Whole-genome mutational burden analysis of three pluripotency induction methods

Kunal Bhutani1,†, Kristopher L. Nazor2, Roy Williams2, Ha Tran2, Heng Dai3,†, Željko Džakula3, Edward H. Cho3, Andy W.C. Pang3, Mahendra Rao4,†, Han Cao3, Nicholas J. Schork1,† & Jeanne F. Loring2

There is concern that the stresses of inducing pluripotency may lead to deleterious DNA mutations in induced pluripotent stem cell (iPSC) lines, which would compromise their use for cell therapies. Here we report comparative genomic analysis of nine isogenic iPSC lines generated using three reprogramming methods: integrating retroviral vectors, non-integrating Sendai virus and synthetic mRNAs. We used whole-genome sequencing and de novo genome mapping to identify single-nucleotide variants, insertions and deletions, and structural variants. Our results show a moderate number of variants in the iPSCs that were not evident in the parental fibroblasts, which may result from reprogramming. There were only small differences in the total numbers and types of variants among different reprogramming methods. Most importantly, a thorough genomic analysis showed that the variants were generally benign. We conclude that the process of reprogramming is unlikely to introduce variants that would make the cells inappropriate for therapy.
Cell replacement therapies using cells derived from human pluripotent stem cells (embryonic stem cells (hESCs) and induced pluripotent stem cells (iPSCs)) have been approved for clinical trials for macular degeneration, spinal cord injury and type-1 diabetes. The limited data available to date indicate that there are no adverse events in patients receiving these cells. However, there continues to be discussion about the theoretical chance that these transplanted cells may develop into tumours or cause other pathologies. The discussion has come to focus on iPSCs, largely due to concerns that the massive epigenetic remodelling that occurs during reprogramming might cause genomic mutations that could make the cells tumorigenic. These concerns have led multiple groups to study the genomic integrity of iPSCs using methods that include single-nucleotide polymorphism (SNP) genotyping, CGH, karyotyping and exome sequencing (reviewed in refs 6, 7). In each case, the focus has been exclusively on a single type of genomic alteration, rather than considering the combined effects of single-nucleotide variants (SNVs), structural variants (SVs) and copy-number variations. Further, detailed comparative genomic analyses of iPSC lines that have been generated via distinct reprogramming methodologies have yet to be reported.

In this study, we assessed genome-wide mutation rates from replicate isogenic cell lines generated by three distinct methods. We used integrating viral (retrovirus), non-integrating viral (Sendai virus) and non-integrating non-viral (messenger RNA (mRNA)) reprogramming strategies to introduce exogenous expression of POU5F1, SOX2, KLF4 and MYC in separate fractions of a single fibroblast population (Fig. 1a). Three clonal lines were established from iPSCs generated by each method and determined to be pluripotent by standard measures. To detect SNVs, SVs and copy-number variations within each line and the parental fibroblast population, we generated whole-genome sequencing data for each iPSC line and the parental fibroblasts at an average read depth of 39-fold, with 93.7% of the autosomal genome covered by at least 10 reads. In addition, to assess chromosomal rearrangements and large SVs with high resolution, we performed whole-genome mapping using the recently developed Irys Technology (BioNano Genomics, San Diego, CA). We detected subtle differences in the numbers of variants depending on the method, but rarely found mutations in genes that have any known association with increased cancer risk. We conclude that mutations that have been reported in iPSC cultures are unlikely to be caused by their reprogramming, but instead are probably due to the well-known selective pressures that occur when hPSCs are expanded in culture.

Results
Identification of single-nucleotide variants. To characterize the mutational burden in iPSCs, we identified SNVs that were unique to each iPSC cell line by integrating results from HaplotypeCaller and MuTect, as described in the Methods section. For variant calling with HaplotypeCaller, we treated all 10 samples (the parent fibroblast population and three biological replicates for each reprogramming method) as part of a single population using the multisample option. This pipeline was tuned for the identification of recurrent variations in population studies, and therefore enabled us to have higher specificity in classifying reprogramming-induced mutations by accounting for mosaicism in the parental fibroblast population. To gain a more sensitive assessment of the SNV landscape across the iPSC samples, we also called variants using MuTect, wherein each iPSC line was compared with the parental fibroblast population in an analogous manner to which tumour samples are compared with normal tissue in oncogenic studies (Fig. 1c; Supplementary Fig. 1). Taken together, the results from these two distinct variant calling pipelines gave us higher confidence in our ability to identify true variants through the moderation of type I and type II errors, respectively.

The identified set of putative unique variants was split into three groups according to our confidence in the variant calls: Variant Set 1 was called unique by both MuTect and HaplotypeCaller; Variant Set 2 had coverage between 20–60 ×, and allele frequency distribution between 0.4 and 0.6 but was called only by MuTect, and Variant Set 3 comprised those with allele frequencies between 0.2–0.4 and 30–50 × coverage.

