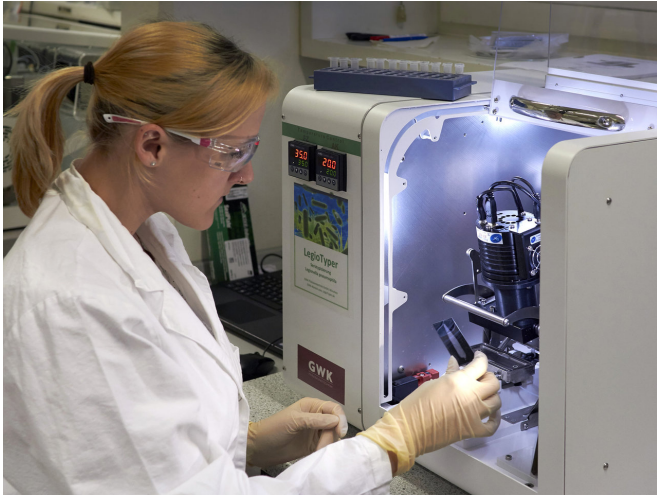


Microarray rapid test speeds up detection during a Legionella pneumophila outbreak

23 March 2018



In an outbreak of Legionnaires' disease, finding the exact source as quickly as possible is essential to preventing further infections. Researchers at the Technical University of Munich have now developed a rapid test that detects *Legionella pneumophila* in about 35 minutes. The picture shows first author Catharina Kober with a LegioTyper-chip at the MCR microarray analysis platform. Credit: Jonas Bemetz / TUM

In an outbreak of Legionnaires' disease, finding the exact source as quickly as possible is essential to preventing further infections. To date, a standard analysis takes days. Researchers at the Technical University of Munich have now developed a rapid test that achieves the same result in about 35 minutes.

Legionella are rod-shaped bacteria that can cause life-threatening pneumonia in humans. They multiply in warm water and can be dispersed into the air via cooling towers, evaporative recooling systems and hot water systems. The most dangerous among the almost 50 species of Legionella is Legionella pneumophila. It is responsible for 80 percent of all infections. When an outbreak occurs, the source of the germs must

be identified as soon as possible to prevent further infections.

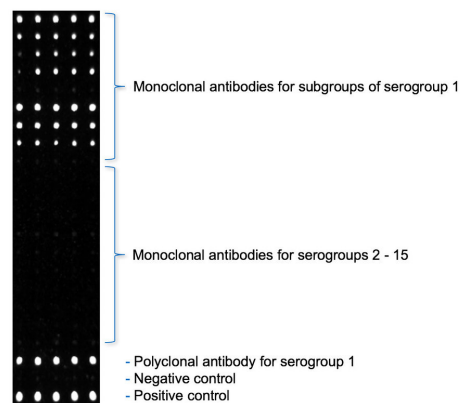
Similar to a paternity test, the origin of the outbreak is confirmed when the germs in the process water of a technical system exactly match those identified in the patient. However, often numerous systems must be tested in the process, and the requisite cultivation for the test takes around ten days.

Faster detection with antibodies

Meanwhile, there is a rapid urine test for detecting the Legionella pathogen in the clinic.

"Unfortunately, this quick [test](#) serves only as a first indication and is not suitable for screening the [water](#) of technical systems," says PD Dr. Michael Seidel, head of the research group at the Chair of Analytical Chemistry and Water Chemistry of the Technical University of Munich.

The team of scientists thus developed a measuring chip in the context of the "LegioTyper" project funded by the German Federal Ministry of Education and Research. This chip not only detects the dangerous pathogen Legionella pneumophila but also identifies which of the approximately 20 subtypes is present.



The most dangerous among the almost 50 species of *Legionella* is *Legionella pneumophila*. It is responsible for 80 percent of all infections. The picture shows the sensor image of *Legionella pneumophila* serogroup 1, strain Bellingham-1. Credit: Catharina Kober / TUM

Fast, inexpensive and versatile

The foil-based measuring chip uses the microarray analysis platform MCR of the Munich company GWK GmbH. Using 20 different antibodies, the system provides a complete analysis within 34 minutes. "Compared to previous measurements, the new method not only provides a huge speed advantage," says Michael Seidel, "but is also so cheap that we can use the chip in one-time applications."

The system can be deployed for environmental hygiene as well as clinical diagnostics. In combination with a second, DNA-based method, the system can even distinguish between dead and living *Legionella* pathogens. This allows the success of disinfection measures to be monitored. The project participants will present their [system](#) to the public for the first time at the Analytica 2018 trade fair in Munich (Hall 3, Booth 315).

More information: Catharina Kober et al, Quantification of viable and non-viable *Legionella* spp. by heterogeneous asymmetric recombinase polymerase amplification (haRPA) on a flow-based chemiluminescence microarray, *Biosensors and Bioelectronics* (2017). [DOI: 10.1016/j.bios.2017.08.053](https://doi.org/10.1016/j.bios.2017.08.053)

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