Empirical Evidence for Latitude and Altitude Variation of the In Situ Cosmogenic $^{26}$Al/$^{10}$Be Production Ratio

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Abstract: We assess if variations in the in situ cosmogenic $^{26}$Al/$^{10}$Be production ratio expected from nuclear physics are consistent with empirical data, knowledge critical for two-isotope studies. We do this using 313 samples from glaciated transported boulders or scoured bedrock with presumed simple exposure histories in the Informal Cosmogenic-nuclide Exposure-age Database (ICE-D) from latitudes between 53°S to 70°N and altitudes up to 5000 m above sea level. Although there were small systematic differences in Al/Be ratios measured in different laboratories, these were not significant and are in part explained by differences in elevation distribution of samples analyzed by each laboratory. We observe a negative correlation between the $^{26}$Al/$^{10}$Be production ratio and elevation ($p = 0.0005$), consistent with predictions based on the measured energy dependence of nuclear reaction cross-sections and the spatial variability in cosmic-ray energy spectra. We detect an increase in the production ratio with increasing latitude, but this correlation is significant only in a single variate model, and we attribute at least some of the correlation to sample elevation bias because lower latitude samples are typically from higher elevations (and vice versa). Using 6.75 as the $^{26}$Al/$^{10}$Be production ratio globally will bias two-isotope results at higher elevations and perhaps higher latitudes. Data reported here support using production rate scaling that incorporates such ratio changes, such as the LSDn scheme, to minimize such biases.

Keywords: cosmogenic nuclides; nuclide production; burial dating

1. Introduction

Paired-nuclide, in situ cosmogenic nuclide analyses are valuable tools for investigating complex landscape histories, including burial after and/or during exposure. In situ cosmogenic nuclides are formed in minerals at the Earth’s surface when exposed to the high-energy particle cascade produced during interactions between cosmic radiation and atmospheric gases [1]. Differences in production and decay ratios between multiple in situ cosmogenic radionuclides are used to estimate burial/exposure durations and erosion histories—work applications ranging from non-erosive glacier histories [2–4], to long-term fluvial incision [5,6] and archaeological investigations [7–9].

An essential component of this methodology is knowing with certainty the production ratio between measured in situ cosmogenic radionuclides. While the decay rates of cosmogenic radionuclides have been empirically constrained [10–12], the production rates of cosmogenic nuclides, and thus their production ratios, are estimated using models of the relevant physics [13] and validated with calibration studies that rely on independent age constraints of landscape features [14,15].

Two of the most-used in situ cosmogenic radionuclides in dual-nuclide studies are $^{26}$Al and $^{10}$Be. Both are produced in quartz, and their ratio has been measured since the 1980s [16]. The near-ubiquity of quartz across the world and the improved analytical precision for $^{10}$Be—and more recently $^{26}$Al [17]—measurements make these nuclides...
the “go to” for dual-nuclide studies. The improvement in measurement precision will allow for more useful interpretation of the data, but only if the production ratio is well constrained [18].

Physics-based nuclide production models suggest that the $^{26}\text{Al}/^{10}\text{Be}$ surface production ratio should decrease with elevation and increase with latitude [19,20]; however, most analyses of $^{26}\text{Al}/^{10}\text{Be}$ data assume a globally constant surface production ratio. Empirical evidence from high-latitude sites [21] suggest that this assumption may not be valid. A surface production ratio that changes with latitude and/or elevation would mean that many of the studies using this dual-nuclide methodology contain systematic biases in their results because the assumed surface production ratio may differ from the actual ratio at the sampling site. As cosmogenic nuclide measurements become more precise, better constraining of the production ratio becomes more important.

Here, we assess if changes in the $^{26}\text{Al}/^{10}\text{Be}$ production ratio with latitude and/or elevation are detectable in empirical data using a compilation of 313 previously published in situ $^{26}\text{Al}/^{10}\text{Be}$ ratios from samples spanning a wide range of latitudes and elevations. We applied selection criteria to increase the chances that these glacially related samples have experienced simple exposure histories—that is, only one short (<25 kyr) period of exposure and no burial, so that the measured ratio ($^{26}\text{Al}/^{10}\text{Be}$) represents the surface production ratio. We apply deductive statistical analyses to this compilation to determine if $^{26}\text{Al}/^{10}\text{Be}$ ratio variations with altitude, latitude, and sample processing laboratory are present at a statistically significant level. This analysis allows us to test whether there are detectable production ratio variations, thus necessitating application of physics-based nuclide production models in dual-nuclide studies, or if the nominal ratio of 6.75 is suitable at all latitudes and elevations.

