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Identification of Balanced Chromosomal Rearrangements Previously Unknown Among Participants in the 1000 Genomes Project: Implications for Interpretation of Structural Variation in Genomes and the Future of Clinical Cytogenetics

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Supplemental Data

Supplemental Data include four tables.

Conflict of interest

The authors declare no conflict of interest.

List of URLs

DGAP: http://dgap.harvard.edu

 $1000\ Genomes\ Project: http://www.internationalgenome.org/data/$

USCS (University of California, Santa Cruz) genome browser: https://genome.ucsc.edu/

ENCODE: https://www.encodeproject.org

DECIPHER (DatabasE of genomic variation and Phenotype in Humans using Ensembl Resources): https://decipher.sanger.ac.uk Integrative Genomics Viewer: http://software.broadinstitute.org/software/igv/

CyDAS (Cytogenetic Data Analysis System): http://www.cydas.org/OnlineAnalysis/

GTEx: http://www.gtexportal.org/home/ Primer3 Web: http://primer3.ut.ee/

NCBI (National Center for Biotechnology Information) Primer-Blast: http://www.ncbi.nlm.nih.gov/tools/primer-blast/

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Abstract

Purpose—Recent studies demonstrate that whole-genome sequencing (WGS) enables detection of cryptic rearrangements in apparently balanced chromosomal rearrangements (also known as balanced chromosomal abnormalities, BCAs) previously identified by conventional cytogenetic methods. We aimed to assess our analytical tool for detecting BCAs in The 1000 Genomes Project without knowing affected bands.

Methods—The 1000 Genomes Project provides an unprecedented integrated map of structural variants in phenotypically normal subjects, but there is no information on potential inclusion of subjects with apparently BCAs akin to those traditionally detected in diagnostic cytogenetics laboratories. We applied our analytical tool to 1,166 genomes from the 1000 Genomes Project with sufficient physical coverage (8.25-fold).

Results—Our approach detected four reciprocal balanced translocations and four inversions ranging in size from 57.9 kb to 13.3 Mb, all of which were confirmed by cytogenetic methods and PCR studies. One of DNAs has a subtle translocation that is not readily identified by chromosome analysis due to similar banding patterns and size of exchanged segments, and another results in disruption of all transcripts of an OMIM gene.

Conclusions—Our study demonstrates the extension of utilizing low-coverage WGS for unbiased detection of BCAs including translocations and inversions previously unknown in the 1000 Genomes Project.

Keywords

balanced chromosomal rearrangement; the 1000 Genomes Project; G-banded chromosome analysis; low-pass whole-genome sequencing

Introduction

A balanced chromosomal rearrangement (or abnormality, BCA) is a type of chromosomal structural variant (SV) involving chromosomal rearrangements (e.g., translocations, inversions and insertions) without cytogenetically apparent gain or loss of chromatin. The incidence of balanced translocations has been estimated to range from 1/500 to 1/625 in the general population^{1–3} and the prevalence is well known to be increased in individuals with clinical anomalies^{4–7}. Based on the association of increased prevalence with abnormal

clinical phenotypes, studies of BCA such as in the Developmental Genome Anatomy Project (DGAP)^{5,6,8,9} among others^{7,10} have a high yield in identification of genetic disease due to gene disruption or dysregulation.

Current high-resolution methods (*i.e.*, chromosomal microarray analysis and whole exome sequencing) are generally insensitive to BCA that are unaccompanied by sizable genomic imbalances. Thus, detection of BCA relies on conventional cytogenetic methods (*i.e.*, Gbanded karyotyping), which are limited to microscopic resolution (~3–10 Mb). More recently, whole-genome sequencing (WGS) using paired-end analysis has enabled molecular delineation of the breakpoints of BCA at base-pair resolution but has been tested and validated only in DNA samples harboring previously recognized BCAs.

By utilizing WGS (mean 7.4-fold base-coverage) and orthogonal techniques (*i.e.*, long-read single-molecule sequencing), the 1000 Genomes Project establishes the most detailed catalogue of human genetic variation, which in turn can be used for association studies relating genetic variation to disease. It provides an unprecedented integrated map of SVs from 2,054 individuals, including copy-number variants, inversions (< 50 kb) and insertions ^{11,12}, and serves as an indispensable reference for geographic and functional studies of human genetic variation. However, no information was available on the frequency of balanced translocations or inversions (> 50 kb in size) in this resource of participants who were healthy at the time of enrollment. Our previous pilot study has shown the feasibility of detecting BCA with low-pass (or low-coverage) paired-end WGS in a blinded fashion ¹³. In the present study, we apply our analytical tool to WGS data released by the 1000 Genomes Project ¹¹.

Materials and Methods

WGS data from the 1000 Genomes Project

Alignment files from 2,504 presumably healthy individuals were downloaded from the 1000 Genomes Project. Assessment of data quality and further analysis was processed for each individual independently.

