PCRedux package - an overview

The PCRedux package authors 2019-10-31

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1 Aims of the Project

A review of the literature (PubMed, Google Scholar; 1984-01-01 - 2019-10-31) and discussion with peers revealed that there is no open source software package to calculate predictors from quantitative PCR amplification curves for machine learning applications. A predictor is a quantifiable *informative* property of an amplification curve. In particular, there is no information available about predictors that can be used from amplification curves apart from measures that describe quantification points, amplification efficiencies and signal levels. Although several amplification curve data sets are available, no curated labeled data sets labels are described in the literature or repositories such as GitHub¹, Bitbucket², SourceForge³ or Kaggle⁴.

Therefore, the aim of the study was to:

- 1. create a collection of classified amplification curve data,
- 2. propose algorithms that can be used to calculate predictors from amplification curves,
- 3. evaluate pipelines that can be used for an automatic classification of amplification curves based on the curve shape and
- 4. to bundle the findings in a public repository open source software and open data package.

2 Introduction to the Detection and Quantification of Nucleic Acids and the Relation to Machine Learning

In subsection 2.1 the reader finds a concise introduction to nucleic acids, including nucleic acid detection methods for the analysis of forensic sample material. Special intention is paid to the quantitative Polymerase Chain Reaction (qPCR), since this method is the *de facto* standard for the detection and high-precision quantification of nucleic acids.

The focus of this study is the development of statistical and bioinformatical algorithms for the PCRedux software (version 1.0.2). This software can be used to automatically calculate putative predictors (features) from qPCR amplification curves. A predictor herein refers to a quantifiable informative property of an amplification curve, employable for data mining, machine learning applications and classification tasks.

¹https://github.com/

²https://bitbucket.org/

³https://sourceforge.net/

⁴https://www.kaggle.com/

The subsection 3.1 deals with elements of the software engineering (e. g., continuous integration, Donald Knuth's *Literate Programming*, unit testing) used within the PCRedux software. The subsection 3.2 gives an introduction into qPCR data, their analysis and explains why there is a need for the PCRedux software. In addition, the data analysis using machine learning is concisely described, after which the work focuses on the analysis of the measured data.

The proposed algorithms were partially tested with machine learning methods. For this purpose, a brief introduction to the subject *machine learning* is given in subsubsection 3.2.1, and results from the belonging chapters are applied.

The details for statistical analyses of qPCR amplification curves are presented in subsubsection 3.2.4 ff. This covers the description of the curvature and the challenges of the calculations.

All scientific and engineering work depends on data. In particular, open data are becoming a cornerstone in science. As data sets of classified amplification curves were not available anywhere else, subsection 3.5 summarizes the aggregation, maintenance, and distribution of classified qPCR amplification curve data sets. The manual classification of amplification curves is a time-consuming and error prone task when working with large data sets. To facilitate the manual analysis procedure, helper tools are presented in subsection 3.4. In particular, a novel approach for curve-shape based group classification is shown.

An achievement of this study is the extensive portfolio of statistical algorithms for predictor calculation. Central findings of the research are presented in subsection 3.3.

It is expected that these implementations will allow the automatic analysis of large data sets for machine learning applications. The expectations of the findings are critically discussed in section 4.

2.1 Bioanalytical Methods for the Detection and Quantification of Nucleic Acids

Nucleic acids are biopolymers composed of nucleotides and allow organisms to transfer genetic information. The nucleotides (e. g., G for guanine, G for cytosine) are arranged in a specific order (sequence). Strands of complementary nucleotide pairs (e. g., G:::C) form stable hybrids. These can be differentiated based on their sequences by the analytical methods presented below. There are two types of nucleic acids:

- deoxyribonucleic acid (DNA) and
- ribonucleic acid (RNA).

The latter has further subtypes such as micro-RNA (Berg, Tymoczko, and Stryer 2002). Jeffreys, Wilson, and Thein (1985) set a milestone for analytical forensics by showing how to create DNA profiles (*DNA fingerprints*) from different sequences in forensic trace analysis, human identification and parenthood testing. This was made possible since short detection probes made of DNA were used to detect sets of hypervariable mini-satellites, which are specific for individuals and even twins.

Beyond any doubt is the detection and quantification of nucleic acids a cornerstone in trace analysis. There are plenty of methods to detect and quantify nucleic acids. Nowadays, nucleic acids can be used

- to differentiate between human and non-human,
- to distinguish individuals or
- to differentiate body fluids

as exemplary shown by Vennemann and Koppelkamm (2010); Halpern and Ballantyne (2011); Geng, Novak, and Mathies (2014); Silva et al. (2015) and Sauer, Reinke, and Courts (2016). Provided, that the nucleic acid is in solution, methods described in subsubsection 2.1.1 ff. are commonly used.

2.1.1 Photometric Determination and Fluorescence Determination of Nucleic Acids

DNA has a maximum absorbency of light at a wavelength of 254 - 260 nm. The DNA can be quantified photometrically, using Lambert-Beer's law (Beer 1852), in a quartz or plastic cuvette. This quantitative method gives the total DNA content in the sample. Herewith, differentiation of specific sequences or the determination of sequence copy numbers is impossible.

There are several fluorescent dyes that bind specifically to nucleic acids for fluorescence determination (e. g., ethidium bromide, SYTO-9, PicoGreen and SYBR® Green (Karsai et al. 2002)). For quantification, the DNA is mixed with the fluorescence dye that binds to the DNA backbone. DNA without the dye binding has no fluorescence. The fluoresce signal intensity is measured upon binding of the dye to the nucleic acid, where the intensity is proportional to the quantity of DNA in the sample. This method gives information about the total DNA content in the sample but allows no differentiation of specific sequences, or the determination of sequence copy numbers.

Fluorescent labeled detection probes have been developed that specifically bind to a target sequence. Examples include *hybridization probes*, *hydrolysis probes* and *molecular beacons* (Rödiger et al. 2014; Halpern and Ballantyne 2011). Once calibrated with standards of known quantities, this enables a precise (sequence-specific) quantification (Singer et al. 1997).

Both principles are used in melting curve analysis, which involves the monitoring of the temperature-dependent dissociation of double stranded nucleic acids (subsubsection 2.1.2).

2.1.2 Melting Curve Analysis of Nucleic Acids

Melting curve analysis is an analytical method to determine the melting temperature of a double-stranded nucleic acid. The melting temperature (T_m) is the temperature at which 50% of nucleic acid is denatured. The temperature dependent denaturation of the nucleic acids is continuously monitored during the melting curve analysis, for which either photometric determination or fluorescence determination are used. The energy required to break hybrids of double-stranded nucleic acids depends on the length of the sequence, the

GC content and the reaction environment (e. g., salt concentration, pH) (Rödiger, Böhm, and Schimke 2013; Rödiger, Burdukiewicz, et al. 2015).

2.1.3 Solid-phase Determination of Nucleic Acids

Solid-phase technologies such as **microbead assays** and **microarrays** have been developed to capture specific target sequences from a solution. In this case, target specific nucleic acid capture probes (e. g., DNA, RNA) are immobilized on a solid phase such as a glass slide or microbeads. The target sequences need to be labeled by a biochemical reaction (e. g., by a fluorescence dye) and are subsequently hybridized to the corresponding capture probe. This binding event can be quantified with specific readers or fluorescence microscopy (Rödiger et al. 2014).

2.1.4 Methods based on Nucleic Acid Amplification

The most common used method for nucleic acid detection and quantification is the **polymerase chain** reaction (PCR). PCR is a biochemical reaction that is used to enzymatically synthesize multiple identical copies of a DNA sequence by a DNA polymerase (e. g., *Taq* polymerase). This is possible by annealing two DNA primer molecules, single stranded (18 - 35 nucleotide long) DNA sequences, which are complementary to the target stand. The primers bind under temperature controlled conditions specific to the target sequence and function as starter molecule for the DNA synthesis by the DNA polymerase.

A DNA polymerase is a transferase (e. g., Taq polymerase) that covalently attaches (phosphodiester bonds) nucleotides (e. g., guanine and cytosine) at the 5'-end starting from the primer. This results in a complementary strand of the template DNA. This process is repeated multiple times to synthesize the DNA copies. The product formation can be analyzed, for example, by agarosegel electrophoresis, or capillary electrophoresis. In both cases, the DNA is separated by size. To visualize specific detection probes, DNA labeling dyes or combination of detection probes and dyes, can be used. (Halpern and Ballantyne 2011; Westermeier 2004)

The PCR is the recommended procedure for short tandem repeat (STR) analysis. Short tandem repeats play a pivotal role in forensics, since they are a gold standard for DNA profiling in forensic casework (Oorschot, Ballantyne, and Mitchell 2010; Geng, Novak, and Mathies 2014).

With the **quantitative PCR** (qPCR) technology, the number of start copies of a target nucleic acid sequence in a genomic or complementary DNA sample can be quantified. qPCR uses the same biochemical principles as PCR. This is possible because the amplification reaction is continuously monitored in each thermo-cycle (Higuchi et al. 1993; Rutledge and Côté 2003). The resulting kinetics of the amplification process are described in subsubsection 3.2.4. To quantify the DNA copies, the amplification reaction of the unknown DNA sample is compared with a sample whose concentration is known (e. g. internal standard, dilution series).

Applications of qPCR include the quantification of human nuclear DNA and mitochondrial DNA from forensic evidence (Alonso and García 2007), analysis of gene expression (Pabinger et al. 2014; George et al. 2016) or the relationship between (poly)pharmacy and clinical consequences (Maher, Hanlon, and Hajjar 2014; Caudle et al. 2019; Rao et al. 2018; Mestdagh et al. 2016; Giuliano et al. 2017; Meyer et al. 2017; Lee et al. 2017). qPCR was used in this work to generate amplification curve data sets. In the later chapters, this work will only focus on qPCR technology.

In the Life Science sector, qPCR has become indispensable. For one, qPCR is used to investigate biological relationships by gene expression analysis. The qPCR is also used for the preparation and post-processing of NGS experiment (Nassirpour et al. 2014; Cristino et al. 2011). The RNA-Seq assumes that all transcripts can be sequenced unbiased. In reality, certain DNA sequences (e. g., GC-rich regions) are more poorly processed and so their quantity is often underestimated. Normalization of RNA-Seq data is based on the assumption that each sample contains the equal total amount of expressed mRNA, which does not necessarily have to be the case. NGS data are derived by mapping the sequenced DNA fragments to a genome reference. Here, mutations and single nucleotide polymorphisms (SNPs) are sometimes ignored. As a consequence, results vary depending on which mismatches are allowed. The qPCR is used for qualitative and quantitative characterization of the material to be sequenced. Moreover, qPCR is a highly sensitive method for checking

the "library DNA" of NGS reactions. Analysis of sequencing data is not the end of an NGS experiment, and sequence variations or expression changes need to be verified reliably. qPCR is used to confirm found differences in the follow-up. There are undisputed qualities of qPCR in terms of sensitivity and specificity. The above described problems play only a minor role in qPCR. For this reason, qPCR is usually used for validation of NGS experiments. In particular, the technical progress of commercially available qPCR technologies has led to the development of easy-to-use devices that enable users to compile large volumes of data on their specific research questions. This involves the requirement to evaluate data objectively and reproducibly (Rödiger, Burdukiewicz, et al. 2015).

The digital PCR (dPCR) is a variation of the PCR which predates qPCR. Due to the technical complexity of dPCRs, it was predominantly used in academia. Technical advances make it widely available to the mass market since 2008. The biochemical foundation of dPCR is similar to qPCR but the reaction mix is partitioned into thousands of small partitions (nanoliter volume). These partitions separate the amplification reactions ("clonal amplification"), which in combination with Poisson statistics, enables an absolute quantification of DNA (Pabinger et al. 2014; Burdukiewicz et al. 2016).

An alternative to PCR is **isothermal amplification**. Isothermal amplification is a continuous reaction at a constant temperature (isothermal) and involves enzymes that differ from the conventional PCRs. An example is the loop-mediated amplification (LAMP) (Rödiger et al. 2011) that is gaining interest in forensics for point-of-care testing and microfluidic devices. Recent advances in the development of technologies for bioanalytical microfluidic devices, medium-throughput and high-throughput technologies were achieved. A device for forensic applications was shown for the analysis of STRs, but also for quantitative assays (Horsman et al. 2007).

Modern **DNA sequencing**, often referred to as next generation sequencing (NGS), is a technology that provides the exact sequence of large and complex mixtures of DNA sequences. NGS is also a quantitative method since the number of target sequences can be counted. As of completion of this study, this method is still laborious and not ubiquitously available in forensic laboratories (Yang, Xie, and Yan 2014).

All the above methods are used for DNA typing, to distinguish individuals according to their DNA profile, to determine the origin of body fluids or to analyse the temporal course of expression of inflammatory mediators in wound tissue (Curran 2009; Alonso and García 2007; Swango et al. 2007; Sauer, Reinke, and Courts 2016; Martins et al. 2015; Wurmb-Schwark et al. 2002). One further application of PCR is the analysis of short tandem repeat (STR) loci (Halpern and Ballantyne 2011).

3 Concepts and Methods

The following sections describe the biostatistical approaches and working principles

- for data management,
- literate programming,
- open data and
- reproducible research

as recommended by Wilson et al. (2017). The proposed work-flow for the study is to:

- 1. define the biostatistical problem exemplified by the curvature of amplification curve data,
- 2. collect and prepare the amplification curve data,
- 3. develop tools for the manual amplification curve classification,
- 4. develop algorithms that can calculate predictors of amplification curve data,
- 5. apply machine learning algorithms from predictors of the curvature,
- 6. improve the results and finally
- 7. present the results as part of an open source software package for reproducible research.

For the generation of negative amplification curves, a set of experiments was conducted to amplify sequences of the Human papillomaviruses (HPV). The availability of negative amplification curves is limited, since in most cases only a few negative controls are included in qPCR experiments. The findings and data set used in

this study were made available as open data bundled in the open source software package called PCRedux (https://CRAN.R-project.org/package=PCRedux).

All statistical analyses were performed using RKWard (Rödiger et al. 2012) v. $\geq 0.7.0z+0.7.1+devel1$. RKWard is an integrated development environment and graphical user interface for the R (v. 3.6.1) statistical computing language. It has been shown that R is a powerful environment for reproducible and transparent data analysis in a highly customizable cross-platform environment (Rödiger, Burdukiewicz, et al. 2015).

3.1 Development, Implementation and Installation

PCRedux is an open source software package (MIT license⁵) for the statistical computing language R. Data with sigmoid curves are common in bioanalytical methods, such as in the widely used real-time PCR (qPCR), applied in human diagnostics, life sciences and forensics (Martins et al. 2015; Sauer, Reinke, and Courts 2016). qPCR amplification curves are an example for sigmoid shaped curves, for which PCRedux contains functions to calculate predictors and classify data sets for machine learning applications.

All technical and experimental aspects should be performed under principles that follow good practices of reproducible research. Here, numerous authors addressed the matter for experimental design and data report. Examples are the *Minimum Information for Publication of Quantitative PCR Experiments* guidelines (MIQE) and the *Real-time PCR Data Markup Language* (RDML). MIQE is a recommended standard of the minimum information for publication of quantitative real-time PCR experiments guidelines and RDML is a data exchange format (Bustin 2017; Rödiger, Burdukiewicz, et al. 2015; Rödiger et al. 2017).

The development of scientific software is a complex process, in particular, when a developer team works in different time zones with no face-to-face meetings. End users need releases with stable software that delivers reproducible results and developers need well documented software to modify the software to their needs.

Under the umbrella Agile Software Development and Extreme Programming, several principles were proposed to deliver high quality software that meet the needs of end users and developers. This includes version control, collaborative editing, unit testing and continuous integration (Lanubile et al. 2010; Myers et al. 2004; Rödiger, Burdukiewicz, et al. 2015). The core contributors are listed in the DESCRIPTION file of the PCRedux package. The following paragraphs describe methods applied for the PCRedux package.

3.1.1 Version Control and Continuous Integration

The development of the PCRedux package started 2017 with the submission of a functional, yet immature source code, to GitHub (GitHub, Inc.). GitHub is a web-based version control repository hosting service. Both distributed version control and source code management are based on Git (Lanubile et al. 2010). Additional functionality of GitHub includes the administration of access management, bug tracking, moderation of predictor requests, task management, some metrics for the software development, and wikis. The source code of PCRedux is available at:

https://github.com/devSJR/PCRedux/

Continuous integration development team members (incl. coders, artists, translators) can commit and integrate their contributions several times a day. An automated build and test system verifies each integration and gives the development team members a timely feedback about the effect of their commit. In contrast to deferred integration, this leads to a reduced number of integration problems and less workload as most errors are solved shortly after they were integrated (Myers et al. 2004).

TravisCI was chosen as continues integration service for PCRedux. The TravisCI server communicates with the GitHub version control system and manages the PCRedux package building process. Continuous interaction is available for the R releases oldrel, release and devel. The history of the build tests are available at:

https://travis-ci.org/devSJR/PCRedux

⁵https://opensource.org/licenses/MIT

3.1.2 Naming Convention and Literate Programming

PCRedux is an R (\geq v. 3.3.3) package, written as an S3 object system. S3 has characteristics of object orientated programming, but eases the development due to the use of the naming conventions (Brito 2008). In most places functions and parameter names are written as underscore separated (underscore_sep), which is a widely used style in R packages (Bååth 2012). This convention had to be violated in coding sections where functionality from other packages was used.

By convention, it was specified that functions from the PCRedux package shall be reported in the form functionname() (e. g., qPCR2fdata()) and functions from other packages in the form functionname() [packagename] (e. g., pcrfit() [qpcR]).

Literate programming, as proposed by Knuth (1984), is a concept where the logic of the source code and documentation is integrated in a single file. Markup conventions (e. g., '#') define in literate programming on how to typeset the documentation. This produces outputs in a typesetting language such as the lightweight markup language **Markdown**, or the document preparation system LATEX.

The roxygen2, rmarkdown and knitr packages were used to write the documentation in-line with code for the PCRedux package.

3.1.3 Installation of the PCRedux Package

The development version of PCRedux can be installed using the devtools package.

```
# Install devtools, if not already installed.
install.packages("devtools")

# Install PCRedux
devtools::install_github("devSJR/PCRedux")
```

PCRedux is available as stable version from the Comprehensive R Archive Network (CRAN) at https://CRAN. R-project.org/package=PCRedux. Package published at CRAN undergo intensive checking procedures. In addition, CRAN tests whether the package can be built for common operating systems and whether all version dependencies are solved. To install PCRedux first install R (\geq v. 3.3.3). Then start R and type in the prompt:

```
# Select your local mirror
install.packages("PCRedux")
```

The PCRedux package should just install. If this failed make sure that write access is permitted to the destination directory and that all package dependencies are met.

```
# The following command points to the help for download and install of packages
# from CRAN-like repositories or from local files.
?install.packages()
```

If this fails try to follow the instructions given by De Vries and Meys (2012).

R CMD check

Results from CRAN check can be found at

http://cran.us.r-project.org/web/checks/check results PCRedux.html.

3.1.4 Unit Testing of the PCRedux Package

Module testing, better known as unit testing, is an approach to simplify the refactoring of source code during software development. Unit testing is not a guarantee for error-free software. The goal is to minimize errors and regressions. It is also intended to ensure that the numerical results from the calculations are reproducible

and of high quality. An unintended behavior of the software should be detected at the latest during the package building process (Myers et al. 2004).

Checkpoints are used to check whether the software performs calculations and data transformations correctly for all builds. For this, numerous (logical) queries have to be defined by the developer in advance, referred to as expectations. It should be ensured that as many errors as possible are covered. A logical query can be, for example, whether the calculation has a numeric or Boolean value as output. If the data type is incorrect during output, this is a sufficient termination criterion. Or it can be checked whether the length of the result vector is correct after the calculation. There are different approaches for unit tests in R, including the packages RUnit, covr, svUnit and testthat (Wickham (2011)).

The package testthat was used in PCRedux because it could well be implemented and its maintenance is relatively simple. The logic is that an *expectation* defines how the result, class or error in the corresponding unit (e. g., function) should behave. Unit tests can be found in the /test/testthat subdirectory of the PCRedux package. They are run automatically during the creation of the package. The following example for the qPCR2fdata() function (for details see subsubsection 3.6.2) uses the test_that() [testthat] function with the *expectations* that:

- an object of class *fdata* is created (see Febrero-Bande and Oviedo de la Fuente (2012) for details of the class *fdata*),
- the parameter rangeval has a length of two,
- the second value of parameter rangeval is 49 (last cycle number) and *whether the object structure of the qPCR2fdata() function does not change if the parameter preprocess=TRUE is set.

```
# Expectations used for the unit testing of the qPCR2fdata() function.
library(PCRedux)

context("qPCR2fdata")

test_that("qPCR2fdata gives the correct dimensions and properties", {
    library(qpcR)
    res_fdata <- qPCR2fdata(testdat)
    res_fdata_preprocess <- qPCR2fdata(testdat, preprocess = TRUE)

expect_that(res_fdata, is_a("fdata"))
    expect_that(length(res_fdata$rangeval) == 2 &&
    res_fdata$rangeval[2] == 49, is_true())

expect_that(length(res_fdata_preprocess, is_a("fdata")))
    expect_that(length(res_fdata_preprocess$rangeval) == 2 &&
    res_fdata_preprocess$rangeval[2] == 49, is_true())
}</pre>
```

Further unit tests were implemented for all functions of the PCRedux package. The coverage of the PCRedux package can be calculated by the package_coverage() [covr] function (Hester 2018) or visually analyzed via web-interface at:

https://codecov.io/gh/devSJR/PCRedux/list/master/.

3.2 Analysis of Sigmoid-Shaped Curves for Data Mining and Machine Learning Applications

The following sections describe PCRedux regarding the analysis, numerical description and predictor calculation from a sigmoid curve. A predictor herein refers to a quantifiable *informative* property of a sigmoid curve. The predictors (subsection 3.3), sometimes referred to as descriptors, can be used for applications such as data mining, machine learning and automatic classification (e. g., negative or positive amplification).

Machine learning is a scientific discipline that deals with the use of simple to sophisticated algorithms to learn from large volumes of data. A number of approaches to machine learning exist. Supervised learning algorithms are trained with data which contain correct answers (Zielesny 2011; Walsh, Pollastri, and Tosatto 2015; Fernandez-Delgado et al. 2014). This allows to create models that assign the data to the answers and use these for further processing and predictions (Tolson 2001). Unsupervised algorithms learn from data without answers. They use large, diverse data sets for self-improvement. Neural networks or artificial neural networks are a type of machine learning that roughly resembles the function of human neurons. They are computer programs that use several levels of nodes (neurons), work in parallel for learning, recognize patterns and make decisions in a human-like manner (Günther and Fritsch 2010). Deep Learning uses a deep neural network with many neuronal layers and an extensive volume of data (Shin et al. 2016). They solve complex, non-linear problems and are responsible for groundbreaking innovations through artificial intelligence, such as the processing of a natural language or images (Tolson 2001). Applications in the life sciences have already been described for each of these methods. Up to now there appears to be no study that uses machine learning for the classification of amplification curves in a scientific setting.

The determination of quantification points such as the Cq value is a typical task during the analysis of qPCR experiments. This is briefly described in dedicated sections (subsubsection 3.2.3ff.).

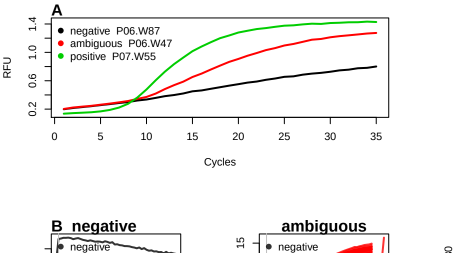
Characteristics of amplification curves that can be used for the statistical and analytical description are discussed (subsubsection 3.3.1) more in detail. The examples described focus on the concepts for binary (dichotomous) classification (Kruppa et al. 2014) as negative or positive. The mere binary classification into classes "positive" or "negative" is not necessarily the aim of the PCRedux package. Instead, it is aimed to provide a tool set for automatic multicategory (polychotomus) classification of amplification curves by any class conceivable. Such classification could be used for the quality of an amplification curve as negative, ambiguous and positive (Figure 1A & B). A definition for binary (dichotomous) classification and multicategory (polychotomus) classification is presented in Kruppa et al. (2014).

3.2.1 Relation of Machine Learning to the Classification of Amplification Curves

Data mining and machine learning can be used for descriptive and predictive tasks during the analysis of complex data sets. Data mining uses specific methods from statistical inference, software engineering and domain knowledge to get a better understanding of the data, and to extract *hidden knowledge* from the preprocessed data (Kruppa et al. 2014; Herrera et al. 2016). All this implies that a human being interacts with the data at the different stages of the whole process as part of the workflow in data mining. Elements of the data mining process are the preprocessing of the data, the description of the data, the exploration of the data and the search for connections and causes.