2. Background

2.1. In Situ Cosmogenic $^{26}\text{Al}$ and $^{10}\text{Be}$

In this study, we focus on the production of in situ cosmogenic $^{26}\text{Al}$ and $^{10}\text{Be}$ in quartz. $^{26}\text{Al}$ and $^{10}\text{Be}$ are produced primarily through spallation reactions in quartz (>95% at sea level and high latitude) with minor production from muon interactions [1,11,12]. The ratio of muonic to spallation production increases with depth below Earth’s surface [1] and decreases at higher elevations. $^{26}\text{Al}$ has a half-life of 0.705 million years [10] and $^{10}\text{Be}$ has a half-life of 1.39 Ma [11,12].

The accuracy of measured $^{26}\text{Al}/^{10}\text{Be}$ ratios as a proxy for the $^{26}\text{Al}/^{10}\text{Be}$ ratio at production is controlled by both laboratory procedures and the geologic history of sample sites. Nuclides inherited from prior periods of exposure both at the surface and at depth influence the concentration of $^{26}\text{Al}$ and $^{10}\text{Be}$ in surface samples and thus the measured $^{26}\text{Al}/^{10}\text{Be}$ ratio. Muon-induced production produces relatively few nuclides but does so at a ratio higher than surface production, which is dominated by neutrons [22]. Storage of previously exposed material at depths below the penetration depth of most neutrons can lower measured ratios as $^{26}\text{Al}$ decays more quickly than $^{10}\text{Be}$ [10].

Laboratory concerns include the measurement of cosmogenically produced isotopes ($^{26}\text{Al}$ and $^{10}\text{Be}$) by Accelerator Mass Spectrometry (AMS), which often sets the limit on precision, and quantification of stable isotopes ($^{27}\text{Al}$ and $^{9}\text{Be}$), which is critical for accuracy. Low-energy AMS machines may be unable to completely reject isobaric interferences encountered in $^{26}\text{Al}$ analyses, thus artificially increasing calculated $^{26}\text{Al}$ concentrations and the $^{26}\text{Al}/^{10}\text{Be}$ ratio [23]. When complexed with fluoride during HF digestion, Al can be difficult to get back into solution, thus leading to underestimation of $^{26}\text{Al}$ and consequently low $^{26}\text{Al}/^{10}\text{Be}$ ratios [24]. Stable beryllium is added as a carrier (isotope dilution) but stable aluminum is native to the quartz being digested, meaning that full retention and recovery of that aluminum is critical to accurately quantifying the concentration of $^{26}\text{Al}$. Stable aluminum quantification errors can arise from chemical processing steps, including aliquot measurements after, rather than before, drying dissolved samples, adding sulfuric acid to digestion solutions, and systematic offsets in calibration of inductively coupled plasma
optical emission spectrometers for measuring $^{27}$Al concentrations [25]. Low recovery of $^{27}$Al will result in lower than actual $^{26}$Al/$^{10}$Be ratios [24]. Significant variation in measured $^{26}$Al between chemistry labs has been observed during inter-lab comparisons, and much of this variation is attributed to differences in methodology for quantifying $^{27}$Al [26–28]. A small number of quartz mineral separates contain significant amounts of stable $^9$Be, which, if unaccounted for, would result in spuriously low $^{10}$Be concentrations and high $^{26}$Al/$^{10}$Be ratios, and few laboratories routinely measure beryllium in quartz [29].

2.2. Applications

Paired in situ cosmogenic $^{26}$Al and $^{10}$Be are used in a wide variety of studies seeking to understand burial and erosion histories. Early applications included the history of enigmatic Libyan desert glass [16], the age of ancient glacial deposits in the Sierra Nevada mountains [30], and the glacial history of Antarctica [31]. Bierman et al. [2] demonstrated the utility of the paired-nuclide approach for calculating minimum total durations of exposure and burial for complex glacial histories in temperate regions—those involving multiple periods of advance and retreat. Investigations of complex glacial histories continue to be a common application of $^{26}$Al/$^{10}$Be [32,33], including cases with minimal glacial erosion due to cold-based ice cover [4,34] and glacial histories inferred from marine sediment records [35,36]. $^{26}$Al/$^{10}$Be has also been used to evaluate long-term erosion rates in arid regions [37], histories of tectonic uplift [38], river incision [5,6], and paleosol burial [39,40]. $^{26}$Al/$^{10}$Be is also used for age control in archaeological investigations of hominin evolution [7–9,41], providing burial ages of bones and artifacts.

