Genomic integrity of human induced pluripotent stem cells across nine studies in the NHLBI NextGen program

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Author contributions
KK, KI, LRY, MAT, CM, IR and RAM analyzed data, KK, KI, LRY, MAT, IR, RAM, KCB, KAF, JWK, CEJ helped with the interpretation of the data and critically drafted the manuscript, RAM, LCB, KAF, JWK, KB, UB, LC, CC, MD, IC-O, TQ, GM, GM, MR, DJR, MHS, ET, and WY all helped in the generation of the iPSC lines, and participated in the interpretation of the results. All authors reviewed and approved the final manuscript.

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None.

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Abstract

Human induced pluripotent stem cell (hiPSC) lines have previously been generated through the NHLBI sponsored NextGen program at nine individual study sites. Here, we examined the structural integrity of 506 hiPSC lines as determined by copy number variations (CNVs). We observed that 149 hiPSC lines acquired 258 CNVs relative to donor DNA. We identified six recurrent regions of CNVs on chromosomes 1, 2, 3, 16 and 20 that overlapped with cancer associated genes. Furthermore, the genes mapping to regions of acquired CNVs show an enrichment in cancer related biological processes (IL6 production) and signaling cascades (JNK cascade & NFκB cascade). The genomic region of instability on chr20 (chr20q11.2) includes transcriptomic signatures for cancer associated genes such as ID1, BCL2L1, TPX2, PDRG1 and HCK. Of these HCK shows statistically significant differential expression between carrier and non-carrier hiPSC lines. Overall, while a low level of genomic instability was observed in the NextGen generated hiPSC lines, the observation of structural instability in regions with known cancer associated genes substantiates the importance of systematic evaluation of genetic variations in hiPSCs before using them as disease/research models.

Keywords

hiPSCs; Structural integrity; GWAS, VanillaICE; Oncogenes; Tumor suppressor genes

1. Introduction

Stem cells are primal cells with the capability of unlimited self-renewal and potential to differentiate into specific cell types. Two major types of human pluripotent stem cells are human embryonic stem cells (hESCs) and human induced pluripotent stem cells (hiPSCs); hESCs are derived from the undifferentiated inner cell mass of an embryo, whereas hiPSCs are reprogrammed from somatic cells by introducing defined pluripotency factors (Takahashi et al., 2007). Recently, hiPSCs have been evolved as a tool for disease modelling, drug development and cytotoxicity studies and individualized regenerative cell therapy (Shi et al., 2017). hiPSC lines reprogrammed from selected donors represent their genetic background and can provide a strong prototype to study the functional link between genetic variants and disease phenotype (Chamberlain, 2016; Warren and Cowan, 2018).
Numerous genetic variants associated with diseases have been identified by genome-wide association studies (GWAS) (Mahajan et al., 2014; Nikpay et al., 2015; Visscher et al., 2017; Tam et al., 2019). However, the functional validation of identified variants remains a challenge. Thus, the National Heart, Lung, Blood Institute (NHLBI) launched the Next Generation Genetic Association (NextGen) program to understand the causal effects of human genetic variations on heart, lung and blood disease phenotypes using hiPSC models (Warren et al., 2017). The NextGen program used iPSC technology to generate cell types relevant to heart, lung and blood disease processes from a large number of donors ascertained on the basis of heart, lung and blood phenotypes. The nine individual studies of the NextGen program integrate the results of genetic association studies (mainly GWAS) with hiPSC based functional studies to interrogate NHLBI-relevant phenotypes (see Supp. Table S1). The cell lines derived within the NextGen program are publicly available through WiCell, stem cell bank (www.wicell.org; NHLBI NextGen collection). The unique features of the NextGen generated hiPSCs include that the lines are derived from different cell types and from donors of several race/ethnicities using several reprogramming methods and then differentiated into various functional cell types.

Prior studies have documented genomic instabilities such as karyotypic abnormalities, single nucleotide variations (SNVs), copy number variations (CNVs) and insertions/deletions (INDELs) in hiPSCs (Mayshar et al., 2010). The abnormalities may occur via three different routes: (i) pre-existing variations in donor cells may be inherited by hiPSCs, (ii) the reprogramming procedure induced mutations, and (iii) passaging induced mutations. Studies suggest that genetic abnormalities in hiPSCs affect their clinical applicability and reliability as research models (D’Antonio et al., 2018; Martins-Taylor and Xu, 2012; Oliveira et al., 2014; Peterson and Loring, 2014; Yoshihara et al., 2017). Earlier genomic characterization studies using a small number of hiPSC lines (range:22–66 cell lines) have reported high mutational loads on the pluripotent cells (Gore et al., 2011; Hussein et al., 2011; Laurent et al., 2011; Mayshar et al., 2010). A recent large scale study of genome-wide profiling of hiPSC lines (N = 711) generated within Human Induced Pluripotent Stem Cells Initiative (HipSci) also reported that 41% of lines had one or more CNVs (Kilpinen et al., 2017).

