume granites, and Pikes Peak granite (figure modified after Smith et al., 1999).

Figure 8. A, Temperature (°C) versus salinity (wt % NaCl equivalent) in fluid inclusions from three quartz crystals in the core of the Wellington pegmatite. B, Type A two-phase inclusions with >80% liquid (L) and <20% vapor (V) found in all three samples. C, Type C inclusions containing cubic crystals of what we interpret to be halite (H) found in quartz sample W-26 from the center of the core and the large quartz crystal W-3 from the edge of the core, but not in the doubly terminated crystal W-29 associated with the REE-mineralization. D, Type D CO$_2$-bearing inclusions found in all three quartz samples. Corresponding data of quartz-hosted secondary L-V and L-V-H inclusions from the Oregon #3 pegmatite are shown for comparison (Gagnon et al., 2004).
that the parental pegmatite melt is principally derived from the PPG, consistent with the Nd-isotopic composition of fluocerite in the pegmatite.

However, and well within the uncertainty of the approach, the pegmatite overall is depleted in Fe, Mg, Mn, the HFSE (Ti, Hf, Zr, Th, Nb), along with P and Ca, as well as the alkaline earths Sr and Ba (Fig. 9). The pegmatite is further depleted in REE relative to the PPG, possibly slightly more for the HREE (Fig. 9). Therefore, despite the presence of the LREE minerals fluocerite and bastnäsite, the Wellington Lake pegmatite should not be considered as LREE-rich. We conclude that the fluocerite and bastnäsite crystallized from a small amount of LREE-enriched residual melt produced by the internal crystal-liquid fractionation processes associated with the solidification of the pegmatite. The early crystallization of the large volume of silicate minerals, with relatively low REE contents (Figs. 3 and 4), that form the wall, intermediate, and core zones of the pegmatite, combined with the progressive inward decrease in LREE/HREE in these zones, generated this LREE-enriched residual melt. Although the albite, hematite, thorite, and columbite pods reverse the inward trend and have higher REE than the IIZ (Figs. 3C and 4), the low LREE/HREE of the small volume of these pods further increased the LREE/HREE of the residual F, CO₂ and LREE-enriched melt from which the fluocerite and bastnäsite crystallized.

The depletion relative to the PPG in all REE for the Wellington Lake pegmatite contrasts with the reported total REE enrichment of a factor of ~10 relative to the host PPG for the Oregon #3 pegmatite in the South Platte district (Simmons and Heinrich, 1980). However, reanalyzing the original data, Simmons (personal communication, 2020) lowered the enrichment factor to at most 2.5, adding uncertainty whether the NYF-type pegmatites in the South Platte district are indeed REE enriched relative to the PPG. It has been proposed that NYF-type pegmatites, such as those enriched in HREE and F in the central South Platte pegmatite district, would have developed their enhanced HREE concentration due to the high concentration of F in their parental melts (London, 2016). For the Wellington Lake pegmatite, which lacks significant F, the overall depletion compared to the PPG in all REE, in HREE relative to LREE, as well as HFSE, suggests that these elements were transferred from the PPG host into the pegmatite parental magma by an originally F-poor, H₂O-rich silicate melt within which these elements are relatively insoluble compared to an F-rich melt.

Crystal-liquid fractionation involving feldspars, zircon, apatite, and magnetite could result in a pegmatite melt depleted in Sr, Ba, Fe, HREE, Zr, and Hf. Zircon is a common accessory phase in the potassic series Pikes Peak granite, and Zr is the most depleted of any of the HFSE in the Wellington Lake pegmatite (Fig. 9; Table 6). However, the HFSE- and HREE-enriched South Platte NYF-type pegmatites also formed from the same Pikes Peak granite as the Wellington Lake pegmatite, and the Nd-isotopic compositions of their REE minerals are consistent with the derivation of the pegmatite magma from this granite (Fig. 7; Table 4), but without the HFSE and HREE depletion that would result from crystal-liquid fractionation involving zircon. For the formation of these pegmatites, Simmons and Heinrich (1980) invoked the segregation from the crystallizing granite of a hydrous fluid with a higher concentration of HFSE and HREE than the remaining silicate melt, due to its high F content (London, 2016), thus fractionating additional quantities of these elements into the pegmatite parental magma. We suggest that formation of the Wellington Lake pegmatite also involved an H₂O-rich silicate melt. This melt included components from a residual melt generated by crystal-liquid fractionation of the crystallizing PPG host. It also contained elements transferred into this melt by an
Table 6. Estimated bulk composition of the Wellington Lake pegmatite compared to the Pikes Peak Granite and corresponding fractional differences.

