This study assesses whether internal U, Th and Rb concentrations of single grains of alkali-feldspar can impact upon luminescence dating. Internal alpha dose-rates determined for two sedimentary samples calculated from mean U and Th concentrations accounted for ~10% of the total dose-rates, while internal beta dose-rates from mean Rb concentrations accounted for ~2%. Depth profiles of measurements penetrating into each grain show that internal U and Th concentrations varied between grains, where some grains had effective internal alpha dose-rates up to 1.8 Gy/ka (32% of the total dose-rate). K concentrations inferred from Rb (K$_{Rb}$) for individual grains suggest that internal U and Th concentrations are related to the feldspar composition. Grains with K$_{Rb}$ > 6% had low U and Th, and grains with K$_{Rb}$ < 6% had higher U and Th concentrations. Internal alpha and beta dose-rates of the perthitic and Na-rich grains contaminating the density-separated K-feldspar fractions had a significant impact upon the single-grain $D_e$ distributions, which was estimated to be equivalent to overdispersion values of ~10 – 15%. The scatter in the $D_e$ distributions arising from internal alpha and beta dose-rates for both samples could be reduced by selecting only the brightest 20% of grains for luminescence dating.

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1. Introduction

Luminescence dating is a versatile geochronological tool that can directly determine the timing of sediment deposition throughout the Late Quaternary. Two principal minerals are used for luminescence dating: quartz and K-feldspar. A major challenge for luminescence dating of K-feldspar in comparison to quartz is the more complex internal chemistry and crystal structure of feldspar. This is because alkali feldspars form a complete solid solution series under high-temperature magmatic conditions (e.g. in volcanic rocks) between K-rich (sanidine) and Na-rich (high albite) compositions. However, at lower temperatures, an immiscibility gap exists between the Na-rich (low albite) and K-rich (orthoclase, microcline) end-members, which can manifest itself either as perthitic textures or as separate Na- and K-feldspars in plutonic rocks. The presence of K, Rb, U and Th within a feldspar grain provides an internal beta dose-rate, while U and Th also provide an internal alpha dose-rate; thus, it is important that this internal dosimetry is accounted for in the environmental dose-rate used to determine luminescence ages.

Many studies have assessed the internal K-content of single grains of K-feldspar (e.g. Huot and Lamothe, 2012; Neudorf et al., 2012; Smedley et al., 2012; Trauerstein et al., 2014; Gaar et al., 2014) and have shown that some density-separated K-feldspar fractions used for dating can contain perthitic and Na-rich grains. Perthitic structures within alkali feldspar grains form after crystallisation of a homogeneous feldspar by a process of exsolution, where K- and Na-rich areas form within the feldspar to accommodate structural changes in the crystal lattice during cooling. The extent of development of perthitic structures in feldspar is dependent upon the initial K/Na composition of the grain (thus enabling the exsolution of different proportions of K- and Na-rich phases) and history of the crystal (particularly the cooling rate, interaction with aqueous liquids and deformation) (Deer et al., 2013). Where sedimentary samples are dominated by end-member feldspar grains, it is easier to perform a physical separation of detrital grains to isolate the K-rich end member for luminescence analysis on the basis of density. However, density separation is more difficult for a sedimentary sample containing perthites (e.g. Smedley et al., 2012). Although detecting the luminescence emission of feldspar grains in blue wavelengths can further isolate the peak emission at ~400 nm of density-separated K-feldspar (Huntley...
et al., 1991), it is possible to detect measurable infra-red stimulated luminescence (IRSL) signals (e.g. Huot and Lamothe, 2012; Smedley et al., 2012; Gaar et al., 2014) and determine equivalent dose ($D_E$) values (e.g. Huot and Lamothe, 2012; Gaar et al., 2014) from grains which are perthitic or Na-rich. This is because perthitic grains of feldspar have wide emission bands (Prescott and Fox, 1993; Spooner, 1992), and some Na-rich grains that have bright peak emissions in yellow wavelengths can also emit a weaker, but detectable signal in blue wavelengths (e.g. Krbetschek et al., 1997).

The range of partition coefficients published for K-feldspar and plagioclase feldspar show that the uptake of U, Th and Rb during crystallisation leads to lower concentrations of U and Th in K-feldspar than plagioclase feldspar (Table 1 and references therein). Therefore, it is usually assumed that the internal dose-rate of K-rich feldspars will mostly originate from K within the grain and the associated U, Th and Rb contributions are negligible. However, potential exists for perthitic and Na-rich grains to have different concentrations of minor elements (i.e. U, Th and Rb) to the K-rich grains (e.g. Zhao and Li, 2005). Published data of internal U, Th and Rb concentrations within feldspar grains used for luminescence dating are limited, especially for grains of different feldspar compositions. Natural abundances of Rb are suggested to be related to K-concentrations between different samples (Mejdahl, 1987; Duller, 1992) and grains (Zhao and Li, 2005). However, Duller (1992) suggested that effective alpha dose-rates from such samples would be minimal as U and Th concentrations are low in naturally-occurring K-feldspar, and would be located in zones or along cracks within the crystalline structure of the grain (e.g. Mejdahl, 1987). As a consequence, it has become routine practise to either assume a negligible internal alpha dose-rate when calculating environmental dose-rates for K-feldspar (e.g. Duller, 1992; Reimann et al., 2011; Rhodes, 2015), or to apply an internal alpha dose-rate of 0.10 $\pm$ 0.05 Gy/ka after Mejdahl (1987) (e.g. Wallinga et al., 2001; Törnqvist et al., 2000; Buylaert et al., 2008, 2011; Sohbatí et al., 2012). However, these assumptions may not be appropriate if some of the feldspar grains used to determine ages for some samples are Na-rich or perthitic, or if the grains contain any inclusions (e.g. micron-sized zircon). Internal U, Th and Rb concentrations taken throughout a grain of feldspar used for luminescence dating have not been quantified on a single grain scale. If it was found that significant concentrations of U, Th and Rb were evenly distributed throughout the grain, the effective internal alpha and beta dose-rates would provide a significant contribution to the environmental dose-rate according to concentrations reported for samples in previous studies (e.g. Duller, 1992; Mejdahl, 1987).

