

# Chapter 2

## Fundamental Concepts of Compositional Data Analysis

**Abstract** Compositional data is considered a statistical scale in its own right, with its own natural geometry and its own vector space structure. Compositional data analysis and this book cannot be understood without a basic knowledge of these issues and how they are represented in **R**. Therefore, this chapter introduces the basic geometric concepts of compositional data and of the **R**-package “compositions”: the relative nature of compositional data; how to load, represent, and display compositional data in **R**; the various compositional scales and geometries and how to select the right geometry for the problem at hand; and how to specify the geometry in “compositions” using the basic classes “acomp”, “rcomp”, “aplus”, “rplus”, “ccomp”, and “rplus”. A concise guide to the most important geometry for compositional data, the Aitchison geometry, is also included. The whole book relies on these basic principles, and the reader should make him or herself familiar with them in Sects. 2.1–2.5 before going on.

### 2.1 A Practical View to Compositional Concepts

#### 2.1.1 Definition of Compositional Data

A *composition* is often defined as a vector of  $D$  positive components  $\mathbf{x} = [x_1, \dots, x_D]$  summing up to a given constant  $\kappa$ , set typically equal to 1 (portions), 100 (percentages), or  $10^6$  (ppm). We already mentioned that this definition is misleading, because many compositional datasets do not *apparently* satisfy it. Let us see how.

Consider, for instance, this simple example of compositional data collected from typical nutrition tables of foodstuff in any kitchen cupboard:

	Fat	Sodium	Carbonates	Protein	Total
soy	7	14	10	13	100
peas	3	12	61	22	100

wheat	1	15	60	23	100
corn	1	180	15	2	82
beans	0.5	680	30	12	NA

This data can be loaded from a text file<sup>1</sup> into **R** with the command `read.table`:

```
> library(compositions)
> kitchen <- read.table("SimpleData.txt", sep="")
```

All the amounts are given in grams g, except for sodium which is given in milligrams mg. A naive interpretation of these raw numbers would tell us that soy and corn are poor in protein and carbonates, whereas beans have seemingly 4 times more sodium and twice more carbonates than corn. However, this naive (i.e., non-compositional) direct interpretation contradicts everything we know about soy or corn. For a good compositional analysis, it is important to understand the fundamental operations and laws governing compositional analysis to avoid these pitfalls. This chapter introduces this methodological background step by step.

### 2.1.2 Subcompositions and the Closure Operation

A first inspection shows that the entries do not sum up to the total 100 %, since these products do not exclusively contain the nutrients reported. We thus only get some parts of a bigger composition, which would include water, vitamins, and further minerals. For instance, these non-reported components represent  $100 - (7 + 14/1000 + 10 + 13) \sim 70$  g out of 100 g of soy.

As people with a cholesterol problem and an office job, we may be interested in getting the most protein for the least amount of fat while maintaining a reasonable medium amount of carbohydrates. Our water and sodium intake is much more influenced by drinking and salting than by a diet choice between beans and peas. Thus, we are going to focus our attention to the composition of fat–protein–carbohydrates. It is typical to *close* the amounts in the parts of interest to sum up to 100 %. In “compositions,” this can be done with the command `clo` (for close composition):

```
> (coi <- clo(kitchen, parts=
              c("Fat", "Carbonates", "Protein"), total=100))
```

	Fat	Carbonates	Protein
soy	23.333	33.33	43.33
peas	3.488	70.93	25.58
wheat	1.190	71.43	27.38

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<sup>1</sup>The “SimpleData.txt” file can be downloaded from <http://www.stat.boogaart.de/compositionsRBook>.

corn	5.556	83.33	11.11
beans	1.176	70.59	28.24

These are now portions given in mass%. The command `clo` has an optional parameter `parts` taking the names or the numbers of columns to be selected and an optional argument `total` specifying the new reference total. If `parts` is not specified, implicitly all columns are used. If `total` is not specified, a default of 1 is assumed.

Note that a naive interpretation of these numbers would now tell us that soy is the richest in protein (whereas before, it was rather in the middle of the ranking for protein amount), and that corn is now richer in carbonates than beans. This illustrates the main risk of interpreting raw portions: our conclusions can be opposite, depending on which composition we are observing.

A composition only representing some of the possible components is called a *subcomposition*. Most of real compositional data is actually representing a subcomposition, as we never analyze each and every possible component of our samples. [Aitchison \(1986\)](#) therefore introduced the principle of *subcompositional coherence*: any compositional data analysis should be done in a way that we obtain the same results in a subcomposition, regardless of whether we analyzed only that subcomposition or a larger composition containing other parts. The interpretation we have been doing of the kitchen example up to now was not coherent, as our consideration of which food is richer in carbonates or protein changes between the original composition and the subcomposition of interest. This concern affects the whole direct application of all traditional multivariate statistics: none obey the principle of subcompositional coherence.

Reclosure is often a controversial *manipulation*, as by closing a subcomposition, it seems that we lose some information on the total mass present in the parts considered, and *apparently* we replace some measured values by some computed ones. However, we must consider that some of the cases in the kitchen dataset are dry and other cooked conserves, some even pre-salted, such that I do not need to add salt anymore. Some of the sodium (and of the other components, as well as water content) are thus artifacts of the form of delivery: they are related neither to the crop properties nor to the final cooked product we might eat. The only relevant quantities unaltered throughout these manipulations are the ratios between nutrients, and they are preserved by the closure operation.

### 2.1.3 Completion of a Composition

An alternative way to recast a vector of amounts to a composition is by adding the complementary variable. We first select the compositional variables using usual vectorized indexing<sup>2</sup> with `[]` :

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<sup>2</sup>Call `help("[")` in **R** for a detailed explanation.

```
> amounts <- kitchen[,c("Fat", "Carbonates", "Protein")]
> amounts
```

	Fat	Carbonates	Protein
soy	7.0	10	13
peas	3.0	61	22
wheat	1.0	60	23
corn	1.0	15	2
beans	0.5	30	12

and then compute the total grams of each column by

```
> (sumPresent <- totals(rplus(amounts)))

    soy  peas wheat  corn beans
30.0  86.0  84.0  18.0  42.5
```

This last line marks the dataset as positive real numbers by applying the `rplus` class<sup>3</sup> and then computes the sum of the individual amounts, by rows with the command `totals`.

We can now create a new dataset with an additional column `Other` giving the remaining mass:

```
> Other <- kitchen[, "total"] - sumPresent
> Amounts <- cbind(amounts, Other=Other)
> clo(Amounts)
```

	Fat	Carbonates	Protein	Other
soy	0.07000	0.1000	0.13000	0.7000
peas	0.03000	0.6100	0.22000	0.1400
wheat	0.01000	0.6000	0.23000	0.1600
corn	0.01220	0.1829	0.02439	0.7805
beans	0.01176	0.7059	0.28235	NA

The `cbind` command (for *column bind*) binds the columns of its arguments to a joint dataset containing all the columns.

Of course, this procedure is only applicable when we know how much sample was analyzed in order to subtract from it the sum of the present components. This completion procedure has an advantage over the reclosure of the observed subcomposition: the reclosure loses all information about how much we had in total of the observed components, whereas in the completion procedure this total present is inherited by the “Other” part.

A look at the last line reveals a practical problem of both the closure and the completion procedures in the presence of missing values. To close a composition with missing elements, we need to guess a value for the missing elements. The computer cannot add up unknown values and thus will close the rest of the

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<sup>3</sup>The different compositional classes will be discussed in Sect. 2.4.

composition to 1, such as if this last components would be absent. However, it is very unlikely that the canned beans do not contain a considerable amount of water.

### 2.1.4 Compositions as Equivalence Classes

Either by reclosing a subcomposition or by completing the composition, the dataset is now compositional in nature, since its columns sum up to a constant. In any way, we homogenized the totals to the same quantity, that is to say: we removed the information on the total amount of each sample. This is licit, because that total is in our kitchen example not relevant to evaluate the nutritive character of the different crops: it is an arbitrary choice of the producer or due to the local unit system. In fact, we could rescale each row of the dataset by multiplying it with a positive value, and the information they convey would not change. We can thus say that two vectors are *compositionally equivalent* (Aitchison, 1997; Barceló-Vidal, 2000) if one is simply some multiple of the other:

$$\mathbf{a} =_A \mathbf{b} \Leftrightarrow \text{exists } s > 0 \text{ for all } i : a_i = sb_i$$

In other words, all vectors with positive entries that are scaled versions of one another represent the same composition. For instance, the following three vectors are the same composition

```
> Amounts[4,]*100/82

      Fat Carbonates Protein Other
corn 1.22      18.29    2.439 78.05

> Amounts[4,]

      Fat Carbonates Protein Other
corn   1         15         2    64

> acomp(Amounts[4,])

      Fat      Carbonates Protein Other
corn 0.0122 0.1829      0.02439 0.7805
attr(,"class")
[1] acomp
```

first giving the composition of corn as portions of 100 g or mass %, then as portions of 82 g, and finally, as relative portions of 1. Note that in the last case, we used the command `acom`, which tells the system to consider its argument as a compositional dataset, implicitly forcing a closure to 1. The different compositional classes (like `acom`, or `rplus` before) are the grounding blocks of “compositions” and will be discussed in detail in Sect. 2.4.

### 2.1.5 Perturbation as a Change of Units

Indeed mass percentage is a quite irrelevant type of quantification for us: we are more interested in the energy intake. But the different nutrients have a different energy content (in kJ/g):

```
> (epm <- acomp(c(Fat=37,Protein=17,Carbonates=17)))

      Fat      Protein Carbonates
0.5211    0.2394      0.2394
attr(,"class")
[1] acomp
```

Since our dataset is not anymore in grams but scaled to unit sum, it is likewise irrelevant in which units the energy contribution is given: the energy per mass can actually be seen again as a composition.

We can now transform our mass percentage subcomposition  $coi$  of nutrients to a energy composition  $ec$  by scaling each entry with the corresponding energy per mass composition and reclosing. For example, for the first line of the dataset:

$$\begin{aligned}
 (ec_{1i})_{i=1,\dots,n} &= \mathcal{C}(coi_{1i}epm_i)_{i=1,\dots,n} \\
 &= \frac{(23.33 \cdot 0.52, 33.33 \cdot 0.24, 43.33 \cdot 0.24)^t}{23.33 \cdot 0.52 + 33.33 \cdot 0.24 + 43.33 \cdot 0.24} \\
 &= (0.4, 0.26, 0.34)^t
 \end{aligned}$$

This operation is called *perturbation*. It is denoted by a  $\oplus$  sign:

$$ec = coi \oplus epm$$

In R, it is done simply by *adding* two `acom` objects:

```
> (ec <- acomp(coi)+epm)

      Fat      Carbonates Protein
soy    0.39846 0.2615      0.3400
peas   0.07293 0.6813      0.2457
wheat  0.02555 0.7044      0.2700
corn   0.11350 0.7822      0.1043
beans  0.02526 0.6962      0.2785
attr(,"class")
[1] acomp
```

Now,  $ec$  contains the composition of my ingredients measured in terms of portion of energy rather than in mass. Perturbation is considered as the fundamental operation in compositional data analysis, equally important as the vector addition is in the usual real vector space  $\mathbb{R}^D$ . In an abstract mathematical view, perturbation is

indeed the *addition* in a vector space structure. Section 2.5 formally introduces this operation and the properties of this vector space structure.

