Visualization of mapped reads

Integrative Genomics Viewer (IGV)

University of Cambridge

Cambridge, UK 30th September 2014



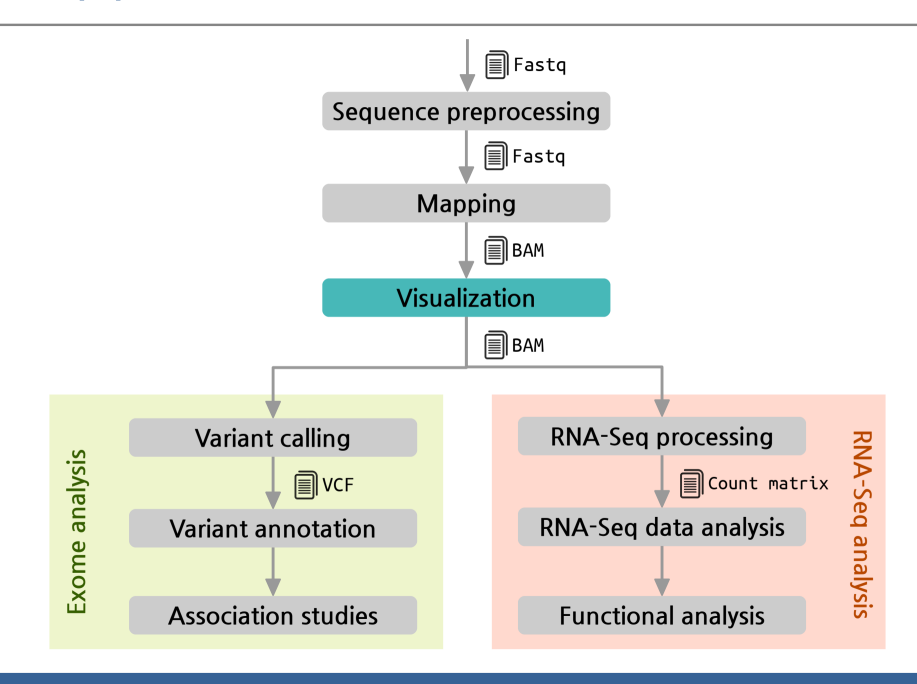


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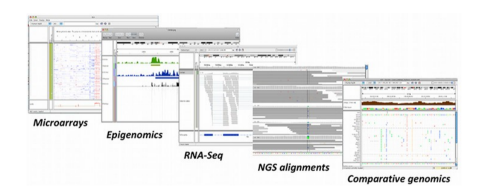
The pipeline



Introduction

- Large quantities of genomic data (NGS, array based methods, etc)
- Human interpretation and judgment using visualization can help complex biological relationships
- Integrative Genomics Viewer (IGV)
 - Integrate different data types simultaneously
 - View large datasets easily
 - Faster navigation or browsing
 - Runs **locally** on your desktop
 - Used by large-scale projects
 - Open source and freely available











Helga Thorvaldsdóttir, James T. Robinson, and Jill P. Mesirov Integrative Genomics Viewer (IGV): high-performance genomics data visualization and exploration
Brief Bioinform (2013) 14 (2): 178-192

Data types

- Any data related to genome coordinates
- Sample annotations or attributes
- Genome annotations

Recommended file formats

| Source data | Recommended File Formats |
|-------------------------|--|
| Sequence alignment data | SAM (must be sorted/indexed) BAM (must be indexed) |
| Genome annotations | GFF or GFF3 format BED format |
| Variant data | VCF |
| Any numeric data | IGV format, TAB format WIG format |
| Gene expression data | GCT format RES format |

Indexing a BAM file

- BAM format: Binary SAM file → Reduces disk space and time
- BAM/SAM files need to be indexed (using samtools) → SAM files will be sorted by start position and indexed
- Index files must reside in the same directory as the BAM or SAM file

Index the example BAM file

samtools index igv1.bam

Registration and download

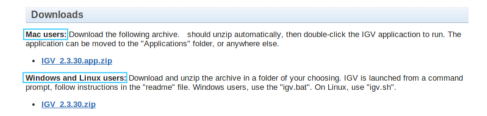
- 1. Be sure that Java 6 or later is installed on your machine
- 2 Go to the IGV website:

http://www.broadinstitute.org/igv/home

- 3. Click **Downloads** at the left panel
- 4. Click to register and fill the form



