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<td><strong>Authors(s)</strong></td>
<td>Barros, Renata, Menuge, Julian</td>
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THE ORIGIN OF SPODUMENE PEGMATITES ASSOCIATED WITH THE LEINSTER GRANITE IN SOUTHEAST IRELAND

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ABSTRACT

Rare-element pegmatites have diverse chemical signatures and are important sources of strategic metals such as Li, Cs and Ta. The two main hypotheses to explain rare-element pegmatite formation are 1) residual magmas from granitic rocks’ crystallization, and 2) partial melts from a relatively rare-element-rich source. In southeast Ireland, spodumene and spodumene-free pegmatite dykes occur along the eastern margin of the S-type Leinster Granite batholith. With indistinguishable emplacement ages around 400 Ma, the origin of the Li-rich pegmatitic fluids has been suggested to have resulted from extreme fractional crystallization of Leinster Granite granodiorite magma. To test this hypothesis, we used whole-rock geochemistry of pegmatite and granodiorite samples from drill cores and geochemical modeling of in situ crystallization and batch melting to investigate which process better explains the formation of pegmatites. Chemical signatures for pegmatites and granodiorite do not indicate a direct comagmatic relationship, as granodiorite has higher concentrations of many incompatible elements than the pegmatites (e.g. concentrations of Zr, Ti and Y). Concentrations of Li, Rb, Cs, Sr and Ba show no clear fractionation trends from granodiorite to pegmatite. The in situ crystallization model using the average granodiorite composition as initial magma generates a range of compositions that does not include pegmatites, so it is unlikely that they represent residual granitic magmas. Modeling of partial melting indicates that Leinster Granite granodiorite and pegmatite magmas could have been formed in separate events and from chemically different source rocks, with pegmatite magmas presumably formed in a younger event because pegmatites intrude granodiorite.

KEYWORDS
Spodumene pegmatite; Leinster Granite; geochemical modeling; in situ crystallization; batch melting; petrogenesis.

INTRODUCTION
Rare-element pegmatites display very diverse chemical signatures and are enriched in elements that are usually present in low concentrations in other rocks. These features make them important sources of a wide range of commodities, including industrial minerals, gemstones and strategic metals. Two alternative petrogenetic processes have been proposed to explain the formation of rare-element pegmatites: 1) extensive fractionation of fertile parental granitic magma, with rare-element pegmatites representing the residual phase in this process; or 2) partial melting of rare-element-rich source rocks. Pegmatites geographically associated with granitic batholiths have often led to support for the first hypothesis that granitic rocks and pegmatite magmas have a comagmatic relationship. Several studies have successfully demonstrated the link between parent granite and pegmatites, including examples in Canada (Černý et al. 2012) and Spain (Roda-Robles et al. 2012). However age and/or geochemical incompatibilities, such as pegmatites being significantly older than granite, or an apparent absence of parental granite, are often documented (e.g. Walker et al. 1989, Muller et al. 2015).

One way to approach and resolve the petrogenetic problem is to use quantitative trace element modeling and mathematically test different models. We use this approach for a pegmatite belt in southeast Ireland, which is associated with the peraluminous Leinster Granite. A genetic link between them has been suggested based on age and geochemical similarities (O’Connor et al. 1991, Whitworth & Rankin 1989), but the exact relationship between them remains uncertain. As a result of new lithium exploration drilling being carried by Blackstairs Lithium Ltd., an extensive lithogeochemical database is available that provides drill core of the entire thickness of several pegmatite dykes. Coupled with textural and mineralogical observations on the drill core, these are used for geochemical modeling to better constrain the origin of these rare metal pegmatites.

GEOLOGICAL SETTING AND PREVIOUS STUDIES

The geology of the southeast of Ireland is dominated by Paleozoic basement and igneous intrusions (Fig. 1). The basement is composed of sedimentary and associated volcanic rocks regionally metamorphosed to lower greenschist facies and is subdivided into the following units and lithologies: greywacke and shale of the Cambrian Bray and Cahore Groups; mudstone, siltstone and sandstone of the Ordovician Ribband Group; deep marine shale, now commonly slate, with abundant intermediate to acidic volcanic rocks of the Ordovician Duncannon Group; and the deep marine turbidite sequence of the Silurian Leinster Inlier (Graham & Stillman 2009, Holland 2009).

