



A regolith lead isoscape of Australia

Candan U. Desem¹, Patrice de Caritat², Jon Woodhead¹, Roland Maas¹, Graham Carr³

¹School of Geography, Earth and Atmospheric Sciences, The University of Melbourne, Melbourne, VIC 3010, Australia

²Geoscience Australia, GPO Box 378, Canberra, ACT 2601, Australia

5 ³Commonwealth Scientific and Industrial Research Organisation, North Ryde, NSW, Australia

Correspondence to: Candan Desem (candandesem@gmail.com)

Abstract. We present the first national-scale lead (Pb) isotope maps of Australia based on surface regolith for five isotope ratios, ²⁰⁶Pb/²⁰⁴Pb, ²⁰⁷Pb/²⁰⁴Pb, ²⁰⁸Pb/²⁰⁴Pb, ²⁰⁷Pb/²⁰⁶Pb, and ²⁰⁸Pb/²⁰⁶Pb, determined by single collector Sector Field-Inductively

10 Coupled Plasma-Mass Spectrometry after an Ammonium Acetate leach followed by Aqua Regia digestion. The dataset is underpinned principally by the National Geochemical Survey of Australia (NGSA) archived floodplain sediment samples. We analysed 1219 'top coarse' (0-10 cm depth, <2 mm grain size) samples, collected near the outlet of 1119 large catchments covering 5.647 million km² (~75% of Australia). The samples consist of mixtures of the dominant soils and rocks weathering in their respective catchments (and possibly those upstream) and are therefore assumed to form a reasonable representation of the average isotopic signature of those catchments. This assumption was tested in one of the NGSA catchments, within which 15 12 similar 'top coarse' samples were also taken; results show that the Pb isotope ratios of the NGSA catchment outlet sediment sample are close to the average of the 12 sub-catchment, upstream samples. National minimum, median and maximum values reported for ²⁰⁶Pb/²⁰⁴Pb were 15.558, 18.844, 30.635; for ²⁰⁷Pb/²⁰⁴Pb 14.358, 15.687, 18.012; for ²⁰⁸Pb/²⁰⁴Pb 33.558, 38.989, 48.873; for ²⁰⁷Pb/²⁰⁶Pb 0.5880, 0.8318, 0.9847; and for ²⁰⁸Pb/²⁰⁶Pb 1.4149, 2.0665, 2.3002, respectively. The new dataset was 20 compared with published bedrock and ore Pb isotope data, and was found to dependably represent crustal elements of various ages from Archean to Phanerozoic. This suggests that floodplain sediment samples are a suitable proxy for basement and basin geology at this scale, despite various degrees of transport, mixing, and weathering experienced in the regolith environment, locally over protracted periods of time. An example of atmospheric Pb contamination around Port Pirie, South Australia, where a Pb smelter has operated since the 1890s, is shown to illustrate potential environmental applications of this new dataset. Other 25 applications may include elucidating detail of Australian crustal evolution and mineralisation-related investigations. The new regolith Pb isotope dataset for Australia is publicly available (Desem et al., 2023; <http://dx.doi.org/10.26186/5ea8f6fd3de64>).

Commented [BG1]: TOS line 69

Commented [BG2]: See my concern about reporting these ratios to 3 decimal places given the precision reported in Methods and Desem et al. 2023

1 Introduction

Isoscapes – isotopic maps of landscapes – are increasingly used as tools to address a wide range of research questions in fields as diverse as hydrochemistry (e.g. Bowen et al., 2009), forensic studies (e.g. Chesson et al., 2014), and tracking animal 30 migrations (e.g. Hobson et al., 2010). Isotopic maps of the element strontium (Sr) in particular, often constructed at large scale,



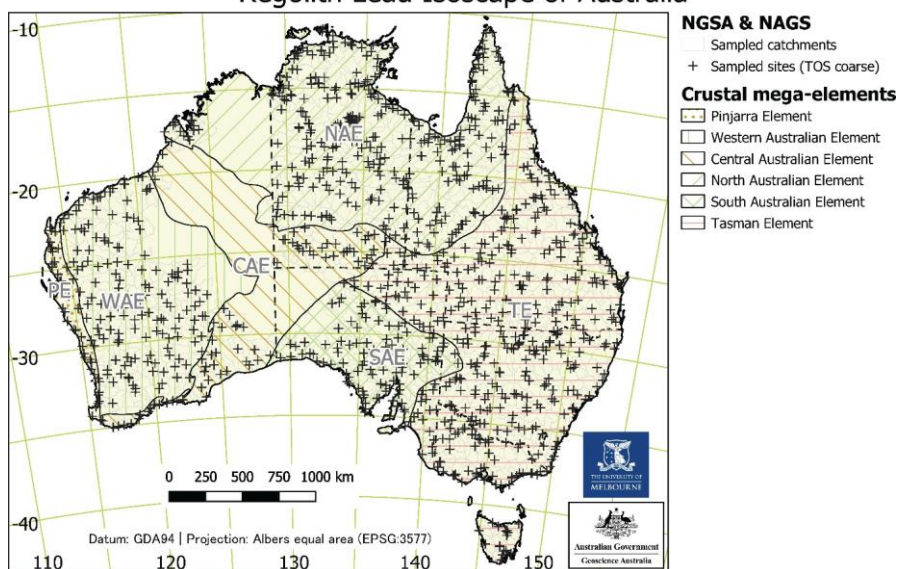
are finding increasing utility in provenance studies (e.g. Adams et al., 2019; Willmes et al., 2018; Bataille et al., 2020; Scaffidi and Knudson, 2020; de Caritat et al., 2022, 2023). The utility of Sr in this regard stems from its high bioavailability, coupled with its relative ease of isotopic determination in soil, water, and animal and plant tissue.

One limitation of Sr as a tracer is that it has only one radiogenic isotope ratio ($^{87}\text{Sr}/^{86}\text{Sr}$), hence a single isotopic determination on a target material may match multiple natural occurrences of that value across an isoscape. In contrast, the element lead (Pb) offers greater resolving potential; its isotopic composition is the result of three independent radioactive decay chains, producing effectively ‘three tracers in one’ and, as a result, much greater potential for accurate source attribution. Pb isotopes have in fact been used for many decades in provenance determination where metallic archaeological objects such as coins, shipwreck anchors, lead ingots, etc., have been traced to the likely sources from which their ores were mined (e.g. Gale and Stos-Gale, 2000). A similar methodology has been applied to track the origin of basaltic stone tools (e.g. Weisler and Woodhead, 1995). Pb is also a relatively bioavailable element, with well-known adverse effects on human health due to its cumulative toxicity and widespread use, and Pb isotopes have been used extensively to track the sources of Pb in humans (e.g. Gulson, 2008). These existing Pb-isotope studies, however, typically rely on matching samples to known point sources and, as such, do not employ the full predictive power of isoscapes, for example allowing the estimation of likely regions that do not have high sampling density. While Sr isoscapes are now in widespread use, the major impediment to the construction of continental-scale Pb-isotope maps is primarily an analytical one: Pb-isotope analysis traditionally requires exacting clean-room chemistry and specialised mass spectrometry procedures, and is correspondingly more expensive and time consuming than Sr-isotope analysis. As a result, very few large-scale, empirical Pb-isotope isoscapes have been constructed – an agricultural soil map of Europe (Reimann et al., 2012) and a tooth enamel study in Britain (Evans et al., 2022) being major exceptions, with ongoing but smaller-scale studies in various other countries aimed at tracking anthropogenic contaminants (e.g. Bing-Quan et al., 2002; Zuluaga et al., 2017), ~~and~~ provenancing cultural materials (e.g. Hsu and Sabatini, 2019), and isotopic changes in blood and teeth associated with migration between countries (Gulson et al., 1997; 2003, 2008).

In this study we release the first regolith Pb isoscapes constructed on a large scale for the Australian continent. Based upon surface regolith samples collected during the National Geochemical Survey of Australia (NGSA; www.ga.gov.au/ngsa; de Caritat and Cooper, 2011, 2016; de Caritat, 2022), these isoscapes are underpinned by a relatively dense and homogeneous distribution of sampling sites across the continent (Figure 1). This work was made possible by technological developments allowing both rapid and precise Pb-isotope analysis of large sample suites, using an analytical method recently described in Desem et al. (2022) and summarised below. The major advantage of this protocol over traditional methodologies for Pb-isotope analysis is it does not require matrix separation, thus greatly streamlining the analysis of large sample suites.



