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Article in Earth and Planetary Science Letters • October 2015
DOI: 10.1016/j.epsl.2015.07.032

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New perspectives on the Li isotopic composition of the upper continental crust and its weathering signature

Lucie Sauzéat a,b,*, Roberta L. Rudnick a, Catherine Chauvel b,c, Marion Garçon d, Ming Tang d

a Department of Geology, University of Maryland, College Park, MD 20742, USA
b University of Grenoble Alpes, ISTerre, F-38041 Grenoble, France
c CNRS, ISTerre, F-38041 Grenoble, France
d Carnegie Institution of Washington, Department of Terrestrial Magnetism, Washington, DC 20015, USA

A R T I C L E   I N F O

Article history:
Received 11 May 2015
Received in revised form 1 July 2015
Accepted 14 July 2015
Available online 3 August 2015
Editor: M. Bickle

Keywords:
Li isotopes
loess
upper continental crust
chemical weathering

A B S T R A C T

Lithium isotopes are increasingly used to trace both present-day and past weathering processes at the surface of the Earth, and could potentially be used to evaluate the average degree of past weathering recorded by the upper continental crust (UCC). Yet the previous estimate of average δ7Li of the UCC has a rather large uncertainty, hindering the use of Li isotopes for this purpose. New δ7Li for desert and periglacial loess deposits (windblown dust) from several parts of the world (Europe, Argentina, China and Tajikistan) demonstrate that the former are more homogeneous than the latter, and may therefore serve as excellent proxies of the average composition of large tracts of the UCC. The Li isotopic compositions and concentrations of desert loess samples are controlled by eolian sorting that can be quantified by a binary mixing between a weathered, fine-grained end-member, dominated by phyllosilicates and having low δ7Li, and an unweathered, coarse-grained end-member, that is a mixture of quartz and plagioclase having higher δ7Li. We use correlations between insoluble elements (REE, Nd/Hf and Fe2O3/SiO2), Li concentrations (henceforth referred as [Li]), and δ7Li to estimate a new, more precise, average Li isotopic composition and concentration for the UCC: [Li] = 30.5 ± 3.6(2σ) ppm, and δ7Li = +0.5 ± 0.6(2σ). The δ7Li for desert loess deposits is anti-correlated with the chemical index of alteration (CIA). Using this relationship, along with our average δ7Li, we infer that (1) the present-day CIA of the average UCC is δ16.5(2σ), higher than the common reference value of 53, and (2) the average proportion of chemically weathered components is as high as 37 ± 17(2σ)% at the surface of the Earth.

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1. Introduction

Weathering is an ubiquitous process that occurred in the past to form (meta-)sedimentary rocks (“past weathering”) and still occurs at present to create sediments and soils (“present-day weathering”). It shapes the continental crust and modifies its chemical composition by producing detrital sediments and releasing ions into the hydrosphere over geological timescales (Taylor and McLennan, 1985; Rudnick, 1995; Lee et al., 2008; Liu and Rudnick, 2011). Chemical weathering also indirectly controls the evolution of climate because carbonates, precipitating with Ca released during weathering of the continental crust, sequester CO2 (Gaillardet et al., 1999; Amiotte Suchet et al., 2003; Jin et al., 2014). Constraining the degree of weathering experienced by the uppermost part of the crust in the past is thus important in providing a framework that can be used to understand both the compositional evolution of the upper crust and climate variation through time.

It is now well established that lithium isotopes are fractionated by low-temperature processes due to preferential partitioning of 7Li (the heavy isotope) into water while 6Li (the light isotope) is incorporated into the weathered products of silicate rocks such as clays (e.g., Pistiner and Henderson, 2003; Vigier et al., 2008). Li isotopes are thus excellent tracers of fluid-rock reactions and have been used to trace weathering processes (Kisakürek et al., 2005, 2004; Liu et al., 2013; Millot et al., 2010; Rudnick et al., 2004), estimate the mass of continental crust lost by chemical weathering through geological times (i.e., Li dissolved in river water) (Liu and Rudnick, 2011) and constrain the present-day erosion cycle of mountain ranges (Dellinger et al., 2014). In addition, Li isotopes are not significantly fractionated by high-temperature...
processes such as metamorphic dewatering (Qiu et al., 2011, 2009; Teng et al., 2007) or igneous differentiation (Teng et al., 2006; Tomascak et al., 1999), hence they are also useful proxies for studying crustal recycling in subduction zones (Elliott et al., 2006; Tang et al., 2014) or beyond (e.g., Vlastelic et al., 2011). Here, we propose to use Li isotopes to quantify the importance of chemical weathering experienced by the upper crust in the past.

To date, the published average lithium concentrations for the upper continental crust differ by more than 40%, ranging from 20 ppm (Taylor and McLennan, 1985) to more than 40 ppm (Teng et al., 2004; Hu and Gao, 2008). The average Li isotopic composition of the UCC ($^{6}Li = 0 \pm 4$ (2$\sigma$, Teng et al., 2004) is estimated to be lower than that of fresh, mantle-derived basalts, suggesting that the upper crust records a weathering signature, but the uncertainty on this value overlaps with that of the mantle ($^{6}Li \approx +4 \pm 2$ (2$\sigma$), average derived from Chan et al., 1992; Seitz et al., 2004, and Tomascak et al., 2008).

