CPgeneProfiler: A lightweight R package to profile the Carbapenamase genes from genome assemblies

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Summary

“Carbapenems” are a specific subset of antibiotics considered to possess a higher spectrum of antimicrobial activity (Papp-Wallace, Endimiani, Taracila, & Bonomo, 2011) against Gram-positive and Gram-negative bacteria. Even so, there are pathogens which are resistant to carbapenems due to the presence of carbapenemase genes (CP genes) which have the ability to hydrolyze carbapenems.

Studies show that those cases infected by carbapenem-resistant pathogens have a higher morbidity and mortality rate compared with those who are infected by non-carbapenem-resistant pathogens (Cai et al., 2017; Duin, Kaye, Neuner, & Bonomo, 2013). Therefore, early discerning of the CP genes and their resistance mechanisms are considered crucial to aid in infection control as well as lessen the likelihood of mortality, duration of hospitalization stay, and related medical costs (Duin et al., 2013; Nordmann & Poirel, 2019). Further, it is understood that the cocarriage of genes encoding different classes of carbapenemases could confer higher resistance to carbapenem antibiotics, which may promote further spread of the disease (Wang et al., 2019).

The detection of the resistance genes from various bacterial strains using techniques such as polymerase chain reaction (PCR) and microarrays in real-time is very time consuming and costly. With the advancement in whole-genome sequencing (WGS) technologies and decreased costs, this is more accessible and WGS provides an alternative method for detection of resistance genes, given that the relevant analysis tools are available.

To this end, several freely available bioinformatics tools such as ABRicate (https://github.com/tseemann/abricate), AMRPlusPlus (Doster et al., 2020), ARG-ANNOT (Gupta et al., 2014), ARIBA (Hunt et al., 2017), Comprehensive Antibiotic Resistance Database – Resistance Gene Identifier (CARD-RGI) (Alcock et al., 2020), NCBI AMRFinderPlus (https://ncbi.nlm.nih.gov/pathogens/antimicrobial-resistance/AMRFinder/), KmerResistance (Clausen, Aarestrup, & Lund, 2018; Clausen, Zankari, Aarestrup, & Lund, 2016), PointFinder (Zankari et al., 2017), Resfinder (Bortolaia et al., 2020), sraX (Panunzi, 2020), and SRST2 (Inouye et al., 2014) assist in finding the antimicrobial resistance genes from the sequence data (Hendriksen et al., 2019).

Statement of Need

Undeniably, all the above-mentioned tools are focused around the antimicrobial-resistant genes, and tools such as ABRicate and CARD-RGI can even generate comparative tables.
across genomes, and sraX can help in visualization of comprehensive AMR gene complement. Nevertheless, they do not readily generate a genetic profile for the presence of CP genes, and extract and visualize the set intersections of cocarriage of CP genes. Achieving this currently necessitates a restructuring and transformation of the output from these tools. Furthermore, in the research settings where it is crucial to quickly examine the transmission of CP genes, it is useful to have a tool that is catered to the CP gene dataset that provides easily interpretable visualizations and statistics. Therefore, to address this need, we describe here a lightweight R package, **CPgeneProfiler**, that scans multiple bacterial genome assemblies to detect and visualize the presence of CP genes and their cocarriage using the R framework. Additionally, this package also allows one to assess the size of CP contigs to check if the CP genes are distributed on the particular sequence size by generating the contig length distribution plots.

**Implementation**

In order to detect CP genes from the genome assemblies, NCBI Bacterial Antimicrobial Resistance Reference Gene Database (2020-07-16.2) ([https://www.ncbi.nlm.nih.gov/bioproject/PRJNA313047](https://www.ncbi.nlm.nih.gov/bioproject/PRJNA313047)) was used for generating a CP gene database. Only those resistant genes whose subclass is categorized as “CARBAPENEM” in the reference gene catalog were considered for database preparation. This excluded the possibility of having resistant gene variants which are beta-lactamases but do not show carbapenem-resistant activity. For example, although OXA-48 is a carbapenemase gene, other OXA variants such as OXA-163 and OXA-405 have been determined to be devoid of any carbapenemase activity (Dortet & Naas, 2017), and therefore their subclass is not categorized as “CARBAPENEM” in the NCBI Bacterial Antimicrobial Resistance Reference Gene Database. Therefore, OXA variants OXA-163 and OXA-405 were not included in the CP gene database.