Regolith Lead Isoscape of Australia



60

Figure 1. Map showing the location of the National Geochemical Survey of Australia (NGSA) and Northern Australia Geochemical Survey (NAGS) sampling sites (black crosses) and NGSA catchments (grey polygons) (de Caritat and Cooper, 2011; Main et al., 2019) overlain by crustal mega-elements (hatched polygons) (Shaw et al., 1998). The mega-elements are labelled as (west to east) PE: Pinjarra Element; WAE: Western Australian Element; CAE: Central Australian Element; NAE: North Australian Element; and TE: Tasman Element.

65

2 Materials and methods

2.1 Materials

Our study principally utilises 'catchment outlet sediment' samples originally collected from two depths (top outlet sediment, or TOS, from 0 to 10 cm depth, and bottom outlet sediment, or BOS, from, on average, 60 to 80 cm depth) during the NGSA project (de Caritat and Cooper, 2011, 2016; de Caritat, 2022), which covered ~80% of the Australian continent (for details on sample collection, see Lech et al., 2007). At Geoscience Australia, freshly collected samples were air dried at 40 °C for a minimum of 48 hrs (or to constant mass), homogenised, and reduced by riffle splitting, with half of each sample set aside in an archive for future investigations, and the other half prepared for various analyses (for details on sample preparation, see de

70



75 Caritat et al., 2009). A ‘bulk’ split was retained for mineralogical analyses, another was sieved to a ‘coarse’ (<2 mm) grain-
size fraction, and yet another was sieved to a ‘fine’ (<75 µm) grain-size fraction. The latter two fractions of both depths were
then further prepared for the comprehensive geochemical analysis program of the NGSA (for details on sample analysis, see
de Caritat et al., 2010). Here, 1204 NGSA TOS coarse samples from 1098 catchments were used, with three of them analysed
twice for a total of 1207 analyses. Twelve additional TOS coarse samples collected in a similar manner to the NGSA samples
(from 0 to 10 cm depth and sieved to <2 mm) during the Northern Australia Geochemical Survey, or NAGS (Main et al.,
80 2019), were included in this study, giving a total of 1219 TOS coarse analyses underpinning the present isoscapes. In addition
to the above, 16 NGSA TOS fine (<75 µm) and 16 NGSA BOS coarse (<2 mm) were also analysed; although these data are
released herewith for the sake of completeness, they are not discussed further.

2.2 Methods

For the purpose of the Pb-isotope analyses, conducted at the University of Melbourne, the TOS coarse fractions from the
85 NGSA project were utilised. All samples to be analysed were subjected to a two-step sequential leaching protocol designed to
minimise and isolate any anthropogenic overprints on the primary Pb-isotope data. In the first step, an ammonium acetate
(AmAc) leach, developed at CSIRO and described in Carr et al. (2011), was applied to extract and remove any labile or loosely
bound/adsorbed components. Splits of ~1.2 g of sample were mixed with 6 mL of a 1:1 mixture of ultrapure water and
ammonium acetate buffer solution (AmAc; pH ~5). The soil/leach solution mix was shaken and left to react at 20 °C for 15
90 hours. Following centrifuging (4.5 min at 3000 rpm), a clear supernatant solution – the ‘AmAc leach’ or ‘A’ sample – was
pipetted off and dried in a high-efficiency particulate air (HEPA)-filtered fume-hood. In the second step, the remaining,
undigested sample was subjected to an aqua regia (AR, 3:1 HCl:HNO₃) acid attack to digest most (though not all) of the more
refractory components of the samples. Following an ultrapure-water rinse of the residual soil, 3 mL AR solution was added,
and the material was shaken and again left to react at 20 °C for 15 hours. After centrifuging, the clear supernatant solution –
95 the ‘AR digest’ or ‘B’ sample – was removed and dried in the HEPA-filtered fume-hood. The lead isotope analyses discussed
in this study were performed on the AR digest, although additional AmAc results are also provided for a subset of samples,
but not discussed further.

Lead isotope analyses followed procedures described in Desem et al. (2022) and are briefly outlined below. Importantly, the
methodology allows for analysis of samples without prior matrix removal, greatly improving sample throughput. Analyses
100 were performed using a Nu Instruments Attom SF-ICP-MS. Dried soil digests were redissolved in 2% HNO₃ run solutions
containing admixed high-purity thallium (1 ppb Tl), and diluted to provide ~1 ppb Pb in solution. Following the method of
Woodhead (2002), addition of natural, Pb-free Tl (with a nominal ²⁰⁵Tl/²⁰³Tl of 2.3871) allowed correction of instrumental
mass bias effects during Pb-isotope analyses. Analyses of the National Institute of Standards & Technology (NIST) Common
Lead Isotopic Standard Reference Material SRM 981 (certificate of analysis available at
105 <https://tsapps.nist.gov/srmext/certificates/archives/981.%20April%2010,%201973.pdf>; last access 5 September 2023)
interspersed throughout the unknown analyses were used to update the long term Pb vs Tl master correlations. Pb blanks for



the combined leaching and chemical procedures were typically <100 pg and are considered negligible relative to the amount of Pb being processed (typically hundreds of ng); as a result no blank corrections have been made. A small number of samples with low Pb concentrations exhibited very low signal sizes during analysis, resulting in correspondingly high analytical uncertainties. Samples producing within-run uncertainties of <1% relative (measured on the $^{207}\text{Pb}/^{204}\text{Pb}$ ratio) were discarded as being insufficiently precise to contribute meaningfully to the dataset.

2.3 Quality assessment

Although previous studies using the Attom SF-ICP-MS technique (e.g. Newman and Georg, 2012) used sample-standard-bracketing techniques to correct for instrumental mass bias during Pb-isotope analysis, in this study Tl doping was found to produce more precise, accurate and reproducible results. As the NIST SRM 981 Standard Reference Material was used to establish the Pb-Tl calibration (see above) SRM 981 values could not be used to assess analytical accuracy. Averages obtained for a variety of other, secondary reference materials measured during the course of this study, however, are consistent with accepted values (see Table 1), providing confidence in the analysis of unknowns.

Table 1. Pb isotope data obtained for geological reference materials run concurrently with the analyses reported in this paper and employed as secondary standards. Nominal values are derived from GeoREM (Jochum et al., 2007).

| Standard | Value | $^{206}\text{Pb}/^{204}\text{Pb}$ | $^{207}\text{Pb}/^{204}\text{Pb}$ | $^{208}\text{Pb}/^{204}\text{Pb}$ | $^{207}\text{Pb}/^{206}\text{Pb}$ | $^{208}\text{Pb}/^{206}\text{Pb}$ |
|-------------------|------------|-----------------------------------|-----------------------------------|-----------------------------------|-----------------------------------|-----------------------------------|
| BCR-2 (n = 39) | Nominal | 18.754 | 15.622 | 38.726 | 0.8329 | 2.0649 |
| | Average | 18.739 | 15.608 | 38.712 | 0.8328 | 2.0665 |
| | %2SD | 0.43 | 0.47 | 0.42 | 0.23 | 0.32 |
| | %deviation | -0.08 | -0.09 | -0.04 | -0.01 | 0.08 |
| BR (n = 11) | Nominal | 19.215 | 15.606 | 39.135 | 0.8122 | 2.0367 |
| | Average | 19.229 | 15.691 | 39.157 | 0.8164 | 2.0372 |
| | %2SD | 0.50 | 0.62 | 0.88 | 0.26 | 0.26 |
| | %deviation | 0.07 | 0.54 | 0.06 | 0.52 | 0.02 |
| AGV-2 (n = 13) | Nominal | 18.870 | 15.616 | 38.554 | 0.8275 | 2.0431 |
| | Average | 18.899 | 15.643 | 38.597 | 0.8279 | 2.0420 |
| | %2SD | 0.27 | 0.35 | 0.40 | 0.27 | 0.17 |
| | %deviation | 0.15 | 0.17 | 0.11 | 0.05 | -0.05 |
| JB-2 (n = 9) | Nominal | 18.342 | 15.561 | 38.274 | 0.8484 | 2.0867 |
| | Average | 18.416 | 15.654 | 38.456 | 0.8493 | 2.0871 |
| | %2SD | 0.68 | 0.60 | 0.71 | 0.29 | 0.34 |
| | %deviation | 0.40 | 0.60 | 0.48 | 0.11 | 0.02 |

