

## The challenge: interpretation of whole-rock chemical data



- Lunar program in the late 1960's came with the requirement of precise and accurate chemical, and then isotopic, analyses of small samples.
- In 1970's appeared novel analytical techniques for trace-element determinations (e.g., XRF; INAA), later ICP-OES, ICP-MS.
- Advancement of radiogenic isotope methods (originally TIMS, then SIMS, ICP-MS).

 Downside: the current flood of precise geochemical data needs to be interpreted by a potent and widely available software tool.

# A revolution? The Language



- Originally designed by Ihaka and Gentleman (1996)
- Version 1.0 published on 29 Feb 2000
- Based on syntax of the S language (Becker et al. 1988) which was developed at Bell Laboratories (formerly AT&T, now Lucent Technologies) by John Chambers and colleagues.
- The commercial version, S-PLUS, is currently being distributed by TIBCO Software (<u>www.tibco.com</u>)
- Since 1997 the development of the R project is overseen by an open group of experts, the R Development Core Team (<a href="http://www.r-project.org">http://www.r-project.org</a>)
- R is Free Software distributed under the Free Software Foundation's GNU General Public License
- R is available for all main OS. It runs on a wide variety of UNIX platforms and similar systems (including FreeBSD and Linux), Windows and MacOS.

# A revolution? The Language

Frequently updated, supported by a large and still growing community



- R is highly extensible by numerous additional packages
- Large collection of statistical (linear and nonlinear modelling, classical statistical tests, time-series analysis, classification, clustering, ...) and database tools
- Data import in many formats, also via SQL and ODBC
- Graphical facilities for data exploration and plotting
- High-level graphical output, well-designed publication-quality plots can include mathematical symbols and formulae where needed
- Graphics can be exported into many formats (*PostScript, WMF, PDF, TIFF, PNG...*), for incorporation into DTP programs, word processors or further editing
- Excellent control over individual functions [= power]
- R allows interactive as well as batch use (as a true programming language).
- Effective object-oriented programming language
- Most of the R system is written in R; for computationally intensive tasks, C,
   C++, and FORTRAN code can be linked and called at run time.

