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Master Thesis

*Benchmarking machine learning performances with compositional* *data*

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# Abstract

Machine Learning in microbiome studies is widely used and the interest is growing. However, there is no universal understanding of the algorithmic approaches that can best utilize the information present in the microbiome data. Thus, this is an interesting and widely discussed topic that can have a great impact on the potential applications leveraging microbiome data. A key topic in microbiome research is the sample space of the input data. The sequencing data appears as count data, but, only relative abundance of the microbial features can be observed, commonly called “compositional data”. Thus, transforming the read counts to relative abundances is usually the first step and Machine Learning methods are usually applied on relative abundances. However, relative abundances raise several limitations, which can have an impact on the performance of the prediction models. Therefore, log-ratio transformations are a proposition made by several studies now, however their impact on Machine Learning performances has never been tested in large-scale studies. The goal of this benchmarking project is to rectify that and apply several log-ratio transformed data in Machine Learning models and compare the performances to *CoDaCoRe*, an algorithm specifically made with microbiome analysis in mind. This way it will become clearer how log-ratio transformations impact Machine Learning performances with microbiome data.

# Introduction

Working with mathematical concepts is always a bit out of the comfort zone for most biologists. The field of statistics looms like a big black cave which is only lightened in a few places, but we have to traverse it anyways. Unfortunately, with Machine Learning encroaching also into biological research questions, this cave just became much bigger, without really bringing any additional light.

Machine Learning provides a useful tool to exploit information from biological data sets. It brings the possibility of predicting host-phenotypes, finding associations between features, and using microbial communities to characterize patients. Several papers (Jasner et al. 2021; Kubinski et al. 2022; Coenders and Greenacre 2021; Zhang and Shi 2019) already gave some practical advice on how to use Machine Learning with microbiome data, however a vital concept concerning the mathematical characteristics of microbiome data has been almost completely ignored in the last few years.

It has been made abundantly clear in the last years that sequencing data is of compositional nature, which means it has distinct mathematical characteristics than other data types (Greenacre et al. 2021b; Gloor et al. 2017; Quinn et al. 2018). While some papers already confirmed that log-ratio transformations do increase Machine Learning performances (Jasner et al. 2021; Kubinski et al. 2022; Quinn and Erb 2020; Zhang and Shi 2019), it has never been tried to assess how other important parameters (like sample sizes, feature sizes or test set methodologies) might be affected by the application of log-ratio transformations in Machine Learning models. Or if perhaps these parameters demand a choice in log-ratio transformation.

The goal of this master thesis is to look a bit deeper into the combination of log-ratio transformations and Machine Learning models. For that, the properties of compositional data will be introduced, and the idea of log-ratio transformations explained. Afterwards, Machine Learning performances will be compared with

comparing how they apply in practical Machine Learning pipelines and if they result in improved and beneficial performances. ~~the information around compositional data more approachable, summarizing the achieved solutions, and in a practical part, trying to assess if these achievements are also applicable when combining compositional data and machine learning concepts.~~

~~As this master thesis uses microbiome sequencing data and was created in a microbiome research group, this text will mostly focus on this field and its papers. All results should be applicable to other high-throughput sequencing data, as well as any data that is in some way confined by an arbitrary sum. Such data is found for example in geochemistry, ecology, sociology, political sciences, etc., and therefore ultimately spans the problematic into various different fields (Greenacre et al. 2021a). However, sequencing data is quite unique as it usually contains way more features than samples, which comes with its own problems in Machine Learning.~~

## Characteristics of Compositional Data

In order to define and illustrate the concept of compositional data, let’s assume a classic biological example. The following Figure (Fig.1A) shows two different ecological fields: A and B. In field A, four rabbits, seven birds, eight bees and one wolf have been counted, whereas field B contains two rabbits, four birds, four bees and one wolf. It becomes clear that, as similar as the diversity may be, field B seems to have only half of the population of field A. This in itself is already valuable information. The total counts per field can be easily preserved in our data collection and therefore, the absolute count of each organism in this field matters.



Figure 1: Information Loss of Normalized Data

(A) Illustration of the number of animals found in two different samples. Field A contains four rabbits, eight bees, seven birds and one wolf, whereas field B contains two rabbits, four bees, four birds and one wolf. In (B) the absolute counts have been plotted as a stacked bar plot, with each animal in a different color. (C) shows the stacked bar plot as normalized counts, e.g., here percentages.

When using absolute counts, the difference between both fields is easily visible (Fig.1B). However, when we really want to compare both fields, we need to transform the samples to a common scale. This is called normalization and we can see the effect in (Fig.1C). As soon as the data is normalized, the particular information of absolute counts gets lost. When collecting ecological data ourselves, we can preserve the fact that field B only contained eleven individuals and field A contained twenty, by saving that number somewhere in an Excel sheet. However, the problem with sequencing data is: we get the data in the form of Fig.1C.

To demonstrate how a sequencing machine cannot preserve absolute counts, imagine the following situation: We want to sample field A multiple times a day, but in order to be more efficient, we buy a machine to do the counting for us. Three times a day this machine transmits the number of all the different animals coming to this field. However, this machine has one flaw: it can only count to twenty. As soon as the twenty-first animal on this day comes to the field, it is just simply not counted.

Consequently, the overall number twenty carries no meaning. Every sample has this exact total number, so it carries no valuable information. Of course, a limit of twenty is weird for us to understand, but sequencing machines do a very similar same thing. They are limited in their capacity on e.g., the flow cells and even the most effective sequencing machines could never fully sequence the entirety of the organism’s DNA contents. But not only the technology limits our counting capabilities, but the sampling procedures to acquire microbiome data can only entail a small fraction of the actual bacterial content. Thus, the total number of sequences measured by sequencing machines ultimately depends on the chemistry of the assay, not the input material (Quinn et al. 2018).

The consequence of this sampling problem is, that we have to accept the fact that the sum of counts in sequencing data are irrelevant. This is particularly problematic as we cannot compare groups of data as we are used to do it. A lot of tools in microbiome data analysis stem from the field of ecology and contain methods like richness or diversity to characterize samples. If we only had the information of Fig.1C, then we would perhaps assume that field A and B share the same ecology. Or perhaps, if this were microbiome data, we would maybe falsely assume that these samples came from patients of the same group.

Data that is constraint in that way is called “Compositional Data”. It was first introduced by John Aitchison, roughly 40 years ago and since then steadily improved and further developed. Thanks to this mathematical theory, we can still draw information from sequencing data. We just have to adjust for the fact that the absolute counts are non-informative (Quinn et al. 2018; Greenacre et al. 2021b; Aitchison 1982). We can instead use relative abundances, sometimes also called proportions, between features in a sample.

At first glance, this does not sound as if it would have serious consequences. However, in the next section we will a look at the mathematical concepts behind compositional data and its properties and demonstrate, that the concept of compositional data changes the whole standard downstream analysis (Gloor et al. 2017).

## The Simplex Space

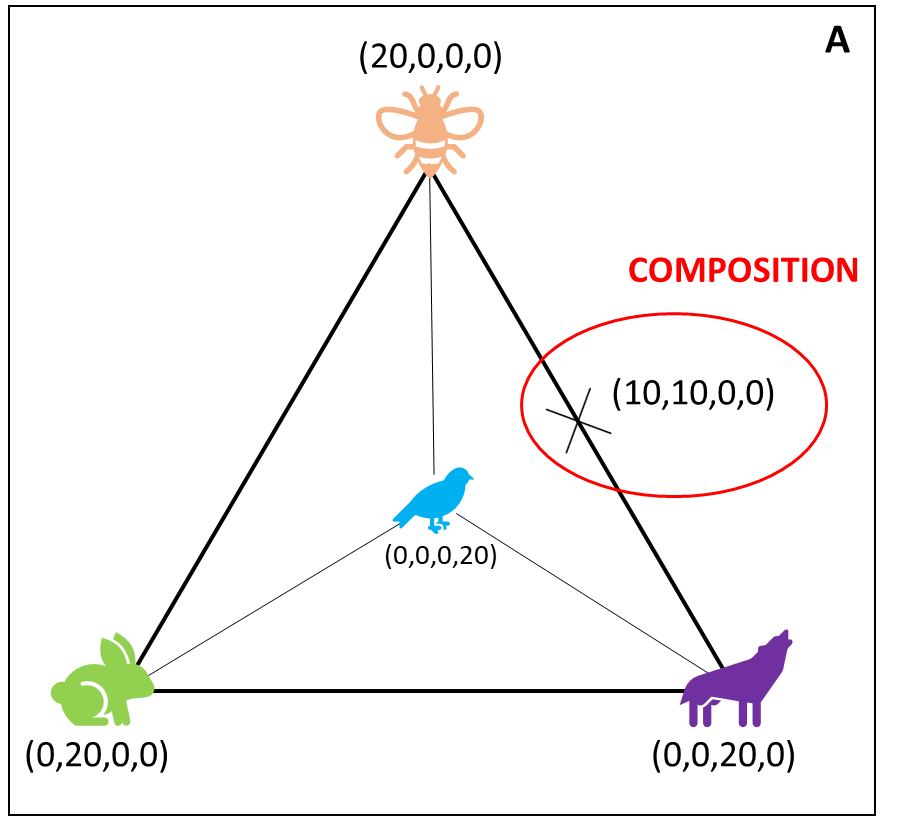
As mentioned, absolute counts in compositional data are unavailable and only relative abundances are of interest. This puts the data in the so-called “Simplex Space”, instead of the for us more common Euclidean space. The following Figure 2 shows how the data from field A in section 2.1 would look like in the Simplex Space:

Figure 2: Biological Example in the Simplex Space

Assuming the collected ecological data from Figure 1 is compositional, it can be visualized in a S3-Simplex space. Geometrically, a tetrahedron is created with all different components (here animals) placed on the four corners of a polytope. A composition is one possible combination of components confined in the Simplex Space.

We stick with the ecological example and place all animals as one corner in a geometrical space. With four features, we are able to create a geometric figure called tetrahedron that represents the Simplex Space visually.

We use the flawed machine, and one day, we sample twenty rabbits in field A. This would lead to a point in the simplex space that sits directly in the left corner, with the coordinates (0,20,0,0), because we only have rabbits, no other animal. Another day, we sample only twenty bees, and no other animals, then we would find our data point where the bee is, at the very top. Marked in red is a sample where the machine counted ten bees, ten wolves and no birds and no rabbits. Every sample round produces one “composition” and the examples show, that the distance between any two variables is sensitive to the presence or absence of other components (Quinn et al. 2018). If a composition is moved from one corner of the animal-simplex, it means that the other values in the composition are directly changed. Consequently, that makes all variables *mutually dependent* on one another. In literature this data is also called “spurious” because it appears as if the data points have a causal relationship when the perceived correlation is only due to data properties. When a composition is moved from bees in the direction of wolves it seems like there is a causal relationship because the increase in the number of wolves, directly decreases the number of bees.

To describe this a bit more mathematically, the problem described above is formally known as “the negative bias problem” (other names are also the constant-sum problem, the closure problem, or the null correlation difficulty) (Aitchison 2003).

