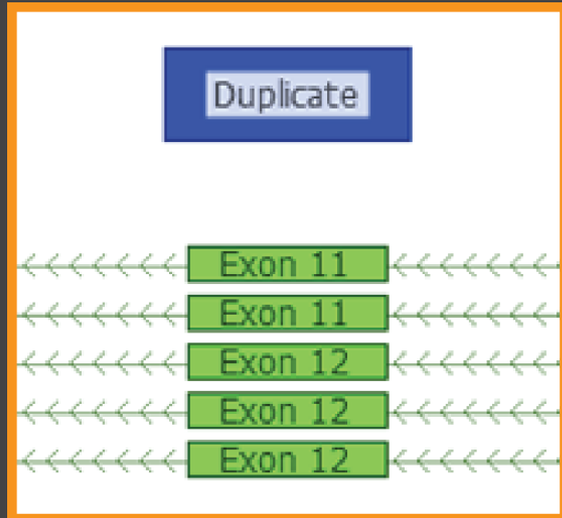


# CNV Analysis in VarSeq



December 7, 2016

Dr. Nathan Fortier  
Senior Software Engineer  
& Field Application Scientist



**1** Overview Golden Helix

**2** Why Call CNVs in NGS

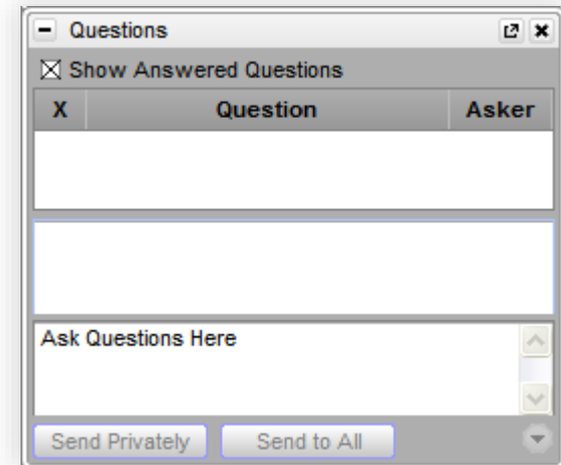
**3** Method and Demo

**4** Availability and Roadmap



# Questions during the presentation

Use the Questions pane in your GoToWebinar window



# Golden Helix – Who We Are



Golden Helix is a global bioinformatics company founded in 1998.



Filtering and Annotation  
Single Sample CNV-Analysis  
Clinical Reports  
Pipeline: Run Workflows

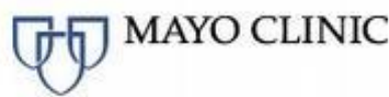


Variant Warehouse  
Centralized Annotations  
Hosted Reports  
Sharing and Integration

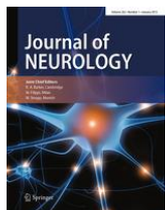
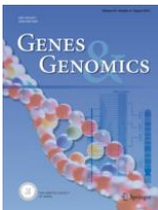
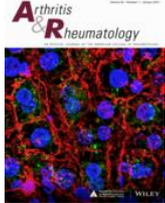
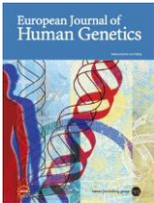
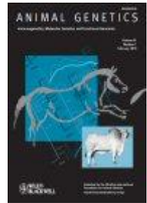
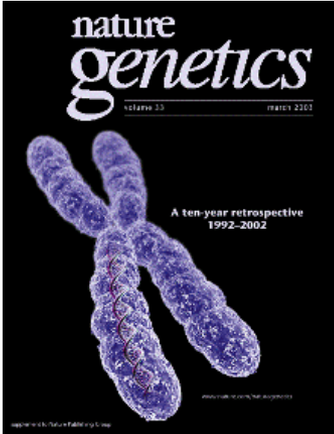
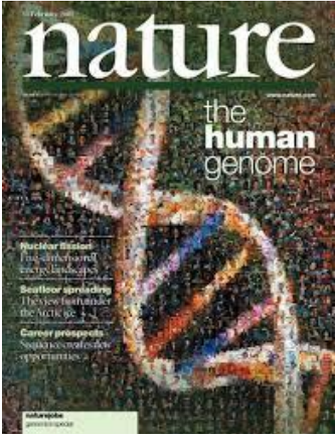
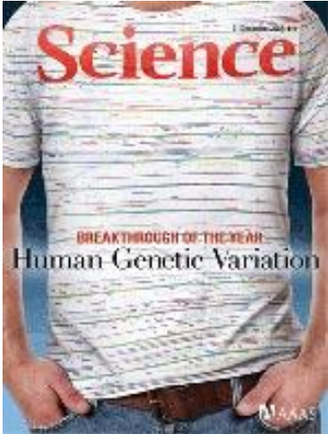


