

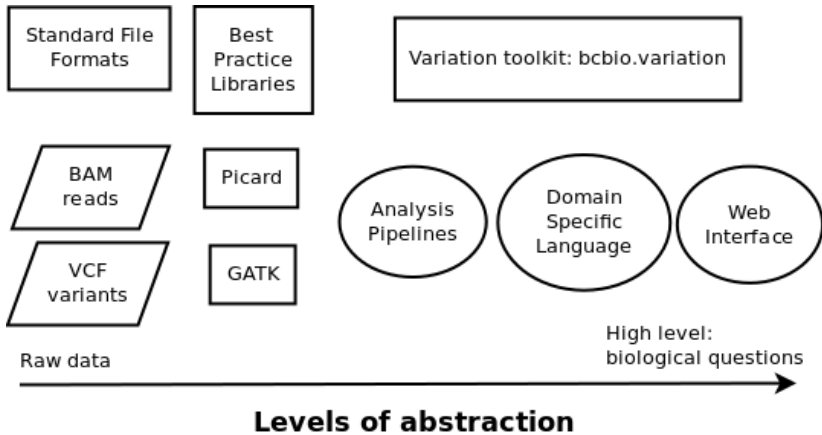
Toolkit for variation comparison and analysis

Brad Chapman, Bioinformatics Core at Harvard School of
Public Health

Bioinformatics Open Source Conference, 13 July 2012

Variation

```
1 1156131 rs2887286 T C 1714.07 PASS
AB=0;ABP=0;AC=1;AC1=2;AF=1.0;AF1=1;AN=1;AO=56;
BVAR;CIGAR=1X;DB;DP=59;DP4=0,0,27,26;DPRA=0;Dels=0.00;
EPP=6.88793;EPPR=0;FQ=-187;FS=0.000;GC=55.45;HRun=0;
HaplotypeScore=0.0000;LEN=1;MEANALT=2;MQ=70.00;MQ0=0;
MQM=70;MQMR=0;NBQ=27.76;NS=1;NUMALT=1;ODDS=308.76;
PAIRED=0.928571;QD=30.61;RO=0;RPP=12.937;RUN=1;
SAP=3.16541;SNPEFF_EFFECT=INTRON;SNPEFF_GENE_NAME=SDF4;
SNPEFF_FUNCTIONAL_CLASS=NONE;SNPEFF_IMPACT=MODIFIER;
SNPEFF_GENE_BIOTYPE=protein_coding;
SNPEFF_TRANSCRIPT_ID=ENST00000263741;
Samples=NA19239_illumina;TYPE=snp;VDB=0.0416;
VQSLOD=16.7453;XAI=0.00018797;XAM=0.00154116
GT:AO:DP:GL:GQ:QA:QR:RO
1:56:58:-141.12,-3.61:99:1527:0:0
```



Answer biological questions; help people

by ERIK CORONA • JULY 9, 2012

The goal of this blog is to catalog the process of finding the genetic cause of an inherited disease. Stacy was diagnosed with CMT4 (Charcot-Marie-Tooth Type 4) in the 1980s following a nerve biopsy which showed the characteristic onion-bulb myelin sheath around peripheral nerves and after measuring nerve conduction velocities along the length of her leg. There are many different causes of CMT, many known and many unknown. Stacy recently had her exome sequenced. My goal is to find the genetic basis of this disease. My name is Erik Corona and I am PhD student in the Stanford Medical school studying the genetics of complex disease. I will be making all results public and I will also post all code and/or scripts I use to generate these results. I will be posting screenshots of candidate structural variants and making every step along the way transparent and clear. This blog will represent a detailed case study of how we now have the technology and ability to use sequence data to find the genetic cause of a rare inherited disease. I welcome all comments, advice, and suggestions.

<http://cmtproject.blogspot.com/>

Comparisons

- Multiple callers: GATK, FreeBayes, samtools
- Multiple technologies: Illumina, SOLiD, IonTorrent

Identify real variants

- Summarize associated metrics
- Remove false positives

Scale

- Millions of variants
- Thousands of samples



<http://genomics.xprize.org/>

Clinical grade genome

- 98 percent genome coverage
- 1 error per million bases (SNPs + small indels)
- Full haplotype phasing
- Structural variations

Sequencing for patients

[My son Bertrand](#) has a new genetic disorder.

Patient 0.

To find it, a team of scientists at Duke University used whole-exome sequencing (a protein-focused variant of whole-genome sequencing) on me, my wife and my son.

We discovered that my son inherited two *different* (thus-far-unique) mutations in the same gene--the NGLY1 gene--which encodes the enzyme N-glycanase 1. Consequently, he cannot make this enzyme.

My son is the only human being known to lack this enzyme.

Below, I'm documenting our journey to the unlikeliest of diagnoses.

This is a story about the kind of hope that only science can provide.

