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New genetic biomarkers to differentiate pathogenic and clinically relevant *Bacillus cereus* strains

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Keywords

Bacillus cereus, pathogenicity, genetic biomarkers

Abstract

Objectives. *Bacillus cereus* is responsible for food poisoning in France and rare but severe clinical infections. The pathogenicity of strains varies from harmless to lethal strains. However, there are currently no markers, either alone or in combination, to differentiate pathogenic from non-pathogenic strains. The objective of the study was to identify new genetic biomarkers to differentiate pathogenic from clinically relevant *Bacillus cereus* strains.

Methods. A first set of 15 *B. cereus* strains were compared by RNAseq. A logistic regression model with lasso penalty was applied to define combination of genes whose expression was associated with strain pathogenicity. The identified markers were checked for their presence/absence in a collection of 95 *B. cereus* strains with varying pathogenic potential (FBO, clinical and non-pathogenic). ROC-AUC analysis determines the combination of biomarkers, which best differentiate between the “disease” versus ‘non-disease’ groups.

Results. 7 genes were identified during the RNAseq analysis with a prediction to differentiate between pathogenic and non pathogenic strains. The validation of the presence/absence of these genes in a larger collection of strains coupled with AUC prediction showed that a combination of 4 biomarkers was sufficient to accurately discern clinical strains from harmless strains, with an AUC of 0.955, sensitivity of 0.9 and specificity of 0.86.

Conclusions. These new findings help in the understanding of *B. cereus* pathogenic potential and complexity and may provide tools for a better assessment of the risks associated with *B. cereus* contamination to improve patient health and food safety.

Introduction

Bacillus cereus is the third causative agent of food-borne-outbreaks (FBO) in Europe [1]. *B. cereus* can induce two types of gastrointestinal diseases, leading to generally mild and self-limiting emetic or diarrhoeal syndromes, although several cases of severe infections have been reported [2]. *B. cereus* also induces systemic infections leading to patient death in approximately 10% of cases [3-7]. *B. cereus* is also a source of central nervous system infections and other systemic infections especially in newborns [3, 8]. Recent epidemiological studies show that the number of cases of serious *B. cereus* infections is largely underestimated [9]. The pathogenic potential of *B. cereus* is extremely variable, with some strains being harmless and others lethal.

B. cereus possesses several toxin genes, such as *nhe*, *hbl* and *cytK* [2, 10]. These toxins provide an indication of the strain toxicity potential but are not sufficient, alone, to discriminate hazardous from harmless strains [9, 11-13]. Indeed, several studies have shown that *Nhe* production by hazardous strains is variable and that non-pathogenic strains can also produce it in large quantities [1, 12]. Moreover, these toxins do not appear to be suitable markers for strains causing non-gastrointestinal infections [9]. *B. cereus* produces other toxins such as haemolysin II (HlyII), the metalloproteases *InhA1*, *InhA2* and the cell wall peptidase FM (CwpFM), which may also be involved in pathogenicity [14-18]. The emetic form of *B. cereus* food poisoning is caused by the peptide cereulide [19], which represent less than 1% of the FBO strains of *B. cereus* [1, 19, 20].

To date, the above described determinants were not sufficient to completely explain the virulence of *B. cereus* [21] and there are currently no markers, either alone or in combination, to differentiate pathogenic from non-pathogenic strains. In this work, we took advantage of a well characterized collection of 95 *B. cereus* strains and compared pathogenic (FBO and clinical) with non-pathogenic strains. We identified a combination of four as yet undescribed biomarkers, wherein their presence/absence allows an accurate identification of clinical *B. cereus* strains. These findings constitute a huge step in the understanding of the *B. cereus* pathogenic potential and complexity and may provide tools to better assess the risks associated with *B. cereus* contamination.

Methods

Isolate information

This study includes 39 *B. cereus* strains associated with foodborne illness [1], 35 strains isolated from human patients following systemic or local infections [9] and 21 non-pathogenic strains [11, 22] (Sup Table 1). We have previously shown a correlation between cytotoxicity and virulence [21]. Nevertheless, although these strains had previously been shown to be weakly cytotoxic to human cells and to have reduced virulence in an insect infection model, this does not rule out their potential ability to produce symptoms in specific vulnerable populations.

