

Establishing a protocol for the selection of zircon inclusions in garnet for Raman thermobarometry

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ABSTRACT

The structural and chemical properties of zircon inclusions in garnet megablasts from the Dora Maira Massif (Western Alps, Italy) were characterized in detail using charge contrast imaging, Raman spectroscopy, and laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS). The aim of this work is to determine to what extent the degree of metamictization, metamorphic recrystallization, inherent structural heterogeneity, chemical composition, and zoning, along with the elastic stress imposed by the host mineral, can influence the Raman peak position of the zircon inclusion and hence, the residual pressure estimated via Raman geo-thermobarometry. We show and confirm that metamictization and inherent structural heterogeneity have a major influence in the Raman spectra of zircon in terms of peak position and peak width. We suggest that, for spectral resolution of 2 cm^{-1} , the peak width of the B_{1g} mode near 1008 cm^{-1} of reliable grains must be smaller than 5 cm^{-1} . The method can be applied to both inherited igneous and newly formed Alpine metamorphic crystals. By coupling structural and chemical information, we demonstrate that there are no significant differences between the Raman spectra of zircon with oscillatory-zoned texture, formed during magmatic crystallization, and those formed by fluid-induced Alpine (re)crystallization. The discrimination between magmatic and metamorphic zircon based only on micro-textural constraints is not robust. Finally, our results allow establishing a protocol devoted to the selection of reliable buried zircon inclusions, relying only on Raman spectroscopic measurements, to use for elastic thermobarometry applications.

Keywords: Dora Maira Massif, elastic thermobarometry, zircon, metamictization


INTRODUCTION

Zircon is one of the most common accessory minerals in various igneous, sedimentary, and metamorphic rocks. Furthermore, due to its large stability field and its physical robustness, zircon often hosts ultrahigh pressure (UHP) metamorphic minerals such as coesite and diamond (Parkinson and Katayama 1999; Ye et al. 2000b; Hermann et al. 2001; Rubatto and Hermann 2007; Katayama and Maruyama 2009). Zircon can be used for a wide range of applications, including U-Pb geochronology, and potential use as a host phase for the disposal of excess weapons-grade Pu because of its capacity to incorporate radioactive elements, (e.g., Ewing et al. 1995). Further, radiation-damaged zircon has been successfully applied as a model system to elucidate the behavior of partially ordered structures at high pressure and temperature conditions (e.g., Colombo et al. 1999; Binvignat et al. 2018).

A recent development in zircon petrology is its exploitation as a mineral inclusion in the frame of Raman elastic thermobarometry (Campomenosi et al. 2018; Stangarone et al. 2019; Zhong et al. 2019). The main idea is that the geochemical and isotopic information that can be obtained from zircon is complemented by elastic thermobarometry of zircon and its host to retrieve *P-T-time-fluid-deformation* paths of metamorphic and igneous

rocks. For this new application, however, zircon composition and structural metamictization due to the decay of radioactive elements such as U and Th within its crystal structure has to be taken into account, because the radiation-induced structural changes can lead to large variations in the Raman spectrum. Indeed, the Raman peaks of radiation-damaged crystals shift toward lower wavenumbers, while the peak width increases with respect to pristine crystals taken as reference (Colombo et al. 1999; Ríos et al. 2000; Nasdala et al. 2001; Geisler and Pidgeon 2002; Binvignat et al. 2018). In this regard, useful diagrams such as peak width (Γ), in terms of *full-width at half maximum*, vs. peak position of the major Raman peak near 1008 cm^{-1} (B_{1g} phonon mode related to antisymmetric SiO_4 stretching) can be exploited to recognize the zircon crystals whose Raman spectra is unaffected by metamictization processes and that can be safely used for elastic thermobarometry purposes (e.g., Zhong et al. 2019). However, even though the degree of metamictization is the main factor affecting the Raman spectra of zircon, it is not the only one. Cathodoluminescence (CL) studies of zircon internal zoning (e.g., Rubatto and Gebauer 2000; Corfu et al. 2003) reveal that single crystals of zircon, particularly in metamorphic rocks, commonly show a complex internal chemical and structural heterogeneity at the micrometer-scale. Furthermore, variations in the Raman peak positions can be caused also by chemical substitution: a typical example is Hf replacing Zr at the dodeca-

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hedral crystallographic site (e.g., Hoskin and Rodgers 1996).

To address these problems, we have chosen a particularly well-known sample suite from the Dora Maira UHP unit (Western Alps, Italy), where garnet megablasts host abundant zircon inclusions. We report micrometer-scale structural and chemical information of partially exposed zircon crystals studied by complementary Raman spectroscopy, charge contrast (CC) imaging, and laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS). The purpose is (1) to determine quantitatively how the structural and chemical heterogeneities of zircons can affect their Raman spectrum and (2) to propose a best-practice protocol for the selection of zircon inclusions suitable for elastic thermobarometric estimations.

SAMPLE DESCRIPTION

We selected exposed and buried zircon inclusions in garnet megablasts from the UHP whiteschists of the Dora Maira Massif. The investigated samples come from the two major outcrops of the Brossasco-Isasca Unit (e.g., Chopin 1984): Vallone Gilba and Vallone Martiniana close to the Case Fapina and Case Parigi localities, respectively. Several polished sections of about 350 μm in thickness have been prepared from three different garnet megablasts (up to 15–20 cm in size). Zircon crystals are included in garnet and kyanite and range in size from ~ 10 to 200 μm ; they are either single, isolated crystals or clusters of two or more inclusions. The shape of the inclusions varies from rounded to elongated and almost idiomorphic with sharp corners and edges. Some of the zircon crystals also host mineral inclusions such as rutile, coesite, sheet silicates, and apatite, in agreement with previous observations (e.g., Gautiez-Putallaz et al. 2016).

Previous results (Gebauer et al. 1997; Gautiez-Putallaz et al. 2016) suggested the existence of two main zircon generations that can coexist within the same crystal in these rocks. The first one records the crystallization of the Permian protolith (~ 275 Ma), while the second domain is Eocene in age (35.1 ± 0.8 Ma) and related to metamorphism during the Alpine subduction. Chopin (1984) was the first to constrain this metamorphic event at UHP conditions by finding coesite inclusions within garnet. Optically single crystals of coesite inclusions have been found also in this study, especially within the garnet neoblasts (SiO_2 saturated whiteschists). Within the garnet megablasts, rare coesite inclusions have been found only at the external rim. More recent studies have constrained the metamorphic peak conditions in the diamond stability field at about 4–4.3 GPa and 730–750 $^\circ\text{C}$ (e.g., Hermann 2003; Ferrando et al. 2009; Gautiez-Putallaz et al. 2016).

The selection of completely buried zircon inclusions follows the protocol given by Campomenosi et al. (2018) for elastic thermobarometry applications (i.e., small, similar in size, non-fractured, and isolated inclusions within the center of a non-fractured host). On the other hand, the selection of the exposed grains (about 60% of the total number of zircon inclusions present in the samples) was based on criteria such as variable size of the inclusion and of the surrounding intact host (the *effective* host), exposure degree of the inclusions (optically estimated in terms of the ratio between the exposed and buried inclusion surface) and presence of fractures or any other kind of discontinuity in both the host and the inclusion.

ANALYTICAL METHODS

Exposed zircon inclusions were investigated by CC imaging first and Raman spectroscopy later to obtain textural and structural information, and finally by LA-ICP-MS for the chemical composition.

CC images were acquired with a ZEISS EV050 scanning electron microscope at the Institute of Geological Sciences, University of Bern, at low vacuum conditions (18 Pa), 12 kV, a beam current of 100 mA and a working distance of 9.5 mm. CC images were obtained from well-polished, uncoated samples. It has been demonstrated that CC images correlate exactly to cathodoluminescence images and result from the complex interaction between the electron beam, the positive ions generated by electron-gas interactions in the chamber, a biased detector, and the sample (Griffin 2000; Watt et al. 2000). Internal check in the Bern laboratory confirmed that CC images are identical to panchromatic cathodoluminescence images.

