**Zeitschrift:** Schweizerische mineralogische und petrographische Mitteilungen =

Bulletin suisse de minéralogie et pétrographie

**Band:** 83 (2003)

Heft: 3

**Artikel:** Strontium isotope systematics in two glaciated crystalline catchments:

Rhone and Oberaar Glaciers (Swiss Alps)

Autor: Arn, Kaspar / Hosein, Rachel / Föllmi, Karl B.

**DOI:** https://doi.org/10.5169/seals-63149

# Nutzungsbedingungen

Die ETH-Bibliothek ist die Anbieterin der digitalisierten Zeitschriften auf E-Periodica. Sie besitzt keine Urheberrechte an den Zeitschriften und ist nicht verantwortlich für deren Inhalte. Die Rechte liegen in der Regel bei den Herausgebern beziehungsweise den externen Rechteinhabern. Das Veröffentlichen von Bildern in Print- und Online-Publikationen sowie auf Social Media-Kanälen oder Webseiten ist nur mit vorheriger Genehmigung der Rechteinhaber erlaubt. Mehr erfahren

## **Conditions d'utilisation**

L'ETH Library est le fournisseur des revues numérisées. Elle ne détient aucun droit d'auteur sur les revues et n'est pas responsable de leur contenu. En règle générale, les droits sont détenus par les éditeurs ou les détenteurs de droits externes. La reproduction d'images dans des publications imprimées ou en ligne ainsi que sur des canaux de médias sociaux ou des sites web n'est autorisée qu'avec l'accord préalable des détenteurs des droits. En savoir plus

# Terms of use

The ETH Library is the provider of the digitised journals. It does not own any copyrights to the journals and is not responsible for their content. The rights usually lie with the publishers or the external rights holders. Publishing images in print and online publications, as well as on social media channels or websites, is only permitted with the prior consent of the rights holders. Find out more

**Download PDF:** 17.08.2025

ETH-Bibliothek Zürich, E-Periodica, https://www.e-periodica.ch

# Strontium isotope systematics in two glaciated crystalline catchments: Rhone and Oberaar Glaciers (Swiss Alps)

Kaspar Arn<sup>1</sup>, Rachel Hosein<sup>1</sup>, Karl B. Föllmi<sup>1</sup>, Philipp Steinmann<sup>1</sup>, Dominique Aubert<sup>2,3</sup> and Jan Kramers<sup>4</sup>

#### Abstract

We studied the strontium (Sr) isotope composition of runoff and particulate material in two glaciated catchments, the Oberaar (OA) and the Rhone (RH) in the Swiss Alps. Both areas are contained within the crystalline rocks of the Aar Massif, but a zone of highly deformed Variscan basement gneisses and schists that may contain up to 9 wt% calcite is present in the Oberaar catchment.

We analysed meltwaters and precipitation as well as bulk suspended sediment and local bedrock. Ca/Na ratios of 5.0–13.3 and 1.8–2.4 in Oberaar and Rhone meltwaters, respectively, indicate that meltwaters in both catchments are enriched in Ca relative to Na, compared to suspended sediments.

<sup>87</sup>Sr/<sup>86</sup>Sr ratios of Rhone meltwaters (0.7251–0.7258) are lower than those of Rhone bulk suspended sediment (0.7279–0.7306). The Rhone suspended sediment composition is interpreted as a mixture of Grimsel Granodiorite (0.7106–0.7161) and Central Aar Granite (0.7449). We explain the lower meltwater Sr ratio by the preferential dissolution of disseminated calcite, which has a relatively low <sup>87</sup>Sr/<sup>86</sup>Sr ratio (estimated 0.720–0.730). This interpretation is supported by the enrichment of Ca in the meltwater compared to the suspended sediment. Early and nonstoichiometric cation release exerts another impact on the meltwater signal.

Oberaar meltwaters have a higher  $^{87}$ Sr/ $^{86}$ Sr isotopic ratio (0.7137–0.7174) than the Oberaar suspended sediment (0.7130–0.7148) which reflects the preferential dissolution of the calcite contained within the Variscan gneisses (estimated 0.7160  $\pm$  0.001).

The <sup>87</sup>Sr/<sup>86</sup>Sr ratio measured in rain is 0.7104 (n = 4). An influence of atmospheric precipitation on Rhone meltwater <sup>87</sup>Sr/<sup>86</sup>Sr ratio is not excluded. The high Sr isotopic signature of Oberaar meltwaters compared to that of the corresponding suspended sediment and also to the low Sr isotopic ratio of rainwater, however, suggests that the atmospheric contribution is not visible in the Oberaar meltwaters.

The results discussed here suggest that the presence of disseminated calcite within the crystalline rocks of the Aar massif exerts a major impact not only on the meltwater major ion composition but also on its Sr isotope systematics.

Keywords: Subglacial chemical weathering, Swiss Alps, glacial meltwaters, suspended sediment, disseminated calcite, strontium isotopes.

#### Introduction

Fluxes and isotopic ratios of Sr have been analysed in stream waters in order to trace mineral weathering reactions and the weathering rates of individual silicate minerals and to compare them with the isotopic characteristics of catchment bedrocks (e.g., Blum et al., 1993; Blum and Erel, 1995; Taylor and Blum, 1995; Clow et al., 1997). The chemical weathering of Sr from minerals is not necessarily stoichiometric, as it is dominated by the rapid dissolution of Sr-rich, or highly reactive minerals or mineral inclusions, such as dis-

seminated calcite (Clow et al., 1997; Blum et al., 1998; Jacobson and Blum, 2000; Jacobson et al., 2002) or apatite (Aubert et al., 2001). Biotite releases considerable amounts of highly radiogenic Sr during the initial stages of weathering and initial weathering rates may exceed those of feldspar (Blum and Erel, 1995, 1997; Arn, 2002). In laboratory dissolution experiments, Brantley et al. (1998) show that early Sr release from feldspars is nonstoichiometric and not isotopically identical to the starting material. The nonstoichiometric release is attributed to secondary phases present in minute proportions, and/or to leaching of cations

<sup>&</sup>lt;sup>1</sup> Institut de Géologie, Université de Neuchâtel, Rue Emile-Argand 11, CH-2007 Neuchâtel, Switzerland. <a href="mailto:kaspar1@bluewin.ch"><a href="mailto:kaspar1@bluewin.ch"><a href="mailto:karl.foellmi@unine.ch">karl.foellmi@unine.ch</a>

 <sup>&</sup>lt;sup>2</sup> EOST, ULP/CNRS, Centre de Géochimie de la Surface, UMR 7517, 1 rue Blessig, 67084 Strasbourg Cedex, France.
<sup>3</sup> present address: Laboratoire des Mecanismes de Transfert en Géologie, Equipe "Eau Sol Erosion", Université Toulouse, 38 rue des 36 Ponts, 31400 Toulouse, France.