RNA extraction

The transcriptome study by RNAseq was carried out on 15 strains representative of the three collections (Sup Table 2) in triplicates. Bacterial cultures were incubated in BHI medium at 30°C in microaerophilic condition (5% O₂–15% CO₂–80% N₂) at pH 7 until entry into stationary growth phase. Samples were centrifuged at 12,000 g for 3 min at 4°C and placed immediately at -80°C until processing. The bacterial pellets were re-suspended with 200 µl of 10 mM Tris-HCl at pH 8 + 4 µl of lysozyme at 50 mg/ml and incubated at 37°C. Total RNA was extracted with the HPRNA kit (High Pure RNA Isolation Kit; Roche) as previously described [23]. The RNA integrity was measured by the RIN (RNA Integrity Number) and were between 7 and 10. The mRNA were enriched with the RiboZero Kit (Illumina). The sequencing of the mRNA was carried out by the I2BC platform (CNRS, Gif-sur-Yvette). Directional and paired libraries were prepared with the Illumina scriptseq kit and the sequencing was performed on an Illumina Nextseq machine.

Transcriptome sequencing analysis

Sequencing quality was assessed using FastQC, and adapter sequences and low-quality base pairs were removed using cutadapt (version 1.9) [24]. Reads were further trimmed in 3' using sickle (version 1.33, option “-x” and default values for all other parameters, implying a Phred quality cutoff of 20). In absence of whole genome sequences for the 15 strains, the cleaned reads were mapped against a repertoire of allelic variants for 23,815 genes aiming at accounting for the pangenome of *B. cereus* group. This repertoire was obtained by single-linkage clustering based on the results of an all-against-all blastn comparison (version 2.2.26, e-value cut-off 1e-5) [25] of 519,931 CDSs extracted from the 91 annotated complete genomes available at the time of

analysis for *B. cereus* group in Genbank. Pairs of CDSs that aligned over at least 70% of the length of the shortest sequence and with at least 75% nucleotide sequence identity were grouped in the same cluster, which resulted in 23,815 clusters representing distinct genes. Reads were mapped using bowtie2 (version 2.2.6, options “-N 1 -L 16 -R 4”) [26] whose results were converted to bam format using SAMtools version 1.9 [27]. Read counts on each allelic variant were obtained using HTSeq-count (version 0.6.1) [28] and summed over allelic variants to obtain a single read count per gene per sample. To cope with sequence similarity between allelic variants of a same gene and fragmentation of the reference according to gene boundaries, R1 and R2 reads were aligned independently and use of HTSeq-count option “-a 0” allowed to count reads that aligned equally well on several allelic variants of a same gene. Of note, since bowtie2 mapped each read on a single allelic variant, reads could not be counted more than once in the sum. Expression levels expressed as \log_2 scaled rpkm (reads per kilobase per million mapped reads) were produced by the R package “edgeR” (version 3.11) using the mean length of the genes in the cluster and a prior count of 1.

Raw transcriptomic data and differential expression analysis are accessible through GEO Series accession number GSE168681

(<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE171128>).

Statistical model

The strategy for statistical analysis of RNAseq data was to select genes to predict whether a strain is pathogenic $y=1$ or not $y=0$ and evaluate the prediction accuracy. We considered the logistic regression model with lasso penalty implemented in the R-package “glmnet”, which allows the selection of a limited subset of genes whose expression is associated with strain pathogenicity [29]. The package glmnet provides an interval cross validation procedure to select the penalty constant, which determines the number of selected genes.

The prediction accuracy of the procedure was evaluated in a cross-validation framework where splitting in training and validation sets preserves the matching of the three replicates of each strain. For each replicate, the model provides a probability \hat{z}_i to be pathogenic, and we considered the average value over the three replicates as the prediction probability of the strain. The predicted pathogenicity status is set to zero if the prediction probability is smaller than 0.5 and 1 otherwise.

Biomarker screen by PCR

The 7 marker genes were retrieved from at least 20 sequenced *B. cereus* strains from NCBI databases and aligned by CLC Main workbench7 software to identify two regions conserved across the strains. Within these regions, 20 bp primers were designed using the Beacon Designer software. For the majority of the selected genes there were no perfectly conserved sequence and some bases had to be replaced with R (A/T), Y (C/T) or W (A/T) for primer design (Sup Table 3). For all the strains of the collection, a single colony was picked, resuspended in 100 µL Tris-EDTA NaCl buffer (TEN) and incubated at 98°C for 10 min. After centrifugation, 1 µL of supernatant was used as DNA matrix. The PCR mixture contained 1 µL DNA matrix, 0.5 µM primer (forward and reverse), 10 µL DreamTaq Green PCR Master Mix (2X) (Thermo Scientific) in a final volume of 20 µL. PCR fragment sizes were revealed on 1.5% agarose gels containing Midori Green, and visualised by a UV imaging device.