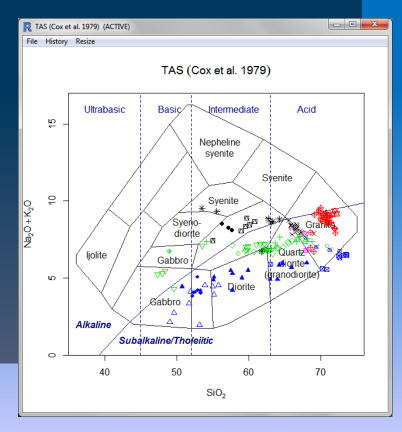


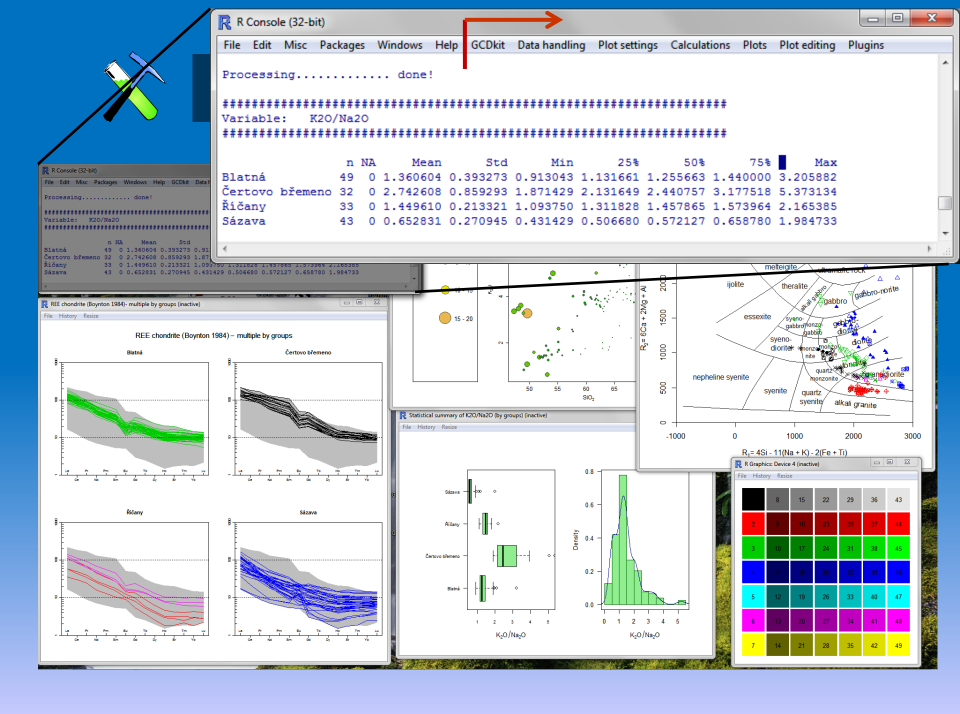
## Geochemical Data Toolkit = GCDkit

### **MAIN FEATURES:**

http://www.gcdkit.org

- A more human (less inhuman) interface to the wealth of functions in R
- Windows-like GUI = no programming necessary!!
- Data ready for further handling under R (dot prompt veterans)
- Standard geochemical calculations involving whole-rock major-, traceelement data and Sr-Nd isotopes
- Effective data management (searching, subsetting, grouping)
- Common plots (binary, ternary, spider, classification, geotectonic...)
- Publication quality graphic output







## Main 'milestones'



- 2000 launched graduate-level courses on interpretation of geochemical data using R (Masaryk University in Brno & Charles University in Prague)
- 2003 Goldschmidt Conference, Kurashiki, Japan GCDkit 1.0 released
- 2006 key publication in *Journal of Petrology* (141 hits on WOS)
- October 2015 Monograph on Geochemical modelling in R/GCDkit (Springer Verlag)
- February 10, 2016 last stable version (4.1, Chlopskie jadlo) released for current R (ver. 3.2.1)
- October 2016 brand new graduatelevel course on data analysis using R and Python (Charles University in Prague)

JOURNAL OF PETROLOGY VOLUME 47 NUMBER 6 PAGES 1255-1259 2006 doi:10.1093/petrology/eg013

#### TECHNICAL NOTE

Interpretation of Whole-rock Geochemical Data in Igneous Geochemistry: Introducing Geochemical Data Toolkit (GCDkit)

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## Invited workshops on GCDkit and/or R modeling

- Czech Geological Survey, Prague, (11 June 2004),
- TU Bergakademie Freiberg, Germany (16 Oct 2005),
- CGS/EOST, Université Louis Pasteur Strasbourg, France (23–24 Oct 2008),
- University of Tromsø, Norway (16–17 June 2010),



- Université Jean Monnet, Saint-Etienne, France (9–11 May 2011),
- University of Helsinki, Finland (7–11 Nov 2011),
- University of Stellenbosch, South Africa (19–23 Mar 2012),
- National Geophysical Research Institute, Hyderabad, India (12–15 Jan 2013),
- University of Arba Minch, Ethiopia (2–6 April 2015),
- Polish Academy of Sciences, Kraków, Poland (23–27 Nov 2015),
- University of Catania, Sicily, (26–30 Sept 2016),...

## "The R Book" – philosophy



Geochemical Modelling of Igneous Processes – Principles and Recipes in R Language

Bringing the Power of R to a Geochemical Community





- Provides basics of R language and its application to geochemical problems,
- Gives the first comprehensive introduction to the GCDkit system,
- Explains fundamentals of numerical modelling of igneous processes,
- Shows not only formulae, but also the successful modelling strategies,
- Includes numerous worked examples how geochemical modelling helps us to understand geological problems.

