

Simultaneous Detection of Copy Number and Copy-Neutral LOH Using a Single Microarray

Application Note

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Abstract

The current SurePrint G3 Human Comparative Genomic Hybridization (CGH) platform has been extended to include a set of Single Nucleotide Polymorphism (SNP) probes on the same microarray. This allows for the simultaneous, high-resolution detection of copy number and copy-neutral variations and eliminates the need to run two separate microarray experiments. The same straightforward Agilent CGH workflow is used to prepare the samples for hybridization to the new SurePrint G3 Human CGH+SNP microarrays. Restriction digestion of genomic DNA allows genotyping of SNPs located in the enzymes' recognition sites. After labeling and hybridization, CGH+SNP microarray data are analyzed using novel algorithms contained within Agilent's Genomic Workbench software. In this application note, we measure SNP genotypes to discover copy-neutral lack or loss of heterozygosity (LOH), as well as copy number variations, to discover cytogenetic abnormalities associated with examples of known syndromes. We conclude that simultaneous high-quality, high-resolution copy number and LOH information can be obtained with the new SurePrint G3 Human CGH+SNP microarrays, available in various catalog and custom formats.



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Introduction

Accurate determination of copy number aberrations is important for understanding developmental disorders. Since array CGH was first described in 1998 by Pinkel et al.¹, many technical improvements have been made resulting in higher specificity and sensitivity. Recently, oligo microarrays, which have a 1000-fold greater resolution than karyotypes, have revolutionized the field of cytogenetics. Array CGH is now commonly used for investigating genomic DNA aberrations associated with various diseases in many cytogenetic labs.

Until recently, Agilent's SurePrint G3 CGH Microarrays have provided ultra-high resolution copy number analysis but could not detect copy-neutral aberrations such as LOH or uniparental disomy (UPD). In LOH events, several mechanisms can be responsible for the loss of the wild-type allele at a heterozygous locus and the unmasking of

a recessive mutant allele.² UPD occurs when both members of a chromosome pair or segments of a chromosome pair are inherited from one parent and no chromosomes are inherited from the other parent.³ If the chromosomes involved are imprinted such that the genes on these chromosomes are monoallelically active (i.e. only the maternal or paternal allele of the pair is expressed), the resulting phenotype will be abnormal. We now have developed a novel approach that provides high-resolution copy number analysis and LOH or UPD data on a single microarray. A subset of probes is used to measure SNPs in parallel with copy number by performing the standard CGH enzymatic workflow. In addition to high-quality copy number aberration discovery, the new Agilent SurePrint G3 CGH+SNP Microarrays also enable detection of copy-neutral LOH, UPD, consanguinity, and parental or sample origin in genetic studies.

Table 1. SurePrint G3 CGH+SNP 2x400K and 4x180K microarray specifications

Format	2x400K	4x180K
Design ID	028081	029830
Total features	420,288	180,880
Control feature count	6,700	8,121
Distinct biological CGH probes	292,097	110,712
Distinct biological SNP probes	118,955	59,647
Distinct SNPs	64,484*	59,647
Replicated probes (5x)	600	600
Genome build	hg19	hg19
Non-unique probes	506	1,467
Homology filtered probes	3,950	2,931
Intragenic probes	210,298	49,963
Intergenic probes	81,799	60,749
Exonic probes	113,770	6,574
Median probe spacing bp		
- <i>intragenic</i>	4,453	21,264
- <i>intergenic</i>	15,875	27,323
- <i>overall</i>	7,170	25,288
Average probe spacing bp		
- <i>overall</i>	10,257	27,066
Refseq coverage (26,855)		
- <i>at least one probe</i>	24,639	16,867
- <i>at least one probe (+/-5kb)</i>	25,245	20,847

*92% of SNPs use two SNP probes per SNP

Methods

Microarray design

The catalog 2x400K SurePrint G3 CGH+SNP array used in this study contains approximately 300,000 CGH probes and 120,000 SNP probes. The genotypes are measured by using two SNP probes per SNP and result in ~5–10 Mb resolution for LOH detection across the entire genome. The CGH probes are gene- and exon-biased (Table 1). The catalog 4x180K SurePrint G3 CGH+SNP array contains approximately 120,000 CGH probes and 60,000 SNP probes. The genotypes on this array are measured by using one SNP probe per SNP and also result in ~5–10 Mb resolution for LOH detection across the entire genome. The CGH probes consist of the entire ISCA (International Standards for Cytogenomic Arrays) Consortium 8x60K version probe set and an additional 60,000 backbone probes. The ISCA 8x60K has an even backbone probe coverage of 60 Kb and high-density coverage of ~500 targeted regions with the spacing of 5 Kb per probe or at least 20 probes per gene region. These targeted regions include telomere and unique centromere FISH clone regions, microdeletion/duplication regions, genes of known haploinsufficiency, and X-linked mental retardation regions. Custom SurePrint G3 CGH+SNP microarrays can be readily designed in eArray, a free web-based application, or eArray XD, the desktop version of eArray (Table 2). The eArray database houses more than 28 million CGH probes and probes for approximately 60,000 SNPs.

