Iterative Refinement Unit Tests

Hint: Many test values are taken from project Algorithms for Bioninformatics of Alexander Mattheis or the lectures.

Test 1 (used **Feng-Doolittle** implementation and **Sum-of-Pairs** implementation)

Input

Sequence a: ACGT Sequence b: AT Sequence c: GCT Sequence d: GC

Gap opening: -1 Enlargement: -3

Match: 1 (and 0 for placeholder #)

Mismatch: 0

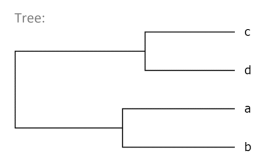
Output (Computation: Global Primary Library)

Feng-Doolittle Output

Alignment: Distance-Table:

ACGT					
Α	T				
G	_CT				
G	_C_				

D	а	b	С	d
а	0	1.8	1.7	2.9
b	1.8	0	2.2	7.6
С	1.7	2.2	0	1.5
d	2.9	7.6	1.5	0



Round-Robin – Minimum Distance Pair Output

Post-Order-Traversal: badc

removed

sequence (Hint: is cleaned from gaps)

b:
$$\underset{x}{\operatorname{argmin}}(D_{b,x}) = a \text{ with } D_{b,a} = 1.8$$

b~a: AT~ACGT

A___T

ACGT

Nothing changes!

a: $\underset{x}{\operatorname{argmin}}(D_{a,x}) = c \text{ with } D_{a,c} = 1.7$

a~c: ACGT~GCT

ACGT |* * GC_T A_T GCT

remaining

alignment

ACGT

G_CT G_C_

GC_

(gap-only columns removed)

after realignment:

GC_ Score: -26

Hint: This alignment is kept.

d:
$$\underset{x}{\operatorname{argmin}}(D_{d,x}) = c \text{ with } D_{d,c} = 1.5$$
 ACGT

d~c: GC~GC#T

GC_T

GC___

GC#T

Nothing changes!

Feng-Doolittle Alignment: Refined Realignment:

 ACGT
 ACGT

 A__T
 A__T

 G_CT
 GC_T

 G_C_
 GC__

Round-Robin – Pairwise Best Pairs Output

(analogous - but it is searched for the best and not the nearest alignment)

Round-Robin - One-Vs-All Output

(analogous - but it is searched for the best and not the nearest alignment)

The other used procedures are already have been already tested during T-Coffee and Feng-Doolittle.