~~When the sum of a composition is constant, then it can be mathematically shown that the covariances between any two compositions equals 0. (Aitchison 2003; Pawlowsky-Glahn and Egozcue 2016). This has the consequence that the calculate variance would be negative, which is problematic, as variances are by definition always positive.~~

It means, that negative covariances are presupposed by the limitation of the sum, instead of produced by stochastic factors (Pawlowsky-Glahn and Egozcue 2016; Aitchison 2003). Covariances are similar to correlation coefficients, only that the latter is constraint between -1 and 1, but otherwise they can be interpreted similarly. Translated, a negative covariance means that two factors are negatively correlated, i.e., if one factor goes up, another goes down. Exactly as has been demonstrated by the ecological example before.

Thus, using any form of statistical test or machine learning tool seems redundant, as errors in correlations, univariate and multivariate tests are almost preconditioned, and we easily would make false assumptions about the correlation of the data. Thus, a correct handling of compositional data and the Simplex Space is not optional (Gloor et al. 2017).

## Mapping the Simplex Space into Euclidean Space

In the ecology sample, the Simplex could have been easily overcome by e.g., normalizing to a field size. This indirectly preserves information about the total number of animals. Similarly, it has been tried for sequencing data to calculate an “effective library size” and therefore recover this way the original scale of data. For that, normalization methods like trimmed mean of M-values (TMM) have been introduced, as well as Reads per Kilobase Million (RPKM) and Transcripts per Kilobase Million (TPM) (Quinn et al. 2018). However, all of those methods involve rescaling counts by the library size and these normalizations come with the drawback that some of these methods are sensitive to the removal of low abundant counts, as well as to data symmetry (Quinn et al. 2018).

Furthermore, Aitchison already criticized very early that there is no “magic to open up closed data” (Aitchison 2003), which is what normalization tries to do. Moreover, since information provided in compositional data is essentially about ratios of the components, it seems logical to also think in terms of ratios. Methods that are able to deal with the Simplex Space when analyzing compositional were introduced by Mateu-Figueras et al., (2011) with the “staying-in-the-simplex” approach or Greenacres (2017) “pragmatic approach”. However, these methods require a technical understanding of the algebraic-geometric structure of the Simplex and are therefore relatively complex.

No, the best way forward in Aitchison’s mind was to transform the data in a way that allows us to use it with Euclidean Space rules. The general idea is, that the Simplex Space is endowed with a Euclidean space structure. This has several mathematical advantages: if one can map the Simplex Space into Euclidean Space, then we can use all the to us more familiar methods and don’t have to rely on more complicated equations to e.g., calculate linear combinations and analyze data (Pawlowsky-Glahn and Egozcue 2016). Aitchison (1982) proposal to achieve that was to use log-ratio transformations.

Here, the focus will be on these log-ratio transformations, as they have been more heavily favored in the last decades due to their practicability (Greenacre et al. 2022). They also come with the advantage that they can be easily added to pre-existing Machine Learning pipelines that have been proposed and created over the last years. The next section introduces them more in detail.

## Log-Ratio Transformations

Add more context to the problem of “interpreting log-ratio transformations”

Missing link if those properties are relevant for ML

When Aitchison (1982) first tried to overcome the “bounded sum problem” he defined some principals to lay down as fundamentals to the good practice of compositional data. These are important as they paved the way that data could be mapped into Euclidean space correctly.

The most agreed upon properties are scale invariance (compositions do not change with e.g., sequencing depth), perturbation invariance (i.e., converting a composition between equivalent units will not change the results), and permutation invariance (i.e., changing the order of the components within a composition will not change the results) (Quinn et al. 2018; Greenacre et al. 2022)

It’s clear why these concepts are commonly accepted. Imagine a composition of the ecology example with composition (8,8,2,2), measuring eight bees, eight wolves, two swallows and two rabbits. Scale invariance means that a transformation should ensure that we get the same ratios even if we had a machine that was able to count to forty instead of twenty and when we would get the composition (16,16,4,4), the ratios between the components are stable. Similarly, if we would not count to the base of ten, but rather to the base of hundred, the transformation should not change the ratios. Lastly, the ratios should not be affected even if we sampled data in another order (e.g., two rabbits, eight wolves, two swallows, eight bees).

Three more properties exist that are more heavily discussed in their importance and if they are strictly necessary: isometry (i.e., the distances between ratios should be exactly the same after mapping into Euclidean space), sub-compositional coherence (i.e., overlapping components in two compositions of the same measurement should have the same ratios), and sub-compositional dominance (i.e., using a complete composition carries less information than using the whole (Quinn et al. 2018; Greenacre et al. 2021a; Greenacre et al. 2022).

Imagine two different scientists observed the same field but scientist A finds eight bees, two birds, two rabbits and eight wolves, whereas scientist B finds two birds, eight wolves and two rabbits. Both were not able to find one animal from the original composition and if one would normalize the results to 1, one would get completely different result (Scientist A: (0.4,0.1,0.1,0,4) and Scientist B: (0.16,0.6,0.16)), respectively. However, if ratios are studied, the overlapping components give the same ratios (e.g., the ratio birds/rabbits in both cases is 1). Given the same example the idea of isometry can also be explained. The distance between eight wolves and two rabbits in the raw compositional data of scientist A is 8-2 = 6, whereas in a log-transformed data set the distance is ln(8)-ln(2) = 1.3. This would mean that in this example applying the natural log does not conform to isometry, as 1.3 ≠ 6 and some advocate that these distances should be preserved in log-ratio transformations.

One could argue that all mentioned properties are important, unfortunately adhering to all of them has practical implications. Aitchison proposed log-transformations, because they already adhere to most of the properties inherently, like scale variance, perturbation invariance and permutation invariance. However, properties like isometry and sub-compositional coherence are more difficult.

The log-ratio transformation that imparts all those properties is called isometric log-ratio (ILR). ILRs are considered the gold standard of log-ratio transformations, as they engender exactly the same multivariate geometric structure of the sample points as that of the formerly mentioned Aitchison geometry (Greenacre et al. 2021b). ILR map a composition in the D-part Aitchison-Simplex isometrically to a D-1 dimensional Euclidian vector, and the procedure how that is done is quite difficult to understand (Greenacre et al. 2021a; Greenacre et al. 2022). Additionally, ILRs are particularly problematic when the numbers of components are high as the computational power to calculate ILRs increases significantly (Greenacre et al. 2021b). Considering that microbiome data is usually very high-dimensional would make ILR already not a good solution.

Text, whiteboard

Description automatically generatedThankfully, there are more types of log-ratio transformation, that are easier to understand and therefore interpret. As a result of this, it was decided to use two types of log-ratio transformation here: ALR (additive log-ratio) and CLR (centered log-ratio).

Figure 3: Equation for CLR

The equation describes the calculation of CLR, with xj as vector of sample features, Dj the total number of features, and g(x) the geometric mean of sample vector x. Log-ratio transformations are applied within a sample (i.e., row-wise).

In CLR, log-ratios are computed between each component and the geometric mean of all components (Gloor et al. 2017). It has the advantage that it is computationally easy to do, which is an advantage compared to ILRs. Furthermore, it reproduces the log-ratio geometry perfectly. However, it is not sub-compositionally coherent, because the whole composition (i.e., sample) is used to calculate the geometric mean and every sample will therefore use a different geometric mean, as well as every sub-composition. Unfortunately, it is not very easy to interpret and it is not very useful in sparse data containing a lot of zeroes (Gloor et al. 2017).

A picture containing text, watch

Description automatically generatedThe second log-ratio transformation is ALR. Here, the log-ratio is taken of each feature within a composition and divided by a chosen reference feature.

Figure 4: Equation ALR

The equation describes the calculation of ALR, with xj as vector of sample features, D the total number of features, and xref the reference feature. Log-ratio transformations are applied within a feature (i.e., column-wise).

Thus, the interpretation of ALR log-ratios is very straight-forward, as one can interpret a change in all features with respect to the reference. ALR transformed values are also sub-compositional coherent, which is traded for a small loss in isometry. The biggest problem with ALR is the choice of reference. A reference can simply be chosen by the scientist, for example when they want to compare one bacterium to all others. However, if one is unsure about the choice of reference, Greenacre et al. (2021) proposed to use three criteria to find a good reference: (i) the reference component should maximize the Procrustes correlation between the additive log-ratio geometry and the exact log-ratio geometry, (ii) the reference should minimize the variance of log-transformed components, and (iii) it should be a well-populated component.

This means the optimal reference can be determined computationally. Using these guidelines produces additive log-ratios close to being near-isometric, which would make them a favorable log-transformation. The obvious drawback is the computational complexity when a Procrustes analysis is used, which increases especially in higher-dimensional data.

## The Problems

Machine Learning models are of great interest for microbiome analysis, as they allow detection of biomarkers, phenotypes or microbial taxa, as well as other interesting tasks, that are not possible with the standard microbiome tool kit (Marcos-Zambrano et al. 2021). In supervised Machine Learning a model is learned that makes predictions on new data and can identify genomic features that underlie a predictive model (Whalen et al. 2022). The hopes are that with the help of AI-based techniques we can transform clinical decision-making and extract more knowledge from the amount of data available to us (Quinn et al. 2022).

However, while modern machine learning algorithms are very powerful, they come with a great lack of transparency. It is hard to understand how and why the model makes predictions. Especially deep neural networks are considered “black boxes” as the inner workings remain incomprehensible to the outsider and the lack of quality assurances goes against good scientific practice (Quinn et al. 2022). As it has been shown that well-specified machine learning models outperform deep learning models (Bailly et al. 2022), and that they allow easier interpretation, more focus is currently put on supervised Machine Learning models like generalized linear models (GLM), support vector machines (SVM) or random forests algorithms (RF). Still, a correct application and assessment of Machine Learning models is key to reproducible and interpretable research results.

Now, with compositional data and log-ratio transformations, a new step stone should be introduced to Machine Learning. Some problems become apparent when combining microbiome data, Machine Learning, and log-ratio transformations. It has been foreshadowed in the previous section that microbiome data itself is a problem for both log-ratio transformation and machine learning. Microbiome sequencing data is usually very sparse, which means it contains a lot of zeroes and therefore it is left-skewed in its distribution (Gloor et al. 2017).

The zero values are especially problematic in the context of log-ratio transformation. The problem is two-fold. First, the logarithm of zero for any type of log is undefined. Secondly, as also ratios are used, placing a zero in the denominator also leads to errors. Simply removing all features containing zeros, as Aitchison suggested (Aitchison 1982) is not practical, especially in a biological context, as it is quite possible to have features with zero values that have biological relevance, for example when comparing gene expressions between samples.

For Machine Learning, zeros itself are not that problematic, still the sparse nature of microbiome data affects machine learning models on several technical levels: the most problematic is the increasing complexity of models, as it needs to fit more coefficients or have greater depth to account for all features. It has also been shown that too many features produce noise in the training data which can easily lead to overfitting and negatively impacts the predictive power. Similarly, a machine learning model can also not differentiate between biological or technical zeros and therefore may underestimate the importance of denser features.

