

# **Genome Variant Analysis: Overview**

1. Types of genomic variation

2. Visualization using IGV

3. File Formats for Variation Data



# Genome Variant Analysis: Types of Genomic Variation

## Variant or Mutation or Alteration or Polymorphism

- Changes in the genome sequence of a sample compared to a reference sequence
- Chromosomes: 22 autosomal pairs + 1 sex pair
  - Each set inherited from maternal and paternal germline cells

## **Germline Variant**

- Variant inherited from one or both parental chromosomes
- Source of genetic differences between ancestral populations and individuals
- Polymorphism: >1% frequency in a population

## **Somatic Variant**

- Mutation acquired during individual's lifetime
- Important to identify in sporadic cancers and other non-familial diseases



# Genome Variant Analysis: Types of Genomic Variation

## a. Single nucleotide base substitutions

- Germline single nucleotide polymorphism (SNP)
- Somatic single nucleotide variant (SNV)

## b. Small insertions or deletions

Germline or somatic insertion or deletion (INDEL)

## c. Copy number changes

- Germline copy number variant (CNV) or polymorphism (CNP)
- Somatic copy number variant (CNV) or alterations (CNA)

## d. Structural rearrangements

Germline or Somatic structural variant (SV)



# Genome Variant Analysis: Single Nucleotide Polymorphism

- ~1.5 to 2 million SNPs per individual
- Identify SNPs from normal peripheral blood mononuclear cells (PBMC)



Heterozygous SNP with 37 reads containing the variant and having depth 79 reads

37/79 (47%) variant allele fraction (VAF)

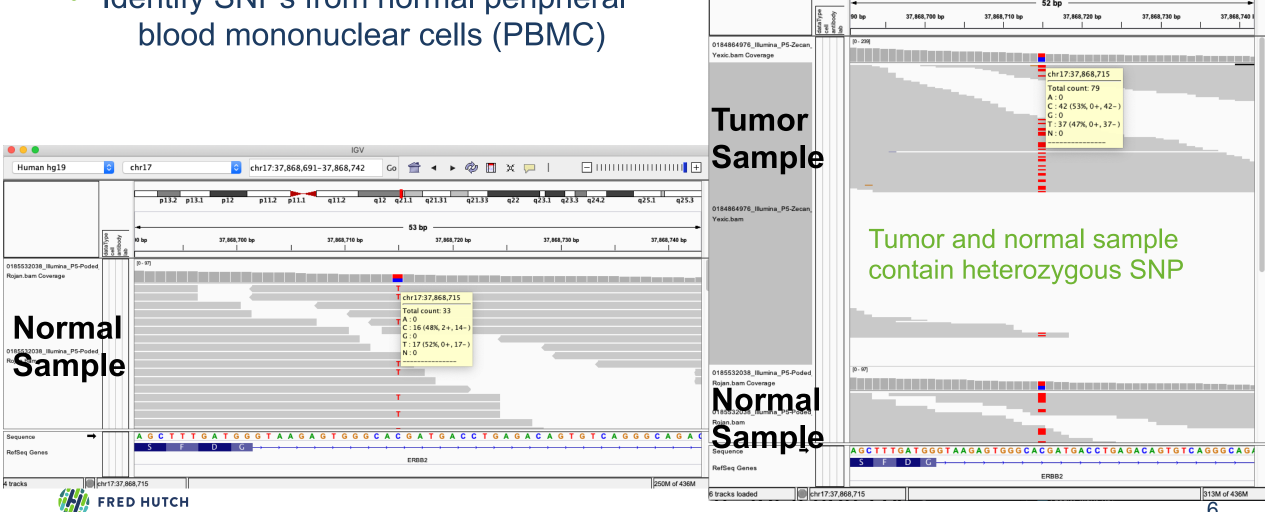
# Genome Variant Analysis: Single Nucleotide Polymorphism

Human hg19

chr17

chr17:37,868,691-37,868,742

- ~1.5 to 2 million SNPs per individual
- Identify SNPs from normal peripheral blood mononuclear cells (PBMC)



