

Zircon luminescence dating revisited

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Abstract. Luminescence dating plays a pivotal role in Quaternary science, yet ongoing methodological challenges persist in refining the temporal range, accuracy, and precision of luminescence methods. Our contribution revisits zircons as potential alternative dosimeters to quartz, feldspar, or calcite for routine dating applications. The essential advantage of zircons over other minerals is the time-invariant and high internal dose rate due to high radionuclide contents, dominating over the external contribution, which is more challenging to assess. Reported drawbacks are low zircon abundance, laborious sample preparation, signal instabilities, unknown optical signal resetting rates, and low signal intensities. Our present study uses modern luminescence detection equipment and analytical methods to investigate mineral separation, mineral characteristics, bleachability, signal spectra and intensities, and the potential to auto-regenerate signals. We present results for two zircon samples that are different in provenance, trace element composition, and luminescence characteristics, each of them containing a couple of hundred grains. Optically stimulated luminescence (OSL) signal resetting rates of zircon in response to simulated sunlight exposure are orders of magnitudes faster than for feldspar and slightly slower than for quartz. The recorded thermoluminescence (TL) spectra confirm previously published results with luminescence emissions in the UV/violet and red wavelength range, supplemented by narrowband emissions associated with rare earth element (REE) dopants. Storage experiments of single zircon grains for auto-regenerated measurements over 1.5 years yielded very low OSL signals. At the same time, after only 3 weeks, we measured acceptable

TL signal intensities at the cost of lower bleaching rates. To date, the auto-regeneration approach seems to be a promising and accurate approach to dating zircon light exposure events, especially when combining the natural OSL with auto-regenerated TL. However, further studies are required to optimise signal intensities and establish zircons as viable targets for routine dating applications.

1 Introduction

Luminescence dating of natural quartz and feldspar minerals is one of the most important and influential geochronological techniques in geosciences and archaeology on Quaternary timescales. Direct dating of a mineral's last sunlight exposure (via optically stimulated luminescence, OSL; Huntley et al., 1985) or heating (via thermoluminescence, TL; Aitken, 1985; Mercier et al., 1995) allows temporal constraints of geomorphological events and processes and of archaeological site formation and human action. However, in best-case scenarios, the method's full age uncertainty is $4\% - 5\%$ but normally ranges from $8\% - 10\%$ (Rhodes, 2011; Wallinga and Cunningham, 2015; Lamothe, 2016; Lancaster et al., 2016). These values are appreciably greater than uncertainties associated with radiometric techniques (e.g. radiocarbon dating; Hajdas et al., 2021) or methods of incremental counting (varve chronology, dendrochronology; Vandergoes et al., 2018; Żarczyński et al., 2018). Consequently, such uncertainty margins sometimes complicate the unequivocal interpretation of luminescence data with regard to the research question. For instance, OSL ages of loess deposits do not generate the temporal resolution necessary to assign millennial-scale climate fluctuations (e.g. Greenland interstadials; Moine et al., 2017; Fischer et al., 2021), and Holocene OSL chronologies of dune re-activation phases as a proxy for droughts in the Great Plains (USA) are not precise enough to be reconciled with multi-year climate forcing mechanisms (Forman et al., 2001; Buckland et al., 2019). In archaeology, the uncertainty assigned to luminescence ages often hinders the unambiguous temporal separation of lithic technocomplexes (Guérin et al., 2013).

Since the early 1970s, zircon (tetr. $ZrSiO₄$) has been identified as a potential TL dosimeter (Zimmerman, 1971). Zircon can be found ubiquitously in sand-rich environments (e.g. van Es, 2008) – albeit in lower quantities than quartz and feldspar – and exhibits some unique features concerning luminescence-based dosimetry. Firstly, its internal U and Th contents are highly variable but typically amount to 10^2 – 10³ µg g−¹ for Th and U (Zimmerman, 1971; Sutton and Zimmerman, 1976). If these internal U and Th levels far exceed the radioelement contents in the sedimentary matrix, the short-range (in particular α -radiation) internal dose rate generated within the zircon dominates the total dose rate. External factors such as water content or spatial distribution of β - and γ -radiation, which usually add additional uncertainty, become far less important or negligible (Sutton and Zimmerman, 1976). The accrued dose in the zircons can be determined by comparing the natural TL with regenerated signals induced by α -irradiation in the laboratory (because of the dominating internal α-dose rate; classical approach), where the dose rate is derived from analyses of the U and Th content of individual grains (Sutton and Zimmerman, 1976). Despite the clear advantage of a dominant internal dose rate, the luminescence signal of zircons commonly suffers from anomalous fading, i.e. loss of signal over time in the hightemperature glow curve region that should be characterised by sufficient signal stability for dating (Wintle, 1973; Sutton and Zimmerman, 1976; Amin and Durrani, 1985; Templer, 1985b; Van Es et al., 2002a). Although Sutton and Zimmerman (1976) obtained TL ages of single zircon grains from ceramics largely in agreement with their known ages, they found all of the zircon ages were systematically underestimated, suggesting problems with anomalous fading. Besides, radioelements in many natural zircon grains are not uniformly distributed but are concentrated in layers resulting from zoning that occurred during initial crystal growth, where zones of high U abundance are anti-correlated with TL sensitivity (Vaz and Senftle, 1971; Sutton and Zimmerman, 1976; Templer and Walton, 1985). The negative relationship between TL sensitivity and U content (and equally Th content) is thought to arise from α -recoil damaging the crystal structure, a process that can be reversed by high-temperature (950 °C) annealing (Vaz and Senftle, 1971; Amin et al., 1983). This anti-correlation is difficult or even impossible to reproduce in the laboratory and leads to erroneously young ages because, in zoned zircon crystals, regions with higher U content produce a lower TL signal than in the U-poor zone, despite having the same age. As a result, the TL signal from U-poor areas (with lower dose rates) dominates the overall measurement, influencing the natural dose calculation. In conventional TL dating, the internal dose rate is estimated based on the average U and Th content without considering that most of the TL signal is emitted from U-poor zones. This leads to an overestimated dose rate and, consequently, an underestimation of the sample's age (Sutton and Zimmerman, 1976). Furthermore, natural zircon grains vary greatly in their optical properties, and Van Es et al. (2002b) recommend using only transparent grains for TL dating due to internal light absorption and radiation damage having substantially altered the dark and opaque grains. Especially for larger zircon grains, administering an external α-dose might result in underestimation of the natural dose, and hence age, because internal absorption of luminescence is stronger in the case of natural signals originating from the entire grain as compared to signals stemming solely from the α -irradiated outer shell of the grain. Irrespective of zoning and optical absorption in zircon, luminescence models predict inaccuracies in dose determination by comparing natural TL/OSL signals with those induced in the laboratory (Turkin et al., 2006). Appropriate heat treatments during or after irradiation might systematically improve inaccurate age estimates (Turkin et al., 2005, 2006), but confirmation by follow-up studies is pending.