Variant validation by quantitative PCR from each of these three groups indicated that the mutations from Variant Set 1 had the highest likelihood of being true somatic variants (Supplementary Data 1; Methods). Therefore, subsequent analyses were restricted to these variants, which likely occurred during the initial doublings of the founder populations of the iPSCs compared with variants that are in lower allelic fractions that could arise as a result of proliferation of the cells. Our filtering strategy focuses on these high-confidence variants and also removes variants that are found in low-complexity regions and dbSNP (Fig. 1c; Supplementary Data 2).

Identification of insertions and deletions. Due to the specificity of multisample HaplotypeCaller in identifying high-confidence SNVs, HaplotypeCaller was also used for identifying insertions and deletions (indels).

Identification of structural variations. Structural variations (SVs), 10 kbp to 1 Mbp in size, are common in the human genome, but challenging to assess by sequencing alone. Genome mapping in nanochannel arrays provides a single-molecule platform complementary to DNA sequencing for accurate genome assembly and SV analysis (Irys System, BioNano Genomics). Unamplified genomic DNA, fluorescently labelled at a seven-base sequence motif and by the intercalating dye YOYO-1, was linearized by electrophoresis into 50-nm channels. An array of approximately 12,000 nanochannels was imaged for each sample, and the process was repeated multiple times, producing data at a throughput of about 1.5 Gbp h⁻¹. Only molecules 150 kbp and longer were used to create a de novo assembly of the complete genome. We analysed the fibroblast control and one iPSC line chosen at random from each of the three reprogramming methods.

We collected 160 Gbp (~50 ×) of high-molecular-weight DNA for each sample, ~500,000 single-molecule maps with the minimal length of 150 kbp. Data from each sample were assembled using an in-house developed de novo assembler based on Overlap-Layout-Consensus paradigm. The N50 length is the length for which the collection of genome maps of that length or greater cover >50% of the total genome length. The genome maps in this study had a N50 of >0.9 Mbp, overlapping close to 90% of the Genome Reference Consortium Human genome (GRch37; Methods; Supplementary Fig. 2; Supplementary Table 1).

Assessing pathogenicity of unique variants. To assess the functional consequences of the variants, the sets of high-confidence unique SNVs and indels were annotated using the SGAdiver and Oncotator program suites. They were further characterized based on overlaps with ENCODE annotated genomic regions. As expected, most mutations fell within intergenic or intronic regions, with the rate of coding mutations in the range of 2–10 mutations per cell line (Fig. 2d). The potential for variants to be oncogenic was assessed by
measuring their overlap with cancer genes, transcription factor-binding sites (TFBS), MutSig genes and Familial Syndrome Cancer Genes (Methods; Supplementary Data 3). We then compared the frequency of the annotated somatic mutations across the three reprogramming methods using an exhaustive permutation testing procedure and one versus all contrasts in analysis of variance (ANOVA). All of the iPSC lines had hundreds of variants compared with the parental fibroblasts, with the mRNA-derived lines having fewer high-confidence mutations on average than the other methods (Fig. 2a; Tables 1 and 2; Supplementary Table 2). We did not find evidence that any one of the methods was more likely than the others to cause oncogenic or deleterious mutations, but we identified trends based on one versus all ANOVA contrasts that linked reprogramming strategies to certain mutations. The cell lines reprogrammed using mRNA had several variants that overlapped binding sites for a transcription factor (EZH2; enhancer of zeste 2 polycomb repressive complex 2 subunit), although it is unknown if these would disrupt transcription factor binding. The retroviral-induced cell lines harboured more potentially damaging mutations than the other methods (Table 3). In addition, the three retrovirally reprogrammed cell lines contained integrated vectors in 7, 8 and 12 sites in the genome, and while the majority of integration sites were intergenic, some integrations mapped to coding regions (Supplementary Fig. 3). In Sendai virus reprogrammed cell lines, there was a trend towards fewer coding mutations. However, we want to emphasize that based on the variance and means of the aggregated statistics, this study was not sufficiently powered to assess the differences among the different reprogramming methods for some variant classifications (Table 1).

We also characterized the context of the mutations, as well as the overall transition/transversion rates for the different samples (Fig. 2b). Mutational context analysis revealed no realizable difference among the different reprogramming methods and identified no links to known cancer-related mutational signatures19. Finally, we looked at the Combined Annotation-Dependent Depletion (CADD)20 score distribution of the variants, and although we saw statistically significant differences among the three methods, the scores for all methods fell mostly within the non-deleterious range of less than 15 CADD Score (Supplementary Fig. 4).

