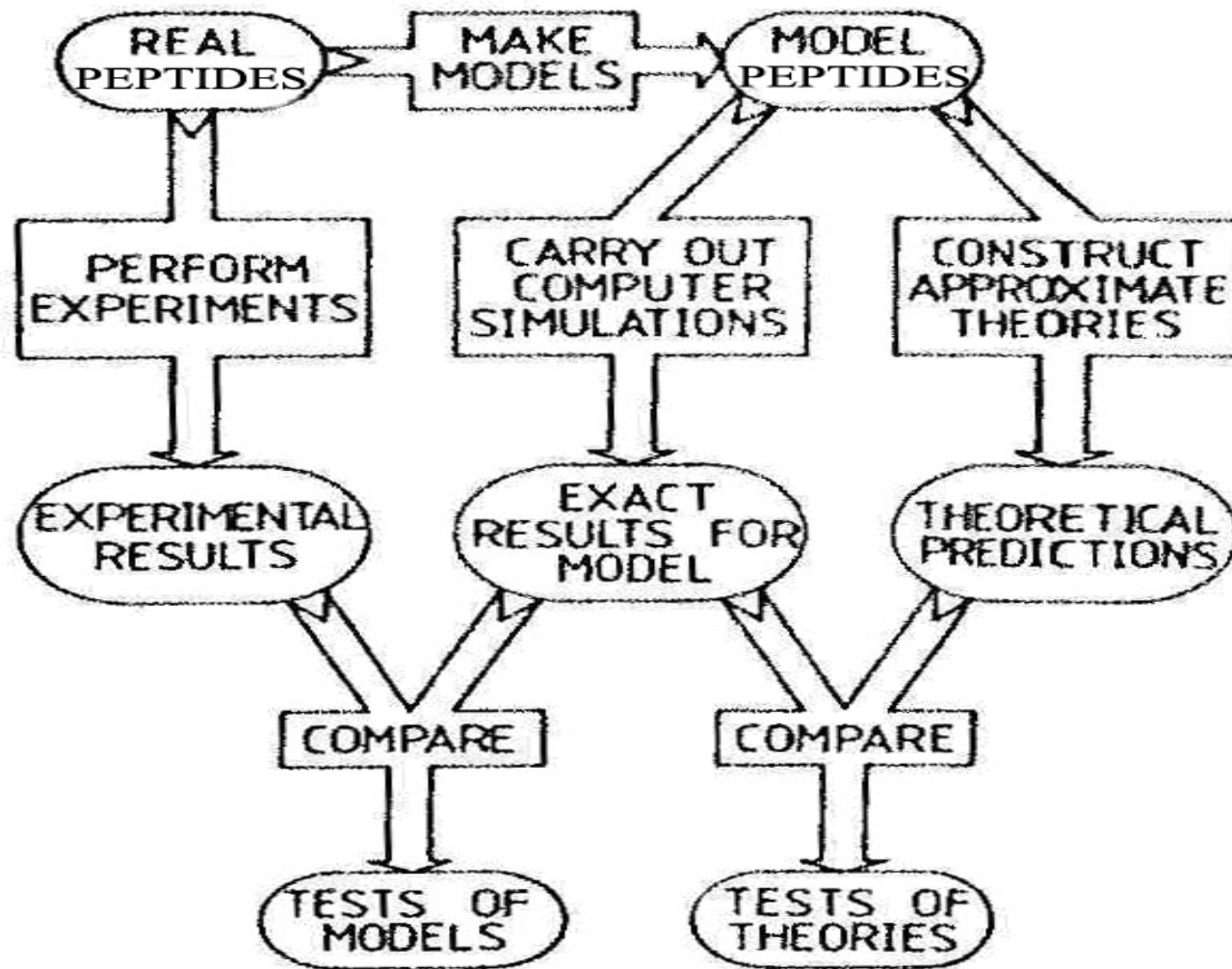
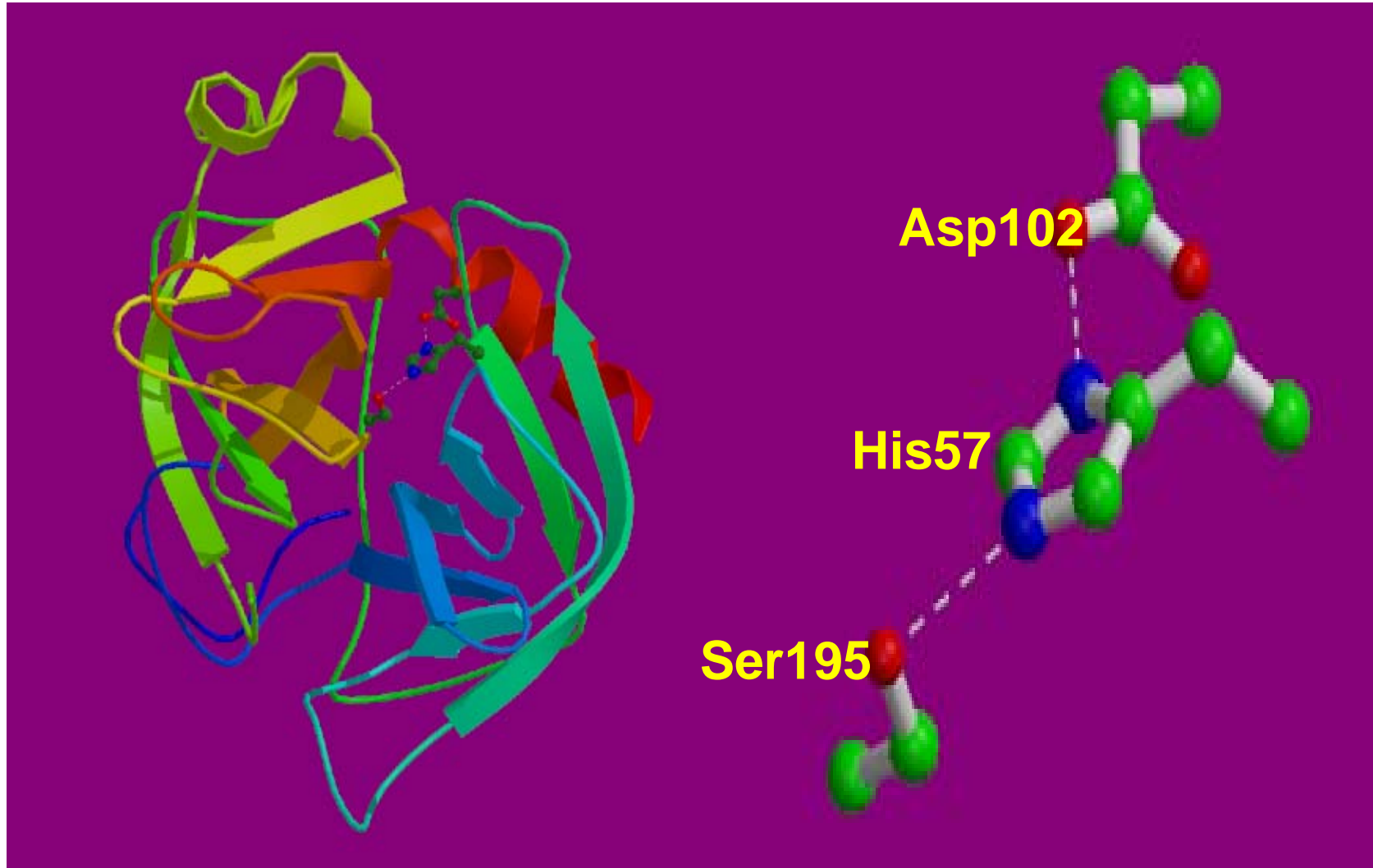