2.3. Previous Constraints on $^{26}$Al/$^{10}$Be Production Ratio

The production ratio of $^{26}$Al/$^{10}$Be at Earth’s surface has been constrained in two ways: experimental measurements—sampling surfaces with ‘known’ exposure histories—and physics-based models that simulate interactions between cosmic radiation, atmospheric atoms, and terrestrial atoms in target minerals. See Corbett et al. ([21] and Table S1 therein) for a summary of these studies and the production ratios they calculated and measured. Studies published prior to 1991 estimated an $^{26}$Al/$^{10}$Be production ratio of ~6.1 [30,42], which was updated to 6.75 following updates to accelerator standards and refinements of the $^{10}$Be half-life [11,12,43]. Early models of $^{26}$Al and $^{10}$Be production indicated a spallation production ratio of 6.05 that did not change with elevation (latitude changes in in situ $^{26}$Al production were not modelled [44]). Without robust empirical evidence to support the use of more complex numerical models for nuclide production, an $^{26}$Al/$^{10}$Be production ratio of 6.75 is typically assumed to be constant over all latitudes and elevations in many scaling schemes used for cosmogenic nuclide data interpretation [45].

2.4. Indication of Spatial Variability in the $^{26}$Al/$^{10}$Be Production Ratio

Cross-sections for nuclide production from spallation reactions suggest changes in the $^{26}$Al/$^{10}$Be production ratio with latitude and elevation [19,20,46–48]. Although $^{26}$Al and $^{10}$Be are both produced in quartz at Earth’s surface primarily through spallation reactions, the cross section—or likelihood of reaction—for spallation production is different for each nuclide. $^{26}$Al has a lower energy threshold for production than $^{10}$Be, so neutron fluxes with different energy spectra produce $^{26}$Al and $^{10}$Be at different ratios [47,48]. Earth’s geomagnetic field deflects lower-energy components of the primary cosmic ray flux more readily at lower latitudes [46]. The energy spectrum of the secondary neutron flux that reaches Earth’s surface therefore differs with latitude, implying a lower $^{26}$Al/$^{10}$Be production ratio at low latitudes, where the more energetic neutron flux favors $^{10}$Be production, and a higher production ratio at high latitudes, where the less energetic neutron flux favors $^{26}$Al production [19].

A similar dynamic is expected with changes in elevation due to changes in the neutron flux energy spectrum with depth in the atmosphere. The cosmogenically derived neutron flux loses energy with increasing atmospheric depth due to interactions with atmospheric
Thus, the $^{26}\text{Al}/^{10}\text{Be}$ production ratio should be lower at high elevations, where the neutron flux has higher energy and favors $^{10}\text{Be}$ production, and highest at low elevations, where the lower energy neutron flux favors $^{26}\text{Al}$ production [46].

3. Materials and Methods

3.1. Data Sources and Sample Selection

To determine if the assumption of a constant $^{26}\text{Al}/^{10}\text{Be}$ production ratio is an oversimplification, we test for spatial heterogeneity in the $^{26}\text{Al}/^{10}\text{Be}$ production ratio using previously published samples in the Informal Cosmogenic-nuclide Exposure-age Database (ICE-D; ice-d.org, n = 313; ref. [49]). Within the ICE-D database, we extracted data from ICE-D: Alpine, data from alpine glacial landforms (n = 243), and ICE-D: Calib, samples used to calibrate cosmogenic-nuclide production rates by assuming exposure ages based on other geologic constraints (n = 70). $^{26}\text{Al}$ and $^{10}\text{Be}$ concentration measurements for all samples were normalized to the KNSTD and 07KNSTD standards, respectively [10,43]. In both sub-databases, we targeted samples that likely experienced simple exposure histories, such that all $^{26}\text{Al}$ and $^{10}\text{Be}$ are from a single exposure extending to the present day with no nuclides remaining from periods of prior exposures. In other words, we presume that the measured $^{26}\text{Al}/^{10}\text{Be}$ concentration ratios equal the $^{26}\text{Al}/^{10}\text{Be}$ production ratios for the samples we selected.

