

#### TRAINING MODULE - MOLECULAR DOCKING AND DYNAMICS

COURSE DURATION: 7 DAYS FEE: 3500 INR

#### **DAY 1: Introduction to Molecular Docking and Computational Biology Tools**

- Fundamentals of Molecular Docking and Drug Discovery (MDD)
- Protein structure and functions insights
- Database overview
- Tools and software walkthrough

# **DAY 2: Protein and Ligand Structure Preparation**

- Preparation of the protein and ligand structure using tools like PyMOL or Chimera to optimize protein structures for docking
- Quality control and structure verification for errors or clashes in prepared protein and ligand structures to ensure accuracy in docking studies

# **DAY 3: Blind Docking and Preliminary Analysis**

- Perform Docking with the prepared protein and ligand structure using AutoDock Vina
- Analysis and interpretation of the docked protein and ligand structure using Chimera-X or Discovery Studio

# **DAY 4: Targeted Docking and Detailed Analysis**

- Perform targeted docking with an emphasis on specific binding sites
- Comparing results with blind docking outcomes to assess pose consistency and energy profiles
- Analysis and interpretation of the docked protein and ligand structure
- Visualizing and interpreting protein-ligand complexes using Chimera-X or Discovery Studio and analysing binding site residues



### **DAY 5: Running MD Simulation**

- Preparing the protein-ligand complex for simulation, solvation, ionization, and energy minimization
- Executing the simulation and observing the stability of the protein-ligand interaction over time using GROMACS

### **DAY 6: Data Analysis and Report Generation**

- Analyzing simulation trajectories: RMSD, RMSF, hydrogen bonds, and interaction stability
- Final interpretation of MD simulation results to assess complex stability and binding reliability

## **DAY 7: Report Submission and Certificate Distribution**