<sup>&</sup>lt;sup>4</sup> Gruppe Isotopengeologie, Geologisches Institut, Universität Bern, Erlachstrasse 9a, CH-3012 Bern, Switzerland.

from damaged sites. They propose that glacially abraded feldspar particles will exhibit this initial nonstoichiometric release.

Detailed studies of meltwater chemistry have been performed in glaciated catchments draining a variety of subglacial rock (e.g., Fairchild et al., 1994, 1999a, 1999b; Anderson et al., 2000; Sharp et al., 2002; Tranter et al., 2002). These studies show that carbonate weathering dominates bulk meltwater chemistry, even in crystalline catchments (Clow et al., 1997; Blum et al., 1998; White et al., 1999; Anderson et al., 2000; Jacobson and Blum, 2000; Jacobson et al., 2002; Tranter et al., 2002).

Strontium fluxes and isotopic ratios in stream waters also reflect the weathering intensity. Elevated Sr isotopic budgets for Himalayan rivers reflect the increased erosion rates in this mountain range (Galy et al., 1999; Galy and France-Lanord, 2001). They also indicate silicate weathering rates and the relative importance of carbonate and silicate weathering in different Himalayan settings (Harris, 1995; English et al., 2000).

The Sr isotopic ratios in continental riverine runoff are used as a proxy for the rate of chemical weathering on continents. They in turn influence the Sr isotopic signature of deposited marine carbonates, together with other Sr sources such as submarine hydrothermal sources and deep-sea sediment pore waters (Blum and Erel, 1995; Palmer and Edmond, 1989; Hodell et al., 1990; Hodell and Woodruff, 1994). The sharp rise in radiogenic Sr in marine carbonates in the late Eocene–Holocene is influenced by the intensified weathering of continental source rocks associated with the uplift of the Tibetan plateau and Himalayan mountain range (e.g. Raymo et al., 1988; Raymo and Ruddiman, 1992; Edmond, 1992).

Subglacial and proglacial weathering processes increase the chemical weathering rates of some primary minerals (Sharp et al., 1995; Hallet et al., 1996; Anderson et al., 1997; Tranter et al., 2002; Hosein et al., 2004). Glacial weathering also enhances the release of Sr by: (1) grinding and abrasion of rock, generating ultrafine particles (< 63 µm) with increased total reactive surface area;

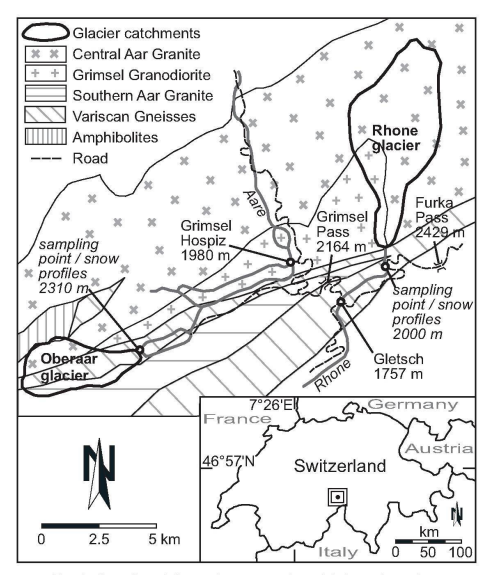


Fig. 1 Location of the catchment together with the main geology.

(2) selective weathering of minerals with relatively high Sr isotopic ratios; (3) providing a continuous supply of these minerals and thereby increased flushing of glacial sediments and soils; (4) producing mechanically damaged surfaces which enhance nonstoichiometric dissolution; and (5) producing large volumes of dilute meltwaters.

We studied subglacial chemical weathering processes and the Sr isotope composition of bedrock, runoff and particulate material in the glaciated catchments of the Oberaar (OA) and Rhone (RH) Glaciers (Swiss Alps), which are both located in the crystalline rocks of the Aar massif. Our goal was to estimate the influence of preferential dissolution of major and trace minerals on the bulk meltwater signal. We suggest that lithological differences between the two areas exert a major influence on bulk meltwater chemistry, meltwater Sr isotope ratios, and the mineralogy of the particulate flux. We show that the dissolution of disseminated calcite and the nonstoichiometric release of Sr from silicates are important sources of Sr and that they strongly influence the Sr isotopic ratios in runoff. A novel aspect of this work is that the glaciated terrains are purely crystalline. Such terrains have not previously been studied in the same detail as mixed crystalline-sedimentary or sedimentary terrains.

#### Field areas

The Rhone and Oberaar Glaciers are located in the central Swiss Alps, to the east and west of the Grimselpass (2100 m above sea level), respectively (Fig. 1 and Table 1). Climatic conditions are therefore comparable (Schwab et al., 2001). Both glaciers are presently retreating. Both areas are contained within the crystalline rocks of the Aar Massif (Stalder, 1964; Abrecht, 1994; Fig. 1), and the lithologies are comparable, except for the presence of a zone of highly deformed Variscan basement gneisses and schists in the Oberaar area (Oberhänsli et al., 1988).

The Central Aar Granite and the Southern Aar Granite contain quartz, K-feldspar (microcline), plagioclase (oligoclase), and micas (mainly biotite). The Grimsel Granodiorite is richer in mica (biotite) and contains several meter- to decameter-sized basic lenses in the higher parts of the Oberaar catchment.

The Variscan Gneisses consist of recrystallized quartz, plagioclase (20–50 vol%; albite), and 8–20% biotite, and may contain some K-feldspar (microcline). They are heterogeneous on a small scale and contain up to 9 wt% calcite (compare also Stalder (1964), who found up to 5 vol% calcite). Sulfide mineralization is evident in these rocks, as there are many rusty stains on weathered surfaces and the SO<sub>4</sub> flux is 3.8 to 4.5 times higher in the Oberaar catchment compared to the Rhone (Hosein et al., 2004).

# Sampling and analytical methods

Weekly (summer) to monthly (winter) bulk meltwater samples (1000 ml) were taken at both glaciers between 6 July 1999 and 1 October 2001 and a representative collection was used for Sr-analyses. The samples were immediately filtered in the field using  $0.45~\mu m$  cellulose nitrate filters, acidi-

*Table 1* Comparison of the field areas.

|                             | Oberaar              |                      | Rhone             |                      |  |  |
|-----------------------------|----------------------|----------------------|-------------------|----------------------|--|--|
| Geogr. Position             | 46° 32' N 8°         | ° 14' E              | 46° 35′ N 8° 3    | 23' E                |  |  |
| Surface of catchment        | $11.2~\mathrm{km}^2$ |                      | $23 \text{ km}^2$ |                      |  |  |
| Altitude                    | 2310 - 3631          | 2310 – 3631 m a.s.l. |                   | 1750 – 3630 m a.s.l. |  |  |
| Glacier cover today (%)     | 57%                  |                      | 73%               |                      |  |  |
| Annual mean temperature †   | -1 °C §              |                      | + 1.2 °C §        |                      |  |  |
| Annual mean precipitation † | 2100 mm              |                      | 2200 mm           |                      |  |  |
| Geology of catchments:      | total*               | ice covered**        | total*            | ice covered¥         |  |  |
| Central Aar Granite         | 38%                  | 26%                  | 90%               | $76 \pm 1\%$         |  |  |
| Grimsel Granodiorite        | 7%                   | 3%                   | 10%               | $24 \pm 1\%$         |  |  |
| Variscan gneisses           | 42%                  | 60%                  | -                 | -                    |  |  |
| Ultramafic Inclusions       | 1%                   | -                    | 7 <u>—</u>        | _                    |  |  |
| Southern Aar Granite        | 12%                  | 11%                  | -                 | _                    |  |  |