*Springer Geochemistry* series, vol. 1 345 pp., 332 illus., 86 illus. in colour D 83,20 € | UK £72.00 | US \$99.00

#### Exercise 2.7: Classification using factors

· Classify samples in the Sázava set according to SiO2 contents (wt. %) in four groups, U (< 45), B (45-52), I (52-63) and A (> 63).



```
(B)
```

```
> sazava <- read.table("sazava.data",sep="\t")
> silica <- cut(sazava[, "SiO2"],breaks=c(0,45,52,63,100),
      labels=c("U", "B", "I", "A"))
[1] IIIBIIBBBIIAAA
Levels: U B I A
```

Note that the levels that do not occur in the data at all (here the ultrabasic rocks, U) are not dropped. If we want to know the classification of individual samples, we convert the factor silica to a character vector:

```
> names(acidity) <- rownames(sazava)
> acidity
Sa-1
Gbs-2 Gbs-3 Po-
```

> acidity <- as.vector(silica)

Sa-3 Sa-4 Sa-7 SaD-1 Gbs-1 Gbs-20



# GCDkit solution GCDkit-> loadData GCDkit-> cutMy("S SiO2 Inte

59.98 Sa-2 55.17 Sa-3 55.09 50.72

#### 2.4.3 Frequency

A nifty application of f

3.6 Spidemlots

a) Write a function that would normalize REE concentrations in the sample by chondritic contents (stored in a comma-delimited file boynton.data).

b) Calculate the normalized REE concentrations for analyses of the Sázava suite and display-using the functions plot, axis, points and linesspiderplots for two Požáry trondhjemites, Po-1 and Po-4.

```
a) > x <- read.table("boynton.data", sep=",")
> chondrite <- as.numeric(x) # conversion to numeric vector
> names(chondrite) <- names(x)
> norm <- function(x,chon){
                                      # normalizing function
      z <- t(x[,names(chon)])/chon
> }
b) > sazava <- read.table("sazava.data",sep="\t")
```

```
> y <- norm(sazava,chondrite)
> plot(y[,"Po-1"],type="0",log="y",axes=FALSE,xlab="",
+ ylab="REE/chondrite",ylim=c(0.1,100),col="darkgreen")
> axis(1,1:length(chondrite),labels=names(chondrite),
      cex.axis=0.75)
> axis(2,cex.axis=0.75)
> points(y[,"Po-4"],col="blue")
> lines(y[,"Po-4"],col="blue")
> abline(h=(10^(-1:3)),lty="dashed") # grid
                                                      # bounding box
```

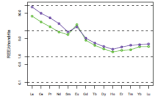


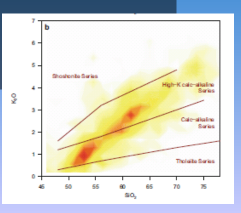
Fig. 3.7 Chondrite-normalized REE patterns for two Požáry trondhjemites (Exercise 3.4).

Plotting spiderplots in GCDkit The preceding exercise can be reproduced easily, using the mother-of-all spiderplot functions in GCDkit, spider. To see normalization schemes available in GCDkit, type ?selectNorm. Specialized spiderplots are dealt with in Chap. 4.2. You can try some of the examples below (see ?spider for more, including a complete list of arguments and different ways to define the norm to use).

## "The R Book" – contents

### Part I: R/GCDkit at work

- Loading and manipulating data
- Linking Whole-Rock Chemistry with Mineral Stoichiometry
- **Statistics**
- Classification and Grouping
- Classical Plots (binary, Harker, ternary, spider)
- Specialized Plots (log-log, specialized spiderplots, contour plots, anomaly plots...)
- Radiogenic isotopes (initial ratios, epsilon values, model ages, isochrons...)



