

Centre of Excellence in Pharmaceutical Sciences (CEPS)

Guru Gobind Singh Indraprastha University (GGSIPU) Sector 16-C, Dwarka, New Delhi.

F.15/GGSIPU/CEPS/2022/ 480

Dated: 24/11/2022

NOTICE

Subject: Invitation for Two day workshop on "Cloud-based Hands-on Workshop: Computational Aided Drug Design Workflow"

Centre of Excellence in Pharmaceutical Sciences (CEPS), Guru Gobind Singh Indraprastha University, Dwarka, Delhi, in collaboration with Schrödinger is going to organize two-day workshop "Cloud-based Hands-on Workshop: Computational Aided Drug Design Workflow" for masters and PhD students.

Requirements: Own laptops are required for the workshop and with appropriate internet speed will be provided by CEPS, GGSIPU. Users can get connected to their laptops by their mobile hotspots as well as Cloud space will be provided and your login credentials will be used for hands-on experiments.

Interested participants would have to fill the registration form for the workshop latest by 28th November, 2022. Participants will be selected on first come first serve basis.

https://forms.gle/nyLAKG7uWy9aSd2UA

Note: Event Brochure (Copy Attached)

Date and Day: 1st (Thursday) and 2nd (Friday) December 2022

Time: 9:30 onwards

Venue: CEPS, GGSIP University

Prof. A.K. Narula (Director, CEPS)





Workshop on

"Cloud-based Hands-on Workshop:

Computational Aided Drug Design Workflow"

Date: 01-02December 2022

Venue: Centre of Excellence in Pharmaceutical Sciences (CEPS), Guru Gobind Singh Indraprastha University (GGSIPU)

Deadline for Registration: 28th November 2022

Research Scholars and postgraduate students are eligible to register for the workshop.

Registration Fee: N/A

Requirements: Laptop/Desktop with at least 10 Mbps Internet connection (mobile hotspots can also be used), and Google Chrome browser

Workshop Highlights: 2-day cloud-based hands-on workshop targeting structure-based drug designing. Participants will get practical experience and in-person guidance in using the Maestro GUI, covering organic molecule sketching, protein selection, preparation, and screening for hit identification of molecules against therapeutic targets. The workshop will also include a brief recap of background theory for Molecular mechanics, Molecular Docking, and Molecular Dynamics, etc. via case studies on the real-time industrial projects.

Coordinators:

Dr A.K. Narula, Director CEPS, GGSIPU and Dr. Parul Sharma, Assistant Professor, CEPS GGSIPU.

Workshop Speakers from Schrödinger:

Dr. Prajwal Nandekar, Dr. Koushik Kasavajhala





Program Details

Day 1				
Time	Workshop Topics			
9:30-10:00 AM	Technical set-up, Audio & Visual Check and			
9:40 AM	Introduction of Speakers Inauguration			
10:00 AM	1. Opening-Molecular Modelling Introductory Presentation			
10:30 AM	2. Logging into Cloud instance			
10:45 AM	3. Maestro GUI 1: 2D Sketcher and 3D Builder			
11:15 AM	4. Maestro GUI 2: Building DNA and Importing SMILES			
11:45 AM	5. Maestro GUI 3: Protein Visualization 6B2Q			
12:15 PM	6. Protein Preparation and Grid Generation 6B2Q			
1:00 PM	Lunch Break			
2:00 PM	7. Ligand Preparation for 6B2Q			
2:10 PM	8. Molecular Docking 6B2Q			
2:40 PM	9. Molecular Docking Analysis 1 – Pose visualization and evaluation			
3:15 PM	10. Molecular Docking Analysis 2 – Ligand Interaction Diagram and Calculation of Interaction Fingerprints			
3:45 – 4:00 PM	Review Day activities and Finish			





Day 2				
Time	Workshop Topics			
10:00AM	1.Opening-Molecular dynamics theory presentation			
10:30AM	2.Logging into cloud instance			
10:45AM	3.Protein Preparation			
11:15AM	4. Desmond Introduction and building your MD			
	simulation system			
11:45AM	5.Desmond Molecular Dynamics Submission			
12:15PM	Break-Continue to uses of software during the break			
2:00PM	Welcome Back			
2:10PM	6.Desmond Molecular Simulation Analysis1–			
	Visual Analysis			
2:40PM	7. Desmond Molecular Simulation Analysis 2 –			
	Quantitative Analysis using Simulation Interaction			
	Diagram			
3:30PM	8.Ligand Designer			
3:50 – 4:00 PM	Review Day 2 Activities and Finish with Concluding			
	Remarks			



DR. B. C. Roy Awardee



Centre of Excellence in Pharmaceutical Sciences (CEPS), GGSIPU, New Delhi

in

Collaboration with Schrödinger

Organising
Two days Workshop on "Computer Aided Drug Design Workflow"
For Masters / Ph.d Students

on

01st & 02nd December 2022

Time: 09.30 a.m. to 04.30 p.m

Venue: CEPS, GGSIPU Campus.

Patron:	Speakers:	Coordinators	
Prof. (Dr.) Mahesh Verma	Dr. Prajwal Nandekar Senior Scientist -I	Prof. A.K Narula,	
Vice Chancellor		Director, CEPS	
Padma Shri Awardee			
National Science and	Dr. Koushik Kasavajhala	Dr. Parul Sharma,	
Technology Awardee	Senior Scientist -II	Assistant Professor, CEPS	