Raman measurements were collected with two different Horiba Jobin-Yvon spectrometers: a T64000 triple-monochromator system operating in a subtractive mode and an Explora_Plus single-monochromator spectrometer at the University of Hamburg and Genova, respectively. The first one was equipped with a symphony LN₂-cooled CCD detector, 1800 gr/mm holographic gratings, an Olympus BH-41 optical microscope and a Coherent Ar⁺ laser, whereas the second with a Peltier-cooled CCD detector, a 2400 gr/mm grating, an Olympus BH-41 optical microscope, and Nd:YAG solid state laser. In both cases, using a slit width of 100 μm , the spectral resolution was approximately 2 cm^{-1} (determined using the photoluminescence line of Sm: SrB₄O₇ for the T64000 and a Ne lamp for the Explora_Plus spectrometers), and therefore apparatus corrections of the peak widths were not required (Nasdala et al. 2001). Sufficient acquisition times and a surface density of the laser power were used to avoid sample overheating during the Raman measurements (Zhong et al. 2019). The instrumental accuracy in the peak position determination was about 0.35 cm^{-1} for the T64000 spectrometer and about 0.55 cm^{-1} for the Explora_Plus. These values refer to the half of the spectral pixel-pixel distance, which results from several instrumental factors, including the CCD detector array, groove density of gratings, and excitation-light wavelength.

However, both the instruments were manually calibrated to the silicon Raman peak at 520.5 cm^{-1} at each measurement session, and the stability of the instrumental setup was double-checked measuring reference crystals every 4 h. The resulting standard deviation in the determination of the silicon peak along different days of measurements was approximately 0.04 cm^{-1} . Furthermore, a crosscheck of the two data set measured with the two different spectrometers has been performed on several zircon grains to verify their consistency. Finally, the lateral spatial resolution of the two instruments is approximately 1 μm and the probed volume at the sample surface is of about 1 μm^3 .

The Origin Lab-Pro 2018 software package was used for data evaluation. The collected spectra were baseline corrected for the continuum luminescence background when necessary, temperature-reduced to account for the Bose-Einstein occupation factor (Kuzmany 2009), and normalized to the acquisition time (7 s). Peak positions, FWHMs, and integrated intensities were determined from fits with pseudo-Voigt functions [PV = (1 - q) Lorentz + q Gauss, q is the weight coefficient]. The criterion for the maximum number of fitted peaks was $\Delta I < I/2$, where I and ΔI are the calculated magnitude and uncertainty of each peak intensity, respectively.

Trace element analyses were obtained with an ASI-RESOLUTION 193 nm laser system coupled to an Agilent 7900 quadrupole Inductively Coupled Plasma mass spectrometer (ICPMS) at the Institute of Geological Sciences, University of Bern. The laser was tuned to a repetition rate of 5 Hz and an energy output of 4 mJ (corresponding to an HV of about 26–27 kV). The ICPMS was tuned for maximum sensitivity and minimum production of molecular species, maintaining ThO⁺/Th⁺ at <0.2%. The spot size on the zircon crystal was 20 μm . Analyses were standardized to glass NIST 612, and zircon 91500 was run as a secondary standard to monitor accuracy. Stoichiometric Si was employed as an internal standard for zircon (SiO₂: 31.6 wt%). Reproducibility and accuracy were within 10% or less across all analyzed elements. The data were reduced with the free-ware Iolite (Paton et al. 2011) and its data reduction scheme for trace elements (Woodhead et al. 2007).

RESULTS

Partially exposed zircon inclusions

Charge contrast imaging. CC imaging of 64 inclusions from samples DM17-13 and DM17-49 (Martiniana locality) and 83 grains from samples DMG4-6 and DM17-35 (Gilba

locality) shows large variability in zircon internal texture. By combining the data on various crystals, it was possible to recognize at least four major domains (Fig. 1): (1) a *dark* (i.e., low CC-emission) domain, commonly corresponding to the crystal cores, (2) an *oscillatory-zoned* domain (medium-high CC emission), which usually corresponds to entire crystals or the core of elongated crystals and are further subdivided into *oscillatory-dark* and *oscillatory-bright* domains based on CC emission, (3) a transition (or *undefined*) domain, whose appearance is between the darker and a brighter domains and without a defined internal texture, and (4) a *bright* domain (i.e., high CC-emission), which usually belongs to the external rim of the crystals. About 70% of zircon inclusions from the

Gilba locality show a more elongated shape with a typical oscillatory-zoned core surrounded by 1 or 2 thin brighter rims. On the other hand, about 80% zircon inclusions from the Martiniana locality present more sub-idiomorphic shapes and more homogeneous internal texture. Furthermore, as already reported by Gautiez-Putallaz et al. (2016) for zircon hosted in Si undersaturated whiteschists, some of the analyzed inclusions in Martiniana, where garnet is more abundant, display oscillatory dark domains.

Raman spectroscopy. Zircon has tetragonal symmetry with space group $I4_1/amd$. According to group-theory analysis, the optical phonons at the Brillouin-zone center of zircon are (Kroumova et al. 2010):

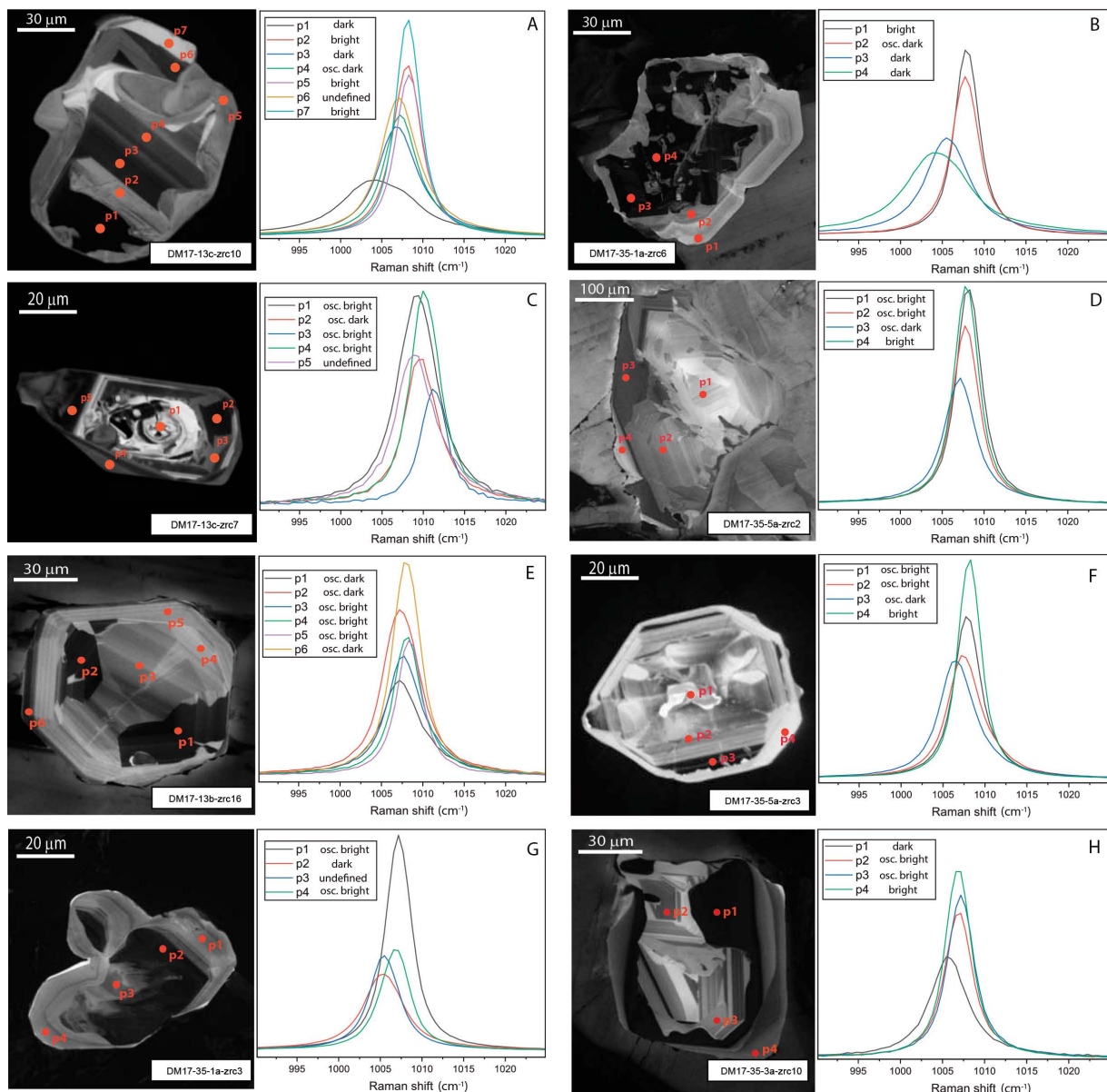


FIGURE 1. Relationship between the CC emission domain and the corresponding Raman spectra (B_{1g} mode near 1008 cm^{-1}) in partially exposed zircon inclusions. Samples DM17-35 are from Gilba locality while sample DM17-13 comes from the Martiniana locality.