Studies in the last years contributed to streamlining Machine Learning pipelines and establishing a good-scientific-practice in order to ensure reproducibility and usability, also specifically for microbiome data (Jasner et al. 2021). Additionally, studies showed how to reduce bias and error by recommending specific cross-validation procedures (Tsamardinos et al. 2015) and the influence of sample size has been tried to estimate for various Machine Learning models (Beleites et al. 2013; Bailly et al. 2022; van der Ploeg et al. 2014).

To this day, it is still unclear if and how log-ratio transformation impact the performance in a prediction task. In general, log-ratio transformations seem to outperform raw proportions for classification tasks, but it is not clear how log-ratio transformations relate to the changes in predictive performance and if other factors in a Machine Learning pipeline are affected by log-ratio transformations.

A solution that tries to solve some of these problems is to tailor algorithms to the specifications of microbiome data. One such solution is *CoDaCoRe*. Published by Gordon-Rodriguez et al in 2021 in the paper “Learning Sparse Log-Ratios for High-Throughput Sequencing Data”, *CoDaCoRe* is a novel learning algorithm for finding balances (**Co**mpositional **Da**ta via **Co**ntinuous **Re**laxations). Balances are defined as the log-ratios between geometric means of two or more features of the input variables. Translated, *CoDaCoRe* finds ratios between groups of features that are explanatory for the given classification task. Such ratios are commonly used as biomarkers of gut health e.g., the Firmicutes-to-Bacteroidetes ratio (Crovesy et al., 2020; Magne et al.; 2020).

As *CoDaCoRe* is able to combine more than two features in its log-ratios, it leads to a richer set of features and therefore more flexible models, compared to e.g., ALR, which only uses one reference. Finding balances is usually very computationally taxing, however, in *CoDaCoRe* Gordon-Rodrigues et al. use a deep learning technology called “continuous relaxation” and only approximate the optimization problem, which has the advantage of greatly reducing the runtime.

Thus, *CoDaCoRe* identifies a sequence of balances, in decreasing order of importance, each of which is sparse and interpretable. This makes it a promising algorithm that is created to also work efficiently on big data sets with a lot of features and could be a good alternative to counter the problems that microbiome data brings into Machine Learning and log-ratio transformation. In their paper, the authors compare *CoDaCoRe* against several machine learning models (Lasso, RF and XGBoost) and show that their algorithm does not sacrifice interpretability nor predictive accuracy.

# Implementation/Assessment

Recent years made it clear that machine learning is a tool that should be available to all biologists, but comes with high complexity and its own pitfalls, even without the addition of mathematical characteristics of compositional data and log-ratio transformations. Solutions like *CoDaCoRe* have been introduced by the scientific community, where they showed that they supposedly work more efficient than Machine Learning models (Gordon-Rodriguez et al. 2021). The goal of this project focuses on collecting insights on the performances of Machine Learning models in direct comparison with *CoDaCoRe*, but also assessing the influence of log-ratio transformations on Machine Learning performances in more detail. Additionally, some practical guidelines will be created on how to effectively combine Machine Learning and compositional data.

As mentioned in the introduction, two log-ratio transformations seem to be the most promising, in terms of applicability and interpretability: ALR and CLR. Additionally, it has been decided to compare ALR transformation in three ways: (i) a random reference will be picked as denominator, to assess the average performance of machine learning models for ALR (random ALR), (ii) find the most optimal denominator (optimal ALR) and (iii) worst ALR denominator (worst ALR).

A standard normalization for raw relative abundances is total sum scaling transformation (TSS). In theory, it removes technical bias related to different sequencing depth by dividing each feature count with the total library size. As mentioned in the introduction, Aitchison criticized these normalization techniques very early on and therefore its impact on Machine Learning performances will be directly compared to chosen log-ratio transformations. Finally, all Machine Diagram

Description automatically generatedLearning performances are compared to *CoDaCoRe* in the following conceptual framework:

Figure 5: Pipeline to compare model performances

The figure shows the proposed pipeline for effective comparison of machine learning performances. Data sets used are chosen by their characteristics large/small, high/low correlations and continuous/discrete variables. Afterwards, data sets will be filtered by abundances and imputed. Microbiome-native methods will be employed and compared to the data being log-transformed and used in machine learning models.

The general pipeline will be constructed of the following building blocks: Pre-processing, Imputation, Transformation, and Machine Learning Models/Microbiome Approaches. Usually, model performances are averaged after several repeats with random training and test splits. However, it was decided here to use the same training/test split for all transformations to directly compare the difference.

As representation for Machine Learning models, Generalized Linear Model (*glmnet*) and Support Vector Machine (*svmRadial*) are used. *Glmnet* represents a common linear machine learning model and *svmRadial* a non-linear one, as several papers already showed different performances for microbiome data depending on the choice of model (papers). -> svm does not care about data size

For *CoDaCoRe*, it was decided to train two models per run, one with lambda = 0 and the other with lambda = 1. In *CoDaCoRe* the parameter lambda controls the regularization strength of the model. Lambda = 1 applies the 1-standard-error rule in the discretization step of the log-ratio. This is typically a good choice, leading to models that are both sparse and predictive. Lambda = 0 corresponds to a 0-standard-error rule, in other words choosing the log-ratio that minimizes cross-validation score (Gordon-Rodriguez et al. 2021). Using both models allows for a direct comparison in how *CoDaCoRe* handles different types of microbiome data sets, that are typically very sparse. Additionally, *CoDaCoRe* allows in prediction tasks to include all found log-ratios or just the most descriptive one. It was decided to use both to see how influential one found log-ratio can be for the chosen data sets or if it is better to use several log-ratios.

Performance assessment scores are necessary to compare all models directly. For discrete parameters (e.g., two binary classes like “healthy” and “non-healthy”) AUROC will be used to assess model performances. AUROC tells about the model’s ability to discriminate between cases (positive examples) and non-cases (negative examples). An AUROC above 0.8 generally means excellent discriminatory ability for the model, whereas an AUROC of 0.5 corresponds to a coin flip (i.e., a useless model).

For continuous parameters (e.g., BMI, which can have any real number) a root mean squared error (RMSE) is used for performance evaluation. It is the [standard deviation](https://www.statisticshowto.com/probability-and-statistics/standard-deviation/) of the [residuals](https://www.statisticshowto.com/residual/). Residuals are a measure of how far from the regression line data points are and a RMSE consequently measures how spread out these residuals are. A value of 0 would indicate a perfect fit to the data. Contrary to the AUROC, the lower the RMSE, the better the model.

As several authors pointed out (Quinn and Erb 2020; Gloor et al. 2017), machine learning performance is influenced by data size. Therefore, three data sets were chosen accordingly to include direct comparison of performances of small and large data sets, as well as high and low known correlations between microbiome and host. There is no explicit definition for small datasets in the literature. However, several studies classified data sets ranging from 18 to 1030 samples as small data sets (Althnian et al. 2021) and this guideline is used here, too.

This leads to two small (CRC and PCOS) and one large data set (EstMB) for this project. The CRC data set was first used and described by (Wirbel et al. 2019) in their meta-analysis for colorectal cancer. This data set is well known and contains 7727 features with 695 samples. It shows clear correlations between gut microbiota and colorectal cancer and is therefore helpful to show the behavior of transformations and machine learning algorithms on small but highly specific data sets. The second data set is the Polycystic Ovary Syndrome (PCOS) data set described by Kreete et al. (2020). It observed 312 individuals, with two-thirds of them being healthy, and 72738 features. It is a valuable addition as it is a small data set that shows no correlation between the disease and microbiome structure (paper Kreete). Lastly, is the Estonian Biobank microbiome cohort (EstMB). This data set includes 2509 individuals with several phenotypical markers collected over time and 17180 features overall, which classifies it as a big data set and it contains two phenotypes that are of value for this project: Hypertensive heart disease (HHD) which has a lower to moderate correlation with gut microbiota and Diabetes Type 2 (DT2) which is known to show a higher correlation to gut microbiome compositions (<https://doi.org/10.1038/s41586-022-04567-7>, other papers?).

It has been decided to remove the majority of zeros in all data sets via abundance filtering. This reduced the runtime of computational procedures and also by sub-compositional coherence, this should have no effect on the performance of machine learning models.

* Repeat PAP
* All features Genus in all data sets?

# Methodology

For the data analysis and model pipelines, the script language R (v4.1.3) in combination with RStudio (v2022.02.1+461) has been used. For data cleaning and filtering the main library is “tidyverse” (1.3.1). Imputation was conducted with “zCompositions” (1.4.0.1) (cmultRepl), and transformations were mostly done with “easyCODA” (0.34.3). Models were constructed with “mikropml” (1.2.2), “tidymodels” (0.2.0) and “codacore” (0.0.3).

Additionally, scripts were created for convenience purposes. All scripts can be found on Github JenniferNeumaier/ml\_coda.

## Pre-Processing

First, all data sets were cleaned in order to remove NAs in predictor columns or patients that have no sequencing data. In EstMB data set, 21 rows removed in metadata due to NA and 21 patients respectively cut out of abundance table. This leads to 2485 final sample-size. In CRC, 128 rows were removed due to NA in feature “BMI”, leading to 567 samples overall. In PCOS, 6 rows were removed in the abundance table because no matching patient has been found in metadata, reducing the number of samples to 304.

As discussed, the sparse nature of microbiome data leads to several problems. Therefore, filters were applied to all three data sets. In this benchmarking project, taxa with ≤10% abundance in samples will be discarded. Additionally, a filter of ≤50% abundance in samples will be applied, as well as a mean relative abundance filter for 0.001. For 10% abundance filters CRC keeps 650 features and PCOS 1154 features. Respectively, for 50% abundance filters CRC keeps 189 features and PCOS 120 features. For EstMB data, 90% abundance was used, as the data was otherwise not practically usable without heavy computational power. Even with 90% filtering, EstMB keeps 3062 features and with 95% filtering 2589 features.

## Imputation

One of the first steps after filtering and before log-ratio transformation is zero-imputation. Introduced by (Palarea-Albaladejo and Martín-Fernández 2015) is the library “zCompositions”. All three data sets were imputed with Geometric Bayesian Multiplicative (GBM) and output form “p-counts”.

## Transformations

Greenacre et al. (2021) proposed a computational solution of finding an ALR reference. Included in the package “easyCODA” is the function ALR() that assesses the abundances and variances of features in a data matrix, followed by a Procrustes analysis to assess their geometry. This leads to a list of possible good denominators for the respective data set if the top results are chosen or worst denominators, if the bottom results are selected. Similarly, “easyCODA” also contains the function CLR() to compute the centered log-ratio. For TSS() a custom function has been created.

## Machine LEarning models

The focus will be on two standard machine learning models, that are already incorporated in easy-to-use packages in R: generalized linear models (GLMs) and support vector machines (SVM) as non-linear approach. Both are available in the function run\_ml() from the package “mikropml” by Schlosslab (paper). This package nicely compacts the use of standard machine learning models to a few lines of code and supports the use of GLMs (*glmnet*), as well as SVMs (*svmRadial*).