The largest of the younger igneous intrusions is the Leinster Granite, a Caledonian batholith composed at least in part of sheeted intrusions (Grogan & Reavy 2002) of S-type two-mica granitic rocks (predominantly granodiorite) intruded into the metasedimentary basement. The batholith is part of the Trans-Suture Suite (Brown et al. 2008), which spans the trace of the Iapetus suture in Britain.
and Ireland. According to the same authors, the origin of these granitic magmas is related to orogen-wide sinistral transtension during the early Devonian, triggering lamprophyre magma generated by melting of Avalonian lithospheric mantle to rise and transfer heat to the lower part of the Avalonian crust, forming partial melts with S-type granitic composition. The Leinster Granite comprises five plutons aligned with the NE-SW regional strike of the Lower Paleozoic supracrustal country rocks (McConnell & Philcox 1994). Despite its size, the batholith is poorly exposed. Most available data are from the better exposed Northern pluton, including a U-Pb monazite age of 405 ± 2 Ma (O’Connor et al. 1989). It has been assumed that all plutons intruded and crystallized simultaneously (Brindley 1973) and therefore that this age applies to the whole batholith. However a wide variability in initial Sr and Nd isotope ratios, both between and within plutons, has been interpreted as resulting from the generation of different magma batches by partial melting of an isotopically heterogeneous sedimentary source (Mohr 1991). In addition, disequilibrium textures in plagioclase phenocrysts indicate a multi-stage and multi-scale acid-acid mixing of magma batches during ascent and emplacement (Grogan & Reavy 2002).

A network of lithium pegmatite dykes, metres to tens of metres thick, is known from at least nine localities along the eastern margin of the Tullow Lowlands pluton and within the schist septum that separates it from the Blackstairs Unit to the south (Fig. 1). There are almost no outcrops of these dykes and the localities have been identified by concentrations of glacial boulders, in several cases confirmed by mineral exploration drilling (Steiger & von Knorring 1974, Steiger 1977). Most of the dykes so far detected roughly follow the NE-SW regional structure of the Caledonian East Carlow Deformation Zone (McArdle & Kennedy 1985). They intrude the Tullow pluton, minor granitic bodies related to the Leinster Granite and quartz-mica schist formed by contact metamorphism of the Leinster Granite with the Ribband Group. Rb-Sr isotope studies of the pegmatites (O’Connor et al. 1991) yielded a whole-rock errorchron age of 396 ± 7 Ma (MSWD = 5.54), constructed using spodumene pegmatites, associated lepidolite greisens and barren pegmatites. This corresponds to an age of 402 ± 7 Ma when recalculated using the recent IUGS recommendation on 87Rb half life (Villa et al. 2015). The pegmatite errorchron initial Sr ratio of 0.705 ± 0.005 (O’Connor et al. 1991) lies within the Leinster Granite range obtained by Mohr (1991). Moreover, results from fluid inclusion analysis (Whitworth & Rankin 1989) and mineral chemistry (Whitworth 1992) suggest a genetic link between the Leinster Granite and lithium pegmatites. Despite the indistinguishable Caledonian ages of granitic rocks and pegmatites and their similar initial Sr isotope ratios, it is debatable whether lithium pegmatites represent the residual magma after crystallization of the Leinster Granite, or were formed from separate partial melts of similar or different metasedimentary sources. Here we aim to better constrain the petrogenesis of the pegmatite belt and its relationship with the Leinster Granite.
Fig 1: Spodumene pegmatite occurrences in southeast Ireland. Dykes are restricted to the eastern margin of the Tullow Lowlands pluton, spatially associated with the East Carlow Deformation Zone (ECDZ). Pegmatites intrude both the Tullow pluton and metasedimentary rocks of the Ribband Group.

CHARACTERIZING THE LEINSTHER RARE-ELEMENT PEGMATITE BELT AND COUNTRY ROCKS

A recent mineral exploration drilling programme has provided an opportunity to study the Leinster spodumene pegmatites, barren (spodumene-free) pegmatites and their immediate country rocks. The rock types identified in the drill cores in the study areas of Aclare and Moylisha (Fig. 1) are:

1. Spodumene pegmatite: coarse-grained (crystals up to 10 cm), composed of spodumene (10-40%), albite (25-35%), quartz (15-20%), Li-muscovite (10-15%), spessartine (5%) and minor K-
feldspar, apatite, cassiterite and sphalerite (all less than 5%). Spodumene, albite and Li-muscovite are present as well-formed subhedral crystals, while quartz is interstitial. The dykes can be unzoned, or present zoning when thicker than 10 meters, with an increase in modal spodumene towards the quartz and feldspar core (Fig. 2). In some dykes, crystals of spodumene are aligned perpendicular, and sometimes parallel, to the dyke margins. These observations suggest that each pegmatite dyke has formed by the progressive \textit{in situ} crystallization of a single body of magma. Fine grained albite-rich (85-90%) aplitic patches are common within all zones and include around 10% combined quartz and muscovite, with accessory apatite, cassiterite, spessartine, beryl and Mn,Fe-phosphate.

