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## Title Page

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Title: Differentiation defective phenotypes revealed by large scale analyses of human pluripotent stem cells

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#### Abstract

We examined the gene expression and DNA methylation of 49 hiPSCs and 10 hESCs and found overlapped variations in gene expression and DNA methylation in the two types of human pluripotent stem cell lines. Comparisons of the in vitro neural differentiation of 40 hiPSCs and 10 hESCs showed that seven hiPSC clones retained a significant number of undifferentiated cells even after neural differentiation culture, and formed teratoma when transplanted into mouse brains. These differentiation-defective hiPSC clones were marked by higher expression levels of several genes, including those expressed from long terminal repeats of specific human endogenous retroviruses. These data demonstrated a subset of hiPSC lines that have aberrant gene expression and defective potential in neural differentiation, which need to be identified and eliminated prior to applications in regenerative medicine.


## Significance statement

In the past few years, the findings have been controversial in regard to whether human induced pluripotent stem cells (hiPSCs) are distinct from human embryonic stem cells (hESCs) in their molecular signatures and differentiation properties. In this study, hiPSCs and hESCs have overlapping variations in molecular signatures such as RNA expression and DNA methylation. On the other hand, some hiPSC clones retained a significant number of undifferentiated cells even after neural differentiation culture, and formed teratoma when transplanted into mouse brains. These differentiation-defective hiPSC clones were marked by higher expression levels of several genes, including those expressed from long terminal repeats of specific human endogenous retroviruses. They need to be identified and eliminated prior to applications in regenerative medicine.

## INTRODUCTION

Human pluripotent stem cells possess a robust potential for proliferation and provide useful sources of cells for regenerative medicine and drug discovery. Two types of human pluripotent stem cells have been generated: human embryonic stem cells (hESCs) derived from blastocysts (1), and induced pluripotent stem cells (hiPSCs), which are generated from somatic cells by factor-mediated reprogramming $(2,3)$.

In the past few years, the findings have been controversial in regard to whether hESCs and hiPSCs are distinct cell types. Some reports have argued that they could not be clearly distinguished (4-6), whereas others have reported that they have differences in their gene expression (7-10), DNA methylation (10-13), and capacity for differentiation (14). In the latter papers, relatively small numbers of cell lines were generally compared. In addition, most comparisons used pluripotent cell lines from various laboratories, such that the observed differences may be attributable to lab-specific variations due to technical differences (15).

To overcome these problems, we compared the mRNA and microRNA (miRNA) expression levels and DNA methylation between 10 hESCs and 49 hiPSCs that had been cultured under the same conditions. Furthermore, we compared the in vitro directed neural differentiation of these pluripotent stem cells.

## RESULTS

## Overlapped variations of the mRNA expression and DNA methylation in hiPSCs and hESCs

We analyzed a total of 49 hiPSCs derived from four types of somatic cells, including human dermal fibroblasts (HDFs), dental-pulp stem cells (DP), cord blood cells (CB) and peripheral blood mononuclear cells (PBMN), generated using three gene delivery methods, including those employing retroviruses, non-integration episomal plasmids and Sendai viruses (Table 1 and Dataset S1). Most clones were generated in our own laboratory, except for three clones that were established in another laboratory (16). Prior to the analyses of the gene/miRNA expression (Figs. 1A, B) and DNA methylation (Fig. 1C), we cultured these hiPSCs, as well as 10 hESCs , under the same culture conditions for at least three passages. In addition, we analyzed the original somatic cells, two human embryonic carcinoma cell (ECC) lines (NTera2 cloneD1 and 2102Ep 4D3) and three cancer cell lines (HepG2, MCF7 and Jurkat).

The mRNA microarray analyses (Fig. 1A) identified 61 probes with significant differences in expression between hESCs and hiPSCs ( $t$-test: FDR<0.05). Each of the 61 probes showed variable expression among both the hESCs and hiPSCs, and the distributions of the expression levels in the two groups overlapped (Fig. 1D). Of note,
hESCs established at Kyoto University (Kyoto hESCs) were more similar to hiPSCs than to the remaining hESCs (other hESCs) in their expression of 15 probes, that were differentially expressed between hESCs and hiPSCs (FDR < 0.05 and FC > 3) (Fig. S1A). In contrast, hierarchical clustering using all probes showed no clear cut separation among Kyoto hESCs, other ESCs, and iPSCs, indicating that the similarity of Kyoto ESCs to iPSCs are confined to a small set of genes (Fig. S1B). In addition, the miRNA array analyses (Fig. 1B) did not find any significant differences between hESCs and hiPSCs ( $t$-test: FDR $<0.05$ ). The expressions of hsa-miR-886-3p and hsa-miR-142-3p tended to be higher in hiPSCs, but the expression levels of these miRNAs showed overlapped variations among hiPSCs and hESCs (Fig. S1A).