### 2.1.6 Amalgamation

A typical operation when dealing with compositional data is to amalgamate some of the variables into a single component. For instance, in the kitchen data reported before, fat was given as a single component. Though the original information gave the proportions of saturated and unsaturated fats,

```
> fatDataset
      total.Fat  sat.Fat  unsat.Fat
soy           7      1.40      5.60
peas          3      0.60      2.40
wheat         1      0.25      0.75
corn          1      0.32      0.68
beans         0.5      0.10      0.10
```

we amalgamated them into the total fats. Amalgamation, though very commonly done, is a quite dangerous manipulation: a fair amount of information is lost in a way that we may find ourselves unable to further work with the amalgamated dataset. For instance, if saturated and unsaturated fats have different energy content, we cannot compute the total fat mass proportion from the total fat mass energy content, or vice-versa. Amalgamation thus should only be applied in the “definition of the problem” stage, when choosing which variables will be considered and in which units. It should be meaningful in the units we are dealing with, because afterwards, we will not be able to change them. And it should have a deep connection with our questions: to amalgamate apples and pears may be meaningful when investigating the nutrition habits of one population (where an apple may be equivalent to a pear) and catastrophic when studying the importance of some fruits in several cultures (where we just want to see different patterns of preference of fruits). Once the parts are defined, the units chosen, and the questions posed, we should not amalgamate a variable anymore.

The easiest way to amalgamate data in **R** is to explicitly compute the amalgamated components as totals of an unclosed amount dataset (of an amount class, “rplus” or “aplus”) containing only these components:

```
> fatDataset$total.Fat = totals(plus(fatDataset,
                                     c("sat.Fat", "unsat.Fat")))$
```

The `plus` command works in the same way as `comp` but explicitly states that the total is still relevant and no reclosure should be performed. It is used to assign the scale of a dataset of amounts in relative geometry. Amalgamation is further explained in Sect. 3.3.1.

### 2.1.7 Missing Values and Outliers

Note that the total amount for beans is missing (the nutrition table was reported for a 250 mL can of this product). We could have more missing values in the dataset by considering potassium, which is only given for some of the products, probably those for which it is relevant. However, it would be naive to assume that there is no potassium in any of these products.

Much more than in classical multivariate analysis, missing values in compositional data need a careful treatment, especially because a missing value in a single component makes it impossible to close or complete the data, such that finally, the actual portion of no single component is known. It will be thus very important to understand which procedures are still valid in the presence of missing values and how others might be (partially) mended. The same applies to many other kinds of irregularities in compositional data, like measurements with errors, atypical values, or values below the detection limit: in all these cases, the total sum will either be unknown or affected by an error, which will propagate to all the variables by the closure. The solution comes by realizing that whichever this total value might be, the (log)ratios between any regular components are unaffected.

The “compositions” package will thus close the non-missing parts as if the missing parts are 0, knowing that this value will not affect any proper analysis.

## 2.2 Principles of Compositional Analysis

Any objective statistical methodology should give equivalent results, when applied to two datasets which only differ by irrelevant details. The following four sections present four manipulations that should be irrelevant for compositions and that thus generate four invariance principles, with far-ranging consequences for the mathematics underpinning compositional data analysis (Aitchison, 1986).

### 2.2.1 Scaling Invariance

For the theoretical approach taken in this book, we will consider a vector as a composition whenever its components *represent the relative weight or importance of a set of parts forming a whole*. In this case, the *size* or total weight of that whole is irrelevant. One can then remove that apparent influence by forcing the data vectors to share the same total sum  $\kappa$  with the *closure operation*,

$$\mathcal{C}(\mathbf{x}) = \frac{\kappa \cdot \mathbf{x}}{\mathbf{1}' \cdot \mathbf{x}}. \quad (2.1)$$

The components  $x_i$  of a closed composition  $\mathbf{x} = (x_1, x_2, \dots, x_D)$  are values in the interval  $(0, 1)$  and will be referred to as *portions* throughout this book. The term *proportions* will be reserved for ratios and relative sizes of components.



By applying the closure, we can compare data from samples of different sizes, e.g., the beans, corn, and wheat in our simple kitchen example (even though their compositions were respectively reported for 100 g, 82 g, and 250 mL of product); a sand/silt/clay composition of two sediment samples of 100 g and 500 g; and the proportion of votes to three parties in two electoral districts of  $10^6$  and  $10^7$  inhabitants. For instance, the vectors  $\mathbf{a} = [12, 3, 4]$ ,  $\mathbf{b} = [2400, 600, 800]$ ,  $\mathbf{c} = [12/17, 3/17, 4/17]$ , and  $\mathbf{d} = [12/13, 3/13, 4/13]$  represent all the same composition, as the relative importance (the *ratios*) between their components is the same. A sensible compositional analysis should therefore provide the same answer independently of the value of  $\kappa$  or even independently of whether the closure was applied or else the data vectors sum up to different values. We say the analysis will be then *scaling invariant*. Aitchison (1986) already showed that *all* scaling-invariant functions of a composition can be expressed as functions of log ratios  $\ln(x_i/x_j)$ .

### 2.2.2 Perturbation Invariance

Compositional data can be presented in many different units, and even when given in portions, it is still relevant in which physical quantities the components were originally measured: g, tons, mass %,  $\text{cm}^3$ , vol. %, mols, molalities, ppm, ppb, partial pressure, energy, electric loading, money, time, persons, events, cells, mineral grains, etc. Even if it is clearly defined what substance we are measuring, how it is quantified is still a choice of the experimenter. It is extremely rare to find a composition where only one single type of quantification is meaningful: in most of the cases, several units could be equally chosen. And since different components are typically also qualitatively different, they might have different interesting properties. For instance, to change a mass proportion to a volume percentage, we need the densities of all components, which will be typically different. The same applies to changing prices between different valuations (from market price to book value, or to buying price, etc.), passing nutrition masses to energy intake, or cell type to energy consumption or biomass production. Thus, different analysts would end up having different values representing exactly the same compositions just due to a different choice of units.

It is evident that the statistical analyses we can apply should thus give the same qualitative results regardless of the units chosen, as long as they contain the same information, i.e., as long as we can transform the units into each other, e.g., by rescaling each component by its density. This operation is done by a perturbation with a composition containing as entries the conversion factors (e.g., the densities) for each component. We would thus require a meaningful compositional analysis to give the same results (obviously, up to a change of units) when applied to a dataset perturbed by this vector. But since we never know which type of quantifications might else be considered, we should request invariance of the analysis with respect to perturbation with all possible weighting factors.

This principle has the same level as the principle of translation invariance for real data. There, we would expect that, e.g., when we represent electric (or gravity) potentials with respect to a different reference level, we would still get the same results (up to that choice of reference level).

This principle is particularly critical when dealing with datasets of mixed units. Often different components are measured in different units. For instance, trace elements are often measured in ppm, while others are given in mg, or even kg, and fluids might be quantified in volume proportions or fugacities. Unfortunately, any method not honoring perturbation invariance would give completely arbitrary results when the units do not match. On the other hand, when we demand perturbation invariance, all such data could be meaningfully analyzed in a common framework, as long as there exists a perturbation bringing them into the same system of units, even if we do not know it.

### 2.2.3 *Subcompositional Coherence*

Sometimes, only a subset of the initial parts is useful for a particular application, and one works with the corresponding *subcomposition*, by reclosing the vector of the chosen components. Subcompositions play the same role with respect to compositions as marginals do in conventional real multivariate analysis: they represent subspaces of lower dimension where data can be projected for inspection. This has several implications jointly referred to as *subcompositional coherence* [Aitchison \(1986\)](#): three examples of these implications follow.

First, if we measure the distance between two  $D$ -part compositions, this must be higher when measured with all  $D$  components than when measured in any subcomposition.

Second, the total dispersion of a  $D$ -part compositional dataset must be higher than the dispersion in any subcomposition.

Finally, if we fitted a meaningful model to a  $D$ -part compositional dataset, the result should not change if we include a new non-informative (e.g., random) component and work with the resulting  $(D + 1)$ -part composition.

*Remark 2.1 (Incoherence of classical covariance).* The classical definitions of covariance and correlation coefficient are not subcompositionally coherent. This is connected to two problems: the *spurious correlation* and the *negative bias*, first identified by [Pearson \(1897\)](#) and later rediscovered by [Chayes \(1960\)](#). The spurious correlation problems states that the correlation between ratios with common denominators (e.g., two components of a closed composition) is arbitrary to an uncertain extent. The negative bias problem appears because each row or column of the covariance matrix of a closed composition sums up to zero: given that the variances are always positive, this implies that

some covariances are forced towards negative values, not due to any incompatibility process but because of the closure itself. For instance, if we compute the correlation coefficient between MnO and MgO content in a compositional dataset of glacial sediment geochemistry,<sup>4</sup> we may obtain 0.95 if we use the whole 10-part composition or  $-0.726$  if we use only the subcomposition of elements not related to feldspar (P–Mn–Mg–Fe–Ti). Given the fundamental role of covariance in statistics, it is not a surprise that there exists a full series of papers on detecting, cataloging, and trying to circumvent the spurious correlation. An account can be found in [Aitchison and Egozcue \(2005\)](#).

### 2.2.4 *Permutation Invariance*

Last but not least, results of any analysis should not depend on the sequence in which the components are given in the dataset. This might seem self-evident, but it is surprising how many methods happen to violate it. For instance, one of the apparent bypasses of the constant sum of covariance matrices (Remark 2.1) was to remove the last component of the dataset: in that way, the dataset was not summing up to a constant anymore, and the covariance matrix was apparently free. That “solution” was obviously not permutation invariant (and even more, it was not a solution, as correlations had exactly the same values, being thus equally forced towards negative spurious correlations).

For the log-ratio approach, it is also a very important principle, when, e.g., we ask which methods can be meaningfully applied to coordinates of compositional data. For instance, a naive Euclidean distance of alr transformed data<sup>5</sup> is not permutation invariant and should thus not be used, e.g., for cluster analysis. We would otherwise risk to have different clusterings depending on which was the last variable in the dataset. This will be further discussed in Remark 2.2.

## 2.3 Elementary Compositional Graphics

Compositional datasets are typically represented in four different ways: as Harker diagrams (scatterplots of two components), as ternary diagrams (closed scatterplots of three components), as scatterplots of log ratios of several parts, or as sequences of bar plots or pie plots. By far, the most common ways are the first two.

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<sup>4</sup>This dataset can be found on <http://www.stat.boogaart.de/compositionsRBook>.

<sup>5</sup>The alr (additive log ratio) transformation was the fundamental transformation in [Aitchison \(1986\)](#) approach, and it is discussed in detail in Sect. 2.5.7.