We applied the following criteria: 1. We selected samples in ICE-D: Alpine with reported $^{26}\text{Al}$ and $^{10}\text{Be}$ concentration measurements and with exposure ages under 25 ka. Querying for ages under 25 ka ensures that measured concentrations are from a single, short period of near-surface exposure. We do not include samples from Antarctica, where prolonged burial that alters measured $^{26}\text{Al}/^{10}\text{Be}$ ratios is evident in many samples (e.g., [50]). 2. We extracted all sample data from ICE-D: Calib and calculated exposure ages using the reported $^{10}\text{Be}$ concentrations, LSDn scaling, and the default exposure age calculator settings in version three of the online exposure age calculator described by Balco et al. [45] (i.e., without the reference production rate from the calibration site). We kept samples in our analysis if their calculated ages using these exposure age calculator settings matched the expected ages from nearby geologic calibration sites, indicating little inherited $^{10}\text{Be}$ (and by association $^{26}\text{Al}$) was present in these samples.

To avoid samples affected by geologic and/or laboratory processes that can skew ratios, we discarded samples from our initial query with physically unreasonable $^{26}\text{Al}/^{10}\text{Be}$ ratios. To account for the inevitable scatter in ratios due to analytical uncertainty of $^{26}\text{Al}$ and $^{10}\text{Be}$ measurements, we first fit a normal distribution to the $^{26}\text{Al}/^{10}\text{Be}$ ratio uncertainties in our compilation (Figures S2 and S3) and calculate the ratio uncertainty mean and standard deviation. We use the uncertainty mean (10.1%) plus one standard deviation (9.0%) as a threshold, beyond which we deemed the ratios physically unreasonable. Applying this 19% analytical uncertainty threshold to the canonical $^{26}\text{Al}/^{10}\text{Be}$ production ratio value of 6.75 gives a range of 5.47 to 8.03 for accepted $^{26}\text{Al}/^{10}\text{Be}$ ratios (details in Supplement). We assume that outlier samples were affected by geologic and/or laboratory processes that skew measured $^{26}\text{Al}/^{10}\text{Be}$ concentration ratios such that they do not reflect the surface production ratio. Our final tally of samples used in statistical analyses (n = 313) does not include outliers (n = 48) removed from the original ICE:D query (Figure S1).

3.2. Statistical Analyses

We use single and bi-variate linear models, Monte Carlo simulations, and analysis of variance (ANOVA) tests to determine if variations in the measured $^{26}\text{Al}/^{10}\text{Be}$ concentration ratios in our compilation are correlated with elevation and/or latitude. We first divide the sample population into three latitude transects and five elevation transects to isolate latitude and elevation as variables and create sample groups for ANOVA testing. Each elevation transect is a bin of samples from similar latitudes but spanning a range of elevations, while each latitude transect is a bin of samples with similar elevations but varying latitudes (Figure 1).
We use single and bi-variate linear models to determine if there are statistically significant correlations between the $^{26}\text{Al}/^{10}\text{Be}$ production ratio, latitude, and elevation. We run single regression linear models for elevation and latitude (as absolute latitude) vs. measured $^{26}\text{Al}/^{10}\text{Be}$ ratios first with the entire compilation and then with samples separated into transects, calculating 95% confidence intervals, correlation coefficients ($r$), and $p$-values against a null model (no variation in production ratio) for each iteration. We assume no uncertainty on latitude or elevation measurements. The bi-variate regression model includes elevation and latitude variables and is run for the entire compilation of samples.

We run Monte Carlo analyses to assess the influence of $^{26}\text{Al}/^{10}\text{Be}$ concentration ratio uncertainties on linear regressions. In each Monte Carlo analysis, we run 1000 iterations of linear regression with samples randomly adopting a $^{26}\text{Al}/^{10}\text{Be}$ concentration ratio value from within their uncertainty bounds (assuming a Gaussian uncertainty distribution) in each iteration. Monte Carlo analyses allow us to constrain a population of regressions using the uncertainties on each data point, producing another type of confidence interval that incorporates data uncertainty.

To assess if $^{26}\text{Al}/^{10}\text{Be}$ concentration ratios differ between transects, we perform ANOVA testing with transects as groups. If ANOVA testing indicates that one or more transect $^{26}\text{Al}/^{10}\text{Be}$ concentration ratio means are different at a 5% significance level, we perform multiple pairwise comparison of the concentration ratio means [51] to determine which transects differ and the statistical significance of differences. To assess if different cosmogenic nuclide sample preparation labs have an influence on measured $^{26}\text{Al}/^{10}\text{Be}$ concentration ratios, we also perform ANOVA testing with the five labs that processed the greatest number of samples in this compilation. The five labs are located at Lawrence Livermore National Laboratory (LLNL), the University of Washington (UW), the Swiss Federal Institute of Technology in Zürich (ETH), the Australian Nuclear Science and Technology Organization (ANSTO), and the Purdue Rare Isotopes Measurement Laboratory (PRIME). Together, these labs are responsible for the $^{10}\text{Be}$ and $^{26}\text{Al}$ extraction of 70% of the samples in our compilation.
4. Results