Therefore, the aim of this study is to quantify internal U, Th and Rb concentrations of K-feldspar grains used for luminescence dating with laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS) and assess any potential impacts upon ages.

2. Methods

2.1. Sample details

Two sedimentary samples with independent age control were used in this study: sample GDNZ13 and LBA12OW1. Sample GDNZ13 was taken from dune sand from North Island, New Zealand (Duller, 1996) underlying the Kawakawa tephra, which has a radiocarbon age of 25.4 $\pm$ 0.2 cal. ka BP (Vandergoes et al., 2013). Sample LBA12OW1 was taken from glaciofluvial sediment associated with the deposition of the Fenix II moraine ridge at Lago Buenos Aires in Patagonia with a terrestrial cosmogenic nuclide (TCN) age of 19.4 $\pm$ 1.5 ka (Kaplan et al., 2004, 2011). Luminescence ages for samples GDNZ13 (Smedley, 2014) and LBA12OW1 (Smedley et al., 2016) have previously been published, where the individual grains within the sediments were deemed to have been well bleached prior to burial.

Samples GDNZ13 and LBA12OW1 were chosen for these investigations as the sedimentary samples contain a mixture of end member and perthitic feldspar grains. This is reflected by the intermediate K-contents of the density-separated K-feldspar fractions determined using a 0.1 g sub-sample of the separated material on a Risø GM-25-5 beta counter for samples GDNZ13 (6% K) and LBA12OW1 (5% K). In addition, measurements of the internal K-content of individual grains of samples GDNZ13 (Smedley et al., 2012; Smedley, 2014) and LBA12OW1 (Smedley et al., 2016) have shown the variability between grains in internal K-content, ranging from 0 to 14%. It has been suggested that this variability in internal K-content can explain the relationship between the signal-intensity emitted in response to a fixed test-dose and the $D_E$ values determined for luminescence dating, where K-rich grains emit brighter signals (Reimann et al., 2012; Smedley, 2014; Smedley et al., 2016).

Samples GDNZ13 and LBA12OW1 therefore provide an opportunity to assess the variability in internal U, Th and Rb concentrations for grains in a density-separated K-feldspar fraction containing perthites and Na-feldspar due to the nature of the feldspar grains in the source rocks.

2.2. Luminescence dating

2.2.1. Environmental dose-rates

External dose-rates for samples GDNZ13 and LBA12OW1 (Table 2) were determined from milled portions of the bulk sediment using thick source alpha and beta counting on Daybreak and Risø GM-25-5 measurement systems, respectively. Although K-concentrations of 6% (GDNZ13) and 5% (LBA12OW1) were determined for the bulk density-separated K-feldspar fractions, such low K-contents are unlikely to characterise those individual grains used for analysis as K-rich grains are reported to emit brighter IRSL signals when detected in blue wavelengths in comparison to Na-

---

### Table 1

<table>
<thead>
<tr>
<th>Range of partition coefficients ($K_d$)</th>
<th>Typical concentrations of granite (ppm)</th>
<th>Estimated concentrations in grains (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>K-feldspar</td>
<td>Plagioclase feldspar</td>
</tr>
<tr>
<td>U</td>
<td>0.002 – 0.07</td>
<td>0.05 – 0.13</td>
</tr>
<tr>
<td>Th</td>
<td>0.01 – 0.03</td>
<td>0.004 – 0.382</td>
</tr>
<tr>
<td>Rb</td>
<td>0.11 – 2.85</td>
<td>0.06 – 0.34</td>
</tr>
</tbody>
</table>
rich and perthites (e.g. Krbetschek et al., 1997). Internal K-contents of 12.5 ± 0.5% are routinely applied for dating density-separated K-feldspar fractions after Huntley and Baril (1997). However, an internal K-content of 10 ± 2% was used in this study to calculate the internal beta dose-rate unless otherwise stated after the observations of by Smedley et al. (2012) determined from a more extensive dataset. The dose-rates (Table 2) were calculated using the conversion factors of Guerin et al. (2011), an alpha efficiency value (α-value) of 0.01 ± 0.03 (Balescu and Lamothe, 1993), and alpha (Bell, 1980) and beta (Guerin et al., 2012) dose-rate attenuation factors. The water contents are expressed as a percentage of the mass of dry sediment and were estimated considering the field and saturated water contents, in addition to the environmental history for each sample. Cosmic dose-rates were determined in accordance with Prescott and Hutton (1994).

