New array approaches to explore single cells genomes

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Microarray analysis enables the genome-wide detection of copy number variations and the investigation of chromosomal instability. Whereas array techniques have been well established for the analysis of unamplified DNA derived from many cells, it has been more challenging to enable the accurate analysis of single cell genomes. In this review, we provide an overview of single cell DNA amplification techniques, the different array approaches, and discuss their potential applications to study human embryos.

Keywords: single cell, array analysis, BAC array, oligoarray, SNP array, chromosomal instability

INTRODUCTION
Chromosomal abnormalities are inherent to human in vitro fertilized (IVF) preimplantation embryos. FISH studies on normally developing, good quality cleavage stage embryos from IVF patients have shown that 30–65% are aneuploid in at least one cell (Iwarsson et al., 1999; Munne et al., 2004). A similar proportion of aneuploidies was detected in embryos derived from normal fertile couples (Rubio et al., 2003; Munne et al., 2004; Baart et al., 2006). However, aneuploid numbers of locus-specific FISH probe signals were in general interpreted as whole-chromosome imbalances thereby neglecting the possibility of structural chromosomal aberrations. It was with the development of metaphase comparative genomic hybridization (CGH) using DOP-amplified single cell DNA that the extent of whole-chromosome imbalances could be probed genome-wide. For the first time also segmental chromosomal imbalances were reported in approximately 7–32% of the embryos (Voulaire et al., 2000, 2002; Wells and Delhanty, 2000; Wilton, 2005; Daphnis et al., 2008; Rius et al., 2011). Mosaicism for whole-chromosome aneuploidies was detected in up to 75% of human cleavage stage embryos (Wells and Delhanty, 2000; Voulaire et al., 2002). With the advent of array CGH, the resolution within a single cell, whole genome amplification (WGA) is required. WGA approaches can be subdivided into PCR- or non-PCR-based (isothermal) methods.

WHOLE GENOME AMPLIFICATION
One cell from an embryo contains approximately 6 pg of DNA. Array methodologies require hundreds of nanograms of input DNA. Hence, to determine genome-wide copy number variation within a single cell, whole genome amplification (WGA) is required. WGA approaches can be subdivided into PCR- or non-PCR-based (isothermal) methods.

A number of PCR-based WGA methods have been developed including degenerate oligonucleotide primed PCR (DOP-PCR; Fiegler et al., 2003), primer extension PCR (PEP; Zhang et al., 1992; Sermon et al., 1996), and ligation-mediated PCR (Saunders et al., 1989). All those methods suffer from random amplification artifacts and incomplete coverage of loci which may result in the drop out (ADO) or preferential amplification (PA) of one of both alleles (Spits and Sermon, 2009). Recently, a new generation of PCR-based methods has been developed. Genomplex (Sigma-Aldrich, St. Louis, MO 63103, USA) and Picoplex/Sureplex (Rubicon Genomics Inc., MI 48108, USA/BlueGnome Ltd., Mill Court, Great Shelford, Cambridge, UK) kits are based upon a semi-random, non-enzymatic fragmentation of genomic DNA followed by the addition of specific adaptor sequences to both ends, forming an in vitro molecular library that can be amplified by PCR utilizing flanking universal priming sites. The size of the DNA fragments ranges from 100 to 1000 bp, with a median size of 400 bp. Based on the company brochures, an ADO rate of 10% (Picoplex) to 30% (Genomplex) can be expected, which is an improvement over previous PCR-based methods.

Multiple displacement amplification (MDA) is a non-PCR-based isothermal reaction using bacteriophage ϕ29 DNA polymerase.
polymerase (Repli-G, Qiagen Inc., Valencia, CA 91355, USA; GenomiPhi, GE Healthcare, Roosendaal, The Netherlands). MDA is based on a strand displacement amplification at constant temperature which results in DNA products of high molecular weight (up to 70 kb; Blanco et al., 1989; Spits et al., 2006a). The ADO rate of the single cell amplified loci varies between 0 and 60% (25.8% on average), while the 25% PA rate was found much more stable between individual cells (Spits et al., 2006a,b).