Using the variant calling algorithms for the data from de novo whole-genome mapping of four of the samples, we called 259
Figure 2 | Characterization of variants caused by reprogramming method. (a) Overall counts for the number of high-confidence SNVs and indels per sample. (b) The relative percentage of mutational subtypes for the SNVs in each sample. (c) A violin plot and box plot for the indel size distributions in the sample, a positive length indicates an insertion, whereas a negative one is a deletion. (d) Variant classifications based on their relative locations in the genome. The error bars indicate the low, median and high replicate for each reprogramming method. Introns and IGR variants are plotted on a different scale.

Table 1 | P values from the permutation-based ANOVA test for variant-type differences across the three reprogramming methods.

| SNV raw counts | SNV relative rates | Insertions raw counts | Deletions raw counts | Indel raw counts | Combined | Samples for 80% power |
|----------------|--------------------|-----------------------|----------------------|-----------------|----------|-----------------------|
| CADD Phred>15  | 0.12               | 0.36                  | ND                   | ND              | 0.12     | 4                     |
| Coding         | 0.03               | 0.18                  | 0.34                 | 0.23            | 0.24     | 0.16                  |
| Damaging       | 0.09               | 0.30                  | ND                   | ND              | 0.09     | 4                     |
| Near cancer gene | 0.10             | 0.94                  | 0.11                 | 0.20            | 0.14     | 0.39                  |
| Total          | 0.01               | ND                    | 0.18                 | 0.16            | 0.16     | 0.20                  |

ANOVA, analysis of variance; CADD, Combined Annotation-Dependent Depletion; ND, not determined; indels, insertions and deletions; SNV, single-nucleotide variant.

Rates were determined by dividing the number of variants by the total number of variants. ND either because it is not consistent with the calculations or there were too few variants to analyse. Combined is the sum of SNVs and indels. The last column lists the sample size estimates necessary based on 80% power for an ANOVA statistic given the current mean and variance for the grouping by reprogramming methods. This is based on the combined counts.
addition, the only large structural variation (a 228.8-kbp deletion) of mutations in binding sites for a transcription factor (EZH2). In reprogramming methods, mRNA-iPSCs had a greater number of mutations than either retrovirus or Sendai virus-based mRNA reprogramming technique resulted in fewer total mutations among the methods. Although we found that the non-integrating mutational load of cells, but there were subtle differences that indicates that all of the reprogramming approaches add to the widely used reprogramming methods for generating iPSCs.

**Discussion**

Our assessment of the mutation profiles associated with three widely used reprogramming methods for generating iPSCs indicates that all of the reprogramming approaches add to the mutational load of cells, but there were subtle differences among the methods. Although we found that the non-integrating mRNA reprogramming technique resulted in fewer total mutations than either retrovirus or Sendai virus-based reprogramming methods, mRNA-iPSCs had a greater number of mutations in binding sites for a transcription factor (EZH2). In addition, the only large structural variation (a 228.8-kbp deletion) we detected was in an mRNA-reprogrammed iPSC line. In contrast, the Sendai virus samples had fewer coding mutations than the other methods.

Our results using retroviral vectors showed that this method caused a similar number of mutations as the other methods, but was slightly more likely to introduce mutations that are classified as deleterious. But the main concern with retroviral vectors is the fact that they insert into the genome. Genomic insertions have a low but finite chance of disruption of active genes or regulatory regions, and can cause cancers if they activate endogenous oncogenes. Retroviral vectors were used in the first methods developed for reprogramming but were reported to be capable of reactivation, which resulted in tumours in mice. Use of retroviral vectors for reprogramming has become less popular as delivery methods for transiently expressing the reprogramming factors without being inserted, such as the Sendai viral vectors and mRNA used here, have become more efficient.

Given the potential practical significance of our findings to clinical applications for stem cells, it is important to appreciate some of the biological context surrounding our experiments. We focused on the high-confidence variants that differed from the parental fibroblast population. These variants likely arose during the initial doublings of the founder population of iPSCs, and as such, the Sendai viral vectors and mRNA used here, have become more efficient.

**Table 2 | High-confidence variants in coding regions.**

| Sample | Synonymous | Nonsense and non-synonymous | Coding regions affected by high-confidence nonsense and non-synonymous variants |
|--------|------------|-----------------------------|--------------------------------------------------------------------------------|
| M1     | 0          | 2                           | Nonsynonymous: BC068088 and C14orf51S9                                         |
| M2     | 2          | 2                           | Nonsynonymous: BPI8F1 and MACROD1                                             |
| M3     | 2          | 1                           | Nonsynonymous: RPAP2                                                           |
| R1     | 2          | 7                           | Nonsynonymous: SALL1; Nonsynonymous: C2orf91, CCDC150, SSCS5, SYT7, UTRN and WDR72 |
| R2     | 0          | 7                           | Nonsynonymous: PRR12; Nonsynonymous: ADAM18, CATSPERG, IKBIP, NCR3Li1, PRR12 and SPTA1 |
| R3     | 3          | 7                           | Nonsynonymous: ALPL, HIST1H2BD, ITGB8, ORSAP2, PKP2, RNF10 and TMPRSS5         |
| S1     | 0          | 5                           | Nonsynonymous: KCNC3, OVO52, SDR16CS, XPR1 and ZNF660                         |
| S2     | 3          | 4                           | Nonsynonymous: CLEC5A, FAM208B, MMMEL1 and PCDHB16                            |
| S3     | 3          | 2                           | Nonsynonymous: FSCN3; Nonsynonymous: FSCN3 and PRDM4                           |

The number of high-confidence synonymous and nonsynonymous coding mutations identified with high-confidence SNVs in each sample. Nonsynonymous variants in protein-coding regions are listed.

