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SpeciesPrimer: A bioinformatics pipeline dedicated to the design of qPCR primers for the quantification of bacterial species

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Background. Quantitative real-time PCR (qPCR) is a well-established method for detecting and quantifying bacteria, and it is progressively replacing culture-based diagnostic methods in food microbiology. High-throughput qPCR using microfluidics brings further advantages by providing faster results, decreasing the costs per sample and reducing errors due to automatic distribution of samples and reactants. In order to develop a high-throughput qPCR approach for the rapid and cost-efficient quantification of microbial species in a given system (for instance, cheese), the preliminary setup of qPCR assays working efficiently under identical PCR conditions is required. Identification of target-specific nucleotide sequences and design of specific primers are the most challenging steps in this process. To date, most available tools for primer design require either laborious manual manipulation or high-performance computing systems.

Results. We developed the SpeciesPrimer pipeline for automated high-throughput screening of species-specific target regions and the design of dedicated primers. Using SpeciesPrimer specific primers were designed for four bacterial species of importance in cheese quality control, namely Enterococcus faecium, Enterococcus faecalis, Pediococcus acidilactici and Pediococcus pentosaceus. Selected primers were first evaluated in silico and subsequently in vitro using DNA from pure cultures of a variety of strains found in dairy products. Specific qPCR assays were developed and validated, satisfying the criteria of inclusivity, exclusivity and amplification efficiencies.

Conclusion. In this work, we present the SpeciesPrimer pipeline, a tool to design species-specific primers for the detection and quantification of bacterial species. We use SpeciesPrimer to design qPCR assays for four bacterial species and describe a workflow to evaluate the designed primers. SpeciesPrimer facilitates efficient primer design for species-specific quantification, paving the way for a fast and accurate quantitative investigation of microbial communities.
SpeciesPrimer: A bioinformatics pipeline dedicated to the design of qPCR primers for the quantification of bacterial species

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Abstract

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primers were designed for four bacterial species of importance in cheese quality control, namely *Enterococcus faecium*, *Enterococcus faecalis*, *Pediococcus acidilactici* and *Pediococcus pentosaceus*. Selected primers were first evaluated *in silico* and subsequently *in vitro* using DNA from pure cultures of a variety of strains found in dairy products. Specific qPCR assays were developed and validated, satisfying the criteria of inclusivity, exclusivity and amplification efficiencies.

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**Introduction**

Quantitative real-time PCR (qPCR) is a well-established method for the detection and quantification of bacteria in microbiology, for instance in the context of pathogen detection in clinical and veterinary diagnostics and food safety (Cremonesi et al. 2014; Curran et al. 2007; Garrido-Maestu et al. 2018; Ramirez et al. 2009). Culture-based diagnostic methods are progressively being replaced by qPCR due to advantages such as faster results, more specific detection, and the ability to detect sub-dominant populations (Postollec et al. 2011). High-throughput microfluidic qPCR brings further advantages including the fast generation of results, a lower cost per sample and fewer errors due to automatic distribution of samples and reactants. However, in order to work efficiently high-throughput qPCR systems use identical PCR chemistry and PCR conditions for all reactions taking place on a single chip. Therefore, existing qPCR assays are often not suitable and new primers have to be designed (Hermann-Bank et al. 2013; Ishii et al. 2013; Kleyer et al. 2017). The main challenges for the successful development of any qPCR assay are the identification of a specific target nucleotide sequence and the design of primers that bind exclusively to that target sequence. Before microbial draft genomes became widely available, the 16S rRNA gene sequence was frequently used as a target sequence. However, the regions that are targeted in the 16S rRNA gene do not provide sufficient resolution to differentiate between closely related bacterial species (Moyaert et al. 2008; Torriani et al. 2001; Wang et al. 2007). Further,
housekeeping genes such as, for instance, \textit{tuf}, \textit{recA} and \textit{pheS}, were successfully used as target sequences for a variety of bacterial species in fermented foods (Falentin et al. 2010; Masco et al. 2007; Scheirlinck et al. 2009). Today, the steadily increasing number of prokaryotic draft genomes facilitates the identification of new and unique target regions.

Various commercial and open source programs facilitate the design of specific primers for a target sequence, such as the standard tools Primer3 and Primer-BLAST (Untergasser et al. 2012; Ye et al. 2012). Primer3 predicts suitable PCR primers for an input target sequence, while Primer-BLAST combines Primer3 with a BLAST search in a selected nucleotide sequence database to assess the specificity of the primers for the target sequence. Additional tools and pipelines that encompass both the identification of target sequences from bacterial draft genomes and the design of primer candidates include, for instance, RUCS and TOPSI (Thomsen et al. 2017; Vijaya Satya et al. 2010). RUCS is able to identify unique core sequences in a positive set of genomes (target) compared to a negative set of genomes (non-target). It designs primers for the core sequences and validates them with an \textit{in silico} PCR validation method against the positive and negative reference set. TOPSI is an automated high-throughput pipeline for the design of primers, primarily developed for pathogen-diagnostic assays. It identifies sequences present in all input genomes and designs specific primers accordingly.

