Cosmogenic $^3$He in terrestrial rocks: a review

P.-H. Blard*
CRPG, CNRS, Université de Lorraine, 54000, Nancy, France

*Email: blard@crpg.cnrs-nancy.fr

Accepted by Chemical Geology (Elsevier) on September 22nd 2021

 Earlier versions:
Non-peer reviewed preprint posted on EarthArXiv on July 5th 2021
Initial submission to Chemical Geology (Elsevier) on July 3rd 2021

Category: Invited Review Article
Editors:
Donald Porcelli, Email: donald.porcelli@earth.ox.ac.uk
Timothy Horscroft, Email: timothy.horscroft@elsevier.com

17,500 words
19 figures
1 Table
3 Supplementary Tables:
- Supp. Table 1: Magmatic $^4$He concentrations database
  https://doi.org/10.24396/ORDAR-89
- Supp. Table 2: Excel spreadsheet for computing radiogenic $^4$He and nucleogenic $^3$He
  https://doi.org/10.24396/ORDAR-90
- Supp. Table 3: $^3$He/$^{10}$Be cross-calibrated production rates database
  https://doi.org/10.24396/ORDAR-91

Keywords: cosmogenic $^3$He; magmatic $^3$He; isochrons; exposure age; production rates; $^3$He/$^{10}$Be cross-calibration
This review article summarizes the state of the art of cosmogenic $^3$He ($^3$He$_c$), with a focus on the most efficient methods and strategies for measuring this cosmogenic noble gas in terrestrial minerals. Our ability to accurately and precisely measure cosmogenic $^3$He in minerals is mainly constrained by the level of the non-cosmogenic $^3$He background (i.e., magmatic, radiogenic, nucleogenic, and atmospheric $^3$He), and thus by the geological characteristics of the samples. Constructing $^3$He vs $^4$He isochrons by analyzing several aliquots from the same surface sample constitutes a useful and overlooked method. Isochrons can be applied to mineral samples with homogeneous radiogenic $^4$He concentration and variable magmatic helium concentrations. This method also allows the direct and joint determination of the cosmogenic $^3$He concentration and the magmatic $^3$He/$^4$He ratio, precluding the often-complicated step of vacuum crushing. I perform numerical modeling to explore the impact of the non-cosmogenic $^3$He components on the final uncertainties and detection limits of $^3$He$_c$ measurement. Reducing the magmatic component by selecting phenocrysts in the 100–500 µm size fraction improves the precision of cosmogenic $^3$He analyses. Moreover, it is important to measure U, Th, and Li concentrations in the analyzed minerals and their host rocks to ensure proper corrections for radiogenic $^4$He and nucleogenic $^3$He, improving both the accuracy and precision of the method. I discuss the most important aspects of analytical techniques, including the key parameters of noble gas mass spectrometry that result in accurate and precise helium isotopic measurements.

I also review $^3$He$_c$ production rates and their spatial variability: the global database of absolute calibration sites yields a world-wide average $^3$He$_c$ production rate in olivine and pyroxene of 124 ± 11 at g$^{-1}$ yr$^{-1}$ using the LSD scaling and the online CREp calculator (https://crep.otelo.univ-lorraine.fr/#/). Cross-calibrations against $^{10}$Be indicate that the ratio of the production rate of $^3$He$_c$ in olivine/pyroxene to that of $^{10}$Be in quartz is 33 ± 2 and increases by less than 7% between sea level and 5,000 m elevation. This important observation demonstrates that $^3$He in olivine/pyroxene and $^{10}$Be in quartz are accurately scaled with existing models and can hence be considered as synchronized chronometers. However, $^3$He$_c$/$^{10}$Be cross-calibrations based on $^3$He$_c$ in accessory minerals (zircon, garnet, kyanite, apatite) yield unexpectedly high $^3$He/$^{10}$Be production ratios of 40–60 above 3,000 m elevation. As the capture of cosmogenic thermal neutrons by $^6$Li is unlikely to explain this excess, other mechanisms are required, such as $^3$He$_c$ inherited from previous exposure episodes, unrecognized specific reaction pathways, or the impact of snow cover. New cross-calibration data obtained by
measuring $^3$He against other cosmogenic nuclides ($^{14}$C, $^{36}$Cl, $^{21}$Ne, $^{53}$Mn) in different settings will advance our understanding of cosmogenic nuclide production rates and improve the accuracy and precision of applications relying on cosmogenic $^3$He. Other improvements could extend the applicability of the $^3$He geoscientific toolbox; for example, coupling $^3$He with radioactive cosmogenic nuclides ($^{10}$Be, $^{14}$C, $^{36}$Cl, $^{53}$Mn) will allow paleoaltimetry or the determination of burial ages or paleo-depths in intermediate and mafic terrains.
Contents

1. Introduction

2. Basic concepts of in-situ terrestrial cosmogenic \( {^3}\text{He} \)
   2.1. Discovery of terrestrial cosmogenic \( {^3}\text{He} \)
   2.2. Advantages and limitations of \( {^3}\text{He} \): an ideal complement to \( {^{10}\text{Be}} \)
   2.3. A useful geo-chronometer and probe for many applications in Earth surface science
   2.4. None like it hot: why is cosmogenic \( {^3}\text{He} \) mainly measured in olivines, pyroxenes, iron oxides and accessory minerals?

3. \( {^3}\text{He} \) sources and correcting for non-cosmogenic \( {^3}\text{He} \)
   3.1. The global \( {^3}\text{He} \) and \( {^4}\text{He} \) budgets in terrestrial minerals
   3.2. Magmatic \( {^3}\text{He} \) and \( {^4}\text{He} \)
     3.2.1. Impact of the \( {^4}\text{He}_{\text{mag}} \) correction on the final cosmogenic \( {^4}\text{He} \) uncertainty
     3.2.2. Using grains sized 100–500 \( \mu \text{m} \) reduces the magmatic helium contribution
   3.3. Radiogenic \( {^4}\text{He} \)
     3.3.1. The production of radiogenic \( {^4}\text{He} \): general principles and equations
     3.3.2. Removing the implanted \( {^4}\text{He}^* \) component
     3.3.3. Uncertainty arising from the radiogenic \( {^4}\text{He} \) correction
   3.4. Nucleogenic \( {^3}\text{He} \)
     3.4.1. The production of nucleogenic \( {^3}\text{He} \): general principles and equations
     3.4.2. Uncertainty arising from the nucleogenic \( {^3}\text{He} \) correction
   3.5. Strategies to estimate the non-cosmogenic \( {^3}\text{He} \) components
     3.5.1. Case 1, the general situation: \( \text{He} \) closure age > exposure age, but \( {^4}\text{He}^* \) is not significantly larger than \( {^4}\text{He}_{\text{mag}} \)
     3.5.2. Case 2: Samples with similar closure and exposure ages – the R-factor
     3.5.3. Isochrons (adequate for Cases 1 and 2)
       3.5.3.1. \( {^3}\text{He} \) vs \( {^4}\text{He} \) isochrons
       3.5.3.2. \( {^3}\text{He}/ {^4}\text{He} \) vs \( 1/ {^4}\text{He} \) isochrons
     3.5.4. Case 3: Samples with large amounts of radiogenic \( {^4}\text{He}^* \) (\( {^4}\text{He}^* >> {^4}\text{He}_{\text{mag}} \))

4. Analytical methods for measuring \( {^3}\text{He} \) and \( {^4}\text{He} \) in minerals
   4.1. Extraction procedure
     4.1.1. In-vacuo crushing
     4.1.2. Heating furnace systems
     4.1.3. Lasers
   4.2. Purification lines
   4.3. Mass spectrometry
   4.4. Interlaboratory comparisons
   4.5. Potential sources of systematic uncertainties in \( {^3}\text{He} \) and \( {^4}\text{He} \) measurements

5. Discussion: cosmogenic \( {^3}\text{He} \) production rates, the accuracy of spatial scaling, cross-calibration with other cosmogenic nuclides, online calculators and new frontiers
   5.1. Production rates
   5.2. The global \( {^3}\text{He} \) production rate calibration database
   5.3. Online calculators
   5.4. Production of cosmogenic \( {^3}\text{He} \) by muons
   5.5. Cross-calibration of \( {^3}\text{He} \) against \( {^{10}\text{Be}} \) and other cosmogenic nuclides
   5.6. New directions of cosmogenic \( {^3}\text{He} \) coupled with cosmogenic radioactive nuclides (\( {^14}\text{C}, {^{10}\text{Be}}, {^{36}\text{Cl}} \), and \( {^{53}\text{Mn}} \))

6. Concluding remarks
1. Introduction

In-situ cosmogenic nuclides are nuclides in rocks and minerals that are produced through nuclear reactions upon bombardment by high energy (>1 MeV) cosmic particles. Because cosmogenic nuclide concentrations in minerals depend on the amount of time the minerals were exposed at Earth’s surface, they represent a powerful and wide array of geochronometric tools with applications in geomorphology, paleoclimatology and geohazard analysis (Fig. 1). Although a large body of literature is dedicated to radioactive cosmogenic nuclides ($^{10}$Be, $^{26}$Al, and $^{36}$Cl), less attention has been given to cosmogenic noble gases ($^{3}$He and $^{21}$Ne): in 2020, articles relying on cosmogenic $^{3}$He and $^{21}$Ne only represented 12% and 10%, respectively, of the ~2,700 publications involving any type of in-situ cosmogenic nuclides (source: Web of Science). $^{3}$He is nonetheless an interesting alternative to $^{10}$Be for quartz-poor lithologies because it is present in measurable concentrations in minerals that are not suited to $^{10}$Be or $^{26}$Al analyses, such as those in mafic rocks (olivine, pyroxene). $^{3}$He is also measurable in accessory minerals in silica-rich lithologies (zircon, apatite, iron oxides). Moreover, because of their nuclear stability, cosmogenic noble gases can, in theory, be used to analyze events that occurred beyond tens of millions of years ago (e.g. Sartégou et al., 2020) and are thus ideal complements to radioactive cosmogenic isotopes, which decay over shorter durations.

Since the discovery of terrestrial cosmogenic $^{3}$He more than 30 years ago by Mark Kurz and others (Craig and Poreda, 1986; Kurz, 1986a), significant progress following the pioneer studies of Cerling and Craig (1994) has developed cosmogenic $^{3}$He into a pertinent geological probe. Over the last decades, dozens of studies have used cosmogenic $^{3}$He to date and quantify a large variety of Earth processes (Figs. 1, 2), such as the timing of volcanic eruptions (e.g. Kurz et al., 1990), fault kinetics (e.g. Fenton et al., 2001), denudation rates (e.g. Puchol et al., 2017), paleoclimatology (e.g. Martin et al., 2018), paleoaltimetry (e.g. Blard et al., 2006a), dating surfaces that are millions of years old (e.g. Margerison et al., 2005), or tsunami hazard analyses (e.g. Ramalho et al., 2015).

The measurement of cosmogenic $^{3}$He requires specific facilities and apparatuses that are currently available in several noble gas laboratories worldwide (e.g. Caltech, Pasadena, USA; CRPG, Nancy, France; SUERC, Glasgow, UK; ETH, Zurich, Switzerland; GFZ, Potsdam, Germany; BGC, Berkeley, USA; LDEO, Palisades, USA). However, because not all minerals are suitable for cosmogenic $^{3}$He analysis, the reliability of cosmogenic $^{3}$He data depends directly on the nature and geological history of the analyzed rocks. Because these methodological aspects have been disseminated in various specialized papers (Blard et al., 2008; Blard and Farley, 2008; Blard and Pik, 2008; Farley et al., 2006; Goehring et al., 2010;
Kurz, 1986a, b; Larsen et al., 2019, 2021; Martin et al., 2017; Protin et al., 2016; Shuster et al., 2004; Trull et al., 1991), this article summarizes the most important theoretical and methodological aspects of cosmogenic $^3$He analyses by reviewing the current state of the art and the best strategies for accurately and precisely measuring cosmogenic $^3$He concentrations.

I begin with a short history of the scientific breakthroughs that led to the discovery of terrestrial cosmogenic $^3$He ($^3$He$_c$), then describe the different sources and production pathways of $^3$He and $^4$He in minerals. I then present methodological guidelines about the theoretical and practical aspects of $^3$He analyses in mafic minerals (olivine and pyroxene), with the objective of attaining the best precision and accuracy. For a given sample, our ability to detect cosmogenic $^3$He is determined by its non-cosmogenic $^3$He background, and then by its geological characteristics. I hence perform numerical modeling to explore the impact of these non-cosmogenic $^3$He components on the total uncertainties. Then, I present the most important aspects of analytical techniques for helium measurements. Finally, I review and discuss our knowledge of the $^3$He$_c$ production rate and its spatial variability, relying on published calibration sites and cross-calibrations against other cosmogenic nuclides, notably $^{10}$Be, $^{36}$Cl and $^{21}$Ne. I also explore potential new applications based on the coupling of $^3$He$_c$ with radioactive cosmogenic nuclides ($^{10}$Be, $^{36}$Cl, $^{53}$Mn).
Figure 1. Principles of cosmogenic $^3$He production and its main applications. $^3$He, is produced by spallation reactions induced by high-energy cosmogenic neutrons (> 10 MeV), low-energy cosmogenic neutrons (<1 keV) and muons. The three main applications of cosmogenic $^3$He are determined by the geomorphological characteristics of the studied objects: 1) on non-eroded surfaces, cosmogenic $^3$He concentrations measure exposure durations; 2) in eroding landscapes (active versants, rivers) that have reached steady-state ($t > A/pg$), $^3$He concentrations measure denudation rates; 3) when coupled with measurements of radioactive cosmogenic nuclides (e.g., $^{10}$Be, $t_{1/2} = 1.4$ Ma) in the same rock sample that is now buried beyond the penetration depth of cosmic particles, it is possible to determine paleoelevations, paleodepths, or measure burial durations. Picture of Stromboli volcano, Italy: © Jcb-caz-11, CC-BY-SA-4.0.

2. Basic concepts of in-situ terrestrial cosmogenic $^3$He

2.1. Discovery of terrestrial cosmogenic $^3$He

Until the mid-1980's, the presence of cosmogenic $^3$He had only been documented in extraterrestrial materials such as lunar samples (e.g., Megruè, 1971) and meteorites (e.g., Fisher, 1972). These analyses were performed by melting samples under vacuum followed by $^3$He
analyses in a static, high-vacuum Nier-source mass spectrometer (Aldrich and Nier 1946; Kurz 1986a). During the 1970’s and 1980’s, however, noble gas systems switched from glass to fully metal purification lines, lowering instrumental blanks and, hence, detection limits by 5 orders of magnitude, from $10^8$ atoms (Megrue 1967) to $<10^3$ atoms on modern instruments (e.g., Zimmermann et al., 2018).

In 1986, Mark Kurz and others demonstrated that cosmogenic $^3$He was also detectable in terrestrial rocks by analyzing olivines and pyroxenes from basalt samples exposed for about 500 kyr at the summit of Haleakalā volcano, Maui, Hawaii (Kurz, 1986a, 1986b; see also Craig and Poreda, 1986). The compelling evidence for the presence of cosmogenic $^3$He resulted from their two-step analysis of He isotopes in olivine phenocrysts: first crushing the olivines in vacuo, then fusing the remaining powder. The fused olivine yielded a $^{3}\text{He}/^{4}\text{He}$ ratio of 2,000 Ra (where $\text{Ra} = 1.384 \times 10^{-6}$ is the atmospheric $^{3}\text{He}/^{4}\text{He}$ ratio; Kurz, 1986). Such a high value had never been documented on Earth, as it is far above the highest $^{3}\text{He}/^{4}\text{He}$ ratios measured by crushing mantle olivines, which preferentially release trapped mantellic helium ($^{3}\text{He}/^{4}\text{He} = 8–30$ Ra; e.g., Parman, 2007). The observed massive $^3$He excess was thus interpreted as resulting from the build-up of cosmogenic $^3$He in the mantle silicates after they had reached the Earth’s surface and been exposed to cosmic rays. Additionally, relying on the fact that magmatic helium has a homogenous isotopic composition in phenocrysts of a few grams, Kurz (1986a, 1986b) proposed that the amount of $^4$He released by melting can be used as a proxy for the concentration of magmatic $^3$He:

$$^{3}\text{He}_c = ^{3}\text{He}_{\text{tot}} - ^{4}\text{He}_{\text{tot}} \times (^{3}\text{He}/^{4}\text{He})_{\text{mag}}, \quad (1)$$

where $^{3}\text{He}_{\text{tot}}$ and $^{4}\text{He}_{\text{tot}}$ are the total $^3$He and $^4$He concentrations, respectively, and $(^{3}\text{He}/^{4}\text{He})_{\text{mag}}$ is the magmatic $^{3}\text{He}/^{4}\text{He}$ ratio estimated from vacuum crushing.

