

# Identification of isolated or mixed strains from long reads: a challenge met on *Streptococcus thermophilus* using a MinION sequencer

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## Abstract

This study aimed to provide efficient recognition of bacterial strains on personal computers from MinION (Nanopore) long read data. Thanks to the fall in sequencing costs, the identification of bacteria can now proceed by whole genome sequencing. MinION is a fast, but highly error-prone sequencing device and it is a challenge to successfully identify the strain content of unknown simple or complex microbial samples. It is heavily constrained by memory management and fast access to the read and genome fragments. Our strategy involves three steps: indexing of known genomic sequences for a given or several bacterial species; a request process to assign a read to a strain by matching it to the closest reference genomes; and a final step looking for a minimum set of strains that best explains the observed reads. We have applied our method, called *ORI*, on 77 strains of *Streptococcus thermophilus*. We worked on several genomic distances and obtained a detailed classification of the strains, together with a criterion that allows merging of what we termed 'sibling' strains, only separated by a few mutations. Overall, isolated strains can be safely recognized from MinION data. For mixtures of several non-sibling strains, results depend on strain abundance.

## DATA SUMMARY

The authors confirm all supporting data, code and protocols have been provided within the article or through supplementary data files:

The *ORI* code (Oxford Nanopore Reads Identification) is available at <https://github.com/gsiekaniec/ORI>.

All sequencing data used in our experiments (raw Nanopore fastq reads) can be downloaded on the Genouest server (<https://data-access.cesgo.org/index.php/s/IApWiOf1BFYUpQV>).

The interactive ITOL tree of *S. thermophilus* species is available at: <https://itol.embl.de/tree/131254134671311597925585>, and the complete list of associated 'maximal biclusters' (subset of *S. thermophilus* strains and subset of

associated specific genes) is available on <https://github.com/gsiekaniec/GeneTree>.

## INTRODUCTION

For industrial, agri-food and clinically relevant bacteria, rapid identification at the fine level of strains is necessary and remains a great challenge. This is clear in the field of health (e.g. differentiating the non-pathogenic *Escherichia coli* strain MG1655 from enterohemorrhagic strain 278F2 [1]). It is also necessary in many other fields such as ecology, where adaptation to a given niche often leads to the emergence of new strains [2], or in food processes, such as fermentation [3]. Moreover, for newly isolated strains or species, looking for the closest known organism can yield important information for its functional characterization.

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**Abbreviations:** ANI, average nucleotide identity; ASP, answer set programming; HMW, high molecular weight; LCA, lowest common ancestor; MCC, Matthews correlation coefficient; MICFAM, microScope gene families; NGS, next generation sequencing; ONT, Oxford Nanopore Technologies; ORI, Oxford Nanopore reads identification; SD, standard deviation; WGS, whole genome sequencing.

*Streptococcus thermophilus* strains have been provided by the International Centre for Microbial Resources – Food-Associated Bacteria (CIRM-BIA), STLO UMR 1253 INRAE-Agrocampus, France ([https://www6.inrae.fr/cirm\\_eng/Food-Associated-Bacteria](https://www6.inrae.fr/cirm_eng/Food-Associated-Bacteria)). All sequence data were annotated by the MicroScope Platform (1). New genome accession numbers are listed in Table S2.

**Data statement:** All supporting data, code and protocols have been provided within the article or through supplementary data files. Nine supplementary tables, two supplementary text and three supplementary figures are available with the online version of this article.

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The development of next generation sequencing (NGS) technologies in the last two decades has revolutionized the characterization of microbial communities, without prior bacterial culturing. For instance, Illumina platforms that produce short reads (up to 300 bp) are now widely used for bacterial community taxonomic profiling [4–6] and *de novo* bacterial genome assembly [7, 8]. The overlapping reads produced are assembled into larger sequences that are mapped on sequence databases. However, read length makes these processes challenging, due to repetitive elements larger than reads, and this leads to fragmented assemblies. Third-generation sequencing technologies such as PacBio or Oxford Nanopore Technologies (ONT), producing read lengths reaching tens of kilobases or even more, can overcome this issue [9] but are error-prone. The best assemblers to date are hybrid, using both short and long reads [10].

Here we consider the issue of identification from long reads only, without prior assembly of these reads. Currently, identification from long reads hardly goes down to a level finer than genus or species [11]. Problems include the proximity of strains within a species, the prevalence of certain species in complex samples that can blur the recognition process, and databases that are far from cover all microbial diversity. Therefore, there is still a gap between the promises of whole genome sequencing (WGS) for microbial identification, notably at the finer-grained taxonomic levels, and its large application in demanding operational contexts [12]. Among the possible obstacles to its diffusion is the very rapid advance in technology, which makes it difficult to produce stable identification software. While programs are becoming mature for work on short reads, the landscape is still very dynamic and uncertain for long reads.

Commonly used identification software can be broadly divided into two categories, the sequence alignment and the k-mers-based approaches. Sequence alignment, largely popularized by the program BLAST and its numerous variants [13], consists in optimally matching a read sequence with a fragment of a genomic sequence (representative software are MEGAN [14], MG-RAST [15], PanPhlAn [16], Centrifuge [17] and Kaiju [18]). To reduce their practical complexity, these algorithms are often limited to high-scoring alignments and loose sensitivity. Over the last few years, research effort has focused on simplified representations of genomes and reads, which has led to a significant improvement in comparison performance. The principle is to split genomic sequences into overlapping k-mers, short strings of fixed size (typically 30 nt). It is possible to store them compactly in an index for all known genomes and retrieve their presence very efficiently in sequenced reads in order to assign each read to its closest genomes. This powerful approach is followed by major current identification tools such as Kraken [19], Kraken 2 [20], CLARK [21], StrainSeeker [22] and Opal [23].

The majority of these tools use short reads because they have been available for a long time and with good accuracy. Unfortunately, they generally lack long-range genomic information to resolve differences at the strain level [24]. Paired-end reads,

## Impact Statement

This paper describes a new efficient method and the associated software for bacterial strain identification from few long read sequencing data without prior assembly. Very few identification software programs have been designed for long read data, a source that offers more information for the discrimination of strains at the cost of a higher error rate. Our work improves the state of the art by combining sensitivity and robustness, based on an efficient data index and exact optimization for parsimonious explanation of result strains. We have validated the entire process from sample sequencing to sample identification on a species whose strains are difficult to distinguish, *Sreptococcus thermophilus*. It is possible to address any set of bacterial species by changing the index, thanks to a compiler accepting any set of genomic sequences. Our tool can therefore be useful to all microbiologists wishing to control the presence or absence of certain strains possibly mixed within a sample. We chose to focus on data from the Oxford Nanopore MinION device since its low cost and real-time sequencing capacities are well suited for testing in any laboratory or even on site. As all programs can be freely downloaded from our sites, and the databases can be configured for the species of interest of each laboratory, we hope this tool will have considerable impact.

comprising short sequences at the end of a fragment separated with a variable gap, provide slightly more information since they can be extended to larger reads by trying to bridge the gap using other reads [25], but the process may introduce errors and these reads seem to have been used mainly for assembly and comparison purposes [26]. Relatively few identification software programs accept long reads. The most popular is Kraken and its more recent version Kraken 2 [20], since it is easy to install, robust and fast. This software is based on the decomposition of genomes into k-mers and uses a very efficient storage structure. Some k-mers are specific to a unique species and some are linked to the lowest common ancestor (LCA) in terms of genus or family of all species where they occur. Thus, reads are classified at various levels of this hierarchy depending on their k-mer content. Kraken 2 rarely allows assigning a read at a level below the species because it uses a compression process (minimizers) that reduces the differences between genomes. Minimizers represent a sequence by a smaller included one, and serve to save space and time by retaining only part of the information, although they introduce a certain number of false positives during the identification process [27]. StrainSeeker [22] is a tool with a more restricted vocation, but which is of interest for strain identification from short reads. It uses a database of k-mer lists organized along a guide tree of bacteria (user-provided) where each leaf corresponds to a strain-specific set of k-mers. It is compatible with long reads, although we do

**Table 1.** Average percentage error rate in the *S. thermophilus* sequences (calculated from an alignment of the reads against the reference genomes performed with Minimap2)

The filters retain only sequences with a quality greater than 9 and a size greater than 2000 bp.