We aimed to design a series of primers that function with the same qPCR cycling conditions and primer concentrations for later usage in a high-throughput microfluidic qPCR platform. Although TOPSI and RUCS were initially considered for the automated design of primers, TOPSI could not be used because no Linux-based cluster was available. RUCS was easily installed, but we were not able to create primer pairs in initial tests with a small set of positive (target) and negative (non-target) genomes. The example in the original publication of RUCS (using \textit{Escherichia coli} genomes as positive and negative sets) indicates that RUCS works best for very similar genome assemblies in the positive and the negative sets. From this example and the initial test, we inferred that RUCS requires a carefully selected training set of positive and negative genomes to identify target sequences, which is a demanding task in the case of complex microbial systems such as those involved in the production of fermented foods and was therefore not suitable for our high-throughput approach.

This study presents a pipeline for automated high-throughput screening for species-specific target regions combined with the design of primer candidates for these sequences. The process of
primer design is fully automated from the download of bacterial genomes to the quality control of primer candidates. The pipeline runs on a standard computer with a multi-core processor and a minimum of 16 GB RAM. We have applied the SpeciesPrimer pipeline to a set of four bacterial species occurring in cheese and other dairy products and validated the primers in silico and in vitro by performing qPCR experiments with a variety of target and non-target strains.

**Description**

**Overview**

The SpeciesPrimer pipeline consists of three main parts (Table 1). First, genome assemblies are downloaded, annotated and then subjected to quality control. Second, a pan-genome analysis is performed to identify single copy core genes. Conserved sequences of these core genes are then extracted and the specificity for the target species is assessed. Finally, primers are designed for these species-specific conserved core gene sequences and subsequently evaluated in a primer quality control step.

**Part 1: Input genome assemblies**

The minimal command line input for the pipeline is the species name. Further, a list of non-target species names can be specified (e.g., species found in the investigated ecosystem but that should not be detected in the specific qPCR assay). For downloading genome assemblies from the National Center for Biotechnology Information (NCBI) automatically, a valid e-mail address is required for accessing the NCBI E-utilities services (Sayers 2009). The pipeline works with a pre-formatted NCBI BLAST database (nt), containing partially non-redundant nucleotide sequences. A local copy of the nt database is required. It can be downloaded from NCBI using the update_blastdb.pl script from the BLAST+ package (Altschul et al. 1990), via FTP from the NCBI FTP server or with the pipeline script (getblastdb.py).

The user-provided species name is used to search for genome assemblies in the NCBI database. The Biopython Entrez module (Cock et al. 2009) searches the NCBI taxonomic identity (taxid) for the target species in the taxonomy database and downloads the genome assembly summary report. Afterwards, SpeciesPrimer downloads the genome assemblies in FASTA format from the NCBI RefSeq FTP server using the links specified in the summary report. Finally, the downloaded genome assemblies are annotated with Prokka (Seemann 2014).
The quality of the genome assemblies is a crucial factor for the pan-genome analysis. Genome assemblies deposited with the wrong taxonomic label or low-quality assemblies drastically reduce the number of identified core genes and of conserved sequences for primer design. The initial quality control step is intended to remove such assemblies from the subsequent analysis. For the verification of the taxonomic classification, the user can choose one or several genes from five conserved housekeeping genes (16S rRNA, tuf, recA, dnaK and pheS). Genome assemblies without an annotation for the specified conserved housekeeping genes or genome assemblies consisting of more than 500 contigs are removed from the downstream pan-genome analysis. The sequences of the specified conserved housekeeping genes are blasted against the local nt database. Genome assemblies pass the quality control if the best BLAST hit for all sequences is a sequence arising from the target species.

Part 2: Identification of target sequences for primer design
A pan-genome analysis is performed using Roary (Page et al. 2015) to identify the core genes of the target species. Based on the results of the pan-genome analysis, single copy core genes are identified. The gene_presence_absence.csv produced by Roary reports the presence (or absence) of every annotated gene for every input genome assembly. Single copy core genes are the genes for which the number of assemblies harboring the sequence and the number of total identified sequences equals the number of total input assemblies. An sqlite3 database containing all annotated sequences of all assemblies is compiled (https://github.com/EnzoAndree/tutorials/blob/patch-1/DBGenerator.py). This database is queried for single copy core genes and the nucleotide sequences are saved in multi-FASTA format. Each multi-FASTA file contains the sequences of one single copy core gene from each input genome assembly. These sequences are aligned using the probabilistic multiple alignment program Prank (Löytynoja 2014). A consensus sequence with ambiguous bases is then created using consambig from the EMBOSS package (Rice et al. 2000). The alignments and extraction of the consensus sequence are performed in parallel for several core genes using GNU parallel (Tange 2011). Continuous consensus sequences longer than the minimal PCR product length, harboring less than two ambiguous bases in the range of 20 bases are used for the subsequent steps of the pipeline. These conserved consensus sequences are used for a BLAST search against the local nt database using the discontiguous BLAST algorithm and an e-value cutoff of 500. For all hits in the
BLAST results, the species name is extracted from the sequence description and compared with the names in the species list (non-target species). If any species name in the species list matches a hit in the BLAST results the corresponding query sequence is discarded, otherwise the sequence is classified as specific for the target and considered for primer design.