In this paper, we report [Li] and $^{6}Li$ for loess deposits that sample vast regions of the UCC. The main advantage of using loesses instead of other sediments (e.g., shales, mudrocks, etc.) for this purpose is that loesses are deposits of eolian dust produced by mechanical erosion and mixing of silt derived from glacial outwash and/or desert environments (Pye, 1995); thus they generally experienced little chemical weathering during their formation. These two features allow loess deposits to be considered as proxies for the average UCC composition (Chauvel et al., 2014; Gallet et al., 1996, 1998; McLennan, 2001; Peucker Ehrenbrink and Jahn, 2001; Taylor et al., 1983).

2. Samples

Loess samples cover about 10% of the Earth’s surface (Haase et al., 2007; Pécsi, 1990) and formed between the end of the Pliocene and the beginning of the Pleistocene, based on climatological dating (Liu, 1985). These Quaternary deposits can be divided into two distinct types: (1) periglacial loess, derived from glacial outwash and transported by winds over limited distances (Haase et al., 2007; Rousseau et al., 2014), and (2) desert loess, transported over longer distances, typically several hundreds of kilometers (Ding et al., 1999), from desert regions.

In this study, we analyzed previously well-characterized desert loesses from China (Jixian, Xifeng, Xining and Luochuan) (Gallet et al., 1996; Jahn et al., 2001), Tajikistan (Chashmanigar) (Ding et al., 2002; Yang et al., 2006) and Argentina (Buenos Aires) (Gallet et al., 1998), as well as periglacial loesses from Western Europe (France, England and Spitsbergen (Gallet et al., 1998). A map showing the sample locations can be found in Chauvel et al. (2014). Mineralogy, provenance and grain-size distribution for these deposits have been reported in previous studies such as Jeong et al. (2008) and Brügner and Heinkele (1990) for the mineralogy of the Chinese loesses, Smith et al. (2003) and Zárate et al. (2003) for the source location of the Argentinian loesses, and Ding and Ding (2003) for the grain-size distribution of loesses from Tajikistan.

3. Methods

3.1. Major, trace elements, and radiogenic isotopes

Trace element concentrations and Nd–Hf isotopes have been previously published for all the loess samples (Chauvel et al., 2014; Gallet et al., 1996). Major element concentrations are available in the literature for most of the samples (Gallet et al., 1998, 1996; Jahn et al., 2001), except for the Tajikistan loesses. We thus analyzed their major element contents in this study (see Supplementary Table A).

Major element data for the Tajikistan samples were determined using an ICP-AES (Varian 720 ES) at ISTerre (Grenoble) following the same procedure described in Chauvel et al. (2011); only a brief synopsis is provided here. About 50 mg of powder, hand-crushed in an agate mortar were dissolved in a HF–HNO$_3$ mixture in Savillex beakers for about 72 h at 90°C. Boric acid was added to neutralize the HF and the resulting liquid was diluted with milliQ water. The concentrations are calculated using calibration curves based on diluted and doped (Al, Mg, Ca, Na, K and P) BCR-2 rock standard solutions.

Both accuracy and reproducibility of the major element contents of loesses from Tajikistan were monitored by replication of international rock standards. The concentrations obtained for the rock standards are in agreement with the reference values, and reproducibility is, on average, better than 5% (2$\sigma$) (see Supplementary Table B).

3.2. Lithium isotopic compositions

Lithium isotopic compositions were measured at the University of Maryland. For each sample, about 25 mg of rock powder was dissolved in a Savillex screw-top beaker in an HF–HNO$_3$ mixture following the procedure detailed by Teng et al. (2004). For samples containing organic matter, a second step of dissolution using a combination of HF–HNO$_3$–HClO$_4$ was employed to achieve a complete digestion. Separation of lithium was performed by chromatography on four successive columns following the ion resin techniques described by Moriguti and Nakamura (1998a). Lithium was purified using columns filled with 1 ml of Bio-Rad AG 50w-x12 (200–400 mesh) resin, first with an HCl solution, followed by an HCl-ethanol mixture. Lithium isotopic composition of the purified solutions (∼50 ppb Li in 2% HNO$_3$) were measured on a Nu Plasma Multicollector Inductively Coupled Plasma Mass Spectrometry (MC-ICPMS) and calculated by standard bracketing using L-SVEC (Flesch et al., 1973) as the reference standard. Prior to each analysis, the Na/Li ratio of the solution was determined and samples with Na/Li ratio greater than 5 went through further purification by chromatography.

The accuracy of the Li isotopic composition is assessed based on the analysis of two rock reference materials (AGV-1 and G-2). Our measured values (Supplementary File A and Supplementary Table C) are within uncertainty of previously published results, with $^{6}Li_{AGV1} = +/-6.7 \pm 0.7$ (2$\sigma$, n = 3), versus +4.6 to +6.7 (Liu et al., 2013, 2010, 2015; Magna et al., 2004), and $^{6}Li_{G-2} = +0.6 \pm 1.8$ (2$\sigma$, n = 3) versus −1.6 to +2.2; (James and Palmer, 2000; Liu et al., 2010; Pistiner and Henderson, 2003; Tang et al., 2014). The long-term precision of our results is assessed by repeated analyses of pure “in-house” Li standard solutions (UMD-1 and IRMM-016) performed over the course of our analyses. Our average $^{6}Li$ results are +54.9 ± 0.5 (2$\sigma$, n = 8) for UMD-1, and +4.0 ± 0.5 (2$\sigma$, n = 8) for IRMM-016 (Supplementary File A and Supplementary Table C). Given our long-term reproducibility, the 2 sigma analytical uncertainty adopted in this study for the Li isotopic composition is ±1‰.

