

responsible for lens transparency, beta and gamma crystallins possess mainly structural role. As reported previously, the interface of two domains in bovine gamma-B crystallin consisting many residues including Phe56 which is critical for both efficient domain pairing and protein stability. While slight destabilizing outcome has been reported for the Phe56Trp mutation, its replacement with either Ala or Asp resulted in drastic instability impact. In this study, Phe56 was in silico mutated for different residues in both bovine and human gamma-B crystallins and their effects on the protein stability was evaluated using different bioinformatics tools. Bovine gamma-B crystallin and its human counterpart demonstrate 83.9% homology as determined using clustalW software. The results show that all mutations performed at this position, depending on size, polarity and charge of substituted amino acid side chain, leading to range of protein instability in both human and bovine gamma-B crystallins. All substitutions performed resulted in significantly higher destabilizing consequence for human gamma-B crystallin compared to that of bovine protein counterpart, suggesting the impact of fine tertiary structures on stability of these proteins. Furthermore, our results indicated strong destabilizing properties for the substitution of Phe56 with the aliphatic residues of Gly, Ala, Val and Leu, and the degree of destabilization was strongly associated to the size of amino acid side chain. Overall, this study suggests that both polarity and size of amino acid side chains might be considered as the determining factors in fine complementary pairing of two domains which are required for both structural integrity and stability of gamma-B crystallin.

Keywords: Gamma-B crystallin, Stability, Mutation, Domain, Phe56.

Abstract No.138

Engineering of Solvent Exposed Loops of Firefly Luciferase Through Arginine Saturation

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Firefly luciferase (EC 1.13.12.7) catalyses the oxidation of aluciferin in the presence of magnesium ions, ATP and molecular oxygen. The product, oxyluciferin, is generated in an excited state which then decays to the ground state with the emission of a photon. In life science, a wide range of novel invitro and invivo applications of Luciferases has been developed. However, several factors limit further application and development of this technology, including the low

stability of the enzyme both invitro and in vivo, a low turnover number and a high Km for the substrate ATP. Protein engineering studies show that thermostable proteins have higher frequency of Arg, especially in exposed state, which would stabilize the exposed part of protein structure. According to bioinformatics analysis, higher ratios of charged amino acids, especially at the surface, increase ion interactions and enhance occurrence of salt bridges and ion pairs in thermophilic proteins which provide thermal stability to proteins. Here, we have extended this strategy by mutating several hydrophobic residues scattered at the surface of *L. turkestanicus* luciferase to arginine. After production of mutants luciferase, the native and mutants luciferase have overexpressed and purified. Thermostability and other properties were determined and results show that after 40 min incubation of enzymes at 40°C, the relative remaining activity of wildtype was only 5%, whereas was for mutants luciferase was 80% of original activity. Also, half-lives ($t_{1/2}$) of luciferase inactivation are measured and results show that and mutants luciferase are increased to 15 and 19.2 min, which is much higher than wild-type luciferase (2.6 min). We have shown that replacement of uncharged polar and hydrophobic residues by Arg leads to the enhancement of thermostability.

Keywords: Luciferase, Oxyluciferin, Hydrophobic Residues, Thermostability.

Abstract No.139

Tramadol Induces Inhibition and Structural Changes on Kidney Alkaline Phosphatase

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Alkaline phosphatase (ALP) is a hydrolytic enzyme with various type of substrate including proteins, nucleotides and phospholipids. ALP removes phosphate group from the substrate. Tramadol is a narcotic-like pain reliever and uses to treat moderate to severe pain. In this study the effect of tramadol on the alkaline phosphatase activity was investigated. The cell free extract was prepared from balb/c mouse kidney and used for enzyme assay. Our results showed that tramadol inhibited kidney alkaline phosphatase activity. The Km (0.28 mM) of enzyme did not change, while the Vmax of enzyme reduced in the presence of drug. The Ki and IC50 values of tramadol were determined