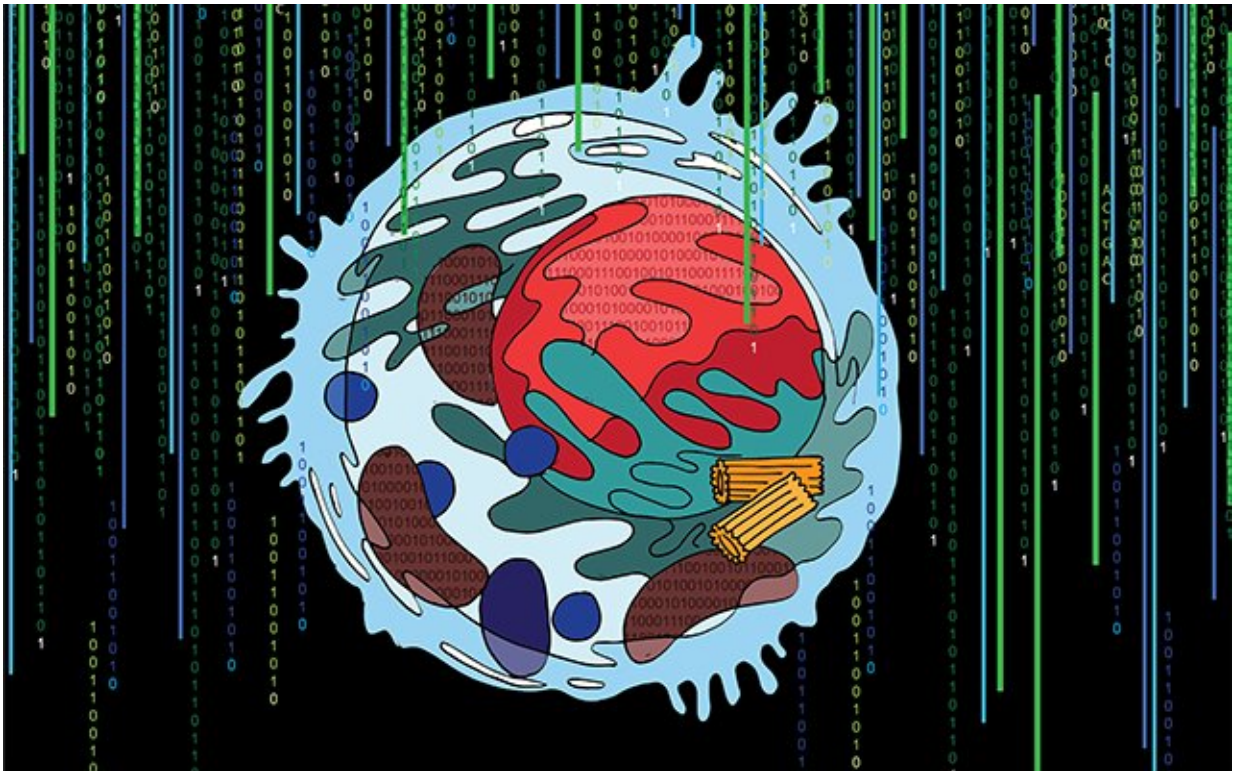


We might not know half of what's in our cells, new AI technique reveals

November 24 2021



UC San Diego researchers introduce Multi-Scale Integrated Cell (MuSIC), a technique that combines microscopy, biochemistry and artificial intelligence, revealing previously unknown cell components that may provide new clues to human development and disease. (Artist's conceptual rendering.). Credit: UC San Diego Health Sciences

Most human diseases can be traced to malfunctioning parts of a cell—a

tumor is able to grow because a gene wasn't accurately translated into a particular protein or a metabolic disease arises because mitochondria aren't firing properly, for example. But to understand what parts of a cell can go wrong in a disease, scientists first need to have a complete list of parts.

By combining microscopy, biochemistry techniques and [artificial intelligence](#), researchers at University of California San Diego School of Medicine and collaborators have taken what they think may turn out to be a significant leap forward in the understanding of human cells.