The auto-regeneration dating technique represents an alternative to the classical zircon dating approach (Sutton and Zimmerman, 1976). It overcomes most of the methodological difficulties outlined above (including anomalous fading and zoning) and could circumvent the general limitations in age precision using quartz and feldspar. The basic principle of this technique, first applied to ceramics, is the zircon grains' capability to restore a measurable luminescence signal within months of storage in the dark (due to the high internal dose rate) after the natural signal's readout. In the linear dose-response range, the luminescence age can hence be calculated as follows (Templer, 1985a, 1986; Templer and Smith, 1988; Smith, 1988):

$$
age = \frac{\text{natural luminescence}}{\text{auto-regenerated luminescence}} \times \text{storage time.} \quad (1)
$$

The internal dose rate does not need to be quantified as a first approximation. With reference to Eq. (1), we note that the calculated final age is expected to vary based on whether the ages of individual grains are averaged or if the integrated luminescence of a set of grains (multi-grain aliquot) is utilised. This bias tends to yield a higher final age estimate when averaging the ages of individual grains, especially in the case of weak auto-regenerated signals (Ogliore et al., 2011). A few studies testified to the general applicability of the autoregeneration method on ceramics with dating accuracies of TL ages of 4 %–45 % (Templer, 1986), which may have the potential for significant improvements. For example, in the simplest scenario with a negligible external dose rate, the age uncertainty is only calculated from the counting statistics of two measured luminescence signals and their instrumental background levels (Templer, 1986), allowing dating precision of a few years. Notably, the standard error of an autoregenerated signal is inversely proportional to the storage time, but the storage time required to attain such precision probably depends on individual sample properties. Notwithstanding these initial achievements, significant obstacles in applying the auto-regeneration technique to zircons persist: (1) The routine separation of transparent zircon grains from bulk sediment in appropriate light conditions is challenging. Several procedures were proposed (Van Es, 2008; Sutton and Zimmerman, 1976; Smith, 1988), but they must be evaluated thoroughly and tested on various samples. (2) Despite some previous experimental work on the role of radioand trace elements and associated defects in the generation of TL and OSL in zircon (Shinno, 1986; Godfrey-Smith et al., 1989; Laruhin et al., 2002), our understanding of the relationship between the trace element composition of zircon and its properties relevant to auto-regeneration dating is still insufficient. (3) OSL ideally targets the signal with the best bleachability. Although some TL ages of detrital zircons, representing the last sunlight exposure event, are in reasonable agreement with independent age control (albeit not obtained with auto-regeneration but by using additive γ -irradiation; Van Es, 2008), to the best of our knowledge, the OSL emission of zircon has not yet been thoroughly tested or applied for dating purposes. (4) Templer (1986) and Smith (1988) reported low TL and OSL signal levels after laboratory storage, precluding OSL from being used in auto-regeneration. Luminescence readers developed in the last 2 to 3 decades now allow single-grain-level stimulation at much higher power densities ($> 50 \,\mathrm{W\,cm^{-2}}$; Duller et al., 1999) than those used in previous studies (\sim 0.002 W cm⁻²; Smith, 1988), with the potential to substantially improve the signal-to-noise ratio for auto-regenerated zircon OSL.

To test the potential of the zircon auto-regeneration approach using state-of-the-art equipment, we address items (2), (3), and (4) listed above by characterising the TL and OSL signal of two zircon samples in terms of signal composition, rates of optical resetting, and the ability to autoregenerate. These data are supplemented by cathodoluminescence images and trace element analytics by laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS) to characterise the distribution and quantify contents of U and Th in zircons and to establish potential links between luminescence behaviour and zircon geochemistry. All results are discussed regarding their implications for advancing the zircon auto-regeneration method towards a competitive, highly accurate, and reasonably precise chronometric tool to investigate land surface changes and archaeological deposits on Mid- to Late Pleistocene and Holocene timescales.

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2.1 Samples and sample preparation

We selected two zircon samples from different environments for the methodological investigations. Sample ZR229 was extracted from Fe–K syenogranite from the Mont Blanc massif (France and Switzerland) and represents a short-lived magmatic event around 303 Ma (Bussy et al., 2000). Sample Can1 is detrital and originates from alluvial sand collected from Rio Carrao close to Canaima (Venezuela). It represents the weathering product of orthoquarzitic sandstone from the Precambrian Roraima Group, which forms the tepuis of the Gran Sabana. The sandstones from this group have a minimum age of 1.5–1.6 Ga (Briceño and Schubert, 1990) or ∼ 1.8 Ga based on zircon U–Pb ages of mafic sills (Santos et al., 2003).

Zircons from sample ZR229 were extracted in large quantities in the 1960s by Ronald Chessex using heavy liquid (bromoform at 2.89 g cm−³ and diiodomethane at 3.32 g cm^{-3}) and magnetic separation at a slope and tilt at 15–20°, and no further grain size selection was applied. While the largest grains reach \sim 150 µm along their c axis, most grains are considerably smaller $\left(< 80 - 100 \,\mu\text{m} \right)$. The sieved (63–100 μ m) Can1 sample was treated with 10 % HCl to dissolve carbonates and run through a Frantz L-1 magnetic separator (slope 15°, tilt 15°, current through magnet 1.7 A; Porat, 2006), and the non-magnetic fraction was then etched in 40 % HF at room temperature for 2 d to dissolve all heavy minerals but zircon while the sample was continuously stirred. After 1 d, HF was replenished. After 2 d of HF etching, inspection of grains indicated the presence of light- and dark-coloured heavy minerals (potentially fluorite or rutile), necessitating an additional HF etch for 3 d under the same conditions to optimise the zircon yield. This way, we could extract a few hundred zircons from an initial sample mass of \sim 500 g.

2.2 Mineral characterisation through cathodoluminescence and LA-ICP-MS

We prepared epoxy resin mounts for 42 and 43 zircon grains of samples ZR229 and Can1, respectively, by polishing them down to their equatorial plane using diamond paste (see Supplement, Figs. S2 and S3). The mounts were carbon coated, after which the polished grain surfaces were imaged using a CamScan MV 2300 SEM operated at an acceleration voltage of 10 kV, with a sample positioned at a working distance of \sim 40 mm. Panchromatic cathodoluminescence (CL) images were taken to visualise the growth structure of zircon grains (zoning) and to optimally select the location of the laser spots for LA-ICP-MS. The carbon coating was removed prior to LA-ICP-MS analyses.