AUC analysis to select combinations of biomarkers

The PCR data were pooled into a presence (1) /absence (0) table, which was then used as input for ROC-AUC analysis facilitated by the web-based suite of tools hosted at www.combiroc.eu. The ROC-AUC analysis determines the combination of biomarkers, which will best differentiate the classes of samples input ('disease' versus 'non-disease' groups). Sets of biomarkers were selected based on their performance in sensitivity or specificity alone, or in combination as the AUC metric. Potential hits were filtered at 85% specificity and 85% sensitivity.

Results

RNAseq analysis

We obtained between 9-15 million reads per samples with 90% correctly paired. The overall alignment rate was over 85%. The analysis enabled the creation of a read counts table based on gene expression levels for each sample (Figure 1). The dispersion of the sample count values was homogeneous and the biological triplicates clustered well together. We identified 3276 genes in the core transcriptome, which represents approximately 65% of the genes in each strain.

Identification of 7 biomarkers by logistic regression analysis

A Mann-Whitney-Wilcoxon nonparametric rank test with a classical 5% of qvalue did not allow the prediction of significant differences in gene expression among the strain collections (not shown). Thus, to identify markers that could potentially differentiate pathogenic from non-pathogenic strains, we performed a penalized conditional logistic regression with the lasso method on the entire counting table to select relevant genes for the prediction of pathogenic potential. By applying the prediction model to the 11,179 genes with the selected penalty constant of 0.01, only 7 genes were selected (Table 1).

With the RPKM values of these 7 genes (Sup Table 4), a prediction in a cross-validation framework among the 15 strains, leads to 13 well classified strains (estimated probability \hat{z}_i value below 0.5 for non-pathogenic and above 0.5 for pathogenic strains) and two misclassified strains, one false positive (NP strain PF predicted as pathogenic) and one false negative (pathogenic FBO strain 12CEB01BAC predicted as NP) (Table 2).

Validation of the biomarkers on a large strain collection

Initially, for the first 15 strains, the presence of the 7 selected genes was further assed by PCR (Table 3). These data revealed that when a gene showed no expression by transcriptomic analysis, the gene was actually absent from the strain. Thus, the identification of these 7 biomarkers was based on gene presence/absence, rather than mRNA expression. As such, an approach centred on gene detection was chosen for the screening of the large bacterial collection with the 7 genes selected (Table 3) and to determine the area under the curve (AUC), specificity, and sensitivity of possible combinations of the selected biomarkers.

1-FBO vs NP

For the FBO strains, the best combination of biomarkers able to differentiate NP from FBO strains was obtained with 4 biomarkers (Figure 2A). With this combination, the best AUC was 0.768, the sensitivity 0.69 and the specificity 0.773. Therefore, we obtained some false positive (NP strains that appear pathogenic), and some false negative (FBO strains that appear NP). Taken together, the general trend for the FBO identification was an overall low AUC among the tested combinations, thus preventing their accurate differentiation.

Nevertheless, we identified that several FBO strains were lacking almost all biomarkers. These FBO strains primarily belong to the phylogeny group IV (table 3). We thus performed an additional

AUC analysis after the removal of all strains of the phylogeny group IV of the collection (FBO and NP). The results were significantly improved and the best combination resulted in an AUC above 0.9 and with significantly improved sensitivity or improved specificity. But a combination resulting in sensitivity and specificity above 0.9 was not determined (Figure 2B).

2-NP vs clinical strains

Regarding the clinical strains, the best results were achieved with a combination of 4 biomarkers with an AUC of 0.955, sensitivity of 0.9 and specificity of 0.86. Therefore, the analysis concludes that an accurate differentiation between clinical and non-pathogenic strains can be obtained by using these biomarkers (Figure 2C). These two combinations allowed the accurate discrimination between the two strain populations. Some markers have the same occurrence within the strain collection (5, 6, 7) and were therefore interchangeable during the AUC analysis. Thus, the best combinations of biomarkers are: 1, 2, 3, 5 (or 6 or 7). The genes are named, *adhB*, *agrC*, *thiJ*, *BCQ_PI180* (or *gshAB* or *BCQ_PI181*).

