We sincerely thank both reviewers for providing valuable feedback on the original version of this manuscript. Page and line numbers in the response below refer to the 'marked changes' manuscript.

**Reviewer 1**

**COMMENT:** I think one of the main issues of the study is the sampling strategy. First, I do not understand why the authors decided to focus on such small catchment areas (a few km² at most) when the aim of the study is to better constrain the composition of river sediment inputs from Svalbard to the oceans (cf. page 2, lines 32-35 and page 3, lines 1-10). What is special about this area?

**REPLY:** The catchments were selected for a number of reasons. First, the smaller the catchment, the easier it is to constrain the geology. Second, we wished to compare a glaciated and an unglaciated catchment with the same lithology, in order to assess the impact of glaciation on chemical fluxes of both the dissolved load and suspended sediments. Third, fieldwork in Svalbard is difficult and these catchments were logistically feasible (close to the nearest settlement). The aim of the study as previously written did not fully reflect the second factor and we have amended the text accordingly (P3 L6-7). (See also comments below regarding representativeness).

**COMMENT:** Are the lithologies exposed in these two catchment areas representative of what can be found elsewhere on Svalbard?

**REPLY:** We have added a panel to Fig. 2 to show the extent of the Central Basin, highlighting that the formations present in the two studied catchments are representative of a large areal proportion (8%) of the archipelago (P3 L4).

**COMMENT:** I found that there was a general confusion on what is considered a sediment source, a sedimentary rock or a sediment all through the manuscript. After several readings, I think I finally understood that the authors had collected random pieces of sedimentary rocks (not in place) in the catchment areas together with river, and supraglacial sediments and that they were trying to determine the nature of the sedimentary formations randomly sampled, the sources of the river and supraglacial sediments, and the sources of the sedimentary formations exposed in the catchment area. Again, if the aim is to constrain the composition of sediments exported from Svalbard to the Arctic ocean, I do not understand the sampling strategy of the authors. Why didn’t they sample the drained formations in place to know the isotopic compositions of the sources? How did they choose the pieces of bulk rock to analyze? The information given in Table 1 and 2 are also not very clear: Which samples are assumed to be the sources and which ones are the sediments for which the sources are unknown? Why are the stream sediments from Fardalen and Dryadbreen (samples L and O) and the sediment collected at the surface of a glacier (sample D) grouped together with bulk rocks collected in the catchment areas? This needs to be clarified. I would suggest that the authors clearly divide their samples into two groups: river sediments and potential sources; or first-cycle sediments and second-cycle sediments; or sedimentary rocks (old consolidated sediments exposed in the catchment area) and sediments (modern river, stream, glacial sediments).

**REPLY:** We are grateful to the reviewer for highlighting the inconsistencies in our use of ‘sediment’ and have corrected usage throughout the manuscript.

The aim of this study was to study the effect of glaciation on dissolved and suspended matter fluxes. In order to interpret the suspended sediment data, we took representative samples from the two catchments to constrain the rock sources supplying sediment to the streams.

We agree that it was not clear which samples were sampled from formations and which were
modern sediments. We have re-organised the samples in Table 1 to clarify which samples are from a specific formation and which are modern sediments. In Table 2, we have added clarification as to which samples are sedimentary rocks and which are modern sediments. In addition, we compare our rock samples to the core samples analysed by Schlegel et al., 2013 on a Herron diagram to provide further confirmation linking the sedimentary rock samples to a specific Formation (Fig. 4). We have re-made Fig. 3 to only focus on the sedimentary rock samples and highlight where our samples plot in relation to the major element composition of the core samples reported in Schlegel et al. (2013). In Fardalen the Frysjaodden Formation is clearly exposed and we took two samples (R01 and G). In Dryadbreen the rocks could not be sampled in outcrop as the outcrops were inaccessible due to the glacier. Here we aimed to take a representative selection of samples. Thus for rock samples collected in Dryadbreen, we identify the Formation that the samples originated from by comparing our data (chemical composition and clay mineralogy) with that of Schlegel et al., 2013. (P6 L15-25).

COMMENT: The notation F and D (Fardalen and Dryadbreen) is also unclear in Table 1 since D is the name of a sample. I would simply add an additional column to mention clearly the name of the catchment area from which the samples were taken in Table 1.

REPLY: The description already states that sample D was collected from the surface of the glacier i.e. Dryadbreen. We do not have space to add another column but as this information is contained in Table 2 we have added a footnote to reference to this in Table 1.

COMMENT: There is a significant isotopic bias between samples from the Fardalen and Dryadbreen catchment areas (see specific comment below), how can the authors be sure that it is not a sampling bias?

REPLY: Please see reply below.

Specific comments
1. What controls the isotopic variability of the samples (grey trend in Fig. 3)?

COMMENT: The question is why are the contributions so variable from one sediment/sedimentary rock to another? Page 9, it is explained that each sedimentary formation may have a distinct isotopic signature. Why not, but I don’t think the arguments are convincing enough. Knowing the composition of a rock sampled in place from each formation would easily solve the problem but since this data is not available I think the authors should develop a little bit more this part of the discussion. Maybe they should add a figure to clearly compare the clay mineralogy and the major and trace element concentrations of each of the formations (published data) to the composition of their sedimentary rocks.

REPLY: We have added a figure (Fig. 4) identifying the origin of the five sedimentary rock samples (two of which were collected directly from the Frysjaodden Formation) and re-made Fig. 3 to only focus on the sedimentary rock samples and clearly highlight the literature data for each formation. The discussion of the major element compositions of the rocks has been moved to the Results section (P6 L15-25).

COMMENT: A striking feature of the author’s dataset is that the Dryadbreen and Fardalen suspended AND catchment rocks form two very distinct groups in the Nd-Sr isotopic space (see red and blue circles in Figure 3 in the attached pdf). This difference is discussed for the suspended sediments only (end of page 8 and page 9) but this actually also concerns ALL the other collected rocks. How do the authors explain this difference given that the same formations are present in the two catchment areas? Did they unexpectedly sample only one formation in each catchment area? Is it a sampling bias? I was wondering whether it could be related to the fact that one catchment is glaciated and not the other. Assuming each rock-type (shale, siltstone, sandstone), not each formation, has its own isotopic signature, would it be possible that sandstones, enriched in Archean erosion products, be more eroded by the glacier in Dryadbreen than they are in the unglaciated catchment? Knowing the sampling locations of
all the studied catchment rocks might help to understand this feature. Alternatively, assuming that the different formations exposed in the catchment area do have distinct Sr-Nd isotopic signatures, why would the presence of the glacier in Dryadbreen favors the erosion of the upper part compared to Fardalen? Is it directly related to the steepness of the eroded area and the location of the glacier? I think more explanations are needed here.

**REPLY:** By definition the formations contain different rock types: Frysjaodden primarily contains shale and Aspelintoppen primarily contains sandstone. Yes, the trends are related to rock-type but this is inextricably linked to the parent formations. The grouping of points referred to by the reviewer is a form of sampling bias: due to the effect of glacial erosion the Frysjaodden Formation has become covered in moraine material derived from the upper formations. The modern-day glacier is cold-based and is not contributing to erosion, but when it was active it would have transported sediment from the top of the catchment to the bottom depositing it in lateral and terminal moraines. Yes, in the end we sampled only the Frysjaodden Formation in the unglaciated catchment and primarily the upper (Battfjellet and Aspelintoppen) formations in the glaciated catchment. We have added text to clarify these points ([P9 L2-6 and L15-25]).

2. Effect of grain-size/mineral sorting

Page 9, lines 17-20: the authors argue that the isotopic difference observed between the clay fractions and the bulk rocks in Figure 3 (black and whites triangles) could be explained by a process similar to the one that creates the general trend (grey lines, Fig.3) in Sr-Nd isotopic space. However, when you link the bulk rocks to their respective clay fractions (see the red arrows in Figure 3 in the attached pdf), the variations point towards different endmembers (much steeper slope than the general trend). To me, this indicates that the intra-sample variability (clay-bulk) cannot be explained with the same processes or the same sources than the inter-sample variability. Initially, I though that the steeper trend was due to the ingrowth of radiogenic Nd since the formation of the Eocene sedimentary rocks but the difference of Sm/Nd ratios between the clay and the bulk fractions is too small to generate such a large variability in εNd in 50 Ma (only 0.1 to 0.4 εNd units according to a quick calculation). One would need to do a mass balance calculation for Nd and Sr but the low εNd value of sample R03 (the only coarse-grained sediment, presumably enriched in REE-rich accessory minerals) suggest that another mineral phase controls the Nd isotopic composition of the sedimentary rocks, probably heavy minerals (monazite, apatite, allanite; see Garzanti et al, EPSL, 2010; 2011 and Garçon et al, Chemical Geology, 2014). This mineral phase would plot below the grey trend. Anyway, my point is that the intra-sample and inter-sample variability are not easily comparable. Trying to understand why bulk and clay are different may be interesting in the scope of the study.

**REPLY:** The purpose of this section was to introduce grain-size variation as a known cause of variation in εNd and we accept this was not clearly done. We completely agree that the clay-sized fraction – bulk pair for a particular sample on a different mixing line to the inter-sample array. We have re-written this part, adding a separate section (Section 5.2.1) to discuss grain-size effects and briefly discuss the offset of the clay array to the bulk rock array ([P10 L1-3]). We do not dwell too much on the latter as the primary focus of the paper is to explain the large-scale inter-sample variations.

3. Effect of weathering/leaching

**COMMENT:** At page 10, the authors speculate on the nature the leached mineral phases based on the REE element pattern of the bulk and residual fractions after leaching (Fig. 7b). My first point is that Sr and Nd are likely not hosted by the same mineral phases, so the authors should address the problem separately for Nd and Sr. Using REE patterns to investigate the carrier phases of leached Nd is OK but this is certainly not a good way for Sr. Secondly, the authors should take advantage of the major and trace element data (Table 1) they measured in the bulk and residual fractions. There, one can see that the residues contain
much less CaO and Mn than the bulk phases. The loss of Ca could be linked to apatite or carbonate leaching but Mn is more difficult to explain, maybe more in favor of a leaching of Mn-oxides?

**REPLY:** We agree that the mineral sources of Sr and Nd should be considered separately and have reworded this section (Section 5.2.2). We discuss the major element data and have added measured P concentrations to Table 1.

**COMMENT:** The leaching experiment and the weathering effects are closely related together. I would discuss these two things in the same part of the discussion for more clarity. The section discussing weathering effect at page 7 is not clear and needs to be rewritten anyway so maybe it would be better to merge it with the leaching section to explain that weathering, in particular the formation of secondary minerals, is responsible for some isotopic variability but cannot account for the large range of isotopic compositions measured in this study.

**REPLY:** We agree that the structure of the paper would be improved by combining the leaching discussion with the discussion on chemical weathering effects. We have rearranged the discussion section accordingly.

4. Seasonal variability

I would be more careful with the effects of seasonality. There are only two samples taken at different times of the year and they do not show a significant variability. For Nd, 20120618F and 20120726F could be slightly different but the difference is barely resolvable if an uncertainty of ±0.6 εNd (2s) is considered, as suggested by the repeated measurements of JNd and the shale SCo-1. For Sr, the effect of seasonality seems to be bigger though.

**REPLY:** We have reworded this start of this section to be more clear which differences are resolvable given the analytical uncertainty. Propagating the 2SD error on 20120618F(0.2) and 20120726F (0.4) gives 0.4 (P12 L10-11). The discussion draws on evidence from previous literature studies which have identified seasonal variations in larger rivers.

5. Implications for Nd as a sediment source tracer (pages 11-12)

I generally agree with this section and I think it is a great idea to discuss the potential consequences of the re-organization of glaciers in catchment areas on short-time (annual) scales. However, the authors need to keep in mind that their studied area is pretty small and the bigger is the drained area, the more homogenous will be the composition of the exported sediments. Therefore, the global river exports from Svalbard to the oceans could be much more homogenous in composition than what they analyzed in this study. Also, this part of the discussion is a bit short and should be more developed/elaborated to strengthen the argument being made. Maybe the authors can try to model the worst-case scenario based on the Nd isotopic variability measured in this study, for example by quantifying by how much such a seasonal variability will change the relative source proportions in the study of Tütken et al. (2002)?

**REPLY:** This section was intended to highlight the impact of changes on glacial-interglacial timescales rather than annual scales. Yes, larger drainage areas would be expected to have a more homogenous sediment composition, but as we highlighted in this paragraph, even large rivers (Madeira River, Amazon and Ganges tributaries) show resolvable seasonal variation in εNd. Over longer timescales, glaciers can significantly redistribute sediment within a basin and reorganize the hydrology, potentially altering the εNd export. We intended to just highlight these ideas and they would require further data to develop further and model. Modelling with the data available is out with the scope of this study.

