JBrowse

- To get started early:
 - Double click VirtualBox on the desktop
 - Click "JBrowse 2016 Tutorial"
 - Click "Start"

JBrowse

PAG 2015

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What is GMOD?

- A set of interoperable open-source software components for visualizing, annotating, and managing biological data.
- An active community of developers and users asking diverse questions, and facing common challenges, with their biological data.

Who uses GMOD?















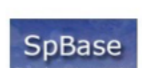












ConiferGDB















mips

































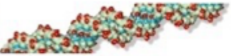


















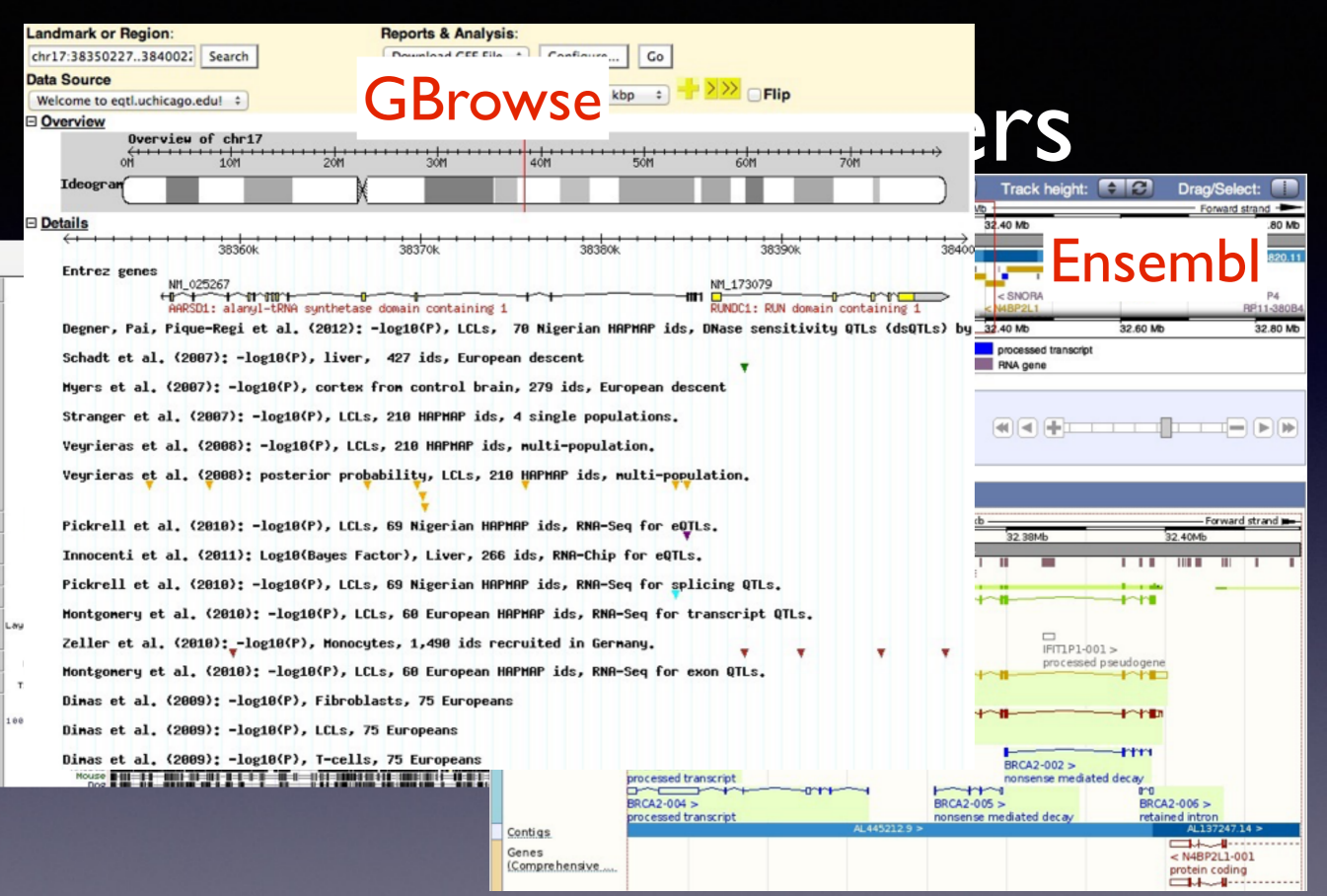






What are Genome Browsers good for?