As a conclusion, a suitable combination of 4 biomarkers has been found to create a robust and accurate test to differentiate clinical from non-pathogenic strains, with an AUC of 0.955, given that test results above 0.9 are considered excellent.

Discussion

The emergence of *B. cereus* as a foodborne pathogen and as an opportunistic pathogen has intensified the need to distinguish strains of public health concern. The pathogenic potential of *B. cereus* is extremely variable, with some strains being harmless and others lethal. Currently, due to the lack of validated and standardized analytical methods, only the presence of *B. cereus* is usually investigated in foods or clinical samples at a species-level. Over the years, new methods have been developed with the leading principle to detect and distinguish *B. cereus* from others *Bacillus* group members by a time-saving and *in-situ* analysis [30], genotyping using high-resolution melting analysis [31], the use of multi-locus sequence (MLST) [32] or the classification of the strains according to their affiliation to a phylogenetic group that offers a first useful indicator of risk [11]. Nevertheless, MLST analysis of the 53 strain sequences included in this study revealed that 21% belonged to the sequence type ST26, and approximately 11% to an undetermined ST (not shown), while >40% of the strains were identified as belonging to PanC

clade III (Table 3). As such, the ST types and PanC classifications were unable to completely explain the grouping of the strains.

Here, we report new markers characteristic of pathogenic *B. cereus* strains, which detection requires only PCR, and is thus independently of growth conditions. We could indeed show that the simple presence/absence of the gene was as discriminant as its expression value by transcriptomic analysis. We further calculated the AUC, specificity and sensitivity obtained using the combination of these 4 biomarkers to discriminate between our large *B. cereus* collection inducing various pathologies. CombiROC results demonstrate that clinical strains were more efficiently separated from the non-pathogenic strains than the FBO strains.

Regarding the FBO strains, to improve the analysis, strains belonging to the phylogenetic group IV were removed, thus allowing a significant improvement in strain differentiation. This might prove very useful for food industries to better communicate the risks of *B. cereus* food contamination and to take the appropriate measures for decontamination while preventing or minimizing economic loss. Nevertheless, this implies a two step-test with a first *panC* phylogenetic attribution followed by a biomarker test.

By contrast, regarding the clinical strains, the combination of 4 biomarkers allowed the identification of a strong differentiation test with an AUC of 0.955, sensitivity of 0.9, and specificity of 0.86. Thus, a global test with a strong AUC (above 0.9) and increased sensitivity (rare false negative) could be proposed to accurately discriminate between clinical and harmless strains. As such, our new findings may be relevant to gain additional knowledge on the strains found in hospitals and healthcare settings.

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COI statement

The authors declare no conflict of interest.

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Author Contributions

DK, BG, RD: performed experiments, analyzed data, manuscript writing; CG, SP, PN: analyzed data; SH, AB: supervision; NR: initial concept, supervision, analyzed data, writing of manuscript, funding sources.

Legends of figures and tables

Figure 1. RNAseq heatmap. Heatmap representation of expression levels (\log_2 rpk) across the pangenomic repertoire of 23,815 genes (rows) and the 45 samples (columns). Dendrograms are built by hierarchical clustering with average-link. The 3,272 genes with signal in all strains are indicated by grey bars. Non-pathogenic strains are indicated in black and pathogenic strains in red.

Figure 2. CombiROC analysis results. The presence/absence matrix resulting from PCR detection of biomarker sequences was analyzed by CombiROC. (A) Foodborne outbreak strains (FBO) versus non-pathogenic; (B) FBO versus non-pathogenic strains, excluding phylogenetic group IV. Links best sensitivity performance, right highest specificity; (C) clinical versus non-pathogenic strains.

Table 1. List of 7 selected biomarkers with gene position (on the reference genome pAH187_270 - NC_011655.1) and putative function.

Table 2. Estimated probability \hat{z}_i for the 15 strains. A logistic regression model with lasso penalty was applied to select the penalty constant, which determines the number of selected genes. Then prediction accuracy of the procedure was evaluated in a cross-validation framework. For each replicate, the model provides a probability \hat{z}_i to be pathogenic, and we considered the average value over the three replicates as the prediction probability of the strain. The predicted non-pathogenicity corresponds to a \hat{z}_i smaller than 0.5 and the predicted pathogenicity corresponds to \hat{z}_i above 0.5.

