Description of LIAYSON

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1 Example

- > library(liayson)
- > ## Load data and map genes onto segments
- > data(epg)
- > data(segments)
- > X=aggregateSegmentExpression(epg, segments, mingps=20, GRCh=38)
- > head(X\$eps[,1:3]); ##Aggregate expression of first three cells

GGACGTCTCTATCCTA-1 AGCTCTCCGCGTTT-1 CAGCATACACC	MUIIN I
1:2360001-28080000 0.5071770 0.3827751 0.	1913876
1:32720001-248900000 0.7212614 0.5564598 0.	3072228
2:1-88940000 0.8058824 0.5558824 0.	3411765
2:91820001-242040000 0.6964657 0.5322245 0.	3659044
3:23580001-60260000 0.4837209 0.3860465 0.	2139535
3:93840001-147160000 0.5737705 0.4426230 0.	2131148

- > ## Calculate number of expressed genes per cell
- > data(epg)
- > gpc = apply(epg>0, 2, sum)
- > ## Calculate copy number from expression
- > cn=segments[rownames(X\$eps),"CN_Estimate"]
- > cnps = segmentExpression2CopyNumber(X\$eps, gpc, cn, nCores=1)
- > head(cnps[,1:3]); ##Copy number of first three cells

	$\tt GGACGTCTCTATCCTA-1$	AGCTCTCTCCGCGTTT-1	CAGCATACACCAGTTA-1
1:2360001-28080000	1	1	1
1:32720001-248900000	2	2	2
2:1-88940000	2	2	2
2:91820001-242040000	2	2	2
3:23580001-60260000	1	1	1
3:93840001-147160000	2	2	2

> outc = clusterCells(cnps, h=0.05)

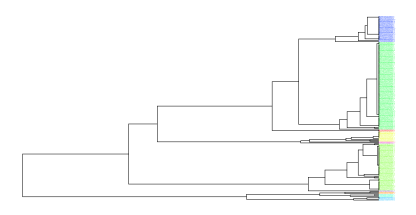


Figure 1: Cells are clustered based on their copy number