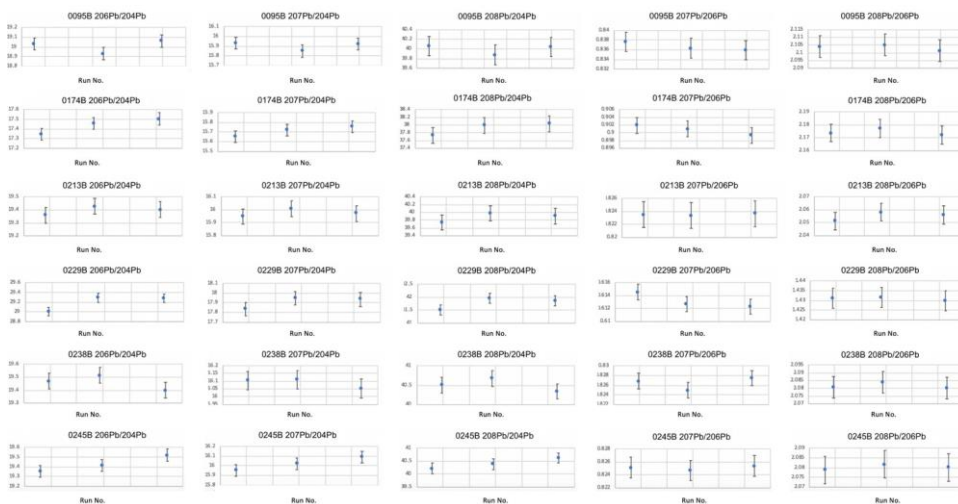
Commented [BG3]: Although useful to show generally good agreement in Nominal (MC-ICP-MS) vs measured (Average SF-ICP-MS) values, given the lower quality of the SF-ICP-MS data (as shown in Figure 4.3 in the Desem thesis and Desem et al. 2023), the Average results for 204 -based ratios should only be reported to 2 decimal places.

We see very good agreement between our SF-ICP-MS data and nominal accepted values for the well-known and described reference materials BCR-2 and AGV-2. Agreement was slightly poorer for the two other reference materials utilised — BR



125 and JB-2 — especially for ^{207}Pb -based ratios but these have been relatively little studied and at least some of this variation may therefore be the result of uncertainty in assigned literature values. Based upon the data for the BCR-2 and AGV-2 reference materials, for which we have the most analyses and for which accepted values are more robust, our accuracy is estimated to be typically $<0.17\%$.

130 A number of samples were also run on different occasions (up to 6 times) in order to provide some measure of long-term reproducibility. A selection of these is shown in Figure 2 and generally they reproduce within (or slightly beyond) the bounds of within-run uncertainties, providing further confidence in our analytical methodology. More detailed assessments of data quality, including comparisons with other instrumental techniques for Pb-isotope analysis, can be found in Desem et al. (2022) and are not re- iterated here.



135 **Figure 2. Laboratory duplicates: examples of single samples run on different occasions.**

140 One hundred and six field duplicate sample pairs (NGSA field duplicates, were collected at a median distance of ~ 100 m from one another on the same landscape unit; (see Cooper et al., 2010), were analysed using the TOS <2 mm sample, and returned a median relative standard deviations for ratios $^{206}\text{Pb}/^{204}\text{Pb}$, $^{207}\text{Pb}/^{204}\text{Pb}$, $^{208}\text{Pb}/^{204}\text{Pb}$, $^{207}\text{Pb}/^{206}\text{Pb}$, and $^{208}\text{Pb}/^{206}\text{Pb}$ of 0.50%, 0.24%, 0.43%, 0.30%, and 0.34%, respectively. The relative standard deviation from field duplicates includes natural variability (mineralogical/chemical heterogeneity of the alluvial deposit), as well as sample collection, preparation, and analysis uncertainties. It is thus expected to be larger than laboratory reproducibility which, as noted above, is usually $<0.17\%$.



Overall, we **consider** that the quality of the data presented herein is adequate for the purpose of regional mapping.

145 2.4 Data presentation

Data management and analysis, including visualisation, were performed using Microsoft Excel (v.2306), IMDEX ioGas (v.8.0), and open software QGIS (v.3.16).

3 Results and discussion

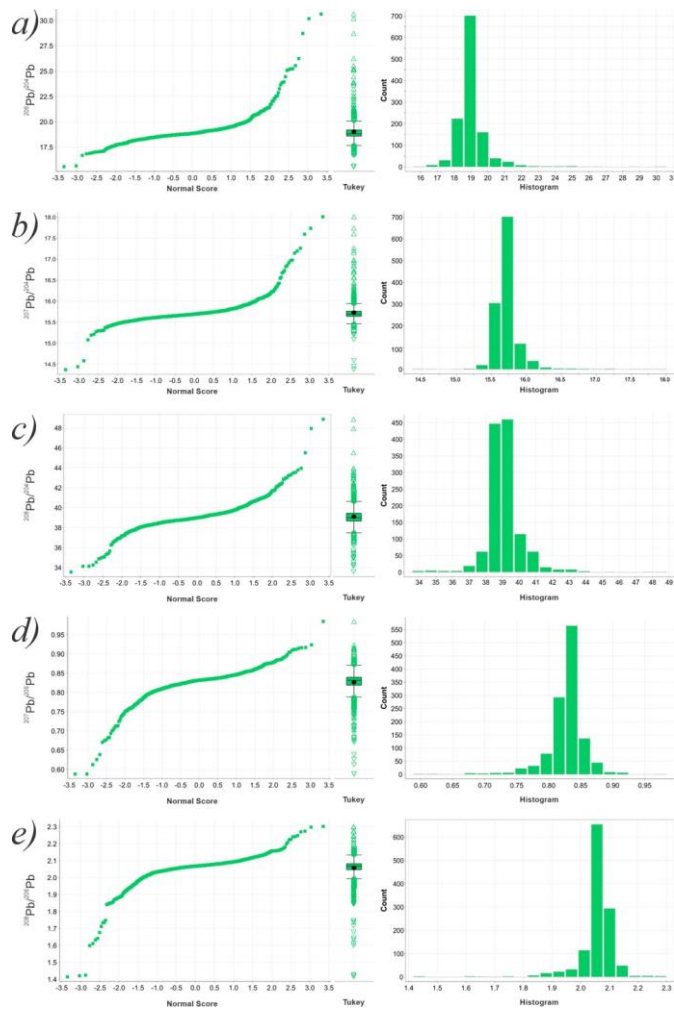
3.1 Statistics

150 The overall results obtained in this study are summarised in Table 2. Normal score, Tukey boxplot and histogram distributions of the $^{206}\text{Pb}/^{204}\text{Pb}$, $^{207}\text{Pb}/^{204}\text{Pb}$, $^{208}\text{Pb}/^{204}\text{Pb}$, $^{207}\text{Pb}/^{206}\text{Pb}$, and $^{208}\text{Pb}/^{206}\text{Pb}$ ratios are shown in Figure 3. It can be seen that these distributions are fairly 'normal' (Gaussian, or 'balanced' about the median), 'tight' (small interquartile ranges, or 'boxes', and widespread lower and upper quartiles), and with variable kurtosis and skewness.