LA-ICP-MS allows us to quantify isotope ratios and elemental abundances along a cylindrical pit created through laser ablation into a solid surface. Here, we used a RESOlution-SE 193 nm ArF excimer laser ablation system equipped with an S155 two-volume ablation cell (Applied Spectra, USA) and interfaced to a sensitive fast-scanning triple-quadrupole ICP mass spectrometer, NexION 5000 (PerkinElmer, USA). The laser repetition rate was set to 8 Hz, and the on-sample energy density was set to 2.9 J cm^{-2} . The beam size was 24 μ m for all analytical spots on zircon grains. The list of analysed isotopes included 27 Al, 29 Si, 42 Ca, 49 Ti, 85 Rb, 88 Sr, 89 Y, 93 Nb, 139 La, 140 Ce, 141 Pr, 143 Nd, 147 Sm, ¹⁵¹Eu, ¹⁵⁷Gd, ¹⁵⁹Rb, ¹⁶³Dy, ¹⁶⁵Ho, ¹⁶⁶Er, ¹⁶⁹Tm, ¹⁷²Yb, ¹⁷⁵Lu, ¹⁷⁸Hf, ¹⁸¹Ta, ²⁰¹Hg, ²⁰⁴Pb, ²⁰⁶Pb, ²⁰⁷Pb, ²³²Th, and ²³⁸U. Their intensities were acquired sequentially using the mass spectrometer in the single-quadrupole regime (Q1 as RF-only ion guide and Q3 as mass filter). Each measurement was reduced twice to quantify the trace element composition and to estimate the ²⁰⁶Pb / ²³⁸U and ²⁰⁷Pb / ²³⁵U ages, respectively. The trace element data were reduced against the NIST SRM 612 soda–lime–silica glass standard (Jochum et al., 2011) as a primary standard for the determination of relative sensitivity factors. It was repetitively ablated at a beam size of $50 \mu m$, using the same repetition rate and energy density as for the sample zircons. Quality control was ensured using the BCR-2G basaltic glass as a secondary standard. For internal standardisation, 29 Si was selected. Its abundance was set up stoichiometrically (32.9 wt $\%$ SiO₂). The U / Pb ages were computed against the GJ-1 reference zircon (Boekhout et al., 2012; Ulianov et al., 2012) as a primary standard that was repetitively analysed at exactly the same ablation settings as for the sample zircons. As secondary standards for samples ZR229 and Can1, respectively, the Plešovice (Sláma et al., 2008) and Harvard 91500 (Wiedenbeck et al., 1995) reference zircons were used. The reference 206 Pb $/$ 238 U ages of these two standards $(337.13 \pm 0.37$ and 1062.4 ± 0.2 Ma at 95 % confidence level) could be reproduced within 0.2 % $(n = 5)$ and 2 % $(n = 7)$, respectively. Both trace element and U / Pb data were reduced in LAMTRACE (Jackson, 2008; detection rules modified after Ulianov et al., 2016). The transient signals were carefully inspected for the presence of intensity spikes and domains that could contain inclusions, cracks, or other sources of chemical alteration and for the consistency of the time-resolved U / Pb spectrum. As monitors of chemical alteration, we used ²⁷Al, ⁸⁵Rb, ⁴²Ca, and, in some cases, ¹³⁹La. The canonical understanding of LA-ICP-MS usually implies the rejection of altered domains. In this study, however, we created two parallel trace element datasets based on the bulk signal (corresponding to the integrated signal across the entire pit) or on specific signal segments selected so as to be devoid of alteration and contamination as much as possible and thus representing the "pure" zircon crystal structure. As the smallest entities relevant for luminescence analyses are single grains, the bulk trace element abundances, of radioelements in particular, cannot be discarded a priori. For Th and especially for U, the bulk and structural values are, however, quite similar (Fig. S1): high contents of these elements usually present in the zircon crystal structure are difficult to change by alteration and contamination. Measured uncertainties amount to \sim 2 %–5 % for values in the mg g^{-1} range, i.e. for elemental abundances well above the critical level of detection. The latter condition was met for most elements and analytical spots. For each zircon grain, either one or two analysis locations (spots) were chosen, guided by inspection of individual grains' zoning as displayed in the CL images.

2.3 Luminescence measurement parameters

For single-grain OSL measurements, zircon grains were placed in the $300 \mu m$ diameter holes of Risø single-grain discs using fine tweezers or a needle. Due to the small average size of the zircons, some holes contained more than one grain due to the electrostatic charge of grains. This information was recorded to retrospectively relate luminescence characteristics to the number of grains in individual holes. The TL signals from the holes were measured simultaneously and hence represent a composite signal for the entire disc.

TL and OSL were recorded with Risø TL/OSL DA-20 readers equipped with a UV-sensitive EMI 9235QB photomultiplier tube (PMT) showing a dark count rate of 35 ± 9 cts s⁻¹. TL signals were not filtered for autoregeneration experiments. To maximise the system's sensitivity and to also detect very dim TL signals close to the instrumental background level, the OSL unit was replaced by the TL flange for these measurements, resulting in a smaller distance between the sample and the PMT and, consequently, a larger solid angle captured by the detector. The heating rate was set to 1 K s^{-1} in all TL experiments. TL spectra were measured with an Andor Kymera 193i spectrograph coupled to a UV-enhanced Andor iXon Ultra 888 electron-multiplying charge-coupled device (EMCCD) camera installed on the DASH of a Risø TL/OSL DA-20 reader. Further technical details on the system are described in Bartz et al. (2023). All spectra reported are energy- and efficiencycalibrated. The background relevant for TL measurements was recorded in a second run, using identical parameters as in the measurement of the sample signal. OSL was stimulated by an Nd:YVO⁴ solid-state laser emitting at 532 nm with a nominal power density at the grain location of $45 \,\mathrm{W \, cm^{-2}}$ and recorded at 125 °C through a 7.5 mm Hoya U340 filter and with the PMT described above. The background relevant for OSL measurements was approximated by the signal recorded by the PMT before the stimulation source was switched on, and no preheat was applied unless stated otherwise. TL and OSL signals were regenerated in the laboratory by irradiation with a ^{90}Sr / ^{90}Y β -source characterised by a dose rate of 0.087 Gy s^{-1} at the time of measurement, as calibrated with Risø calibration quartz batch #200 (Hansen et al., 2015; Autzen et al., 2022). OSL signals were decomposed using the function fit_CWCurve of the R package "Luminescence" (Kreutzer et al., 2012; Kreutzer, 2023; R Core Team, 2024).

3 Results

3.1 Cathodoluminescence

Representative CL images are shown in Fig. 1; a summary of all studied grains can be found in the Supplement. Most grains exhibit zoned growth, as indicated by mostly concentric areas of CL of various intensities. While concentric zones characterise the zircons of sample ZR229, the growth structure of the Can1 zircons is much more complex, with initial (zoned) igneous or metamorphic zircon nuclei overgrown by younger, discordant domains (see Figs. 1c and S4). Considering the zircon typology scheme by Pupin (1980), it appears that both zircon samples differ in the alkalinity of the melt, but, given the multiple growth and abrasional cycles preserved in the Can1 zircons, it remains unclear whether the more rounded shape of the latter relates to a lower crystallisation temperature, metamorphic overprinting and/or to physical abrasion (Markl, 2014). Overall, CL allowed us to visualise the different origins of the two studied samples as indicated by contrasting grain morphology and zoning, reflecting the granitic origin of sample ZR229 on the one hand and the sedimentary transport of zircon grains from their source region in the Trans-Amazon Belt north of the Roraima Basin and the diagenesis experienced by the detrital sample Can1 on the other hand (Santos et al., 2003).

3.2 LA-ICP-MS

Analytical results confirmed that all 42 grains of sample ZR229 are zircons, while 4 out of the 43 grains of sample Can1 were discarded due to their elemental composition not aligning with the characteristic pattern for ZrSiO4. Radioelement (Th, U) contents for the bulk material analysed in one spot (i.e. when integrating over the entire transient signal during ablation) are shown in Fig. 2 and given in detail in Tables S3 and S4. For sample ZR229, Th and U contents are positively correlated and vary by roughly 2 orders of magnitude (100–10 000 µg g^{-1}); about half of the analytical spots yielded values between 100 and 1000 μ g g⁻¹ Th and U. The highest values recorded were 6990 μ g g⁻¹ for Th and 8057 µg g−¹ for U, observed on two different grains, while the lowest contents measured for ZR229 were 97 μ g g⁻¹ for Th and 226 μ g g⁻¹ for U (also two different grains). The average abundances of Th and U for 48 analytical spots totalled 760 and 2456 µg g−¹ , respectively. Figure 2 also suggests that there is no systematic difference in radioelement abundances between the core and the rim of zircon grains.