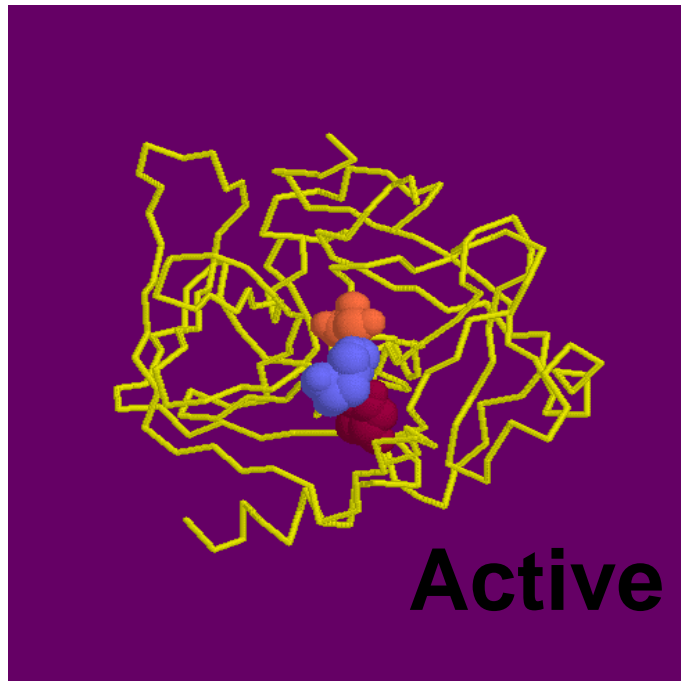