**Technical comments**

**COMMENT:** I found that Figure 1 was not enough detailed. I think you should add more information to help the lecturers understand the context of the study. First, I would enlarge the small map of Svalbard to show the different areas you mention in the text on Svalbard but also around (Central basin, Spritsbergen, Ny Friesland, Longyearbyen, Greenland, Arctic
I’m not a specialist of the Arctic region and Svalbard and had to download several maps to be able to follow your discussion on the sources and the seawater composition, so a general map at the beginning would definitively help. Then, you should show the sampling locations of ALL samples in Figure 1. As it is now, it’s unclear to me which ones are shown and why. Since the suspended sediments were taken from rivers, maybe you can also show the rivers on the figure.

REPLY: We agree that more background information should be provided. We have added a new figure highlighting the circum-Arctic region to show the Paleogene position of Svalbard relative to the proposed source areas (Fig. 1). We have added two panels to the catchment map figure to show Svalbard and the extent of the Central Basin and the stratigraphy of these formations. Further locations have been included in the catchment map (Fig. 2).

COMMENT: P1 L6: Are you talking about the consolidated sediments (drained sources) or modern river sediments here? Given the cited range of values, it looks like you included the drained sources, why?

REPLY: We have changed the text and now only refer to the stream suspended sediments. (P1 L6)

COMMENT: P2 L5: Does this paper mention the Arctic region?

REPLY: No it does not. We have changed the reference to a paper that does focus on the Arctic (Meinhardt et al., 2016). (P2 L3)

COMMENT: P2 L21: Fractionation of Sm from Nd during soil formation should not affect epsilon Nd values unless this soil was formed a few hundred million years ago. A lot of time is needed to see the effect of Sm-Nd fractionation on Nd isotopic compositions...

REPLY: Agree, the soils in the Öhlander et al., 2000 study are formed from 1.9 Ga bedrock, long enough for inter-mineral variation to occur in eNd. We have amended the text for clarification. (P2 L20-21)

COMMENT: P3 L17: Why did you focus your study on such a small area?

REPLY: Please see earlier reply to this question.

COMMENT: Do you think that river sediments from such a small area could be representative of the total river inputs of Svalbard into the Arctic ocean? There is a need for a bit more explanation here.

REPLY: We did not intend to claim that these sediments were representative of the whole of Svalbard, only of the Central Basin. The Central Basin dominates the southern part of Spitsbergen and there have been no prior Sr-Nd measurements in this area. We have amended the text to clarify this point. (P3 L2-7)

COMMENT: P4 L16: Are these from rivers? If yes, then indicate which rivers in Table 1 and Figure 1.

REPLY: Yes, we have amended the text throughout to refer to “Stream suspended sediments” and added a sentence in the method section to describe the sampling location (P4 L23-25). The locations were described in the footnote to Table 1 and locations were shown in Figure 1.

COMMENT: P5 L6: What is the blank level of this technique for Nd and Sr?

REPLY: The procedural blank for the digestion procedure was less than 0.1% of the amount of Sr and Nd in the sample (P5 L19-20).

COMMENT: P5 L10: Please provide data obtained for reference materials analyzed together with the studied samples in a Supplementary File so that the lectors get an idea of the accuracy and precision of your measurements

REPLY: We have added two appendix tables providing this information, one each for major and trace elements. (Tables A3 and A4)
COMMENT: P5 L18: Why?
REPLY: Samples were run in triplicate in order to obtain replicate measurements under different tuning conditions of the instrument in order to properly assess the external reproducibility. Text not amended.

COMMENT: P5 L23: Not that minor!
REPLY: Agree, we have removed the word 'minor'. (P5 L31)

P6 L15: Not necessarily.
REPLY: We have clarified this sentence with reference to the proportion of clay minerals in the samples obtained from XRD measurements of the bulk. (P7 L1-3)

COMMENT: P7 L10: What about mineral sorting during sediment transport? It's likely more important than weathering. I would say "sedimentary processes" rather than "weathering" here.
REPLY: Changed as suggested. (P8 L3)

COMMENT: P7 L16: Again here, this is NOT the fractionation of the Sm/Nd ratio during chemical weathering that generates epsilon Nd variations but the preferential release (or adsorption) of Nd from specific mineral phases (i.e. incongruent weathering).
REPLY: Agree. Text amended. (P10 L16-17)

COMMENT: P7 L20: This part is unclear. You did not measure the composition of individual minerals so how can you be sure that they share the same Nd isotopic compositions? Your results actually suggest the opposite: the fact that you measured distinct Nd and Sr isotopic compositions for clay, bulk and leached residues suggest that Nd and Sr are hosted by different mineral phases with different isotopic compositions in the studied sediments/sedimentary rocks. So the trend could be generated by mineral sorting.
REPLY: We have rewritten this section. We now discuss the effects of chemical weathering using the data from the suspended sediment leaching procedure to examine the effect of a labile phase with a distinct isotopic composition. (Section 5.2.2)

COMMENT: P8 L8: This is discussable since R03 is not on the main trend...
REPLY: We have changed this to -23.3 which is the $\varepsilon$Nd value of D (on the main trend). (P8 L19)

COMMENT: P8 L13: Maybe this can be refined a bit by using the median Sr and Nd concentrations of the two endmembers. Since the two endmembers represent very different lithologies (i.e. basalts vs. likely more felsic lithologies for the Archean terrane), you can at least define whether $r$ is $< 1$ in Figure 6.
REPLY: Using the median values of the same literature data used to define the isotopic composition of the end-members we obtain an $r$ value of 1.8. The basalt source is well defined (little scatter in the data), but the Sr/Nd ratio of individual rock samples from the western source varies from 0.5 to 167! We are not convinced that defining whether $r$ is greater to or less than 1 adds useful information to the discussion when the uncertainty in the felsic source is so great, and have therefore not discussed this further in the text.

COMMENT: P8 L24: Not sure I understand what you say here. Above, you show that the isotopic variability of the studied sediments is related to variable contributions of the two "proto-sources" but we still don't know how this variability was created: is it linked to the type of formation sampled/eroded (you do not provide the information on the origin of the sedimentary rocks so I can't tell)? Is it created by mineral sorting (fine-grained sediments with more basaltic erosion products and coarse-grained with more felsic products)? Is it linked to modern erosion (unglaciated or glaciated catchment)? There is a clear isotopic difference
between the two catchments. For me, this suggests that something is going on at the catchment scale, maybe irrespective of the sedimentary formations eroded.

REPLY: Schlegel et al. 2013 concluded that the geochemical variation between the Frysjaodden and Aspelintoppen Formation was caused by changes in chemical weathering. In Section 5.2.2, we rule out chemical weathering as an explanation for the range in radiogenic Sr and Nd values. This range must therefore be due to end-member mixing between distinct sources. In the re-writing of the discussion this paragraph no longer exists. How the variation might arise is discussed elsewhere in the discussion section.

COMMENT: P9 L1: This suggests that you collected the rocks randomly (rocks not in place) in the catchment areas...is it the case? Why would you do that when you have the geological map to tell you where to go to sample each of the formations exposed in the catchment areas?

REPLY: We have clarified the sampling strategy in the methods section (and please see reply to main comment 1). (Section 3)

COMMENT P9 L2: This is not very convincing. Maybe you can add a figure to show how your samples compare to these published data in terms of clay mineralogy and major elements.

REPLY: We have rewritten this section and added a new figure (please see reply to main comment 1). (P6 L15-25, Figs. 3,4)

COMMENT P9 L17: Ok but when you link the bulk to their respective clay fractions, the variations point towards different endmembers (see arrows on Figure 3 below). This indicates that the intra-sample variability (clay-bulk) cannot be explained with the same processes or the same sources than the inter-sample variability. This is something that should be discussed in more details.

REPLY: We agree and have rewritten this section (please see reply to main comment 2). (Section 5.2.1)

COMMENT: P10 L13: Ok for Nd but carbonate are highly concentrated in Sr, so you probably leached out a lot of carbonate Sr. This is also what the CaO concentrations suggest in Table 1. Acid-treated sediments are 2 to 10 times less concentrated in CaO then the bulk sediments for both the two catchments.

REPLY: Carbonate is leached out, but it is also possible that Sr derives from exchangeable sites on the clay minerals. We have added to text to discuss this (P10 L30 – P11 L3).

COMMENT: P10 L26: The Mn concentrations are divided by at least a factor of 2 between the residue and the bulk so you must have lost a Mn-rich phase.

REPLY: Agree, and this could release adsorbed Sr. We have added to text to discuss this (P10 L30 – P11 L3).

COMMENT: P10 L26: I would say that you can rule out the leaching of clays since the bulk-clay pairs vary following a much steeper trend in the Nd-Sr isotopic space (see red arrows in Figure 3 below).

REPLY: We agree that you can rule out leaching of bulk clay, but the leaching procedure employed could easily release Sr loosely bound in interlayer sites of I/S and this could well fall on a different trend to that defined by the bulk clay samples.

COMMENT: P10 L27: "younger" is not appropriate here. I understand what you want to say but it's better to use the term "more depleted".

REPLY: This sentence no longer exists in the revised manuscript.

COMMENT: P11 L14: I doubt you can create a 12 epsilon Nd variability with diagenetic processes just by mixing seawater and detrital Nd. Your leaching is pretty strong and it's quite
possible that you start to dissolve and/or leach out Nd from primary minerals with such a procedure.
REPLY: Agree. We have rephrased this sentence to more clearly rule out this process as the primary factor causing variation. (P12 L4-7).

COMMENT: P11 L21: Almost not resolvable given the reported external precision (± 0.6 epsilon unit).
REPLY: Please see reply to main comment 4.

COMMENT: P11 L30: You cannot say that when you analyzed only 2 samples AND the seasonal variability is barely resolvable (actually not resolvable for Dryabreen).
REPLY: This sentence was intended to refer to the larger sample size of our data and previously published literature data. We added in the references for the other studies for clarification. (P12 L20).

COMMENT: P12 L15: It's rather unusual to cite a figure in the conclusion for the first time.
REPLY: We have now added references to this figure in the main discussion section of the manuscript.

COMMENT: Fig. 3: When you link the pairs clay-bulk (red arrows), you see that the direction of Sr-Nd isotopic variability generated by grain-size sorting is not the same as the general trend you show in grey.
REPLY: Please see reply to main comment 2.

COMMENT: Fig. 5b: Please explain what the black points are.
REPLY: The black points are the other bulk samples, for which the isotopic composition of the clay-sized separate was not measured. This information is now included in the figure panel. (Fig. 7)

COMMENT: Table 1: Please report in-run errors (2SE) for Nd and Sr isotopic compositions.
REPLY: We have included the 2SD of the 3 replicates. (Table 1)

COMMENT: Table A1: You do not discuss this data in the text. Maybe you can add a few lines in the text to indicate which one of the two dataset is the best and how do they compare.
REPLY: The full sample suite was analysed by ICP-OES only and due to HF digestion we did not obtain Si concentrations. In order to assess whether Si concentrations obtained by difference were accurate we compared ICP-OES data with four samples previously measured by XRF at an earlier data at a different institution. The table shows excellent agreement between the two techniques, giving us confidence that the Si concentrations obtained by difference are accurate. We have added text to refer to this table in the methods section. (P5 L16-20)

Reviewer 2

COMMENT: It is not clear in the current manuscript how representative these data are of the whole Svalbard area. Indeed, the total area of these two catchments corresponds to 0.01% of the total Svalbard surface (61 022 km2), which is very small. In what extent the data from these two catchments can be considered as representative or not of the Svalbard source region for paleo-erosion source-mixing reconstruction?
REPLY: We consider the two studied catchments representative of the Central Basin, and calculate that the Central Basin covers approximately 4800 km², corresponding to 8% of the total Svalbard area. The ‘effective’ proportion contributing sediment may be even larger as
this region is relatively unglaciated compared to other parts of Svalbard e.g. Nordaustlandet. We have added text to clarify this point (P3 L2-7).

COMMENT: In the absence of any information about fluxes, it is difficult to conclude about the integrated contribution of these two catchments and hence whether there is indeed a significant difference between the contribution of glacial and non-glacial areas in Svalbard (as claimed by the authors in lines 30 to 33).
REPLY: The two catchments have different suspended sediment isotopic compositions. As the reviewer correctly states, in the absence of flux measurements we cannot determine the modern contribution of fluxes from the glaciated versus the non-glaciated catchment. However, that there is a difference, implies that for a given point in space, glacial-interglacial cycles can redistribute sediment on land resulting in a non-constant isotopic composition of the suspended sediment flux. We have clarified these sentences (P12 L30).