In contrast, Th and U contents are overall lower in zircon grains of sample Can1, with more than half of the grains giving contents between 10 and $100 \,\mu g \, g^{-1}$. The lowest and highest abundances for Th amount to 1 and $404 \mu g g^{-1}$,

and the lowest and highest abundances for U amount to 15 and 293 μ g g⁻¹. Average values were 71 μ g g⁻¹ for Th and 98 µg g⁻¹ for U, and the dispersion of data (Fig. 2) indicates that grain-to-grain variation in the internal dose rate is smaller in sample Can1 compared to ZR229. Similarly to sample ZR229, grouping the analytical spots into those located in the core of a zircon grain and those located in the rim resulting from a later phase of zircon growth does not reveal any systematic differences in radioelement abundance between these zones (Fig. 2). However, particularly for sample Can1, the differentiation between the core and the rim of zircon grains was often challenging due to frequent abrasion affecting significant portions of the grains. For both samples, the variation in radioelement contents within one grain (i.e. zonation) is smaller than the variation in averaged Th and U contents between grains. For instance, the average variation in the U content within grains of samples ZR229 and Can1 amounts to 75 % and 28 %, whereas the inter-grain scatter of averaged U abundance adds up to 110 % and 52 %, respectively.

Contrasting elemental abundances suggest that the structural Th and U contents in sample Can1 represent by far the most important part of the total, or bulk, Th and U budget of the individual zircon grains, whereas, in sample ZR229, the structural and bulk contents of these elements are less well matched (Fig. S1). There appears to be an indistinct trend in the data for both samples that hints at contaminated or altered (parts of the) grains being enriched in Th and U relative to their structural contents.

Bulk radioelement contents (as the smallest entities relevant for luminescence analyses are single grains) can be translated into internal dose rates that give rise to autoregenerated luminescence. When using the conversion factors of Guérin et al. (2011), α -grain size attenuation factors by Brennan et al. (1991), β -grain size attenuation factors by Guérin et al. (2012), an assumed α-efficiency (*a*-value) of 0.03 ± 0.003 , and a grain size range of 60–120 µm, the average U and Th contents for samples ZR229 and Can1 result in internal α -dose rates of 181 Gy ka⁻¹ and 7.9 Gy ka⁻¹, respectively, as calculated with DRAC (v1.2; Durcan et al., 2015). Internal β -dose rates amount to 34 Gy ka⁻¹ and 1.5 Gy ka−¹ . Assuming a sedimentary matrix with representative radioelement abundances of 1.5 wt % for K, 10μ g g⁻¹ for Th, and $3 \mu g g^{-1}$ for U, the internal dose rate accounts for \sim 99 % and \sim 76 % of the total environmental dose rates for samples ZR229 and Can1, respectively.

Previous studies found that the TL and CL sensitivity of domains in zircon grains characterised by high Th and U abundance was lower than in domains with low radioactivity (Vaz and Senftle, 1971; Sutton and Zimmerman, 1976; Templer and Walton, 1985). This observation was explained by α -recoil damage of the crystal structure in zones of high activity. Here, in case different domains within one zircon grain were analysed separately by two LA-ICP-MS spots, we grouped these two domains into a dim zone and a bright zone

Figure 1. Cathodoluminescence (CL) images of selected zircon grains, with analytical spots from LA-ICP-MS and measured Th and U contents indicated. (a, b) Sample ZR229. (c) Sample Can1. (d) The abanico plots (Dietze et al., 2016) show the U–Pb age distributions for the individual grains dated by LA-ICP-MS.

Figure 2. Radioelement contents (Th, U) for zircon samples ZR229 (a) and Can1 (b), as determined by LA-ICP-MS. Measured uncertainties are in the order of 2 % and not plotted here. Each dot represents one spot analysis located in the core or the rim of grains, while two spots were measured for some grains (in which case both core and rim were analysed).

(relative to each other), based on CL images. The determined radioelement contents for these paired datasets are displayed in Fig. 3 and clearly support a correlation between CL intensity and Th and U abundance: except for a few grains, dim CL corresponds to high radioactivity and vice versa (see also Fig. 1). This trend is notably more pronounced for elevated radioelement abundance and larger U and Th differences between the two analysed domains, while for domains characterised by relatively low radioactivity, factors other than Th and U contents may exert an influence on CL intensity.

LA-ICP-MS analyses facilitated U–Pb dating of the zircon formation age for both samples, which – together with the radioelement abundance – might yield insights into potential metamictisation and its effects on luminescence properties. Altogether, 7 out of 49 analytical spots of sample ZR229 did not yield concordant $^{206}Pb / ^{238}U$ and $^{207}Pb / ^{235}U$ ages (discarded from Fig. 1d). The main reasons for age discordance were the loss of radiogenic lead, the presence of small amounts of common lead, and strong heterogeneity within the ablated volume manifested by inconsistent Pb–U fractionation patterns and strong zoning, including Th / U ratios < 0.1 characteristic for metamorphic zircon grains (Yakym-

Figure 3. Th (a) and U (b) contents measured at two different analytical spots on one zircon grain. A relative comparison of cathodoluminescence (CL) brightness for each grain allowed analytical spots to be grouped into "CL dim" (plotted on the x axis) and "CL bright" (plotted on the y axis); see Fig. 1 for example CL images. The dashed line represents the 1 : 1 line.

chuk et al., 2018) that yield a younger age than initial igneous crystallisation. For sample Can1, 2 out of 49 analytical spots were removed from the U–Pb dataset due to the sampling of domains producing an age much younger than the average age (although their Th / U ratio did not indicate metamorphism). These grains are probably derived from younger source rocks and then mixed with older grains in the Roraima detritus. The 42 accepted measurements from sample ZR229 yield a weighted mean age of 302.4 Ma with an estimated uncertainty range between ± 1.1 Ma (measured) and ± 4.2 Ma (full range including systematic uncertainties) at 68 % confidence level. This age estimate agrees well with the age of 303 ± 2 Ma for the Mont Blanc granite given by von Raumer and Bussy (2004). Considering the full set of spots, we obtain a weighted mean age of 2114 Ma for sample Can1 with an estimated uncertainty ranging between ± 42 Ma (measured) and ± 50 Ma (full range including systematic uncertainties) at 68 % confidence level. This estimate is consistent with the minimum age of 1600–1500 Ma for the Roraima sandstone as obtained for dated diabase intrusions (Briceño and Schubert, 1990) and the U–Pb ages obtained for the Roraima Supergroup by Santos et al. (2003). Spot measurements yielding U–Pb ages different from those in the main age mode (Fig. 1d) would allow further distinction of zircon sources, but, in the context of this study, a bulk age of the sample is sufficient to constrain the time available to produce radiation damage. The age difference and the variance in Th and U contents between the two zircon samples studied here thus cover roughly 1 order of magnitude, with the younger sample (ZR229) producing a much higher internal dose rate than the older sample (Can1).

3.3 TL spectra

Our initial experimentation involved applying various β doses before TL spectrum acquisition, revealing that relatively high doses ($> 2000 \text{ Gy}$) are required to generate a sufficiently intense signal suitable for spectral analyses. Figure 4 shows two typical TL spectra of samples ZR229 (Fig. 4a) and Can1 (Fig. 4c) as contour plots and the spectral composition of TL signals for two different temperature intervals. Both spectra exhibit a broad emission, spanning ∼ 320– 450 nm (centred at ∼ 400 nm), and share an emission centred at ∼ 640–650 nm. The former emission covers almost the entire recorded temperature range (25–400 °C) and displays an increase in intensity for glow temperatures $>$ 350 °C. In contrast, the latter attains its peak intensity within temperatures of 100–200 °C (ZR229) or 150–210 °C (Can1). In addition to these common features, the TL spectrum of sample ZR229 yields narrow emission bands centred at 480 and 580 nm, which can be traced for glow temperatures between 80 and 280 °C. This sample possibly shows an additional emission beyond 650 nm, as indicated by the broad emission plateau in the wavelength region 620–700 nm (Fig. 4b).