To minimize the genomic instability in NextGen hiPSCs, all nine individual studies used non-integrative reprogramming methods and cell line maintenance under feeder free conditions (Warren et al., 2017). Since NextGen is a substantial resource of hiPSCs publicly available to the greater scientific community through WiCell, we performed a thorough analysis for genomic instabilities. The objective of our study was to identify newly acquired CNVs (CNVs that likely occur during reprogramming/passaging or that are sub-clonal in the cell of origin) in the NextGen hiPSCs. In this work, we extend our understanding of the genomic structural integrity of hiPSC lines across a wide range of donor DNA, human diseases and cell engineering techniques. The integrity of hiPSC lines was assessed by automated CNV calling followed with subsequent visual inspection of CNV signals. In this work we provide an in-depth understanding of the genomic stability of NextGen hiPSC lines. In future work, these hiPSCs may serve as surrogates for genetics and functional studies for several different disease phenotypes.
2. Methods

2.1. Generation of hiPSCs

The hiPSC lines within the NextGen program were reprogrammed at nine study sites (Warren et al., 2017) from various cell types including PBMCs, fibroblasts and erythroblasts, and using different cell engineering techniques including nonintegrating episomal vectors, lentivirus and Sendai virus. The culture medium, matrix/feeders and passaging methods used at each study sites are shown in Supp. Table S1. Reprogramming methods and maintenance of the NextGen hiPSC lines have been previously published (Warren et al., 2017 and reference therein) and also available on the Wicell website (www.wicell.org; NHLBI NextGen collection). The subjects include diseased individuals (sickle cell disease, platelet aggregation, coronary disease, pulmonary hypertension, arrhythmias, lipid disorders, left ventricular dynamics and hypertrophy) as well as healthy controls. At some study sites (Table 1) multiple passages were available per donor subject offering up the opportunity to look for acquisition of CNVs between differing passages of the same hiPSC.

2.2. Genotyping, genotype calling and quality control of the genotype data

Genotyping was performed through the NHLBI DNA Resequencing and Genotyping (RS&G) Service Center using the Illumina Infinium MEGA Chip. Genotype calls were made using GenomeStudio (version 2011.1) and Genotyping Module (version 1.9.4) and a total of 2,036,060 SNPs for 2,585 samples were released. The numbers of the genotyped samples and the entire workflow of the analyses are shown in Supp. Fig 1A. We applied a set of quality control (QC) steps to the genotype data and excluded SNPs as follows: 501,246 monomorphic SNPs; SNPs with ≥ 10% missing data; and SNPs showing deviations from Hardy–Weinberg equilibrium at a P value < 10\(^{-6}\). This resulted in 1,321,228 retained SNPs. Furthermore, SNPs with a MAF of <5% and SNPs in high linkage disequilibrium (LD) were also removed from the dataset to achieve 202,042 LD pruned SNPs (Supp. Fig 1B). It should be noted that the SNPs were dropped only for QC steps and not from the CNV analysis where all SNPs were included. Using a set of N = 202,042 LD pruned markers we estimated the identity by state (IBS) for all 2,585 samples by PLINK (Purcell et al., 2007); a total of 3,339,821 pairs for 2,585 samples were generated.

Of the 2,585 samples, we excluded 510 samples as follows: 428 samples with no pair for comparison, 52 pairs with sample swaps at plating where the proportion IBD (PI_HAT) = < 0.6 and 13 pairs that were technical replicates (Supp. Fig 1A). The final pairwise comparisons for IBS were then limited to 1,520 pairs including 1,114 pairs of donor DNA-hiPSC lines and 406 control pairs (Supp. Fig 1A). The IBS plots for all donor and hiPSC line pairs across 1,114 subjects and 406 HapMap controls indicated that all pairs passing IBS QC have Z2 = 1 and Z1 = Z0 = 0 where Z2, Z1 and Z0 are the probabilities of sharing 2, 1, and 0 alleles identical by state between the pair, respectively (Supp. Fig 1C). From the 1,114 donors and hiPSC pairs, 506 pairs were included in the structural integrity analysis because of the plating approach during genotyping, and other QC reasons that are described in detail in the next section (Supp. Fig 1A).
2.3. CNV calling from genotype data and QC measures

GenomeStudio software (Illumina inc.) provided the Log R ratios (LRRs: a normalized measure of the total signal intensity for two alleles of the SNP) and B-allele frequencies (BAFs: a normalized measure of the allelic intensity ratio of two alleles) on all the 2,036,060 genotyped SNPs. We did not exclude any monomorphic SNPs or low MAF SNPs from CNV calling as they provide information about copy number by total intensity. For CNV calling we applied a Hidden Markov Model (HMM) using a threshold of 10 supporting probes on the genotype data to identify CNVs in the samples using R package ‘VanillaICE’ (Scharpf et al., 2008). We used the hg19/GRCh37 reference version of the human genome in the CNV calling pipeline. The X and Y chromosomes were removed due to the complexity of reliably determining copy number by total intensity.

Given that copy number estimation is very sensitive to technological artifacts and differences in the preprocessing and normalization steps (Halper-Stromberg et al., 2011; Scharpf et al., 2011a; Scharpf et al., 2011b) we delineated signal intensities, and subsequent CNV calling was carried out separately for each plate. The main reason behind this sensitivity is that copy numbers are based on total probe A and B intensities (as compared to genotypes, which are based on relative probe intensity). Moreover, the number of CNVs called depends on the quality of the data, which differs between plates. Specifically, the larger the variability in the LRR of a sample (i.e. the lower the quality), the more CNVs are called on average. Since accuracy of CNV calls does not improve with decrease in quality of the genomic data, the increase in CNV calls represents an increase in false positive calls (see Supp. Fig 2A). The top panel shows the median absolute deviation (MAD) of the LRR per sample (x-axis) versus the number of CNVs called in the sample (y-axis). The red line is a loess smoother based on all samples, the blue line is a loess smoother based only on the samples with fewer than 100 CNV calls (which still shows a doubling on average of the number of CNVs called, from 9.8 for the best quality calls, to 18.2 for those with LRR MAD near 0.25). The bottom panel shows boxplots of the MAD LRRs for the samples per plate, showing clear between plate differences.