These compositional graphics will be illustrated with a dataset of geochemistry of glacial sediments (Tolosana-Delgado and von Eynatten, 2010). The dataset is available on the book home page<sup>6</sup>:

```
> GeoChemSed=read.csv("geochemsed.csv",
                      header=TRUE, skip=1)[-c(3,14,31,32)]
> names(GeoChemSed)

[1] "Sample" "GS"      "SiO2"    "TiO2"    "Al2O3"   "MnO"
[7] "MgO"    "CaO"     "Na2O"    "K2O"     "P2O5"    "Fe2O3t"
[13] "Ba"     "Co"      "Cr"      "Cu"      "Ga"      "Nb"
[19] "Ni"     "Pb"      "Rb"      "Sc"      "Sr"      "V"
[25] "Y"      "Zn"      "Zr"      "Nd"
```

### 2.3.1 Sense and Nonsense of Scatterplots of Components

*Harker diagram* is the name given in geochemistry to a conventional scatterplot of two components, *without any transformation applied to them*. For this reason, they may highlight any additive relationship between the variables plotted: e.g., in the case of plotting two chemical components of a dataset evolving in several stages, Harker diagrams visually represent mass balance computations between the several stages (Cortes, 2009). Unfortunately, these diagrams are neither scaling nor perturbation invariant and not subcompositionally coherent (Aitchison and Egozcue, 2005): there is no guarantee that the plot of a closed subcomposition exhibits similar or even compatible patterns with the plot of the original dataset, even if the parts not included in the subcomposition are irrelevant for the process being studied. This is actually the spurious correlation problem; thus, a regression line drawn in such a plot cannot be trusted, in general terms. Figure 2.1 shows two Harker diagrams of the same components in the same dataset, exhibiting the subcompositional incoherence of this representation. One can obtain a Harker diagram by using the standard function `plot(x,y)`.

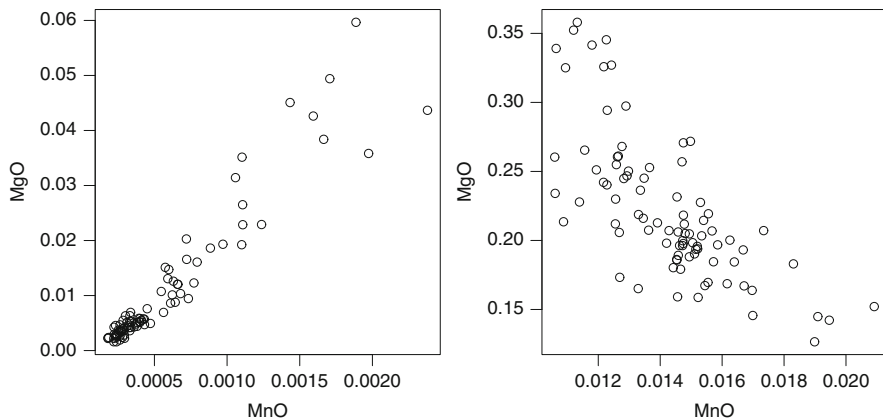
```
> par(mfrow=c(1,2), mar=c(4,4,0.5,0.5))
> plot( clo(GeoChemSed[,3:12])[,c(4,5)])
> plot( clo(GeoChemSed[,c(4,6,7,11,12)])[,c(2,3)])
```

### 2.3.2 Ternary Diagrams

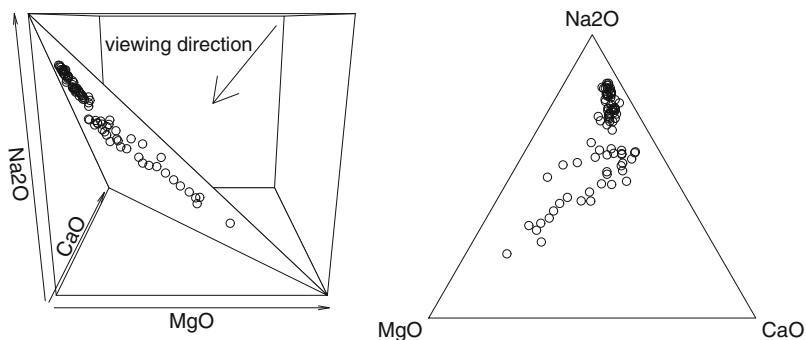
*Ternary diagrams* are similar to scatterplots but display a closed three-part sub-compositions. If we would plot the three components of a closed composition in a three-dimensional scatterplot, all points would lie in a planar triangle spanned

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<sup>6</sup><http://www.stat.boogaart.de/compositionsRBook>.



**Fig. 2.1** Examples of Harker diagrams on the sediment geochemistry dataset. (*Left*) Using the full dataset of major oxides. (*Right*) Using the subcomposition P–Mn–Mg–Fe–Ti

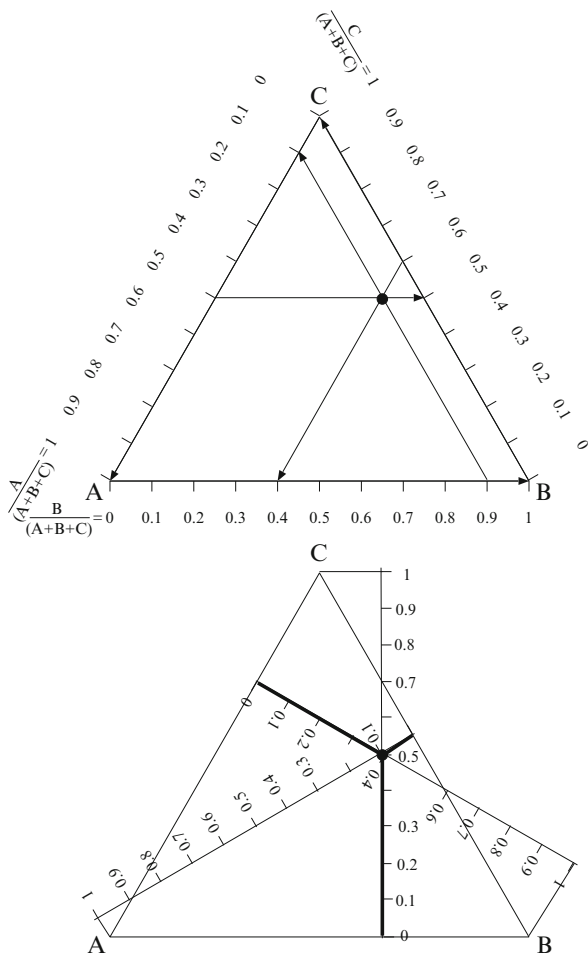


**Fig. 2.2** Example of a ternary diagram embedded in its original three-dimensional space

by the three points  $(1, 0, 0)$ ,  $(0, 1, 0)$ , and  $(0, 0, 1)$  as corners. The ternary diagram displays this triangle in the drawing plane with the corners annotated by the axis on which this corner lies. This is possible since closed datasets of three parts have only 2 degrees of freedom. Figure 2.2 illustrates this with the MgO, CaO, and Na<sub>2</sub>O subcomposition of the example dataset.

For the interpretation of ternary diagrams, we can make use of the property that the orthogonal segments joining a point with the three sides of an equilateral triangle (the *heights* of that point) have always the same total length: the length of each segment is taken as the proportion of a given part. Ternary diagrams have the merit of actually representing the data as what they are: compositional and relative. Geoscientists are particularly used to this kind of representation. Figures 2.3 and 2.4 show several ways in which the relative portions and proportions of the components of a three-part composition can be read from a ternary diagram.

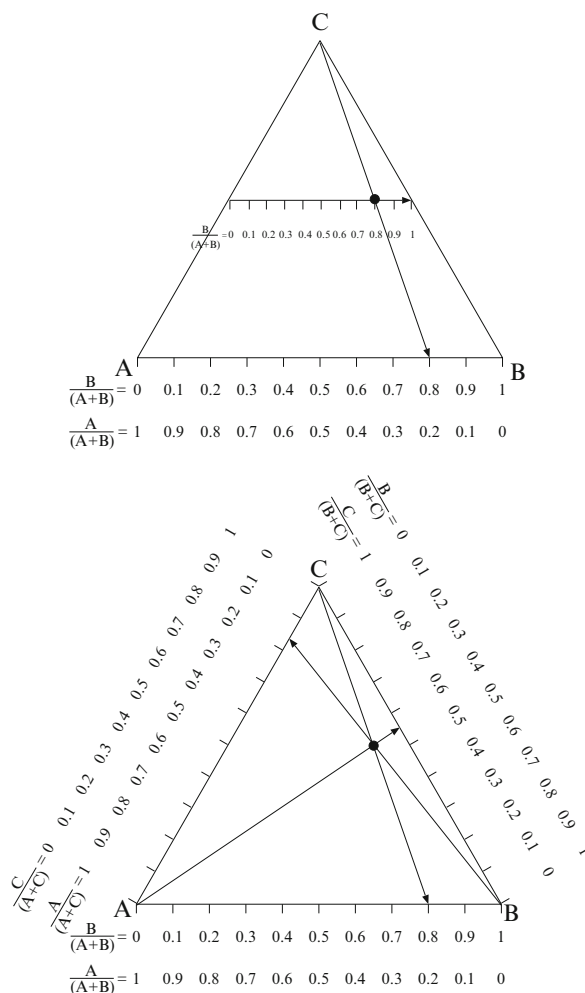
It is worth mentioning that when the three parts represented have too different magnitudes, data tend to collapse on a border or a vertex, obscuring the structure:



**Fig. 2.3** In a ternary diagram, the portion of each part is represented by the portion of the height of the point over the side of the triangle opposite to the part on the total height of the triangle. However, it is difficult to judge height visually. If we assume the three sides of the triangle to form a sequence of axes, we can make a projection of the point onto the axis pointing towards the interesting part parallel to the previous axis. Consequently, a composition on an edge of the triangle will only contain the two parts connected by the side, and a composition in a vertex of the triangle will only contain the part the vertex is linked to. Thus, all compositions along a line parallel to an axis have the same portion of the part opposite to the axis

a typical practice here is to multiply each part by a constant and reclose the result. One can obtain a ternary diagram in **R** by using `plot(x)`, with `x` an object of any compositional class (`acomp` or `rcomp`, see next section for details).