To ensure that additional genotype information from the SNP probes doesn't compromise the quality of the copy number information supplied by the CGH probes and that the results are comparable to legacy CGH arrays, SurePrint G3 CGH microarrays, 4x180K (P/N G4449A) were processed concurrently.

Sample preparation, hybridization, and imaging

All samples were obtained from the Coriell Cell Repository (<http://www.coriell.org/>) and were processed by following the Agilent Oligonucleotide Array-Based CGH for Genomic DNA Analysis, Enzymatic Labeling for Blood, Cells, or Tissues (with a High-Throughput Option) manual, version 6.3 P/N G4410-90010. Genomic DNA was digested with AluI and RsaI in order to enable detection of SNPs located at the enzymes' recognition sites. HapMap samples and samples that have aberrations associated with known cytogenetic disorders were labeled with Cy5 dye. Because the current CGH+SNP assay requires a sample of known genotype as the reference sample, we chose a HapMap sample as a reference, labeled it with Cy3 dye, and hybridized it to the same microarray. The microarray slides were scanned at 3 micron resolution on the Agilent High-Resolution C Scanner and the images were extracted using the Agilent Feature Extraction software (AGW version 6.5).

Table 2. Agilent SurePrint G3 Catalog and Custom CGH+SNP Microarrays

Catalog Kit	1x1M Kit	2x400K Kit	4x180K Kit	8x60K Kit
Number of Arrays/Slide	N/A	2	4	N/A
Number of Slide/Kit	N/A	5	3	N/A
Catalog Kit P/N	N/A	G4842A	G4890A	N/A
Content				
Number of CGH probes	N/A	~300K	~120K*	N/A
Number of SNP probes		~120K	~60K	
Number of probes per SNP		2	1	

Custom Microarray	1x1M	2x400K	4x180K	8x60K
Number of Arrays/Slide	1	2	4	8
Number of Slides	1	1	1	1
Custom Slide P/N	G4882A	G4883A	G4884A	G4885A

*includes ISCA 8x60K

Novel algorithms for measuring total copy number and allele-specific copy number

The SNP probes span variant AluI or RsaI restriction enzyme recognition sites and measure the copy number of the uncut allele at those loci. The total copy number of the region encompassing the SNP site is measured by neighboring CGH probes. The copy number of the cut allele can be inferred from the total copy number and the copy number of the uncut allele.

SNP calls are made from the log ratios of the sample probe signal versus a genotyped internal reference, which compensates for labeling and hybridization bias. Since the reference genotype is known, the raw \log_2 ratios are "reference adjusted" to report absolute allele-specific copy numbers for SNP probes. SNPs with zero copies of uncut allele in the reference sample are ignored, and the algorithm subtracts 1 from the \log_2 ratios of SNPs with one copy in the reference sample. When using eArray to design a custom microarray, and when using one of the supported reference samples (NA12878, NA12891, NA18507, NA18517 or NA18579), the user can choose to avoid SNPs with zero reference copies on the array design. The reference-adjusted log ratios fall into three nearly Gaussian peaks corresponding to the copy numbers of the uncut alleles in the sample, which correspond to the three possible diploid genotypes for the SNPs: AA, AB, or BB (Figure 1).

Regions of copy-neutral LOH or UPD are then located by identifying genomic regions with a statistically significant scarcity of heterozygous SNP calls.