COMMENT: The authors show that the clay mineral fractions have higher εNd and lower 87Sr/86Sr compared to bulk sediments. The implication of this result, which is not discussed in the paper, is that the Sr-Nd isotope composition of detrital sediments is a function of grain-size. This grain-size control is important to take into account for interpretation of Arctic ocean marine sediments paleo-record.
REPLY: We agree that grain-size control is vital to take into account in interpreting the paleo-record from ocean marine sediments. The impacts will depend on distance of the sediment core from source areas and this is currently a topic of active investigation.

COMMENT: Characterization of the leachable phase of Arctic river sediments: the authors do not give any justification for why they have conducted leaching experiment on these sediments (in section 5.3).
REPLY: The leaching experiment provided valuable information of the isotopic composition of labile phases that could be released during modern weathering processes. We have added text (P10 L28) to provide the justification.

COMMENT: Does the presence of an acid-leach labile phase isotopically distinct from the bulk silicate sediments have any implication for studies looking at the isotope composition of the marine authigenic phase (carbonate and oxides fractions recording the marine isotope composition) of marine sediments? In other words, is it possible that part of the signal extracted by leaching methods (the classical Hydroxylamine Hydrochlorid (HH) in Acetic Acid solution extraction) is inherited rather than authigenic (marine)?
REPLY: Yes, we think that an isotopically distinct leachable phase does have implications for the interpretation of Nd isotopes in the traditional HH sediment leach. It would seem to be possible that sediment containing an authigenic phase deposited in e.g. Paleocene could be re-mobilised and deposited in the modern ocean, potentially affecting the extracted εNd value. The impact of inherited phases is not currently well understood but initial work has been conducted e.g. Bayon et al., 2004 EPSL, 224, 477-492 and Charbonnier et al., 2012, 99, 39-56. As this paper focuses on the solid phases we have not amended the manuscript text, but are working on a companion manuscript that addresses this point.

COMMENT: The authors should provide more information about the samples other than just “samples were collected at different times during the melt season” in the main text in section 3. How many times during the melt season? How much different was the runoff between the different sampling dates? What are the suspended sediment concentration? Is there any available information about the fluxes (water and sediments) for these two catchments?
REPLY: Additional information on the collection of the stream suspended sediment samples is now provide in the Methods section (P4 L21-29). Due to difficulties setting up a gauging station during spring when there was still significant snow cover, discharge data is only available for a 9 day period in the summer. With such a limited data set we are unfortunately not able to extrapolate to seasonal fluxes.
Clay mineralogy, strontium and neodymium isotope ratios in the sediments of two High Arctic catchments (Svalbard)

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Abstract. The identification of sediment sources to the ocean is a pre-requisite to using marine sediment cores to extract information on past climate and ocean circulation. Sr and Nd isotopes are classical tools with which to trace source provenance. Despite considerable interest in the Arctic Ocean, the circum-Arctic source regions are poorly characterised in terms of their Sr and Nd isotopic compositions. In this study we present Sr and Nd isotope data from the Paleogene Central Basin sediments of Svalbard, including the first published data of stream suspended sediments from Svalbard.

The stream suspended sediments exhibit considerable isotopic variation (εNd = -20.6 to -13.4; ⁸⁷Sr/⁸⁶Sr = 0.73421 to 0.74704) which can be related to the depositional history of the sedimentary formations from which they are derived. In combination with analysis of the clay mineralogy of catchment rocks and the sediments, we suggest that the Central Basin sedimentary rocks were derived from two ‘proto’ sediment sources. One source is Proterozoic sediments derived from Greenlandic basement rocks which are rich in illite and have high ⁸⁷Sr/⁸⁶Sr and low εNd values. The second source is Carboniferous to Jurassic sediments derived from Siberian basalts which are rich in smectite and have low ⁸⁷Sr/⁸⁶Sr and high εNd values. Due to a change in depositional conditions throughout the Paleogene (from deep-sea to continental) the relative proportions of these two sources varies in the Central Basin formations. The modern stream suspended sediment isotopic composition is then controlled by modern processes, in particular glaciation, which determines the present-day exposure of the formations and therefore the relative contribution of each formation to the stream suspended sediment load. This study demonstrates that the Sr and Nd isotopic composition of stream suspended sediment from the continents exhibits significant seasonal variation, which likely mirrors longer-term hydrological changes, with implications for source provenance studies based on fixed sources through time.

1 Introduction

Since the Miocene, the Arctic has been subject to the repeated advance and retreat of ice sheets, a record of which is preserved in ocean sediments (Svendsen et al., 2004; Knies and Gaina, 2008). Thus, the Arctic Ocean and its surrounding seas are a key region for developing our understanding of past ice sheet dynamics and climate. A considerable number of ocean cores have been drilled in this region, allowing us to access the sedimentary archive (e.g. Vogt et al., 2001; Spielhagen et al., 2004; Knies and Gaina, 2008; Hillaire-Marcel et al., 2013; Fagel et al., 2014; Meinhardt et al., 2016). They provide information on past
ocean chemistry through analysis of foraminifera (e.g. Knies et al., 2014), on iceberg abundance through analysis of ice-rafted debris (IRD, e.g. Spielhagen et al., 2004) and on past sediment sources through analysis of the mineralogy and geochemistry of the sediment (e.g. Meinhardt et al., 2016).

Clay mineralogy in ocean sediment cores is often used to reconstruct paleoclimate and paleoceanography (e.g. Winkler et al., 2002). However, both the source rock and weathering conditions on the continents affect which clay minerals are formed. For example, kaolinite is more likely to form in tropical climates with intense chemical weathering, whereas illite is more likely to form where physical weathering dominates (Singer, 1984). The weathering of basalt will likely lead to the formation of smectite, whereas the weathering of granite will likely lead to the formation of illite or kaolinite (Essington, 2004). The Arctic Ocean has a wide variety of source regions ranging from young basalts in Siberia to the Precambrian rocks of Greenland and it is therefore imperative for paleoclimate studies to identify which changes in clay mineralogy are related to a change in climate and which are related to a change in source region.

The radiogenic isotope tracers $^{143}\text{Nd}/^{144}\text{Nd}$ (expressed as $\varepsilon$Nd) and $^{87}\text{Sr}/^{86}\text{Sr}$ are often used together to understand where and how sediment is generated and weathered, enabling source regions to be characterised (Goldstein and Jacobsen, 1988; Cameron and Hattori, 1997; Tricca et al., 1999; Peucker-Ehrenbrink et al., 2010; Lupker et al., 2013; Clinger et al., 2016). $\varepsilon$Nd is particularly suited to being a source tracer because, unlike Sr and Rb which are fluid mobile, Sm and Nd are immobile and behave very similarly during chemical weathering such that the Sm/Nd ratio does not fractionate during weathering (McCulloch and Wasserburg, 1978) and therefore variations in $\varepsilon$Nd are predominantly controlled by age (Goldstein and Jacobsen, 1988) rather than the weathering of specific minerals which can affect the Rb-Sr system (e.g. Bullen et al., 1997).

Although weathering effects on the neodymium isotopic composition of sediments are considered essentially negligible, it has been shown that in certain circumstances preferential leaching of minerals with a different Sm/Nd ratio to bulk rock can lead to variations in $\varepsilon$Nd in soil profiles (Öhlander et al., 2000; Aubert et al., 2001), which could affect the $\varepsilon$Nd value of the derived sediments. Variations within soil profiles are either attributed to dust inputs (Viers and Wasserburg, 2004; Ma et al., 2010) and/or the dissolution of accessory phases such as phosphate minerals or Fe-Mn oxyhydroxides (Goldstein and Jacobsen, 1987; Öhlander et al., 2000; Aubert et al., 2001; Babechuk et al., 2015). Additionally, $\varepsilon$Nd variations in ocean sediments are often interpreted in terms of changing source regions with the isotopic composition of individual source regions remaining constant though time. However, recent studies have hinted at seasonal variations in $\varepsilon$Nd in river sediments (Viers et al., 2008; Garçon et al., 2013; Lupker et al., 2013), raising the possibility that the $\varepsilon$Nd value of sediment exported from individual source regions may not remain constant over time. Thus, although $\varepsilon$Nd is a reliable tracer of source, one region may contain multiple sources, whose relative contributions vary over time.

The source regions to the Arctic Ocean are in general relatively poorly characterised in terms of coupled $\varepsilon$Nd and $^{87}\text{Sr}/^{86}\text{Sr}$ measurements with only a few samples from the major rivers and shelf sediments (Eisenhauer et al., 1999; Hillaire-Marcel et al., 2013). Svalbard (Fig. 1) is particularly important to characterise owing to its location by the Fram Strait, which is the site of deep water formation essential to the functioning of the global thermohaline circulation, and has therefore been the target of studies seeking to understand the formation of the Atlantic-Arctic gateway (Jakobsson et al., 2007). However, there is neither $^{87}\text{Sr}/^{86}\text{Sr}$ nor $\varepsilon$Nd data on river sediments from Svalbard. Previous studies (Tütken et al., 2002; Maccali et al., 2013)
have taken the bedrock data of granitoids from Ny Friesland in the north-east of Spitsbergen (the largest island in the Svalbard archipelago, Fig. 2, Johansson et al., 1995; Johansson and Gee, 1999) to represent the Svalbard source region. Svalbard has a wide range of rocks from different ages and in this study we characterise sediments from two catchments located in the Paleogene Central Basin, which comprises 8% of the land area of Svalbard, and from which no prior Sr-Nd measurements exist. Being relatively sparsely glaciated and more easily eroded, these sedimentary formations could constitute an important sediment source from Svalbard. The studied catchments have nominally identical lithology but one is glaciated, allowing us to examine the effect of glaciation on stream suspended sediment composition. As clay minerals are the main constituent of the rocks in the study area, we combine the geochemical data ($\varepsilon$Nd and $^{87}$Sr/$^{86}$Sr) with clay mineralogy in order to identify the factors influencing the unexpectedly large variation in radiogenic isotope compositions observed in the catchment sediments.

2 Field site

Svalbard is located in the Arctic Ocean (Fig. 1) and has an Arctic climate. In 2012 (the year samples were collected) the mean temperature was -2.0 °C and precipitation was 268 mm, as recorded at Longyearbyen Airport (Nordli et al., 2012). Permafrost is continuous throughout the islands (Humlum et al., 2003). Two catchments were studied which are situated adjacent to each other approximately 5 km south of Longyearbyen in central Svalbard (Fig. 2a).

The first catchment is a permafrost-affected valley called Fardalen (Fig. 2c) which is likely to have been unglaciated for at least the last 10 kyr (Svendsen and Mangerud, 1997). The catchment area is 3.4 km$^2$ and ranges in elevation from 250 - 1025 m.a.s.l. The second catchment contains a glacier called Dryadbreen and ‘Dryadbreen’ will be used hereafter to refer to the whole catchment and not just the glacier. Between 1936 and 2006 the area of the glacier decreased from 2.59 to 0.91 km$^2$ leaving large terminal and lateral ice-cored moraines and a sandur in front of the glacier (Ziaja and Pipała, 2007). The catchment area is 4.8 km$^2$ and ranges in elevation from 250 - 1031 m.a.s.l.

2.1 Geological background

The two studied catchments are situated in the Paleogene sedimentary Central Basin of Svalbard (Fig. 2a). The sedimentary formations exposed in the catchments are from the Van Mijenfjorden group which is Paleocene to Eocene in age (66 - 33.9 Ma) and contains sandstones, siltstones and shale (Fig. 2b, Major et al., 2000). The formations exposed in the two catchments have been relatively well studied on account of the fact they cover the Paleocene-Eocene Thermal Maximum (PETM) and formed as a consequence of sedimentation which commenced upon the separation of Greenland from Eurasia (e.g. Helland-Hansen, 1990; Müller and Spielhagen, 1990; Cui et al., 2011; Dypvik et al., 2011; Elling et al., 2016; Jones et al., 2016).

The Central Basin sediments are divided into six formations (Major et al., 2000) which were deposited in a series of transgressive-regressive cycles. The youngest four are exposed in the studied catchments. It is thought that the source for the oldest sediments (mid-Paleocene to early Eocene) was from the east (Carboniferous to Jurassic Siberian basalts, Helland-Hansen, 1990), but this changed to the west (Proterozoic Greenlandic/Canadian High Arctic Shield) during early to mid-Eocene
with the erosion of the uplifted West Spitsbergen Fold and Thrust Belt whose formation is linked to rifting of the North Atlantic and the separation of Svalbard from Greenland.

