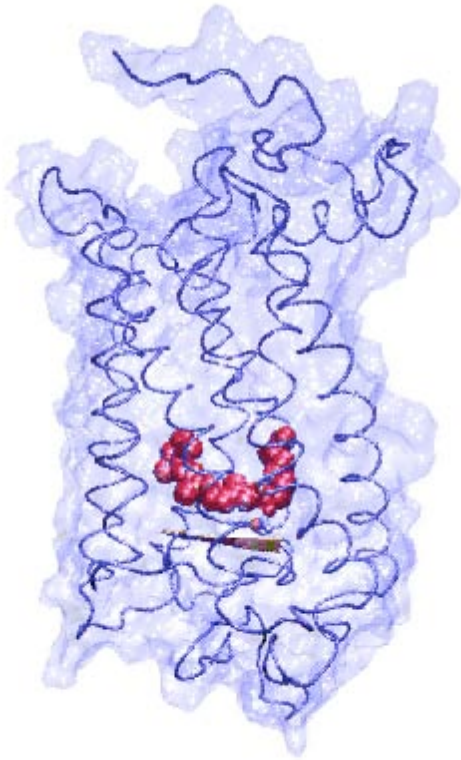
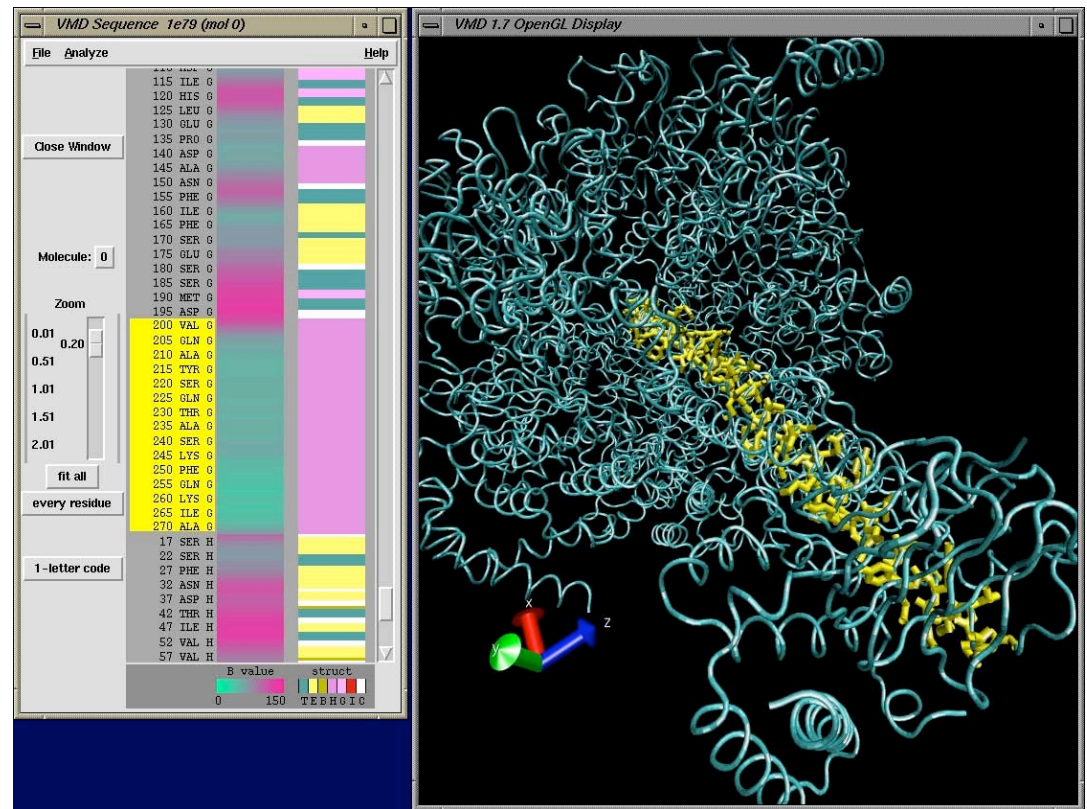