**Part 3: Primer design**

Primer3 is used to design primers for the unique single copy core gene sequences. As pipeline default the optimum primer melting temperature is set to 60 °C and the maximal primer length is set to 26 bases, all other settings are the default settings of the primer3web version ([http://primer3.ut.ee](http://primer3.ut.ee), accessed November 29, 2018). The minimal and maximal amplicon size of the PCR product can be specified individually for every target species through the command line options. The other parameters for primer3 cannot be changed individually, but the general primer3 settings can be changed by modifying the primer3 settings file.

The primer quality control consists of three parts, an in silico PCR to evaluate the specificity of the primer for the template, an estimation of secondary structures of the amplicon sequence and the estimation of the potential to form primer dimers. The specificity check for each primer pair is performed with MFEprimer 2.0 (Qu et al. 2012). For the evaluation of the specificity, three indexed databases are generated: the target template database, the non-target sequence database and the target genome database. The target template database consists of the unique conserved core gene sequences used as template for primer design. The non-target sequence database is compiled from sequences of non-target species, which show similarities to the primer sequences. To identify these sequences, a BLAST search with all primers against the local nt database is performed. BLAST hits with a species name in the description matching a name in the user-specified non-target species list are selected. These selected sequences and 4000 base pairs up- and downstream are extracted from the nt database using the blastdbcmd tool. The target genome database is composed of maximal 10 of the input genome assemblies. If the assembly summary report from the automatic download of genome assemblies from NCBI is available the genome assemblies as complete as possible are preferred (assembly status: complete > chromosome > scaffold > contig). The target sequence database is used to evaluate the maximum primer pair coverage (PPC), a value used by MFEprimer 2.0 to score the ability of the primer pair to bind to a DNA template. The maximum value of the PPC is 100, all primer pairs with a PPC value lower than the specified threshold (mfethreshold, default = 90) for their template are excluded. Next,
MFEprimer 2.0 is used to score the binding of the primer pairs to the sequences of the non-target sequence and the target genome database. The difference of the PPC for the DNA template and the specified threshold ($\Delta$threshold = PPC – mfethreshold) is used as a threshold for the maximum PPC value a primer pair is allowed to have for a non-target sequence. Strong secondary structures at the 5’- or the 3’- end of the PCR product could impair efficient primer binding. Therefore, the PCR products of the primer pairs are submitted to mfold (Zuker et al. 1999) to exclude PCR products with strong secondary structures at the annealing temperature of 60 °C. Moreover, as primer dimers can yield unspecific signals during the qPCR run, the 3’- ends of the primer pairs are checked for their potential to form homo- or hetero-dimers using a Perl script (MPprimer_dimer_check.pl) from MPprimer (Shen et al. 2010).

The pipeline output is a list containing the primer name, primer pair coverage (MFEprimer) and penalty values, primer and template sequences and melting temperatures (Primer3). Further, a report of the genome assembly quality control, a file containing the pipeline run statistics, the core gene alignment and the phylogeny in newick format can be found in the output directory.

**Materials & Methods**

**Primer design**

SpeciesPrimer pipeline runs were performed on a virtual machine (Oracle VM VirtualBox 5.2.8) with Ubuntu 16.04 (64-bit) and docker installed, using 22 of 24 logical processors from two Intel Xeon E5-2643 CPUs and 32 GB of RAM. The used docker image is available from https://hub.docker.com/r/biologger/speciesprimer.

The species list consisted of 259 species and subspecies names detected in dairy products, namely from species names collected from data of 16S rRNA meta-genome sequencing studies in milk and cheese varieties (Marco Meola Agroscope, pers. comm.) and dairy-related bacteria from the list of bacterial species and subspecies with technological beneficial use in food products (Almeida et al. 2014).

The SpeciesPrimer pipeline was run with the input genome assemblies, parameters and the species list specified in the supplemental information (Data S1). Genome assemblies from the strain collection of Agroscope were included for the Pediococci.
**In silico validation**

For the in silico validation, PCR products for the designed primer pairs were used for an online BLAST search against the RefSeq Genomes Database (refseq_genomes) limited to bacterial genomes. The search was performed by qblast (biopython), using blastn, the maximum hitlist size was set to 5000 and the expect threshold (e-value) was set to 500.

Primer pairs were tested for specificity using online Primer-BLAST (Ye et al. 2012). The primers were blasted against the nucleotide collection BLAST database (nr) limited to sequences from bacteria. Default settings were used, except for the primer specificity stringency that was set to ignore targets that have nine or more mismatches to the primer.