4. Results

New major element concentrations for the Tajikistan loesses and previously published data for all other samples are provided in Supplementary Table A. The reader is referred to the studies of Gallet et al. (1998, 1996) and Jahn et al. (2001) for a full description of major element concentrations in loess samples. Here, we briefly compare loess deposits as a function of their formation mechanism (i.e., periglacial vs. desert loesses) and as a function of their sampling locations. The first noticeable difference is
that periglacial samples have higher SiO₂ concentrations (average of 76 ± 9(2σ) wt% vs. 57 ± 12(2σ) wt% for desert loesses) and lower contents of all the other major elements compared to the desert loesses (see Supplementary Table A). Among desert loesses, samples from Tajikistan, China, and Argentina form three distinct groups. Loesses from Tajikistan are generally more concentrated in Fe₂O₃, CaO, Al₂O₃, MnO and MgO than Chinese loesses, which are themselves more enriched in these elements than Argentinian loesses. The only exception is the high Al₂O₃ contents of the Argentinian loesses (see Supplementary Table A). By contrast, loesses from Tajikistan are, on average, depleted in SiO₂, K₂O and Na₂O compared to Chinese loesses, which are themselves lower than the Argentinian loesses. Some of these features are illustrated in Fig. 1 where we plot Al₂O₃/Na₂O vs. Fe₂O₃/SiO₂ showing that desert loesses follow a well-defined linear trend whereas periglacial loesses scatter, in part due to their low Fe₂O₃ and high SiO₂ contents that translates into low Fe₂O₃/SiO₂ ratios. Among desert loesses, Tajikistan samples have higher Al₂O₃/Na₂O and Fe₂O₃/SiO₂ ratios than Chinese samples while Argentinian samples are somewhat lower but more variable.

Lithium concentrations and isotopic compositions are reported in Table 1 for all loesses and are plotted in Figs. 2 and 3, respectively. As for major element concentrations, periglacial loesses differ from desert loesses in their Li compositions. Periglacial deposits show a wide range of δ⁷Li values, between −2.9 and +4.7, with an average value of +0.1 ± 5.6(2σ) (Fig. 3). This variability is also seen in the Li concentrations, which range from 17 to 61 ppm, with an average of 40 ± 39(2σ) ppm (Fig. 2). Such variability is comparable to that seen in periglacial loess deposits investigated by Teng et al. (2004) from New Zealand, Germany, and the midwestern USA (δ⁷Li: −3.1 to +4.8; [Li]: 17 to 41 ppm). Desert loesses have less variable Li concentrations and isotopic compositions (on average 37 ± 13(2σ) ppm and +0.9 ± 3.0(2σ) respectively; Fig. 2 and Fig. 3) and each locality also has distinct averages: the Tajikistan samples have the highest [Li] and the lowest δ⁷Li values ([Li] = 41 ± 15(2σ) ppm and δ⁷Li = −0.3 ± 1.2(2σ); Fig. 3), the Chinese samples are intermediate ([Li] = 37 ± 6(2σ) ppm and δ⁷Li = +1.0 ± 2.0(2σ); Fig. 3) while the Argentinian samples show the lowest [Li] but the heaviest Li isotopic composition ([Li] = 30 ± 9(2σ) ppm and δ⁷Li = +2.7 ± 3.5(2σ); Fig. 3).

5. Discussion

5.1. Use of desert loess deposits to establish the Li isotopic composition of the upper continental crust (UCC)

Several studies have previously used loess deposits to estimate the average composition of the UCC (Taylor et al., 1983; Gallet et al., 1996, 1998; McLennan, 2001; Peucker Ehrenbrink and Jahn, 2001; Chauvel et al., 2014). Most of these studies focused on periglacial loesses. However, as can be seen in Figs. 1 to 3, the two types of loess do not carry the same information. Periglacial loess deposits appear to be enriched in quartz (as seen in their high SiO₂ and low Fe₂O₃/SiO₂, Fig. 1) and have a large range of Li concentrations and Li isotopic compositions (Figs. 2 and 3), while desert loesses are more homogeneous. Such variability can result either from the presence of metamorphic and igneous rocks.

![Fig. 1. Plot of grain-size proxies (Al₂O₃/Na₂O vs. Fe₂O₃/SiO₂) illustrating the differences between periglacial and desert loess samples. Periglacial loesses include data from this study, as well as literature data (light blue) from Teng et al. (2004). Fine-grained particles such as clays have high Al₂O₃/Na₂O and Fe₂O₃/SiO₂. Desert loesses plot on a well-defined mixing array, whereas periglacial loesses show greater scatter. Data for the Chinese loess come from Gallet et al. (1996) and Jahn et al. (2001) and from Gallet et al. (1998) for the Argentinian and Western Europe loess. The correlation coefficients R² are calculated for desert loesses only. Error bars represent 2σ. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)](image-url)

![Fig. 2. Stacked histogram of [Li] in desert (red) and periglacial (blue) loess deposits. Periglacial loesses include data from this study, as well as data from Teng et al. (2004). Li concentrations for the samples analyzed in this study are from Chauvel et al. (2014). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)](image-url)
fragments derived from the source (Swineford and Frye, 1955; Garçon et al., 2014); a “nugget effect” i.e. the over-concentration of heavy minerals associated to quartz grains or reflect the preservation of inherited isotopic variability from the source rocks due to short transport distances (Roussseau et al., 2014).