In conclusion, a variety of single cell WGA methods are commercially available but none is producing a linear representation of the original single cell genome. Furthermore the obtained array results may depend significantly on the single cell WGA method used (see below). ADO or the random loss of alleles can result in false-positive copy number calls as well as the over- or under-amplification of certain loci in the genome. These amplification artifacts complicate reliable genetic variant deduction from the signals obtained with the single cell WGA-DNA on the microarray platform and require specialized interpretation algorithms.

**BAC BASED ARRAY CGH**

Array comparative genomic hybridization (aCGH) enables the detection of gains or losses of DNA in all 24 chromosomes. DNA fragments with known positions in the genome are immobilized on a glass slide. Subsequently, DNA from a test and reference sample are differentially labeled and hybridized onto the slides. Copy number changes are determined by differences in hybridization intensities between test and reference DNA. The resolution of the array depends on the size of the probes and the distance between the probes in the genome. With bacterial artificial chromosomal (BAC) based aCGH, BAC DNA fragments are immobilized on a glass surface. Each clone contains a known chromosomal locus of about 100–300 kb (Oostlander et al., 2004). Besides the homemade BAC arrays provided by several laboratories (Le Caignec et al., 2006; Fiegler et al., 2007) at least two BAC arrays are currently commercially available: (1) the constitutional Chip 4.0 (Perkin Elmer) is comprised of approximately 5000 BAC clones and thereby a theoretical resolution of ~600 kb is achieved. (2) The 24Sure BAC array (Blue Gnome Ltd., Mill Court, Great Shelford, Cambridge, UK) contains 3000 clones spaced 1 Mb apart across the genome. Both commercial platforms have the potential to enable the detection of whole-chromosome aneuploidies in single cells in a 24-h protocol as required in preimplantation genetic testing and provide custom software tools for copy number variation calling.

**VALIDATION OF BAC ARRAYS**

Le Caignec et al. (2006) validated a strategy that combined GenomiPhi MDA amplification and a home-made 1 Mb resolution BAC array for single cell analysis. By averaging all intensity ratios per chromosome across multiple single cell experiments the chromosome-specific threshold for variation was determined. Subsequently, they demonstrated the accurate detection of trisomy 13, 18, 21 and monosomy X in different aneuploid fibroblast lines by chromosome average signals that surpass the chromosome-specific threshold (Le Caignec et al., 2006). By applying an optimized version of the amplification method (GenomiPhiV2) and a new algorithm the resolution to detect de novo imbalances was improved to 10 Mb, when combined with SNP-array technology (Vanneste et al., 2009a; Konings et al., 2012). The mixture model that was used not only allowed the calculation of BAC-probe specific copy number probabilities, but also provided a quality assessment allowing the exclusion of un-interpretable single cell WGA-samples (Vanneste et al., 2009a; Konings et al., 2012). Fiegler et al. (2007) developed a home-made tilling BAC array (26,574 clones) which enabled both the detection of aneuploidies and a de novo deletion of 8.3 Mb in GenomePlex amplified HCT116 single cells. To detect segmental imbalances, the average intensity ratio across 10 clones was calculated for each chromosome and plotted against the midpoint position of the 10 clones used for analysis. Each averaged data point was then further normalized. Substantial gains and losses comprised at least three consecutive data points 1.5 times above or below the estimate of the experimental variability. Cells from a renal carcinoma, a colorectal tumor, a trisomy 21, and from a Prader–Willi microdeletion cell line were used for validation. A false-negative rate of 3.0% was reported, while the false-positive rate was estimated between 2 and 3% (Fiegler et al., 2007). Gutierrez-Mateo et al. (2011) used the 24Sure BAC array (Blue Gnome Ltd., Mill Court, Great Shelford, Cambridge, UK) following Sureplex amplification for cleavage stage embryo genome copy number profiling. However, a thorough validation is lacking. Sensitivity and specificity of single cell DNA copy number typing must be evaluated using a set of cell lines with known DNA imbalances as exemplified by Fiegler et al. (2006), Vermeesch et al. (2007), Vanneste et al. (2009a), and Konings et al. (2012).