**In vitro validation**

The inclusivity of the primer pairs was assayed by performing qPCR with 2 ng DNA of 20 to 25 strains of the target species in technical duplicates. The linear amplification of genomic DNA and PCR efficiency was examined by ten-fold dilution series of the type strain DNA in a range from \(10^6\) to \(10^1\) genome copies per reaction. DNA concentration for the corresponding number of genome copies was estimated by taking the genome size of the type strain (https://www.ncbi.nlm.nih.gov/genome) and an average weight of \(1.096 \cdot 10^{-21}\) g per base pair.

The exclusivity of the primer pairs was assayed by performing qPCR with 2 ng DNA from various bacterial species in technical duplicates found in dairy products in four qPCR runs including three strains of the target species as positive control.

**Bacterial strains**

Strains stored within the Agroscope Culture Collection at -80 °C in sterile reconstituted skim milk powder (10 %, w/v), were reactivated and cultivated according to the conditions specified in Data S2.

**DNA extraction**

Unless otherwise noted, all reagents were purchased from Merck, Darmstadt, Germany.

Bacterial pellets harvested from 1 ml culture by centrifugation (10000 x g, 5 min, room temperature) were used for DNA extraction. For a pre-lysis treatment, the bacterial cells were incubated in 1 ml of 50 mM sodium hydroxide for 15 min at room temperature. Afterwards cells were collected by centrifugation (10000 x g, 5 min, room temperature) and then treated with
lysozyme (2.5 mg/ml dissolved in 100 mM Tris(hydroxymethyl)aminomethane, 10 mM ethylenediaminetetraacetic acid (Calbiochem, San Diego, USA), 25 % (w/v) sucrose, pH 8.0) for 1 hour at 37 °C. After the pre-lysis treatment, the bacterial cells were collected by centrifugation (10000 x g, 5 min, room temperature). Cell lysis and genomic DNA extraction was performed using the EZ1 DNA Tissue kit and a BioRobot® EZ1 workstation (Qiagen, Hilden, Germany) according to the manufacturer's instructions and eluted in a volume of 100 µl. The DNA concentration was measured using a NanoDrop® ND-1000 Spectrophotometer (NanoDrop Technologies, Thermo Fisher Scientific, Waltham, MA, USA).

Quantitative real-time PCR
The qPCR assays were performed in a total reaction mix volume of 12 µl containing 6 µl 2x SsoFast™ EvaGreen® Supermix with low ROX (Biorad, Cressier, Switzerland), 500 nM of forward and reverse primers, respectively, and 2 µl of DNA. Each sample was measured in technical duplicates. The qPCR cycling conditions were an initial denaturation at 95 °C for 1 minute followed by 35 cycles of 95 °C for 5 seconds and 60 °C for 1 minute. For the melting curve analysis, a gradient from 60 – 95 °C with 1 °C steps per 3 seconds was performed. All qPCR assays were run on a Corbett Rotor-Gene 3000 (Qiagen). The analysis was performed using Rotor-Gene 6000 Software 1.7 with dynamic tube normalization and a threshold of 0.05 for quantification cycle (Cq) value calculation, the five first cycles were ignored for the determination of the Cq values. The peak calling threshold for the melt curve analysis was set to -2 dF/dT and a temperature threshold was set 2 °C lower than the positive control peak.

Average nucleotide identity calculations
Average nucleotide identity (ANI) calculations were performed with OrthoANIu (Yoon et al. 2017)). All Enterococcus faecium genome assemblies were compared to the E. faecium reference sequence (NC_017960.1), while all Pediococcus acidilactici genome assemblies were compared to the P. acidilactici reference sequence (NZ_CP015206.1). The genome assemblies were grouped based on the phylogeny tree of the core gene sequences built with FastTree (Price et al. 2010), the ANI values were collected and the average, minimum and maximum was calculated for each group.
Results