M1: mRNA vector; R1: Retrovirus; S1: Sendai virus.

*The CATSPER gene has two non-synonymous high-confidence mutations in this sample.*

**Table 3 | Functional impacts as calculated by ANOVA contrasts using a one versus all approach.**

|          | M1 | M2 | M3 | R1 | R2 | R3 | S1 | S2 | S3 | P value |
|----------|----|----|----|----|----|----|----|----|----|---------|
| Non-integrating mRNA versus all others—transcription factor EZH2-binding sites | 13 | 11 | 6  | 1  | 7  | 2  | 3  | 0  | 3  | 0.008   |
| Retrovirus versus all others—damaging mutations assessed by Condel, Polyphen or SIFT | 5  | 3  | 1  | 5  | 9  | 13 | 5  | 3  | 2  | 0.014   |
| Sendai virus versus all others—mutations in coding regions | 27 | 19 | 9  | 16 | 20 | 24 | 12 | 9  | 5  | 0.044   |

ANOVA, analysis of variance; mRNA, messenger RNA

ANOVA contrasts were set up to compare each reprogramming method against the other two. For each reprogramming method, the most significant difference is presented. The EZH2-binding site overlaps were determined by ENCODE annotations (Methods).

insertions and deletions in the parental fibroblast sample, 239 in the retroviral sample, 248 in the Sendai sample and 268 in the mRNA sample. The size of the variants ranged from 2.8 kbp to 4.9 Mbp. Using a 50% reciprocal size overlap cutoff, we identified variants that were shared among the samples. After manual inspection to eliminate false positives and false negatives, we found no cell line-specific variants in the retroviral and Sendai samples, but one deletion in the mRNA-reprogrammed sample. The deletion in this line was a heterozygous 228.8-kbp deletion at Xp22.11, which removed one copy of the PHEX gene (phosphate-regulating neutral endopeptidase) and one copy of Mir_548. This deletion is illustrated in Fig. 3, which shows the assembly of part of the X chromosome for each cell line, as well as the single-molecule data supporting two haplotypes for this region in the mRNA sample. We subsequently looked for this deletion in the sequencing data and identified it in the same sample; it was not present in any of the other samples. This suggests that this iPSC line was derived from a rare fibroblast that we detected was in an mRNA-reprogrammed iPSC line. In contrast, the Sendai virus samples had fewer coding mutations than the other methods.

Our results using retroviral vectors showed that this method caused a similar number of mutations as the other methods, but was slightly more likely to introduce mutations that are classified as deleterious. But the main concern with retroviral vectors is the fact that they insert into the genome. Genomic insertions have a low but finite chance of disruption of active genes or regulatory regions, and can cause cancers if they activate endogenous oncogenes. Retroviral vectors were used in the first methods developed for reprogramming but were reported to be capable of reactivation, which resulted in tumours in mice. Use of retroviral vectors for reprogramming has become less popular as delivery methods for transiently expressing the reprogramming factors without being inserted, such as the Sendai viral vectors and mRNA used here, have become more efficient.

Given the potential practical significance of our findings to clinical applications for stem cells, it is important to appreciate some of the biological context surrounding our experiments. We focused on the high-confidence variants that differed from the parental fibroblast population. These variants likely arose during the initial doublings of the founder population of iPSCs, but we cannot rule out their origin in a minority population in the heterogeneous parental fibroblasts that was undetectable by sequencing. We note that there were sequencing reads that support several additional lower variant allele frequency mutations in the cell lines as well. It should be noted that the cells we analysed were cultured for a relatively short time, and that variants in an early-arising small subpopulation could become more dominant over time if they give the cells a selective advantage in culture.
To move forward towards applying stem cell-based therapies for human disease, it is important to focus our efforts on improving the likelihood that there will be no adverse effects of these therapies. Our study was more extensive than previous analyses, but larger studies of this depth are still needed. While results of our study do not rule out the possibility that reprogramming cells could introduce oncogenic mutations that compromise the safety of iPSCs, they should alleviate some of the concern about how likely it is that reprogramming itself would cause dangerous genomic changes that could lead to harm to transplant recipients. It is important to note that genomic aberrations, some of which could be oncogenic, are known to occur during the considerable expansion of cells that is required for clinical applications. While development of new methods for reprogramming will make the process simpler and less expensive, it is critical at this stage that we concentrate on monitoring the appearance and potential consequences of mutations that arise during cell division and differentiation in culture and are selected for by the culture conditions.