Minimum physical coverage requirement used in this study

As shown in our previous study¹³, the minimum read-pair count was used to avoid false negative detection of BCA. This minimum number of read-pairs in a small-insert library (400 to 600 bp) was estimated as 120 million (50 bp), which is equivalent to 4-fold base coverage from whole-genome sequencing. However, read length (35 to 100 bp) and insert size (200 to 600 bp, Figure 1A) were varied among samples from the 1000 Genomes Project¹¹. Therefore, physical coverage¹⁴ was used as the required selection criteria instead of the number of read-pairs (Figure 1B).

We defined a chimeric read-pair if two ends aligned to different chromosomes (interchromosomal) or to the same chromosome (intrachromosomal) with an aligned distance larger than 10 kb^{13,15}. Physical coverage was estimated by counting the aligned distances from the non-chimeric and uniquely mapped read-pairs¹³. In the present study, the minimum physical coverage of 8.25-fold, estimated based on 90 million read-pairs (data

from our previous pilot study)¹³, was set to maximize inclusion of 1,166 out of 2,504 samples (Supplementary Table 1). This was based on: (1) only 616 out of 2,504 samples available for this study with 11-fold physical coverage (estimated based on 120 million readpairs), and (2) an increase in the false negative detection rate in our previous study from 11.1% (1/9) with 90 million read-pairs to 33.3% (3/9) with only 60 million read-pairs¹³.

Data quality control and BCA detection

We filtered out low-quality reads (4% of mismatch rates) and extracted uniquely aligned reads in both ends for further analysis. Detection of chromosomal rearrangement is based on a four-step procedure described in our previous study¹³: briefly, (1) Event clustering: chimeric read-pairs were clustered by sorting the aligned coordinates (GRCh37/hg19) and any two read-pairs were considered to represent two distinct events if they were separated by a distance of >10 kb; (2) Systematic error filtering: Each event was filtered against a control dataset, which was built up by using the events from all the 2,504 samples, and a false positive was filtered out if it was identified in more >5% subjects; (3) Random error filtering: Event was filtered with a cluster property matrix (*i.e.*, supporting read-pair amount and the average number of mismatches) with the reported parameters; and (4) Aligned orientations: each event was filtered based on q/p arm genetic exchange (joining type). As some of the samples were with short read lengths (*i.e.*, 35 bp), we further used Sanger sequencing results to fine map the ligated sequences ate the breakpoints.

Chromosome analysis and FISH validation

Epstein-Barr virus (EBV)-transformed B lymphoblastoid (EBV-B) cell lines were obtained from the Coriell Institute (Camden, NJ) for validation. G-banded chromosome analysis was performed using standard protocols for more than 100 cells in each EBV-B cell line¹⁶. Fluorescence in situ hybridization (FISH) was performed for NA18612 using standard procedures with BAC clones labeled by nick translation with SpectrumOrange or SpectrumRed, SpectrumGreen dUTP (Abbott Molecular, Des Plaines, IL)^{16,17}. BAC clones were selected from the UCSC Genome Browser.

Molecular validation of balanced rearrangements

For samples with translocations and available EBV-B cell lines, genomic DNA was extracted using a commercial DNA extraction kit (Puregene; Qiagen, Hilden, Germany). For samples identified with submicroscopic inversions, DNAs were obtained from the Coriell Institute (Camden). Each DNA was quantified subsequently with the Qubit dsDNA HS Assay Kit (Invitrogen, Life Technologies, Waltham, MA) for DNA quality measurement.

Genomic reference sequences (GRCh37/hg19) at a 1 kb distance from each putative breakpoint region (both upstream and downstream) were used for primer design with Primer3 Web and NCBI Primer-Blast (Supplementary Table 2). PCR amplification was performed simultaneously in cases and control (DNA from YH, a well-characterized normal EBV-B cell line¹⁸). PCR products were sequenced by Sanger sequencing on an ABI 3730 machine (Applied Biosystems, Thermo Fisher Scientific Inc, Wilmington, DE)^{8,13,19} and sequencing results were aligned with BLAT for further confirmation of the balanced rearrangement and for mapping breakpoints at single nucleotide level.

RNA preparation, library construction and sequencing

Total RNA was extracted from each EBV-B cell line with a balanced translocation using TRIzol Reagent (Invitrogen) according to the manufacturer's instructions, and subsequently treated with DNase I (Invitrogen)²⁰. For each RNA sample, purity was evaluated with a Nano-Photometer spectrophotometer (Implen, Westlake Village, CA), concentration measured in a Qubit 2.0 Fluorometer (Life Technologies), and RNA integrity verified using an Agilent 2100 BioAnalyzer (Agilent Technologies, Santa Clara, CA).

