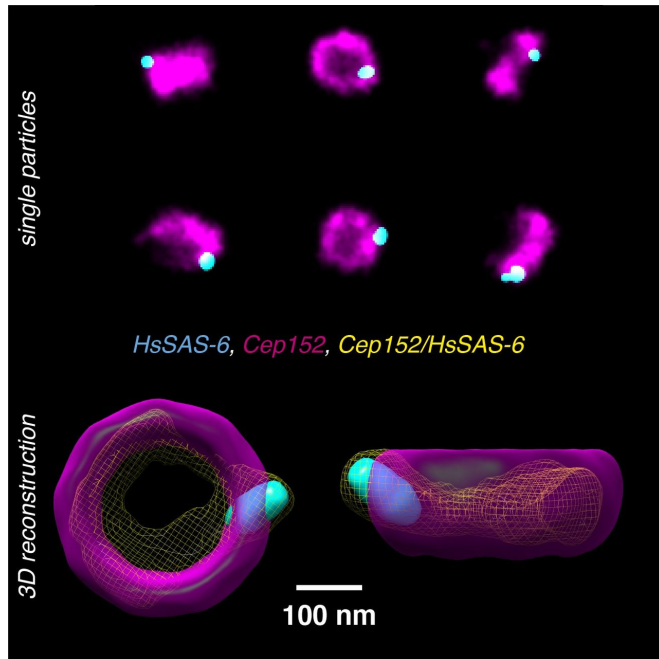


Super-resolution microscopy builds multicolor 3-D from 2-D

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Human centrioles labelled with antibodies against two proteins (Cep152, HsSAS-6) and imaged using super-resolution microscopy. From many individual particles showing projections of the centriole complex in various orientations (upper panel), by using a fused intermediate (yellow, lower panel), the newly developed method allows now to reconstruct a multicolor 3D model (lower panel). Credit: Christian Sieben/EPFL

Super-resolution microscopy is a technique that allows researchers to see beyond the diffraction limit of light. The technique has garnered increasing interest, especially since its developers won the Nobel Prize in Chemistry in 2014. By exploiting fluorescence, super-resolution microscopy now allows scientists to observe cells and their interior structures and organelles in a way never before possible.

Many of the molecular complexes inside cells are made up of multiple proteins. Since current

techniques of super-resolution [microscopy](#) typically only use one or two fluorescent colors, it is difficult to observe different proteins and decipher the complex architecture and underlying assembly mechanisms of the cell's interior structures. An even greater challenge is to overcome the noise inherent to the super-resolution methods and fluorescent labeling, to achieve the full resolution potential.

Scientists from the lab of Suliana Manley at EPFL have now solved both problems by developing a new method to analyze and reconstruct super-resolution images and re-align them in a way that multiple proteins can be placed within a single 3-D volume. The method works with images taken with large field-of-view [super-resolution microscopy](#), with each image containing hundreds of two-dimensional projections of a labeled structure in parallel.

Each 2-D view represents a slightly different orientation of the structure, so that with a dataset of thousands of views, the method can computationally reconstruct and align the 2-D images into a 3-D volume. By combining information from a large number of single images, the noise is reduced and the effective resolution of the 3-D reconstruction is enhanced.

With the help of Pierre Gönczy's lab at EPFL, the researchers tested the method on human centriole complexes. Centrioles are pairs of cylindrical molecular assemblies that are crucial in helping the cell divide. Using the new multicolor super-resolution reconstruction method, the researchers were able to uncover the 3-D architecture of four proteins critical for centriolar assembly during organelle biogenesis.

The new approach allows for unlimited multiplexing capabilities. "With this method, if the proteins in the [structure](#) can be labeled, there is no limit to the number of colors in the 3-D reconstruction," says

Suliana Manley. "Plus, the [reconstruction](#) is independent of the super-resolution method used, so we expect this analysis method and software to be of broad interest."

More information: Christian Sieben et al, Multicolor single-particle reconstruction of protein complexes, *Nature Methods* (2018). DOI: [10.1038/s41592-018-0140-x](https://doi.org/10.1038/s41592-018-0140-x)

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