The performance of CGH array for the detection of cryptic constitutional chromosome imbalances

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Gene dose alterations can cause mental retardation (MR), congenital malformations and miscarriages. Standard chromosome analysis by G-banding has a limited resolution, but molecular cytogenetic techniques, such as multi-subtelomeric FISH, microdeletion FISH, multi-colour FISH and comparative genomic hybridisation (CGH), have played an important role for the diagnosis of MR during the past decade.1 A complete set of subtelomeric FISH probes was presented in 1996 and updated in 2000.2 Consequently, screening for subtelomeric abnormalities has become a diagnostic test that is offered by diagnostic laboratories, and a number of studies reporting new subtelomeric rearrangements have been published.3–16 However, these probes only reveal chromosome rearrangements located in the subtelomeric region. To cover the whole genome, genome wide screening for chromosomal imbalances using micro-satellite markers has been reported,17 18 as well as metaphase CGH.19–21 Yet none of these techniques is able to offer a high resolution screening of the whole genome for chromosome imbalances. The development of accurate and sensitive genome wide screening methods would facilitate the clinical diagnosis of patients with very small or subtle rearrangements. Screening for chromosomal imbalances using array CGH, whether using cDNA19 or BAC clones,20 has mainly been performed on cancer samples,25–30 which usually contain large gene dose alterations. Although array CGH has provided a higher resolution compared to conventional CGH, it has not yet become a widely applied method for the analysis of gene dose alterations in individuals with idiopathic mental retardation. It has been a challenge to achieve the adequate performance needed for the reliable detection of single copy losses or gains of very small regions. Chromosome specific micro-arrays have however been used in a few cases to determine the critical regions in microdeletion disorders.31–32

In this report we used a cDNA micro-array and two BAC clone micro-arrays on 10 previously identified cases containing 16 cryptic chromosome aberrations, to test their performance for the detection of small imbalances.

MATERIALS AND METHODS

Material selection

Ten cases with known cryptic aberrations were selected (seven cases with subtelomeric rearrangements, one case with an interstitial deletion and two cases with microdeletion syndromes: one case with DiGeorge syndrome and one case with a microdeletion of 17p11 (Smith Magenis syndrome) in 60% of the lymphocytes). The 10 cases contained altogether 16 aberrations, with sizes ranging from 1.3 to 20.5 Mb, and they were located on 15 different chromosome arms. Routine chromosome analysis (450–500 bands) had failed to detect the rearrangements in all cases. Cases 2–8 (table 1) were identified using subtelomeric FISH probes. Cases 3–6 and 8 were previously published.31 Case 7, with an interstitial deletion of 15q24, was serendipitously detected during subtelomeric screening, as the 15q control probe (LSI PML) in the subtelomeric screening kit (Vysis Inc, Downers Grove, IL) was deleted. Cases 9 and 10 were detected using microdeletion FISH probes. Case 1 was identified by spectral karyotyping. Despite the large chromosome fragments involved in the unbalanced translocation in this case, repeated standard G-band analysis could not reveal this rearrangement due to the similarity in the banding pattern of the chromosome fragments involved. All chromosome imbalances were size mapped using BAC and PAC clones based on clone mapping databases (www.ensembl.org, July 2003), except for case 9, which was size mapped using a chromosome 22 specific micro-array.31

FISH mapping


Abbreviations: BAC, bacterial artificial chromosome; CGH, comparative genomic hybridisation; FISH, fluorescence in situ hybridisation; MR, mental retardation; PAC, P1 derived artificial chromosome

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206 BAC and P1 derived artificial chromosome (PAC) clones were selected for breakpoint mapping of the chromosome aberrations. The clones were obtained from Resources for Molecular Cytogenetics (Bari, Italy), The Wellcome Trust Sanger Institute (Cambridge, UK) and BACPAC Resource Center Children’s Hospital, (Oakland Research Institute, Oakland, CA). Dual colour FISH was performed by hybridising each of the BAC clones to metaphase slides prepared from the selected patients as previously described.\(^3\)

DNA preparation, digestion, and labelling

Genomic DNA was extracted from blood samples or Epstein-Barr virus transformed lymphocytes using Puregene blood kit (Gentra systems Inc, Minneapolis, MN) according to the manufacturer’s protocol. Genomic DNA from the patients was digested into fragments of 100–2000 bp with DpnII (New England Biolabs Inc, Beverly, MA) and purified using Qiaquick PCR purification kit (Qiagen GmbH, Hilden, Germany). 0.8–2 µg of digested patient DNA, as well as reference DNA, was labelled by random priming with Cy5-dCTP and Cy3-dCTP (Amersham Biosciences, Buckinghamshire, UK) respectively, using Bioprime labelling kit (Invitrogen, Carlsbad, CA).