“Tidymodels” was used split the data into 80% training set and 20% test set. A seed was used to ensure that the data splits were the same for every repeat and following test. The split data has then been fed into run\_ml() and codacore(), respectively. Both functions then split the training set into 5 folds and assessed via 10-fold repeated cross-validation the best model. Here, run\_ml() also tunes hyperparameters for glmnet and svmRadial, respectively. In *CoDaCoRe* lambda is not tuned, but instead two models are trained, one with lambda = 0 and the other with lambda = 1.

The best model then is used for predictions on the test set and its AUROC/RMSE scores saved. The functions run\_ml() and codacore() were repeated 10 times, which allows to assess statistical fluctuations in a specific data split.

For svmRadial the number of repeats had to be lowered, otherwise the training of the model would have taken too long. It has been decided to use a 2-fold repeated cross-validation and only 5 repeats overall.

# Results

## Influence of Transformations on ML Performances

The most important research question of this thesis was if and how log-ratio transformations influence machine learning performances. The in section () introduced pipeline has been applied to all three data sets and the results separated into binary classification and regression setting results.

### Model performances in a Binary Classification Setting

Figure 6 shows the culminated results of the presented pipeline and combined in one figure to show the differences of machine learning and transformations on several data sets directly.

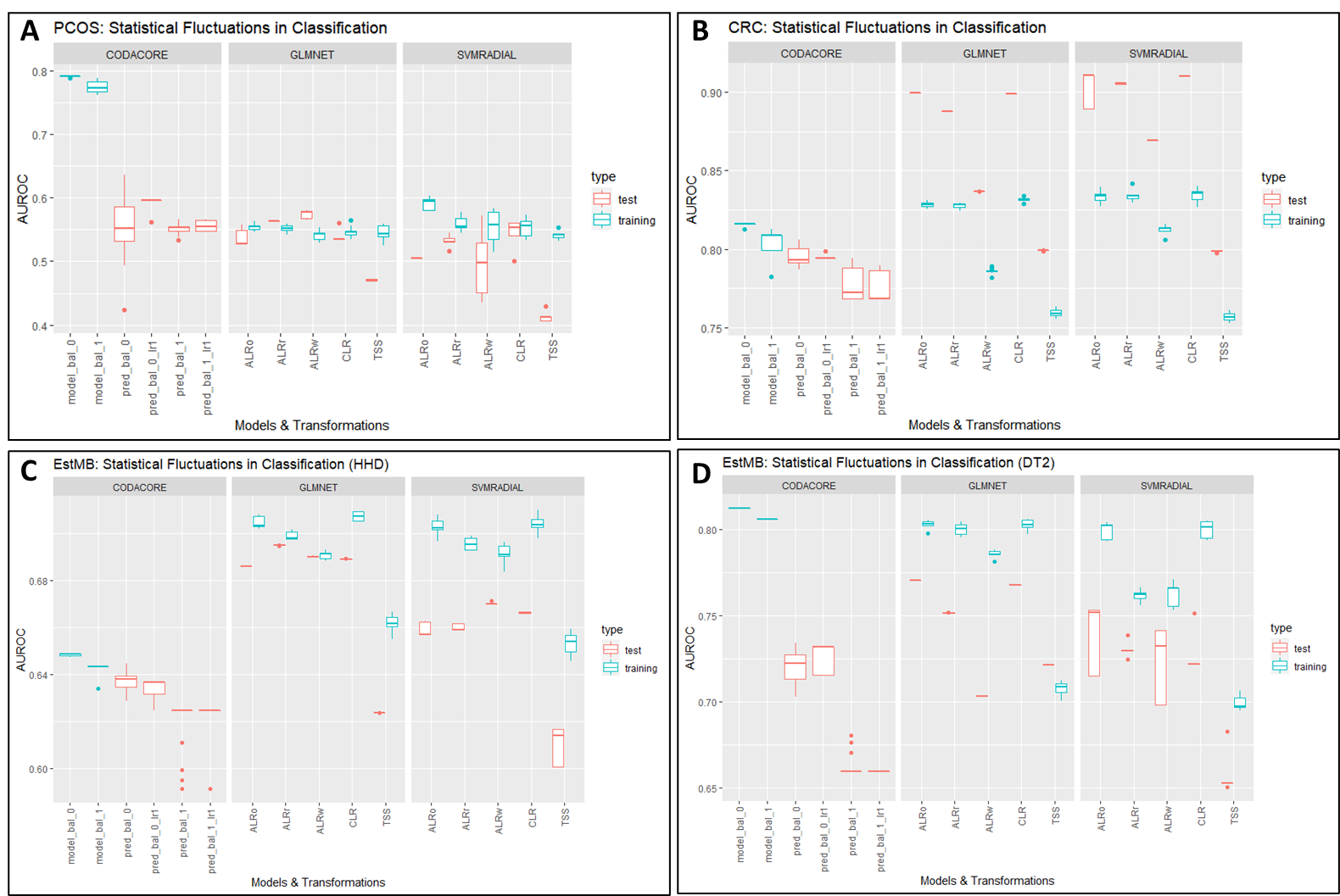


Figure 6: Statistical Fluctuations in Classification

CRC and PCOS use 10% abundance filtered sets, EstMB uses a 90% abundance filtered set. The y-axis contains AUROC scores. The x-axis contains models and transformations, with the most left always being *CoDaCoRe*, the middle portion displaying performances of glmnet and the right svmRadial. In red test performances are plotted and in blue training performances. model\_bal\_0 and model\_bal\_1 correspond to *CoDaCoRe* models with lambda 0 and lambda 1, respectively. Consequently, pred\_bal\_0 and pred\_bal\_0\_lr1 correspond to predictions with all found log-ratios and only the most descriptive log-ratio for model\_bal\_0. The same nomenclature applies to pre\_bal\_1 and pred\_bal\_1\_lr1. ALRo, ALRw and ALRr correspond to the types of ALR transformation applied in this pipeline (optimal, worst and random). A contains data from PCOS data set predicting PCOS phenotype. B contains data from CRC data set predicting CRC phenotype. C and D both contain data from EstMB data set, with C predicting for hypertensive heart disease (HHD) and D for diabetes type 2 (DT2).

For the glmnet models, training data was split into 5 folds, a 10-fold repeated cross-validation has been applied and the models repeated 10 times. For the svmRadial models, training data was split into 5 folds, a 2-fold repeated cross-validation has been applied and the models repeated 5 times. For *CoDaCoRe*, training data was split into 5 folds, a 2-fold repeated cross-validation has been applied and the models repeated 5 times.

As expected for PCOS (Fig.6A), the AUROC scores for test and training performances vary from 0.5 to 0.6 and correspond to the low correlation between microbiome and PCOS. *CoDaCoRe* severely overfits the data set with the test performances corresponding to the performances of both *glmnet* and *svmRadial*.

Contrary to that, Fig. 6B shows the performance of models in a data set with a high correlation between the microbiome and colorectal cancer. All models show good to excellent performances, with ALRo and CLR showing the best performances. ALRr shows surprisingly good results, which is repeated in Fig. 6C and D for the EstMB data set. Although log-ratio transformation not necessarily improve the predictive performance in low correlation data set (Fig. 6A), TSS still shows the lowest performances in all three data sets, independent of correlation or sample size. This trend is also visible in regression settings for all three data sets (see Supplementary).

It is surprising, that *CoDaCoRe* shows lower test and training performances, comparable to ALRw glmnet (Fig. 6B) or TSS (Fig. 6C). In general, its results are very inconclusive throughout different types of data. In the PCOS data set (Fig. 6A) it visibly overfits the data, although its test performances are comparable to glmnet and svmRadial. This trend is also visible in regression settings (see Fig. 10 Supplementary). In high correlation data sets, independent of sample size, it does not manage to perform better than Machine Learning models, performances are still considered good to excellent with an AUROC of 0.8 (Fig. 6B and 6D).

In the CRC data set, all machine learning models severely underfit the data, which usually indicates a bias in the data. A possible reason could be the “leakiness” in the pipeline. It is considered good practice to not conducted pre-processing on the whole data set (Whalen et al. 2022). Although imputation and transformation are handled independently from pre-processing in this pipeline, both can be considered part of pre-processing. Indeed, section () shows that splitting the data set before imputation and transformation solves the underfitting in CRC.

Still, the trend depicted in CRC with ALR and CLR performing best, is also found in EstMB data sets, for both moderate correlation (HHD) and high correlation (DT2) between microbiome and disease, and to a small degree in PCOS. However, linear models seem to perform better than non-linear, although that difference is smaller in high-correlation data sets.

~~For HHD~~ *~~CoDaCoRe~~* ~~shows lower performances, corresponding to TSS performances in glmnet and svmRadial. In DT2~~ *~~CoDaCoRe~~* ~~visibly overfits and also shows a big discrepancy between lambda 0 and lambda 1 predictions, with lambda 0 models showing better performances.~~

~~Interestingly, in all data sets and all models that use different transformations, TSS seems to be clearly underperforming, closely followed by ALRw. This is especially visible in high correlation data sets like CRC and EstMB DT2. However, also in low (PCOS) and moderate correlation (EstMB HHD) data sets, models using TSS normalized data significantly underperforms~~. Considering the effect of transformations even in PCOS, it seems to be a clear indication, that log-ratio transformations can indeed affect the performance in machine learning models. Additionally, *CoDaCoRe* only performs as good as log-ratio transformed data, but never better.

## INFLUENCE OF Transformations on Data Leakage

As mentioned in the section above, data leakage is an easy to overlook problem and already lead to performance biases in the section before, seemingly more in smaller data sets. Information leakage occurs when information from the test set is used to pre-process the training set. This is for example happening when the imputation algorithm uses the whole data set, which includes the test set, to replace zeroes (Whalen et al. 2022). To assess the impact of data leakage when adding log-ratio transformations, this section directly compares the model performances of *glmnet* and *CoDaCoRe* of a leaky and non-leaky pipeline.

In theory, the chosen log-ratio transformations ALR and CLR should not lead to any kind of data leakage. CLR transformations are conducted row-wise and therefore do use information between samples. For ALR, if the analysis for the best reference is conducted on the training set and applied on the test set, then data leakage should also not occur. The same applies if a random reference is chosen.

In the non-leaky procedure, processed data is split into train and test set at the start and imputation and transformations are applied separately on the split data. In the case of ALR, the reference was found on the training set and directly applied to the test set. Finally, train and test set are fed into *glmnet* model. The non-leaky procedure for *CoDaCoRe* works similarly but the transformation step is skipped.

In the leaky procedure, imputation and transformations are conducted on the whole data set and afterwards the data is split into train and test set and fed into the *glmnet* model. Again, for *CoDaCoRe* the transformation step is skipped.

