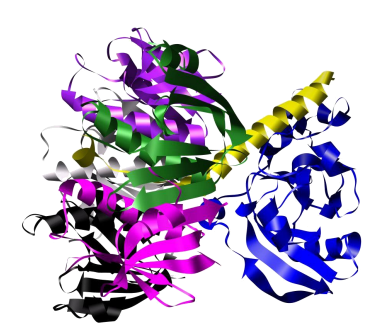
Protein Domains

Two weeks for this topic

many proteins have separate chains



enterotoxin (1lts) 7 chains

a haemoglobin (1h97)

what about units within one chain?

4 domain protein

1cxl has 686 residues Cleaves carbohydrate bond

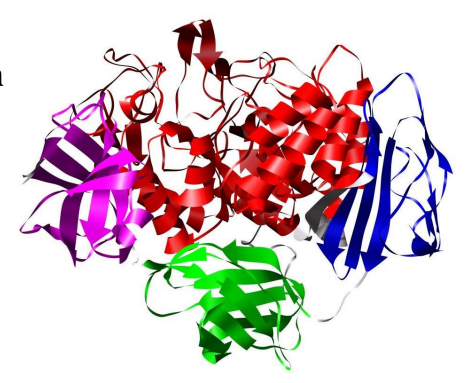
• one solid lump but...



4 domain protein

- 1. α -amylase catalytic
- 2. α-amylase C-terminal
- 3. immunoglobulin like domain
- 4. starch binding

• even clearer example

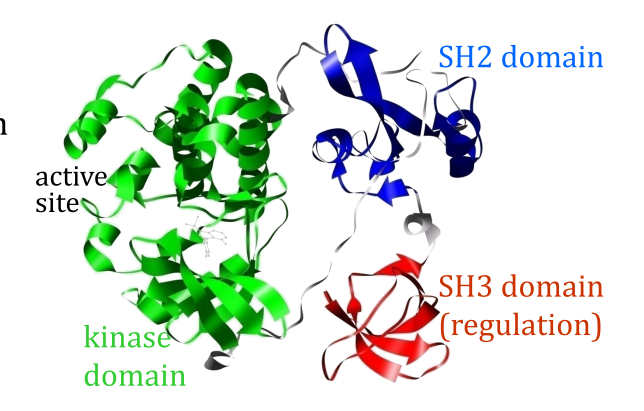


3 domain protein

1qcf "src tyrosine kinase"

The domains really are common to other proteins

Number of domains is not absolutely defined



Plan

- chemistry, examples
- methods to automatically recognise domains (examples)
- chemistry how common are domains of different sizes, types, ...

Earlier history

Term "domain" used before there were many structures

- Invented example: protein that
 - joins ADP + $P_i \rightarrow ATP$
 - performs some oxidation
 - responds to some regulator
 - take protein + protease (splits protein in a few places)
 - cleave / break protein get a few pieces (2, 3, 4..)
 - purify pieces
 - pieces found that
 - can bind ADP/ATP
 - bind sugars, some regulators

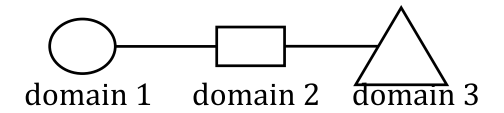
Earlier history

Appeared that for some proteins

- different functions associated with different pieces
- refer to as "functional domains"

Belief / claim

• bigger proteins are made from units, combined over evolutionary time scales



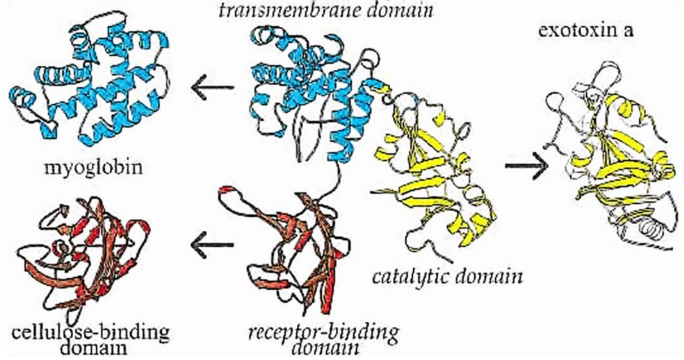
an example...

