### Lung Precancer Analysis

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#### Overview

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- Methods
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### Introduction

### Lung Cancer

- Squamous cell carcinoma
- Adenocarcinoma

#### Precancer

#### Introduction

Study Objectives

## Study Objectives

- Find different mutations
  - between WES
  - between WTS
  - from cancer
  - from precancer
- Pathway examine from the mutations
  - of WES
  - of RNA-seq
- Ultra-deep sequencing to find an infinitesimal quantity of Non-Circulating Tumor DNA
  - from blood
  - from urine
  - frrom bronchus
- Diagnostic performace

#### Materials

### Lung Cancer Data

- WES + WTS
- Normal + {Primary, CIS + AIS, AAH, Dysplasia, MIA}
- Total 112 samples

## Materials

Cancer Types

#### CIS + AIS

• Carcinoma in situ + Adenocarcinoma in situ

#### **AAH**

• Atypical adenomatous hyperplasia

## Dysplasia



#### MIA

• Minimally invasive adenocarcinoma

## Materials

Sample Count

## Sample Count in WES

## Sample Count in Transcriptome

## Methods

#### Methods

Workflows

### Data pre-processing for variant discovery

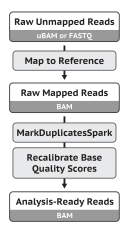


Figure: Data pre-processing for variant discovery (Van der Auwera et al., 2013; DePristo et al., 2011)

#### Somatic short variant discovery

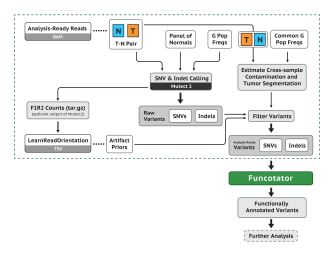


Figure: Somatic short variant (SNVs + Indels) discovery workflow (Van der Auwera et al., 2013; DePristo et al., 2011)

## Germline short variant discovery

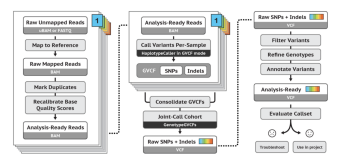


Figure: Germline short variant (SNVs + Indels) discovery workflow (Van der Auwera et al., 2013; DePristo et al., 2011)

## RNA-seq short variant discovery

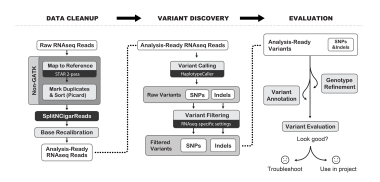


Figure: RNA-seq short variant (SNVs + Indels) discovery workflow (Van der Auwera et al., 2013; DePristo et al., 2011)

#### Methods

Miscellaneous

#### **Used Bioinformatics Tools**

- FastQC (Andrews et al., 2012)
- Sequenza (Favero et al., 2015)
- BWA (Li & Durbin, 2009; Li, 2013)
- STAR (Dobin et al., 2013)
- Bowtie2 (Langmead & Salzberg, 2012)
- Samtools (Li et al., 2009)
- GATK (Van der Auwera et al., 2013; DePristo et al., 2011)
- Picard (Picard toolkit, 2019)
- VCF2MAF (Kandoth et al., 2018)
- VEP (McLaren et al., 2016)

## Python Packages

- Pandas (pandas development team, 2020; Wes McKinney, 2010)
- Sequenza-utils (Favero et al., 2015)
- Matplotlib (Hunter, 2007)
- Seaborn (Waskom & the seaborn development team, 2020)
- CoMut (Crowdis, He, Reardon, & Van Allen, 2020)

#### Results

#### Results

Quality Checks with FastQC

## FastQC?

#### FastQC on WES

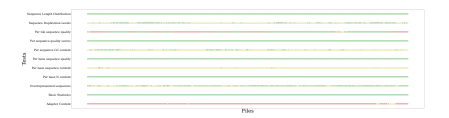


Figure: FastQC with WES data

- ... Only 33P1 has more than 3 failures: 6 FAILs.
- $\therefore$  33P1 is excluded at further analysis.

## FastQC on WTS



Figure: FastQC with WTS data

- ... No sample has more than 5 failures.
- ... All sample are good to analysis.

#### Results

Quality Checks with Sequenza

# Sequenza?

## Cellularity & Ploidy on WES

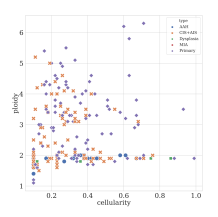


Figure: Cellularity and Ploidy from Sequenza

## Copy Number Variation on WES

Results

Mutect2

### Mutect2?

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