$$\Gamma_{\text{opt}} = 2A_{1g} + A_{1u} + A_{2g} + 3A_{2u} + 4B_{1g} + B_{1u} + B_{2g} + 2B_{2u} + 5E_g + 4E_u$$

The A_{1g} , B_{1g} , B_{2g} , and E_g modes are Raman-active and therefore a total of 12 Raman peaks can be observed in the spectrum of a randomly oriented zircon. According to previous studies (Knittle and Williams 1993; Nasdala et al. 2001; Binvignat et al. 2018), both pressure and metamictization affect most strongly the B_{1g} mode near 1008 cm^{-1} , originating from anti-symmetrical SiO_4 stretching. For this reason, our discussion is mainly focused on this phonon mode.

The Raman spectroscopic analysis is based on 250 spectra collected from 60 zircon inclusions in different CC domains. As an example, Figure 1 shows the variation of the internal SiO_4 antisymmetric stretching mode (B_{1g}) near to 1008 cm^{-1} of representative zircon inclusions exhibiting heterogeneous CC emission. Generally, dark CC domains correspond to Raman peak broadening and shift toward lower wavenumbers when compared to those collected across brighter or oscillatory-zoned domains. Furthermore, transition CC domains show transitional average features also for the Raman spectra.

A more quantitative evaluation of the structural state of zircon crystals can be done plotting the relationship between the phonon wavenumber ω and the corresponding peak width Γ (Geisler and Pidgeon 2002; Zhong et al. 2019). The data measured on partially exposed zircon inclusions (Fig. 2) show two different trends that correlate with the four different zircon domains (*bright*, *oscillatory-zoned*, *undefined*, and *dark*).

The first trend is characterized by $\omega < 1008 \text{ cm}^{-1}$, $4.5 \text{ cm}^{-1} \leq \Gamma < 12 \text{ cm}^{-1}$, and $\omega(\Gamma)$ with a negative slope (inverse correlation). The data points on this trend mostly correspond to zircon domains with dark and oscillatory-dark CC emission.

The second trend, at higher wavenumbers, is characterized by $1008 \text{ cm}^{-1} < \omega < 1011 \text{ cm}^{-1}$, $3.5 \text{ cm}^{-1} < \Gamma < 5.5 \text{ cm}^{-1}$, and $\omega(\Gamma) \approx \text{constant}$. This trend is mostly defined by zircon domains with *bright* and *oscillatory(-bright)* CC emission.

Note that, on average, *oscillatory-dark* domains display the largest Γ and lowest wavenumber with respect to *oscillatory-bright* domains. On the other hand, *undefined* domains are difficult to classify since they scatter over the entire data.

LA-ICP-MS. The different types of zircon domains also show differences in trace-element composition. Tables 1a–1b report representative major, minor, and trace elements composition of analyzed zircon crystal inclusions as determined

by LA-ICP-MS with the associated CC-emission domains. Hf concentration is usually around $11\,000 \mu\text{g/g}$ and only few analyses show higher values up to $14\,000 \mu\text{g/g}$. U content varies significantly between oscillatory-bright zoned, oscillatory-dark, and dark CC domains, with the last showing, usually, the highest concentration (see for example DM17-35-3a-zrc10-p1 and p2 or even DM17-35-3a-zrc14-p3 and p2 in Table 1a). U contents vary from a few hundred $\mu\text{g/g}$ in the bright domains to $5000 \mu\text{g/g}$ for the darker domains and generally show an inverse relationship with CC emission, indicating that U suppresses luminescence (e.g., Rubatto and Gebauer 2000).

In line with previous REE data sets (e.g., Gautiez-Putallaz et al. 2016), zircon cores with oscillatory or convolute zoning have a steep HREE-enriched pattern with a pronounced negative Eu anomaly (e.g., Fig. 3c p1). This REE pattern is characteristic of conditions where plagioclase was present and garnet was absent, i.e., during the crystallization of the granitic protholith in Permian times (e.g., Gautiez-Putallaz et al. 2016).

The rims and many crystals with oscillatory zoning have lower REE contents and a flat HREE pattern with no or weak Eu-anomaly (Fig. 3). This REE pattern is diagnostic for zircons formed during prograde to peak metamorphism, where feldspar is no longer stable (e.g., Gautiez-Putallaz et al. 2016). The variable slope of the HREE patterns ranging from slightly positive to even negative (Fig. 3c) indicates zircon growth together with garnet, as both minerals compete for the HREE. Therefore a first distinction between pre-Alpine inherited igneous zircon and Alpine metamorphic zircon can be obtained from the trace element analysis. Notably, several large crystals that display a fine oscillatory to sector zoning more or less surrounded by a dark or light rim present a depletion in HREE across the entire grain (e.g., Figs. 3d and 3e). This demonstrates that oscillatory zoning occurs also in Alpine metamorphic zircon and is not restricted to inherited igneous zircon, demonstrating that both CC and trace elements are required for the correct classification of the zircon.

Previous analyses on the same locality demonstrated that inherited zircon cores are ca. 275 Ma old and metamorphic rims are $35.1 \pm 0.8 \text{ Ma}$ old (Gebauer et al. 1997; Gauthiez-Putallaz et al. 2016). In our case, a rough estimate of the age can also be obtained from the LA-ICP-MS data, even if proper age standardization and common Pb correction have not been performed. The $^{206}\text{Pb}/^{238}\text{U}$ values estimated in such a way are in general agreement with those obtained before on zircon from the same locality and correlate with the distinction made from the REE patterns (Gauthiez-Putallaz et al. 2016). This rough age estimate allows distinguishing between Alpine domains (usually lower U content and bright or oscillatory-bright CC domains) and pre-Alpine domains (higher U content and dark or oscillatory-zoned CC domains at the zircon cores) especially in cases where REE patterns might be ambiguous (see Table 2). Notably the large crystals with oscillatory-sector zoning and low HREE content are of Alpine age.

Assuming an age of 275 Ma for the pre-Alpine and 35 Ma for the Alpine domains (Gebauer et al. 1997; Gauthiez-Putallaz et al. 2016) and based on the measured U-Th composition of the zircon domains, their accumulation radiation doses D_a was calculated as follows (e.g., Nasdala et al. 2001):

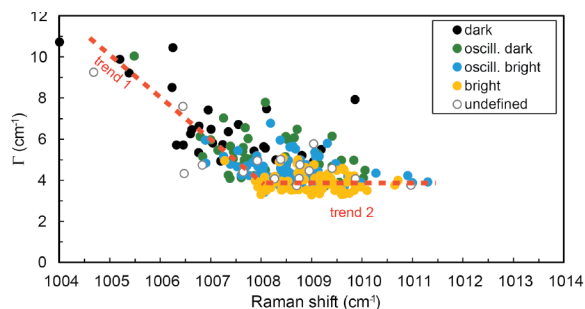


FIGURE 2. Γ vs. Raman shift of the antisymmetric SiO_4 stretching B_{1g} mode near 1008 cm^{-1} of partially exposed crystals with highlighted CC-emission domains.

