

HELSINGIN YLIOPISTO HELSINGFORS UNIVERSITET UNIVERSITY OF HELSINKI

MAATALOUS-METSÄTIETEELLINEN TIEDEKUNTA AGRIKULTUR-FORSTVETENSKAPLIGA FAKULTETEN **FACULTY OF AGRICULTURE AND FORESTRY**



Mobility potential of latent antibiotic resistance genes: novel mobile colistin resistance genes

Antti Karkman github.com/karkman @anttikarkman.bsky.social

Melina Markkanen Svjetlana Dekić Rozman Johanna Muurinen Department of Microbiology, University of Helsinki, Finland

Research Council

of Finland

BACKGROUND

After their recent discovery, several different mobile colistin resistance (mcr) genes have been identified. These genes are closely related to bacterial chromosomal eptA genes. So the question is, are all mobile or plasmid-borne eptA/mcrgenes mobile colistin resistance genes? And could these genes pose a potential risk to human health? Our aim was to characterize the genetic diversity of eptA/mcr-genes in influent sewage and determine their genetic context to infer their mobility potential.

MATERIALS AND METHODS

Influent sewage was sampled on three consecutive days and sequenced with PacBio Sequel II. Metagenomes were assembled with with hifiasm-meta. HMM-models were build from known *mcr*-gene variants and queried against the assembled metagenomes. Plasmid contigs were predicted with PlasX and mobile genetic elements within 5 kb up- or region from the identified gene with downstream MobileElementFinder. Phylogenetic tree was built with FastTree.