<http://matt.might.net/articles/my-sons-killer/>

Technology overview

- Clojure

<http://clojure.org/>

- Genome Analysis Toolkit

[http://www.broadinstitute.org/gsa/wiki/index.php/
Home_Page](http://www.broadinstitute.org/gsa/wiki/index.php/Home_Page)

- GenomeSpace

<http://www.genomespace.org/>

<https://github.com/chapmanb/bcbio.variation>

Why Clojure?

- Hosted on Java Virtual Machine
 - Interoperability with existing libraries: GATK, GenomeSpace
 - Wonderful build, deployment and testing tools
- Functional and immutable
 - Easier to write correct code
 - Small functions: concise and refactorable
- Community
 - Smart people working on hard problems
- Ecosystem
 - Multiple backends: ClojureScript

Example analysis pipeline

- Variant files from two different callers
 - GATK UnifiedGenotyper
 - FreeBayes: <https://github.com/ekg/freebayes>
- Compare, identifying:
 - Identical variants
 - Different variants in each caller
 - Metrics to help discriminate

High level description

experiments:

- sample: Test1

ref: test/data/GRCh37.fa

intervals: test/data/target-regions.bed

align: test/data/aligned-reads.bam

calls:

- name: gatk

file: test/data/gatk-calls.vcf

- name: freebayes

file: test/data/freebayes-calls.vcf

prep: true

annotate: true

filters:

- QD < 2.0

- MQRankSum < -12.5

Simple to run

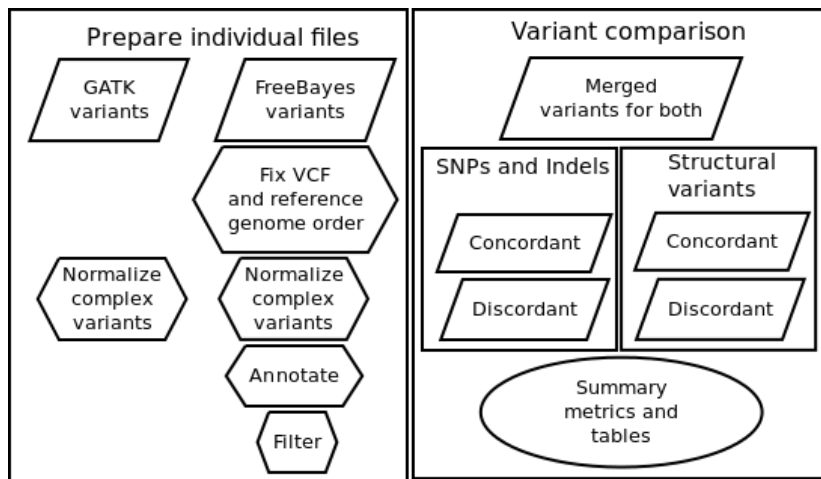
With a new blank machine, get Java and automated Clojure build tool:

```
$ sudo apt-get install openjdk-7-jdk openjdk-7-jre
$ wget https://raw.githubusercontent.com/technomancy/leiningen/\
    preview/bin/lein
$ chmod 755 lein && sudo mv lein /usr/local/bin
```

Get code and dependencies, then run:

```
$ git pull git://github.com/chapmanb/bcbio.variation.git
$ cd bcbio.variation
$ lein deps
$ lein variant-compare comparison_description.yaml
```

What happened?