For library construction, mRNA enrichment was performed with Oligo(dT)25 Dynabeads (Thermo Fisher Scientific) twice and purification was carried out with the Dynabeads® mRNA Purification Kit (Invitrogen, No. 61006). The eluted mRNA was fragmented with Fragmentation Buffer Mix at 94°C for 10 min. Reverse transcription (RT) was performed with RT Buffer Mix and RT Enzyme Mix followed by double strand cDNA (dscDNA) synthesis with Second Strand Buffer Mix and Second Strand Enzyme Mix. End repair, adaptor (with barcode) ligation and PCR amplification were performed after dscDNA purification. Then, the purified double stranded PCR products were heat denatured to single strand and circularized with Splint Oligo Mix and Ligation Enzyme. The single strand circle DNA (ssCirc DNA) library was rolling circle amplified for constructing the DNA nanoball (DNB), which was substantially loaded into a patterned nanoarray. Paired-end sequencing with 50 bp in each end (PE50) was carried out in a BGISeq-500 platform (BGI, Wuhan, China)²¹.

RNA-seq data analysis

Paired-reads that passed standard quality control ^{13,15} were simultaneously aligned to the human genome (GRCh37/hg19) using HISAT (Hierarchical Indexing for Spliced Alignment of Transcripts)²², and aligned to human transcriptome (RefSeq) via Bowtie²³. One base-pair mismatch was set in each alignment. Paired-end aligned reads were used for further analysis. Alignment files were transformed into Pileup files for determination of coverage with Samtools (mpileup). Expression of each gene in each sample was determined based on alignment files from the human transcriptome (RefSeq).

Gene expression of each sample was compared to data reported for 13 EBV-B controls present in the Genotype-Tissue Expression (GTEx) project²⁴.

Validation of cryptic deletions

Quantitative PCR (qPCR) was performed for validation of the two cryptic deletions. Genomic reference sequences (GRCh37/hg19) of each deleted region were used for primer design with Primer3 Web and NCBI Primer-Blast (Supplementary Table 2). Melting curve analysis was carried out for each pair of primers to ensure specificity of the PCR amplification, and the standard curve method was used to determine PCR efficiency (within a range from 95% to 105%).

Each reaction was performed in quadruplicate in $10 \,\mu l$ of reaction mixture simultaneously in cases and control (DNA from YH EBV-B cell line¹³) on a StepOnePlus Real-Time PCR System (Applied Biosystems) with SYBR Premix Ex Taq Tli RNaseH Plus (Takara

Biotechnology Co., Ltd., Dalian, Liaoning, China) with the default setting of the reaction condition. The number of copies in each sample was determined by using the Ct method that compared the Ct (cycle threshold) in case to control²⁵. Two independent pairs of primers (Supplementary Table 2) were used in quintuplicate for validation of each deletion.

Accession Number

The accession number for the RNA-seq data reported in this paper is GSE94043 (NCBI Gene Expression Omnibus).

Code availability

All the programs relevant to this pipeline are available at https://sourceforge.net/projects/bca-analysis/files/BCA.tar.gz/download.

Results

We assessed 1,166 samples with at least 8.25-fold physical coverage (Supplementary Table 1) using our reported approach with the same parameters 13. Four samples (HG02260, HG03729, NA18612 and NA20764) were identified to harbor balanced translocations (Figures 1A), and four samples (NA20759, HG04152, NA18959 and NA21133) were with inversions, the size of which ranged from 57.8 kb to 13.3 Mb (Figures 1A). Among the four cases with balanced translocations, two are female and two are male and they originate from different ethnic populations (Table 1)11. For the four cases with inversions, all are males and also originate from different ethnic populations (Table 2).

G-banded karyotypes were observed to be directly consistent with the WGS data for samples HG02260, HG03729, NA20764 and NA20759 (Figure 1B–E), and those of NA18612 were consistent but much less obvious (Figure 2A, described below). Sanger results confirmed each rearrangement in all eight samples with BCAs (Tables 1, 2, next-generation cytogenetic nomenclature²⁶ shown in Supplemental Table 3). Microhomology sequences were identified in eight of 16 breakpoints suggesting the rearrangements were mediated by microhomology-mediated end joining (MMEJ)²⁷ (Tables 1, 2). The remaining eight breakpoints represented non-homologous end joining (NHEJ)²⁸ (Tables 1, 2).

Subtle balanced translocation identified by WGS

Breakpoints of the t(16;17)(q23.1;q24.2) (NA18612, Figure 2B) were located in bands 16q23.1 and 17q24.2, representing translocated segments of 15.0 Mb and 16.2 Mb, respectively. Due to similarity in the G-banding pattern and size of the exchanged segments, chromosome analysis did not readily identify the translocation (Figure 2A). Therefore, metaphase FISH¹⁷ was performed using BAC probes (SpectrumOrange: RP11-7D23 at 16q24.3, SpectrumGreen: RP11-526M7 at 17q25.1 and SpectrumRed: RP11-135N5 at 17p13.3) in more than 100 cells confirming the t(16;17) (Figure 2C).

