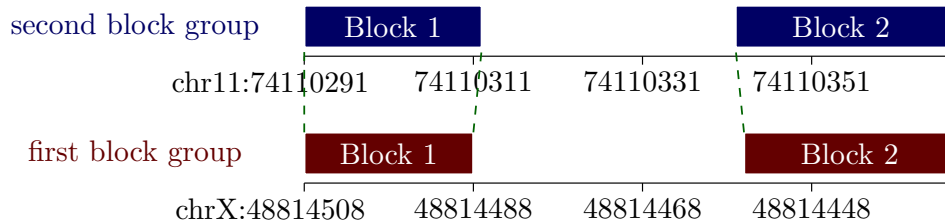


deepBlockAlign v1.0

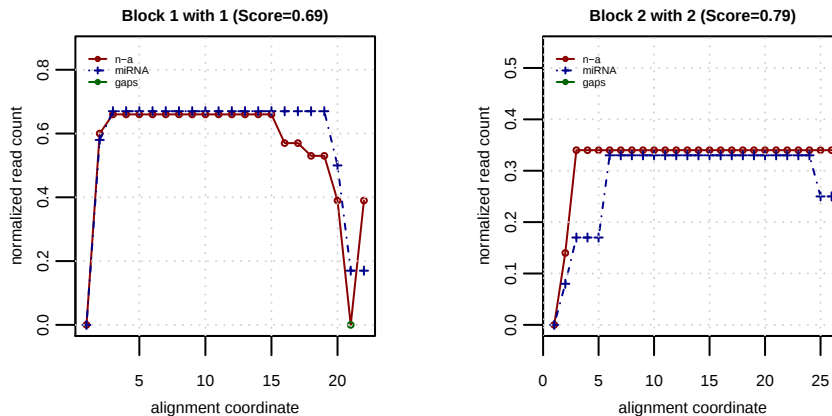
Alignment parameters:

[Input]	
date and time	Sun, 03 Feb 2013 01:09:36 +0100
first block group	BloodGm12878Rep2_cluster.19321
first file	/home/projects/users/sachin/frontier/putative_mirna//tmp1500query.tmp
second block group	bg_125
second file	/home/projects/users/sachin/frontier/bgs/mirbase.all.map.bgs.flagged.sig
[Block group alignment]	
distance weight	1
block weight	1
gap penalty	-1
[Block alignment]	
gap initialization penalty	-2
gap extension penalty	-1
threshold for expression difference	1
match score	1
mismatch score	-1
profile shape difference penalty	1
[Block group information]	
Entropy (first block group)	1.53
Reads (first block group)	14.66
% Unique reads (first block group)	82
Entropy (second block group)	1.78
Reads (second block group)	12.00
% Unique reads (second block group)	100

Block group alignment: BloodGm12878Rep2-cluster-19321-n-a(-) with bg-125-hsa-mir-548al-MI0016851(+) [Score: 0.82]



Block alignment:



Citation: Langenberger, D*, Pundhir, S*, Ekstrøm, C.T., Stadler, P.F., Hoffmann, S., and Gorodkin, J.(2012). deepBlockAlign: A tool for aligning RNA-seq profiles of read block patterns. *Bioinformatics*, **28**(1),17-24. (*equal contribution)