The availability of classified amplification curve data sets and technologies for the classification of amplification curves is of high importance to train and validate models. This is dealt with in subsection 3.5 and subsection 3.4, respectively.

For machine learning, the type of learning task is the first thing that needs to be defined. The learning task can be a classification, clustering or regression problem. Next, suitable algorithms can be selected depending on the task. In the case of classification problems, it is attempted to predict a discrete valued output. The labels (y) are usually categorical and represent a finite number of classes (e. g. "negative", "positive" \rightarrow binary classification). With regression tasks, it is attempted to predict a continuously valued output. Clustering is primarily about forming groups (clusters) based on their similarities. Examples are presented in the following and following chapters.



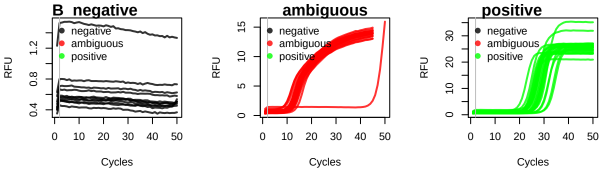


Figure 1: Examples of negative, ambiguous and positive amplification curves. A) A negative (black), ambiguous (red) and positive (green) amplification curve were selected from the 'htPCR' data set. The negative amplification curve is non-sigmoid and has a positive trend. The ambiguous amplification curve is similar to a sigmoidic amplification curve, but shows a positive slope in ground phase (cycle $1 \rightarrow 5$). The positive amplification curve (green) is sigmoid. It starts with a flat baseline (cycle $5 \rightarrow 25$). This is followed by the exponential phase (cycle $5 \rightarrow 25$) and ends in a flat plateau phase (cycle $26 \rightarrow 35$). B) Amplification curves of the 'vermeulen1' data set were divided into groups with negative, ambiguous and positive classification. Negative amplification curves have a low signal level. Interesting is the spontaneous increase (probably due to a sensor calibration) in cycles 1 to 2 followed by a linear signal decrease. In principle, the ambiguous amplification curves have a sigmoid curve shape. However, the plateau phase is fairly broad. One of the ambiguous amplification curves begins to rise sharply at Cycle 45. The positive amplification curves have a characteristic sigmoid curve shape.

In contrast, machine learning uses instructions and data in software modules to create models that can be used to make predictions on novel data. In machine learning, the human being is much less necessary in the entire process. Processes (algorithms) are used to create models with tunable parameters. These models automatically adapt their performance to the information (predictors) from the data. Well-known examples of machine learning technologies are Decision Trees (DT), Boosting, Random Forests (RF), Support Vector Machines (SVM), generalized linear models (GLM), logistic regression (LR) and deep neural networks (DNN) (Lee 2010). The three following concepts of machine learning are frequently described in the literature:

Supervised learning: These algorithms (e. g., SVM, DT, RF) learn from a training data set of labeled and annotated data (e. g., "positive" and "negative"). Classified training data can be created by one or more individuals. It is used for building a generalized model of all data. These algorithms use error or reward signals to evaluate the quality of a found solution (Bischl et al. 2010; Greene et al. 2014; Igual and Seguí 2017). Binomial logistic regression⁶ (also referred to as logit regression or logit model) is used to gain knowledge about a binary relationship, by fitting a regression model y = f(x). y is a categorical variable with two states (negative $\rightarrow 0$, positive $\rightarrow 1$). Typically, this model is used for predicting y with a mixture of n continuous and categorical predictors (features) $x_{i1}, \ldots, x_{k1}, (i = 1, \ldots, n)$.

The logit model is a robust and versatile classification method to explain a dependent binary variable. Their codomain of real numbers is limited to [0,1], hence probabilities can be employed. The logistical distribution function $F(\eta)$, also known as the response function, is strictly monotone increasing and limited to this range.

 η_i establishes the link between the probability of the occurrence and the independent variables. For this reason, η_i is referred to as a link function. The distribution function of the normal distribution is an alternative to the logistical distribution function. By using the normal distribution, the Probit model is obtained. However, since this is more difficult to interpret, it is less widely used in practice. Since probabilities are used, it is possible to make a prediction about the probability of occurrence of an event.

For example, when analyzing amplification curves, diagnosis can be made whether a reaction was unsuccessful (0) or successful (1). For the prediction, independent metric variables (predictors) are used. The metric variables have interpretable distances with a defined order. Their codomain is $[-\infty,\infty]$. The logistic distribution function on the independent variables determines the probability for $Y_i = 0$ or $Y_i = 1$. A logistic regression model can be formulated as follows:

$$F(\eta) = \frac{1}{1 + exp(-\eta)}$$

The logistic regression analysis is based on the maximum-likelihood estimation (MLE). In contrast to linear regression, the probability for Y = 1 is not modeled from explanatory variables. Rather, the logarithmic chance (logit) is used for the occurrence of Y = 1. The term *chance* refers to the ratio of the probability of occurrence of an event (e. g., amplification curve is positive) and the counter-probability (e. g., amplification curve is negative) of an event.

Unsupervised learning: Algorithms, such as k-means clustering, kernel density estimation, or Principal Component Analysis learn from training data sets of unlabeled or non-annotated data to find hidden structures according to geometric or statistical criteria (Bischl et al. 2010; Greene et al. 2014; Igual and Seguí 2017).

Reinforcement Learning: The algorithms learn by reinforcement from *criticism*. The criticisms inform the algorithm about the quality of the solution found but nothing about how to improve. These algorithms search by iterations the improved solution in the entire solution space (Bischl et al. 2010; Igual and Seguí 2017).

 $^{^6}$ Logistical regression can also be used to predict a dependent variable that can assume more than two states. In this case, it is called a multinomial logistic regression. An example would be the classification y of amplification curves as *slightly noisy*, *medium noisy* or *heavily noisy*.

3.2.2 Why is there a need for the PCRedux software?

The binary classification of an amplification curve is feasible using bioanalytical methods such as melting curve analysis (Rödiger, Böhm, and Schimke 2013) or electrophoretic separation (Westermeier 2004). However, this is not always possible or desirable.

- Melting curve analysis is used in some qPCRs as a post-processing step to identify samples which contain the specific target sequence (positive) based on a specific melting temperature. However, some detection probe systems like hydrolysis probes do not permit such classification. Moreover, nucleic acids with similar biochemical properties but different sequences may have the same melting temperature.
- An electrophoretic separation (classification of target DNA sequences by size and quantity) often requires too much effort for experiments with high sample throughput.
- There are mathematical qPCR analysis algorithms such as linreg (Ruijter et al. 2009) that require information on whether an amplification curve is negative or positive for subsequent calculation.
- Raw data of amplification curves can be fitted with sigmoid functions. Sigmoid functions are non-linear, real-valued, have an S-shaped curvature (Figure 3) and can be differentiated (e. g., first derivative maximum, with one local minimum and one local maximum). With the obtained model, predictions can be made. For example, the position of the second derivative maximum can be calculated from this (subsubsection 3.2.4). In the context of amplification curves, the second derivative maximum is commonly used to describe the relationship between the cycle number and the PCR product formation (subsubsection 3.2.4). All software assume that the amplification resembles a sigmoid curve shape (ideal positive amplification reaction), or a flat low line (ideal negative amplification reaction). For example, Ritz and Spiess (2008) published the qpcR R package that contains functions to fit several multiparameter models. This includes the five-parameter Richardson function (Richards 1959) (Equation 3). The qpcR package (Ritz and Spiess 2008) contains an amplification curve test via the modlist() function. The parameter check="uni2" offers an analytical approach, as part of a method for the kinetic outlier detection. It tries to checks for a sigmoid structure of the amplification curve. Then, modlist() tests for the location of the first derivative maximum and the second derivative maximum. However, multi-parameter functions fit "successful" in most cases including noise and give false positive results. This will be shown in later sections. This is exemplary shown in later sections in combination with the amptester() [chipPCR] function (Rödiger, Burdukiewicz, and Schierack 2015), which uses fixed thresholds and frequentist inference to identify amplification curves that exceed the threshold $(\mapsto$ classified as positive). However, the analysis can also lead to false-positive classifications as exemplified in the example below and in Figure 2. Therefore, additional classification concepts would be beneficial.

```
# Apply the amptester() function from the chipPCR package to the amplification
# curve data and write the results to the main of the plots.
for (i in 2:3) {
    res.ampt <- suppressMessages(amptester(amp_data[, i]))</pre>
    # Make a logical connection by two tests (shap.noisy, lrt.test and
    # tht.dec) of amptester to decide if an amplification reaction is
    # positive or negative.
    decision <- ifelse(!res.ampt@decisions[1] &&</pre>
   res.ampt@decisions[2] &&
   res.ampt@decisions[4],
    "positive", "negative"
    # The amplification curves were fitted (17 model) with pcrfit() function.
    # The Cq was determined with the efficiency() function.
   fit <- pcrfit(data = amp_data, cyc = 1, fluo = i, model = 17)</pre>
   res <- efficiency(fit, plot = FALSE)</pre>
   lines(predict(fit), pch = 19, lty = 1, xlab = "Cycles", ylab = "RFU",
         main = "", col = i - 1)
    abline(h = res[["fluo"]], col = "grey")
    points(res[["cpD2"]], res[["fluo"]], pch = 19)
   legend(1, hight[i-1], paste0(colnames(amp data)[i],
                                 " curve -> Decision: ",
                                 bty = "n", cex = 1, col = "red"
}
```

- The analysis and classification of sigmoid data (e. g., quantitative PCR) is a manageable task if the data volume is low, or dedicated analysis software is available. An example for a low number of amplification curves is shown in Figure 4A. All 65 curves exhibit a sigmoid curve shape. It is trivial to classify them as positive by hand. In contrast, the vast number of amplification curves in Figure 4B is barely manageable with a reasonable effort by simple visual inspection. These data originate from a high-throughput experiment that encompasses in total 8858 amplification curves of which only 200 are shown. A manual analysis of the data is time-consuming and prone to errors. Even for an experienced user, it is difficult to classify the amplification curves unambiguously and reproducible as will be later shown in subsection 3.5.
- qPCRs are performed in thermo-cyclers, which are equipped with a real-time monitoring technology. There are numerous commercial manufactures producing thermo-cyclers (Table 5). An example for a thermo-cycler that originated in a scientific project is the VideoScan technology (Rödiger, Schierack, et al. 2013). Most of the thermo-cyclers have a thermal block with wells at certain positions. Reaction vessels containing the PCR mix are inserted into the wells. There are also thermo-cyclers that use capillary tubes that are heated and cooled by air (e. g., Roche Light Cycler 1.0). The thermo-cycler raises and lowers the temperature in the reaction vessels in discrete, pre-programmed steps so that PCR cycling can take place. Instruments with a real-time monitoring functionality have sensors to measure changes of the fluorescence intensity in the reaction vessel. All thermo-cycler systems use software to processes the amplification curves. Plots of the fluorescence observations versus cycle number obtained from two different qPCR systems are shown in Figure 4A and B. The thermo-cyclers produce different amplification curve shapes even with the same sample material and PCR mastermix because of their technical design, sensors, and software. These factors need to be taken into account during the development of analysis algorithms.

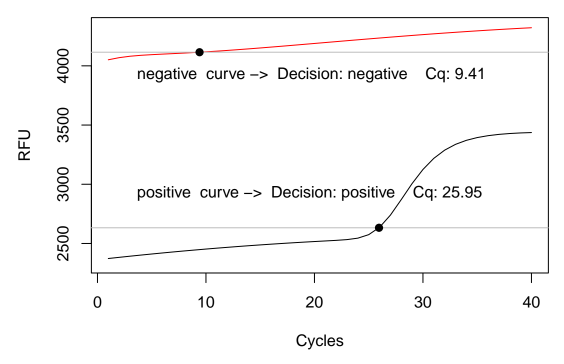


Figure 2: Incorrect model adjustment for amplification curves. A positive (black), and a negative (red) amplification curve were randomly selected from the 'RAS002' data set. The positive amplification curve has a baseline signal of about 2500 RFU and has a definite sigmoidal shape. The negative amplification curve has a baseline signal of approx. 4200 RFU, but only moderately positive slope (no sigmoidal shape). A logistic function with seven parameters ('17') has been fitted to both amplification curves. A Cq value of 25.95 was determined for the positive amplification curve. The negative amplification curve had a Cq value of 9.41. However, it can be seen that the latter model fitting is not appropriate for calculating a trustworthy Cq value. An automatic calculation without user control would give a false-positive result.

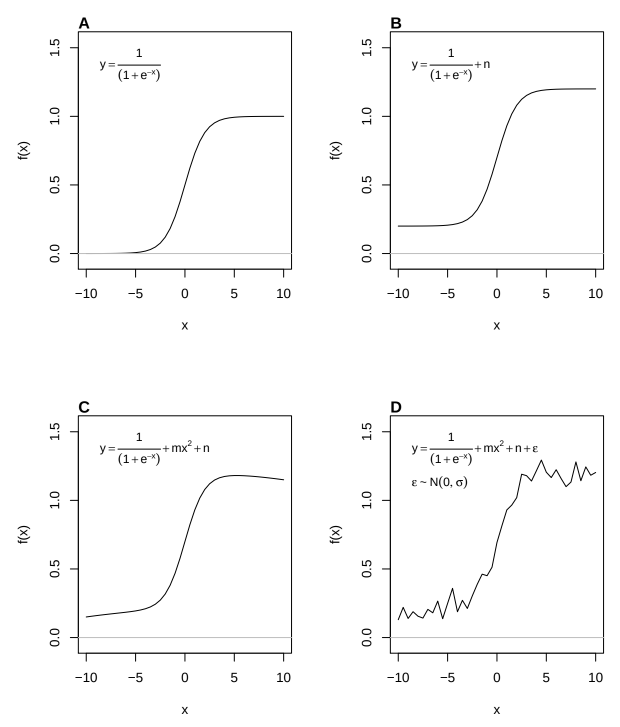
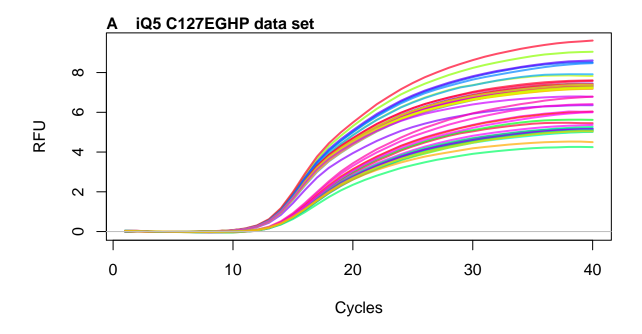


Figure 3: A) Model function of a one-parameter sigmoid function. B) Model function of a sigmoid function with an intercept n=0.2 RFU (shift in base-line). C) Model function of a sigmoid function with an intercept ($n\sim0.2$ RFU) and a square portion $m*x^2, m=-0.0005, n=0.2$ RFU (hook-effect-like). D) Model function of a sigmoid function with an intercept (n) and a square portion of $m*x^2$ and additional noise ϵ (normal distributed, $\mu=0.01, \sigma=0.05$).



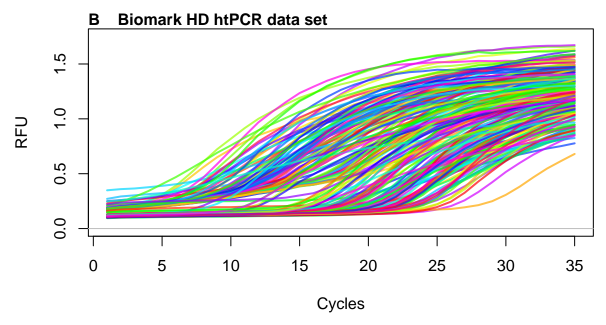


Figure 4: Amplification curve data from an iQ5 (Bio-Rad) thermo-cycler and a high throughput experiment in the Biomark HD (Fluidigm). A) The 'C127EGHP' data set with 64 amplification curves was produced in a conventional thermo-cycler with a 8 x 12 PCR grid. B) The 'htPCR' data set, which contains 8858 amplification curves, was produced in a 95 x 96 PCR grid. Only 200 amplification curves are shown. In contrast to 'A)' have all amplification curves in 'B)' an off-set (intercept) between 0.09 and 0.40 RFU.

3.2.3 Software for the Analysis of Amplification Curve Data

There are several open source and closed source software tools for the analysis of qPCR data (Pabinger et al. 2014). The software packages deal for example with

- missing values and non-detects (McCall et al. 2014),
- noise and artifact removal (Rödiger, Burdukiewicz, and Schierack 2015; Rödiger, Burdukiewicz, et al. 2015; Spiess et al. 2015, 2016),
- inter-run calibration (Ruijter et al. 2015),
- normalization (Rödiger, Burdukiewicz, and Schierack 2015; Ruijter et al. 2013; Feuer et al. 2015; Matz, Wright, and Scott 2013),
- quantification cycle estimation (Ritz and Spiess 2008; Ruijter et al. 2013),
- amplification efficiency estimation (Ritz and Spiess 2008; Ruijter et al. 2013),
- data exchange (Lefever et al. 2009; Perkins et al. 2012; Rödiger et al. 2017),
- relative gene expression analysis (Dvinge and Bertone 2009; Pabinger et al. 2009; Neve et al. 2014) and
- data analysis pipelines (Pabinger et al. 2009; Ronde et al. 2017; Mallona, Weiss, and Egea-Cortines 2011; Mallona et al. 2017).

However, a bottleneck of qPCR data analysis is the lack of predictors and software to build classifiers for amplification curves. A classifier herein refers to a vector of predictors that can be used to distinguish the amplification curves only by their shape. A predictor, also referred to as *feature*, is an entity that characterizes an object. A few potential predictors for amplification curves are described in the literature. These include:

- the starting point (takeoff) of the amplification curve,
- the Cq value and amplification efficiency, and
- the signal level (e. g., slope and intercept of the ground phase).

These alone are presumably not enough to describe amplification curves sufficiently. The number of predictors should be large enough to describe the object accurately and small enough to not interfere with the learning process with redundant information ("overtraining"). There are no references of algorithms in the scientific literature for the calculation of additional predictors from amplification curves. This makes studies on machine learning and modeling difficult.

3.2.4 Principles of Amplification Curve Data Analysis and Predictor Calculation

The shape of a positive amplification curve is in most cases sigmoidal. Many factors such as the sample quality, qPCR chemistry, and technical problems (e. g., sensor errors) contribute to various curve shapes (Ruijter et al. 2014). The curvature of the amplification curve can be used as a quality measure. For example, fragmentation, inhibitors and sample material handling errors during the extraction can be identified. The kinetic of fluorescence emission is proportional to the quantity of the synthesized DNA. Typical amplification curves have three phases.

1. Ground phase: This phase occurs during the first cycles of the PCR, where the fluorescence emission is in most cases flat. Here, noise but no product formation is detected by the sensor system and the PCR product signal is an insignificantly small component of the total signal. This is often referred to as base-line or background signal. Apparently, there is only a phase shift or no signal at all, primarily due to the limited sensitivity of the instrument. Even in a perfect PCR reaction (double amplification per cycle), qPCR instruments cannot detect the fluorescence signal from the amplification. Fragmentation, inhibitors and sample handling errors would result in a prolonged ground phase. Nevertheless, this may indicate some typical properties of the qPCR system or probe system. In many instruments, this phase is used to determine the base-line level for the calculation of the Cycle threshold (Ct). The Ct value is considered statistically relevant as an increase outside of the noise range (threshold) when coming from the amplicon. In some qPCR systems, a flat amplification signal is expected in this phase. Slight deviations from this trend are presumably due to changes (e. g., disintegration of probes) in the fluorophores. Background correction algorithms are often used here to ensure that flat amplification curves without slope are generated. However, this can result in errors and inevitably leads to a loss of information via the waveform of the raw data (Nolan, Hands, and Bustin 2006). The slope, level and

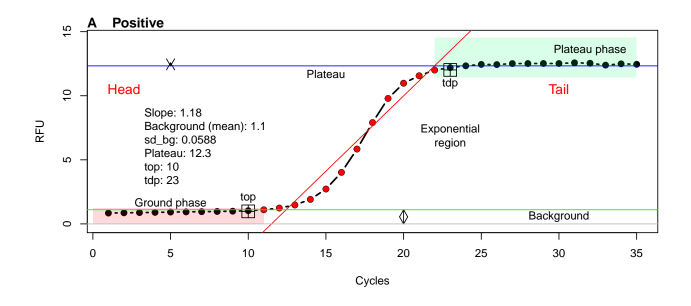
variance of this phase can serve as predictors.

- 2. Exponential phase: This phase follows the ground phase and is also called *log-linear phase*. It is characterized by a strong increase of the emitted fluorescence as the DNA amount roughly doubles in each cycle under ideal conditions and when the amount of the synthesized fluorescent labeled PCR product is high enough to be detected by the sensor system. This phase is used for the calculation of the quantification point (Cq) and curve specific amplification efficiency. The most important measurement from qPCRs is the Cq, which signifies the PCR cycle for which the fluorescence exceeds a threshold value. However, there is an ongoing debate as to what a significant and robust threshold value is. An overview and performance comparison of Cq methods is given in Ruijter et al. (2013). There are several mathematical methods to calculate the Cq.
 - The 'classical' threshold value (cycle threshold, Ct) is the intersection between a manually defined straight horizontal line with the quasi-linear phase in the exponential amplification phase (Figure 6A & B). This simple to implement method requires that amplification curves are properly baselined prior to analysis. The Ct method makes the assumption that the amplification efficiency (~ slope in the log-linear phase) is equal across all compared amplification curves (Ruijter et al. 2013). Evidently, this is not always case as exemplified in Figure 5C. The Ct method is widely used presumably due to the familiarity of users with this approach (e. g., chemical analysis procedures). However, this method is statistically unreliable (Ruijter et al. 2013; Spiess et al. 2015, 2016). Moreover, the Ct method gives no stable in predictions if different users are given the same data set to be analyzed. Therefore, this method is not used within the PCRedux package.
 - Another Cq method uses the maximum of the second derivative (SDM) (Rödiger, Burdukiewicz, et al. 2015) (Figure 6C). In all cases, the Cq value can be used to calculate the concentration of target sequence in a sample (low Cq →high target concentration). In contrast, negative or ambiguous amplification curves loosely resemble noise. This noise may appear linear or exhibit a curvature similar to a specific amplification curve (Figure 1). This however, may result in faulty interpretation of the amplification curves. Fragmentation, inhibitors and sample handling errors would decrease the slope of the amplification curve (Spiess, Feig, and Ritz 2008; Ritz and Spiess 2008). The slope and its variation can be considered as predictors. Since the Cq depends on the initial template amount and amplification efficiency, there is no immediate use of the Cq as an predictor.
 - 3. Plateau phase: This phase follows the exponential phase and is a consequence of the exhaustion of limited reagents (incl. primers, nucleotides, enzyme activity) in the reaction vessel, limiting the amplification reaction, so that the theoretical maximum amplification efficiency (doubling per cycle) no longer prevails. This turning point, and the progressive limitation of resources, finally leads to a plateau. In the plateau phase, there is in some cases a signal decrease called *hook effect* (subsubsection 3.2.2 and (Barratt and Mackay 2002; Isaac 2009; Burdukiewicz et al. 2018)). The slope (*hook effect*), level and variation can be considered as predictors.

If the amplification curve has only a slight positive slope and no perceptible/measureable exponential phase, it can be assumed that the amplification reaction did not occur (Figure 5B). Causes may include poor specificity of the PCR primers (non-specific PCR products), degraded sample material, degraded probes or detector failures. If a lot of input DNA is present in a sample, the amplification curve starts to increase in early PCR cycles (1 - 12 cycles). Some PCR devices have a software that corrects this feature without rechecking, resulting in an amplification curve with a negative trend.

The discussed phases are considered as regions of interest (ROI). As an example, the *ground phase* is in the head area, while the *plateau phase* is in the tail area. The *exponential phase* is located between these two ROIs.