GWAS  
Genomic Prediction  
Large-N-Population Studies  
RNA-Seq  
Large-N CNV-Analysis

Over 300 customers globally



# Cited in over 1000 peer-reviewed publications



# Golden Helix – Who We Are



When you choose a Golden Helix solution, you get more than just software

- REPUTATION
- TRUST
- EXPERIENCE



- INDUSTRY FOCUS
- THOUGHT LEADERSHIP
- COMMUNITY

- TRAINING
- SUPPORT
- RESPONSIVENESS



- TRANSPARENCY
- INNOVATION and SPEED
- CUSTOMIZATIONS

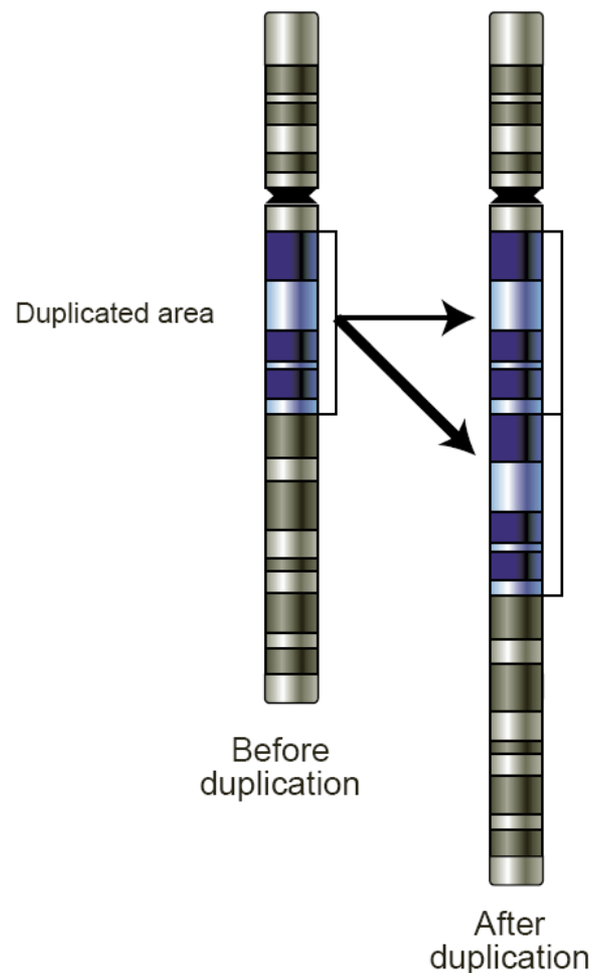


- **Chromosomal microarray**

- Current best practice
- Slow
- Additional expense
- Only detects large events

- **CNV calling from NGS data**

- Calls from existing coverage data
- Detects small single-exon events
- Provides faster results



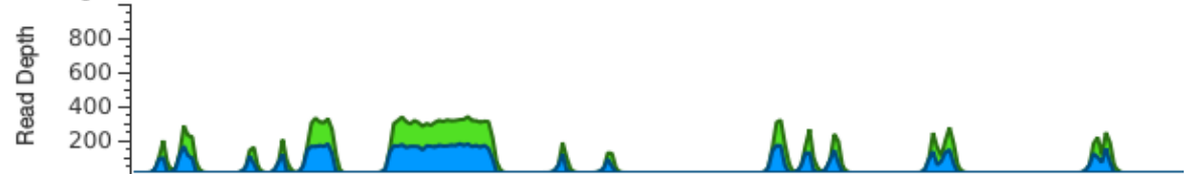
# CNV Detection via NGS



- **CNVs are called from coverage data**
- **Challenges**
  - Coverage varies between samples
  - Coverage fluctuates between targets
  - Systematic biases impact coverage
- **Solutions**
  - Data Normalization
  - Reference Sample Comparison

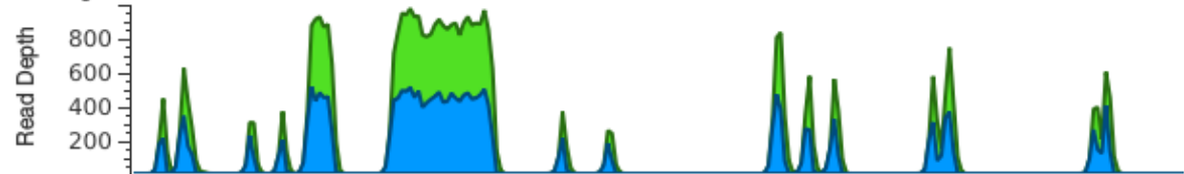
Current Sample: RD-NGSPROGENITYCANCER-SAMPLE11