Errors	Mismatches	Deletions	Insertions	Total
All sequences	1.50%	2.16%	1.40%	5.06%
With filters	1.44%	2.08%	1.37%	4.89%

not know of any application using it in this context. It was designed primarily for the identification of isolates but can handle mixtures of a few strains. Another tool, *Centrifuge* [17], follows a read alignment approach and is at the core of the species identification workflow in the platform EPI2ME (ONT). Unfortunately, its index size can become very large if it has to include strain-level genomes. *Metamaps* [28], specifically developed for the analysis of metagenomic datasets, also uses alignments to identify species/strains, is designed for long reads and, according to its authors [28], seems more precise than *Centrifuge*. However, the current version is not simple to instal and lacks robustness (users may suffer from ‘core dump’ errors).

This paper describes the use of long read sequencing technologies for the identification of bacteria at the strain level. We chose *Streptococcus thermophilus* as a model species given its importance in the food industry and the health sectors as a dairy starter and a probiotic [29] and because it encompasses closely related strains (see Text S1, available in the online version of this paper) with a low genomic diversity [30, 31]. We have developed a new strain identification method called ORI (Oxford Nanopore Reads Identification) and compared it to *Kraken 2* and *StrainSeeker*. This approach dedicated to long reads has a technical background similar to *Kraken*, but differs in its goal to recognize at the strain level. The software for ORI can be found and downloaded from the following site: <https://github.com/gsiekaniec/ORI>. This work demonstrates that despite their significant error rate, the length of reads from MinION (ONT) allows fine-scale taxonomic assignment, even with mixtures and with closely related strains such as those from *S. thermophilus*. ORI can be easily adapted to other genera or species.

## METHODS

### Bacterial strains, growth conditions and DNA extraction

A set of 46 genomes of *S. thermophilus* (Table S1) from public databases and 31 strains (Table S2) from the collection of the CIRM-BIA resource centre were used in this study. *Streptococcus macedonicus* PA (CIRM-BIA2314) and *Lactobacillus delbrueckii* subsp. *bulgaricus* ATCC 11842 (CIRM-BIA658) from the CIRM-BIA collection were added as controls (different species/genus).

The 31 *S. thermophilus* strains and *S. macedonicus* PA were precultured anaerobically in LM17 medium (M17 medium supplemented with 2%, w/v, lactose) [32] initially inoculated at 1% (v/v), and incubated at 42 °C for 9 h. LM17 medium was further inoculated at 1% (v/v) with the LM17 preculture and grown overnight at 42 °C. *L. delbrueckii* subsp. *bulgaricus* ATCC 11842 was grown in MRS medium [33] under the same conditions. To avoid excessive fragmentation, DNA extraction was performed using the Genomic-tip 100/G from Qiagen, according to the manufacturer's protocol with slight modifications. Briefly, lysozyme, proteinase K and RNase A quantities were doubled and lysis incubation times were increased to 1.5 h. After DNA elution from the column, DNA was precipitated with isopropanol and spooled using a glass rod. It was immediately transferred in a clean microcentrifuge Eppendorf tube containing 200 µL EB buffer from Qiagen (10 mM Tris-Cl, pH 8.5) and allowed to dissolve overnight at 4 °C. Finally, DNA integrity and size (>10 kb) were assessed by electrophoresis on an agarose gel (0.7%, w/v, TBE 0.5×, migration under 100 V). DNA purity was estimated by Nanodrop measurements (ND-1000 Spectrophotometer). DNA quantification was done using a 1× dsDNA high-sensitivity assay kit on a QUBIT 4 Fluorometer (Thermo Fisher Scientific).

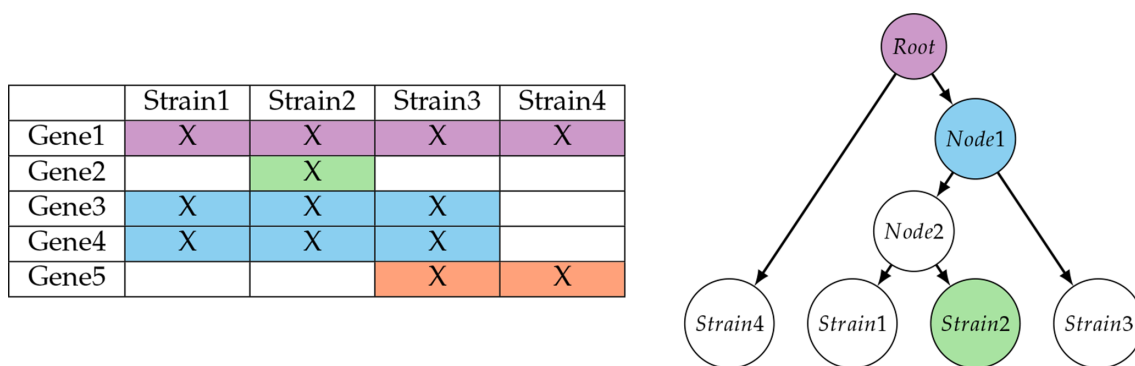
### MinION library preparation and sequencing

Libraries were prepared for MinION (ONT) sequencing according to the manufacturer's protocol ‘Rapid Barcoding Sequencing (SQK-RBK004)’ (version RBK\_9054\_v2\_revF\_23 Jan2018), starting with 400 ng of DNA per *S. thermophilus* strain, using the 12 barcodes to multiplex 12 strains at each sequencing experiment. No additional purification step on AMPure beads XP was performed. Barcoded DNAs were pooled together. Finally, a flow cell (R9.4.1, FLO-MIN106D) was loaded with 200–500 ng of the library and run for 48 h, generating around 7–9 Gbp of sequencing data. The MinKNOW software (version release 19.05.0) was used to monitor the run and generate the fast5 files. Fastq files were obtained after basecalling with Guppy (version 4.4.1, default parameters) in high-accuracy mode.

DNA of the 31 CIRM-BIA strains has previously been sequenced with NGS Illumina technology. Illumina and Nanopore data allowed us to fully and correctly assemble their genomes, using the Unicycler hybrid assembler (version 0.4.7) [10]. Complete annotated genomes have been made available on the NCBI database (accession numbers listed in Table S2).

### Filtering the raw reads

The fastq files generated by MinION sequencing were demultiplexed and adapter-trimmed using qcat (<https://github.com/nanoporetech/qcat>; version 1.1.0). MinION reads have a high error rate, in particular in homopolymers. In our sequencing data, basecalled with version 4.4.1 of Guppy in high-accuracy mode, the error rate was around 5%, including around 3% insertions/deletions. A selection step of best reads was performed in order to limit their global sequence error



**Fig. 1.** Biclusters in a strain  $\times$  gene matrix and associated labelling of nodes in a classification tree.

rate: each read must have an average quality of at least 9 and a length greater than 2000 bp (see details in Table 1). All data (raw Nanopore fastq reads and associated assembled genomes) can be downloaded on the Genouest server (see 'Data summary' section).