modular protein

Diptheria toxin (1ddt) middle of picture

• 3 domains

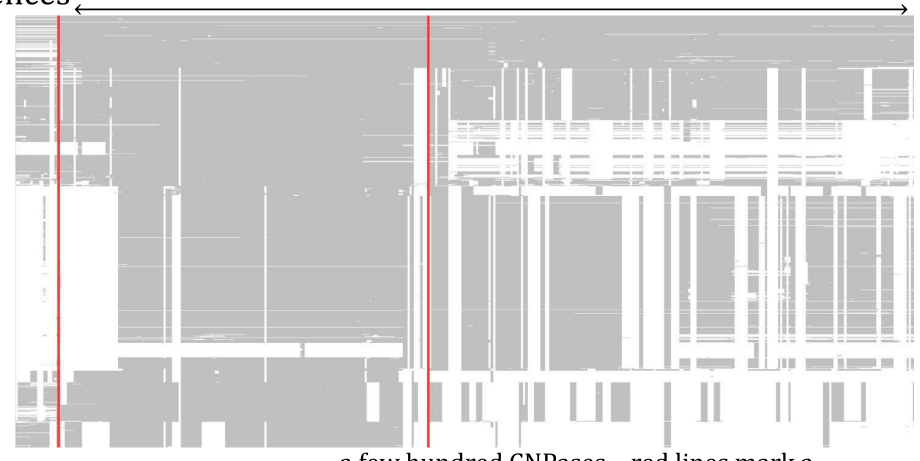
• each similar to some different protein transmembrane domain



- appears as if modules are mixed together
- should be visible at sequence level...

Sequence level domains

Align a group of sequences 450 residues



a few hundred CNPases – red lines mark a domain

schematically..

Sequence level domains

Align a group of sequences



- appears to have 3 or 4 domains
- no reference to structures or function

Domain definitions summary

	structure	sequence	biochemistry
functional	not necessary	not necessary	yes
sequence-based	not necessary	yes	no
structure	yes	usually known	no

How important?

- > $\frac{2}{3}$ proteins have 2 or more domains
- part of definition
 - a piece of a protein which can fold and is stable

Now

methods based on structure

Finding Domains

A definition leads to methods

• domain is a compact unit

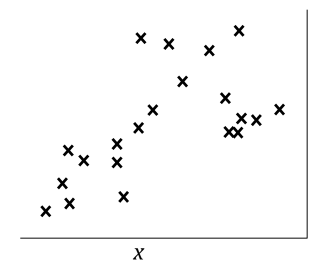
Objective way to look for dense units?

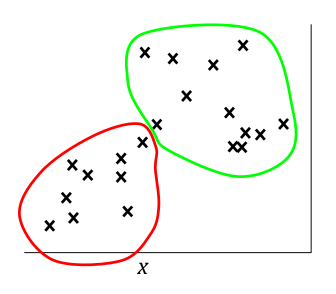
cluster analysis

Philosophy in cluster analysis

• look for dense groupings

Leads to dendrogram

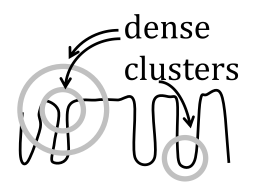




Clustering

Approach

- need a (dis)similarity matrix between every object
- here: distance between C^{α} atoms



residue	1	2	• • •	N
1	0		•••	•••
2		0	•••	
•••			•	
N				0

• does this work?

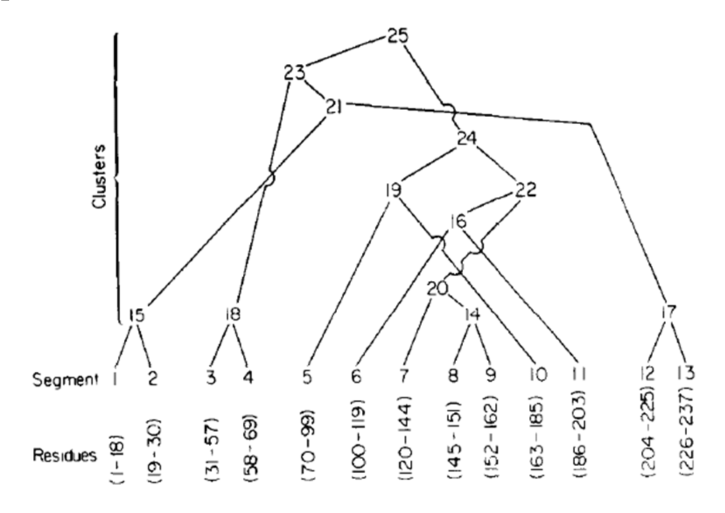
Clustering

Clustering applied to concanavalin A

- bottom small compact pieces
- higher compact units
- looks like natural3 domains