The oldest formation exposed in the studied catchments is the Grumantbyen Formation (Fig. 2b), comprising shallow marine sandstones. The Grumantbyen sediments are part of a regressive trend with sediment derived from the east and possibly the north (Helland-Hansen, 1990). The youngest three formations comprise a regressive sequence with (from oldest to youngest) the Frysjaodden formation comprising fine-grained shales deposited offshore in an open basin; Battfjellet Formation comprising shallow marine sandstone; and Aspelintoppen comprising continental deposits (Helland-Hansen, 1990; Müller and Spielhagen, 1990). The mountain belt is thought to have eroded rapidly (Cui et al., 2011) based on the immaturity of the sandstones (Helland-Hansen, 1990). Detection of pre-Caledonian metamorphic detritus indicates that the mountain belt was eroded down into the basement rocks (Helland-Hansen, 1990). The PETM boundary is near the base of the Frysjaodden Formation (Charles et al., 2011).

3 Methods

A selection of 18 representative solid samples, including sedimentary rocks, bedload sediments and glacial sediment, were sampled from both catchments. The sedimentary rock samples were first crushed (jaw crusher) and were subsequently ground to fine powders (rotary disc mill and planetary ball mill). For the sediment samples, only the latter step was required. A subset of samples (Fig. 2c) were selected for element and isotopic analysis (Table 1) and they are further described here. For all other samples a brief description is included in Table 2. Samples ‘G’ and ‘R01’ are two separate samples of frost-shattered angular pieces of shale, 1 - 4 cm in length collected from the Frysjaodden Formation in the unglaciated Fardalen catchment. Samples ‘R02’ (frost-shattered wacke), ‘R03’ (litharenite), ‘R04’ (shale) and ‘D’ (coarse sediment from glacier surface) were collected from the glaciated Dryadbreen catchment where transport and physical erosion by the glacier has combined rocks from different formations. Additionally, sediment from the stream channels of each catchment were collected (samples ‘L’ and ‘O’). Stream suspended sediment (>0.2 µm) was retrieved from nylon filter papers during water sample collection (Hindshaw et al., 2016) by washing the filter paper with deionized water and then freeze drying the sample. Stream suspended sediment samples were collected from each of the catchment streams on alternate days during June (snow melt) and July/August (summer) 2012, just before their confluence with the main valley stream (Fig. 2c). Part of each sample was treated with 5% HCl to remove carbonates. The leachates were not retained. A subset of four samples, one from each season and each catchment were analysed for this study (Table 1). Discharge measurements are only available for summer; the discharge at the time of sample collection for the two summer samples was 0.24 m s⁻¹ (Fardalen) and 0.35 m s⁻¹ (Dryadbreen) (Hindshaw et al., 2016). Suspended sediment concentrations at the time of sampling are reported in Table 1.

3.1 Semi-quantitative determination of clay mineral abundances

A <2 µm size-fraction was separated from bulk sediment by repeatedly rinsing and re-suspending the sample in de-ionized water with sodium phosphate as a dispersal agent, followed by sonication. The <2 µm fraction was separated by centrifugation
(Moore and Reynolds Jr., 1997) and was then transferred to a clean glass slide in preparation for XRD using a filter-peel technique to orientate the sample (Moore and Reynolds Jr., 1997). XRD analysis was performed on a PANalytical PW1050 X-ray diffractometer with a Hiltonbrooks DG2 X-ray generator (Co-Kα radiation) at the University of St. Andrews. Data were collected between 5 and 40° with a step size of 0.02° and a counting time of 3 s per step. Spectra were collected on air-dried, glycerol-treated, 450°C heated and 550°C heated samples. The glycerol-treated and 450°C spectra were used to obtain semi-quantitative clay mineral abundances using the method outlined in Griffin (1971) which uses the peak heights of kaolinite 001 and illite 001 in the 450°C spectrum and the peak heights of chlorite 004, kaolinite 002, kaolinite 001 and illite 001 in the glycerol treated spectrum to give the relative abundances of kaolinite, chlorite, illite (mica) and expandable layer clay minerals (e.g. smectite and mixed-layer minerals containing smectite).

3.2 Chemical and isotopic composition

A selection of bulk solid samples and separated clay fractions were analysed for major and trace element chemistry using the following method: approximately 100 mg of material was ashed at 950°C for 120 minutes. The sample was then digested in a mixture of concentrated hydrofluoric and nitric acids and repeatedly dried down and re-dissolved in 6M HCl. In the final step, the dried down sample was re-dissolved in 2% HNO₃. Major and trace element concentrations were measured at the University of Cambridge by inductively-coupled plasma optical emission spectrometry (ICP-OES, Agilent Technologies 5100) and quadrupole inductively-coupled plasma mass spectrometry (Q-ICP-MS, Perkin Elmer 63 Nexion 350D), respectively. Si concentrations were calculated by difference assuming 100% recovery. The accuracy of the Si concentrations obtained in this way was confirmed by comparison with XRF obtained for four samples (Table A1). The accuracy of the concentration measurements was verified by repeated measurements of seven different USGS rock standards (Tables A3 and A4), including two shale standards (SCo-1 and SGR-1b). The procedural blanks through the digestion procedure were negligible (<0.1% sample) for Sr and Nd.

Neodymium was separated from the matrix using the method described in Piotrowski et al. (2009). This method uses two columns. The first column contained Eichrom TRUspec resin which separates out REE from the matrix and the second column contains Eichrom LNSpec resin to isolate Nd. The radiogenic neodymium isotopic composition was measured on a Nu plasma (Nu Instruments, University of Cambridge) multi-collector inductively coupled plasma mass spectrometer (MC-ICP-MS). Samples were run at 50-75 ppb with an APEX ACM sample introduction system. Samples were run in triplicate (three measurements on different days) with each measurement comprising 30 cycles with 10 s integration. Samarium interferences were monitored by measuring mass 149. No interferences were detected and oxides were monitored during tuning to ensure they were well below 0.5% of the beam size. The exponential law was applied to correct for instrument mass fractionation and all $^{143}\text{Nd}/^{144}\text{Nd}$ ratios were normalised to $^{146}\text{Nd}/^{144}\text{Nd} = 0.7219$. Standard-sample were bracketing was employed in order to correct for the minor offset with the accepted JNd-1 value: 0.512060±0.000024 (2SD, n=119) compared with the accepted value of 0.512115 (Tanaka et al., 2000). The USGS shale standard SCo-1 was measured and the $^{143}\text{Nd}/^{144}\text{Nd}$ value of 0.512086±0.000029 (n=3, 2SD) is in agreement with a previously published value of 0.512117±0.000010 (2σ, n=20 Krogstad
et al., 2004). Neodymium ratios are reported as deviations relative to the chondritic uniform reservoir (CHUR, $^{143}\text{Nd}/^{144}\text{Nd} = 0.512638$, Jacobsen and Wasserburg, 1980).

Strontium was separated from the matrix using Biorad microp bio-spin columns with Eichrom SrSpec resin (Hindshaw, 2011). The radiogenic strontium isotopic compositions were measured on a Neptune MC-ICP-MS (Thermo, University of Cambridge) and were run at 50 ppb using an APEX sample introduction system. Samples were run in triplicate (three measurements on different days) with each measurement comprising 30 cycles with 8 s integration. $^{85}\text{Rb}$ was monitored to correct for rubidium interferences on $^{87}\text{Sr}$ and data were additionally corrected for Kr interferences by measuring $^{83}\text{Kr}$. The exponential law was applied to correct for instrument mass fractionation and all $^{87}\text{Sr}/^{86}\text{Sr}$ ratios were normalised to $^{86}\text{Sr}/^{88}\text{Sr} = 0.1194$. Measurements of NBS 987 gave a $^{87}\text{Sr}/^{86}\text{Sr}$ value of $0.710249 \pm 0.000029$ (2SD, n=27) and the seawater value was $0.709188 \pm 0.000024$ (2SD, n=9), which is within error of the accepted value of $0.709179 \pm 0.000008$ (2σ, n=17 Mokadem et al., 2015).

### 4 Results

The major and trace element concentrations of the solid samples are provided in Table 1. The measured values are typical for shales (Taylor and McLennan, 1985) and the rare earth element (REE) chondrite normalised element profile of these samples closely follows that of the Post Archaean Australian Shale (PAAS, Table A2). Strontium and neodymium concentrations varied from 78 to 139 and 25 to 49 ppm, respectively. The two rock samples collected from the Frysjaodden Formation (R01 and G) have very similar major element compositions to core samples from this formation (Fig. 3, Schlegel et al., 2013). The formations in the glaciated catchment were not clearly exposed due to the presence of moraine material. However, the core sample from the Asplintoppen Formation is classified as a litharenite (Fig. 4) and as sample R03 is a litharenite and has similar major element chemistry (Fig. 3), we infer that this sample is also derived from the Aspelintoppen Formation. Sample R02 plots close to the wacke-shale boundary in Fig. 4 and in terms of major element chemistry it plots on the edge of the area defined by the Frysjaodden samples (Fig. 3). We therefore infer that this sample originated close to the boundary between the Frysjaodden and Battfjellet formations. Sample R04 is classified as a shale (Fig. 4) but its major element composition is distinct from the Frysjaodden Formation samples, in particular for $\text{Fe}_2\text{O}_3$ (Fig. 3). Given that this sample was collected on the surface of the glacier, it is likely that it is a shale derived from either the Battfjellet or Aspelintoppen Formation. These assignments are corroborated by the clay mineralogy (see below).

There was a large range in both the strontium and neodymium isotopic compositions of the bulk solid samples: $^{87}\text{Sr}/^{86}\text{Sr} = 0.72449$ to 0.75243 and $\varepsilon_{\text{Nd}} = -24.2$ to -11.9 (Table 1). Radiogenic Sr and Nd values are inversely correlated (Fig. 5) and in general, samples collected from the unglaciated Fardalen catchment e.g. R01 and G have higher $^{87}\text{Sr}/^{86}\text{Sr}$ and lower $\varepsilon_{\text{Nd}}$ values than those samples collected in the glaciated Dryadbreen catchment e.g. D and O (Table 1). The clay-sized fraction forms a parallel array to the bulk rock samples in $\varepsilon_{\text{Nd}}-^{87}\text{Sr}/^{86}\text{Sr}$ space (Fig. 5), with the clay-sized samples having higher $\varepsilon_{\text{Nd}}$ and lower $^{87}\text{Sr}/^{86}\text{Sr}$ values (except for R01). The $\varepsilon_{\text{Nd}}$ values of clay fractions were 2.1 to 3.2 epsilon units higher than the corresponding bulk sample and $^{87}\text{Sr}/^{86}\text{Sr}$ values were 1030 to 2030 ppm lower in the clay compared to the bulk, apart from sample R01 where the clay was 1100 ppm higher in the clay than in the bulk. Rubidium,
strontium, neodymium and samarium concentrations in clay samples are comparable to, or higher than, bulk values (Table 1). Given that clay minerals constitute >88% of the non-quartz minerals in these samples (Hindshaw et al., 2016), clays are the main host of these elements. It has been observed that in a compilation of river sediments from all over the world that $\varepsilon$Nd in the clay fraction is greater than in the silt-sized fraction by an average of 0.8 epsilon units (Bayon et al., 2015). Fine sediments (as measured by Al/Si ratio) from the Mackenzie River have also been observed to have higher values than coarser sediments (Vonk et al., 2015). The offset in $\varepsilon$Nd between fine and coarse fractions has been interpreted to reflect the preferential transport of basalt and volcanics in the fine fraction (McLennan et al., 1989; Garçon and Chauvel, 2014; Bayon et al., 2015). A volcanic signal is typically only observed in the first sedimentary cycle, due to the rapid chemical weathering of volcanic particles (McLennan et al., 1989) and therefore, if volcanics are present, they must have been deposited at the same time as the Central Basin sediments. Potential volcanic sources for this period could be the volcanic provinces of North Greenland and Ellesmere Island (58-61 Ma, Jones et al., 2016).

The residues of suspended sediments (collected on 0.2 $\mu$m filter paper) after treatment with 5% HCl had higher $^{87}$Sr/$^{86}$Sr (3528-4832 ppm) and lower $\varepsilon$Nd values (1.0 to 4.6 epsilon units) than the corresponding unleached sediment.