### Creating a pan-genomic index for the sequence characterization of species

To compare reads and genomes efficiently, they were cut into small fragments of fixed size, called k-mers, which are the basic unit of comparison. However, given the high error rate of MinION sequences, we replaced k-mers with a more sensitive pattern, the spaced seeds [34], which introduce *don't care* positions in k-mers that accept substitution errors [35]. Although genomes do not have this error rate problem, they have been indexed like the reads in order to make the two types of sequences comparable. We used a spaced seed pattern of size 15, **111111001111111**, where **1** denotes a perfect nucleotide match and **0** a *don't care* position. This means that the genomes and reads are cut into fragments (k-mers) of size 15, which are compared on the basis of the positions set to 1 in the pattern. This pattern was selected for optimal classification of the reads of JIM8332, using the iedera software [36, 37] on all the spaced seed patterns of size in the range [9:21] (see Fig. S1). The matching positions form a word called qgram. The index containing all these qgrams was built as follows: for each strain, its reference genome was cut into k-mers on which the spaced seed pattern was applied, which yielded qgrams. Each qgram was then inserted into a compact probabilistic data structure of fixed size (about  $5 \times 10^8$  bits) based on Bloom filters [38, 39].

### Data and index used for other methods

In our experiments, we adjusted the index choices for *Kraken 2* and *StrainSeeker* in order to compare the results fairly. The genome databases were exactly the same for the three methods. *StrainSeeker* requires another input, a classification tree on the whole set of genomes guiding the identification process. This tree was generated on the MicroScope platform (see next section) in our experiments.

For *Kraken 2* (version 2.0.9-beta), we used the default k-mer length of 35 and minimizer length 32. For the *StrainSeeker* (version 1.5) index, we used a k-mer length 16.

### Comparing strains

Several measures were used to estimate the similarity between two genomes, the average identity between nucleotides (ANI) in their shared coding regions (at least 70% of identity and 70% coverage of the shorter gene), and two other distances taking the whole genome into account, Jaccard and Hamming distances. We used FastANI (ManyToMany mode) to estimate the ANI distance [40]. The Jaccard distance was computed on sets of qgrams (see Equation S1). The Hamming distance  $H$  was computed based on the proportion of positions that differ between the Bloom filters of the two genomes (i.e. a Bloom filter is a vector of 0/1 and the Hamming distance is the number of positions with two different values in the vector divided by the vector length) (see Equation S1).

The MicroScope platform [41] (<https://mage.genoscope.cns.fr>; access date May 2020, v3.14.0) was used to annotate the genome of the 77 *S. thermophilus* strains, to generate a strain classification tree and to compute the pangenome. Briefly, MicroScope uses the AMIGene software to predict protein-coding genes [42] and combines the results of several tools to assign molecular functions [41]. A strain classification tree is computed using the neighbour-joining algorithm with pairwise genome distances obtained from Mash software [43]. Three sets of genes corresponding to the core, variable and strain-specific genes were downloaded from the pangenome computed by MicroScope. It is based on MICFAM gene families, which are computed using a single linkage clustering algorithm of homologous genes sharing an amino acid alignment coverage and identity above 80%. After standardizing the gene names of the pangenome families for all strains, we created a strain  $\times$  gene matrix (see Fig. 1).

### Clustering strains by gene content

In addition to the strain tree produced by MicroScope, we produced a biclustering of strains and genes, using formal concept analysis [44]. Once biclusters are determined, they



are added to nodes of the strain classification tree, showing how close the strains are with respect to their gene content. The root of the tree contains the core genome genes and the leaves contain the genes specific to this leaf (strain). Note that some nodes may stay unlabelled because they cannot be characterized in terms of a subset of genes. In our example, this is the case for *Node2* since any characterization of it would also cover *Strain3* (Fig. 1). Conversely, *{Strain3, Strain4}* is not present in the tree although it can be uniquely characterized by the presence of *Gene5*. The tree can be displayed using an online tool for the presentation of annotated phylogenetic trees called iTOL [45] and biclusters can be recovered through popups associated with each node.

## Experimentation design

To test our method and compare it to *Kraken 2* [20] and *StrainSeeker* [22], 180 strain identification experiments were performed (see Table S3). They were based on the construction of various sets of reads extracted from the MinION sequencing data by random draws performed uniformly across the filtered sequenced reads. The distribution of read lengths in the sets had the following characteristics: median 6436 bp, mean 8842 bp, standard deviation 7861 bp, for a value in the range [2000–189000]. Each identification result for a fixed set of parameter values is an average that has been calculated on five different sets of strains randomly selected from the whole set of possible strains. The parameters were the number of reads (1000, 4000 or 16000 reads), the number of strains to be identified (four or six strains), the proximity of the strains (distant, moderately close or close strains) and finally the distribution of strain abundances in the experiments (uniform distribution or distribution with dominant and subdominant strains). The set of experiments has been split into two parts to present the results, 90 with a uniform distribution and 90 with low-abundance strains. For close proximity experiments each strain was required to be very close to at least one other strain but could be distant from the others (see Table S4).

The percentage of subdominant strains was 12.5% of the reads for each of the two strains in mixtures of four strains and respectively 6.25, 3.12 and 3.12% for three strains in mixtures of six strains. For the validation of identification methods, we used the sum of Hamming distances between each predicted strain and the closest real, target one in the sample. We also calculated the Matthews correlation coefficient (MCC) to measure the precision/sensitivity trade-off of the method (see Equation S2).

Similar to *Kraken 2*, *StrainSeeker* does not always identify a single target but proposes a group of strains as an identification result. To take this behaviour into account, we counted in this calculation one true positive for each correct strain and one false positive for each incorrect strain. As a correlation coefficient, MCC ranges from 1 for a perfect prediction to -1 for complete disagreement between the observation and reality and 0 indicates no relationship. It is balanced with what we called an ambiguity ratio, which was calculated as

the ratio of the number of predicted to number of actual strains.