(2) Barren (spodumene-free) pegmatite: coarse-grained (typically 4 cm, maximum 10 cm), consisting of K-feldspar (10-20%), albite (20-30%), quartz (20-35%), muscovite (15-20%) and spessartine (2-5%). K-feldspar and albite occur as subhedral prismatic crystals, with later quartz and muscovite. Quartz-feldspar intergrowth (graphic texture) is common.

(3) Granodiorite (Leinster Granite): medium-grained granodiorite consisting of quartz (30-45%), oligoclase (30-40%), microcline (10-20%), biotite (5-10%) and muscovite (2.5%), locally porphyritic with occasional 2 cm plagioclase phenocrysts. Sparse tourmaline is observed in some of the contact zones with spodumene pegmatite and its veinlets.

(4) Quartz-mica schist (Ribband Group): fine- to medium-grained schist consisting of biotite (40-50%), muscovite (20-30%), quartz (20-25%) and minor arsenopyrite, titanite and apatite, with a foliation defined by mica; late tourmaline porphyroblasts with poikiloblastic texture are present up to 20 cm from the contact with spodumene pegmatite.

Spodumene pegmatite dykes crosscut both granodiorite and quartz-mica schist in Aclare, and granodiorite only in Moylisha. The contacts between spodumene pegmatite and Leinster Granite are defined by irregular planes and narrow (up to 10 cm) zones of interaction between the two rock types. Contacts between spodumene pegmatite and mica schist are sharp and parallel to the foliation in the mica schist. Spodumene-free pegmatites occur only within granodiorite, against which their contacts are diffuse.
Fig. 2: Mineral and chemical profiles of zoned spodumene pegmatite (drill core ACL13-04). Modal spodumene (open rectangles) increases towards the quartz-feldspar core. Albite-rich aplite occurs as patches throughout the dyke. Whole-rock chemical compositions measured for homogeneous intervals of rock correlate with the mineralogical variation.

METHODS

Sampling and geochemical analysis

Eight mineral exploration drill cores with close to 100% recovery from two localities, Aclare (6 cm diameter) and Moylisha (4 cm diameter), were split in half and divided into homogeneous rock parts, between 7 cm and 3.05 m long, resulting in 281 samples. These parts were then crushed, decomposed using a four acid digestion and analysed for 48 elements by ICP-MS by ALS Minerals (Loughrea, Co. Galway, Ireland). Routine practices were used to ensure data quality control: sample duplicates (1 in every 20 samples), homogeneous quartz pebbles (1/40) and certified standards (1/20). Results showed reproducibility between duplicates within 15% for most elements, except Ta, Zr and Ce, and no contamination problems. Detection limits for the elements analyzed range between 0.002 and 100 ppm. Whole-rock geochemical analysis of pegmatites can be problematic because of their large crystal size, demanding large volume samples for representative results. However, the pegmatites sampled from drill core in the present study have a typical grain size of around 2 cm, with
the largest, rare, crystals having grain sizes less than 10 cm, which facilitates representative whole-
rock sampling.

Pegmatite drill cores were subdivided for whole-rock analysis by lithology into aplite, spodumene-
rich pegmatite, pegmatite without spodumene, granodiorite and quartz-mica schist. Weighted mean
compositions of each rock type were then calculated for each drill core. For example, in a drill core in
which granodiorite constituted a total length of 4.8 m, three samples of lengths 1.6 m, 1.2 m and 2.0 m
were analysed. Mean concentrations were then calculated by weighting analyses proportional to the
rock volumes in these three core lengths. In the case of pegmatite dykes, the calculation comprised all
the intervals from the upper to the lower contact with the country rocks. Contact zones with visible
and/or chemical haloes in country rocks and granitic lenses within pegmatites were excluded from the
estimated bulk compositions. The method assumes that the drill core samples are representative of the
pegmatite bodies as a whole and therefore does not allow for possible chemical variation along strike
or down dip within the dykes. It is also assumed that each pegmatite dyke crystallized from a single
batch of magma and that bulk pegmatite concentrations of the key elements remained constant during
and after crystallization. To highlight possible heterogeneities within rock types between localitites,
whole-rock data is presented as the weighted mean of each rock type per drill core, for Aclare and
Moylisha separately.