**CLINICAL USE OF BAC ARRAYS**

The first use of BAC aCGH (following GenomiPhiV2 WGA amplification) in PGD was for the selection of unbalanced chromosomes in embryos derived from a patient carrying a complex chromosomal rearrangement (Vanneste et al., 2011). Based on the analysis of 16 embryos, the sensitivity was 100%, while the specificity was 89% (Vanneste et al., 2011). At the same time, Alfarawati et al. (2011) described the first clinical application of BAC array-based screening applied to polar bodies, blastomeres, and trophectoderm cells from patients carrying chromosome rearrangements. They simultaneously screened for unbalanced translocation derivatives and aneuploidy of all 24 chromosomes. They used the Sureplex WGA together with the 24sure Version 2.0 or Cytochip Version 3.0 BlueGnome arrays (BlueGnome Ltd., Mill Court, Great Shelford, Cambridge, UK). A total of 16 couples received genetic diagnosis of their embryos in 20 cycles (Alfarawati et al., 2011). The reported diagnostic efficiency was 91.5% (Alfarawati et al., 2011). Fiorentino et al. (2011) used the same set-up and technology in cleavage stage embryos. Finally, Fishel et al. (2010) reported the birth of the first baby after genome-wide aneuploidy screening using single cell microarray technology and polar body biopsy. Overall, those studies lack validation (Fishel et al., 2010; Alfarawati et al., 2011; Fiorentino et al., 2011).

**OLIGONUCLEOTIDE ARRAYS**

The oligonucleotide aCGH procedure is similar to that of BAC arrays. The first arrays contained 44,000 (Agilent Technologies, Inc., Santa Clara, CA, USA) to 72,000 oligo probes (Nimblegen, Roche NimbleGen Systems, Reykjavik, Iceland), while the latest contain 1–2.1 million targets respectively. Oligonucleotide
probes contain between 45 and 85 nucleotides. The higher probe density allows a higher resolution for copy number evaluation. Moreover, oligonucleotide array construction tends to have a better reproducibility and less batch-to-batch variation compared to BAC arrays (Shearer et al., 2007). The main disadvantage for the detection of (single cell) chromosomal imbalances is the significant noise that becomes more obvious due to the random WGA artifacts and the higher probe density (Geigl et al., 2009).

VALIDATION OF OLIGO ARRAYS
Geigl et al. (2009) performed GenomePlex amplified single cell analyses on high-density oligo tiling arrays including the 380K (chromosome 22), the 2.1-M (Whole Genome) Nimblegen (Roche NimbleGen Systems, Reykjavik, Iceland), and the 240K (Chromosome 22) Agilent (Agilent Technologies, Inc., Santa Clara, CA, USA) arrays. These array analyses revealed that low amounts of template DNA do not result in a completely unbiased WGA but that stochastic amplification artifacts cause significant noise (Geigl et al., 2009). The platforms were validated using different cell lines and a new algorithm enabling the identification of small gains and losses in noisy ratio profiles was developed. They showed imbalances as small as 3.0 Mb could be accurately detected using high resolution oligo arrays and concluded that probe density may have an important impact on the resolution limits (Geigl et al., 2009). To enhance the resolution further and reduce biases in the single cell amplification, we developed a single channel based normalization method to preprocess Agilent 244K human microarray data (Cheng et al., 2011).

CLINICAL USE OF OLIGO ARRAYS
Hellani et al. (2008) reported the birth of the first baby after genome-wide aneuploidy screening of cleavage stage blastomeres using single cell oligo aCGH. Two blastomeres per biopsied embryo derived from eight couples suffering recurrent IVF failures were MDA amplified and analyzed on the Agilent Human Genome CGH 44B Oligo Microarray: The DNA from two blastomeres was pooled allowing for more reproducible data [data not shown (Hellani et al., 2008)]. Overall validation of the platform was limited (Hellani et al., 2008).