## RESULTS

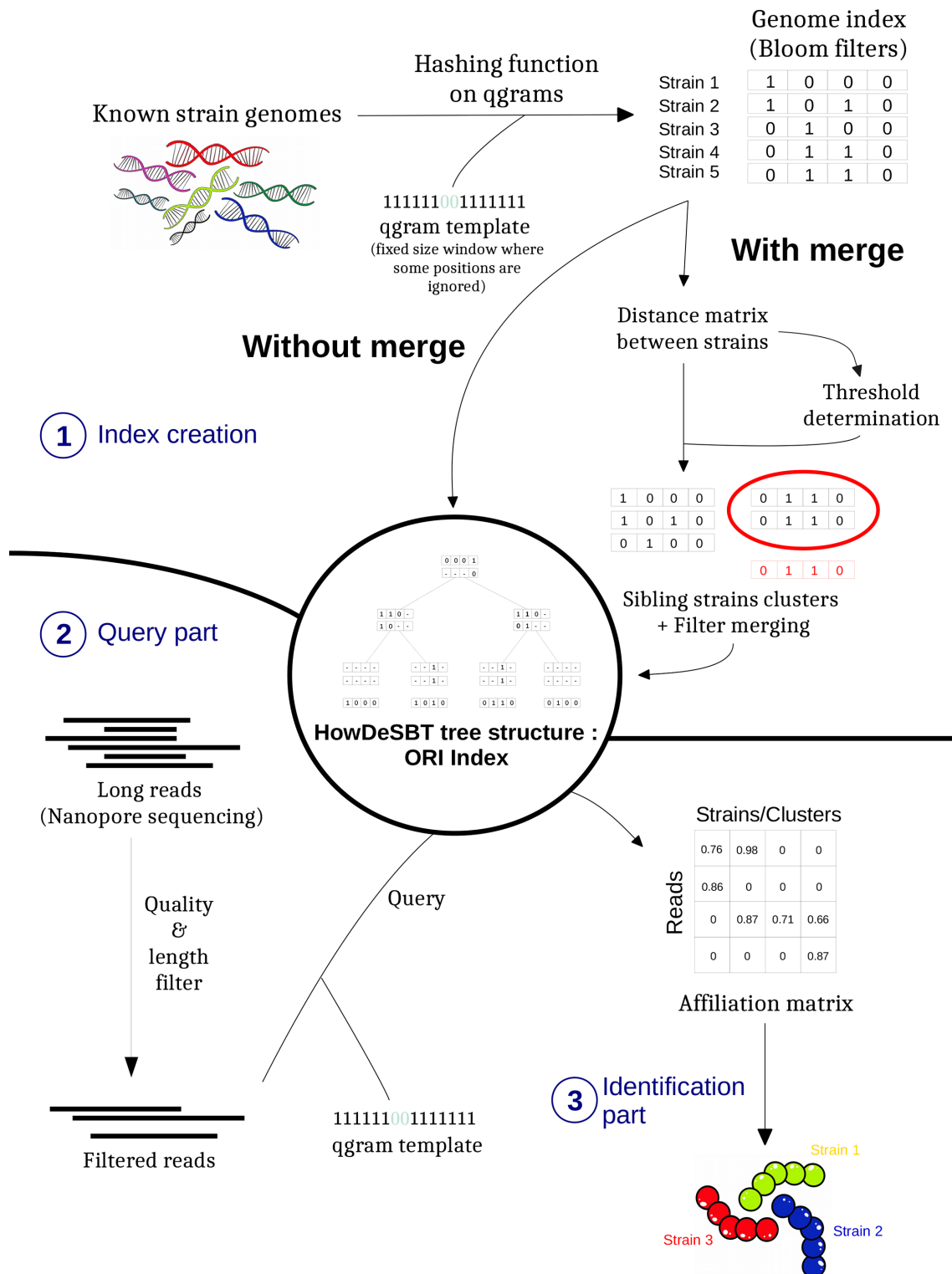
We first describe the new method we propose for the identification of strains, then the identification results themselves, which are compared to those obtained by two other methods, *Kraken 2* and *StrainSeeker*.

### ORI, a new method for the identification of bacterial strains from long reads

We propose a new method and software, *ORI*, based on indexing fragments of a set of known bacterial genomes in a compact data structure. *ORI* is therefore in line with the principles of a method such as *Kraken*, although it differs in many ways. The first step (index creation in Fig. 2) takes as input the sequences of known genomes and builds a data index based on a hierarchical structure of Bloom filters [38, 39], each one being of fixed size ( $5 \times 10^8$  bits). They code for the presence in each genome of small fragments of fixed size, called qgrams, which are similar to k-mers but allow for mismatched characters at some positions. There is an optional but recommended operation during this step, the merging of very closely related strains (distance below a given threshold) in a single pangenome (see sections below). The second step (Query part in Fig. 2) takes as input the sequenced reads of an unknown sample and consists of filtering (for length and quality), qgram extraction, and finally querying against the index. It results in a large score matrix crossing known genomes and reads, estimating for each read its probability of originating from a given genome and thus for the genome to be present in the sample. The last step is the identification process itself, which takes as input the genome  $\times$  read matrix and looks for a minimal set of strains that best explain the reads in the observed sample (see below for more details).

### Compact pan-genomic indexes of bacterial sequences

In all our experiments, two indexes were produced. The first index has been built with all *S. thermophilus* genomes (77 strains) plus two strains from other species/order, *S. macedonicus* and *L. delbrueckii* subsp. *bulgaricus*, for a total size of 23.4 Mb. Note that we have excluded plasmids in the current version since plasmids are considered part of the mobilome (horizontal transfer) and not specific of a strain. The second index was designed for experiments where very closely related isolates (such as mutant strains or variants that are indistinguishable) were merged under a single strain identifier (we call such strains ‘sibling’ strains, see section below). This has been achieved by merging all Bloom filters from closely related strains. This index had a similar size of 22.7 Mb. Both index sizes were comparable to the index of *Kraken 2* (18.1 Mb for exactly the same strains), and much smaller than *StrainSeeker*’s index (2.1 Gb).



**Fig. 2.** Overview of the ORI method in three steps: (1) genome indexing, (2) query the index from filtered reads, and (3) identification of strains.

## Affiliation of MinION reads to strain genomes by index queries

Once an index has been created, our method identifies the bacteria present in a sample by querying the index with the sample reads. As with genomes, each MinION read is cut into k-mers on which a spaced seed is applied, which provides the qgrams of the read. The index is then requested using all these qgrams. A minimum number of qgrams from a read (half of them; parameter `--threshold=0.5` in *ORI*) must be found in a strain genome before it can be retained for identification purposes. This threshold allows efficient filtering of reads that concern the species of interest. It takes full advantage of the length of the reads and allows in particular to remove contaminants, which would be difficult on short reads [46]. The output of a request is a list for each read of possible strains, weighted by their number of qgrams, summarized in a read  $\times$  strain matrix.

## Identification of one or several strains in samples

Starting from a set  $X$  of reference bacterial strains and a sequenced sample, the goal of identification is to find a subset of  $X$  that best explains the observed reads. We say that a read is explained by a strain if this strain is a possible answer to the read query in the index, i.e. it matches with a number of qgrams in the read greater than the threshold (50%).

From a theoretical point of view, this issue may be seen as a *set cover optimization problem*: find a minimal subset to fully explain the reads, a difficult problem that can be solved if the number of operational taxonomic units in the sample is of moderate size (see Text S2).

We followed a declarative problem solving approach called Answer Set Programming (ASP) for this [47]. A quick preprocessing step was applied beforehand to filter reads and strains that provided little information: (1) the reads found in too many strains were not taken into account during the identification (by default, if a read can be affiliated to 18 strains or more, they are assumed to be part of the core genome of the species) and (2) only the best strains (12 by default, parameter `--nbchoices`) were taken into account for each remaining read, according to the proportion of qgrams matching them in the read.

## Measuring the proximity between strains

To evaluate the difficulty of identification and assess its accuracy, it is necessary to measure the proximity of genomes. This has been achieved at the coarse level of genes and at the finer level of genomic content.

### The gene level

We have computed the complete list of maximal biclusters (see section *Comparing and clustering strains* in the Methods) comprising a subset of *S. thermophilus* strains and a subset of associated specific genes (available on <https://github.com/gsiekaniec/GeneTree>). Genes specific to the set of strains under each node of the strain classification tree produced by MicroScope are available in an interactive version of this

tree via clickable links for each node (see <https://itol.embl.de/tree/131254134671311597925585>). All the maximal biclusters cannot be represented by a node of the tree because they follow a different topology. It is already possible to perceive in this tree that some strains are very close if not identical and differ only by few unknown genes.