Gene disruptions by the breakpoints of balanced translocations

Among the four cell lines with balanced translocations, the eight breakpoints disrupted six genes (Table 1, Figures 2D, 2E, 3A, and 3B), four of which resulted in disruption of all

transcripts in the derivative chromosomes of the breakpoints. In contrast, none of the breakpoints from the four cases with inversions disrupted any gene (Table 2).

The breakpoint in seq[GRCh37/hg19] 16q23.1(75,336,134_75,336,138) (NA18612, Figure 2D) disrupts the gene encoding craniofacial development protein 1 (*CFDP1*, NM_006324), resulting in aberrant splicing of intron 6 and absence of expression of exon 7. This disruption is supported by observation of RNA-seq reads mapping in the non-exonic region (Figure 2D) and decreased expression of exon 7 (Figure 2E). Although *CFDP1* has been reported to be necessary for cell survival and differentiation during tooth morphogenesis in organ culture²⁹, it is unlikely to be haploinsufficient [Haploinsufficiency score (HI)=14.9%]³⁰.

In contrast, the seq[GRCh37/hg19] 14q31.1(79,839,173_79,839,174) breakpoint of 46,XX,t(9;14)(q34.2;q31.1) (HG02260) (Figure 3A) disrupts all transcripts of neurexin 3 (*NRXN3*), which is likely to be haploinsufficient (HI=0.3%)³⁰. However, expression of *NRXN3* was not detectable among any of the EBV-B cell lines including cases and controls (Figure 3C)²⁴.

Cryptic deletions

The 3q24 breakpoint of 46,XY,t(3;17)(q24;p13.3) (HG03729) was found to include a 5.2 kb deletion, seq[GRCh37/hg19] 3q24(143,817,430_143,822,651)x1, while the 17p13.3 breakpoint has a 4.4 kb deletion, seq[GRCh37/hg19] 17p13.3(2,910,366_2,914,751)x1 (Table 1, Figure 3D and 3E). Neither deletion was reported previously 11 and both were confirmed by quantitative PCR (Figure 3D and 3E).

Positional effects

Previous studies show that genes in proximity to the breakpoints of a structural variant (*i.e.*, balanced translocation) may be mis-expressed, which is defined as a positional effect⁵. One mechanism for a positional effect is the disruption of topological associated domains (TADs) by the SV's breakpoints^{6,9,31}. Here, we used boundaries predicted from the human IMR90 fibroblast cell line (GRCh37/hg19)³¹ for our study, as TADs are highly conserved across different cell types and across species³².

Eight TADs were disrupted by the breakpoints from the four translocations. Thirty-four genes are located in these eight disrupted TADs, and expression was observed in 16 of these genes in normal EBV-B cell lines (Supplemental Table 4). However, mis-expression was not observed in any of these 34 genes from our RNA-seq data (Supplemental Table 4), even though two of these genes are predicted to be likely haploinsufficient (Supplemental Table 4). By using the published ChIP-seq data from Encyclopedia of DNA Elements (ENCODE)³³ in EBV-B cell line GM12878, 22 of the 34 genes have a candidate promoter (indicated by H3K4Me3) near to a potential active regulatory element (indicated by H3K27Ac)^{31,33}. In addition, from the accessible chromatin landscape³⁴, 19 of the 34 genes have highly associated DNA I hypersensitive sites (DHSs) and each of them has at least one DHS located in the same partial TAD as the gene and the predicted promoter.

The breakpoint in seq[GRCh37/hg19] 17q24.2(64,953,078_64,953,079) is likely to be located between the *CACNG4* promoter (indicated by H3K4Me3) and its potential enhancer [indicated by H3K27Ac^{31,33} and DHSs³⁴, Figure 3F] in a human embryonic stem cell line (H1-hESC)³³. These data suggest that the translocation would likely result in disruption of the interaction between the promoter and enhancer for *CACNG4*³² in this particular cell line. However, mis-expression of *CACNG4* was not observed in our RNA-seq data from EBV-B cell line (Figure 3G). Both the candidate promoter and enhancer for *CACNG4* were likely located downstream of the breakpoint³³ in EBV-B cell line GM12878 (control EBV-B cell line, Figure 3F).

Discussion

Balanced chromosomal abnormalities including translocations and inversions, known to cause reproductive problems and/or an abnormal phenotype, currently are mainly detected by G-banded chromosome analysis. However, subtle or cryptic BCAs are not detectable by current methods but may contribute to birth defects in offspring of the carriers due to unbalanced segregations³⁵. In the present study, by utilizing existing genomic data from the 1000 Genomes Project, we demonstrate the feasibility of using WGS in the detection of BCAs in samples without prior knowledge of their existence.

