# Using savR

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> library(savR)

```
> fc <- savR(system.file("extdata", "MiSeq", package="savR"))</pre>
```

> fc

savProject instance with 1 lanes, 82 total cycles and 1 sequencing runs.

> pfBoxplot(fc)

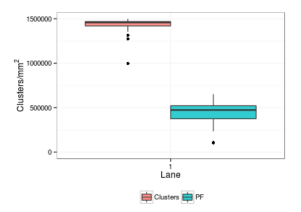


Figure 1: Boxplot of total vs. PF clusters

## Introduction

The Illumina Sequence Analysis Viewer (SAV) is a Windows application provided by Illumina that presents graphs made in real time from data collected over the course of basecalling. This data was previously also made available in HTML format for inspection after the run; however, it is now preserved in binary format and not simply parsed by users who wish to perform automated quality assessment. Here is presented *savR*, an R package to parse the binary output, generate QC assessment plots and make the data available to users of Illumina sequencing instruments. For more information about Illumina SAV, please consult the Illumina iCom website and the Sequencing Analysis Viewer User's Guide, available *online*.

# Description

The savR function is passed a path to an Illumina HiSeq or MiSeq run, and returns a savProject object, containing the parsed data. Accessor methods are available for information in the RunInfo.xml file as well as the parsed SAV Metrics files. These include corrected intensities, quality metrics, tile metrics, and extraction metrics. The *savR* package comes with an example MiSeq data set which can be loaded thusly:

```
> fc <- savR(system.file("extdata", "MiSeq", package="savR"))</pre>
```

### RunInfo.xml

The RunInfo.xml file is parsed and stored in the slots of the savProject object. There are accessor methods for the project's location, reads, number of "ends" or directions, the run ID, the number of cycles, and a description of the flowcellLayout.

```
> directions(fc)
[1] 1
> reads(fc)
[[1]]
An object of class "illuminaRead"
Slot "number":
[1] 1
Slot "cycles":
[1] 76
Slot "index":
[1] FALSE
[[2]]
An object of class "illuminaRead"
Slot "number":
[1] 2
Slot "cycles":
[1] 6
Slot "index":
[1] TRUE
> cycles(fc)
[1] 82
> flowcellLayout(fc)
An object of class "illuminaFlowCellLayout"
Slot "lanecount":
[1] 1
Slot "surfacecount":
[1] 2
```

Slot "swathcount":
[1] 1

Slot "tilecount":
[1] 19

#### Corrected intensitites

Corrected intensity metrics (obtained from CorrectedIntMetricsOut.bin) can be inspected by the correctedIntestites accessor method:

```
> head(correctedIntensities(fc), n=1)
  lane tile cycle avg_intensity avg_cor_A avg_cor_C avg_cor_G avg_cor_T
2
                                        72
     1 1101
                1
                              80
                                                  17
                                                            116
                                                                      101
  avg_cor_called_A avg_cor_called_C avg_cor_called_G avg_cor_called_T num_none
2
               212
                                 266
                                                  283
                                                                    277
                                                                             339
  num_A num_C num_G num_T sig_noise
2 97572 17051 136607 127150
                             6.704378
```

This is a data.frame of intensity metrics; one line for each set of lane, tile and cycle measurements. Reported statistics include average intensity, corrected intensity (for cross-talk between bases and phasing/pre-phasing), called corrected intensities, number of called bases and signal to noise ratio. There are methods which act upon savProject objects to produce QC plots, for example plotIntensity to assess signal intensity for each channel as in figure 2.

#### > plotIntensity(fc)

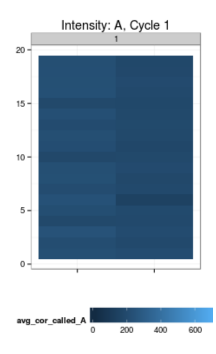


Figure 2: Corrected intensity plot: cycle 1, base "A".

### **Quality Metrics**

The quality metrics (QMetricsOut.bin) file contains per-lane/tile/cycle metrics for the number of clusters with quality at each PHRED value from 1-50.

```
> head(qualityMetrics(fc), n=1)
```

lane tile cycle Q1 Q2 Q3 Q4 Q5 Q6 Q7 Q8 Q9 Q10 Q11 Q12 Q13 Q14 Q15 Q16 Q17 1 1101 0 1 1 0 0 0 0 0 0 0 0 0 0 17087 0 0 0 0 0 Q18 Q19 Q20 Q21 Q22 Q23 Q24 Q25 Q26 Q27 Q28 Q29 Q30 Q31 Q32 Q33 1 0 0 0 6562 0 5323 5901 0 0 13362 0 0 250 22276 13477 12757 Q34 Q35 Q36 Q37 Q38 Q39 Q40 Q41 Q42 Q43 Q44 Q45 Q46 Q47 Q48 Q49 Q50 1 281724 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0

```
> qualityHeatmap(fc,1,1)
```

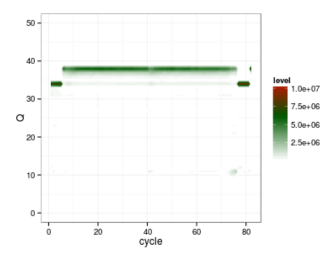


Figure 3: Quality heatmap: lane 1, read 1.

## Tile Metrics

The tile metrics (TileMetricsOut.bin) file contains coded information about per-lane/cycle/tile cluster density, pass-filter clusters, phasing and pre-phasing data. Consult the tileMetrics help page for more information.

```
> head(tileMetrics(fc), n=4)
   lane tile code
                       value
39
      1 1101
              100 1271891.8
41
      1 1102
              100 1455525.0
43
      1 1103
               100
                    995404.6
45
      1 1104
              100 1463691.8
```

## Extraction Metrics

The extraction metrics (ExtractionMetricsOut.bin) file contains per-lane/cycle/tile information about per-base FWHM (full width pixel size of clusters at half maximum) and 90th %-ile intensity of signal intensity.

```
> head(extractionMetrics(fc), n=1)
lane tile cycle FWHM_A FWHM_C FWHM_G FWHM_T int_A int_C int_G int_T
2 1 1101 1 2.235387 2.308783 1.86132 2.174398 180 326 357 400
```

# Coda

There is a convenience function (buildReports), which partially reconstructs the Illumina reports folder that was previously generated by the Illumina instrument software and which was superseded by SAV and InterOp files.