### The genomic fragment level

The proximity between genomes is usually measured with the ANI distance. Our genome index offers other ways to evaluate this proximity. We have produced a slightly more precise Jaccard distance matrix using spaced seeds. Fig. 3 shows an extract of this matrix for 28 strains in the form of a heatmap (eight strains from public genome database and 20 strains from the CIRM-BIA collection). The complete heatmap, containing the 77 *S. thermophilus* strains, can be found in Fig. S2. Two genomes, one from *S. macedonicus* and the other from *L. delbrueckii* subsp. *bulgaricus*, have been added as an external group. As expected, strains of different species or order are easily distinguished. The group of six CIRM-BIA strains at the bottom left of the heatmap that is distant from other strains clearly forms a new subgroup comprising strains that were mostly isolated from traditional Italian dairy products.

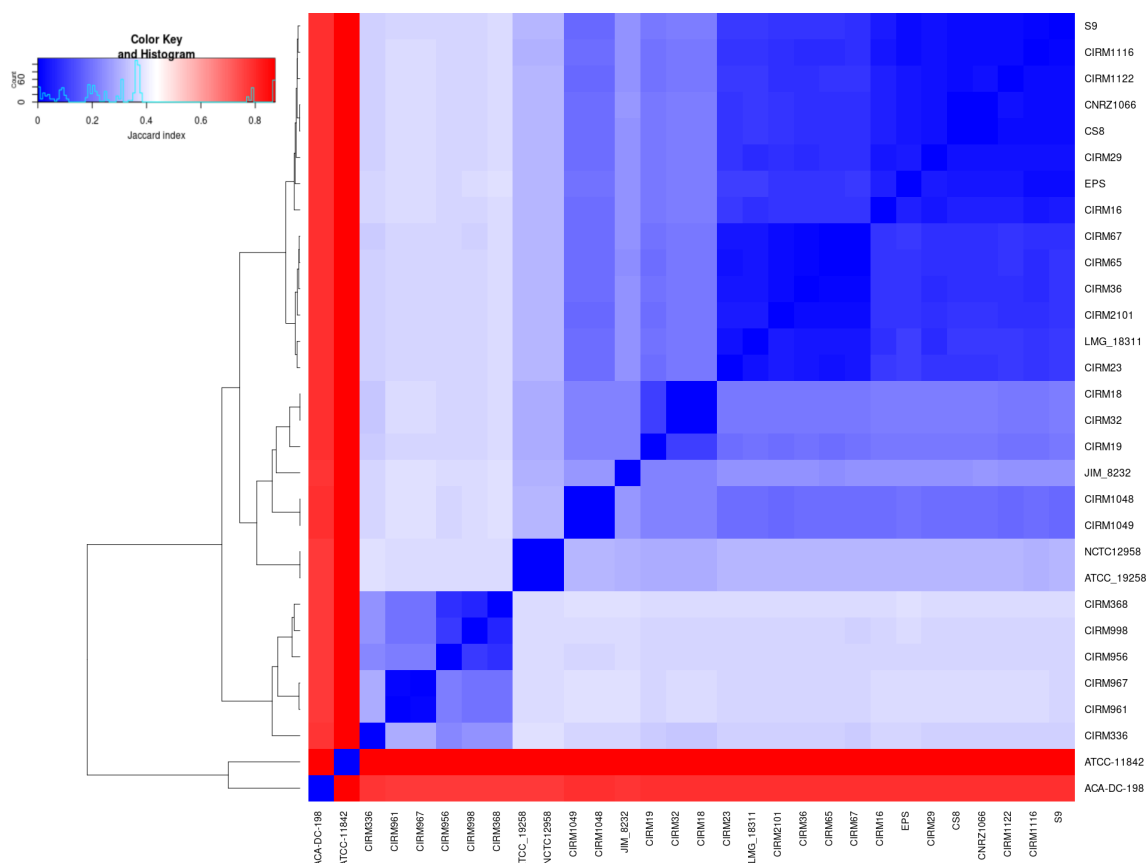
We measured the coherence of Jaccard and ANI distance with respect to the Mash distance used by MicroScope on our strain set. The observed Pearson correlation is very good ( $r=0.967$ ,  $P=1e-04$  for ANI, and  $r=0.987$ ,  $P=1e-04$  for Jaccard distance). As expected, Jaccard is slightly better than ANI distance (ANI only compares orthologous genes).

In our work, it is more efficient to compute the Hamming distance between the Bloom filters rather than the Jaccard distance (see sections *Comparing* and *Clustering strains* in the Methods). We checked that this distance corresponded to the Jaccard distance for strain comparison. There was indeed a very good correlation between the two distances ( $r=0.99$ ,  $P=1e-04$ ). Using quadratic regression, we obtained the equation  $H=3.81e^{-03}(J^2+J)$  between Hamming  $H$  and Jaccard  $J$  distance values (see Fig. S3). For small values ( $J<0.15$ ), a linear relationship fitted the values very well:  $H=4.26e^{-03}J$ .

## Sibling strains: a threshold to establish two isolates as indistinguishable

To improve detection accuracy, we grouped closely related strains into groups called 'sibling strains'. Identification results depend on their definition, which involves some threshold  $\tau$  for the maximum distance between the genome of sibling strains, much lower than that between species. Fortunately, we have shown that the choice of the distance used seems not to be crucial for this task.

In the remainder, we will compare results without or with clustering. The first results are obtained with a Hamming distance threshold of 0 ( $\tau=0$ ), and the second results are those that regroup all isolates based on a Hamming distance less than  $\tau=2e^{-4}$  (0.05 for Jaccard). This default threshold has



**Fig. 3.** Heatmap of the Jaccard distance for 28 *S. thermophilus* strains + *S. macedonicus* ACA-DC 198 + *L. delbrueckii* subsp. *bulgaricus* ATCC 11842.

been determined empirically for *S. thermophilus* by trying different threshold values and looking for a minimum value for which the identification error rate remains low on isolated strains. Below this distance, the sequencer error level prevents reliable affiliation of reads. ORI produces a graph that shows the distribution of distances between strains and helps to select a reasonable maximum distance threshold, which leads to grouping of a limited number of strains with close genomes. Clusters of sibling strains were formed as follows: (1) a graph is created whose vertices are the strains and edges connect those less than  $\tau$  away from each other; (2) maximum cliques of this graph are computed and intersecting cliques are merged; and (3) Bloom filters of the genomes belonging to these cliques are merged, corresponding to the pangenome of the sibling strains (see index creation in Fig. 2 and Table S5, Fig. S2).

### Identification results

We present in this section the results of ORI in various contexts and compare them on the same database with the leading method *Kraken 2* [20] and the program dedicated to strain identification, *StrainSeeker* [22]. As an indication, we have provided tables showing the precision of identification

of *S. thermophilus* strains in all 180 experiments (see Table S6) and the precision of identification of the subdominant *S. thermophilus* strains (see Table S7).

### Identification of isolated strains: ORI is the most robust method

The first test was to identify a single strain, *S. thermophilus* JIM 8232, from a moderate quantity of MinION reads (4000), randomly selected among all reads sequenced for this strain (around 200000 reads). This strain is quite distant from the others (see Fig. 3): it is not closely related to any other known strain. Almost all reads were retained by the three tested methods. For *Kraken 2*, 50.10% of reads were classified as JIM 8232, 26.56% of reads were classified as *S. thermophilus* (it stops at the species level) and the rest had an incorrect strain classification. For ORI, 99.7% of reads were classified as JIM 8232, the rest being unclassified. For *StrainSeeker*, JIM 8232 was recognized as the sole strain of the sample (100% of JIM 8232). Although *StrainSeeker* was developed specifically for Illumina reads, it provided excellent results with ONT reads on this test.