In the present study, we set a cutoff of 8.25-fold physical coverage to maximize the inclusion of 1,166 samples out of 2,504 based on the evaluation of the false negative rate in our previous study (11.1% with 90 million read-pairs with insert sizes ranging from 400 to 600 bp)¹³. The exclusion of more than half of all samples (n=1,338) is because of the smaller insert size generated (259.1±93.5 bp, Figure 1A); the number of non-chimeric and uniquely mapped read-pairs was 97.0±40.2 millions, although the base-coverage reached 7.4-fold on average. This indicates that better performance of detecting BCA can be achieved by using larger insert sizes to increase physical coverage, thus, increasing the number of supporting read-pairs for the potential BCAs.

The prevalence of reciprocal balanced translocations in this dataset is one in 291.5 (0.34%, 4/1,166), which is higher than the rate reported estimated by G-banded chromosome analysis ^{1–3}. This estimate may be biased due to the limited sample size (N=1,166). However, the reported incidences may be underestimated as cryptic or subtle rearrangements, such as that observed for 46,XY,t(16;17)(q23.1;q24.2) (NA18612) may not be readily identified by conventional G-banded chromosome analysis (Figure 3). Another explanation might be that the detection of rearrangements was based on WGS of EBV-B cell line-derived DNAs, which might have EBV-B specific genomic variants owing to the introduction of genomic instability by EBV infection or the conditions of cell culture³⁶. However, as the EBV-B specific genomic variants frequently exist as mosaics³⁶, giving the 100% consistency of more than 100 metaphases in each sample in the present study and the WGS data of these samples used for our analysis were generated from early batches of EBV-Bs in the 1000 Genomes Project, balanced translocations detected probably represent the true events in the subjects' peripheral blood samples. Nonetheless, our approach reports the true events existing in the tested EBV-B cell lines.

In addition to the detection of balanced translocations, the microscopic inversion and three submicroscopic inversions identified (Table 2) were each unique to a single subject among all 1,166 samples analyzed. One explanation for not identifying common or recurrent inversions is that they may be mediated by repetitive elements²⁸, for which sequencing with small-insert library might not be able to detect³⁷. Sequencing with mate-pair library (or large insert library) might be able to overcome such challenge and can also largely reduce the sequencing cost by reducing the read-pair amount required ^{13,19}. Nonetheless, the identification of both balanced translocations and inversions underscores the importance of using low-pass WGS for nucleotide level precision of chromosomal rearrangements in cytogenetic diagnoses, and brings the future of implementing sequencing a step closer as the first tier test.

Gene disruptions were observed in six out of the eight breakpoints in four cases with balanced translocations, and *NRXN3*, a likely haploinsufficient gene, was disrupted (HG02260). Heterozygous deletion of *NRXN3* is reported in autism spectrum disorder (ASD)³⁸. Although this participant in the 1000 Genomes Project is assumed to be healthy at the time of enrollment, a possible explanation for the absence of ASD in this presumably normal individual would be lack of penetrance³⁸ in the absence of a positive comprehensive medical assessment or some technical failure in the process.

Two cryptic deletions involving both breakpoints were identified in t(3;17)(q24;p13.3) (HG03729) and neither of them were reported previously ¹¹. Two possible reasons for missed detection in the previous study ¹¹ are: (1) only a limited number of reads mapping in these regions (Figure 3D and 3E) resulting in read-depth differences insufficiently sensitive for identification, and (2) absence of intra-chromosomal aligned read-pairs supporting these two deletions. Genomic imbalance commonly involves the breakpoint of balanced translocations and some of them are known to be pathogenic or likely pathogenic ⁶, thus, indicating the importance of identification.

No aberration in gene expression resulted from a positional effect, such as disruption of TADs as observed in our EBV-B cell line-derived RNA-seq data from four cases with balanced translocations. One explanation is that expression was only observed in 16 out of the 34 genes in normal EBV-B (Supplemental Table 4), and an effect of dysregulated lower expression cannot be detected for genes without detectable expression in the EBV-B cell lines²⁴. In addition, another reason would be the proximate interaction between promoter and enhancer: (1) 22 of the 34 genes have a candidate promoter near to a potential active regulatory element^{31,33}; and (2) 19 of the 34 genes have highly associated DHSs³⁴, and each of them has at least one DHS located in the same partial TAD as the gene and the predicted promoter, indicating some residual interactions remains between promoter and regulatory elements, thus, the disruption of TADs is likely insufficient to alter the gene expression. As data from RNA expression provides evidence for confirming potential effects attributed to a chromosomal rearrangement, it indicates the importance of combining RNA expression analysis with identification of BCAs based on DNA samples for clinical interpretation.

We observed a potential disruption of an interaction between the promoter and an enhancer for the 17q24.2 breakpoint (NA18612) in H1-hESC, which serves as a reference for disease

association prediction³¹. However, mis-expression of *CACNG4* was not observed in our RNA-seq data from the EBV-B cell line (Figure 3G). One explanation is that both the candidate promoter and enhancer for *CACNG4* are likely located downstream of the breakpoint³³ in the EBV-B cell lines (GM12878, Figure 3F). As another sample type from this subject is not obtainable for further validation, this argues the common usage of peripheral blood as a valuable sample type for disease studies beyond its simple availability.