Molecular Graphics Perspective of Protein Structure and Function



animation

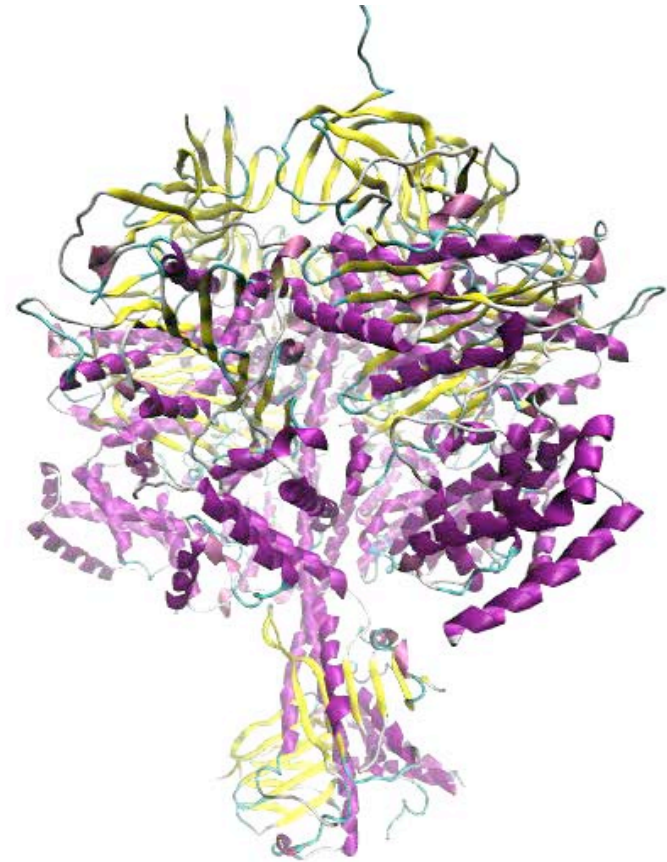


sequence

structure

VMD Highlights

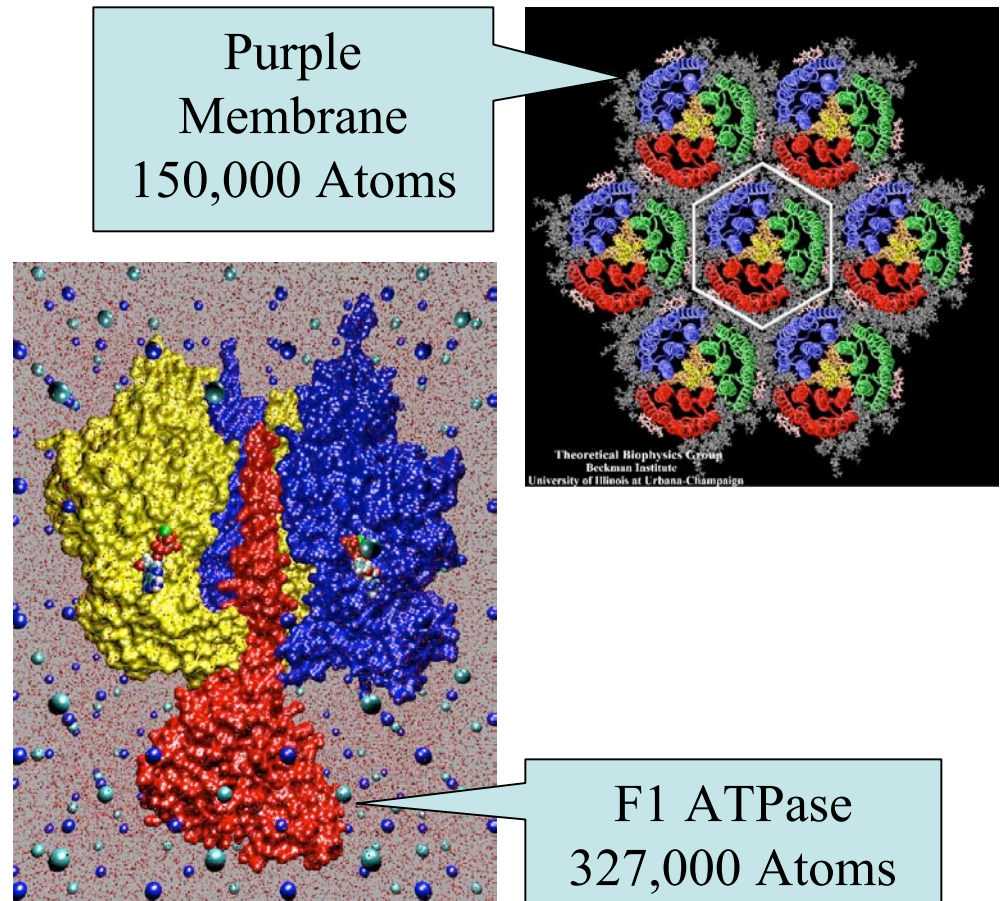
- > 40,000 registered users
- Platforms:
 - Unix (16 builds)
 - Windows
 - MacOS X
- Display of large biomolecules and simulation trajectories
- Sequence browsing and structure highlighting
- Multiple sequence - structure analysis
- User-extensible scripting interfaces for analysis and customization



The program is used today more for preparation and analysis of modeling than for graphics

VMD Permits Large Scale Visualization

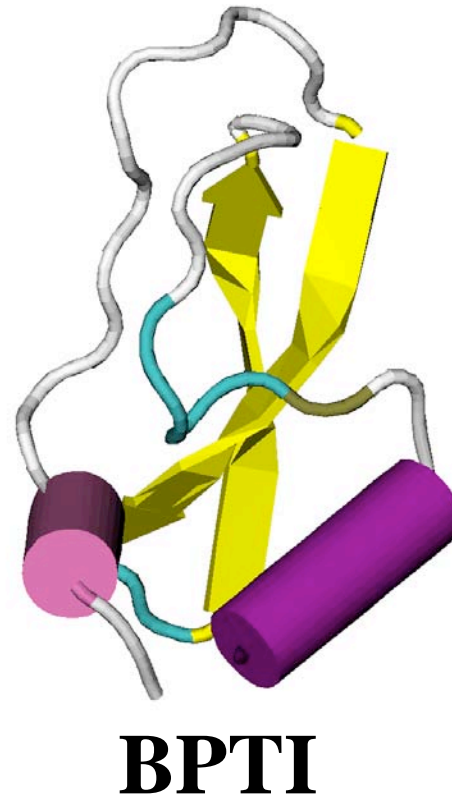
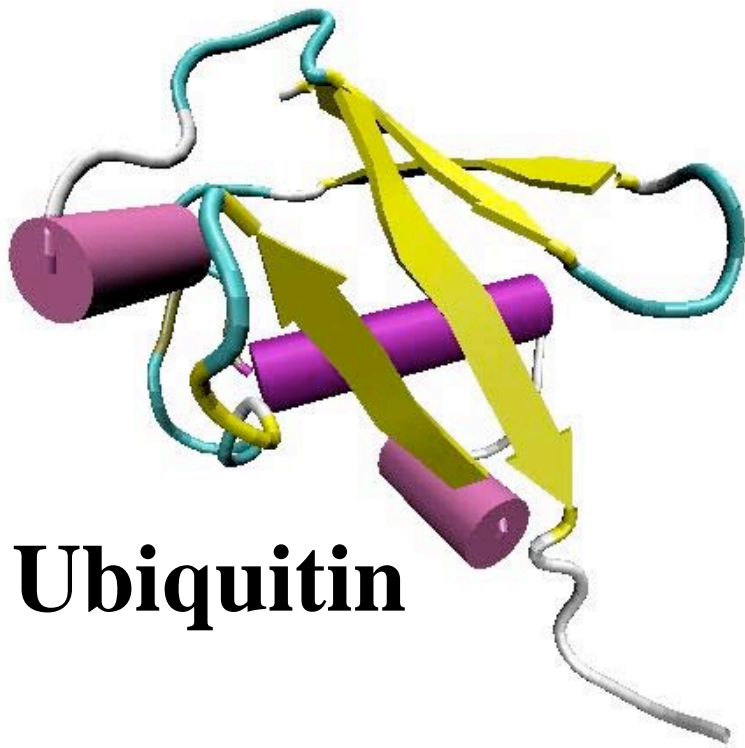
- Large structures: 300,000 atoms and up
- Complex representations
- Long trajectories: thousands of timesteps
- Volumetric data
- Multi-gigabyte data sets break 32-bit barriers
- Handles large data sets, e.g., GlpF: each 5 ns simulation of 100K atoms produces a 12GB trajectory



