

Apr 05, 2022

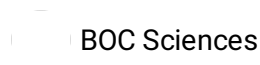
Virtual Screening

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[dx.doi.org/10.17504/protocols.io.n92ldzr6ov5b/v1](https://doi.org/10.17504/protocols.io.n92ldzr6ov5b/v1)

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Virtual screening has been widely applied in early-stage drug discovery. As an alternative or complementary approach to high-throughput screening (HTS) assays with high cost and low hit rate, [virtual screening](#) is an efficient computational method to identify drug candidates in silico from large chemical compound databases. Its usefulness has been verified by current applications that successfully retrieved hit and lead identifications against various disease targets.

DOI

[dx.doi.org/10.17504/protocols.io.n92ldzr6ov5b/v1](https://doi.org/10.17504/protocols.io.n92ldzr6ov5b/v1)<https://www.solutions.bocsci.com/virtual-screening.htm>BOC Sciences 2022. Virtual Screening. **protocols.io**<https://dx.doi.org/10.17504/protocols.io.n92ldzr6ov5b/v1>

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