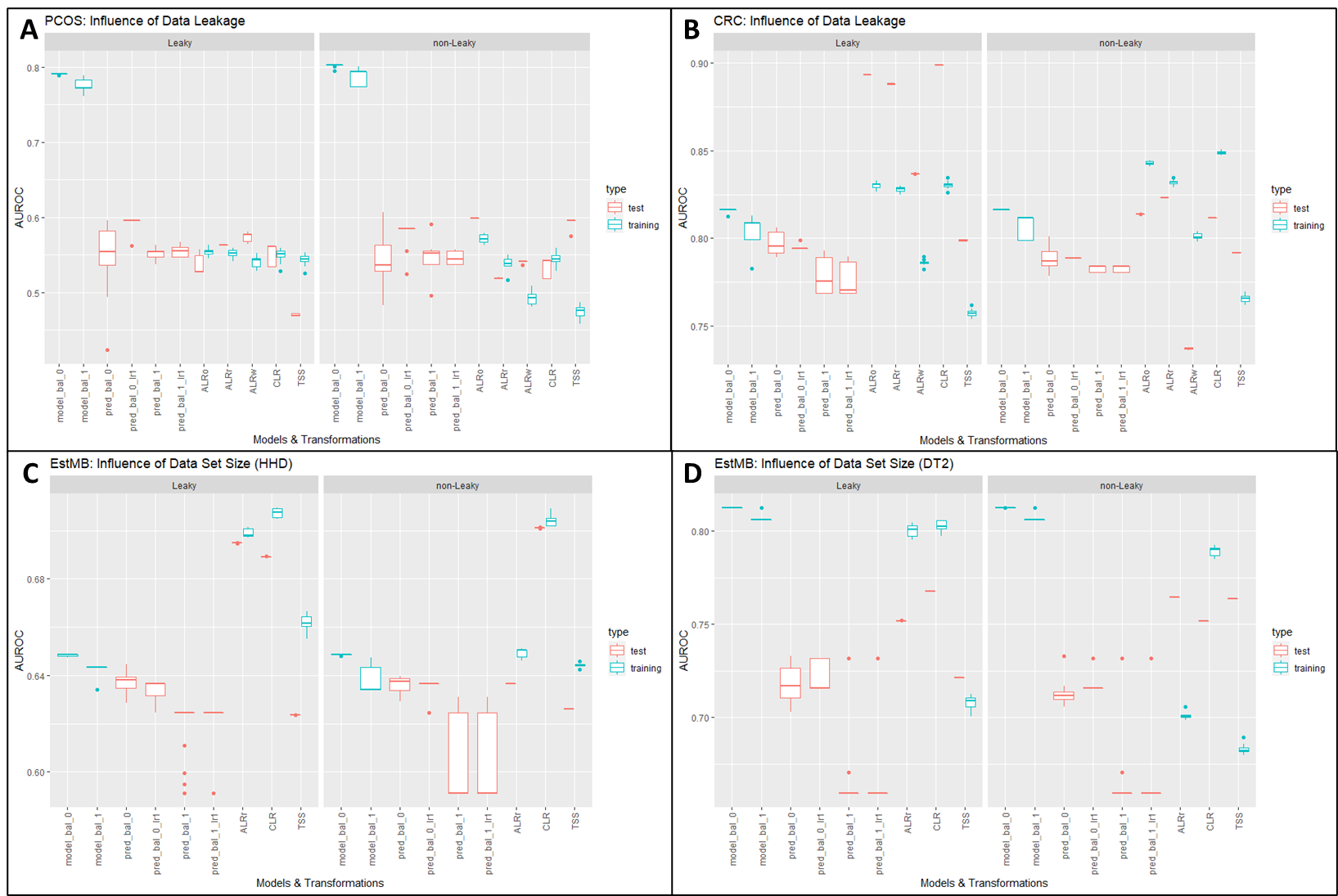


Figure 7: Influence of leaky and non-leaky pipelines on log-ratio transformation

CRC and PCOS use 10% abundance filtered sets, EstMB uses a 90% abundance filtered set. The y-axis contains AUROC scores. The x-axis contains models and transformations, with the most left always being *CoDaCoRe,* followed by *glmnet* results. For EstMB ALRo and ALRw have been skipped because it was too computationally taxing to repeat a Procrustes Analysis. In red test performances are plotted and in blue training performances. model\_bal\_0 and model\_bal\_1 correspond to *CoDaCoRe* models with lambda 0 and lambda 1, respectively. Consequently, pred\_bal\_0 and pred\_bal\_0\_lr1 correspond to predictions with all found log-ratios and only the most descriptive log-ratio for model\_bal\_0. The same nomenclature applies to pre\_bal\_1 and pred\_bal\_1\_lr1. ALRo, ALRw and ALRr correspond to the types of ALR transformation applied in this pipeline (optimal, worst and random). A contains data from PCOS data set predicting PCOS phenotype. B contains data from CRC data set predicting CRC phenotype. C and D both contain data from EstMB data set, with C predicting for hypertensive heart disease (HHD) and D for diabetes type 2 (DT2).

For the glmnet models, training data was split into 5 folds, a 5-fold cross-validation has been applied and the models repeated 5 times.

The results for the leaky pipelines were not repeated and the description for the results can be read in section ().

It becomes apparent that the results for *CoDaCoRe* do not change significantly if imputation is conducted on the whole data set or not. For CRC (Fig. 7B), imputation on the whole data set potentially leads to higher variances in test performances compared to splitting the data before imputation. However, a non-leaky pipeline does not increase performances for *CoDaCoRe* and does not solve the overfitting.

The results for Machine Learning models are a bit more mixed. The results in Fig. 7B show that the underfitting in the CRC data set was most likely due to biases in the pipeline. As soon as a non-leaky was applied, the Machine Learning models are not underfitting the data anymore. The training performances for CRC are very similar to the leaky pipeline, but the test performances have been adjusted. A contrary picture is shown for TSS in PCOS and EstMB DT2 (Fig. 7A and 7D). There, the model is now heavily underfitting the data, whereas no significant changes are seen in Fig. 7B (CRC) and 7C (EstMB HHD) for TSS.

Some trends can be observed for all three data sets: CLR shows very stable performances (sans the underfitting in CRC), which is especially visible in EstMB performances. ALR and TSS are the most impacted by the changed pipeline. In all data sets TSS increases the difference between its test and training performance, independent if it has been under- or overfit the data. In Fig. 7A and B ALRo marginally underfits the data but otherwise shows the highest performances out of all transformations.

Even though a leaky pipeline has been applied in the section before, the trends applied by log-ratio transformations still holds. In both pipelines, log-ratio transformations outperform standard normalization techniques and *CoDaCoRe* is not outperforming standard Machine Learning models. Nevertheless, in all further tests, the non-leaky pipeline will be applied.

## Do log-ratio transformations influence test set methodology?

The pipeline used to compare all data sets and their performances includes a standard data set split of 80/20, meaning 80% of data goes into the training set, 20% of data into the test set. However, the CRC data set has been designed with holdout sets in mind and the original paper uses this technique (Wirbel et al. 2019).

In Machine Learning theory, a holdout differs from a test set as it provides a final estimate of the machine learning model’s performance after it has been trained and validated by a test set. In the CRC paper they called it leave-one-study-out (LOSO) validation, where data from one study was set aside as an external validation set, while the data from the remaining 4 studies was pooled as a training set (Wirbel et al. 2019).

The study design gives a good opportunity to see if the influence of log-ratio transformations is impacted by the choice of test set methodology. The following figure compares *glmnet* and *CoDaCoRe* performances with a standard 80/20 or Holdout set.

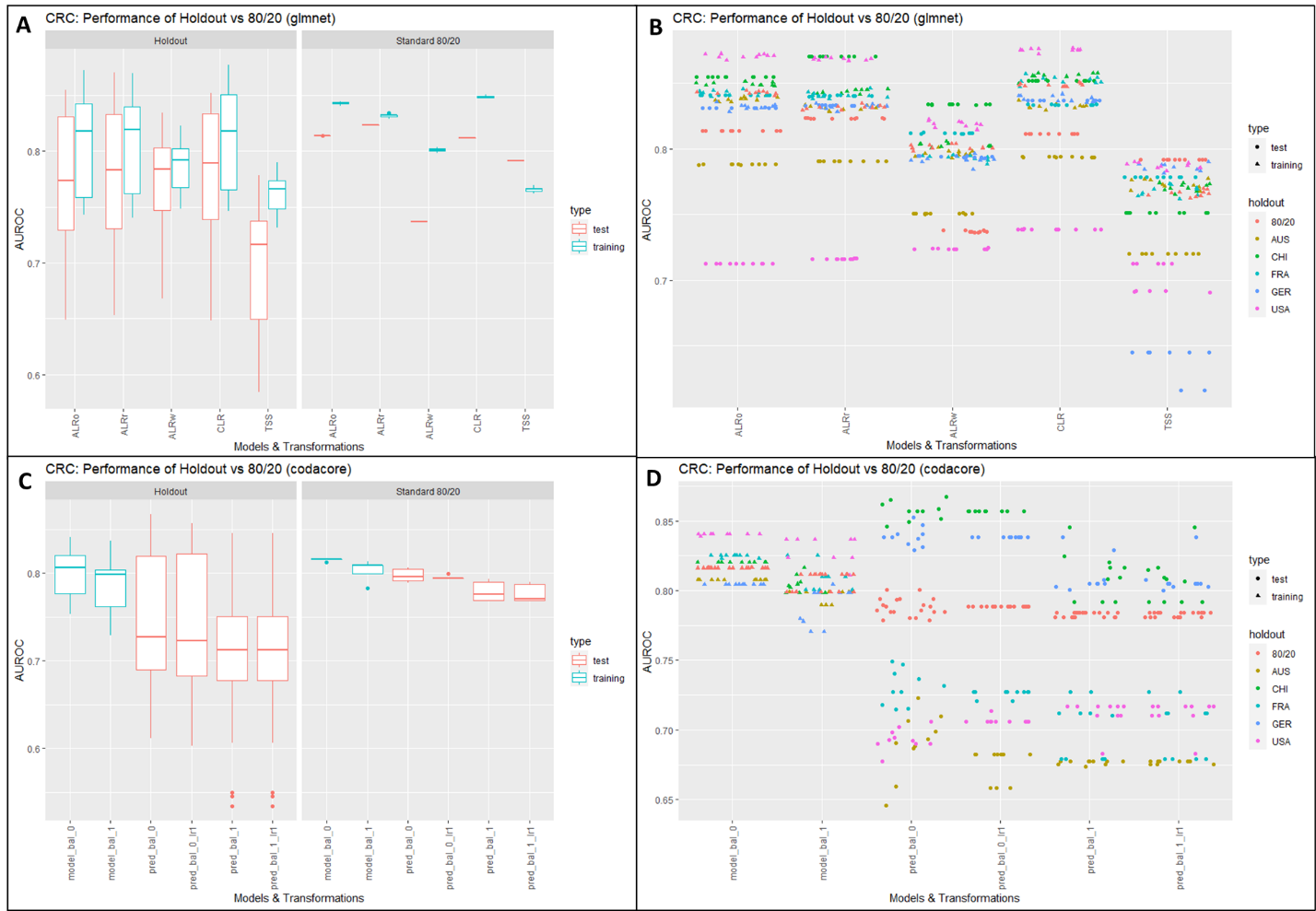
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Figure : Behavior of log-ratio transformations with test set methodology

CRC uses 10% abundance filtered set. The y-axis contains AUROC scores. The x-axis contains models and transformations, with A and B belonging to glmnet models and C and D belonging to *CoDaCoRe*. In red test performances are plotted and in blue training performances. model\_bal\_0 and model\_bal\_1 correspond to *CoDaCoRe* models with lambda 0 and lambda 1, respectively. Consequently, pred\_bal\_0 and pred\_bal\_0\_lr1 correspond to predictions with all found log-ratios and only the most descriptive log-ratio for model\_bal\_0. The same nomenclature applies to pre\_bal\_1 and pred\_bal\_1\_lr1. ALRo, ALRw and ALRr correspond to the types of ALR transformation applied in this pipeline (optimal, worst and random). A contains data from CRC data set predicting CRC phenotype and B plotting all test and training performances in detail. The same applies in C and D for CRC *CoDaCoRe* performances.