Furthermore, technical artifacts across samples within a plate tend to be more alike than technical artifacts across plates. However, false positive calls due to technical artifacts can be greatly reduced (at no change of sensitivity) if samples to be compared are plated together (Scharpf et al., 2012). Thus, we have restricted our structural integrity analysis to the pairs that were plated together. In addition, we also compared MAD of LRR and total number of called CNVs within each sample. The plot suggests that any sample with more than 50 CNVs should not be included in the analysis (Supp. Fig 2B). Based on the aforementioned reasons we dropped 524 pairs that were not plated together and 84 pairs where either donor or hiPSC had >50 CNVs. As a final dataset, 506 hiPSC-donor DNA pairs were analyzed for their structural integrity (Supp. Fig 1A).

2.4. Structural integrity analysis

Given sensitivities in calling CNVs using an automated pipeline (Halper-Stromberg et al., 2011), we attempted to minimize false positives by generating sub-chromosomal plots from the LRRs and BAFs for manual inspection. Since we were specifically interested in CNVs
that were acquired in hiPSC lines relative to the donor DNA, we contrasted the CNVs in
donor DNA and hiPSC lines to identify the CNVs that were called in the hiPSC line but not
the donor DNA, or those that were called in both but were of differing lengths between the
hiPSC line and donor DNA. Then we manually inspected the LRR and BAF plots for each
genomic region harboring the CNV, to examine the structural integrity of our 506 hiPSC
lines (Supp. Fig 1C). Only CNVs that passed dual manual inspection (by two independent
readers) were confirmed as the newly acquired CNVs.

2.5. Gene ontology and gene set enrichment analysis (GSEA)

The Gene List Analysis tool of PANTHER (Protein ANalysis THrough Evolutionary
Relationships) classification system (http://www.pantherdb.org) was used for gene ontology
annotation, functional classification and GSEA. Briefly, genes mapping to the regions of
acquired CNVs were uploaded to PANTHER user interface, and functional classification and
statistical over-representation analyses were performed. Both functional classification and
statistical over-representation analyses were done using the default Homo sapiens reference
list (20,996 genes) in the database. Statistical overrepresentation test finds functional
classes that are statistically over- (or under-) represented in the input list, compared to
the default (reference) list. We used the PANTHER GO-Slim annotation dataset for analysis.
Significance was determined with the Fisher’s exact test and the False Discovery Rate (FDR)
was calculated using the Benjamini-Hochberg procedure. We performed the statistical over­
representation test in a sequential manner based on the number of acquired CNVs per line:
(i) Input set1: genes mapping to 258 CNVs (149 hiPSC lines that acquired ≥1 CNV); (ii)
Input set2: genes mapping to 165 CNVs (56 hiPSC lines that acquired ≥2 CNVs); (iii)
Input set3: genes mapping to 107 CNVs (27 hiPSC lines that acquired ≥3 CNVs); (iv) Input
set4: genes mapping to 59 CNVs (11 hiPSC lines that acquired ≥4 CNVs) and (v) Input
set5: genes mapping to 31 CNVs (4 hiPSC lines that acquired ≥5 CNVs). The sequential
GSEA was similarly done stratified for acquired amplifications (Amps) and deletions (Dels)
separately.

2.6. Transcript expression analysis

Among the 15 hiPSC lines that were found to harbor a chr20q11.2 CNV, 9 hiPSC lines were
reprogrammed at one study site (site5). We relied on RNA sequencing (RNA-seq) previously
generated to validate the CNVs testing for differential expression at genes mapping to this
region for these lines. The detailed protocol for RNA-seq and pre-processing is published by
Carcamo-Oribe et al., (Carcamo-Orive et al., 2017). Briefly, we used log2 counts per million
(CPM) following TMM normalization (Robinson and Oshlack, 2010) implemented in edgeR
(Robinson et al., 2010) RNA-seq data. Data were adjusted for the effects of 8 sequencing
batches and 2 RNA preparation kits and genes with CPM ≥ 1 for at least 30% of the samples
were retained.

The hiPSC lines harboring chr20q11.2 CNV were termed as carrier hiPSC lines (N = 8) and
not harboring the CNV were termed as non-carrier hiPSC lines (N = 8). The core overlap
region of Amps among eight carrier hiPSC lines is 0.87 Mb (chr20:29810887-30682979).
Mapping transcripts were ascertained by overlap between the CNV region and transcript's
start site (i.e. if the start site of the transcript lies within CNV region, they were considered
as mapped transcripts). We calculated fold changes and \( p \)-values for differential expression using moderated t-statistics using the BioConductor package limma, and calculated \( p \)-values using the Bioconductor q-value package. Confidence intervals were calculated in R based on log fold changes and moderated standard errors.

### 2.7. Statistical analysis

Linear regression models were used to determine the effects of cellular reprogramming approaches and passaging on the structural variations of hiPSC lines. Briefly, we generated a matrix of hiPSC with newly acquired CNVs and their phenotypic characteristics and performed regression analysis in R (version 3.3.3). We estimated the significance using three different linear models, first for all CNVs combined, second for only amplifications and third for only deletions. The beta estimates were used to interpret the results and the \( p \)-values from the regression models are reported.

Statistical enrichment of oncogenes in the acquired Amps and tumor suppressor genes (TSG) in the acquired Dels region was determined using Fisher's exact test. We obtained the list of cancer related genes from the COSMIC (https://cancer.sanger.ac.uk/cosmic) database. A \( 2 \times 2 \) contingency table was set up for amplifications in oncogenes and deletions in TSG. Fisher's exact test was done in R (version 3.3.3) for both tables, and ORs and \( p \)-values from the test are given.