Number of domains is not absolutely determined

Very very very old method



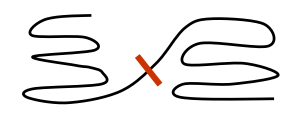
Cuts / Surface area / volume

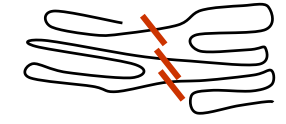
- Simple idea cut chain in two pieces
- density of part 1 / versus part 2



cut so as to maximise density

Problems - one cut is not enough





A method should be able to split with 1, 2, 3, ... cuts

• For 3 cuts with N_{res} positions: $N_{res} \times N_{res} \times N_{res}$

really
$$(N_{res})^{N_{cut}}$$

Problems - density

I want to maximise density

- density of protein ?
 - number of residues in a volume?
 - volume? not sphere

Contacts are easier than density

- within a domain there are many contacts
- between domains few contacts
- an approximation

Counting contacts

Do I have many contacts compared to the number of atoms?

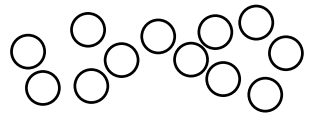
- calculate distance between each $C_i^{\alpha}C_j^{\alpha}$ atoms = d_{ij}
- if d_{ij} < 4 Å, set p_{ij} =1 else p_{ij} =0
- for a given set of N_{res} atoms (not whole protein)

$$\frac{\sum_{i=1}^{N_{res}} \left(\sum_{j>i}^{N_{res}} p_{ij}\right)}{N_{res}}$$

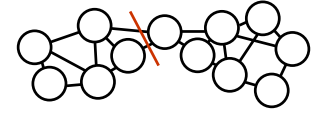
not accurate, but easy to calculate

Cutting / contacts

A protein



Find close contacts

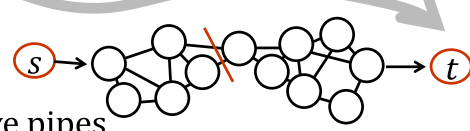


How can one find the best place(s) to cut?

Feed water into *s* (to *t*)

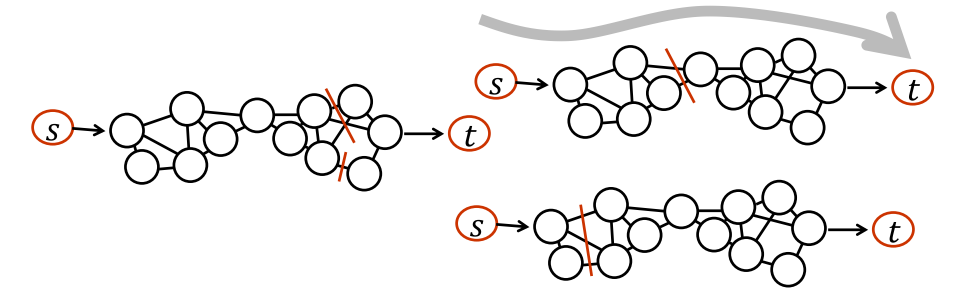


not one, but all that are restrictive



Cutting / contacts

- Flow problem
- Many ways to cut the flow from s to t

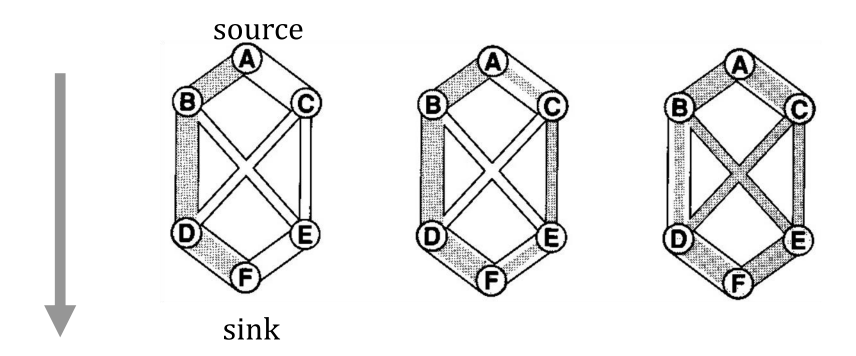


- of all these "st cuts" find the one with smallest capacity (flow)
- more interesting make the pipes different flow capacity
 - how are the residues really touching ? $C^{\alpha} C^{\alpha}$ or C^{α} sidechain

