Computer–based screening for novel inhibitors of human topoisomerase I with FlexiDock docking protocol

Choi InHee¹, Kim Choonmi
College of Pharmacy, Ewha Womans University, Seoul, Korea

DNA topoisomerases I (topo I) and II are essential enzymes that relax DNA supercoiling and relieve torsional strain during DNA processing, including replication, transcription, and repair. Topo I relaxes DNA by cleaving one strand of DNA by attacking a backbone phosphate with a catalytic tyrosine (Tyr723, human topo I). This enzyme has recently been investigated as a new target for antineoplastic drugs. Inhibitors to the enzyme intercalate between the DNA base pairs, interfering religation of cleaved DNA, therefore inhibit the activity of topo I. To search novel inhibitors of topo I from existing chemicals, molecules that are known to act as antitumor, anti-inflammatory or anti–invasion drugs have been selected and docked into the human topo I–DNA complex. Among 13 molecules that have been tested, seven molecules showed intercalative binding modes like known inhibitors. Their structures are mainly composed of multi–rings and planar which are characteristics required to intercalate DNA. Although these results should be substantiated by further biological activity study with human topo I, the docking results suggest the possibility of these molecules being novel topo I inhibitors.

Binding modes of artemisinin to malarial TCTP demonstrated by computer modeling

Chai Jinsun², Kim Choonmi
College of Pharmacy, Ewha Womans University, Seoul, Korea

The translationally controlled tumor–associated proteins (TCTPs) are a highly conserved and abundantly expressed family of eukaryotic proteins that are implicated in both cell growth and human acute allergic response but whose intracellular biochemical function has remained elusive. There are reports that antimalarial drug, artemisinin, binds to Plasmodium falciparum TCTP; however, its 3D structure has not been known. To illustrate the action mechanism of artemisinin, 3D structure of P. falciparum TCTP was constructed by homology modeling using NMR structure of Schizosaccharomyces pombe TCTP (PDB code 1H6Q) as a template. Whose sequence is 39% identical and 56% similar to P. falciparum TCTP. The final model was chosen out of 5 models obtained after evaluation by PROSAIL and PROCHECK. With this structure, docking experiment was carried out with Flexidock to determine the binding modes between the protein and the ligand. Since the TCTP has been shown to react with dihydroartemisinin in the presence of heme, docking simulation of artemisinin with heme was first performed and then the activated artemisinin was docked into the P. falciparum TCTP. The results show that the activated C⁴ of artemisinin interacts with CYS14 of the TCTP, conforming the experimental data reported.

β–ketoacyl–acyl carrier protein synthases for fatty acid biosynthesis in bacteria

Seoul Women’s University

A universal set of genes encodes the components of dissociated, type II, fatty acid synthase system that is responsible for producing the multitude of fatty acid structures found in bacterial membranes. We examined the biochemical basis for the production of fatty acids by bacteria. Several genes from Haemophilus influenzae Rd