The amplification curve shape, the amplification efficiency and the Cq value are important measures to judge the outcome of a qPCR reaction. In all phases of PCR, the curves should be smooth. Possible artifacts in the curves may be due to unstable light sources from the instrument or problems during sample preparation, such as the presence of bubbles in the reaction vessel, incorrectly assigned dye detectors, errors during the calibration of dyes for the instrument, errors during the preparation of the PCR master mix, sample degradation, lack of a sample in the PCR, too much sample material in the PCR mix or a low detection



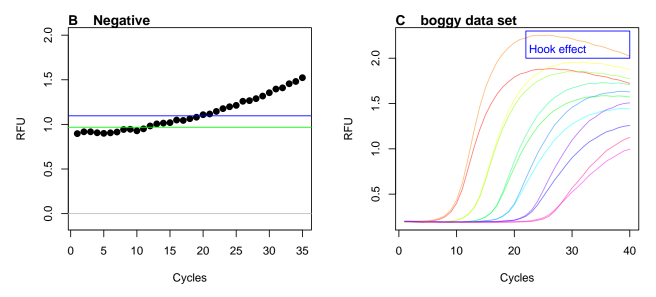


Figure 5: Phases of amplification curves as Region of Interest (ROI). For amplification curves, the fluorescence signal (RFU, relative fluorescence units) of the reporter dye is plotted against the cycle number. Positive amplification curves possess three ROIs: ground phase, exponential phase and plateau phase. These ROIs can be used to determine predictors such as the takedown point ('tdp') or the standard deviation within the ground phase ('sd_bg'). The exponential range (red dots) is used to determine the Cq values and amplification efficiency (not shown). A linear regression model (red) can be used to calculate the slope in this region. B) PCRs without amplification reaction usually show a flat (non-sigmoides) signal. C) The exponential phase of PCR reactions can vary greatly depending on the DNA starting quantity and other factors. Amplification curves that appear in later cycles often have a lower slope in the exponential phase.

probe concentration (Ruijter et al. 2009, 2014; Spiess et al. 2015). Smoothing and filtering cause alterations to the raw data that affects the Cq value and the amplification efficiency.

Most commercial qPCR systems do not display the raw data of the amplification curves on the screen. Instead, raw data are often processed by the instrument software to remove fluorophore-specific effects and noise in all ROIs. Commonly employed preprocessing step of qPCR is smoothing and filtering to remove noise, where the latter can have different causes (Spiess et al. 2015).

The ordinate often does not display the measured fluorescence, but rather the change in fluorescence per cycle ($\Delta RFU = RFU_{cycle+1} - RFU_{cycle}$). Some qPCR systems display periodicity in the amplification curve data, thereby exposing the risk of introducing artificial shifts in the Cq values (Spiess et al. 2016).

In particular the cycle threshold method (Ct method) (subsubsection 3.2.4) is affected by these factors (Spiess et al. 2015, 2016). Therefore, it is advisable to clarify in advance, which processing steps the amplification curves have been subjected to. Failure to do so may result in misinterpretations and incorrect amplification curve fitting models (Nolan, Hands, and Bustin 2006; Rödiger, Burdukiewicz, et al. 2015; Rödiger, Burdukiewicz, and Schierack 2015; Spiess et al. 2015).

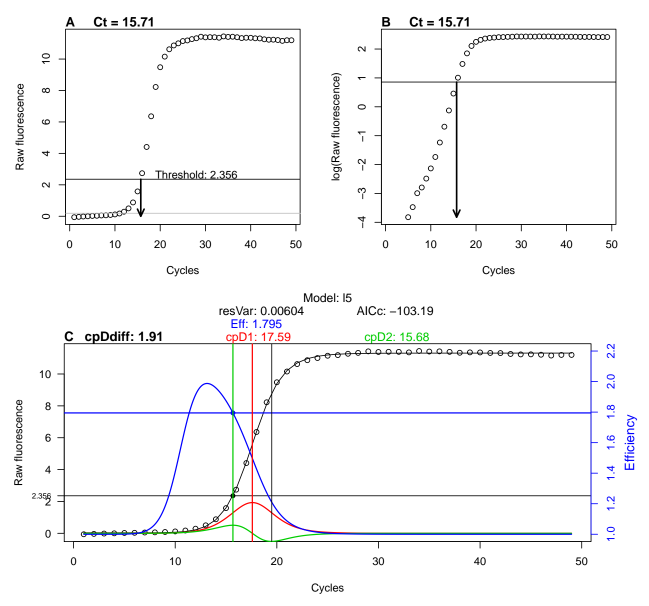


Figure 6: Frequently used methods for the analysis of quantification points. A) The amplification curve is intersected by a gray horizontal line. This is the background signal (3σ) determined from the 68-95-99.7 rule from the fluorescence emission of cycles 1 to 10. The black horizontal line is the user-defined threshold (Ct value) in the exponential phase. Based on this, the cycle at which the amplification curve differs significantly from the background is calculated. B) The amplification curve can also be analyzed by fitting a multi-parametric model (black line, five parameters). The red line is the first derivative of the amplification curve with a maximum of 17.59 cycles. The first derivative maximum ('cpD1') is used as a quantification point (Cq value) in some qPCR systems. The green line shows the second derivative of the amplification curve, with a maximum at 15.68 cycles a minimum at 19.5 cycles. The maximum of the second derivative ('cpD2') is used as the Cq value in many systems. The blue line shows the amplification efficiency estimated from the trajectory of the exponential region. The 'Eff' value of 1.795 means that the amplification efficiency is approximately 89%. 'cpDdiff' is the difference between the first and second derivative maximum (cpDdiff = cpD1 - cpD2).

3.3 Data Analysis Functions of the PCRedux Package

The PCRedux package contains functions for analyzing amplification curves. In the following, these are distinguished into helper functions (subsection 3.6) and analysis functions (subsubsection 3.3.1).

3.3.1 Amplification Curve Analysis Functions of the PCRedux package

On the basis of these observations, **concepts** for predictors (*features*) were developed and implemented in algorithms to describe amplification curves. The functions described in the following are aimed for experimental studies. It is important to note that the concepts for the predictors proposed herein emerged by a *critical reasoning* process and *domain knowledge* of the PCRedux package creator. The aim of the package is to propose a set of predictors, functions and data for an independent research.

3.3.2 pcrfit_single() and encu()- Functions to Calculate Predictors from an Amplification Curve

The following sections give a concise description of the algorithms used to calculate predictor vectors by the pcrfit_single() function. Based on considerations and experience, the algorithms of the pcrfit_single() function are restricted to ROIs (Figure 5) to calculate specific predictors.

The encu() function is a wrapper for the pcrfit_single() function. encu() can be used to process large records of amplification curve data arranged in columns. The progress of processing is displayed in the form of a progress bar and the estimated run-time. Additionally, encu() allows to specify which monitoring chemistry (e. g., DNA binding dye, sequence specific probes) and which thermo-cycler was used. Ruijter et al. (2014) demonstrated that the monitoring chemistry and the type of input DNA (single stranded, double stranded) are important when analysing qPCR data, because they have an influence on the shape of the amplification curve. For simplicity, the documentation will describe the pcrfit single() only.

The underlying hypotheses and concepts of the predictors are formulated and supported by *exemplary applications*. Different representative data sets were used to support a concept or predictors. For example, the RASO02 data set represents a typical qPCR. This means that the positive amplification curves start with a flat plateau phase and then transition into the sigmoid shape with a plateau. The negative amplification curves display no significant peculiarities. For both positive and negative amplification curves, there is a shift from the origin. The htPCR data set serves as a problem example in several places, since it contains many observations (amplification curves from high-throughput experiments). In addition, the amplification curves have a high diversity of curve shapes that cannot be uniquely and reproducibly classified even by experienced users. Other data sets are used in the documentation, but these are not discussed in detail.

To underscore the usability of the algorithms and their predictors, 3302 observations (471 negative amplification curves, 2831 positive amplification curves) from the batsch1, boggy, C126EG595, competimer, dil4reps94, guescini1, karlen1, lievens1, reps384, rutledge, testdat, vermeulen1, VIMCFX96_60, stepone_std, RAS002, RAS003, HCU32_aggR and lc96_bACTXY were analyzed with the encu() function and the results (predictors) were combined in the file data_sample.rda. Users of this function should independently verify and validate the results of the methods for their own applications.

A new data set called data_sample_subset_balanced has been compiled from the data_sample data set for some of the applications. Selection criteria included:

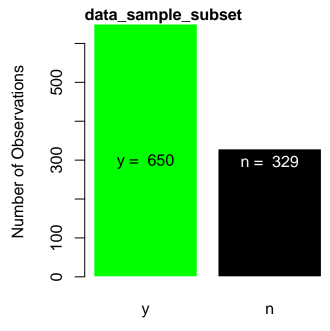
- both positive and negative amplification curves had to be included in a similar ratio,
- there should not be a dominating thermal cycler platform,
- the amplification curves should represent typical amplification curves (subjective criterion). The
 compilation of the data sets batsch1, HCU32_aggR, 1c96_bACTXY, RASO02, RASO03 and stepone_std
 met this requirement satisfactorily.

```
data_sample_subset_balanced <- data_sample[data_sample$dataset %in%
c("batsch1", "boggy", "C126EG595", "HCU32_aggR", "lc96_bACTXY",
    "RAS002", "RAS003", "stepone_std", "testdat"), ]</pre>
```

data_sample_subset_balanced y = 322 n = 329

For the comparison of predictors, the data set was enlarged. Selection criteria for the data sets were comparatively less stringent.

[1] 979 62



The goal is to demonstrate the basic functionality of the algorithms for predictor calculation. Similar concepts are presented in groups. The algorithms are divided into the following broad categories:

- algorithms that determine slopes, signal levels,
- algorithms that determine turning points and
- algorithms that determine areas.

The algorithms in

- earlyreg() (subsubsection 3.3.9),
- head2tailratio() (subsubsection 3.3.10),
- hookreg() & hookregNL() (subsubsection 3.3.11) and
- mblrr() (subsubsection 3.3.12),
- autocorrelation_test() (subsubsection 3.3.13)

were implemented as standalone functions to make them available for other applications.

The output below shows the predictors and their data type (num, numeric; int, integer; Factor, factor; logi, boolean) that were determined with the pcrfit_single() function.

```
library(PCRedux)
# Calculate predictor vector of column two from the RASOO2 data set.
str(pcrfit_single(RASOO2[, 2]))
```

```
1 obs. of 57 variables:
## 'data.frame':
## $ cpD1
                         : num 28.1
## $ cpD2
                         : num 25.9
## $ cpD2_approx
                         : num 26
## $ cpD2 ratio
                         : num 0.997
## $ eff
                         : num 1.02
## $ sliwin
                         : num 1.04
## $ cpDdiff
                         : num 2.19
                         : num 0.0343
## $ loglin_slope
## $ cpD2_range
                         : num 4.62
                         : num 25
## $ top
## $ f.top
                         : num 0.748
## $ tdp
                         : num 33
## $ f.tdp
                         : num 1.62
## $ bg.stop
                         : num 15
## $ amp.stop
                        : num 40
## $ b slope
                        : num -13.6
## $ b_model_param
                         : num -20.6
## $ c_model_param
                         : num 0.712
## $ d_model_param
                         : num 0.996
## $ e_model_param
                         : num 30.6
## $ f model param
                         : num 0.433
## $ f intercept
                          : num 3.17
## $ convInfo iteratons
                         : int 14
                          : Factor w/ 1 level "17": 1
## $ qPCRmodel
                         : Factor w/ 1 level "17": 1
## $ qPCRmodelRF
## $ minRFU
                         : num 0.682
## $ maxRFU
                         : num 1
## $ init2
                         : num 0.419
## $ fluo
                         : num 0.765
## $ slope_bg
                         : num 0.00658
## $ intercept_bg
                        : num 0.675
## $ sigma_bg
                         : num 0.00455
## $ sd bg
                         : num 0.0801
## $ head2tail ratio
                        : num 0.704
## $ mblrr_slope_pt
                         : num 0.00586
## $ mblrr_intercept_bg : num 0.693
                         : num 0.00202
## $ mblrr_slope_bg
## $ mblrr cor bg
                         : num 0.91
## $ mblrr_intercept_pt
                         : num 0.774
## $ mblrr cor pt
                         : num 0.942
## $ polyarea
                         : num 0.0409
## $ peaks_ratio
                         : num 0.00922
## $ autocorellation
                         : num 0.752
                         : int 2
## $ cp_e.agglo
## $ cp_bcp
                          : int 5
## $ amptester_shapiro
                          : logi FALSE
## $ amptester_lrt
                          : logi TRUE
## $ amptester_rgt
                          : logi TRUE
## $ amptester_tht
                          : logi TRUE
## $ amptester slt
                          : logi TRUE
## $ amptester_polygon
                          : num 4.5
```

```
## $ amptester_slope.ratio: num 0.0384
## $ hookreg_hook : num 0
## $ hookreg_hook_slope : num 0
## $ hookreg_hook_delta : num 0
## $ central_angle : num -0.999
## $ number_of_cycles : int 40
```

3.3.3 Amplification Curve Preprocessing

The pcrfit_single() function performs preprocessing steps before each calculation, including checking whether an amplification curve contains missing values. Missing values (NA) are measuring points in a data set where no measured values are available or have been removed arbitrarily. NAs may occur if no measurement has been carried out (e. g., defective detector) or lengths of the vectors differ (number of cycles) between the observations. Such missing values are automatically imputed by spline interpolation as described in Rödiger, Burdukiewicz, and Schierack (2015).

All values of an amplification curve are normalized to their 99% quantile. The normalization is used to equalize the amplitudes differences of amplification curves from thermo-cyclers (sensor technology, software processing) and detection chemistries. To compare amplification curves from different thermo-cyclers, the values should always be scaled systematically using the same method. Although there are other normalization methods (e. g., minimum-maximum normalization, see Rödiger, Burdukiewicz, and Schierack (2015), the normalization by the 99% quantile preserves the information about the level of the background phase. A normalization to the maximum is not used to avoid strong extenuation by outliers. The data in Figure 16D show that the maxRFU values after normalization are approximately 1. There is no statistical significant difference between maxRFU values of positive and negative amplification curves.

Selected algorithms of the pcrfit_single() function use the CPP() [chipPCR] function to preprocess (e. g., base-lining, smoothing, imputation of missing values) the amplification curves. Further details are given in Rödiger, Burdukiewicz, and Schierack (2015). Until package version 0.2.6-4 was the visdat_pcrfit() part of the package. visdat_pcrfit() was used for visualizing the content of data from an analysis with the pcrfit_single() function. There are other more powerful packages such as visdat by Tierney (2017), assertr by Fischetti (2019) and xray by Seibelt (2017).

During the analysis, several values are determined to describe the amplitude of an amplification curve. The resulting potential predictors are minRFU (minimum of the amplification curve, which is determined at the 1% quantile to minimize the influence of outliers), init2 (the initial template fluorescence from an exponential model) and fluo (raw fluorescence value at the second derivative maximum). The minRFU, init2 and fluo values differ significantly between negative and positive amplification curves (Figure 16C, E & F).

3.3.4 Handling of Missing Predictors

Missing values (NA) can occur if a calculation of a predictor is impossible (e. g., if a logistic function cannot be adapted to noisy raw data). The lack of a predictor is nevertheless an useful information (no predictor calculate → amplification curves deviate from sigmoid shape). The NAs were left unchanged in the PCRedux package up to version 0.2.5-1. Since version 0.2.6 the NAs are replaced by numerical values (e. g., total number of cycles) or factors (e. g., lNA for non-fitted model). Under the term "imputation", there are a number of procedures based on statistical methods (e. g., neighboring median, spline interpolation) or on user-defined rules (Williams 2009; Cook and Swayne 2007; Hothorn and Everitt 2014). Rules are mainly used in the functions of PCRedux to relieve the user from the decision as to how to deal with missing values. For example, slope parameters of a model are set to zero when it cannot be determined. The disadvantage is that rules do not necessarily concur to real world values.

3.3.5 Multi-parametric Models for Amplification Curve Fitting

Both the pcrfit_single() function and the encu() function use four multi-parametric models based on the findings of Spiess, Feig, and Ritz (2008) and Ritz and Spiess (2008). The pcrfit_single() function starts

by adjusting a seven-parameter model since this adapts easier and more frequent to a data set (Figure 7).

17:

$$f(x) = c + k1 \cdot x + k2 \cdot x^2 + \frac{d - c}{(1 + exp(b(log(x) - log(e))))^f}$$
(1)

From that model, the pcrfit_single() function estimates the variables b_slope and c_intercept, describing the slope and the y-intercept. The number of iterations required to adapt the model is also stored. That value is returned by the pcrfit_single() function as convInfo_iteratons. The higher the convInfo_iteratons value, the more iterations are necessary to converge from the start parameters (Figure 12L). A low convInfo_iteratons value is an indicator for

- a sigmoid curve shape or
- close start parameters.

High iterations numbers imply

- noisy amplification curves or
- non-sigmoid amplification curves.

The amplification curve fitting process continues with the four-parameter model (l_4 , Equation 2). This is followed by a model with five parameters (l_5 , Equation 3) and six parameters (l_6 , Equation 4).

• l4:

$$f(x) = c + \frac{d - c}{1 + exp(b(log(x) - log(e)))}$$

$$\tag{2}$$

• l5:

$$f(x) = c + \frac{d - c}{(1 + exp(b(log(x) - log(e))))^f}$$

$$\tag{3}$$

• l6:

$$f(x) = c + k \cdot x + \frac{d - c}{(1 + exp(b(log(x) - log(e))))^f}$$

$$\tag{4}$$

The optimal model is selected on the basis of the Akaike information criterion and used for all further calculations. The pcrfit_single() function returns qPCRmodel as a factor (14, 15, 16, 17). In case no model could be fitted, an lNA is returned.

The model is an indicator of the amplification curve shape. Model with many parameters deviate more from an ideal sigmoid model. For instance, a four-parameter model, unlike the six-parameter model, does not have a linear component. A negative linear slope in the plateau phase is an indicator of a *hook effect* (Burdukiewicz et al. 2018).

3.3.6 winklr() - A function to calculate the central angle based on the first and the second derivative of an amplification curve data

winklR() is a function to calculate the in the trajectory of the first negative and the second negative derivatives maxima and minima (Figure 9) of an amplification curve data from a quantitative PCR experiment. For the determination of the angle, the origin is the maximum of the first derivative. On this basis, the vectors to the approximate minimum and maximum of the second derivatives are determined. The vectors result from the relation of the maximum of the first derivative to the minimum of the second derivative and from the maximum of the first derivative to the maximum of the second derivative. In a simple trigonometric approach, the scalar product of the two vectors is formed first. Then the absolute values are calculated and

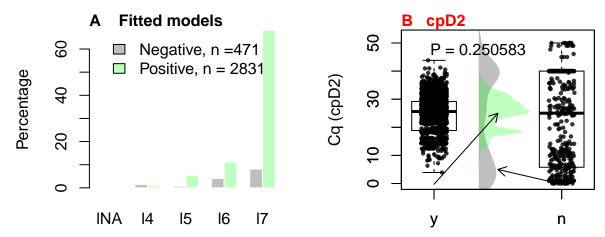


Figure 7: Frequencies of the fitted multiparametric models and Cq values. The amplification curves (n=3302) of the 'data_sample' data set were analyzed with the encu() function. The amplification curves have been stratified according to their classes (negative: grey, positive: green). A) The optimal multiparametric model was selected for each amplification curve based on the Akaike information criterion. lNA stands for 'no model' and l4 . . . l7 for a model with four to seven parameters. B) All Cq values were calculated from optimal multiparametric models. Cqs of positive amplification curves accumulate in the range between 15 and 30 PCR cycles (50%). For the negative amplification curves, the Cqs are distributed over the entire span of the cycles. Note: The Cqs of the negative amplification curves are false-positive!

multiplied by each other. Finally, the value is converted into an angle with the cosine. The assumption is that flat (negative amplification curves) have a large angle and sigmoid (positive amplification curves) have a smaller angle. Another assumption is that this angle is independent of the rotation of the amplification curve. This means that systematic off-sets, such as those caused by incorrect background correction, are of no consequence. The cycles to be analyzed is defined by the user. The output contains the angle and the coordinates of the minima and maxima.

```
# Calculate the central angles for amplification curves from the RAS002 data set.
library(PCRedux)
# Load the amplification curves from the RASOO2 data set.
DATA <- PCRedux::RAS002
# Load the RAS002_decisions data set.
dec <- RAS002_decisions
# Give tabular output of classes
table(dec)
## dec
##
     У
         n
    42 150
# Assign colors to the classes (n: black, y: green).
colors <- factor(dec, levels = c("y", "n"),</pre>
                 label = c("black", "red"))
# Test for the angles via the winklR() function on the RASO02 data set
res_winklR <- sapply(2L:ncol(DATA), function(i) {</pre>
```

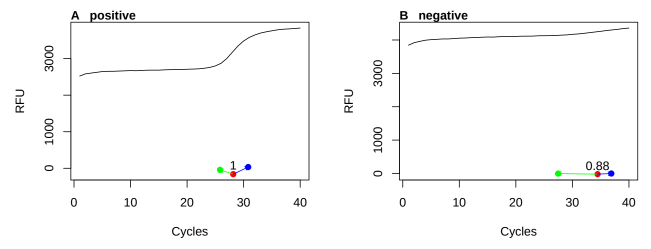


Figure 8: Concept of the winklR() function. Analysis of the amplification curves of the 'RAS002' data set with the winklR() function. Two amplification curves (A: positive, B: negative) were used. The red point shows the origin (first negative derivative maximum) while the green and blue points show the minimum and maximum of the second negative derivative. The angle is calculated from these points. Positive curves have smaller angles than negative curves.

3.3.7 Quantification Points, Ratios and Slopes

The pcrfit_single() function calculates cpD1 and cpD2 and uses them for further analysis. Both the cpD1 and cpD2 value are used to describe the amplification reaction quantitatively. For example, low cpD1 and cpD2 values (< 5 cycles) indicate that the PCR reaction was negative or that the amount of input DNA was to high Figure 7B. Since the pcrfit_single() function gives the parameters of all models (subsubsection 3.3.5) they are part of the feature set for completeness (Figure 10). In particular, the results of the five-parameter function and of the seven-parameter function are reported.

Further predictors from the pcrfit_single() function are:

• eff is the optimized PCR efficiency found within a sliding window (Figure 6C). A linear model of cycles versus log(Fluorescence) is fit within a sliding window (for details see sliwin() [qpcR] function).

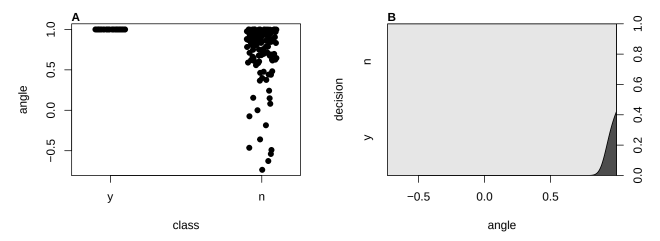


Figure 9: Analysis of the amplification curves of the 'RAS002' data set with the winklR() function. All amplification curves of the data set 'RAS002' were analyzed. Negative amplification curves are shown in red and positive amplification curves in black. The winklR() function was used to analyze all amplification curves. B) The stripchart of the analysis of positive and negative amplification curves shows separation. C) The cdplot calculates the conditional densities of x based on the values of y weighted by the boundary distribution of y. The densities are derived cumulatively via the values of y. The probability that the decision is negative (n) when the angle equals 30 is approximately 100%.

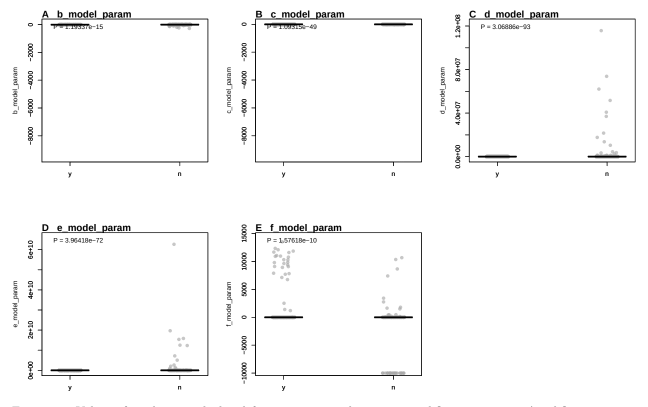


Figure 10: Values of predictors calculated from negative and positive amplification curves. Amplification curve predictors from the 'data_sample_subset' data set were used as they contain positive and negative amplification curves, as well as amplification curves that exhibit a *hook effect* or non-sigmoid shapes. A) 'c_model_param', is the c model parameter of the seven parameter model. B) 'd_model_param', is the d model parameter of the seven parameter model. C) 'e_model_param', is the e model parameter of the seven parameter model. D) 'f_model_param', is the f model parameter of the seven parameter model. The classes were compared using the Wilcoxon Rank Sum Test.