4.1. Compilation Statistics

$^{26}\text{Al}/^{10}\text{Be}$ concentration ratios in our sample compilation approximate a normal distribution with $\mu = 6.57$ and $\sigma = 0.52$ (Figure 2). However, our compilation has spatial bias. While samples are present at most elevations between sea level and 5000 m asl (Figure 3B), they are biased towards the mid-latitudes in both hemispheres, with particular density around the northern mid-latitudes (Figure 3B). There are no samples from low latitude/low elevation or high latitude/high elevation locations (Figure 1). Sample processing year does not have an observable impact on the measured $^{26}\text{Al}/^{10}\text{Be}$ ratio (Figure S10).

Figure 2. (A) Histogram and normal distribution approximation of $^{26}\text{Al}/^{10}\text{Be}$ concentration ratios in our sample compilation analyzed here. Dashed vertical line shows the currently accepted production ratio value of 6.75. (B) Residuals of bi-variate regression model predicting $^{26}\text{Al}/^{10}\text{Be}$ variations due to elevation and latitude.

Figure 3. Measured $^{26}\text{Al}/^{10}\text{Be}$ concentration ratios plotted against elevation (A) and latitude (B). Dashed horizontal line shows the currently accepted production ratio value of 6.75. Samples from the ICE:D—Alpine database are colored blue, samples from ICE:D—Calib are red.
4.2. Regression Statistics

A simple linear regression of measured $^{26}$Al/$^{10}$Be ratios vs elevation is consistent with lowering of the $^{26}$Al/$^{10}$Be production ratio with elevation. The elevation regression shows a statistically significant ($p = 0.000028$) negative correlation ($r = -0.23$) between elevation and measured $^{26}$Al/$^{10}$Be ratios (Figure 4A). Although large, the residuals from the elevation/ratio regression are normally distributed; there is no evidence of heteroscedasticity. The regression exhibits good fit to the data, as indicated by a reduced chi-squared test, which accounts for scatter caused by uncertainty in the data ($\chi^2 = 1.27$). The Monte Carlo regressions support a negative correlation, with every regression exhibiting a negative slope and with the 95% confidence interval (95% of regressions) overlapping the ratio change expected with elevation in nuclide production models (change in ratio = −0.083 per km elevation, calculated from [20]).

![Figure 4](image_url)

**Figure 4.** Linear regressions correlating measured $^{26}$Al/$^{10}$Be concentration ratios to elevation (A) and latitude (B). Gray data points are the same as Figure 3. Central, solid black line in each figure is the most likely regression. Thin, blue lines are individual Monte Carlo regressions. Gray horizontal line shows a ratio of 6.75. Dashed yellow line in (A) shows the expected change in ratio with elevation from LSDn scaling at 5° latitude. Solid red line in (A) is the same but at 60° latitude.

A linear model with measured $^{26}$Al/$^{10}$Be ratios and latitude (as absolute latitude) supports an increase in the production ratio with increasing latitude. The latitude/ratio regression shows a statistically significant ($p = 0.0025$) positive correlation ($r = 0.17$) between latitude and $^{26}$Al/$^{10}$Be ratios (Figure 4B). This latitude/ratio regression has normally distributed residuals with no evidence of heteroscedasticity and exhibits good fit to the data ($\chi^2 = 1.24$). The 95% confidence interval of the change in ratio with latitude as provided by Monte Carlo regressions overlaps the change expected from nuclide production models (change in ratio = 0.0053 per degree latitude, calculated from [20]).

Our bi-variate linear model with elevation and latitude as variables (Table 1) is a statistically significant improvement over a null model ($p = 0.00016$) and supports a negative correlation between elevation and $^{26}$Al/$^{10}$Be production ratio but does not support a positive correlation between latitude and $^{26}$Al/$^{10}$Be production ratio (Table 1). The bi-variate model has normally distributed residuals (Figures 2B and S5), no evidence of heteroscedasticity (Figure S7), and fits the data well ($\chi^2 = 1.24$), offering a marginal improvement over the elevation-only regression. The change in $^{26}$Al/$^{10}$Be ratio with elevation in the bi-variate model is within the 95% Monte Carlo confidence range of the elevation-only model, close to the change predicted by a nuclide production model [20], and is statistically significant ($p = 0.004$). The change in $^{26}$Al/$^{10}$Be ratio with latitude in the bi-variate model is not statistically significant ($p = 0.947$). This model indicates that...
there is variation in measured $^{26}\text{Al}/^{10}\text{Be}$ concentration ratios over space, and that elevation differences appear to have the strongest correlation to these changes.