For decades, this two-step crushing and fusion protocol was used with Equation (1) to correct for the magmatic $^3$He component (e.g. Kurz et al., 1990; Licciardi et al., 1999). However, several authors latter proposed that Equation (1) be modified to account for the presence of radiogenic $^4$He (Blard and Farley, 2008; Dunai and Wijbrans, 2000) to avoid overestimating the magmatic $^3$He contribution (see section 3).
2.2. Advantages and limitations of $^3$He: an ideal complement to $^{10}$Be

During the 1980’s, progress in accelerator mass spectrometry (AMS) made it possible to measure in-situ cosmogenic $^{10}$Be in hand-sized quartz samples (e.g., Klein et al., 1982; Litherland, 1980; Raisbeck et al., 1987). The nearly simultaneous discovery of cosmogenic $^3$He in terrestrial samples thus ideally completed the spectrum of applications of these new geochronological tools because $^3$He is suited to mafic minerals found in volcanic environments, whereas $^{10}$Be is suited to silicic, quartz-rich environments. Although cosmogenic $^3$He is produced in quartz, it is not retained in this mineral at Earth’s surface temperatures (see section 2.4).

Moreover, $^3$He has several specific advantages compared to other cosmogenic nuclides:

i) Because $^3$He is a light isotope, the cross section of spallation reactions producing cosmogenic $^3$He are less sensitive to the masses of target elements compared to other nuclides (Dunai, 2010). Consequently, compared to $^{21}$Ne, $^{36}$Cl, and $^{10}$Be, the rate of $^3$He$_c$ production by high-energy particles (>10 MeV) is less dependent on mineral chemical composition; indeed, empirical data show that olivine and pyroxene have the same $^3$He$_c$ production rates (Cerling and Craig, 1994; Dunai, 2010; Fenton et al., 2009; Martin et al., 2017). That said, Fe-rich minerals, such as ilmenite and hematite, have lower spallogenic production rates (e.g. Larsen et al., 2019; Shuster et al., 2012), whereas in Li-rich (> 10 ppm) minerals, $^3$He production may be enhanced by the capture of cosmogenic thermal neutrons (Dunai et al., 2007).

ii) $^3$He has one of the largest production rate / detection limit ratios, which theoretically enables the detection of exposure episodes shorter than 100 years in the most favorable conditions, i.e. in the case of rocks poor in magmatic helium that erupted less than 10 ka (Niedermann, 2002).

iii) The nuclear stability of $^3$He allows very old (>10 Ma) surfaces to be dated (e.g. Balter-Kennedy et al., 2020), whereas $^{10}$Be and $^{36}$Cl reach saturation after only ~5 Myr and ~1 Myr of exposure, respectively. Moreover, $^3$He may be used to document geological events that occurred in the deep past, such as ancient exposure episodes that occurred several million years ago.
Nonetheless, the stability of $^3$He also somewhat limits its utility: any exposure episode may be recorded by cosmogenic $^3$He, complicating the interpretation of scenarios with multiple burial-exposure episodes. Moreover, the stability of $^3$He implies that phenocrysts, especially ones that crystallized several million years ago, may also accumulate significant amounts of $^3$He through other processes (e.g., Farley et al., 2006). These two properties may be considered the “Achilles’ heel” of cosmogenic $^3$He, and it is fundamental that users take into account the geological context of their samples, and notably the closure (U-Th-Sm)/$^4$He age of the analyzed phenocrysts (see section 3): as closure age increases, so does the minimum measurable exposure age.

2.3. A useful geo-chronometer and probe for many applications in Earth surface science

Cosmogenic $^3$He is now one of the most commonly used cosmogenic nuclides because it can date geological events and quantify numerous surface processes occurring in mafic volcanic environments. Indeed, $^3$He covers a rather wide range of Earth science timescales, from dating historical lava flows (<1 ka; e.g. Heineke et al., 2016) to the oldest known surface exposure ages on Earth’s surface (>10 Ma, e.g. Balter-Kennedy et al., 2020).

Thus, the questions addressed using cosmogenic $^3$He have been numerous and intriguing (Fig. 2), ranging from dating volcanic eruptions (e.g. Kurz et al., 1990; Marchetti et al., 2020) reconstructing paleoglacial dynamics (e.g. Blard et al., 2007; Bromley et al., 2011; Cerling and Craig, 1994; Martin et al., 2018), fault kinematics (e.g. Fenton et al., 2001; Medynski et al., 2016; Ritz et al., 2016), determining fluvial erosion rates (e.g. Ferrier et al., 2013; Litty et al., 2021), and paleoaltimetry (e.g. Blard et al., 2005) to even more exotic and intriguing applications, such as identifying paleo-tsunami deposits (e.g. Ramalho et al., 2015) or dating ancient human footprints (e.g. Heineke et al., 2016).
Figure 2. Geological questions addressed using cosmogenic $^3$He: (A) dating lava infilling and fault scarps in the Afar rift (Medynski et al., 2016), (B) determining denudation rates in rivers of Santo Antao island, Cabo Verde (Litty et al., 2021), (C) dating 11-ka volcanic ash with preserved human footprints (Heineke et al., 2016), (D) moraine geochronology in the Tropical Andes (Martin et al., 2020), and (E) dating of a giant tsunami 73 ka due to flank collapse on the volcanic island of Fogo, Cabo Verde (Ramalho et al., 2015).

As for any cosmogenic nuclide, the validity and the strength of a cosmogenic $^3$He data is directly dependent on the quality of the geomorphological analysis and the representativeness of the collected samples. For example, in the case of surface exposure dating, it is essential to minimize the potential erosion or temporal cover. This can be achieved by selecting the flattest and highest surfaces, and by checking for clues of erosion, weathering or ash/soil cover (e.g. Cerling and Craig, 1994). Moreover, several mandatory field observations and sample characteristics must be noted, such as sample location (elevation, altitude, longitude), the topographic shielding (Dunne et al., 1999), sample thickness and density (Gosse and Phillips, 2001; Dunai, 2010). These data are key to computing accurate sample specific production rates, and, hence, consistent exposure ages or erosion rates.
2.4. None like it hot: why is cosmogenic $^3$He mainly measured in olivines, pyroxenes, iron oxides and accessory minerals?

To record a complete exposure history, cosmogenic nuclides must be quantitatively retained in the mineral of interest. Because of its small size (atomic radius = $31 \times 10^{-12}$ m), helium has relatively high diffusion rates in silicates: at 15 °C, its diffusivity is $\sim 10^{-14}$ to $10^{-13}$ cm$^2$ s$^{-1}$ in quartz (Shuster et al., 2004; Tremblay et al., 2014) and $\sim 10^{-22}$ to $10^{-21}$ cm$^2$ s$^{-1}$ in olivine (Blard et al., 2008; Delon et al., 2020). Figure 3 shows the remaining fraction of cosmogenic $^3$He in 100-µm to 1-cm olivine and quartz grains continuously exposed at Earth’s surface. Even if a 1-cm quartz grain is sufficiently large to retain enough $^3$He to record exposure durations up to $10^4$ years, $^3$He is not quantitatively retained in submillimetric quartz at typical surface temperatures on Earth (Fig. 3C). This is confirmed by measured cosmogenic $^3$He concentrations in < 1 mm quartz that are systematically lower than expected from other age constraints (Brook and Kurz, 1993; Cerling, 1990; Trull et al., 1991). Nonetheless, this partial retentivity can be exploited to determine paleotemperatures (Tremblay et al., 2014).
Figure 3. Modeled cosmogenic $^3$He retention in minerals continuously exposed to cosmic rays: (A) olivine at 50 °C, (B) olivine at 15 °C, and (C) quartz at 15 °C. Note different $y$-axis scales in each plot. These curves were computed by solving the production-diffusion equation for spheres (Wolf et al., 1998) of various diameters (100 µm, 500 µm, and 1 cm). $^3$He diffusion parameters are from low-temperature (<200 °C) diffusion experiments on quartz (Tremblay et al., 2014) and olivine (Delon et al., 2020).
In contrast, olivine behaves as a nearly closed system for exposure durations up to $10^5$ years (Fig. 3A, B): even 100-µm grains continuously exposed at 50 °C lose less than 10% of the $^3$He produced. However, for exposure or paleoexposure ages older than $10^6$ years, mineral sizes and thermal histories should be carefully evaluated to avoid bias due to thermal losses.

Other minerals with helium diffusivities low enough to evolve as closed systems over geological timescales at Earth’s surface include pyroxene, hematite, magnetite, goethite, zircon, and apatite (Amidon et al., 2008; Blard et al., 2008; Farley, 2018; Kober et al., 2005; Shuster et al., 2012; Vasconcelos et al., 2019). In contrast, calcites demonstrated nearly open system behavior at Earth’s surface temperatures at the 10 ka time scale (Amidon et al., 2015; Cherniak et al., 2015). The few available data moreover showed that helium diffusivity varies greatly from one calcite to another (Amidon et al., 2015).

Because olivine and pyroxene have high helium retentivity and are rather abundant in intermediate and mafic rocks (generally 1–10% in andesites and basalts), they are considered to be the most convenient mineral phases for $^3$He$_c$ analyses, and are thus the most commonly used (e.g. Blard et al., 2006; Bromley et al., 2014; Cerling and Craig, 1994; Goehring et al., 2010; Kurz 1986b; Martin et al., 2017). These phases are also relatively poor in U and Th (<10 ppm, (Blard and Farley, 2008)), and Li (<10 ppm, (Amidon et al., 2009; Blard et al., 2013)), implying the limited production of radiogenic $^4$He and nucleogenic $^3$He, respectively.

3. $^3$He sources and correcting for non-cosmogenic $^3$He

Helium has two stable isotopes: $^3$He and $^4$He. $^3$He is by several order of magnitudes less abundant than $^4$He. Moreover, the $^3$He/$^4$He ratio may significantly vary among geological reservoirs. Various natural processes control the incorporation and production of $^3$He and $^4$He in terrestrial minerals such as olivine and pyroxene (Fig. 4). It is thus fundamental to understand the budgets of these isotopes to ensure accurate and precise estimates of cosmogenic $^3$He concentrations in minerals. Thus, in this section, I describe the different production pathways of He isotopes and the most suited strategies to correct for non-cosmogenic $^3$He sources. These strategies are valid for measuring cosmogenic $^3$He in any type of mineral sample, whatever the application (dating surfaces, measuring erosion rates or burial ages; Fig. 1).
3.1. The global $^3$He and $^4$He budgets in terrestrial minerals

In terrestrial minerals, cosmogenic $^3$He is mainly produced by spallation reactions involving high-energy (>1 MeV) cosmic particles (Lal 1987), but also by low-energy (<1 keV) thermal cosmogenic neutrons through $^6$Li(n,α)$^3$He reactions (Dunai et al., 2007) and muons (Larsen et al., 2021; Nesterenok and Yakubovich 2016). These processes produce equivalent amounts of $^3$He and $^3$H, but because $^3$H decays into $^3$He with a half-life of 12 years, secular equilibrium can be assumed at the geological timescale, and the production of $^3$H can be taken as that of $^3$He (Lal, 1987). For simplicity, I hereafter assume that $^3$He$_c$ includes all these "cosmogenic" production pathways.

$^3$He$_c$ is produced at a rate of about 125 at g$^{-1}$ yr$^{-1}$ at sea level and high latitude (Martin et al., 2017) and this rate increases with elevation, reaching ~2,500 at g$^{-1}$ yr$^{-1}$ at 4,000 m elevation and high latitudes (Lal, 1991; Stone, 2000). Hence, geomorphological materials exposed at Earth’s surface for $10^3$-10$^6$ years have $^3$He$_c$ concentrations in the range of 10$^5$ to 10$^9$ at.g$^{-1}$. Modern analytical systems connected to the most recent Nier-source mass spectrometers (see sections 2.1 and 4) can measure such $^3$He concentrations to an analytical precision of a few percent in mineral samples of ~10 mg to ~1 g. Nonetheless, the minimum detectable concentration of cosmogenic $^3$He in a particular sample is determined by our ability to accurately and precisely determine the non-cosmogenic $^3$He contributions in that sample, which become increasingly difficult as the He closure age of a mineral increases.

$^3$He has four known origins in minerals, each contributing variable $^3$He concentrations (Fig. 4; Table 1):

i) Cosmogenic $^3$He ($^3$He$_c$) is hosted in the matrix and its concentration increases with exposure age;

ii) Magmatic inherited $^3$He ($^3$He$_{mag}$) is hosted in fluid/melt inclusions and the matrix, and its concentration is independent of time;

iii) Nucleogenic $^3$He ($^3$He$_{nuc}$) is produced by neutron capture in $^6$Li nuclei and their subsequent disintegration, is matrix-hosted, and its concentration increases with closure age (eruption and closure ages are similar in the case of volcanic rocks), and;

iv) Atmospheric $^3$He ($^3$He$_{atm}$) is present as a contaminant on the surfaces of analyzed silicates and its concentration is time-independent.
In contrast, $^4\text{He}$ is $10^6$ times more abundant than $^3\text{He}$ in the atmosphere and $10^5$ times more abundant in the mantle (e.g., Marty and Jambon 1987). Hence, although cosmogenic $^4\text{He}$ is also produced in terrestrial rocks, the average non-cosmogenic $^4\text{He}$ concentrations in minerals are so large that they hamper the detection of any cosmogenic $^4\text{He}$ excess. It is thus reasonable to consider cosmogenic $^4\text{He}$ contributions as negligible. Thus, only three $^4\text{He}$ origins are relevant to the $^4\text{He}$ budget of terrestrial minerals that have been exposed at the Earth’s surface (Fig. 4; Table 1):

i) Magmatic $^4\text{He}$ ($^4\text{He}_{\text{mag}}$) is fluid/melt inclusion- and matrix-hosted, and its concentration is time-independent;

ii) Radiogenic $^4\text{He}$ ($^4\text{He}^*$) is produced by the decay of $^{238}\text{U}$, $^{235}\text{U}$, and $^{232}\text{Th}$, is matrix-hosted, and its concentration increases with helium closure age, and;

iii) Atmospheric $^4\text{He}$ ($^4\text{He}_{\text{atm}}$) is present as a contaminant on the surfaces of analyzed silicates and its concentration is time-independent.
Figure 4. (A) Inventory of \(^3\)He and \(^4\)He species \((^3\text{He}_c, \text{cosmogenic } ^3\text{He}; ^3\text{He}_\text{mag}, \text{magmatic } ^3\text{He}; ^3\text{He}_\text{nuc}, \text{nucleogenic } ^3\text{He}; ^4\text{He}_\text{mag}, \text{magmatic } ^4\text{He}; ^4\text{He}_*, \text{radiogenic } ^4\text{He})\) and production pathways in an olivine phenocryst. (B) Ranges of \(^3\)He and \(^4\)He concentrations in phenocrysts, assuming cosmogenic \(^3\)He production at sea level, a nucleogenic \(^3\)He production rate of 0.01 at g\(^{-1}\) yr\(^{-1}\) and a radiogenic \(^4\)He production rate of \(10^5\) at g\(^{-1}\) yr\(^{-1}\) (typical values for olivine phenocrysts in basalt) (C) Stopping distances of cosmogenic \(^3\)He and \(^3\)H in basalt (Larsen et al., 2019; Ziegler et al., 2010).

In practice, diverse laboratory procedures are used to selectively release these different helium components, depending on their location in the studied mineral (Table 1; Fig. 4). The two main categories of extraction methods are: i) *in vacuo mineral crushing*, which preferentially releases the magmatic helium components hosted in fluid and melt inclusions; and ii) *in vacuo heating or melting (using a furnace or a laser)*, which extracts all matrix and inclusion-hosted helium components from the mineral. The heating step is mandatory to extract the totality of the cosmogenic \(^3\)He from a sample, but inconveniently releases all other varieties (magmatic \(^3\)He and \(^4\)He, radiogenic \(^4\)He, nucleogenic \(^3\)He, and atmospheric \(^3\)He and \(^4\)He), which may complicate the detection of cosmogenic \(^3\)He. Thus, this section presents the most suited
strategies to correct for these non-cosmogenic $^3$He components, and section 4 details extraction procedures and helium measurement techniques.

