Lung Cancer

Jaewoong Lee 2021-01-22

Contents

1	Intr	oduction	3
2	Mat	erials	3
3	Met	hods	3
	3.1	Genome Analysis Toolkit	3
	3.2	Alignment	3
		3.2.1 Burrows-Wheeler Aligner	3
		3.2.2 Bowtie2	3
		3.2.3 STAR	3
		3.2.4 Samtools	3
	3.3	Quality Check	3
		3.3.1 FastQC	3
		3.3.2 Sequenza	3
	3.4	Workflow	3
4	Resi	ults	3
	4.1	FastQC	3
5	Disc	eussion	3
,	D.£		1
6	Keie	erences	3
L	ist o	of Tables	
L	ist o	of Figures	
	1	Somatic short variant (SNVs + Indels) discovery workflow (Van der Auwera et al., 2013; DePristo et al.,	
		2011)	4
	2	FastQC with WES data	4
	3	FastOC with WTS data	4

1 Introduction

2 Materials

3 Methods

3.1 Genome Analysis Toolkit

Genome analysis toolkit (GATK) is a software package for variant discovery among sequencing data (Van der Auwera et al., 2013; DePristo et al., 2011).

3.2 Alignment

3.2.1 Burrows-Wheeler Aligner

Burrows-Wheeler Aligner (BWA) is a software package for aligning short-read sequences unto a large reference genome (Li & Durbin, 2009). BWA-MEM is one of the contained algorithms in BWA software package, is a novel algorithm for mapping sequence reads on a large reference genome (Li, 2013).

3.2.2 Bowtie2

Bowtie 2 is an efficient and fast software package for aligning sequencing reads against long reference sequences (Langmead & Salzberg, 2012).

3.2.3 STAR

STAR is a swift universal RNA-seq alignment tool (Dobin et al., 2013).

3.2.4 Samtools

Samtools is a suite software packages for discovering in high-throughput sequencing data (Li et al., 2009).

3.3 Quality Check

3.3.1 FastQC

FastQC is a software package which aims to provide a productive method to do quality control check on raw sequence data (Andrews et al., 2012).

3.3.2 Sequenza

Sequenza is a software package to investigate genomic sequencing data, such as cellularity and ploidy estimation, from paired normal-tumor samples (Favero et al., 2015).

3.4 Workflow

- 4 Results
- 4.1 FastOC
- 5 Discussion

6 References

- Andrews, S., Krueger, F., Segonds-Pichon, A., Biggins, L., Krueger, C., & Wingett, S. (2012, January). *FastQC*. Babraham Institute. Babraham, UK.
- DePristo, M. A., Banks, E., Poplin, R., Garimella, K. V., Maguire, J. R., Hartl, C., ... others (2011). A framework for variation discovery and genotyping using next-generation dna sequencing data. *Nature genetics*, 43(5), 491.
- Dobin, A., Davis, C. A., Schlesinger, F., Drenkow, J., Zaleski, C., Jha, S., ... Gingeras, T. R. (2013). Star: ultrafast universal rna-seq aligner. *Bioinformatics*, 29(1), 15–21.
- Favero, F., Joshi, T., Marquard, A. M., Birkbak, N. J., Krzystanek, M., Li, Q., ... Eklund, A. C. (2015). Sequenza: allele-specific copy number and mutation profiles from tumor sequencing data. *Annals of Oncology*, 26(1), 64–70.

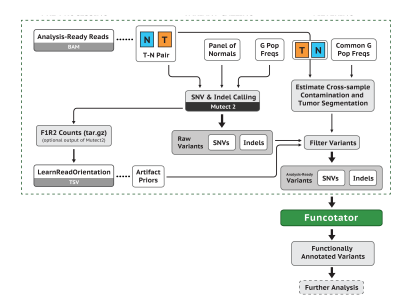


Figure 1: Somatic short variant (SNVs + Indels) discovery workflow (Van der Auwera et al., 2013; DePristo et al., 2011)

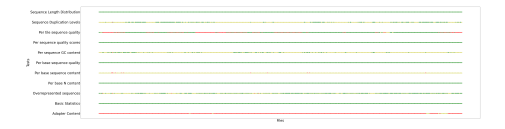


Figure 2: FastQC with WES data



Figure 3: FastQC with WTS data

- Langmead, B., & Salzberg, S. L. (2012). Fast gapped-read alignment with bowtie 2. Nature methods, 9(4), 357.
- Li, H. (2013). Aligning sequence reads, clone sequences and assembly contigs with bwa-mem. *arXiv preprint* arXiv:1303.3997.
- Li, H., & Durbin, R. (2009). Fast and accurate short read alignment with burrows—wheeler transform. *bioinformatics*, 25(14), 1754–1760.
- Li, H., Handsaker, B., Wysoker, A., Fennell, T., Ruan, J., Homer, N., ... Durbin, R. (2009). The sequence alignment/map format and samtools. *Bioinformatics*, 25(16), 2078–2079.
- Van der Auwera, G. A., Carneiro, M. O., Hartl, C., Poplin, R., Del Angel, G., Levy-Moonshine, A., ... others (2013). From fastq data to high-confidence variant calls: the genome analysis toolkit best practices pipeline. *Current protocols in bioinformatics*, 43(1), 11–10.