Table 3. Presence/absence of biomarkers among non-pathogenic (green), FBO (blue) and clinical (beige) strains. The presence of each biomarker gene was assessed by PCR in all strain of the collection. If the gene was present, a score of 1 was attributed (green boxes), if the gene is absent, a score of 0 is attributed (red boxes).

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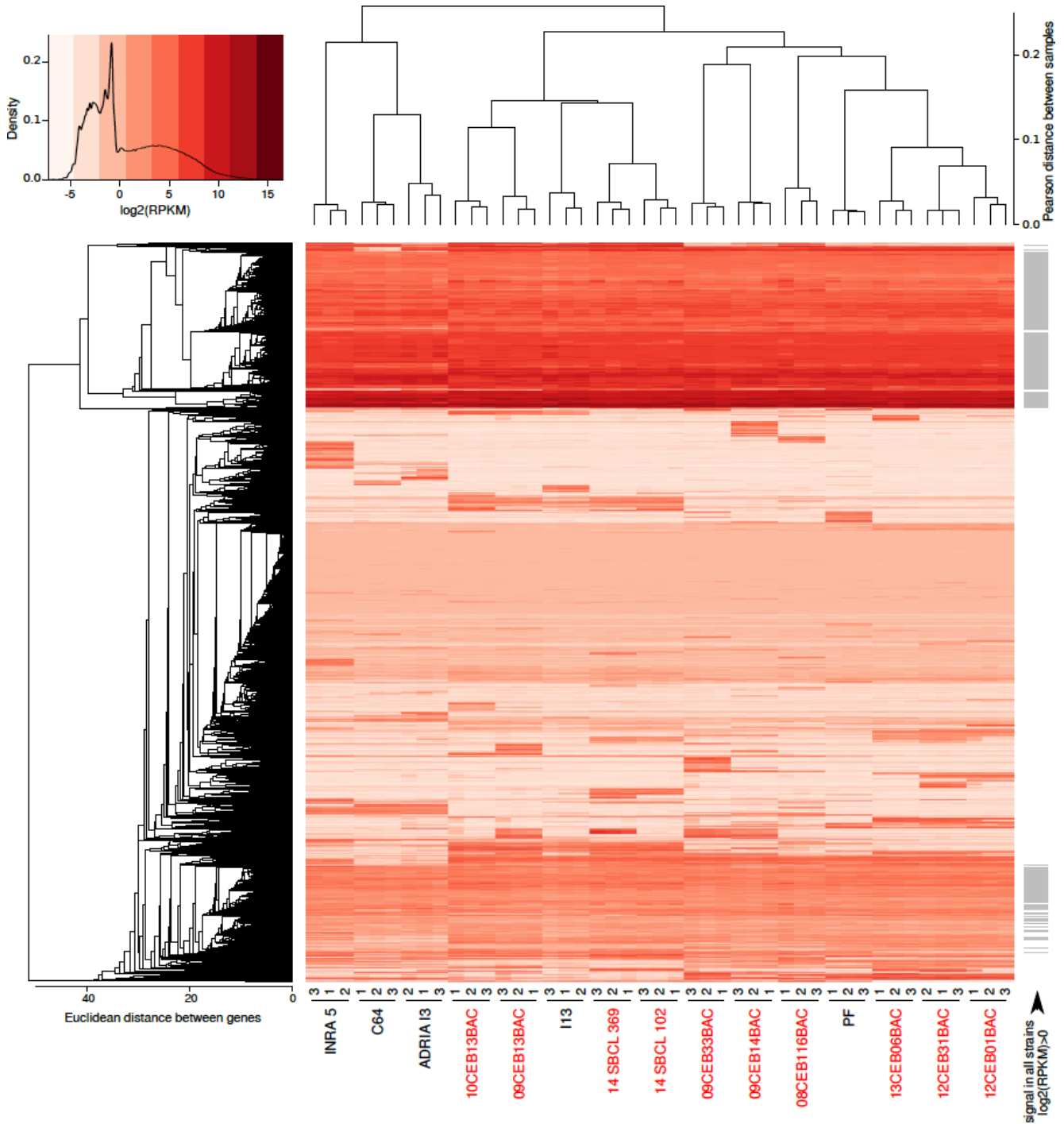
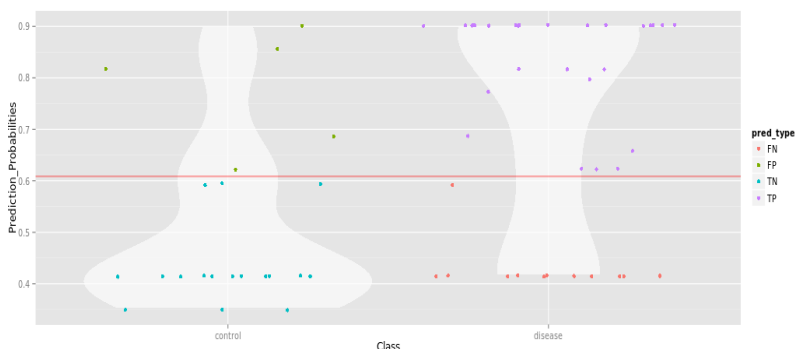