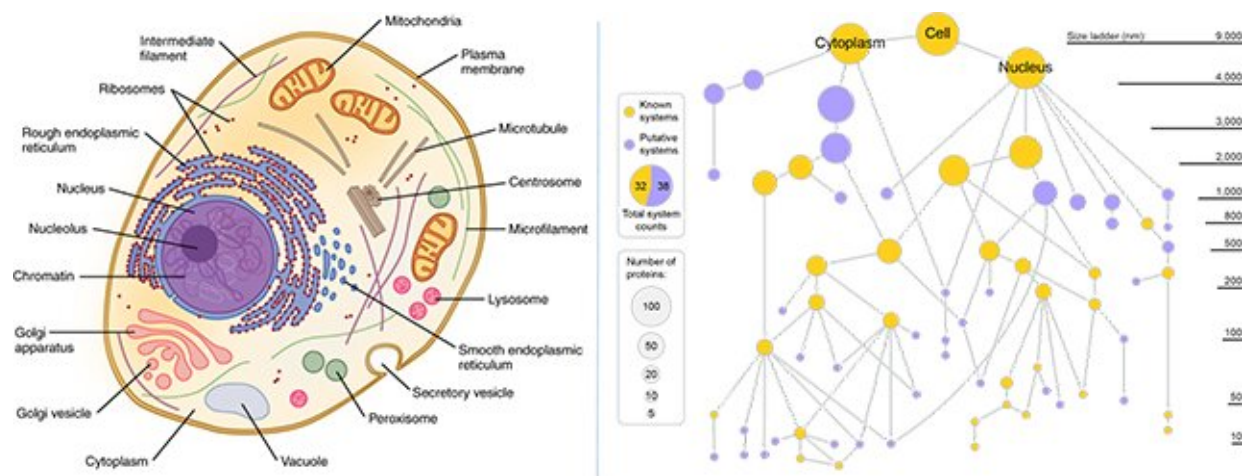
The technique, known as Multi-Scale Integrated Cell (MuSIC), is described November 24, 2021 in *Nature*.

"If you imagine a cell, you probably picture the colorful diagram in your cell biology textbook, with mitochondria, endoplasmic reticulum and nucleus. But is that the whole story? Definitely not," said Trey Ideker, Ph.D., professor at UC San Diego School of Medicine and Moores Cancer Center. "Scientists have long realized there's more that we don't know than we know, but now we finally have a way to look deeper."

Ideker led the study with Emma Lundberg, Ph.D., of KTH Royal Institute of Technology in Stockholm, Sweden and Stanford University.

In the [pilot study](#), MuSIC revealed approximately 70 components contained within a human kidney cell line, half of which had never been seen before. In one example, the researchers spotted a group of proteins forming an unfamiliar structure. Working with UC San Diego colleague Gene Yeo, Ph.D., they eventually determined the structure to be a new complex of proteins that binds RNA. The complex is likely involved in splicing, an important cellular event that enables the translation of genes to proteins, and helps determine which genes are activated at which times.

The insides of cells—and the many proteins found there—are typically studied using one of two techniques: microscope imaging or biophysical association. With imaging, researchers add florescent tags of various colors to proteins of interest and track their movements and associations across the microscope's field of view. To look at biophysical associations, researchers might use an antibody specific to a protein to pull it out of the cell and see what else is attached to it.



Left: Classic textbook cell diagrams imply all parts are clearly visible and defined. (Credit: OpenStax/Wikimedia). Right: A new cell map generated by MuSIC technic reveals many novel components. Gold nodes represent known cell components, purple nodes represent new components. The size of node reflects number of distinct proteins in that component. Credit: UC San Diego Health Sciences

The team has been interested in mapping the inner workings of cells for many years. What's different about MuSIC is the use of deep learning to map the cell directly from cellular microscopy images.

"The combination of these technologies is unique and powerful because

it's the first time measurements at vastly different scales have been brought together," said study first author Yue Qin, a Bioinformatics and Systems Biology graduate student in Ideker's lab.

Microscopes allow scientists to see down to the level of a single micron, about the size of some organelles, such as mitochondria. Smaller elements, such as individual proteins and protein complexes, can't be seen through a microscope. Biochemistry techniques, which start with a single [protein](#), allow scientists to get down to the nanometer scale.

"But how do you bridge that gap from nanometer to micron scale? That has long been a big hurdle in the biological sciences," said Ideker, who is also founder of the UC Cancer Cell Map Initiative and the UC San Diego Center for Computational Biology and Bioinformatics. "Turns out you can do it with artificial intelligence—looking at data from multiple sources and asking the system to assemble it into a model of a cell."

The team trained the MuSIC artificial intelligence platform to look at all the data and construct a model of the cell. The system doesn't yet map the cell contents to specific locations, like a textbook diagram, in part because their locations aren't necessarily fixed. Instead, component locations are fluid and change depending on cell type and situation.

Ideker noted this was a pilot study to test MuSIC. They've only looked at 661 proteins and one cell type.

"The clear next step is to blow through the entire human cell," Ideker said, "and then move to different cell types, people and species. Eventually we might be able to better understand the molecular basis of many diseases by comparing what's different between healthy and diseased [cells](#)."

More information: Trey Ideker, A multi-scale map of cell structure

fusing protein images and interactions, *Nature* (2021). [DOI: 10.1038/s41586-021-04115-9](https://doi.org/10.1038/s41586-021-04115-9).
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