2.2.2. OSL analysis
To isolate the coarse grains of feldspar for luminescence analysis, the bulk sediment sample was treated with a 10% volume concentration (v/v) dilution of 37% HCl and 20% v/v H2O2 to remove carbonates and organics, respectively. Dry sieving was then used to provide the 180 – 212 μm diameter grains, and density-separation using sodium polytungstate and a centrifuge isolated the <2.58 g cm−3 (K-feldspar-dominated) fraction. To prevent the non-isotropic removal of the grain surface, the K-feldspar grains were not etched in hydrofluoric acid prior to measurement (Duller, 1992; Porat et al., 2015). The K-feldspar grains were finally mounted into 10 by 10 grids of 300 μm diameter holes in a 9.8 mm diameter aluminium single-grain disc for analysis. All luminescence measurements were performed using a Risø TL/OSL DA-15 automated single-grain system equipped with an infrared laser (830 nm) fitted with an RG-780 filter (3 mm thick) to remove any shorter wavelengths (Batter-Jensen et al., 2003; Duller et al., 2003). A blue detection filter pack containing a BG-39 (2 mm), GG-400 (2 mm) and Corning 7-59 (2.5 mm) filter was placed in front of the photomultiplier tube. The system was equipped with a 85Sr/90Y beta source delivering ~0.04 Gy/s and the variability of the beta dose as this improves the reproducibility of the background: (2) the uncertainty in the luminescence measurement of the test dose was greater than 10%; (3) the recycling ratio was outside the range 0.9 – 1.1, taking into account the uncertainties on the individual ratios; and (4) recuperation was greater than 5% of the response from the largest regenerative dose.

2.3. LA-ICP-MS
LA-ICP-MS analyses (U, Th and Rb) were conducted using a Coherent GeoLas ArF 193 nm Excimer laser coupled to a Thermo Finnegan Element 2 sector field high resolution ICP-MS. Concentrations of U, Th and Rb determined for an ATHO-G reference material using LA-ICP-MS were within error of accepted concentrations published by GeoReM (Table 4). The individual feldspar grains analysed using LA-ICP-MS in this study were kept in the single-grain holders after Dc values were determined for each grain and secured using an organic glue solution (1% PVA) to prevent the grains from moving during ablation. A large beam size with diameter of 80 μm was used for laser ablation at an energy density of 3 J/cm2, firing at 3 Hz to create an aerosol for analysis using the ICP-MS. Analyses were performed in low mass resolution (ΔM/M = 300) to provide sufficient sensitivity. Table 4 details the detection limits of the elements from LA-ICP-MS analyses. The intensities of the minor elements (determined from the following isotopes: 238U, 232Th, 85Rb) were calibrated by normalisation to 29Si as an internal standard because Si has a near constant stoichiometry within all alkali feldspars. This removes the associated effects of changes in ablation yield between standards and grains, and is standard practise in LA-ICP-MS analysis (see Perkins and Pearce, 1995; Pearce et al., 2004). The NIST-612 glass reference material was used for external calibration (Pearce et al., 1997) by comparing the analyte to 29Si count rates and correcting for the differences in internal standard concentrations between the feldspars and the reference material. For the ICP-MS operating conditions used in this study see Pearce et al. (2011).

To monitor the variability of minor elemental concentrations within individual grains, depth profiles were determined by repeating ablation cycles on the same spot. Between 7 and 10 spectra were acquired from the same crater for each individual grain over a period of 105 – 150 s, this is estimated to have produced craters up to –150 μm deep and –80 μm wide in grains of 180 μm diameter.
Due to the LA-ICP-MS set-up and nature of the analyses, major and minor elements could not be easily and quickly analysed during the same ablation cycle. This restricted the analysis of both K (a major element which requires determination in high mass resolution mode, \(D_M/M > 3000\); see Smedley et al., 2012) and U, Th and Rb concentrations (minor elements which require determination in low mass resolution mode, \(D_M/M = 300\)) for each individual depth profile. Alternatively, the Rb concentrations were used to infer an approximate K concentration (termed \(K_{Rb}\)) using the regression provided from a correlation of Rb with K for 27 feldspar samples, where \(Rb = 9.17 \times 10^{-4} + 38.13 \times \% \) (Mejdahl, 1987); thus, 28.96 ppm Rb was equivalent to 1% K. This gave an indication as to whether the grain composition was K-rich, Na-rich or perthitic. The mean of the \(K_{Rb}\) concentrations determined for single grains from samples GDNZ13 (6%; \(n = 21\) grains) and LBA12OW1 (4%; \(n = 14\) grains) were similar to the respective K-contents of 6 and 5% which were determined for the density-separated fraction of each sample using beta counting (Section 2.1); this improves confidence in the reliability of these estimations of \(K_{Rb}\).

### Table 4

<table>
<thead>
<tr>
<th>ATHO-G GeoReM accepted values (Jochum et al. 2005)</th>
<th></th>
<th></th>
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</thead>
<tbody>
<tr>
<td>Accepted ppm</td>
<td>2.37</td>
<td>7.40</td>
</tr>
<tr>
<td>95% C.L. ppm</td>
<td>0.12</td>
<td>0.27</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>ATHO-G this study ((n = 5))</th>
<th></th>
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</tr>
</thead>
<tbody>
<tr>
<td>Avg. ppm</td>
<td>2.37</td>
<td>7.20</td>
</tr>
<tr>
<td>St. dev. ppm</td>
<td>0.16</td>
<td>0.31</td>
</tr>
<tr>
<td>RSD %</td>
<td>6.8</td>
<td>4.3</td>
</tr>
</tbody>
</table>

| Blank cps \(\pm 1\) s.d.                      | 0 ± 0 | 3 ± 3 | \(-600 \pm 20\) |

Typical sensitivity cps/ppm:

- 1st quartile: \(0.004 \pm 0.10\) ppm
- 2nd quartile: \(0.11 \pm 0.53\) ppm
- 3rd quartile: \(0.65 \pm 1.77\) ppm
- 4th quartile: \(1.84 \pm 3.95\) ppm

### Analytical error estimates

<table>
<thead>
<tr>
<th>U range (ppm)</th>
<th>U avg. 1 s.d. error, ppm (RSD %)</th>
<th>U avg. 1 s.d. error, ppm (RSD %)</th>
<th>U avg. 1 s.d. error, ppm (RSD %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1st quartile</td>
<td>(0.004 \pm 0.10) ppm</td>
<td>(0.009 (38.2%)) ppm</td>
<td>(0.02 (0.7%)) ppm</td>
</tr>
<tr>
<td>2nd quartile</td>
<td>(0.11 \pm 0.53) ppm</td>
<td>(0.023 (11.7%)) ppm</td>
<td>(0.30 (12.5%)) ppm</td>
</tr>
<tr>
<td>3rd quartile</td>
<td>(0.65 \pm 1.77) ppm</td>
<td>(0.093 (7.97%)) ppm</td>
<td>(1.66 \pm 6.46) ppm</td>
</tr>
<tr>
<td>4th quartile</td>
<td>(1.84 \pm 3.95) ppm</td>
<td>(0.14 (5.42%)) ppm</td>
<td>(6.54 \pm 32.98) ppm</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Th range (ppm)</th>
<th>Th avg. 1 s.d. error, ppm (RSD %)</th>
<th>Th avg. 1 s.d. error, ppm (RSD %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1st quartile</td>
<td>(0.004 \pm 0.10) ppm</td>
<td>(0.009 (38.2%)) ppm</td>
</tr>
<tr>
<td>2nd quartile</td>
<td>(0.11 \pm 0.53) ppm</td>
<td>(0.023 (11.7%)) ppm</td>
</tr>
<tr>
<td>3rd quartile</td>
<td>(0.65 \pm 1.77) ppm</td>
<td>(0.093 (7.97%)) ppm</td>
</tr>
<tr>
<td>4th quartile</td>
<td>(1.84 \pm 3.95) ppm</td>
<td>(0.14 (5.42%)) ppm</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Rb range (ppm)</th>
<th>Rb avg. 1 s.d. error, ppm (RSD %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1st quartile</td>
<td>(0.004 \pm 0.10) ppm</td>
</tr>
<tr>
<td>2nd quartile</td>
<td>(0.11 \pm 0.53) ppm</td>
</tr>
<tr>
<td>3rd quartile</td>
<td>(0.65 \pm 1.77) ppm</td>
</tr>
<tr>
<td>4th quartile</td>
<td>(1.84 \pm 3.95) ppm</td>
</tr>
</tbody>
</table>

| Avg 1 s.d. for all measurements (ppm) | 0.07 | 0.14 | 2.02 |

- 212 µm. Due to the LA-ICP-MS set-up and nature of the analyses, major and minor elements could not be easily and quickly analysed during the same ablation cycle. This restricted the analysis of both K (a major element which requires determination in high mass resolution mode, \(\Delta M/M > 3000\); see Smedley et al., 2012) and U, Th and Rb concentrations (minor elements which require determination in low mass resolution mode, \(\Delta M/M = 300\)) for each individual depth profile. Alternatively, the Rb concentrations were used to infer an approximate K concentration (termed \(K_{Rb}\)) using the regression provided from a correlation of Rb with K for 27 feldspar samples, where \(Rb = -9.17 \times 10^{-4} + 38.13 \times \% \) (Mejdahl, 1987); thus, 28.96 ppm Rb was equivalent to 1% K. This gave an indication as to whether the grain composition was K-rich, Na-rich or perthitic. The mean of the \(K_{Rb}\) concentrations determined for single grains from samples GDNZ13 (6%; \(n = 21\) grains) and LBA12OW1 (4%; \(n = 14\) grains) were similar to the respective K-contents of 6 and 5% which were determined for the density-separated fraction of each sample using beta counting (Section 2.1); this improves confidence in the reliability of these estimations of \(K_{Rb}\).

### 3. Distribution of trace elements within different feldspar grains

Depth profiles determined using LA-ICP-MS assessed the distribution of U and Th within K-feldspar grains used for luminescence dating. The concentrations in the depth profiles determined for grains from sample GDNZ13 (\(n = 21\) grains) and LBA12OW1 (\(n = 14\) grains) demonstrated that the distribution of U and Th varied between individual grains. The grains could be categorised into three types according to the depth profiles (Fig. 1): Type 1 contain low U (<0.1 ppm) and Th (<0.5 ppm) concentrations throughout the depth profile; Type 2 contain high U (>0.5 ppm) and low Th (<0.2 ppm) concentrations, and Type 3 contain high U (>0.5 ppm) and high Th (>0.2 ppm) concentrations.
and Th (>3 ppm) concentrations throughout the depth profile; and Type 3 contain moderate but variable U (0.1 – 0.5 ppm) and Th (0.5 – 3 ppm) within the depth profile. These depth profiles suggest that the effective internal alpha dose-rates of U and Th will vary between these different grain types.