<sup>†</sup> Schwab (2001)

<sup>§</sup> within proglacial area; 2300 m at Oberaar, 1757 m at Rhone

<sup>\*</sup> from geological maps of areas, see text for references

<sup>#</sup> Ultramafic inclusions neglected, see text for explanations

<sup>¥</sup> calculated in this work, see text for explanations

Table 2 Results of the different bedrock and suspended sediment analyses.

| 100        |             |             |                            |                                    |             | •               |                            |              |              |       |
|------------|-------------|-------------|----------------------------|------------------------------------|-------------|-----------------|----------------------------|--------------|--------------|-------|
| Sample     |             |             | Sr<br>µmol g <sup>-1</sup> | <sup>87</sup> Sr/ <sup>86</sup> Sr | 2σ<br>error | Calcite*<br>wt% | Ca<br>µmol g <sup>-1</sup> | Ca/(1000×Sr) | Na<br>µmol g | Ca/Na |
| Bedrock    |             |             |                            |                                    |             |                 |                            |              |              |       |
| Central A  | ar Granit   | e           |                            |                                    |             |                 |                            |              |              |       |
| R17        | TD          | SB          | 0.422                      | 0.7449                             | ± 0.000026  | 0.11            | 52                         | 0.122        | 526          | 0.098 |
| CAGr       |             | ¥           | 1.050                      |                                    | ± 0.000109  |                 | 159                        | 0.151        | 660          | 0.240 |
| Aar10      |             | #           | 0.902                      |                                    | ± 0.000028  |                 |                            |              |              |       |
|            | Bio         | #           | 1.142                      |                                    | ± 0.000167  |                 |                            |              |              |       |
|            | Kfs         | #           | 0.711                      |                                    | ± 0.000058  |                 |                            |              |              |       |
| Aar12      | 7           | #           | 0.799                      |                                    | ± 0.000032  |                 |                            |              |              |       |
|            | Bio         | #           | 0.652                      |                                    | ± 0.000043  |                 |                            |              |              |       |
| Aar17      |             | #<br>#<br># | 0.811                      |                                    | ± 0.000056  |                 |                            |              |              |       |
|            | Bio         | #           | 0.148                      |                                    | ± 0.000098  |                 |                            |              |              |       |
|            | Plagio      | #           | 0.822                      |                                    | ± 0.000079  |                 |                            |              |              |       |
| Grimsel C  | 12-00       |             |                            |                                    |             |                 |                            |              |              |       |
| R6         | TD          | SB          | 1.941                      | 0.7115                             | ± 0.000024  | 0.27            | 185                        | 0.096        | 874          | 0.212 |
| R1         | ענו         | SB          | 3.139                      |                                    | ± 0.000024  | 0.27            | 285                        | 0.090        | 597          | 0.212 |
| R9         |             | SB          | 3.630                      |                                    | ± 0.000024  |                 | 503                        | 0.138        | 911          | 0.552 |
| Aar31      | Bio         | #           | 0.594                      |                                    | ± 0.000026  |                 | 303                        | 0.136        | 911          | 0.332 |
| Aaisi      | Kfs         | #           | 3.311                      |                                    | ± 0.000045  |                 |                            |              |              |       |
| AD20       | Пуз         | π<br>#      | 2.686                      |                                    | ± 0.000039  |                 |                            |              |              |       |
| 711)20     | Plagio      | #<br>#      | 1.498                      |                                    | ± 0.000075  |                 |                            |              |              |       |
| ACIIh      | 1 iugio     | #           | 2.573                      |                                    | ± 0.000073  |                 |                            |              |              |       |
| 7 CHI      | Bio         | #           | 0.726                      |                                    | ± 0.000074  |                 |                            |              |              |       |
| 37         |             | 31          | 0.720                      | 0.7151                             | 2 0.000071  |                 |                            |              |              |       |
| Variscan g |             | CD          | 1.700                      | 0.7101                             | 0.000024    | 0.15            | 1010                       | 1.014        | 450          | 4.040 |
| R8         | TD          | SB          | 1.792                      |                                    | ± 0.000024  | 0.15            | 1818                       | 1.014        | 450          | 4.040 |
| R15        | TD          | SB          | 4.121                      | 0.7078                             | ± 0.000022  | 9.01            | 1230                       | 0.298        | 321          | 3.832 |
| Suspende   | d sedimen   | ıt          |                            |                                    |             |                 |                            |              |              |       |
| Oberaarg   | lacier      |             |                            |                                    |             |                 |                            |              |              |       |
| OAP6       | TD          | SB          | 1.838                      | 0.7148                             | ± 0.000013  |                 | 241                        | 0.131        | 569          | 0.423 |
| OA1        | SA          | BE          | 2.717                      |                                    | ± 0.000014  |                 | 355                        | 0.131        | 453          | 0.783 |
| OA2        | SA          | BE          | 2.705                      |                                    | ± 0.000008  |                 | 351                        | 0.130        | 500          | 0.702 |
| Rhonegla   |             |             |                            |                                    |             |                 |                            |              |              |       |
| RH3.8.     | TD          | SB          | 1.164                      | 0.7302 -                           | ± 0.000015  |                 | 141                        | 0.121        | 639          | 0.220 |
| RH1        | SA          | BE          | 1.104                      |                                    | ± 0.000013  |                 | 162                        | 0.121        | 621          | 0.220 |
| RH2        | SA          | BE          | 1.176                      |                                    | ± 0.000021  |                 | 168                        | 0.142        | 597          | 0.281 |
|            |             |             |                            |                                    | ± 0.00000∠₹ |                 | 100                        | 0.173        | 531          | 0.201 |
|            | e crusts (s |             | ially precipit             |                                    |             |                 |                            |              |              |       |
| OK1        |             | BE          | 1.725                      |                                    | ± 0.000013  |                 |                            |              |              |       |
| RH1        |             | BE          | 0.160                      | $0.7192 \pm$                       | ± 0.000024  |                 |                            |              |              |       |

SB measured at Centre de Géochimie de la Surface in Strasbourg; TD bulk samples

fied using suprapure HNO<sub>3</sub>, and stored in prewashed (10% HNO<sub>3</sub>) polypropylene bottles at 4 °C. Suspended sediment was collected from both glaciers during the ablation seasons (1999 to 2001) by filtering meltwater in the field. The sediment samples were oven-dried at 40 °C. Rain samples were taken during the 2000 and 2001 ablation seasons: A polypropylene bottle was connected to a plastic funnel via a polypropylene tube (all equipment was prewashed with 10% HNO<sub>3</sub>). The bottle was left for a maximum of 24

hours before the rainwater was filtered and stored like the meltwater samples. 17 samples of bedrock were taken randomly within the proglacial areas and on the valley walls beside the glaciers. The bedrock samples include the major lithologies of the two catchment areas. Subglacially precipitated carbonate deposits were also sampled in both catchments, they appear quite widespread in front of the retreating glaciers. Whole-rock and carbonate samples were crushed and ground to a fine powder using an agate mill.