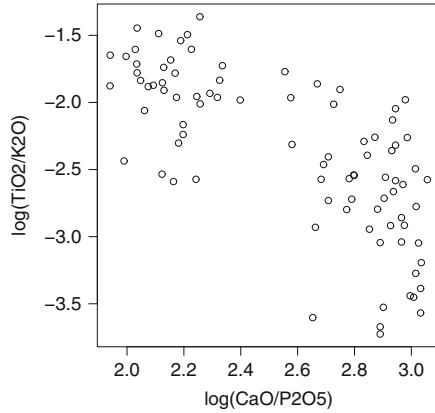
```
> xc = acomp(GeoChemSed, c("MgO", "CaO", "Na2O"))
> plot(xc)
```



**Fig. 2.4** The two part subcompositions can be read in two ways: (1) by projecting the composition to an edge of the triangle along a ray coming from the vertex of the part that should be ignored. Thus, all compositions along such rays have the same subcomposition, when the part where the ray is coming from is removed. (2) By drawing a scale between 0 and 1 on a segment parallel to the axis opposite to the part we want to remove, passing through our datum

### 2.3.3 Log-Ratio Scatterplots

*Log-ratio scatterplots* are just scatterplots of the log ratio of two parts against that of two other parts, though sometimes the denominators are the same. This representation is fully coherent with the description of compositions given in Sect. 2.1 but on the other side, does not capture any mass balance between the represented parts. In fact, it is “independent” of such mass balances: whenever we



**Fig. 2.5** Example of a log-ratio scatter plot obtained using the formula interface. This example was obtained with `plot(log(TiO2/K2O) ~ log(CaO/P2O5), GeoChemSed)`. Recall that the formula syntax is  $y \sim x$

cannot ensure that the studied system is not shrinking or growing, then log ratios of scatterplots display almost all the information we really have. For instance, if we are working with transported sediments, we do not know if any part can be dissolved by water and removed from the system; or when dealing with party vote shares between two different elections, we do not know if exactly the same people took part in the election. In fact, most of the datasets obtained in natural systems (meaning, not in the lab under strictly controlled situations) will be in such situations. Despite these issues, scatterplots of log ratios are seldom encountered in applications. To generate a log-ratio scatterplot, one can use the formula interface to `plot(formula, data)`, providing the data set where the variables involved will be extracted, and the desired formula describing the log ratios:

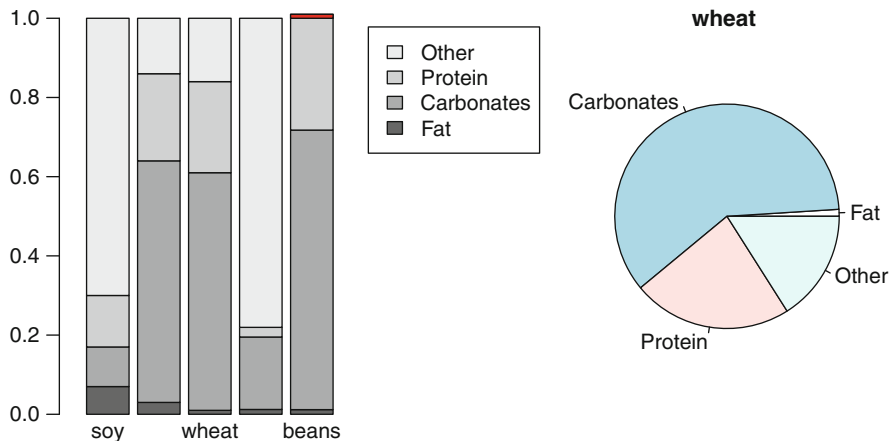
```
> plot(log(var1/var2) ~ log(var3/var4), dataset)
> plot(I(var1/var2) ~ I(var3/var4), dataset, log="xy")
```

Figure 2.5 shows an example obtained with the first syntax. These two syntax examples would only differ in the way axes are labeled, with the transformed values in the first case, with the original values of the ratios in the second case (using the command `I` is needed to tell **R** that the quotient inside it are not symbolic formulae, but actual ratios).

### 2.3.4 Bar Plots and Pie Charts

A *bar plot* is a classical representation where all parts may be simultaneously represented. In a bar plot, one represents the amount of each part in an individual as a bar within a set. Ideally, the bars are stacked, with heights adding to the total of the composition (1 or 100 %). Then, each individual is represented as one of such





**Fig. 2.6** Examples of bar plot (*left*) and pie plot (*right*). Note the presence of an extra segment on top of the bars for beans, showing the presence of a missing value

stacked bars, possibly ordered if that is possible (e.g., in time series). Stacked bars are provided in **R** by the command `barplot(x)`, when `x` is a compositional dataset. Individual compositions can also be displayed in the form of pie charts. Pie charts are produced by the `pie(x)` command, but now `x` must be a single composition (as only one pie diagram will be generated). Pie charts are not recommended for compositions of more than two parts, because the human eye is weak in the comparison of angles if they are not aligned (Bertin, 1967). Figure 2.6 is created by

```
> barplot( acomp(Amounts), xlim=c(0,11) )
> pie( acomp(Amounts["wheat",]), main="wheat")
```

## 2.4 Multivariate Scales

A fundamental property of each variable (or set of variables) in a dataset is its scale. The scale of a measurement describes which values are possible outcomes and which mathematical operations on these values are meaningful in the context of the application. Classical statistical practice distinguishes several scales for single variables, like (e.g., Stevens, 1946): categorical (several values are possible, but they are not comparable), ordinal (several values are possible, and they can be compared by ordering them), interval (any value in a subset of the real line is a possible outcome, absolute differences between them are meaningful), and ratio (any value in a subset of the *positive* real line is a possible outcome, relative differences between them matter). Other authors have criticized this list for being too limited: for instance, it does not include a separation between finite, infinite countable, and infinite uncountable scales (in any case, with interval or ratio differences).

Although a scale is above all a set of possible values together with some meaningful mathematical operations (and not a statistical model), each scale has

typical statistical models associated to them: e.g., the multinomial distribution is the most adequate and general model for categorical data, and the normal distribution appears as a central distribution for the real interval scale.

Compositional data add a further complexity to the scale issues as they are multivariate objects on themselves. In a composition, we cannot treat one isolated variable because its values are only meaningful in relation to the amounts of the other variables. The whole composition has thus to be considered as a unique object with some scale with joint properties, such as summing to 1 or 100 %, being positive, and obeying a ratio law of difference. A compositional scale is thus a new multivariate scale, not just the concatenation of several separate scales for each variable.

Compositional data has historically been treated in several ways according to different considerations on the underlying scale, and there was a lot of discussion on which is the *right* way of doing so. The “compositions” package provides several of these alternative approaches, with the aim of facilitating the comparison of results obtained following each of them. However, from our experience and the considerations presented in the first sections of this chapter, one of these scales should be the preferred one, as it is the one grounded in the mildest hypotheses: this is the [Aitchison \(1986\)](#) geometry for compositions, discussed in Sect. 2.5. Therefore, this book focuses on this scale, and the next section will introduce it in detail. But depending on our view of the problem, in some case, other scales might be more appropriate. They are available in the package and thus all presented and briefly discussed in this section for the sake of completeness. A comprehensive treatment of the multiple scale mechanisms in the package can be found in [van den Boogaart and Tolosana-Delgado \(2008\)](#).

To highlight the relevance of the choice of a scale, in “compositions”, you assign it to the dataset *before any analysis*. This is done by a command like (e.g., `acomp` for Aitchison **Composition**):

```
DataWithScale <- acomp(DataWithoutScale)
```

The scale of each compositional dataset is stored as its S3-class.<sup>7</sup> The package “compositions” then automatically treats the dataset according to the scale chosen. Each row of the dataset is considered as one observation in a compositional scale. The following multivariate scales are available, for both vectors of amounts and vectors of components (or actual compositions):

- “**rmult**”—real **multivariate** scale
- “**rplus**”—real interval restricted to the positive (**plus**) real orthant,  $\mathbb{R}_+^D$
- “**aplus**”—Aitchison (i.e., ratio) geometry in  $\mathbb{R}_+^D$
- “**rcomp**”—real (i.e., interval) **compositional** scale
- “**acomp**”—Aitchison (i.e., ratio) **compositional** scale
- “**ccomp**”—count **compositional** scale

---

<sup>7</sup>An S3-class is a concept in the object model of **R**. The class of an object determines how it is treated by generic functions, such as `plot` or `summary`. Use `help(class)` for further details on the concept in **R**.

Again, consider that the default choice of scale for compositional data should be the “acomp” Aitchison compositional scale.

### 2.4.1 *Classical Multivariate Vectorial Data (rmult)*

The real multivariate scale associated to the class “rmult” is the Cartesian product of  $D$  classical real scales, with values in a vector space  $\mathbb{R}^D$ . *This is neither a compositional nor an amount scale, as here negative values are fully meaningful.* The multivariate normal distribution is its central statistical model. Most well-known multivariate procedures were devised for this scale, such as cluster analysis, mean vector and variance–covariance matrix, principal component analysis, factor analysis, multivariate regression, and multivariate analysis of variance. Because it is the “conventional” scale, the package “compositions” only provides a limited functionality within it. It is mainly intended as a back-end for procedures based on treating the other scales through transformations. An example dataset displaying this scale might be the locations of earthquakes near Fiji Islands, data included in a standard **R** distribution:

```
> data(quakes)
> MultivariateData <- rmult(quakes[,c("lat", "long", "depth")])
```

### 2.4.2 *Positive Data with Absolute Geometry (rplus)*

Historically, the oldest approach of working with vectors of amounts is to see them as real multivariate data restricted to the positive orthant  $\mathbb{R}_+^D$ . Methods based on this approach are typically not scaling invariant, since each data is weighted proportional to its total amount (if nothing nastier happens). An associated distribution could be a truncated multivariate normal distribution. The way in which values of our sample can be changed is by simple addition or subtraction of some components, as long as the results remain positive. Analysis within the “rplus” framework is quite simple: apply the standard procedures from a multivariate scale to a dataset of positive values and hope for the best. It is thus easiest to analyze data in the “rplus” scale, as all statistical packages may be used for that end. However, several problems arise when interpreting results: for instance, in regression analysis, we will likely obtain negative predictions for an amount; if any procedure is based on assuming a normal distribution and our data have low means and high variances, truncation will be a source of problems; and last, but not least, if we are working with a vector of amounts with bounded sum (for instance, an incomplete composition), then correlation coefficients cannot be interpreted in the way we are used to, as they are spurious to an unknown extent (see Remark 2.1). In summary, the “rplus” scale is probably the most dangerous of all, because it is easily accessible in a conventional approach, but interpreting the results is a path full of pitfalls. An example from **R** illustration datasets might be the phosphor contents separated according to the

inorganic and organic part of the soil in an experiment analyzing the nutrient content of the resulting crop:

```
> data(phosphor)
> x <- rplus(phosphor, c("inorg", "organic"))
```

### 2.4.3 Positive Data with Relative Geometry (*aplus*)

[Stevens \(1946\)](#) introduced the ratio scale to capture the notion that for a positive observational variable, only its relative variations might be meaningful. For several reasons, it is common to analyze such data after a log transformation. In this case, the reference distribution of statistical analysis becomes the lognormal distribution, and the natural way of describing how a variable evolves is through products/divisions. Later on, this idea has been extended to multivariate positive datasets (e.g. [Pawlowsky-Glahn et al., 2003](#)), giving rise to the “*aplus*” scale. The idea behind the approach is that the amount of each component can be individually analyzed in a relative scale, in the sense of ratio scale ([Stevens, 1946](#)), i.e., with component-wise product/division and distances computed based on the log-transformed data. Therefore, the natural central distribution in this case should be the multivariate lognormal. Though this is a common alternative to a full compositional approach, the analyst should be aware that most methods based on this approach are not scaling invariant.