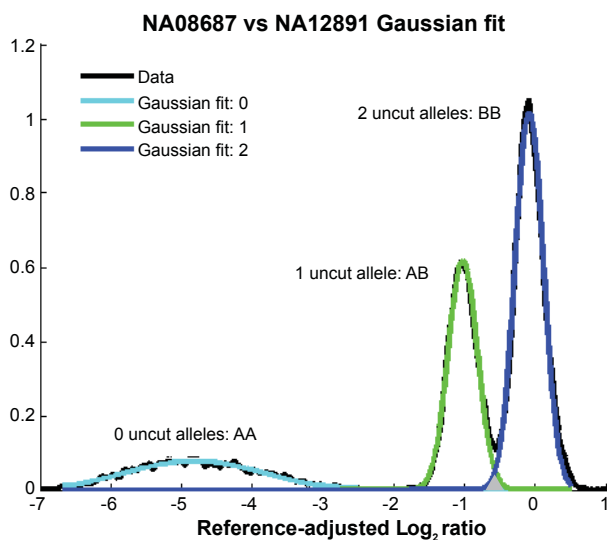


Figure 1. Reference-adjusted SNP \log_2 ratio distributions (probability density function). The black line is the distribution of measured log ratios. The cyan, green, and blue lines are 3 different Gaussian fits to the population of SNPs with 0 (AA), 1 (AB), and 2 (BB) copies of the uncut allele (A represents the cut allele and B represents the uncut allele).

Results

SNP call rate and accuracy

To determine the SNP call rate, defined as the fraction of the SNPs called by the algorithm at 95% confidence, and the SNP call accuracy, several HapMap samples with known genotypes were hybridized against another HapMap sample. In this experiment >95% of SNPs were called at a confidence level of >95% (Figure 2), with >99% accuracy (Figure 3).

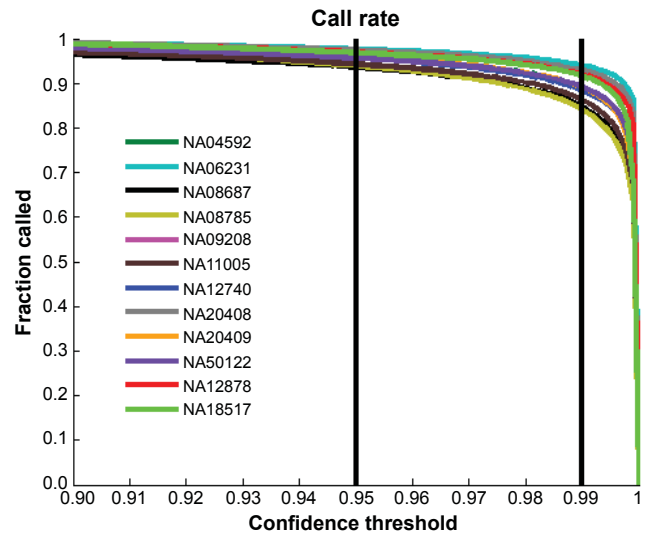


Figure 2. Call rate for 12 samples hybridized against a single HapMap reference sample. More than 95% of SNPs were called at a confidence level of >95% in all 12 samples. The black vertical lines correspond to p-value thresholds (confidence levels) of 0.95 (95%) and 0.99 (99%).

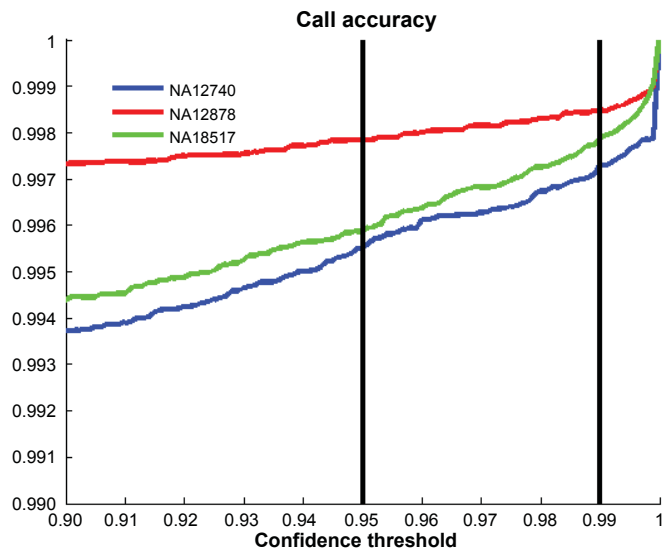


Figure 3. Call accuracy for 3 HapMap samples hybridized against a single HapMap reference sample. The call accuracy was >99% in all 3 samples. The black vertical lines correspond to p-value thresholds (confidence levels) of 0.95 (95%) and 0.99 (99%).