Cutting / contacts

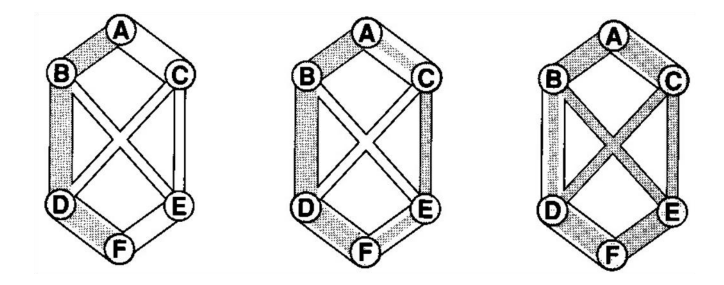
Two steps

- 1. find the maximum flow from *s* to *t*
- 2. cut *s* from *t* at the few most filled pipes



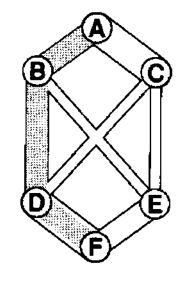
rule

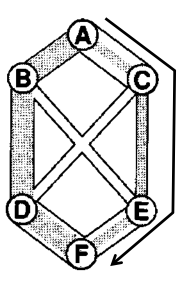
- if every path from source to sink has one full edge
 - flow is maximum

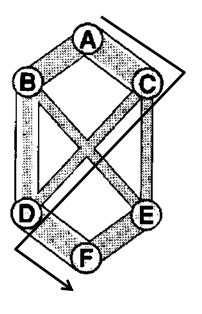


- keep trying every possible path, look to see if there is unused capacity
- we can go backwards

- A B D F note DF is not quite full
- add some A C E F AC, EF are not full
- look at C, switch some capacity to CD (DF)



