Cell3[™] Target: Cell Free DNA Target Enrichment



De-multiplexing Cell3TMTarget Data - Illumina Sequencing

This document outlines an example procedure for sample de-multiplexing of pooled Cell3TMTarget libraries following Illumina sequencing. The i7 indexing read contains a 9bp molecular tag in addition to the unique 8bp sample index. The i5 index contains 8bp unique sample index only. The final read structure is Y*, I8Y9,I8,Y*.

Input requirements

Protocol Guide v1.1.1

- Illumina bcl2fastq software a Base Call (BCL) Files to Fastq conversion software
 - http://emea.support.illumina.com/sequencing/sequencing_software/bcl2fastq-conversion-software.html
- 2. Basecalls directory
- 3. Samplesheet.csv ensure there are no N's present in the sample sheet (examples provided on request)
- 4. Check RunInfo.xml to ensure the correct number of cycles have been performed for the indexes
 - a. i7 = 17 cycles
 - b. i5 = 8 cycles

Expected file outputs per sample

- 1. Read 1 fastq file (R1)
- 2. Read 2 fasta file (R3)
- 3. fastq file containing information on a per fragment molecular tag (R2)
- 4. Index 1 and 2 fastq files (these are not required)

Procedural steps

- 1. Open a command terminal
- 2. Move into the basecalls directory of the data to be de-multiplexed

* replace with the number of cycles performed for read 1 and 2

 Run the following bcl2fastq command bcl2fastq --create-fastq-for-index-reads --mask-short-adapter-reads 0 -use-bases-mask Y*,l8Y9,l8,Y* --no-lane-splitting (optional)