Modeling of pegmatite origin by in situ crystallization of granitic magma

Calculated bulk concentrations of key elements were used to test whether the Leinster pegmatites
could have been derived as a residual phase of continuous crystallization of Leinster Granite
granodiorite magma. The elements Ba, Sr, Li, Rb and Cs were chosen as petrogenetic tracers, the first
two as compatible elements in granitic minerals and the others as incompatible in granodioritic
magma but enriched in pegmatites. To estimate the mean concentrations of these key elements for the
Tullow Lowlands pluton, the overall mean of all granitic intervals in all drill cores was considered,
weighted by volume as core diameters are different for drill core from Aclare and Moylisha. The same
was done for pegmatites and schist, to obtain representative concentrations assuming intrusion of one
batch of Li-pegmatite magma and one batch of barren pegmatite magma per dyke.

The equation for Rayleigh fractionation has frequently been used for trace element modeling of the
formation of pegmatites from granitic magmas (e.g. Shearer et al. 1992, Roda-Robles et al. 2012).
However, the idea of crystals separating from melt through gravity is most likely only applicable to
ultramafic melts (Rollinson 1993). In granites and pegmatites solidification most likely starts from the
intrusion margins inwards and in pegmatites experimental data suggest that this is a rapid process (e.g.
is the quantitative in situ crystallization model proposed by Langmuir (1989), in which a solidification
zone occurs between the crystallized margins and the central convecting magma batch in a chamber.
Langmuir (1989) considers the case where once solidification starts, all liquid remaining in the solidification front is ultimately separated from the crystal mush and returns to the interior of the magma chamber, which is a reasonable approximation of the current understanding of granite and pegmatite crystallization, and from this assumption the derived equation is 

\[
\frac{C_L}{C_0} = \left(\frac{M_L}{M_0}\right)^{f(D-1)/[D(1-f)+f]}
\]

where \(C_L\) = concentration of the element in the differentiated magma; \(C_0\) = initial concentration of the element in the magma; \(M_L\) = mass of the differentiated magma; \(M_0\) = initial mass of the magma chamber; \(f\) = fraction of magma allocated to the solidification zone which returns to the unFractionated magma chamber; and \(D\) = bulk partition coefficient. The bulk partition coefficient was calculated through the equation 

\[
D = \Sigma (X_A K_{dA} + X_B K_{dB} + X_C K_{dC} + \ldots),
\]

where \(X_A\) = weight fraction of mineral \(A\) in the rock and \(K_{dA}\) = mineral \(A\) / liquid partition coefficient. This \textit{in situ} crystallization equation also yields the result for simple Rayleigh fractionation, for the condition \(f = 1\).

\textbf{Modeling of pegmatite origin by partial melting}

Direct partial melting of a sedimentary source has previously been proposed as an alternative mechanism to form rare element pegmatite magma (e.g. Jolliff \textit{et al.} 1992, Shearer \textit{et al.} 1992). The most suitable process to generate the felsic melts discussed is batch melting (Shaw 1970), that can be modelled by the equation 

\[
\frac{C_L}{C_0} = \frac{1}{[D_{RS} + F(1 - D_{RS})]},
\]

where \(C_L\) = concentration of the element in the generated melt; \(C_0\) = concentration of the element in the unmelted source; \(D_{RS}\) = bulk partition coefficient of the residual solid; and \(F\) = weight fraction of melt produced. The bulk partition coefficient is calculated in the same way as for the \textit{in situ} crystallization calculations. Minimum PT conditions of partial melting can be estimated based on the minimum temperature of 700 °C to produce granitic composition melts (MacRae & Nesbitt 1980) and the spodumene stability field, yielding a minimum pressure around 350 MPa. Conditions considered are in agreement with the thermal model for generation of S-type granitic magmas within a transtensional pull-apart zone proposed by Brown \textit{et al.} (2008).

\textbf{RESULTS}

\textbf{Whole-rock geochemistry}

Ranges and means of whole-rock trace and major element concentrations for the four rock types and the relevant study areas are given in Table 1. The granodiorite is classified as peraluminous using the index of Shand (1943) and carries the signature of syn-collisional granites according to the trace element discrimination diagram of Pearce \textit{et al.} (1984) (Fig. 3). Granodiorite in Aclare has lower alkali contents thus higher A/CNK and slightly lower Y and Nb contents when compared to the same rock type in Moylisha. Both types of pegmatites are also peraluminous and syn-collisional, but
spodumene-free pegmatites exhibit the greater scatter in Rb, Y and Nb values. Spodumene pegmatites in Aclare present a high scatter of A/CNK values and higher than Moylisha.