A second test concerned *S. thermophilus* CIRM-BIA67, which is much more difficult to identify than JIM 8232



**Table 2.** Identification of reads from *S. thermophilus* strain CIRM-BIA67 from various numbers of reads

Method	100 reads	1000 reads	10000 reads	20000 reads
<i>Kraken 2</i>	C67 2%	C67 1.70%	C67 1.31%	C67 1.23%
<i>StrainSeeker</i>	One group of many indistinguishable strains (C67 and 66 other strains) 100%	C67 100%	C67 13.13% +4 groups (other strains)	C67 2.2% +C65 1.86% +22 other strains +2 groups
<i>ORI</i>	C67 100%	C67 100%	C65 100%	C65 100%

C67, CIRM-BIA67; C65, CIRM-BIA65.

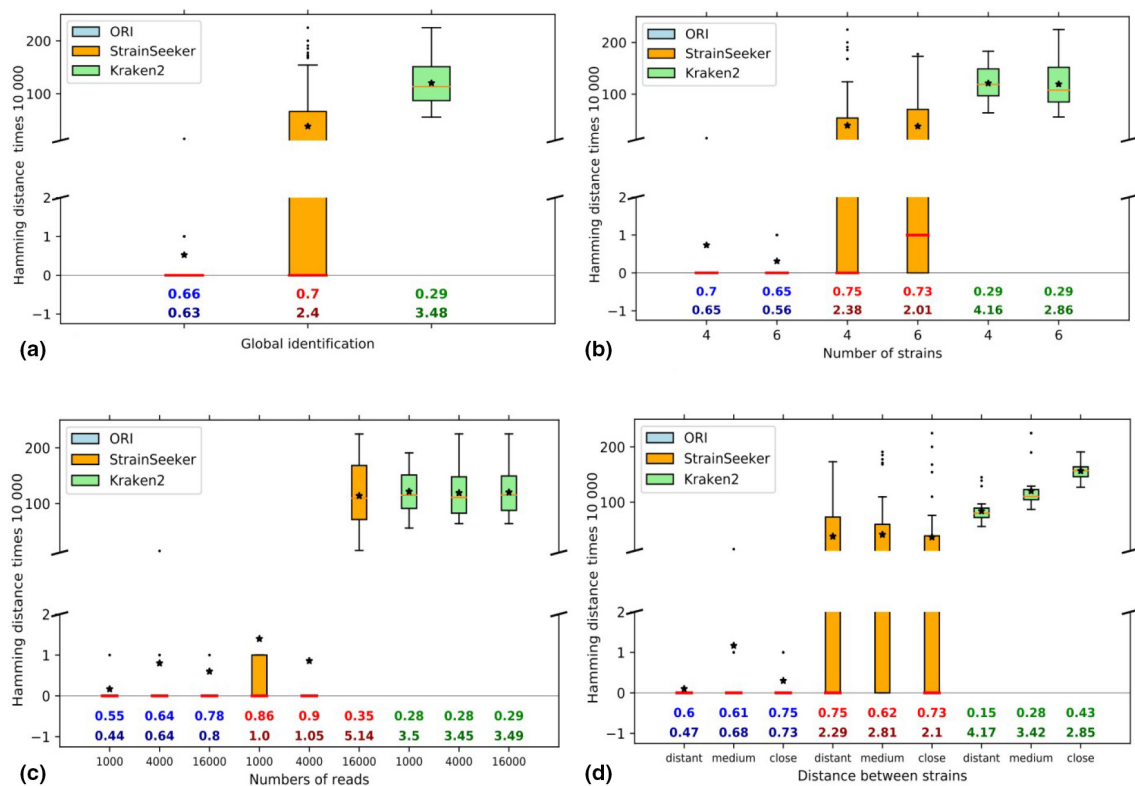
because of its proximity to other strains (see Fig. 3). In particular, CIRM-BIA67 and CIRM-BIA65 are sibling strains. The differences stem mainly from a contraction in CIRM-BIA67 of two tandem repeats from a 13.5 kb fraction of the genome. Table 2 shows detailed results for an increasing number of reads.

*Kraken 2* assigned most of the reads to the species *S. thermophilus* and less than 2% to strain CIRM-BIA67. *StrainSeeker* recognized the species level (a large group of strains that cannot be distinguished) for 100 reads, found a perfect result for 1000 reads, and then a decreasing proportion of CIRM-BIA67 for more reads. It is thus very sensitive to the

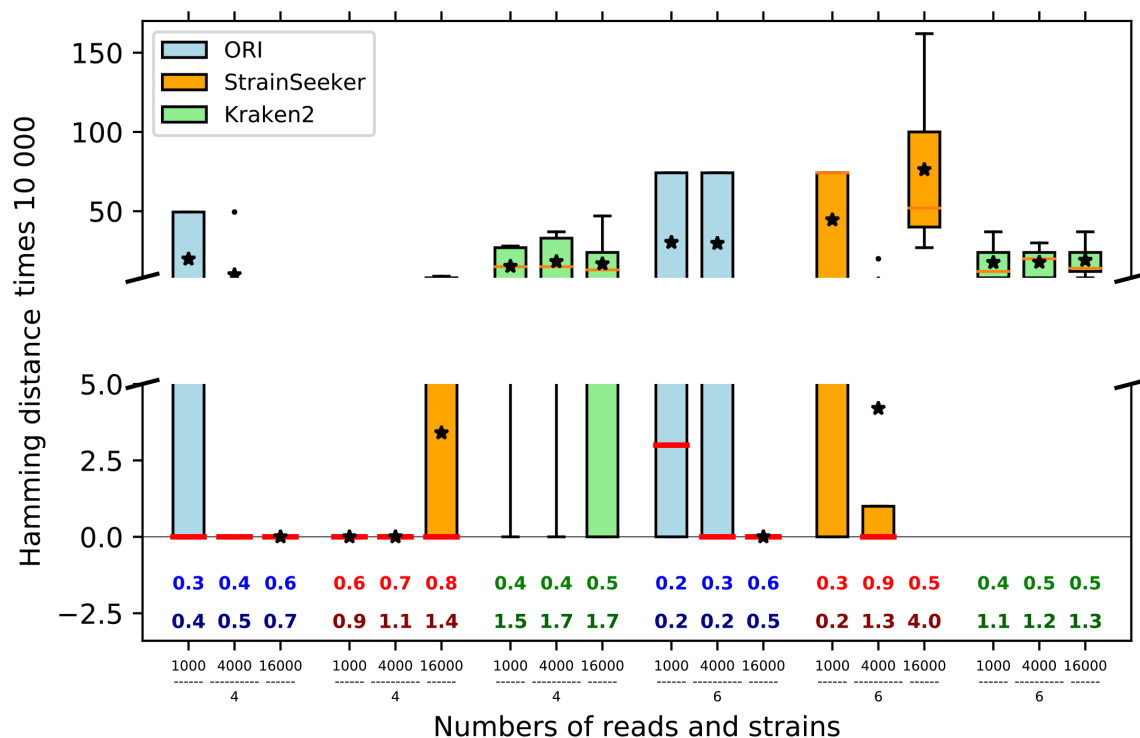
number of reads. *ORI* identification is almost perfect in each case, with identification of the correct CIRM-BIA67 strain up to 1000 reads and the sibling strain CIRM-BIA65 with more reads. For all methods, the error level of reads introduces limitations in the identification of closely related strains. For *ORI*, this can be corrected by merging them (see *Identification of strain mixtures after merging*).