### 3. Results

#### 3.1. Genomic structural integrity of NextGen hiPSC lines

Out of 506 hiPSC lines compared to donor DNA, 357 lines showed no detectable difference in CNV calls compared to the paired donor DNA. We observed differences in CNVs between donor and hiPSC in 149 lines: 93 lines had one CNV difference, 29 lines had two and 27 lines had three or more CNVs different from the paired donor DNA. One hiPSC line had 15 CNVs, while the remaining 148 lines had six or fewer (Fig 1A). We did not observe any instance of aneuploidy in our hiPSC collection. The summary of the hiPSC lines examined and newly acquired CNVs are shown in Table 1. Overall, we identified 258 newly acquired CNVs are shown in Table 1. Overall, we identified 258 newly acquired CNVs (Supp. Fig 3) across the 149 hiPSC lines. The newly acquired CNVs include 111 Amps and 147 Dels. Size of the acquired CNVs ranged from 0.02 Mb to 26.77 Mb (mean = 0.98 Mb, median = 0.28 Mb). For each of the 149 lines harboring one or more CNVs, we calculated the cumulative size of the gained CNVs (mean = 1.09 Mb; median = 0.26 Mb) and the cumulative impact per hiPSC line was small. The majority of hiPSC lines (\( N = 132, 85\% \)) had cumulative CNV coverage of less than 2 Mb. The cumulative size of CNVs in the remaining 17 lines, ranged from 2 to 4 Mb in fourteen lines, a deletion of 12 Mb in one line, and Amps of 27 Mb in two lines (Fig 1B). The distribution of acquired CNVs was relatively even across all the chromosomes, with a few exceptions (Fig 1C). First, we did not observe any CNVs on chr21. Second, we observed a large region of Amp on chr17 in two hiPSC lines that overlaps 240 protein coding genes. Third, we observed a set of six regions on five chromosomes (chr1, chr2, chr3, chr16 and chr20) with recurrent CNVs (defined as more than one line with a CNV mapping to the region), affecting 30 hiPSC lines (Fig 1C).
3.2. Functional enrichment of genes mapping to acquired CNVs

A total of 1,649 genes mapped to the regions of acquired CNVs and we were able to map 1,395 of these genes to PANTHER IDs in the PANTHER database (Mi et al., 2019). On stratifying the acquired CNVs into Amps and Dels 1,010 genes mapped to the regions of acquired Amps and 700 genes mapped to the regions of acquired Dels. Of these 847 genes in Amps category and 585 genes in Dels category were mapped to PANTHER IDs. Using the biological process GO annotation dataset, we identified functional classification of the mapped genes (genes mapping to PANTHER IDs) under 16 GO terms. A descriptive overview of all GO terms, and gene sets within each GO term are represented in Supp. Table S2.

We detected enrichment in 22 biological processes gene sets at an FDR of 5% (q < 0.05) including a total of 470 genes (Fig 2A). To further understand if there were differences in regions with repetitive CNV acquisition over the 149 lines, we performed GSEA sequentially: Input set1/Input set2/Input set3/Input set4/and Input set5 included genes mapping to regions of acquired CNVs in 149 (≥1CNVs), 56 (≥2CNVs), 27 (≥3CNVs), 11 (≥4CNVs) and 4 (≥5CNVs) lines. This consisted 1,649/1,016/630/308/and 202 input genes, corresponding to 1,395/866/532/246/and 163 mapped genes, respectively. There were total 55 gene sets with a significant enrichment (q < 0.05) for at least one of the five input sets of genes Supp. Table S3. Interestingly, the fold enrichment (FE) of the identified gene sets increased markedly as we went from Input set 1 through to Input set 5, suggesting stronger enrichment in pathways for the more recurrent regions of acquired structural variation.

Of these, we highlight two cancer related signaling cascades that were over-represented at an FDR of 5% in all five input sets: “I-kappaB kinase/NF-kappaB signaling cascade” (GO:0007249) and “JNK cascade” (GO:0007254). For the I-kappaB kinase/NF-kappaB cascade FE was 3.10 (q = 4.32E-02) for the genes in Input set1, 4.64 (q = 2.51E-03) for the genes in Input set2, 6.96 (q = 9.7E-05) for the genes in Input set3, 13.81 (q = 3.26E-07) for the genes in Input set4 and 18.94 (q = 6.18E-08) for Input set5. For the JNK cascade, FE increased from 4.10 (q = 1.68E-02) to 5.51 (q = 5.31E-03), 8.07 (q = 6.07E-04), 17.46 (q = 1.18E-06), and 26.35 (q = 4.56E-08), respectively. In addition, we also identified the over-representation of “Cytokine production” (GO:0001816) among five input sets of genes with a FE of 3.50 (q = 9.12E-02) in Input set1, 5.64 (q = 4.81E-03) in Input set2, 8.26 (q = 5.5E-04) in Input set3, 17.86 (q = 1.07E-06) in Input set4 and 23.96 (q = 5.3E-07) in Input set5. Specifically, GO-term defining “IL-6 production” (GO:0032635), a pro-tumorigenic cytokine was identified to be overrepresented with FE raising from 8.03 (q = 6.48E-03) to 12.93 (q = 3.8E-04) to 21.05 (q = 1.05E-05), 45.52 (q = 2.74E-08) and 68.72 (q = 1.39E-09) from Input set 1 through to Input set 5 (Fig 2B).