155 Table 2. Summary statistics of the Pb isotope data obtained from 1207 National Geochemical Survey of Australia (NGSA) and 12 Northern Australia Geochemical Survey (NAGS) TOS coarse analyses by Aqua Regia digestion following Ammonium Acetate leach (n = 1219). See text for further details.

| Value | $^{206}\text{Pb}/^{204}\text{Pb}$ | $^{207}\text{Pb}/^{204}\text{Pb}$ | $^{208}\text{Pb}/^{204}\text{Pb}$ | $^{207}\text{Pb}/^{206}\text{Pb}$ | $^{208}\text{Pb}/^{206}\text{Pb}$ |
|-------------------------------------|-----------------------------------|-----------------------------------|-----------------------------------|-----------------------------------|-----------------------------------|
| Minimum | 15.558 | 14.358 | 33.558 | 0.5880 | 1.4149 |
| Maximum | 30.635 | 18.012 | 48.873 | 0.9847 | 2.3002 |
| Range | 15.077 | 3.654 | 15.315 | 0.3968 | 0.8854 |
| Mean | 19.047 | 15.720 | 39.116 | 0.8270 | 2.0568 |
| Standard Deviation | 1.07 | 0.22 | 1.09 | 0.03 | 0.07 |
| Coefficient of Variation (%) | 5.63 | 1.41 | 2.80 | 3.80 | 3.28 |
| Median | 18.844 | 15.687 | 38.989 | 0.8318 | 2.0665 |
| Robust Standard Deviation | 0.45 | 0.09 | 0.59 | 0.02 | 0.03 |
| Robust Coefficient of Variation (%) | 2.41 | 0.58 | 1.50 | 1.86 | 1.27 |
| Kurtosis | 33.17 | 30.12 | 13.26 | 12.95 | 27.41 |
| Skewness | 4.30 | 3.45 | 1.26 | -2.22 | -3.74 |

Commented [BG4]: Comment as above



160 **Figure 3.** Normal score, Tukey boxplot and histogram distributions for (a) $^{206}\text{Pb}/^{204}\text{Pb}$, (b) $^{207}\text{Pb}/^{204}\text{Pb}$, (c) $^{208}\text{Pb}/^{204}\text{Pb}$, (d) $^{207}\text{Pb}/^{206}\text{Pb}$, and (e) $^{208}\text{Pb}/^{206}\text{Pb}$ isotope ratios obtained for Australian TOS coarse samples by Aqua Regia digestion following Ammonium Acetate leach ($n = 1219$). Mean, outlier and far outlier values are shown on the Tukey boxplots as dots, circles, and triangles, respectively. See text for further details.



3.2 Validation

165 3.2.1 Intra-catchment variation

One NGSA catchment was also sampled at higher resolution by several NAGS samples, providing an opportunity to test the fundamental assumption underpinning the catchment-based sampling strategy of the NGSA, namely that one catchment outlet sediment sample fairly represents an average value for the whole catchment. In this case NGSA sample 2007190096 is from the Newcastle Creek catchment (TS0715), approximately 100 km northeast of Elliott in the NT, which was also sampled by 12 NAGS samples. The mean of these 12 NAGS samples is within 0.62 standard deviation of the singular catchment-outlet NGSA value for $^{206}\text{Pb}/^{204}\text{Pb}$, $^{207}\text{Pb}/^{204}\text{Pb}$, $^{208}\text{Pb}/^{204}\text{Pb}$, and $^{207}\text{Pb}/^{206}\text{Pb}$; for the isotopic ratio $^{208}\text{Pb}/^{206}\text{Pb}$ it is within 1.11 standard deviation (Table 3). All samples are TOS coarse fractions digested in AR after an AmAc leach as described above. The comparison supports the premise of the NGSA sampling strategy, namely that catchment outlet sediments are geochemically and mineralogically representative of their overall catchment, yet of course recognises that intra-catchment variation is occurring and can be significant.

180 **Table 3. Pb isotope data obtained from 12 Northern Australia Geochemical Survey (NAGS) samples in catchment TS0715, compared to the National Geochemical Survey of Australia (NGSA) catchment outlet sample for the same catchment. The Difference (Average NAGS – NGSA) is shown as absolute values and normalised to the standard deviation (SD) of the NAGS values. See text for further details.**

| Samples in catchment TS0715 | $^{206}\text{Pb}/^{204}\text{Pb}$ | $^{207}\text{Pb}/^{204}\text{Pb}$ | $^{208}\text{Pb}/^{204}\text{Pb}$ | $^{207}\text{Pb}/^{206}\text{Pb}$ | $^{208}\text{Pb}/^{206}\text{Pb}$ |
|-----------------------------|-----------------------------------|-----------------------------------|-----------------------------------|-----------------------------------|-----------------------------------|
| 20173120170 (NAGS) | 17.021 | 15.071 | 34.541 | 0.8853 | 2.0284 |
| 20173120227 (NAGS) | 19.108 | 15.740 | 39.043 | 0.8239 | 2.0432 |
| 20173120413 (NAGS) | 18.967 | 15.655 | 38.682 | 0.8249 | 2.0399 |
| 20173120558 (NAGS) | 18.994 | 15.650 | 38.768 | 0.8242 | 2.0396 |
| 20173120577 (NAGS) | 18.902 | 15.652 | 38.765 | 0.8284 | 2.0508 |
| 20173120699 (NAGS) | 19.049 | 15.633 | 38.923 | 0.8209 | 2.0438 |
| 20173120722 (NAGS) | 19.091 | 15.674 | 38.889 | 0.8206 | 2.0359 |
| 20173120774 (NAGS) | 19.095 | 15.650 | 38.777 | 0.8196 | 2.0309 |
| 20173120957 (NAGS) | 18.981 | 15.657 | 38.650 | 0.8246 | 2.0362 |
| 20173120982 (NAGS) | 18.928 | 15.725 | 39.070 | 0.8306 | 2.0637 |
| 20173121029 (NAGS) | 18.925 | 15.635 | 38.944 | 0.8261 | 2.0568 |
| 20173121195 (NAGS) | 19.006 | 15.640 | 38.981 | 0.8233 | 2.0517 |
| Average (NAGS) | 18.839 | 15.615 | 38.503 | 0.8294 | 2.0434 |
| SD (NAGS) | 0.577 | 0.175 | 1.255 | 0.0179 | 0.0106 |
| 2007190096 (NGSA) | 19.128 | 15.689 | 39.280 | 0.8210 | 2.0552 |
| Difference | -0.289 | -0.074 | -0.777 | 0.0084 | -0.0118 |
| Difference / SD (NAGS) | -0.50 | -0.43 | -0.62 | 0.47 | -1.11 |

Commented [BG5]: Comment as above



3.2.2 Regional scale isotopic variation

The regolith Pb isotope data, at continental scale, are clearly governed by major crustal boundaries. At the highest level this is reflected in the more radiogenic signatures (e.g., higher $^{206}\text{Pb}/^{204}\text{Pb}$ ratios) in older terranes (i.e., west and northern Australia), compared with less radiogenic signatures of younger terranes (Tasman and New England Fold belts), but is also visible in data averages calculated for each of the ‘crustal elements’ indicated in Figure 1 and see Table 4. At this scale the $^{206}\text{Pb}/^{204}\text{Pb}$ and $^{208}\text{Pb}/^{204}\text{Pb}$ isotope signatures exhibit the greatest variation since most ^{235}U decayed early in earth history and thus changes in $^{207}\text{Pb}/^{204}\text{Pb}$ are more subdued compared to variations in the $^{206}\text{Pb}/^{204}\text{Pb}$ and $^{208}\text{Pb}/^{204}\text{Pb}$ signatures.

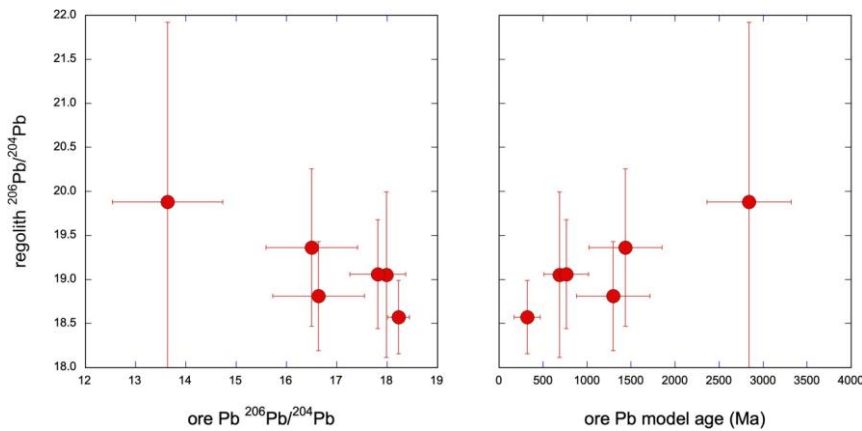
Regolith-derived data averages also broadly correlate with initial Pb signatures derived from a compilation of ore Pb data (Huston et al., 2019, 2021) and the terrane model ages derived from these data (Fig. 4).