Figure 1

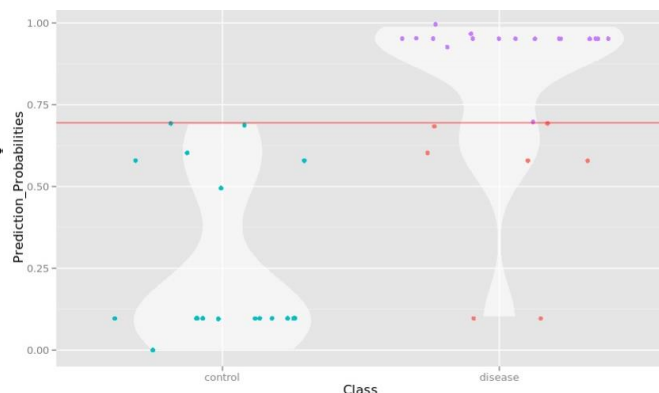
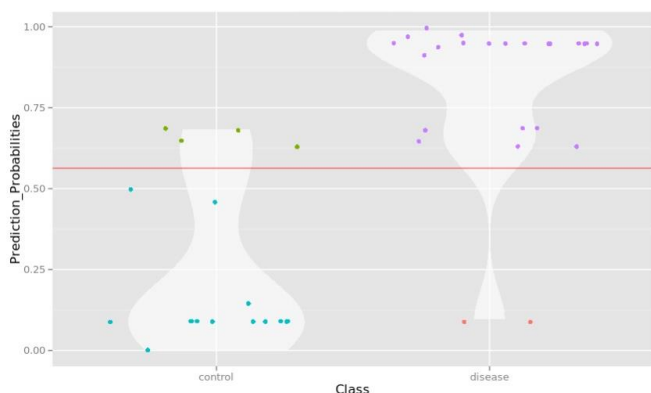
A

Biomarkers	Genes	AUC	SE	SP
Marker2-Marker3-Marker4-Marker6	agrC, thiJ, araC, gshAB	0.768	0.692	0.773



B

Biomarkers	Genes	AUC	SE	SP
Marker1-Marker2-Marker4-Marker5-Marker6	adhB, agrC, araC, BCQ_PI180, gshAB	0.917	0.917	0.778
Marker1-Marker3-Marker4-Marker5-Marker6	adhB, thiJ, araC, BCQ_PI180, gshAB	0.919	0.708	1.000



C

Biomarkers	Genes	AUC	SE	SP
Marker1-Marker2-Marker3-Marker6	adhB, agrC, thiJ, gshAB	0.955	0.909	0.864
Marker1-Marker2-Marker3-Marker5	adhB, agrC, thiJ, BCQ_PI180	0.955	0.909	0.864

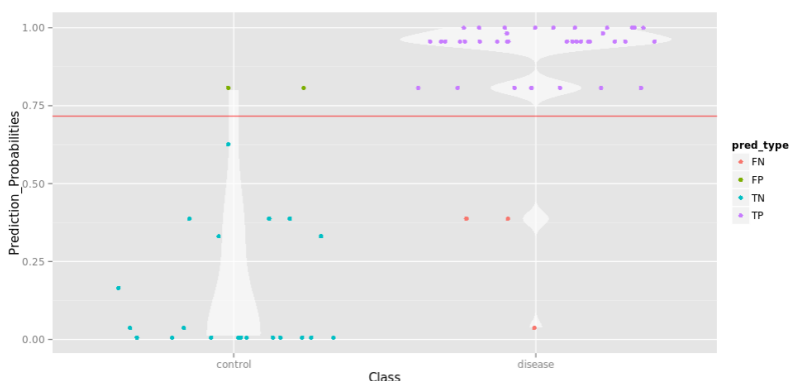


Figure 2

Table 2: Estimated probability \hat{z}_i for the 15 strains. A logistic regression model with lasso penalty was applied to select the penalty constant, which determines the number of selected genes.