For the glmnet models, training data was split into 5 folds, a 10-fold cross-validation has been applied and the models repeated 10 times.

Fig. 8A and 8B show the difference of test performances as boxplots for different transformations of the same data set and for *CoDaCoRe*, respectively. All five holdout sets have been used separately as test set once and the results pooled. It can be seen clearly that for all transformations, the holdout sets show a lot more variance in the test set performances, compared to the standard 80/20 split, which shows almost no variance at all and arguably higher test performances. This is also visible for *CoDaCoRe* performances.

In Fig. 8B and 8D it becomes clear why the test performances of the holdout sets have such a high variance. For all transformations and holdout sets, the test performance is quite stable. Marked in pink dots is the test set performance when USA has been chosen as holdout set. For all ten repeats of the model, the test performance is stable around 0.72. However, choosing CHI as the holdout set produces test set performances of 0.85. This shows that the high variances in Fig. 8A are not due to statistical fluctuations, but the choice of holdout set. Splitting the data set in an 80/20 manner leads to a performance somewhere in the middle. This is also true when using *CoDaCoRe,* although the test set performances fluctuate more strongly.

Additionally, log-ratio transformations show a similar picture as in the sections above. The performances of ALRo, ALRr and CLR are higher than TSS and ALRw, even when using holdout sets. *CoDaCoRe* doesn’t change its performance.

~~The idea to verify a model by gathering extra test data is sensible, however as seen by the results, it easily introduces a bias and for beginners in machine learning it would be more beneficial to stick to 80/20 data splits.~~

## Data set size

As mentioned in the introduction, the dimensionality and sparse nature of microbiome data is problematic for both Machine Learning models and log-ratio transformations. A common technique to reduce the runtime for pre-processing and zeroes contained in the microbiome data, an abundance filter is applied, or the features are selected via ordination techniques (sources). For all tests above, a 10% abundance filtering has been used, as it is very common in microbiome and ecology (source) and the performances especially for highly correlated data seems promising.

However, it would be of great advantage if the number of features could be reduced even further, with Machine Learning performances staying stable. Therefore, the following test will assess how machine learning models perform if a 50% abundance filter is applied. Additionally, *CoDaCoRe* performances will be again compared to *glmnet*.

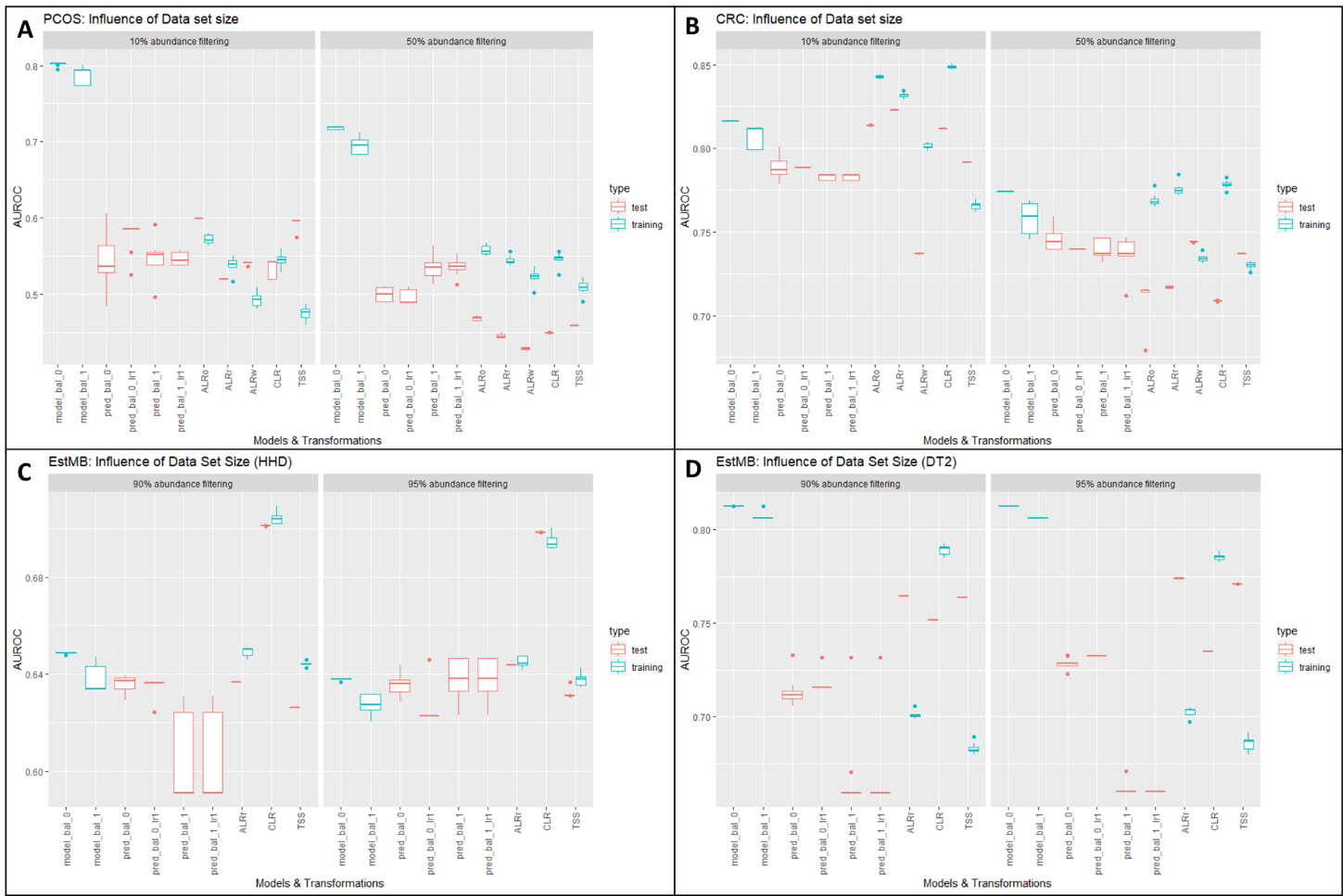


Figure 9: Influence of Data set size

CRC and PCOS use 10% and 50% abundance filtered sets, EstMB uses a 90%/95% abundance filtered sets. respectively. The y-axis contains AUROC scores. The x-axis contains models and transformations, with the most left always being *CoDaCoRe*, the middle portion displaying performances of glmnet and the right svmRadial. In red test performances are plotted and in blue training performances. model\_bal\_0 and model\_bal\_1 correspond to *CoDaCoRe* models with lambda 0 and lambda 1, respectively. Consequently, pred\_bal\_0 and pred\_bal\_0\_lr1 correspond to predictions with all found log-ratios and only the most descriptive log-ratio for model\_bal\_0. The same nomenclature applies to pre\_bal\_1 and pred\_bal\_1\_lr1. ALRo, ALRw and ALRr correspond to the types of ALR transformation applied in this pipeline (optimal, worst and random). A contains data from PCOS data set predicting PCOS phenotype. B contains data from CRC data set predicting CRC phenotype. C and D both contain data from EstMB data set, with C predicting for hypertensive heart disease (HHD) and D for diabetes type 2 (DT2).

For the glmnet models, training data was split into 5 folds, a 10-fold repeated cross-validation has been applied and the models repeated 10 times. For *CoDaCoRe*, training data was split into 5 folds and the models repeated 5 times.

The results for 10% abundance filters were not repeated and the description for the results can be read in section ().

Reducing the number of features has visible impacts on smaller data sets (Fig. 9A and B). For PCOS, *glmnet* is overfitting the data more pronounced compared to 10% filtered data sets and in CRC, the decrease in performance is clearly visible (Fig. 9B), accompanied by heavier overfitting.

The impact on bigger data sets is smaller, however. In Fig. 9C a 95% abundance filter leads to less distance between test and training performances compared to 90% filtering, whereas these distances increase slightly in Fig. 9D. In general. though the performances are very similar.

Again, although the performances in CRC (Fig. 9B) are decreasing, the impact of log-ratio transformations are still visible. ALRw and TSS still perform worse compared to ALRo, ALRr and CLR. Especially the latter is also very stable in EstMB data sets.

The performances of *CoDaCoRe* are not heavily affected, only in CRC do they decrease with a decrease in features.

# Discussion/Conclusion

The goal of this was to assess the influence of log-ratio transformations on Machine Learning performances, specifically in the context of microbiome data. Additionally, it has been tried to assess if some blocks in the Machine Learning pipeline are more affected by the addition of log-ratio transformation.

## Log-ratio Transformations

Overall, it can be stated that log-ratio transformation indeed increases Machine Learning performances, compared to standard normalization. These results are backed by several other studies, which showed similar results and recommended the use of log- or log-ratio transformation (Quinn and Erb 2020; Jasner et al. 2021; Kubinski et al. 2022). Furthermore, it can be stated that any choice of log-ratio transformation increases Machine Learning performances. Although only CLR and ALR have been tested in this study, recent studies have shown that ILR and several balancing models are similarly improving performances (Kubinski et al. 2022; Quinn and Erb 2020).

In low and moderate correlations, the increase in performances is only slight and shows mainly in less distance between test and training performances, which indicates that mapping the data in the Euclidean space does not conjure more correlations.

Furthermore, in high-correlation data the choice of transformation seems almost detrimental as CLR and optimal ALR show similar good performances. Choosing a random feature for ALR shows surprisingly good performances, which would support recent claims that strict-isometry may not be necessary for machine learning purposes (Greenacre et al. 2022). Therefore, choosing a random ALR feature could be very beneficial as it is not as computationally taxing as calculating the optimal ALR, especially in bigger data sets. It should be noted however, that random reference features could lead to performances as indicated by ALR worst, with a chance of 1/number of features. This probability becomes more redundant the bigger the data set and as an additional factor, the impact of a bad reference feature on the performance seems to decrease with sample size. In CRC, ALR worst shows performances close to TSS, whereas in EstMB data sets the performance is closer to the other ALR performances, independent of phenotype correlation. This would support the hypothesis by Greenacre et al. (2022) that deviation from isometry (and sub-compositional coherence) might become more diluted in higher-dimensional data and is not as necessary anymore. If this is not possible because of the sample size of the data set, actively choosing a reference feature (or choosing it randomly) does not need any data and can therefore be a valid option.