### Table 1. Bivariate regression statistics table with elevation and latitude as variables ($y = x_1 + x_2 \times \text{elev} + x_3 \times \text{lat}$).

|        | Estimate | SE     | tStat | p Value |
|--------|----------|--------|-------|---------|
| Intercept | 6.86    | 0.24   | 28.26 | $8.74 \times 10^{-88}$ |
| Elevation | $-8.97 \times 10^{-5}$ | $3.09 \times 10^{-5}$ | $-2.91$ | 0.004 |
| Latitude | $-3.02 \times 10^{-4}$ | 0.005 | $-0.07$ | 0.947 |

F-statistic vs. constant model: 9, $p$-value = 0.00016.

Linear regressions in elevation and latitude transects support $^{26}\text{Al}/^{10}\text{Be}$ ratio variations in only one transect. Measured $^{26}\text{Al}/^{10}\text{Be}$ ratios in elevation transect 1 (spanning latitudes $40^\circ$–$50^\circ$S; Figure 1) exhibit a statistically significant ($p = 0.011$) negative correlation ($r = -0.34$) with elevation. Correlations in every other transect are not significant at the 5% level and exhibit wide 95% confidence intervals (Figure S4).

### 4.3. ANOVA

ANOVA tests indicate that there is a statistically significant difference in measured $^{26}\text{Al}/^{10}\text{Be}$ ratios between samples from the highest and lowest elevations. Both the mean and median $^{26}\text{Al}/^{10}\text{Be}$ concentration ratio from latitude transect 1 (200 to 600 m asl) and 3 (4000 to 5000 m asl) are significantly different ($p = 4.96 \times 10^{-5}$; Figure 5). Moreover, the ratio differences are as predicted by nuclear physics models, with the lower elevations having a higher $^{26}\text{Al}/^{10}\text{Be}$ ratio (mean $\pm$ SE = $6.80 \pm 0.12$, median = 6.95) than higher elevations (mean = $6.49 \pm 0.11$, median = 6.48, Figure 5). The mean and median $^{26}\text{Al}/^{10}\text{Be}$ concentration ratio from latitude transect 2, covering the 1400 to 1800 m asl elevation band, is different than latitude transect 1 at the 5%, but not 1%, significance level ($p = 0.025$) and is not significantly different than latitude transect 3 (Figure 5).

![Figure 5](image-url)
Mean $^{26}$Al/$^{10}$Be concentration ratios from elevation transects 1 (6.76 ± 0.14) and 3 (6.45 ± 0.12), which cover 40° to 50°S and 36° to 39°N, respectively, are significantly different ($p = 0.008$; Figure 6). Every other elevation transect is statistically similar. Aside from transect 3, all elevation transects also have mean $^{26}$Al/$^{10}$Be concentration ratios that overlap the canonical value of 6.75 within the envelope of mean standard errors (Figure 6B).

**Figure 6.** ANOVA boxplots (A) and multiple comparison of means (B) for elevation transects. See Figure 5 for explanation of each figure. Red cross in (A) is an outlier.

Sample preparation lab ANOVA testing revealed that samples from the University of Washington Cosmogenic Nuclide Laboratory (UW) have higher mean and median $^{26}$Al/$^{10}$Be concentration ratios than other labs (Figure 7A). Sample ratios from UW were different than samples processed at Lawrence Livermore National Laboratories (LLNL), but not other labs, at a statistically significant level ($p = 0.006$).

**Figure 7.** (A) ANOVA boxplots for sample preparation labs, see Figure 5 for explanation. Note that the PRIME box appears distorted because the upper limit of the median comparison interval exceeds the 75th percentile value. (B) Elevations of samples processed by five different sample preparation laboratories.
5. Discussion

The $^{26}\text{Al}/^{10}\text{Be}$ production ratio changes with elevation as numerical models of the underlying nuclear physics predict. Our analysis of 313 glacially eroded and exposed samples from around the world, for which we assume simple exposure histories, supports the calculations and conclusions of Lifton et al. [20] and Argento et al. [19]. These numerical models predict a decrease in the $^{26}\text{Al}/^{10}\text{Be}$ production ratio from sea level to 5000 m asl of ~4% and 2.5%, respectively (see Figure 8 in [20]), and our ANOVA results agree well with the model predictions (4.6 ± 0.7% difference between latitude transects 1 and 3). Our bi-variate regression produces a good fit to measured $^{26}\text{Al}/^{10}\text{Be}$ ratios ($\chi^2_\nu = 1.24$), and the negative correlation between elevation and ratio is highly significant (Table 1).