Multi-element diagrams per locality are presented on Fig. 4. Average trace element concentrations for granodiorite show relative abundances decreasing towards the more incompatible elements to the right of the plot and a strong negative Nb anomaly. Spodumene pegmatites and barren pegmatites present a multi-peak signature, with low Ba, Th, La, Ce, Sr and Zr and Y, strong negative Ti anomaly and positive Rb, Ta and Hf anomalies. In comparison with both types of pegmatites, the Leinster Granite granodiorite is enriched in the LIL elements Ba and Sr, which are compatible in feldspars, and in the light rare-earth elements La and Ce.

As Li is a suitable element to differentiate the three granitic rock types (granodiorite, spodumene free pegmatite and spodumene pegmatite), incompatible and compatible elements were plotted against this element (Fig. 5). The three rock types cluster separately without well-defined trends between them. Spodumene pegmatites in Aclare have higher mean Li when compared with the same rock type in Moylisha, while spodumene-free pegmatites in Moylisha present a higher mean Li concentrations than those from Aclare. Granodiorite has the lowest mean Rb concentration and highest mean Sr and Ba, while concentrations of Li and Cs are intermediate between spodumene-free and spodumene pegmatites.

The ratio K/Rb versus Cs is controlled by K-feldspar and micas and is a useful petrogenetic index in granite-pegmatite systems (Černý et al. 1981) and values for the studied rock types are shown on Fig. 6. Granodiorites present the highest K/Rb ratios (50-120) and can be locally enriched in Cs as represented by one sample from Aclare. Spodumene pegmatites and barren pegmatites show lower K/Rb ratios of 20-60 and 20-90, respectively. The data do not fit on the expected evolution trend for the case of a parental granite magma fractionating to residual pegmatitic magmas.
Table 1: Geochemical data for the studied rock types in the localities Aclare and Moylisha. Values in ppm. LOD = limit of detection. Duplicate analysis of 15 of the samples shows maximum percentage differences between the higher and lower concentration presented in the last column of the data table.

<table>
<thead>
<tr>
<th>LOC</th>
<th>Quartz mica-schist</th>
<th>Granodiorite</th>
<th>Spodumene pegmatite</th>
<th>Spodumene-free pegmatite</th>
<th>Granodiorite</th>
<th>Moylisha</th>
<th>Spodumene pegmatite</th>
<th>Spodumene-free pegmatite</th>
<th>Estimate of reproducibility (%)</th>
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Values indicate minimum and maximum concentrations calculated for rock types in each drill core and the mean value is calculated as the mean of all analysed samples from each locality weighted by volume. The volumes of core that contributed to these calculated mean concentrations are: quartz-mica schist – Aclare and total 0.075 m³; granodiorite – Aclare 0.11 m³, Moylisha 0.13 m³, total 0.24 m³; spodumene pegmatite – Aclare 0.15 m³, Moylisha 0.05 m³, total 0.20 m³; spodumene-free pegmatite – Aclare 0.02 m³, Moylisha 0.04 m³, total 0.06 m³.
Fig. 3: A) A/CNK versus A/NK plot (Shand 1943) showing the peraluminous character of granodiorite and pegmatites. B) Trace element discrimination diagram of Pearce et al. (1984) showing...
Fig. 4: Trace element concentrations are normalized to the composition of chondritic meteorites given by Thompson (1982). Samples in black are from Aclare and in grey from Moylisha. The diagrams are for A) quartz-mica schist; B) granodiorite; C) spodumene pegmatites and D) spodumene-free pegmatites. Gaps indicate no available data.
Fig. 5: Bivariate plots of Li versus A) Rb, B) Sr, C) Cs and D) Ba. Samples in black are from Aclare and in grey from Moylisha.
Fig. 6: Cs versus K/Rb as an index of chemical evolution of pegmatites. (*) For comparison, the arrow indicates data presented in London (2008) for variation within a granite-pegmatite group (Red Cross Lake, Manitoba) with a proven petrogenetic link.

Partition coefficients used in modeling

Modeling of both *in situ* crystallization and batch melting requires calculation of bulk partition coefficients. As already debated in other pegmatite modeling studies (e.g. Jolliff *et al.* 1992, Shearer *et al.* 1992), uncertainty of the appropriate values of bulk partition coefficients can arise for several reasons. Several variables can affect mineral/melt partition coefficients, including temperature, confining pressure and melt composition (Rollinson 1993), but none of these are considered to be major issues in the studied granite-pegmatite systems. The partition coefficients and their published sources chosen for the key elements in granitic and pegmatitic minerals are presented in Table 2. Values were chosen to be representative of high-Si crystallizing magmas, consistent with the predominant granodiorite in Leinster Granite, and maximum values available were used for
calculations in cases where the reference presented a range. Bulk partition coefficients were calculated considering the mineral assemblages discussed below.