- The comparison of positive and negative amplification curves in Figure 12A demonstrates that the classes are significantly different from each other. The eff values differ significantly between negative and positive amplification curves (Figure 12A).
- sliwin is the PCR efficiency by the 'window-of-linearity' method (Spiess, Feig, and Ritz 2008) (Figure 12B). The sliwin values differ significantly between negative and positive amplification curves (Figure 12B).
- cpDdiff is the difference between the first (cpD1) and the second derivative maximum cpD2 (cpDdiff = cpD1 cpD2) from the fitted model (Figure 6C). Provided that a model can be exactly fitted, the estimates of the difference are reliable. Higher cpDdiff values indicate a negative amplification reaction or a very low amplification efficiency. The comparison of positive and negative amplification curves in Figure 12C demonstrates that the classes are significantly different from each other. In the event that the cpDdiff value cannot be determined (NA), it is replaced by zero. The cpDdiff values differ significantly between negative and positive amplification curves (Figure 12C).
- cpD2_range is the absolute value of the difference between the minimum and the maximum of the second derivative maximum (cpD2_range = |cpD2m cpD2|) from the diffQ2() function (no model fitted) (Figure 11E). The cpD2_range value does not require an adjustment of a multiparametric model. The approximate first and second derivatives are determined using a five-point stencil (Rödiger, Burdukiewicz, and Schierack 2015). The comparison of positive and negative amplification curves in Figure 12E shows that the classes differ significantly from each other. In the event that the cpD2_range value cannot be determined (NA), it is replaced by zero. The cpD2_range values differ significantly between negative and positive amplification curves (Figure 12E).
- cpD2_approx is the approximate second derivative maximum. In most cases the value should be close to the cpD2. Deviations indicate noise in the data, negative amplification curves or positive amplification curves that deviate from a typical sigmoid amplification curve.
- cpD2_ratio is is the ratio between the the approximate second derivative maximum cpD2_approx and the second derivative maximum cpD2 (cpD2_ratio = cpD2/cpD2_approx). In the event that the cpD2_ratio value cannot be determined (NA, Inf), it is replaced by zero. Provided that a model can be exactly fitted and that the function has little noise the ratio between both values should be close to 1. Note: Empirical data suggest that an interval of 0.85 and 1.1 indicates positive amplification curves. These can be set to 1. Values outside this interval indicate non-sigmoidal (e.g., negative) amplification curves. These can be set to 0.
- bg.stop is the end of the ground phase and amp.stop is the end of the exponential phase estimated by the bg.max() [chipPCR] function (Rödiger, Burdukiewicz, and Schierack 2015). A graphical presentation of the locations in the amplification curve are shown in Figure 11. The bg.stop and bg.stop values differ significantly between negative and positive amplification curves (Figure 12J & K).
- top is the takeoff point as proposed by Tichopad et al. (2003). The top is calculated using externally studentized residuals, which tested to be an outlier in terms of the t-distribution. The top signifies to first PCR cycle entering the exponential phase. tdp is the takedown point. This is an implementation in the pcrfit_single() function, which uses the rotated $f(x) \mapsto f_1(f(x))$ and flipped g(x) = -(x) amplification curve for calculation. Figure 5A describes the location of top and tdp. The position (f.top, f.tdp) on the ordinate is also determined from these points. If an amplification curve is negative or neither top nor tdp can be calculated, then top & tdp will be assigned the number of cycles and f.top & f.tdp the value 1. The distribution of top, tdp, f.top and f.tdp is shown in Figure 12F-I. The top, tdp, f.top and f.tdp values differ significantly between negative and positive amplification curves. Potentially they enable a qualitative classification of the amplification reaction. An interesting aspect is that the positive f.top values are markedly lower than the negative f.top values The same applies inversely to the tdp values. In this way, amplification curves can be classified according to these values.
- peaks_ratio is based on a sequential chaining of functions. The diffQ() [MBmca] function determines numerically the first derivative of an amplification curve. This derivative is passed to the mcaPeaks() [MBmca] function. In the output all minima and all maxima are contained. The ranges are calculated from the minima and maxima. The Lagged Difference is determined from the ranges of the minima and

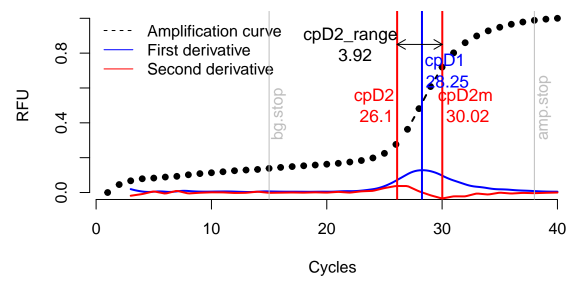


Figure 11: Location of the predictors 'cpD2_range', 'bg.start', 'bg.stop' within an amplification curve. The minimum (cpD2m) and maximum (cpD2) of the second derivative were calculated numerically using the diffQ2() function. This function also returns the maximum of the first derivative (cpD1). The 'cpD2_range' is defined as $cpD2_range = |cpD2-cpD2m|$. Large 'cpD2_range' values indicate a low amplification efficiency or a negative amplification reaction. The predictor 'bg.start' is an estimate for the end of the ground phase. 'bg.start' is an approximation for the onset of the plateau phase.

maxima. Finally, the ratio of the differences (maximum/minimum) is calculated. The peaks_ratio values differ significantly between negative and positive amplification curves (Figure 23B).

• loglin_slope is calculated from the slope determined by a linear model of the data points from the cycle dependent fluorescence at the minimum of the second derivative and maximum of the second derivative (Figure 13), provided that the locations of the minimum of the second derivative and the maximum of the second derivative yield a *suitable* interval. As a precaution, the algorithm checks, for example, whether the distance between the minimum of the second derivative and the maximum of the second derivative is not more than nine PCR cycles. Failing this, the loglin_slope value is set to zero (no slope), as in the example of Figure 13. The loglin_slope values differ significantly between negative and positive amplification curves (Figure 12D).

The predictor loglin_slope is used in the following to test if the slope within this ROI can be used to distinguish positive and negative amplification curves. The hypothesis is that positive amplification curves have a higher loglin_slope than negative amplification curves. As shown in Figure 12D, there is a statistically significant difference between positive and negative amplification curves.

The loglin_slope values from the data_sample_subset_balanced data set (subsubsection 3.3.1) were used to save computing time. A binomial logistic regression (see subsubsection 3.2.1)) was used to analyze the relationship between the loglin_slope value and the class (negative, positive). The data set was split into two chunks. This is an important step during such applications. One chunk is for adapting, i. e. training, the model and the other chunk for testing the model. By convention, 70% to 80% of the data is used for training (Walsh, Pollastri, and Tosatto 2015; Kuhn 2008). The binomial logistic regression model was adapted using the glm() [stats] function by using the parameter family = binomial(link = 'logit'). To objectify the splitting, the sample() [stats] function was used.

```
library(PCRedux)

data <- data_sample_subset_balanced

n_positive <- sum(data[["decision"]] == "y")</pre>
```

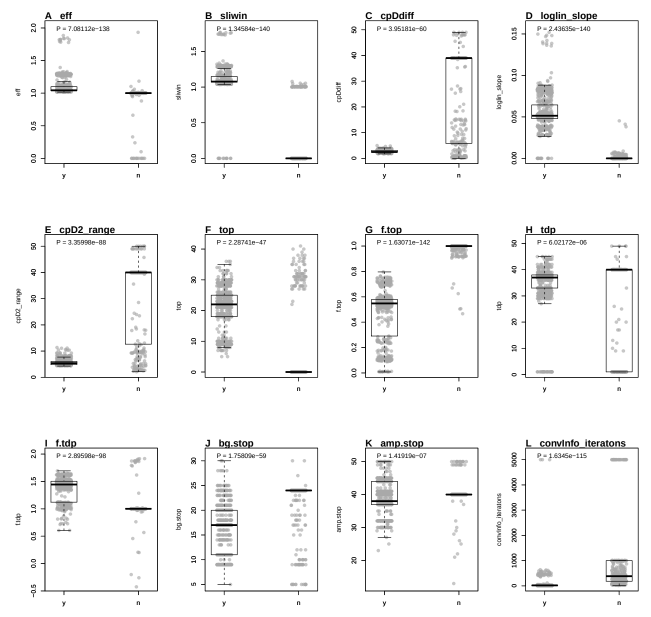


Figure 12: Values of predictors calculated from negative and positive amplification curves. Amplification curve predictors from the 'data_sample_subset' data set were used as they contain positive and negative amplification curves, and amplification curves that exhibit a hook effect or non-sigmoid shapes. A) 'eff', optimized PCR efficiency found within a sliding window. B) 'sliwin', PCR efficiency by the 'window-of-linearity' method. C) 'cpDdiff', difference between the Cq values calculated from the first and the second derivative maximum. D) 'loglin_slope', slope from the cycle at the second derivative maximum to the second derivative minimum. E) 'cpD2_range', absolute value of the difference between the minimum and the maximum of the second derivative maximum. F) 'top', takeoff point. G) 'f.top', fluorescence intensity at takeoff point. H) 'tdp', takedown point. I) 'f.tdp', fluorescence intensity at takedown point. J) 'bg.stop', estimated end of the exponential phase. L) 'convInfo iterations', number of iterations until convergence.

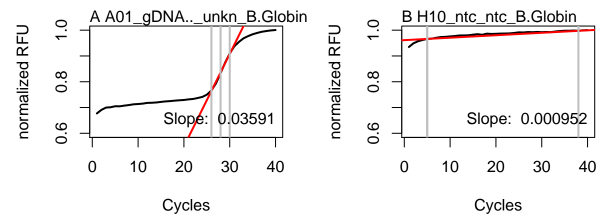


Figure 13: Concept of the 'loglin_slope' predictor. The algorithm determines the fluorescence values of the raw data at the approximate positions of the maximum off the first derivative, the minimum of the second derivative and the maximum of the second derivative, which are in the exponential phase of the amplification curve. The data were taken from the 'RAS002' data set. A linear model is created from these parameter sets and the slope is determined. A) Positive amplification curves have a clearly positive slope. B) Negative amplification curves usually have a low, sometimes negative slope.

```
n negative <- sum(data[["decision"]] == "n")</pre>
dat <- data.frame(loglin_slope = data[, "loglin_slope"],</pre>
                   decision = as.numeric(factor(data$decision,
                                                 levels = c("n", "y"),
                                                 label = c(0, 1)) - 1)
# Select randomly observations from 70% of the data for training.
# n_train is the number of observations used for training.
n_train <- round(nrow(data) * 0.7)</pre>
pasteO("Percentage of observations (", n_train, ") = ",
       signif((n_train/nrow(data)*100),3), "%")
## [1] "Percentage of observations (456) = 70%"
# index test is the index of observations to be selected for the training
index_test <- sample(1L:nrow(dat), size = n_train)</pre>
# index_test is the index of observations to be selected for the testing
index_training <- which(!(1L:nrow(dat) %in% index_test))</pre>
# train_data contains the data used for training
train_data <- dat[index_test, ]</pre>
# test_data contains the data used for training
test_data <- dat[index_training, ]</pre>
# Fit the binomial logistic regression model
```

The summary() function returns the results of the model fitting. This can be analysed and interpreted. summary(model glm)

```
##
## Call:
## glm(formula = decision ~ loglin_slope, family = binomial(link = "logit"),
       data = train_data)
##
##
## Deviance Residuals:
##
       Min
                   1Q
                         Median
                                       3Q
                                                Max
## -3.02640 -0.28970 -0.28970
                                  0.01636
                                            2.52661
##
## Coefficients:
                Estimate Std. Error z value Pr(>|z|)
##
                                    -9.975
                                              <2e-16 ***
## (Intercept)
                -3.1499
                             0.3158
## loglin_slope 170.7028
                            18.5689
                                      9.193
                                              <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
       Null deviance: 632.12 on 455 degrees of freedom
## Residual deviance: 119.21 on 454 degrees of freedom
## AIC: 123.21
##
## Number of Fisher Scoring iterations: 8
```

Based on the results it can be concluded that the parameters (intercept) and loglin_slope are statistically significant (P < 2e-16). This indicates an association between loglin_slope and the probability that an amplification curve is positive.

In order to apply the model to a new data set, further steps are necessary. predict() [stats] is a generic function for predicting the results of a model fitting function (Figure 14A). All previously split test data is passed to the function argument newdata. By setting the type = 'response' parameter, the predict() function returns probabilities in the form of P(y = 1|X). In the case in hand, it was decided that a decision limit of 0.5 is to be applied. If P(y = 1|X) < 0.5 then y = 0 (amplification curve negative), otherwise y = 1 (amplification curves positive).

```
library(PCRedux)

# Create graphic device for the plot(s)
# Plot train_data (grey points) and the predicted model (blue)
par(mfrow = c(1,2))

plot(train_data$loglin_slope, train_data$decision, pch = 19,
```

```
xlab = "loglin_slope", ylab = "Probability",
     col = adjustcolor("grey", alpha.f = 0.9), cex = 1.5)
mtext("A", cex = 1, side = 3, adj = 0, font = 2, las = 0)
abline(h = 0.5, col = "grey")
curve(predict(model_glm, data.frame(loglin_slope = x), type = "resp"),
      add = TRUE, col = "blue")
# Plot test data (red)
points(test_data$loglin_slope, test_data$decision, pch = 19,
       col = adjustcolor("red", alpha.f = 0.3))
legend("right", paste("Positive: ", n_positive,
                      "\nNegative: ", n_negative), bty = "n")
# Plot the sensitivity, specificity and other measures to describe
# the prediction.
position_bp <- barplot(as.matrix(res_performeR), yaxt = "n",</pre>
                       ylab = "Probability", main = "", las = 2,
                       col = adjustcolor("grey", alpha.f = 0.5),
                       border = "white")
par(srt = 90)
text(position bp, rep(0.8, length(res performeR)),
    paste(signif(res_performeR, 2)*100, "%"), cex = 0.6)
axis(2, at = c(0, 1), labels = c("0", "1"), las = 2)
abline(h = 0.85, col = "grey")
mtext("B", cex = 1, side = 3, adj = 0, font = 2, las = 0)
```

Sensitivity, specificity and further parameters for estimating predictions were calculated using the performeR() function (subsubsection 3.6.1). The results indicate that the sensitivity and specificity for the test data set provides a good result. However, in this case, they depend heavily on the computer-aided random sampling of the training data, and the total size of the data set (Figure 14B). Over-fitting and under-fitting and other problems need to be addressed (Walsh, Pollastri, and Tosatto 2015).

To proof the results, further methods such as Likelihood Ratio Test, McFadden's \mathbb{R}^2 , k-fold cross-validation, Receiver Operating Characteristic (ROC) analysis and model interpretation should be used (Arlot and Celisse 2010; McFadden 1974; Sing et al. 2005).

• sd_bg is the standard deviation from the first PCR cycle to the takeoff point (Figure 5A). Manufacturers of thermo-cyclers use different sensors and data processing algorithms. The same applies to the detection chemistry used in experiments subsubsection 3.2.2. The signal variation in the ground phase differs between the different systems (Figure 3D). If no takeoff point can be determined from an amplification curve, the value for sd_bg is calculated from the first to the eighth PCR cycle. The results for the predictor sd_bg were broken down by the thermo-cycler and the output of the amplification reaction (negative, positive). It can be seen that the signal variation between the thermo-cyclers seems to be different. There is also a difference between negative and positive amplification curves Figure 15. The sd_bg values differ significantly between negative and positive amplification curves (Figure 16J).

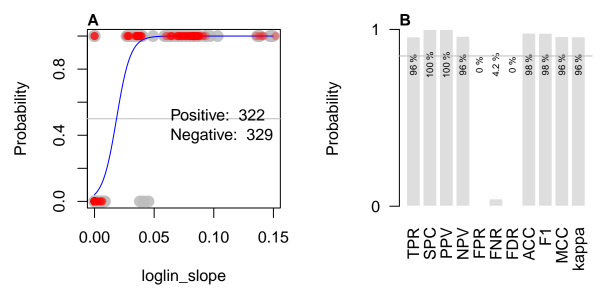


Figure 14: Machine classification by means of binomial logistic regression using the 'loglin_slope' predictor. A) For the calculation of a binomial logistic regression model, the categorical response variable Y (decision with classes: negative and positive) must be converted to a numerical value. With binomial logistic regression, the probability of a categorical response can be estimated using the X predictor variable. In this example, the predictor variable 'loglin_slope' is used. Grey measurement points (70% of the data set) were used for training. Red dots represent the values used for testing. The regression curve of the binomial logistic regression is shown in blue. The grey horizontal line at 0.5 marks the threshold of probability above which it is determined whether an amplification curve is negative or positive. B) The performance indicators were calculated using the performeR() function. Sensitivity, TPR; Specificity, SPC; Precision, PPV; Negative prediction value, NPV; Fall-out, FPR; False negative rate, FNR; False detection rate, FDR; Accuracy, ACC; F1 score, F1; Matthews correlation coefficient, MCC, Cohens kappa (binary classification), kappa (κ).

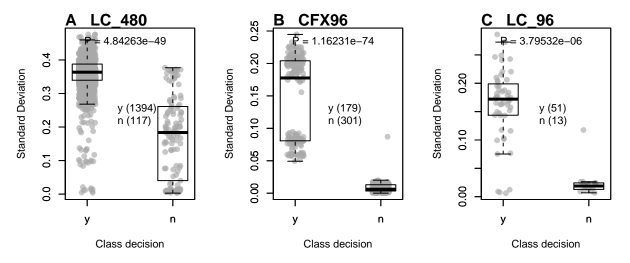


Figure 15: Standard deviation in the ground phase of various qPCR devices. The 'sd_bg' predictor was used to determine if the standard deviation between thermo-cyclers and between positive and negative amplification curves was different. The standard deviation was determined from the fluorescence values from the first cycle to the takeoff point. If the takeoff point could not be determined, the standard deviation from the first cycle to the eighth cycle was calculated. The Mann-Whitney test was used to compare the medians of the two populations (y, positive; n, negative). The differences were significant for A) LC_480 (Roche), B) CFX96 (Bio-Rad) and C) LC96 (Roche).

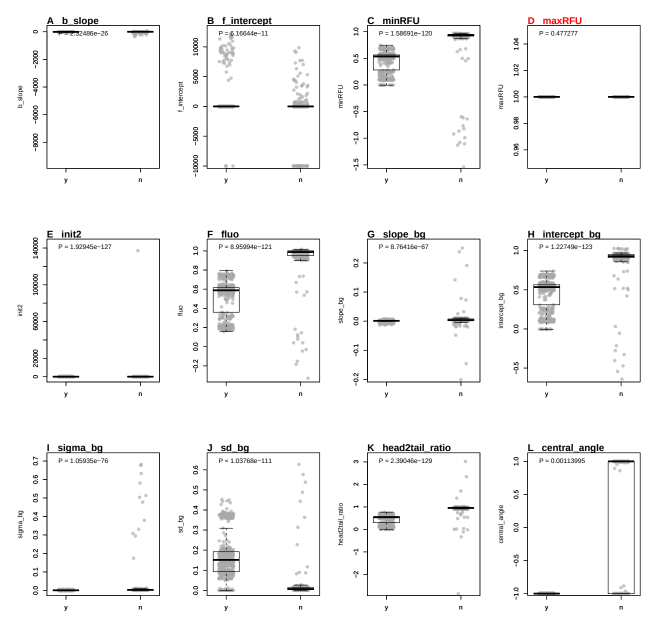


Figure 16: Values of predictors calculated from negative and positive amplification curves. Amplification curve predictors from the 'data_sample_subset' data set were used as they contain positive and negative amplification curves, as well as amplification curves that exhibit a hook effect or non-sigmoid shapes. A) 'eff', optimized PCR efficiency in a sliding window. B) 'sliwin', PCR efficiency according to the window-of-linearity method. C) 'cpDdiff', difference between the Cq values calculated from the first and the second derivative maximum. D) 'loglin_slope', slope from cycle at second derivative maximum to second derivative minimum. E) 'cpD2_range', absolute difference between the minimum and maximum of the second derivative. F) 'top', takeoff point. G) 'f.top', fluorescence intensity at takeoff point. H) 'tdp', takedown point. I) 'f.tdp', fluorescence intensity at the takedown point. J) 'bg.stop', estimated end of the ground phase. K) 'amp.stop', estimated end of the exponential phase. L) 'convInfo_iteratons', number of iterations until convergence when fitting a multiparametric model. The classes were compared using the Wilcoxon Rank Sum Test.

3.3.8 Integration of the amptester() function in PCRedux

amptester_polygon is another method to calculate the area under an amplification curve. amptester_polygon⁷ is part of the amptester() [chipPCR] package (Rödiger, Burdukiewicz, and Schierack 2015). In contrast to the implementation in amptester(), amptester_polygon has values normalized to the total number of cycles, thereby allowing comparable predictions (Figure 23Ds).

⁷This predictor is determined from the points in an amplification curve (like a polygon, in particular non-convex polygons) in a 'clockwise' order. The sum over the edges result in a positive value if the amplification curve is 'clockwise' and is negative if the curve is 'counter-clockwise'.

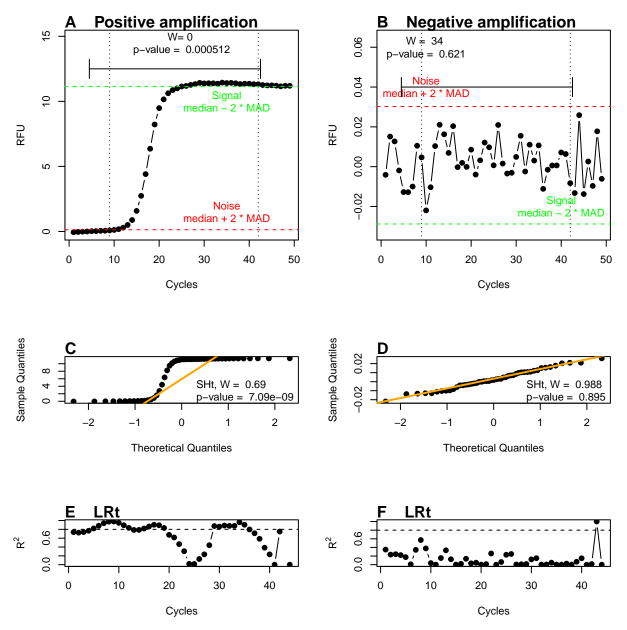


Figure 17: Analysis of amplification curves with the "amptester()" function. A & B) The threshold test (THt) is based on the Wilcoxon ranksum test and compares 20% of the fluorescence values of the ground phase with 15% of the plateau phase. In the example, a significant difference (p=0.000512) was found for the positive amplification curve. However, this did not apply to the negative amplification curve (p=0.621). C & D) A Q-Q diagram is used to graphically compare two probability distributions. In this study the probability distribution of the amplification curve was compared with a theoretical normal distribution. The orange line is the theoretically normal quantil-quantile plot that passes through the probabilities of the first and third quartiles. The Shapiro-Wilk test (SHt) of normality checks whether the underlying measurement data of the amplification curve is significantly normal distributed. Since the p-value of $7.09e^{-9}$ of the positive amplification curve is $\alpha \le 5e^{-4}$, the null hypothesis is rejected. However, this does not apply to the negative amplification curve (p=0.895). E & F) The linear regression test (LRt) calculates the coefficient of determination (R^2) using an ordinary least square regression where all measured values are integrated into the model in a cycle-dependent manner. Experience shows that the non-linear part of an amplification curve has a R^2 smaller than 0.8, which is also shown in the example.

3.3.9 earlyreg() - A Function to Calculate the Slope and Intercept in the Ground Phase of an Amplification Curve

The signal height and the slope in the first cycles (1 - 10) of amplification curves are potentially useful because some qPCR systems calibrate themselves by fluorescence intensity of the first cycles. This is noticeable as strong signal changes which appear spontaneously between the first and second cycle (e. g., Figure 1B). Furthermore, the signal level can be used to determine which background signal is present and whether the ground phase already has a slope. Moreover, characteristics of the detection probe system are noticeable (see subsubsection 3.2.4). From the slope, it may be deduced whether amplification has already started (see subsubsection 3.2.4).

Consequently, the earlyreg() function was developed. This function uses an ordinary least squares linear regression within a limited number of cycles. As ROI, the first 10 cycles were defined. This restriction is based on the developers experience, suggesting that during the first ten cycles only a significant increase in signal strength can be measured within few qPCRs. However, earlyreg() does not ignore the first cycle, as many thermo-cyclers use this cycle for sensor calibration. Extreme values are therefore included. As standard, the next nine amplitude values are used for the linear regression. The number of cycles can also be adjusted via the parameter range. Since all amplification curves are normalized to the 99%-percentile, comparability between the background signals and the slopes is ensured. The output of the earlyreg() function is:

- slope_bg, which is the slope of the ordinary least squares linear regression model,
- intercept_bg, which is the intercept of the linear model and
- sigma_bg, which is the square root of the estimated variance of the random error.

The slope_bg, intercept_bg and sigma_bg values differ significantly between negative and positive amplification curves (Figure 16G-I).