Although heavy minerals, mainly zircons, have been shown to significantly influence neodymium, hafnium and lead isotopic compositions of fine-grained sediments, including loesses (Garçon et al., 2014; Chauvel et al., 2014), such effects are unlikely to be responsible for the Li isotopic variability we observe. Indeed, periglacial loesses are enriched in heavy minerals such as zircons (Chauvel et al., 2014; Taylor et al., 1983; Taylor and McLennan, 1985), as reflected by their very low Nd/Hf ratio (Fig. 3a), which is, on average, lower than 3.5 (Chauvel et al., 2014). Zircons control the Hf but not the Nd budget of sediments (Garçon et al., 2014), therefore, an excess of zircon in the sediments generates a low Nd/Hf ratio. Although zircons have variable Li contents (0.5 to 250 ppm) and Li isotopic compositions, ranging from −24 to +14 (Bouvier et al., 2012; Ushikubo et al., 2008), characteristics that could explain some of the Li variations seen in periglacial loesses, a mass balance calculation demonstrates that heterogeneous distribution of zircon cannot be the cause of the Li isotopic variability. Here we consider two extreme cases for Li in zircon ([Li] = 250 ppm; δ7Li = −24 and [Li] = 250 ppm; δ7Li = +14) and
we calculate that the presence of <0.5% zircon by mass (the average proportion of zircons in periglacial loesses as suggested by Bronger, 2003 and Rousseau et al., 2007) can shift the Li isotopic composition by no more than 1%. Therefore, the Li variations in periglacial loesses cannot be explained simply by an excess of zircons and a similar conclusion can probably be drawn for the other heavy minerals (e.g. epidote, goethite and hematite).

Thus, preservation of Li isotopic variability inherited form the source is likely responsible for the Li heterogeneity observed in periglacial loesses. Such results are consistent with the conclusions of Chauvel et al. (2014) who focused on trace elements and Nd–Hf isotopic characteristics of loess and concluded that periglacial loess samples, in addition to being extremely variable regarding their trace element patterns, are likely too heterogeneous to be useful in deriving upper crustal averages for Nd and Hf. We conclude that the same is likely true for determining the average Li composition of the UCC. By contrast, desert loesses are transported over greater distances (Ding et al., 1999), are less enriched in heavy minerals such as zircons (mostly Nd/Hf > 3.5, Chauvel et al., 2014; Fig. 3), and are well homogenized. As a consequence, their chemical composition is less biased, and they are better proxies for constraining the average composition of the UCC. We will thus only focus on the composition of the desert loesses in the following discussion.

5.2. Eolian mineralogical sorting controls Li within desert loess

It is essential to understand which processes control the variations of Li concentrations and isotopic compositions observed in the desert loess deposits and why each locality has a different average δ7Li (Fig. 3). Because Li is a soluble element (Brenan et al., 1998) it is affected by weathering processes and Li concentration may vary due to post-depositional alteration. However, in desert loesses, positive correlations between Li contents and immobile elements such as the rare earth elements (REE) (Fig. 4) suggest that Li has not been significantly remobilized by post-depositional chemical weathering, which we refer to as present-day weathering, i.e., transformation into soils (Dellinger et al., 2014).

Another process that can lead to Li isotopic variability is source rock heterogeneity within the provenance (Dellinger et al., 2014; Qiu et al., 2009), i.e., mafic versus felsic or juvenile versus old crustal material. Based on the observation that Argentinian loesses are enriched in clasts of volcanic rocks (andesites, basalts, dacites, rhyolites) (Imbellone and Teruggi, 1993; Teruggi, 1957) and that young volcanogenic sediments tend to have higher δ7Li (Bouman et al., 2004), one could link the observed variations in δ7Li in desert loesses to the provenance of the dust. This assumption can be evaluated using source proxies that are not significantly affected by sedimentary processes, such as the ratio of insoluble elements Th/La (Plank, 2005) that traces the contribution of felsic and mafic materials, or the Nd isotopic composition (Goldstein et al., 1984) that is controlled by the proportion of young to old mantle-derived igneous rocks in the source region. If we consider the less evolved volcanic rocks (i.e., basalts) that should create the largest variability, we can see that the average Th/La ratio of desert loess samples of 0.37 ± 0.04 (2σ), including the Argentinian samples, is similar to that of the UCC (~0.34 after Rudnick and Gao, 2014) but sig-

| Table 1 |
| Li isotopic compositions and concentrations of loess samples. |
| Sample name | Location | Longitude | Latitude | δ7Li (%e) | [Li] (ppm) |
| Western Europe (Periglacial loess) | | | | | |
| PR RT | Port Racine, Normandy, France | 1°53'W | 49°43'N | 2.1 | 17.9 |
| SAB 1a ~160 mm | Sables d’or, France | 2°24'W | 48°37'N | 4.7 | 16.6 |
| LO94 | Spitbergen (Svalbard) | 20°43'E | 77°40'N | −1.2 | 61.3 |
| PZ1E | Spitbergen (Svalbard) | | | −0.3 | 59.9 |
| SCIL | Scilly Island, England | 6°20'W | 49°55'N | −2.9 | 41.1 |
| SCIL dup | Scilly Island, England | | | −1.5 | |
| China (Desert loess) | | | | | |
| JX-4 | Jixian | 110°39'E | 36°06'N | 0.8 | 37.1 |
| JX-6 | Jixian | | | 0.2 | 37.4 |
| JX-10 | Jixian | | | | 0.5 | 39.9 |
| XF-10 | Xifeng | 107°42'E | 35°45'N | | |
| XF-6 | Xifeng | 101°48'E | 36°36'N | 1.1 | 37.5 |
| XN-4 | Xining | | | 2.6 | 36.3 |
| XN-10 | Xining | | | 2.6 | 38.1 |
| L9 | Luochuan | 109°26'E | 35°28'N | −0.1 | 37.9 |
| L6 | Luochuan | | | −0.2 | no data |
| L2 | Luochuan | | | 1.1 | no data |
| Tajikistan (Desert loess) | | | | | |
| TJK2772 | Tajikistan | 69°49'57''E | 38°23'32''N | −1.1 | 44.3 |
| TJK2773 | Tajikistan | | | −0.5 | 43.0 |
| TJK2930 | Tajikistan | | | −0.4 | 35.6 |
| TJK3012 | Tajikistan | | | −0.4 | 46.7 |
| TJK3070 | Tajikistan | | | 0.9 | 35.4 |
| TJK3148 | Tajikistan | | | −0.7 | 37.8 |
| TJK3179 | Tajikistan | | | −0.4 | 55.2 |
| TJK3198 | Tajikistan | | | 0.0 | 32.5 |
| Argentina (Desert loess) | | | | | |
| 12–14 | Argentina | 59°22'W | 34°38'S | 2.5 | 26.3 |
| 24–26 | Argentina | | | 2.2 | 26.1 |
| LUJA | Argentina | | | 0.2 | 36.7 |
| LUJA dup | Argentina | | | 4.1 | 30.6 |