3.4 OSL signal and its composition

We show typical OSL decay curves of samples ZR229 and Can1 in Fig. 5. The sensitivity (in cts s⁻¹ Gy⁻¹) and the optical decay rate vary substantially from grain to grain, covering about 2 orders of magnitude. The background-subtracted integrated OSL signal over the entire stimulation period (0.25– 2.75 s) is plotted against the number of grains in each hole of the single-grain disc in Fig. 6. Comparing the signals obtained for holes without grains with those induced by one single grain suggests that the fraction of grains that do not give rise to any significant OSL is higher in sample Can1 than in ZR229, in which case all grains seem to produce measurable OSL following a β -dose of \sim 200 Gy. Furthermore, the counterintuitive observation that measurements involving single grains yield, on average, larger OSL signals than the simultaneous measurement of several grains can be best explained by the fact that placing just one grain in the hole of a singlegrain disc was much easier to realise for comparatively large zircons, while it was harder for smaller zircon grains that in many cases could not be separated from each other easily due to electrostatic charging. Figure 6 hence implies that the OSL signal from large zircon grains tends to be more intense than that from smaller grains and that there is a positive correlation between grain size/volume and OSL produced. This trend is more pronounced for ZR229, while this sample also produces single-grain OSL signals with a median more than 4 times larger than for sample Can1. This observation and the overall larger scatter in OSL intensities of sample ZR229 between holes are consistent with the grain size spectrum of both samples, which is well constrained to $63-100 \,\mu m$ for Can1, while no grain size separation was done for ZR229. The integrated OSL intensity, along with the information on the number of grains in each hole, allows the average OSL sensitivity of one zircon grain for each of the two samples to be estimated. It amounts to \sim 1.7 cts Gy⁻¹ for both samples for the summed OSL signal over the stimulation period of 2.5 s, with estimated minimum standard deviations (due to simultaneous measurement of multiple grains in some holes) at 1σ of 1.2 cts Gy⁻¹. However, these averaged figures completely mask the observation that larger grains give rise to the strongest OSL signals (see Fig. 6). Normalising OSL sensitivity to sample mass might be a better alternative.

Previous studies suggested that the zircon OSL signal comprises several components characterised by different optical decay rates or photoionisation cross-sections (Smith, 1989), similar to later observations for quartz OSL (Bailey et al., 1997). To gain further insights into zircon OSL signal properties, we decomposed the regenerated (\sim 200 Gy) continuous-wave OSL decay curves obtained from measuring one or more grains in the holes of a single-grain disc. Optical decay rates were translated into photoionisation crosssections, assuming the accuracy of the green laser's stimulation power density. About 25 %–34 % of OSL decay curves could be fitted with one exponentially decaying component, while 59 %–65 % required two components (see examples shown in Fig. S7). Only 6 %–13 % of the decay curves necessitated three components for a satisfactory fit; exact numbers are provided in Table 1, and a summary of fitted components and their photoionisation cross-sections is shown in Fig. S8. The distribution of photoionisation cross-sections of the two "fastest" components is displayed in Fig. 7 for samples ZR229 and Can1 as a histogram and probability density function (PDF). Calculated values for the photoionisation crosssection span a range from 2.4×10^{-22} to 3.2×10^{-19} cm². It appears that heating sample ZR229 to 180 °C prior to OSL measurement does not substantially change the bimodality and values of photoionisation cross-sections (see Fig. 7a and b). Furthermore, whenever just one component could be fitted, its photoionisation cross-section is, in the great majority of cases, smaller than $\sim 5 \times 10^{-20}$ cm², suggesting that, for those grains, the fastest component (with a photoionisation cross-section in the order of 10^{-19} cm²) is absent in the signal.

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Figure 4. TL spectra of samples ZR229 (a, b) and Can1 (c, d) following a regenerative β-dose of 2000 and 3200 Gy, shown as contour plots (a, c) and cross-sections for glow temperature intervals $100-175$ and $300-350$ °C (b, d). All data shown are background-subtracted and energy- and efficiency-calibrated.

Table 1. Proportion of the components required to fit the OSL decay curves of the studied samples satisfactorily. The ZR229-PH sample was preheated to 180 °C prior to measurement.

Sample	ZR ₂₂₉	ZR229-PH	Can ₁
	(%)	(%)	(%)
1 component	0.29	0.34	0.25
2 components	0.65	0.59	0.62
3 components	0.06	0.07	0.13

3.5 Bleaching of zircon OSL

The zircon OSL decay curve decomposition results suggest that the reduction rate of zircon OSL under sunlight exposure is slower than that of quartz OSL. To test this assumption and the suitability of the zircon OSL signal to date light

exposure events, we conducted a bleaching experiment on sample ZR229. A single-grain disc populated with one or more grains per hole was repeatedly irradiated with a β dose of ∼ 220 Gy and exposed to the light from a Hönle UVACUBE 400 (see emission spectrum in Fig. S8), and the residual OSL signal was measured. The light exposure times were set to 0, 10, 30, 60, 120, and 300 s. To distinguish OSL signal reduction due to fading from the reduction induced by light exposure, we ensured that the delay between irradiation and measurement was always kept at 16 min, irrespective of the bleaching duration. In this way, fading was identical in all bleaching cycles, and the net effect of bleaching could be isolated. Following the reading of the residual signal, a test dose cycle (administering a β -dose of 200 Gy) was inserted to correct for any sensitivity changes. This experiment was carried out once with sample ZR229 receiving a cutheat

Figure 5. (a) OSL decay curves resulting from green laser stimulation of zircon grains in the holes of a single-grain disc following \sim 200 Gy (Can1) or \sim 540 Gy (ZR229) regenerative β -irradiation, demonstrating that OSL sensitivities of individual grains of one sample (ZR229) and of grains from different samples vary by almost 2 orders of magnitude. Some grains did not yield a signal above the background (not shown here). The first five channels (0– 0.25 s) and the last five channels (2.75–3 s) represent the instrumental background recorded before the stimulation light was switched on. (b) Normalised OSL decay curves of grains from both samples indicate that the OSL signal decays at very different rates on a grain-to-grain level.

using an arbitrarily selected temperature of 180 °C directly following β -irradiation and once without preheating.

Experimental results include only positions in the singlegrain disc yielding monotonic decay with bleaching time and giving integrated signals at 0 s bleaching time larger than 100 cts (∼ 3 times the standard deviation of the background; Fig. 8). Although some scatter between residual signals of the same bleaching time can be observed, the boxplot in Fig. 8a confirms the reduction in the average residual signal by $> 50\%$ following 10 s light exposure. The zircon OSL signal decayed to negligible values $\left(< 3\% \right)$ of the initial signal) after 300 s bleaching. Although the trend in OSL signal reduction with bleaching time is the same for the nonheated and the preheated samples, it appears that preheating removes part of the most light-sensitive OSL components.