**Methods**

**Overview of experimental approach.** The design of our study was to evaluate mutation profiles associated with the three different reprogramming strategies (see below for details on each method) we considered, as well as the characterization of different forms of variation and the analysis of the variation within and across the different reprogramming strategies.

**Retroviral reprogramming.** PLAT-A-packaging cells (Cell Biolabs, Inc.) were plated on six-well plates coated with poly-L-lysine at a density of $1.5 \times 10^6$ cells per well without antibiotics and incubated overnight. Cells were transfected with 4 μg of Moloney murine leukaemia-based retroviral vectors (pMXs) containing the human complementary DNA of SOX2, KLF4, or MYC (Addgene catalog number 17217, 17218, 17219, and 17220, respectively) by Lipofectamine 2000 (Life Technologies, Carlsbad, CA) according to the manufacturer’s instructions. Viral supernatants were collected at 48 and 72 h post transfection, filtered through a 0.45-μm pore-size filter, 200,000 Human dermal fibroblasts (Science cell Catalog number 2300) were seed onto each well of a six-well plate overnight prior transfection. Equal volumes of fresh 48 and 72 h viral supernatants containing each of four retroviruses supplemented with 6 μg ml$^{-1}$ of Polybrene (Sigma) were added onto the cells on day 1 and day 2, respectively. On day 5, the transduced cells were split onto mouse embryonic fibroblasts (MEFs) at a density of $10^6$ cells per well of a six-well plate in hESC medium supplemented with 0.5 mM valproic acid (VPA; Stemgent). Cells were fed every other day with VPA-supplemented hESC medium for 14 days before VPA was withdrawn. Individual iPSC colonies were manually picked and clonally expanded 3 weeks post transduction and transferred onto MEF plates.

**Sendai virus reprogramming.** Human dermal fibroblasts (Science cell Catalog number 2300) were reprogrammed according to the manufacturer’s instructions (CytoTune-iPS 2.0 Sendai Reprogramming Kit, Life technology catalog number A1378001). Cells were transfected with Sendi viruses containing the Yamanaka factors, and individual iPSC colonies were identified by morphology. The colonies were manually picked, expanded for 3 weeks post transduction and transferred onto a feeder layer of irradiated MEFs.

**mRNA reprogramming.** Human dermal fibroblasts (Science cell Catalog number 2300) were reprogrammed according to the manufacturer’s instructions (The Stemgent mRNA Reprogramming Kit catalog number 00–0071). Individual iPSC colonies were manually picked and clonally expanded three weeks post transfection and transferred onto MEF feeder layers.

**Cell culture.** PLAT-A-packaging cells (Cell Biolabs, Inc.) were maintained according to the manufacturer’s instructions. Human dermal fibroblasts (Science cell Catalog number 2300) were cultured in DMEM, 2 mM GlutaMax, 10% fetal bovine serum and 0.1 mM nonessential amino acids (Life Technologies), iPSCs were generated and maintained in standard hESC medium containing DMEM/F12 supplemented with 20% Knockout Serum Replacement (Life Technologies), 2 mM GlutaMAX, 0.1 mM nonessential amino acids, 0.1 mM 2-mercaptoethanol and 12 ng ml$^{-1}$ of Human Recombinant Fibroblast Growth Factor-basic (bFGF, Stemgent), HDFiPS cells were cultured on MEF feeder layers in hESC medium and mechanically passaged once a week. The hESC medium was changed daily. All cultures were tested and were negative for mycoplasma.

**DNA extraction and sequencing.** The Qiagen DNeasy Blood and Tissue Kit (catalog number 69504) was used to prepare genomic DNA from ~2 million cells of each cell line, as recommended by the manufacturer. Template DNA fragments (3 μg) were hybridized to the surface of flow cells HiSeq Paired-End cluster Kits (v2.5 or v3) and amplified to form clusters using the Illumina cBot. Paired-end libraries were sequenced for 2 × 101 cycles of incorporation and imaging using TruSeq SBS kits. Sequencing was performed at Illumina, Inc. (San Diego).

**Realignment of illumina reads and recalibration.** Reads were extracted from Illumina Casava aligned BAM files using the HTSLib by first shuffling the reads
and then extracting interleaved reads; see http://www.broadinstitute.org/gatk/guide/article?id=2908. These reads were then processed through the GATK Best Practices workflow for Variant Calling v2.6, which included first aligning the reads using BWA 0.7 with the ‘mem’ option, marking duplicates, pursuing local realignment, considering base quality recalibration and finally using the ‘reduce reads’ options. The BAM files generated from this process were used with the different variant calling methods.