# Theozymes - Compuzymes

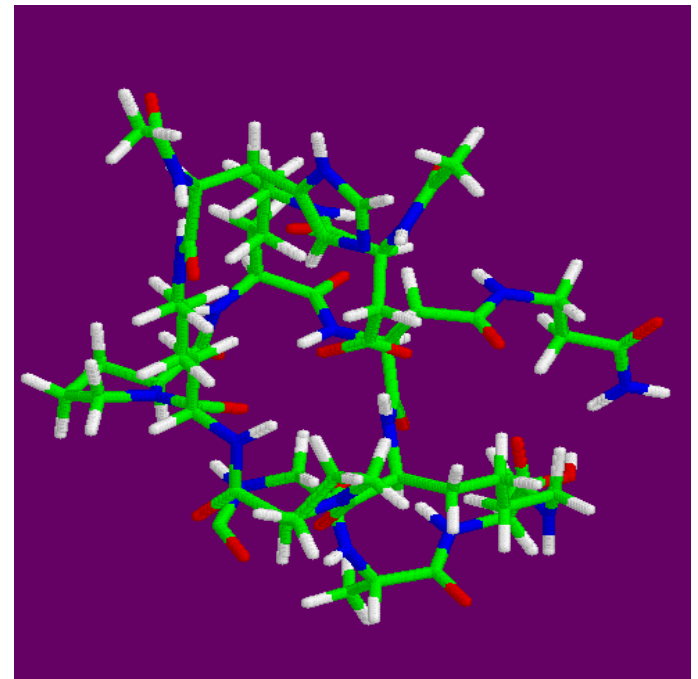


# chymotrypsin



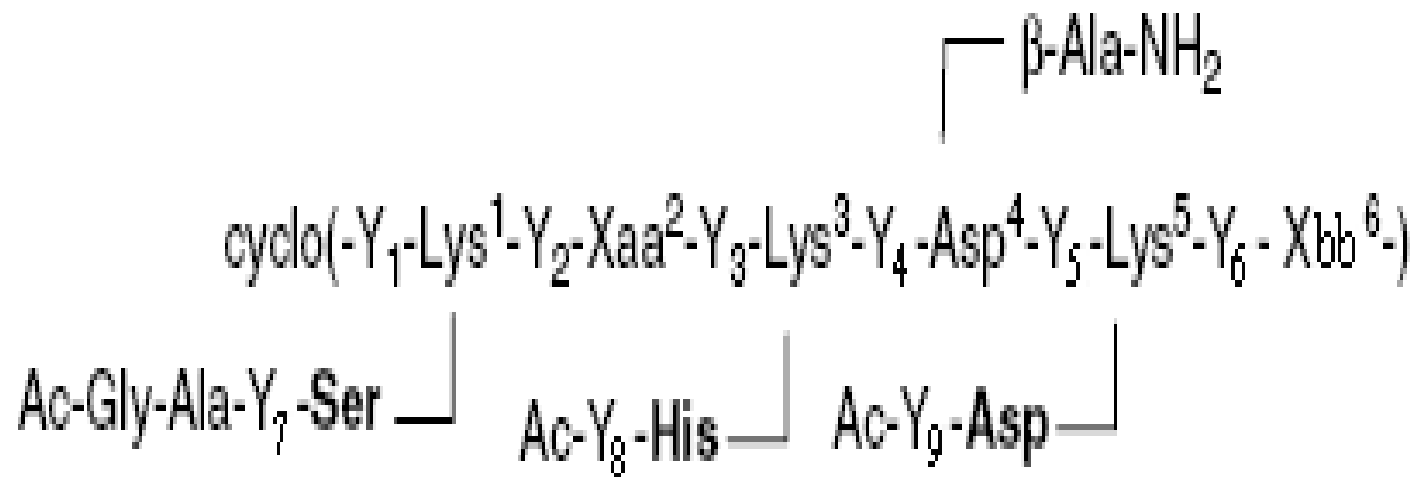


**Chymotrypsin  
245 aminoacids**

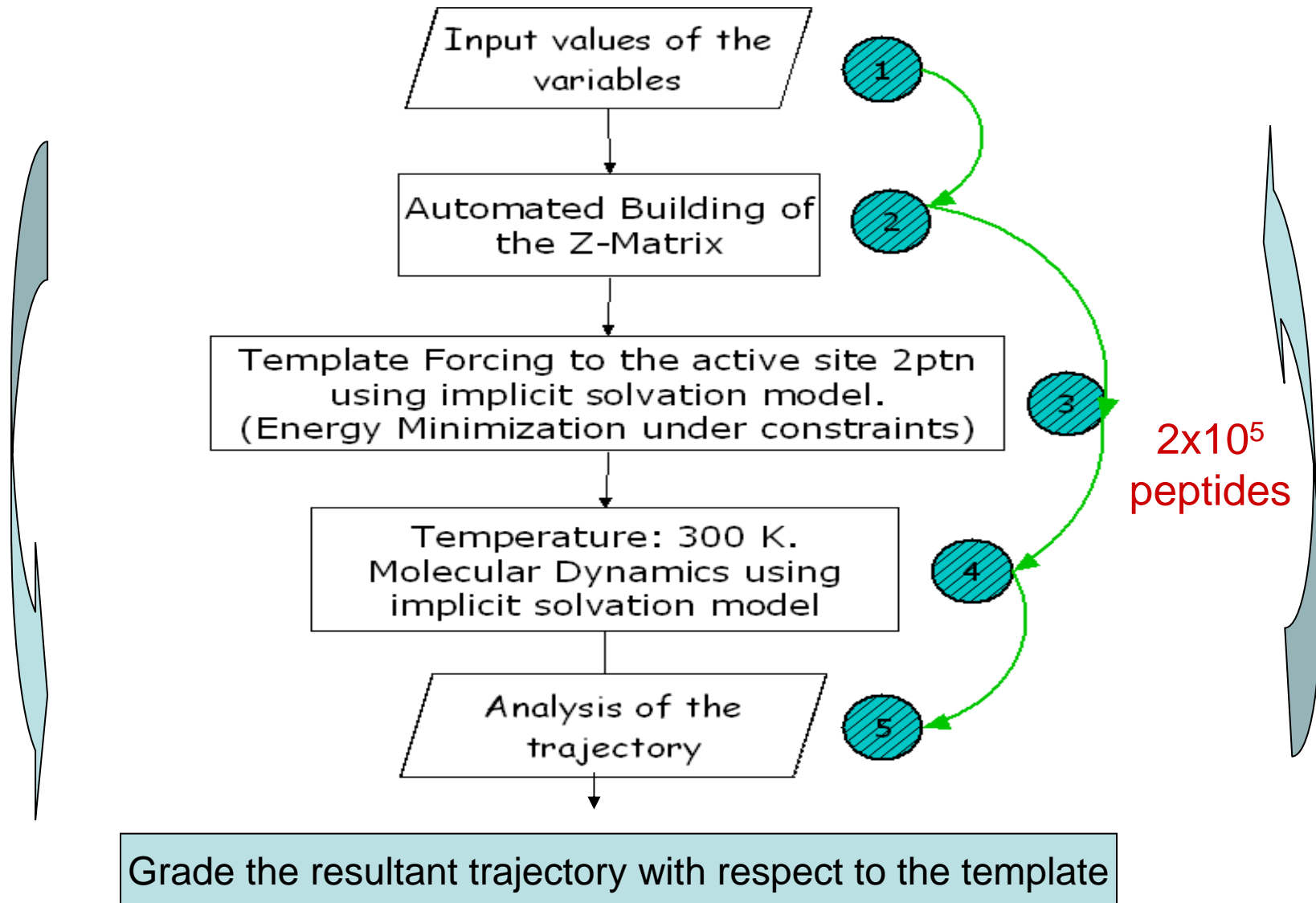


**Peptide-template  
12 aminoacids**

# Molecular template for chymotrypsin



# Flowchart



# The application (TAS) (1/5)

- **TrypsinActSite(TAS)** is a new tool that provides an integrated framework to build and predict the best mimetic for the serine protease, chymotrypsin.

# The application (TAS) (2/5)

- An extension of the well-known software **TINKER** (Ponder, J.W) (<http://dasher.wustl.edu/tinker/>) that:
- (a) constructs branched cyclic peptides of a certain pattern

# The application (TAS) (3/5)

- (b) guides the peptides to adopt a conformation similar to the active site



