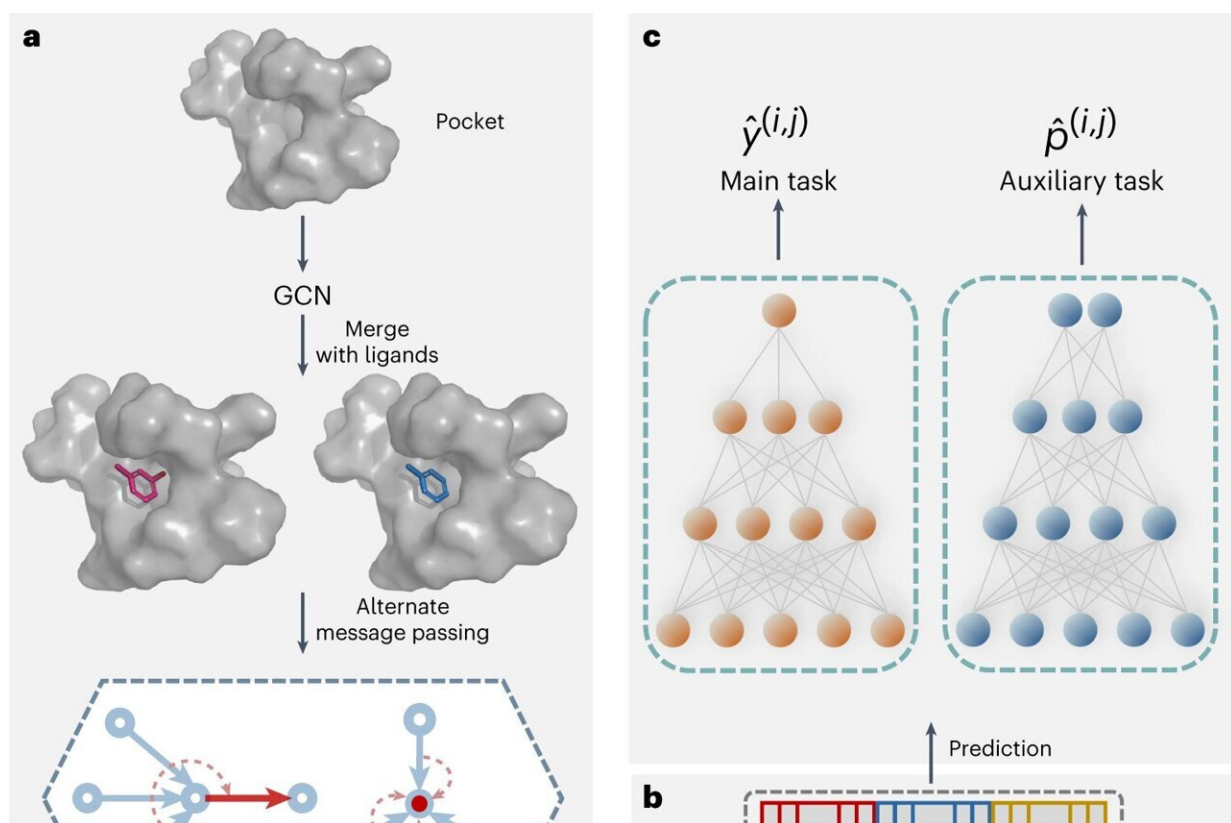


New tool developed to efficiently predict relative ligand binding affinity in drug discovery

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The framework of PBCNet. Credit: *Nature Computational Science* (2023). DOI: 10.1038/s43588-023-00529-9

Lead optimization in drug discovery is a challenging process that heavily

relies on hypotheses and the experience of medicinal chemists. This often leads to uncertain outcomes and inefficiency. Furthermore, the process is time-consuming and requires significant resources. Therefore, the introduction of artificial intelligence (AI) predictive tools to accelerate this process would be highly valuable in the field of drug discovery.

Silico methods such as [free energy](#) perturbation (FEP) and molecular mechanics generalized born [surface area](#) (MM-GB/SA) have proven useful in lead [optimization](#) by calculating binding free energy. However, their complex preparation process, limited molecule throughput, and restricted allowance for changes between molecules hinder their routine usage. There is an urgent need to develop an efficient and accurate in silico predictive tool to guide lead optimization.

In a study [published in *Nature Computational Science*](#), a team of researchers led by Prof. Zheng Mingyue from the Shanghai Institute of Materia Medica (SIMM) of the Chinese Academy of Sciences developed a pairwise binding comparison network (PBCNet).

This network predicts the relative binding affinity among congeneric ligands by using a physics-informed graph attention mechanism with a pair of protein pocket-ligand complexes as input. PBCNet demonstrates practical value in guiding structure-based drug lead optimization with its speed, precision, and ease-of-use.

To validate the performance of PBCNet in terms of ranking ability and accuracy, Zheng's group used two held-out sets provided by Schrodinger, Inc. and Merck KGaA. These sets comprised over 460 ligands and 16 targets. Transfer learning was applied in their work, involving pretraining models on large-scale datasets and fine-tuning them for tasks with limited data. This approach specifically improved the performance of the models on the tasks.

Benchmarking results obtained from the test data showed that pretrained PBCNet outperformed Schrodinger's Glide, MM-GB/SA, and four recently reported deep learning models (DeltaDelta, Default2018, Dense, and PIGNet). Furthermore, with a small amount of fine-tuning data (2–10 ligands with known binding activity), PBCNet achieved performance comparable to that of Schrodinger's FEP+, which is considered the standard computational lead optimization method in the pharmaceutical industry.

The researchers also tested whether PBCNet could efficiently identify key high-activity compounds in a real-world lead optimization scenario. They used a benchmark consisting of nine recently-published chemical series and compared the order of model selection to the experimental order of synthesis.

The evaluation demonstrated that after using PBCNet, the tested lead optimization projects were accelerated by approximately 473% while resource investment was reduced by an average of 30%.

This study showed that PBCNet has immediate practical value in guiding lead optimization projects. Additionally, there is a free academic [web service](#) available that utilizes PBCNet to predict ligand binding affinity.

AI has become increasingly important in solving scientific problems by incorporating domain-specific knowledge into its models. PBCNet exemplifies this approach by integrating physical and a priori knowledge into its modeling process.

More information: Jie Yu et al, Computing the relative binding affinity of ligands based on a pairwise binding comparison network, *Nature Computational Science* (2023). [DOI: 10.1038/s43588-023-00529-9](https://doi.org/10.1038/s43588-023-00529-9)

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