Comparing the depth profiles of U and Th with the Rb and KRb concentrations (Fig. 1) provides information on the composition of each feldspar grain. Fig. 1 shows a K-rich (>10% KRb) feldspar grain with homogeneously low U and Th throughout the depth profile (Type 1), whereas the KRb Concentrations determined for grain Type 2 (6 – 9%) and Type 3 (4 – 6%) are representative of intermediate feldspar, likely with perthitic inclusions. The Type 1 and Type 3 grains (Fig. 1) fulfill the expectations of previous studies that U and Th concentrations of K-feldspar are either low or are variable throughout the grain and so the internal alpha dose rate is less effective. However, the higher U and Th concentrations determined throughout the depth profiles for Type 2 grains suggest that internal alpha dose-rates can contribute significantly to the environmental dose-rate and introduce scatter into single-grain D\textsc{e} distributions, in addition to the variability caused by the internal beta dose-rate arising from K and Rb.

Mean U, Th and Rb concentrations were then calculated for each depth profile (Fig. 2). The ranges in U, Th and Rb concentrations determined for both samples GDNZ13 and LBA12OW1 are consistent with estimates provided for feldspar grains from reported partition coefficients and typical concentrations in granite (Table 1). Mean Rb concentrations ranged from ~10 ppm to 630 ppm, which suggests that the composition of these feldspar grains was variable. This was also reflected by the KRb concentrations from 1 – 14% for sample GDNZ13 and 0 – 11% for sample LBA12OW1. A large proportion of the grains had <0.4 ppm of U and ≤2 ppm of Th for samples GDNZ13 (81% and 67% of the grains, respectively) and LBA12OW1 (67% and 64% of the grains, respectively). This is similar to reported estimates of U and Th for 16 individual feldspar grains, where 15 out of the 16 grains determined qualitative estimates of U and Th from 0.0 – 0.5 ppm and 0 – 0.6 ppm, respectively, and a single grain in the population had significantly higher respective estimates of 1.4 ppm and 1.7 ppm (Zhao and Li, 2005).

The relationship between the U and Th concentrations and the composition of the feldspar grains was assessed by comparing the mean U, Th and KRb concentrations determined for each depth profile (Fig. 3). The results show that only one grain categorised as Type 3 had KRb concentrations >6%. In contrast, the majority of the grains categorised as Type 1 had KRb concentrations >6%. The comparison between U and Th concentrations and KRb is consistent with the suggestion that the K-rich end member of the alkali feldspar phase has lower concentrations of U and Th than intermediate feldspar, but higher Rb (Deer et al., 2013). Determining accurate environmental dose-rates will be difficult for a sample that contains K-rich feldspar grains with low internal alpha dose-rates, in addition to perthitic and/or Na-rich feldspar which have a lower K-content, but effective internal alpha dose-rates (i.e. Type 2 grains). Therefore, it is important for single-grain dating of density-separated K-feldspar fractions containing variable K-contents to determine: (1) whether the internal alpha dose-rate can have a significant contribution to the overall dose-rate; (2) whether variability between grains can impact upon the single-grain D\textsc{e} distributions; and (3) whether the effects of this can be reduced for luminescence dating.

4. Implications for environmental dose-rates

Mean Rb concentrations were calculated from the depth profiles

---

**Fig. 2.** Histograms of the mean U (a, d), Th (b, e) and Rb (c, f) concentrations calculated from the depth profiles from individual grains from samples GDNZ13 and LBA12OW1 determined using LA-ICP-MS.
obtained for all of the grains from samples GDNZ13 (293 ppm; Fig. 2c) and LBA12OW1 (218 ppm; Fig. 2f) and equate to an internal beta dose-rate of 0.07 ± 0.01 Gy/ka and 0.06 ± 0.01 Gy/ka, respectively. These internal beta dose-rates from Rb are similar to those determined for a variety of sedimentary samples from multiple grains (e.g. Mejdahl, 1987) and account for a similarly small proportion of the total dose-rate (i.e. from 1 – 4%). While Rb can provide an internal beta dose-rate during burial of the grains, it is low in comparison to the internal beta dose-rate from K.

Mean internal U and Th concentrations were also calculated from the depth profiles determined for the grains analysed from samples GDNZ13 (U 0.3 ppm, Th 1.7 ppm) and LBA12OW1 (U 0.8 ppm, Th 2.8 ppm) (Fig. 2). The U and Th concentrations were broadly consistent with semi-quantitative estimates ranging from 0.2 – 0.5 ppm for U and 1.7 – 3.0 ppm for Th which were determined for samples of similar material to GDNZ13 from dune sand in New Zealand using a spiked internal standard for calibration (Duller, 1992). Internal alpha and beta dose-rates were then calculated for samples GDNZ13 (U 0.3 ppm, Th 1.7 ppm) and LBA12OW1 (U 0.8 ppm, Th 2.8 ppm) using the mean U and Th concentrations determined from LA-ICP-MS analysis and assuming an internal K-content of 10 ± 2% (see Section 2.2.1). When combining the beta dose-rate arising from U and Th with that from K, the internal beta dose-rates for samples GDNZ13 (0.68 ± 0.12 Gy/ka) and LBA12OW1 (0.70 ± 0.12 Gy/ka) increased by 3% and 6% respectively from the internal beta dose-rate determined using only the internal K-content of 10 ± 2% (0.66 ± 0.12 Gy/ka). The internal alpha dose-rates arising solely from U and Th inside the grain were 0.22 ± 0.04 and 0.45 ± 0.07 Gy/ka for samples GDNZ13 and LBA12OW1, respectively. These internal alpha dose-rates are similar to those previously determined for sedimentary samples used for luminescence dating (e.g. Kolstrup and Mejdahl, 1986; Mejdahl, 1992; Edwards, 1993) and suggest that it may not be appropriate to assume an internal alpha dose-rate of 0.10 ± 0.05 Gy/ka (after Mejdahl, 1987) for luminescence dating of these samples when all of the grains are used to determine a D_e value.