Table 4. Regolith Pb isotope ratios from this study, averaged for each of the mega-crustal elements (Shaw et al., 1998), compared with ore Pb signatures and Stacey-Kramers model ages from Huston (2019, 2021).

| Mega-crustal element | $^{206}\text{Pb}/^{204}\text{Pb}$ | | $^{207}\text{Pb}/^{204}\text{Pb}$ | | $^{208}\text{Pb}/^{204}\text{Pb}$ | | Model age (Ma) |
|----------------------|-----------------------------------|----------------------|-----------------------------------|----------------------|-----------------------------------|----------------------|----------------|
| | This study | Huston et al. (2019) | This study | Huston et al. (2019) | This study | Huston et al. (2019) | |
| Pinjarra | 19.053 | 17.985 | 15.791 | 15.710 | 39.502 | 38.677 | 688 |
| West Australian | 19.879 | 13.637 | 15.936 | 14.698 | 39.722 | 33.481 | 2840 |
| South Australian | 18.811 | 16.637 | 15.684 | 15.479 | 38.939 | 36.350 | 1298 |
| Central Australian | 19.059 | 17.814 | 15.748 | 15.684 | 39.562 | 38.129 | 765 |
| North Australian | 19.363 | 16.501 | 15.740 | 15.500 | 39.477 | 36.247 | 1436 |
| Tasman | 18.578 | 18.230 | 15.630 | 15.611 | 38.572 | 38.233 | 321 |

Commented [BG6]: For non experts

Commented [BG7]: Comment as above re decimal places



195

Figure 4. Regolith vs ore Pb isotope averages for each mega-crustal element (left), and vs model ages (right) derived for these from ore Pb data (Huston et al., 2019, 2021).

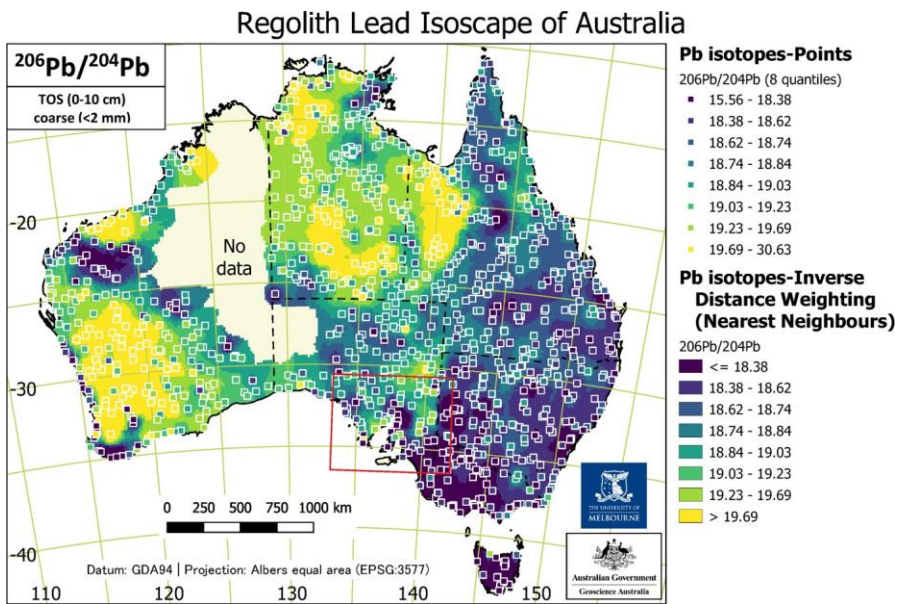
3.3 Isoscapes

The Pb isotope maps (isoscapes) for the ratios $^{206}\text{Pb}/^{204}\text{Pb}$, $^{207}\text{Pb}/^{204}\text{Pb}$, $^{208}\text{Pb}/^{204}\text{Pb}$, $^{207}\text{Pb}/^{206}\text{Pb}$, and $^{208}\text{Pb}/^{206}\text{Pb}$ are presented below (Figures 5 to 9). Each map includes a series of ‘Points’ coloured according to eight quantile classes for binning overlain on a raster surface coloured in the same way. The raster is an ‘Inverse distance weighting’ (IDW) interpolation produced with the **Grid (IDW with Nearest Neighbor Searching)** or **invdistnn** GDAL tool in QGIS. The ‘inverse distance to a power’ gridding method is a weighted average interpolator. Sample points are weighted during interpolation such that the influence of one point relative to another declines with distance from the unknown pixel to be estimated. Here, a weighting power of 2, minimum/maximum nearest neighbouring points of 6/12, and grid cells of $0.25^\circ \times 0.25^\circ$ resolution are the parameters used. The rectangular rasters thus produced were subsequently clipped to a custom polygon combining the Australian coastline with the area of ‘No data’ in the NGS coverage using the **Clip Raster by Mask Layer** or **gdalwarp -cutline** GDAL tool in QGIS.

200

205

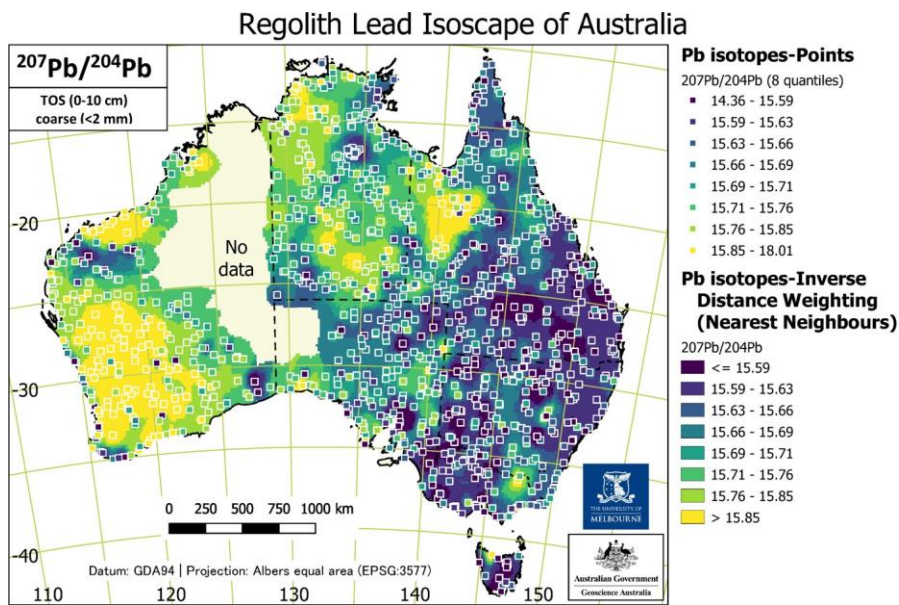
Commented [BG8]: Why bold?



210 Figure 5. Regolith Pb isoscape of Australia for $^{206}\text{Pb}/^{204}\text{Pb}$ with data points (AR digestion of TOS coarse samples) classed by quantiles and overlain on an IDW interpolation raster classed identically. The red rectangle indicates the location of Figure 10. See



text for further details.



215 **Figure 6.** Regolith Pb isoscape of Australia for $^{207}\text{Pb}/^{204}\text{Pb}$ with data points (AR digestion of TOS coarse samples) classed by quantiles and overlain on an IDW interpolation raster classed identically. See text for further details.