BE measured at Geologisches Institut, University of Berne; SA strong acid leachates

NE measured at GEA Lab, University of Neuchâtel

<sup>¥</sup> Schaltegger et al. (1990)

<sup>#</sup> Challandes et al. (2001)

<sup>\*</sup> From RockEval analyses, GEA Lab, University of Neuchâtel

Table 3 Results of the different meltwater and rain water analyses.

| Sample   |         | Sr*<br>µmol l <sup>-1</sup> | <sup>87</sup> Sr/ <sup>86</sup> Sr | 2σ<br>error | Ca<br>µmol l⁻¹ | $Ca/(1000 \times Sr)$ | Na#<br>µmol l <sup>-1</sup> | Ca/Na    |
|----------|---------|-----------------------------|------------------------------------|-------------|----------------|-----------------------|-----------------------------|----------|
| Meltwate | er      |                             |                                    |             |                |                       |                             |          |
| Oberaar  | glacier |                             |                                    |             |                |                       |                             |          |
| O51      | BE      | 0.403                       | 0.7156 ±                           | 0.000024    | 203.74         | 0.51                  | 23.04                       | 8.84     |
| O65      | BE      | 0.197                       | 0.7137 ±                           | 0.000023    | 86.78          | 0.44                  | 6.52                        | 13.31    |
| O75      | BE      | 0.146                       | 0.7154 ±                           | 0.000020    | 90.27          | 0.62                  | 9.57                        | 9.44     |
| O80      | BE      | 0.109                       | 0.7174 ±                           | 0.000046    | 67.48          | 0.56                  | 16.96                       | 3.97     |
| P6       | SB      | 0.125                       | $0.7160 \pm$                       | 0.000013    | 69.98          | 0.56                  | 14.00                       | 5.00     |
| P4       | NE      | 0.218                       |                                    |             | 136.90         | 0.63                  | -                           | _        |
| P7       | NE      | 0.079                       |                                    |             | 83.40          | 1.05                  | 8 <del></del>               |          |
| K8       | NE      | 0.301                       |                                    |             | 136.77         | 0.45                  | 14.35                       | 9.53     |
| P8       | NE      | 0.240                       |                                    |             | 108.87         | 0.45                  | -                           |          |
| P9       | NE      | 0.479                       |                                    |             | 231.07         | 0.48                  | 9 <u></u>                   |          |
| K25      | NE      | 0.238                       |                                    |             | 99.52          | 0.42                  | 6.52                        | 15.26    |
| Rhone g  | lacier  |                             |                                    |             |                |                       |                             |          |
| A111     | BE      | 0.013                       | $0.7256 \pm$                       | 0.000159    | 9.48           | 0.75                  | 3.91                        | 2.42     |
| A142     | BE      | 0.018                       |                                    | 0.000192    | 12.22          | 0.67                  | 6.09                        | 2.01     |
| RH3.8.   | SB      | 0.011                       | $0.7251 \pm$                       | 0.000014    | 9.00           | 0.84                  | 5.00                        | 1.80     |
| P046     | NE      | 0.008                       |                                    |             | 23.65          | 2.82                  | 10.00                       | 2.37     |
| A067     | NE      | 0.007                       |                                    |             | 20.18          | 2.77                  | 7.83                        | 2.58     |
| P093     | NE      | 0.006                       |                                    |             | 9.10           | 1.51                  | 4.35                        | 2.09     |
| P071     | NE      | 0.009                       |                                    |             | 19.31          | 2.09                  | 8.26                        | 2.34     |
| P142     | NE      | 0.010                       |                                    |             | 27.77          | 2.87                  | 6.09                        | 4.56     |
| A169     | NE      | 0.027                       |                                    |             | 45.06          | 1.69                  | 15.65                       | 2.88     |
| Rain     |         |                             |                                    |             |                |                       |                             |          |
| RR1      | BE      | n.a.                        | 0.7100 ±                           | 0.000108    | 2.49           | _                     | 2.17                        | 1.15     |
| RR2      | BE      | n.a.                        | 0.7097 ±                           | 0.000024    | 11.72          | _                     | 3.91                        | 3.00     |
| RR3      | BE      | n.a.                        | 0.7098 ±                           | 0.000013    | n.a.           | =                     | n.a.                        | -        |
| RR4      | BE      | n.a.                        | 0.7100 ±                           | 0.000033    | n.a.           | _                     | n.a.                        | <u> </u> |

BE measured at Geologisches Institut, University of Berne

A first series of samples was analysed at the Geological Institute of the University of Berne. Approximately 100 ml of water were evaporated and the residue dissolved in 1 ml of 3M HNO<sub>3</sub>. The evaporated samples were taken into solution with 6M HCl and H<sub>2</sub>O (ultra-pure deionized), evaporated again and subsequently dissolved in 1 ml of 3M HNO<sub>3</sub>. 4 suspended sediment samples (approx. 0.06 g each) containing the sediment of several subsequent samplings and a carbonate deposit sample were digested for 72 hours at 120 °C in closed Savilex vessels with 14M HNO<sub>3</sub> (strong acid samples, SA). The supernatant was centrifuged, evaporated and subsequently dissolved in 1 ml of 3M HNO<sub>3</sub>.

All samples were spiked with approximately 0.05–0.06 mg of an <sup>84</sup>Sr spike. All acids used were purified by double distillation. All samples were centrifuged prior to analysis using columns with

Sr spec-pure resin. Strontium samples were stored in 0.01M HNO<sub>3</sub> and some of them were diluted with 0.35M HNO<sub>3</sub> prior to analyses. The samples were analysed on a Nu Instruments® Multiple Collector Inductively Coupled Plasma-Mass Spectrometer (MC-ICP-MS) in static mode. Mass fractionation corrections were applied by normalising the average <sup>87</sup>Sr/<sup>86</sup>Sr ratios to the <sup>86</sup>Sr/<sup>88</sup>Sr ratio of 0.1194. A total of 19 standards (NBS 987) were measured with an average <sup>87</sup>Sr/<sup>86</sup>Sr of 0.71030 (2 s.d. = 0.00002).