# Genome Variant Analysis: Single Nucleotide Variant (SNV)

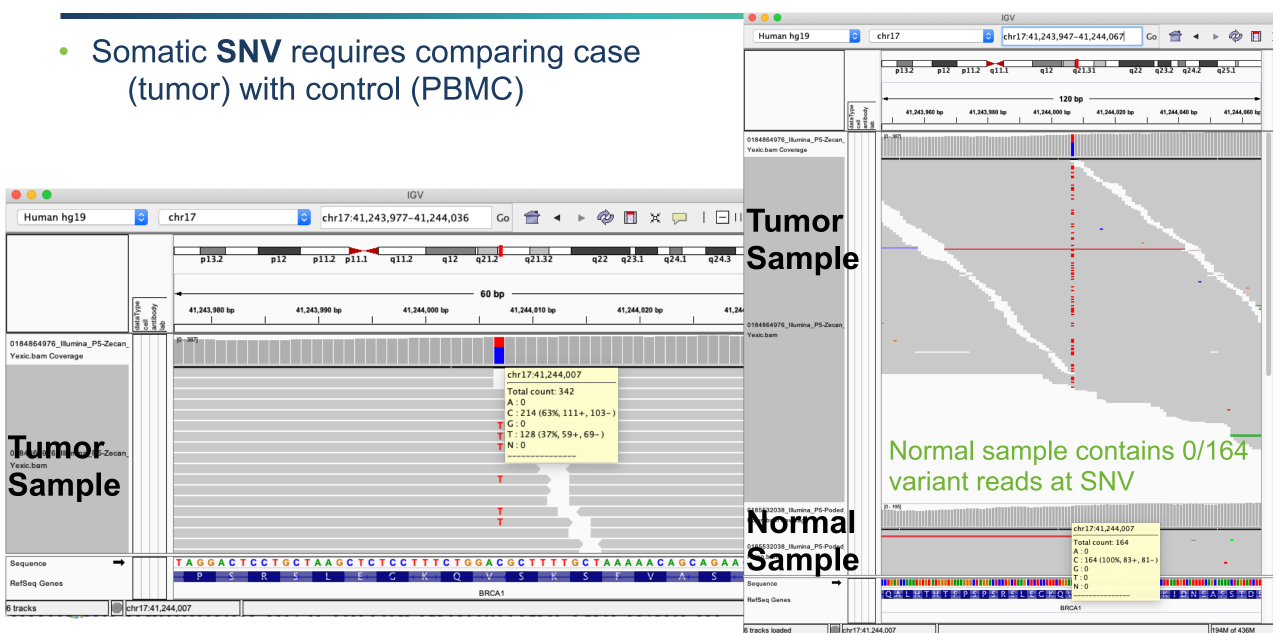
 Somatic SNV requires comparing case (tumor) with control (PBMC)