## Data Leakiness

The increase in Machine Learning performances after applying log-ratio transformations persists throughout other results in this thesis, and their influence seems to be impervious to changes in the pipeline. It has been hypothesized, that two factors could influence the data leakage in the first pipeline: imputation and log-ratio transformations. In theory, CLR-transformed data should not be affected is it is conducted sample-wise by nature. The results how that Machine Learning performances with CLR-transformed data shows very similar results compared to the leaky-pipeline, which supports the mathematical theory. In the case of ALR, if the reference feature is calculated on the training set, no information leakage should occur, however the results do not show conclusive results in that regard. Independent if a reference feature is calculated or randomly chosen, Machine Learning performances vary heavily between leaky and non-leaky pipelines, sometimes switching from underfitting to overfitting. More repeats for all three data sets with ALR transformations would be helpful in determining if this log-ratio transformation can be as stable as CLR in a non-leaky pipeline.

CLR results indicate that log-ratio transformations do not explain the significant underfitting in CRC data, which leads imputation seemingly to be the most impacting factor for data leakage in this pipeline. This is supported when comparing Machine Learning performances of data split before imputation and transformation with data split only before imputation, but transformations are conducted on the whole data set (see Supplementary). These results suggest that conducting log-ratio transformation on the whole data set was not the main driver of underfitting in the CRC data set.

Incidentally, although EstMB data sets were conducted with the same pipeline as CRC, the models did not overfit the data, especially when comparing both high correlation data sets. An explanation could be that in bigger data sets the bias introduced by assumptions from the imputation algorithm on the whole data set is significantly reduced, as more information are available for more detailed assumptions.

However, the scientific community is consistently discussing over the impact of imputation for any type of data analysis and this discussion transferred to the application of log-ratio transformations (Quinn et al. 2018; Greenacre et al. 2022; Gloor et al. 2017; Talwar et al. 2018). With the R library “zCompositions” (Palarea-Albaladejo and Martín-Fernández 2015) a tool has been created to impute zeroes with compositional data in mind. Alternative imputation algorithms exist, that facilitate Autoencoders like in (Talwar et al. 2018) or are designed specifically for sparse microbiome data like (Jiang et al. 2021). The goal of mbImpute is to provide an approach that tries to decide between technical and biological zeroes by leveraging additional information from sample covariates and taxon phylogeny. Additionally, recent papers also suggested that it could be possible to circumvent the problem of zero replacement altogether by applying Box-Cox power transformation or chi-square distances (Greenacre et al. 2022). Furthermore, data-driven alpha-transformations are currently investigated that enables one to deal with the presence of 0s in the compositions directly.

Further studies would be necessary to assess if there is a “best choice” in how to deal with zeroes in microbiome data, especially in smaller data sets.

## Machine Learning

Classification in Machine Leaning is a challenging task by itself, and it becomes more challenging when dealing with small datasets, as the limited size of training data can lead to unreliable and biased classification models (Althnian et al. 2021). This is important in biological contexts, as sample sizes are usually limited, and it should be accounted for when Machine Learning models are planned.

The CRC data set shows that high sample-sizes are not strictly necessary in high-correlation data sets, which is supported by several other studies in recent years (Althnian et al. 2021; van der Ploeg et al. 2014; Bailly et al. 2022). Especially for logistic regression models (like *glmnet*), smaller data set sizes are sufficient to achieve quick and good performances (Bailly et al. 2022). Experts are in agreement that an overall performance of classifiers depends how well the data set represents the original distribution rather than its size (Althnian et al. 2021) and that the most robust classifier is not necessarily the best one (Althnian et al. 2021; Bailly et al. 2022). However, it can be approximated that a number between 75–100 samples will usually be needed to test a good but not perfect classifier (Beleites et al. 2013). These results are supported by comparing PCOS and CRC Machine Learning model performances. The variances in training and test set performances in PCOS are very low, although the test set contains only 60 samples, which indicates that the algorithm is quite sure that it cannot find a correlation.

For non-linear models, it has been shown that SVM shows instabilities in smaller data sets and mentions that SVM may need over 10 times as many features per phenotype to achieve a stable AUC compared to linear models (van der Ploeg et al. 2014). Additionally, when comparing several Machine Learning models, non-linear models usually perform better than linear ones (Kubinski et al. 2022; Zhang and Shi 2019). This trend is not depicted in the results here, however further tests with e.g., XGBoost could correct the results. Considering that non-linear models are better equipped to deal with more features, they could be an answer to the sparse nature of microbiome data.

The impact of sample size on log-ratio transformation is hard to assess. Mathematically speaking, they shouldn’t, as compositions that are sampled from the same simplex space, e.g., disease group or control group, are equivalence classes.

Comparing CRC and EstMB (DT2) results it is clear that excellent Machine Learning performances in high-correlation data sets can be achieved independent of data size. But these results do not show if the AUROC scores could be higher with a higher sample size. An additional test with the EstMB data set could give more insight, by e.g., creating several differently sized data sets out of it and comparing the performances. If the performances increases with the sample size it could be an indicator that log-ratio transformations are also affected by sample sizes.

## What is codacore doing??

Although *CoDaCoRe* shows good results in high-correlation data like CRC and EstMB (DT2), it does not outperform Machine Learning models. In data sets with low to moderate correlation it severely overfits the data. These results set a contrast to the results in the paper that published *CoDaCoRe* (Gordon-Rodriguez et al. 2021)*.* Of course, it should be noted that here the performances with the same data split are directly compared, in contrary to the original paper, that evaluated the performances over 20 random train/test splits. However, the performances are also not corrected after applying a non-leaky pipeline as seen in section() and no bias has been introduced with 80/20 split (see section()).

A possible explanation for the different performances could be explained by another important variable for good Machine Learning performances: the number of features. As can be seen in section (), the performances for all models in small data sets show a decrease in performance after 50% abundance filtering, compared to 10% abundance filtering. Large data sets do not seem to be affected.

Sub-compositional coherence is – from a scientific point of view – a very important factor in compositional data. For the sake of reproducibility, it should be possible to get comparable results, even if only a sub-composition is chosen. This property is applied when data is filtered in pre-processing. Sub-compositional coherence should make it possible that statistical analysis results are similar. Greenacre et al. (2022) confirmed as much when they used several 50% sub-compositions from the same data set and showed that they had very low variability. In the case of abundance filtering, both data sets are sub-compositions of the original, with only keeping features that show up more abundant and should therefore inherit similar results when analyzing ratios. Contrary to that argument, when analyzing the calculated top 20 ALR references for CRC, the selected features differ greatly between 10% and 50% filter of the data set (see Fig.12A, Supplementary). This would indicate that perhaps the Procrustes analysis is skewed when applying it to different sub-compositions.

The drop in model performance is also true for *CoDaCoRe*, which is especially interesting as a qualitative analysis (see Fig.12B, Supplementary) shows that in both 10% and 50% filtering for CRC data sets *CoDaCoRe* finds very similar and overlapping log-ratios. This suggests that the algorithm for *CoDaCoRe* is indeed quite stable, independent of feature size and also contrary to ALR references. Additionally, calculating correlation coefficients (see Fig.11A, Supplementary) shows that only a small number of correlation coefficients show significantly different values between 10% and 50% filtering, which suggests that compositional data properties are still adhered to.

It raises however the question if these found log-ratios are predictive for the phenotype. Throughout high-correlating data sets the model applying a 1-standard-error rule did not perform significantly better than its counterpart, which means that the found log-ratios were not very predictive to begin with. In theory, not many samples are needed to achieve good predictive performances and in computational sciences Machine Learning models are frequently trained on only a few features.

However, the results hint that feature sizes perhaps need to be assessed as an independent parameter in microbiome data and are a very Machine Learning specific problem. Some rules of thumb have been proposed uncorrelated features, the optimal feature size may be N−1 (with N = sample size) and for highly correlated features (Hua et al. 2005). However, these recommendations come from the computer science community and may not be applicable to biological models.

But more concretely, 10% abundance filters leave 650 features for CRC and 1154 features for PCOS. Respectively, for 50% abundance filters CRC keeps 189 features and PCOS 120 features. Applying a 90% abundance filter on EstMB data set leaves 3062 features and applying 95% filtering leaves 2589 features.

The data sets that perform well (CRC, EstMB) both have more symmetric data sets after filtering than PCOS, with the number of samples comparably close to feature sizes. Furthermore, 10%/50% abundance filters lead to a loss of over 90%/97% of features for CRC and over 98%/99% for PCOS, respectively. In EstMB the loss is only 83%85% for 90%/95% filtering, respectively. Maybe those parameters should be tested more in detail to achieve stable Machine Learning performances.

In summary, filtering has a practical approach, as it reduces computational time drastically and without filtering, the discussion around imputation becomes more complicated, as imputation algorithms fail when features contain only zeros and using microbiome-specific tools is also not possible, as e.g., *CoDaCoRe* cannot work with zeros.

Suggestions for a more targeted feature selection besides abundances before Machine Learning have been proposed, also in compositional data. Most of them target the search for main factors that maximally explain log-ratio variances (Greenacre et al. 2022). However, an argument can be made that such techniques have been used in this project with *CoDaCoRe* and also ALR reference selection. As shown, *CoDaCoRe* does not improve performances although it supposedly found predictive balances and calculating ALR references is indicated to be affected by the sub-composition.

~~These findings suggest that the drop in performance in CRC may be of quantitative nature, instead of qualitative. Machine Learning needs a lot of data, and this seems to hold true even if they are highly predictive. Furthermore, all conducted tests also suggest that simply applying compositional data theory correctly does not solve classic Machine Learning problems.~~

# Conclusion

Regardless of the mentioned problems in the section before some practical key notes on transformations in the context of Machine Learning can be extracted from this thesis: (1) Transformations are not able to conjure correlations when there are none, (2) any log-ratio transformation is better than none.

This comes with several advantages when combining compositional data with Machine Learning:

Log-ratio transformations can be easily integrated into existing pre-processing procedures and if the data set is too small to conduct log-ratio transformations separately on training and test set, choosing ALR or CLR circumvents this problem easily. Otherwise, any type of log-transformation works, as several studies have already shown (Jasner et al. 2021; Greenacre et al. 2022), and the library “easyCODA” makes many log-ratio transformations easily applicable. Furthermore, even in pooled data sets and additional steps like batch corrections log-ratio transformed data performs better than classically normalized one, which suggests that log-ratio transformations could also work in highly individualized pipelines (Kubinski et al. 2022).

The previous section indicated that the majority of inconsistencies found in this thesis was due to the character of Machine Learning and microbiome data and not necessarily introduced by log-ratio transformations.

As Imputation and Filtering methodology seem to be the most prevalent problems in Machine Learning, further tests would be necessary, if those steps perhaps change the geometry of compositional data. Calculating correlation coefficients before and after imputation (see Fig.11B, Supplementary) does not suggest that imputation changes sub-compositions drastically, but perhaps other properties are affected, and new weight has recently been placed on distributional equivalence. Furthermore, it has already been shown to improve Machine Learning performances when genus level representation was used and all features belonging to the same genus merged through PCA before Machine Learning (Jasner et al. 2021). Additionally, data-driven alpha-transformations are currently investigated that enables one to deal directly with the presence of 0s in the compositions (Clarotto et al. 2022).