A second series of samples was analysed in the Centre de Géochimie de la Surface in Strasbourg. Representative bedrock samples, two suspended sediment samples, and meltwater samples were analysed for their Sr isotopic composition and chemistry. The chemistry, including Sr, was measured on an ICP-MS. For the analyses of the Sr isotopic composition, powders of the samples were

SB measured at Centre de Géochimie de la Surface in Strasbourg

NE measured at GEA Lab, University of Neuchâtel

<sup>\*</sup> measured using quadrupole ICP-MS at GEA Lab, University of Neuchâtel

<sup>\*</sup> measured using ion chromatography at GEA Lab, University of Neuchâtel

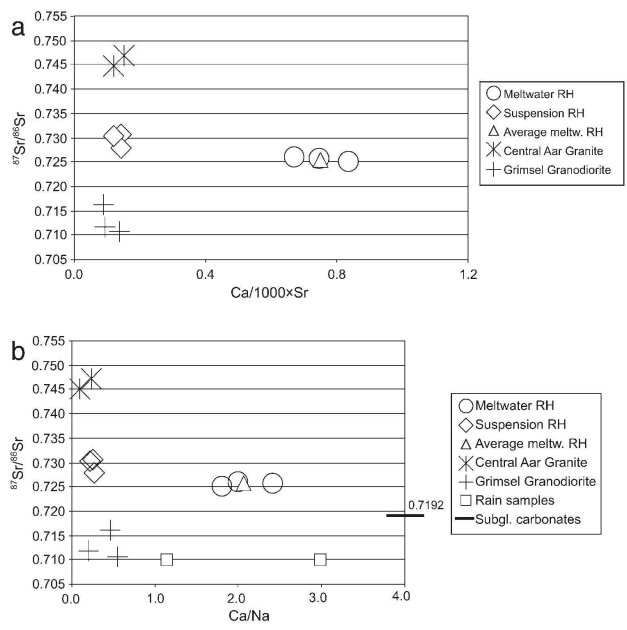


Fig. 2 (a)  ${}^{87}\text{Sr}/{}^{86}\text{Sr}$  vs. Ca/(1000×Sr) and (b)  ${}^{87}\text{Sr}/{}^{86}\text{Sr}$  vs. Ca/Na for water, bedrock and suspended sediment samples from Rhone catchment.

digested for 7 days in closed Savilex vessels containing a mixture of HF, HNO<sub>3</sub>, and HClO<sub>4</sub> (bulk samples, TD). Strontium was separated from the other ions using a cation exchange resin with 1.5 and 4M HCl as eluents. The total procedural blank was < 0.5 ng. The  $^{87}$ Sr/ $^{86}$ Sr ratios were determined on a VG Sector® multicollector thermal ionisation mass spectrometer (TIMS). Strontium was deposited on a W filament with Ta<sub>2</sub>O<sub>5</sub> as activator. Corrections for mass fractionation were made as above. The accuracy of the  $^{87}$ Sr/ $^{86}$ Sr analyses was evaluated by determining the NBS 987 standard. A mean value of  $^{87}$ Sr/ $^{86}$ Sr = 0.71030 (2 s.d. = 0.00003, n = 9) was obtained.

A third series of samples was analysed at the Geochemical and Environmental Analyses (GEA) Laboratory at the University of Neuchâtel: 10 fur-

ther meltwater samples were measured on a quadrupole ICP-MS. The detection limit of this quantitative method is < 100 ppt for Ca and < 1 ppt for Sr.

Total inorganic carbon was determined on about 100 mg of ground rock powder, using Rock Eval 6 standard bulk rock pyrolysis (Espitalié et al., 1986; Lafargue et al., 1996). The inorganic or mineral carbon contents (MINC) were calibrated with marble standards and expressed as wt% calcite. For low concentrations, the error is 20% (RSD); for higher concentrations (≥0.5 wt%), the error is 5% (RSD) or less.

Major cations in the meltwater samples were measured by ion chromatography (Dionex DX 500, Hosein et al., 2004). The error is  $\pm$  5% for concentrations of < 100 ppb and  $\pm$  2% for concentrations between 100 ppb and 10 ppm.

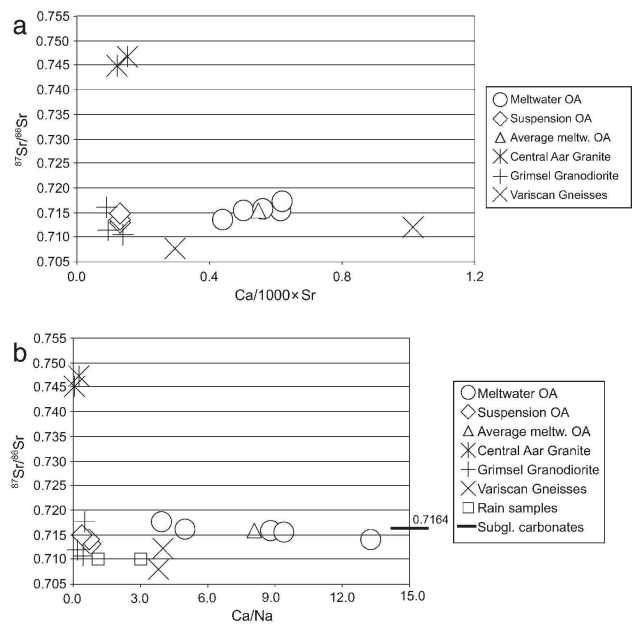


Fig. 3 (a) 87Sr/86Sr vs. Ca/(1000×Sr) and (b) 87Sr/86Sr vs. Ca/Na for water, bedrock and suspended sediment samples from Oberaar catchment.

# Results

Our bedrock <sup>87</sup>Sr/<sup>86</sup>Sr ratios generally correspond well with published data for the same rocks (Table 2). The <sup>87</sup>Sr/<sup>86</sup>Sr ratio for Central Aar Granite (0.7449) concurs with the data from Challandes (2001). Our <sup>87</sup>Sr/<sup>86</sup>Sr ratios for the Grimsel Granodiorite (0.7106 and 0.7115) are also comparable with Challandes (2001; 0.7088–0.7114). For the Variscan Gneisses two <sup>87</sup>Sr/<sup>86</sup>Sr ratios (0.7121 and 0.7078) were measured. There are no published values for this lithology. The <sup>87</sup>Sr/<sup>86</sup>Sr ratios of the Rhone suspended sediments are between 0.7279 and 0.7306, whereas the Oberaar suspended sediments have <sup>87</sup>Sr/<sup>86</sup>Sr ratios between 0.7130 and 0.7148.

The meltwaters from both glaciers contain similar concentrations of Sr: 0.006–0.027 and 0.079–0.479 µmol 1<sup>-1</sup> for Rhone and Oberaar, respectively (Table 3). The <sup>87</sup>Sr/<sup>86</sup>Sr ratios of the meltwaters sampled throughout the ablation season show little variation (0.7251–0.7258 for Rhone and 0.7137–0.7174 for Oberaar samples), and those of the rain samples measured are quite similar (Table 3).