Concurrent analysis of CGH and SNP data

Data from samples containing known cytogenetic aberrations are displayed in Figure 4. For each image, the Genomic Workbench view is composed of three horizontally stacked panels—the top one displaying the number of uncut alleles for each SNP probe, the middle the CGH \log_2 ratios for the CGH+SNP microarrays, and the bottom one the \log_2 ratios for the CGH-only microarrays. For a normal diploid region of the genome, one expects the 0, 1, and 2 SNP copy numbers of the uncut allele to be randomly distributed, as represented by the three states (red dots) in the SNP panel. In a diploid genome carrying a copy-neutral LOH or UPD aberration, the SNP probes will only report alleles that are homozygously cut and uncut (0 and 2 uncut alleles) and therefore only two states are found. A region of the genome affected by a hemizygous deletion is also seen as two states that, in this case, represent SNPs that only have one copy of the allele, either cut or uncut (0 or 1 copy of the uncut allele). The amplification of a region or entire chromosome adds a state that corresponds to the probes reporting the additional uncut allele (0, 1, 2, 3 ... copies of the uncut allele). Table 3 summarizes the relationship of genotype to SNP status.

The SurePrint G3 CGH+SNP microarrays correctly identified both known copy number and copy-neutral variations. The CGH data was not compromised by the presence of SNP probes on the microarrays—expected gains and losses were consistently and confidently identified based upon published information. All QC metrics passed the Agilent recommended thresholds (Table 4). Copy number aberrations detected on the G3 CGH+SNP microarrays and computed by Agilent Genomic Workbench 6.5 (Figure 4) are comparable to those for G3 CGH microarrays.

Table 4. QC results of the SurePrint G3 CGH+SNP 2x400K microarrays

QC Metric	HapMap/Reference
DLRSpread	0.14 ± 0.005
SignalToNoiseExp	126 ± 4
SignalToNoiseRef	150 ± 2
SignalIntensityExp	381 ± 13
SignalIntensityRef	300 ± 4
BGNoiseExp	3.03 ± 0.03
BGNoiseRef	2.00 ± 0.01
ReproducibilityExp	6.33 ± 0.52
ReproducibilityRef	5.98 ± 0.44

Average scores are provided along with standard deviations across 3 samples. All QC metrics passed, including an excellent DLR Spread (derivative log ratio spread, measure for probe-to-probe log ratio noise, below 0.2).

Table 3. Relationship of genotype to SNP status (number of uncut alleles)

Genomic status	Genotype	No. uncut allele
Normal diploid genome	AA, AB, BB	0, 1, 2
Diploid genome with copy-neutral LOH or UPD	AA, BB	0, 2
Hemizygous LOH	A, B	0, 1
Amplification: e.g. trisomy	AAA, AAB, ABB, BBB	0, 1, 2, 3