To understand if the observed enrichment is driven either by acquired Amps or Dels, we performed sequential GSEA for gene sets mapping to the regions of acquired Amps and Dels separately. At an FDR of 5%, the FE for both signaling cascades, cytokine and IL6 production was higher in the gene sets mapping to regions of acquired Dels than gene sets mapping to all CNVs combined. There was only 1 hiPSC line that had ≥5 Dels, therefore here we had four input sets including 700/428/350/andl39 input genes corresponding to...
585/352/285/andl07 mapped genes. The FE for GO:0007249 was 6.78 (q = 4.83E-05) for the genes in Input set1, 10.4 (q = 1.77E-06) for the genes in Input set2, 12.73 (q = 1.78E-07) for the genes in Input set3, and 25.49 (q = 2.25E-08) for the genes in Input set4. For GO:0007254 FE increased from 8.06 (q = 2.31E-04) to 12 (q = 2.62E-05), 14.76 (q = 4.62E-06), and 39.4 (q = 1.23E-09), respectively. GO:C0001816 also represented the similar pattern as the FE went from 6.6 (q = 6.49E-03) in Input set1, to 10.9 (q = 1.76E-04) in Input set2, 13.42 (q = 3.80E-05) in Input set3, and 55.84 (q = 2.18E-08) in Input set4. The most dramatic effect was observed in over-representation of GO:0032635 with FE raising from 18.92 (q = 2.08E-05) to 31.3 (q = 5.83E-07) to 38.48 (q = 1.02E-07), and >100 (q = 5.36E-11) from Input set 1 through to Input set 4 (Fig 2C). Of note, all these GO-terms were the same as those seen in the combined analysis above; they were found to be statistically significant in the gene sets mapping to regions of acquired Dels and not Amps from the stratified analysis.

3.3. Enrichment of cancer related genes in the repetitive regions of CNVs in hiPSCs

Given the increasing FE of biological process in regions with repetitive CNV acquisition we performed an in-depth exploration of the genomic regions with instability. The sub-chromosomal region chr20q11.2 was the most recurrent region. Acquired CNVs in this region were observed in 16 hiPSC lines (15 Amps and 1Del) and the core overlap among the lines harboring amplifications was 0.60 Mb. This recurrent region of Amp encompasses 15 genes including known cancer associated loci ID1, BCL2L1 and TPX2 (Fig 3A). In addition to chr20q11.2, we identified an additional region on the p arm of the chr20. This region affects MACROD2 (MACRO Domain Containing 2) which is expressed at low levels in hiPSCs (Fig 3B). A region on chr16 includes two acquired Dels and one Amp. This region overlaps with the tumor suppressor gene WWOX (WW domain containing oxidoreductase) (Fig 3C). Three recurrent regions of Dels were observed on chr3, chr2 and chr1. The chr3 region overlaps FHIT (Fragile histidine triad) gene, that acts as a tumor suppressor and plays a role in apoptosis (Fig 3D). The region on chr2 overlaps a tumor suppressor gene RPRM (Reprimo, TP53 dependent G2 arrest mediator candidate) that is involved in the regulation of p53-dependent cell-cycle arrest (Xu et al., 2012) and the p53 pathway is linked to reprogramming (Krizhanovsky and Lowe, 2009)(Fig 3E). The chr1 region affects 10 genes including three cancer associated genes ITGB3BP, ROR1 and EFCAB7. This region also encompasses the FOXD3 gene, required for maintenance of pluripotent cells in the pre-implantation and peri-implantation stages of embryogenesis (Fig 3F).

The genomic positions, sizes, recurring scores (number of lines with the acquired CNV) and the genes affected by the recurrent regions of CNVs are given in Supp Table S4. We queried these recurrent CNVs against the Database of Genomic Variants (dgv.tcag.ca) (MacDonald et al., 2014) and the DGV structural variants overlapping with CNV regions are shown in Supp. Fig 4. The average size of the DGV structural variants was greater than our recurrent CNVs, except for the chr1 and chr20q arm CNVs.

3.4. Transcriptomic integrity of hiPSC lines harboring chr20 amplification

We had a total of 1,280 assembled transcripts on chr20 for 16 hiPSC lines (including both carriers and non-carriers) in our RNA-seq dataset. Among those, 631 transcripts passed the
CPM QC threshold (CPM > 1 in more than 30% of all samples) and 18 mapped to the core Amp region (the intersection of the chr20 Amp region among all eight carriers). We observed increased expression of four genes clustering at one end of the core amplification region: XKR7, CCM2L, RP1-310O13.7 and HCK. Fig 4 shows the fold changes (FCs) versus the genomic location of all 18 mapped genes on chr20, and the genes with significant up-regulation are shown in red. The FCs for the up-regulated genes are: XKR7 (FC = 2.13, P = 8.80E-04, CI = 1.44–3.16), CCM2L (FC = 1.97, P = 3.86E-03, CI = 1.29-3.00), RP1-310O13.7 (FC = 1.76, P = 3.36E-02, CI = 1.05–2.95), and HOC (FC = 2.16, P = 4.84E-02, CI = 1.01–4.62). Among the significantly up-regulated genes HCK gene is a known cancer associated gene. The FCs for other known cancer associated genes in the region are ID1 (0.89), BCL2L1 (1.17), TPX2 (1.16) and PDRG1 (1.21), but these were not statistically significant.

3.5. CNV patterns in sibling hiPSC lines

In our hiPSC collection there were 106 (21%) sibling hiPSC lines (two hiPSC lines reprogrammed from the same donor that have different passage numbers) reprogrammed from 53 donors. Among sibling hiPSCs, 81 lines showed no detectable difference in CNV calls compared to the paired donor DNA. We observed differences in CNVs between donor and hiPSC in 25 lines: 17 lines had one CNV difference, 6 lines had two and 2 lines had three CNVs different from the paired donor DNA (Table 2). In those with acquired CNVs, we compared the LRR and BAF plots between parent DNA and both early and late passage hiPSC lines. The CNV patterns were identical for both early and late passage hiPSCs (see Supp. Fig. 5) for 56% of the lines where we noted an acquired CNV. The passage numbers and CNVs acquired by each sibling hiPSC line are given in Supp. Table S5.