An example might be the measures of morphology of iris flowers (length and width of petal and sepal), which are positive quantities with a relative geometry but where total size matters (since it discriminates between species):

```
> data(iris)
> x <- aplus(iris[, -1])
```

### 2.4.4 Compositional Data with Absolute Geometry (*rcomp*)

The most widely used way of treating compositional data is *as if they were* just multivariate real datasets that incidentally happened to be positive and sum to 1 or 100%, but where these circumstances were irrelevant. Methods based on this approach are typically neither scaling nor perturbation invariant and definitively not subcompositionally coherent. A useful reference distribution for a  $D$ -part composition would be then a degenerate,  $(D - 1)$ -dimensional multivariate normal distribution restricted to have all components positive and summing up to the constant. In this context, all changes between samples must be characterized by vectors of components summing up to *zero*, thus representing pure transference of “mass” between the parts.

Even though this naive approach was dominant, some special procedures were developed apart from the classical multivariate statistical toolbox, like the representation in ternary diagrams, and it was long warned that classical methods gave

meaningless results. Awareness of these problems stemmed from the Mathematical Geosciences community: problems linked to the spurious correlation (as it appeared already in the “rplus” scale) were reported for correlation and covariance (Butler, 1979; Chayes, 1960), principal components (Chayes and Trochimczyk, 1978), cluster analysis (Butler, 1978), autocorrelation (Pawlowsky-Glahn, 1984), etc. In fact, it was even shown that this kind of analysis does not obey the important principles of a meaningful statistical analysis of Sect. 2.2 (Aitchison, 1986). Nevertheless, the “rcomp” scale is still probably the most used of all scales, because of the same reasons that make the “rplus” scale dominant: the availability of software doing the job and the lack of awareness on the pitfalls of its interpretation.

But to be fair, we must also take into account that this scale honors a very important law of natural sciences: *mass balance* (Cortes, 2009; Shurtz, 2003), i.e., the fact that in a closed system (one where mass neither enters nor leaves the system), the total mass amount we have before and after any process must be preserved. This is most adequately captured by the vectors of zero sum describing change in this scale. Note nevertheless that the hypothesis of system closeness is a very strong one, that cannot come from the compositional datasets themselves, but from external, complementary information (Aitchison, 1986; Cortes, 2009). Furthermore, in the case that one wants to apply mass, matter, or volume preservation, the units of the dataset should be consistently masses, moles, or volumes. In this context, most changes of units would be nonsense.

### 2.4.5 Compositional Data with Aitchison Geometry (*acomp*)

In the early 1980s, Aitchison (1981, 1986) realized that in order to obtain meaningful results in a general framework, some basic principles had to be honored by any statistical analysis of compositional data. These principles have been stated in Sect. 2.2: scaling invariance, perturbation invariance, permutation invariance, and subcompositional coherence. He then showed that these principles imply a relative geometry, in which only the ratios of different components actually matter. In this context, a meaningful change between compositions has to be measured as a *perturbation* (as introduced in page 18, and more formally defined in Sect. 2.5), and the reference distribution becomes the additive-logistic-normal distribution, or normal distribution on the simplex (Chap. 3).

It was not until some years later that it was realized that the mathematical structure Aitchison had defined is indeed a vector space structure in its own right, isometrically equivalent to  $\mathbb{R}^{D-1}$  (Barceló-Vidal, 2000; Billheimer et al., 2001; Pawlowsky-Glahn and Egozcue, 2001). A logical consequence of this equivalence is that we can convert any statistical problem involving compositions of  $D$  parts onto a classical multivariate problem involving real vectors of  $(D - 1)$  coordinates. This is what Pawlowsky-Glahn (2003) called “the principle of working in coordinates”: whenever your scale is described by an Euclidean space structure, you automatically honor that scale if you statistically analyze the coordinates of the dataset with respect to any orthonormal basis (instead of treating the raw data). Section 2.5

presents in more detail this Euclidean space structure and the implications of the principle of working on coordinates for compositions. Note that the same principle of working on coordinates justifies analyzing log-transformed vectors of amounts if their scale is “aplus” (Pawlowsky-Glahn et al., 2003).

### 2.4.6 Count Compositions (*ccomp*)

Sometimes it is not possible to directly observe the “true” composition of a system but rather how many individuals belong to each of the groups in which this system is divided. We assume that the counts are proportional to the sizes of the groups. We call the result a *count composition*, because its elements are counts. A typical example are species compositions, such as how many insects of each taxon are found in an insect trap or how many people with blue, green, brown, and gray eyes are found in a classroom. The most apparent difference between both “*acomp*” or “*rcomp*” against “*ccomp*” scales is that the former are vectors of real numbers (thus can be called *real compositions*), whereas the latter are vectors of integers.

A typical model for count compositions is that conditioned to a fix total number of counts, the actual counts of each class are the outcomes of multinomial distribution with probabilities equal to the unknown, “true” composition, which we call the *underlying composition* (most probably, of “*acomp*” type).

Count compositions can be seen as an indirect observation of Aitchison compositions. However, this scale has some particular properties unlike the Aitchison compositional scale, as it has a dual nature. On one side, count compositions follow a relative scale, given that the actual count total (the number of insects in the trap, the number of people in the classroom) is irrelevant to understand the underlying process or distribution. But on another side, the statistical sampling error strongly depends on this total number of counts. Grossly simplifying, we can say that the total is irrelevant for the mean behavior of our system, but it matters when characterizing the variability of our observations and the uncertainty on our estimations. For example, in count compositions, a value of zero is easily possible as a random result if the corresponding component in the underlying composition is small (but not necessarily zero).

There are some practical problems with count compositions. For instance, plotting should take care to somehow represent how many times each of the possible count compositions was observed. There exist some competing models for count compositions (being log-linear models the most relevant). However, in our view, count compositions can be better fundamentally understood based on the properties of Aitchison compositions.

### 2.4.7 Practical Considerations on Scale Selection

This book is mainly devoted to the Aitchison composition scale (“*acomp*”). The other scales in the package are only discussed for background information.

However, the choice of a scale should not be driven by the goals of this book, but for an assessment of which is the meaningful scale for a problem at hand. This is necessarily a subjective decision that the analyst should thoroughly ground. Here are some guidelines.

One can select the right scale for the analysis based on the answers to three questions:

- *Are the components counts?*

Many compositions are physically measured as counts, integer values, even if that is not relevant. For instance, many geochemical compositions are derived from counting how many atoms of a given type hit the sensor: but there are so many of them that we may safely consider that composition as a real composition. The same happens to the number of votes given in large electoral districts to the parties entering the parliament. On the other hand, when counting mineral or insect species, or the votes of all parties in a small district, the total is typically small, and we may be in a count composition case. Thus, the question must be stated more precisely: if this composition is formed by counts, *is their total so small that the discrete random selection error in the counts is important?* If the answer is *yes*, we are within a count composition problem.

- *Is the total mass in each observation an important aspect of the process under investigation?*

If the answer is *yes*, the dataset should be treated with an amount scale, i.e., “*aplus*”, “*rplus*” (if all variables are positive), or “*rmult*” (if they form a real multivariate dataset). If the answer is *no*, the dataset should be considered of a compositional scale, i.e., “*acomp*”, “*rcomp*”, or “*ccomp*”.

- *Is the question of our focus only meaningful if the composition is expressed in a specific, common set of units? Does the composition contain all possible, potentially interesting components?*

If the answer to *both* questions is *yes*, then the principles of scaling or perturbation invariance and subcompositional coherence are not necessary, and scales of an absolute type would be reasonable, e.g., “*rplus*” or “*rcomp*”. This would imply that the structure under investigation is expected to be linear in real values of the components (e.g., a law of mass conservation). If the answer is *no* to any of these questions, then at least one of the principles is relevant, and we should only use relative scales (“*aplus*”, “*acomp*”, or “*ccomp*”) to honor them.

*Example 2.1 (Discussing the scale of a problem).* The package “*compositions*” contains several example datasets for illustration that can be loaded with the command `data`. One of them contains chemical analyses of samples of water from rivers and streams of a medium-sized Mediterranean river basin from western Spain, including major (Na, Ca, Mg, K, Cl, SO<sub>4</sub>, HCO<sub>3</sub>) and minor ions (Ba, Sr) coming from the geological background, as well as other linked to urban sewage, industrial, or fertilizer pollution (total organic Carbon, H, NO<sub>3</sub>, PO<sub>4</sub>). Note that the data do not sum

up to a constant value. Now we can ask ourselves several questions about this dataset, leading to different choices of scales.

- An obvious question is which water samples are cleaner, or in other words, which carry a lower ionic concentration. It is obvious that in this case the total mass in each observation matters. It is also important which units do we choose to tackle with the problem, because we are going to obtain different answers when expressing our components in mol/L or in g/L. But, on the other hand, there are potentially thousands of pollutants that this dataset did not report. Thus, we should work with a scale of amounts, but the choice between “rplus” and “aplus” is open.
- A second question in this line would be which are the most important pollutants. In this case, the total amount still matters, but the units in which we are working (whether mol/L or g/L) do not: now it is more important to know the enrichment of each component on each sample in relation to a background value (for instance, the average); ammonia, for instance, being always present in small proportions, would not qualify as important for the first question but becomes a key element to answer the second question, because in polluted samples, it can be up to 150 times larger than in the background. Thus, we need a relative scale for amounts, the “applus” one.
- Finally, we can also ask ourselves which are the geological factors conditioning the water chemistry. In this case, the total amount does not actually matter, as it is rather a function of dilution (how much water circulates) instead of the geology of our basin. Moreover, it is not good that our answer depends on the units we use. We should therefore treat the samples with an “acomp” scale. In fact, this sort of questions is usually answered by hydrogeochemists using the so-called Piper plot, a joint representation of four ternary diagrams representing the subcomposition of major ions in a “rcomp” scale.

In any case, once we have clear in mind our question and selected a scale, e.g., that of an Aitchison composition (“acomp”), you have to assign it to the dataset, as its S3-class:

```
> data(Hydrochem)
> dat <- acomp(Hydrochem[,6:19])
> class(dat)

[1] "acomp"
```

Further analysis of `dat` will then be automatically performed in a way consistent with the chosen compositional scale.



## 2.5 The Aitchison Simplex

### 2.5.1 The Simplex and the Closure Operation

A composition which elements sum up to a constant is called a *closed* composition, e.g., resulting from applying the closure operation (2.1). With “compositions”, you can close a vector or a dataset using the command `clo`. The set of possible closed compositions

$$\begin{aligned}\mathbb{S}^D &:= \left\{ \mathbf{x} = (x_i)_{i=1,\dots,D} : x_i \geq 0, \sum_{i=1}^D x_i = 1 \right\} \\ &= \left\{ \mathbf{x} = (x_1, \dots, x_D) : x_i \geq 0, \sum_{i=1}^D x_i = 1 \right\}\end{aligned}$$

is called the *D*-part *simplex*. Throughout the book, notations  $\mathbf{x}$ ,  $(x_i)_i$ ,  $(x_1, \dots, x_D)$  and  $(x_i)_{i=1,\dots,D}$  will equivalently be used to denote a vector, in the two last cases showing how many components it has. The first two notations are reserved for cases where the number of components is obvious from the context or not important.

