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PointFinder: a novel web tool for WGS-based detection of antimicrobial resistance associated with chromosomal point mutations in bacterial pathogens

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Background: Antibiotic resistance is a major health problem, as drugs that were once highly effective no longer cure bacterial infections. WGS has previously been shown to be an alternative method for detecting horizontally acquired antimicrobial resistance genes. However, suitable bioinformatics methods that can provide easily interpretable, accurate and fast results for antimicrobial resistance associated with chromosomal point mutations are still lacking.

Methods: Phenotypic antimicrobial susceptibility tests were performed on 150 isolates covering three different bacterial species: Salmonella enterica, Escherichia coli and Campylobacter jejuni. The web-server ResFinder-2.1 was used to identify acquired antimicrobial resistance genes and two methods, the novel PointFinder (using BLAST) and an in-house method (mapping of raw WGS reads), were used to identify chromosomal point mutations. Results were compared with phenotypic antimicrobial susceptibility testing results.

Results: A total of 685 different phenotypic tests associated with chromosomal resistance to quinolones, polymyxin, rifampicin, macrolides and tetracyclines resulted in 98.4% concordance. Eleven cases of disagreement between tested and predicted susceptibility were observed: two C. jejuni isolates with phenotypic fluoroquinolone resistance and two with phenotypic erythromycin resistance and five colistin-susceptible E. coli isolates with a detected pmrB V161G mutation when assembled with Velvet, but not when using SPAdes or when mapping the reads.

Conclusions: PointFinder proved, with high concordance between phenotypic and predicted antimicrobial susceptibility, to be a user-friendly web tool for detection of chromosomal point mutations associated with antimicrobial resistance.

Introduction

Horizontal gene transfer among bacterial isolates is often considered the main mediator of acquired antimicrobial resistance. However, mutational resistance is another important way to confer resistance.

It has previously been shown that WGS can be an alternative to phenotypic susceptibility testing of bacterial isolates for detection of horizontally acquired resistance.1,2 Databases for mapping to chromosomal mutations have also been developed for Mycobacterium tuberculosis.3 However, at present there is a lack of suitable bioinformatics methods to provide easily interpretable results for antimicrobial resistance associated with chromosomal point mutations for most bacterial species.

In this study, a novel web tool, PointFinder, was developed for detection of chromosomal point mutations associated with antimicrobial resistance, in bacterial WGS data. PointFinder may be run in parallel and become an extension to the already existing web server tool ResFinder,1 which detects horizontally acquired resistance genes in WGS data. The performance was compared with that of an in-house mapping method for detecting point mutations and both results were compared with phenotypic antimicrobial susceptibility tests in order to validate the possibilities of using these methods as alternatives to standard phenotypical testing.
Materials and methods

Chromosomal mutation database

Information regarding mutations in chromosomal genes associated with antimicrobial resistance was collected from published papers (Table 1). The reference sequences were selected from WT *Escherichia coli* strain K-12 (MG1655) for the *E. coli* database, *Salmonella* Typhimurium strain LT2 for the *Salmonella enterica* database and *Campylobacter jejuni* NCTC 11168 for the *C. jejuni* database.

Bacterial isolates and WGS data

In total, 150 isolates covering three species were included in the study: *E. coli* (*n* = 50) and *Salmonella* (*n* = 50) isolates from the in-house strain collection at the National Food Institute and *C. jejuni* (*n* = 50) isolates from the in-house strain collection at Statens Serum Institut. The isolates were selected on the basis of having both WGS data and phenotypes available. The *Salmonella* isolates included strains from 10 different serovars (Tables S1 to S3, available as Supplementary data at JAC Online). All bacterial isolates were sequenced using the Miseq platform (Illumina) to obtain paired-end sequences and assembled de novo using Velvet (reference software).

Bacterial strains were screened for phenotypic resistance using MIC determinations interpreted according to EUCAST (www.eucast.org). Only the susceptibility tests relevant for antimicrobial resistance associated with chromosomal point mutations for each species were analysed (Table 2). As resistance to some of the antimicrobial agents can be caused by either acquired genes or chromosomal point mutations, ResFinder-2.1 (www.genomicepidemiology.org) was used to detect known acquired resistance genes in the WGS data, using a threshold of 98% identity (%ID) and 60% length (minimum percentage length of the resistance gene to be covered). All isolates with disagreement between the phenotypic and predicted susceptibility were re-tested.