3.6. Effect of reprogramming method, and donor phenotypes on structural instability

To test the hypothesis that the choice of cell reprogramming and passaging might induce instability in the hiPSC lines we evaluated structural integrity of the different hiPSC lines among the cell reprogramming approaches and passage number. We also looked at the effects of donor cell source, sex, and age-related effects on structural integrity in the hiPSC lines. A detailed overview of all the different hiPSC lines, donor cell type, reprogramming method, passage number, genomic position of CNV, and CNV status is given in Supp Table S6. Given that our hiPSC lines are derived using non-integrating methods we did not observe any effect of cell reprogramming method on the stability of the lines. However, we observed a significant association between the structural variations of different hiPSC lines as it relates to passage number (p = 0.01). Notably, the hiPSC lines with higher passage number acquired more amplifications (p = 0.001). We also observed the highest number of acquired CNVs among the lines that were reprogrammed from fibroblast (p = 0.01) as compared to PBMCs. On stratifying the CNVs into Amps and Dels, the significant relationship was observed with Dels only (p = 0.04). In context of donor phenotypes, we did not observe any relationship based on the donor sex, but we observed age-related effects (p = 0.004). Lines derived from older individuals gained more Dels (p = 0.02) as compared to Amps.
4. Discussion

Genomic instability in hiPSC lines has been well recognized. To date there is only one large scale study that has assessed the phenotypic consequences of recurrent genomic CNVs from various standpoints including epigenome, transcriptome and proteome to cell differentiation and morphology (Kilpinen et al., 2017). This large scale genomic characterization study (Kilpinen et al., 2017) including 711 iPSC lines derived from 301 healthy individuals in the HipSci consortium used genotyping arrays to detect CNVs. The hiPSC lines in the HipSci collection (http://www.hipsci.org) are derived mostly from fibroblasts using Sendai virus. The authors identified trisomies in 4% of cell lines and 41% of lines harbored at least one or more CNVs of 7.15 Mb on average. They observed 35 recurrent regions of CNVs including 20 Amps, 11 Dels, three regions with both gain and loss, and one full chromosome duplication. Three most frequent CNVs identified in their study are X trisomy, chromosome 17 and 20. Genomic instability in our NextGen hiPSC lines was 29% with average size of 1.09 Mb, which is lower than the genomic instability reported in the HiPSci collection. The hiPSC lines in our collection weren't affected from any aneuploidies, and we observed 6 recurrent regions of CNVs on chromosomes 1, 2, 3, 16 and 20 (Fig 3). The Amp on chr20 and Del on chr1 have been previously observed in pluripotent cells but others have not been identified previously, to our knowledge. In addition to hiPSCi study, there are two other studies that evaluated the integrity of hiPSCs (Panopoulos et al., 2017) and hiPSC derived megakaryocyte cells (Kammers et al., 2017) from a genetic and transcriptomic perspective. However, the cell lines analyzed in both these studies are N = 222 and N = 14, respectively and is smaller than our NextGen collection.

Earlier reports included small-scale studies that evaluated chromosomal (Mayshar et al., 2010) and sub-chromosomal integrity (Gore et al., 2011; Hussein et al., 2011; Laurent et al., 2011). The first comprehensive evaluation of the chromosomal integrity or aneuploidy of hiPSC lines was performed through transcriptional profiling and reported the identification of a substantial number of abnormal cell-lines (Mayshar et al., 2010). Laurent et al., 2011 compared CNVs among normal somatic cell lines and hiPSC lines and reported that hiPSC lines contain more deletions than normal somatic cell lines. This work also found that the reprogramming process is associated with selection for deletions that affect tumor-suppressor genes, whereas maintenance of the cell lines is associated with selection for duplications in oncogenic genes (Laurent et al., 2011). Later, Hussein et al., 2011 observed a higher frequency of CNVs compared to the donor fibroblast cells. However, the number of CNVs in hiPSCs decreased with passaging (Hussein et al., 2011). They also concluded that the reprogramming process is associated with high mutation rates, causing increased levels of CNVs that lead to genetic mosaicism in early hiPSC lines (Hussein et al., 2011). In the aforementioned studies, the numbers of hiPSC lines were small (22–66 hiPSC lines) and the cells were reprogrammed using both integrating methods and non-integrating method (Mayshar et al., 2010; Hussein et al., 2011; Laurent et al., 2011). Our NextGen collection is very diverse in terms of source cells including PBMCs, fibroblasts and erythroblasts, and the hiPSC lines were reprogrammed using non-integrating methods only.

Another aspect of genomic instability was studied by Gore and colleagues (Gore et al., 2011). They explored the point mutations in hiPSCs using exome sequencing and validated
the ones that were fixed in the hiPSCs. Their study identified many missense mutations in the hiPSC lines that are predicted to alter protein function, and point mutations were enriched in cancer associated genes. We used SNP arrays to study structural instabilities in our NextGen hiPSC collection. Given that SNP arrays have limited resolution we were able to detect the larger CNVs but not point mutations in the NextGen hiPSC lines. It should be noted that every procedure that requires a "call" needs to consider the trade-off between sensitivity and specificity. In a typical CNV analysis, the false positive CNV calls substantially outnumber the true CNV calls depending on various factors such as the size of the CNV call and the data quality in that sample. Thus, it is imperative to use stringent requirements for data quality such as the MAD for the LRRs and a minimum number of probes supporting a CNV in an array-based approach. Obviously, this comes at the expense of possibly missing some true CNVs, specifically smaller CNVs. Compared to sequencing, array-based CNV calling has some advantages such as its high-throughput capacity, but also disadvantages such as the CNV resolution and the size detection limits. In our study, we identified 258 newly acquired CNVs across the 149 hiPSC lines, ranging from 0.02 Mb to 26.77 Mb. There could be CNVs smaller than 20 kB in truth, but these would be extremely hard, if not impossible, to detect with array-based technology. We recognize these limitations with respect to the CNV calling from the array data. In contrast, CNV calling based on short reads from sequencing has the capability to detect much shorter CNVs. In addition, sequencing also has a much higher resolution for break point detection, and sometimes even yields break points at base pair resolution.