There is some discussion on whether the parts should be allowed to have a zero portion ( $x_i = 0$ ) or not. For the scope of this book, zero values will be considered special cases, pretty similar to missing values (see Chap. 7).

The simplex of  $D = 3$  parts is represented as a ternary diagram, where the three components are projected in barycentric coordinates (Sect. 2.3.2). In the same way, the simplex of  $D = 4$  parts can be represented by a solid, regular tetrahedron, where each possible 3-part composition is represented on one side of the tetrahedron. To represent a given 3-part subcomposition of a 4-part dataset, we can project every 4-part datum inside the tetrahedron onto the desired side with projection lines passing through the opposite vertex, corresponding to the removed part. This is exactly the same as closing the subcomposition of the three parts kept. With help of that image, we can conceive higher-dimensional simplexes as hyper-tetrahedrons which projections onto 3- or 4-part sub-simplexes give ternary diagrams and tetrahedral diagrams.

### 2.5.2 Perturbation as Compositional Sum

The classical algebraic/geometric operations (addition/translation, product/scaling, scalar product/orthogonal projection, Euclidean distance) used to deal with conventional real vectors are neither subcompositionally coherent nor scaling invariant. As an alternative, [Aitchison \(1986\)](#) introduced a set of operations to replace these conventional ones in compositional geometry.

*Perturbation* plays the role of sum or translation and is a closed component-wise product of the compositions involved:

$$\mathbf{z} = \mathbf{x} \oplus \mathbf{y} = \mathcal{C}[x_1 \cdot y_1, \dots, x_D \cdot y_D]. \quad (2.2)$$

With “compositions”, one can perturb a pair of compositions in two ways: by using the command `perturbe(x,y)` or by *adding or subtracting* two vectors of class `acomp`, i.e.,

```
> (x = acomp(c(1,2,3)))
[1] 0.1667 0.3333 0.5000
attr("class")
[1] acomp

> (y = acomp(c(1,2,1)))
[1] 0.25 0.50 0.25
attr("class")
[1] acomp

> (z = x+y)
[1] 0.125 0.500 0.375
attr("class")
[1] acomp
```

The neutral element of perturbation is the composition  $\mathbb{1} = [1/D, \dots, 1/D]$ . It is an easy exercise to show that for any composition  $\mathbf{x}$ , the neutral elements fulfill  $\mathbf{x} \oplus \mathbb{1} = \mathbf{x}$ . The neutral element is thus playing the role of the zero vector. Any composition  $\mathbf{x} = [x_1, \dots, x_D]$  has its inverse composition: if one perturbs both, the result is always the neutral element. The inverse is  $\ominus \mathbf{x} = \mathcal{C}[1/x_1, \dots, 1/x_D]$  and it holds  $\mathbf{x} \oplus \ominus \mathbf{x} = \mathbb{1}$ . Perturbation by the opposite element plays the role of subtraction, hence the notation with  $\ominus$ . Perturbation with an inverse composition can as usually also be denoted with a binary  $\ominus$  operator:  $\mathbf{x} \ominus \mathbf{y} := \mathbf{x} \oplus \ominus \mathbf{y}$ . In “compositions”, inverse perturbation can be obtained subtracting two compositions,  $\mathbf{x}-\mathbf{y}$  or with `perturbe(x,-y)`.

*Example 2.2 (How and why do we perturb a whole dataset?).* To perturb a whole compositional dataset by the same composition, we have to give both objects an `acomp` class and just ask for the dataset plus (or minus) the composition, for instance,

```
> data(Hydrochem)
> Xmas = Hydrochem[,c("Cl", "HCO3", "SO4")]
> Xmas = acomp(Xmas)
> mw = c(35.453, 61.017, 96.063)
> mw = acomp(mw)
> Xmol = Xmas-mw
```

`Xmas` contains the subcomposition  $\text{Cl}^- - \text{HCO}_3^- - \text{SO}_4^{=}$  in mass proportions, and this is recasted to molar proportions in `Xmol` by dividing each component (*inverse perturbation*) with its molar weight (in `mw`). Most changes of compositional units may be expressed as a perturbation (between molar proportions, mass percentages, volumetric proportions, molality, ppm, etc.), as explained in Sect. 2.2.2. This operation is also applicable in the centering procedure of Sect. 4.1. Note that the two involved objects must have either the same number of rows (resulting in the perturbation of the first row of each dataset, the second of each dataset, etc.) or one of them must have only one row (resulting in all rows of the other object perturbed by this one).

In statistical analysis, it is often necessary to perturb or “sum up” all the compositions in the dataset. This is denoted by a big  $\oplus$ :

$$\bigoplus_{i=1}^n \mathbf{x}_i := \mathbf{x}_1 \oplus \mathbf{x}_2 \oplus \cdots \oplus \mathbf{x}_n$$

### 2.5.3 Powering as Compositional Scalar Multiplication

*Powering* or *power transformation* replaces the product of a vector by a scalar (geometrically, this is equivalent to scaling) and is defined as the closed powering of the components by a given scalar:

$$\mathbf{z} = \lambda \odot \mathbf{x} = \mathcal{C}[x_1^\lambda, \dots, x_D^\lambda]. \quad (2.3)$$

Again, the package offers two ways to power a composition by a scalar: with the generic function `power(x, y)` or by *multiplying* a scalar by a vector of class `acomp`.

```
> 2*y
```

```
[1] 0.1667 0.6667 0.1667
attr(,"class")
[1] acomp
```

### 2.5.4 Compositional Scalar Product, Norm, and Distance

The *Aitchison scalar product* for compositions

$$\langle \mathbf{x}, \mathbf{y} \rangle_A = \frac{1}{D} \sum_{i>j}^D \ln \frac{x_i}{x_j} \ln \frac{y_i}{y_j} \quad (2.4)$$

provides a replacement for the conventional scalar product. The generic function `scalar(x, y)` has been added to “compositions” to compute scalar products: if  $\mathbf{x}$  and  $\mathbf{y}$  are `acomp` compositions, the result will be that of (2.4). Recall that the scalar product induces a series of secondary, useful operations and concepts.

- The norm of a vector, its *length*, is  $\|\mathbf{x}\|_A = \sqrt{\langle \mathbf{x}, \mathbf{x} \rangle_A}$ . From an `acomp` vector  $\mathbf{x}$ , its norm can be obtained with `norm(x)`. If  $\mathbf{x}$  is a data matrix of class `acomp`, then the result of `norm(x)` will be a vector giving the norm of each row.
- The normalized vector of  $\mathbf{x}$  (or its *direction*, or a unit-norm vector) is  $\mathbf{v} = \|\mathbf{x}\|_A^{-1} \odot \mathbf{x}$ . The generic function `normalize` provides this normalization, either for a single composition or for the rows of a data matrix.
- The angle between two compositions,

$$\alpha(\mathbf{x}, \mathbf{y}) = \cos^{-1} \frac{\langle \mathbf{x}, \mathbf{y} \rangle_A}{\|\mathbf{x}\|_A \cdot \|\mathbf{y}\|_A} = \cos^{-1} \langle \|\mathbf{x}\|_A^{-1} \odot \mathbf{x}, \|\mathbf{y}\|_A^{-1} \odot \mathbf{y} \rangle_A,$$

which allows to say that two compositions are *orthogonal* if their scalar product is zero, or their angle is  $90^\circ$ . This is not implemented in a single command, but you get it with

```
> acos(scalar(normalize(x), normalize(y)))
```

- The (orthogonal) projection of a composition onto another is the composition

$$P_{\mathbf{x}}(\mathbf{y}) = \frac{\langle \mathbf{y}, \mathbf{x} \rangle_A}{\langle \mathbf{x}, \mathbf{x} \rangle_A} \odot \mathbf{x},$$

or in case of projecting onto a unit-norm composition,  $P_{\mathbf{v}}(\mathbf{y}) = \langle \mathbf{y}, \mathbf{v} \rangle_A \odot \mathbf{v}$ . This last case appears next when discussing coordinates with respect to orthonormal bases. A related concept is the (real) projection of a composition on a given direction,  $\langle \mathbf{y}, \mathbf{v} \rangle_A$ , which is a single real number.

- The *Aitchison distance* (Aitchison, 1986) between two compositions is

$$d_A(\mathbf{x}, \mathbf{y}) = \|\mathbf{x} \ominus \mathbf{y}\|_A = \sqrt{\frac{1}{D} \sum_{i=1}^D \sum_{j>i}^D \left( \ln \frac{x_i}{x_j} - \ln \frac{y_i}{y_j} \right)^2}. \quad (2.5)$$

The distance between two compositions can be directly computed by `norm(x-y)`.

In “compositions”, the generic function `dist` automatically computes the Aitchison distances between all rows of an `acomp` data matrix. The result is an object of class `dist`, suitable for further treatment (e.g., as input to hierarchical cluster analysis, see Sect. 6.2). Note that other compositional

distances, like Manhattan or Minkowski distances, are also available through the extra argument `method`:

```
> a = acomp(c(1,2,4))
> b = acomp(c(3,3,1))
> mycomp = rbind(a,b)
> mycomp = acomp(mycomp)
> dist(mycomp)

      a
b 1.813

> norm(a-b)

[1] 1.813

> dist(mycomp,method="manhattan")

      a
b 2.851

> sum(abs(clr(a-b)))

[1] 2.851
```

### 2.5.5 The Centered Log-Ratio Transformation (*clr*)

The set of compositions together with the operations perturbation  $\oplus$ , powering  $\odot$  and Aitchison scalar product  $(\cdot, \cdot)_A$  build a  $(D - 1)$ -dimensional Euclidean space structure on the simplex. This means that we can translate virtually anything defined for real vectors to compositions, as an Euclidean space is always equivalent to the real space. This equivalence is achieved through an *isometry*, i.e., a transformation from the simplex to the real space that keeps angles and distances. The first isometric transformation we use is the *centered log-ratio transformation* (*clr*)

$$\text{clr}(\mathbf{x}) = \left( \ln \frac{x_i}{g(\mathbf{x})} \right)_{i=1,\dots,D} \quad \text{with} \quad g(\mathbf{x}) = \sqrt[D]{x_1 \cdot x_2 \cdots x_D}, \quad (2.6)$$

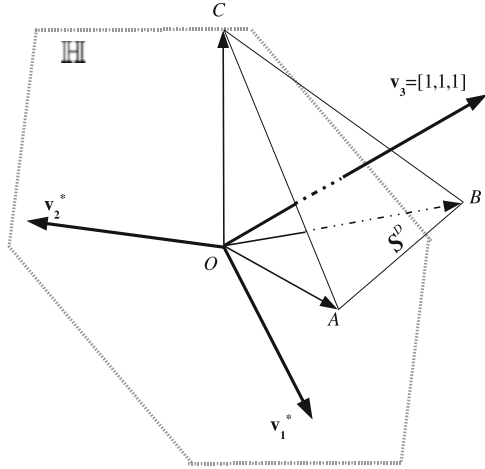
or in a compact way  $\text{clr}(\mathbf{x}) = \ln(\mathbf{x}/g(\mathbf{x}))$ , where the log ratio of the vector is applied component-wise. The inverse *clr* transformation is straightforward: if  $\mathbf{x}^* = \text{clr}(\mathbf{x})$ , then  $\mathbf{x} = \mathcal{C}[\exp(\mathbf{x}^*)]$ , where the exponential is applied by components. By its definition, the *clr* transformed components sum up to zero: in fact, the image of

**Fig. 2.7** Representation of a 3-part simplex ( $ABC, \mathbb{S}^3$ ) and its associated clr-plane ( $\mathbb{H}$ ), with an orthogonal basis  $\{\mathbf{v}_1^*, \mathbf{v}_2^*, \mathbf{v}_3^*\}$  of  $\mathbb{R}^3$ . The reader can find illustrative to play with a 3-D representation of the clr-transformed data of a 3-part composition, like

```

> data(Hydrochem)
> idx = c(14,17,18)
> x = Hydrochem[,idx]
> x = acomp(x)
> plot3D(x, log=TRUE)

```



the clr is a hyperplane (and a vector subspace, denoted with  $\mathbb{H}$ , called the *clr-plane*) of the real space  $\mathbb{H} \subset \mathbb{R}^D$  orthogonal to the vector  $\mathbf{1} = [1, \dots, 1]$ , i.e., the bisector of the first orthant (Fig. 2.7). This may be a source of problems when doing statistical analyses, as e.g., the variance matrix of a clr-transformed composition is singular.