TABLE 1a. Major, minor, and trace elements (in $\mu\text{g/g}$) of zircon inclusions (spots are referred to Fig. 3)

Sample name	CC domain	Ti	Y	Nb	Hf	Pb	Th	U
DM17-35-3A-zrc8-p1	osc. dark	2.28	1639	3.16	10300	154	158	1436
DM17-35-3A-zrc8-p2	osc. bright	4.70	2651	8.73	7710	390	793	1241
DM17-35-3A-zrc10-p1	osc. bright	35.3	492	34.2	10880	10.0	84.9	378
DM17-35-3A-zrc10-p2	dark	5.03	197	5.98	13690	52.4	84.9	378
DM17-35-3A-zrc10-p3	bright	7.90	183	4.01	10840	24.3	6.85	883
DM17-35-3A-zrc14-p1	osc. bright	2.91	850	1.94	12450	138	40.7	730
DM17-35-3A-zrc14-p2	osc. dark	6.90	3140	8.05	10300	326	696	1900
DM17-35-3A-zrc14-p3	osc. bright	4.90	1820	4.50	11660	237	158	1320
DM17-35-3A-zrc14-p4	osc. bright	2.67	1353	1.42	8680	66.1	73.4	333
DM17-35-3A-zrc18-p1	osc. bright	6.85	1850	1.64	9100	308	179.6	474
DM17-35-3A-zrc13-p2	osc. bright	1.69	920	3.05	9570	130	89.0	673
DM17-35-3A-zrc13-p1	osc. bright	1.79	1051	2.80	10960	154	87.0	828
DM17-35-3A-zrc13-p3	bright	3.04	1150	4.26	7070	242	336	635
DM17-35-5A-zrc2-p1	osc. bright	10.3	211	5.86	11770	9.65	22.4	367
DM17-35-5A-zrc2-p2	osc. bright	8.81	83.3	9.95	11190	17.6	34.3	695
DM17-35-5A-zrc2-p3	osc. dark	8.30	83.4	8.75	11270	31.4	34.9	1208
DM17-35-5A-zrc2-p4	osc. dark	4.60	58.5	3.10	11030	32.3	11.3	1148
DM17-35-5A-zrc2-p5	bright	5.10	198	2.85	11170	28.6	7.17	1092
DM17-35-5A-zrc3-p1	osc. bright	2.02	930	0.76	9950	51.5	44.1	253
DM17-35-5A-zrc14-p1	osc. bright	48.3	1682	26.8	9768	8.94	87.1	380
DM17-35-5A-zrc14-p2	dark	10.1	490	5.49	11520	16.4	15.7	530
DM17-35-5A-zrc14-p3	bright	10.8	430	6.06	10840	3.80	12.3	112
DM17-35-5A-zrc14-p4	bright	22.5	632	20.1	11400	8.96	32.7	370
DM17-35-5A-zrc14-p5	osc. dark	7.10	262	6.38	12790	10.5	8.84	407
DM17-35-1a-zrc6-p2	osc. dark	14.3	720	8.31	14570	65.8	41.1	3147
DM17-35-1a-zrc6-p3	dark	20.2	826	7.62	12610	77.1	32.8	2949
DM17-35-1a-zrc3-p1	osc. bright	13.3	204	1.63	12930	19.7	9.89	452
DM17-35-1a-zrc3-p2	dark	9.90	103.3	2.98	11690	73.7	7.82	3154
DM17-35-1a-zrc3-p3	osc. bright	13.1	161.8	3.09	12280	80.7	12.1	3700
DM17-35-1a-zrc9-p1	osc. bright	6.38	179.3	2.47	12190	43.7	16.8	1640
DM17-35-1a-zrc10-p1	osc. bright	10.8	587	2.42	11220	9.10	10.7	249.8
DM17-35-1a-zrc10-p2	osc. bright	7.90	536	8.53	10980	8.71	13.0	261
DM17-13-b-zrc13-p1	osc. bright	17.0	1479	21.6	11960	31.1	51.5	1307
DM17-13-b-zrc13-p2	osc. bright	11.8	1209	11.5	10820	22.5	26.6	755
DM17-13-b-zrc15-p2	osc. bright	6.30	649	4.80	10180	16.9	11.5	604
DM17-13-b-zrc15-p3	osc. bright	6.10	589	5.86	10310	9.37	7.11	346
DM17-13-b-zrc16-p1	osc. bright	16.8	1817	13.7	10840	34.7	35.2	1342
DM17-13-b-zrc16-p2	osc. bright	8.50	383	3.96	11030	16.8	14.9	562
DM17-13-b-zrc16-p3	osc. dark	20.9	1504	29.9	10550	43.3	56.8	1538
DM17-13-b-zrc16-p4	osc. bright	5.66	227.8	4.54	11550	10.3	18.5	435.3
DM17-13-c-zrc7-p1	osc. bright	16.5	2140	2.44	10600	125	149	719
DM17-13-c-zrc7-p2	osc. dark	10.2	1790	7.65	11900	21.1	13.2	950
DM17-13-c-zrc7-p3	undefined	10.1	1982	12.5	12280	25.4	17.8	1037
DM17-13-c-zrc4-p1	bright	27.0	3250	24.3	12380	32.4	35.2	1335
DM17-13-c-zrc4-p2	undefined	11.8	2100	13.6	11850	56.1	39.8	931
DM17-13-c-zrc10-p1	dark	20.9	2680	33.7	11080	102	33.0	5060
DM17-13-c-zrc10-p2	bright	7.30	582	4.56	11860	28.0	7.63	1175
DM17-13-c-zrc10-p3	osc. dark	10.7	1540	8.24	11480	46.2	6.67	2004

Note: Precision plus accuracy of measurements is within 10%.

$$D_{\alpha} = \frac{0.9928N_A c_U n_{238}}{10^6 m_{238}} (e^{\lambda_{238} t} - 1) + \frac{0.0072N_A c_U n_{235}}{10^6 m_{235}} (e^{\lambda_{235} t} - 1) + \frac{N_A c_{Th} n_{232}}{10^6 m_{232}} (e^{\lambda_{232} t} - 1) \quad (1)$$

where λ_n and m_n are, respectively, the nuclear decay constant and mass of the corresponding isotope (^{238}U , ^{235}U , and ^{232}Th) (Steiger and Jaeger 1977), the coefficient n represents the number of α decays per nucleus ($n_{238} = 8$, $n_{235} = 7$, and $n_{232} = 6$) and N_A is the Avogadro's number c_U and c_{Th} are the measured concentrations of U and Th, respectively, and t the time. Note that Equation 1 presumes a U isotopic composition of 99.28% ^{238}U and 0.72% ^{235}U . The resulting D_{α} values are reported in Table 2.

Completely buried zircon inclusions

The Raman spectra collected on the completely buried zircon crystals have higher wavenumbers (from 1010 cm^{-1} up to 1013.5 cm^{-1}) with respect to the partially exposed grains, with a

Γ mostly between 3.5 and 5.5 cm^{-1} . Only a few analyses reach values up to 6.5 cm^{-1} (Fig. 4). The overlap between the two sets of inclusions is limited to about 10% of the analyses.

The data from the exposed and fully enclosed zircon inclusions can also be analyzed in terms of the wavenumber difference ($\Delta\omega$) between the B_{1g} mode near 1008 cm^{-1} and the A_{1g} mode near 440 cm^{-1} rather than the *absolute* wavenumber. In this way, since the data were collected along different sessions of measurements, we can avoid eventual effects due to instrumental drift. As shown previously by Knittle and Williams (1993) and more recently by Binvignat et al. (2018) both phonon modes show an increase in the phonon wavenumber as the hydrostatic pressure increases. However, as reported from the same authors, the A_{1g} mode shows a weaker pressure dependence with respect to the B_{1g} mode and, therefore, to an increase of the pressure acting on the crystal should correspond an increase in the value of $\Delta\omega$ (Fig. 5).