The positive correlation between the $^{26}\text{Al}/^{10}\text{Be}$ ratio and latitude is less robust in our data than the negative correlation with elevation. The statistically significant correlation observed in the latitude/ratio regression (Figure 4B) is not replicated in the bi-variate model (Table 1) and ANOVA tests are inconclusive, with only two latitude bands of the five elevation transects exhibiting a statistically significant difference in $^{26}\text{Al}/^{10}\text{Be}$ (Figure 6).

These two elevation transects are not substantially different in terms of absolute latitude, with transect 1 covering 40°–50°S and transect 3 covering 36°–39°N (Figure 1), and we attribute at least some of the difference in ratios to sample elevation differences between the transects. Despite the intention for transects to isolate latitude and elevation as variables, these two transects contain samples from different elevations. Samples in elevation transect 1 range from sea level to ~1500 m asl; samples in elevation transect 3 range from ~1500 to 5000 m asl. Numerical model predictions and the statistically significant elevation/ratio correlation in our analyses suggest that $^{26}\text{Al}/^{10}\text{Be}$ ratios in elevation transect 1 should be several percent higher than ratios in elevation transect 3 just due to elevation differences. Thus, the 1 to 5% difference in mean $^{26}\text{Al}/^{10}\text{Be}$ ratios observed between these transects in ANOVA results is at least partially due to elevation-related differences in production.

The higher ratios observed in samples processed at UW do not skew our interpretation of elevation and latitude influences on the $^{26}\text{Al}/^{10}\text{Be}$ production ratio, and we attribute the higher ratios partially to differences in the elevation of samples processed in these two labs. To assess the leverage of the higher-ratio UW samples, we created a bi-variate regression model with these samples removed and found no significant difference in our results. Both the elevation/ratio correlation and the model itself were still statistically significant, although the elevation/ratio correlation was not as robust as when the UW samples are included ($p = 0.02$ vs. $p = 0.004$; Table S1). Samples processed at UW are from low elevations, with more than 50% from below 500 m asl, while the only lab with significantly different ratios, LLNL, has samples from predominantly high elevation locations (Figure 7B). The negative correlation between elevation and $^{26}\text{Al}/^{10}\text{Be}$ ratios demonstrated here and predicted by Lifton et al. [20] could thus be partially responsible for the observed difference between UW and LLNL results.

Differences in sample processing techniques may also explain some of the difference in measured ratios between labs. Data from our initial query (i.e., before setting cutoff values to constrain “reasonable” ratios) show that UW has less variance in measured $^{26}\text{Al}/^{10}\text{Be}$ ratios than any other chemical processing lab (relative standard deviation, RSD = 9.6% compared to RSD > 11% at other labs and 17.7% at LLNL), and all but two UW samples were within the cutoff ratio bounds. LLNL ratios from the initial query are skewed low, indicating perhaps an underestimation of native $^{27}\text{Al}$ in samples and thus the calculated $^{26}\text{Al}$ concentrations. We fit our bi-variate model to the measured ratios from each of the major chemical processing labs to assess this hypothesis and indeed found a more left-skewed residuals distribution from LLNL and a tighter fit (smaller residuals) from UW (Figure 8). Thus, we attribute the difference in measured $^{26}\text{Al}/^{10}\text{Be}$ ratios between LLNL and UW to both elevation differences between samples and more variable and low-skewed $^{26}\text{Al}/^{10}\text{Be}$ measurements from LLNL.
Figure 8. Residual histograms demonstrating the fit of our bi-variate model to samples from each of the major chemical processing labs.
6. Implications

Our analysis suggests that dual-nuclide studies that assume a spatially invariant 
$^{26}$Al/$^{10}$Be production ratio of 6.75 contain small but systemic biases in their data interpretations. The change in the $^{26}$Al/$^{10}$Be production ratio (5–6% between the equator and the poles and between sea level and mountain landscapes) is similar to the current analytical uncertainty of well-measured $^{26}$Al/$^{10}$Be ratios. Using the nominal ratio (6.75) at high latitudes and low elevations will underestimate burial times at higher latitudes and lower elevations.

