### **Protein Threading**

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#### **Goals for Lecture**

Key concepts

- threading prediction task
- threading search task
- template models
- branch and bound search for threading

## **Protein Threading**

- Generalization of homology modeling
  - homology modeling: align sequence to sequence
  - threading: align sequence to structure (templates)
- Key ideas
  - limited number of basic folds found in nature
  - amino acid preferences for different structural environments provide sufficient information to choose among folds

#### A Core Template



Figure from R. Lathrop et al. Analysis and Algorithms for Protein Sequence-Structure Alignment, in *Computational Methods in Molecular Biology*, Salzberg et al. editors, 1998.

## Components of a Threading Approach

- Library of core fold templates
- Objective function to evaluate any particular placement of a sequence in a core template
- Method for searching over space of alignments between sequence and each core template
- Method for choosing the best template given alignments

### Task Definition: Prediction Via Threading

- Given:
  - a protein sequence
  - a library of core templates



 Return: the best alignment of the sequence to a template

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## **Threading Objective Functions**

- Possible sequence-template alignments are scored using a specified objective function
- Objective function scores the sequence-structure compatibility between
  - sequence amino acids
  - their corresponding positions in a core template
- It takes into account factors such as
  - amino acid preferences for solvent accessibility
  - amino acid preferences for particular secondary structures
  - interactions among spatially neighboring amino acids

## **Core Template with Interactions**

Figure from R. Lathrop et al. Analysis and Algorithms for Protein Sequence-Structure Alignment.



Figure from R. Lathrop et al. Analysis and Algorithms for Protein Sequence-Structure Alignment.

 A threading can be represented as a vector t , where each element indicates the index of the amino acid placed in the <u>first position</u> of each core segment



- there are variable length gaps between the core segments, <u>and</u>
- the objective function includes interactions between neighboring (in 3D) amino acids

## A General Pairwise Objective Function

• General objective function with pairwise interactions is:

$$f(\vec{t}) = \sum_{i} g_{1}(i, t_{i}) + \sum_{i} \sum_{j > i} g_{2}(i, j, t_{i}, t_{j})$$

scores for scores for segment interactions individual segments

• We wish to minimize this function

# Searching the Space of Alignments

- If interaction terms between amino acids are <u>not</u> allowed
  - dynamic programming
    - will find optimal alignment efficiently

#### If interaction terms allowed

- heuristic methods
  - fast
  - might not find the optimal alignment
- exact methods (e.g. branch & bound)
  - if they return an alignment, it will be optimal
  - might take exponential time
  - might fail due to time or space limits

## **Branch and Bound Abstractly**

- 3 components
  - A data structure for compactly representing a set of potential solutions
    - May be a very large set
  - An algorithm for computing a lower bound on the score obtained by any member of a set
    - In general, should not explicitly examine all members of the set

- An algorithm for splitting a set into subsets

### **Branch and Bound Search**

initialize Q with one entry representing the set of all threadings repeat

 $l \leftarrow \text{set in } Q$  with lowest lower bound

if *l* contains only 1 threading

return l

else

split *l* into smaller subsets compute lower bound for each subset put subsets in *Q* sorted by lower bound

## **Branch and Bound Illustrated**



- A hypothetical branch and bound search
  - each circle illustrates the space of possible threadings
  - solid lines indicate splits made in previous steps
  - dashed lines indicate splits made in current step
  - numbers indicate lower bounds for newly created subsets
  - arrows show the set that has been split

Figure from R. Lathrop and T. Smith. Global Optimum Protein Threading with Gapped Alignment and Empirical Pair Score Functions. *Journal of Molecular Biology* 255:641-665, 1996.

## **Branch and Bound Search**

- Two key issues
  - how to compute the lower bound for a set of threadings
  - how to split a threading set into subsets
- These determine the expected efficiency

## A Simple Lower Bound

$$\begin{split} \min_{\vec{t} \in T} f(\vec{t}) &= \min_{\vec{t} \in T} \sum_{i} \left[ g_1(i, t_i) + \sum_{j > i} g_2(i, j, t_i, t_j) \right] \begin{cases} \text{objective function} \\ \text{function} \end{cases} \\ &\geq \sum_{i} \left[ \min_{b_i \leq x \leq d_i} g_1(i, x) + \sum_{j > i} \min_{\substack{b_i \leq y \leq d_i \\ b_j \leq z \leq d_j}} g_2(i, j, y, z) \right] \end{cases} \end{split}$$

- Calculate minimum over each term separately
- Can choose different starting positions for the same segment (e.g. x and y for segment i)

#### A Better Lower Bound

- Lower bound is closer approximation to actual score
- Constrain more of the starting indices to be shared instead of choosing them all independently
- Same t<sub>i</sub> appears in multiple terms

# Splitting a Threading Set

- A threading set is split by choosing
  - a single core segment
  - a split point  $s_i$  in the segment
- A simple method
  - split the segment having the widest interval,
    - i.e.  $\max_{i} [d_i b_i]$
  - choose the split point  $s_i$  as the value that results in the lower bound for the set

