cfDNAKit

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5/13/2021

 $This\ rmarkdown\ to\ demonstrate\ how\ cfdnak it\ work.\ First,\ load\ cfdnak it\ package\ to\ environment$

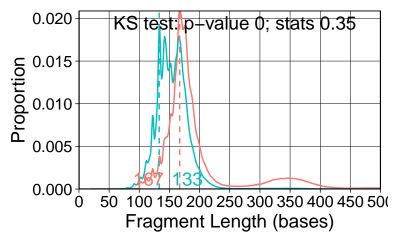
```
library(cfdnakit)
```

Let cfdnakit read sequence alignment file (.bam) with function read_bamfile. This function will split sequence reads into equal-size non-overlapping windows. Available size of bin are 100, 500, and 1000 KB.

[1] "Number of bin : 2888"

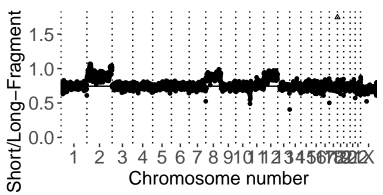
Extract cfDNA fragment length information.

Healthy Control — plasma-01-01_OE0290-P



fragment length profile distribution-1.pdf

```
plot_sl_ratio(sample_profile)
```



genome-wide SLRatio-1.pdf

Save fragment profile as RDS file for later use or for creating Panel-of-Normal

```
save_fragment_profile(sample_profile,
```

output_dir = "/icgc/dkfzlsdf/analysis/0E0290_projects/pediatric_tumor/whole_genom

[1] "Saving RDS : Done"

Making a Panel-of-Normal is necessary for downstream analysis as we want to compare fragment profile between a cfDNA from patient with pooled of healthy individuals. First, we create a text file where each line is a full path to rds file created by aforementioned function.

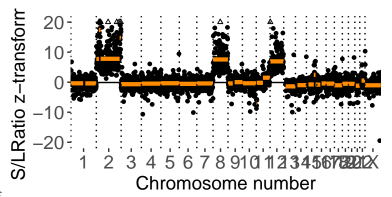
```
path_to_PoN_txt = "/icgc/dkfzlsdf/analysis/0E0290_projects/pediatric_tumor/whole_genome_sequencing/proc
create_PoN(path_to_PoN_txt,output_dir = "/icgc/dkfzlsdf/analysis/0E0290_projects/pediatric_tumor/whole_
```

[1] "Done"

We rescale the SLRatio by the median of PoN samples.

```
PoN_rdsfile = "/icgc/dkfzlsdf/analysis/0E0290_projects/pediatric_tumor/whole_genome_sequencing/processissample_zscore = get_zscore_profile(sample_profile,PoN_rdsfile)
```

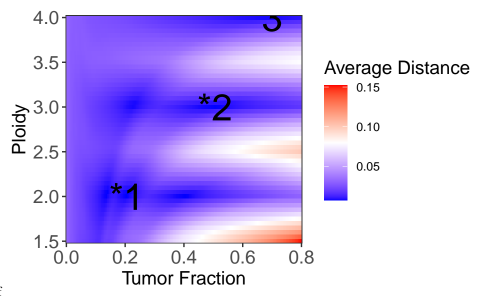
```
## Reading PoN profile /icgc/dkfzlsdf/analysis/0E0290_projects/pediatric_tumor/whole_genome_sequencing/
sample_zscore_segment = segmentByPSCB(sample_zscore)
plot_transformed_sl(sample_zscore,sample_zscore_segment)
```



Panel-of-Normal samples and plot scaled value-1.pdf

Performing CNV Calling and plot distance matrix.

```
sample_cnv = call_cnv(sample_zscore_segment,sample_zscore)
plot_distance_matrix(sample_cnv)
```

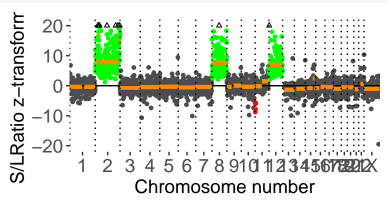


calling and plot distance matrix-1.pdf

```
solution_table = get_solution_table(sample_cnv)
solution_table
```

We select a solution and plot the CNV Calling result.

plot_cnv_solution(sample_cnv,selected_solution = 1)



cnv-calling first solution-1.pdf

Calculate PCA Score from the segmentation result

calculate_PCA_score(sample_zscore_segment)

[1] 169.017