# The application (TAS) (4/5)

- (c) subjects the molecules to unconstrained molecular dynamics (implicit solvation environment)

# The application (TAS) (5/5)

- (d) grades the resultant trajectories with respect to the actual trypsin spatial arrangement of the active site.

# Results

## INPUT

```

Comments
/opt      # Installation path of TINKER
1        # Version of TINKER (4.1)
Gly      # Xaa peptide (Fig. 1)
Gly      # Xbb peptide (Fig. 1)
1        # Choice 1 (specific D/L chirality)
LLDLLLDLDD # Peptide 1 (Y1- ... -Y9)
LDLLDLDD  # Peptide 2 (Y1- ... -Y9)
DLLDLLLLL # Peptide 3 (Y1- ... -Y9)
DLDDLDDL  # Peptide 4 (Y1- ... -Y9)
          # Leave this line blank to continue
220000   # The total time of MD (fs)
1        # Time step length (fs)
2        # Time between dumps (ps)
300      # Temperature (K)
SASA     # Implicit Solvation model
amber99  # Force Field
```

## OUTPUT

Peptide	RMSd(%)	d1(%)	d2(%)	d3(%)
1	97	4	0	27
2	12	4	0	2
3	1	0	0	0
4	11	10	1	15
<b>Amber99/GBSA</b>				
1	9	10	4	7
<b>Charmm27/GBSA</b>				
1	4	0	0	0
<b>Charmm27/SASA</b>				
1	100	99	0	15
<b>Amber99/ONION</b>				
1	100	100	0	100

# TAS + GRIDS

- TAS is especially suited for distributed computing.
- The package can screen over  $2 \cdot 10^5$  substances, exhibiting the pattern (template) for possible mimetics of chymotrypsin.
- This computer process can be easily distributed.

# Future Enhancements of TAS

Next editions will take into account

- a. the simulation of other active sites
- b. the choice of the number of the amino acids being in the circle
- c. hydrogen bond directionality constraints
- d. the binding site topology