We next compared the global DNA methylation status between hiPSCs and hESCs by the Illumina Infinium Human Methylation27 BeadChip assay. Among 27,445 CpG dinucleotides examined, we did not identify significantly differentially methylated CpG regions (CG-DMR) between the hESCs and hiPSCs (Mann-Whitney's U test, FDR<0.05) (Fig. 1C).

We then validated the CG-DMRs reported in previous studies. Three studies identified a total of 205 regions as CG-DMRs, including 130, 71 and four regions identified by comparing five hiPSCs and two hESCs (13), three versus three (12) and
nine versus three (10) cell lines, respectively. Of the 205 regions, 46 regions containing 66 CpG dinucleotides were covered by the Infinium platform used in our study (Table S1A). Based on the methylation levels in our hiPSCs and hESCs, these CpGs were clustered into three groups (Fig. 1E). Two-thirds of these CpGs belonged to group A: they tended to be highly methylated in hESCs, ECCs and cancer cell lines, whereas they tended to be hypomethylated in hiPSCs, as well as somatic cells. However, they were also hypomethylated in Kyoto ESCs (17). The methylation status of the upstream region of the $P O N 3$ gene, a representative example of CpGs in a Group A, was confirmed by pyrosequencing (Fig. 1F). Thus, the CpG methylation status in Group A may distinguish some, but not all, hESCs from hiPSCs. Seventeen CpGs belonged to group B, which showed similar methylation levels in both hESCs and hiPSCs. Five CpGs, representing three genes, belonged to group C and showed higher methylation levels in some, but not all, hiPSCs compared to hESCs. The remaining hiPSCs showed low methylation levels, comparable to those in hESCs. We confirmed the methylation status of a representative example of CpGs in a group C , the upstream region of the TCERG1L gene, by pyrosequencing; the methylation levels were low in 21 of the 49 hiPSCs (Fig. 1G). Therefore, the CpGs in group C may distinguish some, but not all, iPSCs from hESCs.

A previous report (12) showed that many CG-DMRs were located in CpG shores, rather than CpG islands. Since only a few (9 out of 71) of their CG-DMRs were covered by our Infinium platform, we analyzed five CG-DMRs in CpG shores by pyrosequencing, including A2BP1, IGF1R, POU3F4, PTPRT and ZNF184 (Fig. S1C). We detected significant differences in the averaged DNA methylation levels between hESCs and hiPSCs in four out of five genes. However, variations of DNA methylation levels in hESCs and hiPSCs are overlapped. They may distinguish some, but not all, hiPSCs from hESCs.

## A subset of hiPSC clones retain undifferentiated cells after neural differentiation

To examine whether hESCs and hiPSCs have comparable differentiation potential, we performed in vitro directed differentiation into neural stem and progenitor cells using the modified serum-free floating culture of embryoid body-like aggregates (SFEBq) method (Fig. 2A) (18). We initially performed the neural induction of two hESCs and 21 hiPSCs. Fourteen days after induction, the differentiation efficiency was evaluated based on the expression of an early neural marker, polysialylated neural cell adhesion molecule (PSA-NCAM). We found that all hESCs and hiPSCs differentiated into PSA-NCAM ${ }^{+}$cells with more than $80 \%$ efficiency (Fig. 2B). We also quantified the expression levels of the early neural marker, $P A X 6$, and the late neural marker, $M A P 2$, in
neurospheres by qRT-PCR (Fig. S2A). All the examined hES/iPSCsexpressed PAX6 at $>100$-fold higher levels, and MAP2 at $>20$-fold higher levels in comparison to undifferentiated H9 hESC. However, in some hiPSCs, we noticed slightly lower differentiation efficiency than the remaining hiPSCs and hESCs (Fig. 2B). This lower efficiency in neural differentiation was inversely correlated with a higher proportion of OCT3/4 ${ }^{+}$and TRA1-60 ${ }^{+}$undifferentiated cells (Fig. 2C). We also detected residual undifferentiated cells after a different neural differentiation protocol using adhesion culture (19) (Fig. S2B).

We then increased the number of clones and examined the proportions of OCT3/4 ${ }^{+}$ undifferentiated cells after neural induction from 10 hESCs and 40 hiPSCs . The 50 clones were ranked according to their proportions of OCT3/4 ${ }^{+}$cells on day 14 (Dataset S1 and Fig. 2D). The proportions of OCT3/4 ${ }^{+}$cells varied from $0 \%$ to $\sim 20 \%$. Thirty-eight clones, including nine hESCs and 29 hiPSCs, showed less than $1 \%$ OCT3 $/ 4^{+}$cells in all experiments. We designated these clones as "good" clones. On the other hand, seven hiPSCs contained more than $10 \%$ OCT3/4 ${ }^{+}$cells after neural differentiation in at least one experiment. We designated these clones as "differentiation-defective" clones. Clones which were not "good" or "defective" were categorized as "intermediate".

