

Course Presentation









Presentation

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Introduction Who we are

- Teachers:
 - David Montaner: Head of the Biostatistics Unit at CIPF (Valencia, Spain)
 - Marta Bleda: Computational Biologist and Data Analyst at Department of Medicine, Addenbrooke's Hospital (Univ. Cambridge)
 - F. Javier Lopez: Bioinformatician and Software Engineer at EMBL-EBI Variation (Cambridge, UK)
 - Ignacio Medina: Head of Computational Biology Lab, HPCS University of Cambridge, UK. EMBL-EBI Scientific Collaborator at EBI Variation team (Cambridge, UK)
 - Joaquin Dopazo: Head of the Computational Genomics Department at CIPF (Valencia, Spain)
- Everything started at Joaquin Dopazo's group at CIPF:
 - http://bioinfo.cipf.es/
- More than 8 years of experience in microarrays and NGS data analysis, and also developing metholodogies and bioinformatics tool for data analysis. Many suites and tools developed: GEPAS, Babelomics, Genome Maps, BierApp, VARIANT, ...
- More than 60 papers in the last 8 years in peer reviewed journals: NAR, Bioinformatics, Nat. Biotech., ...
- · Many collaborations with experimental and clinic groups
- Many international courses run last years: Massive Data Analysis (MDA)

Introduction Goals, ambitious

- To learn the basics to understand and be able to conduct a standard NGS data analysis from scratch in a Linux environment
- To know and understand the different analysis pipelines and data formats (FASTQ, SAM/BAM, VCF)
- To preprocess and perform QC of raw and processed data
- To learn and use the most widely used tools to perform NGS data analysis and visualization
- To learn the basics of the functional interpretation of variant (DNA resequencing) and RNA-seq analysis
- Optionally, learn how to install NGS software in Linux and how to tune up data analysis pipelines by simulating data

ProgramFirst day

- 09:30 Introduction to NGS Technologies
- 10:30 Course presentation and introduction to NGS data analysis & GNU/Linux shell
- 11:00 Coffee Break
- 12:00 FastQ Quality Control for NGS Raw Data (theory)
- 12:45 Lunch Break
- 14:00 FastQ Quality Control for NGS Raw Data (hands-on)
- 14:45 Mapping NGS Reads for Genomic and Transcriptomics Studies (theory)
- 15:30 Tea Break
- 15:45 Mapping NGS Reads for Genomic and Transcriptomics Studies I (hands-on)
- 17:00 Finish
- 17:00 Optional: A more advanced Linux session. NGS software Installation (1h)

Presentation

ProgramSecond day

- 09:30 Mapping NGS Reads for Genomic and Transcriptomics Studies II (hands-on)
- 10:30 Visualization of NGS data (theory)
- 11:00 Coffee Break
- 11:15 Visualization of NGS data (hand-on)
- 11:45 Variant Calling (SNPs & INDELs) and Variant Visualization (VCF) I
- 12:30 Lunch Break
- 14:00 Variant Calling (SNPs & INDELs) and Variant Visualization (VCF) II
- 14:45 Variant Annotation (theory)
- 15:30 Tea Break
- 15:45 Variant Annotation (hands-on)
- 16:15 Variant prioritization
- 17:00 Finish

