

NGS TARGET ENRICHMENT HALOPLEX DISEASE RESEARCH PANELS

Premium Performance with Content You Can Trust

Benefits

Confidence in Genomic Content

- Comprehensive content tailored to specific disease research
- Developed in conjunction with top clinical research leaders

Ease of Use From Sample to Analysis

- Simple, automatable protocol with no library preparation
- SureCall software for intuitive variant analysis and report formats

Results You Can Trust

- Multiple amplicon coverage of targets providing better coverage, less PCR artifacts and more accurate mutation calls
- Premium performance including high sensitivity, specificity, and mutation detection

The ability to detect sequence level variations through next generation sequencing in a fast, cost effective application has revolutionized the field of genetics. In order to harness the power of next generation sequencing for specific disease states, it is critical to target these specific genomic regions. Developed in conjunction with clinical research industry leaders, Agilent Technologies offers HaloPlex next generation sequencing target enrichment panels for researching specific genetic disorders.

From Sample to Result in Less than 2 Days

1

Order HaloPlex kit

Ready to Order

Cancer
Cardiomyopathy

or

Made to Order

Arrhythmia
Connective Tissue Disorder
Noonan Syndrome
ICCG
X Chromosome



Available on Illumina MiSeq
and Ion PGM

2

Prepare Samples and Sequence

3

Analyze data, print/export results

SureCall



CATALOG PANELS

HaloPlex Cancer

HaloPlex Cancer is a comprehensive next generation sequencing target enrichment panel designed specifically for genetic anomalies in known cancer hotspots. This NGS application targets a set of 47 genes found in previous research to be associated with a broad range of cancer types as well as with published drug targets. The COSMIC database was the primary reference in the design process.

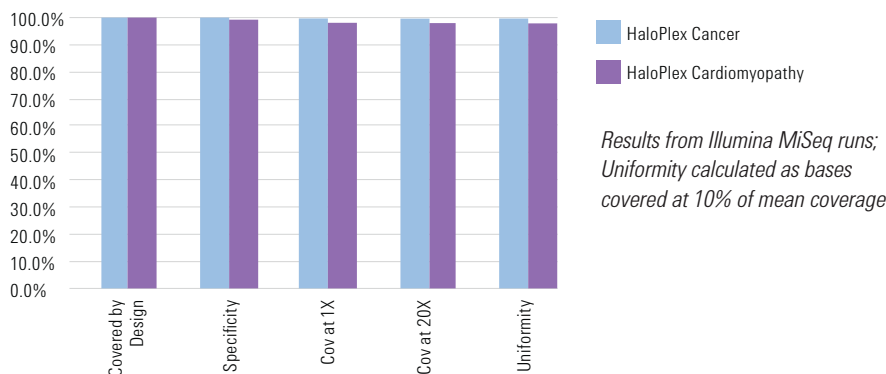
HaloPlex Cancer is uniquely suited for high performance with cancer research samples, which are commonly preserved as formalin fixed and paraffin embedded (FFPE). This FFPE process often results in highly fragmented DNA, resulting in insufficient sequencing target coverage. FFPE also commonly produces small changes in single bases, cytosine to thymine, in DNA sequences. Unlike competitive technologies, HaloPlex covers each base with several amplicons, and smaller fragments function as a backup for longer fragments that may fail. This allows for high sequencing target coverage even in highly degraded FFPE samples.

HaloPlex Cancer Gene List

Targeting solid tumors, hematological cancer and actionable mutations

| | | | | |
|--------|-------|--------|--------|-------|
| ABL1 | ERBB2 | HRAS | MET | SMAD4 |
| AKT1 | ERBB4 | IDH1 | NOTCH1 | SMO |
| ALK | FANCA | IDH2 | NPM1 | SRC |
| AR | FANCC | JAK2 | NRAS | STK11 |
| ATM | FANCF | JAK3 | PDGFRA | TP53 |
| BRAF | FANCG | KIT | PIK3CA | VHL |
| CDKN2A | FGFR1 | KRAS | PIK3R1 | WT1 |
| CSF1R | FGFR2 | MAP2K1 | PTEN | |
| CTNNB1 | FGFR3 | MAP2K2 | RET | |
| EGFR | FLT3 | MAP2K4 | RUNX1 | |

Premium Target Enrichment Performance



HaloPlex Cardiomyopathy

HaloPlex Cardiomyopathy is a next generation sequencing target enrichment panel designed specifically for inherited forms of cardiomyopathy. Following a careful review of cardiomyopathy publications as well as information available from GeneReviews, an NIH online resource, 34 genes known to be associated with hypertrophic cardiomyopathy, dilated cardiomyopathy, and arrhythmogenic right ventricular cardiomyopathy have been included.