The following example illustrates possible usage of earlyreg(). For that purpose, amplification curves from the C127EGHP data set were analysed (Figure 18A). In figure Figure 18A the amplification curves for all cycles are shown. Next, the earlyreg() function was used to determine the slope_bg, intercept_bg and sigma_bg in the range of the first ten PCR cycles. The results were used in a cluster analysis using k-means clustering, demonstrating that the slope seems to be an indicator of differences between the amplification curves. The first 8 cycles were colored according to their cluster (Figure 18B). After cluster analysis, the same could also be observed (Figure 18D-F). Hence, it can be postulated that the slope in the background phase is useful for the amplification curve classification.

```
library(PCRedux)
# box_cox() function for the Box-Cox transformation of data
box cox <- function(x, lambda = 1, offset = 0) {
    if (lambda == 0) {
        log(x + offset)
    } else
    {
        ((x + offset)^lambda - 1)/lambda
    }
}
# Load the C127EGHP data set
data <- chipPCR::C127EGHP[, -1]</pre>
# Normalize each amplification curve to their 0.99 percentile and use the
# earlyreg() function to determine the slope and intercept of the first
# cycles 'user_range'
user range <- 8
```

```
res_earlyreg <- do.call(rbind, lapply(2L:ncol(data), function(i) {</pre>
  earlyreg(x = data[, 1], y = data[, i], range = user_range, normalize = TRUE)
})) %>% box cox(.)
# Label the observation with their names
rownames(res_earlyreg) <- substr(colnames(data)[-1], 1, 10)
# Show the first five lines of the res earlyreg data matrix
head(res_earlyreg)
##
        intercept
                       slope
                                   sigma
## EG1 -0.9996982 -1.0000257 -0.9994170
## EG2 -0.9999187 -0.9999724 -0.9994349
## EG3 -1.0001748 -0.9999236 -0.9995357
## EG4 -1.0000410 -0.9999487 -0.9994961
## EG5 -1.0001467 -0.9999259 -0.9995627
## EG6 -1.0003437 -0.9998825 -0.9995230
# Perform k-means clustering on the res_earlyreg data matrix
cl <- kmeans(res_earlyreg, centers = 2)</pre>
# Plot the results
# Use x roi (cycles) and rfu range (RFU values) to limit the
# range for the detailed plot of first cycles.
x_roi <- 1:(user_range + 1)</pre>
rfu_range <- range(data[x_roi, -1])
# Create graphic device for the plot(s)
layout(matrix(c(1, 2, 1, 2, 3, 3), 3, 2, byrow = TRUE))
# Plot of raw amplification curves
matplot(
    data[, 1], data[, -1], ylim = range(data[, -1]), pch = 19, lty = 1,
        type = "l", xlab = "Cycles", ylab = "RFU", main = "", col = "grey"
mtext("A", cex = 1, side = 3, adj = 0, font = 2)
abline(v = c(1,user_range))
text(3, range(data[, -1])[2], "ROI")
# Detailed plot of the first cycles and the clusters according
# to the k-means clustering
# Define some user colors (blue: EvaGreen, orange: Hydrolysis probes)
colors <- c(</pre>
    adjustcolor("blue", alpha.f = 0.5),
            adjustcolor("orange", alpha.f = 0.8)
)
matplot(NA, NA, xlim = range(x_roi), ylim = rfu_range, xlab = "Cycles",
        ylab = "RFU", main = "")
for(i in 1L:length(unique(cl$cluster))) {
```

```
cl_id <- which(cl[["cluster"]] == i) + 1</pre>
par(new=TRUE)
matplot(data[x_roi, 1], xlab = "", ylab = "", xaxt = "n", yaxt = "n",
        data[x_roi, cl_id], ylim = rfu_range, pch = 19, lty = 1, type = "l",
        col = colors[i])
}
mtext("B", cex = 1, side = 3, adj = 0, font = 2)
abline(v = c(1,user_range))
text(3, rfu_range[2], "ROI")
legend("bottomleft", c("Cluster 1", "Cluster 2"), pch = 15, cex = 1.2,
       col = colors, bty = "n")
# Overview of clusters and corresponding detection chemistry
eghp <- rep(0.7, length(cl$cluster))
names(eghp) <- names(cl$cluster)</pre>
barplot(eghp, las = 2, col = colors[cl[["cluster"]]],
        border = "white", xlab = "", ylab = "",
       yaxt = "n", ylim = c(0,2.2), cex.axis = 0.7)
mtext("C", cex = 1, side = 3, adj = 0, font = 2)
legend("topleft", c("Cluster 1", "Cluster 2"), pch = 15, col = colors,
       bty = "n", box.col = "white", cex = 0.9)
arrows(0.5, 2, 38, 2, angle = 90, code = 3)
arrows(39, 2, 76.5, 2, angle = 90, code = 3)
text(c(18.75, 57.5), c(1.6, 1.6), c("EvaGreen", "Hydrolysis probes"))
```

3.3.10 head2tailratio() - A Function to Calculate the Ratio of the Head and the Tail of a Quantitative PCR Amplification Curve

The ratios from the ground and plateau phase can be used to search for patterns in amplification curves. Positive amplification curves have different slopes and intercepts at the start (head, background region) and the end (tail, plateau region) of the amplification curve. Hence, these regions are potentially useful to extract a predictor for amplification curve classification. Negative amplification curves (no slope) are assumed to have a ratio of about 1. In contrast, positive amplification curves should have a ratio of less than 1.

The n-dimensional space of all predictor variables $X_{1,2...n}$ is also called feature space. In the present study the feature space was extended by domain knowledge using known features. The head2tailratio()-function is an example for this. Here the feature $X_3 \equiv head2tail_ratio$ could be calculated by determining the ratio of the $X_1 \equiv fluorescence$ intensity in the head region and $X_2 \equiv fluorescence$ intensity in the tail region of a quantitative PCR amplification curve $(X_3 = \frac{X_1}{X_2})$. As ROI, the areas in the ground phase (head) and plateau phases (tail) are used (Figure 5A). For the calculation, the median from the first six data points of the amplification curve and the median from the last six data points are used. The determination of six data points in both regions was made on the basis of empirical experience. As a rule, no significant increase in amplification signals can be measured in the first six cycles and in the last six cycles (where the amplification curve usually transitions into the plateau). This assumption is sometimes violated (e. g., hook effect) and might lead to false estimates.

```
library(PCRedux)
# Load the RAS002 amplification curve data set and assign it to the object data
```

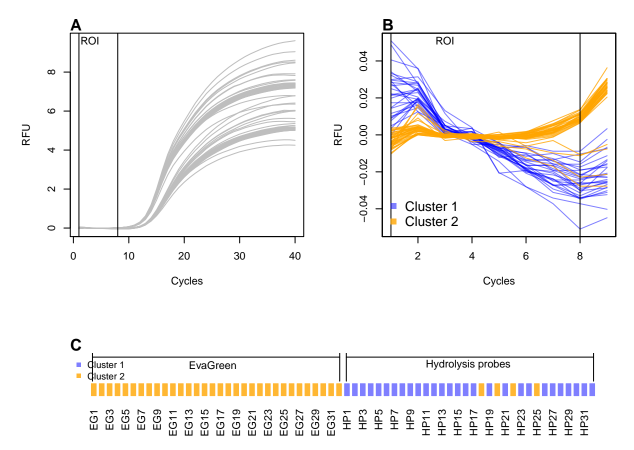


Figure 18: Analysis of the ground phase with the early reg() function and the 'C127EGHP' data set (n = 64 amplification curves). This data set consists of 32 samples, which were simultaneously monitored with the intercal ator EvaGreen or hydrolysis probes. A) All amplification curves possess slightly different slopes and intercepts in the first cycles of the ground phase (ROI: Cycles 1 to 8). Both the slope and the intercept of each amplification curve were used for cluster analysis (k-means, Hartigan-Wong algorithm, number of centers k = 2). B) The amplification curves were assigned to five clusters, depending on their slope and their intersection (red, black). C) Finally, the clusters were associated to the detection chemistries (EvaGreen (EG) or hydrolysis probes (HP)).

```
data <- RAS002
# Load the RASOO2 decision data set and assign it to the object data decisions
data_decisions <- RAS002_decisions
# Calculate the head2tailratio of all amplification curves
res_head2tailratio <- lapply(2L:ncol(data), function(i) {</pre>
 head2tailratio(
    y = data[, i], normalize = TRUE, slope_normalizer = TRUE,
    verbose = TRUE
  )
})
# Fetch all values of the head2tailratio analysis for a later comparison
# by a boxplot.
res <- sapply(1L:length(res_head2tailratio), function(i)</pre>
  res_head2tailratio[[i]]$head_tail_ratio)
data_normalized <- cbind(</pre>
  data[, 1],
  sapply(2L:ncol(data), function(i) {
    data[, i] / quantile(data[, i], 0.99)
  })
)
# Assign colors to the classes (n: black, y: green).
colors <- as.character(factor(</pre>
  data_decisions, levels = c("y", "n"),
  labels = c(
      adjustcolor("green", alpha.f = 0.5), adjustcolor("black", alpha.f = 0.5))
))
res_wilcox.test <- stats::wilcox.test(res ~ data_decisions)
```

The amplification curves in Figure 19 show a signal increase within the first three cycles, and those in Figure 5C have a negative slope in the tail. The median is used to minimize the influence of outliers.

```
# Plot the results of the analysis
#
# Position and plot parameters
h <- max(na.omit(res))
h_text <- rep(h * 0.976, 2)

# Create graphic device for the plot(s)
layout(matrix(c(1,1,2), 1, 3, byrow = TRUE))

matplot(
   data_normalized[, 1], data_normalized[, -1],
   xlab = "Cycles", ylab = "normalized RFU", main = "",
   type = "l", lty = 1, lwd = 2, col = colors
)
for (i in 1L:(ncol(data_normalized) - 1)) {</pre>
```

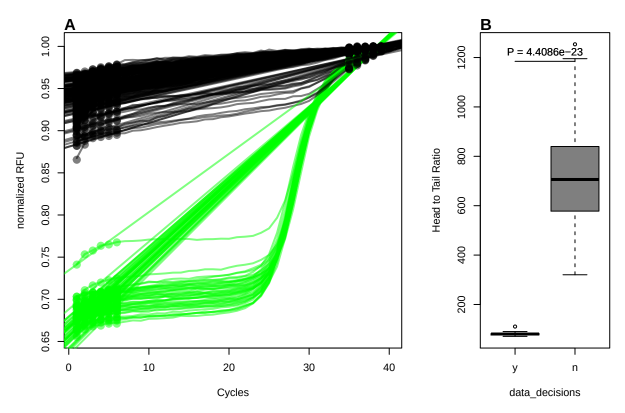


Figure 19: Ratio between the head and the tail of a quantitative PCR amplification curve. A) Plot of quantile normalized amplification curves from the 'RAS002' data set. Data points used in the head and tail are highlighted by circles. The intervals for the Robust Linear Regression are automatically selected using the 25% and 75% quantiles. Therefore, not all data points are used in the regression model. The straight line is the regression line from the robust linear model. The slopes of the positive and negative amplification curves differ. B) Boxplot for the comparison of the head/tail ratio. Positive amplification curves have a lower ratio than negative curves. The difference between the classes is significant.

```
points(
    res_head2tailratio[[i]]$x_roi, res_head2tailratio[[i]]$y_roi,
    col = colors[i], pch = 19, cex = 1.5
)
    abline(res_head2tailratio[[i]]$fit, col = colors[i], lwd = 2)
}
mtext("A", cex = 1, side = 3, adj = 0, font = 2)

# Boxplot of the head2tail ratios of the positive and negative
# amplification curves.

boxplot(res ~ data_decisions, col = unique(colors), ylab = "Head to Tail Ratio")

lines(c(1, 2), rep(h * 0.945, 2))
text(1.5, h_text, paste0("P = ", signif(res_wilcox.test[["p.value"]])),
    cex = 1)

mtext("B", cex = 1, side = 3, adj = 0, font = 2)
```

In subsubsection 3.2.4 and in Figure 1, it was shown that negative amplification curves may have a slope with a positive or negative sign. There is no consent in the literature and among peers how to deal with this during

the processing. One solution is to include the slope as factor in the ratio calculation. The head2tailratio() function uses a linear model that calculates the slope between the ground and plateau phases. If the slope of the model is significant, then the ratio from the head and tail is normalized to this slope. This requires setting the slope_normalizer parameter in the head2tailratio() function. By default, this parameter is not set.

The head2tail_ratio values differ significantly between negative and positive amplification curves (Figure 16K).

3.3.11 hookreg() and hookregNL() - Functions to Detect Hook Effect-like Curvatures

hookreg() and hookregNL() are functions to detect amplification curves bearing a hook effect (Barratt and Mackay 2002) or negative slope at the end of the amplification curve. Both functions calculate the slope and intercept of an amplification curve data. The assumption is that a strong negative slope at the end of an amplification curve is indicative for a hook effect. hookreg() and hookregNL() are part of a peer-reviewed publication (Burdukiewicz et al. 2018). For this reason, the functions will not be discussed here.

3.3.12 mblrr() - A Function to Perform the Quantile-filter Based Local Robust Regression

mblrr() is a function to perform the median based local robust regression (mblrr) from a quantitative PCR experiment. In detail, this function attempts to break the amplification curve in two ROIs (head (~background) and tail (~plateau)). As opposed to the earlyreg() function, the mblrr() function does not use a fixed interval. Instead, the mblrr() function dynamically determines cut points for each amplification curve. It was defined that:

- the 25% quantile is the value for which 25% of all values are smaller than this value.
- the 75% quantile is the value for which 75% of all values are greater than this value.

Subsequent, a robust linear regression analysis (lmrob()) is preformed individually on both regions of the amplification curve. The rationale behind this analysis is that the slope and intercept of an amplification curve differ in the background and plateau region. This is also shown by the simulations in Figure 3A-C. In the example shown below, the observations "P01.W19", "P06.W35", "P33.W66", "P65.W90", "P71.W23" and "P87.W01" were arbitrarily selected for demonstration purposes Figure 20. Another example is shown in Figure 29A. Those amplification curves have a slight negative trend in the base-line region and a positive trend in the plateau region.

The correlation coefficient 8 is a measure to quantify the dependence on variables (e. g., number of cycles, signal height). The correlation coefficient is always between -1 and 1, with a value close to -1 describing a strong-negative dependency and close to 1 describing a strong-positive dependency; if the value is 0, there is no dependency between the variables. The most frequently used correlation coefficient to describe a linear dependency is the Pearson correlation coefficient r. The correlation coefficient can be used as a predictor. Similar data structures have similar correlation coefficients. However, variables that are not strongly correlated can also be important for modeling.

```
library(PCRedux)

# Select four amplification curves from the RASO02 data set
amplification_curves <- c(2, 3, 4, 5, 44, 45)
data <- RASO02[, c(1, amplification_curves)]

# Load the decision_res_htPCR.csv data set from a csv file.
filename <- system.file("decision_res_RASO02.csv", package = "PCRedux")
decision_res <- read.csv(filename)

# Overview of the amplicon curve classifications
res_decision <- decision_res[amplification_curves - 1, -c(3,4)]</pre>
```

⁸Product moment correlation coefficient (Pearson)

```
res_decision
##
                        RAS002 test.result.1 conformity
## 1 A01_gDNA.._unkn_B.Globin
                                                    TRUE
                                           У
## 2
         A01 gDNA.. unkn HPRT1
                                                    TRUE
                                           n
## 3 A02_gDNA.._unkn_B.Globin
                                                    TRUE
                                            У
         A02_gDNA.._unkn_HPRT1
                                                    TRUE
                                            n
## 43 B10_gDNA.._unkn_B.Globin
                                                    TRUE
                                            n
        B10_gDNA.._unkn_HPRT1
                                                    TRUE
# Plot the regions and the linear regression line in the
# amplification curve plot
colors <- c(
            adjustcolor("blue", alpha.f = 0.5),
            adjustcolor("orange", alpha.f = 0.8)
# Create graphic device for the plot(s)
par(mfrow = c(3, 2))
for (i in 2L:ncol(data)) {
  x <- data[, 1]
  y_tmp <- data[, i] / quantile(data[, i], 0.99)</pre>
  res_q25 <- y_tmp < quantile(y_tmp, 0.25)</pre>
  res_q75 <- y_tmp > quantile(y_tmp, 0.75)
  res_q25_lm <- try(
    suppressWarnings(lmrob(y_tmp[res_q25] ~ x[res_q25])),
    silent = TRUE
  )
  res_q75_lm <- try(
    suppressWarnings(lmrob(y_tmp[res_q75] ~ x[res_q75])),
    silent = TRUE
  )
  plot(x, y_tmp, xlab = "Cycles", ylab = "RFU (normalized)",
    main = "", type = "b", pch = 19)
  mtext(paste0(LETTERS[i - 1], " ", colnames(data)[i]), cex = 1,
        side = 3, adj = 0, font = 2)
  legend("topleft", paste0(ifelse(res_decision[i - 1, 2] == "n",
                                   "negative", "positive")),
         bty = "n")
  abline(res_q25_lm, col = colors[1])
  points(x[res_q25], y_tmp[res_q25], cex = 2.5, col = colors[1])
  abline(res_q75_lm, col = colors[2])
```

Finally, the results of the analysis were printed in a tabular format.

 $points(x[res_q75], y_tmp[res_q75], cex = 2.5, col = colors[2])$

```
# Load the xtable library for an appealing table output
library(xtable)
# Analyze the data via the mblrr() function
```

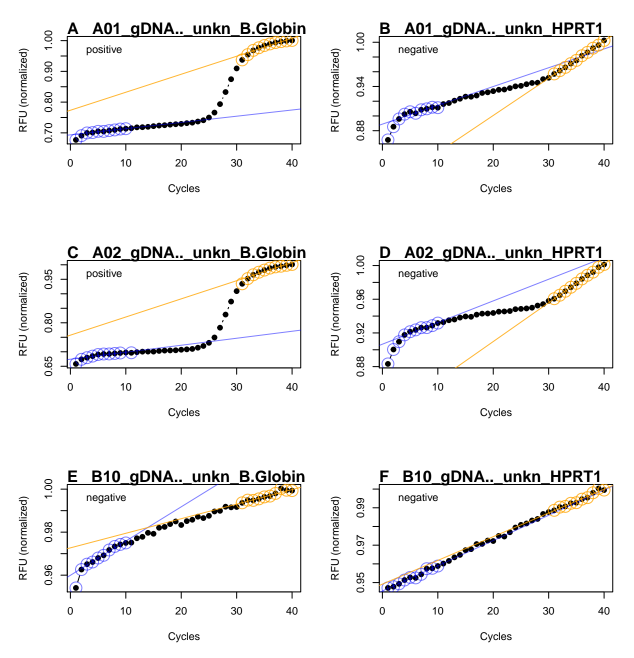


Figure 20: Robust local regression to analyze amplification curves. The amplification curves were arbitrarily selected from the 'RAS002' data set. In the qPCR setup, the target genes beta globin (B. globin) and HPRT1 were simultaneously measured in a PCR cavity using two specific hydrolysis probes (duplex qPCR). Both positive (A, C, E) and negative (B, D, F) amplification curves were used. The amplification curves are normalized to the 99% quantile. The differences in slopes and intercepts (blue and orange lines and dots). The mblrr() function is presumably useful for data sets which are accompanied by noise and artifacts.

```
res_mblrr <- do.call(cbind, lapply(2L:ncol(data), function(i) {</pre>
  suppressMessages(mblrr(
    x = data[, 1], y = data[, i],
    normalize = TRUE
  )) %>% data.frame()
colnames(res_mblrr) <- colnames(data)[-1]</pre>
# Transform the data for a tabular output and assign the results to the object
# output_res_mblrr.
output_res_mblrr <- res_mblrr %>% t()
# The output variable names of the mblrr() function are rather long. For better
# readability the variable names were changed to "nBG" (intercept of the head
# region), "mBG" (slope of the head region), "rBG" (Pearson correlation of head
# region), "nTP" (intercept of the tail region), "mTP" (slope of tail region),
# "rBG" (Pearson correlation of the tail region)
colnames(output_res_mblrr) <- c(</pre>
  "nBG", "mBG", "rBG",
  "nTP", "mTP", "rTP"
print(xtable(
  output_res_mblrr, caption = "Selected results of predictors from the mblrr()
             function. nBG, intercept of head region; mBG, slope of head region;
             rBG, Pearson correlation of head region; nTP, intercept of tail
             region; mTP, slope of tail region; rBG, Pearson correlation of
             tail region",
  label = "tablemblrrintroduction"
), comment = FALSE, caption.placement = "top")
```

Table 1: Selected results of predictors from the mblrr() function. nBG, intercept of head region; mBG, slope of head region; rBG, Pearson correlation of head region; nTP, intercept of tail region; mTP, slope of tail region; rBG, Pearson correlation of tail region

	nBG	mBG	rBG	nTP	mTP	rTP
A01_gDNAunkn_B.Globin	0.69	0.00	0.91	0.77	0.01	0.94
A01_gDNAunkn_HPRT1	0.89	0.00	0.87	0.80	0.01	1.00
A02_gDNAunkn_B.Globin	0.67	0.00	0.88	0.76	0.01	0.95
A02_gDNAunkn_HPRT1	0.91	0.00	0.90	0.82	0.00	1.00
B10_gDNAunkn_B.Globin	0.96	0.00	0.95	0.97	0.00	0.96
B10_gDNAunkn_HPRT1	0.95	0.00	0.99	0.95	0.00	0.98

In another example, the results from the $\mathtt{mblrr}()$ function were combined with human classifications (positive, negative) to apply them in an analysis with Fast and Frugal Trees (FFTrees). FFTrees belong to class of simple decision rules. DT's are a classic approach to machine learning (Quinlan 1986). Here relatively simple algorithms and simple tree structures are used to create a model. A general introduction to decision trees is given in (Quinlan 1986; Luan, Schooler, and Gigerenzer 2011). In many situations, FFTrees make fast decisions based on a few predictors (N = 1 - 5). In this example six predictors were used for the analysis.

The FFTrees package (Phillips et al. 2017) provides an implementation for the R statistical computing language. All that is needed for the present example are:

- the data assessed by the mblrr() function,
- the classification of the amplification curve data by a human,
- and a standard formula, which looks like $outcome \sim var1 + var2 + \dots$ along with the data arguments. The function FFTrees() returns a fast and frugal tree object. This rich object contains the underlying trees and many classification statistics (similar to subsubsection 3.6.1). In the following example, the RASO02 data set was used.

```
# Load the xtable library for an appealing table output
library(FFTrees)
library(PCRedux)
# The RASOO2 amplification curves were analyzed with the mblrr() function
# to save computing time and the.results of this analysis are stored in the
# `data_sample` data set.
data <- data_sample[data_sample$dataset == "RAS002", c("mblrr_intercept_bg",</pre>
                                                         "mblrr_slope_bg",
                                                         "mblrr_cor_bg",
                                                         "mblrr intercept pt",
                                                         "mblrr_slope_pt",
                                                         "mblrr_cor_pt")]
# The output variable names of the mblrr() function are rather long. For better
# readability the variable names were changed to "nBG" (intercept of head
# region), "mBG" (slope of head region), "rBG" (Pearson correlation of head
# region), "nTP" (intercept of tail region), "mTP" (slope of tail region),
# "rBG" (Pearson correlation of tail region).
res_mblrr <- data.frame(</pre>
    class = as.numeric(as.character(factor(RAS002_decisions,
                                            levels = c("y", "n"),
                                            label = c(1, 0))),
  data
)
colnames(res_mblrr) <- c("class", "nBG", "mBG", "rBG", "nTP", "mTP", "rTP")</pre>
res mblrr.fft <- suppressMessages(</pre>
            FFTrees(formula = class ~., data = res_mblrr)
```

Figure 21 shows the Fast and Frugal Trees by using the predictors nBG (intercept of head region), mBG (slope of head region), rBG (Pearson correlation of head region), nTP (intercept of tail region), mTP (slope of tail region), and rBG (Pearson correlation of tail region).

R offers several packages like party (Hothorn, Hornik, and Zeileis 2006), rpart (Therneau, Atkinson, and Ripley 2017) and Rattle (Williams 2009) for creating and visualizing decision trees.

3.3.13 autocorrelation_test() - A Function to Detect Positive Amplification Curves

Autocorrelation analysis is a technique that is used in the field of time series analysis. It can be used to reveal regularly occurring patterns in one-dimensional data (Spiess et al. 2016). Autocorrelation measures the correlation of a signal f(t) with itself shifted by some time delay $f(t - \tau)$.