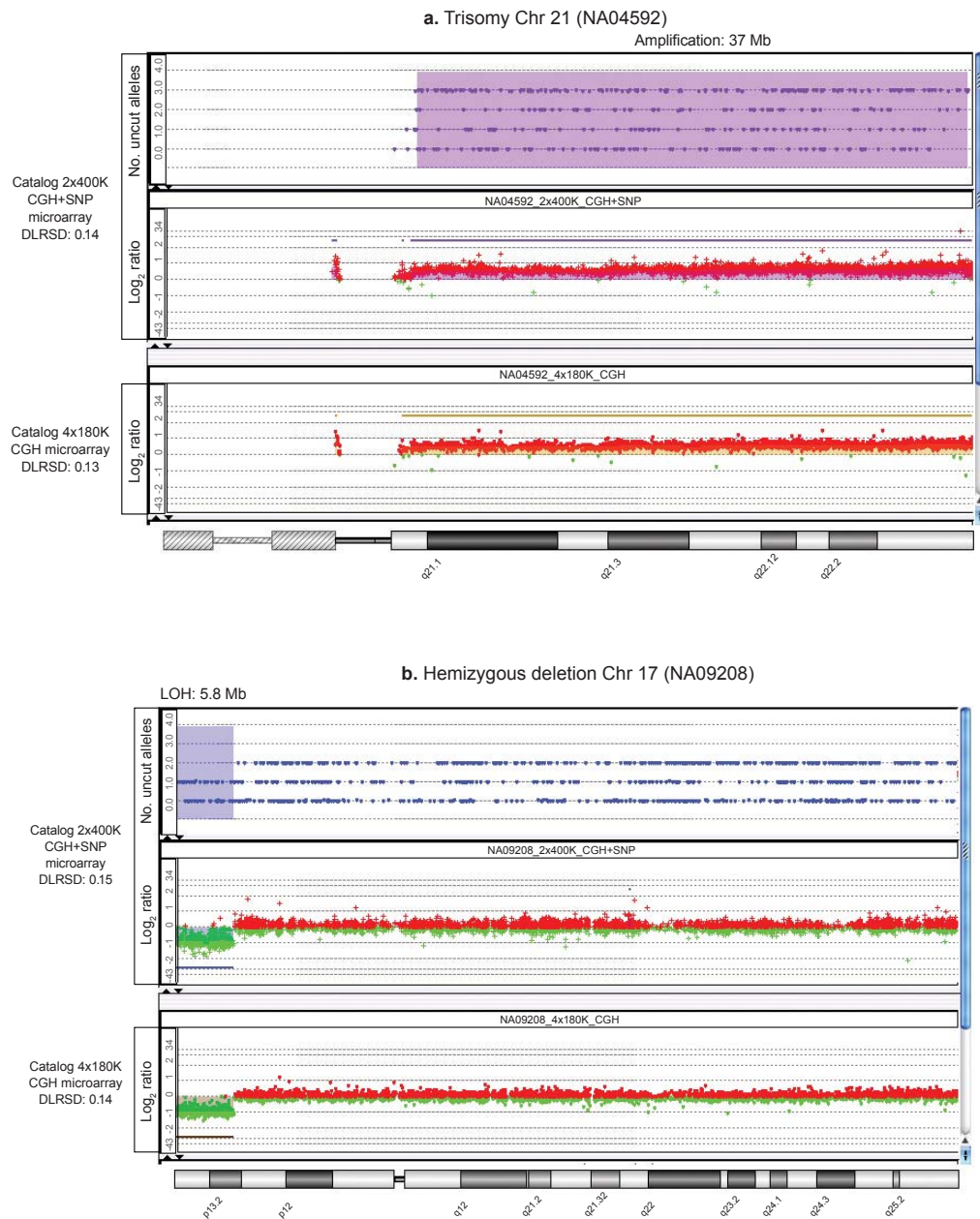


Figure 4. Agilent Genomic Workbench view of the SNP data (number of uncut alleles, top panels), the CGH data (\log_2 ratios, middle panels) for the CGH+SNP arrays, and the CGH data for the CGH-only arrays (\log_2 ratios, bottom panels) in affected chromosomes: a. trisomy of chromosome 21, b. hemizygous deletion in chromosome 17 (continued on opposite page).

A trisomy of chromosome 21 was identified in one sample (Figure 4a). Several LOH regions as small as ~5 Mb were detected. One sample with a hemizygous deletion was accurately profiled by the CGH and SNP probes (Figure 4b). In one case, extended segments of homozygosity were

found throughout the genome as a consequence of parental consanguinity (Figure 4c). In an additional case, the UPD observed was associated with an individual known to have genomic aberrations associated with Angelman syndrome (Figure 4d).