| Oxide     | WL Model Wt.% | Granite Wt.% | Diff. in % |
|-----------|----------------|--------------|------------|
| SiO₂      | 77.3           | 75.7         | +2.8       |
| TiO₂      | 0.0334         | 0.201        | -83.7      |
| Al₂O₃     | 12.2           | 11.9         | +2.1       |
| Fe₂O₃     | 0.859          | 2.50         | -66.4      |
| MnO       | 0.00515        | 0.0453       | -88.8      |
| MgO       | 0.0159         | 0.0235       | -33.3      |
| CaO       | 0.256          | 0.859        | -70.7      |
| Na₂O      | 2.75           | 2.99         | -8.8       |
| K₂O       | 6.42           | 5.47         | +16.8      |
| P₂O₅      | 0.00432        | 0.0101       | -58.4      |
| Element   |                |              |            |
| Ga        | 10.8           | 10.2         | +4.5       |
| Cs        | 0.94           | 1.39         | -33.2      |
| Ba        | 75.1           | 210          | -64.8      |
| Rb        | 406            | 246          | 63.5       |
| Sr        | 8.35           | 18.1         | -54.5      |
| Zr        | 72.5           | 532          | -86.7      |
| Y         | 26.7           | 132          | -80.2      |
| Sm        | 3.36           | 5.71         | -42.2      |
| Nb        | 23.0           | 56.4         | -60.2      |
| Th        | 12.9           | 32.2         | -60.9      |
| Hf        | 3.21           | 16.2         | -80.6      |
| U         | 4.34           | 6.41         | -34.1      |
| La        | 70.1           | 148          | -54.1      |
| Ce        | 133            | 315          | -59.1      |
| Pr        | 14.3           | 36.7         | -62.3      |
| Nd        | 46.6           | 134          | -66.3      |
| Sm        | 7.82           | 26.1         | -70.9      |
| Eu        | 0.38           | 0.84         | -55.5      |
| Gd        | 5.05           | 23.8         | -79.4      |
| Tb        | 0.78           | 3.92         | -80.5      |
| Dy        | 4.51           | 24.3         | -81.9      |
| Ho        | 0.92           | 4.80         | -81.3      |
| Er        | 2.85           | 13.6         | -79.4      |
| Tm        | 0.49           | 2.03         | -76.5      |
| Yb        | 3.42           | 12.7         | -73.6      |
| Lu        | 0.54           | 1.84         | -71.1      |

H₂O-rich, but in this case F-poor fluid phase, depleted in elements such as HFSE and REE that are less soluble in an F-poor compared to an F-rich aqueous fluid. This would account for the depletion in the Wellington Lake pegmatite of, for example, Ti and Nb, elements that are not sequestered by the crystallization of zircon or other accessory phases in the PPG.

In contrast to the Wellington Lake pegmatite, the nearby large McGuire pegmatite is enriched in F, Ti, and Nb with abundant fluorite and topaz, and lesser monazite, ilmenite, and niobian rutile (Černý et al., 1999). This pegmatite exhibits a less-developed concentric zoning and lacks the miarolitic cavities in the core that characterizes the Wellington Lake pegmatite, and its bulk composition is more similar to the large NYF-type pegmatites of the South Platte district. For the McGuire pegmatite, an inward crystallization model of progressively more F- and HFSE-rich melt can explain the late crystallization of the ilmenite and niobian rutile. In the Wellington Lake pegmatite, the observed progressive inward decrease in LREE/HREE of the intermediate zone and so-called “replacement” associations suggest an opposite trend. The trend in the Wellington Lake
pegmatite is of a continuous LREE enrichment during fractional crystallization of an H$_2$O-rich and F-poor melt. This results in the final and highly localized crystallization of the LREE minerals fluorite and bastnasite from only a small amount of residual F- and CO$_2$-rich solution remaining at the top of the pegmatite near the contact of the core and intermediate zones. The late-stage transition of fluorite to bastnasite growth in the Wellington Lake pegmatite suggests an increase in pH and/or decrease in F/CO$_2$ activity ratio (William-Jones and Wood, 1992). The REE zonation seen in the bastnasite may involve selective complexation of certain REE with CO$_2$. The oscillatory zonation observed in the bastnasite has been seen in late-stage rare-earth element minerals in pegmatites around the world, and may be indicative of locally buffered, rapidly changing chemical environment at the final stages of crystallization (William-Jones and Wood, 1992; Gysi et al., 2016).