4.1 Semi-quantitative clay abundance

Illite, chlorite and kaolinite were present in all the samples analysed. In addition, the presence of an expandable layer clay mineral is also evident in the collapse of the XRD signal around 12.7 Å ($8^\circ2\theta$ Co radiation) between the air-dried and glycerol-treated spectra (Fig 6). Additionally, the asymmetry of the illite 001 peak (Fig. 6) suggests that this expandable layer clay mineral is an illite-smectite mixed-layer phase (Moore and Reynolds Jr., 1997) and this is in agreement with the interpretation by Dypvik et al. (2011) of XRD spectra from core samples from the same formations exposed in the studied catchments. This mixed layer expandable phase will be referred to as ‘I/S’ in the following discussion. The relative proportions of illite, chlorite, kaolinite and I/S are given in Table 2.

The solid samples collected from the glaciated Dryadbreen catchment tend to have higher illite abundances and lower I/S abundances than those samples collected from the unglaciated Fardalen catchment (Table 2). For all samples, there is an inverse relationship between the relative abundances of I/S and illite (Fig. 7b). The relative abundances of kaolinite and chlorite were similar in both catchments (Table 2). We are not able to distinguish between authigenic and detrital clay minerals.

The clay-sized fraction of the sedimentary rock samples from the Frysjaodden Formation (R01, G) have a lower relative proportion of illite than the samples inferred to be derived from the Battfjellet (R02) and Aspelintoppen (R03) formations (Fig. 7). This is in agreement with the decrease in the relative proportion of illite observed in the clay-sized fraction of core samples from the Aspelintoppen Formation (64%) to the Frysjaodden Formation (51%) (Schlegel et al., 2013). Samples with a high relative proportion of illite had high $^{87}$Sr/$^{86}$Sr and low $\varepsilon$Nd values compared to those with a low relative proportion (Fig. 7c,d).
Discussion

The variation in clay mineralogy, $^{87}\text{Sr}/^{86}\text{Sr}$ and $\varepsilon$Nd could be caused by three scenarios: mixing between two or more sediment sources, sedimentary processes (either modern or in the Eocene), or a combination of source and process effects.

5.1 Sediment sources

The linear trend between $^{87}\text{Sr}/^{86}\text{Sr}$ and $\varepsilon$Nd is suggestive both of a common regional process affecting both isotope systems (Goldstein and Jacobsen, 1988) and of mixing between two end-members. The sediments deposited in the Central Basin during the Eocene were themselves derived from Mesozoic sediments. Based on zircon dating, it is thought that during the Mesozoic, the sediment source to Svalbard alternated between an older (Proterozoic) western component comprised of reworked sediments from Greenland and Canada and a younger (Carboniferous-Jurassic) eastern component from Siberian fold-and-thrust belts (Fig. 1, Bue and Andresen, 2014; Elling et al., 2016; Petersen et al., 2016). The erosion of the Siberian Traps which formed within the same time period (Permian to Triassic) would also have contributed sediment to the ocean. We will refer to these two sources as ‘East’ and ‘West’.

The eastern source (Fig. 1, Lightfoot et al., 1993; Wooden et al., 1993; Spadea and D’Antonio, 2006) is relatively well-defined since the samples are essentially mono-lithologic (basaltic) and were deposited over a relatively short time-period. Based on zircon dating, both the Uralides and the Verkhoyansk Fold-and-Thrust Belt (Fig. 1) have been identified as potential sources to Svalbard Mesozoic sediments and the Paleocene Basilika and Firkanten formations (Fig. 2b, both underlie the Grumantbyen Formation) (Bue and Andresen, 2014; Elling et al., 2016; Petersen et al., 2016). The western end-member is much harder to characterise as it consists of Precambrian rocks which have undergone extensive metamorphism. An end-member with $\varepsilon$Nd values lower than -23.3 is required (D, Table 1) and therefore only data from western Greenland is considered (Fig. 1). The East coast was affected by the Caledonian orogeny (Henriksen, 1999) and later by the rifting of the North Atlantic (Bernstein et al., 1998) and therefore has higher (younger) $\varepsilon$Nd values (Jeandel et al., 2007). The range in $^{87}\text{Sr}/^{86}\text{Sr}$ and $\varepsilon$Nd from literature data of Archaean rocks from western and northern Greenland (Jacobsen, 1988; Collerson et al., 1989; Weis et al., 1997; Kalsbeek and Frei, 2006; Friend et al., 2009) is 0.70153 to 2.33356 and -56 to -2.75. By changing the Sr/Nd mass ratio of these two end-members, mixing lines can be drawn which encompass all the data, with the majority of points falling on a mixing line with an r value of 1 (i.e. the Sr/Nd mass ratio of both end-members is the same, Fig. 9).

The variation of $^{87}\text{Sr}/^{86}\text{Sr}$ and $\varepsilon$Nd in the clay-sized fraction forms a parallel trend to the bulk samples and can be explained by the same two end-members, but with a lower r value (Fig. 9). Variation in the r value between the bulk and the clay-sized fraction is to be expected given their different mineralogical compositions.

The variation in clay mineralogy (Fig. 7) can be explained by the different lithological sources of the two end-members (Fig. 9). Basalt typically weathers to smectite group minerals (e.g. Curtin and Smillie, 1981; Parra et al., 1985) and modern sediments originating from Siberia (basaltic) are enriched in smectite (Nürnberg et al., 1994; Wahsner et al., 1999). Any volcanic particles present will also tend to weather to smectite (Bayon et al., 2015). The western source is dominated by granitic rocks where the mica and K-feldspar typically weather to illite and kaolinite, respectively (Essington, 2004). Illite has high Rb/Sr ratios
and detrital illite is resistant to weathering, as the western source is older, this results in high $^{87}\text{Sr}/^{86}\text{Sr}$ and low $\varepsilon_{\text{Nd}}$ values. By contrast the younger eastern source will have lower $^{87}\text{Sr}/^{86}\text{Sr}$ and higher $\varepsilon_{\text{Nd}}$ values (Fig. 10). The lower $^{87}\text{Sr}/^{86}\text{Sr}$ and higher $\varepsilon_{\text{Nd}}$ values of the sedimentary rocks from the Frysjaodden Formation (G, R01) compared to those of the Battfjellet Formation (R04, Table 1) implies that the Frysjaodden Formation contains a greater proportion of the eastern end-member (Fig. 9). Indeed, zircon dating of samples from the Battfjellet Formation was consistent with a western, Greenland/Canadian Shield source, with an almost complete lack of Uralide aged grains (Petersen et al., 2016). The distinct differences between the two sources leads to the observed correlations between clay mineralogy, $^{87}\text{Sr}/^{86}\text{Sr}$ and $\varepsilon_{\text{Nd}}$ values (Figs. 7 and 10).

Schlegel et al. (2013) concluded that, on the basis of microscopic observations, the geochemical changes observed between the different formations arose as a result of increased chemical weathering during the late Paleocene and not as a change in source rock provenance, which remained from the west. However, that western source in the Paleocene-Eocene was itself comprised of two sources deposited in the Mesozoic. Chemical weathering during the Paleocene cannot be reconciled with the wide variation in $^{87}\text{Sr}/^{86}\text{Sr}$ and $\varepsilon_{\text{Nd}}$ values. Therefore, the overall trend observed between $^{87}\text{Sr}/^{86}\text{Sr}$ and $\varepsilon_{\text{Nd}}$ is more likely caused by two ‘proto-sources’ of different ages mixing: an illite-rich end-member with high $^{87}\text{Sr}/^{86}\text{Sr}$ and low $\varepsilon_{\text{Nd}}$ and an illite-poor end-member with low $^{87}\text{Sr}/^{86}\text{Sr}$ and high $\varepsilon_{\text{Nd}}$.

Mixing of two sediment sources can explain the difference between the stream suspended sediments collected in the two catchments (Fig. 5). Stream suspended sediments from the glaciated Dryadbreen catchment have lower $\varepsilon_{\text{Nd}}$, higher $^{87}\text{Sr}/^{86}\text{Sr}$ values and a greater relative proportion of illite, compared to those from the unglaciated Fardalen catchment. These observations can be readily accounted for if the stream suspended sediments from Fardalen receive a greater relative contribution from the Frysjaodden Formation (enriched in the eastern end-member) than stream suspended sediments from Dryadbreen (Figs. 2 and 10). The lower relative contribution of the Frysjaodden Formation to stream suspended sediments from the glaciated Dryadbreen catchment can be explained by the moraine material being predominantly derived from sedimentary rocks once located in the upper reaches of the catchment (Aspelintoppen Formation), and the modern day sandur plain, containing the products of this physical erosion, essentially burying the lower down Frysjaodden Formation. Changes in erosion caused by glaciation, which conveys sediment the head to the toe of the glacier, could therefore influence the Sr and Nd isotopic of sediments exported to the ocean (von Blanckenburg and Nägler, 2001).

5.2 Sedimentary processes

5.2.1 Grain-size sorting

Sediments are sorted as a function of particle size as they travel through the water, such that coarser particles (typically primary minerals such as feldspar and quartz) will settle faster than finer particles (clay minerals). A size-sorting effect is observed in the difference between the $^{87}\text{Sr}/^{86}\text{Sr}$ and $\varepsilon_{\text{Nd}}$ values of the bulk and the clay-sized (<2 µm) fraction. This effect is observed at a global scale and is interpreted to reflect the preferential transport of volcanics and basalt in the fine fraction (Bayon et al., 2015). However, mineral sorting between a clay-sized fraction and a coarser fraction cannot account for the overall trend in the bulk samples (Fig. 5) as the clay minerals plot on a parallel array to the bulk samples (Fig. 5) indicating that the clay-sized
fraction is not an end-member of the bulk sample trend. The offset between the clay-sized fraction and bulk is consistent with
the clay-sized fraction being enriched in authigenic phases with higher $\varepsilon_{\text{Nd}}$ and lower $^{87}\text{Sr}/^{86}\text{Sr}$ compared to the bulk which
is more likely to contain detrital minerals with a larger grain size.

Grain-size sorting could occur within the clay-sized fraction as illite will settle faster than smectite in the marine environment
(Sionneau et al., 2008). The Frysjaodden Formation, being furthest away from shore, could become enriched in smectite-
enriched particles derived from the basaltic eastern end-member whereas the Aspelintoppen Formation, deposited in a near-
shore environment, could become enriched in relatively coarser illite-enriched particles derived from the granitic western
end-member. As clays form $>88\%$ of the non-quartz mineral fraction of the bulk (excluding R03 which does not plot on
the main trend, Fig. 5, Hindshaw et al., 2016)), the trends observed between illite, smectite (I/S) and the radiogenic isotope
compositions (Fig. 7) should be reflected in the bulk samples (Fig. 5).

5.2.2 Preferential leaching of a labile phase

There are examples from previous studies where chemical weathering has been identified as the cause of an inverse correlation
between the relative proportions of illite and smectite (Fig. 7b, e.g. Setti et al., 2004). However, whilst modern day weathering
processes can induce large variations in $^{87}\text{Sr}/^{86}\text{Sr}$ primarily as a result of large inter-mineral variations in the Rb/Sr ratio (e.g.
Bullen et al., 1997), it is much harder to induce large variations in the Sm/Nd ratio of minerals and this ratio is often assumed
to remain constant once a rock has been formed (e.g. McCulloch and Wasserburg, 1978). Preferential release of minerals with
different Sm/Nd ratios during chemical weathering has been implicated in the generation of small $\varepsilon_{\text{Nd}}$ offsets of around 2
epsilon units (Rickli et al., 2013) and larger variations in $\varepsilon_{\text{Nd}}$ were observed in a soil profile developed on granitic till in
northern Sweden (Öhlander et al., 2000). In the latter study, a 7.7 epsilon unit variation was observed between the E horizon
and the humic horizon, which was attributed to the preferential weathering of minerals enriched in Nd over Sm e.g. allanite.

In these catchments there is essentially no soil development due to recent glaciation and at the bulk scale the mineralogy
of the rocks is broadly similar (Hindshaw et al., 2016) as are their Sm/Nd ratios (Table 1). Additionally, the major element
chemistry is very similar to that observed in core samples drilled through the same formations (Fig. 3, Schlegel et al., 2013),
confirming that weathering processes since the Paleocene have had minor impact on bulk element and, by inference, $^{87}\text{Sr}/^{86}\text{Sr}$
and $\varepsilon_{\text{Nd}}$ values. The geochemical changes observed between formations in the core samples (Schlegel et al., 2013) were
attributed to increased chemical weathering during the late Paleocene. In the given time period of this study (50 Ma) preferential
weathering of minerals with Sm/Nd ratios significantly different from bulk would be required in order to generate the 14 epsilon
unit variation in $\varepsilon_{\text{Nd}}$. In order to test this hypothesis, we applied a 5% HCl leach to remove easily leached mineral phases.

The chemical and isotopic composition of leached suspended sediment is distinct from bulk suspended sediment (Fig. 8a).