Coverage



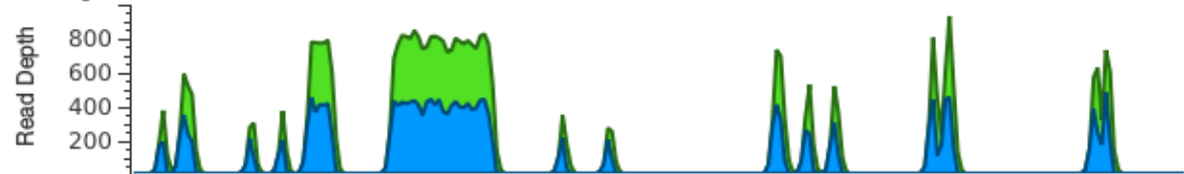
Current Sample: RD-NGSPROGENITYCANCER-SAMPLE12

Coverage

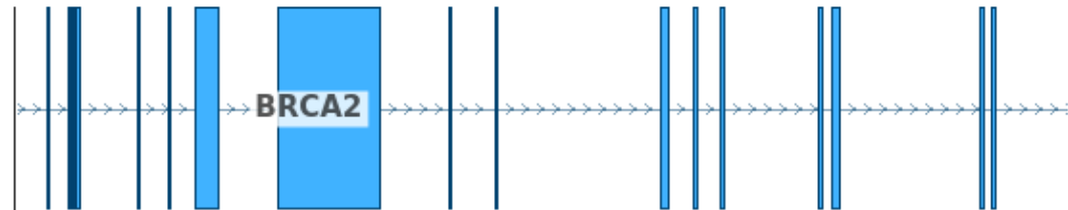


Current Sample: RD-NGSPROGENITYCANCER-SAMPLE13

Coverage

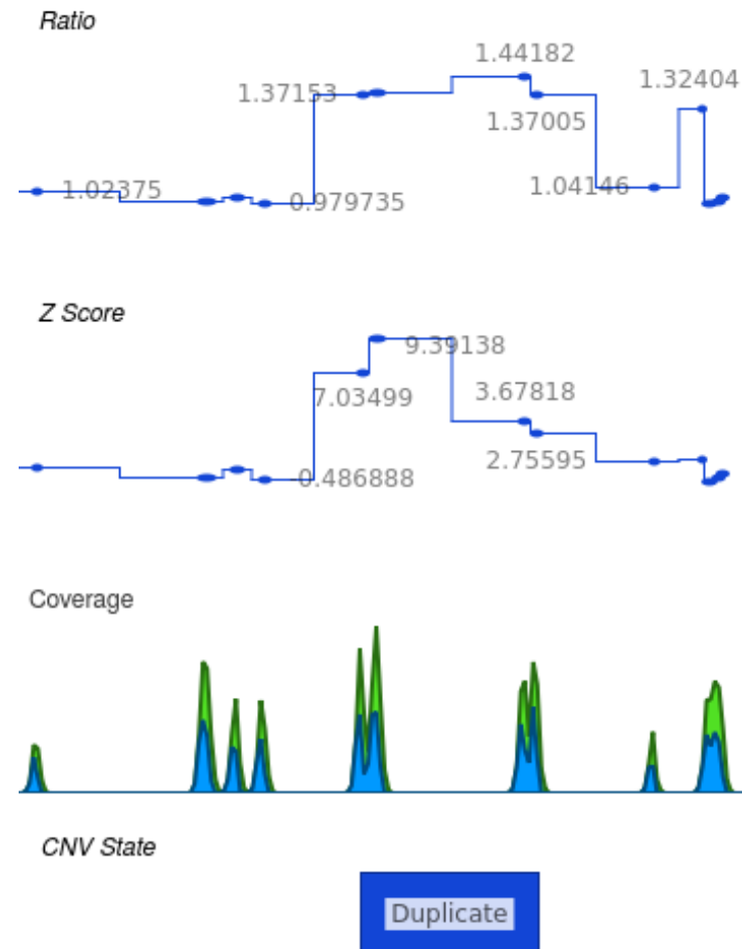


RefSeq Genes





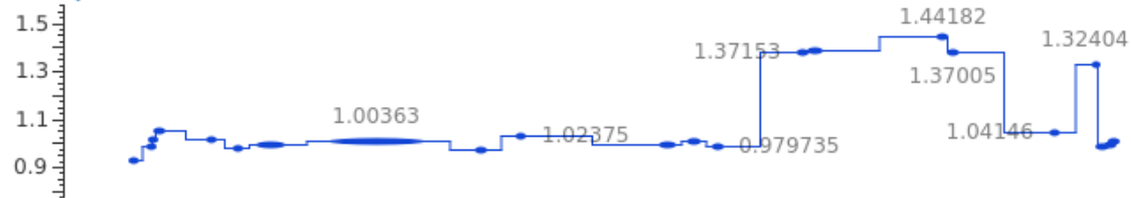
- Reference samples used for normalization
- Probabilistic model used to call CNVs
- Metrics
  - Z-score: number of standard deviations from reference sample mean
  - Ratio: sample coverage divided by reference sample mean
  - VAF: Variant Allele Frequency



# Ratio and Z-score



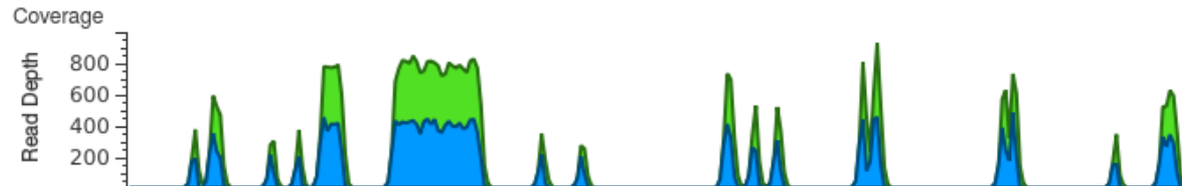
Current Sample: RD-NGSPROGENITYCANCER-SAMPLE13 - Ratio



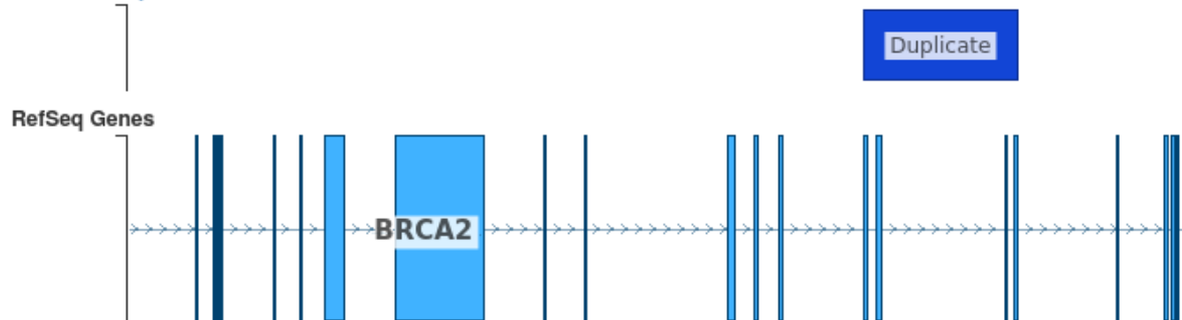
Current Sample: RD-NGSPROGENITYCANCER-SAMPLE13 - Z Score



Current Sample: RD-NGSPROGENITYCANCER-SAMPLE13



Current Sample: RD-NGSPROGENITYCANCER-SAMPLE13 - CNV State





- **VAF provides supporting evidence**

- Values other than 0 or 1 are evidence against het. Deletions
- Values of 2/3 and 1/3 are evidence for duplications

Variants - RD-CRExome-14



Sample 14 (no VAF) - Z score



Sample 14 (with VAF) - Z score





- **Low quality events can be flagged if**
  - Event targets have low coverage
  - There is high variation between samples at event targets
  - Event cannot be differentiated from noise at a region
  
- **Samples can be flagged if**
  - The sample does not match the references
  - The sample has extremely low coverage
  - There is high variance across the target regions

# Reference Samples



- **Matched references are chosen for each sample**
- **Samples with lowest percent difference are chosen**
- **Performance affected if controls don't have matching coverage profile**
- **Samples are flagged if the average percent difference is above than 20%**



- **100x Coverage**
- **Reference samples**
  - Recommend at least 30 references
  - Minimum of 10
  - From same platform and library preparation
  - Gender matched references required for Non-autosomal calls





# Questions or more info:

- Email [info@goldenhelix.com](mailto:info@goldenhelix.com)
- Request an evaluation of the software at [www.goldenhelix.com](http://www.goldenhelix.com)