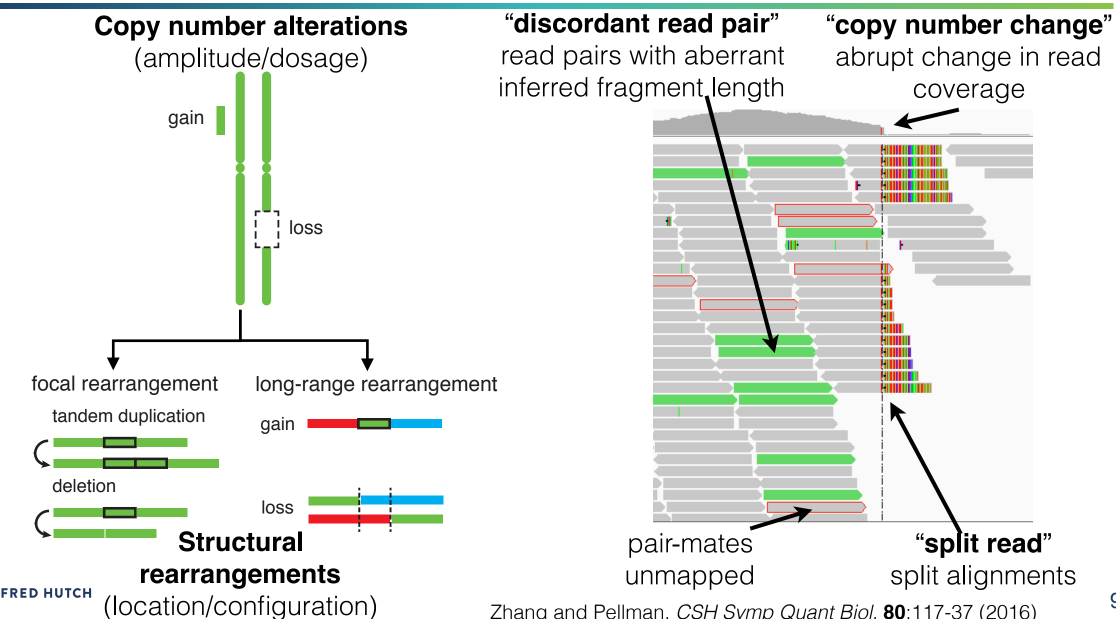
Potential SNV with 128/342 (37%) VAF

p.V1181I

# Genome Variant Analysis: Single Nucleotide Variant (SNV)



# Genome Variant Analysis: Copy Number and Structural Variation



Zhang and Pellman. CSH Symp Quant Biol. 80:117-37 (2016)

# **Genome Variant Analysis: Copy Number Variation**





# Genome Variant Analysis: Common Variant File Formats

## a. Variant Call Format (VCF)

- http://samtools.github.io/hts-specs/VCFv4.2.pdf
- Used mostly for SNV/SNP, INDEL, and SV

## b. Mutation Annotation Format (MAF)

- https://docs.gdc.cancer.gov/Data/File\_Formats/MAF\_Format/
- http://software.broadinstitute.org/software/igv/MutationData
- Tab-delimited format containing columns for mutation information and annotations
- Used primarily for SNV/SNP and INDEL data

## c. Browser Embedded Data (BED)

- a. https://bedtools.readthedocs.io/
- b. Used for any genomic features/region and annotations, including CNV and SV (BEDPE)

#### d. Others

- a. <a href="http://genome.ucsc.edu/FAQ/FAQformat">http://genome.ucsc.edu/FAQ/FAQformat</a>
- b. GFF, WIG/bigWIG, etc.



# Genome Variant Analysis: Variant Call Format (VCF)

## http://samtools.github.io/hts-specs/VCFv4.2.pdf

#### a. Header information

```
##fileformat=VCFv4.2
##GATKCommandLine=<ID=HaplotypeCaller,CommandLine="HaplotypeCaller">
##INFO=<ID=AC,Number=A,Type=Integer,Description="Allele count in genotypes, for each ALT allele">
##INFO=<ID=AF,Number=A,Type=Float,Description="Allele Frequency, for each ALT allele, in the same order as listed">
##INFO=<ID=AN,Number=1,Type=Integer,Description="Total number of alleles in called genotypes">
##INFO=<ID=DP,Number=1,Type=Integer,Description="Approximate read depth; some reads may have been filtered">
##FORMAT=<ID=AD,Number=R,Type=Integer,Description="Allelic depths for the ref and alt alleles in the order listed">
##FORMAT=<ID=DP,Number=1,Type=Integer,Description="Approximate read depth">
##FORMAT=<ID=DP,Number=1,Type=Integer,Description="Genotype Quality">
##FORMAT=<ID=GQ,Number=1,Type=Integer,Description="Genotype">
##FORMAT=<ID=CT,Number=1,Type=Integer,Description="Normalized, Phred-scaled likelihoods for genotypes as defined in the VCF specification">
##FORMAT=<ID=PS,Number=1,Type=Integer,Description="ID of Phase Set for Variant">
##FILTER=<ID=PASS,Description="All filters passed">
##FILTER=<ID=DLowQual,Description="Low quality">
##FILTER=<ID=LowQual,Description="Low quality">
##FILTER=<ID=LowQual,Description="Low quality">
##FILTER=<ID=LowQual,Description="Low quality">
##FILTER=<ID=DLowQual,Description="Low quality">
##FILTER
```