#### Identification of strain mixtures: *ORI* needs at least 500 reads for each strain

We ran the 90 experiments containing a uniform number of reads for each experiment (see *Experimental design* in Methods).



**Fig. 4.** Identification results on a balanced mix of *S. thermophilus* strains. The Hamming distance between observed and expected strains, on the y-axis, has been multiplied by 10000 (in blue for *ORI*, orange for *StrainSeeker* and green for *Kraken 2*). Stars represent mean values. Matthews correlation coefficient (MCC) values are given on the first line just above the x-axis at the bottom of the diagrams, followed by the ambiguity ratio (number of strains identified/number of strains present).



**Fig. 5.** Identification of subdominant strains in a mixture of *S. thermophilus* strains using various numbers of reads. The Hamming distance between observed and expected strains, on the y-axis, has been multiplied by 10000 (in blue for *ORI*, orange for *StrainSeeker* and green for *Kraken 2*). Matthews correlation coefficient (MCC) values are given on the first line just above the x-axis at the bottom of the diagrams, followed by the ambiguity ratio (number of strains identified/number of strains present).

The diagrams in Fig. 4 display boxplots for the sum of the Hamming distances showing how close the predicted strains are from real ones. Below each boxplot, two numbers are given, MCC and the Ambiguity ratio. The MCC, which measures the adequacy of the binary decision on the presence/absence of strains, shows the balance between specificity and sensitivity. The ambiguity ratio is necessary because *Kraken 2* and *StrainSeeker* propose ambiguous answers with more strains than actually exist, and it tends to artificially increase the MCC, whereas *ORI* tries to minimize the number of predicted strains.

We computed the Hamming distance, the MCC and the ambiguity measures over all experiments (global identification, Fig. 4a), then detailed the results along three parameters: the number of strains, the number of reads and the proximity of the strains (Fig. 4b–d). Overall, identification results are very good for *ORI* (Fig. 4a), from the point of view of both mean and standard deviation, showing the robustness of the method. *StrainSeeker* also gives good results and has the best MCC but is penalized by the production of multiple solutions and wider variations than *ORI*. *Kraken 2* is clearly less well adapted to the recognition of strains.

Fig. 4b shows the effect of increasing the number of strains in the mixture. Results are very similar for four or six strains.

For Fig. 4c, different amounts of read data were used. No difference was observed for *Kraken 2*, using 1000, 4000 or 16000 reads. By contrast, *ORI* seems to be sensitive to read number: the more data, the better strain identification with regard to MCC and ambiguity ratio. For *StrainSeeker*, using a large number of reads clearly leads to a drop in accuracy and results are optimal with 4000 reads.

Finally, the resolution power of methods was measured for mixtures containing increasingly close target strains (increasing the difficulty of identification). Fig. 4(d) shows that *Kraken 2* is sensitive to this parameter and provides poor results for a mixture of closely related strains. With *ORI* and *StrainSeeker*, which are more sensitive, the effect is not marked but *ORI* shows a slight degradation of MCC and the ambiguity ratio. This behaviour, due to the existence of *S. thermophilus* genomes with high proximity, motivated the introduction of a merging procedure prior to strain identification (see next section).

We end with the most difficult context, the identification of subdominant strains in a mixture of four or six strains (Fig. 5). The mixture of four strains contains two dominant strains (representing respectively 50 and 25% of the total reads each) and two subdominant strains (12.5% of the reads each), while the mixture of six strains contains three dominant strains (50/25/12.5% respectively) and three

**Table 3.** *S. thermophilus* strain identification by *ORI*, with and without merge index, in a balanced mixture of four or six strains more or less genetically close, by using 1000, 4000 or 16000 sequencing reads

Best results are in bold type. Values of Hamming distance (0=perfect identification); MCC: Matthews correlation coefficient (1=perfect correlation); Ambiguity: number of strains identified/number of strains present.

(a) Global identification results (mean over all 90 experiments):						
Method	ORI		ORI_merge			
Distance	0.52	0.41				
(MCC/Ambiguity)	0.66/0.63	0.92/0.91				
(b) Heterogeneity, mean results (variable number of strains mixed):						
Method	ORI		ORI_merge			
Number of strains	4	6	4	6		
Distance	0.73	0.31	0.53	0.29		
(MCC/Ambiguity)	0.70/0.65	0.65/0.56	0.94/0.93	0.96/0.96		
(c) Data quantity, mean results (variable number of .fastq reads):						
Method	ORI		ORI_merge			
Number of reads	1000	4000	16000	1000	4000	16000
Distance	0.17	0.8	0.6	0	0.43	0.8
(MCC/Ambiguity)	0.55/0.44	0.64/0.64	0.78/0.80	0.86/0.77	0.93/0.92	0.98/1.05
(d) Resolution power, mean results (variable proximity between strains within the mixture):						
Method	ORI		ORI_merge			
Proximity	Distant	Medium	Close	Distant	Medium	Close
Distance	0.10	1.17	0.30	0	0.90	0.33
(MCC/Ambiguity)	0.75/0.73	0.61/0.68	0.6/0.47	0.93/0.89	0.87/0.85	0.97/1

subdominant strains (6.25%/3.125/3.125% respectively). Of note is that *Kraken 2* behaves better in this situation, reducing the difference compared to *ORI* and especially *StrainSeeker*. *ORI* continues to perform well for strains with at least 500 reads but its results are degraded below this. For *StrainSeeker*, the number of reads used for identification is crucial: it works well with 4000 reads (even better than *ORI*) but works surprisingly poorly with 16000 reads.

#### Identification of strain mixtures after merging highly similar strains: best results for *ORI*

As previously observed in Fig. 5, the main issues for *ORI* come from very closet related isolates. We have tested the impact of preprocessing the data by merging them as one strain type. The results obtained with this new merged index were compared to the previous results (Tables 3 and 4).

**Table 4.** Subdominant *S. thermophilus* strain identification by *ORI*, without/with merge, in a mixture of four or six strains, by using 1000, 4000 or 16000 Nanopore sequencing reads

Best results are in bold type. Values of Hamming distance: in all experiments, minimum value is 0 (perfect identification); MCC: Matthews correlation coefficient (1=perfect correlation); Ambiguity ratio: number of strains identified/number of strains present; sd: standard deviation

No. of strains	4 (ORI/ORI_merge)			6 (ORI/ORI_merge)		
Number of reads	1000	4000	16000	1000	4000	16000
Distance	19.8/19.8	9.9/0	<b>0/0</b>	30.3/15.4	26.7/0	<b>0/0</b>
MCC	0.28/0.38	0.42/0.78	0.57/0.9	0.22/0.38	0.34/0.65	0.63/0.8
Ambiguity	0.4/0.4	0.5/0.8	0.7/1	0.2/0.33	0.2/0.47	0.53/0.67

Overall, merging leads to almost perfect identification results, with a Hamming distance that decreases with respect to the version of ORI without merging and it is mostly null, except for experiments with an insufficient number of reads. The MCC increases with the number of available reads, and is close to 1 on average, showing that ORI with merge combines high accuracy with high sensitivity. Note that it is possible to get a null average distance and a MCC less than 1; this points to perfect accuracy but loss of sensitivity: strains are identified but some are lacking.

The method is more robust in the sense that the variations in parameter values have less influence on the results.

