

## The 10-foot-tall microscopes helping combat world's worst diseases

October 30 2018



A section of a b2m amyloid fibril, with the atomic model for 8 copies of the b2m protein built into the EM density. The fibril is built from two 'protofilaments' that wrap around each other (blue and pink). This section is around 4 billionths of a metre long Credit: University of Leeds

The century old mission to understand how the proteins responsible for amyloid-based diseases such as Alzheimer's, Huntingdon's and Parkinson's work has taken major steps forward in the last 12 months, thanks to a revolution in a powerful microscopy technique used by scientists.



High-powered microscopes using electrons instead of light to 'see' the actual shape of samples put under them, at near atomic-levels of detail, have only recently become available to UK scientists.

The UK has invested heavily in the 'game-changing' cryo-electron microscopes, but there are still fewer than 25 of the multi-million pound instruments in UK universities and research institutes.

The two instruments at the University of Leeds, funded by the University itself and Wellcome, are the only ones of their kind in the north of England.

They have already proved their worth as a key tool for scientists who have used them in a number of research projects, but have just delivered their biggest success yet: to reveal the structure of amyloid—a build-up of abnormal proteins in the body that causes disease.

There are less than 10 good quality images and structures of these kinds of proteins available to study in the world, so the Leeds' research makes a significant contribution to scientists' understanding of how proteins form aggregates and how they might contribute to amyloid disease.

The images and 3-D structures of the <u>protein</u> aggregates—which the Leeds scientists showed formed long, twisted fibres—has been published in the journal *Nature Communications*. The protein involved- $\beta$ 2-microglobulin—is normally involved in a healthy immune system, but can assemble into the pain-causing amyloid fibres in people who undergo long term dialysis for kidney failure. When they lodge in people's joints they can cause osteoarthritis

It is anticipated the findings will be used by drug manufacturers and research groups internationally who strive to fund cures for <u>amyloid</u> <u>diseases</u> of all types.



Professor Sheena Radford FMedSci, FRS and Professor Neil Ranson from the University's Astbury Centre for Structural Molecular Biology, led the five year programme to image the protein fibres and show their 3-D structure.

The pair were supported by colleagues at Leeds, Josh Boardman, who at the time was an undergraduate student in Biochemistry.



Dr Rebecca Thompson, the University of Leeds' Cryo-Electron Microscopy Centre Manager loads a sample into one of the microscopes. Credit: University of Leeds

The study also involved a long-standing collaboration with Professor Bob Griffin, from the Massachusetts Institute for Technology, who specialises in another method of advanced biological analysis of biological matter- solid- state nuclear magnetic resonance.



Professor Radford said: "Over the past six decades since the first electron microscopy pictures of amyloid were created, scientists have progressed from working with blurred low-resolution images to our razor sharp 3-D images and structures, thanks to modern advances in <u>cryo-electron microscopy</u>.

"Now we know exactly where each kink and point is on the protein, we may be able to develop compounds which lock tightly to it, or disrupt it, and find out how the fibres contribute to disease. It's the equivalent of going from trying to make two balloons stick together to having two cogs rotating perfectly with each other.

She added: "We've used cryo-electron microscopy not only to uncover the shape and structure of <u>amyloid proteins</u>, but also how they grow and intertwine with each other like the stands in a rope to form larger assemblies. This knowledge is going to be crucial for knowing how to deal with them."

Professor Ranson said: "Until a year or so ago, scientists knew the structure looked more or less like a ladder, but we have now shown it is much more complex than that. We're now beginning to see how different proteins folded up into different shapes and how those vary with every disease they cause.

"The extra detail we have uncovered means we can start to understand these proteins' disease-causing abilities.

He added "Amyloid fibres are also known to have the strength of steel, and, now we understand their structures.we might be able to make new biomaterials inspired by their structures. This is a great example of where cryo-electron microscopy can have added advantages."

Knowing the structure of the protein in the level of detail the Leeds



researchers have provided, and measuring those differences in different types of amyloid disease and different patients, could also allow doctors to show who would be most at risk, meaning treatment can be targeted to those who need it most.

The next step for the science community is to begin identifying and developing inhibitors' - compounds which can control protein assembly into amyloid. Professor Radford has secured almost £2million from Wellcome to carry out this stage of development.

Further lab trials, clinical trials, regulatory approval and the involvement of a drug developer would still be required before drugs could be brought to market, but the significant steps forward in image clarity and understanding of the amyloid folding structure mark a major leap forward.

The full research paper The <u>structure</u> of a  $\beta$ 2microglobulin fibril suggests a molecular basis for its <u>amyloid</u> polymorphism is published in *Nature Communications*.

**More information:** Matthew G. Iadanza et al, The structure of a  $\beta$ 2-microglobulin fibril suggests a molecular basis for its amyloid polymorphism, *Nature Communications* (2018). <u>DOI:</u> <u>10.1038/s41467-018-06761-6</u>

## Provided by University of Leeds

Citation: The 10-foot-tall microscopes helping combat world's worst diseases (2018, October 30) retrieved 28 June 2024 from <u>https://phys.org/news/2018-10-foot-tall-microscopes-combat-world-worst.html</u>



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