The tool first uses the `cpblast` command, by which each fasta file is searched against the CP gene database using NCBI BLAST+ (Camacho et al., 2009) (version 2.9.0+), which is pre-installed in the local system as a dependency. The presence of a CP gene in an assembled genome is confirmed if the CP gene meets the identity and coverage thresholds (default: 100%) when aligned with the genome sequence. The genome sequences that meet the thresholds are extracted from the BLAST results using the `filt_blast` command.

Visualizing the presence of CP genes and their corresponding counts across all the genome assemblies in a simple heatmap enables one to find CP gene variants that are found across the samples and aids in exploring the pattern of CP gene presence with reference to species or sequence type (ST). In order to facilitate this, the `cpprofile` command generates a profile of CP genes (Figure 1A) across the genome assemblies, while the cocarriage command finds cocarriage of CP genes in the genome assemblies. In addition to this, the tool also generates plots to visualize the set intersections of CP genes across all the input genome assemblies using the command `upsetR_plot` (Figure 1B). It is understood that isolates that harbour multiple carbapenemase genes are considered to produce high resistant phenotypes, and running the commands `cocarriage` and `upsetR_plot` provides an overview of the CP genes as well as their cocarriages present in all the genomes.

Given a set of bacterial genomes that are of same species, it would be useful to explore if the CP genes are found on specific plasmids or scattered across multiple plasmids/chromosomes of different sequence lengths. This can be achieved by plotting the number of contigs across the contig length by using the `plot_conlen` command (Figure 2).

Lastly, CPgeneProfiler can also generate the N50, N90, and assembly size statistics for each of the genome assemblies and also plots the assembly size against N50 and N90 using the `assembly_stat` command (Figure 3A, 3B). This helps in quickly assessing and comparing the quality of the assembled genomes provided as an input. All the generated output files from various commands of the package are arranged accordingly into their respective folders using the `cp_summarize` command.

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Availability and Implementation

The R package CPgeneProfiler (version 2.1.1) is supported on UNIX/Linux machines. The source code, guide and datasets are currently available on Github repository (https://github.com/ramadatta/CPgeneProfiler).

Step 1: Download CP gene database using R console

```r
# Specify CP gene database URL
url <- "https://raw.githubusercontent.com/ramadatta/CPgeneProfiler/master/testData/db/NCBI_BARRGD_CPG_DB.fasta"

# Specify destination where CP gene database file should be saved
path <- "/home/user/db"  # Can be changed to preferred location
setwd(path)
destfile <- "NCBI_BARRGD_CPG_DB.fasta"

# Download the CP gene database file to the folder set in "path"
download.file(url, destfile)
```

Step 2: Install CPgeneProfiler package

The R package CPgeneProfiler can be installed by typing the following in R:

```r
devtools::install_github("ramadatta/CPgeneProfiler")
```

Figures

Figure 1. (A) CP gene profile obtained by ‘cpprofile’ command (B) Set intersection plot of the available CP genes across genome assemblies, obtained by the ‘upsetR_plot’ command

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Conclusion

CPgeneProfiler can be used to understand the CP gene profile of a set of bacterial genome assemblies. It generates a simple heatmap for visualization of the CP gene profile and reports details on cocarriage of CP genes within the genomes. The capability to identify and visualize the presence of CP genes across multiple genomes would have useful applications, for example, in a dataset of outbreak samples, and the CPgeneProfiler could aid researchers in obtaining an overview of the samples and their CP gene carriage.

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