δ7Li = (7Li/6Li)sample/(7Li/6Li)L-SVEC − 1) × 1000 in %e. L-SVEC data comes from Flesch et al. (1973).

* Li concentrations are from Chauvel et al. (2014). “dup” stands for complete duplicate analysis.
significantly different from the range known for Andean arc basalts (Fig. 5a). This suggests that the presence of mafic components in the provenance, such as Andean basalts in the Argentinian loess samples, is probably minimal and has had no effect on the Li isotopic compositions, as shown by the absence of correlation between Th/La and δ²⁷Li (Fig. 5a). Even though the Argentinian loesses tend to have higher δ²⁸Nd (∼−2), which is explained by the presence of juvenile material in their source (Chauvel et al., 2014; Gallet et al., 1998; Smith et al., 2003), the lack of correlation between the Nd and the Li isotopic compositions of our samples (Fig. 5b) suggests that the Li isotopic compositions and concentrations of desert loesses are not mainly controlled by source heterogeneity.

When sediments are transported by wind, the finest and lightest particles are preferentially transported over longer distances compared to coarse and dense particles that accumulate close to the source regions (Pye, 1995). This is manifest as differences in the average grain size (Yang and Ding, 2004), mineralogy (Eden et al., 1994; Yang et al., 2006), and chemical composition (Feng et al., 2009, 2010, 2011) as a function of distance from their source regions. This grain-size dependence of the chemical composition has also been observed in other types of sediments (Bouchez et al., 2011; Carpentier et al., 2014; Dellinger et al., 2014; Garçon et al., 2014, 2013; Garzanti et al., 2011). Indeed, as minerals have different chemical and isotopic compositions (Garçon et al., 2014, 2013, 2011), sediments with different grain-sizes, composed of different minerals in different proportions, will have different chemical compositions. For example, quartz is preferentially enriched in coarse-grained fractions (Garzanti et al., 2011, 2010), has very high δ²⁷Li of ∼+30 due to preferential enrichment of ⁷Li in the 2- or 4-fold sites in quartz (Dennenn, 1966; Maloney and Nabelek, 2008; Teng et al., 2006) and has relatively low [Li] of ∼10 ppm (Dennenn, 1966; Garçon et al., 2014; Lynton et al., 2005; Monecke et al., 2000; Teng et al., 2006). By contrast, clay minerals

Fig. 4. [Li] versus rare earth elements (REE) in desert loess samples. For each correlation we used a weighted linear regression (York et al., 2004) followed by a Monte-Carlo simulation to interpolate [Li] and its uncertainty at the average REE concentration of the UCC published by Rudnick and Gao (2014). By averaging all of these values, we infer the average [Li] of the UCC (see Supplementary File B for more details). Gray field represents uncertainties on the linear regressions derived from Monte Carlo simulations. Errors bars represent 2σ uncertainty.
The major minerals of desert loesses: quartz, plagioclase and phyllosilicates (Eden et al., 1994; Gallet et al., 1996; Jeong et al., 2008, 2011), can be mixed to produce the observed range of Li compositions (Fig. 6). Both Li concentrations and Li isotopic compositions are controlled by mineral sorting between a fine-grained end-member, enriched in phyllosilicates (clays and micas) – with a high [Li] and a low δ7Li – and a coarse-grained end-member, enriched in quartz and plagioclase – with a low [Li] and a high δ7Li (see caption of Fig. 6 for more details). Isotopic mixing calculations allow us to quantify the amount of unweathered, coarse-grained vs. weathered, fine-grained particles (Fig. 6). The results of the mixing calculations match the modal mineralogical proportions estimated from previous studies of the Chinese and Argentinian loess deposits (Camilón, 1993; Jeong et al., 2011, 2008; Teruggi, 1957) (Fig. 6). For example, the relative average proportions of phyllosilicates, quartz, and plagioclase given by our mixing calculations for the Argentinian loess are 40 wt.%, 36 wt.%, and 24 wt.% respectively, while the average modal proportions reported by Camilón (1993) and Teruggi (1957) are 40 wt.%, 33 wt.%, and 27 wt.% respectively. There are no estimated mineral proportions for the Tajikistan samples. Nevertheless, their average grain-size is smaller than the other loess deposits (~12 μm, Ding and Ding, 2003), which is consistent with the higher phyllosilicate proportion estimated from our isotopic mixing calculations (Fig. 6). Significant contributions from mafic minerals such as amphibole, which can contain appreciable Li (Marks et al., 2008), can be ruled out. Indeed, if we assume that our Li variations are partly controlled by mafic minerals (mainly amphibole), we cannot reproduce the Li variations observed in the desert loess deposits. Moreover, because the abundance of mafic minerals in desert loess samples is very low (proportions <4 wt.%, Eden et al., 1994; Jeong et al., 2008; Teruggi, 1957) relative to quartz, plagioclase and phyllosilicates, they cannot control the Li variations. Based on these observations, we conclude that the variations of δ7Li and [Li] in desert loesses can be understood in terms of relatively simple mixing between minerals of different grain size fractions, as shown in Fig. 6.