Focus on two proteins

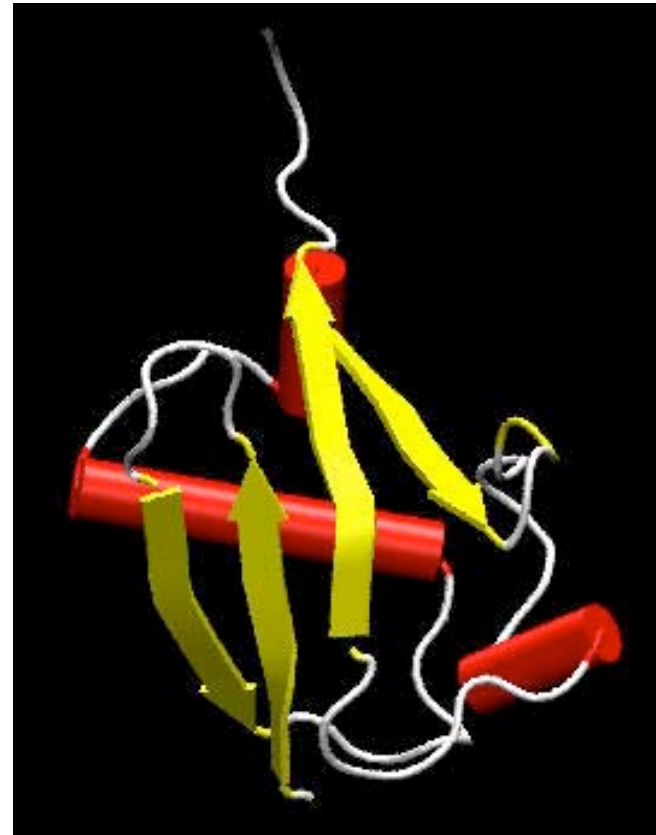
Ubiquitin

Bovine Pancreatic Trypsin Inhibitor (BPTI)



Ubiquitin

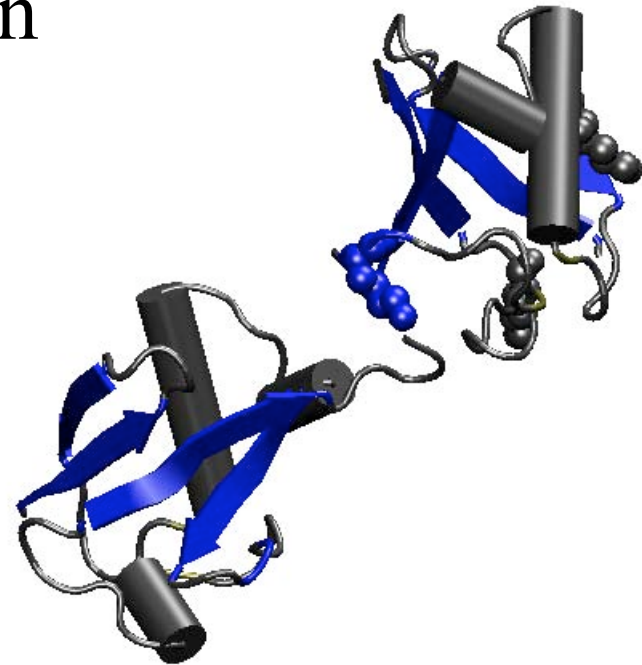
- 76 amino acids
- highly conserved
- covalently attaches to proteins and tags them for degradation
- other cell trafficking



- Glycine at C-terminal attaches to the Lysine on the protein by an isopeptide bond.

- it can attach to other ubiquitin molecules and make a polyubiquitin chain.

There are 7 conserved lysine residues in ubiquitin.



Two ubiquitins attached together through LYS 48. LYS 63 and LYS 29 are also shown there.

Ubiquitination Pathway



The Nobel Prize in Chemistry 2004

"for the discovery of ubiquitin-mediated protein degradation"

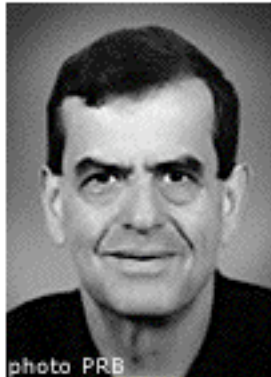


photo PRB

Aaron Ciechanover

🕒 1/3 of the prize
Israel

Technion – Israel
Institute of
Technology
Haifa, Israel

b. 1947

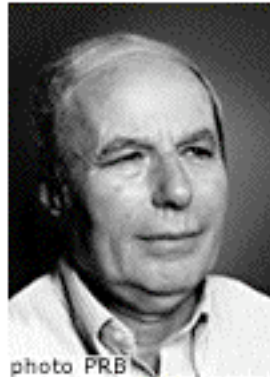


photo PRB

Avram Hershko

🕒 1/3 of the prize
Israel

Technion – Israel
Institute of
Technology
Haifa, Israel

b. 1937
(In Karcag, Hungary)