Using a nuclide-specific production rate spatial scaling model (such as the LSDn scaling scheme from Lifton et al. [20]) will improve the accuracy of dual-nuclide studies. The LSDn scaling scheme fits the $^{26}$Al/$^{10}$Be ratio data in this compilation nearly as well as the bi-variate regression ($\chi^2 = 1.46$; Figures 9, S6, S8 and S9) and predicts ratio variations with elevation that are consistent with the empirical data (Figure 4A). The LSDn scaling scheme also predicts an increase in the $^{26}$Al/$^{10}$Be production ratio with latitude, which agrees with our single variate latitude/ratio regression but is not observed in our bi-variate regression.

![Figure 9. Residuals histogram showing differences between production ratios predicted by the LSDn scaling scheme for samples in the ICE:D compilation and measured $^{26}$Al/$^{10}$Be concentration ratios. The similarity in residuals distribution seen here compared to Figure 2B, the residuals from our bi-variate model based solely on measured ratios, demonstrates the good fit of LSDn scaling to empirical data.](image)

Our analysis is limited by spatial gaps in the data, particularly at low latitude/low altitude and high latitude/high altitude locations and thus may be biased by elevation-dependence of sites from different latitudes. Filling these gaps is essential to improving our understanding of $^{26}$Al/$^{10}$Be production ratio variations, but will be challenging. Glaciers did not occupy low latitude/low altitude sites, but other episodically exposed surfaces, such as those from rock falls, could be useful. Sampling high latitude/high altitude sites is logistically difficult, and many of these sites which have been sampled show evidence for significant concentrations of nuclides inherited from prior periods of exposure.

Analysis of these compiled data indicate the need for improving the precision and accuracy of $^{26}$Al/$^{10}$Be measurements and thus their application to geochronology and understanding landscape dynamics. Of particular concern are measurements of stable $^{27}$Al. Use of internal laboratory standards for quality control can help assure the quality of both $^{26}$Al and $^{10}$Be data. These are available as liquid standards [53], homogenized glass...
sand powder [25], and as purified quartz [26,28]. Improved precision of $^{26}\text{Al}$ concentration measurements will also be critical to constraining $^{26}\text{Al}/^{10}\text{Be}$ ratios [17].

**Supplementary Materials:** The following are available online at https://www.mdpi.com/article/10.3390/geosciences11100402/s1, Extended methods and results, Figures S1–S10 and Table S1. Figure S1: Distribution of $^{26}\text{Al}/^{10}\text{Be}$ concentration ratios from the initial ICE:D query. Vertical dashed line marks a ratio of 6.75. Figure S2: Distribution of concentration ratio uncertainties (%) from the initial ICE:D query. Figure S3: Standard boxplot of concentration ratio uncertainties from the initial ICE:D query. Box limits are the 25th and 75th percentile values, center red line is the median, whiskers are the high and low values not considered outliers, red crosses show outliers. Figure S4: Linear regressions (central lines) and 95% confidence intervals (upper and lower lines) for elevation and latitude transects. Note: 95% confidence interval lines for latitude band 2 are outside the y-axis bounds. Figure S5: Model check for normality in bi-variate linear regression correlating the measured $^{26}\text{Al}/^{10}\text{Be}$ concentration ratios to elevation and latitude. X-axis shows fitted values (ratios) from this model, while y-axis shows the measured ratios. Blue line is the 1:1 reference line. Figure S6: Model check for homoscedasticity in the bi-variate regression model. X-axis is fitted ratio values from the model, y-axis is residuals of the model compared to the data. No clear pattern in residuals is observed, indicating homoscedasticity. Figure S7: Model check for normality in LSDn scaling model against the measured $^{26}\text{Al}/^{10}\text{Be}$ concentration ratios. Axes are same as in Figure S4, but x-axis now shows fitted ratio values from LSDn scaling. Figure S8: Model check for homoscedasticity in the LSDn scaling model compared to concentration ratio data. Axes are same as in Figure S5, but x-axis shows fitted values from the LSDn scaling model. Figure S9: Comparison of the bi-variate linear model from this study (bottom, gray) and the LSDn scaling model (top, multi-colored) against the data in this compilation (blue dots). The decrease in concentration ratio with increasing elevation is nearly identical between the two models, but the LSDn model shows a more pronounced increase in ratio with latitude. Figure S10: Exploration of the influence of sample collection year on measured ratios and ratio uncertainties. Table S1: the statistics table for the bivariate regression run without UW samples.

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