5.3. A new estimate of the δ7Li and [Li] in the average upper continental crust

Estimating the composition of the average UCC using sediments dates back to the work of Goldschmidt in the 1930s. This can be done either using an average composition of sediments, assuming that no processes had biased their composition compared to their source rocks, or using correlations observed in the concentrations of insoluble elements (McLennan, 2001). Because of the mineralogical sorting observed in our desert loess samples, it is not advisable to use their average composition as a representative value for the UCC and we focus here on the second method.

Fig. 4 shows well-defined correlations observed for desert loesses between Li and immobile elements, in this case, the REE. As REE are preferentially enriched in fine-grained sediments (Garçon et al., 2014; Taylor and McLennan, 1985), such trends reflect mixing lines between a fine-grained end-member (enriched in REE and Li) and a coarse-grained end-member (depleted in REE and Li). Using the correlations between Li and REE that are assumed to have the same behavior as Li during magmatic differentiation (i.e., from Sm to Er) (Ryan and Langmuir, 1987), we estimate a new average Li concentration for the UCC (Fig. 4). For each of the seven correlations (i.e., Li vs. Sm, Li vs. Eu, Li vs. Gd, Li vs. Tb, Li vs. Dy, Li vs. Ho and Li vs. Er; Fig. 4), we start with the raw data and first run a Monte-Carlo routine to estimate the uncertainties on the slopes and intercepts of the linear regressions. For each individual data point, we thus randomly sample a value within the

are enriched in the finest fractions (Garzanti et al., 2011, 2010) and have low δ7Li of ~−1 and high [Li] of ~63 ppm (Qu et al., 2009; Romer et al., 2014; Tsai et al., 2014). Consequently, one could expect fine-grained sediments, preferentially enriched in clays, to have higher Li concentrations and lower δ7Li than coarse-grained sediments that are preferentially enriched in quartz.

In desert loesses, the well-defined correlations between both [Li] and REE (Fig. 4), δ7Li and Nd/Hf (Fig. 3a), δ7Li and Fe2O3/SiO2 (Fig. 3b), as well as between Al2O3/Na2O and Fe2O3/SiO2 (Fig. 1), suggest that the bulk composition and most Li variations are related to the sediment grain-size. Indeed, Fe2O3, Al2O3 and REE are preferentially enriched in the fine-grained fraction of the loesses because phyllosilicates (clays and micas) are rich in iron, aluminium and REE (Garçon et al., 2014; Taylor and McLennan, 1985). By contrast, Na2O, SiO2 and Hf are mostly hosted in plagioclase, quartz and zircons, respectively, three mineral phases that are abundant in the coarse fraction (Eden et al., 1994; Garzanti et al., 2011; Yang et al., 2006). As a consequence, Nd/Hf, Al2O3/Na2O and Fe2O3/SiO2 ratios are excellent proxies for grain-sizes and reflect transport-driven compositional changes. This observation is consistent with the average grain-size of the samples as determined by previous studies. The Tajikistan samples are the finest loess samples (6 to 12 μm, Ding and Ding, 2003; Fig. 1) and have the highest [Li] and the lowest δ7Li values, while the Argentinian samples are the coarsest samples (>40 μm, Teruggi, 1957; Fig. 1) and have the lowest [Li] and the highest δ7Li values.

Fig. 5. a) δ7Li versus Th/La and b) δ7Li versus εNd in desert loess samples. The range of Th/La ratio for the UCC is from Rudnick and Gao (2014) (excluding data from Eade and Fahrig (1973) due to XRF-determined La leading to anomalously high values). The average εNd for the UCC is from Chauvel et al. (2014). Arrays represent the range of estimates for the UCC (gray) given by Rudnick and Gao (2014) for Th/La and by Chauvel et al. (2014) for εNd. Data for Andean basalts (green) are from the GEOROC database. Error bars are 2σ. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
uncertainty of the measured data point to generate a synthetic dataset. We then fit a straight line through all of the synthetic datasets using weighted linear regressions following the algorithm of York et al. (2004) that has been implemented in MATLAB™ by Thirumalachai et al. (2011). Within this first Monte-Carlo routine, we perform a second Monte-Carlo simulation to interpolate [Li] at the REE concentrations published by Rudnick and Gao (2014) for the UCC, taking into account the uncertainties on the REE concentrations published by Rudnick and Gao (2014) (see Supplementary File B for more details). We then compile all the interpolated [Li] and calculate both the average and the standard deviation of the distribution. Using this method, we obtain an average Li concentration of 30.5 ± 3.6 (2σ) ppm for the UCC, in agreement with the previously published values of Teng et al. (2004); 35 ± 11 (2σ) ppm and Rudnick and Gao (2014; 24 ± 10 ppm) but with a significantly lower uncertainty. Our value is somewhat lower than the derived by Hu and Gao (41 ± 6 (2σ) ppm) on the basis of correlations between Li and In observed in various sediments and sedimentary rocks.