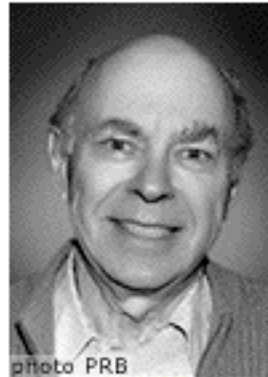


photo PRB

Irwin Rose

🕒 1/3 of the prize
USA

University of
California
Irvine, CA, USA

b. 1926

Ubiquitin-mediated protein degradation

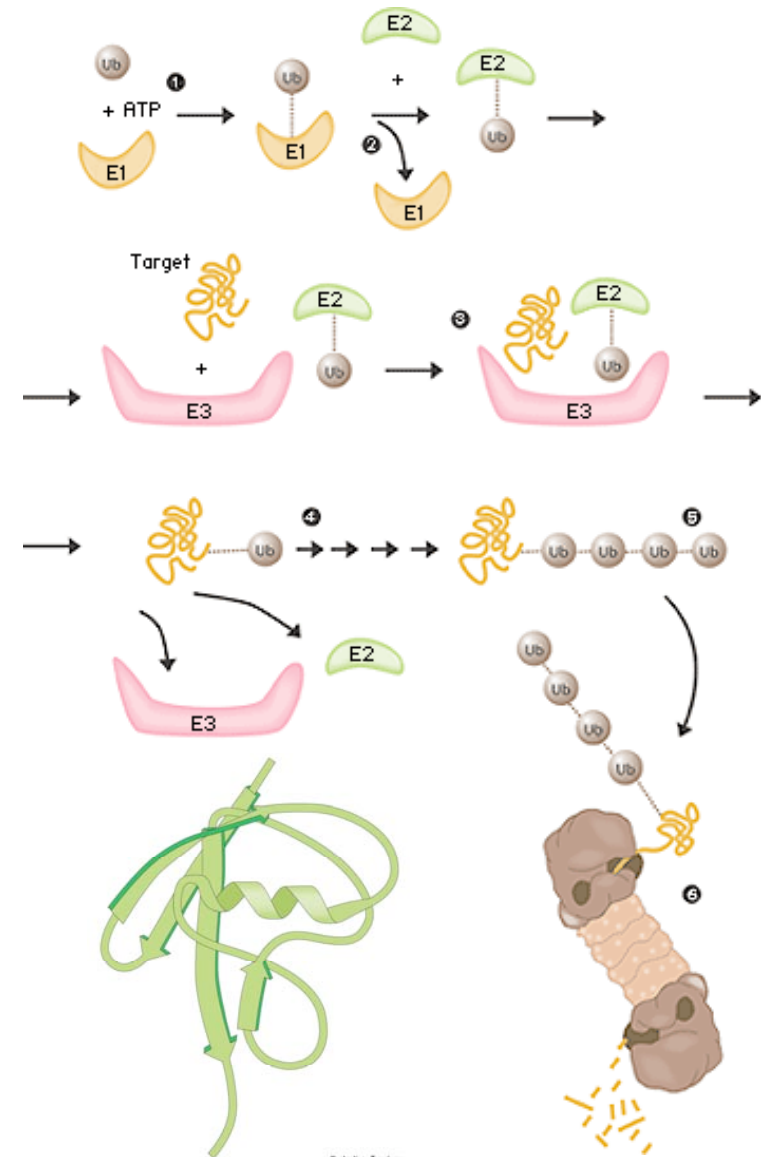
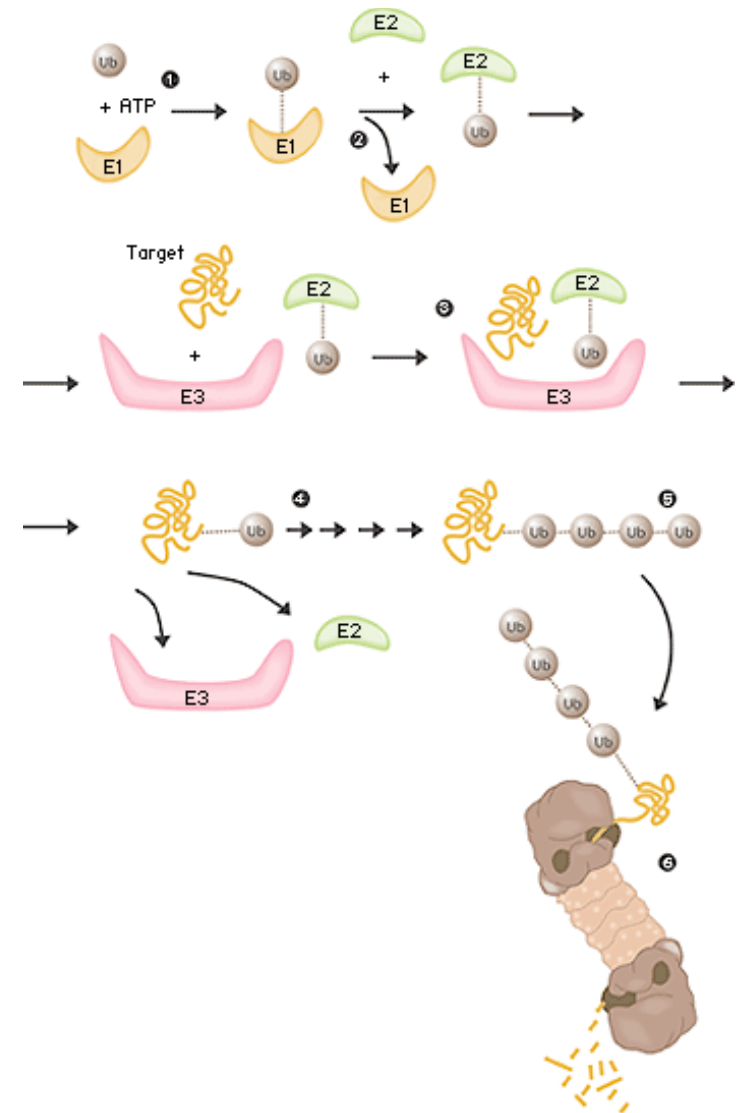


Illustration: Tgatom

Ubiquitination Pathway

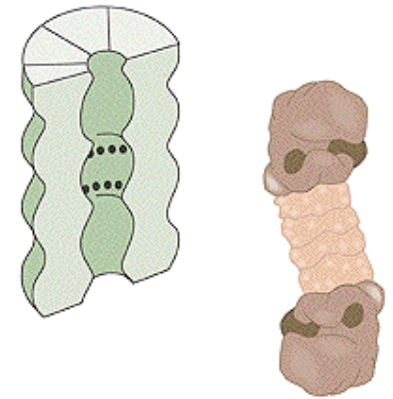
- Activation by E1 (ATP dependent process)
(thiol-ester linkage between a specific cysteine residue of E1 and Glycine on ubiquitin)
- Transfer to a cysteine residue on E2
(ubiquitin conjugation enzyme)
- Transfer of ubiquitin by E3 to the substrate lysine residue.
- E3 recognizes the ubiquitination signal of the protein.



Ubiquitin Functions

- tagging misfolded proteins to be degraded in the proteasome (kiss of death).
- regulates key cellular processes such as cell division, gene expression, ...

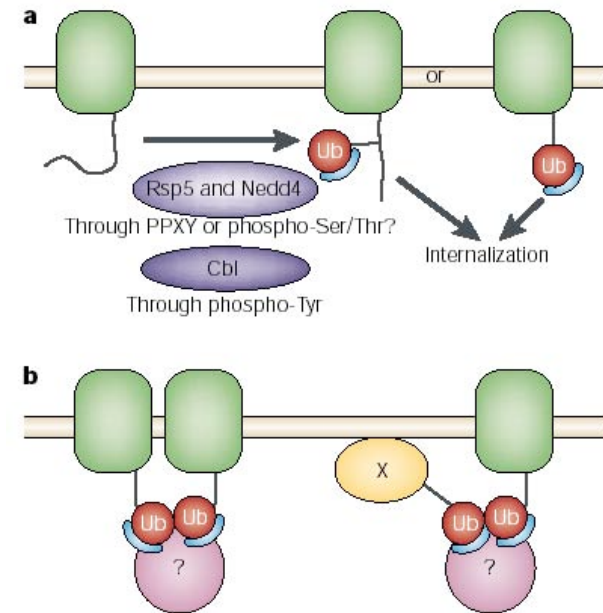
A chain of at least four ubiquitins is needed to be recognized by the proteasome.



The cell's waste disposer, the proteasome. The black spots indicate active, protein-degrading surfaces.