The proportion of the overall dose-rate (Section 2.2.1) for each component has been calculated when the contributions from internal U and Th are included with the calculations in Table 2 (Fig. 4). The external dose-rates account for a large proportion of the total dose-rates for samples GDNZ13 (64%) and LBA12OW1 (71%), respectively. The majority of the internal dose-rate arises from beta
provided by K and Rb within the grain: 28% of the total dose-rate for GDNZ13 and 18% for LBA12OW1. However, based on the LA-ICP-MS analyses in this study, the mean internal alpha dose-rates determined for samples GDNZ13 and LBA12OW1 account for 9% and 11% of the total dose-rates, respectively (Fig. 4). This contribution is larger than the external alpha dose-rate and the cosmic dose-rate for both samples; both of these contributions are routinely accounted for when calculating environmental dose-rates for luminescence dating. It is therefore important to address the contributions of internal alpha dose-rates arising from U and Th within the grains, especially given the potential for large variability between different grains for samples containing variable types of feldspar grains.

5. Impact upon luminescence characteristics

To assess whether luminescence dating of K-feldspar from samples GDNZ13 and LBA12OW1 could be affected by the analysis of grains with variable internal U and Th concentrations, the internal dosimetry of each individual grain from sample GDNZ13 and LBA12OW1 is compared with the $D_i$ value determined for dating (Fig. 5). Previous studies have suggested that the variability in internal K-content for both samples GDNZ13 (Smedley, 2014) and LBA12OW1 (Smedley et al., 2016) can be reduced by selecting the brightest 20% of grains for luminescence dating. For sample GDNZ13 and LBA12OW1, this included the grains that emitted signal-intensities in response to a fixed 52 Gy test-dose of

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**Fig. 5.** Comparing the results determined from geochemical and luminescence analyses.
>10,000 cts/0.3 s and >4000 cts/0.3 s, respectively. The grains used for geochemical analysis in this study were taken as a subsample of the dataset used for luminescence dating of both these samples and so the sub-sampled grains could be categorised into those grains that were part of the brightest 20% of the population and those that were not (Fig. 5). A total of eight out of the 21 grains for sample GDNZ13 were categorised as the brightest grains (e.g. Fig. 5a), whereas only three out of the total 14 grains were categorised as the brightest grains for sample LBA12OW1 (e.g. Fig. 5b).

5.1. Feldspar composition

Internal alpha dose-rates are compared to the $K_{rb}$ concentrations for samples GDNZ13 (Fig. 5a) and LBA12OW1 (Fig. 5b). For sample GDNZ13, the results suggest that the larger internal alpha dose-rates (>0.4 Gy/ka) were determined from grains with $K_{rb}$ concentrations that are <6%, and that these grains were not part of the brightest 20% of the grains. Moreover, by selecting only the brightest 20% of the grains for sample GDNZ13, the range in internal alpha dose-rate was reduced from 0 – 0.63 Gy/ka for all grains to 0 – 0.25 Gy/ka. Larger internal alpha dose-rates (>0.25 Gy/ka) were also determined for the grains of sample LBA12OW1 that had $K_{rb}$ concentrations of <6%. However, one grain was an exception to this trend as it had ~11% $K_{rb}$ and an internal alpha dose-rate of ~1.7 Gy/ka. The range in internal alpha dose-rates reduced from 0 – 1.8 Gy/ka for all grains to 0 – 0.35 Gy/ka when only the brightest 20% of grains were selected from sample LBA12OW1. The results shown in Fig. 5 for both samples provide evidence to suggest that previous studies (Smedley, 2014; Smedley et al., 2016) have reduced the variability between grains in internal K-contents and reduced the influence of internal alpha dose-rates upon luminescence dating by selecting the brightest 20% of grains for both samples GDNZ13 and LBA12OW1.

5.2. Internal K-contents

The geochemical data determined in this study provide a rare opportunity to determine internal alpha and beta dose-rates arising from $K_{rb}$, U and Th for grains that are used to determine $D_e$ values for dating. If either or both internal dose-rates have a dominant control on the $D_e$ distributions, it is expected that there will be a relationship between the two parameters. The $K_{rb}$ concentrations determined for each grain from samples GDNZ13 (Fig. 5c) and LBA12OW1 (Fig. 5d) have been compared with the corresponding $D_e$ values used for dating. The results show that there is a broad positive relationship approaching significance ($R = 0.41$ with $p = 0.06$) between $K_{rb}$ concentrations and $D_e$ values for all the individual grains from sample GDNZ13 (Fig. 5c). An R value of 0.41 ($p = 0.06$) suggests that the internal K-content is likely to have had a control on the single-grain $D_e$ distribution for this sample. However, for sample LBA12OW1 there is no relationship ($R = 0.02$ with $p = 0.94$) between the $D_e$ values and $K_{rb}$ (Fig. 5d). This difference between the data shown for GDNZ13 (Fig. 5c) and LBA12OW1 (Fig. 5d) may be related to the fact that the internal dose-rate accounts for a smaller proportion of the total dose-rate for sample LBA12OW1 (17%) in comparison to GDNZ13 (28%) and/or the larger contributions provided by the internal alpha dose-rate for sample LBA12OW1 (Fig. 5b) in comparison to sample GDNZ13 (Fig. 5a). Unfortunately, the datasets for the brightest 20% of the grains for both samples are too small to provide a reliable assessment of any relationship between $D_e$ and $K_{rb}$.