Table 2: Crystal / melt partition coefficients used for quantitative modeling.

<table>
<thead>
<tr>
<th>Mineral/melt</th>
<th>Albite</th>
<th>Muscovite</th>
<th>Quartz &amp; aluminosilicate</th>
<th>K-feldspar</th>
<th>Biotite</th>
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<tr>
<td>Li</td>
<td>0.1</td>
<td>1.67</td>
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<td>0.05</td>
<td>1.65</td>
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</tr>
<tr>
<td>Rb</td>
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<td>1.75</td>
<td>0.016</td>
<td>0.74</td>
<td>5.3</td>
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<tr>
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<td>3</td>
<td>7</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Cs</td>
<td>0.44</td>
<td>0.24</td>
<td>0.044</td>
<td>0.13</td>
<td>3.1</td>
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<td>7</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Sr</td>
<td>3.31</td>
<td>0.5</td>
<td>0.01</td>
<td>5</td>
<td>0.06</td>
</tr>
<tr>
<td>reference</td>
<td>4</td>
<td>3</td>
<td>assumed</td>
<td>5</td>
<td>3</td>
</tr>
<tr>
<td>Ba</td>
<td>0.19</td>
<td>5.5</td>
<td>0.015</td>
<td>6.7</td>
<td>7</td>
</tr>
<tr>
<td>reference</td>
<td>6</td>
<td>3</td>
<td>7</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

References: 1 Mahood & Hildreth (1983); 2 Icenhower & London (1996); 3 Icenhower & London (1995); 4 Drake & Weill (1975); 5 Long (1978); 6 Bea et al. (1994); 7 Nash & Crecraft (1985); 8 Jollif et al. (1992); 9 Walker et al. (1989).

In situ crystallization modeling

In situ crystallization calculations test the hypothesis that pegmatites and granodiorite are genetically linked through a continuous evolution model in a closed system (Fig. 7). Whole-rock concentrations of Li, Rb, Cs, Sr and Ba were estimated for granodiorite, spodumene pegmatite, barren pegmatite and schist as means weighted by total drill core volume of all analyses from all drill cores, for each rock type. These means for granodiorite are then considered to represent the initial magma from which residual pegmatitic fluids might have evolved in the late stages of crystallization, and subsequently escaped to be intruded as dykes; concentrations for schist are presented for comparison only in this scenario. Fractionation paths with various combinations of parameters were calculated, to show whether it is possible to reach bulk concentrations for the residual magma that are similar to mean spodumene pegmatite and barren pegmatite chemical compositions. The crystallizing assemblage considered is plagioclase (40%), K-feldspar (25%), quartz (25%), biotite (5%) and white mica (5%).

In in situ crystallization, the enrichment of a trace element in the residual melt relative to the parent melt \((C_L/C_0)\) varies according to the mass of residual melt relative to the total mass of the magma chamber \((M_L/M_0)\), the fraction of liquid returned to the main magma body from the
solidification zone \((f)\) and the bulk partition coefficient \((D)\). From lithogeochemical data, the values of \(C_0\) were considered to be the bulk trace element concentration calculated for granodiorite as the initial magma. The fraction \(M_L/M_0\) is equivalent to the fraction of residual magma and values between 0.01 (99% residual magma, start of crystallization) and 0.99 (1% residual magma, end of crystallization) were considered. Values of \(f\) range between 0.01 (1% liquid returned from solidification zone, representing very rapid crystallization) and 1 (all liquid returns from the solidification zone, representing Rayleigh fractionation). The bulk partition coefficient, and therefore different values of partition coefficients for the elements studied, can contribute substantially to variations in the final result and this will be discussed later.