Ubiquitin acts independent of proteasome degradation

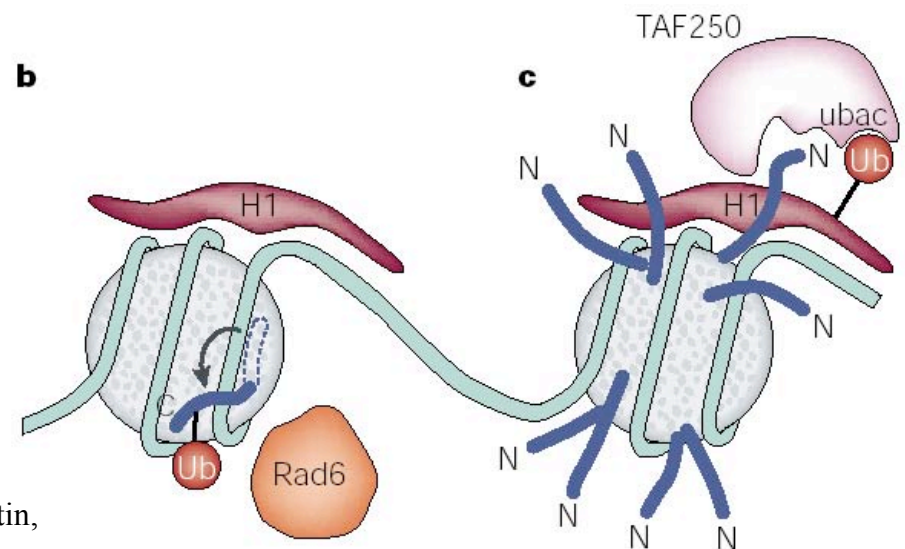
- Controlling the traffic in the cell
- Directing the traffic in the cell, i.e., determining where the newly synthesized proteins should go
- Tagging membrane proteins for internalization



Ubiquitine Regulates gene expression:

(indirectly, by destruction of some of the involved proteins)

- Recruiting Transcription Factors (proteins needed for gene expression)
- Conformational changes in Histone, necessary before gene expression



Hicke, L., Protein regulation by monoubiquitin,
Nat. Rev. Mol Cell Biol., 2, 195-201 (2001)

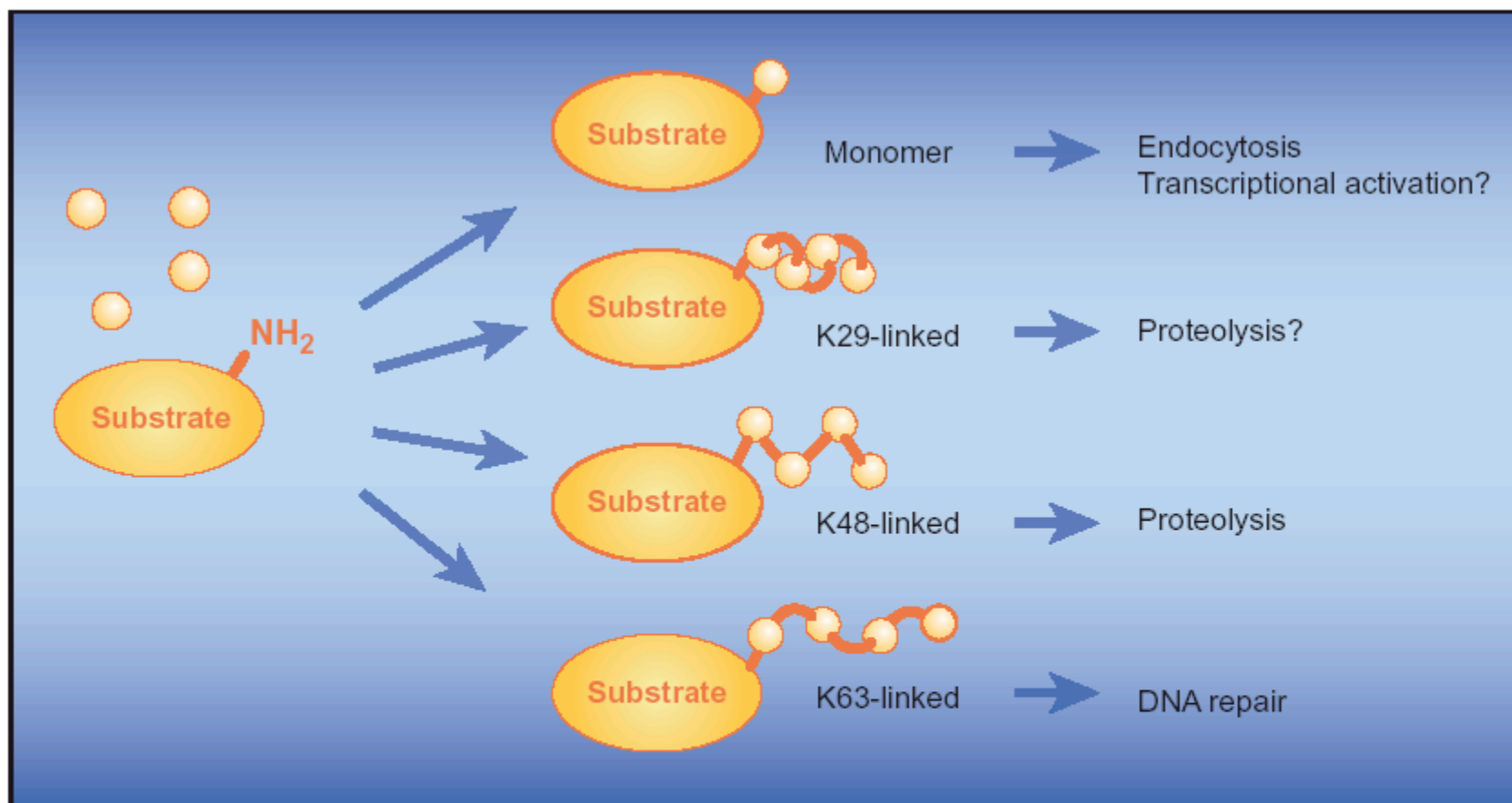
Different types of ubiquitin signals arise from

- Length of the ubiquitin chain
- How ubiquitins are attached together
- Where the signals are read

Examples:

- multi-ubiquitin chains, linked through Lysine 48, target protein for proteasome degradation
- K63 linkages direct DNA repair

Mono-ubiquitylation versus multi-ubiquitylation



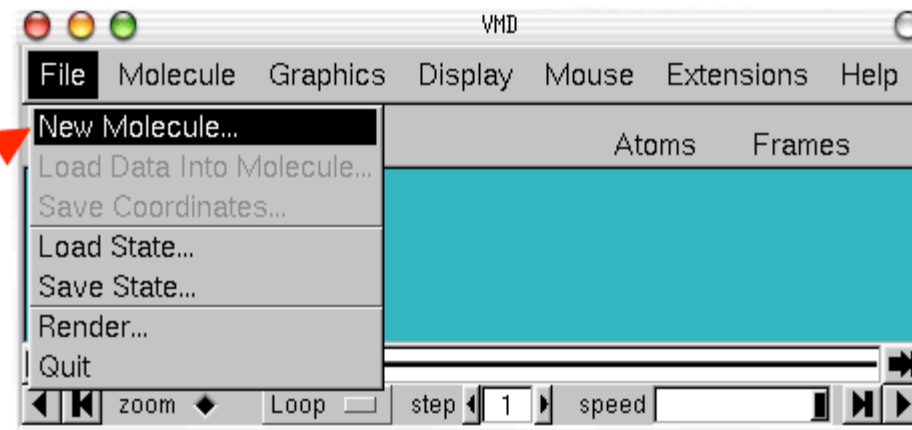
Multifaceted. Ubiquitin can attach to its various substrate proteins, either singly or in chains, and that in turn might determine what effect the ubiquitination has. (K29, K48, and K63 refer to the particular lysine amino acid used to link the ubiquitins to each other.)