Calcite was detected in all bedrock samples of all lithologies (Table 2). The calcite content of the Grimsel Granodiorite (0.11 wt%) is lower than that of the Central Aar Granite (0.27 wt%). The Variscan Gneisses contain 9 and 0.15 wt% of calcite, respectively; this large difference reflects the heterogeneity of these rocks (Arn, 2002).

The strong acid attack (14 M HNO<sub>3</sub>) of Oberaar suspended sediment yielded smaller <sup>87</sup>Sr/<sup>86</sup>Sr ratios (SA samples: 0.7130 and 0.7138) than the bulk sample (TD, 0.7148). We attribute this to differences in sample size. The SA samples contain the sediment of several subsequent sampling sessions, whereas the TD sample was taken on one single day. All Oberaar meltwater samples lie above the two <sup>87</sup>Sr/<sup>86</sup>Sr ratios of the Variscan Gneisses (bulk samples, TD, 0.7078 and 0.7121). The strong acid attack of Rhone suspended sediment yielded values comparable to those of the bulk sample (Table 2).

#### Discussion

# General trends of meltwater

Suspended sediment consists of bedrock that was recently mechanically abraded at the glacier base (Benn and Evans, 1998; Drewry, 1986), and its mineral content represents an average of the different bedrock compositions of todays' glacier catchments (Arn, 2002). The composition of suspended sediment is affected by subglacial geology, by abrasion and transport, and by the relative susceptibility (i.e. hardness) of lithologies and their constituent minerals to mechanical erosion. Minerals leaving a glacier may be leached subglacially, and further proglacial leaching may not correspond to the initial leach.

The Sr isotopic signal of the meltwaters is usually dominated by disseminated calcite, due to its high dissolution rate and its preferential weathering in subglacial environments, as reported from other glaciated areas (e.g., Tranter et al., 1993; Fairchild et al., 1994; Anderson et al., 1997; Fairchild et al., 1999a; White et al., 1999; Tranter et al., 2002; Hosein et al. 2004; this study). The Rhone and Oberaar meltwaters show elevated Ca/Na ratios supporting the hypothesis that Ca-rich minerals are weathered preferentially (Figs. 2b, 3b). The granitic rocks in the studied areas contain measurable amounts of disseminated calcite, but the Variscan Gneisses (found solely in Oberaar catchment) contain up to 9 wt% calcite (Table 2). This is the most likely source of Ca enrichment in the Oberaar catchment.

A daily trend in subglacial meltwater regime was not identified from our samples unlike in the studies of Sharp et al. (2002) or Anderson et al. (2000). The variable Ca enrichment, however, allows us to discuss the different sources of the Sr present in our meltwaters.

We suggest the main sources of Sr to be the preferential weathering of disseminated calcite,

contained within the different bedrock lithologies, together with easily hydrated ions derived from freshly cleaved silicate surfaces, whose early release is nonstoichiometric and may not correspond to the bulk mineral signature (Brantley et al., 1998). The relative importance of these sources is not the same in the two studied catchments (see below).

The concentrations of PO<sub>4</sub> measured in meltwaters from both glaciers are comparable even though the suspended sediment from the Oberaar catchment contains significantly more apatite than that from the Rhone catchment (average concentrations of total PO<sub>4</sub> are 0.07 ( $\pm$  0.03) and  $0.10 (\pm 0.04) \mu \text{mol } l^{-1}$  for Rhone and Oberaar ablation period runoff, respectively. Mean detrital PO<sub>4</sub> concentrations in suspended sediment are 0.32 and 0.58 ( $\pm$  0.05) mg g<sup>-1</sup> for Rhone and Oberaar, respectively (Hosein, 2002). Thus, we propose that apatite may only act as a minor source of Sr in the meltwaters studied here, and therefore, that its influence on the Sr signal is negligible. The lower 87Sr/86Sr ratio of the Rhone meltwaters (Fig. 2b), compared to bulk suspended sediment, may imply that the atmospheric input (with its generally low 87Sr/86Sr ratios) also has an impact on the meltwater signal. However, looking at the Ca/Na ratios of two of the rain samples, it appears that this influence is most probably negligible for Rhone meltwaters. Oberaar meltwater 87Sr/86Sr and Ca/Na ratios are higher than the suspended sediment and rain values, therefore atmospheric inputs are clearly negligible in the Oberaar catchment (Fig. 3b).

#### Rhone systematics

a) Suspended sediment vs. catchment lithologies. Using a mass balance approach, we calculated the portions of the two bedrock lithologies within the catchment (Central Aar Granite and Grimsel Granodiorite). The 87Sr/86Sr ratios and Sr concentrations of the two main lithologies in the Rhone catchment are listed in Table 2. Considering the suspended sediment to be a mixture of the two recently mechanically eroded bedrock lithologies (bulk suspended sediment (TD) 87Sr/86Sr ratio and Sr concentration) we calculated the portions of these lithologies within the suspended sediment. This calculation yields approximately  $76 \pm 1\%$  of Central Aar Granite and  $24 \pm 1\%$  of Grimsel Granodiorite. Assuming comparable overall hardness and erosion rates for both areas, this result further indicates that the subglacial exposure of the two lithologies are approximately 76 and 24% (Table 1). The values are quite different from an estimated distribution using different maps, which give approximately 90 and 10% for Central Aar Granite and Grimsel Granodiorite, respectively (Table 1). However, the available maps are not precise for the glacier-covered parts of the catchment, hence our calculated distribution of the two lithologies is probably more reliable.

b) Rhone meltwater signal. At the Rhone glacier, the meltwater 87Sr/86Sr ratio is smaller than that of the suspended sediment. We propose two reasons for this fact: (1) The Ca enrichment of the Rhone meltwaters is significant relative to Na compared to the Rhone suspended sediment, therefore we assume that the dissolution of disseminated calcite exerts a dominant impact on the meltwater chemistry (Hosein et al., 2004). The presence of subglacially precipitated carbonate crusts in front of the Rhone glacier suggests that they may also occur beneath the ice. Their subglacial dissolution exerts another lowering influence on the meltwater Sr signal (87Sr/86Sr ratio of 0.7192, Fig. 2b). (2) The nonstoichiometric Sr release from silicates during initial subglacial weathering seems to further lower the <sup>87</sup>Sr/<sup>86</sup>Sr ratios in our meltwaters. The feldspars from Grimsel Granodiorite have a higher Sr concentration (1.498–3.311 µmol g<sup>-1</sup>) than those from the Central Aar Granite (0.711– 0.822 µmol g<sup>-1</sup>). They may therefore dominate the nonstoichiometric Sr release and according to their lower 87Sr/86Sr ratios (0.7101–0.7105 compared to 0.7338–0.7434 for Central Aar Granite feldspars), they may decrease the meltwater 87Sr/86Sr ratios. The importance of nonstoichiometric Sr release from biotites is difficult to estimate, as the available Sr concentrations are similar but show a high degree of scattering. The Sr concentrations in biotite are all lower than in the feldspars from the Grimsel Granodiorite.