Contrasting the bleaching characteristics of other commonly used luminescence signals for dating (Fig. 8b), zircon OSL bleaches at least 1 order of magnitude faster than

Figure 6. Integrated OSL signal (0.25–2.75 s) as a function of the number of grains per hole of a single-grain disc, as counted under an optical binocular, for samples ZR229 (a) and Can1 (b).

any IRSL signal of feldspar but requires a few minutes more of sunlight exposure to be fully reset as compared to quartz OSL (Colarossi et al., 2015).

3.6 Auto-regeneration of zircon OSL and TL

Following the experiment on the optical resetting rate of zircon OSL, we investigate its signal's potential for autoregeneration dating. One crucial requirement with this is the ability of zircon OSL to build up a measurable signal within a reasonable amount of time, ideally not exceeding 1–2 a. We prepared one single-grain disc for which all holes were filled with merely one zircon grain of sample ZR229 and a second one with each hole containing more than one grain of the same sample. All holes were initially stimulated for 4 s with the green laser (see Sect. 2.3) to erase inherited signals, and the two single-grain discs were then stored in the dark at room temperature (\sim 20 °C). After storage periods of 64, 127, 291, and 540 d, the auto-regenerated signal of the grains in holes 1–20, 21–40, 41–60, and 61–80 were measured, respectively. Although sample ZR229 was prepared and handled under daylight conditions, some grains yielded an initial OSL signal significantly above instrumental background, indicating that green laser stimulation is more effective in resetting the OSL signal than daylight. To ensure that all zircon

Figure 7. Distribution of photoionisation cross-sections obtained by fitting single exponential decays to OSL transients of samples ZR229 (a), ZR229 preheated to 180° C (b), and Can1 (c).

grains were set to zero before starting the auto-regeneration experiment and that we could also detect weak induced OSL signals, we excluded holes from the analyses that showed an integrated OSL signal (0–2.5 s of the decay curve) larger than 40 cts s^{-1} during the initial screening of the bleached sample. The results of the OSL auto-regeneration experiment are shown in Fig. 9, with the measured OSL signals for each hole represented by boxplots. It is evident that, over a storage period of 540 d (∼ 1.5 a), the OSL signal did not increase

Figure 8. Results from the bleaching experiment with zircon grains of sample ZR229 placed in a single-grain disc. (a) Boxplots of all residual signals measured following various bleaching times. (b) Comparison of zircon OSL residual signals as a function of bleaching time with those of other commonly used luminescence signals for dating. Zircon OSL data points represent the average of between 63 and 90 individual signals; the preheat temperature was 180 °C. The data for quartz OSL and K-feldspar IRSL signals were taken from Colarossi et al. (2015).

significantly but remained barely distinguishable from the instrumental background.

Due to these dim auto-regenerated OSL signals and previously reported higher TL sensitivity of zircon, we investigated the potential to exploit the TL auto-regeneration signal. To this end, one single-grain disc with one or multiple zircon grains of samples ZR229 and Can1 was heated to 400 °C to erase all previously accrued doses and then stored in the dark for 21 d. The auto-regenerated signals following this period are displayed in Fig. 10. While sample Can1 shows a negative net signal for glow temperatures $< 240 \degree C$, indicating background reproducibility issues, sample ZR229 displays a notable positive TL signal of up to $400-500$ cts s⁻¹. It rises sharply for glow temperatures $> 100 °C$ and flattens out at 170–250 °C. The integrated net signal from the temperature interval 0–280 °C is larger than 3σ of the corresponding background (which was measured four times following

Figure 9. Boxplot of OSL auto-regeneration results for sample ZR229. Different colours represent analyses for single grains ("SG") and multiple grains ("MG") loaded into the holes of a singlegrain disc. A total of 20 holes in a single-grain disc were analysed for each storage period.

Figure 10. Auto-regenerated TL (not wavelength-filtered) of 505 zircon grains of sample ZR229 and 138 grains of sample Can1 following 21 d of storage without light exposure. Net TL signals are shown with their background (measured by a second glow) already subtracted.

the readout of the auto-regenerated signal), confirming that it is not an artefact caused by background fluctuations. We repeated the TL auto-regeneration experiment for a storage period of 28 d with a very similar outcome for both samples.

4 Discussion

4.1 Geochemistry of zircons in relation to their luminescence

Amin and Durrani (1985) suggested that the TL spectrum of zircon changes as a function of radiation damage (metamictisation), with highly damaged zircons exhibiting only an orange-red emission, while less damaged zircons show an additional emission in the blue-green range (480 nm). This observation is consistent with spectrally resolved TL from the two samples studied here. Sample Can1 is about 1 order of magnitude older than sample ZR229 and had more time to accumulate radiation damage. Indeed, this sample emits predominantly in the orange-red part of the spectrum, while ZR229 features additional emissions at shorter wavelengths. However, when the radioelement abundance in these two samples is taken into account, the situation becomes less clear. Averaged across all grains analysed by LA-ICP-MS, Th and U contents are higher by a factor of \sim 10 and \sim 20 for sample ZR229. This difference in radioelement abundance between the two samples should have roughly equilibrated the integrated radiation damage effects and led to qualitatively more or less similar TL spectra. Therefore, other factors, such as the different geochemical composition of the samples, likely exert an additional influence on the TL properties (see Fig. 3b). It should also be noted that the mineralogical history of the two samples is strikingly different, with the detrital zircons of the older sample, Can1, exhibiting a much more complex pattern of grain evolution, possibly including several erosional and depositional cycles and associated moderate (but long-term) heating through diagenesis. These divergent sample histories might also have left their imprint in the TL spectra of both samples beyond the alteration induced by radiation damage. Vaz and Senftle (1971), Jain (1978), Amin et al. (1983), and Amin and Durrani (1985) found that high-temperature annealing above ∼ 500 °C increases the luminescence sensitivity and may restore the TL spectrum as observed for less damaged zircons. Similarly, Godfrey-Smith et al. (1989) noted enhanced luminescence sensitivity following annealing, accompanied by increased crystallinity, as detected by X-ray diffraction. Here, we annealed samples only to temperatures of 400 °C, in the course of which we could not detect any pronounced TL spectral and sensitivity changes. On the sub-grain spatial scale, our data confirm the previous finding that both CL and TL intensity are negatively correlated with the radioelement content in different domains of zircon crystals, especially for high U and Th abundances (Vaz and Senftle, 1971; Amin et al., 1983; Templer and Walton, 1985; Nasdala et al., 2003).

The TL spectra of samples ZR229 and Can1 yielded a broad emission centred at ∼ 400 nm across a wide range of glow temperatures (room temperature to 400 °C) and a red emission at \sim 640 nm, peaking at 170–180 °C. In addition, sample ZR229 shows superimposed sharp emission lines at \sim 480 and \sim 580 nm that occur between 80 and 400 °C; these lines are not present in the spectrum of sample Can1. Iacconi et al. (1980) explained the broad blue-violet emission observed in both our samples as an intrinsic emission of the SiO4[−] 4 ionic group in the ZrSiO⁴ host lattice. An electron is detached from an oxygen atom through irradiation and then trapped around a central Si atom. Recombination of the electron causes an emission at ∼ 365–380 nm (Iacconi et al., 1980; Godfrey-Smith et al., 1989), which is slightly shifted towards a shorter wavelength relative to our observations. However, it is hard to identify a well-defined central wavelength due to the overall low intensity of this emission in both