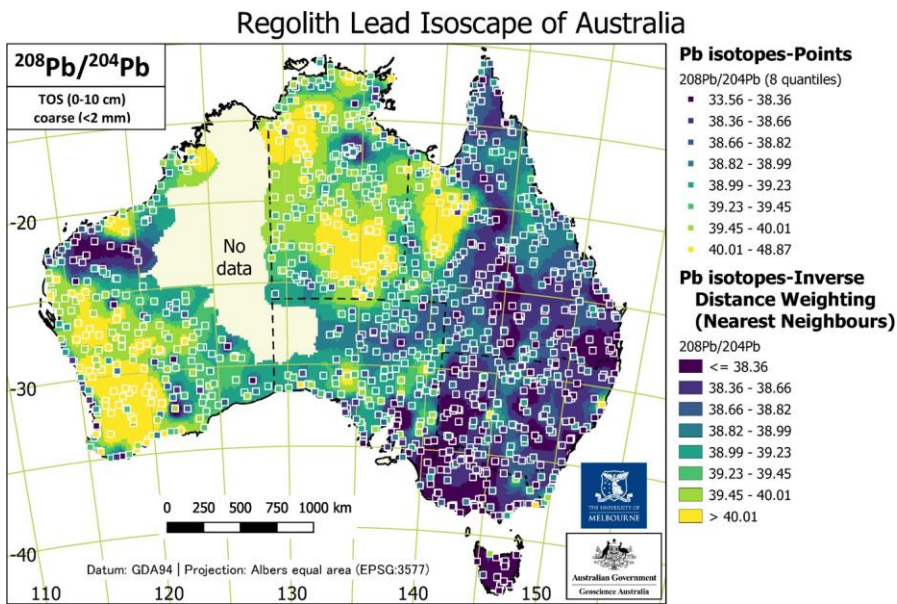
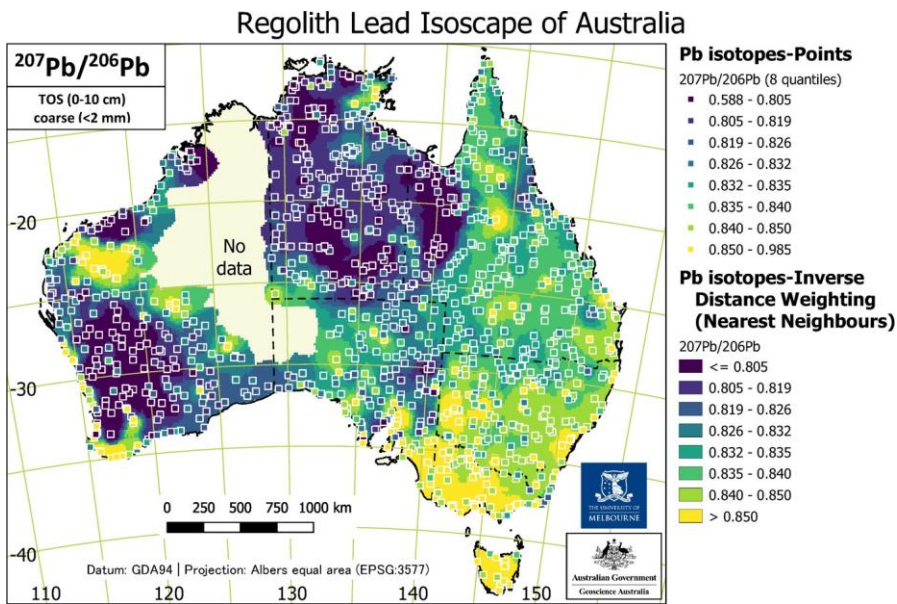


Figure 7. Regolith Pb isoscape of Australia for $^{208}\text{Pb}/^{204}\text{Pb}$ with data points (AR digestion of TOS coarse samples) classed by quantiles and overlain on an IDW interpolation raster classed identically. See text for further details.



220 Figure 8. Regolith Pb isoscape of Australia for $^{207}\text{Pb}/^{206}\text{Pb}$ with data points (AR digestion of TOS coarse samples) classed by quantiles and overlain on an IDW interpolation raster classed identically. See text for further details.

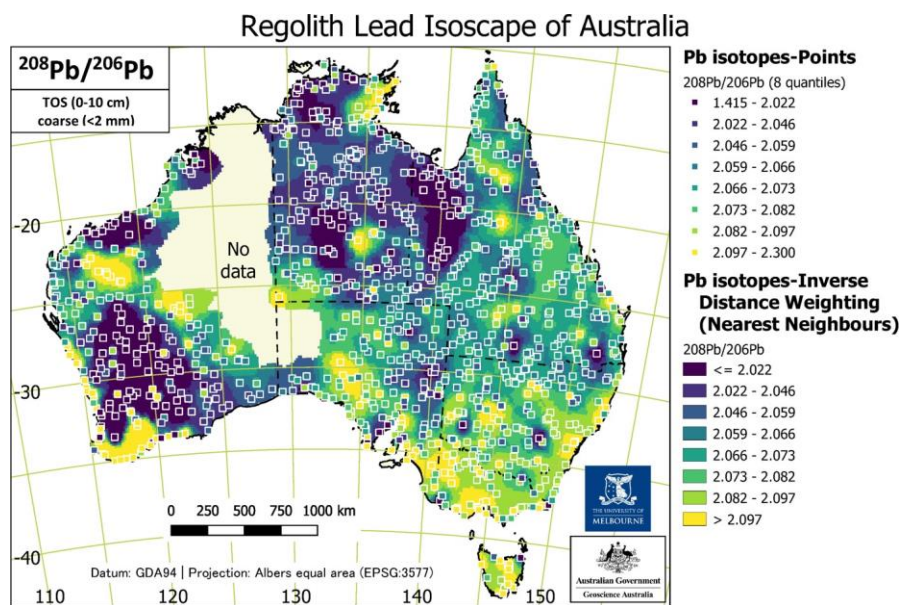


Figure 9. Regolith Pb isoscape of Australia for $^{208}\text{Pb}/^{206}\text{Pb}$ with data points (AR digestion of TOS coarse samples) classed by quantiles and overlain on an IDW interpolation raster classed identically. See text for further details.

225 All isoscape IDW-NN geotiff rasters are downloadable as per the Data subsection.

3.3 Applications

The present Pb isoscapes can be applied to studies of the evolution of the Australian crust, regional mineral exploration, and baselines for environmental investigations. The former two will be developed elsewhere, but the latter is illustrated below with an example from data obtained close to the Port Pirie smelter in South Australia. Port Pirie is the locality of the largest Pb smelter and refinery in the southern hemisphere; a Pb smelter has been active there over 130 years (SA EPA). The widespread contamination of the area surrounding the smelter by means of windblown Pb dust is well documented in the literature (e.g. Gulson et al., 1981). Our TOS regolith data (Figure 10) show pronounced unradiogenic signatures (e.g., lower $^{206}\text{Pb}/^{204}\text{Pb}$ ratios) adjacent to the area, which likely reflect a profound overprint from the ores processed in the facility, despite application of the AmAc pre-leach to these samples. During its >100 year history, the dominant source of the feedstock for the smelter was from the geologically ancient (~1600 million years; Gulson 1984) These ores originate from both the Broken Hill

<https://doi.org/10.5194/essd-2023-357>
Preprint. Discussion started: 19 October 2023
© Author(s) 2023. CC BY 4.0 License.



deposit in New South Wales ([Body et al., 1988](#)) and, [later](#), the Teutonic Bore mine in Western



235 Australia. [The low \$^{206}\text{Pb}/^{204}\text{Pb}\$ ratios in the smelter emissions are consistent with the low ratios of 16.00 in the Broken Hill ore](#)
(Gulson 1984). For two NGS sites — 2007190995 located south of the Port Pirie Pb smelter and 2007190228 from north of
Port Pirie location — we analysed both TOS and BOS sample aliquots, with the TOS coarse fraction producing less
radiogenic Pb isotope signatures compared to the BOS fraction samples. This further suggests that most of the anthropogenic
contamination resides in the surface layer (recent deposits). The bottom fraction samples for both locations are very similar
240 to one another and likely reflect the signature of floodplain deposits formed prior to the initiation of smelting activities
started and therefore provide a more reliable geogenic signature.

[The isotopic profiles in this study are consistent with those identified by Gulson et al. \(1981\).](#)

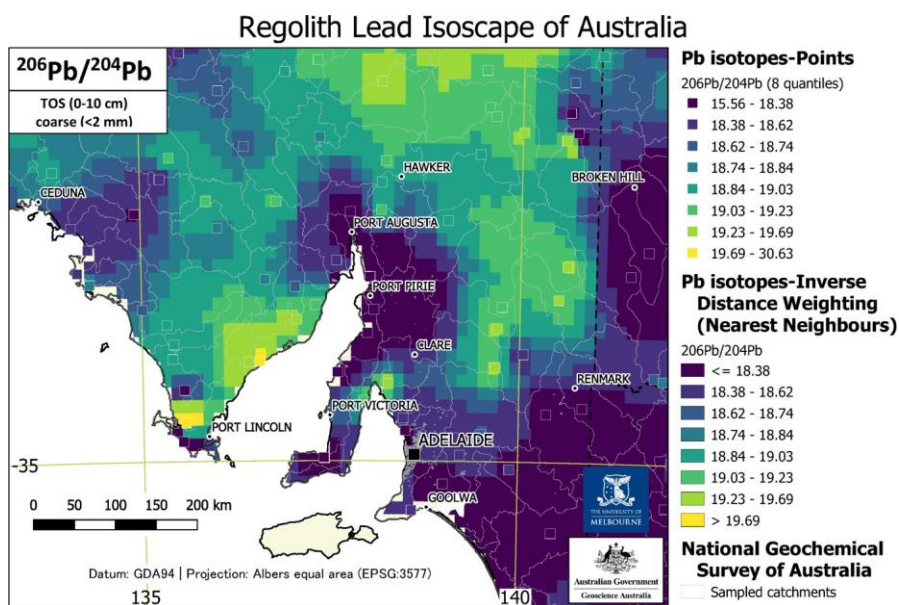


Figure 10. Regional detail of the regolith Pb isoscape of Australia in the vicinity of the Pt Pirie smelter, South Australia, for $^{206}\text{Pb}/^{204}\text{Pb}$ with data values (AR digestion of TOS coarse samples) classed by quantiles and overlain on an IDW interpolation raster classed identically. The NGS catchments are shown by the thin grey polygons. See text for further details.