The ranges of possible enrichment and depletion of the five key elements, with granodiorite as model initial magma, are plotted on Fig. 8. Average concentrations for key elements in pegmatites are not always within the ranges of possible concentrations reached by \textit{in situ} crystallization and the maximum limiting Rayleigh fractionation. The magnitude of depletion in Ba and Sr and enrichment in Rb and Li by \textit{in situ} crystallization is sufficient to reach the mean composition of spodumene pegmatite. However, very different crystallization conditions are required to individually explain the elements analysed; for example ~95% crystallization at \(f\) ~0.8 can produce mean spodumene pegmatite composition on the Ba-Li plot, whereas at least 99% crystallization is required at \(f\) ~0.3 to produce mean spodumene pegmatite on the Rb-Li plot; there are no conditions which can account for the mean Cs concentration of the spodumene pegmatites. Spodumene-free pegmatite compositions are even less well reproduced by in situ crystallization (Fig. 8).
Fig. 7: *In situ* crystallization model (Langmuir 1989) applied to the Leinster Granite granodiorite - pegmatites system, depicting continuous granodiorite-to-pegmatites evolution within the same magma batch. A) Solidification starts from the margins of the magma chamber inwards, with a fraction of magma returning to the main magma body. The initial magma’s composition is equivalent to the bulk composition of the crystallized granodiorite. B) After extensive crystallization, the residual magma would have become concentrated in incompatible elements and have a composition equivalent to the most evolved bulk spodumene pegmatite dykes. The same process could also generate the (less evolved) spodumene-free pegmatites as an intermediate member or as another end-member.
Fig. 8: Bivariate plots with the results of *in situ* crystallization modeling with the average Leinster Granite granodiorite as the initial magma. The arrows show the range from minimum (10%) to maximum (100%) fractions of liquid returned (f) to the main magma body from the solidification zone considered in this system – 100% returned is equivalent to Rayleigh fractionation. The grey area shows the range of values with other values of f between 0.1 and 1.

**Batch melting modeling**

The range of suitable initial compositions of sedimentary sources that underwent batch partial melting forming the three rock types were also calculated to assess whether a common origin for them is possible. In batch melting, the concentration of an element in the generated melt compared with the concentration of the element in the unmelted source \( \frac{C_f}{C_0} \) varies with the weight fraction of melt.
produced (F) and the bulk partition coefficient of the residual solid in the source (D_{RS}). The weighted mean concentrations measured in the three rock types were considered as different values for C_L, allowing calculation of the range of necessary concentrations in the source C_0 to generate melts with the measured compositions. The range of initial concentrations was calculated from 1% melting (F = 0.01) to 99% melting (F = 0.99). In order to calculate D_{RS}, the source was estimated to be quartz feldspar-rich sediment, since partial melting of pelitic sediments would result in Li, Rb and Cs being retained in the mica-rich residual solid as partition coefficients are high for these elements (Jolliff et al. 1992 and references therein). The residual solid considered would be mainly composed of quartz (25%), feldspars (50%) and aluminosilicate (25%), thus allowing for all mica to be melted. Variations in the percentages of these minerals in the residual solid and from the partition coefficients chosen do not have major impact on the final results.

Possible initial concentrations in sources that can form melts with the compositions of average granodiorite and pegmatites are featured on Fig. 9. Considering a range of 1 to 40% of partial melting to form the three rock types (1 to 10% for pegmatites and 10 to 40% for granodiorite), the lines of initial concentrations converge to a restricted area representing little variation in the source for Rb (50 to 130 ppm), Li (30 to 430 ppm) and Cs (around 10 ppm). For Ba and Sr, initial concentrations converge to restricted areas for pegmatites (30 to 90 ppm Ba, 30 to 60 ppm Sr), but indicate a source more enriched in those elements for granodiorite (around 600 ppm Ba and 400 ppm Sr).
Fig. 9: Bivariate plots with the results of partial melting modeling. The arrows show the possible initial concentrations of the magma source rocks for spodumene pegmatite (line 1), spodumene-free pegmatite (line 2) and granodiorite (line 3) if they underwent 1 to 99% melting. Average composition of the schist is plotted for comparison, representing metasedimentary rocks of the Ribband Group.

IMPLICATIONS FOR SPODUMENE PEGMATITE FORMATION

The strong negative Nb anomaly in Leinster Granite granodiorite indicates that a crustal component is involved in its formation, which can also be inferred for pegmatites, but with a less pronounced negative anomaly. Pegmatites do not always carry higher concentrations of incompatible elements than granodiorite (e.g. concentrations of Zr, Ti and Y) and no clear fractionation trends are observed among incompatible or compatible elements, or ratios that indicate chemical evolution.
Plotted geochemical data is scattered rather than representing a chemically closed or semi-closed continuous evolution system. Although the key elements Li and Cs have not been analysed in samples from elsewhere in the Leinster Granite, the chemical signature of granodiorites within the batholith varies between localities, probably reflecting the multi-intrusion character of the batholith and some variation in source rock composition (Mohr 1991, Sweetman 1987, Grogan & Reavy 2002).