5. Download the most suitable file for your system

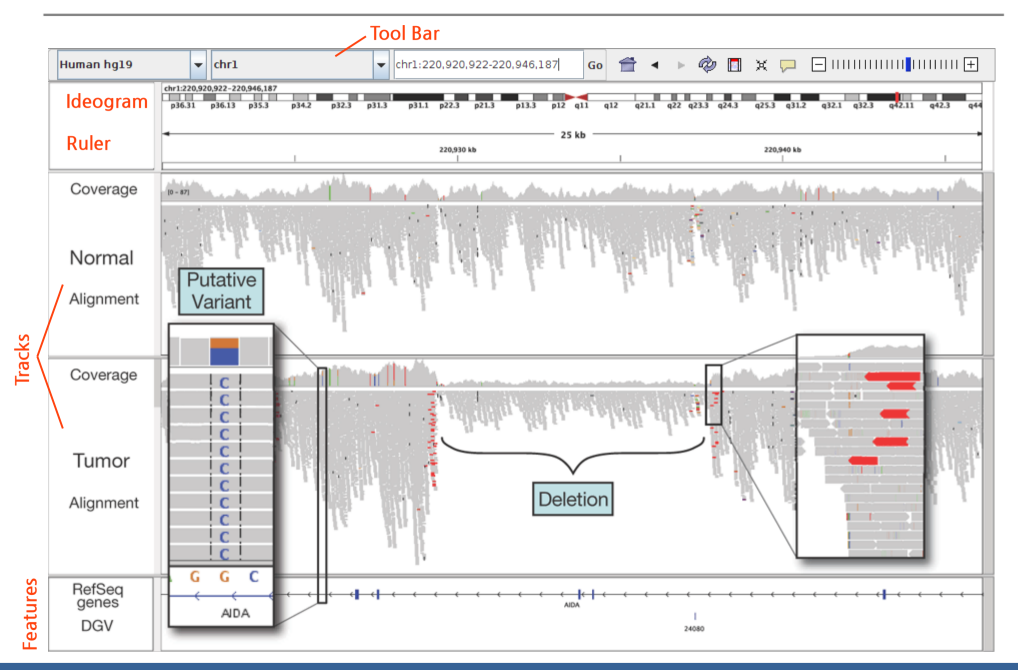


6. Run IGV

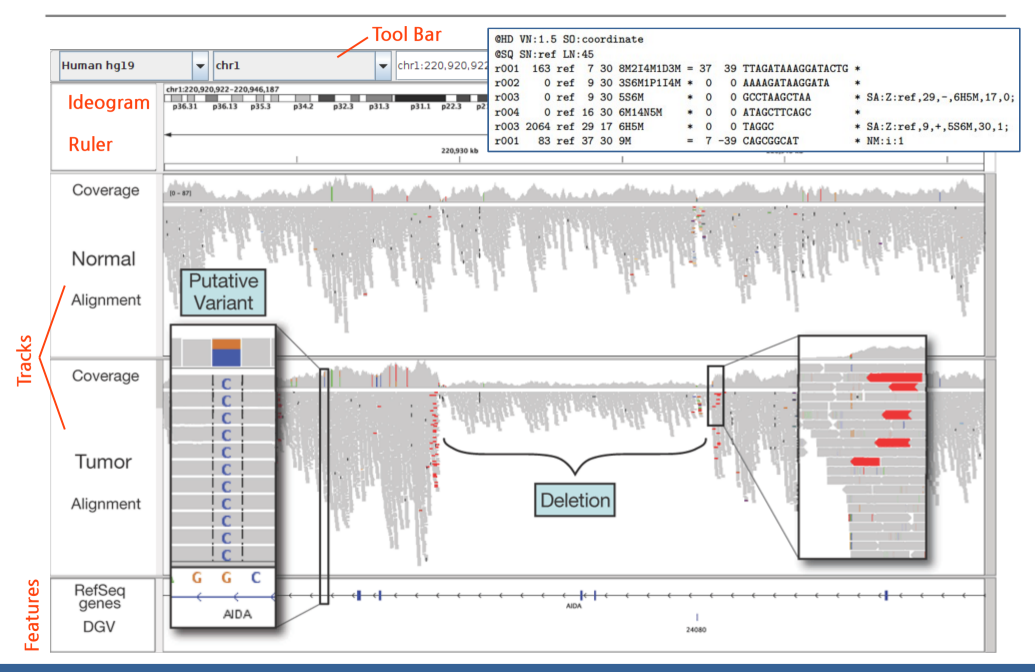
igv



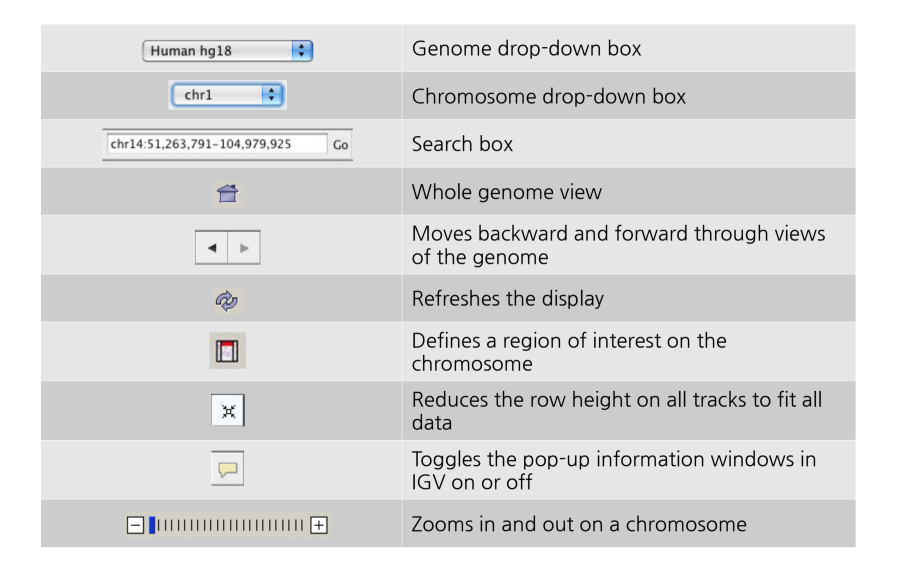
IGV interface



IGV interface



IGV Tool Bar



Download genome

Current version of IGV has "Human hg18" and "Human hg19" already loaded IGV provides a number of genomes that are hosted on a server at the Broad Institute List of genomes hosted: http://www.broadinstitute.org/software/igv/Genomes

Genomes → Load genome from server...

Select Human hg19