DISCUSSION

Effect of minor and trace elements on zircon Raman shift

A large number of non-formula elements can be incorporated in the zircon crystal structure, however, most of them are usually far below 1 wt% (e.g., Hoskin and Schaltegger 2003), with the notable exception of

Hf. Raman data measured on isomorphic series ZrSiO_4 – HfSiO_4 , suggest that even if as much as 25% of all Zr ions are replaced by Hf the frequency variation of the main peaks does not exceed 3 cm^{-1} (Hoskin and Rodgers 1996). Furthermore, Nasdala et al. (2002) suggested that positions and Γ of the main peaks in annealed, metamict gemstone-quality zircon, containing up to 6000 $\mu\text{g/g}$ of U and 16 300 $\mu\text{g/g}$ of Hf, deviate less than 1 cm^{-1} from the data of pure well-crystalline ZrSiO_4 . Therefore, since our exposed zircon inclusions contain impurities and trace elements below these values (maximum U content is 5060 $\mu\text{g/g}$ and maximum Hf content is 14 570 $\mu\text{g/g}$, see Table 1), the effect of chemical variations on zircon Raman spectra is negligible.

Zircon variation in HREE composition (i.e., depletion in the Alpine domains) has been previously interpreted as the result of growth zoning during metamorphism in a fractionating bulk composition where REE are largely incorporated in garnet (e.g., Gautiez-Putallaz et al. 2016). Indeed, the garnet host shows a similar REE pattern along a core-to-rim line profile (e.g.,

TABLE 1b. REE composition (in µg/g) of zircon inclusions (spots are referred to Fig. 3)

Sample name	CC domain	La	Ce	Pr	Nd	Sm	Eu	Gd	Tb	Dy	Ho	Er	Tm	Yb	Lu
DM17-35-3A-zrc8-p1	Osc. dark	0.04	5.90	0.21	2.65	5.07	0.15	28.9	10.5	143	54.9	273	60.7	553	112
DM17-35-3A-zrc8-p2	Osc. bright	0.46	18.7	0.40	6.43	12.1	0.26	65.8	21.8	267	91.0	411	79.6	653	128
DM17-35-3A-zrc10-p1	Osc. bright	b.d.l.	2.80	0.04	1.06	6.80	2.23	70.5	16.0	86.2	12.3	29.5	4.80	38.9	6.95
DM17-35-3A-zrc10-p2	dark	b.d.l.	0.59	b.d.l.	b.d.l.	0.51	0.11	4.78	2.01	20.1	4.93	15.8	2.73	22.2	3.98
DM17-35-3A-zrc10-p3	bright	b.d.l.	1.03	b.d.l.	0.14	0.54	0.14	4.54	1.83	19.6	4.77	17.0	2.67	19.3	2.80
DM17-35-3A-zrc14-p1	Osc. bright	b.d.l.	0.44	b.d.l.	0.17	1.06	0.19	10.05	4.41	63.0	26.4	143	35.7	351	73.0
DM17-35-3A-zrc14-p2	Osc. dark	12.0	37.0	3.10	14.6	11.5	0.55	61.0	22.7	291	104	497	101	839	156
DM17-35-3A-zrc14-p3	Osc. bright	0.49	3.40	0.21	1.19	3.14	0.19	22.0	9.90	142	57.1	305	71.7	670	135
DM17-35-3A-zrc14-p4	Osc. bright	0.04	1.54	0.04	1.13	3.51	0.15	20.8	8.45	115	44.7	219	46.1	408	79.5
DM17-35-3A-zrc18-p1	Osc. bright	1.12	5.21	0.85	8.20	10.7	0.97	51.6	16.9	192	62.1	269	53.6	454	84.8
DM17-35-3A-zrc13-p2	Osc. bright	b.d.l.	2.78	b.d.l.	0.29	1.49	0.12	9.77	4.76	73.3	30.2	161	37.3	350	71.9
DM17-35-3A-zrc13-p1	Osc. bright	b.d.l.	2.38	0.02	0.33	1.08	0.10	9.02	4.55	75.7	32.9	182	43.8	424	88.1
DM17-35-3A-zrc13-p3	bright	0.34	14.1	0.17	1.78	3.15	0.73	16.5	6.16	85.0	34.4	185	44.1	442	97.0
DM17-35-5A-zrc2-p1	Osc. bright	0.08	1.08	0.04	0.42	3.30	1.36	53.7	10.8	48.2	5.31	11.1	1.39	10.0	1.69
DM17-35-5A-zrc2-p2	Osc. bright	b.d.l.	0.75	b.d.l.	0.34	4.74	1.41	43.7	6.35	22.2	2.26	4.55	0.61	5.21	0.90
DM17-35-5A-zrc2-p3	Osc. dark	b.d.l.	0.99	b.d.l.	0.12	3.47	1.13	34.8	5.44	19.9	2.26	4.60	0.61	5.82	1.03
DM17-35-5A-zrc2-p4	Osc. dark	b.d.l.	0.57	b.d.l.	b.d.l.	0.91	0.37	13.3	2.64	12.1	1.45	3.48	0.61	4.62	0.88
DM17-35-5A-zrc2-p5	bright	b.d.l.	0.81	b.d.l.	b.d.l.	0.62	0.14	5.83	1.92	20.3	5.27	18.7	3.33	23.5	3.37
DM17-35-5A-zrc3-p1	Osc. bright	0.06	1.27	0.11	0.78	2.50	0.07	14.1	5.45	76.9	30.7	154	33.6	301	59.3
DM17-35-5A-zrc14-p1	Osc. bright	b.d.l.	3.06	0.02	1.10	10.1	3.72	164	46.6	288	47.8	117	16.7	118	19.3
DM17-35-5A-zrc14-p2	dark	0.40	2.11	0.19	1.35	2.51	0.86	31.0	9.74	70.3	12.9	32.5	4.76	31.7	5.42
DM17-35-5A-zrc14-p3	bright	b.d.l.	0.75	b.d.l.	0.27	2.44	0.65	31.7	9.81	68.9	11.6	30.1	4.66	34.8	6.21
DM17-35-5A-zrc14-p4	bright	b.d.l.	1.89	0.04	0.68	4.14	1.39	54.2	15.9	104	16.4	40.5	5.53	38.8	6.39
DM17-35-5A-zrc14-p5	Osc. dark	b.d.l.	0.72	b.d.l.	0.10	1.19	0.40	16.3	5.58	38.4	6.47	16.4	2.28	15.6	2.60
DM17-35-1a-zrc6-p2	Osc. dark	b.d.l.	1.42	0.02	0.66	6.17	1.75	64.7	18.2	118	18.4	43.4	5.31	30.5	4.17
DM17-35-1a-zrc6-p3	dark	0.08	2.06	0.07	1.08	7.30	1.84	72.4	20.2	131	21.2	49.2	6.34	36.5	5.05
DM17-35-1a-zrc3-p1	Osc. bright	0.06	1.83	0.06	0.86	1.35	0.27	8.60	2.45	21.8	5.43	19.8	3.36	24.6	3.52
DM17-35-1a-zrc3-p2	dark	b.d.l.	0.45	b.d.l.	0.13	0.74	0.23	7.75	2.34	15.4	2.60	7.27	1.19	8.46	1.46
DM17-35-1a-zrc3-p3	Osc. bright	b.d.l.	0.95	0.02	0.26	1.50	0.44	14.4	4.10	25.8	4.42	12.0	2.04	14.0	2.42
DM17-35-1a-zrc9-p1	Osc. bright	b.d.l.	0.53	b.d.l.	0.19	1.40	0.46	16.1	4.63	27.3	4.51	12.9	2.03	14.5	2.48
DM17-35-1a-zrc10-p1	Osc. bright	b.d.l.	0.56	b.d.l.	0.23	3.06	0.93	38.1	11.7	89.1	18.0	51.8	7.30	47.6	7.10
DM17-35-1a-zrc10-p2	Osc. bright	0.05	0.70	b.d.l.	0.20	2.45	0.91	36.0	10.4	81.6	16.8	47.4	6.71	42.7	6.59
DM17-13-b-zrc13-p1	Osc. bright	b.d.l.	1.14	0.01	0.44	7.78	3.36	110	32.1	219	42.4	115	16.5	96.9	14.3
DM17-13-b-zrc13-p2	Osc. bright	b.d.l.	0.64	b.d.l.	0.40	5.57	2.46	86.6	25.6	177	35.2	95.6	14.4	85.9	12.7
DM17-13-b-zrc15-p2	Osc. bright	b.d.l.	0.67	b.d.l.	0.12	2.26	1.08	39.2	11.2	90.3	18.9	58.5	9.16	62.8	10.9
DM17-13-b-zrc15-p3	Osc. bright	b.d.l.	0.44	b.d.l.	0.09	1.87	0.97	33.6	10.1	80.4	17.2	54.2	8.52	55.6	8.56
DM17-13-b-zrc16-p1	Osc. bright	b.d.l.	0.88	b.d.l.	0.58	9.34	4.24	147	40.6	288	54.3	149	21.4	133	19.4
DM17-13-b-zrc16-p2	Osc. bright	b.d.l.	0.41	b.d.l.	0.16	2.34	0.96	32.0	8.26	58.5	11.0	30.3	4.72	33.8	6.03
DM17-13-b-zrc16-p3	Osc. dark	b.d.l.	1.28	b.d.l.	0.34	7.11	3.05	116	33.2	233	44.0	116	16.2	99.6	13.2
DM17-13-b-zrc16-p4	Osc. bright	b.d.l.	0.35	b.d.l.	0.14	1.32	0.54	15.3	4.54	32.1	6.00	16.2	3.04	26.4	5.43
DM17-13-c-zrc7-p1	Osc. bright	0.84	3.40	0.31	2.32	5.24	0.18	31.2	13.0	175	69.7	334	71.6	647	122
DM17-13-c-zrc7-p2	Osc. dark	b.d.l.	0.57	b.d.l.	0.15	2.20	1.22	54.0	23.0	230	55.0	164	25.0	160	24.0
DM17-13-c-zrc7-p3	undefined	b.d.l.	0.73	b.d.l.	b.d.l.	3.23	1.36	54.2	24.5	250	63.5	197	30.1	210	32.9
DM17-13-c-zrc4-p1	bright	b.d.l.	1.64	b.d.l.	0.53	6.90	2.72	97.7	39.0	397	104	335	54.3	407	69.1
DM17-13-c-zrc4-p2	undefined	b.d.l.	1.21	b.d.l.	0.28	3.64	1.53	54.4	21.5	230	67.1	247	45.7	377	68.5
DM17-13-c-zrc10-p1	dark	b.d.l.	1.97	b.d.l.	0.47	6.10	2.69	90.3	33.8	310	84.0	326	64.0	536	94.8
DM17-13-c-zrc10-p2	bright	b.d.l.	0.35	b.d.l.	b.d.l.	0.95	0.44	18.9	7.34	69.3	19.0	74.4	14.6	126	23.0
DM17-13-c-zrc10-p3	Osc. dark	b.d.l.	0.62	b.d.l.	0.32	2.79	1.19	40.9	16.8	170	47.9	191	39.1	347	65.0