- Visualizing dense data from a multitude of sources (genes from a GFF file, RNASeq data from a BAM file, variation data from a VCF file)
- Homology and gene expression support for gene models



 Many "specialty" browsers (eg, Biodalliance, Savant)

Why Install Your Own?

- You have data no one else has
- You want to be able to share it with your group, community, the world (potentially with "less savvy" users)
- You want to have control over how it looks

Why JBrowse?

- (Fairly) Easy install
- Good user experience (getting close to a browser-desktop hybrid)
- Good community support (mailing lists, tutorials, software updates)

Installation

- Only requires:
 - Web server (apache, lighttpd, nginx, etc)
 - Perl/make/standard unix-y tools

JBrowse Attributes

- Do everything possible on the client side, in JavaScript.
- Fast, smooth navigation.
- Supports GFF3, BED, Bio::DB::*, Chado, WIG, BAM, BigWig, VCF, and UCSC import (intron/exon structure, name lookups, quantitative plots).
- Is stably funded by NHGRI.
- Is open source, of course.
- Did I mention it's fast?

The JBrowse Project

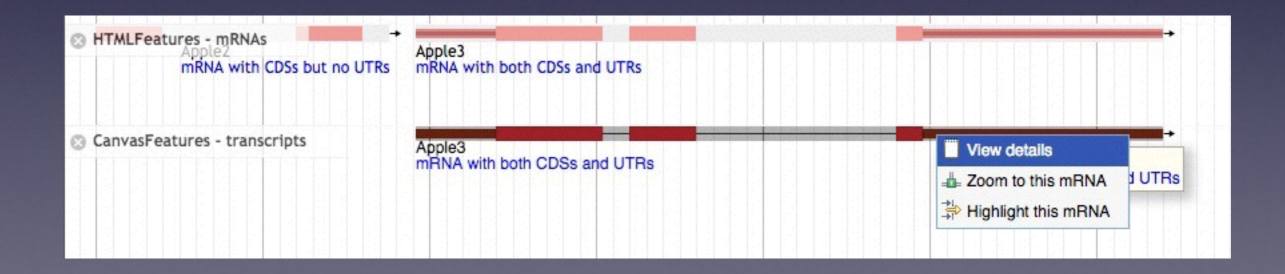
- free and open source (license: LGPL / Artistic)
- a GMOD project
 - http://gmod.org
- developed using git, hosted on GitHub
 - http://github.com/GMOD/jbrowse
- Pls most involved: Ian Holmes, Lincoln Stein, Suzi Lewis

New in 1.12.0

- Ability to start a new genome by uploading FASTA (indexed or not)
- Created stand-alone desktop version of JBrowse using the Electron platform for OSX, Windows, Gnome
- Several new plugins

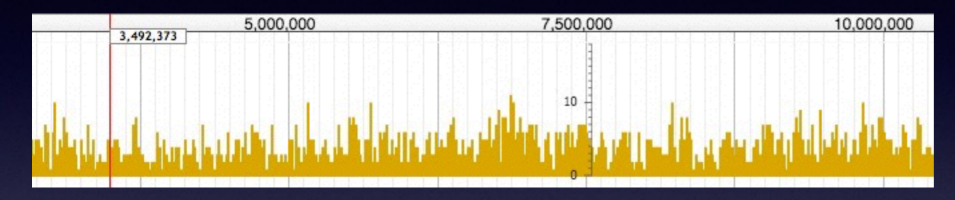
Feature Tracks

- HTMLFeatures: Rectangles (<div>s) with various fills and heights to represent the feature spans (more compatible)
- CanvasFeatures: Much prettier, more configurable glyphs
- Super-configurable left clicking and right-click