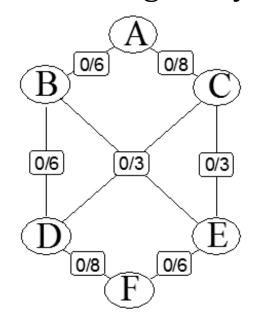


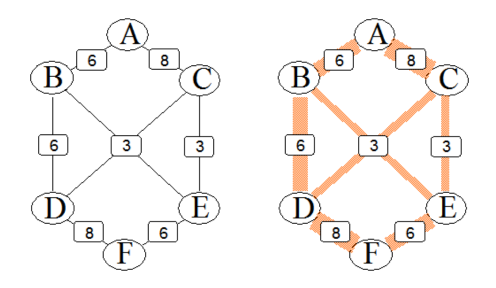


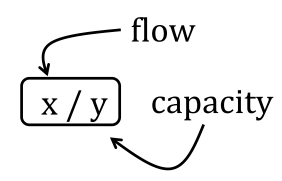
with some numbers on the edges

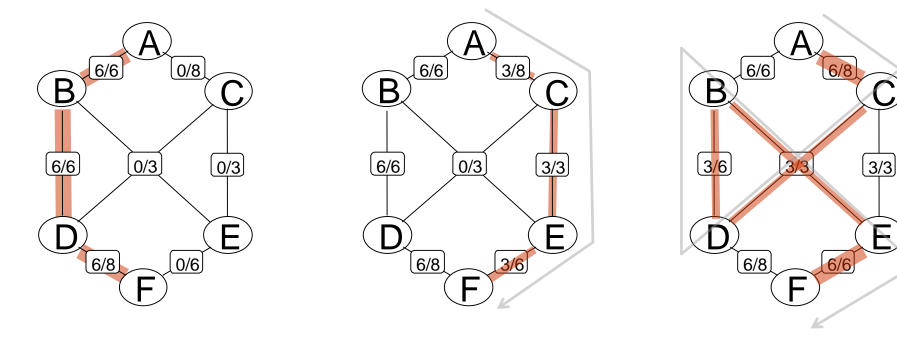
Define an example system

- flow into A
- out of F
- capacities at each edge vary





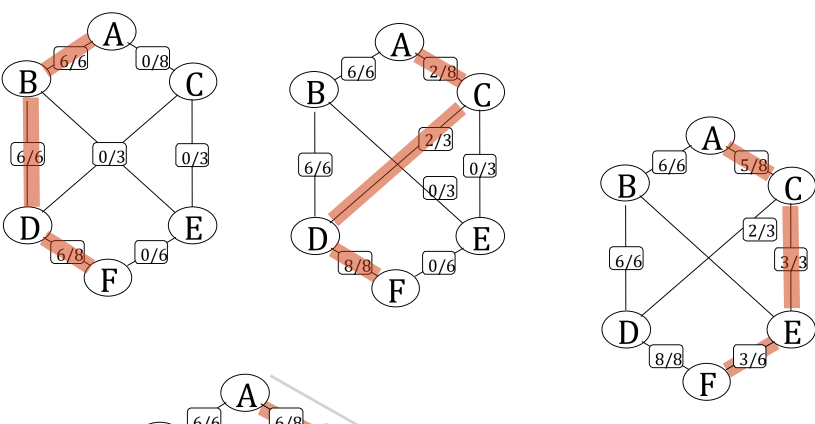




Find a new path (possibly with backwards flow)

- what is the smallest unused capacity on the path ? Δf
 - > 0? send flow Δf in this path

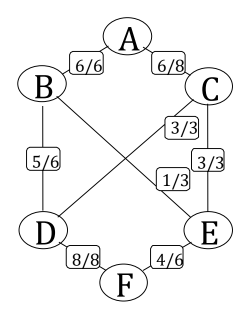
Alternative

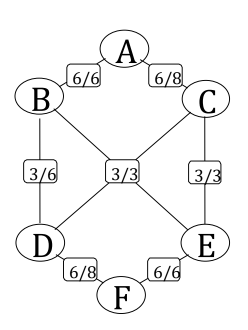


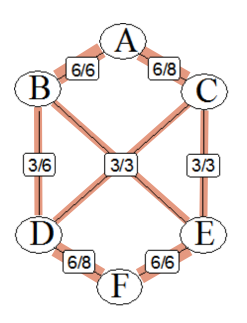
- also ends with flow of 12
- look why this is definitely time to stop ...

Alternative

- are there any paths with unfilled pipes?
- start at A
 - to the left is filled
 - try $A \rightarrow C$
 - both routes out of C are filled
- more solutions?
- definitely different ways to find solutions
 - different order of visiting paths







- path = any route from A to F
- is there any path where all edges have extra capacity?
 - finished flow is maximum
- algorithm (not optimal)

```
while (flow not maximum / path found)
  add flow to path
```

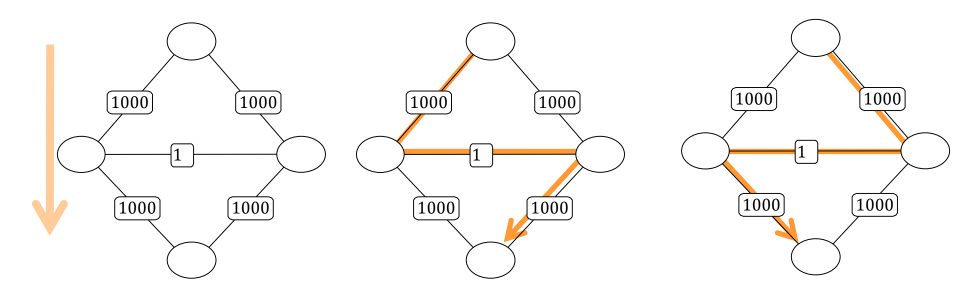
Our definition - finished when

 every path from source to sink has at least one edge (pipe) which is full

Is this efficient?

Efficiency

Worst possible selection of path order would require 2000 iterations



First part of procedure finished

- flow is maximum
- next
 - where to cut graph

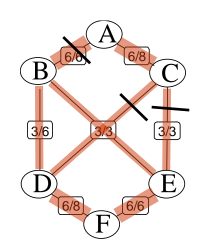
Cutting graph

Find ways to cut network, max flow = 12

- AB, AC capacity =14
- BD, BE, AC capacity = 17
 - both bigger than flow (12)

Better

- for each path
 - find first full pipe cut
 - AB, CD, CE capacity = 12
 - = max flow
 - best cut



Cutting graph

If the capacity across our set of cuts = maximum flow

- it is a "minimum cut"
- smallest connection between two parts of graph
- graph / network / protein is broken into two parts / domains

Useful yet?