The autocorrelation_test() function coerces the amplification curve data to an object of class "zoo" (zoo

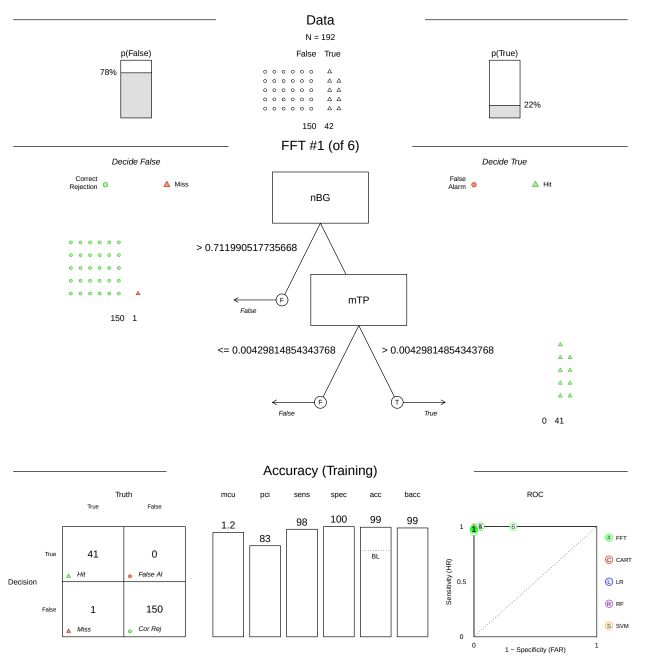


Figure 21: Visualization of decisions in Fast and Frugal Trees after data analysis of amplification curves via the mblrr() function. Top row 'Data') Overview of the data set, stating the total number of observations (N = 192) and percentage of positive (22%) and negative (78%) amplification curves. Middle row 'FFT #1 (of 6)') Decision Tree with the number of observations classified at each level of the tree. For the analysis, six predictors (nBG, intercept of head region; mBG, slope of head region; rBG, Pearson correlation of head region; nTP, intercept of tail region; mTP, slope of tail region; rBG, Pearson correlation of tail region) have been used for the analysis. After two tree levels (nBG, nTP), the decision tree is created, where all positive amplification curves (N = 40) are correctly classified. Two observations are classified as false-negative in the negative amplification curves. Lower row 'Performance') The FFTrees() [FFTrees] function determines several performance statistics. For the training data, there is a classification table on the left side showing the relationship between tree 'decision' and the 'truth'. The correct rejection ('Cor Rej') and 'Hit' are the right decisions. 'Miss' and false alarm ('False Al') are wrong decisions. The centre shows the cumulative tree performance in terms of mean of used cues ('mcu'), Percent of ignored cues ('pci'), sensitivity ('sens'), specificity ('spec'), accuracy ('acc') and weighted Accuracy ('wacc'). The receiver operating characteristic (ROC) curve on the right-hand side compares the performance of all trees in the FFTrees object. The system also displays the performance of the fast frugal trees ('#', green), CART ('C', red), logistical regression ('L', blue), random forest ('R', violet) and the support vector machine ('S', yellow).

package) as indexed totally ordered observations. Then follows the computation of a lagged version of the amplification curve data. The shifting of the amplification curve data is based on the number of observations (number of cycles 'c') with the following τ .

Number of Cycles (c)	τ
$c \le 35$	8
$35 > c \le 40$	10
$40 < c \le 45$	12
c > 45	14

This is followed by a significance test for correlation between paired observations (original & lagged amplification curve data). The hypothesis is that paired observations of positive amplification curves exhibit significant correlation (stats::cor.test, significance level is 0.01) in contrast to negative amplification curves (noise). The application of the autocorrelation_test() function is shown in the following example.

In addition, the decisions file decision_res_RASO02.csv from the user was analyzed for the most frequent decision (modus) using the decision_modus() function (subsubsection 3.5.3).

```
# Test for autocorrelation in amplification curve data. The amplification
# curve data from the `htPCR` data set was used.
library(PCRedux)
# Load the decision_res_htPCR.csv data set from a csv file.
filename <- system.file("decision res htPCR.csv", package = "PCRedux")
decision_res_htPCR <- read.csv(filename)</pre>
# Select only amplification curves (obs) were all
# classifications were in concordance ("conformity == TRUE" ).
# This subset of the htPCR data set contains the classes
# n: negative
# a: ambiquous
# p: positive
obs_number <- which(decision_res_htPCR[["conformity"]] == TRUE)</pre>
dec <- unlist(decision_res_htPCR[obs_number, "test.result.1"])</pre>
# Give tabular output of classes
table(dec)
## dec
##
     a
         n
##
     2 202 280
# Since the number of ambiguous is low (n = 2), they were re-assigned to the
# class negative
dec[dec == "a"] <- "n"
table(dec)
## dec
##
     a
        n
##
    0 204 280
```

```
# Load only the amplification curves from the htPCR data set that were
# uniquely assigned to one class (e. g., eight out of eight positive).
data <- qpcR::htPCR[, c(1, obs_number + 1)]</pre>
# Assign colors to the classes (n: black, y: green).
colors <- factor(dec, levels = c("y", "n"), label = c("black", "green"))</pre>
# Test for autocorrelation in the subset of the htPCR data set
res_ac <- sapply(2:ncol(data), function(i) {</pre>
    autocorrelation_test(data[, i], ns_2_numeric = TRUE)
})
# Plot curve data as overview
# Names of the observations
# Create graphic device for the plot(s)
layout(matrix(c(1, 2, 3, 1, 4, 4), 2, 3, byrow = TRUE))
matplot(
    data[, 1], data[, -1], xlab = "Cycles", ylab = "RFU",
    main = "", type = "1", lty = 1,
    col = colors, lwd = 2
)
legend("topleft", c("positive", "negative"), pch = 19, col = c(1, 2), bty = "n")
mtext("A
            RAS002 data set", cex = 1, side = 3, adj = 0, font = 2)
# Convert the n.s. (not significant) in 0 and others to 1.
# Combine the results of the aromatic autocorrelation_test as variable "ac",
# the human classified values as variable "hc" in a new data frame (res_ac_hc).
cutoff <- 0.85
res ac hc <- as.matrix(data.frame(</pre>
    ac = ifelse(res_ac > cutoff, 1, 0),
                        hc = ifelse(dec == "y", 0, 1)
))
res_performeR <- performeR(s = res_ac_hc[, "ac"], r = res_ac_hc[, "hc"])</pre>
plot(density(res_ac), xlab = "Autocorrelation", ylab = "Density", main = "")
rug(res_ac)
abline(v = cutoff)
mtext("B", cex = 1, side = 3, adj = 0, font = 2, las = 0)
cdplot(res_ac, droplevels(dec), xlab = "Autocorrelation", ylab = "Class decision")
```

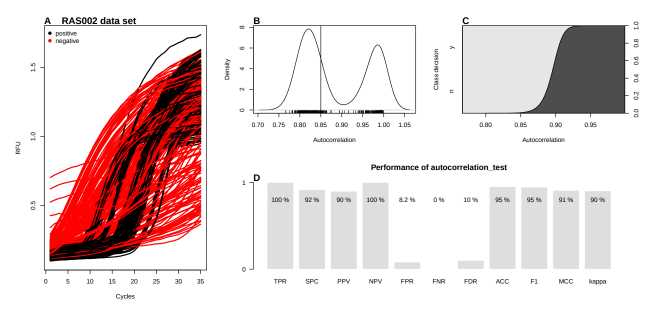


Figure 22: Autocorrelation analysis of the amplification curves of the 'RAS002' data set. A) Display of all amplification curves of the data set 'RAS002'. Negative amplification curves are shown in red and positive amplification curves in black. The autocorrelation_test() function was used to analyze all amplification curves. B) The density diagram of the autocorrelation of positive and negative amplification curves shows a bimodal distribution. C) The cdplot calculates the conditional densities of x based on the values of y weighted by the boundary distribution of y. The densities are derived cumulatively via the values of y. The probability that the decision is negative (n) when autocorrelation equals 0.85 is approximately 100%. D) Performance analysis using the performeR() function (see subsubsection 3.6.1 for details).

```
mtext("C", cex = 1, side = 3, adj = 0, font = 2, las = 0)

barplot(
    as.matrix(res_performeR[, c(1:10, 12)]), yaxt = "n", ylab = "",
        main = "Performance of autocorrelation_test",
        col = adjustcolor("grey", alpha.f = 0.5), border = "white"
)

text(position_bp, rep(0.8, length(res_performeR[, c(1:10, 12)])),
    paste(signif(res_performeR[, c(1:10, 12)], 2)*100, "%"))

axis(2, at = c(0, 1), labels = c("0", "1"), las = 2)

mtext("D", cex = 1, side = 3, adj = 0, font = 2, las = 0)
```

As shown in this example, the autocorrelation_test() function is able to distinguish between positive and negative amplification curves. Negative amplification curves were in all cases non-significant. In contrast, the coefficients of correlation for positive amplification curves ranged between 0.766 and 0.999 at a significance level of 0.01 and a lag of 3.

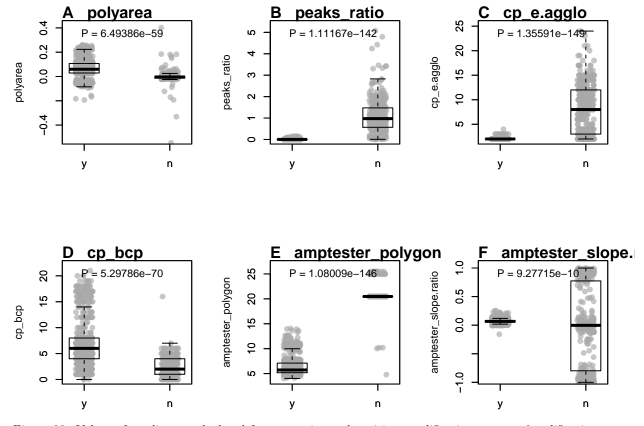


Figure 23: Values of predictors calculated from negative and positive amplification curves. Amplification curves predictors from the 'data_sample_subset' data set were used as they contain positive and negative amplification curves and amplification curves that exhibit a hook effect or non-sigmoid shapes. A) 'polyarea', is the area under the amplification curve determined by the Gauss polygon area formula. B) 'peaks_ratio', is the ratio of the local minima and the local maxima. C) 'cp_e.agglo', makes use of energy agglomerative clustering. Positive amplification curves have fewer change points than negative amplification curves. These two change point analyses generally separate positive and negative amplification curves. D) 'cp_bcp', analyses change points by a Bayesian approach. Positive amplification curves appear to contain more change points than negative amplification curves. Nevertheless, there is an overlap between the positive and negative amplification curves in both methods. This can lead to false-positive or false-negative classifications. E) 'amptester_polygon' is the cycle normalized order of a polygon. F) 'amptester_slope.ratio' is the slope (linear model) of the raw fluorescence values at the approximate first derivate maximum, second derivative minimum and second derivative maximum.

3.3.14 Frequentist and Bayesian Change Point Analysis

Change point analysis (CPA) encompasses methods to identify or estimate single or multiple locations of distributional changes in a series of data points indexed in time order. A change herein refers to a statistical property. CPA is used for example in econometrics and bioinformatics (Killick and Eckley 2014; Erdman, Emerson, and others 2007). Several change point algorithms exist, such as the binary segmentation algorithm (Scott and Knott 1974). In change point analysis one assumes independent ordered observations $x_1, x_2, \ldots, x_n \in \mathbb{R}^d$ (N. A. James and Matteson 2013). In the case of qPCR, this is simply the cycle-dependent fluorescence, used to create k homogeneous subsets of unknown size (Erdman, Emerson, and others 2007). While frequentist methods make an estimation of the parameter at the location (e. g., mean) of the change points at specific points, change point analysis using Bayesian method produces a probability for the occurrence of a change point at certain points. For the analysis of the amplification curves, it was hypothesized that the number of change points differs between positive (sigmoidal) and negative (noise) amplification curves.

The pcrfit_single() function uses two independent approaches for change point analysis. These are the bcp() [bcp] (Erdman, Emerson, and others 2007) and the e.agglo() [ecp] function (N. A. James

and Matteson 2013). The e.agglo() function performs a non-parametric change point analysis based on agglomerative hierarchical estimation and is useful to "detect changes within the marginal distributions" (N. A. James and Matteson 2013). Measurements from the qPCR systems typically show noise that has rapidly changing components. Differentiators amplify these rapidly changing noise components (Rödiger, Böhm, and Schimke 2013). Therefore, the first derivation of the amplification curve was used for both change point analyses. It was assumed for the change point analysis of amplification curves, that this leads to larger differences between positive and negative amplification curves. An example is shown on Figure 24. In contrast the bcp() [bcp] function performs a change point analysis based on a Bayesian approach. This method can detect changes in the mean of independent Gaussian observations. As a result, the analysis returns the posterior probability of a change point at each x_i . An example is shown on Figure 24. Both the change point analysis methods provide additional information to distinguish positive and negative amplification curves Figure 23E & F).

3.3.15 Frequentist Approaches to Test the Class of an Amplification Reaction and Application of the amptester() Predictors

A part of pcrfit_single() is the amptester() [chipPCR] function, which contains tests to determine whether an amplification curve is positive or negative. The input values for the function differ due to the different preprocessing steps in the pcrfit_single() function. Therefore, the concepts of the tests are briefly described below.

- The first test, designated as SHt, is based on this Shapiro-Wilk test of normality. This relatively simple procedure can be used to check whether the underlying population of a sample (amplification curve) is significantly ($\alpha \le 5e 04$) normal distributed. The name of the output of the pcrfit_single() function is amptester_shapiro.
- The second test is the *Resids growth test* (RGt), which tests if the fluorescence values in linear phase are stable. Whenever no amplification occurs, fluorescence values quickly deviate from linear model. Their standardized residuals will be strongly correlated with their value. For real amplification curves, the situation is much more stable. Noise (meaning deviations from linear model) in background do not correlate strongly with the changes in fluorescence. The decision is based on the threshold value (here 0.5). The output is binary coded (negative = 0, positive = 1). The output name of the pcrfit_single() function is amptester_rgt.
- The third test is the Linear Regression test (LRt). This test determines the coefficient of determination (R^2) by an ordinary least squares linear (OLS) regression. The R^2 are determined from a run of $\sim 15\%$ range of the data. If a sequence of more than six R^2 s larger than 0.8 is found, a nonlinear signal is plausible. This is somewhat counter-intuitive, because R^2 of nonlinear data should be low. The output is binary coded (negative = 0, positive = 1). The output name of the pcrfit_single() function is amptester_lrt.
- The fourth test is called *Threshold test* (THt), based on the Wilcoxon rank sum test. As a simple rule the first 20% (head) and the last 15% (tail) of an amplification curve are used as input data. From this, a one-sided Wilcoxon rank sum tests of the head versus the tail is performed ($\alpha \le 1e 02$). The output is binary coded (negative = 0, positive = 1). The output name of the pcrfit_single() function is amptester_tht.
- The fifth test is called Signal level test (SLt). The test compares the signals of the head and the tail by a robust "sigma" rule (median + 2 * MAD) and the comparison of the head/tail ratio. If the returned value is less than 1.25 (25 percent), then the amplification curve is likely negative. The output is binary coded (negative = 0, positive = 1). The output name of the pcrfit_single() function is amptester_slt.
- The sixth test is called *Polygon test* (pco). The pco test determines if the points in an amplification curve (like a polygon) are in a "clockwise" order. The sum over the edges result in a positive value if the amplification curve is "clockwise" and is negative if the curve is counter-clockwise. Experience states that noise is positive and "true" amplification curves are "highly" negative. In contrast to the implementation in the amptester() function, the result is normalized by a division to the number of PCR cycles. The output is numeric. The output name of the pcrfit_single() function is amptester_polygon.
- The seventh test is the Slope Ratio test (SIR). This test uses the approximated first derivative maximum,

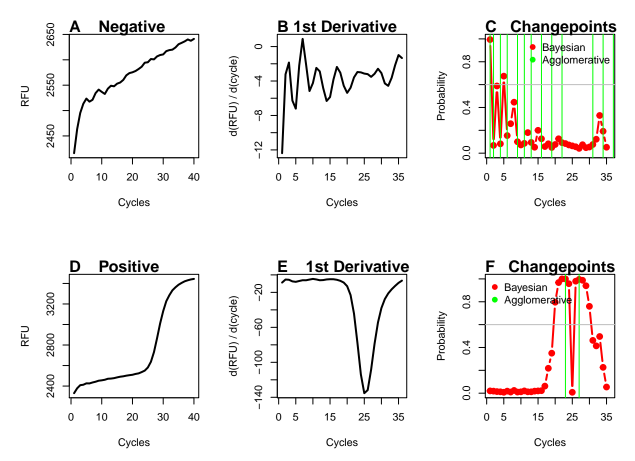


Figure 24: Bayesian and energy agglomerative change point analysis on negative and positive amplification curves. An analysis of a negative and a positive amplification curve from the 'RAS002' data set was performed using the pcrfit_single() function. In this process, the amplification curves were analysed for change points using Bayesian change point analysis and energy agglomerative clustering. A) The negative amplification curve has a base signal of approximately 2450 RFU and only a small signal increase to 2650 RFU. There is a clear indication of the signal variation (noise). B) The first negative derivative amplifies the noise so that some peaks are visible. C) The change point analysis shows changes in energy agglomerative clustering at several positions (green vertical line). The Bayesian change point analysis rarely exceeds a probability of 0.6 (grey vert line). D) The positive amplification curve has a lower base signal (~ 2450 RFU) and increases up to the 40th cycle (~ 3400 RFU). A sigmoid shape of the curve is visible. E) The first negative derivation of the positive amplification curve shows a distinctive peak with a minimum at cycle 25. F) The change point analysis in energy agglomerative clustering shows changes (green vertical line) only at two positions. The Bayesian change point analysis shows a probability higher than 0.6 (grey horizontal line) at several positions.

the second derivative minimum and the second derivative maximum of the amplification curve. Next, the raw fluorescence at the approximated second derivative minimum and the second derivative maximum are taken from the original data set. The fluorescence intensities are normalized to the maximum fluorescence of this data and then employed in a linear regression, using the estimated slope. The output is numeric and the output name of the pcrfit_single() function is amptester_slope.ratio.

Random Forest is an enhancement of decision tree algorithms. Random Forest uses n random data subsets, by creating an ensemble consisting of n small decision trees. Each decision tree contains a biased classifier. Only classes that provide reliable prediction to the outcome are then selected for classification. Compared to a single tree classifier, Random Forests display a high robustness against noise, outliers and over-fitting (Williams 2009; Breiman 2001).

In the following example, the randomForest() [randomForest] function (Liaw and Wiener 2002) was used for classification. In classification problems, one tries to predict a discrete number of values, in this case a binary classification. The aim of this **proof-of-concept** in silico experiment was to find the most important predictors for the classification of positive and negative amplification curves. As response vector (y) (in R type vector) the decision served with its possible finite classes labeled as "positive" and "negative".

```
• amptester_shapiro,
```

- amptester lrt,
- amptester_rgt,
- amptester_tht,
- amptester_slt,
- amptester_polygon and
- amptester_slope.ratio served as a matrix of predictors describing the model to be adapted. The data_sample_subset_balanced data set (subsubsection 3.3.2) was used for the analysis to save computing time. The data_sample_subset_balanced data set contains similar proportions of observations (positive and negative amplification curves.

```
library(randomForest)
library(PCRedux)
set.seed(1999)
# Dimensions of the data_sample_subset_balanced object
dim(data_sample_subset_balanced)
## [1] 651 62
# Show proportions of positive and negative amplification curves in
# data_sample_subset_balanced
table(data sample subset balanced[["decision"]])
##
##
     У
## 322 329
data <- as.matrix(cbind(data sample subset balanced[, c("amptester shapiro",</pre>
                            "amptester_lrt",
                            "amptester rgt",
                            "amptester_tht",
                            "amptester slt",
                            "amptester_polygon",
                            "amptester slope.ratio")],
                  decision = as.numeric(
                      factor(data_sample_subset_balanced$decision,
                                          levels = c("n", "y"),
```

```
label = c(0, 1)) - 1)
                )
# Summary of data
summary(data)
   amptester_shapiro amptester_lrt
                                      amptester_rgt
                                                       amptester_tht
## Min.
          :0.0000
                     Min. :0.0000
                                      Min.
                                            :0.0000
                                                       Min.
                                                              :0.0000
                                                       1st Qu.:1.0000
## 1st Qu.:0.0000
                     1st Qu.:1.0000
                                      1st Qu.:1.0000
## Median :0.0000 Median :1.0000
                                      Median :1.0000
                                                      Median :1.0000
         :0.4117 Mean
                           :0.9785
                                                             :0.9708
## Mean
                                      Mean
                                            :0.9462
                                                       Mean
## 3rd Qu.:1.0000 3rd Qu.:1.0000
                                      3rd Qu.:1.0000
                                                       3rd Qu.:1.0000
          :1.0000 Max.
## Max.
                           :1.0000
                                      Max.
                                             :1.0000 Max.
                                                              :1.0000
## amptester slt
                    amptester_polygon amptester_slope.ratio
                                                               decision
## Min.
         :0.0000 Min. : 4.018
                                      Min.
                                            :-0.999888
                                                            Min.
                                                                   :0.0000
## 1st Qu.:0.0000
                    1st Qu.: 5.505
                                      1st Qu.:-0.008099
                                                            1st Qu.:0.0000
## Median :1.0000
                    Median :14.053
                                      Median : 0.050291
                                                            Median :0.0000
## Mean
         :0.5008
                    Mean
                           :13.765
                                      Mean
                                            : 0.034182
                                                            Mean
                                                                  :0.4946
## 3rd Qu.:1.0000
                    3rd Qu.:20.475
                                      3rd Qu.: 0.128971
                                                            3rd Qu.:1.0000
## Max.
          :1.0000
                    Max.
                           :25.480
                                      Max.
                                             : 0.999209
                                                            Max.
                                                                   :1.0000
# Select randomly 70% of the observations data for training (\rightarrow n_train).
# n_train is the number of observations used for training.
# First determine the number of observations that would cover 70% of all data.
n train <- round(nrow(data) * 0.7)</pre>
pasteO("Percentage of observations (n = ", n_train, ") -> ",
       signif((n_train/nrow(data)*100), 3), "%")
## [1] "Percentage of observations (n = 456) -> 70%"
# index_test is the index of observations to be selected for the testing
index_test <- sample(1L:nrow(data), size = n_train)</pre>
# index_test is the index of observations to be selected for the training
index_training <- which(!(1L:nrow(data) %in% index_test))</pre>
# The randomForest() function was used to train a forest of 200 trees.
# Convert decision into factor, since randomForest() uses type="regression",
# instead of type="classification" by default. Each tree is trained on 63.2 %
# of the training data. Each observation is drawn at random with replacement
# from the original data.
# The predictor variables are drawn at random out of the feature space
# "amptester_shapiro", "amptester_lrt", "amptester_rgt", "amptester_tht",
\hbox{\# "amptester\_slt", "amptester\_polygon" and "amptester\_slope.ratio"}.
model_rf <- randomForest(as.factor(decision) ~ ., data = data,</pre>
                        subset = index_training,
                        ntree = 200, importance = TRUE)
# The prediction accuracy of the random forest is summarized
```

The number of variables randomly selected at each split is mtry = 2 (as starter, the square root of total number of all predictors is used). The out-of-bag (OOB) error (= misclassification rate) is 0.516%. To fine tune a random forest model, the number of ntree values must be varied and plotted against the OOB rate. It recommended to select a mtry value with minimum OOB error, which was not done in this example.

Confusion matrix:

0 96 1 0.01030928 ## 1 0 98 0.00000000

##

0 1 class.error

```
# Determine variable importance
res_importance <- importance(model_rf)

print(xtable::xtable(res_importance, caption = "Results of the random forest
classification.",
label = "randomforstresults"
), comment = FALSE, caption.placement = "top")</pre>
```

Table 3: Results of the random forest classification.