From mass balance constraints, this points to the existence of a labile pool containing Ca, P, Mn and Fe, with low $^{87}\text{Sr}/^{86}\text{Sr}$ and
high $\varepsilon_{\text{Nd}}$ (Table 1, Fig. 8a). Over 50% Ca is leached from the sediments with a Sr/Ca mass ratio of 0.002 to 0.007, consistent
with carbonate (Veizer, 1983), which would be expected to dissolve readily in 5% HCl (Tessier et al., 1979). A loss of P is
also observed (39-56%, Table 1) suggesting the dissolution of apatite (containing Ca and Sr), and the decrease in Fe and Mn
concentrations (Table 1) could be indicative of leaching Fe-Mn oxyhydroxides. In addition to apatite and carbonate, Sr could
be derived from exchange sites within the clay minerals or adsorbed onto Fe-Mn oxyhydroxide surfaces. Further, the leaching procedure applied in this study is relatively aggressive and could have dissolved part of the clay mineral structure (Chester and Hughes, 1967).

REE (rare earth element) patterns in leachates are commonly used to identify which mineral phases have been leached (e.g. Haley et al., 2004). The residual phase is depleted in MREE (middle REE) (Fig. 8b), implying the leachate is MREE enriched. A MREE enrichment is consistent with diagenetic apatite and Fe/Mn oxyhydroxides (Ohr et al., 1994; Tricca et al., 1999; Johannesson and Zhou, 1999; Su et al., 2017) and inconsistent with a carbonate phase, which is typically HREE enriched (Byrne and Kim, 1990; Millero, 1992; Byrne and Sholkovitz, 1996), suggesting that Nd in the leachate is derived from the former phases. Volcanic ash also has a MREE enriched REE pattern (Tepe and Bau, 2014) and would have high εNd and low 87Sr/86Sr. However, the amount of a volcanic component is expected to be minor in the studied sediments as the volcanic ash component of particulates readily leaches upon contact with seawater (Pearce et al., 2013; Wilson et al., 2013) and therefore may already have been leached during deposition in the Paleocene-Eocene. Additionally, volcanic ash in these layers has been diagenetically altered to bentonites (Cui et al., 2011; Elling et al., 2016; Jones et al., 2016) which are unlikely to be readily leached.

The shales in the Frysjaodden Formation were deposited in the marine environment and any authigenic minerals which were formed at that time are likely to have incorporated fluids with the Eocene seawater composition and deep-sea clays are most susceptible to incorporating seawater (Dasch, 1969). Furthermore, smectite is the only clay mineral which forms in significant amounts in seawater (Griffin et al., 1968) and therefore it is very likely that the deep-sea Frysjaodden Formation contains authigenic smectite in addition to smectite derived from continental weathering, increasing the relative proportion of smectite in this Formation. In addition to diagenetic changes, adsorption may also occur. Samples containing a greater relative proportion of I/S, have a greater cation exchange capacity and are therefore more likely to contain a greater proportion of ions from seawater, increasing the difference between 87Sr/86Sr and εNd in the residue and bulk (Dasch, 1969; Ohr et al., 1991). Adsorption of Nd from seawater was also implicated in a study by von Blanckenburg and Nägler (2001) where leachates of marine sediments had higher εNd than bulk and terrestrial sediments which had had no contact with seawater showed the reverse pattern (lower εNd in leachate compared to bulk). The greater difference between εNd in bulk and residue in the Fardalen (unglaciated) stream suspended sediments compared to Dryadbreen (glaciated, Fig. 8b) is therefore consistent with the greater relative proportion of the Frysjaodden Formation to the Fardalen stream suspended sediments.

Assuming the leached phase is comprised of a mixture of authigenic minerals such as apatite and cations readily leached from clay minerals, then the leachate should have a seawater isotopic composition. Radiogenic Sr in seawater in the past is relatively well constrained given that it has a uniform value across the worlds oceans. Radiogenic neodymium, on the other hand, varies between ocean basins. A study based on fish, provides some constraints on the εNd and 87Sr/86Sr isotopic composition of the Arctic Ocean during the Eocene (Fig. 1, Gleason et al., 2009), with 87Sr/86Sr ranging from 0.7078 to 0.7088 and εNd ranging from -7.5 to -5.5. If we assume that the location of this study and the area of the future Central Basin were connected, then this end-member would be within error of the eastern end-member and therefore could not be distinguished (Fig. 9). This is the most likely reason why the two trends (residue-bulk-leachate and east-west bulk) appear to fall on a common mixing line.
(Fig. 8a). Awwiller (1994) concluded that provenance information based on Nd-Sr isotopes could be obscured by the partial incorporation of Sr and Nd from seawater. Diagenetic alteration has been implicated in shales which give unrealistically old Nd model ages (Arndt and Goldstein, 1987; Awwiller and Mack, 1991; Bock et al., 1994; Cullers et al., 1997; Krogstad et al., 2004). However, although the leached phase is isotopically distinct from bulk it cannot account for the isotopic difference observed between the residual phase isotopic compositions (Fig. 8a). Further, although the potential for diagenetic processes to modify bulk $\varepsilon$Nd and $^{87}\text{Sr}/^{86}\text{Sr}$ values cannot be ruled out, it is a subordinate effect to the primary trend of mixing between distinct lithological end-members (Fig. 9).

5.3 Implications for Nd as a sediment source tracer

The stream suspended sediments observed in this study have highly heterogeneous Nd isotopic compositions with a difference between the two catchments of up to 6.8 epsilon units. Additionally, seasonal variation is observed in the unglaciated Fardalen catchment (1.6±0.4 epsilon units), but is not resolved in the glaciated Dryadbreen catchment (0.6±0.5 epsilon units). A similar magnitude of seasonal variation in $\varepsilon$Nd has previously been reported in the suspended sediments of much larger rivers. A 1.3 seasonal variation has been reported in suspended sediments from the Madeira River (Amazon, Viers et al., 2008) and a 2 epsilon unit range was observed in suspended sediments from two tributaries of the Ganges (Kosi and Narayani, Garçon et al., 2013). The seasonal variation in both of these studies was attributed to the seasonal variation in hydrology which affects how efficient the mixing of tributaries draining different geological units is. The role of hydrology and geology was recently demonstrated at a Canadian glacier where seasonal variations in $\varepsilon$Nd in a geologically heterogeneous catchment were attributed to the changes in subglacial hydrology (distributed to channelised) which altered where erosion occurred (Clinger et al., 2016). In contrast, glacial catchments with more homogenous lithology have little seasonal variation in $\varepsilon$Nd (Clinger et al., 2016; Rickli et al., 2017). From this small dataset (this study, Viers et al., 2008; Garçon et al., 2013; Clinger et al., 2016) it would appear that seasonal variations of $\varepsilon$Nd in suspended sediments are present where rivers drain mixed geology and have a pronounced seasonality to their hydrological cycle.

For the purposes of using Nd in ocean sediment cores as a tracer for past sediment sources it is assumed that the Nd isotopic composition of sediments is constant for broad source regions (e.g. Jeandel et al., 2007), and this will not be affected by seasonal variations. However, seasonal cycles give an insight into weathering and erosion conditions under different hydrological regimes that are an analogue for longer term trends. It is entirely plausible that an intensified or weakened hydrological cycle could change the Nd isotopic composition of sediment export for a given region (Burton and Vance, 2000). Of particular relevance to the Arctic region is the re-organisation of drainage basins as the ice sheets waxed and waned and the attendant changes in magnitude and location of discharge to the ocean (e.g. Teller, 1990; Wickert, 2016). Therefore, it should not necessarily be assumed that the continental regions have had a constant $\varepsilon$Nd export to the oceans over glacial-interglacial timescales. For example, the 5.7 epsilon unit range in $\varepsilon$Nd (which is of similar magnitude to difference between catchments observed in this study) in an Arctic sediment core (PS1533, Tütken et al., 2002) over the last 140 ka was attributed to changes in the relative proportion of sediment derived from two isotopically distinct sources (Svalbard and Siberia) over glacial-interglacial cycles.
Although the broad end-member identification will unlikely be affected, the calculated proportions of each end-member at different points in time would change if the those end-members were not constant over glacial-interglacial cycles.

6 Conclusions

The large variations in $^{87}$Sr/$^{86}$Sr and $\varepsilon$Nd observed in sediments from two small catchments in Svalbard can be explained as a result of two isotopically and geochemically distinct sediment sources mixing during the Meoszoic and subsequently forming the Paleocene-Eocene sedimentary formations which are eroding today (Fig. 10). The two original sources are an eastern sediment source derived from basaltic rocks from Siberia and a western sediment source derived from basement rocks from Greenland. The original geology of the sources controls the initial geochemistry, Sr and Nd isotope values and subsequently determines the type of clay minerals formed during weathering, susceptibility to later diagenesis and particle-size transport effects.

Changes in erosion caused by the glaciation of Dryadbreen has led to material from the upper (younger) formation, which contains a higher proportion of material derived from the western source, being moved lower down in the catchment where it is present in the moraines and sandur plain. In contrast, the lower (older) formation, which contains a higher proportion of material derived from the eastern source, is fully exposed in the unglaciated catchment, having not been covered by sediment from the upper formations of the catchment. This leads to a marked difference in the suspended sediment export from the two catchments and suggests that changes in continental erosion during glacial-interglacial cycles could have a pronounced effect on the Sr and Nd isotopic composition of sediment exported from sedimentary catchments where those sediments have a complex history of multiple sources and sedimentary cycles. Given that the majority of the main rivers in the circum-Arctic region drain shale, the temporal variation of $^{87}$Sr/$^{86}$Sr and $\varepsilon$Nd exported to the ocean from a given region over glacial-interglacial periods, may not have been constant. Further changes on the continents occurring during glacial-interglacial cycles (hydrology, basin configuration) should also be considered as factors affecting $\varepsilon$Nd variation in ocean sediments.

Data availability. All data used in this manuscript is contained in the included tables apart from XRD data files. For data related queries, please contact the corresponding author.

Competing interests. The authors declare no competing interests.

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Figure 1. Overview of the circum-Arctic region. The red cross indicates the position of Svalbard during the Paleogene (Jones et al., 2016). The purple shaded area indicates the western granitic source whilst the green circles indicate potential basaltic source regions in the east. The yellow star indicates the location of the Gleason et al. (2009) study to recover Eocene seawater compositions referred to in the text.
Figure 2. (a) Map of Svalbard indicating the extent of the Paleogene Central Basin. The location of the study area in relation to Longyearbyen is also indicated. (b) The stratigraphy of the Van Mijenfjorden Group adapted from Cui et al. (2011). The catchments are located at the NE side. (c) Topographic map of the sediment sampling locations with the geology of the catchments superimposed. Glaciers and their moraines are shown in blue and orange respectively and contours are displayed at 50 m intervals. The red dashed lines demarcate the catchment boundaries. Dryadbreen is on the left and Fardalen on the right. The coloured circles indicate where samples referred to in Table 1 were collected: R01 and G (green circle); supraglacial stream suspended sediment sample, R03, R04 and D (yellow); Fardalen stream suspended sediment sample and L (blue); Dryadbreen stream suspended sediment sample and O (red); R02 (purple). Other rock and sediment samples referred to in Table 2 were collected at various locations within the two catchments. The red dot in the inset shows the location of the study area (Latitude, 78°08’N; Longitude, 15°30’E) in relation to the rest of Svalbard. Geological information is taken from (Major et al., 2000) and the topographic information is based on GIS data from the Norwegian Polar Institute.
### Figure 3