Overall, this study is the first reported investigation utilizing low-pass WGS to explore detection of BCAs among samples from the 1000 Genome Project without prior knowledge of a chromosomal abnormality. In addition, disruption of gene, cryptic imbalances and potential disruption of promoter and enhancer interaction were observed in the four cases with balanced translocations, demonstrating the advantage of detecting the breakpoints in BCAs by molecular methods via paired-end sequencing and Sanger sequencing, and has important implications for a new dawn of improved diagnostics in clinical cytogenetics.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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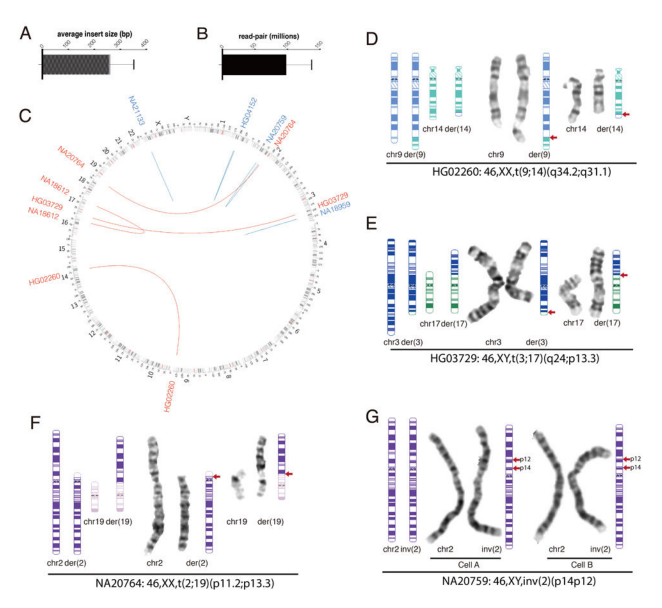


Figure 1. Spectrum of inter- and intra-chromosomal balanced rearrangements and cytogenetic validations

Distributions of the average insert sizes and read-pair amounts of 2,504 samples from the 1000 Genomes Project are shown in (**A**) and (**B**), respectively. Insert size and read-pair amounts were calculated based on non-chimeric and uniquely mapped read-pairs. (**C**) Spectrum of BCAs. Balanced translocations are indicated with red lines and the corresponding sample IDs are shown in red font in each affected chromosome in the outmost circle. Inversions are indicated in blue lines and sample IDs are shown in blue font. Chromosomal nucleotide positions and bands are shown according to the UCSC Genome Viewer Table Browser. In figures (**D**), (**E**) and (**F**), validation of balanced translocations and inversion (**G**) by G-banded chromosome analysis are shown. Ideograms of the balanced rearrangements are shown on the left, while the karyogram images are to the right with the corresponding ideogram of the derivative chromosomes for reference. Breakpoint regions

are indicated with red arrows. Sample name and the International System for Human Cytogenomic Nomenclature (ISCN) description are shown below each.

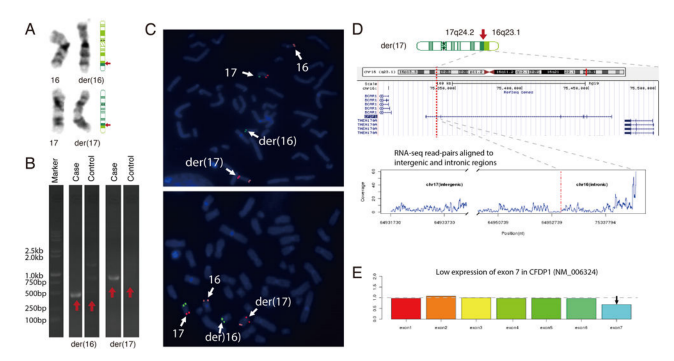


Figure 2. A subtle translocation t(16;17)(q23.1;q24.2) (NA18612) and the aberrant splicing of intron 6 of CFDPI (NM_006324)

(A) Validation from chromosome analysis. Ideograms of the derivative balanced translocation chromosomes are shown with the corresponding G-banded chromosome pairs. Breakpoint regions are indicated with red arrows. (B) PCR validation of the junction of DNA sequences from the two derivative chromosomes indicated by red arrows while absent in the negative control. (C) Validation from FISH. BAC probes are shown with the targeted bands (16q24.3 in SpectrumOrange, 17p13.3 in SpectrumRed and 17q25.1 in SpectrumGreen, respectively). Derivative chromosomes and normal chromosomes are designated with arrows. (**D**) In the der(17), the genomic location of anti-sense gene *CFDP1* (NM_006324) is shown with the breakpoint mapping in intron 6 (red dotted line). RNA-seq read-pairs align to the region (expressed as coverage) proximal to the breakpoint in seq[GRCh37/hg19] 16q23.1(75,336,134 75,336,138). It includes the intergenic region in 17q24.2 and the partial intron 6 of *CFDP1* (NM_006324) in 16q23.1 (two grey dotted lines), indicating the aberrant splicing of intron 6. (E) Transcript coverage was plotted with the paired-end aligned reads (RNA-seq). The coverage of each coordinate is divided by the average coverage in this transcript, and subsequently normalized coverage with the average coverage from the other three EBV-B cell lines from the 1000 Genomes Project. Black arrow indicates low expression in exon 7 that is the absence of exon 7 in the disrupted transcript.