Note: Precision plus accuracy of measurements is within 10%.

Gautiez-Putallaz et al. 2016), indicating equilibrium conditions with the associated zircon grains. The qualitative $^{206}\text{Pb}/^{238}\text{U}$ measurements confirm that zircon crystals showing a flat HREE pattern throughout the entire grain are completely metamorphic in origin. In this regard, it is important to note that, although REE chemical zonation does not influence the main Raman scattering features, such as peak broadening and position, it can give rise to heterogeneous photoluminescence, and hence to different background levels of the Raman spectra collected from zones. Besides, depending on the excitation laser wavelength, additional photoluminescence peaks may be observed next to the fundamental Raman peaks of zircon.

Metamorphic vs. inherited zircon domains

Charge contrast or cathodoluminescence imaging of metamorphic and inherited zircons usually show notable differences in terms of the corresponding internal texture (e.g., Rubatto and Gebauer 2000; Corfu et al. 2003). Oscillatory-zoned grains have been chemically interpreted as the result of alternating deple-

tion and enrichment in trace elements (i.e., U and Y) during crystal growth at the crystal-melt interface. On the other hand, bright domains at the rim of the crystals, usually show higher and homogeneous CL-emission, presenting irregular shape that often overgrows the pre-existing crystals whose texture can sometime be evident as a relict (Rubatto and Gebauer 2000). Our results indicate that metamorphic zircon can also include dark, bright, oscillatory, and undefined CC domains. This observation warns against using internal zircon zoning alone for distinguishing magmatic vs. metamorphic zircon.

As portrayed in Figure 2, bright and oscillatory-bright domains (yellow and blue spots, respectively), have the same Raman spectral features and define the “non-metamict” domain in the diagram (see details below). Therefore, despite their magmatic or metamorphic origin, from a structural point of view, these two domains are effectively equivalent and there will be no difference in the calculated residual stress even if the two domains coexist within the same crystal.

Finally, if we plot our data in a Γ of the B_{1g} mode near 1008 cm^{-1}

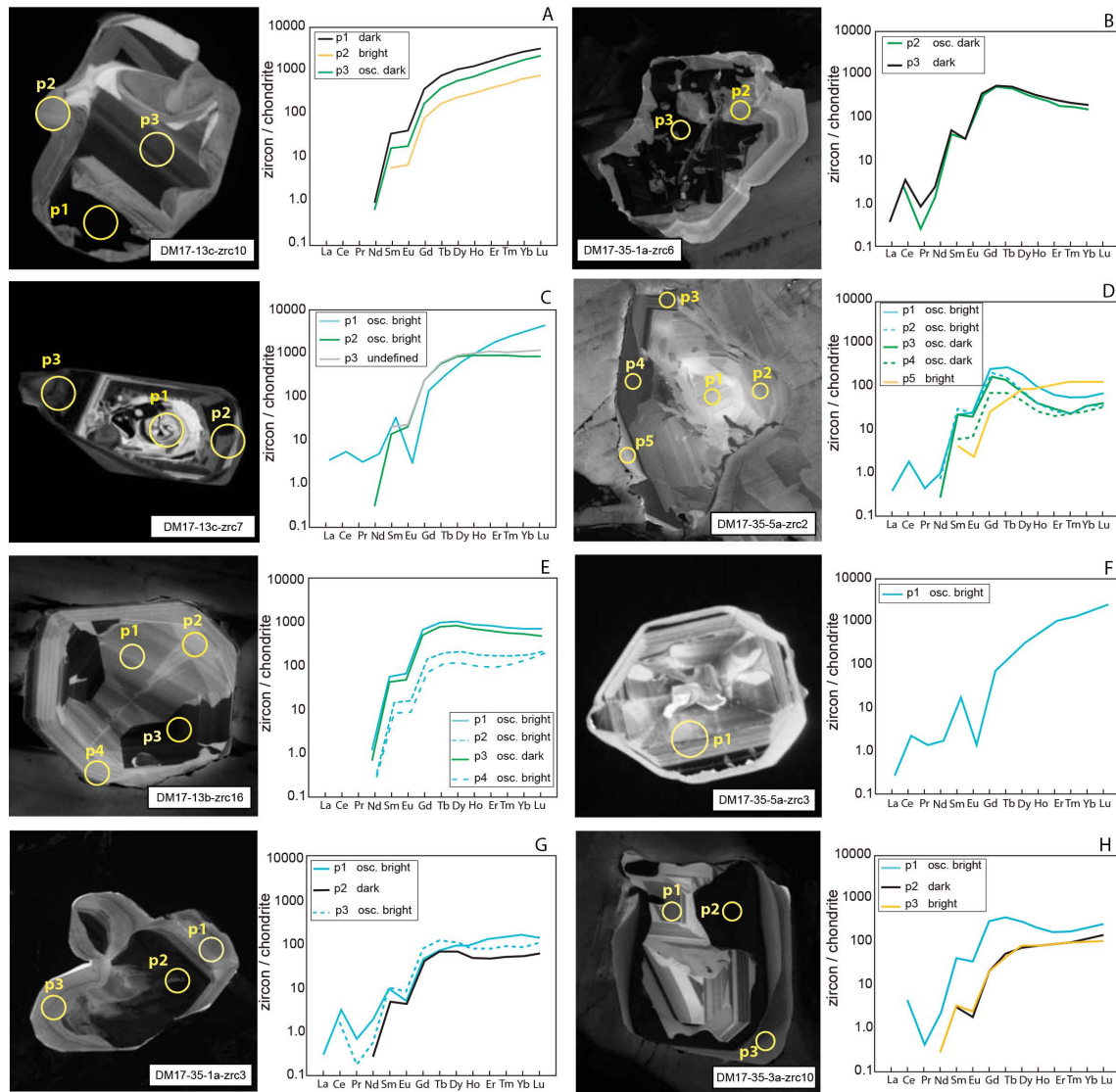


FIGURE 3. REE pattern in selected partially exposed zircon inclusions. Color coding follows that in Figure 2: black = dark, green = oscillatory-dark, blue = oscillatory-bright, orange = bright, gray = undefined. Yellow circles are scaled with respect the figure and are 20 μm across (laser spot).