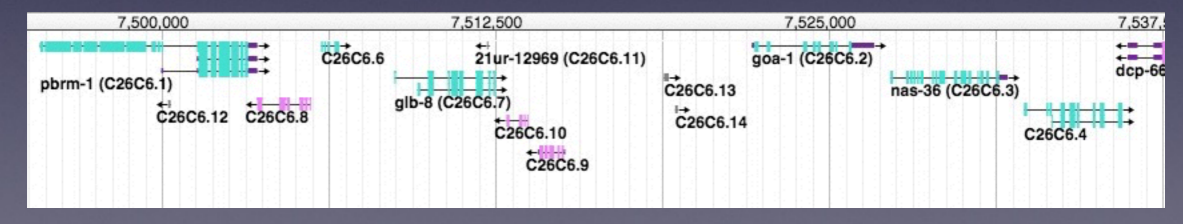


Feature Density Plots

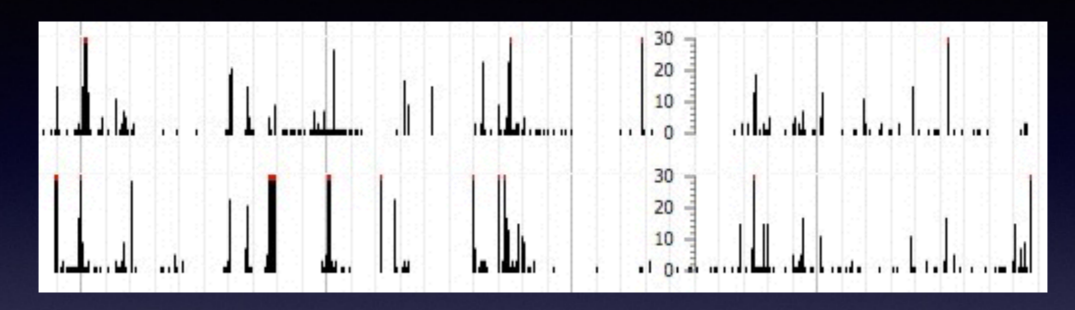
Zoomed out



Zoomed in

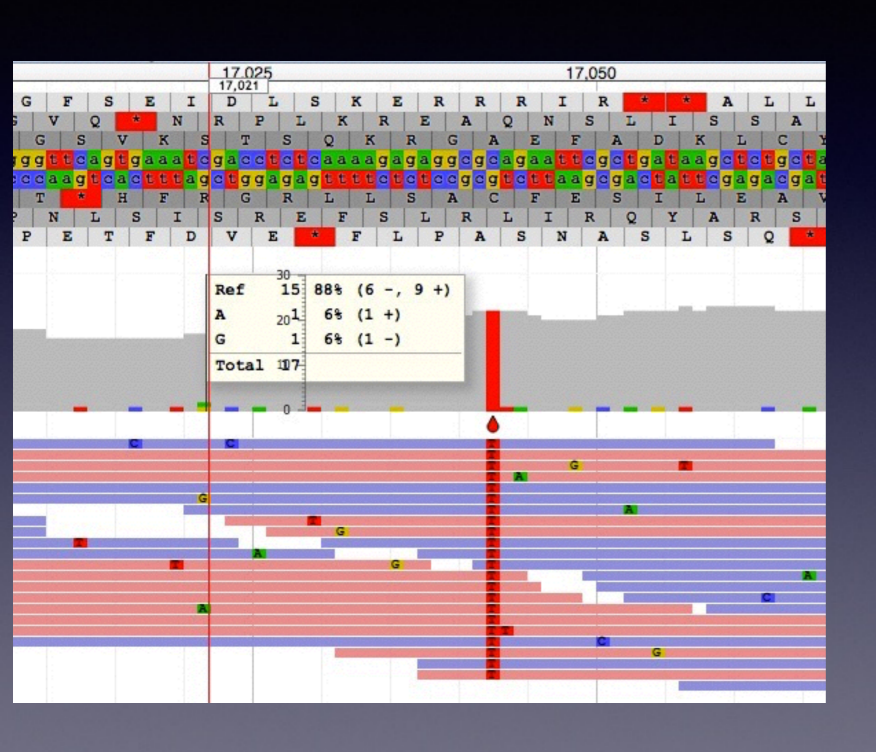


Wiggle/BigWig Tracks



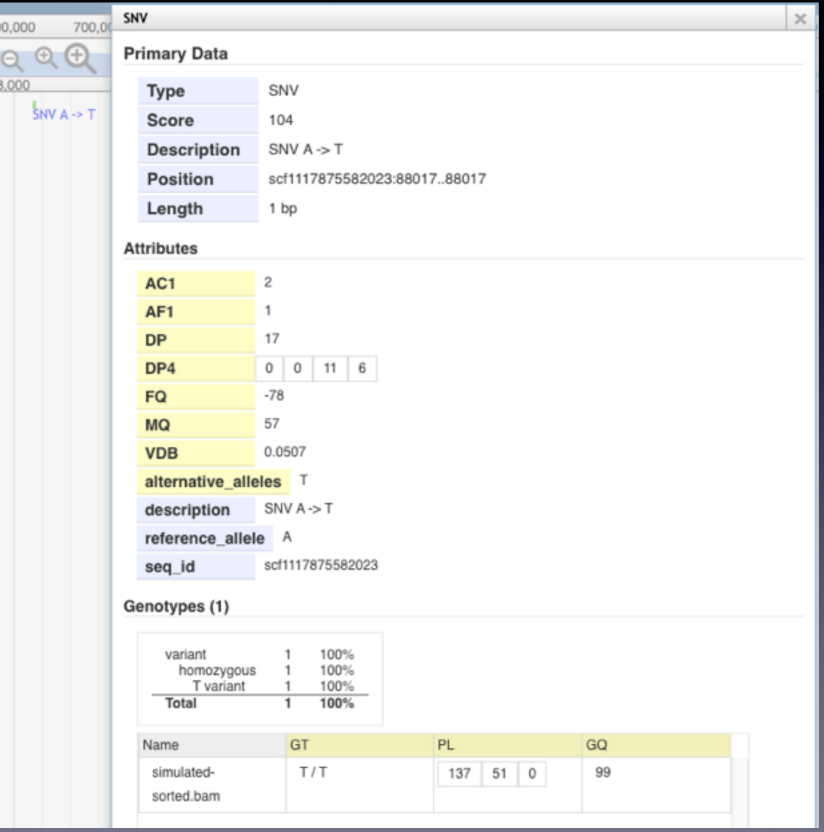
- Reads small chunks directly from BigWig file.
- Needs only a not-super-old (< 5 yrs) browser,
 except for Internet Explorer.
- IE requires version 10.

BAM Alignment Tracks



- Reads small chunks directly from BAM file.
- Coverage and mismatches.

VCFTracks



- Reads directly from VCF files compressed and indexed with bgzip and tabix.
- Shows all VCF data: alleles, genotypes, quality, etc.

Particular Strengths

- Web-based, but fast and smooth easy to set up
- Compressed NGS data: direct-to-browser BAM, BigWig, and VCF
- Optional faceted track selector efficiently search thousands of tracks
- Open local files directly on client, no data transfer required
- Highly customizable, embeddable, integratable, programmable

JBrowse as an Application

- Server-free JBrowse on your desktop
- Just provide a FASTA file and some annotations (GFF, BAM, BigWig, etc) and start browsing
- Download from http://jbrowse.org/ jbrowse-I-I2-0/

Apollo