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samples. This emission occurs over a large range of temperatures and may be related to the observation and theoretical prediction that the distance between oxygen and silicon atoms in $ZrSiO₄$ can vary, resulting in a (quasi-)continuous distribution of activation energies. This effect was calculated and observed for the TL of synthetic zircon in the temperature range of $100-350 \text{ K } (-173 \text{ to } 77 \text{ °C})$ (Iacconi et al., 1980). For the photoluminescence of zircon, Shinno (1986) explained broad emissions in the spectrum occurring at various energies with centres created through α -decay of U and Th. The intensity of such emissions was proposed to relate to the integrated radiation damage and hence to mineral formation age (or re-crystallisation or annealing age), which, in the case of the red TL emission at ∼ 640 nm, is consistent for the two samples studied here. In comparison, such a relationship cannot be identified for the broad TL emission at ∼ 400 nm. The narrowband emissions in the TL spectrum of sample ZR229 appearing at ∼ 480 and ∼ 580 nm are in agreement with many previously published TL spectra of zircon (Jain, 1978; Kirsh and Townsend, 1987; Huntley et al., 1988; Van Es et al., 2002a) and are generally attributed to rare earth elements (REEs) in the crystal lattice, with Tb^{3+} (Jain, 1978), Eu^{3+} (Nicholas, 1967), or a combination of Dy^{3+} and Tb^{3+} (Shinno, 1986; Laruhin et al., 2002) as the primary responsible activators. Laruhin et al. (2002) and Van Es et al. (2002a) argue that Dy^{3+} gives rise to narrowband and intense TL emissions at lower glow temperatures (100–175 °C), while Tb^{3+} -related emissions dominate the TL spectrum at higher temperatures (300–400 °C). Based on these previous observations, one might expect a clear difference in Dy and Tb contents between samples ZR229 and Can1, given their different TL spectra regarding the narrowband emissions. Indeed, LA-ICP-MS results averaged across all analytical spots of a sample (see Tables S3 and S4) demonstrate higher contents of both elements in ZR229 (Tb: $17 \pm 12 \mu g g^{-1}$; Dy: $212 \pm 140 \,\mu g \,g^{-1}$; uncertainty range indicates the standard deviation at 1σ) as compared to Can1 (Tb: $6 \pm 4 \,\mu$ g g⁻¹; Dy: $67 \pm 44 \,\mu g \, g^{-1}$). Although these results potentially explain the absence of narrowband emissions at 480 and 580 nm in sample Can1, they do not provide definitive information on whether Dy^{3+} or Tb³⁺ accounts for those. By contrast, Godfrey-Smith et al. (1989) did not detect any clear correlation between Dy (or any other REE) content and specific emissions and overall luminescence sensitivity. Given the geochemical and structural differences between the two studied zircon samples, it is interesting that they yield very similar OSL sensitivities and OSL signal composition (see Sect. 3.4). As OSL was detected in the region ∼ 270–380 nm, this could imply that both OSL and TL share the same recombination centres, i.e. those giving rise to the broad band centred around 400 nm in the TL spectra of both samples (Fig. 4). However, to address this question in detail, element analytical results of individual grains should be linked to their specific TL and OSL characteristics, which was not feasible in the scope of this study.

4.2 Auto-regeneration dating

4.2.1 Optical signal resetting

One of the main aims of this study was to test if the OSL signal of zircon is a promising candidate for auto-regeneration dating, given the technological advances made in measurement equipment and data analysis since the initial development of the method in the 1980s. An essential requirement for a mineral to successfully date a sunlight exposure event is the OSL signal's fast optical resetting. Our experiments demonstrate that, for the sample studied, the initial portion of the zircon OSL signal is reduced to negligible levels af ter > 300 s of exposure to simulated sunlight, in line with the bleaching rate of zircons in response to a 60 W tungsten filament lamp, resulting in 10 % residual signal after 120 s exposure (Smith, 1988). Future studies should investigate the bleaching rate for more samples, including coloured and opaque zircons. In terms of dating light exposure events of short duration or attenuated photon flux (e.g. slope wash, fluvial transport), this provides a distinct advantage over the TL signal of zircon, which was reported to show residuals of 35 %–65 % of the initial signal after 30 min exposure to a solar simulator (Smith, 1988) and over the infraredstimulated luminescence signals of K-feldspar, which require much more prolonged light exposure to be fully bleached or that can be reduced only to an unbleachable residual (Chen et al., 2013; Kars et al., 2014). However, the comparatively rapid bleaching of zircon OSL starkly contrasts with the photoionisation cross-section values determined from laserstimulated decay curves. Comparing published values for quartz with the values obtained in this study for zircon suggests the presence of signals with decay rates similar to the s1 and s3 components in quartz (Singarayer and Bailey, 2003); at least the s3 component would not reduce to zero following 300 s of daylight exposure. Most probably, this mismatch relates to the stimulation power density of the green laser used in our experiments, which might vary both over time and spatially, depending on the exact position of the grain in the hole of the single-grain disc, and that might deviate from its nominal value used to calculate the photoionisation cross-section. Consequently, a systematically wrong stimulation power density would lead to invalid photoionisation cross-section values.

4.2.2 Signal intensities

A second requirement for OSL auto-regeneration dating of zircons is a sufficiently high OSL sensitivity to induce wellquantifiable signals – i.e. signals significantly above the instrumental background – within a reasonable amount of time, say 1–2 a at maximum. In this regard, our results are somewhat disappointing, at least for sample ZR229, which exhibited no noticeable OSL signal increase after 1.5 a of storage, corresponding to a self-dose of about 0.17 Gy based on the internal dose rate derived in Sect. 3.2. This finding applies to "pure" single grains and multiple grains stored together and measured simultaneously, although there seems to exist a tendency of large grains to produce stronger OSL than small grains (Fig. 6). Furthermore, it is consistent with observations of the low OSL sensitivity of zircon samples Smith (1988) studied. Thus, an increase in optical stimulation power density of 1–4 orders of magnitude compared to experimental setups used previously (Smith, 1988, 1989) cannot sufficiently improve the signal-to-background ratio to render zircon OSL auto-regeneration practical. The low OSL signal yield might be explained by the fact that OSL selectively detects only a fraction of all emitted signals (those with energies higher than the stimulation energy), while TL allows the exploitation of all emissions in the energy range for which the PMT is sensitive, thus maximising the TL yield. Smith (1989) argues that only a tiny portion of electrons evicted from their traps by light exposure recombine and produce OSL, while the majority of charge is re-trapped in shallow traps, giving rise to photo-transferred TL upon heating. Assuming an inherent similarity in TL and OSL emission spectra of zircons, our experimental data (Fig. 4) indicate that green laser stimulation, along with a detection window in the UV range, certainly captures only a small fraction of all photons emitted upon charge recombination. Moreover, for autoregeneration measurements, care should be taken to physically separate individual zircon grains, e.g. in the holes of single-grain discs. The regenerated signals of multiple grains in a hole, as compared to "pure" single-grain measurements, are probably affected by cross-irradiation of grains during storage. Vaz and Senftle (1971) have shown that zones with high radioelement contents in zircon grains yield low TL signals and vice versa, an observation supported by our data (see Sect. 3.1 and 3.2). On a grain-to-grain level, this anticorrelation would induce comparatively high luminescence signals in grains with low U and Th contents that are irradiated by neighbouring grains with higher activity, thus boosting the overall auto-regenerated luminescence output of a cluster of grains stored next to each other as compared to single zircon grains left for auto-regeneration in isolation from other grains. Although we found individual zircon grains are sometimes tricky to handle, future studies should thus follow the recommendation by Smith (1988) to store zircon grains separately for auto-regeneration.