Primer design

Primer design for four bacterial species commonly found in cheese was performed with the SpeciesPrimer pipeline. The pipeline runs were completed in two to eight hours, excluding the time required for downloading and annotation of the genome assemblies. Depending on the number of genome assemblies, downloading and annotation of the genome assemblies took from 24 minutes (27) to 12 hours 27 minutes (575). The average time for downloading and annotation was two seconds and one minute six seconds, respectively. The analysis of the *Enterococcus faecalis*, *Enterococcus faecium*, *Pediococcus acidilactici* and *Pediococcus pentosaceus* assemblies resulted in 15, 2, 2 and 160 identified primer pair candidates, respectively (Table 2). The primer pair candidates for *E. faecalis* and *P. pentosaceus* were filtered for the highest primer pair coverage score (*E. faecalis*: 2; *P. pentosaceus*: 29); for *P. pentosaceus* only the two primer pairs with the lowest primer pair penalty values were selected. The phylogeny trees of the core gene alignments from *E. faecium* and *P. acidilactici* were created using Roary and FastTree (Figure 1). The unrooted tree from the concatenated core genes of *E. faecium* shows the phylogenetic distance of two distinct groups of sequences, a main cluster with 531 sequences and a subcluster with 44 sequences. The tree made with the concatenated core gene sequences of *P. acidilactici* shows the phylogenetic distance of one sequence from all other sequences. From this observation, the existence of different species or subspecies was suspected. Calculation of the average nucleotide identity (ANI) has been proposed as a valuable tool to determine species boundaries (Richter & Rossello-Mora 2009). Therefore, we performed ANI calculations for the genome assemblies and the reference sequence for *E. faecium* (NC_017960.1) using the tool OrthoANIu. The genome assemblies of the *E. faecium* subcluster have an average ANI of 94.67 %. The ANI between the genome assembly of the *P. acidilactici* strain FAM 18987 and the NCBI reference sequence for *P. acidilactici* (NZ_CP015206.1) was only 89.21 %. The OrthoANI values (Table 3) of the assemblies in the subclusters of *E. faecium* (94.15 - 95.60 %) are at the border and the value for the *P. acidilactici* strain FAM 18987 (89.21%) is below the proposed species threshold cutoffs (95 - 96 %) (Kim et al. 2014; Richter & Rossello-Mora 2009). *P. acidilactici* strain FAM 18987 should therefore probably be assigned to a new species or subspecies. However, for certain species also lower boundary cutoffs might be reasonable (Ciufò et al. 2018). According to the
current taxonomic classification, we proceeded with the assumption that these genome assemblies reflected the actual diversity of strains and thus included the assemblies for the primer design.

Two test cases were generated to exemplify the influence of the input genome assemblies on the pipeline results. Firstly, a single genome assembly with a wrong taxonomic label was used as input in addition to the correctly labelled genome assemblies. Introducing a genome assembly with a wrong taxonomic label (GCF_000415325.2, *E. faecalis*) into the pool of *E. faecium* genome assemblies resulted in a decrease of identified core genes (from 1131 to 43) and provided no species-specific sequence. Secondly, the genome assembly of the *P. acidilactici* strain (FAM 18987) that was distinct from the other assemblies in the phylogenetic tree with an ANI to the reference sequence below 90% was excluded from the pipeline run. This resulted in an increased number of identified core genes (from 921 to 1238), of species-specific sequences (from 54 to 516) and of reported primer pairs (from 2 to 53). The results of the two test cases illustrate that the SpeciesPrimer pipeline performs best on closely related genome assemblies with a good overall quality.

**In silico validation**

Two parameters were selected as criteria for the primer validation using web-based BLAST. First, the BLAST hits for the predicted PCR product sequence should only match the target species. If sequences of other bacterial species matched to parts of the sequence, the corresponding primer pairs were discarded, unless more than three mismatches were found in each primer-binding region for the forward and reverse primers. Second, the primer binding sites in the target sequences were not allowed to have mismatches in the 3’-end region. The criterion for the primer validation by Primer-BLAST was that no predicted PCR products for other bacterial species had been reported by Primer-BLAST. Primer pairs exclusively binding to the target sequence of the target species were classified as specific. The results of the in silico validation are summarized in Table 4. With the exception of Ec_faeca_g3060_1_P0 and Ec_faeci_cysS_3_P1, all primer pairs showed a perfect match to their target sequences. For primer pair Ec_faeca_g3060_1_P0, the first three nucleotides of one sequence out of 690, are missing in the forward primer-binding region. For Ec_faeci_cysS_3_P1, only one sequence out of 1058 aligned sequences showed a single nucleotide transition in the reverse primer-binding region (Data S3).
**In vitro validation**

The specificity of the qPCR assays was assessed with 21 to 25 strains of the target species (inclusivity) and 121 non-target bacterial strains found in dairy products (exclusivity). The qPCR assay performance was assessed by 10-fold dilution series of type strain DNA from $10^6$ to $10^1$ copies per reaction. The results of the qPCR runs were interpreted as positive if both qPCR reactions (duplicates) reached the fluorescent threshold before quantification cycle 35 and the peak of the melting curve analysis was above the peak calling threshold (-2 dF/dT). A summary of the results is shown in Table 5. The primer sequences can be found in Table S1. The inclusivity of the qPCR assays was 100 % for the assays Ec_faeca_acuI, Ec_faeca_g3060, Ec_faeci_cysS, Pd_acidi_asnS, Pd_acidi_g1164, Pd_pento_nagK and Pd_pento_g4364. Only one qPCR assay, Ec_faeca_purD was negative for one of the tested target strains.

Out of the 121 non-target strains analyzed to determine the exclusivity of the qPCR assays (Figure 2), all strains were negative for Ec_faeca_acuI and Pd_acidi_asnS. Both assays targeting *E. faecium* were positive solely for one *L. fermentum* strain (FAM 20347). Later it was found that the stock culture of this strain was contaminated with an *E. faecium* strain (data not shown).