- no mention of finding source *s* and sink *t*
- details efficiency not mentioned

Network flow and proteins

Source

- find a surface residue
- connect an s
- connect to nearby surface residues

Sink

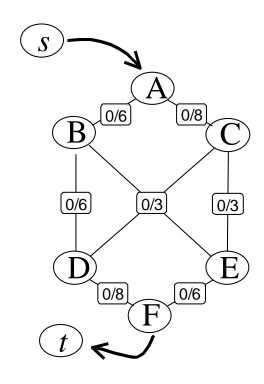
• find a surface residue far away, connect to *t*

Ad hoc? arbitrary? optimal?

maybe not critical

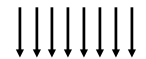
Multiple domains?

while (domains not too small)
 keep trying to split



Background story - Ising spin model

- energy of spin i depends on i, i + 1



For lots of spins

- islands of same spin
- can be generalised to 2D, 3D

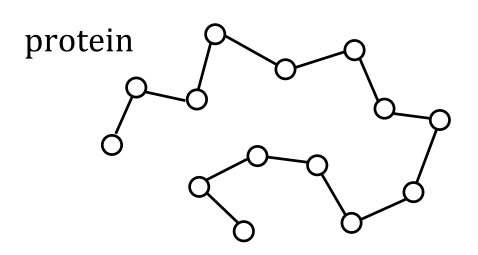
Finding low energies? Simplest method

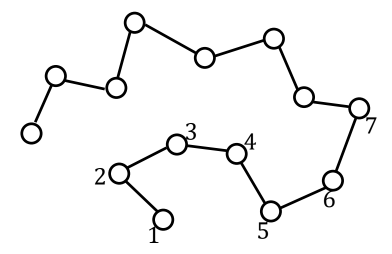
- try to flip a random spin
 - accept flip if energy improves
 - sometimes accept if energy goes up (probabilistic)

Slightly better method

Protein version

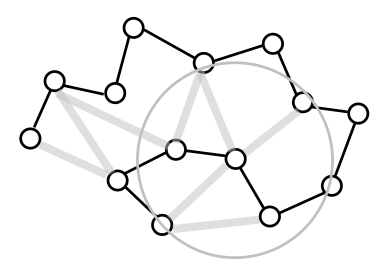
- for any known structure
 - easy to make list of neighbours of each residue
- residues close in space should be in similar domains





label all points with a number

make a list of neighbours for each residue

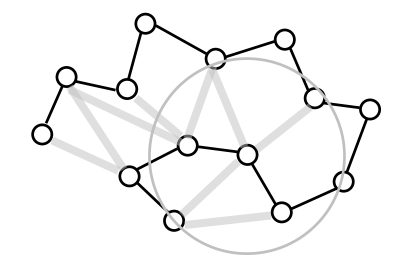


label of a residue is m_i while (labels changing) for each residue j

$$m_{av} = \frac{\sum_{i \in neighbours} m_i}{n_{neighbour}}$$
if $(m_{av} > m_j)$

$$m_j (new) = m_j (old) + 1$$
else if $(m_{av} < m_j)$

$$m_j (new) = m_j (old) - 1$$



step	residue number								
0	1	2	3	4	5	6	7	8	
1	2	3	3	4	6	7	8	8	
2	2	2	4	3	5	8	9	9	

Properties of Taylor / Ising spin-inspired method

Optimism

will converge and become stable

Requires threshold - what is a neighbour

- can use sophisticated averaging distance dependent
- may converge to 2, 3, ... domains

Methods so far

- 1. simple look for single cut points and maximise density
- 2. Crippen / hierarchical clustering
- 3. Network flow
- 4. Ising spin / Taylor
- All methods have arbitrary numbers

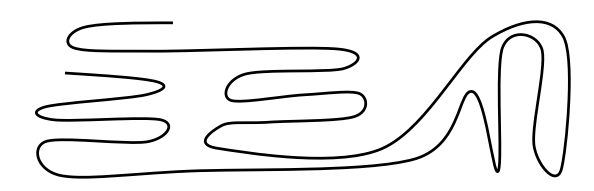
Why are methods so complicated?