**SNVs and indel variant calling.** HaplotypeCaller, as bundled with GATK v2.7, was used to call all 10 samples together. The variant calls were recalibrated using files in the GATK bundle that included data from HapMap, Omni, dbSNP, Mills Indels and 1,000 Genomes Indels databases. Variants falling in tranches 0–90 and 90–99.00 were used for downstream analysis. Unique variants were identified by looking at positions where only one sample had a non-reference allele. It should be noted that the HaplotypeCaller multisample calling pipeline described sacrifices specificity for sensitivity, as it is meant for population scale studies. The pipeline may favour the identification of variants common across the samples rather than variants or mutations unique to each. To gain a more sensitive assessment of the somatic SNV’s landscape across the iPSC samples, we also ran MuTect by treating the parent fibroblast population as the normal, and each iPSC cell line as a different derived cell (in an analogous manner to which tumours are treated with respect to germline samples in oncogenomics studies). MuTect was run with default settings and the calls were filtered using the judgment ‘KEEP’ option. Unique variants from the analysis were determined by intersecting the calls based on chromosomal coordinates and the variant calls.

**SNVs and indel validation and filtering.** To validate the identified variants, we split them into confidence intervals of high-confidence variants that had coverage between 40–60 × with the variant allele frequency in the range of 0.4–0.6; moderate-confidence variants were classified as having coverage 20–40 × with variant allele frequency between 0.4–0.6; and subclonal variants were those with variant allele frequencies between 0.2–0.4 and 40–60 ×. We validated these results using quantitative PCR. The ABL files associated with the variants were read in using the ‘abiply’ package (https://github.com/bow/abiply/), and looking at the calls made by the SOLID software. For any variants that appeared heterozygous, the relative amount of noise in the 10 upstream and downstream bases was evaluated based on the assumption that they should be homozygous. Any amplitude values that were two s.d. higher than the mean background noise were called as variant calls. Most of the calls made in this manner were validated through manual inspection (Supplementary Table 1).

**ENCODE TFBSs annotation.** Genomic coordinates (hg19) for transcription factor–bound regions of DNA, as curated by ENCODE (http://genomics.scripps.edu/ENHancer/) for nucleosome positioning. A SNV was considered damaging if it had a harmful designation by Condel, Polyphen or SIFT. For a SNV to be considered damaging, it had to be considered damaging if it had a harmful designation by Condel, Polyphen or SIFT. We also ran the variants through the OncoNator annotation web service (http://www.broadinstitute.org/oncnotator/) and found overlaps with MutTgen and Familial Syndrome Cancer Genes. Finally, the variants were fed through CADD annotation service through their online web service.

**PERMUTATION TESTING.** We leveraged permutation testing in an ANOVA setting to find the probability of observing the differences in mutation and variant rates across the three reprogramming methods. We exhaustively relabeled all the reprogramming types, recomputing the ANOVA statistic each time we relabeled the cell lines, thus determining exactly how likely it would be to observe our mutation profile differences across the reprogramming cell types given the three replicates for each of the three reprogramming methods (a total of 280 permutations). We tested each genomic feature separately using this permutation strategy. To calculate the relative rates of different types of mutations (for example, coding, non-coding and so on), we divided by the total number of mutations.

**ANOVA contrasts.** We also tested whether one method was different from the other two methods using contrast tests. Three different tests were employed for mRNA expression: retroviral versus all and Sendai virus versus all, using the contrasts (–2, 1, 1), (–1, –2, 1), (1, –1, –2), respectively. We employed an ANOVA fit to estimate the impact of the reprogramming strategy on the aggregated counts of the different variant classifications. We employed this strategy to look at the difference in the nominal counts for the different variant classifications, but we also subdivided by the type of variant (SNV, insertion or deletion), as well as the relative rates of each variant type. We further filtered reported results to classifications that had at least 10 variants.

**Comparison of CADD score distributions.** To compare the distribution of CADD Scores across the different samples, we employed a Kruksal–Wallis test. First, we looked at the differences between the replicates of the same reprogramming method to ensure that the variance within a group was not high. Next, we looked at pair-wise comparisons between the different reprogramming methods by pooling all the variants for the replicates into one distribution for the reprogramming method. This analysis revealed that the synthetic mRNA had a distribution skewed towards lower scores compared with the retroviral vectors and non-integrating Sendai virus. There was no viable difference between Sendai virus and non-integrating Sendai virus. However, further inspection of the distributions revealed that most of the variants still fell in the non-damaging designation of CADD scores, indicating that the results were not biologically significant.

**BioNano DNA labelling.** DNA was labelled following the IrisPrep Reagent Kit protocol (BioNano Genomics). Briefly, 900 ng of DNA was digested with 10 U of Nt.BspQI nicking endonuclease (New England Biolabs, Ipswich, MA) for 2 h at 37°C. Nick-digested DNA was then incubated for 1 h at 72°C with fluorescently labelled dUTP and Taq Polymerase (New England Biolabs). Taq ligase (New England Biolabs) was used in the presence of dNTPs for ligation of nicks. DNA was solubilized with GE-Lase (Epigenic, Madison, WI) and extracted DNA was drop dialyzed for 4 h. DNA concentrations were measured using the Quant-iT dsDNA Assay Kit (Life Technologies).

