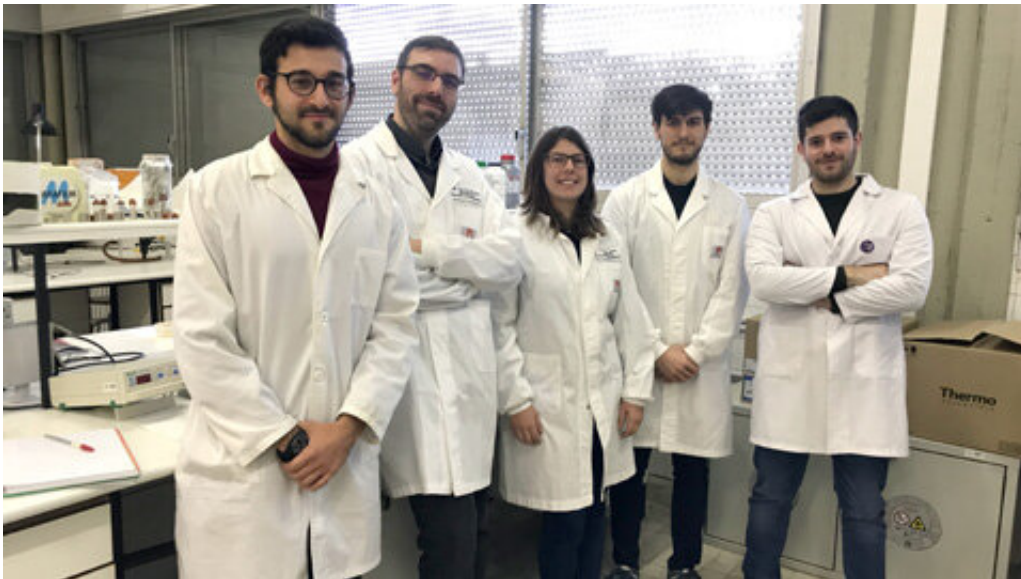


Researchers develop a database to aid in identifying key genes for bacterial infections

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A team of scientists from the UAB and the CRG have created the BacFITBase database, which characterises bacterial genes relevant to the infection process in live organisms. The new database will make it easier to identify new therapeutic targets for the creation of antibiotics. Credit: Autonomous University of Barcelona

Infectious diseases are caused by pathogenic microorganisms capable of entering, colonizing and growing within a host organism, thus producing an infection. Bacterial infections have been on the rise worldwide in recent years, but many mechanisms underlying bacterial pathogenesis are still poorly understood. This is highly relevant given the fact that the

development of new antimicrobial therapies is largely based on current knowledge of the mechanisms behind these infections. The proteins coded by the bacterial genes are responsible for the thousands of biochemical processes essential for the efficient propagation of the pathogen. Many studies demonstrate, however, that in order to identify these genes, *in vivo* information is needed on what happens with the bacteria in a real case of an infected host. The *in vitro* studies, i.e., those recreated in laboratories with cell and bacterial cultures, later do not always correlate with data from *in vivo* studies. This is due to the fact that pathogenic bacterial genes essential for producing the infections depend on the environment of the colonized organism.

A team of researchers from the Department of Biochemistry and Molecular Biology of the Universitat Autònoma de Barcelona and from the Centre of Genomic Regulation (CRG) has created the [BacFITBase database](#). Based on the results of the *in vivo* experiments, researchers systematically characterized the bacterial genes relevant for host cell invasion and infection. All the experiments performed were based on a technique named transposon mutagenesis, where the DNA fragments called transposons are transferred to the organism's pathogenic genes, thereby inactivating them. By doing so, their role in the infection can be observed directly and researchers can determine which are essential for a specific host organism to become infected. Therefore, this database will make it easier to identify target proteins that can help in fighting [infectious diseases](#) and accelerate the development of new antimicrobial agents.

The database contains over 90,000 entries with information on specific pathogenic [bacterial genes](#) and their contribution to *in vivo* infectious conditions in five different host species. It includes information on a total of 15 bacteria (two variants of *Salmonella enterica*, *Haemophilus influenzae*, *Streptococcus pyogenes*, *Porphyromonas gingivalis*, *Mycobacterium avium*, three variants of *Escherichia coli*, *Vibrio*

cholerae, *Campylobacter jejuni*, *Klebsiella pneumoniae*, *Acinetobacter baumannii*, *Serratia marcescens* and *Vibrio parahaemolyticus*) and five model vertebrates (cow, pig, hen, mouse and rabbit), with information across 10 different tissues.

More information: Javier Macho Rendón et al. BacFITBase: a database to assess the relevance of bacterial genes during host infection, *Nucleic Acids Research* (2019). [DOI: 10.1093/nar/gkz931](https://doi.org/10.1093/nar/gkz931)

Provided by Autonomous University of Barcelona

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