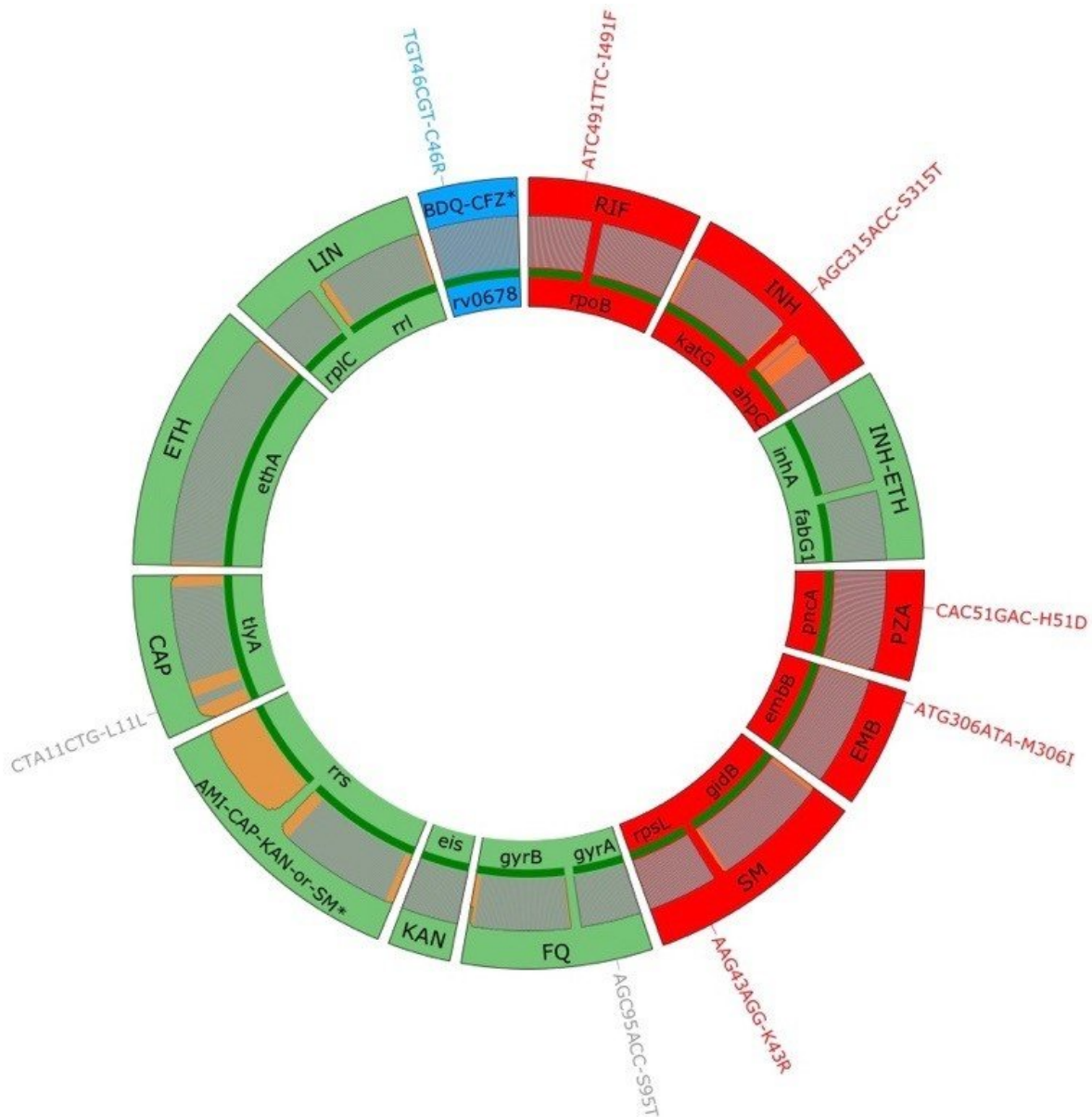


Outbreak of multidrug-resistant tuberculosis undetected by standard tests

October 17 2018



Detection of multi-resistance to antibiotics in a South African strain of tuberculosis using the Deeplex-MycTB® assay. Red and blue sections indicate confirmed or probable mutations causing resistance to first-line antibiotics and bedaquiline, respectively. Credit: Genoscreen, Lille

On 26 September, the United Nations announced a plan to raise \$13 billion annually for the fight to eradicate tuberculosis by 2030. With 10 million new cases and 1.6 million deaths in 2017, it is the most common infectious disease in the world, ahead of HIV. In over 450,000 new cases of antibiotic-resistant tuberculosis that likely appeared, only 25% were detected. A study by an international research team co-directed by Philip Supply, a CNRS researcher at the Center of Infection and Immunity of Lille (CNRS/Inserm/Institut Pasteur de Lille/Université de Lille), has underlined this serious problem of under-detection, in South Africa in particular.

The findings, published in *The Lancet Infectious Diseases*, show that certain South African isolates of *Mycobacterium tuberculosis* (the bacterium which causes the disease) carry a specific combination of mutations which make them resistant to the two primary first-line antibiotics prescribed: rifampicine and isoniazide. This combined resistance goes undetected by the standard tests endorsed by the World Health Organization: the gene region carrying a particular mutation causing rifampicine resistance is not included in the DNA test, and the resistance to the treatment due to this mutation is not detected in cultures.

This omission leads to unsuccessful first-line treatments in patients, increased mortality and contagion, and the development of additional antibiotic resistances. Researchers especially detected the presence of mutations probably causing decreased sensitivity to bedaquiline, the

newest molecule used to treat cases of multidrug-resistant (MDR) tuberculosis. These mutations appeared immediately following its launch in the country from 2013 on.

This was discovered thanks to a new MDR screen test developed by Genoscreen together with P. Supply. Unlike standard DNA tests, this one analyses a wide panel of target genes in the bacteria and can identify resistance to over a dozen antibiotics simultaneously. These results are obtained in as little as one to three days, compared to the weeks needed for cultures. The test will help solve the problem of under-detection of MDR [tuberculosis](#).

It will benefit from a new algorithm for the detection of resistance mutations, the effectiveness of which has been detailed in an article published in The New England Journal of Medicine by another consortium (CRyPTIC) in which Dr. Supply and Genoscreen took part. This study was based on an analysis of 10,000 genomes, making it one of the biggest microbial genome sequencing projects conducted to date.

More information: *The Lancet Infectious Diseases* (2018). [DOI: 10.1016/S1473-3099\(18\)30496-1](https://doi.org/10.1016/S1473-3099(18)30496-1)

Provided by CNRS

Citation: Outbreak of multidrug-resistant tuberculosis undetected by standard tests (2018, October 17) retrieved 23 May 2024 from <https://medicalxpress.com/news/2018-10-outbreak-multidrug-resistant-tuberculosis-undetected-standard.html>

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