If we cut protein chain once

• methods are easy - use density criterion

Cut protein twice? more? remember $(N_{res})^{N_{cut}}$



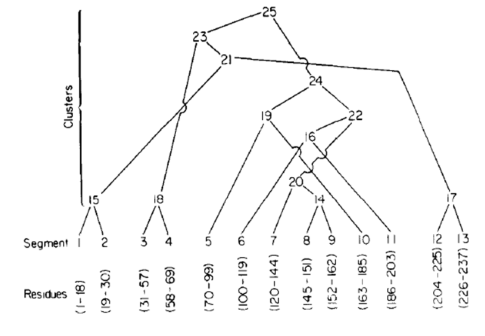


How many domains?

- Crippen / clustering method whatever you want
- Network flow repeatedly split and eventually stop
- Taylor / Ising may converge to > 2 domains

Crippen / hierarchical clustering

At what level of hierarchy do I cut tree?



Network flow

- what constitutes a contact ? (any 2 atom < 4 Å?)
- give pipes (edges) more weight for different kinds of contacts
- are solutions unique?
 - probably in practice
- when do we stop splitting domains?

Taylor / Ising spin method

- what constitutes a contact? how many A?
- type of averaging to get m_{av} ?
- when does one converge?

Elegance

Do methods work as described? not really

All authors report problems - example

- Taylor finds different results for α -helical and β -sheet regions
- simple explanation? distances within / between secondary structure are very different

Do methods work?

With many fixes and tuning - yes

• distance criteria, thresholds

Do methods agree?

Only ask question if you agree to think in terms of structure

answer will be different in terms of evolution or sequences

Criteria

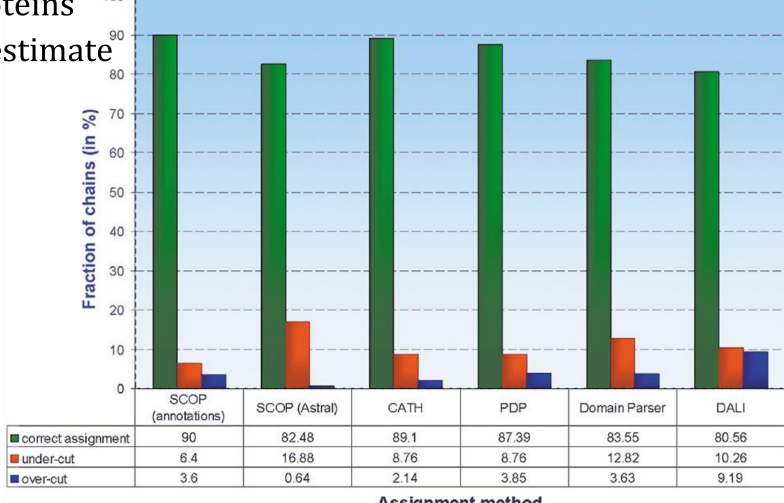
- how many domains inside a protein?
- where are the domain borders?

