

INSTRUCTIONS digitalMLPA Data Demo Set

SALSA[®] digitalMLPA[™] Probemix D007-A1 Acute Lymphoblastic Leukemia

1. Purpose

This digitalMLPA data demo set serves to provide an idea of possible results that can be obtained with SALSA® digitalMLPA[™] Probemix D007-A1 Acute Lymphoblastic Leukemia. It contains data files of ten samples with different genomic profiles and five reference samples. This data demo set also allows you to familiarise yourself with Coffalyser digitalMLPA[™] and the way digitalMLPA results are presented by the software.

The Sample Results PDF reports that are included in this data demo set offer you the possibility to have a look at the sample results without analysing the data.

2. **Description of samples included**

DNA Sample	Barcode	Expected Results
REF-Male-1	BP02-33	Reference sample: normal ratios for all probes.
Coriell NA10925	BP02-34	7p : Heterozygous deletion affecting probes for <i>ADCY1</i> , <i>ABCA13</i> and <i>IKFZ1</i> .
		Xp : Heterozygous duplication affecting the probes for <i>CRLF2</i> and <i>CSF2RA</i> exon 1–10.
Coriell NA03184	BP02-35	15q : Trisomy chromosome 15 affecting the probes for <i>ATP10A</i> , <i>OCA2</i> , <i>SPRED1</i> , <i>VPS13C</i> , <i>USP3</i> , <i>CHSY1</i> and <i>TM2D3</i> .
Coriell NA04371	BP02-36	5q : Heterozygous gain affecting the probes for <i>EBF1</i> , <i>COL23A1</i> , <i>SQSTM1</i> , <i>MAPK9</i> and <i>FLT4</i> .
		Xp : Heterozygous gain affecting the probes for CSF2RA.
REF-Male-2	BP02-37	Reference sample: normal ratios for all probes.
Coriell NA07106	BP02-38	22q : Trisomy chromosome 22 affecting the probes for GAB4, CECR2, VPREB1, IGLL1, ZMAT5, SFI1, TRMU and BRD1.
Coriell NA00501	BP02-39	2p : Heterozygous deletion affecting the <i>TMEM18</i> probe.
		NR3C2, CYP4V2 and TRIML1.
Coriell NA00782	BP02-40	4q : Heterozygous gain affecting the probes for <i>TECRL</i> , <i>UGT2A1</i> , <i>LEF1</i> , <i>INPP4B</i> and <i>ZNF827</i> .
		Xp : Heterozygous gain affecting all probes for CSF2RA.
Coriell NA02030	BP02-41	8p : Trisomy chromosome 8 affecting the probes for <i>FBXO25</i> , <i>CLN8</i> ,
		9 Trisomy chromosome 8 affecting the probes for SNTC1 TOX CHD7
		<i>RMDN1, CPNE3, TSNARE1</i> and <i>SLC39A4.</i>
REF-Male-3	BP02-42	Reference sample: normal ratios for all probes.
Coriell NA03563	BP02-43	3q : Heterozygous gain affecting the probes for CD200, BTLA, XRN1, SI C9A9, TBI 1XR1, LINC00501, OPA1 and ACAP2.

9p: Heterozygous deletion affecting the probes for *DOCK8* and *KANK1*.

11p: Heterozygous deletion affecting the probes for *LMO2*, *CD44*, *SLC1A2*

Та

Coriell NA06803

BP02-45

3. Analysis in Coffalyser digitalMLPA software

You can analyse the samples with Coffalyser digitalMLPA using either the FASTQ file or the Coffa files provided.

3.1. Analysis using FASTQ file

The FASTQ file *FASTQ digitalMLPA Data Demo Set D007-A1.fastq.gz* is generated by the Illumina[®] NGS platform. It is provided in a zip archive, which can be imported directly in Coffalyser digitalMLPA without the need to unpack the archive first. It is recommended to use this file if you wish to do the complete analysis from start to finish. Follow the procedure for creating a new experiment as described in the *Coffalyser digitalMLPA User Manual*.

In several steps described in the User Manual, you are required to enter specific information into Coffalyser digitalMLPA:

- Step 6: Select the barcode lot on 03-xxx-yymmdd; Illumina standard.
- Step 8: Select the barcodes listed in Table 1.
- Step 11: Select *digitalMLPA* from the experiment type drop-down menu.
- Step 12: Select *D007* from the product drop-down menu.
- Step 13: Select D007-A1-0922 default tube C599CB 12-09-22 [default sheet] from the sheet drop-down menu.
- Step 15: Select *Reference* from the drop-down menu in the column *type* for the reference samples (see Table 1). All other samples can be defined as *Test*.
- Step 16: Select *Pooled DNA source* from the drop-down menu in the column *options* for the reference samples. Leave this on *Default* for all other samples.

3.2. Analysis using Coffa files

Alternatively, you can perform an analysis using the Coffa files included ¹. A Coffa file is a file (*.coffa) containing the sequencing output and several analysis properties of a single digitalMLPA reaction. Analysis using the Coffa files is faster than using the FASTQ file.

For instructions, consult the *Coffalyser digitalMLPA User Manual* for the procedure on reanalysis of an experiment using Coffa files.

¹ The Coffa files have been generated with Coffalyser digitalMLPA version 221020.1234. They may not be compatible with newer versions of the software.