HaloPlex Cardiomyopathy Gene List

| | | | |
|--------|-------|-------|--------|
| TTR | ACTC1 | DES | RBM20 |
| MYL2 | TNNI3 | LMNA | TGFB3 |
| MYL3 | TPM1 | SGCD | DSP |
| MYOZ2 | TTN | VCL | PKP2 |
| NEXN | ACTN2 | LDB3 | DSG2 |
| MYH6 | CSRP3 | ABCC9 | DSC2 |
| MYH7 | PLN | SCN5A | TMEM43 |
| MYBPC3 | TNNC1 | TAZ | JUP |
| TNNT2 | TCAP | | |

MADE TO ORDER PANELS

HaloPlex Arrhythmia

Targeting genomic regions known to be associated with four inherited arrhythmia-related heart disorders, HaloPlex Arrhythmia is a next generation sequencing target enrichment panel. Included are 20 genes known to correlate with long QT syndrome, short QT syndrome, Brugada syndrome, and catecholaminergic polymorphic ventricular tachycardia, developed from information gathered after a thorough review of publications for arrhythmia and from GeneReviews, an NIH resource. The genes associated with different types of arrhythmia are overlapping in some cases, and using HaloPlex Arrhythmia, a comprehensive arrhythmia profile can be assembled for clinical research samples.

HaloPlex Arrhythmia Gene List

| | | |
|---------|--------|-------|
| KCNQ1 | CAV3 | SCN1B |
| KCNH2 | SCN4B | KCNE3 |
| KCNJ2 | AKAP9 | SCN3B |
| ANK2 | SNTA1 | RYR2 |
| KCNE1 | SCN5A | CASQ2 |
| KCNE2 | GPD1L | |
| CACNA1C | CACNB2 | |

HaloPlex Noonan Syndrome

HaloPlex Noonan Syndrome is a next generation sequencing panel designed using information from published literature and the NIH resource GeneReviews. This panel is designed to detect genetic mutations known to be associated with Noonan syndrome and related disorders such as LEOPARD, cardio-facio-cutaneous syndrome, and Costello syndromes.

HaloPlex Noonan Syndrome Gene List

| | | |
|--------|--------|-------|
| BRAF | MAP2K2 | RAF1 |
| CBL | NRAS | SHOC2 |
| HRAS | PTPN11 | SOS1 |
| MAP2K1 | KRAS | NF1 |
| SPRED1 | | |

HaloPlex Connective Tissue Disorder

The HaloPlex Connective Tissue Disorder focuses on inherited forms of connective tissue disorders, specifically targeting genetic variations associated with Marfan syndrome, Ehlers-Danlos syndrome, Loeys-Dietz syndrome, thoracic aortic aneurysm and dissection (TAAD), Stickler syndrome, Osteogenesis imperfecta and other related disorders.

HaloPlex Connective Tissue Disorder Gene List

| | | |
|---------|--------|--------|
| AMPD1 | COL6A2 | TCAP |
| LMNA | DES | SGCB |
| SEPN1 | DYSF | TPM2 |
| TPM3 | COL6A3 | FKTN |
| ACTA1 | EMD | POMT1 |
| POMGNT1 | DMD | TRIM32 |
| ANO5 | FHL1 | FKRP |
| PYGM | ITGA7 | TNNT1 |
| TNNI2 | ISPD | MYOT |
| CAPN3 | SGCE | SGCD |
| CAV3 | LAMA2 | SIL1 |
| CHKB | POMT2 | PLEC |
| LARGE | SGCA | SGCG |
| COL6A1 | | |

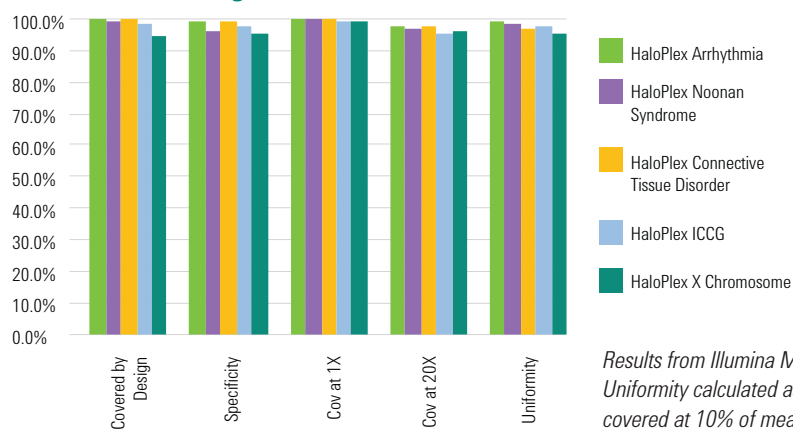
HaloPlex ICCG

ICCG, International Collaboration for Clinical Genomics, is the organization formerly named ISCA, International Standards for Cytogenomic Arrays. In the HaloPlex ICCG gene panel, 180 genes as defined by ICCG have been incorporated into a novel next generation sequencing application, following the associated ICCG recommendations for design and resolution.