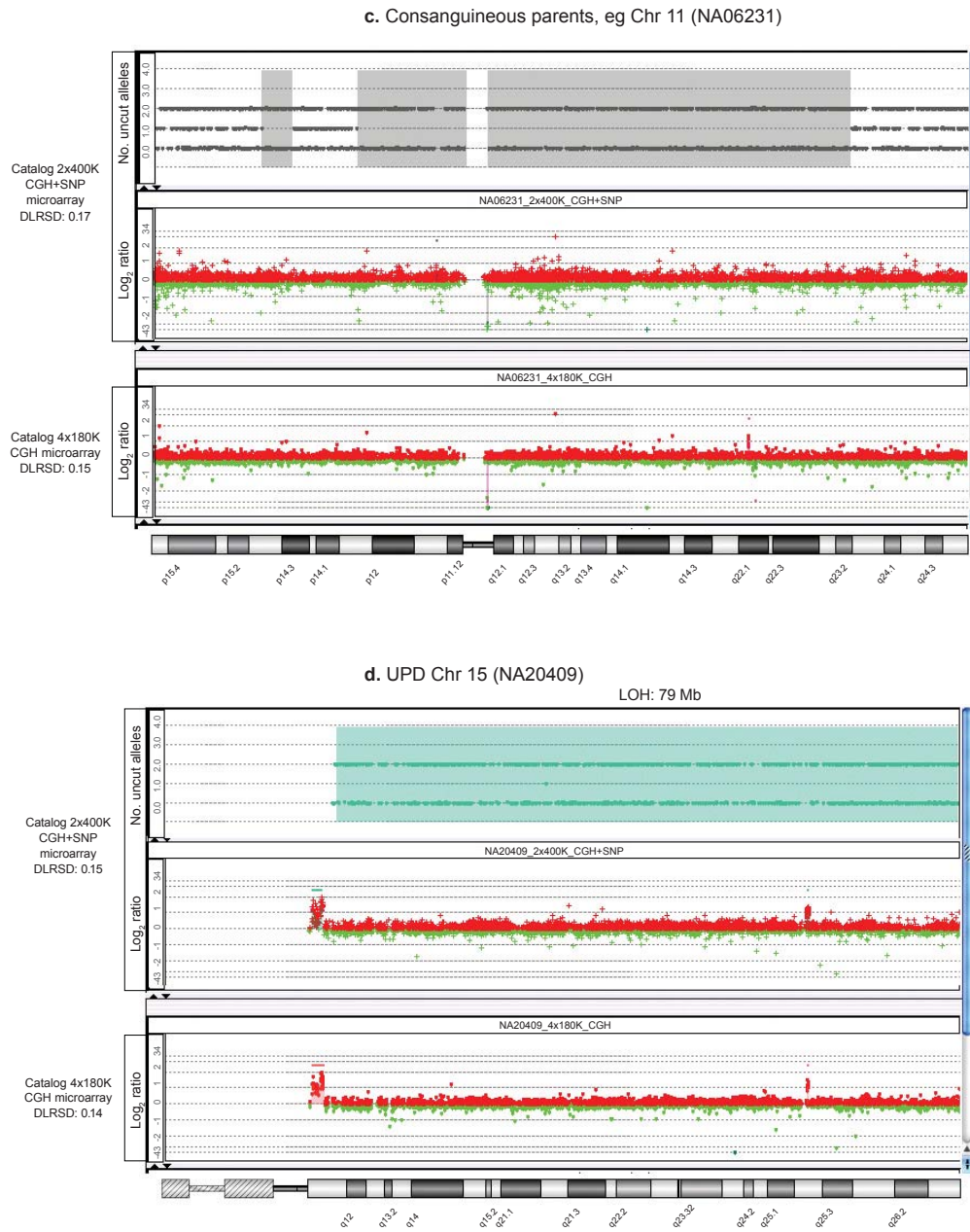


Figure 4 (continued). c. continuous segments of homozygosity due to parental consanguinity, d. UPD of the entire chromosome 15. Settings for CGH aberration calling: ADM-2, threshold 5, minimum of 3 probes, $\geq 0.25 \log_2$ ratio. Purple: amplification, blue: LOH due to deletion, grey and aqua: copy-neutral LOH.

Conclusion

We have shown that the addition of SNP probes to the Agilent SurePrint G3 CGH Microarrays enables the detection of blocks of LOH that could result from UPD or consanguinity. Copy-neutral LOH and UPD are confidently and accurately identified in samples known to have these aberrations. Moreover, SNP calls match copy number calls in regions of hemizygous deletions and amplifications. The addition of SNP probes does not compromise the performance of CGH probes,

and CGH aberration calls are not affected by the addition of SNP content to the microarrays. In conclusion, we have demonstrated that the new Agilent SurePrint G3 CGH+SNP Microarrays can simultaneously provide ultra-high-resolution copy number analysis and can detect significant blocks of LOH or UPD. As a result, researchers no longer need to choose between high-resolution copy number data and UPD/LOH or alternatively run two separate microarrays. Now one microarray can detect both.

References

1. Pinkel *et al. Nat Genet.* 1998;20(2):207–11.
2. Cavanee *et al. Nature.* 1983;305(5937):779–84.
3. Engel. *Am J Med Genet.* 1980;6(2):137–43.

Required Agilent CGH Processing Components

Description	Part Number
SureTag Complete DNA Labeling Kit	5190-4240
Human Cot-1 DNA	5190-3393
Agilent Oligo aCGH Hybridization Kit (25) or (100)	5188-5220 or 5188-5380
Agilent Oligo aCGH Wash Buffer 1 and 2 Set	5188-5226
Hybridization Chamber, stainless	G2534A
Hybridization Chamber Gasket Slides	Varies by array format and quantity
Hybridization Oven	G2545A
Hybridization Oven Rotator Rack	G2530-60029
SureScan Microarray Scanner	G4900DA
Agilent CytoGenomics	G1662AA–G1667AA

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