#### b. Variant record

#CHROM	POS	ID	REF	ALT	QUAL	FILTER	INFO	FORMAT	Sample_1
chr1	11542		A	Т	49.77	PASS	AC=1; AF=0.5; AN=2; DP=4	GT:AD:DP:GQ:PL:PS	0 1:2,2:4:78:78,0,78



# Genomic Data Analysis in R

# Lecture 16

Tuesday, November 124, 2020 @ 1pm

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## **Overview**

## I. R and Bioconductor Packages for Genomic Data

## **II. Tutorials**

- Analyzing Genomic Data
- Analyzing and Annotating Variants
- Analyzing Sequence Data



# **Overview: Learning Objectives**

- I. R Bioconductor Packages for Genomic Data
  - GenomicRanges, Rsamtools, VariantAnnotation
- II. Tutorials
  - 1. Genomic Data Analysis (GenomicRanges)
    - i. Load, inspect, query a BED/SEG file
    - ii. Genomic regions overlap
  - 2. Sequence Data Analysis (Rsamtools)
    - i. Load, inspect, query a BAM alignment file
    - ii. Extract sequences and qualities
    - iii.Compute "pile-up" statistics at genomic loci
  - 3. Genomic Variants and Annotations (VariantAnnotation)
    - i. Load, inspect, query a VCF file



# **Tutorial #1: Genomic Data Analysis**

- 1. Loading and querying BED/SEG text files
  - a. Use packages GenomicRanges
- 2. Download the VCF and SEG files for this tutorial
  - https://www.dropbox.com/sh/zoitjnobgp7l7c2/AABBIpTQcNA4lWYOFnV5dlMKa?dl=0
  - BRCA.genome\_wide\_snp\_6\_broad\_Level\_3\_scna.seg
- 3. R Markdown file for tutorial on GitHub: Lecture16 GenomicData.Rmd



# **Tutorial #2: Sequence Data Analysis**

- 1. Loading and querying a BAM file using Rsamtools
  - a. Define the genomic coordinates and components to query (ScanBamParam)
  - b. Scanning the BAM file (scanBam)
- 2. We will use the example from Lecture 15: Slides 19-22.
- 3. Download the BAM file for this tutorial
  - https://www.dropbox.com/sh/zoitjnobgp7l7c2/AABBIpTQcNA4lWYOFnV5dlMKa?dl=0
  - BRCA\_IDC\_cfDNA.bam
  - BRCA IDC cfDNA.bam.bai
- 4. R Markdown file for tutorial on GitHub: Lecture16 Rsamtools.Rmd



# **Tutorial #3: Variant Call Format (VCF)**

- 1. Loading and querying VCF files in R
  - a. Use packages VariantAnnotation
  - b. Download the VCF files for this tutorial
  - https://www.dropbox.com/sh/zoitjnobgp7l7c2/AABBIpTQcNA4lWYOFnV5dlMKa?dl=0
  - GIAB\_highconf\_v.3.3.2.vcf.gz
  - GIAB\_highconf\_v.3.3.2.vcf.gz.tbi
- 2. R Markdown file for tutorial on GitHub: Lecture16 VariantCalls.Rmd



# Homework #7: Genomic Data Analysis in R

#### Problem Set in R Markdown file

- Contains 4 problems with some code to prepare you for the questions.
- Please complete the assignment within the markdown file
- You will be evaluated on
  - i. the results and outputs
  - ii. your code and documentation
  - iii. Partial points awarded for code with correct logic/function even if the final answer may be incorrect