Summarizing, the Sr signal of Rhone meltwaters shows the influence of dissolution of disseminated calcite as well as the influence of nonstoichiometric Sr release from silicates.

#### Oberaar systematics

a) Comparing Oberaar lithologies and suspension. At Oberaar the lithological architecture is visible from the broad geomorphological framework: the main valley incision occurs in the zone of the highly deformed Variscan Gneisses, representing approximately 60% of the total glacier-covered area (Table 1). The remaining 40% at the glaciers' sides are composed of granitic rocks, which are harder and less erodable. The lithologies are elongated parallel to the main valley geometry from the glacier front to the top of the catchment (the Oberaar Joch). The average sus-

pended sediment flux (kg km<sup>-2</sup> yr<sup>-1</sup>) of the Oberaar glacier is 2 times higher than that of the Rhone glacier (Hosein et al., 2004). Assuming that the Variscan Gneisses represent 60% of the total glacier-covered area and that the erodability of granitic rocks is equal to 1, we calculate the erodability of the Variscan Gneisses to be 2.7 to obtain an overall flux that is 2 times higher. This corresponds to an influence of the gneiss on the bulk suspended sediment of 73%.

b) Oberaar meltwater signal. In accordance with the values of the two samples of Variscan Gneisses, the 87Sr/86Sr ratios of the Oberaar suspended sediment (TD) and meltwaters are lower than those of Rhone (Fig. 3a and 3b, Table 2 and 3). The Oberaar meltwater shows a greater Ca/ Na-enrichment compared to the suspended sediment than that of the Rhone glacier (Fig. 3b), pointing to the clear dominance of the dissolution of disseminated calcite and its impact on the meltwater chemistry and Sr signal in the Oberaar catchment. The Oberaar subglacial weathering regime is highly influenced by the presence of Variscan Gneisses with their higher erodability and calcite content compared to the granitic lithologies. However, a comparison of the varying 87Sr/86Sr ratios, as well as of the different calcite contents of the two samples of Variscan Gneisses (R8 = 0.15 wt%, R15 = 9 wt%) clearly shows the *highly* heterogeneous nature of these gneisses: they are not adequately represented by the two samples measured here. According to the Sr ratio of the meltwaters, an average gneiss calcite 87Sr/86Sr ratio would be expected to be close to  $0.716 \pm 0.001$ .

The general Ca enrichment of Oberaar meltwaters may partly be explained by the dissolution of subglacial carbonate precipitates, which are quite widespread in this catchment, as observed in front of the retreating glacier front. We assume that they also exist beneath the ice, where they would participate in the subglacial weathering processes. The <sup>87</sup>Sr/<sup>86</sup>Sr ratio of one sample (0.7164) and its Ca/Na ratio, which is higher than all the meltwater samples measured, seems to have a visible impact on the meltwater isotopic signal.

Even though single-mineral data are lacking for the gneisses, the strong Ca enrichment observed points to a minor impact of nonstoichiometric Sr release from silicates on Oberaar meltwaters compared to the calcite sources.

### Conclusions

This is the first report on Sr isotope systematics in purely crystalline glaciated catchments in the

Alps. In general, meltwaters draining crystalline glaciated watersheds reflect the high 87Sr/86Sr ratios of the underlying bedrock. Meltwater composition is influenced by different short-term subglacial processes: The dominant impact of calcite dissolution not only on major ion composition of the meltwater, but also on its Sr isotope systematics was confirmed in the Oberaar catchment (where the Variscan Gneisses contain more calcite and are more easily eroded than the other granitic rocks) by comparing the 87Sr/86Sr ratio of the meltwaters to the 87Sr/86Sr ratios of disseminated and subglacially precipitated calcite.

The Ca-enrichment of the Rhone meltwaters that drain a purely granitic catchment also suggests that the dissolution of disseminated calcite strongly influences the major ion composition of the runoff as well as its Sr isotope systematics. It was shown that the nonstoichiometric Sr release from silicates may exert another dominant impact.

# Acknowledgements

We thank Virginie Matera (Neuchâtel) and Ingeborg Hebeisen (Berne) for laboratory assistance. We also gratefully acknowledge the support of the Swiss National Science Foundation (Grants 21-53997.98 and 20-61485.00). This paper benefited from the constructive reviews of A. Galy and F. Chabaux.

#### References

Abrecht, J. (1994): Geologic units of the Aar massif and their pre-Alpine rock associations: a critical review. Schweiz. Mineral. Petrogr. Mitt. 74,5-27.

Anderson, S.P., Drever, J.I., Frost, C.D. and Holden, P. (2000): Chemical weathering in the foreland of a retreating glacier. Geochim. Cosmochim. Acta 64, 1173-1189

Anderson, S.P., Drever, J.I. and Humphrey, N.F. (1997): Chemical weathering in glacial environments. Geology **25**, 399–402.

Arn, K. (2002): Geochemical weathering of two glaciated crystalline catchments in the Swiss Alps (Oberaar- and Rhoneglacier). PhD thesis Université de Neuchâtel, 331 pp.

Aubert, D., Stille, P. and Probst, A. (2001): REE fractionation during granite weathering and removal by waters and suspended loads: Sr and Nd isotopic evi-

dence. *Geochim. Cosmochim. Acta* **65**, 387–406. Benn, D.I. and Evans, D.J.A. (1998): Glaciers and glaciation. Arnold Publishers, London, 734 pp.

Blum, J.D. and Erel, Y. (1995): A silicate weathering mechanism linking increases in marine 87Sr/86Sr with global glaciation. Nature 373, 415-418.

Blum, J.D. and Erel, Y. (1997): Rb-Sr isotope systematics of a granitic soil chronosequence: The importance of biotite weathering. Geochim. Cosmochim. Acta 61, 3193-3204.

Blum, J.D., Erel, Y. and Brown, K. (1993): 87Sr/86Sr ratios of Sierra Nevada stream waters: implications for relative mineral weathering rates. *Geochim Cosmo-chim. Acta* **58**, 5019–5025. Blum, J.D., Gazis, C.A., Jacobson, A.D. and Chamberlain, C.P. (1998): Carbonate versus silicate weathering in the Raikhot Watershed within the High Himalayan Crystalline Series. Geology 26, 411–414.

Brantley, S.L., Chesley, J.T. and Stillings, L.L. (1998): Isotopic ratios and release rates of strontium measured from weathering feldspars. *Geochim. Cosmochim. Acta* **62**, 1493–1500.

Challandes, N. (2001): Comportement des systèmes isotopiques <sup>39</sup>Ar-<sup>40</sup>Ar et Rb-Sr dans les zones de cisaillement: Exemples du massif de l'Aar (Massifs cristallins externes) et de la nappe de Suretta (Alpes centrales suisses). PhD thesis Université de Neuchâ-

tel, 268 pp. Clow, D.W., Mast, M.A., Bullen, T.D. and Turk, J.T. (1997): <sup>87</sup>Sr/Sr <sup>86</sup> as a tracer of mineral weathering reactions and calcium sources in an alpine/subalpine watershed, Loch Vale, Colorado. Water Resources Research 33, 1335–1351.