Plots of major elements in the collected rock samples against the Al₂O₃/SiO₂ ratio compared with core data (Store Norske Well 11-2003) covering the same formations (Schlegel et al., 2013). Sample R03 contains 10% carbonate, accounting for the high Ca concentration (Hindshaw et al., 2016).
**Figure 4.** Herron classification (Herron, 1988) of the sedimentary rock samples collected in this study and from the core analysed by Schlegel et al. (2013). Samples G and R01 were collected in the unglaciated catchment (Fardalen) directly from the Frysjaodden Formation. Samples R03, R02 and R04 were collected in the glaciated catchment (Dryadbreen) but could not be linked to a specific formation in the field due to the heterogeneous nature of moraine material. Based on this classification and the major element chemistry (Fig. 3) we infer that R03 was derived from the Aspelintoppen Formation and R02 close to the Battfjellet/Frysjaodden boundary. Sample R04 is classified as a shale but has a markedly different iron content (Fig. 3) to the shale samples from the Frysjaodden Formation (R01, G). Based on its sampled location on the surface of the glacier we infer that it is a shale from either the Battfjellet or Aspelintoppen Formation (Fig. 2c).
Figure 5. A plot of $^{87}$Sr/$^{86}$Sr vs $\varepsilon$Nd. The grey lines highlight the parallel trends of the bulk and the clay-sized fraction data. Sample R03 is a litharenite whereas the other samples are predominantly shale (Table 1).
Figure 6. XRD patterns from two different samples: one with a high relative illite abundance (D) and one with low relative illite abundance and the clear presence of an expandable clay, likely an illite-smectite (I/S) mixed phase mineral (L).
Figure 7. (a) Part of the orientated clay XRD spectrum (air-dried) for five samples. I/S stands for a mixed layer clay containing smectite and illite. Note the decrease in the I/S peak from R01 to D. (b) The relative abundances of illite and I/S in the clay fraction are inversely correlated. (c,d) There is a positive correlation between the relative abundance of illite and the radiogenic strontium isotope ratio in the clay fraction and an inverse correlation with the radiogenic neodymium isotope ratio.
Figure 8. (a) $\varepsilon$Nd and $^{87}\text{Sr}/^{86}\text{Sr}$ data for bulk suspended sediments and residual suspended sediments after leaching with 5% HCl. In the background is the data from Fig. 5. (b) Rare earth element (REE) abundances of the residue relative to the bulk phase. The greater the depletion in middle REE (MREE), the greater the difference in $\varepsilon$Nd between bulk and residual phases and this difference is most pronounced in the unglaciated catchment (Fardalen) where relative I/S abundances are higher compared to the glaciated catchment (Dryadbreen, Table 2). The residual phases have a lower $\varepsilon$Nd value compared to the bulk.
Figure 9. Plot of $\varepsilon_{Nd}$ vs $^{87}\text{Sr}/^{86}\text{Sr}$. The two points ‘East’ ($^{87}\text{Sr}/^{86}\text{Sr} = 0.70626, \varepsilon_{Nd} = -0.4$) and ‘West’ ($^{87}\text{Sr}/^{86}\text{Sr} = 0.78059, \varepsilon_{Nd} = -37.1$) are averages of the interquartile range of literature data and the error bars indicate the interquartile range (for the ‘East’ source the error in $^{87}\text{Sr}/^{86}\text{Sr}$ is too small to see). ‘East’ data (n=99): Siberian Traps (Lightfoot et al., 1993; Wooden et al., 1993) and Uralides (Spadea and D’Antonio, 2006). ‘West’ data (n=65): Archaean rocks (predominantly gneisses, Jacobsen, 1988; Collerson et al., 1989; Kalsbeek and Frei, 2006; Friend et al., 2009), basal ice particles from GISP 2 and GRIP and granite bedrock samples from GISP 2 (Weis et al., 1997). The ‘r’ values are the Sr/Nd ratio of the ‘East’ source divided by the Sr/Nd ratio of the ‘West’ source. The star indicates the isotopic composition of Eocene seawater (Gleason et al., 2009, see text for details).
Figure 10. Summary of the processes leading to the variation in clay mineralogy, $^{87}\text{Sr}/^{86}\text{Sr}$ and $\varepsilon\text{Nd}$ observed in the two studied catchments. The thickness of the lines gives an indication of the relative contribution of Mesozoic sources to Eocene formations and the Eocene formations to suspended sediment export from the two catchments. For example, the Frysjaodden Formation receives a greater proportion of sediment from the eastern source as compared to the western source. (a) The contribution of Mesozoic sediment sources to Eocene formations is determined by the depositional location (far-shore vs near-shore), particle size and susceptibility to authigenic phase formation. (b) The contribution of the Paleogene formations to the present-day suspended sediment load is determined by the present-day exposure of the formations in each catchment (Fig. 2) which is determined by the recent erosional history of the catchments.
Table 1. Major and selected trace metal concentrations, $^{87}$Sr/$^{86}$Sr and $\varepsilon$Nd values for solid samples. Major elements (in wt%) were measured by ICP-OES and trace elements (mg/kg) were measured by ICP-MS. REE data are presented in Table A2.

| Name and description | SSC $^1$ mg/L | SiO$_2$ $^2$ wt% | Al$_2$O$_3$ wt% | Fe$_2$O$_3$ wt% | TiO$_2$ wt% | MgO wt% | CaO wt% | Na$_2$O wt% | K$_2$O wt% | LOI wt% | Mn mg/kg | Ba mg/kg | Sr mg/kg | Rb mg/kg | Nd mg/kg | Sm mg/kg | $^{87}$Sr/$^{86}$Sr 2SD $^4$ | $\varepsilon$Nd 2SD $^4$ | 2SD $^4$ |
|----------------------|----------------|------------------|-----------------|----------------|-------------|---------|---------|------------|------------|--------|---------|---------|--------|--------|--------|--------|----------|-------------|--------|
| Sedimentary rock samples | | | | | | | | | | | | | | | | | | | |
| R01 = shale (pieces) | 65.2 | 15.7 | 7.07 | 0.79 | 1.36 | 0.36 | 1.13 | 2.38 | 0.21 | 79 | 497 | 438 | 98 | 103 | 32 | 6 | 0.72490 | 17 | -11.9 | 0.0 |
| R04 = shale (rock) | 58.3 | 18.5 | 5.50 | 0.88 | 1.34 | 0.17 | 0.74 | 3.16 | 0.09 | 133 | 190 | 542 | 107 | 147 | 39 | 7 | 0.74368 | 32 | -19.8 | 0.7 |
| Sediment sample | | | | | | | | | | | | | | | | | | | |
| D = sediment (surface of glacier) | 54.2 | 18.3 | 6.71 | 0.75 | 1.47 | 0.53 | 0.93 | 3.28 | 0.11 | 13.8 | 735 | 706 | 99 | 135 | 36 | 6 | 0.75243 | 36 | -23.3 | 0.2 |
| Stream sediment samples | | | | | | | | | | | | | | | | | | | |
| L = stream sediment (Fardalen) | 62.3 | 15.3 | 7.68 | 0.74 | 1.35 | 0.32 | 0.99 | 2.42 | 0.21 | 8.9 | 613 | 425 | 100 | 103 | 32 | 6 | 0.72596 | 28 | -12.1 | 1.0 |
| Stream suspended sediments (Bulk) | | | | | | | | | | | | | | | | | | | |
| 20120801SG | 47.7 | 16.4 | 5.72 | 0.74 | 1.35 | 0.34 | 0.90 | 3.00 | 0.14 | 19.1 | 482 | 687 | 98 | 123 | 34 | 6 | 0.74300 | 11 | -20.6 | 0.5 |
| 20120617D | 62 | 19.1 | 8.82 | 0.60 | 1.68 | 0.47 | 4.04 | 3.03 | 4.13 | 101 | 650 | 427 | 93 | 132 | 33 | 6 | 0.74766 | 27 | -10.0 | 0.2 |
| Stream suspended sediments (Acid-treated) | | | | | | | | | | | | | | | | | | | |
| 20120617D(HCl) | 63.3 | 18.2 | 6.75 | 0.85 | 1.77 | 0.49 | 0.89 | 3.68 | 0.21 | 10.9 | 660 | 756 | 114 | 162 | 37 | 7 | 0.74531 | 22 | -20.2 | 0.5 |

$^1$ Suspended sediment concentration
$^2$ SiO$_2$ concentrations calculated assuming 100% recovery. For a comparison of XRF and ICP-OES concentrations see Table A1
$^3$ Samples names are YYYYMMDD and the subsequent letters are D=Dryadbreen (glaciated), F= Fardalen (unglaciated) and SG = supraglacial
$^4$ n=3
Table 2. Relative proportions of clay minerals as determined from XRD patterns for the solid samples collected in the two catchments. Samples are ordered from highest to lowest relative illite abundance.

| Sample | Description                  | Catchment | Illite (%) | I/S (%) | Kaolinite (%) | Chlorite (%) |
|--------|------------------------------|-----------|------------|---------|---------------|--------------|
| I*     | fossil-rich rock             | Dr        | 82         | 0       | 9             | 9            |
| D      | sediment (surface of glacier)| Dr        | 72         | 12      | 7             | 9            |
| R03*   | litharenite                  | Dr        | 67         | 14      | 8             | 10           |
| R04*   | shale (rock)                 | Dr        | 59         | 26      | 7             | 7            |
| A      | sandur sediment              | Dr        | 52         | 28      | 8             | 12           |
| F      | moraine sediment             | Dr        | 55         | 28      | 8             | 9            |
| O      | stream sediment              | Dr        | 51         | 32      | 8             | 9            |
| R02*   | wacke                        | Dr        | 48         | 33      | 5             | 14           |
| H      | soil                         | F         | 49         | 35      | 9             | 7            |
| M      | stream bank sediment         | F         | 51         | 37      | 6             | 7            |
| C      | moraine sediment             | Dr        | 44         | 37      | 9             | 10           |
| E*     | shale rock                   | Dr        | 43         | 40      | 8             | 9            |
| R01*   | shale (pieces)               | F         | 46         | 40      | 7             | 7            |
| N      | stream bedload               | F         | 46         | 41      | 6             | 7            |
| L      | stream sediment              | F         | 46         | 43      | 5             | 6            |
| K      | stream sediment              | F         | 45         | 43      | 6             | 6            |
| B*     | shale (pieces)               | Dr        | 38         | 49      | 6             | 7            |
| G*     | shale (pieces)               | F         | 36         | 50      | 7             | 7            |

* Sedimentary (often frost-shattered) rock. All other samples are modern sediments.

1 Catchment where sample was collected. F=Fardalen, Dr= Dryadbreen.

2 I/S is an illite-smectite mixed phase mineral.
Table A1. Comparison of element concentrations collected by XRF (only available for four samples) and ICP-OES. Data from ICP-OES is used in the paper.

| Name and description | SiO$_2$ | Al$_2$O$_3$ | Fe$_2$O$_3$ | TiO$_2$ | MgO | CaO | Na$_2$O | K$_2$O | P$_2$O$_5$ | LOI | Sum | Mn | Ba | Sr | Rb | Nd | Sm |
|----------------------|---------|-------------|-------------|---------|-----|-----|---------|-------|-----------|-----|-----|----|----|----|----|----|----|
| Data collected by XRF|         |             |             |         |     |     |         |       |           |     |     |    |    |    |    |    |    |
| R01                  | 63.2    | 16.3        | 7.1         | 0.8     | 1.0 | 2.6 | 0.3     | 7.5   | 100.5     | 488 | 403 | 97.4 | 102 | 35 | <10 |
| R02                  | 65.4    | 13.0        | 7.9         | 0.6     | 0.7 | 2.2 | 0.3     | 7.2   | 99.4      | 1038| 357 | 81.9 | 88.8 | 27 | <10 |
| R03                  | 70.9    | 8.9         | 3.2         | 0.6     | 0.6 | 1.6 | 0.1     | 7.0   | 100.5     | 472 | 356 | 142  | 55.1 | 23 | <10 |
| R04                  | 57.1    | 19.1        | 3.5         | 0.9     | 1.3 | 0.2 | 0.6     | 1.29  | 99.3      | 178 | 511 | 116  | 150 | 42 | <10 |
| Data collected by ICP-OES$^1$ |         |             |             |         |     |     |         |       |           |     |     |    |    |    |    |    |    |
| R01                  | 63.2    | 15.7        | 7.1         | 0.8     | 1.1 | 2.4 | 0.2     | 7.9   | 100.0     | 497 | 438 | 98  | 103 | 32 | 6  |
| R02                  | 64.7    | 12.6        | 8.0         | 0.6     | 0.8 | 2.1 | 0.3     | 8.9   | 100.0     | 1056| 371 | 83  | 89  | 26 | 5  |
| R03                  | 70.0    | 8.6         | 3.3         | 0.6     | 1.1 | 1.5 | 0.1     | 7.7   | 100.0     | 480 | 366 | 139 | 54  | 25 | 5  |
| R04                  | 58.3    | 18.5        | 3.5         | 0.9     | 1.3 | 0.2 | 0.7     | 3.2   | 133       | 100.0| 542 | 107 | 147 | 39 | 7  |

$^1$ SiO$_2$ concentrations calculated by assuming 100% recovery.
Table A2. Rare earth element concentrations in mg/kg\(^1\).