The commands `clr` and `clrInv` compute these two transformations: they admit either a vector (considered as a composition), or a matrix or data frame (where each row is then taken as a composition).

### 2.5.6 The Isometric Log-Ratio Transformation (ilr)

It is well known that there is only one  $(D-1)$ -dimensional Euclidean space typically called  $\mathbb{R}^{D-1}$ , up to an isometric mapping. Therefore, there exist an isometric linear mapping between the Aitchison simplex and  $\mathbb{R}^{D-1}$ . This mapping is called the *isometric log-ratio transformation* (ilr).

The isometry is constructed by representing the result in a basis of the  $(D-1)$ -dimensional image space  $\mathbb{H}$  of the clr transformation. This is constructed by taking an *orthonormal basis* of  $\mathbb{R}^D$  including the vector  $\mathbf{v}_D = [1, \dots, 1]$ , i.e., some  $(D-1)$  linearly independent vectors  $\{\mathbf{v}_1^*, \dots, \mathbf{v}_{D-1}^*\} \in \mathbb{H}$  (Fig. 2.7). Then the set of compositions defined as  $\mathbf{v}_i = \text{clr}^{-1}(\mathbf{v}_i^*)$  form a basis of  $\mathbb{S}^D$ . In computational terms, one can arrange the vectors  $\{\mathbf{v}_j^*\}$  by columns in a  $D \times (D-1)$ -element matrix, denoted by  $\mathbf{V}$ , with the following properties:

- It is a quasi-orthonormal matrix, as

$$\mathbf{V}^t \cdot \mathbf{V} = \mathbf{I}_{D-1} \quad \text{and} \quad \mathbf{V} \cdot \mathbf{V}^t = \mathbf{I}_D - \frac{1}{D} \mathbf{1}_{D \times D}, \quad (2.7)$$

where  $\mathbf{I}_D$  is the  $D \times D$  identity matrix, and  $\mathbf{1}_{D \times D}$  is a  $D \times D$  matrix full of ones.

- Its columns sum up to zero, because they represent vectors of the clr-plane,

$$\mathbf{1} \cdot \mathbf{V} = \mathbf{0}. \quad (2.8)$$

Thanks to these properties, we can find simple expressions to pass between *coordinates*  $\xi$  and *compositions*  $\mathbf{x}$ :

$$\text{clr}(\mathbf{x}) \cdot \mathbf{V}^t = \ln(\mathbf{x}) \cdot \mathbf{V}^t =: \text{ilr}(\mathbf{x}) = \xi \quad (2.9)$$

$$\text{ilr}(\mathbf{x}) \cdot \mathbf{V} = \text{clr}(\mathbf{x}) \longrightarrow \mathbf{x} = \mathcal{C}[\exp(\xi \cdot \mathbf{V})]. \quad (2.10)$$

Through these expressions, we also defined implicitly the so-called *isometric log-ratio transformation* (ilr): this is nothing else than a transformation that provides the coordinates of any composition with respect to a given orthonormal basis. There are as many ilr as orthonormal basis can be defined, thus as matrices  $\mathbf{V}$  fulfilling (2.7) and (2.8). The one originally defined by [Egozcue et al. \(2003\)](#) was based on the (quite ugly) Helmert matrix

$$\mathbf{V} = \begin{pmatrix} \frac{D-1}{\sqrt{D(D-1)}} & 0 & \cdots & 0 & 0 \\ \frac{-1}{\sqrt{D(D-1)}} & \frac{D-2}{\sqrt{(D-1)(D-2)}} & \cdots & 0 & 0 \\ \frac{-1}{\sqrt{D(D-1)}} & \frac{-1}{\sqrt{(D-1)(D-2)}} & \cdots & 0 & 0 \\ \vdots & \vdots & \ddots & \vdots & \vdots \\ \frac{-1}{\sqrt{D(D-1)}} & \frac{-1}{\sqrt{(D-1)(D-2)}} & \cdots & \frac{2}{\sqrt{6}} & 0 \\ \frac{-1}{\sqrt{D(D-1)}} & \frac{-1}{\sqrt{(D-1)(D-2)}} & \cdots & \frac{-1}{\sqrt{6}} & \frac{1}{\sqrt{2}} \\ \frac{-1}{\sqrt{D(D-1)}} & \frac{-1}{\sqrt{(D-1)(D-2)}} & \cdots & \frac{-1}{\sqrt{6}} & \frac{-1}{\sqrt{2}} \end{pmatrix}. \quad (2.11)$$

The ilr transformation induces an isometric identification of  $\mathbb{R}^{D-1}$  and  $\mathbb{S}^D$ . For measure and probability theory purposes, this induces an own measure for the simplex, called the *Aitchison measure* on the simplex, denoted as  $\lambda_{\mathbb{S}}$ , and completely analogous to the Lebesgue-measure  $\lambda$ . This Aitchison measure is given by

$$\lambda_{\mathbb{S}}(A) = \lambda(\{\text{ilr}(x) : x \in A\}).$$

The ilr transformation is available through the command `ilr(x)`. An optional argument `V` can be used to specify a basis matrix different from the default one. This is itself available through the function `ilrBase(x,z,D)`, where the arguments represent, respectively, the composition, its ilr transformation, and its number of parts (only one of them can be passed!). An alternative interface to some ilr transformations is provided by the functions `balance(x)` and `balanceBase(x)`, explained in Sect. 4.3. The commands `ilrInv` provide the inverse transformation. Also, if we want to pass from ilr to clr or vice versa, we can use functions `ilr2clr` and `clr2ilr`.

```

> a = c(1,2,4)
> (ac = acomp(a))

[1] 0.1429 0.2857 0.5714
attr("class")
[1] acomp

> ilr(ac)

      [,1]      [,2]
[1,] 0.4901 0.8489
attr("class")
[1] "rmult"

> clr2ilr(clr(ac))

[1] 0.4901 0.8489

> (Vd = ilrBase(x=a))

      [,1]      [,2]
1 -0.7071 -0.4082
2  0.7071 -0.4082
3  0.0000  0.8165

> clr(ac) %%% Vd

[1] 0.4901 0.8489
attr("class")
[1] "rmult"

```

### 2.5.7 The Additive Log-Ratio Transformation (*alr*)

Finally, it is worth mentioning that compositional data analysis in the original approach of [Aitchison \(1986\)](#) was based on the *additive log-ratio transformation* (*alr*),  $\text{alr}(\mathbf{x}) = (\ln(x_1/x_D), \dots, \ln(x_{D-1}/x_D)) = (\ln(x_i/x_D))_{i=1, \dots, D-1}$ . Though we will not use it in this book, the *alr* transformation is available with the command `alr(x)`.

*Remark 2.2 (Why so many log-ratio transformations?).* The answer is quite easy: because none is perfect. All three of them recast perturbation and



powering to classical sum and product,

$$\begin{aligned}\text{clr}(\mathbf{x} \oplus \mathbf{y}) &= \text{clr}(\mathbf{x}) + \text{clr}(\mathbf{y}), \text{ilr}(\mathbf{x} \oplus \mathbf{y}) = \text{ilr}(\mathbf{x}) + \text{ilr}(\mathbf{y}), \text{alr}(\mathbf{x} \oplus \mathbf{y}) \\ &= \text{alr}(\mathbf{x}) + \text{alr}(\mathbf{y}) \\ \text{clr}(\lambda \odot \mathbf{x}) &= \lambda \cdot \text{clr}(\mathbf{x}), \text{ilr}(\lambda \odot \mathbf{x}) = \lambda \cdot \text{ilr}(\mathbf{x}), \text{alr}(\lambda \odot \mathbf{x}) = \lambda \cdot \text{alr}(\mathbf{x}).\end{aligned}$$

The first two, because they are isometric, also preserve the scalar product, but this does not happen with the alr transformation,

$$\langle \mathbf{x}, \mathbf{y} \rangle_A = \text{clr}(\mathbf{x}) \cdot \text{clr}'(\mathbf{y}) = \text{ilr}(\mathbf{x}) \cdot \text{ilr}'(\mathbf{y}) \neq \text{alr}(\mathbf{x}) \cdot \text{alr}'(\mathbf{y})$$

Thus, alr should not be used in case that distances, angles, and shapes are involved, as it deforms them.

On the other side, we already mentioned that the clr yields singular covariance matrices, and this might be a source of problems if the statistical method used needs inverting it. One would need to use Moore–Penrose generalized inverses. As an advantage, the clr represents a one-by-one link between the original and the transformed parts, which *seems* to be helpful in interpretation.

This is exactly the strongest difficulty with the ilr-transformed values or any orthonormal coordinates: each coordinate might involve many parts (potentially all), which makes it virtually impossible to interpret them in general. However, the ilr is an isometry and its transformed values yield full-rank covariance matrices; thus, we can analyze ilr data without regard to inversion or geometric problems. The generic ilr transformation is thus a perfect black box: compute ilr coordinates, apply your method to the coordinates, and recast results to compositions with the inverse ilr. If interpretation of the coordinates is needed, one should look for preselected bases and preselected ilr transformations, as those presented in Sect. 4.3.

### 2.5.8 Geometric Representation of Statistical Results

The fact that the simplex is an Euclidean space has some implications on the way we apply, interpret and represent most linear statistics. The basic idea is summarized in the *principle of working on coordinates* (Pawlowsky-Glahn, 2003), stating that one should:

1. Compute the coordinates of the data with respect to an orthonormal basis (2.9).

2. Analyze these coordinates in a straightforward way with the desired method; no special development is needed, because the coordinates are real unbounded values.
3. Apply those results describing geometric objects to the orthonormal basis used, recasting them to compositions (2.10). Final results will not depend on the basis chosen.

