

Lecture and Practice Proceedings & Objectives

- Tutors are present to help you. Contact them!!!
- · Get a flavor of molecular structure and modeling
- Acquire some basic theoretical background
- · Practice the molecular graphics techniques
- Use them to uncover structure-activity relationship
- Get a detailed knowledge about the 3D structures of pMHC and TCR/pMHC

→ You should be able to perform simple tasks of molecular graphics and analysis

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Installing UCSF Chimera In this lecture/practical you will use the software UCSF Chimera for 3D structure visualisation and analysis. This software is: - free for teaching or academic research - available for the most current platforms (Windows, Mac, Linux) - open source (you can modify it for your needs if you know how to code in python. This is out of the scope of this lecture). totes for a lis Plat You can download the latest production release here: do859a Mar 06, 2025 Doc Doc vs 64-bit Size: 152587563 bytes https://www.cgl.ucsf.edu/chimera/download.html chimera-1.19-mac64.dmg Size: 162110856 bytes MDS: Saha28548che684-4647 ac OS X 64-bit Mar 06, 2025 Documentation Runs on Mac chimera-1.19-linux x86_64.bin Size: 155519902 bytes MD5: 995ae1-05-254-50-40-40-40 Mar 06, 2025 Document ux 64-bi Please, install this software on your machine. It will be mandatory for the practicals, but also useful for the 64-bit B theoretical lectures Pla Aar 06, 20 chimera-alpha-win6 Size: 152606292 bytes Aac OS X 64-b Mar 06, 2 chimera-Size: 1621 x86_64.bin Linux 64-bi Mar 06, 2025 chimera-alpha-linux Size: 155536605 bytes MDE: 900xx2665678000 is Linux 64-bit chimera-alpha-linux x86 64 comesa Size: 149451263 bytes MDI: s0ac9719cd763009c30ee06e7281e3b2 Unil 4

The dedicated web site

This teaching has been conceived to alternate theoretical lectures and practicals, so that you will:

- experiment yourself the visualisation and analysis of protein 3D structures
- get a 3D view of the systems mentioned in the lecture (to prevent being limited by the 2D views in the slides)

To facilitate the process, a web site has been especially conceived for this teaching. You can find it here:

http://www.immunology-and-modelling.ch

- This web site will indicate you when to switch between lecture and practicals. For instance, you will be able to make exercices 1 and 2 just after the prologue regarding molecular representation, while exercises 3 to 5 will be made after the lecture section dedicated to MHC-I and before that of MHC-II
- Videos on how to execute exercices 1 to 5 have been made for your help. They are without sound, but all instructions are detailed in the booklet
- 3. The booklet of the practicals and the PDF of the lecture can be downloaded from the web site too



The dedicated web site – Seeing the lecture in 3D Mail Mole ular Modelling for Immunology 4. Once you will be trained in using UCSF Chimera, you will be able to use this tool to see in 3D, on your computer, the systems that are Videos of exercices displayed in 2D in the lecture. here Links to So, when you see this icon of in the lecture slide, you can check the slide download number, and click on the corresponding link on the web site. lecture and exercices Upon clicking, the molecular system should be automatically displayed here in UCSF Chimera, if your browser is managing it correctly. Otherwise, you can download the linked file, with the .chimerax extension, on your machine. Then, double clicking on the file, or drag&droping it on UCSF chimera will open it in this software. Although you are encouraged to do it, to beneficiate from a full 3D "immersion", this is totally optional. This possibility is provided to you to go beyond 2D images in the slides. But if it does not work for you, or if you don't want to use it, it will not Exercise 4. Superposing and comparing 3D strucures. W ALGIGUTV block better to HLA-A*2201 than AAGIGUTV? prevent you from /IART-1/Melan-A understanding the lecture. Side 83 & Side 90. Structure of pMHC Class Open the "Model Panel" to showhide surfaces. 6







































TechniqueAdvantagesDisadvantagesXray crystallographyHigh resolution (1 to 3 Å)Requires to crystallize the protein Does not allow studying transmembrane or very flexible proteinsNMRDoes not require protein crystallization ~ High resolutionGenerally limited to small proteinsCryo-EMDoes not necessitate to crystallize the protein: possible to study transmembrane proteins, and more flexible proteins than Xray.Generally limited to large proteins Low resolution, 4 to 20 Å (a lot of progresses have been done recently)	xperimental methods - Summary						
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Molecular recognition – Historical models "Lock and key" model. Emil Fischer in the 1890s. Induced fit model Daniel Koshland 1958. The protein has a particular shape into The binding site of the macromolecule is flexible which the ligand fits exactly. and its shape can be modified as the ligand interacts with it. Ligand Receptor Molecular recognition: Collection of interactions between molecules that govern their binding. Qualitative nature of the interactions? Quantitative intensity of the molecular recognition? Unil 42





























































olecular recognition – Summary					
Category	Interaction	Distance	Residues involved	Remarks	
	lonic (charge-charge)	Long range	Arg, Lys, Asp, Glu His (if charged)	Called salt bridge at short distance	
Electrostatic	Hydrogen bond	Short range	Arg, Lys, Asp, Glu His, Tyr Ser, Thr, Asn, Gln Cys	Directionality / locality of interactions Specificity of molecular recognition	
	π interaction	Short range	Phe, Tyr, Trp, His		
Electrostatic/Non- polar	Van der Waals	Short range	Ala, Val, Ile, Leu, Pro, Cys, Met Phe, Tyr, Trp, His	Packing of atoms Shape complementarity	
Non-polar	Hydrophobic effect	-	All	Solute aggregation	
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Structure of pMHC Class II

- _
- Transmembrane glycoprotein 2 chains α (35 kDa) and β (28 kDa), forming 4 domains, α_1 , α_2 , β_1 and β_2 . -
- Deep groove to bind a peptide (12-18 residues) between α_1 and β_1 -
- _ Expressed at cell surface









3D structure of the complex between the T-cell receptor (TCR) and pMHC









TCR 3D structure Exercise 7 of the booklet Load the 2BNR PDB structure into UCSF chimera _ Produce a molecular representation similar to the one you see on the right CDR1: 28 - 32 CDR2: 51 - 55 CDR3: 94 - 101 Residues in the CDR for TCR α : CDR1: 25 - 29 CDR2: 49 - 53 Residues in the CDR for TCR β : • . CDR3: 94 - 100 . Display the atoms of the CDRs for TCR $\!\alpha$ and TCR $\!\beta$ What are the CDRs that make most of the contacts with the peptide epitope? And with the MHC? What are the TCR residues that make contact with peptide Trp5? What types of interactions are taking place? What MHC residues are close to TCR β Ala51? If you wanted to change residue 51 of TCR β , to increase the affinity of TCR . for pMHC, what mutation would you introduce? Why? 97

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