	0	1	MeanDecreaseAccuracy	MeanDecreaseGini
amptester_shapiro	1.72	7.85	7.80	14.84
$amptester_lrt$	0.00	3.67	3.63	0.28
$amptester_rgt$	1.90	-1.00	1.88	0.07
$amptester_tht$	1.85	3.61	4.02	0.57
$amptester_slt$	10.98	12.93	12.64	34.49
$amptester_polygon$	11.76	13.73	13.65	36.18
$amptester_slope.ratio$	1.13	5.56	5.40	7.79

The variables MeanDecreaseAccuracy and MeanDecreaseGini are used to determine the importance of the variables for a classification from a Random Forest model. MeanDecreaseAccuracy tells how much removing each variable reduces the accuracy of the model. Mean Decrease Gini tells how important a variable is based on the Gini impurity index used for the calculation of splits in trees. The higher the values, the more significant they are.

```
# Create graphic device for the plot(s)
par(mfrow = c(1,3))

# Plot the properties if the Random Forest model

plot(model_rf, main = "", las = 2)
mtext("A", cex = 1, side = 3, adj = 0, font = 2, las = 0)

rownames(res_importance) <- substr(rownames(res_importance), 11, 22)</pre>
```

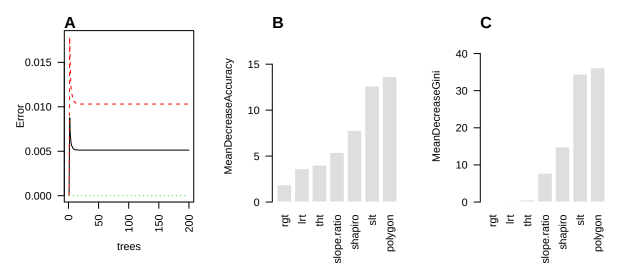


Figure 25: The predictors 'amptester_lrt' (lrt), 'amptester_rgt' (rgt), 'amptester_tht' (tht), 'amptester_slt' (slt), 'amptester_polygon'(polygon) and 'amptester_slope.ratio' (slope.ratio) were used for classification using random forest. A) This plot shows the error depending on the number of trees. The error decreases as more and more trees are added and averaged. B). Mean Decrease Accuracy shown how much the model accuracy decreases if a variable is dropped. C) Mean Decrease Gini shows the importance of a variable based on the Gini impurity index used for the calculation of splits in trees.

The top two important predictors are polygon (MeanDecreaseAccuracy = 13.65, MeanDecreaseGini = 36.18) and slt (MeanDecreaseAccuracy = 12.64, MeanDecreaseGini = 34.49).

3.4 Classified Amplification Curve Datasets

Amplification curves from different sources (e. g., detection chemistries, thermo-cyclers) were manually classified with the humanrater() function (subsubsection 3.5.1) or with the tReem() function (subsubsection 3.5.2). Raw amplification curve data were exported as comma separated values or in the Real-time PCR Data Markup Language (RDML) format via the RDML package. RDML is human readable data exchange format for qPCR experiments. A detailed description can be found in Rödiger et al. (2017). The following code section describes the import of an RDML file from the PCRedux package. The RDML file contains amplification curve data of a duplex qPCR (HPV 16 & HPV 18) performed in the CFX96 (Bio-Rad).

At least the following datasets with a dichotomous decision (positive, negative) are included.

- decision res batsch1.csv
- decision res batsch2.csv
- decision res batsch3.csv
- decision res batsch4.csv
- decision_res_batsch5.csv
- decision_res_boggy.csv
- decision res C126EG595.csv
- decision res C127EGHP.csv
- decision res C316.amp.csv
- decision_res_C317.amp.csv
- decision_res_C60.amp.csv
- decision res CD74.csv
- decision res competimer.csv
- decision res dil4reps94.csv
- decision res guescini1.csv
- decision res guescini2.csv
- decision res HCU32 aggR.csv
- decision_res_htPCR.csv
- decision_res_karlen1.csv
- decision_res_karlen2.csv
- decision res karlen3.csv
- decision_res_lc96_bACTXY.csv
- decision res lievens1.csv
- decision res lievens2.csv
- decision res lievens3.csv
- decision_res_RAS002.csv
- decision_res_RAS003.csv
- decision_res_reps2.csv
- \bullet decision_res_reps384.csv
- decision res reps3.csv
- decision res reps.csv
- decision res rutledge.csv
- decision_res_stepone_std.csv
- decision_res_testdat.csv
- decision_res_vermeulen1.csv
- decision_res_vermeulen2.csv
- decision res VIMCFX96 60.csv

```
library(RDML)
# Load the RDML package and use its functions to import the amplification curve
# data
library(RDML)
filename <- system.file("RAS002.rdml", package = "PCRedux")
raw_data <- RDML$new(filename = filename)</pre>
```

The following example shows the export of the RASO02.rdml file from the RDML format to the csv format.

```
# Export the RDML data from the PCRedux package as the objects RAS002 and RAS003.
library(RDML)
library(PCRedux)

RAS002 <- data.frame(RDML$new(paste0(
    path.package("PCRedux"), "/", "RAS002.rdml"))$GetFData()
    )

# The obbject RAS002 can be stored in the working directory as CSV file with
# the name RAS002_amp.csv.
write.csv(RAS002, "RAS002_amp.csv", row.names = FALSE)</pre>
```

RDML data file	Device	Target gene	Detection chemistry
RAS002.rdml	CFX96	HPV16, HPV18, HPRT1	TaqMan
RAS003.rdml	CFX96	HPV16, HPV18, HPRT1	TaqMan
hookreg.rdml	Bio-Rad	various	TaqMan, DNA binding dyes

3.5 Technologies for Amplification Curve Classification and Classified Amplification Curves

Many machine learning concepts exist. One method is supervised machine learning, where the goal is to derive a property from user-defined (classified) training data. Categories such as negative, ambiguous or positive are assigned depending on the form of the amplification curve. An extensive literature research showed that there are no openly accessible classified amplification curve data sets. Open Data is meant in the sense that data are freely available, free of charge, free to use and that data can be republished, without restrictions from copyright, patents or other mechanisms of control (Kitchin 2014).

Therefore, a large number of records with amplification curves and their classification (negative, ambiguous, positive) were added to the PCRedux package.

For the amplification curves in Table 5, a dichotomous classification was performed (roughly sigmoid or negative amplification reaction with a flat curve shape). Consequently, this does not rule out

- if an unspecific amplification product has been synthesized,
- if a contamination has been amplified or
- if only primer-dimers have been amplified.

To answer this question, other methods such as agarose gel electrophoresis need to be used.

3.5.1 Manual Amplification Curve Classification

For machine learning and method validation, it was important to classify the amplification curves individually. In Rödiger, Burdukiewicz, and Schierack (2015), the humanrater() [chipPCR] function was described. This function was developed to help the user during the classification of amplification curves and melting curves. The user has to define classes, which get assigned to an amplification curve after an expert has entered the class in input mask.

By convention, class labels were specified as e. g., negative ("n"), ambiguous ("a"), positive ("y") in the PCRedux package.

All amplification curve data sets listed in Table 5 were classified in interactive, semi-blinded sessions. humanrater() [chipPCR] was set to randomly select individual amplification curves. All data sets were manually classified at least three times. The htPCR data set (Figure 4B) was classified eight times in total (see Figure 26). Most of the amplification curves are neither unequivocal classifiable as positive or negative.

Table 5: Classified amplification curve data sets. Decision data sets in PCRedux: table with results of manual classification as comma separated values. qPCR data set: name of original amplification curve data set. Package: name of the R package containing the amplification curves. Device: is the device used to measure the amplification reaction. Note: The original data sets contain information about the detection chemistry used within the corresponding qPCR experiments. AB, Applied Biosystems.

Decision Datasets in PCRedux	qPCR Dataset	Package	Device
decision_res_RAS002.csv	RAS002.rdml	PCRedux	CFX96, Bio-Rad
$decision_res_RAS003.csv$	RAS003.rdml	PCRedux	CFX96, Bio-Rad
decision_res_batsch1.csv	batsch1	qpcR	Light Cycler 1.0, Roche
decision_res_batsch2.csv	batsch2	qpcR	Light Cycler 1.0, Roche
decision_res_batsch3.csv	batsch3	qpcR	Light Cycler 1.0, Roche
decision_res_batsch4.csv	batsch4	qpcR	Light Cycler 1.0, Roche
decision_res_batsch5.csv	batsch5	qpcR	Light Cycler 1.0, Roche
decision_res_lc96_bACTXY.csv	$lc96_bACTXY.rdml$	RDML	Light Cycler 1.0, Roche
decision_res_boggy.csv	boggy	qpcR	Light Cycler 96, Roche
$decision_res_C126EG595.csv$	C126EG595	chipPCR	Chromo4, Bio-Rad
decision_res_C127EGHP.csv	C127EGHP	chipPCR	iQ5, Bio-Rad
decision_res_C316.amp.csv	C316.amp	chipPCR	iQ5, Bio-Rad
decision_res_C317.amp.csv	C317.amp	chipPCR	iQ5, Bio-Rad
decision_res_C60.amp.csv	C60.amp	chipPCR	iQ5, Bio-Rad
decision_res_CD74.csv	CD74	chipPCR	iQ5, Bio-Rad
decision_res_competimer.csv	competimer	qpcR	Light Cycler 480, Roche
decision_res_dil4reps94.csv	dil4reps94	qpcR	CFX384, Bio-Rad
decision_res_guescini1.csv	guescini1	qpcR	Light Cycler 480, Roche
decision_res_guescini2.csv	guescini2	qpcR	Light Cycler 480, Roche
decision_res_htPCR.csv	htPCR	qpcR	Biomark HD, Fluidigm
decision_HCU32_aggR.csv	$HCU32_aggR.csv$	PCRedux	VideoScan
decision_res_karlen1.csv	karlen1	qpcR	ABI Prism 7700, AB
decision_res_karlen2.csv	karlen2	qpcR	ABI Prism 7700, AB
decision_res_karlen3.csv	karlen3	qpcR	ABI Prism 7700, AB
decision_res_lievens1.csv	lievens1	qpcR	ABI7300, ABI
decision_res_lievens2.csv	lievens2	qpcR	ABI7300, ABI
decision_res_lievens3.csv	lievens3	qpcR	ABI7300, ABI
decision_res_reps.csv	reps	qpcR	MXPro3000P, Stratagene
decision_res_reps2.csv	reps2	qpcR	MXPro3000P, Stratagene
decision res reps3.csv	reps3	qpcR	MXPro3000P, Stratagene
decision_res_reps384.csv	reps384	qpcR	CFX384, Bio-Rad
decision_res_rutledge.csv	rutledge	qpcR	Opticon 2, MJ Research
decision_res_stepone_std.csv	stepone_std	RDML	StepOne, AB
	_	qpcR	Light Cycler 1.0, Roche
decision res testdat.csv	testdat	9020	
decision_res_testdat.csv decision_res_vermeulen1.csv	vermeulen1	_	Light Cycler 480, Roche
		qpcR	Light Cycler 480, Roche
decision_res_vermeulen1.csv	vermeulen1	_	· · · · · · · · · · · · · · · · · · ·

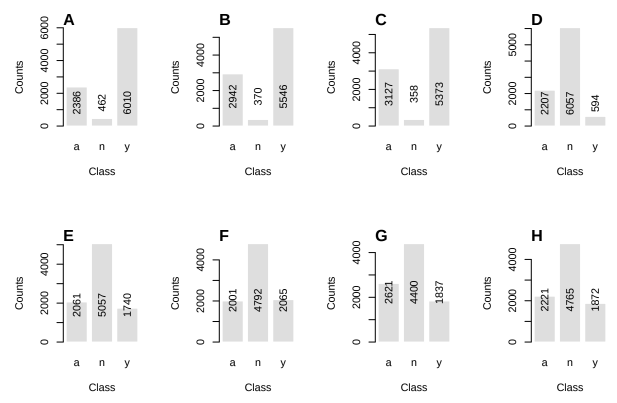


Figure 26: Variations in the classification of amplification curves. A prerequisite for the development of machine-learning models is the availability of manually classified amplification curves. Amplification curves (n = 8858) from the 'htPCR' data set have been classified by one user eight times at different points over time (classes: ambiguous (a), positive (y) or negative (n)). During this process, the amplification curves were presented in random order. The example shows that different (subjective) class mappings may occur for the same data set. While only a few amplification curves were classified as negative in the first three classification cycles (A-C), their proportion increased almost tenfold in later classification cycles (D-H).

```
# Suppress messages and load the packages for reading the data of the classified
# amplification curves.
library(PCRedux)
# Load the decision_res_htPCR.csv data set from a csv file.
filename <- system.file("decision_res_htPCR.csv", package = "PCRedux")
decision_res_htPCR <- read.csv(filename)</pre>
# Create graphic device for the plot(s)
par(mfrow = c(2, 4))
for (i in 2L:9) {
    data_tmp <- table(as.factor(decision_res_htPCR[, i]))</pre>
    barplot(data_tmp, col = adjustcolor("grey", alpha.f = 0.5),
            xlab = "Class", ylab = "Counts", border = "white")
    text(c(0.7, 1.9, 3.1), rep(quantile(data_tmp, 0.25), 3),
         data_tmp, srt = 90)
    mtext(LETTERS[i - 1], cex = 1, side = 3, adj = 0, font = 2)
}
```

This approach is well suited and has been applied to classify a variety of amplification curves during the

development of the PCRedux package. From experience this is time-consuming and tiring for large data sets, especially when the amplification curves are similar in shape. A high similarity between amplification curves exists, for example, in replicates and negative controls.

3.5.2 tReem() - A Function for Shape-based Group-wise Classification of Amplification Curves

The similarity of amplification curves can be used to form groups of similar shapes. The amplification curves in the groups can then be classified in bulk. In this way, a higher throughput can be achieved, a concept yet not described for the analysis of qPCR data in the literature.

The tReem() function was developed to perform a *shape-based group classification*. To use the tReem() function, the first column must contain the qPCR cycles and all subsequent columns must contain the amplification curves. Two measures of similarity are used within the tReem() function.

- In the first measure (default), the Pearson correlation coefficients (r) are determined in pairs for all combinations of the amplification curves. The correlation coefficient is a statistical measure to describe the strength of the correlation between two or more variables. The correlation coefficient r is regarded as distance between the amplification curves. r is a dimensionless value and only takes values between -1 and 1. If r = -1, there is a maximum reciprocal relationship. If r = 0 there is no correlation between the two variables. If r = 1, there is a maximum rectified correlation.
- In the second measure, the *Hausdorff distance* is used to determine the similarity between amplification curves. The Hausdorff distance is "the maximum of the distances from a point in any of the sets to the nearest point in the other set" (Rote 1991; Herrera et al. 2016). The amplification curves are converted within the tReem() function using the qPCR2data() function.

Both methods process the distances in the same steps. This involves the calculation of the distance matrix using the Euclidean distances of all distance measures to determine the distance between the lines of the data matrix.

This is used to perform a hierarchical cluster analysis. In the last step, the cluster is divided into groups based on a user-defined k value. For example, two groups are created for k = 2. If the amplification curves shapes are highly diverse, a larger k should be used. After a chain of processing steps presents the tReem() function a series of plots with grouped of amplification curves. The corresponding classes can then be assigned to the groups of amplification curves by the user using an input mask.

Grouping the amplification curves with the Pearson correlation coefficient as a distance measure is usually faster than the Hausdorff distance. The Hausdorff distance is an approximation of a shape metrics to define similarity measures between shapes. (Charpiat, Faugeras, and Keriven 2003).

```
# Classify amplification curve data by correlation coefficients (r)
data <- qpcR::testdat

classification_result <- tReem(data[, 1:15], k = 3)
classification_result</pre>
```

3.5.3 decision_modus() - A Function to Get a Decision (Modus) from a Vector of Classes

For the systematic statistical analysis of classification data sets, the decision_modus() function has been developed. This allows the most common decision (mode) to be determined. The mode is useful to consolidate large collections of different decisions into a single (most frequent) decision.

Observed: a, a, a, a, a, n, n, n \rightarrow frequencies 5 x a, 3 x n \rightarrow mode: a. Since the class names are known, they only have to be interpreted by the user (e. g., "a",n,"y" -> "ambivalent", "negative", "positive").

A manual classification was performed out for the htPCR data set (for an example plot Figure 4B) with the humanrater() function. The classification of each amplification curve was performed eight times at different time points since many of the amplification curves did not resemble optimal curvatures (e. g., Figure 1). It is

likely that the amplification curve (P06.W47, Figure 1) is considered as ambiguous or even positive (positive \leftrightarrow ambivalent) by the users.

Table 6 shows from a total of 8858 amplification curves the first 25 lines classified as negative (conformity=TRUE) and the first 25 lines classified as positive. Since in total, the curves were classified eight times (test.result.1 ... test.result.8) a whole of 70864 amplification curves was analysed. In this classification experiment the amplification curves have been classified differently in 94.5% of the cases (e. g., line 1 "P01. W01").

Table 6: Results of the 'htPCR' data set classification. All amplification curves of the 'htPCR' data set were classified as 'negative', 'ambiguous' and 'positive' by individuals in eight analysis cycles ('test.result.1' . . . 'test.result.8'). If an amplification curve was always classified with the same class, the last column ('conformity') shows 'TRUE'. As an example, the table shows 25 amplification curves with consistent classes and 25 amplification curves with differing classes ('conformity = FALSE').

htPCR	test.result.1	test.result.2	test.result.3	test.result.4	test.result.5	test.result.6	test.result.7	test.result.8	conformity
P01.W01	у	a	a	n	n	n	a	n	FALSE
P01.W02	a	y	a	n	n	n	n	n	FALSE
P01.W03	У	a	a	n	n	n	n	n	FALSE
P01.W04	У	a	a	n	n	n	n	n	FALSE
P01.W05	У	a	a	n	n	n	a	n	FALSE
P01.W06	a	a	a	n	n	n	n	n	FALSE
P01.W07	a	a	a	n	n	n	n	n	FALSE
P01.W08	У	a	a	n	n	n	n	n	FALSE
P01.W09	у	a	a	n	n	n	n	n	FALSE
P01.W10	n	a	a	n	n	n	n	n	FALSE
P01.W11	У	У	a	у	у	у	a	У	FALSE
P01.W12	a	a	a	n	n	n	n	n	FALSE
P01.W13	У	a	у	n	n	n	n	n	FALSE
P01.W14	y	a	y	n	n	n	n	n	FALSE
P01.W15	У	a	v	n	n	n	n	n	FALSE
P01.W16	У	a	a	n	n	n	n	n	FALSE
P01.W17	y	a	a	n	n	n	n	n	FALSE
P01.W18	a	a	a	n	n	n	n	n	FALSE
P01.W19	у	a	a	n	n	n	a	n	FALSE
P01.W19	y	a	a	У	a	a	у	a	FALSE
P01.W21	a.	a	a	n n	n	n	n n	n	FALSE
P01.W21		a	a	n n	n n	n n	n n	n n	FALSE
P01.W23	У								FALSE
P01.W23	У	a	a	n	n	n	n	n	FALSE
	У	a	a	n	n	n	a	n	
P01.W25	У	a	a	n	n	n	n	n	FALSE
P01.W58	n	n	n	n	n	n	n	n	TRUE
P02.W09	У	У	У	У	У	У	У	У	TRUE
P02.W19	У	У	У	У	У	У	У	У	TRUE
P02.W31	У	У	У	У	У	У	У	У	TRUE
P02.W41	У	У	У	У	У	У	У	У	TRUE
P02.W72	У	У	У	У	У	У	У	У	TRUE
P02.W81	У	У	У	У	У	У	У	У	TRUE
P02.W84	n	n	n	n	n	n	n	n	TRUE
P03.W09	У	У	У	У	У	У	У	У	TRUE
P03.W21	y	У	У	y	У	У	У	У	TRUE
P03.W22	y	y	У	У	У	У	y	y	TRUE
P03.W31	У	y	y	y	y	y	У	y	TRUE
P03.W32	y	У	у	у	у	у	у	У	TRUE
P03.W39	y	У	У	У	У	У	У	У	TRUE
P03.W56	y	y	у	У	у	у	y	y	TRUE
P03.W59	У	y	У	У	у	у	У	У	TRUE
P03.W68	у	y	y	y	y	y	y	y	TRUE
P03.W72	У	У	У	y	y	y	у	y	TRUE
P03.W73	у	y	y	y	y	y	y	y	TRUE
P04.W19	у	y	y	y	y	y	y	y	TRUE
P04.W31	У	y	y	y	y	y	y	y	TRUE
P04.W66	y	y	y	y	y	y	y	y	TRUE
P04.W81	У	y	y	y	y	y	y	y	TRUE
P04.W91	У	y	y	y	y	y	y	y	TRUE
P05.W20	v	v	v	v	v	v	v	v	TRUE

The decision_modus() function was applied to the record decision_res_htPCR.csv with all classification rounds (columns 2 to 9) and the mode was determined for each amplitude curve Figure 27.

```
# Use decision_modus() to go through each row of all classifications done by
# a human.

# Determine the number of observations where all classifications were
# the same (conformity == TRUE).
conformity <- decision_res_htPCR[["conformity"]]

# List all classifications.
dec <- lapply(1L:nrow(decision_res_htPCR), function(i) {
    decision_modus(decision_res_htPCR[i, 2:9])[1]
}) %>% unlist()
```

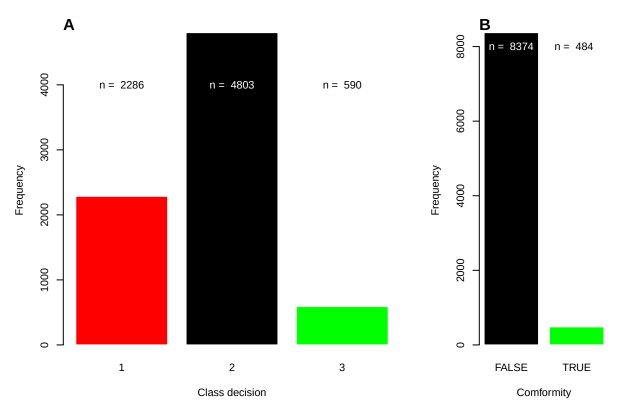


Figure 27: Frequency of amplification curve classes and conformity in the 'htPCR' data set. The 'htPCR' data set was classified by hand eight times. Due to the unusual amplification curve shape and input errors during classification, many amplification curves were classified differently. A) Frequency of negative (black), ambiguous (red) and positive (green) amplification curves in the 'htPCR' data set. The combined number of ambiguous and negative amplification curves appears to be higher, than the number of positive amplification curves. B) The number of observations where all classification cycles made the same decision (conformity == TRUE) accounts for only 5% of the total number of observations. TRUE, all classes of the amplification curve matched. FALSE, at least one in eight observations had a different class.

```
\# Show statistic of the decisions
summary(dec)
##
      Min. 1st Qu.
                     Median
                                Mean 3rd Qu.
                                                  Max.
                                                          NA's
##
     1.000
              1.000
                      2.000
                               1.779
                                        2.000
                                                 3.000
                                                          1179
```

Another usage mode of decision_modus() is to set the parameter as max_freq=FALSE. This option specifies the number of all classifications.

The amplification curve P01. W06 was classified as a=3 times and as n=5 times. Therefore, the decision would turn negative.

3.5.4 PCRedux-app

PCRedux-app is a web server, based on the shiny technology (Chang et al. 2019) wrapped around the encu() function (subsubsection 3.3.2). An user can upload qPCR data and download obtained amplification curve features.

There are different ways to use the function.

- Through RScript (Scripting Front-End for R):
 - Enter the command Rscript -e 'PCRedux::run_PCRedux()' in a console and copy the pasted URL in a browser.
- Through Graphical User Interfaces:
 - The function can be started directly in RStudio or RKWard by:

run the Shiny app
PCRedux::run_PCRedux()

Table 7: Measures for performance analysis for binary classification. TP, true positive; FP, false positive; TN, true negative; FN, false negative

Measure	Formula
Sensitivity - TPR, true positive rate	$TPR = \frac{TP}{TP + FN}$
Specificity - SPC, true negative rate	$SPC = \frac{TN}{TN + FP}$
Precision - PPV, positive predictive value	$PPV = \frac{TP}{TP + FP}$
Negative predictive value - NPV	$NPV = \frac{TN}{TN + FN}$
Fall-out, FPR, false positive rate	$PPV = \frac{TP}{TP}$ $NPV = \frac{TN}{TN + FN}$ $FPR = \frac{FP}{FP \pm TN} = 1 - SPC$
False negative rate - FNR	$FNR = \frac{F'N}{TN + FN} = 1 - TPR$
False discovery rate - FDR	$FDR = \frac{TN + TN}{TP + FP} = 1 - PPV$
Accuracy - ACC	$ACC = \frac{(TP+TN)}{(TP+FP+FN+TN)}$
F1 score - F1	$F1 = \frac{2TP}{(2TP+FP+FN)}$
Matthews correlation coefficient - MCC	$F1 = \frac{2TP}{(2TP+FP+FN)}$ $MCC = \frac{(TP*TN-FP*FN)}{\sqrt{(TP+FP)*(TP+FN)*(TN+FP)*(TN+FN)}}$
Likelihood ratio positive - LRp	$LRp = \frac{TPR}{1 - SPC}$
Cohen's kappa (binary classification)	$\kappa = \frac{p_0 - p_c}{1 - p_0}$

3.6 Helper Functions of the PCRedux Package

3.6.1 performer() - Performance Analysis for Binary Classification

Statistical modeling and machine learning are powerful but expose a risk to the user by introducing an unexpected bias. This may lead to an overestimation of the performance. The assessment of the performance by sensitivity and specificity is fundamental to characterize a classifier or screening test (G. James et al. 2013). Sensitivity is the percentage of true decisions that are identified and specificity is the percentage of negative decisions that are correctly identified (Table 7). An example for the application of the performer() function is shown in subsubsection 3.3.13.