**cDNA micro-array experiment**

cDNA micro-array production was performed within the Van Andel Institute Laboratory of Microarray Technology as described by Takahashi et al.\(^3\) Briefly, cDNA clones of the sequence verified human cDNA library (Research Genetics, Huntsville, AL) were PCR amplified directly from bacterial stock. The purified PCR products were robotically arrayed onto amino silane coated glass slides. Slides were blocked using succinic anhydride under standard conditions (http://cmgm.stanford.edu/pbrown/). The arrays contained 21 632 human cDNAs, generally with insert sizes of 0.25–2.5 kb.

0.8–2 µg of labelled probes were mixed with 40 µg human Cot-1 DNA (Invitrogen, Carlsbad, CA), before 16 ml of 2×hybridisation solution (50% formamide, 10×SSC, 0.2% SDS) was added. The probes were denatured at 100°C for 3 min and prehybridised at 37°C for 30 min. After 16 h hybridisation at 42°C on the micro-array in a hybridisation chamber (Corning Inc, Corning, NY), the slides were washed in 1×SSC/0.1% SDS, 0.2×SSC/0.1% SDS and 0.1×SSC for 5 min each at room temperature and immediately dried by centrifugation.

**BAC micro-array experiments**

We used two different commercially available genomic human BAC arrays (Spectral Genomics Inc, Houston, Texas). The arrays contained 1000 and 2500 BAC and PAC clones respectively, with all clones printed in duplicate. Hybridisation was performed according to the manufacture’s protocol.

**Data analysis**

Arrays were scanned in a fluorescence confocal scanner (Scan Array Lite, GSI Lumonics, Billerica, CA). Images were analysed using GenePix Pro 3.0 software (Axon Instruments, Burlingame, CA). Spots were defined by the automatic grid feature of the software and adjusted manually when necessary. Further data handling was performed using Microsoft Excel. The fluorescence ratios across the array were normalised to achieve an average ratio of 1 for all cDNA or BAC elements on the array. Colour reverse experiments were performed for each patient sample to reduce variations related to labelling and hybridisation efficiencies. The mean Cy5/Cy3 ratios, standard deviation and co-efficient of variation of four replicas for each clone were calculated.

The cDNA micro-array consisted of 6292 known genes and 9944 unknown genes. For approximately 10 000 of these 16 236 different UniGenes, sequence accession numbers for each spot on the micro-array were mapped to UniGene cluster identifications and chromosome location using ftp://ftp.ncbi.nlm.nih.gov/repository/UniGene (June 2002). Average ratios were calculated between fluorescent signals for each cDNA clone that clustered to the same gene and also from the colour reverse experiments.

The BAC clone arrays were analysed by calculating the average ratio between four fluorescent signals (two signals from the duplicated clone on the array and two signals from the colour reverse experiment) and the fluorescence ratios were plotted according to their mapped location (http://www.ncbi.nlm.nih.gov/mapview/ (March 2003)). Clones displaying a standard deviation >0.20 were interpreted as hybridisation failures and were excluded from the BAC array analysis. DNA copy number profiles >1.2 were scored as gained, <0.80 as lost, as previously described.\(^9\),\(^25\)

**RESULTS**

The data for each case are summarised in table 1. Using the cDNA array we detected 50% of the aberrations (7 out of 14). The smallest detected aberration by this method was a terminal deletion of 8.7 Mb on chromosome 4q (fig 1A).

The results from the cDNA arrays were difficult to interpret because of numerous false positive and false negative findings. The aberrations were only detectable when the Cy5/Cy3 ratio values from a large number of clones located next to each other on the physical map deviated from ratio 1 (normal). Fig 1A illustrates the detection of the unbalanced rearrangements of case 5 with a duplication of 2q and a deletion of 4q, barely detectable by the cDNA array.