#### 11.4 Assimilation and Fractional Crystallization (AFC)

The AFC model describes the compositional evolution of a magma that undergoes simultaneous assimilation and fractionation. This is, for instance, the case of a mantle-derived magma contaminated by the continental crust through which it ascends. The model assumes that the extra heat needed for assimilation (which is an endothermic process) is provided by the latent heat of crystallization. Already O'Hara (1977) developed equations describing trace-element (and isotope) behaviour during open-system fractionation in a periodically recharged magma chamber with assimilation. The forward AFC model was elaborated by Allègre and Minster (1978), Taylor (1980) and De Paolo (1981).

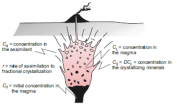


Fig. 11.7 Schematic representation of the AFC process for trace elements (after De Paolo 1981).

The instantaneous mass balance of the process (Fig. 11.7) is:

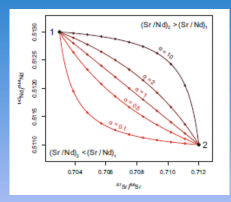
$$dW_L + dW_C + dW_A = 0 (11.31)$$

with  $W_L$ ,  $W_C$  and  $W_A$  referring to the amounts of the given element in the melt, crystallized phases and assimilated material, respectively. The evolution of the magma chemistry is then controlled by interplay of two key parameters: D, the bulk distribution coefficient and r, rate of assimilation to fractional crystallization (i.e., the mass ratio of material assimilated to that crystallized) (De Paolo 1981):

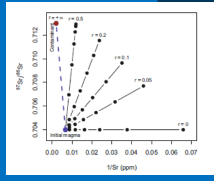
$$r = \frac{\dot{W}_A}{\dot{W}_A}$$
(11.32)

Defining the fraction of the liquid remaining, F:

$$F = \frac{W_0 - W_C + W_A}{W}, \quad (11.33)$$



"The R Book" – contents



14 Forward Modelling in R

GCDkit-> legend("bottomright",legend=rep(ff,2),pch=15,col=col,
+ bg="white",ncol=2,title="PRIMA/DM")

#### Deep melting (15 kbar)

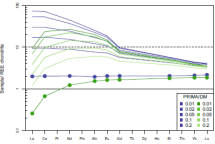


Fig. 14.4 Chondrite-normalized REE patterns for various degrees of a deep (gamet stability field) batch melting of Primitive Mantle (PRIMA, blue circles) and Depleted Mantle (DM, green circles). Magmas resembling OIB are generated from the PRIMA source, but the DM/deep melts resemble no common two of basalts (Exercise 14.3).



Table 14.4 contains selected trace-element contents in an average Mid-Ocean Ridge Basalt (NMORB, Sun and McDonough 1989) and an Upper Continental Crust (UCC, Taylor and McLennan 1995).

#### Table 14.

	Rb	Zr	Nb
NMORB	0.56	74	2.33
UCC	112	190	25

. Draw a mixing hyperbola between the two end members in a Rb vs. Zr/Nb plot.

## Parts II–IV: Majors, traces, radiogenic isotopes

- Core of the book
- Explains fundamentals of each direct and reverse modelling, including the relevant formulae
- Then introduces the numerical solution and its implementation in the R language
- Includes a number of real numerical problems
- Each is presented as a numerical receipt with solution in R (± GCDkit)

#### 21 Semi-Quantitative Geochemical Approach

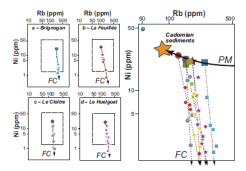


Fig. 21.4 a-d. Logfincompatible = Rb) vs. logfcompatible = Ni) plots for Cadomian and Hereynian granites from Brittany. Six individual granitic suites (four of which are plotted here) define fractional crystallization (FC) trends, specific to each pluton, e The parental melts of each pluton (darker, bigger symbols), however, plot along a batch melting trend (PM), the source being local metasediments (Georget 1986).