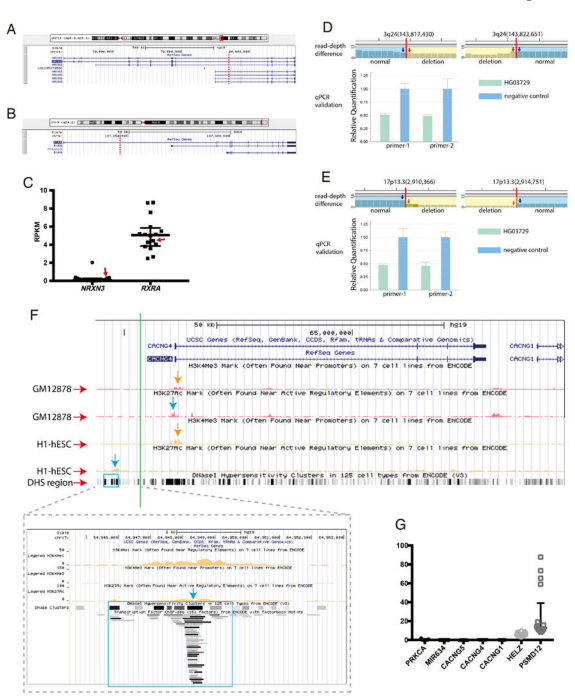


Figure 3. Gene disruption, cryptic deletions and potential disruption of interaction between promoter and enhancer by the breakpoints of balanced translocations Figures (A), (B) and (C) NRXN3 disruption in 46,XX,t(9;14)(q34.2;q31.1) (HG02260). (A)

and **(B)** Genomic locations of *NRXN3* and *RXRA* are shown with breakpoints indicated by red dotted lines. **(C)** *NRXN3* and *RXRA* expression for the four cases from the 1000 Genomes Project and for 13 reported EBV-B normal control cell lines (the GTEx project). Gene expression for *NRXN3* and *RXRA* in HG02260 are indicated with red arrows. Figures **(D)** and **(E)** cryptic deletions involved at the breakpoints in translocation 46,XY,t(3;17)

(q24;p13.3) (HG03729). Two cryptic deletions of seq[GRCh37/hg19] 3q24(143,817,430_143,822,651)×1 and seq[GRCh37/hg19] 17p13.3(2,910,366_2,914,751) ×1 were detected by read-depth difference algorithm and were further confirmed by quantitative PCR. The deleted regions are shown in a yellow background with a red arrow while the normal copy-ratio (diploid) is shown in a blue background with a blue arrow. Two independent pairs of primers (Supplementary Table 2) were used to perform qPCR in quintuplicate for validation of each deletion. The bars in cyan show the relative quantification of HG03729, while the bars in blue indicate the negative control. Figures (F) and (G) potential disruption of interaction between promoter and enhancer from rearrangement in 46,XY,t(16;17)(q23.1;q24.2) (NA18612) in H1-hESC. (F) Genes and the ChIP-seq data from the ENCODE Project are shown in terms of the genomic location. Each cell line with the ChIP-seq data (i.e., H3K4Me3 and H3K27Ac)³³ is labeled with a red arrow. Breakpoint in seq[GRCh37/hg19] 17q24.2(64,953,078_64,953,079) is shown by a green vertical line, while the candidate promoters and enhancers are indicated with orange and blue arrows, respectively. The region of potential enhancer in H1-hESC is highlighted in DNase I Hypersensitivity Clusters³⁴ in a blue rectangle (DHS region). The figure below is zoomed in on the potential enhancer region in H1-hESC. Enrichment of H3K4Me1 and absence of H3K4Me3 support a potential active enhancer in this region^{33,39}, while enrichment of DNA-binding sequence motifs also indicates the candidate region of the interaction for regulatory elements³³. (G) Gene expression level (Read Per Kilobase Million) of the four cases and 13 EBV-B normal control samples (GTEx project)²⁴.