| Sample\(^2\) | La  | Ce  | Pr  | Nd  | Sm  | Eu  | Gd  | Tb  | Dy  | Ho  | Er  | Tm  | Yb  | Lu  |
|-------------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| **Sedimentary rock samples** |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| R01         | 35.7| 71.2| 8.4 | 32.4| 6.0 | 1.2 | 4.5 | 0.7 | 4.0 | 0.7 | 2.0 | 0.3 | 1.9 | 0.3 |
| G           | 35.7| 70.4| 8.3 | 32.3| 6.0 | 1.3 | 4.6 | 0.7 | 4.2 | 0.8 | 2.3 | 0.3 | 2.2 | 0.3 |
| R04         | 49.1| 95.4| 10.7| 39.3| 6.6 | 1.3 | 4.0 | 0.7 | 3.5 | 0.7 | 1.9 | 0.3 | 1.8 | 0.2 |
| R02         | 28.5| 55.5| 6.6 | 25.7| 4.8 | 1.1 | 3.8 | 0.6 | 3.2 | 0.6 | 1.7 | 0.2 | 1.5 | 0.2 |
| R03         | 30.5| 60.3| 6.8 | 25.4| 4.5 | 0.9 | 3.1 | 0.4 | 2.2 | 0.4 | 1.0 | 0.1 | 0.9 | 0.1 |
| **Sediment sample** |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| D           | 43.9| 88.5| 9.9 | 36.5| 6.5 | 1.2 | 4.3 | 0.6 | 3.3 | 0.6 | 1.6 | 0.2 | 1.4 | 0.2 |
| **Stream sediment samples** |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| L           | 37.1| 72.8| 8.3 | 31.6| 5.8 | 1.2 | 4.4 | 0.7 | 4.0 | 0.8 | 2.2 | 0.3 | 2.0 | 0.3 |
| O           | 37.9| 76.3| 8.7 | 33.4| 5.9 | 1.1 | 4.2 | 0.6 | 3.1 | 0.5 | 1.4 | 0.2 | 1.3 | 0.2 |
| **<2 µm fraction fraction of bulk rock and sediment samples** |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| R01-clay    | 37.0| 72.7| 9.4 | 38.1| 8.0 | 1.8 | 7.1 | 1.0 | 5.8 | 1.0 | 2.8 | 0.4 | 2.5 | 0.3 |
| R04-clay    | 38.9| 78.0| 8.6 | 31.9| 5.8 | 1.0 | 3.7 | 0.6 | 3.5 | 0.7 | 2.0 | 0.3 | 2.0 | 0.3 |
| G-clay      | 32.4| 64.3| 8.0 | 32.6| 6.5 | 1.5 | 5.7 | 0.9 | 4.8 | 0.9 | 2.4 | 0.3 | 2.2 | 0.3 |
| D-clay      | 31.5| 64.0| 7.0 | 25.9| 4.8 | 1.0 | 3.8 | 0.6 | 3.4 | 0.7 | 1.8 | 0.3 | 1.7 | 0.2 |
| L-clay      | 30.5| 60.4| 7.1 | 29.0| 6.0 | 1.4 | 5.4 | 0.9 | 4.9 | 1.0 | 2.7 | 0.4 | 2.5 | 0.4 |
| **Suspended sediments (Bulk)\(^2\)** |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| 20120801SG  | 41.8| 82.8| 9.4 | 34.4| 6.2 | 1.2 | 4.2 | 0.6 | 3.3 | 0.6 | 1.6 | 0.2 | 1.5 | 0.2 |
| 20120617D   | 40.3| 79.3| 9.1 | 34.1| 6.2 | 1.2 | 4.3 | 0.6 | 3.5 | 0.6 | 1.6 | 0.2 | 1.5 | 0.2 |
| 20120618F   | 40.9| 84.5| 10.0| 40.3| 8.2 | 1.8 | 7.1 | 1.0 | 5.4 | 1.0 | 2.5 | 0.3 | 2.0 | 0.3 |
| 20120726F   | 44.3| 89.6| 10.4| 41.8| 8.6 | 1.9 | 7.7 | 1.2 | 6.3 | 1.1 | 2.9 | 0.4 | 2.4 | 0.3 |
| 20120729D   | 44.6| 88.9| 9.9 | 37.3| 6.6 | 1.3 | 4.7 | 0.7 | 3.7 | 0.7 | 1.8 | 0.3 | 1.7 | 0.3 |
| **Suspended sediments (Acid-treated)** |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| 20120617D(HCl) | 38.1| 72.8| 7.9 | 28.6| 4.5 | 0.7 | 2.3 | 0.4 | 2.2 | 0.4 | 1.3 | 0.2 | 1.3 | 0.2 |
| 20120618F(HCl) | 36.8| 71.0| 7.8 | 27.8| 4.2 | 0.9 | 2.5 | 0.4 | 2.7 | 0.5 | 1.5 | 0.2 | 1.7 | 0.2 |
| 20120726F(HCl) | 42.2| 81.2| 8.8 | 30.7| 4.7 | 0.9 | 2.7 | 0.5 | 3.0 | 0.6 | 1.7 | 0.3 | 1.8 | 0.2 |
| 20120729D(HCl) | 42.1| 80.4| 8.8 | 31.7| 4.7 | 0.9 | 2.5 | 0.4 | 2.4 | 0.5 | 1.4 | 0.2 | 1.4 | 0.2 |

\(^1\) The hotplate digestion method used does not digest zircons and therefore the high rare earth element (HREE) concentrations may be lower than the true total. Nd and Sm are fully digested by the hotplate method (Rickli et al., 2013).

\(^2\) Please refer to Table 1 for sample descriptions.
Table A3. Measurements of four USGS standards by ICP-OES compared to certificate values.

| Standard | SiO$_2$ wt% | Al$_2$O$_3$ wt% | Fe$_2$O$_3$ wt% | TiO$_2$ wt% | MgO wt% | CaO wt% | Na$_2$O wt% | K$_2$O wt% | P$_2$O$_5$ wt% | LOI wt% |
|----------|-------------|-----------------|-----------------|-------------|---------|---------|-------------|---------|--------------|---------|
| SGR-1b   | 30.7$^1$    | 6.3             | 3.3             | 0.3         | 4.4     | 8.1     | 3.1         | 1.6     | 0.1          | 42.2    |
| Mean (n=8) | 2SD         | 0.4             | 0.2             | 0.0         | 0.3     | 0.3     | 0.1         | 0.1     | 0.0          | 0.5     |
| Ref. value |             | 28.2            | 6.5             | 3.0         | 0.3     | 4.4     | 8.4         | 3.0     | 1.7          | 0.3     |
| G-2      | 69.0$^1$    | 14.9            | 3.0             | 0.6         | 0.7     | 2.0     | 4.2         | 4.2     | 0.1          | 1.4     |
| Mean (n=2) | 2SD         | 0.3             | 0.0             | 0.0         | 0.0     | 0.0     | 0.1         | 0.1     | 0.0          | 0.7     |
| Ref. value |             | 69.1            | 15.4            | 2.7         | 0.5     | 0.8     | 2.0         | 4.1     | 4.5          | 0.1     |
| BCR-2    | 51.5$^1$    | 13.1            | 16.5            | 2.6         | 3.5     | 7.0     | 3.3         | 1.9     | 0.1          | 0.5     |
| Mean (n=4) | 2SD         | 0.3             | 1.1             | 0.1         | 0.1     | 0.1     | 0.1         | 0.1     | 0.1          | 0.0     |
| Ref. value |             | 54.1            | 13.5            | 13.8        | 2.3     | 3.6     | 7.1         | 3.2     | 1.8          | 0.4     |
| AGV-2    | 58.5$^1$    | 16.4            | 7.6             | 1.2         | 1.7     | 5.2     | 4.3         | 2.9     | 0.2          | 2.1     |
| Mean (n=2) | 2SD         | 0.4             | 0.2             | 0.1         | 0.0     | 0.4     | 0.2         | 0.2     | 0.2          | 0.1     |
| Ref. value |             | 59.3            | 16.9            | 6.7         | 1.1     | 1.8     | 5.2         | 4.2     | 2.9          | 0.5     |

$^1$ SiO$_2$ concentrations calculated assuming 100% recovery.

For a comparison of XRF and ICP-OES concentrations see Table A1
Table A4. Measurements of four USGS standards by ICP-MS compared to literature value. Concentrations are in mg/kg\(^1\).

| Standard | Mn  | Rb  | Sr  | Ba  | La  | Ce  | Pr  | Nd  | Sm  | Eu  | Gd  | Tb  | Dy  | Ho  | Er  | Tm  | Yb  | Lu  |
|----------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| **SCo-1**<br>Mean (n=12) | 431 | 115 | 160 | 627 | 30.2 | 59 | 7.0 | 27 | 5.2 | 1.07 | 4.2 | 0.6 | 3.5 | 0.69 | 1.9 | 0.28 | 1.79 | 0.26 |
| 2SD | 69 | 19 | 20 | 72 | 4.4 | 7 | 0.5 | 1 | 0.2 | 0.08 | 0.2 | 0 | 0.2 | 0.06 | 0.1 | 0.03 | 0.14 | 0.03 |
| Govindaraju (1994) | 410 | 112 | 174 | 570 | 29.5 | 62 | 6.6 | 26 | 5.3 | 1.19 | 4.6 | 0.7 | 4.2 | 0.97 | 2.5 | 0.42 | 2.27 | 0.34 |
| **BIR-1**<br>Mean (n=6) | 1419 | 0.22 | 105 | 7 | 0.69 | 1.89 | 0.36 | 2.4 | 1.1 | 0.54 | 1.98 | 0.37 | 2.6 | 0.58 | 1.7 | 0.26 | 1.72 | 0.26 |
| 2SD | 155 | 0.09 | 10 | 1 | 0.09 | 0.19 | 0.02 | 0.0 | 0.0 | 0.04 | 0.05 | 0.01 | 0.2 | 0.03 | 0.1 | 0.01 | 0.09 | 0.02 |
| Govindaraju (1994) | 1324 | 0.25 | 108 | 7 | 0.62 | 1.95 | 0.38 | 2.5 | 1.1 | 0.54 | 1.85 | 0.36 | 2.5 | 0.57 | 1.7 | 0.26 | 1.65 | 0.26 |
| **BHVO-2**<br>Mean (n=7) | 1358 | 9.59 | 372 | 133 | 15.1 | 37.0 | 5.16 | 24.1 | 6.05 | 2.09 | 6.41 | 0.95 | 5.34 | 0.99 | 2.56 | 0.34 | 2.05 | 0.28 |
| 2SD | 89 | 0.98 | 35 | 14 | 1.7 | 4.0 | 0.39 | 1.2 | 0.26 | 0.06 | 0.22 | 0.03 | 0.24 | 0.04 | 0.09 | 0.01 | 0.09 | 0.01 |
| Raczek et al. (2001) | 1317 | 9.08 | 396 | 131 | 15.2 | 37.5 | 5.29 | 24.5 | 6.07 | 2.07 | 6.24 | 0.94 | 5.32 | 0.97 | 2.54 | 0.34 | 2.00 | 0.27 |
| **BCR-2**<br>Mean (n=9) | 1523 | 47.3 | 312 | 657 | 23.7 | 50.2 | 6.39 | 27.4 | 6.34 | 1.88 | 6.62 | 1.04 | 6.29 | 1.27 | 3.61 | 0.53 | 3.37 | 0.50 |
| 2SD | 142 | 5.8 | 31 | 44 | 2.3 | 4.1 | 0.38 | 0.8 | 0.14 | 0.05 | 0.08 | 0.02 | 0.22 | 0.04 | 0.15 | 0.02 | 0.14 | 0.02 |
| Raczek et al. (2001) | 1471 | 46.9 | 340 | 677 | 24.9 | 52.9 | 6.57 | 28.7 | 6.57 | 1.96 | 6.75 | 1.07 | 6.42 | 1.30 | 3.66 | 0.56 | 3.38 | 0.52 |
| **AGV-2**<br>Mean (n=6) | 785 | 70.3 | 616 | 1141 | 37.7 | 68.3 | 7.94 | 30.2 | 5.48 | 1.43 | 4.29 | 0.64 | 3.49 | 0.67 | 1.85 | 0.27 | 1.68 | 0.25 |
| 2SD | 75 | 8.2 | 45 | 121 | 4.1 | 6.8 | 0.54 | 1.4 | 0.22 | 0.05 | 0.23 | 0.01 | 0.15 | 0.04 | 0.10 | 0.01 | 0.08 | 0.01 |
| Raczek et al. (2001) | 774 | 66.3 | 661 | 1130 | 37.9 | 68.6 | 7.68 | 30.5 | 5.49 | 1.53 | 4.52 | 0.64 | 3.47 | 0.65 | 1.81 | 0.26 | 1.62 | 0.25 |

\(^1\)The hotplate digestion method used does not digest zircons and this accounts for the low recovery of the HREE in the sedimentary standard SCo-1.