The assay Pd_pento_nagK targeting *P. pentosaceus* was positive for two out of three tested *Leuconostoc lactis* strains, the fluorescence signal reached the threshold after Cq 26, and the melting curve analysis showed a peak at 85 °C, while the positive control samples for this assay displayed a peak at 83.5 °C. Nine out of the 121 non-target strains were positive for the Ec_faeca_g3060 qPCR assay, for these samples the fluorescence signals reached the threshold after Cq 26 and had a melting curve peak at a higher temperature than the target PCR product.

The assays Pd_acidi_g1164 and Pd_pento_g4364 were positive for five and eight non-target strains, respectively. Notably, all three tested *Lactobacillus paracasei* strains were positive for the Pd_acidi_g1164 assay, the fluorescence signal reached the threshold around Cq 21 and 22 and they showed a distinct melting curve peak at 86 °C.

The qPCR assays displayed linear results between $10^1$ and $10^6$ genome copies per reaction. The calculated efficiency of the qPCR assays was between 92 and 100 %. The linear regression equations ($C_q = slope \cdot \log (copies) + intercept$) had slopes between -3.329 and -3.523 and correlation coefficients of 0.990 or above.

**Discussion**

After setup of the SpeciesPrimer docker container, the download of the local BLAST database
and the selection of the SpeciesPrimer run settings, no further manual handling was required to
get primer pair candidates for all four bacterial species after a total time of 44 hours and 30
minutes. The number of input genomes and subsequently the number of retrieved primer pairs
for the specificity check have the highest impact on speed. During the specificity check, blasting
the primer sequences optimized for short sequences (blastn-short) and the subsequent
compilation and indexing of the non-target sequence database are the most time consuming
steps.

The results of the SpeciesPrimer pipeline for the four target species ranged from two to 160
identified primer pair candidates. Several factors can influence the number of identified primer
pairs, such as the quality of the input genome assemblies, assemblies with wrong taxonomic
labels and the genetic diversity within the species. A low-quality assembly with missing
sequences or contaminations can decrease the number of identified core genes. The initial quality
control helps to minimize the risk that such assemblies are included in the pipeline runs.

However, also an increased sequence diversity, either due to sequencing errors, assembly errors
or real diversity, limits the number and the length of identified conserved sequences.

Subsequently this affects the yield of reported primer pairs, since the pipeline selects highly
conserved sequences for primer design.

The specificity of the designed primers was evaluated in silico by BLAST with a more extensive
database (RefSeq Genome) than the one used for the specificity check during primer design. The
validation showed that the specificity of the tested amplicons was high and no other species than
the target species had an identical sequence. Most target sequences in the database showed a
perfect match for the primers in the primer-binding region. For all tested primer pairs, solely the
expected PCR products for the target species and no amplicons for other sequences were
predicted by Primer-BLAST. The results of Primer-BLAST indicate that the reported primer
pairs were very specific, even though the species list used for the specificity evaluation during
primer design covered only 259 non-target species.

In this work, 21 to 25 target strains for each target species and 121 non-target strains have been
tested to assess inclusivity and exclusivity of the qPCR assays, respectively. The in vitro
validation of primer pairs has shown that the in silico validation is not always able to predict
non-target PCR products. The fluorescence signals occurring at late quantification cycles (Cq >
30) are probably due to PCR products with suboptimal primer binding. Testing the qPCR assays
in mixtures and communities could be interesting to assess if these PCR products also accumulate in presence of target DNA. The specificity could be sufficient in mixtures due to competition for the primers and the difference in primer binding and amplification efficiency. For many research applications, qPCR assays with a low signal in negative samples are acceptable, assuming that low-level signals can be distinguished from low concentrations of target species DNA by the melting curve analysis. Further, for many applications, the annealing temperature can be optimized by empirical determination of a suitable annealing temperature and the primer concentration can be adjusted to improve the specificity of the assay (www.bio-rad.com/en-ch/applications-technologies/qpcr-assay-design-optimization). We did not try to optimize our assays with these measures, because the aim was to design primers for high-throughput qPCR, requiring the exact same PCR conditions. For the tested qPCR conditions the most specific qPCR assays were Ec_faecal_acuI (E. faecalis), Ec_faeciu_cysS (E. faecium), Pd_acidi_ansS (P. acidilactici) and Pd_pento_nagK (P. pentosaceus). Further work will be necessary in order to make these qPCR assays fully operational for the quantification of bacteria in a complex system such as food. For instance, suitable qPCR standards should be designed and validated, so that the limit of detection of each assay can be determined. Primer-BLAST and RUCS allow designing primers for different applications, but demand experience and manual manipulations. Primer-BLAST designs primers and performs specificity checks, but requires a user provided target sequence. In the case of RUCS manual manipulation and experience is needed to prepare the positive and negative reference sets. Compared to primer-BLAST and RUCS, the task SpeciesPrimer performs is really specialized, namely to design primers for species-specific sequences. In contrast, SpeciesPrimer requires no previous knowledge about the input genome assemblies and no manual manipulation of sequences has to be performed. The ability of SpeciesPrimer to run on standard computers with good performance instead of specialized high-performance computers, will hopefully allow primer design for a wide range of scientists. Docker containers simplify the installation procedure and should allow non-bioinformaticians to setup and use the SpeciesPrimer pipeline.