The identification of subdominant strains is shown in Table 4 (full results can be found in Tables S8 and S9). The results are clearly better with merging for samples of 4000 reads; in our experiments, merging sibling strains allowed us to achieve strain identification from as few as 250 reads. For experiments with 1000 reads, which need a recognition from 125 reads or fewer, the method is limited and the gain is smaller.

## DISCUSSION

In this work, we have compared three methods that have a common methodological basis. They split genomes and reads into small fragments that are inserted in an index, a very compact and efficient data structure for making comparisons. However, ORI seems to be better than *Kraken 2* in terms of identifying bacterial strains. *StrainSeeker* behaves correctly with parameters carefully tuned. It has been designed for short reads of high quality and it will return poor results either if there are too few or too many reads. Moreover, *StrainSeeker* does not scale to a large number of genomes, and its index requires 100 times more memory than the other methods. ORI features three major choices adapted to strain-level identification from long read noisy data. A first one is to replace k-mers by qgrams, which tolerate a few substitution errors, and is particularly useful given the higher error level of ONT sequencing. Note that it is effective for relatively short k-mers since indel errors are not taken into account. Another important difference relates to the choice of the tradeoff to be made between the sensitivity and the computational cost of the method. *Kraken 2* is an excellent choice for large-scale species recognition, based on the use of minimizers. This choice leads to a relative loss of sensitivity which does not allow it to recognize closely related strains. ORI uses a first identification step for each read, utilizing the fact that all the k-mers in a long read must belong to a single species. A last difference is that ORI uses an exact optimization step in order to select a minimum number of species/strains that explain most of the reads. This is a unique point of ORI that increases its robustness (very few false positive) and also reduces the list of identified strains, which can be very useful for subsequent analysis of the results.

There is a need to achieve ID resolution of a group of highly similar isolates [48], rather than the exact isolate itself. Clinically it is useful for identifying outbreak strains, which is a major use of MinION sequencing (error-prone Nanopore reads). In a

recent paper [49] the authors worked on almost 2000 samples of *Streptococcus agalactiae*, characterized by different degrees of virulence and preferred host. However, even if the microbiologists are demanding regarding the distinction of strains, they only distinguish a few dozen different types at most. Stopping strain distinction at the individual isolate level is of little practical value. Another recent application of clustering close strains is in the inference of antibiotic resistance and susceptibility [50]. We have presented two versions of ORI, one including a preprocessing step merging in a single pangenome the sequences of known isolates that appear to be very similar. Our results showed that the version that merges seems the most appropriate with regard to the high error rate of current ONT sequencing technology. The identification results are robust and the biologist may conduct additional investigations to discriminate the strain among a limited number of possibilities. Considering these sibling strains in the iTOL tree, they are annotated mostly by specific genes of unknown function, contrary to other strains (e.g. *S. thermophilus* JIM 8232). Two unknown genes labelled differently could nevertheless be related and sibling strains, and therefore could usefully point to groups with a compact pangenome, analysable in fine detail, valuable knowledge for strain selection. An interesting perspective would be to check that merging allows us to distinguish strains for important phenotypic traits such as antimicrobial resistance profiles [51].

Another important parameter is the number of reads to be used for identification. We proposed a set of 4000 reads to be sufficient for a mixture and 500 reads for an isolate, something that helps in fast identification of the content of a sample. In fact, it seems that a good information/noise ratio is needed in practice: either the number of reads is insufficient to detect the accessory genome, or it is too high and noisy reads tend to introduce ghost recognitions. We still need to work on this aspect so that ORI does not decrease its identification quality when the number of reads increases. As identification requires a good information/noise ratio, one way to improve it is to pre-filter the reads and strains in order to obtain the fewest erroneous and most strain-specific reads.

We end this section with a few perspectives that could help extend this work.

### Rapid identification at low cost

In our experiments, DNA has been extracted by a kit specific for long reads, using the ONT flow cell R9.4 for sequencing. Such DNA extraction requires long passage times through the column by gravity (6–8 h, depending on the strain) and is relatively expensive, including all costs. In practice, it is crucial to decrease costs and time as much as possible while maintaining ORI's good performance. Using a DNA extraction kit designed for short reads would lead to fragments three times shorter but save time and money (4 h extraction, 5 € per sample). A good compromise could be a rapid extraction with magnetic beads (e.g. MagAttract HMW from Qiagen). For the library sequencing kit, we used a rapid barcoding sequencing kit, which reduces size by a factor of 3 and time by a factor of 20 (15 min) compared to the classic SQK-LSK109 kit. Once loaded



with 12 samples, the flow cell produced about 3 million reads (9 Gb of sequences) after 48 h of run. In fact, less than 10 min of sequencing is sufficient to produce the 4000 reads necessary for strain identification. Furthermore, ONT suggest the low-cost, low-throughput Flongle flow cell that could be an interesting alternative for producing the required data in about 30 min. Overall, from sample collection (e.g. a fermented food product) to sequence files, for one sample containing four strains, requires less than 6 h (for a total cost of maximum 200 €, using Flongle). Then, depending on the processor used, identification could be achieved within 1 h.

### Quantification of strains in a mixture

We plan to estimate strain abundances in *ORI* by using an expectation-maximization (EM) algorithm similar to the one used in *Centrifuge* [17]. Users already have access to indicative abundance values in conjunction with strain classification to help interpret the identification results. For instance, consider a sample that contains 16 000 reads of six *S. thermophilus* strains equitably distributed and making two clusters of sibling strains: A={CIRM-BIA18, CIRM-BIA32} and B={CIRM-BIA2101, CIRM-BIA23, CIRM-BIA65 and CIRM-BIA67}. Thus, the sample comprises 1/3 of A and 2/3 of B. For this sample *ORI* found 25% of A, 72% of B and 3% of a third cluster C={CIRM-BIA1116, CIRM-BIA1122, CIRM-BIA16, CIRM-BIA29, CNRZ1066, CS8, EPS, S9} (MCC=0.807). Since the abundance of C is quite low and it is quite close to cluster B (see Fig. 3), it is reasonable to assume that the reads originated in fact from B. By increasing the merging threshold  $\tau$ , B and C would indeed be merged, leading to perfect identification.

### Towards a generalization to other species and genera?

The index can be tailored to any set of bacterial genomes by modifying the genome index, as explained on the website. We tested our method on *Streptococcus pyogenes* (around 200 genomes in NCBI; some closely related and mostly different by their phage content). A known *S. pyogenes* emm75 strain of our collection was correctly identified by *ORI*. We also used 1600 complete genomes to create a specific *E. coli* index. Reads were from a piglet gut microbiota. This gave promising results in about 5 h: all the reference strains identified were also from a piglet gut microbiota. The next challenge is to test *ORI* on a larger number of species in order to fit users' study species. As a first step, we plan to increase our database to the whole order *Lactobacillales*. We aim at keeping good accuracy and sensitivity for identification at the strain level while staying efficient with respect to memory and computation time. This will require some modifications in the means of requesting the index which, currently, remains the most time-consuming part.

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### Author contributions

Conceptualization, J.N.; methodology, G.S., J.N. and T.L.; software, G.S. and T.L.; validation, G.S.; resources, E.R.; data curation, G.S. and E.R.; writing—original draft preparation, J.N., G.S. and E.R.; writing—review and editing, J.N., E.R., G.S. and E.G.; supervision, J.N., E.R. and E.G.; project administration, J.N.; funding acquisition, J.N. and E.G. All authors have read and agreed to the published version of the manuscript.

### Conflicts of interest

The authors declare that there are no conflicts of interest.

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