**Table 1 – Characteristics of $^3$He and $^4$He varieties**

<table>
<thead>
<tr>
<th>Component</th>
<th>Cosmogenic</th>
<th>Magmatic</th>
<th>Nucleogenic</th>
<th>Radiogenic</th>
<th>Atmospheric</th>
</tr>
</thead>
<tbody>
<tr>
<td>$^3$He</td>
<td>Matrix-hosted.</td>
<td>Hosted in fluid/melt inclusions and matrix.</td>
<td>Produced by thermal neutron capture in $^6$Li.</td>
<td>Absent.</td>
<td>Contaminant adsorbed at the grain surface.</td>
</tr>
<tr>
<td></td>
<td>Determination by vacuum crushing or isochrons.</td>
<td>Less abundant in fractions finer than 500 µm.</td>
<td></td>
<td></td>
<td>Less abundant in fractions coarser than 100 µm.</td>
</tr>
<tr>
<td>$^4$He</td>
<td>Negligible compared to other $^4$He varieties.</td>
<td>Same as $^3$He.</td>
<td>Absent.</td>
<td>Produced by decay of $^{238}$U, $^{235}$U, and $^{232}$Th.</td>
<td>Same as $^3$He.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Matrix-hosted.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Concentration increases with closure age.</td>
<td></td>
</tr>
</tbody>
</table>

When a mineral sample is fused in vacuo, all $^3$He and $^4$He species are released, whatever their location in the mineral. Hence, the total $^3$He amount measured with a noble gas mass spectrometer is defined by the budget (modified from Farley et al., 2006; Blard and Farley, 2008):
Equation (2) is similar to:

\[ \text{He}_{\text{tot}} = \int_0^{t_e} P_3 \, dt + \text{He}_{\text{mag}} + \int_0^{t_e} P_{\text{nuc}} \, dt + \text{He}_{\text{atm}}, \]  

\[ (3) \]

where \( P_3 \) and \( P_{\text{nuc}} \) (at g\(^{-1}\) yr\(^{-1}\)) are the time-dependent local cosmogenic and nucleogenic \(^3\)He production rates, respectively, and \( t_e \) and \( t_c \) (yr) are the exposure and closure ages of the sample, respectively.

Additionally, the \(^4\)He budget can be written as:

\[ \text{He}_{\text{tot}} = \text{He}_{\text{mag}} + \text{He}_{\text{atm}}, \]  

\[ (4) \]

or:

\[ \text{He}_{\text{tot}} = \text{He}_{\text{mag}} + \int_0^{t_c} P_4 \, dt + \text{He}_{\text{atm}}, \]  

\[ (5) \]

where \( P_4 \) (at g\(^{-1}\) yr\(^{-1}\)) is the time-dependent \(^4\)He\(^*\) production rate and \( t_c \) (yr) is the closure age of the sample.

### 3.2. Magmatic \(^3\)He and \(^4\)He

#### 3.2.1. Impact of the \(^3\)He\(_{\text{mag}}\) correction on the final cosmogenic \(^3\)He uncertainty

Just as it is more difficult to find a golf ball in tall grass than on a well-maintained fairway, the relative uncertainty on the cosmogenic \(^3\)He concentration increases with increasing contribution from magmatic \(^3\)He (Figs. 4, 5). This is a signal-to-noise ratio issue: the higher the ratio, the lower the uncertainty arising from the magmatic \(^3\)He correction. This relationship can be modeled by defining the uncertainty on the total cosmogenic \(^3\)He concentration based on the uncertainties on the other variables, notably the \(^4\)He concentrations and the magmatic \(^3\)He/\(^4\)He ratio: equation (6) is derived from equation (1) by applying the Taylor series approximation and assuming that all involved uncertainties are independent:

\[ \sigma_{\text{He}_c}^2 = \sigma_{\text{He}_{\text{tot}}}^2 + \left[ \sigma_{\text{He}_{\text{mag}}} \cdot \frac{\text{He}}{\text{He}}_{\text{mag}} \right]^2 + \left( \text{He}_{\text{mag}} \cdot \sigma_{3/4} \right)^2, \]  

\[ (6) \]
where \( \sigma_{3He_c} \), \( \sigma_{3He_{tot}} \), \( \sigma_{4He_{mag}} \), and \( \sigma_{3/4} \) are the uncertainties on the cosmogenic \(^3\)He concentration, the total \(^3\)He amount extracted by fusion, the magmatic \(^4\)He concentration, and the magmatic \(^3\)He/\(^4\)He ratio, respectively.

Figure 5 shows the impact of the magmatic helium correction on the final uncertainty of the measured \(^3\)He\(_c\) concentration under different exposure scenarios using Equation (6). This calculation illustrates the importance of the \(^3\)He\(_c\)/\(^3\)He\(_{mag}\) ratio on the final uncertainty on \(^3\)He\(_c\): the larger the magmatic correction, the lower the uncertainty. The impact of the magmatic correction on the \(^3\)He\(_c\) uncertainty is hence larger at lower elevations, where the cosmogenic production rate is smaller. For the median magmatic \(^4\)He concentrations reported in olivine and pyroxene phenocrysts (Fig. 5C), the total relative uncertainty on cosmogenic \(^3\)He is below 5% and 20% for 100 ka and 10 ka surfaces, respectively (Fig. 5A,B). However, the uncertainty may reach 50-100% in the case of 1 ka surfaces.
Figure 5. (A, B) Modeling the total relative uncertainty (1σ) of the measured cosmogenic $^3$He concentration for variable magmatic $^4$He concentrations and exposure durations. Because the $^3$He production rate changes with elevation, this simulation considers a mineral at (A) 2,000 m elevation and (B) sea level. This calculation applies Equation (6) assuming a magmatic $^3$He/$^4$He ratio of 8 Ra with an uncertainty of 20%, an uncertainty of 3% on the total amount of $^3$He extracted by fusion, an uncertainty of 1% on the total amount of $^4$He extracted by fusion, and a negligible contribution from radiogenic $^4$He. (C) The distribution of the total magmatic $^4$He concentrations observed in olivine and pyroxene phenocrysts (Ackert et al., 2003; Blard et al., 2006; Cerling and Craig, 1994; Dunai and Wijbrans 2000; Eaves et al., 2015; Fenton and Niedermann, 2014; Foeken et al., 2012; Goehring et al., 2010; Licciardi et al., 2006; Licciardi et al., 1999). This dataset includes only lavas that are young enough (<200 ka) to ensure that the radiogenic $^4$He correction does not represent a significant systematic uncertainty.
A recent compilation of magmatic helium concentrations measured in olivine and pyroxene phenocrysts in basalts (Puchol et al., 2017 and references therein) indicates that the total magmatic helium concentrations in these phenocrysts are largely variable, both at the scale of a single lava flow and among minerals in a single sample (Fig. 5C, Table S1, https://doi.org/10.24396/ORDAR-89). On average, magmatic $^4$He concentrations measured in 132 olivine and 32 pyroxene phenocrysts (ranging in size from 0.2 to 2 mm and from basalts in different geological settings) display mean $^4$He concentrations of $3 \times 10^{11}$ at g$^{-1}$ (median of $8 \times 10^{10}$ at g$^{-1}$) for both olivine and pyroxene. Note, however, that some samples may be very rich in $^4$He, reaching concentrations above $10^{12}$ at g$^{-1}$, implying a significant intersample variability (standard deviation $> 6 \times 10^{10}$ at g$^{-1}$; Fig. 5C, Table S1). Such variability may also be observed for aliquots of hundreds of grains, even if all grains originate from the same sample. This significant intersample variability can be described as a “nugget effect” that is probably controlled by the abundance and repartitioning of fluid and melt inclusions, which are generally rich in magmatic gas and vary in their abundance in phenocrysts from one geological context to another (e.g. Puchol et al., 2017). However, this dataset is not large enough to establish a systematic or to understand why some volcanic fields present phenocrysts with higher magmatic helium contents than others (Table S1). In any case, an efficient means of minimizing the uncertainty from the magmatic He correction is to select phenocrysts with low magmatic helium concentrations, or to find methods for reducing this initial concentration (see section 3.2.2).

### 3.2.2. Using grains sized 100–500 µm reduces the magmatic helium contribution

Observations show that phenocryst size is a first-order control on helium concentrations, all other things being equal (Puchol et al., 2017). Furthermore, progressive phenocryst crushing experiments have demonstrated that laboratory (Williams et al., 2005) or natural crushing (e.g., by erosion; Puchol et al., 2007) may reduce the magmatic He content by releasing the magmatic component that is preferentially hosted in melt and fluid inclusions. Hence, crushing phenocrysts (in vacuo or in air) before fusing the finest fraction may appear to be a good strategy, because it purges magmatic helium and thus reduces the uncertainty on the computed cosmogenic $^3$He concentration. Although it is sometimes misleading to propose a universal grain size cutoff, experiments have shown that analyzing phenocrysts smaller than 500 µm is generally efficient in significantly reducing the magmatic helium component and intergrain variability (Puchol et al., 2017; Williams et al., 2005).
Reducing the granulometry of the analyzed phenocrysts should however be performed with caution: several studies have reported the suspected loss of matrix-hosted $^3$He during in-vacuo crushing for longer than 1 min (Blard et al., 2006; Hilton et al., 1993; Yokochi et al., 2005) and, more worrisome, Protin et al., (2016) recently demonstrated that significant amounts of atmospheric helium may be adsorbed onto the newly created surfaces when crushing silicates in atmospheric conditions. This effect must be considered seriously because experimental data indicate that the high adsorption energy of atmospheric helium requires heating above 1000 °C to release the contamination, which affects cosmogenic $^3$He (Protin et al., 2016). Because atmospheric helium is adsorbed on surfaces, the magnitude of the contamination is inversely proportional to grain size. Experimental data from olivines show that contamination remains insignificant (compared to matrix-hosted helium varieties) for grains larger than 100 µm (Protin et al., 2016). As both the loss of matrix-hosted cosmogenic $^3$He and unexpected atmospheric contamination result in underestimated cosmogenic $^3$He concentrations, grains smaller than 100 µm should be avoided in cosmogenic $^3$He analyses. Consequently, combining the need to reduce the magmatic helium component while avoiding any atmospheric helium contamination defines the ideal granulometric window for cosmogenic $^3$He analyses: the fusion of minerals ranging in size from 100 to 500 µm should be favored, regardless of whether they were previously crushed (Fig. 5).

Another promising and innovative technique recently tested by Hofmann et al. (2021) is to use X-ray micro-computed tomography (µCT) to identify grains that are free of melt/fluid inclusions and thus have much lower magmatic $^3$He concentrations. Finally, an alternative to crushing is to melt several aliquots from the same sample and build $^3$He vs $^4$He isochrons (section 3.5.3). Using this method, one can distinguish the cosmogenic and magmatic $^3$He components without vacuum crushing, avoiding all the above-mentioned potential issues.

### 3.3. Radiogenic $^4$He

Radiogenic $^4$He is produced by the decay of radioactive isotopes (such as uranium and thorium) present in the mineral of interest (e.g. olivine, pyroxene). It is important to properly estimate the $^4$He* concentration to accurately estimate the magmatic $^4$He and $^3$He contributions (Blard and Farley, 2008). In many settings, radiogenic $^4$He concentrations are indeed non-negligible and may be of the same order of magnitude as the magmatic $^4$He component. In extreme cases, the radiogenic $^4$He concentration may even be several orders of magnitude...
greater than the magmatic $^4$He concentration, requiring specific approaches to estimate the magmatic $^3$He component (e.g. Martin et al., 2018; Balter-Kennedy et al., 2020).

The radiogenic $^4$He concentration, which is constrained by the mineral’s closure age $t_c$ and the chemical composition of the rock, can hence be used to determine the most suitable method and analytical strategies for performing non-cosmogenic $^3$He corrections (see section 3.5).

### 3.3.1. The production of radiogenic $^4$He: general principles and equations

Radiogenic $^4$He is produced by the $\alpha$-decay of $^{235}$U, $^{238}$U, and $^{232}$Th when these elements are present in a studied mineral and its host lithology. $^{147}$Sm and $^{148}$Sm also release alpha particles through radioactive decay, but this contribution is generally negligible in silicates (Zeitler, 2014). $^4$He* production through time is described as:

$$^4\text{He}^* = \int_0^{t_c} P_4 dt,$$  (7)

where $t_c$ is the helium closure age (i.e., the eruption age in the case of volcanic rocks) and $P_4$ is the production rate of $^4\text{He}^*$. If $t_c$ is long enough to have reached secular equilibrium, Equation (7) simplifies to:

$$^4\text{He}^* = P_4 \times t_c.$$  (8)

Note that, in the case of rocks with young crystallization ages (<500 ka), the assumption of secular equilibrium may not be valid, and disequilibrium must be considered when calculating $P_4$ (Aciego et al., 2007; Farley et al., 2002). That said, even for lavas younger than 500 ka and when using the maximum $^{230}$Th/$^{238}$U fractionation reported for pyroxenes and olivines (Aciego et al., 2007; Wood et al., 1999), the obtained value of $P_4$ differs from that obtained under the secular equilibrium assumption by <5% (Farley et al., 2002).

To avoid circular reasoning and the difficulty of estimating the magmatic $^4$He contribution, $t_c$ should be estimated using a different radiochronometer than (U-Th)/$^4$He*, such as K-Ar or $^{40}$Ar/$^{39}$Ar; this is often feasible for lavas because the crystallization age equals the helium closure age. However, in the case of plutonic or metamorphic rocks, in which minerals generally have helium closure ages younger than the crystallization age, (U-Th)/$^4$He* dating is a necessary alternative. In those cases, this should be done using minerals with large (U-
Th)\(^4\)He\(_{\text{mag}}\) ratios to avoid bias and uncertainties in estimating \(^4\)He\(^*\) that result from the magmatic \(^4\)He component.

Since the ejection distance of alpha particles is \(~20\) \(\mu\)m in rocks (Ziegler 1977), some radiogenic \(^4\)He produced in the surrounding matrix is also implanted in minerals (Lal 1989). Conversely, a fraction of the in-situ \(^4\)He\(^*\) may also be ejected from minerals. Thus, one must consider both ejection and implantation when computing \(P_4\). Based on a series development formula, \(P_4\) can be calculated as (Farley et al., 2006):

\[
P_4 = I_4 \left[ 1 - 1.5\left( S/D \right) + 0.5\left( S/D \right)^3 \right] + M_4 \left[ 1.5\left( S/D \right) - 0.5\left( S/D \right)^3 \right],
\]

(9)

where \(I_4\) and \(M_4\) are the \(^4\)He\(^*\) production rates in the mineral of interest and in the surrounding rock, respectively, \(S\) (\(\mu\)m) is the stopping distance of alpha particles (\(~20\) \(\mu\)m), \(D\) (\(\mu\)m) is the crystal diameter, and Equation (9) is thus only valid for minerals having a spherical geometry; for non-spherical phenocrysts, it is necessary to compute an equivalent sphere diameter (Farley and Stockli 2002).

If secular equilibrium is reached, \(I_4\) and \(M_4\) follow (e.g., Wolf et al., 1998):

\[
I_4 \text{ (or } M_4\text{)} = 8\lambda_{238}[^{238}\text{U}] + 7\lambda_{235}[^{235}\text{U}] + 6\lambda_{232}[^{232}\text{Th}],
\]

(10)

where \([^{238}\text{U}], [^{235}\text{U}],\) and \([^{232}\text{Th}]\) are the isotopic concentrations measured in the mineral (for \(I_4\)) or the whole rock (for \(M_4\)) and \(\lambda_{238}, \lambda_{235},\) and \(\lambda_{232}\) are their respective decay constants.

### 3.3.2. Removing the implanted \(^4\)He\(^*\) component

The proportion of in-situ \(^4\)He\(^*\) relative to the implanted component is controlled by phenocryst size and by the relative U and Th concentrations in the phenocrysts and their host rock, i.e., the effective mineral/melt partition coefficients for these elements. Measurements of U and Th in mineral separates and in whole rocks show that these partition coefficients vary greatly across lithologies and geological settings (e.g. Blard and Farley, 2008) and are strongly dependent on the abundance of U-Th-rich melt inclusions. They should thus be called “apparent partition coefficients”. The presence of U-Th-rich melt inclusions may also induce significant spatial heterogeneities in the mineral of interest, implying that spot analyses by secondary ion mass spectrometry could be misleading. Bulk U-Th analyses after mineral dissolution are thus preferable because they represent the mean U-Th concentration.
Correcting for implantation and ejection of $^4$He* may also be complicated by several issues, and thus be a source of significant uncertainties. This is particularly true for detrital minerals in river sediments: because these minerals may have been fragmented, their initial sizes are generally unknown, as well as the exact U and Th concentrations of their source rocks. Moreover, in the case of non-spherical grains, shape estimates and the conversion to equivalent sphere diameter may also induce significant uncertainties.