Marx, J., Ubiquitin lives up its name, *Science* 297, 1792-1794 (2002)

Basics of VMD

Loading a Molecule

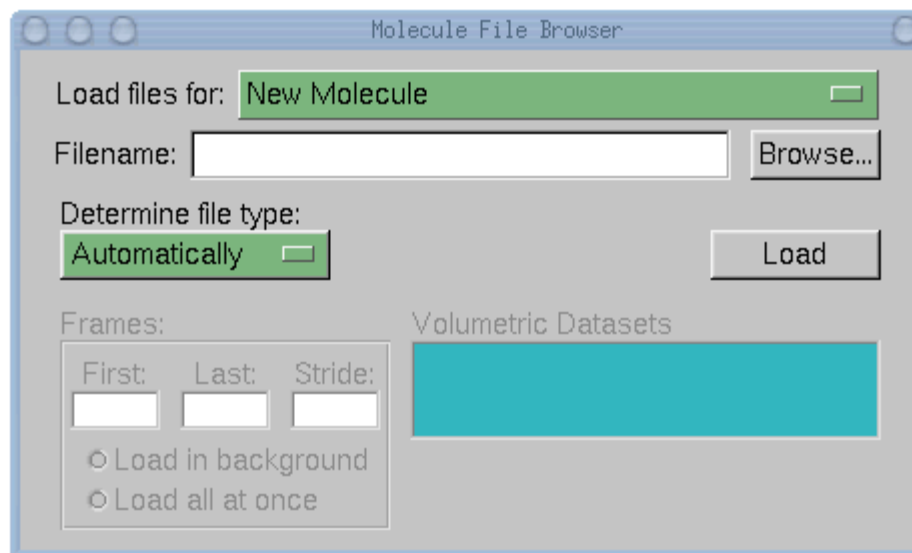
New Molecule (a)



(b) Molecule file browser

(c) Browse

(d) Load



Basics of VMD

Rendering a Molecule

Current graphical representation (a)

Draw style (b)

Coloring (c)

Drawing method (d)

(e) Selected Atoms

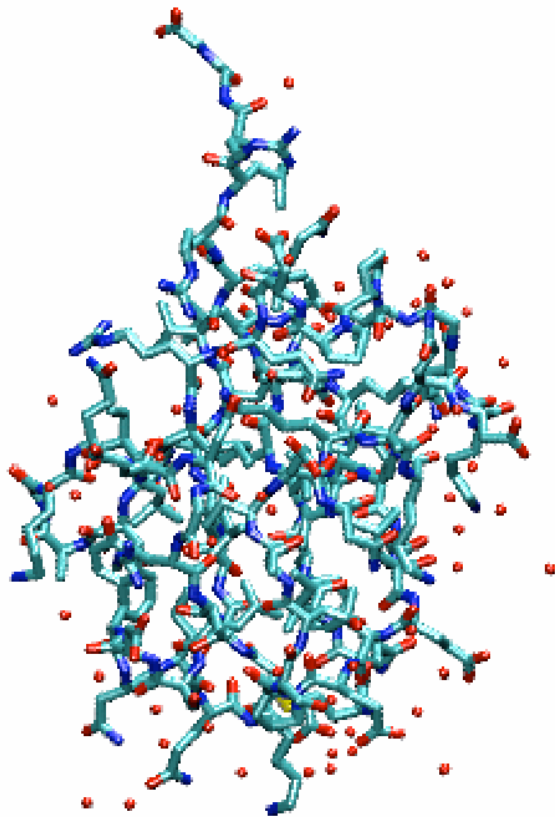
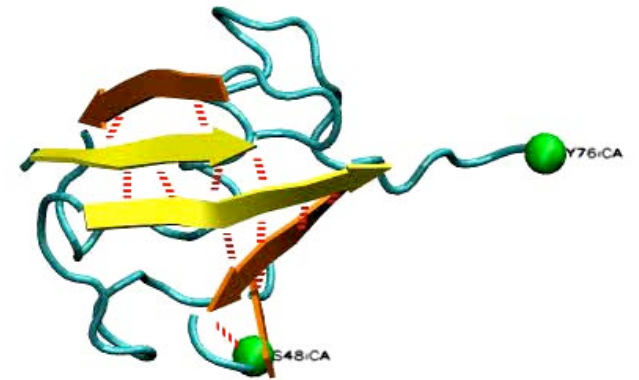
(e) Resolution, Thickness

The screenshot shows the 'Graphical Representations' window in VMD. At the top, the 'Selected Molecule' is '0: 1UBQ'. Below this are 'Create Rep' and 'Delete Rep' buttons. A table lists graphical representations with columns for 'Style', 'Color', and 'Selection'. The first row is highlighted in yellow and contains 'Lines', 'Name', and 'all'. Below the table is a 'Selected Atoms' text box containing 'all'. The 'Draw style' section has tabs for 'Draw style', 'Selections', 'Trajectory', and 'Periodic', with 'Draw style' selected. Under 'Draw style', there are 'Coloring Method' (set to 'Name') and 'Drawing Method' (set to 'Lines'). The 'Material' section has a dropdown set to 'Opaque'. At the bottom, there is a 'Thickness' spinner set to '1' and an 'Apply Changes Automatically' checkbox with an 'Apply' button.