SNP ARRAYS
A different type of oligonucleotide arrays are SNP arrays targeting hundreds of thousands of SNPs. Single cell SNP-array analyses can detect copy number variations and disclose the genotype of the cell (Iwamoto et al., 2007; Vanneste et al., 2009a). Affymetrix (Affymetrix Inc., Santa Clara, CA, USA) and Illumina (San Diego, CA, USA) provide SNP arrays which differ by the number of probes and the SNP typing chemistry (such as the amplification technique, the number of color channels, the labeling procedure, and the time of the procedure). At first Affymetrix arrays targeted only 10,000 SNPs, but in their more recent product this further improved to 900K SNPs supplemented with 900K copy number probes (SNP 6.0 Array). Affymetrix arrays (Affymetrix Inc., Santa Clara, CA, USA) are based on a single color assay which is composed of multiple allele-specific probes that are complementary to SNP loci present in a reduced fraction of the genome. The reduced representation of the genome is achieved by digestion, adapter ligation, and PCR amplification. Following PCR-based amplification and fragmentation, the DNA is incorporated with biotin labeled nucleotides and hybridized onto the array. Following wash and scan, the signal intensity is quantified, and compared to intensities of normal individuals (such as HapMap individuals) in order to analyze the copy number. The total time for performing the assay is 4 days (Affymetrix Inc., Santa Clara, CA, USA)1. Illumina arrays (San Diego, CA, USA) are based on direct hybridization of whole genome-amplified genomic DNA (via a rolling circle replication mechanism) to a bead array of 50 mer locus-specific probes. These probes end one nucleotide before the SNP. Following hybridization each SNP is scored by a single base extension assay using differently labeled nucleotides. These labels are visualized by staining with an immunohistochemistry assay. For Illumina (San Diego, CA, USA) the first platform contained 317,000 probes and the newest 1 million SNP targets. The total time for performing the assay is 3 days (Illumina San Diego, CA, USA)2, however Johnson et al. (2010) published a 24-h protocol.

VALIDATION OF SNP ARRAYS
Iwamoto and co-workers were the first to report single cell SNP-array analyses for copy number detection using Affymetrix technology. They showed that variability in single cell amplification bias affected the genotype and copy number analysis of the cell severely (Iwamoto et al., 2007). Discordant genotypes were mainly localized close to centromeres or telomeres (Iwamoto et al., 2007). Single cell test WGA-samples could be successfully corrected for insufficiently amplified DNA regions by using both non-WGA and single cell WGA products as a reference for SNP-copy number calling (Iwamoto et al., 2007). Treff et al. (2011a) compared the PCR-based GenomePlex with the isothermal GenomiPhi and Repli-g MDA–WGA reaction for amplification reliability, fidelity and accuracy using the 250K NspI Affymetrix SNP microarray. The average genotyping coverage was 74% for GenomiPhi and 78% for Genomempelex, which where both significantly lower than the 88% obtained using Repli-g (Treff et al., 2011a). In our laboratory, an average coverage of 72% via the GenomiPhiV2 method was measured on the 250K NspI Affymetrix SNP (Vanneste et al., 2009a). Significant differences in the accuracy and copy number assignment exist amongst different single cell WGA methods (Treff et al., 2010, 2011a). Using the CNAT software (Affymetrix, Santa Clara, CA, USA) for single cell SNP-copy number typing, 62% of the SNPs had a match with the copy number profile of the genomic DNA sample extracted from many cells when the GenomiPhi method for single cell WGA was used. Copy number concordances of 95 and 99% were reached on the same SNP-platform when single cell genomes were amplified with Repli-G (Qiagen Inc., Valencia, CA 91355, USA) and Genomemplex (Sigma-Aldrich, St. Louis, MO 63103, USA) respectively (Treff et al., 2011a). Johnson et al. (2010) developed a new method for PGS, termed “parental support” which uses Illumina SNP microarray (CytoSNP-12 chips, San Diego, CA, USA) measurements from parental DNA to “clean” single cell microarray measurements on embryonic cells and explicitly computes confidence in each copy number call. A false-negative rate of 2.1% and a false-positive...
rate of 3.9% for copy number detection were estimated (Johnson et al., 2010). To detect segmental copy number aberrations their algorithm was applied to segments spanning one-fifth of a chromosome, limiting the ability to detect segmental abnormalities (Johnson et al., 2010). Handyside et al. (2010) developed a method enabling genome-wide linkage-based inheritance analysis of a broad range of genetic abnormalities including structural chromosomal abnormalities, DNA copy number variants as well as single gene defects identified as pre-existing in one or both parents using the Illumina Human CNV370 Infinium-II Quad and duo. They demonstrated that karyomapping is possible at the single cell level following MDA (Repli-G, Qiagen Inc., Valencia, CA 91355, USA) if only heterozygous informative loci are used (Handyside et al., 2010). We developed a novel algorithm that determines the allelic origin of (aberrant) loci by identifying and visualizing SNPs with a Mendelian error in a parent-specific manner at the genome-wide level (Voet et al., 2011). Therefore, we genotyped both parents and the single blastomeres using Affymetrix 250K SNP arrays (Voet et al., 2011).