Removing the external portion (at least 20 µm) of phenocrysts should therefore be a systematic sample preparation step because it obviates the need to correct for implanted and ejected components (Aciego et al., 2007; Blackburn et al., 2007; Bromley et al., 2014; Gayer et al., 2008; Min et al., 2006). Using this method, geometric corrections are unnecessary and the radiogenic $^4$He production rate equals the in-situ $^4$He* production rate, simplifying Equation (9) to $P_4 = I_4$; this has the potential to reduce the uncertainties associated with calculating the $^4$He* component. For olivine and pyroxene, Bromley et al., (2014) proposed a chemical treatment in which minerals are leached for 2 hours in a 3% HF:10% HCl acid solution in an ultrasonic bath at 40 °C. Physical techniques, such as over-pressurized air abrasion in closed chambers, also efficiently remove the external $^4$He*-enriched rim (Aleinikoff et al., 1990).

If it is not possible to remove the external rim affected by alpha particle mobility, it is of primary importance to measure the U and Th concentrations in both the phenocrysts and their host rock, especially in samples with large closure/exposure age ratios.

### 3.3.3. Uncertainty arising from the radiogenic $^4$He correction

To show the importance of accurately and precisely estimating the radiogenic $^4$He concentration, I modelled the total relative uncertainty on the cosmogenic $^3$He concentration for different exposure ages, closure age scenarios (10 ka to 1 Ga), and different elevations in Figure 6. In this model, the uncertainty on cosmogenic $^3$He is computed from a Taylor series expansion as:

$$
\sigma_{^3\text{He}}^2 = \sigma_{^3\text{He}}^2 + \left[ \sigma_{^4\text{He}_{\text{mag}}} \cdot \left( \frac{^3\text{He}}{^4\text{He}} \right)_{\text{mag}} \right]^2 + \left[ \sigma_{^4\text{He}^*} \cdot \left( \frac{^3\text{He}}{^4\text{He}} \right)_{\text{mag}} \right]^2 + \left[ \left( ^4\text{He}_{\text{tot}} - ^4\text{He}^* \right) \cdot \sigma_{3/4} \right]^2.
$$

(11)
The model results demonstrate that the radiogenic $^4$He correction may induce a significant uncertainty on the final estimated cosmogenic $^3$He concentration, with the error increasing with increasing closure/exposure age ratio and magmatic $^4$He concentration. For helium closure ages older than 10 Ma and exposure ages younger than 100 ka, the total relative uncertainty on $^3$He$_c$ is larger than 5% and may even reach 100% for exposure ages younger than 10 ka (Fig. 6). In these extreme scenarios with 100% uncertainty, $^3$He$_c$ is below the detection limit. The precision and detection limit of $^3$He$_c$ determination can hence be improved by selecting samples with the youngest (U-Th)/$^4$He closure age, when possible, and by reducing the magmatic He component by preferentially melting grains in the 100–500 µm size range or by identifying inclusion free minerals by µCT (Hofmann et al., 2021).
Figure 6. Modeling the impact of the radiogenic $^4$He correction - described by the $(U$-$Th)/^4$He* cooling age - on the total relative uncertainty ($1\sigma$) on the measured cosmogenic $^3$He concentration for variable exposure durations. Two magmatic $^4$He concentrations are considered: $10^{10}$ and $10^{11}$ at g$^{-1}$ (dashed and solid curves, respectively). Because the $^3$He production rate changes with elevation, this simulation considers samples exposed at (A) 2,000 m elevation and (B) sea level. This calculation applies Equation (11) assuming a magmatic $^3$He/$^4$He ratio of 8 Ra with 20% uncertainty, an uncertainty of 3% on the total amount of $^3$He extracted by fusion, an uncertainty of 1% on the magmatic $^4$He concentration, and an uncertainty of 10% on the estimated radiogenic $^4$He concentration.
3.4. Nucleogenic $^3$He

3.4.1. The production of nucleogenic $^3$He: general principles and equations

In minerals, nucleogenic $^3$He is produced by the capture of low-energy neutrons by $^6$Li through the reaction (Andrews 1985; Andrews and Kay 1982; Dunai et al., 2007):

$$^6\text{Li}(n, \alpha)\ ^3\text{H} \xrightarrow{\beta \text{ decay}} ^3\text{He} .$$

(12)

Neutrons involved in this reaction are “thermal”, “epithermal”, and “slow” neutrons with energies ranging between 0.025 and 300 eV, far below those of the cosmogenic “fast” neutrons (>1 MeV) involved in the main cosmogenic $^3$He production pathway (Dunai, 2010; Gosse and Phillips, 2001). Some of this low-energy neutron flux into rocks originates from nuclear reactions between cosmic rays and the terrestrial environment (Lal, 1987). However, $^3$He production through the capture of cosmogenic thermal neutrons is significantly lower than that by cosmogenic spallation, at least in the case of minerals with typical Li concentrations of 2–50 ppm (Dunai et al., 2007).

Low-energy neutrons involved in the production of $^3$He through capture by $^6$Li are also produced by ($\alpha$,n) reactions (Andrews, 1985; Andrews and Kay, 1982) involving elements within rocks and alpha particles emitted by the natural decay of U and T. The total production rate $P_{\text{nuc}}$ of nucleogenic $^3$He thus depends on the Li concentration in the mineral of interest and the bulk U and Th concentrations and chemical composition of the whole rock within ~1 m of the mineral. In a given mineral, $P_{\text{nuc}}$ can be calculated as (modified from Andrews, 1985):

$$P_{\text{nuc}} = f_n \left[ \sigma_{\text{Li}} C_{\text{Li}} \right]_{\text{mineral}} \sum_i \frac{\sigma_i C_i}{[\sum_i \sigma_i C_i]_{\text{rock}}} .$$

(13)

where $f_n$ is the average thermal neutron flux originating from the bulk rock, $\sigma_i$ is the neutron capture cross section specific to element $i$, and $C_i$ is the concentration of element $i$ (in mol g$^{-1}$ or atomic percent). To accurately compute $P_{\text{nuc}}$, it is thus important to measure the Li concentration in the mineral of interest. According to Andrews and Kay (1982), the whole-rock concentrations of 17 elements ($k = 17$) are also needed to calculate the denominator of Equation (13), $[\sum_i \sigma_i C_i]_{\text{rock}}$: nine major elements (Si, Al, Fe, Mn, Mg, Ca, Na, K, Ti) and eight trace elements with cross sections larger than 1 barn (Li, B, Co, Cr, Ni, Gd, Sm, Cl).
The whole-rock concentrations of the two main $\alpha$-emitters, U and Th, also need to be
determined. The $(\alpha,n)$ reactions occurring in rock indeed determine $f_n$, which is computed as
(Andrews and Kay, 1982):
\[
\begin{align*}
    f_n &= \frac{U}{100} \cdot (13.8 \cdot Na + 5.4 \cdot Mg + 5.0 \cdot Al + 1.31 \cdot Si + 0.6 \cdot Ca) + \frac{Th}{100} \cdot (6 \cdot Na + 2.45 \cdot Mg + 2.55 \cdot Al + 0.56 \cdot Si + 0.25 \cdot Ca) + 0.4764 \cdot U, \\
\end{align*}
\]
where U and Th concentrations are in ppm ($10^{-6} \text{ g g}^{-1}$) and the whole-rock major elements
concentrations are in atomic percent (%). Importantly, all these elemental concentrations must
be measured in a rock sample that is representative of the bulk rock. Indeed, the absorption
paths of low-energy neutrons produced through $(\alpha,n)$ reactions in rocks typically range between
50 and 80 cm, implying that the radiogenic thermal neutron flux is spatially averaged and rather
homogenous within a monolithologic unit (Dunai et al., 2007; Lal, 1987).

An Excel spreadsheet is available in the Supplementary Information (Table S2,
https://doi.org/10.24396/ORDAR-90) to compute $P_{nuc}$ and the total $^3\text{He}_{nuc}$ concentration when
the closure age is known. As inputs, this spreadsheet requires the 19 whole-rock major and trace
element concentrations involved in Equation (13), as well as the Li concentration measured in
the mineral of interest.

3.4.2. Uncertainty arising from the nucleogenic $^3\text{He}$ correction

The concentration of nucleogenic $^3\text{He}$ in a mineral is thus mainly controlled by the
chemical composition of the rock and the helium closure age. Figure 7A shows modeled
nucleogenic $^3\text{He}$ concentrations vs helium closure age for various Li contents and lithologies
(granite, andesite, basalt). Compared to basalt phenocrysts, the build-up of $^3\text{He}_{nuc}$ may be
several order of magnitudes higher in Li-rich minerals (e.g. amphiboles) hosted in U-rich
granite (Fig. 7A).

To assess the impact of correcting for $^3\text{He}_{nuc}$ on the precision of the measured
cosmogenic $^3\text{He}$ concentration, I modeled the relative uncertainty arising from the nucleogenic
$^3\text{He}$ correction as a function of helium closure age for variable exposure durations (1 ka, 10 ka,
100 ka, 1 Ma), at 2,000 m elevation or sea level, in the case of a mineral containing 10 ppm Li
and hosted in an andesite (Fig. 7B, C). This model shows that for helium closure ages younger
than 1 Ma, the uncertainty arising from the nucleogenic $^3\text{He}$ correction remains below 20%,
even for exposure ages as young as $\sim$1 ka (Fig. 7B, C). However, for longer helium closure
Whenver possible, it is thus important to select rock samples with the largest possible exposure/closure age ratio to improve the precision on the measured cosmogenic $^3$He concentration.

Figure 7. (A) Nucleogenic $^3$He concentrations vs helium closure age for various mineral Li contents and lithologies (granite, andesite, basalt). (B, C) Uncertainty on the cosmogenic $^3$He concentration due to the nucleogenic $^3$He correction for variable closure ages, at 2,000 m elevation, and sea level, respectively. This modeling assumes a mineral containing 10 ppm Li in a rock of average andesitic composition with bulk U and Th contents of 10 ppm and 30 ppm, respectively, corresponding to $P_{\text{nuc}} = 0.25$ at g$^{-1}$ yr$^{-1}$ (shown in blue in (A)). This model assumes that the nucleogenic production rate is known with a relative uncertainty of 10%.
3.5. Strategies to estimate the non-cosmogenic $^{3}$He components

Correcting for non-cosmogenic $^{3}$He components is key to achieving low detection limits and final uncertainties on cosmogenic $^{3}$He concentrations. Several characteristics of a rock’s geological history must be carefully evaluated for each sample, notably the helium closure age, the amount of magmatic helium in the minerals, and the concentrations of U, Th, and Li in the minerals and their host rocks. Depending on the thermal history and chemical composition of a rock sample, the minimum measurable surface exposure age may vary by several orders of magnitude, from $10^2$ to $>10^6$ years (Figs. 5–7). Hence, when background geological information is available (e.g. from geological maps), field sampling should first target rocks with the youngest cooling history, and select minerals/rocks with the lowest U, Th and Li concentrations, such as olivine and pyroxenes in basalts or andesites.

Different analytical strategies and procedures can be used to correct for the non-cosmogenic $^{3}$He components, and the choice of the best strategy mainly depends on the specific characteristics of a given sample. Figure 8 is an easy-to-use flowchart that summarizes the analytical protocol, from bulk sample crushing to final cosmogenic $^{3}$He calculation, for various sample cases.
Flowchart for cosmogenic \(^3\text{He}\) measurements

1 – Isolation of minerals (olivines, pyroxenes)
- Save a 3 cm x 3 cm piece
- Crush bulk rock
- Sieve: 100-500 µm and 500-1000 µm
- Density and magnetic separation
- Quality check: handpicking under a microscope to remove grains with adhering lava, oxides, clays

2 – Removal of 20 µm external rim
to remove the implanted \(^4\text{He}\)
- Chemical: HF 3% - HCl 5% for 2 hours (20 mL/g) at 40 °C
- HCl 40% rinsing ×3 - H₂O rinsing ×3 - Drying
- Physical: abrasion with compressed air
- Microscope quality check: remove newly formed fluorides

\(^{3}\text{He}_{\text{tot}} = ^{3}\text{He}_{\text{mag}} + ^{3}\text{He}_{\text{eu}}\)

\(^{4}\text{He}_{\text{tot}} = ^{4}\text{He}_{\text{mag}} + ^{4}\text{He}_{\text{eu}}\times (^{4}\text{He}_{\text{mag}} / ^{3}\text{He}_{\text{mag}}) + f(U_{\text{tot}} Th_{\text{tot}} U_{\text{mag}} t_{\text{c}})\)

\(^{4}\text{He}_{\text{tot}} = ^{4}\text{He}_{\text{mag}} + ^{4}\text{He}_{\text{eu}} + f(U_{\text{mag}} Th_{\text{mag}} t_{\text{c}})\)

Measurement Calculation

Thin section: petrographic characteristics (optional)

Major and trace elements in rock
- Li, U-Th-Sm
- Thermal neutron flux in the rock
- Major and trace elements in the mineral
- U-Th-Sm, Li

\(P_{\text{nuc}} = P_{4}\times (^{4}\text{He}_{\text{nuc}} / ^{3}\text{He}_{\text{mag}})\)

\(^{3}\text{He}_{\text{c}} = b + ^{4}\text{He} \times a - ^{3}\text{He}_{\text{nuc}}\)

\(t_{\text{c}}\) is unnecessary

* Particular case of uneroded lava flow:

\(^{3}\text{He}_{\text{c}} = \frac{b}{R}\times (^{3}\text{He}_{\text{mag}} / ^{4}\text{He}_{\text{mag}})\)

Isochrons

**Before any He extraction in a furnace, weigh the samples and wrap in metal foil.**
Figure 8. Flowchart presenting up-to-date protocols for measuring cosmogenic $^3$He, including the estimation of radiogenic $^4$He and nucleogenic $^3$He, which requires independent knowledge of the helium closure age of a rock, $t_c$, except in the special case of uneroded lava flows. See text for detailed equations, notably those for calculating the radiogenic $^4$He production rate, $P_4$ (Equations 9-10) and the nucleogenic $^3$He production rate, $P_{nuc}$ (Equations 13-14). See section 3.5.2 for more details about the R-factor and section 3.5.3 for details about the isochron method.

3.5.1. Case 1, the general situation: $^4$He closure age > exposure age, but $^4$He* is not significantly larger than $^4$He$_{mag}$

For typical U and Th concentrations in olivine and pyroxene (1-100 ppb), the common case in which $^4$He* and magmatic $^4$He concentrations are of comparable orders of magnitudes (i.e., $0 < ^4$He*/$^4$He$_{tot} < 1$) is encountered in minerals with helium closure ages younger than a few tens of millions of years (Blard and Farley, 2008). In such cases, an efficient strategy is to compute $^4$He* using Equations (7)–(10); the magmatic He correction can then be performed accurately using a modified version of Equation (1) (Blard and Farley, 2008):

$$3^\text{He}_c = 3^\text{He}_{tot} - (4^\text{He}_{tot} - 4^\text{He}^*) \times (4^\text{He}/3^\text{He})_{mag} - 3^\text{He}_{nuc}. \quad (15)$$

In this case, the total uncertainty on cosmogenic $^3$He can be computed applying Equation (11).

3.5.2. Case 2: Samples with similar closure and exposure ages – the R-factor

Although rare, the condition that $t_c = t_e$ applies to the case of uneroded lava flows. In such cases, the accumulation of radiogenic $^4$He and cosmogenic $^3$He in the minerals of interest (olivine, pyroxenes) starts synchronously when a lava flow cools at the Earth’s surface (Blard and Farley, 2008). Because $t_c = t_e$:

$$4^\text{He}^* = \frac{P_4}{p_3} \times 3^\text{He}_c. \quad (16)$$

Then, using Equation (16) to substitute $^4$He* into Equation (15), and making the reasonable assumption that $^3$He$_{nuc}$ is negligible in this case, isolating $^3$He$_c$ on the left-hand side gives:

$$3^\text{He}_c = \frac{3^\text{He}_{tot} - 4^\text{He}_{tot} \times (4^\text{He}/3^\text{He})_{mag}}{R}, \quad (17)$$

where
Equation (18) defines the so-called \( R \)-factor (“\( R \)” for radiogenic; Blard and Farley, 2008; Blard and Pik, 2008). This approach has the advantage of including the full impact of the radiogenic \( ^4\text{He} \) correction on the calculation of the cosmogenic \( ^3\text{He} \) concentration without requiring any independent estimate on the helium closure age of the rock.