4.2.3 Practical considerations

Given the difficulties of auto-regenerating zircon OSL signals within practical storage times, several strategies to further pursue this method are conceivable:

1. Apart from newly developed high-sensitivity luminescence readers, time-resolved OSL (TR-OSL) could improve the signal-to-noise ratio because it separates in time the stimulation and detection of OSL and thus potentially enhances the signal-to-noise ratio (Chithambo,

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2007; Bulur et al., 2014). While previous studies demonstrated efficient signal stimulation at 445 nm (Bulur et al., 2014), main OSL emissions are likely to be located at 350–500 nm and at 620 nm for the samples studied here due to shared recombination centres in TL and OSL (Kristianpoller et al., 2006).

- 2. Increasing the OSL measurement temperature (e.g. to 150 °C) can potentially improve the signal-to-noise ratio through thermal eviction of electrons from shallow "photo-transfer" traps (Smith, 1988). However, Smith (1989) noted that, at higher temperatures, a harder-to-bleach OSL component is preferentially sampled, while, in contrast, at lower temperatures, an easyto-bleach component dominates the signal. This might have repercussions on dating poorly bleached samples.
- 3. It was shown that heating a zircon sample to temperatures $> 500 °C$ enhances its TL sensitivity by factors of 3–20 and OSL sensitivity by up to a factor of 200 (depending on heating conditions and sample properties), thereby annealing previously accumulated radiation damage and increasing crystallinity (Jain, 1978; Amin et al., 1983; Amin and Durrani, 1985; Godfrey-Smith et al., 1989). In particular, geologically old zircons in a metamict state show low luminescence sensitivity that might, however, be (partially) restored by annealing. Therefore, Smith (1989) proposed a dating protocol in which the natural luminescence signal is measured first, followed by annealing the sample before auto-regeneration. The enhanced luminescence sensitivity is supposed to reduce auto-regeneration storage periods substantially, possibly to a few months, for most zircon samples. This is compatible with findings by Templer and Smith (1988) and Smith (1989), who state that most heated zircons yield measurable autoregenerated TL within 1 year and others even within periods as short as 2 months. One potential caveat, however, is that heating changes the opacity of zircons, leading to apparent sensitivity changes through internal light absorption effects which, according to Templer (1985b, a), amount to only $< 10\%$. Further experiments on heatinduced sensitisation of zircon OSL are necessary to test the feasibility of this approach for optical resetting events.
- 4. Since we detected a significant auto-regenerated TL signal for one of the studied samples after 3 weeks of storage, combining zircon OSL and TL signals in an autoregeneration approach might offer some pivotal advantages (Smith, 1988). Such an auto-regeneration protocol benefits from rapid bleaching of the natural zircon OSL, minimising the risk of age overestimation due to inherited signal contributions while shortening storage periods by using the more sensitive TL signal for autoregeneration. An annealing step (e.g. at 500 °C) after

reading the natural OSL erases any potential residual signals before storage and increases sample sensitivity. This changing luminescence sensitivity and using two different stimulation modes (OSL and TL) in the protocol necessitates introducing a test dose for normalising OSL and TL signals, respectively. However, zircon luminescence signals induced by laboratory irradiation suffer from anomalous fading (Wintle, 1973; Sutton and Zimmerman, 1976; Templer, 1985b) and exhibit unstable signal components so that Smith (1988) suggests a thermal pretreatment at 125° C for 2d to circumvent these problems and produce a laboratory signal most similar to the natural signal, as judged from the similarity of their TL glow curves.

5. As the efficiency of zircons to produce OSL (using the experimental parameters in this and most previous studies) is much smaller than their TL efficiency, photo-transferred TL (PTTL; Zimmerman, 1979) was proposed as a technique to measure small doses with greater efficiency and avoid anomalous fading (Bailiff et al., 1977) while at the same time benefitting from the better bleachability of the OSL signal (Smith, 1988). PTTL could be used in conjunction with the combined OSL/TL approach and allows a much larger range of stimulation wavelengths, including those shorter than the 532 nm light used here, that are more efficient in completely erasing the OSL signal before regeneration.

5 Conclusions

Using zircon as a luminescence geochronometer offers distinct advantages over other minerals. Its high internal dose rate facilitates the use of auto-regenerated signals and significantly reduces the impact of the external dose rate and associated factors (e.g. gravimetric water content) on age. Here, we re-appraised the auto-regeneration approach, using up-to-date measurement technology, for its practical value by examining key features of the TL and OSL of two zircon samples with contrasting geochemical compositions. We summarise our findings as follows:

- 1. CL and LA-ICP-MS yielded highly contrasting radioelement contents between the two studied samples, spanning more than 1 order of magnitude. In most zircons, U and Th are non-uniformly distributed in zones, with zones low in radioactivity producing brighter luminescence and vice versa.
- 2. The TL spectra of the two samples show as common features a broad band at ∼ 400 nm and a red emission at ∼ 640–650 nm, while the geologically young sample with higher internal α -activity exhibits additional narrowband emissions at ∼ 480 and ∼ 580 nm, usually attributed to rare earth ions acting as charge traps in zircon.
- 3. OSL sensitivity and decay rate vary substantially between grains of a zircon sample; however, both samples appear to share common signal components present in individual grains' signals in variable proportions.
- 4. Zircon OSL can be bleached by sunlight to negligible residuals within 300 s of exposure. Further studies, including opaque zircons, are pending.
- 5. While a storage period of 1.5 a was not long enough to auto-regenerate a zircon OSL signal (for the highactivity sample), a measurable auto-regenerated TL signal could be obtained within 21 d. Special emphasis needs to be placed on the accurate determination of background levels and their reproducibility.
- 6. A protocol combining the natural OSL signal with autoregenerated TL, following Smith (1988), appears to be the most promising one in rendering the methodology practical and fostering a more widespread use in Quaternary geochronology.
- 7. A future avenue is certainly to constrain the lower and upper dating limits of the zircon auto-regeneration technique. The upper dating limit varies significantly between studies, with estimates ranging from 10 ka (Faïn et al., 1988) to 400 ka or more (Vaz and Senftle, 1971; Templer, 1986; Templer and Smith, 1988). Reconciling a linear α -dose range of 20 kGy (Sutton and Zimmerman, 1976) with the dose rates calculated from LA-ICP-MS data of the two samples studied here (see Sect. 3.2) gives an upper dating limit of between ∼ 100 ka and \sim 2 Ma. Van Es et al. (2002a) propose the method could date samples as young as 12 months.

Code and data availability. All raw and partially processed data and R codes are available at https://doi.org[/10.48657/705a-g067](https://doi.org/10.48657/705a-g067) (Schmidt, 2024).

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Author contributions. CS conceived the study; carried out OSL, TL, and auto-regeneration experiments; analysed and visualised the data; acquired funding; provided supervision to TH; prepared the first draft of the article; and organised the writing process with contributions from all authors. TH carried out the bleaching experiments as part of his bachelor's thesis. PRH developed zircon purification procedures, validated the data, and contributed to writing and editing. AU planned and carried out LA-ICP-MS analyses and helped with data processing and interpretation. BP assisted with refining zircon separation procedures and contributed to editing. GEK provided supervision to TH, validated the data, and contributed to writing and editing. SK validated the data, acquired funding, and contributed to writing and editing.

Competing interests. At least one of the (co-)authors is a member of the editorial board of *Geochronology*. The peer-review process was guided by an independent editor, and the authors also have no other competing interests to declare.

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