**Fluid Inclusions**

Fluid inclusions at the Wellington Lake pegmatite indicate the presence of at least two compositionally distinct types of fluids: H$_2$O-NaCl and H$_2$O-CO$_2$. H$_2$O-NaCl fluid inclusions are only rarely saturated with respect to NaCl, as indicated by the few halite-bearing inclusions. It is possible that the rare higher temperature, high salinity halite-bearing type C, and low salinity vapor-rich type B fluid inclusions represent a distinct earlier fluid of higher temperature boiling fluids as suggested by Simmons and Heinrich (1980), who also observed halite-bearing primary fluid inclusions with homogenization temperatures > 400°C in other South Platte pegmatites. The type-A liquid-rich H$_2$O-NaCl fluid inclusions have intermediate salinities and lower temperatures of homogenization.

The inclusions observed in the three quartz samples from Wellington Lake also show some thermometric and compositional similarities and differences with inclusions described by Gagnon et al. (2004) for the Oregon #3 pegmatite in the South Platte region. In both pegmatites, type A two-phase, liquid and vapor, liquid-rich, and also type C three-phase vapor, liquid, and halite inclusions were identified. A third additional type of inclusion rich in CO$_2$ was also recognized in Wellington Lake, which was not described by Gagnon et al. (2004) for the Oregon #3 pegmatite, but was noted by Simmons and Heinrich (1980) in other South Platte pegmatites. Most significantly, a large proportion of the inclusions from Wellington Lake pegmatite were trapped from solutions of the H$_2$O-NaCl ± KCl type as indicated by their eutectic point, which is close to -21°C. In contrast, those from Oregon #3-involved fluids can be characterized as more complex, including both Na + K + Sr + Ba and Na + K + Sr + Ba + Ca solutions as determined by laser ablation-ICP-MS analysis. The Oregon #3 pegmatite also contains fluorite crystals that, according to Gagnon et al. (2004), are associated with REE mineralization, whereas fluorite is not present at Wellington Lake. Fluorite crystals in Oregon #3 also have liquid + vapor inclusions. These contain solutions with Na + K + Sr + Ba as well as REE, Y, Th, and U as measured by LA-ICP-MS analysis, and are, therefore, referred to by Gagnon et al. (2004) as type F4 inclusions.

The two high-temperature types (B and C) of co-existing inclusions are interpreted to be the result of boiling of primary fluids—fluids that precede the ones responsible for lower temperature A-type inclusions (Fig. 8A) as suggested by Simmons and Heinrich (1980) for South Platte district pegmatites. The temperature of all the inclusions in the Wellington Lake pegmatite, with a maximum of 488°C when corrected for pressure, suggests that crystallization took place well below the solidus temperature of an H$_2$O-saturated granitic magma. This is consistent with recent models of pegmatite petrogenesis leading to nucleation-controlled mega-crystal growth resulting from supercooling (London, 2008). The undercooling and associated nucleation delay also may explain the presence of columbite in the Wellington Lake pegmatite, despite its low total estimated Nb content of 22 ppm (Table 6). As suggested by London (2016), a pegmatite fluid with < 50 ppm Nb could become saturated and crystallize columbite only below 525°C.

**CONCLUSIONS**

We postulate, based on geochemical, mineralogical, and textural evidence, that the Wellington Lake pegmatite formed within the PPG from an H$_2$O-rich, but F-poor melt pocket that underwent crystallization associated with supercooling, as suggested to be critical to the development of the characteristic textures and mineralogical zoning of pegmatites (London, 2008, 2014, 2016, 2018). Although the pegmatite melt was derived from the parental PPG magma and had Si, Al, Na, and K contents similar to the PPG magma, it was depleted in other elements such as REE and HFSE, which have low solubility in F-poor silicate melts and aqueous fluids. Processes of zone refinement in the inward-crystallizing pegmatite (London, 2018) could explain the mineralogical zonation observed in the Wellington Lake pegmatite. F and CO$_2$ played a role only at the last stages of crystallization, when the LREE fluoro and carbonate minerals in the pegmatite core zone crystallized from a low-density silicate melt containing the necessary F, CO$_2$, and H$_2$O to form these phases as well as create open void spaces in the quartz core.

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