**BioNano DNA data collection.** Labelled and counterstained DNA samples were loaded into IrysChips (BioNano Genomics) and run on the Iris (BioNano Genomics) imaging instrument. Data were collected for each sample until ≥50-fold coverage of long molecules (>150 kbp) was achieved. The IrisView (BioNano Genomics) software package was used to detect individual linearized DNA molecules using the YOYO-1 counterstain and to determine the localization of labelled nick sites along each DNA molecule. Sets of single-molecule maps, equivalent to ~50 haploid coverage, for each sample were then used to build a full-genome assembly.

**BioNano de novo assembly.** De novo assembly of single molecules is accomplished using BioNano’s custom assembler software program based on an Overlap-Layout-Consensus paradigm. First, we started with pair-wise comparison of all molecules longer than 150 kbp and ≥5 labels to find all overlaps with a P value <5 × 10−10, and then we could construct a draft consensus genome map based on these overlaps. The draft map could be further refined by mapping single molecules to it and recalculating the label positions. Next, the maps were extended by aligning overlapping molecules to the maps and calculating a consensus in the extended regions. Finally, the genome maps were compared and merged where patterns match with a P value <10−15. The process of extension and merge was repeated five times before a final refinement was applied to ‘finish’ all genome maps. The result of this assembly is a genome map set entirely independent of any known reference or external data (Fig. 3). Statistics about N50 and percentage coverage of GRC37 are described in Supplementary Table 1.

**BioNano structural variation calls.** Structural variation was detected by examining the alignment profiles between the de novo assembled genome maps against the GRC37 human reference assembly. Significant discrepancies in (a) the distance or (b) the number of unaligned labels between adjacent aligned labels...
would indicate the presence of insertion and deletion events. We used two algorithms to call SV, and they differ in the way discrepant regions in the alignment (termed outliers) were handled. In the first algorithm, the reference and maps were split at outliers, and split maps were iteratively realigned. The alignments of the newly split maps would then pinpoint the locations of the insertion and deletion variants. We used an alignment \( P \) value of \( 10^{-12} \) and an outlier cutoff of \( 10^{-4} \) to call variants in all four cell line samples. In the second algorithm, the reference and genome maps were not split; instead, the global alignment profiles were kept with insertions and deletion events being intra-alignment gaps. For all four cell line samples, the alignment \( P \) value was \( 10^{-12} \) and the outlier cutoff was \( 10^{-4} \). Alignment \( P \) values are calculated using the algorithm described in Anantharaman et al.14.

BioNano identification of cell line-specific calls. A series of steps were used to identify cell line-specific variants. First, we conservatively selected only insertions and deletions detected by both calling algorithms. Using a 50% reciprocal size overlap cutoff, we cross-compared variants detected among all cell lines to identify those that were putatively cell line specific. Finally, we manually curated the candidate variants to ensure that (a) there were molecules supporting the variant allele in the cell line of interest and (b) there was no molecule supporting the variant allele in all other cell line samples.

