Targeting SARS-CoV-2 with AI- and HPC-enabled Lead Generation

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Small-molecule therapeutics against COVID

The SARS-CoV-2 virus comprises 28 unique proteins

Small-molecule therapeutics seek to disrupt its functioning

Size and Content
Diameter ~100nm

Nucleoprotein
Membrane protein
Envelope protein
Spike trimer
Structures exist for most of the key proteins
Leveraging Argonne resources in the fight

Serial X-Ray Crystallography (SSX)
Solving complex COVID protein structures with high-throughput imaging at near room temperature (structures provide insight into drug and vaccine development)

Extreme-scale machine learning
Applying machine learning models to screen potential therapeutics based on molecule databases and computed molecular features
Enabling Serial Crystallography (SSX) at Scale

- Perform serial imaging of chips with thousands of embedded protein crystals
- Analyze batches of images as collected
- Report statistics and summary images during experiment
- Return crystal structure to scientist

Connecting light sources and leadership computing facilities to enable new science

Ryan Chard et al.
Automating and scaling the analysis of SSX data

Globus Automate flow to batch files, move data to ALCF, perform analysis using funcX/Parsl, and catalog results

Integrates with APS DM to trigger flow, ALCF resources for computing, and ALCF portal for monitoring experiments and reprocessing data

With Andrzej Joachimiak, Darren Sherrell et al. APS Sector 19
“These data services have taken the time to solve a structure from weeks to days and now to hours”
Darren Sherrell, SBC beamline scientist APS Sector 19

ALCF + APS capabilities were used to determine the room temperature structure of 2 viral surface proteins

4 structures are now available in PDB
Challenges screening potential candidates

- Many molecules (>10^9 drug-like molecules in collected databases)
- Testing in the wet lab is very expensive, clinical trials even more so
- Protein docking simulations are computationally expensive

→ apply machine learning methods to predict which molecules have a high likelihood of docking
AI and supercomputers can accelerate drug discovery
Parsl-computed features for AI-based drug screening

23 input datasets, 4.2B molecules, 60 TB of molecular features and representations

Parsl processing pipelines used ~2M core hours on ALCF Theta, TACC Frontera, OLCF Summit

1. Convert each molecule to a canonical SMILES
2. For each molecule, compute:
   a. ~1800 2D and 3D molecular descriptors using Mordred
   b. Molecular fingerprints encoding structure
c. 2D images of the molecular structure

Computed data provide crucial input features to AI models for predicting molecular properties such as docking scores and toxicity

https://2019-ncovgroup.github.io/data/

Canonical SMILES
23 CSV files with 4.2B molecules

Mordred Descriptors
420,130 CSV files, 48.70TB

Molecular Fingerprints
4,221 CSV files with base64 encoded fingerprints, 578.27GB

2D images
420,707 Pickle GZ files, 11.48 TB