Our study outlines the systematic genetic characterization of hiPSC lines at a large scale (>500 hiPSC lines). We observed high genomic structural integrity of the hiPSC lines, with 71% of the hiPSCs in the NextGen consortium showing no detectable CNVs. Among the 29% (N = 149) hiPSC lines that acquired CNVs: 49 lines acquired only Amps, 68 lines acquired only Dels and 32 lines acquired both Amps and Dels. The frequency of acquired Amps (43%) was lower than the frequency of Dels (57%) but the average size of Amps (1.28 Mb) was larger than Dels (0.76 Mb). In addition, to the recurrent CNVs on chromosomes 1, 2, 3, 16 and 20, one of the previously reported recurrent CNV on chr17 (chr17q11.2) was detected in one hiPSC line in our collection. We acknowledge that omission of chrX from our study is a limitation of our study. Our in-house CNV caller (VanillaICE) was developed only for the autosomes, implementing a likelihood in the hidden Markov model that reflects the data structures of LRRs and BAFs in autosomes. We understand that studying the structural instabilities on sex chromosomes will provide a complete structural landscape of all chromosomes of NextGen hiPSC lines, and additional work will be needed to examine sex chromosomal instabilities in these data. Generally, in our collection, hiPSCs acquired Dels included TSG whereas Amps included oncogenes, suggesting that cancer related genes provide additional selective advantage to the hiPSC line. On performing the Fisher’s exact test, we observed a statistically significant enrichment of oncogenes in the Amp regions (OR = 1.80; p = 0.02) and TSG in the Del regions (OR = 2.11; p = 0.01).

We identified enrichment of cancer related signaling cascades and biological process among the genes mapping to the genomic regions of acquired CNVs. The FE for both I-kappaB kinase/NF-kappaB and JNK signaling cascades increased sequentially with repetitive CNV acquisition in our collection. However, it should be kept in mind that inflammatory pathways...
are complicated, involved in many cellular processes, and are detected in almost every perturbation being analyzed. Cytokine production is a crucial biological process in context of cancer as cytokines exert both pro- and anti- cancer effects. Cytokines bind to their cognate receptor on the effector cells and promote or inhibit tumor development and progression in autocrine and paracrine manners (Van Gorp and Lamkanfi, 2019). A number of cytokines including interleukin-1 (IL-1), interleukin-6 (IL-6), interleukin-15 (IL-15), tumor necrosis factor (TNF), transforming growth factor-β (TGF-β) and vascular endothelial growth factor (VEGF) are implicated in tumorigenesis (Dranoff, 2004; Van Gorp and Lamkanfi, 2019). Among these cytokines, IL-6 is highly upregulated in many cancers and promotes several tumorigenic activities such as enhanced cell proliferation, survival, invasion and angiogenesis (Taniguchi and Karin, 2014). Elevated levels of IL-6 stimulate hyperactivation of IL-6/JAK/STAT3 signaling pathway that promotes tumor growth and progression and hinder antitumor immunity (Johnson et al., 2018).

Querying our recurrent CNVs against structural variants in the DGV suggested that these CNVs have been reported in human populations previously. This highlights that CNVs identified in our hiPSC lines are recurrent in the context of comparison to prior publications. Numerous studies have reported the chromosome 20 (20q11.21) Amp as a recurrent mutation in hESCs (Avery et al., 2013; International Stem Cell et al., 2011; Lefort et al., 2008; Narva et al., 2010; Nguyen et al., 2014; Spits et al., 2008; Wu et al., 2008) and in hiPSCs (Kilpinen et al., 2017; Laurent et al., 2011; Martins-Taylor et al., 2011). From our study, we conclude that only 2.96% (15 of 506) of our hiPSC lines were affected by this amplification. This is well below the previously published studies where the percentage of affected hiPSC lines ranged from 13 to 50% (Elliott et al., 2010; Laurent et al., 2011; Martins-Taylor et al., 2011). Previous studies in hESCs and hiPSCs reported that this Amp region contains several genes, including the known pluripotency-associated gene DNMT3B (Laurent et al., 2011), and BCL2L1, which encodes the anti-apoptotic protein BCL-X (International Stem Cell et al., 2011; Laurent et al., 2011). Our data confirmed that the duplicated region of chr20 (20q11.21) in all 15 hiPSC lines harbors three carcinoma associated genes; namely ID1, BCL2L1 and TPX2. ID1 plays role in cell growth, senescence, and differentiation and is associated with the salivary gland (Hu et al., 2017) and breast cancer (Gumireddy et al., 2014). BCL2L1 acts as an apoptotic inhibitor and is related to many types of cancers such as melanoma, glioblastoma (Trisciuoglio et al., 2017), breast carcinoma (Choi et al., 2016) and colorectal carcinoma (Koehler et al., 2013). TPX2 participates in cell cycle progression and this gene is linked to hepatocellular carcinoma (Hsu et al., 2017).

We extended the investigation of the genomic region with instability on chr20 of our hiPSC lines to the transcript level. A total of 18 transcripts mapped to the core overlap region of Amp, of which 13 are protein coding. In particular, there were four transcripts that were significantly different between the carrier and non-carrier Amp lines. Among the differentially expressed transcripts, transcript of HCK gene showed highest log2 FC. HCK is a member of the Src family of tyrosine kinases and is associated with tumor progression in colorectal cancer (Roseweir et al., 2019). The Amp region consists of transcripts for five cancer associated genes including ID1, BCL2L1, TPX2, PDRG1 and HCK. Although we didn't observe the significant p-values for other cancer associated genes except HCK.
the FCs were relatively high, and our limited sample size may impact the power to find these differences statistically meaningful. Confirming that the presence of oncogenes in the genomic regions of instability provide additional selective advantage to the hiPSC lines.