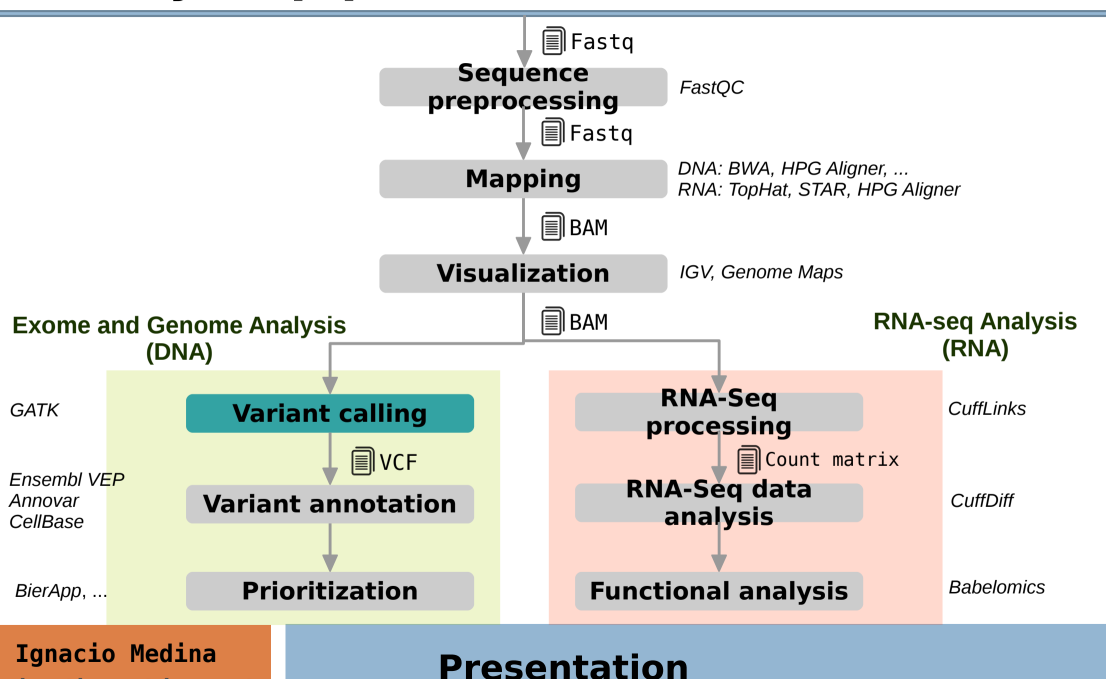
ProgramThird day

- 09:30 RNA-seq data preprocessing
- 11:00 Coffee Break
- 11:15 RNA-Seq Quantification and Isoforms Finding
- 12:30 Lunch Break
- 14:00 Functional Analysis
- 15:00 Tea Break
- 15:15 Exercises and questions
- 17:00 Finish

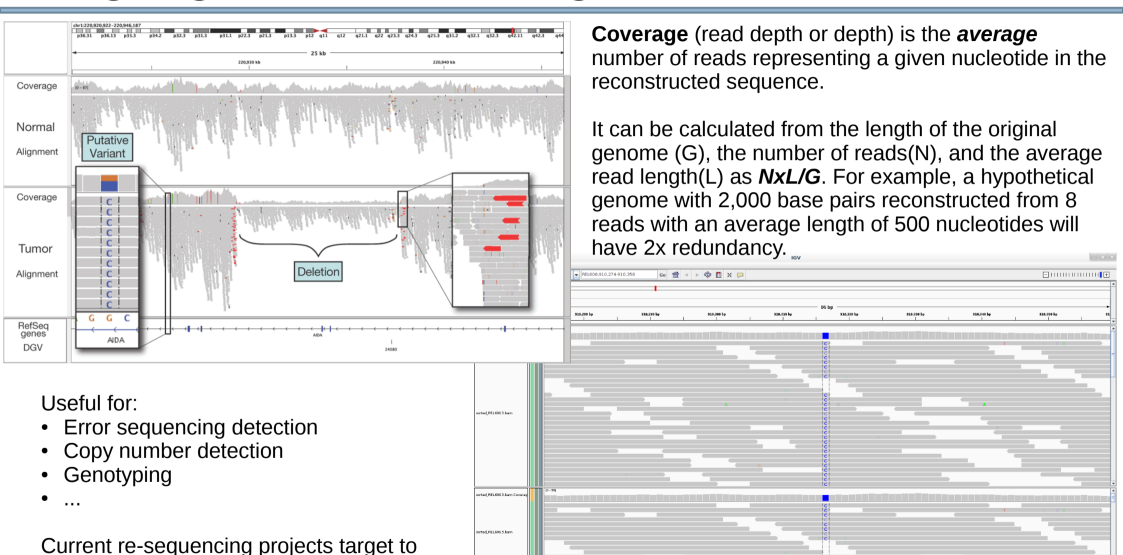
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Analysis pipeline