Regarding the last step, most “geometric objects” obtained with linear statistics are points or vectors, lines/hyperplanes, and ellipses/ellipsoids.

*Points* (like averages and modes or other central tendency indicators, outliers, intersection points, etc.) are identified with their vectors of coordinates in the basis used. Thus, to represent a point as a composition, it suffices to apply its coordinates to the basis, i.e., compute the inverse  $\text{ilr}$  transformation (2.10). To draw points in a ternary diagram (or a series of ternary diagrams), one can use the function `plot(x)` where  $\mathbf{x}$  is already a composition of class `acomp` (or a dataset of compositions). If points have to be added to an existing plot, the optional argument `add=TRUE` will do the job. This function admits the typical **R** set of accessory plotting arguments (to modify color, symbol, size, etc.).

*Lines* (like regression lines/surfaces, principal components, discriminant functions, and linear discriminant borders) are completely identified with a point  $\boldsymbol{\alpha}$  and a vector  $\boldsymbol{\beta}$  in the  $\text{clr}$ -plane. The parametric equation  $\boldsymbol{\xi}(t) = \boldsymbol{\alpha} + t \cdot \boldsymbol{\beta}$  runs through every possible point of the line taking different values of the real parameter  $t$ . Applying this parametric equation to the basis in use, we obtain a compositional line,

$$\mathbf{x}(t) = \mathbf{a} \oplus t \odot \mathbf{b}, \quad (2.12)$$

with  $\mathbf{a} = \text{ilr}^{-1}(\boldsymbol{\alpha})$  and  $\mathbf{b} = \text{ilr}^{-1}(\boldsymbol{\beta})$ . A  $P$ -dimensional plane needs  $P$  compositions of the second kind, i.e.,

$$\mathbf{x}(t_1, \dots, t_P) = \mathbf{a} \oplus \bigoplus_{i=1}^P t_i \odot \mathbf{b}_i. \quad (2.13)$$

Note that a  $P$ -dimensional plane projected onto a subcomposition is still a  $P$ -dimensional plane. In particular, a line projected on a 3-part subcomposition is a line, obtained by selecting the involved parts in the vectors  $\mathbf{a}$  and  $\mathbf{b}$ . Straight lines can be added to existing plots by the generic function `straight(x,d)`, which arguments are respectively the point  $\mathbf{a}$  and the vector  $\mathbf{b}$  (as compositions!). Sometimes only a segment of the line is desired, and two functions might be helpful for that. Function `segments(x0,y)` draws a compositional line from  $\mathbf{x}_0$  to  $\mathbf{y}$ , i.e., along  $\mathbf{y}-\mathbf{x}_0$ ; both *can* be compositional datasets, thus resulting in a set of segments joining  $\mathbf{x}_0$  and  $\mathbf{y}$  row by row. Function `lines(x)` on the contrary *needs* a compositional dataset and draws a piece-wise (compositional) line through all the rows of  $\mathbf{x}$ . Further graphical arguments typical of lines (width, color, style) can also be given to all these functions.

*Ellipses and hyper ellipsoids* (like confidence or probability regions or quadratic discriminant borders) are completely specified by giving a central point  $\boldsymbol{\alpha}$  and a

symmetric positive definite matrix  $\mathbf{T}$ . An ellipse is the set of points  $\xi$  fulfilling

$$(\xi - \alpha) \cdot \mathbf{T} \cdot (\xi - \alpha)^t = r^2, \quad (2.14)$$

namely, the set of points with norm  $r$  by the scalar product represented by the matrix  $\mathbf{T}$ . This can be extended to conics (and hyper quadrics) by dropping the condition of positive definiteness. Any hyper quadric projected onto a subcomposition is still a hyper quadric. In particular, in two dimensions (three parts), an ellipse is obtained with a positive definite matrix, a parabola with a semi-definite matrix, and an hyperbola with a non-definite matrix. All hyper quadrics show up as borders between subpopulations in quadratic discriminant analysis. But by far, their most typical use is to draw elliptic probability or confidence regions around the mean (see Example 4.2). In this case, we can better characterize the ellipses with the original variance matrix, i.e.,  $\mathbf{T} = \mathbf{S}^{-1}$ . This is the approach implemented in the generic function `ellipses(mean, var, r)`, which arguments are a central composition (`mean`), a clr-variance matrix (`var`, described in Sect.4.1), and the radius of the ellipse (`r`). Further graphical arguments for lines are also admitted.

#### Summary of graphical functions:

- `plot(x)` to draw ternary diagrams of the `acomp` composition `x`
- `plot(x, add=TRUE)` to add compositions to existing ternary diagrams
- `straight(x,d)` to draw lines on ternary diagrams along `d` passing through `x`, both `acomp` compositions
  - `segments(x0,y)` to add compositional segments from the rows of `x0` to the rows of `y`
  - `lines(x)` to add piece-wise compositional lines through `x` rows
- `ellipses(mean, var, r)` to draw an ellipse around the `acomp` composition `mean`, defined by a radius `r` and a clr-variance `var`

### 2.5.9 Expectation and Variance in the Simplex

In an Euclidean space, like the simplex, expectation and variance can be defined with regard to its geometry. [Eaton \(1983\)](#) presents the theory for any Euclidean space, and we will here present without proof the most important results in the Aitchison geometry of the simplex. The goal of this section is twofold. On the one hand, we want to see that the actual basis used to compute coordinates and apply statistics is absolutely irrelevant: back-transformed results are exactly the same whichever basis

was used. On the other hand, we will also find that variance matrices may be seen as *representations* of some objects on themselves, with a full meaning: this ensures us that whichever log-ratio transformation we use (clr, ilr or alr), expressions like (2.14) are fully correct as long as all vectors and matrices are obtained with regard to the same log-ratio transformation.

Take the Euclidean space structure of the simplex  $(\mathbb{S}^D, \oplus, \odot, \langle \cdot, \cdot \rangle_A)$  and a random composition  $\mathbf{X} \in \mathbb{S}^D$ . Its *expectation within the simplex* is a fixed composition  $\mathbf{m}$  such that the expected projection of any composition  $\mathbf{v} \in \mathbb{S}^D$  onto  $\mathbf{X}$  is perfectly captured by its projection onto  $\mathbf{m}$ ,

$$E_{\mathbb{S}}[\mathbf{X}] = \mathbf{m} \Leftrightarrow \text{for all } \mathbf{v} : E[\langle \mathbf{v}, \mathbf{X} \rangle_A] = \langle \mathbf{v}, \mathbf{m} \rangle_A. \quad (2.15)$$

This says that knowing  $\mathbf{m}$ , we know the average behavior of  $\mathbf{X}$  projected onto any direction of the simplex. This definition does not make use of any basis, so we can be sure that the concept itself is basis-independent. To compute it, however, we may take the vector  $\mathbf{v}$  successively as each of the  $(D - 1)$  vectors of *any* ilr basis, thus obtaining the mean coordinates with respect to that basis,

$$\text{ilr}(E_{\mathbb{S}}[\mathbf{X}]) = E[\text{ilr}(\mathbf{X})] \in \mathbb{R}^{D-1}$$

This is the result we would obtain by applying the principle of working on coordinates. In other words, the definition given by (2.15) and the one implied by the principle of working on coordinates are equivalent, and both are in fact basis-independent.

For the variance, we may obtain equivalent results by defining it as an endomorphism. An endomorphism within the simplex is an application  $\Sigma : \mathbb{S}^D \rightarrow \mathbb{S}^D$  that preserves the linearity of perturbation and powering, e.g.,  $\Sigma(\mathbf{x} \oplus \lambda \odot \mathbf{y}) = \Sigma(\mathbf{x}) \oplus \lambda \odot \Sigma(\mathbf{y})$ . The *variance within the simplex* of a random composition  $\mathbf{X} \in \mathbb{S}^D$  is an endomorphism  $\Sigma$  such that the expected product of the projections of any pair of compositions  $\mathbf{u}, \mathbf{w} \in \mathbb{S}^D$  onto  $\mathbf{X}$  (centered) is perfectly captured by their scalar product through  $\Sigma$ :

$$\text{var}_{\mathbb{S}}[\mathbf{X}] = \Sigma \Rightarrow E[\langle \mathbf{w}, \mathbf{X} \ominus \mathbf{m} \rangle_A \cdot \langle \mathbf{u}, \mathbf{X} \ominus \mathbf{m} \rangle_A] = \langle \mathbf{w}, \Sigma(\mathbf{u}) \rangle_A = \langle \mathbf{u}, \Sigma(\mathbf{w}) \rangle_A. \quad (2.16)$$

In words, by knowing  $\Sigma$ , we know the expected covariation of  $\mathbf{X}$  projected onto any pair of directions of the simplex. As with expectation, the ilr-coordinate representation of  $\Sigma$  is a  $(D - 1) \times (D - 1)$  matrix  $\Sigma_v = (\sigma_{ij}^v)$  such that  $\sigma_{ij}^v = \text{cov}[\text{ilr}_i(\mathbf{X}), \text{ilr}_j(\mathbf{X})]$ . And equivalently, its clr representation is a  $D \times D$  matrix  $\Sigma = (\sigma_{ij})$  such that  $\sigma_{ij}^v = \text{cov}[\text{clr}_i(\mathbf{X}), \text{clr}_j(\mathbf{X})]$ . Note that, whichever transformation we use,

$$\langle \mathbf{u}, \Sigma(\mathbf{w}) \rangle_A = (\text{ilr}(\mathbf{u}), \text{ilr}(\Sigma(\mathbf{w}))) = \text{ilr}(\mathbf{u}) \cdot \Sigma_v \cdot \text{ilr}'(\mathbf{w}) = \text{clr}(\mathbf{u}) \cdot \Sigma \cdot \text{clr}'(\mathbf{w}).$$

Thus, all these matrices and vectors, expressed as compositions, in clr coefficients or in ilr coordinates, represent exactly the same objects. We can consistently change between them with the following expressions:

$$\begin{aligned}\text{ilr}(\mathbf{m}) &= \text{clr}(\mathbf{m}) \cdot \mathbf{V} = \boldsymbol{\mu}_v & \mathbf{m} &= \text{clr}^{-1}(\boldsymbol{\mu}_v \cdot \mathbf{V}^t) = \text{ilr}^{-1}(\boldsymbol{\mu}_v), \\ \boldsymbol{\Sigma}_v &= \mathbf{V}^t \cdot \boldsymbol{\Sigma} \cdot \mathbf{V} & \boldsymbol{\Sigma} &= \mathbf{V} \cdot \boldsymbol{\Sigma}_v \cdot \mathbf{V}^t.\end{aligned}\quad (2.17)$$

Functions `clrvar2ilr(xvar)` and `ilrvar2clr(zvar)` implement the variance transformations of (2.17), where `xvar` is the clr variance  $\boldsymbol{\Sigma}$  and `zvar` is the ilr variance  $\boldsymbol{\Sigma}_v$ . To transform the mean vector, we may use the self-explaining functions `ilr`, `clr`, `ilrInv`, `clrInv`, `ilr2clr`, and `clr2ilr`.

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