RESULTS

s1.ctg0168641

s14701.ctg0192081

2.5kb

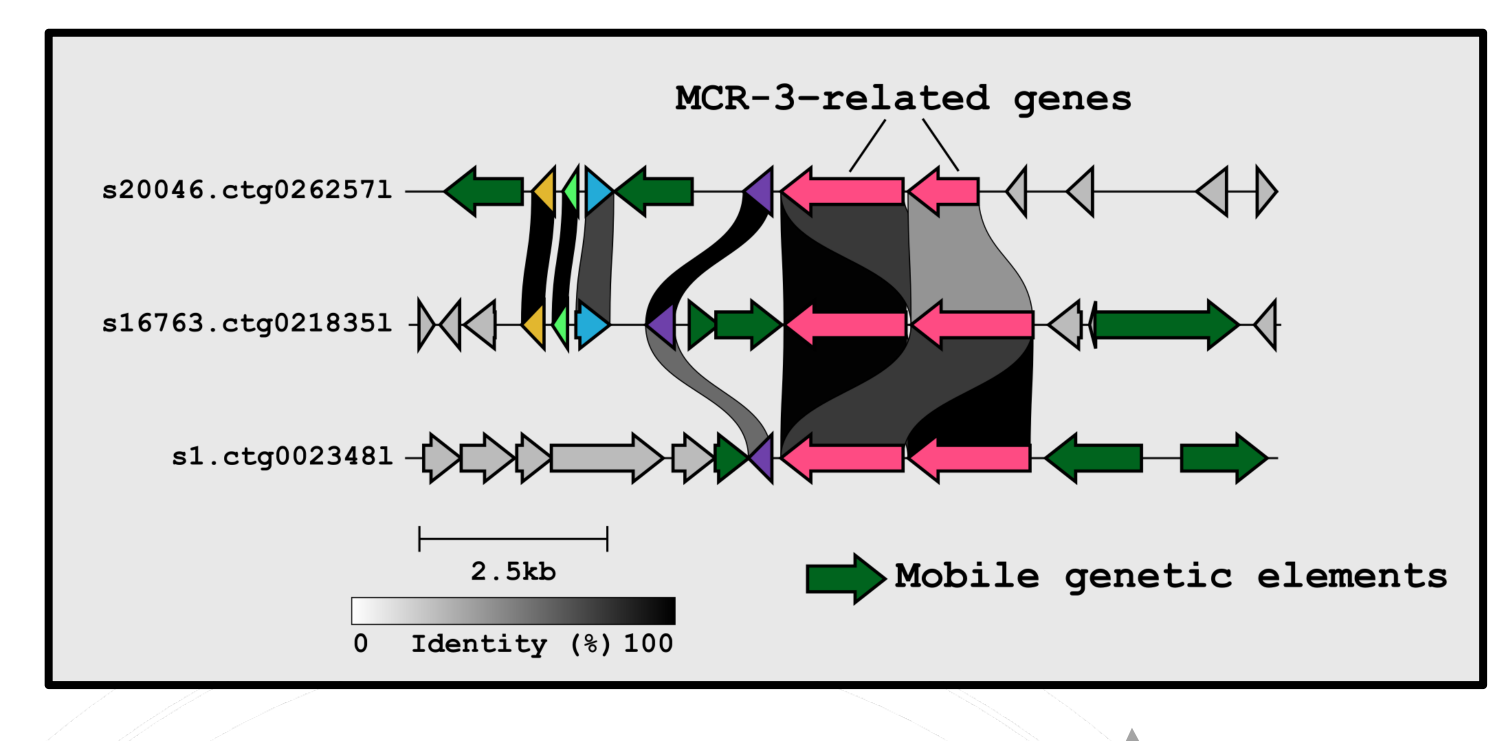
0 Identity (%) 100

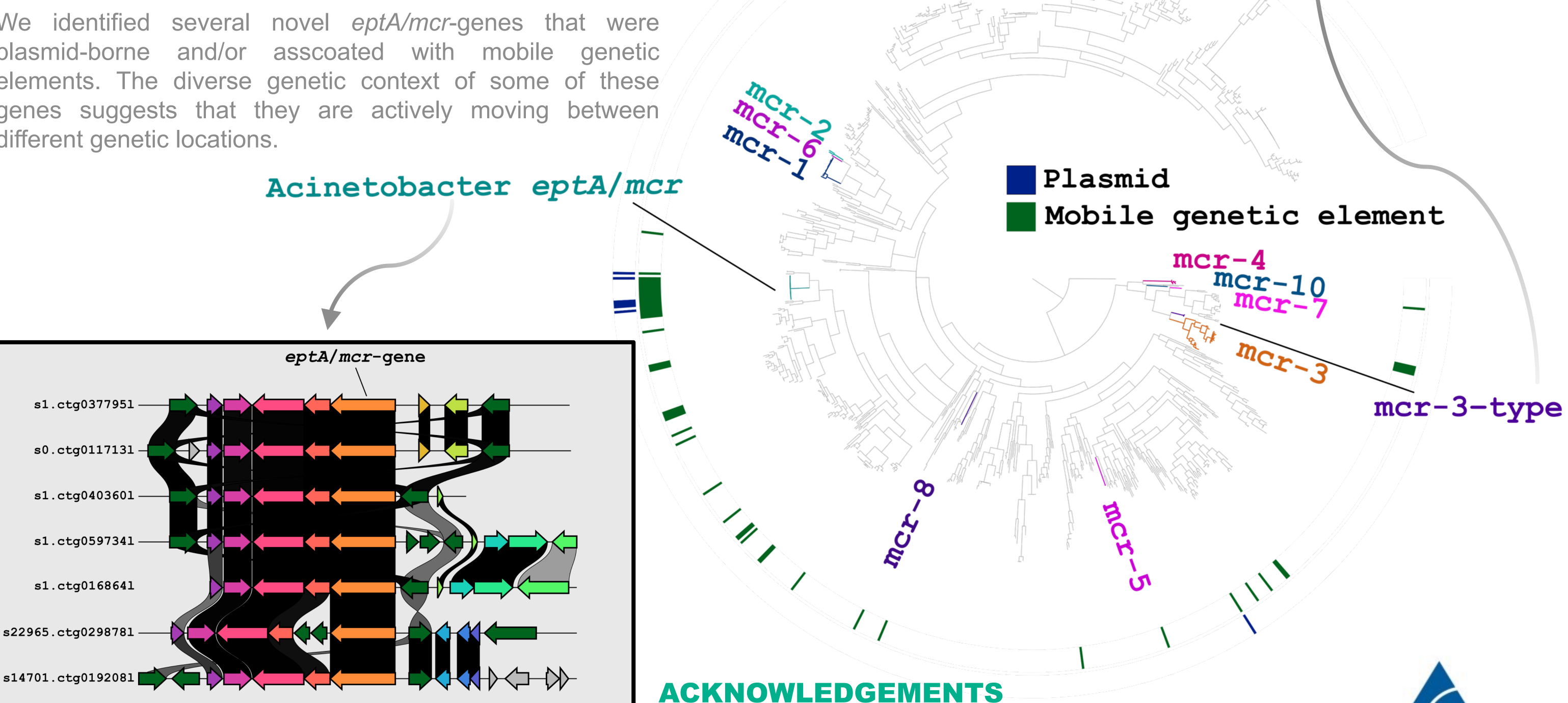
We identified several novel eptA/mcr-genes that were plasmid-borne and/or asscoated with mobile genetic elements. The diverse genetic context of some of these genes suggests that they are actively moving between different genetic locations.

Mobile genetic elements

CONCLUSION

Sewage hosts a diverse pool of eptA/mcr-genes. Some of these genes could be considered novel mobile colistin resistance genes based on their mobiity potential. Further work is needed to link these geens to their hosts, determine their phenotypic resistance patterns and get better insights of the possible horizontal transfer routes of these genes.





computational resources

The authors wish to acknowledge the DNA Sequencing and Genomics

sequencing and the CSC-IT Center for Science, Finland, for providing

Laboratory, Institute of Biotechnology, University of Helsinki, for