

# Project Overview

J.Lundeberg\_14\_01

## Project Information

<b>User Project Name</b>	140117_Rapid_Ventana_TdT
<b>NGI Project Name</b>	J.Lundeberg_14_01
<b>NGI Project ID</b>	P955
<b>User Contact</b>	<a href="mailto:jlundeberg@ki.se">jlundeberg@ki.se</a>
<b>NGI Application Type</b>	Finished Library (No best practice analysis)
<b>Samples &amp; Lanes</b>	1 sample, 2 lanes
<b>Project Status</b>	Sequencing Finished
<b>Order Dates</b>	<i>Order received:</i> 2014-01-10, <i>Contract received:</i> 2014-01-15, <i>Samples received:</i> 2014-01-20, <i>Queue date:</i> 2014-01-23, <i>All data delivered:</i> 2014-04-02, <i>Report Date:</i> 2014-12-02
<b>UPPMAX Project ID</b>	b2011007
<b>UPPNEX project path</b>	/proj/b2011007/INBOX/J.Lundberg_14
<b>Reference Genome</b>	Human, hg19
<b>Minimum ordered reads</b>	200 million

## Methods

### Library construction

- A) Library was prepared using the “650 bp insert standard DNA (Illumina TruSeq DNA)” protocol and clustering was done by cBot.

### Sequencing

- A) All samples were sequenced on HiSeq2500 (HiSeq Control Software 2.0.12.0/RTA 1.17.21.3) with a 2x101 setup. The Bcl to Fastq conversion was performed using bcl2Fastq v1.8.3 from the CASAVA software suite. The quality scale used is Sanger / phred33 / Illumina 1.8+.

## Data Flow

All data (demultiplexed) from the instrument are collected and transferred securely to a storage server with well established pipeline.

## Data Processing:

A set of standard quality checks were performed to assure that all sequenced data meet NGI guaranteed quality / quantity. All analysis are carried out in UPPMAX servers before delivering the raw data.

## Swedac Accreditation

The National Genomics Infrastructure is accredited by [Swedac](#). This means that our services are subject to highly stringent quality control procedures, so that you can be sure that your data is of excellent quality.

Library preparation	<input type="checkbox"/> Not Swedac Accredited
Sequencing data	<input type="checkbox"/> Swedac Accredited
Data flow	<input type="checkbox"/> Swedac Accredited
Data processing	<input type="checkbox"/> Swedac Accredited

## Sample Info

NGI ID	User ID	Index	Lib Prep
P955_101	140117_Rapid_Ventana_TdT	Index 8 (ACTTGA)	A

## Yield Overview

Sample	Lib QC	Avg. FS	≥ Q30	# Reads	Status
P955_101b	Passed	350 bp	59.34%	105.66 M	Passed

- *Lib QC*: Reception control library quality control step
- *Avg. FS*: Average fragment size.
- *≥ Q30*: Percentage of bases above quality score Q30 for the sample.
- *# Reads*: Millions of reads sequenced.

## Run Info

Date	FC id	Lane	Clusters	% PhiX	≥ Q30	% Unique	Method
140123	B-H8A63ADXX	1	66.88 M	0.52%	58.70%	80.51%	A
140123	B-H8A63ADXX	2	65.89 M	0.56%	57.15%	78.32%	A

- *FC id*: Position on flowcell - Flowcell ID.
- *≥ Q30*: Percentage of bases above quality score Q30 on the lane.
- *Unique*: Percentage of reads recovered after demultiplexing.
- *Method*: Sequencing method used. See above for description.

## General Information

### Naming conventions

The data is delivered in FastQ format using Illumina 1.8 quality scores. There will be one file for the forward reads and one file for the reverse reads (if the run was a paired-end run).

The naming of the files follow the convention:

[LANE]\_[DATE]\_[POSITION][FLOWCELL]\_[NGI-NAME]\_[READ].fastq.gz

### Data access at UPPMAX

Data from the sequencing will be uploaded to the UPPNEX (UPPMAX Next Generation sequence Cluster Storage, [uppmx.uu.se](http://uppmx.uu.se)), from which the user can access it. You can find the data in the INBOX folder of the UPPNEX project, which was created for you when your order was placed:

/proj/b2011007/INBOX/J.Lundberg\_14

If you have problems accessing your data, please contact SciLifeLab [genomics\\_support@scilifelab.se](mailto:genomics_support@scilifelab.se). If you have questions regarding UPPNEX, please contact [support@uppmx.uu.se](mailto:support@uppmx.uu.se).

## Acknowledgements

In publications based on data from the work covered by this contract, the authors must acknowledge SciLifeLab, NGI and Uppmax:

*The authors would like to acknowledge support from Science for Life Laboratory, the National Genomics Infrastructure, NGI, and Uppmax for providing assistance in massive parallel sequencing and computational infrastructure.*

## Further Help

If you have any queries, please get in touch at [genomics\\_support@scilifelab.se](mailto:genomics_support@scilifelab.se).