Table 1

Balanced translocations detected in the 1000 Genomes Project

| | | Continental | | , | Bræ | Breakpoint A | Bre | Breakpoint B | Miono | | Deletion | Deletion involving breakpoints | akpoints | Conc | |
|-----------|------------------------------|-------------|--------------------------|--------|-----|-------------------------------|-----|--------------|----------|-----|--------------|--------------------------------|-----------|-----------|--|
| Sample ID | Karyotype | group# | group# Population* der\$ | 8 | chr | chr Coordinate chr Coordinate | chr | Coordinate | homology | chr | Size (bp) | Start | End | disrupted | Genes in TAD |
| 0200011 | 46,XX, t(9;14)(q34.2;q31.1) | AMR | PEL | 6 | 6 | 137230907 14 79839174 | 14 | 79839174 | ı | 6 | 14 13 | 137230908 137230922 | 137230922 | RXRA | WDR5,RNU6ATAC |
| 11002200 | | | | 14 | 14 | 79839173 | 6 | 137230923 | I | 14 | 0 | NA | NA | NRXN3 | ı |
| HG02720 | 46,XY, t(3;17)(q24;p13.3) | SAN | ITI | ю | ю | 143817430 | 17 | 2910366 | I | 8 | 5,219 14 | 143817431 | 143822650 | ı | SLC9A9,C3orf58 |
| 67/500H | | | | 17 | с | 143822651 | 17 | 2914751 | Щ | 17 | 4,383 2 | 2910367 | 2914750 | RAPIGAP2 | MIR1253 |
| C1201 AIV | 46,XY, t(16;17)(q23.1;q24.2) | ASN | CHB | 16 | 16 | 75336134 | 17 | 64953079 | C | 16 | 3 7 | 75336135 | 75336137 | CFDPI | CTRB2,CTRB1,BCAR1,TMEM170A,CHST6,CHST5,TMEM231,GABARAPL2,ADAT1 |
| 10A10012 | | | | 17 | 17 | 64953078 | 16 | 75336138 | Ą | 17 | 0 | NA | NA | 1 | MIR634, CACNG5, CACNG4, CACNG1, HELZ |
| 1350CAIN | 46,XX, t(2;19)(p11.2;p13.3) | EUR | TSI | 2 | 7 | 86491099 | 19 | 424310 | I | 7 | 1 8 | 86491100 | 86491100 | REEPI | KDM3A,CHMP3,RNF103-CHMP3,RNF103,RMND5A,CD8A,CD8B,ANAPC1PI,MRPL35 |
| 1NA20/04 | | | | 19 | 19 | 424308 2 | 2 | 86491101 | TG | 19 | - | 424309 | 424309 | SHC2 | PPAP2C,MIER2,THEG,C2CD4C, ODF3L2,MADCAMI |
| | | | | | | | | | | | | | | | |

"Continental group: AMR, SAN, ASN and EUR refer to American, East Asian, South Asian and European, respectively

*
Population: PEL, ITU, CHB and TSI refer to Peruvian in Lima (Peru), Indian Telugu in the UK, Han Chinese in Beijing (China) and Toscani in Italy, respectively.

\$ der: derivative chromosome

^ TAD: topological associated domain; domain information from the human IMR90 fibroblast cell line (hg19)

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Table 2

Inversions detected in the 1000 Genomes Project

| Sample ID | Karyotype | Continental group# | * | Coord | Coordinate | Microhomology | | Deletion (bp) | Gene disruption | ruption |
|-----------|---------------------------|--------------------|------------|---|----------------------------------|---------------|-------------------------|---|-------------------------|----------------------|
| | | | Population | Break A ^c Break B ^c | Break $\mathbf{B}^{\mathcal{C}}$ | | Break $A^{\mathcal{C}}$ | Break $A^{\mathcal{C}}$ Break $B^{\mathcal{C}}$ Break $A^{\mathcal{C}}$ Break $B^{\mathcal{C}}$ | Break $A^{\mathcal{C}}$ | Break \mathbf{B}^c |
| NA20759 | 46,XY,inv(2) (p14p12) | EUR | ISI | 67974196 81338340 | 81338340 | Ð | 1 | 0 | 1 | 1 |
| | | | | 67974194 | 81338341 | I | | | | |
| HG04152 | 46,XY,inv(1) (q32.1q32.1) | SAN | BEB | 199531693 | 203428169 | I | 99 | 0 | I | I |
| | | | | 199531758 | 203428141 | I | | | | |
| NA18959 | 46,XY,inv(3) (q26.1q26.1) | ASN | JPT | 166243083 | 166459774 | AT | 4 | 0 | I | I |
| | | | | 166243087 | 166243087 166459775 | Ŋ | | | | |
| NA21133 | 46,XY,inv(X) (p22.2p22.2) | SAN | GIH | 15629309 | 15687230 | L | 0 | 0 | I | ı |
| | | | | 15629310 15687231 | 15687231 | I | | | | |

#Continental group: EUR, SAN and ASN refer to European, South Asian and East Asian, respectively

*
Population: TSI, BEB, JPT and GIH refer to Toscani in Italy, Bengali in Bangladesh, Japanese in Tokyo (Japan) and Gujarati Indian in Houston (TX, USA) respectively.

 $^{\mathcal{C}}_{Breakpoint} = breakpoint$