245 4 Data availability

<https://doi.org/10.5194/essd-2023-357>
Preprint. Discussion started: 19 October 2023
© Author(s) 2023. CC BY 4.0 License.



The regolith Pb isotope dataset of Australia is publicly available (Desem et al., 2023; <http://dx.doi.org/10.26186/5ea8f6fd3de64>).



5 Conclusions

New national-scale regolith lead (Pb) isoscapes for Australia are presented for the ratios $^{206}\text{Pb}/^{204}\text{Pb}$, $^{207}\text{Pb}/^{204}\text{Pb}$, $^{208}\text{Pb}/^{204}\text{Pb}$,
250 $^{207}\text{Pb}/^{206}\text{Pb}$, and $^{208}\text{Pb}/^{206}\text{Pb}$. The results of this study suggest that the isotopic signatures obtained from transported regolith in
Australia are dominated by Pb from the catchment bedrock geology. This influence is more easily visible in older (i.e. Archaean
and Proterozoic) terranes, simply because of the greater opportunity for radiogenic ingrowth here. This is demonstrated by the
correlation of Pb signatures with major crustal elements and their respective geological model ages. Exogenous
(anthropogenic) inputs of Pb have been detected at some sample locations and provide further encouragement for the utilisation
255 of the new Pb isoscapes in source attribution for environmental studies. An example from around the Port Pirie Pb smelter is
developed for illustration of this application. Other potential applications of this dataset include studies of the crustal evolution
of Australia, and using Pb isotopes for mineral exploration.

Author contributions

Each author to indicate contribution as sentences, not dot points. Recommended categories are:

260 https://publications.copernicus.org/services/contributor_roles_taxonomy.html

CD provided Investigation (Pb isotope analysis), Validation, and Writing – original draft.

PdC provided Data curation, Formal analysis, Resources (samples), Validation, Visualisation, and Writing – original draft,
reviewing and editing.

JW provided Project administration, Methodology, Supervision, and Writing – original draft, reviewing and editing.

265 RM Supervision, Validation (standards data), Methodology

GC Supervision, Methodology

Competing interests

The contact author has declared that none of the authors has any competing interests.

Disclaimer

270 This paper is published with the permission of the Chief Executive Officer, Geoscience Australia.

Acknowledgements

The National Geochemical Survey of Australia (NGSA) project would not have been possible without Commonwealth funding
through the “Onshore Energy Security Program” (<http://www.ga.gov.au/ngsa>, last access: 5 September 2023), and Geoscience



275 Australia appropriation. The Northern Australia Geochemical Survey (NAGS) project was funded under the “Exploring for
the Future” programme (<https://eftf-production.ga.gov.au/northern-australia-geochemical-survey>, last access: 5 September
2023) and Geoscience Australia appropriation. Collaboration with the geoscience agencies of all states and the Northern
Territory is gratefully recognized. We acknowledge all land owners and custodians, whether private, corporate, and/or
traditional, for granting access to the field sites for the purposes of sampling. We are also grateful to Geoscience Australia
laboratory staff for assistance with preparing and analysing the samples. We thank Geoscience Australia reviewers, XXX and
280 YYY; journal reviewers, XXX, YYY, and ZZZ; and the journal editor, XXX, for their detailed and constructive critique of
our work.

Financial support

Analytical work was conducted as part of Candan Desem’s PhD studies supported financially by University of Melbourne
Research Scholarship, Baragwanath Geology Research Scholarship and the Albert Shimmins Fund, in addition to a PhD
285 Student Research Grant from the International Association of GeoChemists (IAGC).

This research has also been supported by the Australian Government (Exploring for the Future,
<https://www.eftf.ga.gov.au/about>, last access: 5 September 2023). Geoscience Australia’s Exploring for the Future programme
provides pre-competitive information to inform decision-making by government, community, and industry on the sustainable
development of Australia’s mineral, energy, and groundwater resources. By gathering, analysing, and interpreting new and
290 existing pre-competitive geoscience data and knowledge, we are building a national picture of Australia’s geology and resource
potential. This leads to a strong economy, resilient society, and sustainable environment for the benefit of all Australians. This
includes supporting Australia’s transition to net-zero emissions; strong, sustainable resources and agriculture sectors; and
economic opportunities and social benefits for Australia’s regional and remote communities. The Exploring for the Future
programme, which commenced in 2016, is an 8-year, AUD 225 million investment by the Australian Government.

295 References

- Adams, S., Grün, R., McGahan, D., Zhao, J.-X., Feng, Y., Nguyen, A., Willmes, M., Quaresimin, M., Lobsey, B., Collard, M.,
and Westaway, M. C.: A strontium isoscape of north-east Australia for human provenance and repatriation, *Geoarchaeol.*, 34,
231–251, <https://doi.org/10.1002/gea.21728>, 2019.
- Bataille, C. P., Crowley, B. E., Wooller, M. J., and Bowen, G. J.: Advances in global bioavailable strontium isoscapes,
300 *Palaeogeogr. Palaeoclimatol.*, 555, 109849, <https://doi.org/10.1016/j.palaeo.2020.109849>, 2020.



- Bing-Quan, Z., Yu-Wei, C., and Xiang-Yang, C.: Application of Pb isotopic mapping to environment evaluation in China. *Chem. Speciation Bioavail.*, 14, 49–56, <https://doi.org/10.3184/095422902782775335>, 2002.
- Bowen, G. J., West, J. B., Vaughn, B. H., Dawson, T. E., Ehleringer, J. R., Fogel, M. L., Hobson, K., Hoogewerff, J., Kendall, C., Lai, C.-T., Miller, C. C., Noone, D., Schwartz, H., and Still, C. J.: Isoscapes to address large-scale earth science challenges, 305 *EOS Trans. Am. Geophys. Union*, 90, 109–116, <https://doi.org/10.1029/2009EO130001>, 2009.
- Carr, G. R., Korsch, M. J., Denton, G. J., Gatehouse, S., Law, A., Gray, D. R., Andrew, A. S.: AMIRA P618, Isotopic Discrimination of Partial Leach, Geochemical Anomalies in Covered Terrains, Final Report, CSIRO Division of Earth Science and Resource Engineering, CSIRO Report, EP0410117, 2011.
- Chesson, L. A., Tipple, B. J., Howa, J. D., Bowen, G. J., Barnette, J. E., Cerling, T. E., and Ehrlinger, J. R.: Stable isotopes in 310 forensics applications, in: *Treatise on Geochemistry, Second Edition*, edited by: Holland H. D. and Turekian K. K., 14, 285–317, <http://dx.doi.org/10.1016/B978-0-08-095975-7.01224-9>, 2014.
- Cooper, M., Caritat, P. de, Burton, G., Fidler, R., Green, G., House, E., Strickland, C., Tang, J., and Wygralak, A.: National Geochemical Survey of Australia: Field Data, Record, 2010/18, *Geosci. Austral.*, Canberra, <https://doi.org/10.11636/Record.2011.020>, 2010.
- 315 de Caritat, P. and Cooper, M.: A continental-scale geochemical atlas for resource exploration and environmental management: the National Geochemical Survey of Australia, *Geochem. Explo. Env. Anal.*, 16, 3–13, <https://doi.org/10.1144/geochem2014-322>, 2016.
- de Caritat, P. and Cooper, M.: National Geochemical Survey of Australia: The Geochemical Atlas of Australia, Record, 2011/20, *Geosci. Austral.*, Canberra, <http://pid.geoscience.gov.au/dataset/ga/71973>, 2011.
- 320 de Caritat, P., Cooper, M., Lech, M., McPherson, A., and Thun, C.: National Geochemical Survey of Australia: Sample Preparation Manual, Record, 2009/08, *Geosci. Austral.*, Canberra, <http://pid.geoscience.gov.au/dataset/ga/68657>, 2009.
- de Caritat, P., Cooper, M., Pappas, W., Thun, C., and Webber, E.: National Geochemical Survey of Australia: Analytical Methods Manual, Record, 2010/15, *Geosci. Austral.*, Canberra, <http://pid.geoscience.gov.au/dataset/ga/70369>, 2010.
- de Caritat, P., Dosseto, A., and Dux, F.: A strontium isotope of inland southeastern Australia, *Earth Syst. Sci. Data*, 14, 4271–325 4286, <https://doi.org/10.5194/essd-14-4271-2022>, 2022.
- de Caritat, P., Dosseto, A., and Dux, F.: A strontium isotope of northern Australia, *Earth Syst. Sci. Data*, 15, 1655–1673, <https://doi.org/10.5194/essd-15-1655-2023>, 2023.
- de Caritat, P.: The National Geochemical Survey of Australia: review and impact, *Geochem. Explo. Env. Anal.*, geochem2022-032, <https://doi.org/10.1144/geochem2022-032>, 2022.