vs. $D\alpha$ diagram, most of the Alpine domains fall within the broad interpolation line given by Nasdala et al. 2001 (Fig. 6). Nasdala et al. (2001) interpreted the data plotting outside of such interpolation band as the possible effects of thermal annealing in the crystal. In our case, we interpret the few points outside the interpolation band as the result of possible partial annealing of inherited zircon cores (Permian and Caledonian in age) during the Alpine metamorphism.

Effect of metamictization and annealing on Raman shift

Previous studies have established that partial metamictization can have a major influence on zircon Raman spectra (e.g., Zhang et al. 2000; Geisler et al. 2001; Nasdala et al. 2001; Binvignat et al. 2018). In general, depending on the degree of metamictization, zircon crystals can show a significant Raman peak broadening and frequency shift toward lower wavenumbers with respect to pristine samples taken as reference. Furthermore, Geisler et al.

(2001) pointed out the possible heterogeneous (step-evolution) effect that annealing could have on peak broadening and position.

Based on these considerations, the relationship between the peak width and the Raman shift (Fig. 2) can be exploited as a discriminant between partially metamict/annealed crystals and pristine crystals, even when they are completely buried within their mineral host (see also Zhong et al. 2019). Indeed, differently to phonon wavenumber, variation in the Γ of a Raman peak is independent from minor amounts of stress (Binvignat et al. 2018) as those usually recorded in host-inclusion systems. It follows that, when considering zircon inclusions, a buried partially metamict crystal should present Γ comparable to partially metamict crystals exposed at the surface and the same is valid for well-crystalline zircon grains. Therefore, we conclude that measurements with Γ greater than 5 cm^{-1} showing an inverse relationship with the Raman shift, represent partially metamict/annealed domains. On the

TABLE 2. Indicative age and dose of partially exposed zircon inclusions (spots are referred to Fig. 3)

Sample name	CC domain	Indicative ²⁰⁶ Pb/ ²³⁸ U	Indicative Age	D _α (g/events)	Raman shift (cm ⁻¹)	Γ (cm ⁻¹)
DM17-35-3A-zrc8-p1	osc. dark	0.04	Permian	8.3E+17	1007.6	5.6
DM17-35-3A-zrc8-p1	osc. dark	0.04	Permian	8.3E+17	1007.6(2)	5.6(4)
DM17-35-3A-zrc8-p2	osc. bright	0.08	Caledonian	2.2E+18	1008.0(3)	3.4(3)
DM17-35-3A-zrc10-p1	osc. bright	0.01	Alpine	5.3E+16	1007.9(2)	3.9(4)
DM17-35-3A-zrc10-p2	dark	0.01	Alpine	2.6E+17	1006.7(5)	5.3(3)
DM17-35-3A-zrc10-p3	bright	0.01	Alpine	1.2E+17	1007.9(4)	3.6(3)
DM17-35-5A-zrc2-p1	osc. bright	0.01	Alpine	4.9E+16	1009.1(3)	3.6(4)
DM17-35-5A-zrc2-p4	osc. dark	0.01	Alpine	1.6E+17	1008.0(3)	4.5(4)
DM17-35-5A-zrc2-p5	bright	0.01	Alpine	1.4E+17	1008.9(2)	3.5(4)
DM17-35-1A-zrc6-p2	osc. dark	0.01	Alpine	3.3E+17	1008.9(2)	3.5(3)
DM17-35-1A-zrc6-p3	dark	0.01	Alpine	3.9E+17	1006.6(5)	6.2(3)
DM17-35-1A-zrc3-p2	dark	0.01	Alpine	3.7E+17	1006.3(3)	5.7(3)
DM17-35-1A-zrc3-p3	osc. bright	0.01	Alpine	4.1E+17	1007.7(3)	4.2(2)
DM17-13-b-zrc13-p1	osc. bright	0.01	Alpine	1.5E+17	1007.9(2)	4.6(4)
DM17-13-b-zrc13-p2	osc. bright	0.01	Alpine	1.1E+17	1007.7(3)	4.7(4)
DM17-13-b-zrc15-p2	osc. bright	0.01	Alpine	8.6E+16	1008.3(2)	4.5(4)
DM17-13-b-zrc15-p3	osc. bright	0.01	Alpine	4.7E+16	1008.3(2)	4.0(3)
DM17-13-b-zrc16-p1	osc. bright	0.01	Alpine	1.7E+17	1007.9(3)	5.1(3)
DM17-13-b-zrc16-p2	osc. bright	0.01	Alpine	8.5E+16	1008.2(1)	4.0(4)
DM17-13-b-zrc16-p3	osc. dark	0.01	Alpine	2.2E+17	1007.3(2)	5.6(4)
DM17-13-b-zrc16-p4	osc. bright	0.01	Alpine	5.3E+16	1008.3(2)	3.6(5)
DM17-13-c-zrc7-p1	osc. bright	0.04	Permian	6.6E+17	1009.3(1)	5.8(3)
DM17-13-c-zrc7-p2	osci. dark	0.01	Alpine	1.1E+17	1011.3(2)	3.9(3)
DM17-13-c-zrc7-p3	undefined	0.01	Alpine	1.3E+17	1010.2(3)	4.3(4)
DM17-13-c-zrc10-p1	dark	0.01	Alpine	5.2E+17	1006.2(5)	10.6(3)
DM17-13-c-zrc10-p2	bright	0.01	Alpine	1.4E+17	1008.3(2)	3.3(2)
DM17-13-c-zrc10-p3	osc. dark	0.01	Alpine	2.3E+17	1007.4(3)	5.0(4)

Note: Uncertainty on the D_α is within 14 % based on the U and Th concentration uncertainty.

other hand, domains with lower Γ (i.e., 3.5–5.0 cm⁻¹) and with no correlation with the Raman shift are associated to a non-metamict domain. In this regard, it is worth noting that also Alpine zircons with high U contents fall into the metamict domain.

Geisler and Pidgeon (2002) pointed out that possible annealing processes may influence both the Raman shift and the peak width, as well as the relationship between them during secondary geological processes. However, from their results, it is evident that such effects are critical for zircon with moderate to heavy levels of radiation damage (i.e., ω < 1004 cm⁻¹ and Γ > than 11–12 cm⁻¹). In our case, the zircon inclusion displaying a negative slope result all with Γ < 11 cm⁻¹ and ω > 1004 cm⁻¹. This correlation can be tentatively interpreted as an indication that, as previously stated (e.g., Fig. 6), annealing effects were negligible or absent in most of the samples over the relatively short geological evolution (280 or 35 Ma to present) and fast subduction metamorphism (Gebauer et al. 1997;

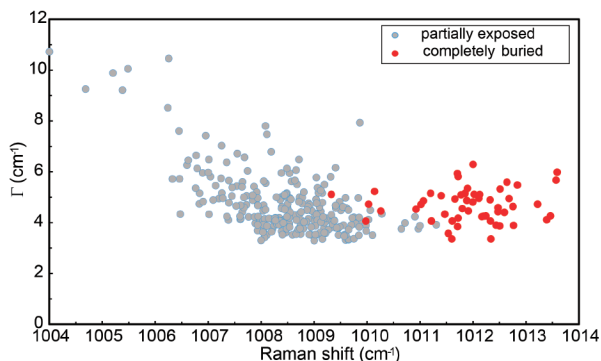


FIGURE 4. Γ and Raman shift of partially exposed vs. completely buried zircon inclusions.