Then prediction accuracy of the procedure was evaluated in a cross-validation framework. For each replicate, the model provides a probability \hat{z}_i to be pathogenic, and we considered the average value over the three replicates as the prediction probability of the strain. The predicted non-pathogenicity corresponds to a \hat{z}_i smaller than 0.5 and the predicted pathogenicity corresponds to \hat{z}_i above 0.5.

NP	Prob mean
INRA 5	0.153328340753618
C64	0.0752423643321016
ADRIA13	0.0437357685829226
I13	0.5
PF	0.599889993544854
FBO	
10CEB13BAC	0.993824252074421
08CEB116BAC	0.675323289631434
14SBCL102	0.953746924319411
14SBCL369	0.950799749333682
12CEB01BAC	0.382731024964747
Clinical	
09CEB13BAC	0.975134675591066
09CEB14BAC	0.890033149139494
09CEB33BAC	0.788491148616572
12CEB31BAC	0.977652814613013
13CEB06BAC	0.986545096552651

Table 3. Presence/absence of biomarkers among non-pathogenic (green), FBO (blue) and clinical (beige) strains. The presence of each biomarker gene was assessed by PCR in all strain of the collection. If the gene was present, a score of 1 was attributed (green boxes), if the gene is absent, a score of 0 is attributed (red boxes).

	Marker 1	Marker 2	Marker 3	Marker 4	Marker 5	Marker 6	Marker 7	PanC group
	adhB	agrC	thiJ	araC	BCQ_PI180	gshAB	BCQ_PI181	
INRA-PF_S09	0	0	1	0	0	0	0	III
I13_S10	1	0	0	0	0	0	0	IV
INRA-5_S11	0	0	0	0	0	0	0	VI
INRA-C64_S12	0	0	0	0	0	0	0	VI
ADRIA-I3_S13	0	0	0	0	0	0	0	VI
INRA-BN_S36	1	1	1	0	0	0	0	II
INRA-PA_S37	0	1	1	1	1	1	1	III
INRA-A3_S38	1	1	1	1	0	0	0	IV
I23_S39	0	0	1	0	0	0	0	IV
SB_S40	0	0	1	0	0	0	0	V
I11_S41	1	1	0	1	0	0	0	V
INRA-C1_S42	0	0	0	0	1	1	1	VI
INRA-C46_S43	0	0	0	0	0	1	0	VI
INRA-SL_S44	0	0	0	0	0	0	0	VI
INRA-SO_S45	0	0	0	0	0	0	0	VI
INRA-BC_S47	1	0	1	0	0	0	0	II
I2_S48	0	0	0	0	0	0	0	IV
INRA-BL_S49	0	0	0	0	0	0	0	VI
ADRIA I21_S50	0	0	0	0	0	0	0	VI
INRA-SV_S51	0	0	0	0	0	0	0	VI
WSBC-10204_S52	0	1	0	0	0	0	0	VI
08CEB116BAC_S1	1	1	1	0	1	1	1	II
10CEB13BAC_S2	1	1	1	1	1	1	1	IV
12CEB01BAC_S3	1	1	1	1	0	0	0	III
14 SBCL 102_S4	1	1	1	1	1	1	1	IV
14 SBCL 369_S5	1	1	1	1	1	1	1	IV
09CEB01BAC_S26	1	1	1	1	1	1	1	III
09CEB04BAC_S27	1	1	1	1	1	1	1	VII
09CEB26BAC_S28	1	0	0	1	1	1	1	II
09CEB40BAC_S29	1	1	1	0	0	0	0	II
10CEB46BAC_S30	0	0	0	0	0	0	0	IV
10CEB88BAC_S31	1	1	1	1	1	1	1	III
14 SBCL 013_S32	1	1	1	1	1	1	1	III
14 SBCL 038_S33	1	1	1	0	0	0	0	IV
14 SBCL 281_S34	1	1	1	1	0	0	0	IV