Following the same procedure (weighted linear regression followed by interpolation using Monte-Carlo simulations), we estimate the average δ7Li of the UCC using the correlations between δ7Li and two independent ratios of immobile elements that are not significantly affected by chemical weathering processes, namely, Nd/Hf (Fig. 3a) and Fe2O3/SiO2 (Fig. 3b). These two ratios are good grain-size proxies (see previous section) and are relatively well known in the UCC (Nd/Hf = 5.1 ± 2.1 (2σ) and Fe2O3/SiO2 = 0.084 ± 0.02 (2σ), Rudnick and Gao, 2014). Interpolating these values on the regression lines, we get two sets of values having consistent average Li isotopic compositions: δ7Li = +0.8 ± 0.4 (2σ) using the correlation between δ7Li and Nd/Hf (Fig. 3a) and δ7Li = +0.4 ± 0.4 (2σ) using the correlation between δ7Li and Fe2O3/SiO2 (Fig. 3b). Combining these two sets of data, we obtain an average δ7Li = +0.6 ± 0.6 (2σ) (see Supplementary File B for more explanations). Here again, this new estimate is within the uncertainties of the previously published value of 0 ± 4 (2σ) (Teng et al., 2004), but with a much smaller uncertainty.

5.4. Determining the average weathering signature of upper continental crust

Chemical weathering is an important process affecting the composition of the continental crust (Dehlinger et al., 2014) because it involves the breakdown of rocks into secondary phases such as clays and hydroxides, and it releases soluble elements to the hydrosphere. It has been suggested that this process progressively modifies the chemical composition of the Earth’s surface (Albarède, 1998; Lee et al., 2008; Liu and Rudnick, 2011; Rudnick, 1995) but the degree of weathering experienced in the past within the average UCC and quantified by the proportion of weathered sedimentary rocks exposed at the Earth’s surface is still poorly known.

Silicate rock weathering is very important in the worldwide atmospheric CO2 consumption occurring today (Amiotte Suchet et al., 2003; Dessert et al., 2003; Gaillardet et al., 1999; Jin et al., 2014). Hence, through the disintegration of pre-existing rocks, chemical weathering acts as sink for atmospheric CO2, and so indirectly controls the evolution of our climate. Shales appear to have a significant influence on global CO2 consumption, accounting for 40% of the total CO2 consumed worldwide (Amiotte Suchet et al., 2003), but only a few studies have tried to quantify their abundances on continents and these studies show rather large discrepancies. Blatt and Jones (1975) estimate that 66% of the rocks exposed on the Earth’s surface are sedimentary rocks, but estimates for the proportion of shales are less well known and differ widely between 13 and 34% (Amiotte Suchet et al., 2003; Condie, 1993; Gibbs and Klump, 1994; Meybeck, 1987). An isotopic approach could potentially provide a more robust estimate of the average degree of weathering experienced by the UCC, and help us to understand the processes controlling the compositional evolution of the upper crust.
Fig. 7 shows how $\delta^7$Li varies as a function of CIA (Chemical Index of Alteration) in the loess deposits. The CIA, calculated as the ratio of major elements ($Al_2O_3/[Al_2O_3 + CaO + Na_2O + K_2O]$; see caption of Fig. 7 for more details), measures the loss of mobile cations ($Ca^{2+}, K^+$ and $Na^+$) present in labile minerals (feldspars, pyroxenes, amphiboles) relative to the amount of $Al^{3+}$ that is preserved in more stable minerals (i.e., clay) under surface conditions (Nesbitt and Young, 1982). Hence, the CIA is a good proxy for the intensity of chemical weathering: the higher the CIA, the greater the degree of weathering the sample has experienced in the past. In Fig. 7, we show that the range of $\delta^7$Li and CIA values obtained for the desert loess deposits follow a binary isotopic mixing trend between unweathered, mantle-derived igneous rocks (i.e., I-type granites) with high $\delta^7$Li and low CIA, and a weathered component (i.e., shales) with low $\delta^7$Li and high CIA. The Li composition ($\delta^7$Li = $-0.9 \pm 4.7$ (2$\sigma$); [Li] = $66 \pm 92$ (2$\sigma$) ppm) and CIA value (CIA = $74 \pm 21$ (2$\sigma$)) of the weathered end-member (shale) is compiled from several studies (Hu and Gao, 2008; Moriguti and Nakamura, 1998b; Qiu et al., 2009; Romer et al., 2014; Tang et al., 2014; Teng et al., 2004, and unpublished data of Su Li and others), while values for the unweathered end-member (Clarence River Super-suite I-type granite) come from Bryant et al. (2004) ($\delta^7$Li = $4.3 \pm 4.1$ (2$\sigma$), [Li] = $17 \pm 18$ (2$\sigma$) ppm and CIA = $54 \pm 8$ (2$\sigma$)). These granites differentiated from mantle-derived magma and have not incorporated any meta-sedimentary materials (Bryant et al., 2004), which is important as the unweathered end-member should not include weathered materials, which many other I-type granites do (Teng et al., 2004). Unfortunately, the compositions of the two end-members are not very well constrained (large uncertainties on the averages) and the large errors can have a non-negligible impact on our isotopic mixing calculation. However, the very good consistency observed between the isotopic modeling and our data suggests these average values of the two end-members are realistic.