PointFinder

PointFinder consists of two databases: a chromosomal gene database, with all reference sequences in fasta format; and a chromosomal mutation database containing information on codon positions and substitutions. PointFinder uses BLASTn for identifying the best match for each gene in the chromosomal gene database, and only hits with an identity of ≥80% are further analysed. The program goes through each alignment comparing each position for the query (sequence found in input sequence) with the corresponding position in the subject (database sequence). All mismatches are saved and compared with the chromosomal mutation database. It is

<table>
<thead>
<tr>
<th>Species</th>
<th>Gene</th>
<th>Chromosomal mutations</th>
<th>Resistance</th>
<th>Reference(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>pmrA</td>
<td>S39I, R81S</td>
<td>colistin</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>pmrB</td>
<td>V161G</td>
<td>colistin</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>folP</td>
<td>P63R, P64L, P64S, P64A, P64H</td>
<td>sulphamides</td>
<td>9</td>
</tr>
<tr>
<td>23S(^a)</td>
<td></td>
<td>A2059G</td>
<td>macrolide</td>
<td>11</td>
</tr>
<tr>
<td>16S rrsB(^a)</td>
<td></td>
<td>A523C, G527T, C528T, G1064T, G1064C, G1064A, C1066T, G1068A</td>
<td>spectinomycin</td>
<td>12–15</td>
</tr>
<tr>
<td>16S rrsB(^a)</td>
<td></td>
<td>A964G, G1053A, C1054T, A1055G, G1058C</td>
<td>tetracycline</td>
<td>13, 16</td>
</tr>
<tr>
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<td></td>
<td>T1406A, A1408G</td>
<td>gentamicin</td>
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</tr>
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<td>kasugamycin</td>
<td>18</td>
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<tr>
<td>16S rrsH(^a)</td>
<td></td>
<td>C1192T</td>
<td>spectinomycin</td>
<td>19</td>
</tr>
<tr>
<td></td>
<td>parC</td>
<td>T66I, G78D, S80R, S80I, E84K, E84G</td>
<td>quinolone</td>
<td>4, 21</td>
</tr>
<tr>
<td></td>
<td>parE</td>
<td>M438E, E464G</td>
<td>quinolone</td>
<td>4, 20, 22</td>
</tr>
<tr>
<td>16S rrsD(^a)</td>
<td></td>
<td>C1065T, C1192T</td>
<td>spectinomycin</td>
<td>24</td>
</tr>
<tr>
<td><em>C. jejuni</em></td>
<td>gyrA</td>
<td>A70T, D85T, T86I, T86A, T86K, T86V, D90A, D90N, D90T, P104S</td>
<td>quinolone</td>
<td>25–28</td>
</tr>
<tr>
<td>23S(^a)</td>
<td></td>
<td>A2074G, A2074T, A2074C, A2075G</td>
<td>macrolide</td>
<td>28</td>
</tr>
<tr>
<td></td>
<td>cmeR</td>
<td>A86G</td>
<td>macrolide</td>
<td>29</td>
</tr>
<tr>
<td></td>
<td>rplV</td>
<td>A103C</td>
<td>macrolide</td>
<td>29</td>
</tr>
<tr>
<td></td>
<td>rpsL</td>
<td>K88E, K88R, K88Q</td>
<td>spectinomycin</td>
<td>30</td>
</tr>
</tbody>
</table>

\(^a\)rRNA gene, mutation shown in DNA.

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2765
possible for users to select whether they want to see all mismatches or only known mismatches found in positions from the chromosomal database. In this study we have only looked at mismatches found in positions known to confer resistance, and thus specified in the database.

Mapping method

The fastq files corresponding to the paired-end reads were mapped against the chromosomal gene sequence database using the assmple.py script described in Joensen et al. In brief, 17mers from the reads were mapped to the reference sequence and extended to ungapped alignments that were considered significant if they had a score of at least 50, using a match score of 1 and a mismatch score of −3. A base was called if Z = (X − Y)/(X + Y) was > 3.29, where X is the number of observations of the most common nucleotide and Y is the number of other nucleotides at that position. Furthermore, nucleotide calls were considered significant only when the most common nucleotide was at least 10 times more abundant than other nucleotides at the position. All mismatches in positions from the chromosomal database were outputted except silent mutations, which were discarded. In cases with disagreement between PointFinder and mapping, the isolates were re-assembled de novo using SPAdes and re-analysed by PointFinder.

Results

MIC and predicted antimicrobial resistance

The 150 isolates were each tested against four to six different antimicrobial agents (Table 2), leading to a total of 684 susceptibility test results (Tables S1–S3). These results were compared with the results from PointFinder, mapping and ResFinder. Resistance to colistin, sulphonamides, tetracycline, erythromycin and spectinomycin can be caused by both chromosomal point mutations and acquired resistance genes; therefore results from both PointFinder and ResFinder were used to explain resistance.

For all Salmonella isolates, complete agreement between tested and predicted susceptibility was observed (Tables S1–S3). Disagreements in E. coli and C. jejuni were observed in five and four cases, respectively (Table 3).

