

# In silico identification of potential inhibitors for human aurora kinase b

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# Key points

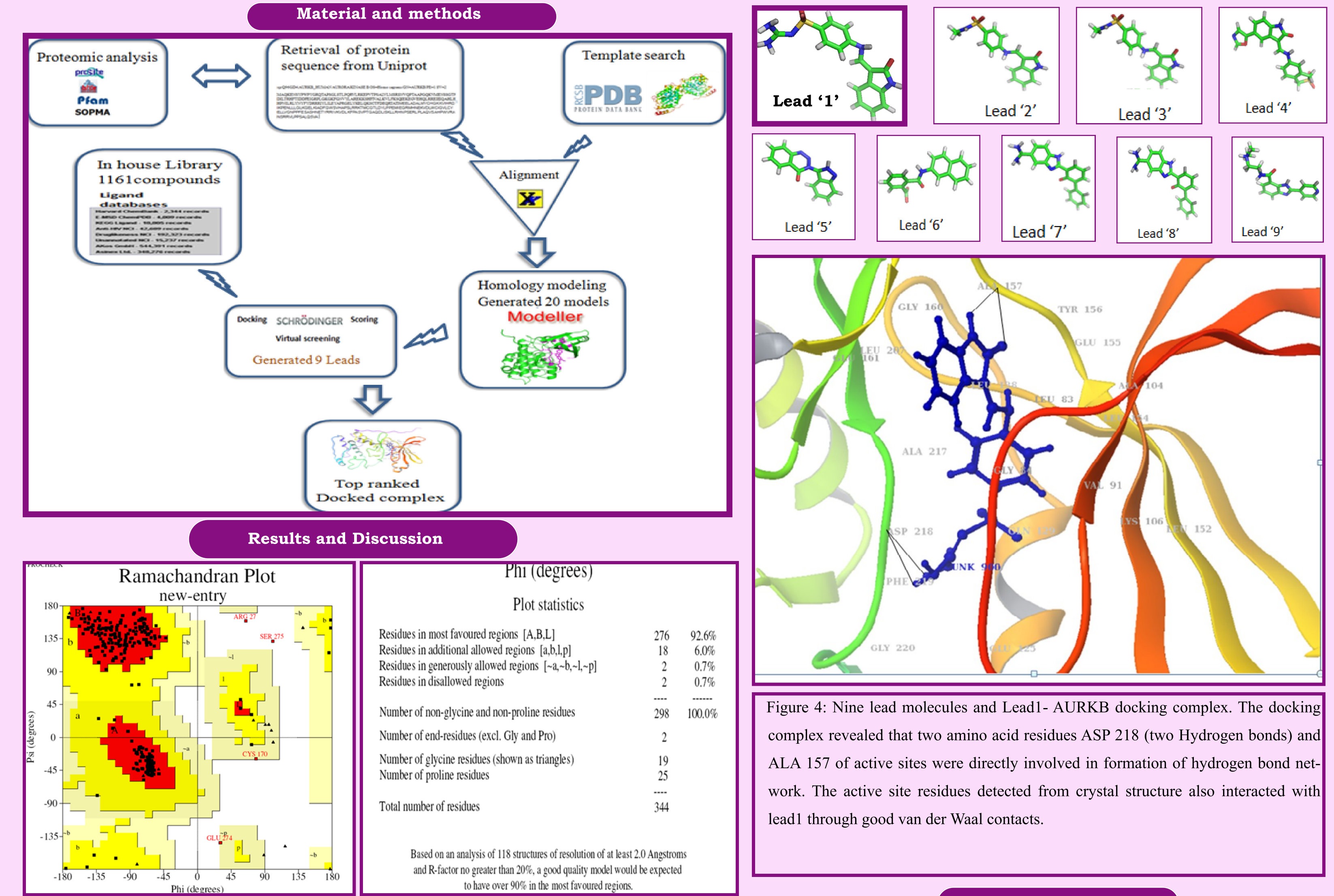
• Cell cycle progression through mitosis and meiosis involves regulation by serine or threonine kinases from aurora family.

• Human aurora kinase b (AURKB) is a protein mainly involved in the proper segregation of chromosomes during mitosis as well as meiosis.

• Over expression of AURKB leads to the unequal distribution of genetic information creating a aneuploid cells, a hallmark of cancer. and this heads to genetic instability is linked on primary nonsmall cell lung carcinoma.