Calculating \( R \) requires only the local time-integrated production rate \( P_3 \), the \( ^4\text{He}^* \) production rate \( P_4 \), and the magmatic \( ^3\text{He}/^4\text{He} \) ratio. The lower the value of \( R \), the greater the correction arising from radiogenic \( ^4\text{He} \). When \( R \approx 1 \), the radiogenic correction is negligible compared to the production of cosmogenic \( ^3\text{He} \), i.e., \( \frac{P_4}{P_3} \cdot \left( \frac{^3\text{He}}{^4\text{He}} \right)_{\text{mag}} \ll 1 \), and the magmatic correction simplifies to Equation (1). However, this situation is rare in nature; modelling \( R \) for different exposure altitudes and U and Th concentrations shows that \( R \) is generally < 1 (Fig. 9).

Importantly, in the case of lava flows used to calibrate the cosmogenic \( ^3\text{He} \) production rate, most actual \( R \)-factors are between 0.80 and 0.95 (Blard and Farley, 2008), meaning that neglecting the radiogenic \( ^4\text{He} \) correction would translate to underestimating \( ^3\text{He} \) by 5–20%.

Because some earlier studies did not perform this radiogenic \( ^4\text{He} \) correction, we have recomputed all reported \( R \)-factors in the updated online production rate database ICE-D (http://calibration.ice-d.org/), which is used as a reference by the CREp calculator (https://crep.otelo.univ-lorraine.fr/#/) to compute cosmogenic \( ^3\text{He} \) exposure ages (Martin et al., 2017).
Figure 9. Modeled values of $R$-factor for uneroded lava flows as a function of U concentration in the phenocrysts and exposure elevation. Here, $R$ is computed for a Th/U ratio of 3 and a magmatic $^{3}$He/$^{4}$He ratio of 8 Ra. Implantation of $^{4}$He* from surrounding lava is neglected. The equations of Stone (2000) were used to scale the sea-level $^{3}$He$_{c}$ production rate of 124 at g$^{-1}$ yr$^{-1}$ (Martin et al., 2017). This simulation shows that $R$ is strongly dependent on the altitude at which a lava flow is emplaced.

### 3.5.3. Isochrons (adequate for Cases 1 and 2)

An alternative method of estimating non-cosmogenic $^{3}$He components is to fuse several aliquots of the same mineral sample and build helium isochrons. The main advantage of this isochron method is to avoid the initial in vacuo crushing step. Measured $^{3}$He and $^{4}$He concentrations obtained by melting different aliquots (of ~0.5 g) with variable magmatic helium concentrations define straight lines in $^{3}$He$_{melt}$ $vs$ $^{4}$He$_{melt}$ space (this study) or ($^{3}$He/$^{4}$He)$_{melt}$ $vs$ $1/^{4}$He$_{melt}$ space (Blard and Pik, 2008; Cerling and Craig, 1994) (Fig. 10); these ‘melt’ concentrations represent total concentrations. The crushing step is thus unnecessary because the magmatic $^{3}$He/$^{4}$He ratio is directly defined by the linear regression of an isochron. Even though the isochron method is powerful for estimating the magmatic $^{3}$He contribution, note that this technique still requires the determination of potential contributions from radiogenic $^{4}$He (by measuring mineral and rock U and Th concentrations and the helium closure age; see section 3.3) and nucleogenic $^{3}$He components (by measuring bulk-rock major and trace elements and mineral Li concentrations; see section 3.4).
Below, I detail the basic equations and outcomes that can be derived from the isochron approach.

3.5.3.1. \( ^3\text{He} \) vs \( ^4\text{He} \) isochrons (this study). The general equation of the "\(^3\text{He} \) vs \(^4\text{He} \) isochron" is:

\[
\begin{align*}
^3\text{He}_{\text{tot}} &= ^4\text{He}_{\text{tot}} \times \left(\frac{^3\text{He}}{^4\text{He}}\right)_{\text{mag}} + \left[ ^3\text{He}_c + ^3\text{He}_{\text{nuc}} - ^4\text{He}^* \times \left(\frac{^3\text{He}}{^4\text{He}}\right)_{\text{mag}} \right]. \\
&= \left(\frac{^3\text{He}}{^4\text{He}}\right)_{\text{mag}} - ^4\text{He}^* \times \left(\frac{^3\text{He}}{^4\text{He}}\right)_{\text{mag}} + ^3\text{He}_c + ^3\text{He}_{\text{nuc}}.
\end{align*}
\] (19)

This corresponds to the equation of a line, \( y = ax + b \), in which the slope \( a \) is \( \left(\frac{^3\text{He}}{^4\text{He}}\right)_{\text{mag}} \) and the \( y \)-intercept \( b \) is the sum of all components that are independent of the magmatic component, including cosmogenic \(^3\text{He} \) (Fig. 10A). \(^3\text{He}_c \) can then be estimated after correcting for the nucleogenic \(^3\text{He} \) and radiogenic \(^4\text{He} \) components as:

\[
^3\text{He}_c = b + ^4\text{He}^* \times a - ^3\text{He}_{\text{nuc}}.
\] (20)

In the specific case of uneroded lava flows, the radiogenic \(^4\text{He} \) correction may be simplified using the \( R \)-factor (Eq. 18) to:

\[
^3\text{He}_c = \frac{b}{R}.
\] (21)
Figure 10. Cosmogenic $^3$He isochrons: (A) $^3$He vs $^4$He isochron (this study) and (B) $^3$He/$^4$He vs $^1$/$^4$He isochron (Blard and Pik, 2008; Cerling and Craig, 1994).
3.5.3.2. \(^{3}\text{He}/^{4}\text{He}\) vs \(^{1}/^{4}\text{He}\) isochrons (Blard and Pik, 2008; Cerling and Craig, 1994).

The general equation of the \(^{3}\text{He}/^{4}\text{He}\) vs \(^{1}/^{4}\text{He}\) isochron" is:

\[
\left( \frac{^{3}\text{He}}{^{4}\text{He}} \right)_{\text{tot}} = \left[ ^{3}\text{He}_c + ^{3}\text{He}_{\text{nuc}} - ^{4}\text{He}^* \times \left( \frac{^{3}\text{He}}{^{4}\text{He}} \right)_{\text{mag}} \right] \times \frac{1}{^{4}\text{He}_{\text{tot}}} + \left( \frac{^{3}\text{He}}{^{4}\text{He}} \right)_{\text{mag}}.
\] (22)

Again, this corresponds to the equation of a line, but in this case the \(y\)-intercept is \((^{3}\text{He}/^{4}\text{He})_{\text{mag}}\) and the slope is the sum of "magmatic-independent" components, including cosmogenic \(^{3}\text{He}\) (Fig. 10B). \(^{3}\text{He}_c\) can then be estimated after correcting for the nucleogenic \(^{3}\text{He}\) and radiogenic \(^{4}\text{He}\) components as:

\[
^{3}\text{He}_c = a + ^{4}\text{He}^* \times b - ^{3}\text{He}_{\text{nuc}}.
\] (23)

Using the R-factor in the specific case of uneroded lava flows, the radiogenic \(^{4}\text{He}\) correction may be simplified to:

\[
^{3}\text{He}_c = \frac{a}{R}.
\] (24)

Note that the mathematical regression of the "\(^{3}\text{He}/^{4}\text{He}\) vs \(^{1}/^{4}\text{He}\) isochron" requires computing the correlation factor \(\rho_{\text{err}}\) between the \((^{3}\text{He}/^{4}\text{He})_{\text{tot}}\) and \(^{1}/^{4}\text{He}_{\text{tot}}\) uncertainties:

\[
\rho_{\text{err}} = \frac{\sigma_{a}/^{4}\text{He}}{\sqrt{(\sigma_{a}/^{3}\text{He})^2 + (\sigma_{a}/^{4}\text{He})^2}}.
\] (25)

To achieve the best isochron regression, statistical parameters (MSWD, York, 1966) should be calculated using modern numerical tools, such as the online version of IsoplotR (http://isoplotr.es.ucl.ac.uk/; Vermeesch, 2018). Although, in its current version, IsoplotR does not include a specific tool for computing cosmogenic \(^{3}\text{He}\), this is easily done using the generic functions of "other x vs y regressions" in this online software. Even if both isochron types yield equivalent results, the "\(^{3}\text{He}/^{4}\text{He}\) vs \(^{1}/^{4}\text{He}\) isochron" is more suited to defining the magmatic \((^{3}\text{He}/^{4}\text{He})\) ratio, but also requires calculating the correlation factor \(\rho_{\text{err}}\).
3.5.4. Case 3: Samples with large amounts of radiogenic $^4$He* ($^4$He* ≫ $^4$He$_{\text{mag}}$)

In this case, the amount of $^4$He extracted by fusion is similar to, and indistinguishable from, the radiogenic $^4$He* concentration within analytical uncertainties. In other words, the magmatic $^4$He component is below the detection limit. This occurs for long closure ages (typically $>$10$^6$ yr) or in minerals with particularly high U-Th concentrations ($>$10 ppm), such as zircons. In such a situation, using the total amount of $^4$He extracted by fusion to estimate the concentrations of magmatic $^4$He and $^3$He is misleading, even when applying Equation (15).

This may induce major uncertainties, and indeed potential inaccuracies, on the final cosmogenic $^3$He concentrations (Fig. 6). However, it is not always straightforward to diagnose this extreme case, notably because, in practice, there exists a continuum between situations where $^4$He* ≈ $^4$He$_{\text{mag}}$ and those where $^4$He* ≫ $^4$He$_{\text{mag}}$. If the closure age and the U-Th concentrations of the mineral of interest are well known, one can determine if $^4$He* ≫ $^4$He$_{\text{mag}}$ by calculating the amount of $^4$He* and comparing it with the total amount of $^4$He extracted by fusion. Another way to pose this diagnostic is the impossibility of building isochrons: fused aliquots will present significant scatter in $^3$He$_{\text{tot}}$ vs $^4$He$_{\text{tot}}$ space and will not align on a straight line, displaying errorchrons. Since this case occurs for closure ages $>$1 Ma, the nucleogenic $^3$He contribution may be significant. This correction must therefore be considered carefully, using an accurate and precise estimate of the helium cooling age.

If it is established that $^4$He* ≫ $^4$He$_{\text{mag}}$, an alternative strategy must be employed to estimate the magmatic $^3$He correction. The most efficient alternatives are:

1) Find and fuse a sample of the same lithology that has been shielded from cosmic rays to estimate the magmatic and nucleogenic $^3$He components (e.g., Amidon et al., 2009). This assumes that the magmatic $^3$He component is homogeneous.

2) Prolonged in vacuo crushing (10 min, 100 strokes) of an aliquot of the same sample that has been dated by cosmogenic $^3$He. This allows the extraction of the majority of the magmatic $^3$He, which can be used to perform the magmatic $^3$He correction (Blard et al., 2009; Martin et al., 2018).

Note that these strategies may be sources of undetected inaccuracies, for example due to unrecognized magmatic or nucleogenic $^3$He components. Thus, when it is determined that $^4$He* ≫ $^4$He$_{\text{mag}}$, it is key to minimize the contribution of magmatic helium by selecting samples in the 100–500 µm size fraction (Puchol et al., 2017; Williams et al., 2005) or use prescreening methods to select inclusion free minerals (Hofmann et al., 2021).
4. Analytical methods for measuring $^3$He and $^4$He in minerals

4.1. Extraction procedure

As mentioned in sections 2.1 and 3.1, different extraction techniques must be used to selectively release the different helium components: measuring magmatic helium requires crushing in vacuo, whereas the complete extraction of cosmogenic $^3$He requires that minerals be heated at high temperatures and melted, using a high-vacuum, high-temperature furnace or a laser.

4.1.1. In-vacuo crushing

This method is suited to selectively releasing the magmatic component from minerals of interest (e.g. olivine, pyroxene) (Kurz, 1986a, 1986b). The several types and sizes of available crushers can accommodate different sample sizes with variable crushing efficiency (Blard et al., 2008; Scarsi, 2000). Typical crushers comprise ~50–100-g stainless steel cylinders that are moved in vacuo by external solenoids (Kurz, 1986a; Yokochi et al., 2005). The associated $^3$He and $^4$He blanks of $\sim 5 \times 10^3$ and $5 \times 10^8$ atoms per minute of crushing, respectively (Blard et al., 2008). Typical sample sizes are 0.5–1 g of solid materials; larger samples require a larger crusher to ensure crushing efficiency. If samples are too poor in magmatic helium to yield high signal/blank ratios, the isochron approach may be a useful alternative because it obviates the magmatic crushing step (e.g., Marchetti et al., 2020).

4.1.2. Heating furnace systems

High-temperature heating (1300-1500 °C for pyroxene, 1500-1800°C for olivine) is necessary to extract the entire cosmogenic $^3$He component present in samples, either by achieving total diffusion or melting. Double-vacuum resistance furnaces are ideal because they can be used to heat sample sizes ranging from 100 mg to 2 g (e.g., Amidon et al., 2009; Kurz, 1986b; Margerison et al., 2005). These apparatuses have typical $^4$He blanks in the range of $10^8$–$10^{10}$ atoms when heating at 1500 °C for 15 minutes (see review by Zimmermann et al., 2018).

For most analyzed concentrations, such amounts of phenocrysts (0.1–2 g) are necessary to ensure sufficiently high signal/blank ratios. It is important to note that all mineral samples must be weighed and wrapped in metal foils (e.g. Sn, Al, Cu) before being loaded into the furnaces. Recently, Zimmermann et al., (2018) developed and patented a new all-metal single-vacuum furnace whose crucible is heated by induction (Fig. 11). As the mass of heated metal is small compared to classical double-vacuum resistance furnaces, this design ensures better blanks ($^4$He
= \( (5 \pm 2) \times 10^8 \) atoms for 15 min at 1500 °C), faster heating and cooling rates (400 °C min\(^{-1}\)), improved robustness, lower functional costs, and, for models equipped with a thermocouple or thermal camera, improved temperature control.

**Figure 11.** Newly patented all-metal single-vacuum induction furnace (Zimmermann et al., 2018) allowing 1-g samples to be heated to 2000 °C in less than 5 minutes. Photo credit: ©Cryoscan/CNRS/Satt-Sayens

### 4.1.3. Lasers

Depending on their wavelength, lasers may either be used for ablation or heating. Although this extraction system is best suited for \( ^3 \)He-rich extraterrestrial materials (e.g., Megru, 1971), the utility of lasers is limited for terrestrial cosmogenic \( ^3 \)He. When they are used to directly heat minerals, laser beams can only heat a maximum of 10 mg of material (less in the case of ablation), with \( ^4 \)He blanks on the order of \( 10^8 \)–\( 10^9 \) atoms, implying lower signal/blank ratios compared to furnaces (Zimmermann et al., 2018). Note that BGC recently developed a laser “microfurnace” system, in which a sample of up to \( \sim 200 \) mg can be analyzed in a crimped tantalum tube heated with a diode laser (Balter-Kennedy et al., 2020).

When the uncertainty attached to the blank correction is propagated to the measured \( ^3 \)He and \( ^4 \)He concentrations, what matters is the absolute standard deviation on the blank level. Hence, to get an accurate estimate of the blank variance, it is important to run several blanks of an extraction apparatus. This must be done under strictly similar duration and temperature conditions to ensure that the blank variability is as low as possible.
4.2. Purification lines

After gas extraction by crushing or heating, it is necessary to remove the undesirable gas before analyzing the $^3$He and $^4$He concentrations. Pressures may indeed reach $10^{-3}$ mbar in hot furnaces, whereas most analyzers must be operated at pressures lower than $10^{-6}$ mbar to preserve their integrity and ensure accurate analysis. In most modern labs, this gas purification consists of sequentially releasing the sample in successive volumes of a stainless steel ultrahigh vacuum line, in which physical (e.g. charcoal cooled by liquid nitrogen at $-196 \, ^\circ$C) and chemical traps (e.g. SAES getters, titanium sponges) remove most hydrocarbons, CO$_2$, H$_2$O, H$_2$, and heavy noble gases (Xe, Kr, Ar) (e.g. Blard et al., 2015). Then, a cryogenic sorber (or “cryopump”) is often used in the final part of the line to trap He and Ne at temperatures <10 K, before being heated above 35 K to selectively release helium into the analyzer. This last purification step both removes neon and maximizes the helium signal.