With the chosen partition coefficients it is not possible to reach spodumene pegmatite mean concentration considering all elements, therefore a single crystallizing system from granodiorite to pegmatite. If, however, the bulk partition coefficient for Ba (D_{Ba}) is double that listed in Table 2, by consequence of higher $K_{Ba}^{\text{biotite/liquid}}$ and $K_{Ba}^{\text{K-feldspar/liquid}}$, the required depletion of this element from granodiorite to pegmatites can be attained. Such high values were obtained experimentally by Icenhower & London (1996), for example. For Rb, a slightly lower bulk partition coefficient makes possible the enrichment observed in spodumene pegmatites. This would be mainly controlled by a lower $K_{Rb}^{\text{biotite/liquid}}$ and such lower values have been obtained experimentally (e.g. Icenhower & London 1995). However, the spodumene pegmatite Sr and Cs concentrations cannot be reached using reasonable partition coefficients for these elements. They could only be reached with a much higher Cs bulk partition coefficient. Extreme Li concentrations found in spodumene pegmatite can be reached towards high fraction of magma return from the solidification front, or Raleigh fractionation, and with less than 5% magma left, but this scenario would require the combination of cumulate-type crystallization followed by effective extraction of the residual magma. Therefore granodiorite-to-pegmatite magmatic evolution seems highly unlikely taking account of all five elements.

One possibility to explain the discrepancy observed when trying to correlate concentrations of key elements through the Langmuir in situ crystallization equation is a significant loss of the mobile elements Rb, Cs and Li at some stage between generation, ascent and emplacement of pegmatite magma. However, very narrow exomorphic haloes (up to 20 cm) into the granodiorite and the permeable quartz-mica schist do not support this hypothesis. Cs mobility does however seem possible as high-Cs granodiorite is observed (Figs. 5 and 6). The mean Cs concentration for the initial magma could be lower if this locally high-Cs granodiorite is considered to be part of an exomorphic halo (and therefore excluded from bulk composition calculations), but still considerably higher D_{Cs} would be needed to reach pegmatite concentrations. Another possibility is contamination from country rocks, especially contamination of spodumene pegmatites by the schist, but its mean composition with low Li (<250 ppm), Rb (<150 ppm) and Cs (<30 ppm) cannot explain the deviations of in situ crystallization trends away from pegmatite compositions.

On the other hand, with a narrow range of high source rock incompatible element concentrations and reasonable assumptions for degrees of partial melting, it is possible to form melts with the average composition of granodiorite, spodumene-free and spodumene pegmatites. However, source compatible element concentrations have to be highly variable if the three lithologies share a common source. As bulk concentrations of Ba and Sr would strongly depend on the amount of feldspars in
metasedimentary rocks, this variation can be explained by the presence of feldspar-rich and feldspar-poor source rocks, generating granodiorite and pegmatites, respectively. If pegmatite magmas were generated at the same time or later than the granodioritic magmas that formed the Leinster Granite, as implied by cross-cutting relationships, the required source rock heterogeneity likely existed on at least a kilometre scale. Otherwise, it is difficult to envisage the pegmatite source rocks remaining unmelted during Leinster Granite granodiorite magma formation.

CONCLUSIONS AND FUTURE WORK

Observing patterns on multi-element diagrams and bivariate plots, the behaviour of incompatible elements and the absence of fractionation trends among elements, the hypothesis that pegmatite magmas formed by the continuous evolution of the same magmatic system as the Leinster Granite granodiorite is not a satisfactory explanation for the origin of these rocks. Specifically, in situ crystallization modeling shows that it is highly unlikely that pegmatites represent the residual magma from crystallization of Leinster Granite granodiorite. This requires that granodiorite and pegmatite magmas were derived in separate partial melting events, the pegmatites presumably representing a later event because they are observed to intrude granodiorite. However the source rocks required to account for the geochemistry of pegmatites and the Leinster Granite granodiorite must be different, with considerably more feldspar required in the granodiorite source rocks than in the pegmatites’ source rocks. An unresolved question is why magmas that formed igneous rocks in close spatial association (granodiorite plutons and pegmatite dykes) were not apparently formed by partial melting of the same source rocks. In an alternative scenario the voluminous Leinster Granite could represent mixing between mantle-derived mafic magma and crustal melts of similar rocks that alone generated pegmatite magmas. Precise dating and radiogenic isotope studies to define the sources are planned towards a better understanding of the story behind the formation of pegmatites in southeast Ireland.

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REFERENCES


