

NGF, BDNF, NT3/4, NT3 and their receptors displayed crucial roles as proliferation, migration, differentiation, survival, apoptosis, In previous studies improved TRK B is as oncogene agent and BDNF bind to trk b and active signaling angiogenesis for tumor proliferation. In the current century using intelligent biomolecule such as antibody and peptides is hopeful therapies in cancers. In this study we design of new peptides by using monte carlo methods and study the effect of them on cancer cell lines. For this aim we use backrub protocol and docking with trkB as receptor. our result showed that this protocol could improve the peptide design for cancer treatment. At the first step of this protocol we designed peptide library by using sequence tolerance method in rosetta3.3 package, then peptide energy optimization performed by backrub protocol for finding peptides with more stability, the five of best peptides selected based on R software and peptide 3D-structure prediction performed by using molecular dynamic in Hyperchem 7 software. the final step is Docking of peptides with receptor trkb in HADDOCK then cyclotraxin and designed peptides.

**Keywords:** TrkB inhibitor, Cancer Treatment, Peptide Desining, Rosetta3.3 Package.

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#### Abstract No.194

##### **A Study on the Interaction of Chickpea Seedling Copper Diamine Oxidases by Tetraethylen Pentamine**

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Copper amine oxidase are soluble dimeric enzymes, each monomer contains one Cu(II) ion and one organic prosthetic group 2,4,5-trihydroxyphenylalanine quinone as cofactors. Inhibitors represent important role in the study of catalytic properties of copper amine oxidase and they also find a wide application in physiological research. They catalyze the oxidative deamination of primary amines to aldehydes with a ping-pong mechanism consisting of a transamination, followed by the transfer of two electrons to molecular oxygen which is reduced to H<sub>2</sub>O<sub>2</sub>. Kinetic parameters K<sub>m</sub> and V<sub>max</sub> of purified enzyme by analysis of Lineweaver - Burk plot was determined 3.3 mM and 0.95 mmol/min/mg, respectively. In this study interaction chickpea diamine oxidase with tetraethylen pentamine was studied. Analysis of kinetic

data indicated considerable inhibitory effects for tetraethylen pentamine. Results showed that in the presence of tetraethylen pentamine reduced apparent K<sub>m</sub> and V<sub>max</sub> i.e. tetraethylen pentamine with K<sub>i</sub>=0.1 mM inhibits the enzyme by linear mixed inhibitory effect. In linear mixed inhibition, the inhibitor can bind to the enzyme at the same time as the substrate. However, the binding of the inhibitor affects the binding of the substrate, and vice versa. This type of inhibition can be reduced, but not overcome by increasing concentrations of substrate.

**Keywords:** Chickpea, Copper-Containing Amine Oxidases, Tetraethylen Pentamine, Linear Mixed.

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#### Abstract No.195

##### **Synthesis of Cyclooxygenase Inhibitor 2-(1-benzyl-alkyl thio-5-imidazolyl)-3-phenyl-1, 3-Thiazolidin-4-one as Anticancer Agent**

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Recently study was shown that enzyme cyclooxygenase have a role in cancer tissue. Since the derivatives of thiazolidine 4-one and other pharmacophore patterns of central ring have a cox-2 inhibitory effect, thus we decide to synthesis of novel derivatives of thiazolidin -4-ones. At first, benzyl amine hydrochloride was produced. Then the mixture of, dihydroxy acetone, thiocyanate potassium was reacting for 72 hours. By alkylation and oxidation formyl imidazole was obtained. The resultant aldehyde was reacted with aniline and thioglicolic in two ways including classic (in one and two step ways) and microwave method until the title compounds were obtained. 1-Classic method: In the one step way, all the reactant (resultant aldehyde, aniline and thioglicolic) were refluxed in dean stark apparatus in dry toluene for 48 hours. B: In the two step way, at first aldehyde and aniline react in dean stark for 24 hours to give imine intermediate, then thioglicolic acid was added and reacted for 24 hours. In two stages: at first aldehyde and aniline react in dean stark for 24 hours after synthesizing dimidiate product, thioglicolic acid was added and react for 24 hours. 2- Microwave method: All of reactant was treated at 800 w, 100 c for 15 minute. TLC was used to evaluate the progress of the reaction and purify material.