References

1. Hussein, S. M. et al. Copy number variation and selection during reprogramming to pluripotency. Nature 471, 58–62 (2011).
2. Laurent, L. C. et al. Dynamic changes in the copy number of pluripotency and cell proliferation genes in human ESCs and iPS cells during reprogramming and in culture. Cell Stem Cell 8, 106–118 (2011).
3. Mayshar, Y. et al. Identification and classification of chromosomal aberrations in human induced pluripotent stem cells. Cell Stem Cell 7, 521–531 (2010).
4. Taapken, S. M. et al. Karotypic abnormalities in human induced pluripotent stem cells and embryonic stem cells. Nat. Biotechnol. 29, 313–314 (2011).
5. Gore, A. et al. Somatic coding mutations in human induced pluripotent stem cells. Nature 471, 63–67 (2011).
6. Peterson, S. E. & Loring, J. F. Genomic instability in pluripotent stem cells: implications for clinical applications. J. Biol. Chem. 289, 4578–4584 (2014).
7. Schlaeger, T. M. et al. A comparison of non-integrating reprogramming methods. Nat. Biotechnol. 33, 58–63 (2015).
8. McKenna, A. et al. The Genome Analysis Toolkit: a MapReduce framework for analyzing next-generation DNA sequencing data. Genome Res. 20, 1297–1303 (2010).
9. Cibulskis, K. et al. Sensitive detection of somatic point mutations in impure and heterogeneous cancer samples. Nat. Biotechnol. 31, 213–219 (2013).
10. Das, S. K. et al. Single molecule linear analysis of DNA in nano-channel labeled with sequence specific fluorescent probes. Nucleic Acids Res. 38, e177 (2010).
11. Haste, A. R. et al. Rapid genome mapping in nanochannel arrays for highly complete and accurate de novo sequence assembly of the complex Aegilops tauschii genome. PloS ONE 8, e58684 (2013).
12. Lam, E. T. et al. Genome mapping on nanochannel arrays for structural variation analysis and sequence assembly. Nat. Biotechnol. 30, 771–776 (2012).
13. Xiao, M. et al. Rapid DNA mapping by fluorescent single molecule detection. Nucleic Acids Res. 35, e16 (2007).
14. Anantharaman, T. S. & Mishra, B. in Proceedings of the First International Workshop on Algorithms in Bioinformatics 27–40 (Aarhus, Denmark; Springer-Verlag, 2001).
15. John, V. N. Genomic Mapping: a Statistical and Algorithmic Analysis of the Optical Mapping System (PhD thesis Univ. Southern California, 2010).
16. Valouev, A., Schwartz, D. C., Zhou, S. & Waterman, M. S. An algorithm for assembly of ordered restriction maps from single DNA molecules. Proc. Natl Acad. Sci. USA 103, 15770–15775 (2006).
17. Erikson, G. A., Deshpande, N., Kesarvani, B. G. & Torkamani, A. SG-ADVISER CNV: copy-number variant annotation and interpretation. Genet. Med. 17, 714–718 (2015).
18. Ramos, A. H. et al. Oncotator: cancer variant annotation tool, Hum. Mutat. 36, E2423–E2429 (2015).
19. Kandoth, C. et al. Mutational landscape and significance across 12 major cancer types. Nature 502, 333–339 (2013).
20. Kircher, M. et al. A general framework for estimating the relative pathogenicity of human genetic variants. Nat. Genet. 46, 310–315 (2014).
21. Baum, C. et al. Chance or necessity? Insertional mutagenesis in gene therapy and its consequences. Mol. Ther. 9, 5–13 (2004).
22. Takahashi, K. & Yamanaka, S. Induction of pluripotent stem cells from mouse embryonic and adult fibroblast cultures by defined factors. Cell 126, 663–676 (2006).
23. Okita, K., Ichisaka, T. & Yamanaka, S. Generation of germ-line-competent induced pluripotent stem cells. Nature 448, 313–317 (2007).
24. Korhonen, J., Martinmaki, P., Pizzi, C., Rastas, P. & Ukkonen, E. MOODS: fast search for position weight matrix matches in DNA sequences. Bioinformatics 25, 3181–3182 (2009).
25. Li, H. et al. The Sequence Alignment/Map format and SAMtools. Bioinformatics 25, 2078–2079 (2009).
26. Li, H. & Durbin, R. Fast and accurate short read alignment with Burrows-Wheeler transform. Bioinformatics 25, 1754–1760 (2009).

Acknowledgements

K.B. was supported by the National Science Foundation Graduate Research Fellowship under Grant No. DGE-1144086. K.L.N. was supported by an Autism Speaks Fellowship. N.J.S. and K.B. were supported in part by NIH Grants U19 AG023122 and R21 AG045789, and the Tanner Foundation. J.F.L., K.L.N., H.T. and R.W. were supported by grants from the California Institute for Regenerative Medicine CIRM (RT1-01108, RT3-07655, TR1-01250, TR3-05603 and CL1-00502) and NIH (R33 MH087925). M.R. was supported by NIHCRM, a NIH common fund program, and New York Stem Cell Foundation unrestricted funds. H.D., Z.D., E.H.C., A.W.C.P. and H.C. were supported by BioNano Genomics, Inc.

Author contributions

K.B., K.L.N. and R.W. analysed the Illumina whole-genome sequencing data. H.D., Z.D., E.H.C., A.W.C.P. and H.C. generated and analysed the BioNano mapping data. H.T. derived iPSC lines using the different reprogramming strategies and prepared samples for Illumina sequencing and de novo mapping. K.B., K.L.N., H.C., M.R., N.J.S. and J.F.L. conceived the study and wrote the manuscript with contributions from all the authors.

Additional information

Accession codes: The whole-genome sequencing data have been deposited in the NCBI database under accession codes: BioProject ID: PRJNA304745; SRA database: SRP067962. The BioNano Genomics mapping data are available at http://bxninstall.com/publicdatasets/ipsc_Bhutani_et_al_2015.zip

Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

Competing financial interests: H.D., Z.D., E.H.C., A.W.C.P. and H.C. are employees of BioNano Genomics, Inc.

Reprints and permission information is available online at http://www.nature.com/reprintsandpermissions/

How to cite this article: Bhutani, K. et al. Whole-genome mutational burden analysis of three pluripotency induction methods. Nat. Commun. 7:10536 doi: 10.1038/ncomms10536 (2016).