

AI speeds up drug design for Parkinson's by ten-fold

April 17 2024



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Researchers have used artificial intelligence techniques to massively accelerate the search for Parkinson's disease treatments.

The researchers, from the University of Cambridge, designed and used an AI-based strategy to identify compounds that block the clumping, or aggregation, of alpha-synuclein, the protein that characterizes Parkinson's.

The team used machine learning techniques to quickly screen a chemical library containing millions of entries, and identified five highly potent compounds for further investigation.

Parkinson's affects more than six million people worldwide, with that number projected to triple by 2040. No disease-modifying treatments for the condition are currently available. The process of screening large chemical libraries for <u>drug candidates</u>—which needs to happen well before potential treatments can be tested on patients—is enormously time-consuming and expensive, and often unsuccessful.

Using machine learning, the researchers were able to speed up the initial screening process by ten-fold, and reduce the cost by a thousand-fold, which could mean that potential treatments for Parkinson's reach patients much faster. The <u>results</u> are reported in the journal *Nature Chemical Biology*.

Parkinson's is the fastest-growing neurological condition worldwide. In the UK, one in 37 people alive today will be diagnosed with Parkinson's in their lifetime. In addition to motor symptoms, Parkinson's can also affect the gastrointestinal system, <u>nervous system</u>, sleeping patterns, mood and cognition, and can contribute to a reduced quality of life and significant disability.

Proteins are responsible for important cell processes, but when people



have Parkinson's, these proteins go rogue and cause the death of nerve cells. When proteins misfold, they can form abnormal clusters called Lewy bodies, which build up within brain cells stopping them from functioning properly.

"One route to search for potential treatments for Parkinson's requires the identification of small molecules that can inhibit the aggregation of alpha-synuclein, which is a protein closely associated with the disease," said Professor Michele Vendruscolo from the Yusuf Hamied Department of Chemistry, who led the research. "But this is an extremely time-consuming process—just identifying a lead candidate for further testing can take months or even years."

While there are currently <u>clinical trials</u> for Parkinson's currently underway, no disease-modifying drug has been approved, reflecting the inability to directly target the molecular species that cause the disease.

This has been a major obstacle in Parkinson's research, because of the lack of methods to identify the correct molecular targets and engage with them. This technological gap has severely hampered the development of effective treatments.

The Cambridge team developed a machine learning method in which chemical libraries containing millions of compounds are screened to identify small molecules that bind to the amyloid aggregates and block their proliferation.

A small number of top-ranking compounds were then tested experimentally to select the most potent inhibitors of aggregation. The information gained from these experimental assays was fed back into the machine learning model in an iterative manner, so that after few iterations, highly potent compounds were identified.



"Instead of screening experimentally, we screen computationally," said Vendruscolo, who is co-Director of the Centre for Misfolding Diseases. "By using the knowledge we gained from the initial screening with our <u>machine learning model</u>, we were able to train the model to identify the specific regions on these <u>small molecules</u> responsible for binding, then we can re-screen and find more potent molecules."

Using this method, the Cambridge team developed compounds to target pockets on the surfaces of the aggregates, which are responsible for the exponential proliferation of the aggregates themselves. These compounds are hundreds of times more potent, and far cheaper to develop, than previously reported ones.

"Machine learning is having a real impact on the drug discovery process—it's speeding up the whole process of identifying the most promising candidates," said Vendruscolo. "For us this means we can start work on multiple drug discovery programs—instead of just one. So much is possible due to the massive reduction in both time and cost—it's an exciting time."

More information: Discovery of Potent Inhibitors of α-Synuclein Aggregation Using Structure-Based Iterative Learning, *Nature Chemical Biology* (2024). DOI: 10.1038/s41589-024-01580-x

Provided by University of Cambridge

Citation: AI speeds up drug design for Parkinson's by ten-fold (2024, April 17) retrieved 21 May 2024 from <u>https://medicalxpress.com/news/2024-04-ai-drug-parkinson-ten.html</u>



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