4.3. Mass spectrometry

Mass spectrometers operated in static mode remain the most precise, accurate, and sensitive instruments for measuring helium isotopes in solid minerals. Since the invention of Nier-type sources (Fig. 12; De Laeter and Kurz, 2006), the main improvements have been in the design, materials, and detector sensitivity. State-of-the-art analytical instruments for measuring $^3$He and $^4$He abundances are stainless steel mass spectrometers with Nier-type sources, such as the Noblesse AN37 or the ThermoFisher SFT (“Split Flight Tube”; Fig. 12). The geometry of the SFT flight tube is designed to minimize the analytical volume (1400 cc) while simultaneously collecting $^4$He on a Faraday cup (with $10^{10}$–$10^{12}$ Ω amplifiers) and $^3$He on an electron multiplier. Whereas mono-collection mass spectrometers require peak jumping, this bi-collection of helium isotopes improves analytical stability and speed. The resolution on the electron multiplier is 700, ensuring the complete separation of $^3$He$^+$ from the isobars HD$^+$ and HHH$^+$. Sensitivities of the Thermo SFT and the Noblesse AN37 are comparable, in the range of 0.1–0.2 A/bar.
The most promising recent development in noble gas mass spectrometry is the creation of the compressed ion source, which can increase instrumental sensitivity for helium by one to two orders of magnitude (Baur, 1999; Matsumoto et al., 2010). For now, however, this advancement remains theoretical because, in practice, measurements are limited by the blank levels of the extraction system and, more importantly, by the presence of non-cosmogenic helium in samples. To fully exploit this progress in the sensitivity of noble gas mass spectrometers, parallel analytical developments must focus on reducing blanks and selecting minerals with the lowest possible non-cosmogenic $^3$He concentrations.

### 4.4. Interlaboratory comparisons

To identify potential sources of systematic and random uncertainties in $^3$He and $^4$He measurements, interlaboratory comparisons performed by analyzing splits of the same standard mineral material provide useful insights (Blard et al., 2015; Vermeesch et al., 2015). Recently, six international laboratories (GFZ, Potsdam; Caltech, Pasadena; CRPG, Nancy; SUERC, Glasgow; BGC, Berkeley; Lamont Doherty Earth Observatory, New York) participated in an intercomparison experiment (Blard et al., 2015), analyzing 5–22 aliquots of the CRONUS-P pyroxene standard, which has been shown to be homogeneous (Schaefer et al., 2016). Adding the 17 analyses recently performed at BGC (with the new “Ohio” system, Balter-Kennedy et al., 2020), this interlaboratory dataset (totaling 83 determinations from the six laboratories; Fig. 13) is characterized by a global weighted mean of $(5.02 \pm 0.05) \times 10^9$ at $^3$He g$^{-1}$ (1σ) with an overdispersion of 2.6% (1σ). $^4$He is characterized by a larger variability than $^3$He, with a global
weighted mean of $(3.60 \pm 0.09) \times 10^{13}$ at $^3$He g$^{-1}$ (1σ) with an overdispersion of 5.2% (1σ) (Blard et al., 2015). This interlaboratory overdispersion probably reflects systematic differences between the six laboratories, since the overdispersion is, by definition, larger than the analytical uncertainties. To further evaluate this and homogenize datasets, thereby improving the accuracy of cosmogenic $^3$He dating, all future studies should measure the CRONUS-P standard as part of their analyses. Potential sources of systematic uncertainties that may explain these previously overlooked interlaboratory offsets are discussed in the next subsection.

**Figure 13. Interlaboratory comparison of $^3$He concentrations measured in the CRONUS-P pyroxene standard (n = 83).** Note that I updated here the initial dataset of (Blard et al., 2015) to incorporate the data recently obtained by the “Ohio” analytical system of BGC (Balter-Kennedy et al., 2020). Plotted individual analytical uncertainties are at 1σ. For each lab are given the weighted means with their standards errors at 1σ (color horizontal bars show the 2σ range), the intra-lab overdispersion (OD, 1σ) and the MSWD. This statistical analysis was performed with the online tool IsoplotR (www.ucl.ac.uk), computing the weighted mean using a random effects model with two sources of uncertainty: the analytical uncertainty and an overdispersion term (Vermeesch 2018). Overdispersion describes the interlaboratory variance that is not included in the analytical uncertainties. Hence, computed overdispersion also depends on the analytical uncertainties reported by each lab. The standard error of the global weighted mean and the inter-lab overdispersion are given at 1σ (grey area show the 2σ range). Early measurements by (Niedermann et al., 2007; Schäfer et al., 1999) are also shown.
4.5. Potential sources of systematic uncertainties in $^3$He and $^4$He measurements

Various analytical issues may lead to potentially overlooked biases and contribute to the systematic differences in the $^3$He and $^4$He concentrations measured in different laboratories. Nonetheless, as some such biases have been recognized in some laboratories, discussing them openly and suggesting possible control benchmarks is a first step toward improving analytical precision and accuracy in all laboratories.

i) **Estimate of the initial pressure in gas standard tanks.** Nearly all laboratories use a gas standard as a primary reference to calibrate the sensitivities of their mass spectrometers. For this, it is vital to properly estimate the initial helium pressure (i.e., the number of $^3$He and $^4$He atoms) in the gas tank. This determination must be as accurate as possible, but depends on the non-linear behavior of capacitance manometers or the use of poorly calibrated volumes during dilution of the calibration gas in the reservoir. When a new gas standard is prepared, a useful experiment is to cross-check the measured initial pressure against a measurement of a mineral standard material, such as CRONUS-P.

ii) **Tank depletion after several standard iterations.** For daily calibration procedures, laboratories generally extract a small aliquot (<1 cc) from a large bottle of the gas standard (typically 2 L). Although the aliquot volume is small compared to the standard reservoir volume (ratio ~$10^{-4}$), it is essential to know the volumes of both the aliquot and the standard bottle to correctly calculate the amount of standard gas remaining in the bottle over time. See Fleck and Calvert (2014) for a complete discussion of the most accurate depletion equation. Modern automated extraction systems can use $>10^3$ standard aliquots per year; hence, a small initial bias on the aliquot/bottle volume ratio may be significantly amplified over several years of standard use. Thus, even if the gas standard remains the most convenient for daily calibrations, regularly cross-checking against a mineral standard material is again a useful and important control routine.

iii) **Volumetric calibration of the extraction and purification line.** Since pressure may reach high values ($>10^{-5}$ mbar) in furnaces operating at $>1000 \, ^\circ$C, it is often necessary to split the extracted gas to remove the gas fraction in the “dirty” part
of the purification line. Depending on the line design, this may imply that dilution routines differ between mineral samples and gas standards. Correcting for this dilution requires precise and accurate knowledge of the volumes involved. Extraction line volumes are generally determined with capacitance manometers at pressures between a few millibars and 1 bar. This measurement may be biased by adsorption onto the internal surfaces and thus overestimate the true volume. It is therefore particularly complicated to estimate volumes in which chemical and physical traps are present. Hence, when possible, splitting and dilution should be avoided or limited to a small fraction of the gas to reduce potential inaccuracies. Ideally, cryofocusing is useful for avoiding this issue and applying the same dilution to samples and standards.

iv) **Overly efficient and unknown He cryotrapping in the presence of heavier noble (Ne) or other gases.** In most laboratories, helium is trapped by a sorber head that is cooled to <10 K before being released into the last part of the extraction line at 35 K, the empirically established vacuum-release temperature for helium in most cryopump systems. This cryotrapping step is useful because it optimizes helium purification and separation from other noble gases. Because the cryopump is generally placed in the last part of the purification line, cryofocusing also maximizes the pressure of the gas being input into the mass spectrometer, improving the sensitivity of the entire analytical system. At CRPG, we tested the impact of the release temperature by measuring the helium signal obtained at various temperatures between 35 and 100 K (unpublished data): when analyzing pure helium gas, 35 K is sufficient to release 100% of the helium that was previously trapped on the cryopump sorber head. However, when analyzing helium extracted from silicates, 5% of the total helium remained trapped at 35 K, and heating to 70 K was necessary to release the totality of the sample’s helium. Hence, it is possible that the release temperature of helium from the cryopump sorbing step depends on the He/Ne ratio, or on the presence of other gases that were not removed in the upstream part of the purification line. These observations are not sufficient to define a clear relationship between the helium trapping and release temperatures and the ratio of He to other gases, and more attention should be given to this issue. New experimental data with
variable He/(other gas) ratios would be useful for improving the accuracy of He isotopic measurements.

v) **Pressure effects on the linearity of mass spectrometer sensitivity.** Empirical observations have shown that the amount of helium present in the mass spectrometer may modify instrumental sensitivity when measuring $^3$He and $^4$He (Fig. 14; Burnard and Farley, 2000; Mabry et al., 2013; Sano et al., 2008). This effect is more pronounced at low helium pressures ($<10^{-9}$ mbar) and may reach 20% in the worst situations (Sano et al., 2008; Zimmermann and Bekaert, 2020). The mechanisms producing this pressure dependence are not well constrained, but such instrumental fractionation could occur in a Nier-type source due to a space-charge effect on the ionization efficiency (Burnard and Farley 2000). Other possible mechanisms might involve the flight tube or the detectors (Sano et al., 2008). Importantly, this pressure effect varies significantly from one mass spectrometer to another (Sano et al., 2008) and is sensitive to the source settings, notably the trap current (Burnard and Farley, 2000; Mabry et al., 2013).

To minimize the potential bias arising from this pressure effect, source settings should be optimized to define the largest possible zone of stability (Fig. 14). Lower trap currents (50–200 µA) minimize this non-linearity. If it is not possible to suppress this instrumental fractionation, it is important to analyze similar amounts of gas standard and sample. Since this procedure often requires splitting gas standards, the volume of the preparation line must be accurately calibrated. Alternatively, this issue could be overcome by using He-poor gas standards: if a single pipette delivers a low-pressure helium signal in the range $10^{-11}$–$10^{-10}$ mbar, the dilution step will be unnecessary. Because the presence of other gases may also affect this instrumental bias (see above), it may be useful to ensure that samples and standards are of similar helium purity, notably He/Ne ratios. This can be achieved either by cryo-separation of neon and helium or by performing a quick scan of the amount of $^{20}$Ne in the sample using a quadrupole installed on the purification line. A neon spike may then be used to adjust the He/Ne ratio of the helium standard to be similar to that of the sample.
Figure 14. (Top) Measured $^3\text{He}/^4\text{He}$ ratios and (bottom) instrumental sensitivity ($A$ Torr$^{-1}$) as a function of the true helium pressure in the mass spectrometer. These data were obtained for a VG5400 with a source-trap current of 400 $\mu$A (Zimmermann and Bekaert, 2020); this effect is less pronounced on the Helix SFT (Sano et al., 2008). Note that source settings may affect this relationship because lower trap currents (<300 $\mu$A) reduce the impact of helium pressure on instrumental sensitivity.

vi) Variations of the absolute $^3\text{He}/^4\text{He}$ atmospheric ratio. Many gas standards are prepared using atmospheric helium that is artificially enriched in $^3\text{He}$ (e.g., Matsuda et al., 2002). Other standards are natural magmatic gases with $^3\text{He}/^4\text{He}$ ratios greater than that of the atmosphere (e.g. Blard et al., 2015). In either case, gas standards must be cross-calibrated against a gas of known composition to determine their absolute He contents and $^3\text{He}/^4\text{He}$ ratios. This is often performed against atmospheric helium, because the $^3\text{He}/^4\text{He}$ ratio in this reservoir is spatially homogeneous with a variability lower than 2‰ (95% confidence interval; Boucher et al., 2018). However, for absolute determinations of $^3\text{He}$, it is vital to know with accuracy the absolute value of the atmospheric $^3\text{He}/^4\text{He}$
ratio. The few studies that have attempted to measure the absolute $^3$He/$^4$He ratio in air (Clarke et al., 1976; Hoffman and Nier 1993; Lupton and Graham 1991; Mamyrin et al., 1970) yield a weighted average of $(1.386 \pm 0.005) \times 10^{-6}$ (1σ; Boucher et al., 2018). However, a few studies have reported values beyond this range; notably, Sano et al., (2008) analyzed air collected in the center of Tokyo (Japan) and obtained a ratio of $1.34 \times 10^{-6}$, ~3.5% lower than the world average, suggestive of a potential $^4$He enrichment by a local anthropogenic source. Moreover, temporal variations of the atmospheric $^3$He/$^4$He ratio are possible (Sano et al., 2010), but this effect is probably smaller than 1% in the Northern Hemisphere over the past 40 years (Mabry et al., 2015).

In summary, all these sources of systematic uncertainties may accumulate, leading in the worst cases to under-/over-estimates and interlaboratory discrepancies larger than 5%. To reduce these potential inaccuracies and improve interlaboratory comparisons, it is highly recommended that future studies analyze solid standard materials alongside each sample batch, such as the CRONUS-P pyroxene mineral standard (Schaefer et al., 2016; Blard et al., 2015).
5. Discussion: cosmogenic $^3$He production rates, the accuracy of spatial scaling, cross-calibration with other cosmogenic nuclides, online calculators and new frontiers

5.1. Production rates

Knowing cosmogenic $^3$He production rates accurately and precisely is important because this parameter is involved in all applications of cosmogenic $^3$He, notably in calculating surface exposure ages. Production rates are spatially variable (Gosse and Phillips 2001; Lal, 1991; Lal and Peters 1967) because the Earth’s magnetic field controls the flux and repartition of incoming primary charged cosmic particles (Dunai 2001; Lal, 1991; Lal and Peters 1967), and because atmospheric shielding results in elevation-dependent production rates (Hess 1912; Lal, 1991; Lal and Peters 1967; Pacini et al., 1911; Stone 2000). Hence, scaling models are necessary to calculate the production rate at a given sampling location by accounting for spatial characteristics (altitude, latitude, longitude) and exposure duration. Such scaling models were initially established from empirical relationships linking production rate to latitude, longitude, and altitude based on physical observations of the reactions triggered by cosmic particles (Balco et al., 2008). Recent scaling models based on ab-initio physical modeling efficiently describe the spatial variability of production rates (Argento et al., 2013; Lifton et al., 2014). However, geological calibration sites for cosmogenic nuclide production rates are necessary to ground truth and refine these models (Borchers et al., 2016; Martin et al., 2017). A production rate calibration site is a continuously exposed geomorphological surface that has gone uneroded and uncovered since its emplacement at the surface, and whose exposure age can be determined independently with accuracy and precision (e.g., by $^{14}$C, K-Ar, or luminescence dating). If a scaling model is accurate, all calibration sites should yield the same normalized production rate, within uncertainties, after the dataset is scaled to sea level and high latitude (>60°; SLHL).

5.2. The global $^3$He production rate calibration database

Figure 15A shows the locations of the 24 production rate calibration sites available in the literature (Ackert et al., 2003; Amidon and Farley 2011; Blard et al., 2013; Blard et al., 2006; Borchers et al., 2016; Cerling and Craig, 1994; Delunel et al., 2016; Dunai and Wijbrans 2000; Eaves et al., 2016; Fenton et al., 2009; 2013; Foeken et al., 2012; Goehring et al., 2010; Licciardi et al., 2006; Licciardi et al., 1999). The SLHL $^3$He production rates presented in Figure 15B–E were computed from this empirical dataset using the CREp calculator (https://crep.otelo.univ-lorraine.fr/#/). Whereas the compilation of Borchers et al., (2016) only considers a limited subset of the data, this up-to-date synthesis integrates all published raw
calibration data from the 24 calibration sites, which are available in the ICE-D online database (http://calibration.ice-d.org/). Contrary to the compilation of (Goehring et al., 2018), this updated synthesis also ensures that all \(^3\)He concentrations are corrected for radiogenic \(^4\)He, which amounts to a 1–5% correction at a few sites (Blard and Farley, 2008). The SLHL and time corrections presented in Figure 15 were performed with the Lifton-Sato-Dunai (LSD) scaling scheme (Lifton et al., 2014), the ERA-40 atmospheric model, and the Virtual Dipole Moment (VDM) dataset, derived from atmospheric \(^10\)Be (Muscheler et al., 2005). This analysis shows that there is no clear correlation between SLHL \(P_3\) and the latitudes, altitudes, or exposure durations of these calibration surfaces, suggesting that this time-dependent scaling model is accurate for the majority of the calibrated production rates (Balco et al., 2008). This dataset defines a worldwide average weighted mean of 124 ± 11 at \(g \cdot yr^{-1}\) with MSWD = 5 (Fig. 15B). For comparison, the Lal-Stone time-dependent scheme (Balco et al., 2008) yields a larger dispersion (MSWD = 7), suggesting that the LSD model performs slightly better, at least for \(^3\)He (Borchers et al., 2016; Martin et al., 2017). When plotted against exposure duration, SLHL \(P_3\) values scaled with the Lal-Stone model present a slight negative correlation (i.e., lower \(P_3\) values for longer exposure times >100 ka); this is not the case with the LSD scaling.