Conclusions

In this work, we presented the SpeciesPrimer pipeline, which is a fully automated pipeline from the download of bacterial genomes, the identification of conserved species-specific core genes to primer design and subsequent quality control of primer candidates. Primers for four bacterial
425 species were designed and validated and have shown to perform adequately under the same
426 qPCR conditions.
427 A standard computer with good performance, good quality genome assemblies, a local copy of
428 the nt BLAST database and a list of non-target bacterial species are the only requirements for
429 primer design with SpeciesPrimer. A complete image, of a Linux OS with all dependencies and
430 the pipeline scripts, is available from Dockerhub. To simplify primer design for users not
431 familiar with command line tools, a graphic user interface is provided in the latest version of
432 SpeciesPrimer. SpeciesPrimer facilitates efficient primer design for species-specific
433 quantification, paving the way for a fast and accurate quantitative investigation of microbial
434 communities.

Acknowledgements
We would like to thank Marco Meola and Remo Schmidt for critically reading the manuscript
and many helpful comments and Daniel Marzohl, Nadine Sidler, Elvira Wagner and Kotchanoot
Srikham for their valuable technical help.

References


Table 1 (on next page)

Overview of the SpeciesPrimer pipeline workflow and the used software.
<table>
<thead>
<tr>
<th>Pipeline workflow</th>
<th>Tools</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Input genome assemblies</td>
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<td></td>
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<tr>
<td>- download</td>
<td>NCBI Entrez (Biopython)</td>
<td>(Cock et al. 2009; Sayers 2009)</td>
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<tr>
<td>- annotation</td>
<td>Prokka</td>
<td>(Seemann 2014)</td>
</tr>
<tr>
<td>- quality control</td>
<td>BLAST+</td>
<td>(Altschul et al. 1990)</td>
</tr>
<tr>
<td>Core gene sequences</td>
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<td></td>
</tr>
<tr>
<td>- identification</td>
<td>Roary</td>
<td>(Page et al. 2015)</td>
</tr>
<tr>
<td>- phylogeny</td>
<td>FastTree 2</td>
<td>(Price et al. 2010)</td>
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<td>- selection of conserved sequences</td>
<td>Prank, consambig (EMBOSS), GNU parallel</td>
<td>(Löytynoja 2014)</td>
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<td>- evaluation of specificity</td>
<td>BLAST+</td>
<td>(Rice et al. 2000)</td>
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<td></td>
<td></td>
<td>(Tange 2011)</td>
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<td></td>
<td></td>
<td>(Altschul et al. 1990)</td>
</tr>
<tr>
<td>Primer</td>
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<tr>
<td>- design</td>
<td>Primer3</td>
<td>(Untergasser et al. 2012)</td>
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<tr>
<td>- quality control</td>
<td>BLAST+, MFEPrimer 2.0, MPprimer, Mfold</td>
<td>(Altschul et al. 1990)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(Qu et al. 2012)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(Shen et al. 2010)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(Zuker et al. 1999)</td>
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Table 2 (on next page)

Pipeline input and run statistics.
<table>
<thead>
<tr>
<th>Species</th>
<th>( E. \ faecalis )</th>
<th>( E. \ faecium )</th>
<th>( P. \ acidilactici )</th>
<th>( P. \ pentosaceus )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pipeline input</td>
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<td></td>
</tr>
<tr>
<td>NCBI genomes</td>
<td>390</td>
<td>575</td>
<td>9</td>
<td>14</td>
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<tr>
<td>ACC genomes</td>
<td>0</td>
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<tr>
<td>Total genome assemblies</td>
<td>390</td>
<td>575</td>
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<td>Download and annotation (h:min)</td>
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<td>12:27</td>
<td>1:55</td>
<td>0:24</td>
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<tr>
<td>Pipeline statistics</td>
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<td></td>
<td></td>
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</tr>
<tr>
<td>Running time (h:min)</td>
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<td>8:05</td>
<td>1:55</td>
<td>4:25</td>
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<td>Core genes</td>
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<td>1131</td>
<td>921</td>
<td>1341</td>
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<tr>
<td>Single copy core genes</td>
<td>632</td>
<td>563</td>
<td>641</td>
<td>889</td>
</tr>
<tr>
<td>Conserved sequences</td>
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<td>624</td>
<td>566</td>
<td>2782</td>
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<tr>
<td>Species-specific sequences</td>
<td>329</td>
<td>36</td>
<td>54</td>
<td>672</td>
</tr>
<tr>
<td>Potential primer pairs</td>
<td>89</td>
<td>4</td>
<td>7</td>
<td>632</td>
</tr>
<tr>
<td>Primer pairs after QC</td>
<td>15</td>
<td>2</td>
<td>2</td>
<td>160</td>
</tr>
</tbody>
</table>