The results of the BAC clone micro-array containing 1000 clones were confusing (data not shown). False positive and false negative results were detected using this array and the smallest aberration detected was a 3.5 Mb deletion on chromosome 6q (case 6), while a large duplication on chromosome 9q (case 3) was not detected. This can be explained by errors in the physical mapping of the clones, which resulted in chromosome regions that were not well covered (9qter had a gap of 5.3 Mb). In addition, one clone on 9qter showed significant homology to chromosome 13 and three clones on 9qter gave false negative results due to weak specific signals.

The 2500 clone array detected all of the aberrations studied and their sizes could also be determined by approximately 1 Mb accuracy (table 1). No false positive results were observed. However, false negative findings were observed. Nine clones located in the subtelomeric regions of the aberrant chromosomes, four clones on 10q (AL392043.1, RP11-90B19, GS-137-E24, GS-261-B16), two clones on 17q (AQ285007, GS-362-K4), one clone on 12q (RP11-1192J, one clone on 6p (AL035696.14), and one clone on 9q (RP11-89P10) showed a ratio close to 1 (normal). In addition, two clones on the X chromosome (CTB-18817, AL031643.1) did not show the expected ratio gain (>1.2) when a female sample was hybridised against a male reference, or loss (<0.8) when a male sample was hybridised against a female reference. Clones AL031643.1 and CTB-18817 were tested for cross-hybridisation by FISH. Clone AL031643.1 hybridised weakly but specifically to chromosome Xp21.1. However, CTB-188117 was not X-specific, since it also hybridised on the Y chromosome (data not shown). All aberrations from cases 1, 4, and 6 to 10 were also correctly detected in a blind analysis performed by Spectral Genomics using a 1400 BAC array (data not shown).
DISCUSSION

Reliable and high-resolution detection of copy number changes of genomic DNA is fundamental in diagnosing patients with mental retardation and chromosome imbalances in cancer. CGH makes it possible to screen the whole genome for genomic imbalances. However, CGH performed on metaphase chromosomes has a comparatively low resolution (5–10 Mb) and the resolution is likely to depend on the chromosome region involved. The use of cDNA micro-arrays was not very successful in delineating chromosome changes and there may be several explanations for these results. Firstly, cDNA micro-arrays were primarily developed for expression profiling. The cDNA sequences are therefore mostly selected based on gene function, not on chromosome location, which results in poor coverage of some chromosome regions. Secondly, we found it difficult to interpret the

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D, detected; ND, not detected; --, not performed
*Mosaic (60% of lymphocytes).

Figure 1  A) Ratio plot from Human cDNA micro-array data from chromosome 2 and 4 (case 5). Each dot on the chart represents the normalised, average ratio between fluorescent signals from colour reverse experiment for each cDNA sequence that cluster to the same gene. The dots are linearly ordered along the chromosome according to their Mb location, starting with pter to the left and qter to the right. The plot shows a duplication of 2qter, and a deletion of 4qter is barely visible. Not all chromosome regions are well represented, especially on chromosome 4, and many ratios deviate from a modal value of 1.0, which complicates the interpretation. B) Ratio plot from Human BAC micro-array data, containing 2500 clones from all chromosomes with a magnification of ratio plot for chromosome 2 and 4 (case 5). The dots are linearly ordered along all chromosomes according to their Mb location, starting with chromosome 1 to the left and the Y chromosome to the right. The plot shows a clear duplication of 2qter, a clear deletion of 4qter, a duplication of the X chromosome and a deletion of Y (the patient DNA is hybridised against reference DNA of the opposite sex). Spots showing ratios >0.8 and <1.2 are considered to be normal.
the use of specifically designed DOP-PCR primers for single ESTs are not reliable targets in array CGH experiments for the detection of single DNA copy changes. 26 Thirdly, the slides were coated to attach the cDNA on the glass surface, a strategy that introduces hybridisation background noise.

Several different strategies to improve the specific signal to noise ratio have been reported during the past year—for example the repeat free and non-redundant strategy27 and the use of specifically designed DOP-PCR primers for amplification of the BAC clones.28 Regarding the BAC micro-arrays used in this study, the DNA of the BAC clones was not PCR amplified but purified in large scale from culture and the nucleic acids were attached onto an uncoated glass surface by using chemical coupling.29 This increases the sensitivity and lowers the background signal.