Our study design offered us an opportunity to study the CNV patterns of sibling hiPSC lines and compare with parent cell. The patterns were almost identical for both early and late passage hiPSCs for all instances where we noted an acquired CNV. Our work also demonstrates that using non-integrating reprogramming method can limit the occurrence of CNVs in the hiPSCs. However, the choice of source cell may affect the occurrence of the structural variations in the hiPSC lines. Among the hiPSC lines that acquired CNVs, the highest number of CNVs was found in lines generated from fibroblasts. It should be noted that these lines were compared to genomic DNA from blood/PBMC sources and not to donor fibroblasts and it is highly plausible that a meaningful number of the apparently 'acquired' CNVs might have originated from the source fibroblast.

5. Conclusions

In conclusion, our data shows the overall high quality of the hiPSC lines generated through the NextGen program. Our results also confirm that some sub-chromosomal regions are particularly susceptible to genomic instability and that in this large set of hiPSCs from NextGen, the acquired CNVs show a specific enrichment for cancer related biological process and signaling cascades. Additionally, the recurrent regions of acquired CNVs are enriched for cancer related genes. Given this cancer related enrichment, the estimation of genomic instability of hiPSCs is necessary for both basic research and hiPSC-based therapies.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Fig. 1.
Characteristics of the Copy Number Variants in NextGen hiPSC lines. [A] Histogram showing the number of hiPSC lines with number of detected CNVs. [B] Histogram showing the cumulative size of CNVs in mega base pairs (Mb) per hiPSC line. Black dashed line shows that cumulative CNV coverage for 85% hiPSC lines was less than 2 Mb. [C] Distribution of 258 CNVs acquired by hiPSC lines across the genome. The chromosomal regions harboring cluster of CNVs are indicated.
Fig. 2.
Gene set enrichment analysis for the set of 1,395 genes mapping to 258 CNVs from 149 hiPSC lines for enrichment with regards to biological processes. [A] Fold enrichment of the 22 GO Terms with an FDR of <5% (q < 0.05) are displayed. q values <0.01 and <0.001 are shown as "*" and "**", respectively. [B] Sequential Gene Set Enrichment of 4 biological process: I-kappaB kinase/NF-kappaB signaling cascade (GO:0007249), JNK cascade (GO:0007254), Cytokine production (GO:0001816) and IL-6 production (GO:0032635). Input set 1/2/3/4 and 5 include gene sets mapping to regions ≥1, ≥2, ≥3, ≥4 and ≥5 acquired CNVs in lines. The FE and FDR-P values increase markedly from input set1 through to input set5. [C] Sequential GSEA for gene sets mapping to regions of acquired Dels. Input set 1/2/3 and 4 include gene sets mapping to regions of ≥1, ≥2, ≥3 and ≥4 acquired Dels in hiPSCs. The FE and FDR-P values are higher for gene sets mapping to regions of Dels than all CNVs combined.
Fig. 3.
Chromosomal regions harboring clusters of CNVs. The chromosomal position of each CNV cluster is shown in a red box over the ideogram. The amplifications are represented as blue bars and deletions are shown in red bars. Dashed vertical lines delineate the core overlap regions, and in some cases nearby CNVs within the same genomic locus are also shown. The bottom panel shows the genes lying in the overlap region that could be potentially affected by the CNV in the hiPSC lines.
Fig. 4.
Expression levels of the transcripts mapping to the core overlap region of chr20 amplification. The chromosomal position of the amplification is shown in a red box over the ideogram. hiPSC lines harboring chr20 amplification are shown as blue bars and dashed vertical lines delineate the core overlap region. The bottom panel shows the fold changes comparing carriers to non-carriers of the genes mapping to the overlap region, and their respective 95% confidence interval. Statistically significant up-regulation is observed for the genes marked in red. The expression of each gene is drawn positionally to the center of the gene.
Table 1
Genomic structural integrity of the NextGen hiPSC lines. Out of 506 hiPSC lines, 149 lines acquired CNVs during reprogramming/passaging.

| Site | Donor DNA (N) | hiPSC Line (N) | Pairs (N) | hiPSC lines with acquired CNV(s) (N) | Acquired CNVs (N) |
|------|---------------|----------------|-----------|-------------------------------------|------------------|
| Site1| 132           | 132            | 132       | 2                                   | 2                |
| Site2| 7             | 7              | 7         | 0                                   | 0                |
| Site3| 1             | 1              | 1         | 1                                   | 1                |
| Site4| 171           | 171            | 171       | 104                                 | 203              |
| Site5| 46            | 83 *           | 83        | 21                                  | 26               |
| Site6| 9             | 9              | 9         | 2                                   | 2                |
| Site7| 49            | 60 *           | 60        | 7                                   | 7                |
| Site8| 0             | 0              | 0         | 0                                   | 0                |
| Site9| 38            | 43 *           | 43        | 12                                  | 17               |
| Total| 453           | 506            | 506       | 149                                 | 258              |

* >1 hiPSC line for some donor DNA.
Table 2

Details of sibling hiPSC lines. 25 hiPSC lines acquired CNVs as compared to their corresponding donor cell.

| Site | Donor DNA (N) | iPSC Line (N) | Pairs (N) | hiPSC lines with acquired CNV(s) (N) | Acquired CNVs |
|------|---------------|---------------|-----------|--------------------------------------|---------------|
| Site5| 37            | 74            | 74        | 17                                   | 22            |
| Site7| 11            | 22            | 22        | 2                                    | 2             |
| Site9| 5             | 10            | 10        | 6                                    | 11            |
| Total| 53            | 106           | 106       | 25                                   | 35            |