Tests of the different atmospheric models have shown that the spatially distributed ERA-40 atmospheric grid (Uppala et al., 2005) performs better than the standard atmosphere model (Stone 2000) in reducing the scatter on SLHL \(P_3\) (Martin et al., 2017). Similarly, the VDM database (Muscheler et al., 2005) yields the lowest dispersion (Martin et al., 2017). However, local inaccuracies persist and require further investigation and additional data and calibration sites: 4 of the 24 calibration sites yield SLHL \(P_3\) values stand beyond 2 weighted standard deviations. For example, in Western Africa, a calibration site on Fogo Island (Cabo Verde) yielded a SLHL \(P_3\) value that is particularly low, 2\(\sigma\) below the weighted average (Foeken et al., 2012). Future calibration studies in these regions will indicate whether this outlier is due to an improper independent K-Ar date or unrecognized erosion, or whether it results from an as yet unrecognized local anomaly such as high atmospheric pressure or a specific magnetic field anomaly (Lifton et al., 2014). In contrast, the Hawaiian calibration site yields a SLHL \(P_3\) more than 2\(\sigma\) above the global average, suggesting either inaccuracy in the independent dating or in the atmospheric or temporal correction.

To reduce potential inaccuracies in the spatial scaling, local calibration sites should generally be favored over the use of a global average, particularly in low-latitude, high-altitude sites. A local calibration site should ideally be as close as possible to the site to be dated. When the horizontal/vertical distance between the object to be dated and the calibration site is less
than 1,000 km/1,000 m, discrepancies between scaling models remain within analytical uncertainties (Martin et al., 2018). However, in practice, many regions do not have any $^{3}$He calibration sites (Fig. 15A), hindering the use of a local calibration to compute exposure ages. The case of the High Tropics is particularly important, since the altitude and magnetic field corrections are extreme in these regions, inducing large differences between the different scaling schemes. The use of a global-averaged production rate there may hence yield large discrepancies between ages computed with different scaling schemes. The sole calibration sites of the High Tropics are located in the Tropical Andes (e.g. Blard et al., 2013; Delunel et al., 2016), whereas none have been established in Africa, Eurasia, or Oceania. Future calibration studies could also focus on all elevation zones of Eurasia, Oceania, Africa, Central America, Greenland, and Antarctica, where calibration sites are scarce or absent (Fig. 15A). High-elevation, high-latitude calibration sites are also poorly represented.
Figure 15. (A) Calibration sites for cosmogenic $^3$He production rates available in the ICE-D calibration database (24 calibration sites, 158 surface samples). (B) The distribution of production rates and (C–E) production rate variations with latitude, elevation, and exposure age, respectively, were computed and plotted using the online CREp calculator (https://crep.otelo.univ-lorraine.fr/). SLHL and time corrections were performed here with the LSD scaling scheme (Lifton et al., 2014) using the ERA-40 atmospheric model and the VDM dataset derived from atmospheric $^{10}$Be (Muscheler et al., 2005). Source articles of the $^3$He calibration site dataset: (Ackert et al., 2003; Amidon and Farley 2011; Blard et al., 2013; Blard et al., 2006; Borchers et al., 2016; Cerling and Craig, 1994; Delunel et al., 2016; Dunai and Wijbrans 2000; Eaves et al., 2016; Fenton et al., 2009; 2013; Foeken et al., 2012; Goehring et al., 2010; Licciardi et al., 2006; Licciardi et al., 1999).
5.3. Online calculators

Three main calculators are available online to compute the spatial scaling factor and convert $^3$He data into $^3$He exposure ages: (i) the BGC calculator, formerly the initial CRONUS calculator (https://hess.ess.washington.edu/; Balco et al., 2008), (ii) the CRONUS calculator (http://cronus.cosmogenicnuclides.rocks/2.1/; Marrero et al., 2016), and (iii) the CREp calculator (https://crep.otelo.univ-lorraine.fr/#/; Martin et al., 2017).

Each calculator requires particular characteristics (spatial position, thickness, topographic shielding) as inputs to compute production rates specific to the collected samples. This scaling is performed using the most accurate available models, i.e., the Lal-Stone time-dependent model (Balco et al., 2008) and/or the LSD model (Lifton et al., 2014). Note that, in each calculator, the input cosmogenic $^3$He concentration data must already be corrected for other $^3$He sources.

The CRONUS calculator uses a unique and non-modifiable $^3$He, production rate and VDM correction. In contrast, the CREp and BGC calculators allow users to choose from among several production rates: world average, regional mean, and local production rates. In CREp, these production rates can easily be selected from a clickable map. Moreover, CREp connects to the ICE-D calibration database daily, which is regularly updated to incorporate the newest literature data. The CREp and BGC calculators also offer the possibility of using different atmospheric models and geomagnetic databases.

5.4. Production of cosmogenic $^3$He by muons

Numerical modeling based on experimental cross sections suggests that both fast muons and negative-muon capture reactions may produce cosmogenic $^3$He (Lal 1987; Nesterenok and Yakubovich 2016). However, geological evidence supporting such reactions has been scarce, which has precluded the precise calibration of the muogenic production of $^3$He (Kurz 1986a; Shuster et al., 2012). Larsen et al., (2021) recently presented a vertical profile of $^3$He concentrations measured in pyroxenes and ilmenites along a 300-m-long drill core in the Columbia River basalts, which were emplaced by successive massive eruptions 16 Ma (Kasbohm and Schoene 2018). After correcting for non-cosmogenic $^3$He components, this dataset displays a clear attenuation profile that precisely defines the muogenic production of $^3$He: assuming a single exponential attenuation model (Braucher et al., 2011), the surficial SLHL muogenic $^3$He production rate is $0.3 \pm 0.1$ at g$^{-1}$ yr$^{-1}$ and the attenuation length is 8,780 g cm$^{-2}$ (Fig. 16; Larsen et al., 2021). Hence, in a massif having an average density of 2.7 g cm$^{-3}$, the muogenic production rate decreases by an exponential factor each 30 m.
At Earth’s surface, the contribution of this muogenic component represents only 3‰ of the spallogenic $^3$He production (Martin et al., 2017), implying that muons can be safely neglected when determining cosmogenic $^3$He exposure ages. However, with these production parameters, muogenic $^3$He becomes dominant at depths $>5$ m and may represent $10^5$–$10^6$ at $g^{-1}$ in the case of exposures longer than 1 Myr. Shielded samples are useful for determining the non-cosmogenic $^3$He correction for surficial samples (e.g., Amidon and Farley, 2011), for analyzing the pristine $^3$He/$^4$He magmatic component (Dodson et al., 1997), for ensuring the preservation of paleo-exposure signals, and for computing accurate burial ages (Blard et al., 2006). Given the muogenic $^3$He production parameters obtained by Larsen et al. (2021), several tens of meters of shielding are required to ensure the absence of cosmogenic $^3$He for these applications.

**Figure 16.** Vertical attenuation of spallogenic and muogenic $^3$He production rates below the atmosphere-rock interface at SLHL. This model assumes surficial spallogenic and muogenic $^3$He production rates of 124 at $g^{-1}$ yr$^{-1}$ (Martin et al., 2017) and 0.3 at $g^{-1}$ yr$^{-1}$ (Larsen et al., 2021), respectively, with respective attenuation lengths of 160 g cm$^{-2}$ (Gosse and Phillips 2001) and 8780 g cm$^{-2}$ (Larsen et al., 2021). In this calculation, rock density is assumed to be 2.7 g cm$^{-3}$.
These muogenic production parameters affect the calculation of denudation rates from cosmogenic $^3$He measurements in basin-wide eroded sediments (e.g., Ferrier et al., 2013; Puchol et al., 2017). Because $^3$He is not radioactive, all muogenic $^3$He nuclides produced at depth will be preserved until they reach the rock surface. Therefore, muon parameters must be included in the equation used to compute the denudation rate ($\varepsilon$) from cosmogenic $^3$He concentrations:

$$\varepsilon = \frac{A_n \bar{p}_n + A_\mu \bar{p}_\mu}{\rho \bar{p}_{^3\text{He}}},$$ (26)

where $\bar{p}_n$ and $\bar{p}_\mu$ are the basin-averaged spallogenic and muogenic production rates (at g$^{-1}$ yr$^{-1}$), respectively, $A_n$ and $A_\mu$ are their respective attenuation lengths (g cm$^{-2}$; Charreau et al., 2019), and $\rho$ is density of the rock (g cm$^{-3}$).

Previous articles by Ferrier et al. (2013), Litty et al. (2021), and Puchol et al. (2017) calculated denudation rates considering only the spallation parameters. Ideally, these published denudation rates should thus be recomputed with Equation (26), employing the new, well-defined muogenic parameters of Larsen et al. (2021). This revision would increase denudation rates by 5% for basins at an average elevation of 2,000 m, whereas denudation rates at sea level, where the contribution of muogenic production is the largest, would increase by 13%. In the case of $^{10}$Be in quartz, incorporating muogenic production may increase calculated denudation rates by 20% at sea level (Lupker et al., 2012).

Even if equation (26) is a reasonable approximation for computing denudation rates, the model of Heisinger et al. (2002a, b) most accurately describes muogenic production at a given depth (Balco, 2017). However, more work is needed to implement the muogenic $^3$He calibration data of Larsen et al. (2021) into the model of Heisinger et al. (2002a, b).

5.5. Cross-calibration of $^3$He against $^{10}$Be and other cosmogenic nuclides

The most commonly used empirical scaling models that were published in the 1990’s and 2000’s (Dunai and Wijbrans 2000; Lal, 1991; Stone 2000) consider the impact of elevation to be similar for all cosmogenic nuclides, including $^3$He and $^{10}$Be. However, ab-initio physical models have recently revisited the use of different spatial scaling factors for each nuclide (Argento et al., 2013; Lifton et al., 2014), after D. Lal initially proposed the idea in his PhD thesis (Lal, 1958). These models predict that the $^3$He/$^{10}$Be production ratio increases with
increasing elevation, with the computed production of $^3\text{He}$ exceeding that of $^{10}\text{Be}$ by 4% at 3,000 m elevation and up to 7% at 5,000 m (Fig. 17). Empirical benchmarks are necessary to test the validity of these theoretical models and better constrain the spatial variations of each nuclide’s production rate. This can be done by measuring the $^3\text{He}/^{10}\text{Be}$ production ratios in minerals that have recently been exposed at the surface: if the possibility of an inherited cosmogenic $^3\text{He}$ component produced during an earlier exposure episode can be discarded, such cross-calibrations are key because they integrate the same exposure history for both nuclides. Only a handful of studies have produced such cross-calibration data: three articles reported cosmogenic $^3\text{He}$ concentrations in pyroxenes and $^{10}\text{Be}$ concentrations in coexisting quartz (Amidon et al., 2009; Blard et al., 2013; Luna et al., 2018), three others documented $^3\text{He}$ in accessory minerals (apatites, zircons, kyanites, garnets, iron oxides) and $^{10}\text{Be}$ in coexisting quartz (Amidon et al., 2008; Gayer et al., 2004; Kober et al., 2007), and another few articles reported cosmogenic $^3\text{He}$ concentrations in olivines/pyroxenes and $^{10}\text{Be}$ in coexisting feldspars (Blard et al., 2013; Zerathe et al., 2017) or olivines/pyroxenes (Blard et al., 2008; Nishiizumi et al., 1990). For this review, I updated and homogenized these $^3\text{He}/^{10}\text{Be}$ cross-calibration datasets by recalculating the $^3\text{He}/^{10}\text{Be}$ production ratios ($R_{3/10}$) using Equation (3) in Blard et al., (2013), to account for the radioactive decay of $^{10}\text{Be}$. All $^3\text{He}$ concentrations in this dataset were only corrected for magmatic and nucleogenic $^3\text{He}$. They thus include all cosmogenic production pathways: spallation, thermal neutron capture, and muogenic production. Note that these $R_{3/10}$ values are raw $^3\text{He}/^{10}\text{Be}$ production ratios that were not scaled for elevation. These updated data are available in Table S3 (https://doi.org/10.24396/ORDAR-91) and plotted in Figures 17–19. In the following, I use this updated dataset to test correlations and review potential causes of the spatial and rock-specific variabilities of the $^3\text{He}/^{10}\text{Be}$ production ratio.

The cross-calibration studies based on cosmogenic $^3\text{He}$ in pyroxenes and $^{10}\text{Be}$ in coexisting quartz (Amidon et al., 2009; Blard et al., 2013; Luna et al., 2018) provide important observations (Fig. 17A). These two nuclide-mineral pairs are among the most commonly used, and their absolute production rates are the best documented (http://calibration.ice-d.org). Using this dataset, the CREp calculator (Martin et al., 2017), and the Lal-Stone time-dependent scaling (Balco et al., 2008) provides a theoretical SLHL $R_{3/10}$ value of 30.1 ± 3.1 (1σ) (Fig. 17). Then, comparing $^3\text{He}$ in pyroxene-$^{10}\text{Be}$ in quartz cross-calibrations measured at 1,333, 4,530, and 4,827 m elevation indicates a slight impact of elevation on the measured $R_{3/10}$ value (Fig. 17A). These data are in agreement, within 1σ uncertainties, with the theoretical models of Argento et al. (2013) and Lifton et al. (2014) that predict a 7–8% increase of $R_{3/10}$ at 5,000 m compared to
the sea-level value. Although this observation supports the use of nuclide-specific scaling, further new cross-calibration and absolute calibration datasets at variable elevations and latitudes will be useful for confirming and refining this relationship. I stress that the use of classical, non-nuclide-specific scaling factors, such as the Lal-Stone (Balco et al., 2008) or LSD scaling (Lifton et al., 2014), do not yield inaccuracies if using locally calibrated production rates, i.e., from calibration sites at similar elevations as the dated objects (Blard et al., 2013; Kelly et al., 2015; Martin et al., 2015).

Figure 17. Comparison of modeled (Argento et al., 2013; Lifton et al., 2014; Lal, 1991) and observed $^{3}$He/$^{10}$Be production ratios vs elevation. Cross-calibrations are from (A) $^{3}$He in pyroxenes and $^{10}$Be in quartz (Amidon et al., 2009; Blard et al., 2013; Luna et al., 2018) and (B) $^{3}$He/$^{10}$Be in any mineral pair (Amidon et al., 2008, 2009; Blard et al., 2008, 2013; Gayer et al., 2004; Kober et al., 2007; Luna et al., 2018; Nishiizumi et al., 1990; Zerathe et al., 2017). The vertical line shows the $^{3}$He/$^{10}$Be production ratio at SLHL calculated using $^{3}$He in pyroxene-$^{10}$Be in quartz data (http://calibration.iced.org), the CREp calculator (https://crep.otelo.univ-lorraine.fr/#; Martin et al., 2017), and the Lal-Stone time-dependent scaling (Balco et al., 2008); the gray shaded bar indicates 1σ uncertainty.

However, considering the global $^{3}$He/$^{10}$Be cross-calibration dataset, reported $R_{3/10}$ values deviate markedly from SLHL production ratio, and even from the theoretical production models of Argento et al. (2013) and Lifton et al. (2014). These high $R_{3/10}$ values reach ~50 above 4,000 m, and are from cross-calibration data based on $^{3}$He concentrations measured in accessory minerals (garnet, apatite, zircon; Amidon et al., 2008; Gayer et al., 2004). By determining the $^{3}$He in garnet/$^{10}$Be in quartz production ratio from Himalayan glacial rocks at 3,000–4,500 m
elevation, Gayer et al. (2004) identified an overproduction of cosmogenic $^3$He compared to the standard scaling model that assumes an elevation-independent $^3$He/$^{10}$Be production ratio (Fig. 17; Lal, 1991). Dunai et al. (2007) attributed this high-elevation $^3$He production excess to the unrecognized impact of cosmogenic thermal neutrons through the $^6$Li(n,$\alpha$)$^3$H reaction. According to Dunai et al. (2007), this reaction could account for a 50% increase of cosmogenic $^3$He production in the Li-rich garnets (20–100 ppm Li) analyzed by Gayer et al. (2004). However, Amidon et al. (2008) later reported $^3$He and $^{10}$Be cross-calibration data on Li-poor (<10 ppm) minerals (zircons, kyanites) from the high Himalayas, which yielded $^3$He/$^{10}$Be production ratios in the same range as those determined in garnet by Gayer et al. (2004) (Fig. 17). This empirical evidence indicates that Li is not the main driver of this “anomalously” high cosmogenic $^3$He production rate. Furthermore, when $^3$He/$^{10}$Be cross-calibration data across all mineral pairs ($n = 88$) are plotted against Li concentration (Fig. 18), no correlation is observed ($R^2 = 0.19, n = 78$). Hence, other mechanisms are required to explain the high $R_{3/10}$ values above 3,000 m. Interestingly, the highest anomalous $^3$He production rates are reported for metamorphic or plutonic lithologies (Gayer et al., 2004; Amidon et al., 2008), whereas materials from recent (<10 Ma) volcanic terrains yield much lower $^3$He/$^{10}$Be production ratios (Amidon et al., 2009; Blard et al., 2013; Luna et al., 2018).