#### 21.3.3 Crystallization vs. Melting Using Incompatible Elements

Minster and Allègre (1978) proposed an alternative treatment, based only on in-

compatible elements and specifically focusing (i.e. basalts). Element  $\alpha$ , as previously  $D_{\alpha} \ll F \ll 1$ . Element  $\gamma$  is moderately incobining Rayleigh's equations for  $\alpha$  and  $\gamma$  yie

 $\frac{C_L^{\alpha}}{C_L^{\gamma}} = \frac{C_0^{\alpha}}{C_0^{\gamma}}$ 

Since  $D_a$  and  $D_v$  are small, Eq. (21.9) becomes

 $\frac{C_L^{\alpha}}{C_L^{\gamma}} \approx$ 

The ratio of two incompatible elements is

22.3 Dealing with Accessory Minerals



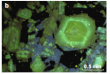
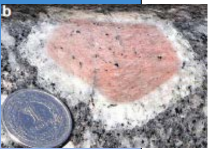


Fig. 22.5 Petrological identification of accessory minerals. a Optical microscope (cross-polarized light) timage of a monet allamic, mineral by pietacie, in *Kankonosez ganital*. Poland (Photo & Salary) b Optical CL image of the Kozinovice granoclierite, Central Bohemian Plutonic Complex (Lanosals et al. 2000). Apart from plagociaes (cohecidal; yellowy and K-fekspar (blos), the technique reveals tiny bright yellow apaties concentrated mainly in homblende (euhodralcadobode) tookunique over air; early



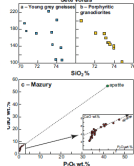


Fig. 226 Identifying the role of accessory minerals fractionation. a-b in the Sete Volas Archaene plution issuites (Brazi) (Martin et al. 1997), young grey genises show no correlation between SO<sub>2</sub> and Zr (a), whereas a strong negative correlation is observed in the prophyritic granocliorites (b), implying zirron fractionation e. P.G. vol. Col diagram discloses the role played by apatite during the differentiation of the Mazury granitoid Complex (Poland) (Duckense et al. 2010), For P.G. v. O'. vt. S<sub>c</sub>, data plot on a trend pointing towards the apatite composition.

## "The R Book" – contents

### Part V: Practical Modelling

- Choosing an Appropriate Model (evidence for crystallization, partial melting, magma mixing and assimilation...)
- Semi-Quantitative Approach

   (assessing the trace-element compatibility, process identification, mixing test...)
- Constraining a Model

   (using appropriate strategy, input parameters for the model, partition coefficients, dealing with accessories...)
- Numerical Tips and Tricks
   (reducing system, colinearity, breaking minerals to end-members, coupling majors and traces...)
- Common Sense in Action

   (thermodynamic, rheological constraints, scale/speed of processes, how well can we distinguish between models, dangerous projections...)

25.4 Summary 259

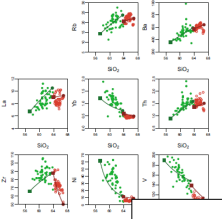


Fig. 25.12 Binary plots of silica (wt. %) vs. selected trace sults of the two-step model for trace elements. Same captionally left empty, to facilitate comparison with Fig. 25.5.

#### 25.4 Summary

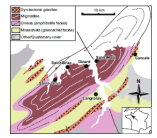
The preferred model is summarized in Figs. 25 diagrams reveals that our model is perhaps not id prove it. One could for instance try different phenocrysts are indeed compositionally variable), (within the range of permissible values). One may portions, estimated with other approaches (e.g., traces). One may also question the choice of

#### Chapter 26

Progressive Melting of a Metasedimentary Sequence: the Saint-Malo Migmatitic Complex, France<sup>1</sup>

#### 26.1 Geological Setting

The Saint-Malo Massif belongs to the high-T belt of the Cadomian Orogen (Brun and Balé 1990). Figure 26.1 shows that it consists of three main lithological and metamorphic units (Brun and Martin 1977), which are, from S to N, (i) greenschist-facies micaschists and gneisses, (ii) amphibolite-facies gneisses, and (iii) migmatities. The metamorphic event, culminating in anatexis, has been dated at ca. 3-40 Ma (Peucat and Martin 1985; Peucat 1986).



ig. 26.1 Simplified geological map of the Saint-Malo Massif showing the geographical distribuon of the main lithologies; the degree of metamorphism increases northwards, i.e. towards the ore of the dome (redrawn after geological map of France and Martin 1980). Insect sketch map of nance for orientation; arrow shows the location of the Saint-Malo Massif.