Number of domains

test set of few hundred proteins

compare against author's estimate 90

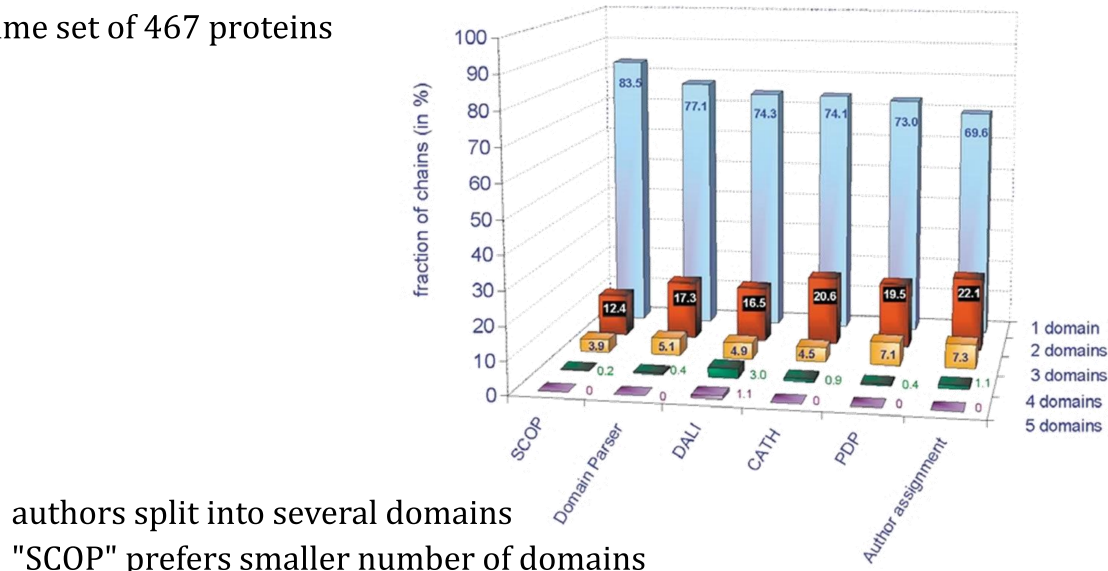
• 80-90 % agreement



Assignment method

How many domains per protein?

Same set of 467 proteins



• "SCOP" prefers smaller number of domains

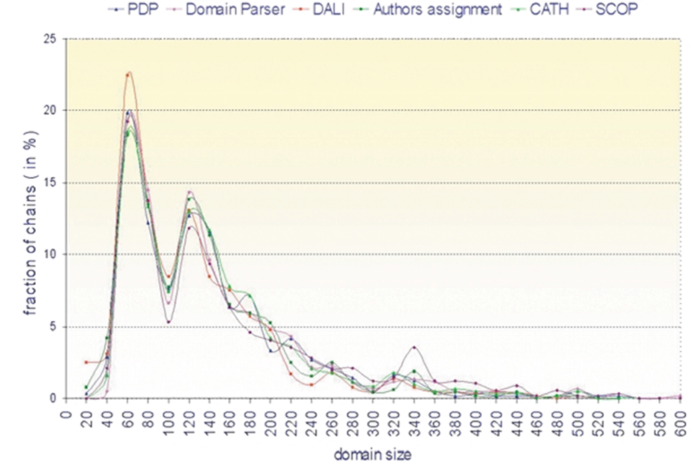
Agreement?

Lots of room for differences

How big is a protein domain?

Peaks near 60 and 130 residues

Some statistics



How complicated are domains?

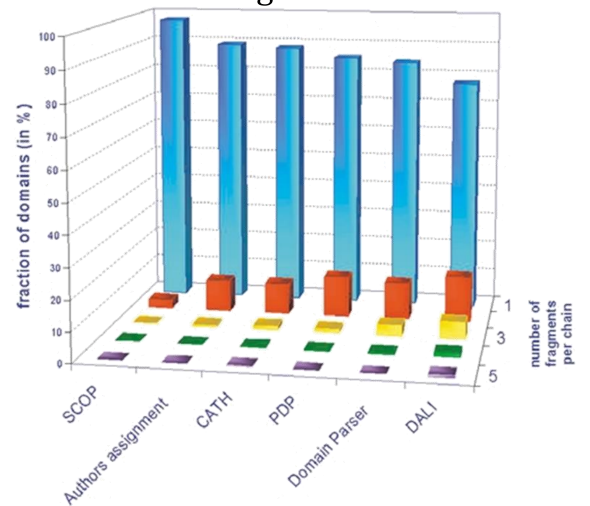
Justification for complicated domain recognition

single cuts, double cuts in chains are not enough

What percentage of domains are built from

- 1 chain?
- 2 chains? ...

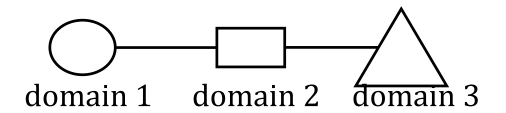
In "DALI", 23 % of domains are not continuous (multiple crosses of chain)



Evolutionary picture

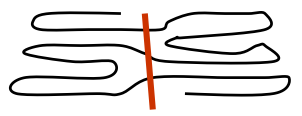
Original claim

domains are units that move as a module in evolution



If we see multiple cuts 10-20 % of time

• picture is much less clear



Summary

Domain definitions

- functional, structural, sequence based
- Finding domains
- relies on contacts, density
- method must be able to handle multiple crossings of chain

We considered

- clustering / hierarchical
- network flow
- Taylor / Ising spin-inspired
- Methods do not agree with each other
- Some trends in size and number of domains
- Real proteins are not as simple as evolutionary picture