Figure 18. $^3$He/$^{10}$Be production ratio plotted against Li concentration (ppm) in the cosmogenic $^3$He-hosting mineral (data sources: Amidon et al., 2008, 2009; Blard et al., 2008, 2013; Gayer et al., 2004; Kober et al., 2007; Luna et al., 2018; Nishiizumi et al., 1990; Zerathe et al., 2017).
Although further evidence is required to reach definitive conclusions, several mechanisms might explain these observations:

i) **Inheritance of previous exposure.** Because $^{3}$He is a stable nuclide, it is more prone than radioactive nuclides to be affected by inheritance from previous exposure episodes that might have occurred several million years ago. Hence, after the radioactive decay of $^{10}$Be during subsequent burial episodes, such inherited $^{3}$He concentrations will appear to be in excess compared to the actual $R_{3/10}$ value. This mechanism was proposed by Zerath et al. (2017) to identify two outliers among a dataset of eight samples from a $^{3}$He/$^{10}$Be cross-calibration based on andesitic boulders. Another observation supporting this explanation is the relationship between observed $R_{3/10}$ values and $^{10}$Be-derived exposure ages for the whole dataset (Fig. 19). I used $^{10}$Be to compute these exposure ages because this radioactive nuclide is less susceptible than $^{3}$He to be biased by inheritance and because its production rate is better constrained. Interestingly, the scatter among observed $R_{3/10}$ values is much greater for younger exposure ages (<50 ka), and the maximum values of $R_{3/10}$ (>50) are only observed in that population. Moreover, exposure ages older than 100 ka ($n = 10$ of 88) systematically yield $R_{3/10} < 40$. If true, the interpretation of $^{3}$He inheritance implies that the anomalously large $^{3}$He/$^{10}$Be production ratios are not real, but a geomorphological artifact. However, the Himalayan samples (Gayer et al., 2004; Amidon et al., 2008) are from a region characterized by rapid exhumation (1–10 km/Myr; Amidon et al., 2008), a context that, in theory, favors the rapid turnover and rejuvenation of the subsurface, which should preclude significant inheritance.
Figure 19. $^{3}$He/$^{10}$Be production ratios plotted against $^{10}$Be exposure ages (data sources: Amidon et al., 2008, 2009; Blard et al., 2008, 2013; Gayer et al., 2004; Kober et al., 2007; Luna et al., 2018; Nishiizumi et al., 1990; Zerathe et al., 2017).

ii) Chemical compositions of rocks and minerals. Larsen et al., (2019) recently underscored that the composition of the whole rock may play a major role in the $^{3}$He production rate, and that this bulk chemistry could be more important that the mineral composition itself because of the long-distance (20–100 µm) ejection of cosmogenic $^{3}$He and $^{3}$H (Fig. 4; Nesterenok and Yakubovich, 2016). The impact of this $^{3}$He redistribution is larger for smaller grains (Larsen et al., 2019). This explanation may be compatible with the fact that the greatest observed $^{3}$He overproduction (i.e., $R_{3/10} > 40$) at high elevation has so far only been documented in silicic lithologies (Amidon et al., 2008; Gayer et al., 2004), and not in intermediate rocks such as dacite or andesite (Blard et al., 2013; Luna et al., 2018). Although potential production pathways may be implicated in this bulk compositional effect (e.g., Lal, 1987), the exact reaction(s) and element(s) that may have induced this unrecognized $^{3}$He production remain elusive and should be explored with additional data.

iii) Impact of water on thermal neutron capture. Liquid water, ice, or snow enhance thermal neutron capture and may modify the balance between spallation and thermal neutron production in the first few meters below the air-rock...
interface. This may affect the \(^3\)He/\(^{10}\)Be production ratio because \(^{10}\)Be is not produced by thermal neutron capture (Delunel et al., 2014; Dunai et al., 2014). Indeed, the largest anomalous \(^3\)He/\(^{10}\)Be production ratios above 3,000 m have only been documented in wet regions of the Himalayas (Amidon et al., 2008; Gayer et al., 2004), and not in the dry Andes (Blard et al., 2013; Luna et al., 2018; Zerathe et al., 2017). However, the global dataset clearly indicates that \(R_{3/10}\) is not correlated with Li content, which contradicts the potential involvement of a \(^6\)Li(n,\(\alpha\))\(^3\)H reaction. If future data confirm the role of water or snow cover in inducing high \(^3\)He/\(^{10}\)Be production ratios, the existence of an unknown production pathway involving thermal neutrons and an element other than Li should be explored. \(^2\)H(n)\(^3\)H and \(^{10}\)B(n,\(^8\)Be)\(^3\)H reactions are potential candidates (Lal, 1987). They could be tested with ad hoc experimental data.

New \(^3\)He/\(^{10}\)Be cross-calibration data and measuring cosmogenic \(^3\)He in different minerals (with variable compositions and grain sizes) from different lithologies (andesite, dacite, granite-like composition) exposed at variable elevations and in different climatic contexts with variable snow cover will be useful in addressing these issues.

Cross-calibrations of \(^3\)He\(_c\) against cosmogenic nuclides other than \(^{10}\)Be may also provide information about nuclide-specific altitudinal scalings, although such observations are rare: few studies have provided intercomparison data with cosmogenic \(^{21}\)Ne, \(^{26}\)Al, and \(^{36}\)Cl measured in the same samples as \(^3\)He (Licciardi et al., 2008; Luna et al., 2018; Schimmelpfennig et al., 2011). The \(^{36}\)Cl production rate is extremely sensitive to mineral composition, notably the K, Ca, and Cl contents of the target minerals (Schimmelpfennig et al., 2009). The \(^3\)He/\(^{36}\)Cl production ratios reported by Licciardi et al. (2008), Luna et al. (2018), and Schimmelpfennig et al. (2011) were obtained by measuring \(^{36}\)Cl concentrations in enstatites and feldspars or whole rocks with variable chemical compositions, notably their Cl contents. As these studies are based on different localities at variable latitudes and altitudes, there are too many unknowns to draw definitive conclusions about \(^3\)He/\(^{36}\)Cl production ratios from these datasets.

That said, Schimmelpfennig et al. (2011) documented the \(^3\)He/\(^{36}\)Cl production ratios by measuring \(^3\)He, \(^{36}\)Cl, and \(^{21}\)Ne in low-Ca pyroxenes (15% Ca) of homogenous composition sampled along a vertical transect between 1,000 and 4,300 m elevation at Kilimanjaro volcano (3°S, Tanzania). This dataset suggests that the \(^3\)He/\(^{21}\)Ne and \(^3\)He/\(^{36}\)Cl production ratios in low-Ca pyroxenes (average values of 5.4 ± 0.2 and 17.2 ± 1.8, respectively) are not elevation-
dependent. Luna et al. (2018) also measured \(^3\)He/\(^{21}\)Ne in low-Ca pyroxenes (En\(_{88-94}\)) from moraines of the Altiplano-Puna plateau (24°S, Argentina) at 4,500 m elevation; they reported a \(^3\)He/\(^{21}\)Ne production ratio of 4.06 ± 0.12. By measuring the \(^3\)He/\(^{21}\)Ne production ratio in olivines and pyroxenes from <200 ka lava flows in Arizona situated at 1,500–2,000 m elevation, Fenton and Niedermann (2014) evidenced that mineral composition has a significant impact on the absolute cosmogenic \(^{21}\)Ne production rate. Hence, the contrast between the \(^3\)He/\(^{21}\)Ne production ratios determined by Luna et al. (2018) and Schimmelpfennig et al. (2011) above 4,000 m probably results from distinct absolute \(^{21}\)Ne production rates due to different pyroxene compositions.

Although many uncertainties remain on the element-specific production pathways of other nuclides, these cross-calibrations against \(^{21}\)Ne and \(^{36}\)Cl support the hypothesis that \(^3\)He in olivines and pyroxenes is probably not characterized by an unrecognized overproduction at high elevation (>2,000 m) (Blard et al., 2013a, b), in contrast to accessory minerals such as garnet (Gayer et al., 2004), kyanite, or zircon (Amidon et al., 2008). New cross-calibrations varying mineralogy, elevation, latitude, and snow cover will be useful for confirming this conclusion.

5.6. New directions of cosmogenic \(^3\)He coupled with cosmogenic radioactive nuclides (\(^{14}\)C, \(^{10}\)Be, \(^{36}\)Cl, and \(^{53}\)Mn)

Beyond cross-calibration issues, measuring two (or more) cosmogenic nuclides with different half lives in the same mineral also has great potential for many geoscience applications, including i) burial dating (Lebatard et al., 2014), ii) measuring paleo-depths of exposure (Hidy et al., 2018) or the thickness of snow cover (Hippe et al., 2014), iii) determining paleo-elevations (Blard et al., 2019a, b), or iv) detecting complex exposure histories such as the dynamics of sediment transfer (Wittmann and von Blanckenburg, 2009). The most commonly used nuclide pair for these applications is \(^{26}\)Al/\(^{10}\)Be in quartz (Granger, 2006), although \(^{10}\)Be/\(^{21}\)Ne in quartz has the potential to probe longer timescales (5–20 Ma, compared to 0.5–6 Ma for \(^{26}\)Al/\(^{10}\)Be) (Balco and Shuster, 2009; Sartégou et al., 2020).

Despite their intriguing potential, these nuclide pairs remain to be tested in quartz-free environments, such as andesitic or mafic terrains. Although measuring \(^3\)He is often possible in such geological settings, measuring radioactive cosmogenic nuclides in the same rock sample remains challenging; only a few studies have reported joint measurements of \(^3\)He and a radioactive cosmogenic nuclide in the same mineral (Blard et al., 2008; Licciardi et al., 2008; Nishiizumi et al., 1990).
Because the measurement of $^{36}$Cl in Ca- and K-rich minerals (feldspars, pyroxene) is quite well established (Schimmelpfennig et al., 2009), the $^{3}$He-$^{36}$Cl pair is, for now, the most accessible nuclide pair for multi-isotopic analyses in mafic terrains (Licciardi et al., 2008). However, given the half-life of $^{36}$Cl ($t_{1/2} = 301$ kyr), this pair can only be used to probe the last few million years in the best cases. Measuring $^{10}$Be in olivine and pyroxene might prove useful in extending this temporal range, although removing atmospheric $^{10}$Be contamination from mafic minerals has long been considered an unsolvable problem (Ivy-Ochs et al., 1998). A new chemical procedure employing a preliminary crushing step has allowed the accurate measurement of $^{10}$Be in olivine and pyroxene (Blard et al., 2008; Eaves et al., 2018). Using these data and the Lal-Stone time-dependent model, Eaves et al., (2018) determined the SLHL $^{10}$Be production rate to be $3.2 \pm 0.8$ at g$^{-1}$ yr$^{-1}$ in olivine and pyroxene, extending the utility of $^{10}$Be burial age or paleoaltimetry measurements in intermediate and mafic rocks to 5 Ma. Recent developments in accelerator mass spectrometry offer the possibility to measure $^{53}$Mn in Fe-rich minerals, such as iron-oxides (Schiffer et al., 2020). With a half-life of 3.74 Myr, $^{53}$Mn is an interesting complement to $^{3}$He; this nuclide pair could extend the range of burial dating to 25 Ma. The ability to measure short-lived in-situ cosmogenic $^{14}$C ($t_{1/2} = 5,730$ yr; Pigati et al., 2010) in the same mineral as $^{3}$He is also an interesting development because this pair could be used to determine paleo-depths of exposure, snow cover thicknesses, landslide volumes, and burial ages from 1 to 40 ka.

All such applications involving exposures below the rock-atmosphere interface, i.e., burial episodes, should carefully integrate the impact of the muogenic contribution using the most recently determined parameters, currently those of Larsen et al. (2021) for $^{3}$He$_c$. See reviews by Balco (2017), Braucher et al. (2011), and Lupker et al. (2015) for the muogenic production of other cosmogenic nuclides.

6. Concluding remarks

In this review article, I have summarized and different strategies that can be applied to accurately and precisely measure cosmogenic $^{3}$He concentrations. Even using the most accurate and precise mass spectrometer, non-cosmogenic $^{3}$He components have a major impact on the detection limit and the precision of the method; correcting for other sources of $^{3}$He in the mineral of interest thus requires particular attention. The choice of the best correction strategy is determined by the respective concentrations of radiogenic and magmatic $^{4}$He:
Case 1: Magmatic and radiogenic $^4\text{He}$ concentrations are of the same order of magnitude.

Measure U, Th, and Li and independently determine the He closure age to compute $^4\text{He}^*$. 

Strategy 1a – Crush and fuse minerals, use Equation (15).

Strategy 1b – Fuse multiple aliquots and build isochrons (see section 3.5.3).

Case 2: Uneroded lavas (a particularity of Case 1). In this case, it is not necessary to calculate the He closure age. However, U, Th, and Li measurements remain necessary to correct for radiogenic He by calculating the $R$-factor.

Strategy 2a – Crush and fuse minerals, use Equations (17-18).

Strategy 2b – Fuse multiple aliquots and build isochrons (see section 3.5.3).

Case 3: $^4\text{He}^* \gg ^4\text{He}_{\text{mag}}$. This is the case for samples with very long closure ages ($>1$ Myr) and/or U-Th-rich minerals ($>1$ ppm). In this situation, the correction for magmatic $^3\text{He}$ is not easy and may hinder the precision on cosmogenic $^3\text{He}$, particularly for short exposure ages ($<10$ ka). Prolonged crushing or the fusion of cosmogenically shielded samples from the same lithology are ways to estimate the magmatic $^3\text{He}$ component.

Future improvements will be to: i) improve our knowledge of the spatial and temporal variability of production rates; ii) develop cosmogenic $^3\text{He}$ measurements in new minerals, such as accessory minerals in Si-rich rocks; iii) better document the impacts of the chemical compositions of minerals and their host rocks on production rates; and iv) better understand exotic $^3\text{He}$ production pathways, notably those involving muons and thermal neutrons. To achieve these objectives, important information will be acquired from new calibrations of $^3\text{He}_c$ production rates measured in non-eroded and well-dated surfaces, as well as from cross-calibrations with other cosmogenic nuclides such as $^{10}\text{Be}$. These future directions have the potential to improve the accuracy and precision of the $^3\text{He}$ geoscientific toolbox and open the door to new applications (burial dating, paleoaltimetry, paleo-depths, durations of volcanic eruptions) and lithologies.

Acknowledgment

I thank Chemical Geology and notably David Hilton, Don Porcelli and Tim Horsercroft, who invited me to write this review article. I am grateful to them for their tremendous patience. Indeed, I started writing
this article about five years ago, meaning that I spent 1,000 nights near the cosmic camel Alexandrin
during the entire writing process. Thorough comments by three anonymous reviewers and by editor D.
Porcelli improved the submitted version of the article. I am also grateful to Robert Dennen (rd-
editing.com) for English language editing. This review greatly benefited from numerous previous works
and discussions with my colleagues who use cosmogenic nuclides in geoscience (J. Charreau, J.-F. Ritz,
C. Litty, S. Zerathe, P. Valla). Previous work by cosmogenic $^3$He experts was also a great inspiration:
Shuster, G. Bromley, N.A. Lifton, J.M. Schaefer, G. Winkler, R.P. Ackert, S. Niedermann, M.
Tremblay, P.M. Vasconcelos, R. Yokochi, S. Mukhopadhyay, B. Marty, H. Baur, I. Leya, R. Wieler, J.
Masarik, and many others. Finally, I of course send a special “thank you” to the helium pioneers, whose
fundamental discoveries and principal publications opened new paths and fields of research: G.H.
Megreue, M.D. Kurz, T.E. Cerling, D. Lal, H. Craig. “Nani gigantum humeris insidentes – We are dwarfs
standing on the shoulders of giants” Bernard de Chartres, 12th century.
References


Nishiizumi, K., J. Klein, R. Middleton, and H. Craig. 1990. “Cosmogenic 10Be, 26Al and 3He