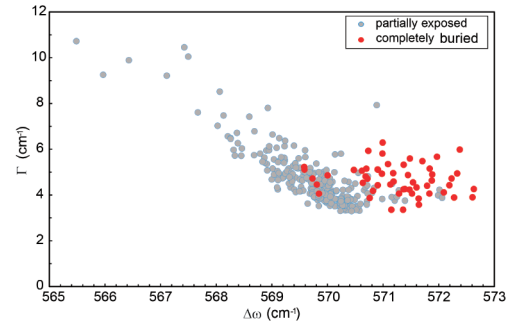


FIGURE 5. Γ of the B_{1g} mode near 1008 cm⁻¹ and Δω = ω₁₀₀₈ - ω₄₄₀ of partially exposed vs. selected and completely buried zircon inclusions.

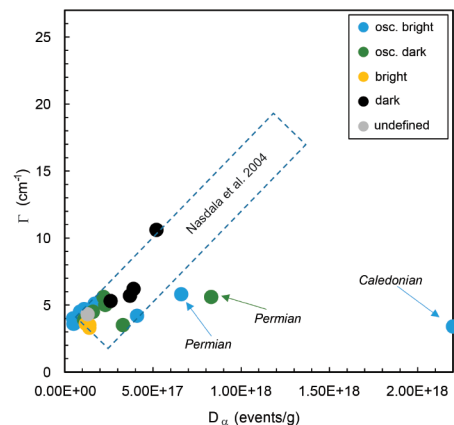


FIGURE 6. Γ of the B_{1g} mode near 1008 cm⁻¹ as function of the effect of increasing metamictization in terms of accumulated radiation dose.

Rubatto and Hermann 2001; Gauthiez-Putallaz et al. 2016).

However, it is difficult to make a rigorous prediction of the effect of metamictization on the determination of the residual pressure of a buried inclusion. For zircon with Γ larger than 5 cm⁻¹ there are at least two additional unknown variables:

- The reference wavenumber ω₀ of an equally metamict free crystal;
- The phonon mode compressibility β = [1/ω₀][dω/dP].

Both variables are very sensitive to the accumulated radiation dose D_α (e.g., Binivignat et al. 2018). Unfortunately, the dispersion of our data in the established Γ–D_α trend is too large to give a reliable value of D_α and consequently of ω₀ and β for the purposes of Raman thermobarometry.

Effect of zircon size on Raman shift for partially exposed inclusions

For completely unstressed pristine crystals, the phonon wavenumber usually is expected not to exceed the values of 1008.5–1009 cm⁻¹ (see Geisler et al. 2001; Binivignat et al. 2018). Nevertheless, our partially exposed grains, even considering only the non-metamict domain in Figure 2, show a larger variation (up to 1011 cm⁻¹).

As reported by Campomenosi et al. (2018), partially exposed

inclusions can still preserve a notable stress state in terms of Raman peak shift as a function of the inclusion exposition degree and size. To better clarify this issue, Figure 7 shows a Γ vs. $\Delta\omega$ diagram of partially exposed inclusions discriminating between grains with different sizes (<50 and >50 μm). Note that, in this case, we show only data from the exposed inclusions showing bright and or oscillatory-bright domains at CC because only for such domains can we safely exclude other effects described above. Data from inclusions of different size largely overlap. However, about 20% of the smaller inclusions have a higher $\Delta\omega$ than the larger one (with one exclusion) and reach a $\Delta\omega$ of 572 cm^{-1} that are never observed in the larger inclusions. This is in agreement with theoretical predictions (Mazzucchelli et al. 2018) for which, within the same host, small inclusions, when partially exposed, tend to retain more stress, due to the larger surface-to-volume ratio, i.e., larger impact of the host upon relatively larger surface of the inclusion. On the other hand, small inclusions should exhibit less dispersion of the degree of exposure. Hence, the quite large spread of our data is most probably due to the superposition of both effects related to the inclusion size: the surface-to-volume ratio and the degree of exposure.

A new protocol for the selection of zircon inclusions in garnet for elastic thermobarometry

Based on the above considerations, we propose a simple protocol for selecting zircon inclusions to get reliable residual pressure estimates for elastic thermobarometry.

- Buried crystals must be isolated from other inclusions, section surfaces, host boundary, cracks, and from any other kind of boundaries. This implies that thicker sections (i.e., 300 μm) are better (see also Campomenosi et al. 2018).
- At least 2–3 measurement spots for each crystal moving from core to rim should be performed: this enables detection of eventual structural heterogeneity within the crystal. For this purpose, it is better to use a spectrometer with a confocal optical microscope to reach the best spatial resolution.
- An instrumental spectral resolution equal or better than 2 cm^{-1} should be used and only those inclusions presenting a Γ of the B_{1g} mode near 1008 cm^{-1} smaller than 5 cm^{-1}

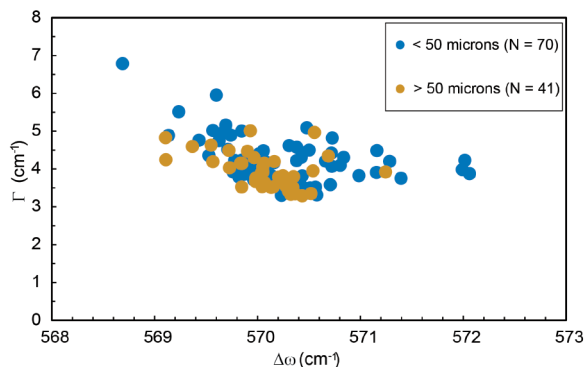


FIGURE 7. Γ of the B_{1g} mode near 1008 cm^{-1} vs. $\Delta\omega = \omega_{1008} - \omega_{440}$ comparing small (<50 μm) and large (>50 μm) partially exposed zircon inclusions. The spots refer only to bright and oscillatory bright domains.

should be considered. Note that this value represents the summation of the analytical uncertainty (i.e., instrumental spectral resolution) to the minimum value of about 3 cm^{-1} that has been measured on several grains by the two different spectrometers (see Fig. 4).

- Only inclusions in which the phonon wavenumber and the Γ is constant across the entire crystal volume should be selected.
- If Γ is constant, but the phonon wavenumber changes across the crystal volume there could be effects due to the shape of the inclusion (Campomenosi et al. 2018). In this case it is recommended to consider the measurement collected at the center of the crystal to calculate the residual pressure of the inclusion.
- Partially exposed crystals should not be used as reference to calculate residual stress in the buried inclusions: they can be still under a notable residual stress state. It is recommended using a completely free crystal or large exposed inclusions for which the residual stress state and the metamictization effects are negligible (see for example Fig. 1d).
- Whenever possible, a statistically significant amount of partially exposed inclusions should be selected to double check their textural complexity by CC or CL imaging and their chemistry. As an alternative, imaging and chemical checks should be performed on inclusions that have been exposed after the Raman measurements.
- Chemical and age measurements should be considered as important corollary information to reconstruct the petrogenesis of the zircon inclusions and the garnet host.

IMPLICATIONS

Zircon inclusions are difficult to manage correctly for elastic thermobarometry applications, and a detailed characterization of the inclusions should be performed before extracting thermobarometric data. In our systematic study, the combination of structural and chemical information obtained by different analytical techniques on partially exposed zircon inclusions allows to reliably predict the structural state of buried crystals for Raman spectroscopic measurements.

Our results provide a solid basis for the selection of reliable zircon inclusions to use for elastic thermobarometry applications. This method, in combination with the already rich tool set that can be applied to zircon [e.g., Ti-in-zircon thermometry (Watson et al. 2006), U-Pb geochronology, oxygen isotopes], will provide an even more detailed characterization of P - T - t fluid and deformation history of metamorphic rocks.

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