Drewry, D. (1986): Gacial Geologic Processes. Edward Arnold Ldt., London, 276 pp.

Edmond, J.M. (1992): Himalayan tectonics, weathering processes, and the strontium isotope record in marine limestones. Science 258, 1594–1597.

English, N.B., Quade, J., DeCelles, P.G. and Garizone, C.N. (2000): Geologic control of Sr and major element chemistry in Himalayan Rivers, Nepal. Geochim. Cosmochim. Acta 64, 2549-2566.

Espitalié, J., Deroo, G. and Marquis, F. (1986): La pyrolyse Rock-Eval et ses applications - III partie. Rev. Ínst. Fr. Pet. **41**, 73–89.

Fairchild, I.J., Bradby, L., Sharp, M.J. and Tison, J.-L. (1994): Hydrochemistry of carbonate terrains in Alpine glacial settings. Earth Surface Processes and Landforms **19**, 33–54.

Fairchild, I.J., Killawee, J.A., Hubbard, B.P. and Dreybrodt, W. (1999a): Interactions of calcareous suspended sediment with glacial meltwater: a field test of dissolution behaviour. Chemical Geology 155,

Fairchild, I.J., Killawee, J.A., Sharp, M.J., Spiro, B., Hubbard, B.P., Lorrain, R.D. and Tison, J.-L. (1999b): Solute generation and transfer from a chemically reactive alpine glacial-proglacial system. Earth Šurface Processes and Landforms 24, 1189–12TT.

Galy, A. and France-Lanord, C. (2001): Higher erosion

rates in the Himalaya: Geochemical constraints on riverine fluxes. *Geology* **29**, 23–26. Galy, A., France-Lanord, C. and Derry, L.A. (1999): The strontium isotopic budget of Himalayan Rivers in Nepal and Bangladesh. Geochim. Cosmochim. Acta 63, 1905–1925.

Hallet, B., Hunter, L. and Bogen, J. (1996): Rates of erosion and sediment evacuation by glaciers: A review of field data and their implications. Global and Planetary Change 12, 213-235

Harris, N. (1995): Significance of weathering Himalayan metasedimentary rocks and leucogranites for the Sr isotope evolution of seawater during the early Miocene. *Geology* **23**, 795–798.

Hodell, D.A., Mead, G.A. and Mueller, P.A. (1990): Variation in the strontium isotopic composition of seawater (8 Ma to present): Implications for chemical weathering rates and dissolved fluxes to the oceans. Chemical Geology **80**, 291–307. Hodell, D.A. and Woodruff, F. (1994): Variations in the

strontium isotopic ratio of seawater during the Miocene; stratigraphic and geochemical implications. *Paleoceanography* **9**, 405–426.

Hosein, R. (2002): Biogeochemical weathering processes in the glacierised Rhone and Oberaar catchments, Switzerland and the Apure catchment, Venezuela. PhD thesis, Université de Neuchâtel, 312 pp.

Hosein, R., Arn, K., Steinmann, P., Adatte, T. and Foellmi, K. (2004): Carbonate and silicate weathering in two presently glaciated, crystalline catchments in the Swiss Alps. *Geochim. Cosmochim. Acta* 68, 1021–1033.

Jacobson, A.D. and Blum, J.D. (2000): Ca/Sr and 87Sr/86Sr geochemistry of disseminated calcite in Himalayan silicate rocks from Nanga Parbat: Influence on river-

water chemistry. Geology 28, 463–466.

Jacobson, A.D., Blum, J.D., Chamberlain, C.P., Poage, M. and Sloan, V.F. (2002): Ca/Sr and Sr isotope systematics of a Himalayan glacial chronosequence: Carbonate versus silicate weathering rates as a function of landscape surface age. *Geochim. Cosmochim. Acta* 66, 13–27.

Lafargue, E., Espitalié, J., Marquis, F. and Pillot, D. (1996): Rock Eval 6 applications in hydrocarbon exploration, production and in soil contamination studies, Latin American Congress on Organic Geochemistry, Cancun.

Oberhänsli, R., Schenker, F. and Mercolli, I. (1988): Indications of Variscan nappe tectonics in the Aar Massif. Schweiz. Mineral. Petrogr. Mitt. 68, 509–520.

- sif. Schweiz. Mineral. Petrogr. Mitt. **68**, 509–520. Palmer, M.R. and Edmond, J.M. (1989): The strontium isotope budget of the modern ocean. Earth Planet. Sci. Lett. **92**, 11–26.
- Raymo, M.E. and Ruddiman, W.F. (1992): Tectonic forcing of late Cenozoic climate. *Nature* **359**, 117–122.
- Raymo, M.E., Ruddiman, W.F. and Froelich, P.N. (1988): Influence of late Cenozoic mountain building on ocean geochemical cycles. *Geology* 16, 649–653.
- Schwab, M., Frei, C., Schär, C. and Daly, C. (2001): Mean annual precipitation throughout the European Alps 1971–1990. In: BWG (Bundesamt für Wasser und Geologie, Editor), The Hydrological Atlas of Switzerland. Bundesamt für Landestopographie, Bern.

- Sharp, M.J., Creaser, R.A. and Skidmore, M. (2002): Strontium isotope composition of runoff from a glaciated carbonate terrain. *Geochim. Cosmochim. Acta* 66, 595–614.
- Sharp, M.J., Tranter, M., Brown, G.H. and Skidmore, M. (1995): Rates of chemical denudation and CO<sub>2</sub> drawdown in a glacier-covered alpine catchment. *Geology* **23**, 61–64.
- Stalder, H.A. (1964): Petrographische und mineralogische Untersuchungen im Grimselgebiet (Mittleres Aarmassiv). Schweiz. Mineral. Petrogr. Mitt. 44, 187–389.
- Taylor, A. and Blum, J. (1995): Relation between soil age and silicate weathering rates determined from the chemical evolutin of a glacial chronosequence. *Geology* 23, 979–982.
- Tranter, M., Brown, G., Raiswell, R., Sharp, M. and Gurnel, A. (1993): A conceptual model of solute acquisition by Alpine glacial meltwaters. *J. Glaciology* **39** (133), 573–581.
- Tranter, M., Sharp, M., Lamb, H.R., Brown, G.H., Hubbard, B.P. and Willis, I.C. (2002): Geochemical weathering at the bed of Haut Glacier d'Arolla, Switzerland A new model. *Hydrological Processes* **16**, 959–993.
- White, A.F., Bullen, T.D., Vivit, D.V., Schulz, M.S. and Clow, D.W. (1999): The role of disseminated calcite in the chemical weathering of granitoid rocks. *Geochim. Cosmochim. Acta* **63**, 1939–1953.

Received 21 December 2003 Accepted in revised form 27 July 2004 Editorial handling: R. Gieré