HaloPlex X Chromosome

A wide variety of genetic disorders have been shown to be correlated to changes in the X chromosome. The HaloPlex X Chromosome panel is designed to interrogate these particular genetic changes on the X chromosome in a new next generation sequencing kit. Together with SureCall analysis software, laboratories are now able to create a comprehensive X chromosome mutation profile for clinical research samples in one efficient, cost effective application.

Premium Target Enrichment Performance



Results from Illumina MiSeq runs; Uniformity calculated as bases covered at 10% of mean coverage

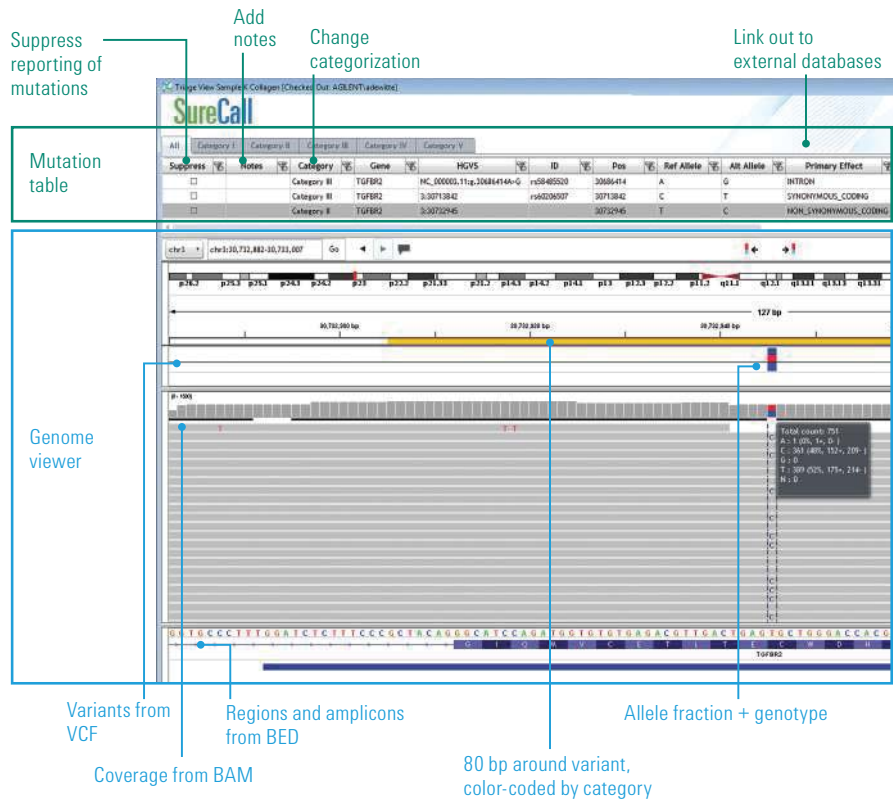


HALOPLEX DISEASE RESEARCH PANELS

SureCall Software

SureCall addresses the critical need of clinical researchers. The free-of-charge software incorporates the most widely accepted open source algorithms.

- Easy-to-use and faster time-to-results
- Streamlined workflow for analysis
- Reduce the need for complex bioinformatics



SureCall display showing mutation found for sample sequenced using the HaloPlex Connective Tissue Disorder Panel. On the left, aligner view highlighting mutation; on the right, final report exported from software.

Catalog, Ready to Order Panels

| Panel | PN | Reactions |
|----------------|-------------------------------|-----------|
| Cancer | G9903A (ILM), G9904A (ION) | 16 |
| | G9903B (ILM), G9904B (ION) | 96 |
| Cardiomyopathy | G9908A (ILM), G9909A (ION) | 16 |
| | G9908B (ILM), G9909B (ION) | 96 |

Pre-Designed, Made-to-Order Panels

| Panel | Design ID (ILM) | Design ID (ION) | Ordering |
|----------------------------|------------------|------------------|---|
| ICCG | 00100-1358263628 | 00100-1360592497 | Order each Design ID using a Custom PN, through SureDesign www.agilent.com/genomics/suredesign |
| Connective Tissue Disorder | 00100-1358243605 | 00100-1360592472 | |
| X-Chromosome | 00100-1358242818 | N/A | |
| Arrhythmia | 00100-1358263563 | 00100-1360592417 | |
| Noonan Syndrome | 00100-1358243073 | 00100-1360592460 | |

References

1. ICCG website, <https://www.iccg.org/>

Request more information or buy online: www.agilent.com/genomics/ngs



Find an Agilent customer center in your country:
www.genomics.agilent.com/contactUs.jsp
 U.S. and Canada: 1-800-227-9770 | cag_sales-na@agilent.com

© Agilent Technologies, Inc. 2014, 2015
 Printed in USA, September 3, 2015
 5991-2526EN
 For Research Use Only. Not for use in diagnostic procedures.