5.3. Internal alpha dose-rates

The internal alpha dose-rates arising from U and Th have also been compared to the $D_e$ values determined for individual grains of both sample GDNZ13 (Fig 5e) and LBA12OW1 (Fig 5f). Any direct relationship between internal alpha dose-rates and $D_e$ values for these samples is difficult to identify because of the large uncertainties determined for some of the grains. However, the grains with large uncertainties in the $D_e$ values also determine larger internal alpha dose-rates, and are the dimmer grains in the $D_e$ distribution. Fig. 5e suggests that it is beneficial to select the brightest 20% of grains as it reduces the impact of those grains on the single-grain $D_e$ distribution, which have large internal alpha dose-rates and determine variable $D_e$ values.

5.4. Combined internal alpha and beta dose-rates

The internal alpha and beta dose-rates are then combined for each grain of samples GDNZ13 (Fig. 5g) and LBA12OW1 (Fig. 5h) and compared with the corresponding $D_e$ values. The data suggests that there is no direct relationship between the internal dose-rates and $D_e$ values for sample GDNZ13 (Fig. 5g) or LBA12OW1 (Fig. 5h). This is likely because other factors also contribute towards the scatter in a single-grain $D_e$ distribution determined for K-feldspar. For example, variability in the external dose-rate received by each individual grain (i.e. external microdosimetry) may have caused additional scatter in the single-grain $D_e$ distributions as 64% (GDNZ13) and 70% (LBA12OW1) of the total dose-rate originates from the external dose-rate. Variability between grains in factors inherent to the analysis of single grains of feldspar using the pIRIR signal may also have contributed towards the scatter in single-grain $D_e$ distributions (e.g. bleaching rate of the pIRIR signal and anomalous fading). Given that a number of different factors influence the scatter in a single-grain $D_e$ distribution determined for K-feldspar, it is unlikely that a single factor will have a direct relationship with the $D_e$ values; this is reflected in the comparison with the internal alpha and beta dose-rates presented here.

6. Implications for luminescence dating

6.1. Simulated $D_e$ distributions

The independent age control provided for the well-bleached samples GDNZ13 (radiocarbon dating) and LBA12OW1 (TCN dating) provides an opportunity to assess the influence of internal dose-rates on the scatter in single-grain $D_e$ distributions. Grain-specific environmental dose-rates have been calculated in three different ways using the external dose-rates (Table 2) and geochemical data determined from LA-ICP-MS: (1) using $K_{rb}$ and Rb concentrations and excluding internal U and Th; (2) using an assumed internal K-content of 10 ± 2% and the measured internal U and Th concentrations; and (3) using combined internal dose-rates derived from $K_{rb}$, Rb, U and Th. These grain-specific dose-rates were then multiplied by the independent age control to determine the expected $D_e$ value for each grain, which in turn simulated single-grain $D_e$ distributions that are the result of internal dose-rates.

Overdispersion values of 11 ± 1% (applying internal $K_{rb}$ and Rb and excluding internal U and Th), 8 ± 1% (applying an assumed internal K-content of 10 ± 2% and the measured internal U and Th) and 10 ± 1% (combining internal dose-rates from $K_{rb}$, Rb, U and Th) were determined from these $D_e$ distributions for sample GDNZ13. The simulated single-grain $D_e$ distributions suggest that variability caused by internal $K_{rb}$ and Rb (~11%) caused similar scatter in the $D_e$ distribution to the internal alpha dose-rates (~8%) for sample GDNZ13. The single-grain $D_e$ distributions simulated from the combined internal alpha and beta dose-rates (~10%) suggest that the internal alpha dose-rates counter-balance some of the scatter
caused by the internal beta dose-rates as those grains with larger internal alpha dose-rates had lower internal beta dose-rates. For sample LBA12OW1, overdispersion values of 9 ± 1% (applying internal K\textsubscript{Rb} and Rb and excluding internal U and Th), 15 ± 1% (applying an assumed internal K-content of 10 ± 2% and the measured internal U and Th) and 15 ± 1% (combining internal dose-rates from K\textsubscript{Rb}, Rb, U and Th) were determined for the simulated D\textsubscript{e} distributions. The overdispersion values suggest that internal alpha dose-rates cause more scatter (~15%) than the internal beta dose-rates (~9%) for sample LBA12OW1.

The overdispersion values determined for the simulated D\textsubscript{e} distributions could then be related to the natural D\textsubscript{e} distributions by comparison with the overdispersion values provided from the original studies of samples GDNZ13 (Smedley, 2014) and LBA12OW1 (Smedley et al., 2016), which were 25 ± 1% and 39 ± 1%, respectively. When only the brightest 20% of these grains from GDNZ13 (14 ± 1%) and LBA12OW1 (26 ± 1%) were used to determine overdispersion, the values were lower. This is consistent with the suggestion that selecting the brighter grains better isolates the K-rich feldspar grains, and the overdispersion values of 10 ± 1% (GDNZ13) and 15 ± 1% (LBA12OW1) determined from the single-grain D\textsubscript{e} distribution simulated using the internal alpha and beta dose-rates can explain the reduction in scatter. Understanding the amount of scatter in a D\textsubscript{e} distribution caused by internal dose-rates of K-feldspar is important as it can inform the parameters that should be used for statistical modelling (e.g. \(\sigma_9\) in the minimum age model) of single-grain D\textsubscript{e} distributions determined from heterogeneously-bleached sediments.