CLINICAL USE OF SNP ARRAYS
Schoolcraft et al. (2010) presented the first clinical application of blastocyst trophectoderm biopsy followed by SNP microarray based 24 chromosome aneuploidy screening, applied to 132 cases with advanced maternal age, recurrent pregnancy loss, or recurrent implantation failure. They achieved a clinical pregnancy rate of 77.8% (Schoolcraft et al., 2010). A case study by Brezina et al. (2011) reported the birth of a healthy child after combination of PGD using DNA sequencing, PCR linkage and the accuracy of single cell analyses are likely to come from different array approaches. The accuracy has been established for different array platforms and the technologies are widely used to study human embryos. Nevertheless, the resolution remains well below the resolution obtained when using unamplified DNA on the same platforms. Further improvements in the resolution as well as the accuracy of single cell analyses are likely to come from further improvements in WGA methods, array platform design as well as algorithmic improvements.

CONCLUSION AND FUTURE PROSPECTIVES
The genome-wide detection of aneuploidies and segmental chromosomal imbalances in single cells can be accurately achieved by different array approaches. The accuracy has been established for different array platforms and the technologies are widely used to study human embryos. Nevertheless, the resolution remains well below the resolution obtained when using unamplified DNA on the same platforms. Further improvements in the resolution as well as the accuracy of single cell analyses are likely to come from further improvements in WGA methods, array platform design as well as algorithmic improvements.

Besides fundamental research questions, the main application of single cell array analysis in the clinic is PGD. PGD is typically performed on embryos from carriers of monogenic disorders or carriers of chromosomal rearrangements affecting the fertility (such as translocations or marker chromosomes). Currently chromosomal abnormalities are selected against using FISH, while monogenic disorders are diagnosed via PCR. Recently, single cell array techniques have been clinically implemented for PGD as well (Fishel et al., 2010; Alfarawati et al., 2011; Fiorentino et al., 2011; Treff et al., 2011b; Vanneste et al., 2011). Single cell microarray analysis could be envisioned as a novel generic diagnostic tool for PGD because it: (1) covers all 24 chromosomes, (2) makes the interpretation of the results more objective, and (3) excludes the family specific preparation compared to the current techniques.

In addition to PGD, PGS may be the prime clinical application. PGS is based on the hypothesis that selection of chromosomally normal embryos for uterine transfer can increase the live birth rate and decrease the spontaneous abortion rate per embryo transferred. Single cell array analysis allows genome-wide screening of de novo whole-chromosome and segmental imbalances in 24 h and can be performed on polar bodies, blastomeres, or trophectoderm cells. The widespread acceptance for PGS, however, will rely on the clinical validation, i.e., that this screening leads to increased baby-take-home-rates (Vanneste et al., 2009b; Geraedts et al., 2010; Harper et al., 2010a). The application of this technology is rapidly increasing (Harper et al., 2010b).

Whereas single cell BAC- and oligoarrays can be used to detect chromosomal imbalances, single cell SNP arrays may be used to find Mendelian disorders. The ability to genotype and haplotype single cells using SNP arrays has opened new avenues of single cell research. First, determining the haplotypes can strengthen the power and thus increase the resolution of single cell CNV detection (Vanneste et al., 2009a; Voet et al., 2011). Extracting the haplotypes enables the analysis of cross-over sites and the identification of copy number neutral abnormalities such as segmental uniparental disomy (UPD) as well as long contiguous stretches of homozygous SNPs (Kotzot, 2008). When parental genotypes are known as well, transmission of haplotypes can be measured. Simple haplotyping of SNPs surrounding and embedded in disease-causing genes may thus allow the selective transfer of genetically and chromosomally normal embryos for patients undergoing IVF with PGD for monogenic diseases (Handyside et al., 2010).

To understand the chromosomal instability in human embryos it may well be necessary to go beyond the inherent limitations of array CGH. With the strongly reduced costs of whole genome sequencing, those sequencing technologies offer the promise to further increase the resolution and provide the ultimate genomic architectural view of single cells for basic genome research purposes. It seems likely that the picture of human embryonic instability will become more complete in the years to come.

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