

How bacteria make syringes: Scientists reconstruct bacterial transport channel in test tube

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Adding needle protein leads to a spontaneous prolongation of some needles. The bar corresponds to 100 nanometers. 1 nanometer corresponds to a millionth millimeter. Credit: Christian Goosmann and Michael Kolbe

For a successful infection, bacteria must outwit the immune system of the host. To this aim, they deliver so-called virulence factors through a transport channel located in the bacterial membrane. In some bacteria this transport channel is formed like a syringe, enabling them to inject virulence factors directly into the host cell.

German scientists from the Max Planck Society and the Federal Institute for Materials Research and Testing have now succeeded for the first



time in elucidating basic principles of the assembly of this transport channel. This is an important starting point for the development of new drugs that might interfere considerably earlier than antibiotics in the course of infection. (*Nature Structural & Molecular Biology*, 13 June 2010)

Every day the human organism is confronted with a huge variety of pathogens. Most of them are fended off by our <u>immune system</u>. To execute a successful infection, bacteria must therefore manipulate the host to ensure their survival. They secrete virulence factors through a transport channel located in the bacterial membrane. Some bacteria, such as the causative agents of dysentery, food poisoning, typhoid fever, and pest, have developed a specialized transport mechanism called the Type three secretion system. Electron microscopy reveals that this structure is formed like a syringe: the base of the syringe is imbedded in the bacterial membrane and the needle protrudes out of the bacteria. With this apparatus bacteria can inject virulence factors directly into the host cell.



Shigella flexneri, the causative agent of dysentery (orange), establishes contact with a human host cell (blue). The bar corresponds to a micrometer or a thousandth millimeter, respectively. Credit: Volker Brinkmann, Diane Schad, and Michael Kolbe



So far, little has been known about how bacteria build this nano-syringe. Scientists from the Max Planck Institute for Infection Biology in Berlin, the Max Planck Institute for Biophysical Chemistry in Göttingen, and the Federal Institute for Materials Research and Testing have now succeeded in elucidating fundamental principles of the needle assembly. This was made possible by reconstitution experiments which allowed them to study the assembly of proteins into a needle in the test tube.

The close observation of these events revealed how the proteins are assembled into a syringe: the bacterium synthesizes the proteins in the cell interior, transports them through the syringe to the outside, and stacks them one after the other onto the tip of the growing needle. The scientists could also show that the proteins change their threedimensional structure during the assembly process. They were able to pinpoint the exact structural changes down to the single amino acid level. These results open new perspectives in the development of medicines that might interfere in the course of infection much earlier than antibiotics.



You can clearly see the two membranes (orange) enclosing the cell interior (blue) and the needles protruding to the outside (orange, marked in blue). The bar corresponds to a micrometer or a thousandth millimeter, respectively. Credit:



Ulrike Abu Abed, Diane Schad, and Michael Kolbe

These so-called anti-infectives could inhibit the assembly of the needle and the injection of <u>virulence factors</u> into the host cell. This would be a major advantage over antibiotics, which have to travel through the membrane into the bacteria to be able to kill it. Furthermore, <u>antibiotics</u> cannot distinguish between good and evil, i.e. disease-causing, <u>bacteria</u>, often leading to unwanted side effects. Lastly, the use of anti-infectives would circumvent the problem of antibiotic resistance development.

The change of the three-dimensional structure of the proteins during the needle assembly was analyzed by X-ray structural experiments at BESSY in Berlin and ESRF in Grenoble and NMR-spectroscopic experiments based on radio waves at the Max Planck Institute for Biophysical Chemistry. The scientists compared the three-dimensional structure of the needle protein before and after the needle assembly.

More information: Ömer Poyraz, et al.: Protein refolding is required for assembly of the Type three secretion needle, Nature Structural & Molecular Biology, 13 June 2010

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