Style	Color	Selection
Lines	Name	all

Basics of VMD

Change rendering style



CPK



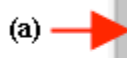
tube



cartoon

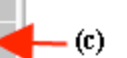
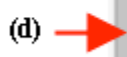
Basics of VMD

Create Representation (a)



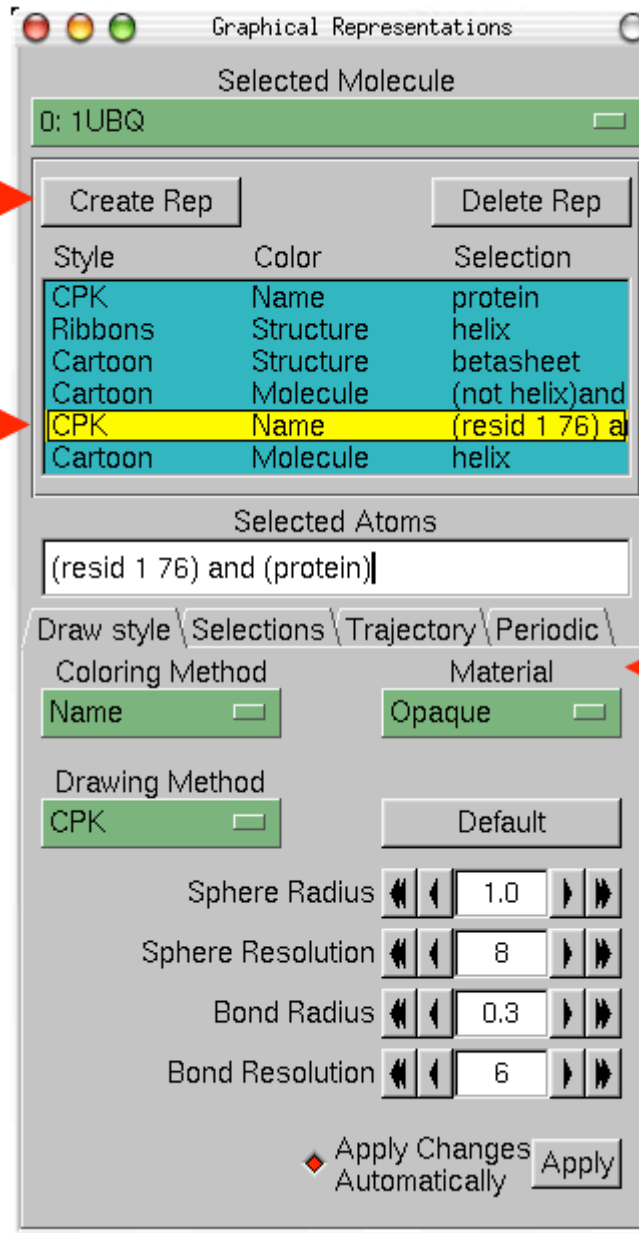
Delete Representation (b)

Current Representation (d)

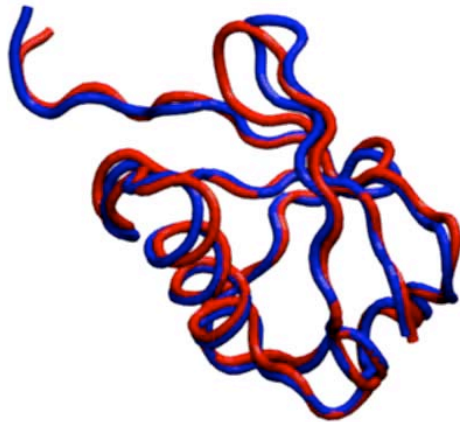


Material (c)

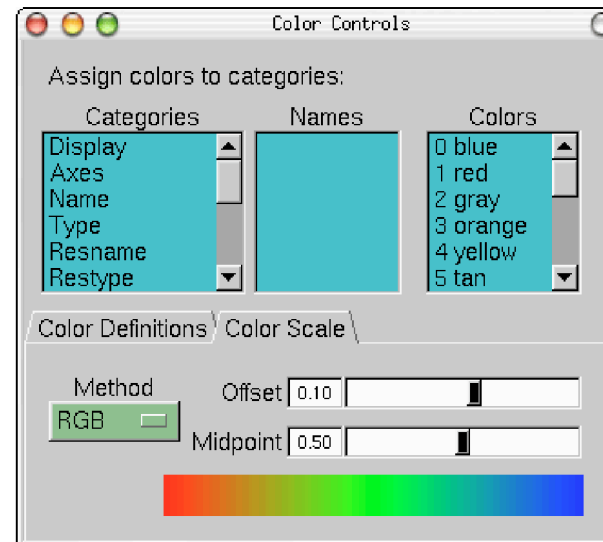
Multiple representations



VMD Scripting



Left: Initial and final states of ubiquitin after spatial alignment
Right (top): Color coding of deviation between initial and final



The Color Controls window showing the Color Scale tab.

VMD Sequence Window

(a)

File Help

23 ILE X
24 GLU X
25 ASN X
26 VAL X
27 LYS X
28 ALA X
29 LYS X
30 ILE X
31 GLN X
32 ASP X
33 LYS X
34 GLU X
35 GLY X
36 ILE X
37 PRO X
38 PRO X
39 ASP X
40 GLN X
41 GLN X
42 ARG X
43 LEU X
44 ILE X
45 PHE X
46 ALA X
47 GLY X
48 LYS X
49 GLN X
50 LEU X
51 GLU X
52 ASP X
53 GLY X
54 ARG X
55 THR X
56 LEU X
57 SER X
58 ASP X
59 TYR X
60 ASN X
61 ILE X
62 GLN X
63 LYS X
64 GLU X

Close Window

Molecule: 0

Zoom

0.01
0.51
1.01 1.00
1.51
2.01

fit all

every residue

1-letter code

B value

0 150

struct

T E B H G I C

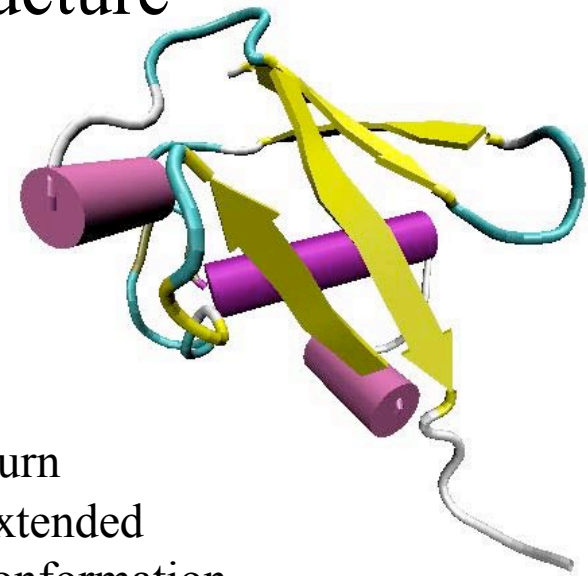
(d)

List of (e) the residues

Zoom (f)