### Branch and Bound: Splitting a Set



$$T = \left\{ \begin{array}{ccc} \vec{t} \mid b_i \leq t_i \leq d_i, \ b_j \leq t_j \leq d_j, \ b_k \leq t_k \leq d_k, \ b_l \leq t_l \leq d_l \right\}$$

$$(choose \\ segment \\ 2 \\ T = \left\{ \begin{array}{ccc} \vec{t} \mid b_i \leq t_i < s_i, \\ \cdots \end{array} \right\}$$

$$2 \\ T = \left\{ \begin{array}{ccc} \vec{t} \mid t_i = s_i, \\ \cdots \end{array} \right\}$$

$$3 \\ T = \left\{ \begin{array}{ccc} \vec{t} \mid s_i < t_i \leq d_i, \\ \cdots \end{array} \right\}$$

## **Threading Example**

Suppose we have three segments (i, j, k), each of which includes three amino acids. For a given sequence there are three possible starting positions for each segment. Suppose that you are given the following values for the scores of the individual segments and the scores for segment interactions.

g1(j,8) = 9g1(i,2) = 5g1(k,13) = 3g1(i,3) = 2g1(j,9) = 7g1(k, 14) = 4g1(j,10) = 6 g1(k,15) = 1q1(i,4) = 8g2(i,j,2,8) = 1g2(j,k,8,13) = 7 $g_{2}(i,k,2,13) = 1$ g2(i,j,2,9) = 2q2(j,k,8,14) = 8 $g_{2}(i,k,2,14) = 2$  $g_{2}(i,j,2,10) = 2$ g2(j,k,8,15) = 7g2(i,k,2,15) = 5g2(i,j,3,8) = 5g2(j,k,9,13) = 1g2(i,k,3,13) = 5g2(i,j,3,9) = 6g2(i,k,3,14) = 6 $g_{2}(j,k,9,14) = 6$ g2(i,j,3,10) = 4g2(j,k,9,15) = 8 $g_{2}(i,k,3,15) = 4$ g2(i,j,

g2(i,j,4,8) = 7	g2(j,k,10,13) = 11	g2(i,k,4,13) =
g2(i,j,4,9) = 3	g2(j,k,10,14) = 12	g2(i,k,4,14) =
g2(i,j,4,10) = 4	g2(j,k,10,15) = 13	g2(i,k,4,15) =

We'll find the optimal threading using the "simple lower bound" and splitting a set on the segment with the minimal g1 value. When splitting the selected segment, we'll divide it into three intervals of length one.

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## **Branch and Bound Efficiency**

#### • 58 proteins threaded against their "native" (i.e. correct) models

Protein number	PDB code	Protein length	Number of core segments	Search Space Size	Number of search iterations	Total (search-only) seconds	Equivalent threadings per iteration	Equivalent threadings per second
1	256b	106	5	6.19e + 3	6	1 (1)	1.03e + 3	6.19e + 3
2	1end	137	3	4.79e + 4	6	1 (1)	7.98e + 3	4.79e + 4
3	1rcb	129	4	5.89e + 4	7	1 (1)	8.41e + 3	5.89e + 4
4	2mhr	118	4	9.14e + 4	7	1 (1)	1.31e + 4	9.14e + 4
5	351c	82	4	1.12e + 5	5	1 (1)	2.24e + 4	1.12e + 5
6	1bgc	174	4	1.63e + 5	6	1 (1)	2.72e + 4	1.63e + 5
7	1ubq	76	5	1.70e + 5	6	1 (1)	2.83e + 4	1.70e + 5
8	1mbd	153	8	1.77e + 5	10	1 (1)	1.77e + 4	1.77e + 5
9	11is	136	5	5.02e + 5	7	1 (1)	7.17e + 4	5.02e + 5
10	1aep	161	5	5.76e + 5	13	1 (1)	4.43e + 4	5.78e + 5
50 51 52	5tmn 1lec 1nar	316 242 290	14 15 17	6.51e + 18 7.01e + 18 2.33e + 19	• 164 320 3984	28 (7) 26 (12) 208 (183)	3.97e + 16 2.19e + 16 5.85e + 15	2.32e + 17 2.70e + 17 1.12e + 17
53	1s01	275	15	4.36e + 19	541	32 (13)	8.05e + 16	1.36e + 18
54	5cpa	307	16	1.22e + 20	1089	72 (50)	1.12e + 17	1.69e + 18
55	9api	384	17	1.95e + 22	290	57 (25)	6.71e + 19	3.41e + 20
56	2had	310	19	2.57e + 22	4027	201 (179)	6.39e + 18	1.28e + 20
57	2cpp	414	20	6.37e + 24	3068	205 (164)	2.08e + 21	3.11e + 22
58	otaa	478	23	9.63e + 31	4917	1409 (1267)	1.96e + 28	6.83e + 28
Table from R. Lathrop and T. Smith, <i>Journal of Molecular Biology</i> 255:641-665, 1996.								25