- 330 Desem, C. U., de Caritat, P., Woodhead, J. D., Maas, R., and Carr, G.: National Geochemical Survey of Australia: Lead Isotopes Dataset, Geosci. Austral., Canberra [data set], <http://dx.doi.org/10.26186/5ea8f6fd3de64>, 2023.
- Desem, C. U., Maas, R., Woodhead, J., Carr, G., and Greig, A.: The utility of rapid throughput single-collector sector-field ICP-MS for soil Pb isotope studies, *Appl. Geochem.*, 143, <https://doi.org/10.1016/j.apgeochem.2022.105361>, 2022.
- Evans, J. A., Pashley, V., Mee, K., Wagner, D., Parker Pearson, M., Fremondeau, D., Albarella, U., and Madgewick, R.:
335 Applying lead (Pb) isotopes to explore mobility in humans and animals. *PLoS ONE*, 17(10), e0274831, <https://doi.org/10.1371/journal.pone.0274831>, 2022.
- Gale, N. H. and Stos-Gale, Z. A.: Lead isotope analyses applied to provenance studies, in: *Modern Analytical Methods in Art and Archaeology*, edited by: Ciliberto, E. and Spoto, G., Wiley, New York, ISBN: 978-0-471-29361-3, 503–584, 2000.
- Gulson, B. L., Tiller, K. G., Mizon, K. J., and Merry, R. H.: Use of lead isotopes to identify the source of lead contamination near Adelaide, South Australia, *Env. Sci. Technol.*, 15, 691–696, <https://doi.org/10.1021/es00088a008>, 1981.
- Gulson, B.: Stable lead isotopes in environmental health with emphasis on human investigations. *Sci. Total Environ.*, 400, 75–92, <https://doi.org/10.1016/j.scitotenv.2008.06.059>, 2008.
- Hobson, K. A., Barnett-Johnson, R., and Cerling, T.: Using isoscapes to track animal migration, in: *Isoscapes*, edited by: West, J. B., Bowen, G. J., Dawson, T. E., and Tu, K. P., Springer, Dordrecht, The Netherlands, 273–298, https://doi.org/10.1007/978-90-481-3354-3_13, 2010.
345
- Hsu, Y.-K. and Sabatini, B. J.: A geochemical characterization of lead ores in China: an isotope database for provenancing archaeological materials. *Plos One*, 14, e0215973, <https://doi.org/10.1371/journal.pone.0215973>, 2019.
- Huston, D. L., Champion, D. C., Ware, B., Carr, G., Maas, R., and Tessalina, S.: Preliminary National-Scale Lead Isotope Maps of Australia, *Record*, 2019/001, Geosci. Austral., Canberra, <http://dx.doi.org/10.11636/Record.2019.001>, 2019.
- 350 Jochum, K.P., Nohl, U., Herwig, K., Lammel, E., Stoll, B. and Hofman, A.W.: GeoREM: a new geochemical database for Reference Materials and Isotopic standards. *Geostandards and Geoanalytical Research* 29 (3), 333–338, <https://doi.org/10.1111/j.1751-908X.2005.tb00904.x>, 2005.
- Lech, M. E., de Caritat, P., and McPherson, A. A.: National Geochemical Survey of Australia: Field Manual, *Record*, 2007/08, Geosci. Austral., Canberra, <http://pid.geoscience.gov.au/dataset/ga/65234>, 2007.
- 355 Main, P. T., Bastrakov, E. N., Wygralak, A. S., and Khan, M.: Northern Australia Geochemical Survey: Data Release 2 – Total (Coarse Fraction), Aqua Regia (Coarse and Fine Fraction), and Fire Assay (Coarse and Fine Fraction) Element Contents, *Record*, 2019/002, Geosci. Austral., Canberra, <http://dx.doi.org/10.11636/Record.2019.002>, 2019.
- Newman, K. and Georg, R.B.: The measurement of Pb isotope ratios in sub-ng quantities by fast scanning single collector sector field-ICP-MS. *Chem. Geol.*, 304–305, 151–157, <https://doi.org/10.1016/j.chemgeo.2012.02.010>, 2012.



- 360 Reimann, C., Flem, B., Fabian, K., Birke, M., Ladenberger, A., Negrel, P., Demetriades, A., Hoogewerff, J., and GEMAS Project Team: Lead and lead isotopes in agricultural soils of Europe – the continental perspective. *Appl. Geochem.*, 27, 532–542, <https://doi.org/10.1016/j.apgeochem.2011.12.012>, 2012.
- Scaffidi, B. K. and Knudson, K.J.: An archaeological strontium isoscape for the prehistoric Andes: understanding population mobility through a geostatistical meta-analysis of archaeological $^{87}\text{Sr}/^{86}\text{Sr}$ values from humans, animals, and artifacts. *J. Archaeol. Sci.*, 117, 10521, <https://doi.org/10.1016/j.jas.2020.105121>, 2020.
- 365 Shaw, R. D., Wellman, P., Gunn, P. J., Whitaker, A. J., and Tarlowski, C.Z.: Australian Crustal Elements (National Geoscience Dataset), Geosci. Austral., Canberra [data set], <http://pid.geoscience.gov.au/dataset/ga/21195>, 1998.
- Weisler, M. and Woodhead, J.D.: Basalt Pb isotope analysis and the prehistoric settlement of Polynesia, *Proc. Nat. Acad. Sci.*, 92, 1881–1885, <https://doi.org/10.1073/pnas.92.6.1881>, 1995.
- 370 Willmes, M., Bataille, C. P., James, H. F., Moffat, I., McMorrow, L., Kinsley, L., Armstrong, R. A., Eggins, S., and Grün, R.: Mapping of bioavailable strontium isotope ratios in France for archaeological provenance studies, *Appl. Geochem.*, 90, 75–86, <https://doi.org/10.1016/j.apgeochem.2017.12.025>, 2018.
- Woodhead, J.D.: A simple method for obtaining highly accurate Pb isotope data by MC-ICP-MS, *Journal of Analytical Atomic Spectrometry.*, 17, 1381–1385, <https://doi.org/10.1039/B205045E>, 2002.
- 375 Zuluaga, M. C., Norini, G., Ayuso, R., Nieto, J. M., Lima, A., Albanese, S., and De Vivo, B.: Geochemical mapping, environmental assessment and Pb isotopic signatures of geogenic and anthropogenic sources in three localities in SW Spain with different land use and geology, *J. Geoch. Explo.*, 181, 172–190, <https://doi.org/10.1016/j.gexplo.2017.07.011>, 2017.

Body, P.E., Inglis, G.R. and Mulcahy, D.E. 1988. *Lead Contamination in Port Pirie South Australia*. SA Department of Environment and Planning, Adelaide.

Gulson, B.L. 1984. Uranium-lead and lead-lead investigations of minerals from the Broken Hill lodes and mine sequence rocks. *Econ. Geol.*, 79, 476-490.

Gulson, B.L., Mizon, K.J., Korsch, M.J., Palmer, J.M., and Donnelly, J.B. 2003. Mobilisation of lead from human bone tissue during pregnancy and lactation – a summary of long-term research. *Science Total Environ.*, 303, 79-104.

Gulson, B.L., Gillings, B.R., and Jameson, C.W. 1997. Stable lead isotopes in teeth as indicators of past domicile – a potential new tool in forensic science. *J. Forensic Sciences*, 42, 787-791.

Formatted: Font: Not Bold

Formatted: Font: Not Bold

Formatted: Indent Left: 1.27 cm, No bullets or numbering

Formatted: Font: Not Bold

Formatted: Indent Left: 1.27 cm, No bullets or numbering