(b) Beta Value

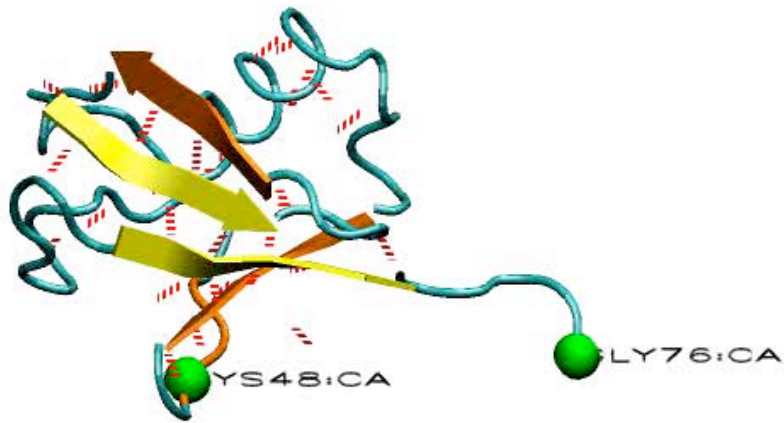
(c) Structure



T: Turn
E: Extended
conformation
H: Helix
B: Isolated Bridge
G: 3-10 helix
I: Phi helix

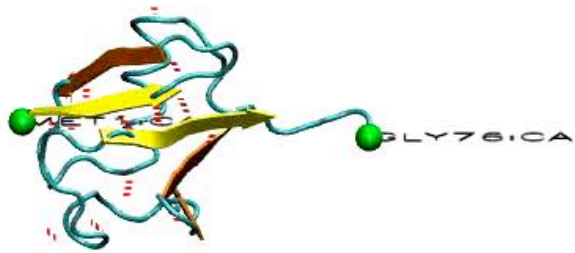
VMD Macros to Color Beta Strands

Use VMD scripting features to color beta strands separately; show hydrogen bonds to monitor the mechanical stability of ubiquitin



Ubiquitin stretched between the C terminus and K48 does not fully extend!

Discovering the Mechanical Properties of Ubiquitin

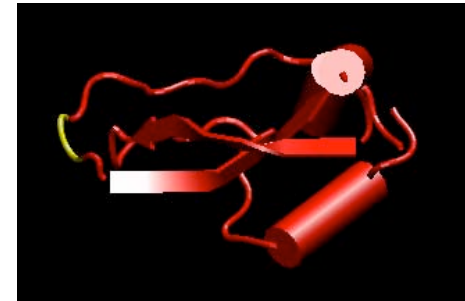


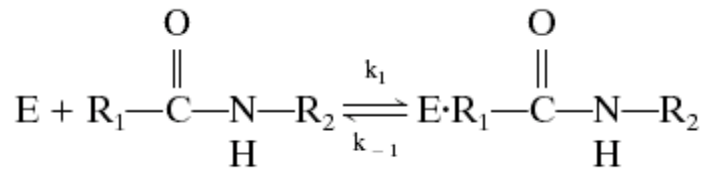
Ubiquitin stretched between the C and the N termini extends fully!

Discover BPTI on your own!

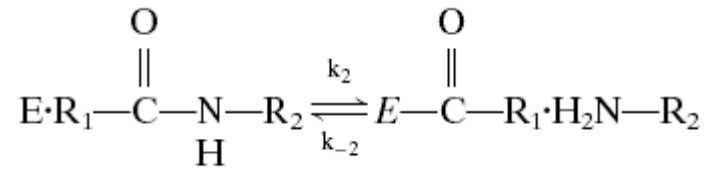
bovine pancreatic trypsin inhibitor

- small (58 amino acids)
- rigid
- binds as an **inhibitor** to Trypsin
(a serine proteolytic enzyme, that appears in digestive system of mammals.)
- blocks its active site.

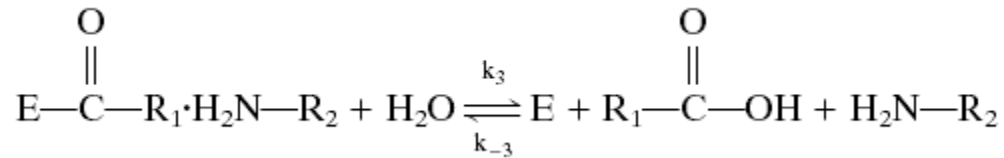




(Michaelis complex)



(Acyl-enzyme with leaving peptide
noncovalently bound)



(Free enzyme and products)

Mechanism of cleavage of peptides with serine proteases.

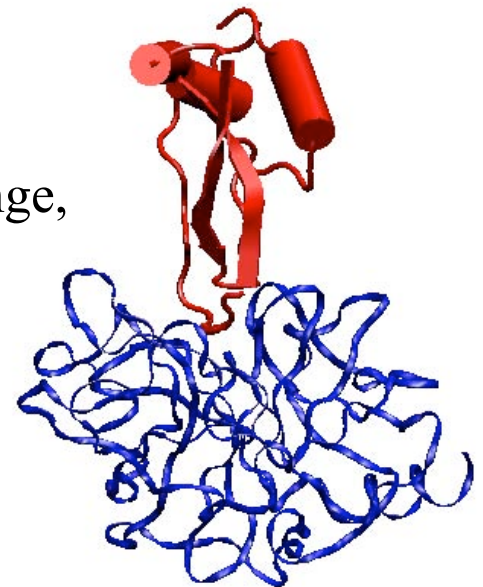
Radisky E. and Koshland D. Jr., Proc. Natl. Acad. Sci., USA, 99, 10316-10321

Trypsin: A proteolytic enzyme that hydrolyzes peptide bonds on the carboxyl side of **Arg** or **Lys**.

BPTI: A “standard mechanism” inhibitor

- Binds to Trypsin as a substrate.
forms an acyl-enzyme intermediate rapidly.
- Very little **structural changes** in trypsin or BPTI.
several H-bonds between backbone of the two proteins change,
little reduction in conformational entropy → binds tightly
- Remains uncleaved.
hydrolysis is 10^{11} times slower than for other substrates

Structures of the **protease binding region**, in the proteins of all 18 families of standard mechanism inhibitors are similar.



Why does Trypsin cleave BPTI so slowly?

- Disruption of the non-covalent bonds in the **tightly bonded** enzyme-inhibitor complex increases the energy of transition states for bond cleavage.
- Water molecules do not have access to the active site, because of the **tight binding** of Trypsin and BPTI.
- After the cleavage of the active-site peptide bond, the newly formed termini **are held in close proximity**, favoring reformation of the peptide bond.
- The **rigidity** of BPTI may also contribute by not allowing necessary atomic motions.

Dance of Ubiquitin