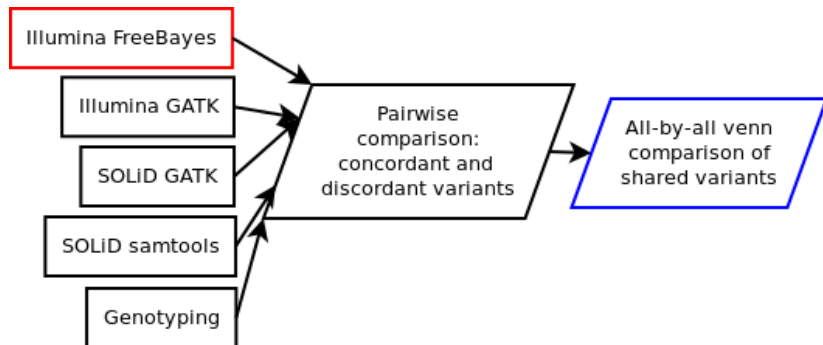


Establishing true variants

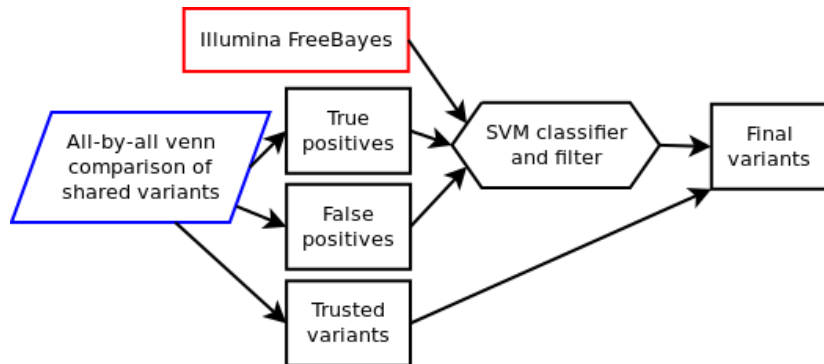
X Prize: haploid gold standard genome

- Public genomes from HapMap/1000 genomes
 - NA12878: Caucasian female from Utah.
 - NA19239: Yoruba male from Ibadan, Nigeria.
- Multiple technologies
 - Illumina
 - SOLiD
 - IonTorrent
- Multiple callers
 - GATK
 - FreeBayes
 - samtools mpileup

True variant workflow – comparisons



True variant workflow – finalize



Comparison web interface

X PRIZE scoring [Home](#) chapmanb

Submit variation file for scoring

Upload method Web upload GenomeSpace

Variations

Sequence differences, in **VCF format**, relative to the [GRCh37 reference genome \(FASTA download\)](#).
[Example file](#)

Scoring regions

Regions to assess for scoring, in **BED format**.
[Example file](#)

Comparison genome

Score

<https://validationprotocol.org/>

Summary web interface

X PRIZE scoring [Home](#)

 chapmanb ▾

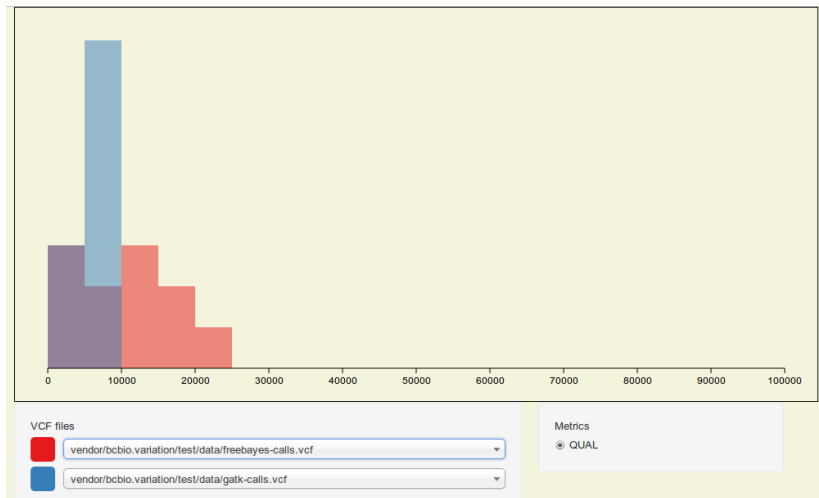
Summary

Metric	Value
Accuracy score	78.261
Accuracy score, including phasing	75.000
Completeness	94.74
Total bases scored	18
Possible evaluation bases	19
Discordant SNPs	3
Discordant indels	1
Discordant structural variants	0
Phasing Error SNPs	1
Phasing Error indels	0
Phased haplotype blocks	5

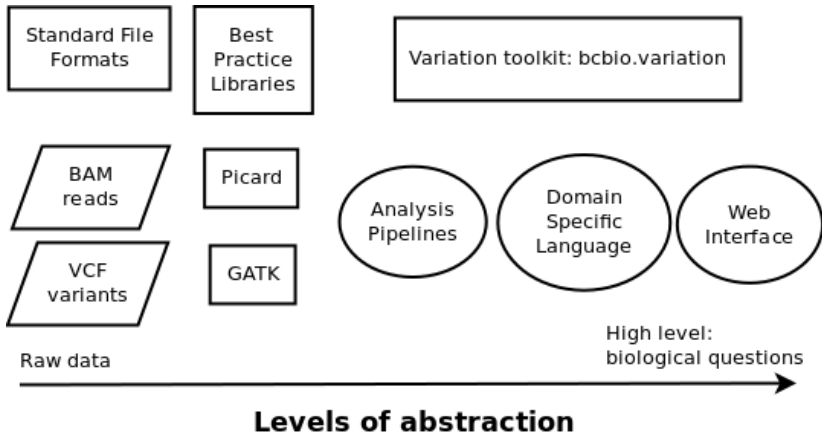
Variant files in VCF format

- [Concordant variants](#)
- [Discordant variants](#)
- [Missing variants](#)
- [Variants with phasing errors](#)

Next web steps – metrics



<http://keminglabs.com/>



Answering biological questions

- Establish set of true variants
 - Understand boundaries of certainty
 - Make patient decisions
- Comparison architecture
 - Cancer: tumor/normal pairs
 - Mendelian inherited diseases: father/mother/child
- Prioritize based on known data
 - Gemini: <https://github.com/arq5x/gemini>
 - 1000 genomes frequencies
 - Mappability
 - Clinical information