Additional tests on specific blocks of the Machine Learning pipeline for microbiome data are necessary, but the results concerning log-ratio transformations inspire confidence and do not seem to add additional difficulties.

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# Supplementary

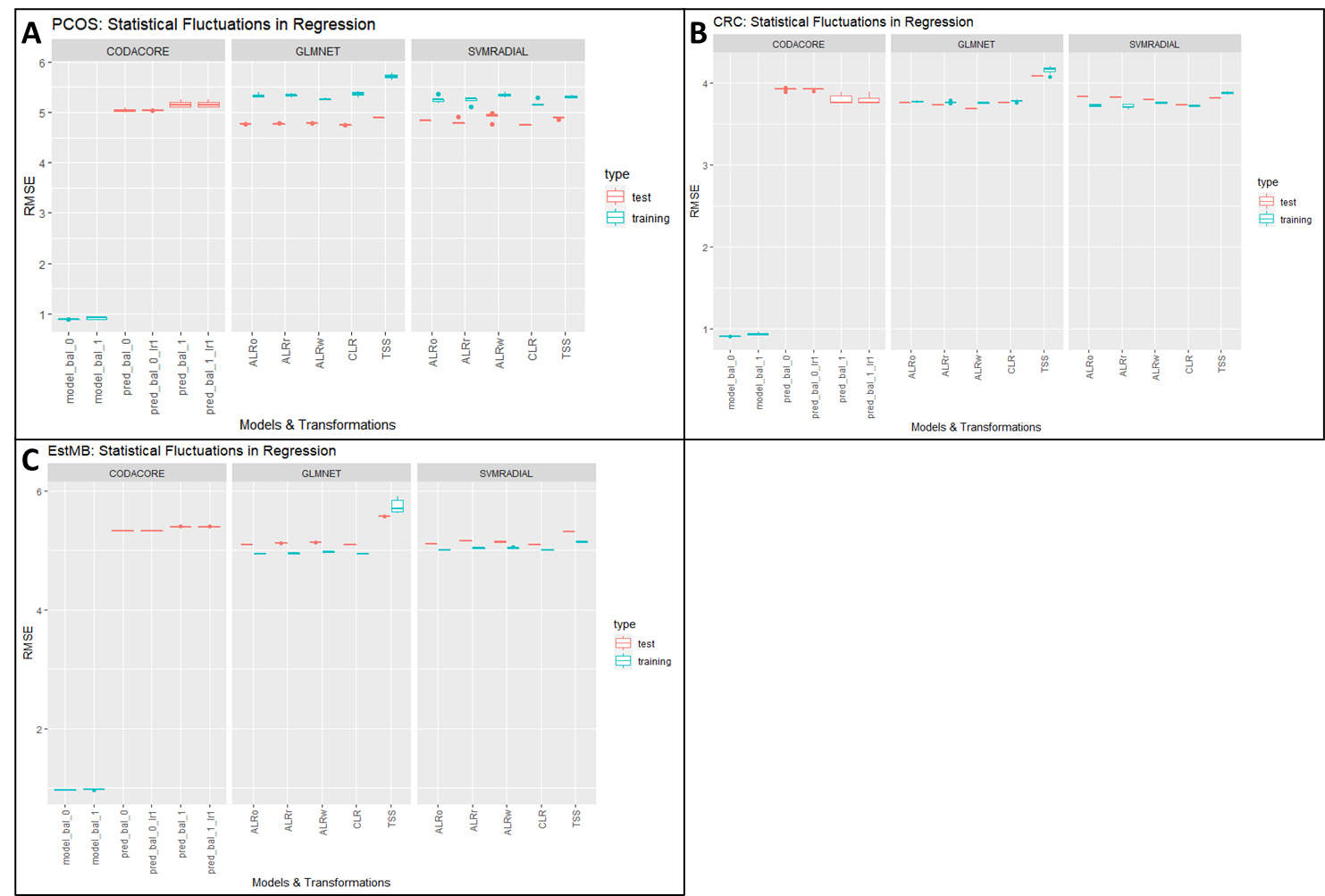


Figure : Statistical Fluctuations in Regression

CRC and PCOS use 10% abundance filtered sets, EstMB uses a 90% abundance filtered set. The y-axis contains AUROC scores. The x-axis contains models and transformations, with the most left always being *CoDaCoRe*, the middle portion displaying performances of glmnet and the right svmRadial. In red test performances are plotted and in blue training performances. model\_bal\_0 and model\_bal\_1 correspond to *CoDaCoRe* models with lambda 0 and lambda 1, respectively. Consequently, pred\_bal\_0 and pred\_bal\_0\_lr1 correspond to predictions with all found log-ratios and only the most descriptive log-ratio for model\_bal\_0. The same nomenclature applies to pre\_bal\_1 and pred\_bal\_1\_lr1. ALRo, ALRw and ALRr correspond to the types of ALR transformation applied in this pipeline (optimal, worst and random). A contains data from PCOS data set predicting PCOS phenotype. B contains data from CRC data set predicting CRC phenotype. C and D both contain data from EstMB data set, with C predicting for hypertensive heart disease (HHD) and D for diabetes type 2 (DT2).

For the glmnet models, training data was split into 5 folds, a 10-fold repeated cross-validation has been applied and the models repeated 10 times. For the svmRadial models, training data was split into 5 folds, a 2-fold repeated cross-validation has been applied and the models repeated 5 times. For *CoDaCoRe*, training data was split into 5 folds, a 2-fold repeated cross-validation has been applied and the models repeated 5 times.

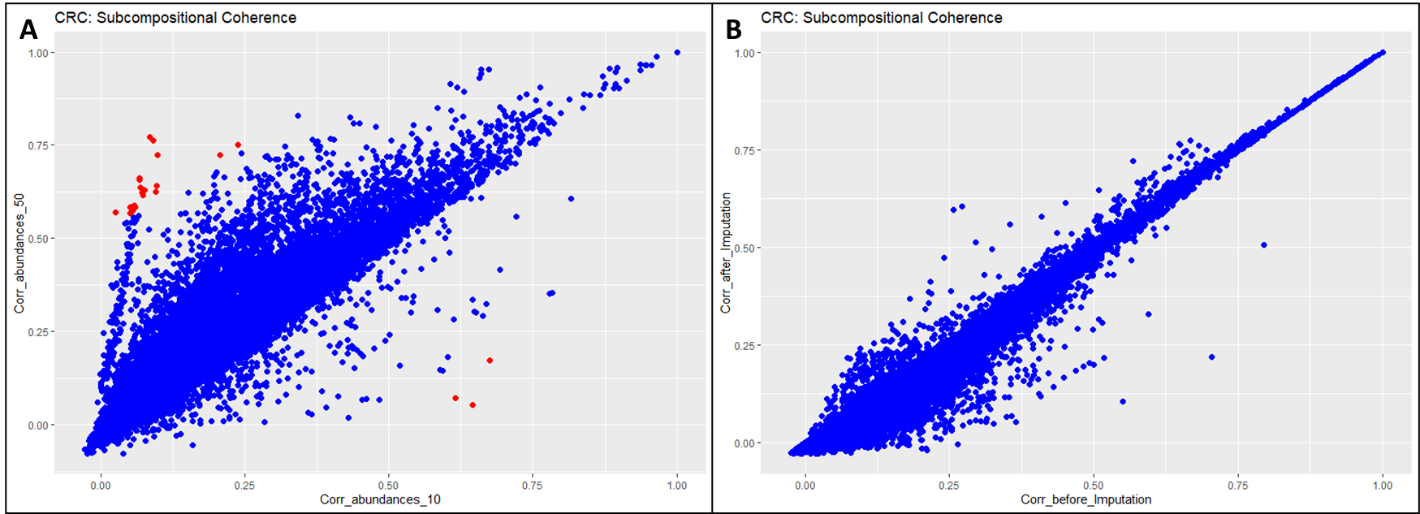


Figure 11: Correlation coefficients for CRC

Adapted from [https://towardsdatascience.com/relative-vs-absolute-understanding-compositional-data-with-simulations](https://towardsdatascience.com/relative-vs-absolute-understanding-compositional-data-with-simulations-fdc15e0c781e?gi=3a154a1a19fc). A shows scatterplot to assess sub-compositional coherence in CRC data set between 10% and 50% abundance filtered data sets. B shows scatterplot with correlation coefficients before and after imputation for 10% abundance filtered data set. Red marked dots indicate a change >0.5 in correlation coefficients.

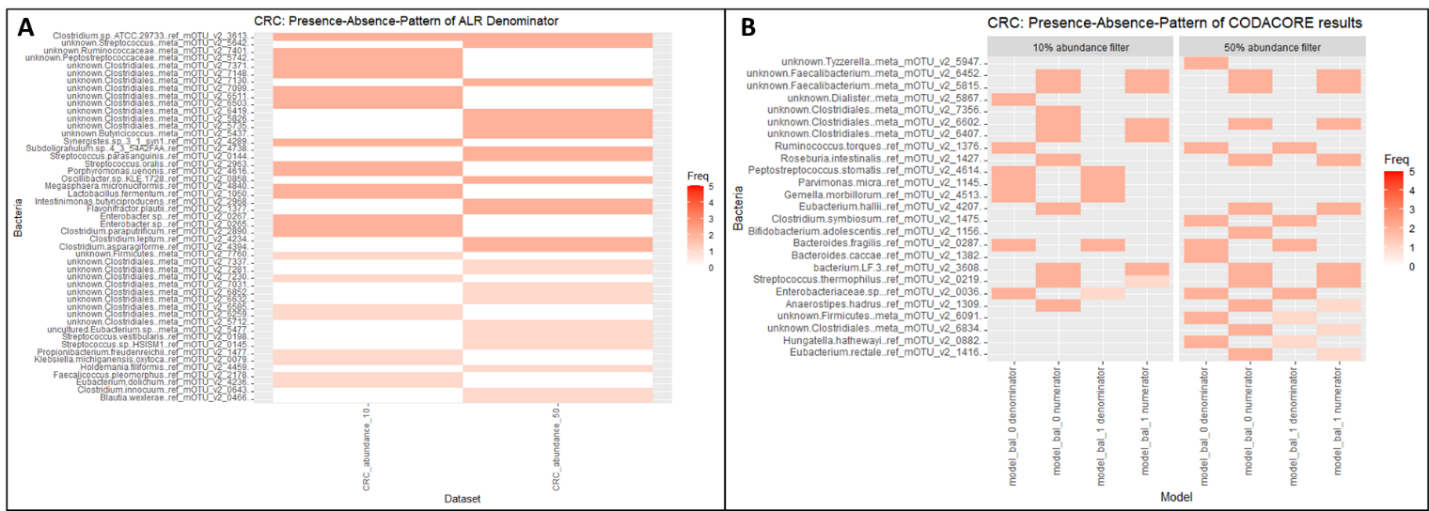


Figure 12: Presence-Absence-Patterns for CRC

A shows Presence-Absence-Pattern to show similarities and differences in calculated top 20 ALR references for CRC data set for differently sized data sets. B shows found numerators and denominators for both model types by CoDaCoRe also for differently sized CRC data sets.