

CNV Analysis in VarSeq

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Agenda



1 Overview Golden Helix

2 Why Call CNVs in NGS

3 Method and Demo

4 Availability and Roadmap







Questions during the presentation

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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













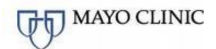




























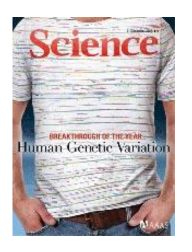




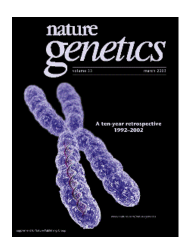


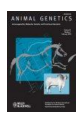
Cited in over 1000 peer-reviewed publications



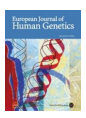


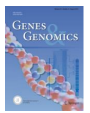


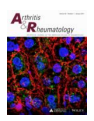




















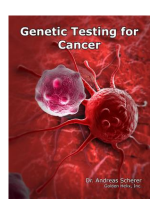
Golden Helix – Who We Are



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



CNV Detection

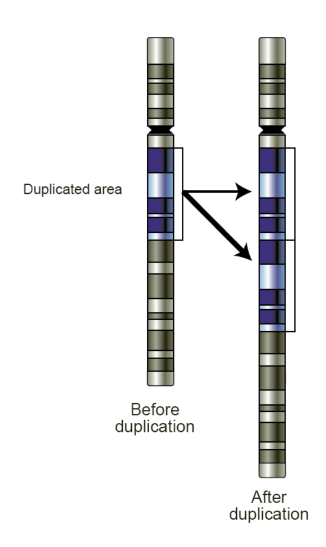


Chromosomal microarray

- 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



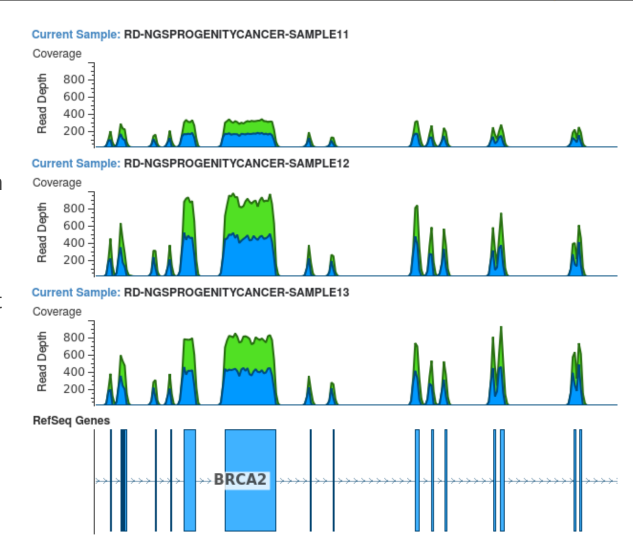
CNVs are called from coverage data

Challenges

- Coverage varies between samples
- Coverage fluctuates between targets
- Systematic biases impact coverage

Solutions

- Data Normalization
- Reference Sample Comparison

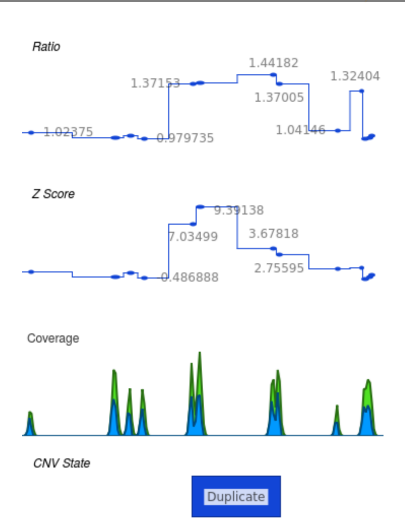




CNV calling in VarSeq



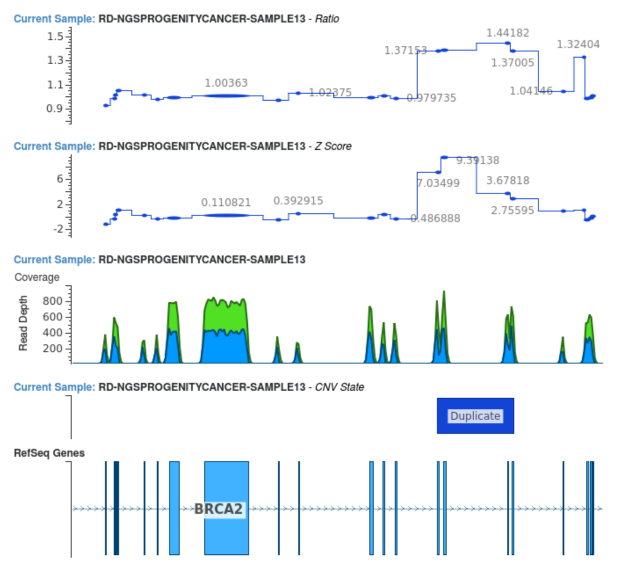
- Reference samples used for normalization
- Probabilistic model used to call CNVs
- Metrics
 - Z-score: number of standard deviations from reference sample mean
 - Ratio: sample coverage divided by reference sample mean
 - VAF: Variant Allele Frequency





Ratio and Z-score





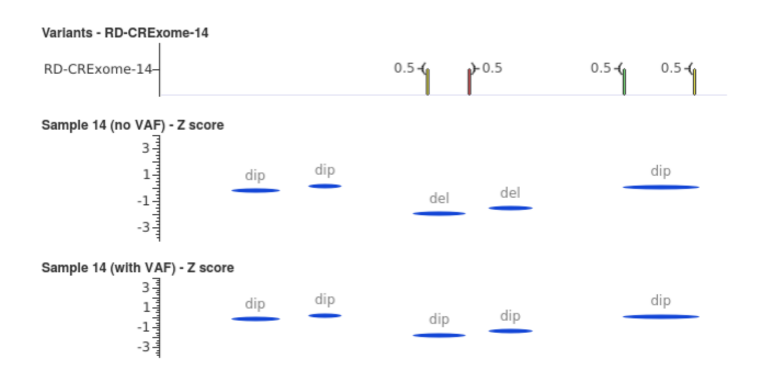


VAF



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





QC Flags



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%



Requirements



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



VarSeq Demonstration







Questions or more info:

- Email info@goldenhelix.com
- Request an evaluation of the software at <u>www.goldenhelix.com</u>