(formerly WebApollo) ((and Apollo before that))

- Client side is a JBrowse plugin
- Clients receive updates in real time (like Google Docs)
- Extensive server-side Java
- http://genomearchitect.org/WebApolloDemo/
- Computer demo 3 (Tuesday 10:50 in this room)
- Poster 371,387

JBrowse Plugins

- Extend JBrowse with your own JavaScript code
- Can do pretty much anything
 - Add your own track visualizations
 - Add your own data backends
 - Add menu items
 - Subscribe to event notifications (pub/sub system)
 - Reach deep into the guts of JBrowse and (carefully!)
 - change anything at all!
- WebApollo client is a JBrowse plugin

Coming in 2.X

- MORE: data types, sorting options, speed
- Graphical configuration
- Multiple views, linked or independent
- Logins, uploading, track sharing
- Circular genomes

Big Thanks

Ian Holms (UC Berkeley)

Rob Buels Mitch Skinner Amelia Ireland Eric Yao

Lincoln Stein (OICR)
Julien Smith-Roberge
Erik Derohanian
Julie Moon
Natalie Fox
Adam Wright

Suzi Lewis (LBNL)
Gregg Helt
Ed Lee
Justin Reese (UofMo)
Colin Diesh (UofMo)

NHGRI Cold, hard cash

The End (on to the workshop)

http://jbrowse.org/

GMOD: http://gmod.org/wiki/JBrowse

Github: http://github.com/GMOD/jbrowse