

Review of: "Design and Molecular Screening of Various Compounds by Molecular Docking as BACE-1 Inhibitors"

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Potential competing interests: No potential competing interests to declare.

Good topic but needs a lot of improvement. Figures of chemical structures need to be uniform - Please use ChemDraw and correct the settings.

Figure 17, B, and Figure 20, B - Please explain why the unfavourable bond is there? And yet you have chosen them.

What about the validation of docking results? And no MD simulation data?