14 SBCL 714 _S35	1	1	1	0	0	0	0	II
07CEB21BAC _S65	1	1	1	1	1	1	1	III
07CEB48BAC _S66	1	1	1	1	0	0	1	III
07CEB53BAC _S67	0	1	1	1	1	1	1	III
08CEB121BAC _S68	0	0	0	0	0	0	0	IV
08CEB145BAC _S69	0	0	0	0	0	0	0	IV
08CEB037BAC _S70	0	0	0	0	0	0	0	IV
08CEB049BAC _S71	1	1	1	1	1	1	1	III
08CEB075BAC _S72	1	1	1	1	1	1	1	III
09CEB03BAC _S73	0	0	1	1	0	0	0	III
09CEB05BAC _S74	1	1	1	1	1	1	1	III
09CEB38BAC _S75	1	1	1	1	1	1	1	III
10CEB06BAC _S76	1	1	1	1	1	1	1	III
10CEB33BAC _S77	1	1	1	1	1	1	1	III
10CEB68BAC _S78	1	1	1	1	1	1	0	III
14 SBCL 008 _S79	0	0	0	0	0	0	0	IV
14 SBCL 016 _S80	0	0	0	0	0	0	0	IV
14 SBCL 020 _S81	0	0	0	0	0	0	0	IV
14 SBCL 022 _S82	0	0	0	0	0	0	0	IV
14 SBCL 049 _S83	0	1	0	0	0	0	0	IV
14 SBCL 175 _S84	0	0	0	1	0	0	0	VII
14 SBCL 180 _S85	0	0	0	0	0	0	0	IV
14 SBCL 266 _S86	0	0	0	0	0	0	0	IV
14 SBCL 374 _S87	0	0	0	0	0	0	0	IV
14 SBCL 566 _S88	0	1	1	1	1	1	0	III
09CEB13BAC _S6	1	1	1	1	1	1	1	IV
09CEB14BAC _S7	1	1	1	1	1	1	1	II
09CEB33BAC _S8	1	1	1	1	1	1	1	III
12CEB31BAC _S14	1	1	1	1	1	1	1	III
13CEB06BAC _S15	1	1	1	1	1	1	1	III
09CEB11BAC _S16	1	1	1	1	1	1	1	III
09CEB16BAC _S17	1	1	1	1	1	1	1	III
12CEB30BAC _S18	1	1	1	1	1	1	1	II
12CEB40BAC _S20	1	1	1	1	1	1	1	III
12CEB46BAC _S21	1	1	1	1	1	1	1	IV
12CEB47BAC _S22	1	1	1	0	0	0	0	IV
12CEB51BAC _S23	1	1	1	1	1	1	1	II
13CEB01BAC _S24	1	1	0	0	0	0	0	III
09CEB12BAC _S53	1	1	1	1	1	1	1	III
09CEB34BAC _S59	1	1	1	1	1	1	1	III
09CEB36BAC _S61	1	1	1	0	1	1	1	III
12CEB34BAC _S64	1	1	1	0	0	0	0	IV
12CEB37BAC _S90	1	0	0	0	0	0	0	IV
12CEB38BAC _S91	1	1	1	1	1	1	1	III
12CEB39BAC _S92	1	1	1	1	1	1	1	III
12CEB42BAC _S94	1	1	1	1	1	1	1	III
12CEB43BAC _S95	1	1	1	1	0	0	0	III
12CEB44BAC _S96	1	1	1	1	1	1	1	IV
12CEB45BAC _S97	1	1	1	1	1	1	1	II
12CEB48BAC _S98	1	1	1	1	1	1	1	II
12CEB49BAC _S99	1	1	0	0	0	0	0	IV
12CEB50BAC _S100	1	1	1	0	0	0	0	IV
12CEB52BAC _S101	0	1	1	1	0	0	0	III

13CEB03BAC_S102	1	1	1	1	1	1	1	1	II
13CEB07BAC_S105	1	1	1	1	1	1	1	1	III
13CEB09BAC_S106	1	1	1	1	1	1	1	1	III
13CEB30BAC_S107	1	1	1	0	0	0	0	0	II
14CEB16BAC_S114	1	1	1	1	1	1	1	1	IV
14CEB17BAC_S115	1	1	1	1	1	1	1	1	III
14SBCL987_S116	1	1	1	0	0	0	0	0	IV