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Analysis pipeline Aligning reads, the coverage



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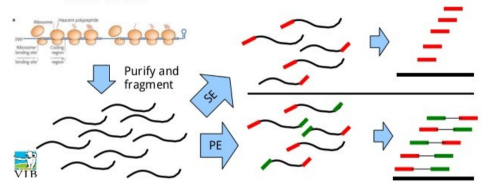
40x depth

Mapping NGS reads for genomic studies

Analysis pipeline paired-end vs single-end alignment

PE versus SE Illumina

- Single end (SE): from each cDNA fragment only one end is read.
- Paired end (PE): the cDNA fragment is read from both ends.



Paired-end sequencing:

- Improves read alignment and therefore variant calling
- Helps to detect structural variation
- Can detect gene fusions and splice junctions
- Useful for de novo assembly
- ...

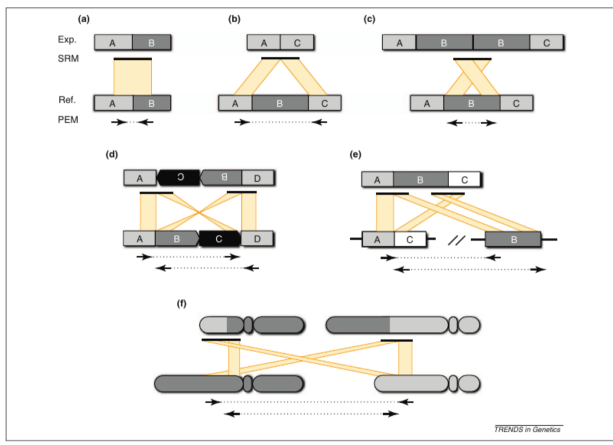


Figure 2. Detecting canonical structural variation (SV) breakpoints through sequencing. When DNA sequences are collected from an experimental (Exp.) genome and aligned to a reference (Ref.) genome, each structural variant class generates a distinct alignment pattern. The patterns observed for paired-end mapping (PEM) and split-read mapping (SRM) are illustrated when both genomes have identical structure (a), and cases where the experimental genome contains a deletion (b), a tandem duplication (c), an inversion (d), a transposon insertion (e) or a reciprocal translocation (f). PEM relies upon readpairs whose unsequenced portion (dotted lines) spans a SV breakpoint. When aligned to the reference genome, the alignment distance and orientation of such readpairs indicate the type of rearrangement that has occurred. Reads that map to the plus strand are shown as right-facing arrows, those that map to the negative strand as leftward-facing arrows. All examples depict Illumina paired-end sequence data, where in the absence of SV the normal concordant orientation is plus for the leftmost read and minus for the rightmost read. Note that the expected orientation is different for Illumina mate-pair libraries and for other sequencing platforms, such as SOLiD. In the case of a deletion (b), the readpairs ends will align much farther apart than expected for the DNA library. In contrast to PEM, SRM depends on contiguous sequences that contain an SV breakpoint. Consequently, the sequences before and after the breakpoint will align to disjoint regions of the reference genome. In contrast to PEM, breakpoints are identified at single-base resolution.

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Mapping NGS reads for genomic studies

Some considerations

- NGS data can be big, very big, huge! Biology is now a Big Data science
 - No too many web or graphical applications to perform analysis yet, sorry.
 - Most tools developed to work on **Linux**, many command line programs
- How to work in NGS?
 - Small datasets (<1TB): workstations
 - Medium sized datasets (<100-200TB): clusters
 - Big datasets (200TB-20PB): big clusters and cloud based solutions
- Exercises during this course will be done using with human **chromosome 21** to speed up analysis and not use too much memory. Under real circumstances using the whole genome the commands are exactly the same
- Software has been already installed to save time so you are not expected to download and install the software we are going to use. However it's usually needed to learn the basics of software installation in Linux, there is an optional session at the end of the first day for those that want to learn how to install NGS software in a standard Linux

What about you? Brief presentation

- Who are you?
- Which is your background?
- Which is your interest?
- What do you expect of this course?