In conclusion, the human BAC clone micro-array works very well for the detection of cryptic chromosome rearrangements. In our hands the method was very robust. It is also very useful for size mapping of the aberrations, which facilitates the phenotype-genotype correlation. It is therefore likely that the CGH array will be offered as a genetic test in clinical diagnostic laboratories in the near future. Regular CGH and cDNA arrays are limited in sensitivity and robustness and are therefore not as reliable as the BAC array. False positive and false negative results have also been a large concern when using BAC arrays for genomic screening for single copy detection. We experienced these problems using the 1000 clone BAC array, but the improved 2500 clone array did not show any false positive results, and only 11 clones showed false negative results. There are several plausible explanations for the fact that these clones showed fluorescence ratios deviating from the expected values for one copy loss or gain. Segments of these clone sequences can be highly similar to sequences present elsewhere in the genome (for example, clone CTB-188177 shown not to be X-specific) and some clones might contain large segments of commonly shared repeat sequences. These sequences are suppressed by Cot-1 DNA during prehybridisation, but it reduces the specific signal intensity while incomplete suppression can result in non-specific signals. Finally, some of the clones might still be incorrectly assigned on the physical map.

In our study, we did not experience any problems in the detection or interpretation of the size of the aberrations studied, using the 2500 clone array despite the fact that 11 clones failed to detect the duplication or deletion, since they all had a size of at least 1 Mb. For the detection of small chromosome aberrations (<1 Mb), the array needs further improvement, by replacing the unstable clones and by adding more clones.

In the very near future, all elements of BAC libraries will be completely characterised and mapped, a fact that will further improve the reliability of BAC micro-arrays. 32 000 selected human BAC clones containing the draft sequence of the human genome can be ordered (http://bacpac.chori.org/phumanMinSet.htm) to manufacture a micro-array that fully covers the genome with a theoretical resolution that is only limited by the size of the BACs. When BAC arrays are introduced in diagnostic laboratories, genetic changes that were previously unknown will be detected and new genomic imbalances and syndromes will be described. In addition, normal variants will probably also be found, as was the case when subtelomeric screening was widely performed.31

ACKNOWLEDGEMENTS
We thank Xiang Guo for his statistical expertise, Isabel Tapia Páez for the chromosome 22 specific array data and the Laboratory of DNA and Protein Microarray technology, Van Andel Research Institute, for providing the cDNA arrays. We also want to thank Resources for Molecular Cytogenetics, Bari, the Wellington Trust Sanger Institute and BACPAC Resource Center, Children’s Hospital Oakland Research Institute for providing BAC and PAC clones.

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This work was supported by funds from the Swedish Medical Research Council, Frimurarna Barnahuset Foundation, and the Ronald McDonald Child Foundation.

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Received 3 September 2003
In revised form 16 September 2003
Accepted 25 September 2003

REFERENCES
13 Joyce CA, Dennis NR, Cooper S, et al. Subtelomeric rearrangements: results from a study of selected and unselected probands with idiopathic mental retardation and control individuals by using high-resolution G-banding and FISH. Hum Genet 2001;109:480–51.

Standardised measures of retinal function pinpoint genotype in X linked congenital stationary night blindness

Eye specialists using standardised electrophysiological criteria for the first time have suggested that these are more reliable indicators of genotype in X linked congenital stationary night blindness (CSNBX) than those used formerly.

Just three measures separated 20 affected males from 11 British families with CSNBX, according to whether they had mutations in NYX or CACNAIF genes. NYX mutations resulted in absent scotopic oscillating potentials, CACNAIF mutations in subnormal OFF response, and each mutation had different wave forms and amplitudes in 30 Hz flicker tests. Other indicators of eye function were not specific enough.

Three of the families had CACNAIF mutations, each with a different mutation, one of which was a novel nonsense mutation in exon 7. The others were a nonsense mutation in exon 24 and a base pair (bp) deletion in exon 9. All were expected to produce a truncated protein product. The eight remaining families had five NYX mutations: a splicing mutation; a missense and a nonsense mutation predicting truncated protein product; a 15 bp in frame deletion; and a 335 bp deletion.

Affected males were prospectively clinically evaluated for visual disorders and by psychophysiological and electrophysiological testing in parallel with genotyping.

Until now, non-standardised testing has subdivided CSNBX phenotype into “complete” and “incomplete” forms, apparently associated with NYX and CACNAIF mutations, respectively. However this distinction has proved unsatisfactory, in the light of reports of a functional rod pathway in patients with NYX mutations and of complete and incomplete forms of the condition in the same family.

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