#### DOI 10.1007/978-3-662-46792-3 26

### "The R Book" – contents

- Part VI: Worked Examples
- Differentiation of a Calc-Alkaline
   Series: Atacazo-Ninahuilca volcanoes,
   Ecuador
- Progressive Melting of a
   Metasedimentary Sequence: the Saint-Malo Migmatitic Complex, France
- Appendix A: R Syntax in a Nutshell
- Appendix B: Introduction to GCDkit
- Appendix C: Solving Systems of Linear Algebraic Equations in R

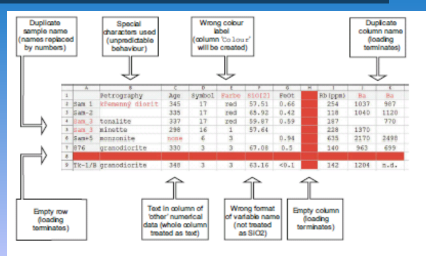


Fig. A2.4 Examples of common formatting mistakes.

Adapted from Brown (1979), Martin (1977, 1979, 1980) and Weber et al. (1985).

Springer-Verlag Berlin Heidelberg 2015
 V. Janoušek et al., Geochemical Modelling of Igneous Processes – Principles And Recines in R Language Springer Geochemistry



### Web links

http://www.r-project.org http://www.gcdkit.org http://blog.gcdkit.org http://book.gcdkit.org

## GCDkit forum

News, hints, comments and more from GCDkit team

Tuesday 27 October 2015

#### Book on geochemical modelling of magmatic processes in R/GCDkit

We break the two years of our silence with some news potentially interesting to anyone involved in interpretation of whole-rock geochemical data from igneous rocks. We are pleased to announce that, after two years of intense writing, we have finally published a

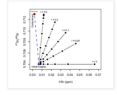
Janoušek, V., Moven, J. F., Martin, H., Erban, V. & Farrow, C. M. (2016). Geochemical Modelling of Igneous Processes - Principles and Recipes in R Language. Bringing the Power of R to a Geochemical Community. Springer-Verlag, Berlin, Heidelberg, 346



doi:10.1007/978-3-662-46792-3

#### Abstract from the publisher:

"The aim of this book is to unlock the power of the freeware R language to advanced university students and researchers dealing with whole-rock geochemistry of (meta-) igneous rocks. The first part covers data input/output, calculation of commonly used indexes and plotting in R. The core of the book then focuses on the presentation and practical implementations of modelling techniques used for fingerprinting processes such as partial melting, fractional crystallization, binary mixing or AFC using major-, trace-element and radiogenic



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#### **Getting Started**

R is a free software environment for statistical computing and graphics. It compiles and runs on a wide variety of UNIX platforms, Windows and MacOS. To download R, please choose your preferred CRAN mirror.

If you have questions about R like how to download and install the software, or what the license terms are, please read our answers to frequently asked questions before you send an

#### News

- The R Journal Volume 8/1 is available.
- The useR! 2017 conference will take place in Brussels. July 4 - 7, 2017, and details will be appear here in due course.
- . R version 3.3.1 (Bug in Your Hair) has been released on Tuesday 2016-06-21.
- R version 3.2.5 (Very, Very Secure Dishes) has been released on 2016-04-14. This is a rebadging of the quick-fix release 3.2.4-revised.
- Notice XQuartz users (Mac OS X) A security issue has been detected with the Sparkle update mechanism used by XQuartz. Avoid updating over insecure channels.
- The R Logo is available for download in high-resolution PNG or SVG formats.
- useR! 2016, hase taken place at Stanford University, CA, USA, June 27 - June 30, 2016.

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## References and further reading