1 QC: primer quality control, ACC: Agroscope culture collection
Table 3 (on next page)

Summarized results of the average nucleotide identity (ANI) calculations.
<table>
<thead>
<tr>
<th></th>
<th>E. faecium main cluster</th>
<th>E. faecium subcluster</th>
<th>P. acidilactici main cluster</th>
<th>P. acidilactici subcluster</th>
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</thead>
<tbody>
<tr>
<td>Assemblies</td>
<td>530</td>
<td>44</td>
<td>125</td>
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<td>ANI (%)</td>
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<td>89.21</td>
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<tr>
<td>average</td>
<td>99.43</td>
<td>94.67</td>
<td>98.28</td>
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<tr>
<td>maximum</td>
<td>99.86</td>
<td>95.60</td>
<td>98.83</td>
<td></td>
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<tr>
<td>minimum</td>
<td>98.19</td>
<td>94.15</td>
<td>96.88</td>
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</tr>
</tbody>
</table>
Table 4 (on next page)

Summary of the *in silico* validation of the selected primer pairs.
<table>
<thead>
<tr>
<th>Target species</th>
<th>Primer pair</th>
<th>PPC</th>
<th>BLAST (perfect/total)</th>
<th>primer-BLAST (perfect/total)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>E. faecalis</em></td>
<td>Ec_faec_acul_1_P0</td>
<td>100</td>
<td>specific (694/694)</td>
<td>specific (24/24)</td>
</tr>
<tr>
<td></td>
<td>Ec_faec_g3060_1_P0</td>
<td>100</td>
<td>specific (689/690)</td>
<td>specific (24/24)</td>
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<tr>
<td><em>E. faecium</em></td>
<td>Ec_faei_cysS_3_P1</td>
<td>96.7</td>
<td>specific (1057/1058)</td>
<td>specific (63/63)</td>
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<tr>
<td></td>
<td>Ec_faei_purD_2_P0</td>
<td>93.3</td>
<td>specific (1083/1083)</td>
<td>specific (63/63)</td>
</tr>
<tr>
<td><em>P. acidilactici</em></td>
<td>Pd_acidi_asnS_2_P0</td>
<td>90.1</td>
<td>specific (19/19)</td>
<td>specific (5/5)</td>
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<tr>
<td></td>
<td>Pd_acidi_g1164_1_P0</td>
<td>93.3</td>
<td>specific (23/23)</td>
<td>specific (5/5)</td>
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<tr>
<td><em>P. pentosaceus</em></td>
<td>Pd_pento_nagK_1_P0</td>
<td>100</td>
<td>specific (15/15)</td>
<td>specific (7/7)</td>
</tr>
<tr>
<td></td>
<td>Pd_pento_g4364_1_P0</td>
<td>100</td>
<td>specific (15/15)</td>
<td>specific (7/7)</td>
</tr>
</tbody>
</table>
Summarized results of the *in vitro* validation of the selected qPCR assays.

Inclusivity: Number of positive DNA samples / total number of target species DNA samples.
Exclusivity: Number of DNA samples showing a fluorescence signal below quantification cycle 35 and a melting curve peak above the threshold / total number of non-target DNA samples.

Calculated efficiency, slope, intercept and correlation coefficient ($R^2$) of the linear regression equation.
<table>
<thead>
<tr>
<th>Species</th>
<th>E. faecalis</th>
<th>E. faecium</th>
<th>P. acidilactici</th>
<th>P. pentosaceus</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target gene</td>
<td>acu</td>
<td>g3060</td>
<td>cysS</td>
<td>purD</td>
</tr>
<tr>
<td>Exclusivity</td>
<td>0/121</td>
<td>9/121</td>
<td>0*/121</td>
<td>0*/121</td>
</tr>
<tr>
<td>Efficiency</td>
<td>98 %</td>
<td>97 %</td>
<td>92 %</td>
<td>97 %</td>
</tr>
<tr>
<td>Intercept</td>
<td>32.107</td>
<td>32.694</td>
<td>32.006</td>
<td>31.051</td>
</tr>
<tr>
<td>R²</td>
<td>0.998</td>
<td>0.997</td>
<td>0.990</td>
<td>0.996</td>
</tr>
</tbody>
</table>

1 * Contamination of stock culture of the strain FAM20347 with E. faecium.
Figure 1

Phylogeny of core gene alignments.

(A) *E. faecium*. (B) *P. acidilactici*. The phylogenies are displayed with SeaView (http://doua.prabi.fr/software/seaview) in circular view.
**Figure 2**

qPCR assay quantification cycle heatmap.

Depicted are all tested non-target strains and their average quantification cycle (technical duplicates). Bars represent results with a melt curve peak above the threshold and a Cq value below Cq 35. The gray shades represent the Cq values from 10 to 35 (if no fluorescent signal was measured the value was set to Cq 35). Abbreviations: A.: *Acidipropionici*; Cl.: *Clostridium*; Lb.: *Lactobacillus*; Ln.: *Leuconostoc*; Pb.: *Propionibacterium*; Pd.: *Pediococcus*; Sc.: *Streptococcus*; NTC: no template control.