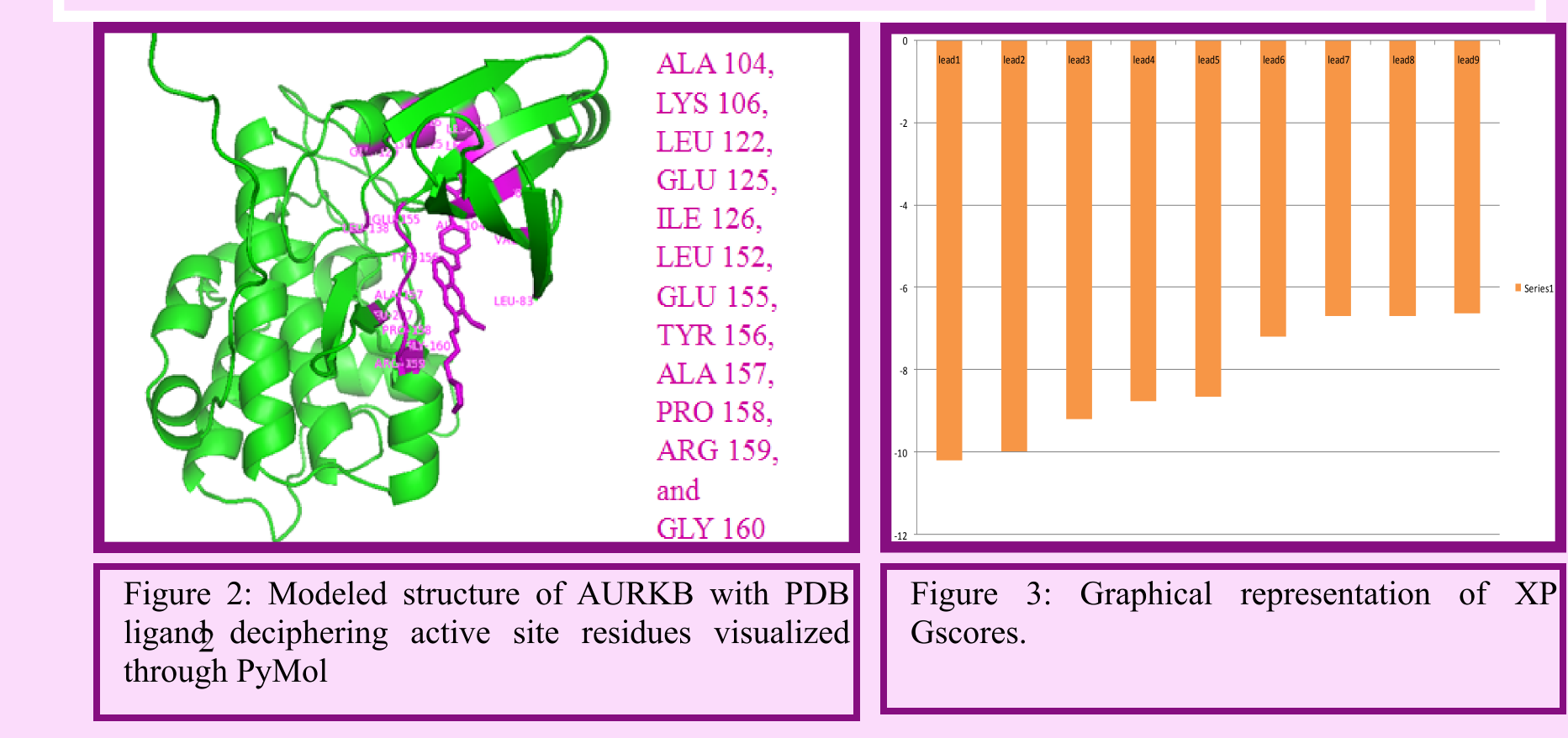
• Inhibition of AURKB results inhibition of cytokinesis (or anticytokinesis), hence is an attractive anticancer strategy.

#### • In silico work was carried out to identify novel potent inhibitors towards human AURKB.



#### Conclusion

## Figure 1: Ramachandran plot for modeled Human AURKB



Analysis of the Aurkb 3D model had revealed that ALA 157 an essential amino

acid for AURKB activity is directly getting blocked by lead 1 by forming hydro-

gen bond and good van der wall interactions.

Thus it would be highly effective as a novel inhibitor towards of human AURKB

protein for treatment of metastasis.

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