The UCC is considered to have a granodioritic composition (Arndt, 2013; Eade and Fahrig, 1973; Rudnick and Gao, 2014; Taylor and McLennan, 1985) and juvenile crust should have the high $\delta^7$Li and low CIA values shown in Fig. 7 for igneous rocks that are devoid of a sedimentary component in their source regions. As its composition is modified by chemical weathering, the Li isotopic composition of the UCC decreases, with a concomitant increase of its CIA, reaching, in extreme cases, the very low $\delta^7$Li and high CIA values found in shales (Fig. 7). It is therefore reasonable to expect that the present-day UCC falls on a mixing line between an unweathered component (granites) and a weathered one (shales). Using our estimate for the average $\delta^7$Li of upper crust and the results of the isotopic mixing, we derive the average CIA of present-day UCC to be between 59 and 65 with an average of 61 (Fig. 7). This new value differs from the average value given by Rudnick and Gao (2014) for the upper crust (CIA = 53, calculated using the average major element concentrations), and which is derived from a compilation of several studies, mainly based on sampling widespread outcrop exposures. This discrepancy may reflect inadequate sampling methods, as a majority of these studies focused on a limited part of the world, e.g., North Craton China (Gao et al., 1998) and Canadian Precambrian shield (Eade and Fahrig, 1973; Shaw et al., 1976, 1967). The low CIA may thus reflect local composition where the proportion of weathered rocks is lower than the average UCC. Or, as these studies rely only on outcrops at the Earth’s surface, the discrepancy can be explained by the presence of under-sampled sedimentary rocks that occur in fold belts and continental platforms, as previously suggested by Amiotte Suchet et al. (2003).

Our new CIA estimate is also significantly lower than the value estimated for Archean UCC (average CIA = 77) and for Paleoproterozoic UCC (average CIA = 67) as determined by Gaschnig et al. (2014) using ancient glacial diamictites. The higher CIA in the past has been interpreted as resulting from more intense weathering (Gaschnig et al., 2014). Condie (1993) reached a similar conclu-
sion on the basis of the depletion in Na, Ca and Sr in Archean shales, which he suggested reflected more intense chemical weathering during the Archean. Therefore, the difference between the present-day and ancient CIA of the UCC may reflect a change in the intensity of chemical weathering through time that modified the chemical composition of the UCC.

Another approach to quantify the importance of past chemical weathering is to determine the amount of weathered products produced by this process. Our novel isotopic approach shows that the chemical composition of the present upper continental crust, having a $\delta^{11}Li$ of $+0.6 \pm 0.6$ and a CIA of 61, can be explained by a mixture of $63^{+10}_{-17}(2\sigma)$% of unweathered igneous rocks (granites) and $37^{+10}_{-16}(2\sigma)$% of rocks produced by weathering (shales) (cf. Fig. 7). The inferred proportion of shales is slightly higher than previous estimates. Amiotte Suchet et al. (2003) have shown that ~25% shales present at the Earth’s surface accounts for about 40% of the CO$_2$ drawdown in the atmosphere. With ~40% shales at the surface, as estimated here, almost 60% of the worldwide CO$_2$ could be consumed, highlighting how important weathering may be for the overall CO$_2$ budget of the Earth. This deserves more attention, particularly from a climate studies point of view. Thus, both means of estimating the average weathering signature of the UCC (average CIA and mixing of shales and granites) demonstrate that weathering has had a profound influence on the average composition of the UCC and, based on the relationship between weathering and CO$_2$ draw-down, the continental crust has, in turn, profoundly influenced climate.

6. Summary and conclusions

Based on analyses of global loess samples, periglacial loess deposits are too heterogeneous to be used as proxies to estimate the Li composition of the upper continental crust. Desert loess deposits appear to be more homogenous and thus more suitable to infer average Li values. The Li isotopic compositions and concentrations of desert loesses are controlled by mineralogical sorting and can be reproduced by mixing between a fine-grained, weathered end-member (i.e., phyllosilicates) and a coarse-grained, relatively unweathered end-member (i.e., a mixture of quartz and plagioclase). Using the correlations between Li and REE concentrations in the desert loesses we derive a new estimate for the average Li concentration of the upper crust: $30.5 \pm 3.6(2\sigma)$ ppm. Similarly, using correlations between $\delta^{11}Li$ and independent ratios of immobile elements, we estimate an average $\delta^{11}Li = +0.6 \pm 0.6(2\sigma)$ for the UCC. These results are similar to previous estimates, but are more precisely constrained, and show that the UCC carries a significantly more weathered signature than fresh mantle-derived rocks. Using an isotopic mixing approach we also quantify the cumulative importance of chemical weathering on the continents over Earth history. We estimate that $37^{+10}_{-16}(2\sigma)$% of the current upper crust is composed of highly weathered sediments (i.e., shales) that are the products of chemical weathering experienced in the past and that the UCC’s present CIA is $61^{+10}_{-17}$, which is lower than what it was in the Paleoproterozoic and Archean periods, but higher than previous estimates of the UCC. Our results thus provide a framework from which to compare the UCC through time.

Acknowledgements

We thank S. Gallet and B.M. Jahn for providing the samples analyzed in this study. We also greatly thank S. Bureau for major element analyses, I.S. Puchtel for his help in the clean laboratory, as well as R.D. Ash for his assistance during isotopic measurements. Thanks also to X.-M. Liu and W.F. McDonough for constructive discussions and advice that helped improve the content of the manuscript. We thank reviewers Mathieu Dellingner and Edward Tipper for their comments that helped improve the manuscript and editor Mike Bickle for his pleasant comments and efficient editorial handling. This work was supported by NSF grant EAR 0948549 and General funds from INSU-CNRS, University of Grenoble and Rhône-Alpes Region.

Appendix A. Supplementary material

Supplementary material related to this article can be found online at http://dx.doi.org/10.1016/j.epsl.2015.07.032.

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