3.6.2 qPCR2fdata() - A Helper Function to Convert Amplification Curve Data to the fdata Format

qPCR2fdata() is a helper function to convert amplification curve data to the functional fdata class (Febrero-Bande and Oviedo de la Fuente 2012). The fdata format is used for functional data analysis to determine the similarity measures between amplification curves shapes by the Hausdorff distance. Similarity herein refers to the difference in spatial location of two *objects* (e. g., amplification curves). Objects with a close distance are presumably more similar. For single objects (e. g., points) one can use a vector distance, such as the Euclidean distance (Herrera et al. 2016).

The qPCR2fdata() function takes a data.frame containing the amplification cycles (first column) and the fluorescence amplitudes (subsequent columns) as input.

Noise and missing values may affect the analysis adversely. Therefore, an instance of the CPP() [chipPCR] function (Rödiger, Burdukiewicz, and Schierack 2015) was integrated in qPCR2fdata(). If preprocess=TRUE in qPCR2fdata(), then all curves are smoothed (Savitzky-Golay smoother), missing values are imputed and outliers in the ground phase get removed as described in Rödiger, Burdukiewicz, and Schierack (2015).

The following example illustrates a hierarchical cluster analysis of the testdat data set. The amplification curves of the testdat data set remained as raw data or were preprocessed (smoothed). Subsequent, the amplification curves were converted by the qPCR2fdata(). The converted data were subjected to cluster analysis (Hausdorff distance). This method uses the elements of a proximity matrix to generate a dendrogram. The dendrogram can then be used to further analyze the clusters. There are methods to determine the number of clusters automatically k (Cook and Swayne 2007). However, for simplicity, the number of clusters was determined visually.

```
# Calculate the Hausdorff distance of the amplification curves
# cluster the curves.
# Load additional packages for data and pipes.
library(fda.usc)

data <- qpcR::testdat

# Convert the qPCR data set to the fdata format
# Use unprocessed data from the testdat data set
res_fdata <- qPCR2fdata(data)

# Extract column names and create rainbow color to label the data
columnames <- data[-1] %>% colnames(.)
colors <- rainbow(length(columnames), alpha = 0.5)

# Calculate the Hausdorff distance (fda.usc) package and plot the distances
# as clustered data.

res_fdata_hclust <- metric.hausdorff(res_fdata)
res_hclust <- hclust(as.dist(res_fdata_hclust))</pre>
```

The distance based on the Hausdorff metric was already calculated in the next steps involving the cutree() [stats] function to split the dendrogram into smaller junks. A priori it was defined that two classes (positive & negative) are expected. Therefore, the group parameter was set to k=2 in the cutree().

```
# Cluster of the unprocessed amplification curves
res_cutree <- cutree(res_hclust, k = 2)
res_cutree <- factor(res_cutree)
levels(res_cutree) <- list(y = "1", n = "2")</pre>
```

The dendrogram shows that

- the observations are correctly assigned to a cluster of positive or negative amplification curves and that
- the shift of the Cq (late increase of the fluorescence) is reflected in the positive cluster (Figure 28).

This workflow can be used to cluster amplification curve data according to similar shape. Classification tasks can be preformed in batches of amplification curves. The calculation of the distances is a computation expensive step dependent on the number of amplification curves.

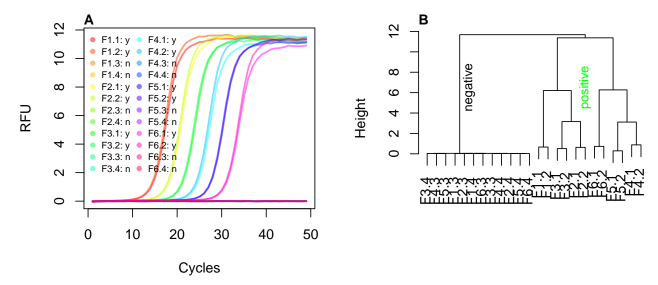


Figure 28: Shape-based clustering of amplification curves. A) The clustering of the amplification curves of the 'testdat' data set (A) was based on the Hausdorff distance. B) The amplification curves were converted with the qPCR2fdata() function, and the Hausdorff distance of the curves was determined by cluster analysis. There were no errors in distinguishing between negative (n) and positive (y) amplification curves.

The following example illustrates the usage for the HCU32_aggR.csv data set from the VideoScan platform with 32 heating and cooling units (equivalent of 32 PCR vessels). In this experiment, the bacterial gene aggR from $E.\ coli$ was amplified in 32 replicate qPCR reactions. Details of the experiment are described in the manual of the PCRedux package. The ambition was to test if the 32 amplification curves of the qPCR reaction are identical. As before, the data were processed with the qPCR2fdata() function and compared by the Hausdorff distance. Ideally, the amplification curves form only few clusters.

```
# Calculate slope and intercept on positive amplification curve data from the
# VideoScan 32 cavity real-time PCR device.
# Use the fda.usc package for functional data analysis
library(fda.usc)

# Load the qPCR data from the HCU32_aggR.csv data set
# Convert the qPCR data set to the fdata format

filename <- system.file("HCU32_aggR.csv", package = "PCRedux")
data_32HCU <- read.csv(filename)

res_fdata <- qPCR2fdata(data_32HCU)
# Extract column names and create rainbow color to label the data
columnames <- data_32HCU[-1] %>% colnames(.)
colors <- rainbow(length(columnames), alpha = 0.55)</pre>
```

In advance the Cq values were calculated by the following code:

```
# Load the qpcR package to calculate the Cq values by the second derivative
# maximum method.
library(qpcR)

res_Cq <- sapply(2L:ncol(data_32HCU), function(i) {
    efficiency(pcrfit(data_32HCU, cyc = 1, fluo = i, model = 16))
})</pre>
```

```
data.frame(
    obs = colnames(data_32HCU)[-1],
           Cq = unlist(res_Cq["cpD2", ]), eff = unlist(res_Cq["eff", ])
)
#
         Results
#
# obs
         Cq
                 eff
# 1
         A1 14.89 1.092963
# 2
         B1 15.68 1.110480
# 3
         C1 15.63 1.111474
# ...
# 30
         F4 15.71 1.109634
# 31
         G4 15.70 1.110373
# 32
         H4 15.73 1.117827
```

Next, the amplification curves (Figure 29A), the differences between base-line region and plateau region (Figure 29B), the correlation between the Cq value and amplification efficiency (Figure 29C) and the clusters based on the Hausdorff distance were taken into account.

Some amplification curves (Figure 29A) had stronger noise, and all curves exhibited a negative non-linear trend and shift in the ground phase. The comparison of the ground phase and the plateau phase showed a difference between the 32 amplification curves. The observations E1, F1 and H1 were most close in the ground and plateau phase. The comparison of Cq values and amplification efficiency showed that most amplification curves are similar. However, there are also amplification curves that show a greater deviation from the median of all Cq values (Figure 29C). The cluster analysis confirmed the shape similarity (Figure 29D).

```
# Use the fda.usc package for functional data analysis
library(fda.usc)
# To save computing time, the Cq values and amplification efficiencies were
# calculated beforehand and transferred as a hard copy here.
calculated_Cqs <- c(</pre>
    14.89, 15.68, 15.63, 15.5, 15.54, 15.37, 15.78, 15.24, 15.94,
    15.88, 15.91, 15.77, 15.78, 15.74, 15.84, 15.78, 15.64, 15.61,
    15.66, 15.63, 15.77, 15.71, 15.7, 15.79, 15.8, 15.72, 15.7, 15.82,
    15.62, 15.71, 15.7, 15.73
)
calculated_effs <- c(</pre>
    1.09296326515231, 1.11047987547324, 1.11147389307153, 1.10308929700635,
    1.10012176315852, 1.09136717687619, 1.11871308210321, 1.08006168654712,
   1.09500422011318, 1.1078777171126, 1.11269436700649, 1.10628580163733,
   1.1082009954558, 1.11069683827291, 1.11074914659374, 1.10722949813473,
   1.10754282514113, 1.10098387264025, 1.1107026749644, 1.11599641663658,
    1.11388510347017, 1.11398547396991, 1.09410798249025, 1.12422338092929,
    1.11977386646464, 1.11212436173214, 1.12145338871426, 1.12180879952503,
    1.1080276005651, 1.10963449004393, 1.11037302758388, 1.11782689816295
)
# Plot the converted qPCR data
# Create graphic device for the plot(s)
layout(matrix(c(1, 2, 3, 4, 4, 4), 2, 3, byrow = TRUE))
```

```
plot(res_fdata, xlab = "Cycles", ylab = "RFU", main = "HCU32_aggR", type = "1",
         lty = 1, lwd = 2, col = colors)
legend("topleft", as.character(columnames), pch = 19, col = colors,
       bty = "n", ncol = 4)
mtext("A", cex = 1, side = 3, adj = 0, font = 2)
# Plot the background and plateau phase.
boxplot(
    data_32HCU[, -1] - apply(data_32HCU[, -1], 2, min),
        col = colors, las = 2, main = "Signal to noise ratio",
        xlab = "Sample", ylab = "RFU"
mtext("B", cex = 1, side = 3, adj = 0, font = 2)
# Plot the Cqs and the amplification efficiencies.
# Determine the median of the Cq values and label all Cqs, which a less 0.1 Cqs
# of the median or more then 0.1 Cqs of the median Cq.
plot(
    calculated_Cqs, calculated_effs, xlab = "Cq (SDM)",
     ylab = "eff", main = "Cq vs. Amplification Efficiency",
     type = "p", pch = 19, lty = 1, lwd = 2, col = colors
)
median_Cq <- median(calculated_Cqs)</pre>
abline(v = median_Cq)
text(median_Cq + 0.01, 1.085, expression(paste(tilde(x))))
labeled <- c(
    which(calculated_Cqs < median_Cq - 0.1),</pre>
             which(calculated_Cqs > median_Cq + 0.1)
)
text(
    calculated Cqs[labeled], calculated effs[labeled],
     as.character(columnames)[labeled]
mtext("C", cex = 1, side = 3, adj = 0, font = 2)
# Calculate the Hausdorff distance using the fda.usc package and cluster the
# the distances.
res_fdata_hclust <- metric.hausdorff(res_fdata)</pre>
cluster <- hclust(as.dist(res_fdata_hclust))</pre>
# plot the distances as clustered data and label the leafs with the Cq values
# and colored dots.
plot(cluster, main = "Clusters of the amplification\n
curves as calculated by the Hausdorff distance", xlab = "", sub="")
```

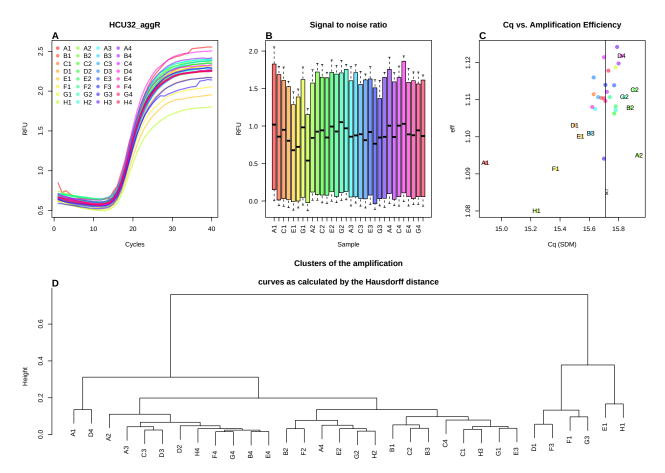


Figure 29: Clustering and variation analysis of amplification curves. The amplification curves of the 32HCU were processed with the qPCR2fdata() function and then processed by cluster analysis (Hausdorff distance). A) Amplification curves were plotted from the raw data. B) Overall, signal-to-noise ratios of the amplification curves between all cavities were similar. C) The Cq values and amplification efficiency were calculated using the efficiency(pcrfit()) [qpcR] function. The median Cq is shown as a vertical line. Cqs greater or less than 0. 1 of Cq \tilde{x} are marked with observation labels. D) The cluster analysis showed no specific pattern with respect to the amplification curve signals. It appears that the observations D1, E1, F1, F3, G3 and H1 differ most from the other amplification curves.

The analysis gives an overview on the variation of the amplification curve data.

4 Summary and Conclusions

Extensive amounts of data represent a serious challenge in the analysis of qPCR amplification curves. In a manual classification (e. g. negative, positive), the result is usually characterized by the subjective perception of the experimenter. In addition, the time required for a manual analysis is high. An automatic system for amplification curve analysis might objectify and generalize the decision process. Interestingly, so far most authors focused on the extraction of single predictors from amplification curves. These are mainly the Cq and the amplification efficiency, which are used for the downstream processing such as expression analysis or genotyping (Pabinger et al. 2014).

Numerous software tools were developed, which deal with theses analytical steps. For example Baebler et al. (2017) published quantGenius and Mallona et al. (2017) published Chainy. However, none of them attempts to make use of characteristics of the amplification curve. Positive amplification curves usually exhibit a sigmoid shape, consisting of a ground phase, exponential phase, and plateau phase. Negative amplification curves resemble flat noisy signal. As a result, the experienced user is usually able to correctly interpret the curve shape. Similarly, outliers and measurement errors can also be identified. When setting up a qPCR assay, manual data analysis is a useful and necessary approach to familiarize oneself with the properties of the qPCR amplification curves.

It is challenging to analyze and classify amplification curves if they deviate substantially from the sigmoid shape or if their number is no longer feasible for manual analysis. Furthermore, the supposed objectivity of the user must also be questioned. In the scientific environment there is often the temptation - or rather the compulsion - to use all data for publications. As a result, amplification curves of rather poor quality are often provided. In Figure 27 it was demonstrated that a reproducible and objective analysis of amplification curves is not always given.

For a novice user, the quality of an amplification curve can be acceptable, for an experienced user, not. Ambiguous amplification curves are a big challenge for the user, as both classes (positive and negative) can be true. In most cases, however, the user is interested in an automatic distinction between positive and negative samples. This is important for screening applications. In subsubsection 3.2.2, further reasons were elaborated on why statistical methods are necessary for the objective and reproducible interpretation of the amplification curves.

A few scientific approaches have previously been shown in which machine learning was used for the analysis of amplification curves. The intention of Gunay, Goceri, and Balasubramaniyan (2016) was to improve the determination of Cq values, without dealing with classification. The authors postulated that they had developed an improved prediction of Cq values using a modified three-parameter model. One assumption of their approach was that their modified three-parameter model could be applied to any amplification curve. However, there are reasons why such an assumption is not valid. In chapters subsubsection 3.2.3 and Figure 27 it was described that a considerable proportion of amplification curves deviate clearly from a three-parameter model. Multiparametric models with more than four parameters are more frequently adapted to amplification curves. In addition, the multiparametric models tend to adapt to noise (Figure 2). Unsurprisingly, a Cq value is calculated for actually negative amplification curves, demonstrating that a three-parameter model alone cannot provide reliable predictions. However, a correct model is important for the extraction of Cq values, for the determination of predictors from the curves and consequently for classification.

In addition to the determination of quantification points, the classification of amplification curves is necessary. For example, a diagnostician is interested in whether a sample is positive or negative. In research using high-throughput screening methods, it is important to classify large data sets quickly and cost-effectively. It is important to bear in mind that in manual classification, the classification result is influenced by the subjective perception of the experimenter and that it is comparatively time-consuming.

Here, an automatic computer-assisted classification of amplification curves is feasible because it renders the entire analysis process faster, more objective and more reproducible. The objectives were therefore to

- 1. create a collection of classified amplification curve data,
- 2. propose algorithms that can be used to calculate predictors from amplification curves,

- 3. to develop pipelines (e. g., machine learning, decision trees) that can be used for automatic classification of amplification curves,
- 4. evaluate pipelines that can be used for an automatic classification of amplification curves based on the curve shape and
- 5. to bundle and distribute the findings in a public repository open source software and open data package with an open data.class

for an automatic analysis of amplification curve data by machine learning.

For this purpose, the PCRedux package was developed. This package contains proof-of-concept algorithms and functions with which predictors (mathematically describable properties) of amplification curves can be calculated. The pcrfit_single() function is an extensible wrapper function for all algorithms and functions developed. PCRedux version 1.0.2 offers concepts to calculate 57 predictors from an amplification curve. In addition, predictors such as the employed chemistry, the qPCR device and further experimental details can be added by the user. Other parameters (e. g. hydrolysis probe, DNA binding dye) can be converted into binary classifiers and be used for modeling. Machine learning requires predictors to train a model (Saeys, Inza, and Larranaga 2007). The model should then be able to put new unknown data into a meaningful context. Predictors can have a significant influence on the accuracy of the prediction model. The predictors of the PCRedux package are a novelty in the literature for the classification of amplification curves, so that the collection probably constitutes the most extensive one at the time of the first release on https://github.com/devSJR/PCRedux (summer 2017).

It can be assumed that not all predictors are suitable or necessary for machine learning. Some **potential** predictors are more likely to be suitable for validation of data integrity (quality management) and data mining. The maxRFU predictors and sigma_bg (Figure 16) are to be mentioned here as examples. The predictor maxRFU should ideally be at a value of 1. If this is not the case, it can be assumed that a preprocessing problem was encountered. The sigma_bg value describes the standard deviation of the ground phase. It should be low ($sigma_bg \leq 0.1$), otherwise it can be assumed that unusually high variations of the intensity values are present. Consequently, the relevance of the predictors must be determined independently by each user on the basis of domain knowledge, the objective, and the given data set.

At this point it should be mentioned that these approaches were deliberately sought and developed to make the approach and implementation more understandable. Self-learning machine-learning methods are expected to be included in the PCRedux package in future releases, from the necessity to design and test additional predictors.

To a modest extent, the usefulness of the predictors was tested on exemplary data sets, with the aim of achieving a high degree of objectivity and reproducibility. It is probably not possible to fulfill this ideal completely, since the algorithms were designed from a limited human perspective, which poses a generic problem in machine learning. In particular, data records can be distorted if the user excludes seemingly problematic data. Hence, it is advisable to perform a more comprehensive analysis of all predictors.

Several ROIs (see Figure 5) can be obtained from amplification curves. Sigmoidal amplification curves have turning points that can serve as an indicator of a positive amplification curve, and are mathematical and statistical starting points for calculating predictors. Amplification curves can have unique trajectories and often deviate drastically from ideal sigmoid models (see Figure 4A and Figure 4B). Some amplification curves have only a slight increase with positive or negative signs lacking a sigmoid curvature.

Almost all real-time thermo-cyclers have built-in software that performs preprocessing steps such as smoothing, baseline correction and normalization on the amplification curves (Rödiger, Burdukiewicz, and Schierack 2015; Spiess et al. 2015, 2016). For this reason, it is nearly impossible to get access to informative raw data, so that the impact on the predictor extraction process cannot be adequately estimated.

The volume and classification of data sets needs to be representative (Herrera et al. 2016). The PCRedux package contains a large number of manually classified amplification curves. Here, data preparation is an important step, as it encompasses data cleansing, data transformation and data integration (Herrera et al. 2016). To assist, the xray package (Seibelt 2017) and assertr package (Fischetti 2019) can be used to analyze the distribution and variables in records for important anomalies such as missing values, zeros,

empty strings (Blank) and infinite numbers (Inf). Users of the PCRedux package should use such tools before proceeding with the analysis. Even though the data records (qPCR runs) in the PCRedux package have a very similar structure, some records contain missing values or have different dimensions. For example, the data set C127EGHP comprises a matrix of 40 x 66 (35 cycles x observations (65 amplification curves)), while the data set htPCR comprises a matrix of 35 x 8859.

Talking about data sets, it is important to make sure that the volume of the data is representative. In this study amplification curves were categorized by hand and processed with the algorithms described in this study. Although this data set is fairly large in comparison to what existed before, there is no numeric evidence how well it reflects amplification curves in general. In particular, not all data sets have comparable case numbers. For example, the htPCR data set (Biomark HD, Fluidigm) encompass in total 8858 amplification curves, while the C127EGHP data set (iQ5, Bio-Rad) encompass in total 64 amplification curves. While most of the amplification curves of the C127EGHP data set have a classical sigmoid curve shape, amplification curves from the htPCR data set largely display noisy curvatures with non-sigmoid shape. One may question if the larger data set introduces a confirmation bias.

Although data sets in the PCRedux package are large compared to what has been previously available, there is no conclusive evidence of how well these represent amplification curves in other settings. Bellman coined the term "Curse of Dimensionality" in 1961 when he dealt with adaptive control processes, vaguely describing the practical difficulties of high-dimensional analysis and estimation. There is only a maximum number of predictors for a sample of a certain size. If there are too many, the performance of an algorithm decreases rather than improves. As a result, many data mining algorithms fail with high dimensionality because the data points are sparse (Herrera et al. 2016). In addition, the following applies:

- The user's knowledge, prejudices, and skills are reflected in the classified data. For example, an amplification can be classified as "ambiguous" by one and "positive" by another user.
- Not all data sets have a comparable number of cases as described above. The most of the amplification curves of the C127EGHP data set have an *ideal* sigmoid waveform. In contrast, the amplification curves from the htPCR data set are noisy and difficult to classify.
- The user decides
 - how the data is preprocessed,
 - which predictors are determined by the amplification curves,
 - which data set is used for machine learning and to what extent,
 - how the models are tested and
 - which results are reported.

Accordingly, the domain knowledge, biases and competences of the human operator are reflected by the software. For example, amplification that are classified by one human as 'ambiguous' might be classified as 'positive' by another human operator. This will affect (bias) the characteristics of the training data set. The model is intended to be in accordance with the human operator. The implications can be problematic. For example, amplification curves rated as false negative might lead to an adverse evidence in a forensic setting. As illustrated in subsection 3.2, every human operator will make an association between the shape of an amplification curve and the class it belongs to.

Measures to minimize errors in manual classification include the implementation of algorithms to detect them. This should preferably be done in the form of open source packages with publicly accessible data sets. In contrast to black box algorithms and hidden data sets, third parties can review and modify all elements. Many qPCR devices have built-in software that performs preprocessing steps like smoothing, base-lining and normalization and the data sets (Rödiger, Burdukiewicz, and Schierack 2015; Spiess et al. 2015, 2016). This will have an impact on the predictor extraction process. Same applies to the preprocessing steps in the PCRedux package, which have not been studied thoroughly.

For the examples in the PCRedux package, the question could not be clarified whether class imbalances in the data sets cause a confirmation distortion. However, a class imbalance may result in loss of prediction strength, because some classifiers make the assumption of similar class distributions (Herrera et al. 2016).

A new concept for the fast group-wise classification of amplification curves was introduced within the PCRedux

package. Based on experience with the collected data sets it can be stated that only a few iterations are necessary to classify a large data set. This part of the software is intended as an assistance tool for the users of the package.

Measures to minimize these sources of error are based on the implementation of algorithms in the form of an open source package, and the publication of the data sets. Unlike black box algorithms and inaccessible data sets, every user can check and correct all steps. However, it is advised that users of the PCRedux package verify independently whether their models are objective. Failure to do so may have severe implications, for instance, amplification curves classified as false negative may lead to misleading inferences in a forensic analysis.

Machine-learning algorithms require careful data preprocessing and quality management. In a first step, relatively large data sets of known characteristic vectors have to be collected and depending on the machine learning approach, predictors calculated. In a second step, these characteristics are used to classify unknown characteristic vectors using the automatic learning algorithm. As an example, the amplification curves would need to be randomly split into training data and test data. Some examples were used in the PCRedux package to show how predictors of an amplification curve data set can be calculated and used for classifications.

The scope of the PCRedux package is wide. The quantitation of nucleic acids by curve parameters such as the Cq and amplification efficiency is meaningful only if the amplification curves have a sigmoid shape (Ruijter et al. 2013, 2014; Ritz and Spiess 2008), which in principle, can now be verified with the PCRedux package.

In this study, it was shown how a given data set of amplification curves with known classifications can be used to build a system that can predict the classification of amplification curves. The algorithms provide means for a sensitive and specific classification of amplification curves from qPCR experiments in both supervised an unsupervised analysis mode. However, this method might be applicable to melting analysis too. This needs to be investigated in further studies.

Importantly, the concepts elucidated in this work may also be applied to other bioanalytical methods (e. g. enzyme kinetics, receptor binding studies, ELISA results, biological growth curves) with sigmoidal structure, however this requires more detailed interrogation. The PCRedux software may also be coupled with other technologies like Next Generation Sequencing. qPCR is used for pretesting (DNA quality) and as a confirmation test for RNA-Seq quantification (Nassirpour et al. 2014). For this purpose, automated quality control and decision support are conceivable.

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