The point mutation pmrB V161G was found by PointFinder in five E. coli isolates (E30–E34), but all tested phenotypically susceptible to colistin (MIC ≤1 mg/L). In C. jejuni, two isolates (C23 and C39) tested phenotypically resistant to ciprofloxacin (MIC 8) and nalidixic acid (MIC >64 mg/L), while two (C8 and C24) tested erythromycin resistant (MIC >128 mg/L), but neither mutations nor acquired genes were found that could explain the resistance.

PointFinder versus mapping

Mapping and PointFinder found the same mutations in all isolates except the five pmrB V161G mutations found by PointFinder in E. coli strains (E30–E34). The five isolates were re-assembled de novo using SPAdes and run through PointFinder, and this time no mutations were found in pmrB in any of the isolates. The codon change detected in the Velvet assembly of the five isolates was GTG→GGG, and when looking further into the sequences, the mapping showed that 28%–37% (Table 4) of the reads mapping to pmrB contained GGG instead of GTG.

Discussion

This study showed a high agreement between phenotypic susceptibility tests and WGS-predicted resistance, with only 11 (1.6%) mismatches. However, since the number of isolates included in the evaluation was very limited and selected, this has to be further verified in future studies. The six disagreements observed in C. jejuni all involved predicted susceptibility, whereas the isolates were phenotypically resistant, which may be due to unknown novel genes or mutations, as neither ResFinder nor PointFinder can detect novel resistance mechanisms.

We found that the BLAST-based method was dependent on the assembly method, which can cause either false-positive or negative results. As the mapping method does not depend on the assembly this method gives a more precise result, which is consistent with a recent study by Clausen et al. Exploring the sequences mapping to pmrB, we found that ~1/3 of the isolate sequences for
The user must therefore have prior knowledge of which mutational regions of genes associated with mutational resistance, as well as information about position and mutation in the corresponding gene. ARG-ANNOT does not automatically detect these mutations, so the user has to manually browse through the alignment to detect potential mutations. CARD’s resistance gene identifier (RGI) protein variant models use curated SNP matrices to detect and report mutations and amino acid codon changes, predicted resistance and links to functional resistance, which is not relevant for the bacteria in question. Both ARG-ANNOT and CARD have tried to incorporate chromosomal point mutations in their databases. ARG-ANNOT has a database with partial sequences for chromosomal mutational regions of genes associated with mutational resistance, as well as information about position and mutation in the corresponding gene. ARG-ANNOT does not automatically detect these mutations, so the user has to manually browse through the alignment to detect potential mutations. CARD’s resistance gene identifier (RGI) protein variant models use curated SNP matrices to detect and report mutations associated with resistance. Unfortunately, neither ARG-ANNOT nor CARD takes the bacterial species into account. This means that both methods also output possible mutations/sequences related to mutational resistance, which is not relevant for the bacteria in question. The user must therefore have prior knowledge of which mutational genes and specific mutations they are looking for in order to use these methods. To cope with some of these problems, we have developed PointFinder, with the purpose of facilitating user-friendly detection of chromosomal point mutations associated with resistance. In addition to being user friendly, the output from the web tool is easily understandable, reporting the detected mutations, nucleotide and amino acid codon changes, predicted resistance and links to papers describing the detected mutations. In the current version this covers mutations conferring resistance to quinolones, macrolides and polymyxin in E. coli, Salmonella and C. jejuni, but will be developed continuously with additional species.

Conclusions

This study showed a high concordance between phenotypic antimicrobial susceptibility and predicted genotype by ResFinder and PointFinder from WGS data. PointFinder is a user-friendly method for detection of chromosomal point mutations associated with antimicrobial resistance.

Funding

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Table 4. Mapped sequences to pmrB position 161 (amino acid position)

<table>
<thead>
<tr>
<th>Isolate ID</th>
<th>Total mapped sequences</th>
<th>No. of sequences mapping to GTG (%)</th>
<th>GGG (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>E30</td>
<td>69</td>
<td>50 (72)</td>
<td>19 (28)</td>
</tr>
<tr>
<td>E31</td>
<td>70</td>
<td>44 (63)</td>
<td>26 (37)</td>
</tr>
<tr>
<td>E32</td>
<td>80</td>
<td>54 (68)</td>
<td>26 (33)</td>
</tr>
<tr>
<td>E33</td>
<td>70</td>
<td>47 (67)</td>
<td>23 (33)</td>
</tr>
<tr>
<td>E34</td>
<td>89</td>
<td>64 (72)</td>
<td>25 (28)</td>
</tr>
</tbody>
</table>

European Union’s Horizon 2020 research and innovation programme under grant agreement no. 643476.

Supplementary data

Table S1 to S3 are available as Supplementary data at JAC Online.

References