### 6.2. Natural D\textsubscript{e} distributions used for luminescence dating

Mean internal alpha dose-rates were calculated for all the grains analysed using LA-ICP-MS, in addition to only those categorised as the brightest 20% of the grains from Section 5 (Table 5). Although the datasets for the brightest grains of sample GDNZ13 (n = 8 grains) and LBA12OW1 (n = 3 grains) are small, comparing the brighter and dimmer grains gives an indication of the potential differences in geochemistry between them. The mean internal alpha dose-rate for sample GDNZ13 reduced from ~0.2 Gy/ka (all grains; 9% of the total dose-rate) to ~0.1 Gy/ka (brightest 20% of grains; 5% of the total dose-rate), and from ~0.4 Gy/ka (all grains; 11% of the total dose-rate) to ~0.1 Gy/ka (brightest 20% of grains; 9% of the total dose-rate) for sample LBA12OW1. The environmental dose-rates for both samples from Table 2 have been adjusted to include the approximate internal alpha dose-rates for all the grains, in addition to only the brightest 20% of the grains (Table 5). The CAM\(_{D_{e}}\) values provided from the original studies for samples GDNZ13 (Smedley, 2014) and LBA12OW1 (Smedley et al., 2016) have been divided by the adjusted dose-rates to calculate new ages (Table 5). The age determined from only the brightest 20% for sample LBA12OW1 still agrees within ±1\(\sigma\) with the independent age control for sample LBA12OW1 (Table 5). However, the age determined for only the brightest 20% of grains still underestimates the independent age control beyond ±2\(\sigma\). Underestimation of pIRIR ages by ~10% determined when using only the brightest 20% of the feldspar grains in comparison to independent age control was identified by the original study (Smedley, 2014) and have also been reported by other studies (e.g. Reimann et al., 2012). At present it is unknown what causes this underestimation, but the geochemical analyses in this study suggest that it cannot be explained by the internal dose-rate.

### 7. Conclusions

Geochemical analyses of U, Th and Rb using LA-ICP-MS in this study have provided unique insights into the internal dosimetry of single grains of feldspar used for luminescence dating. Depth profiles of LA-ICP-MS ablation cycles penetrating through each grain of feldspar showed that the distribution of U and Th within the grains could be characterised into three types: Type 1 had low U (<0.1 ppm) and Th (<0.5 ppm); Type 2 had high U (>0.5 ppm) and Th (>3.0 ppm); and Type 3 had moderate but variable U (0.1 – 0.5 ppm) and Th (0.5 – 3.0 ppm). The depth profiles suggested that effective internal alpha dose-rates from U and Th of these different grain types will likely vary, with Type 2 grains receiving a significant internal alpha dose-rate which is not restricted to zones within the crystal structure.

Comparing mean internal U and Th concentrations calculated from the depth profiles of each grain with internal K\textsubscript{Rb} concentrations suggests that the variability in U and Th is linked with the feldspar composition (i.e. K-rich, Na-rich or perthitic). The majority of K-rich feldspar grains were characterised by lower internal alpha dose-rates (<0.4 Gy/ka), and the grains with <8% K\textsubscript{Rb} had larger internal alpha dose-rates (>0.4 Gy/ka). Variability between grains of internal dose-rates will therefore have a minimal impact upon single-grain D\textsubscript{e} distributions obtained from sedimentary samples containing only K-feldspar that are easily separated on the basis of density. However, grain-to-grain variability in internal dose-rates will incorporate more scatter into D\textsubscript{e} distributions obtained from density-separated K-feldspar fractions contaminated by Na-rich and perthitic grains. Single-grain D\textsubscript{e} distributions simulated using independent age control and grain-specific dose-rates determined from the LA-ICP-MS data suggest that overdispersion values of ~10% and ~15% could be explained by internal dose-rates of feldspar. The geochemical evidence in this study suggests that it is important to select only the brightest 20% of grains from density-separated K-feldspar fractions that are contaminated by Na-rich or perthitic grains to circumvent the scatter in single-grain D\textsubscript{e} distributions that is potentially caused by grain-to-grain variability in internal dose-rates used for luminescence dating.

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### Table 5

Results from LA-ICP-MS analyses and luminescence dating for the samples in this study.

<table>
<thead>
<tr>
<th></th>
<th>Mean internal alpha dose-rate (Gy/ka)</th>
<th>Adjusted dose-rate (Gy/ka)</th>
<th>n</th>
<th>CAM (D_{e}) (Gy)</th>
<th>Age (ka)</th>
</tr>
</thead>
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<td><strong>GDNZ13 25.4 ± 0.2 cal ka BP</strong></td>
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<td></td>
</tr>
<tr>
<td>All grains</td>
<td>0.2</td>
<td>2.42 ± 0.14</td>
<td>597</td>
<td>44.2 ± 0.5</td>
<td>18.3 ± 1.1</td>
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<td>21.6 ± 1.3</td>
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<tr>
<td><strong>LBA12OW1 19.4 ± 1.5 ka</strong></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>All grains</td>
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<td>3.91 ± 0.16</td>
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<tr>
<td>Brightest</td>
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<td>3.61 ± 0.16</td>
<td>36</td>
<td>61.2 ± 2.8</td>
<td>17.0 ± 1.1</td>
</tr>
</tbody>
</table>

Acknowledgments

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