## Activation of specific LTR7 elements in "differentiation-defective" clones

To identify molecular signatures that can predict "differentiation-defective" clones, we compared the global gene expression patterns of 38 "good" clones and seven "differentiation-defective" clones under the culture conditions used for the undifferentiated state. We identified 19 probes (13 putative transcripts) that showed > 5-fold differences in expression, with a FDR <0.05, shown by magenta dots in Fig. 3A and listed in Table S1B.

Of the 19 probes identified, five probes recognized HHLAl (human endogenous retrovirus-H LTR-associating 1). Previous reports have shown that HHLAI is regulated by a long terminal repeat (LTR) of a human endogenous retrovirus-H (HERV-H) (20). The LTR in HHLA1 is classified as LTR7. Moreover, among the genes recognized by the 19 probes, we found that at least two others, $A B H D 12 B$ and C4orf51, also contained LTR7 sequences in their gene bodies. According to a microarray analysis, we confirmed that these three LTR7-containing genes were upregulated in the "differentiation-defective" hiPSCs, as well as the nullipotent hECC line 2102Ep 4D3(21), but they were expressed at lower levels in the "good" hiPSCs, hESCs, pluripotent hECC line NTera2 cloneD1, while they were almost not expressed in the
original somatic cells (Fig. 3B).

The Agilent microarray platform has 12 probes, including two reverse probes $[\mathrm{d}(\mathrm{r})$, $\mathrm{f}(\mathrm{r})$ ], for HHLAl and its neighboring gene, OC90, which is reported to make a fusion transcript with HHLA1 (20) (Fig. 3C). Among them, seven probes located downstream of LTR7 showed higher expression levels in "differentiation-defective" clones than in "good" clones (Figs. 3C, D). Similarly, there are two probes for ABHD12B, designed for exons 4 and 13 (Fig. 3C). Only the exon 13 probe, located downstream of LTR7, showed a higher expression in "differentiation-defective" clones than in "good" clones (Fig. 3D). We also performed an exon array (Affymetrix) of ABHD12B and C4orf51, and found that exons downstream of LTR7 were preferentially upregulated in "differentiation-defective" hiPSC clones (Fig. 3E). We also found that the methylation status of LTR7s in these three genes were lower in "differentiation-defective" hiPSC clones than in "good" clones (Fig. 3F). These results indicate that the three genes are transcribed from activated LTR7.

DNA hypomethylation exists in some, but not all, of LTR7s in "differentiation-defective" clones

According to the Repeatmasker software program, there are 3523 LTR7 elements in
the human genome. In order to extract microarray probes that are potentially affected by LTR7s, we first selected genes containing LTR7s in their gene bodies or regions 2 kb upstream from their transcription start sites. We then retrieved the microarray probes located between each LTR7 and the 3' end of the corresponding gene body. As a result, we selected 763 probes as LTR7-related probes (Fig. S3A and Table S1C). We found that most of these probes showed comparable expression levels in "good" and "defective" lines (Fig. S3B), with the exception of some probes, such as those corresponding to $A R R B 1, F A A H 2$ and $T B C 1 D 23$, that were differentially expressed between "good" and "defective" clones (FDR < 0.05 and FC >2) and showed slightly higher expression in "defective" lines.

We then checked the DNA methylation status of the LTR7 regions in these three genes and three other genes ( $D N M T 3 B, A B C A 1$ and $A P P$ ) whose expression levels were not significantly different between the "good" and "defective" clones. By pyrosequencing and clonal bisulfite sequencing, we found that the LTR7 regions in four genes (ARRB1, $F A A H 2, T B C 1 D 23$ and APP) were hypomethylated in "defective" clones compared to "good" clones. In contrast, the LTR7 regions in two genes (DNMT3B and ABCA1) did not show such hypomethylation (Fig. S3C). Therefore, the activation of LTR7 is not confined to HHLA1, ABHD12B and C4orf51; DNA hypomethylation exists in some, but
not all, of LTR7s in "defective" hiPSCs.

## "Differentiation-defective" hiPSC clones form teratomas in mouse brains

To further evaluate the "defective" hiPSCs, we induced their differentiation into dopaminergic neurons, which were then transplanted into the striata of nonobese diabetic/severe combined immune deficient (NOD/SCID) mouse brains (Fig. 4A). Thirty and sixty days after transplantation, we obtained T2-weighted images of the mouse brains with a magnetic resonance imaging (MRI) scanner to observe the graft sizes at the transplanted sites (Fig. 4B). The quantification of the MR images showed that "defective" hiPSC clones resulted in significantly larger graft sizes than "good" clones (Fig. 4C). Notably, some mice that had received "defective" clones died or developed symptoms that required euthanasia before day 60 (Table S2). Therefore, we could not obtain the graft size data on day 60 in these mice.

To identify the composition of the surviving grafts, we performed a histological analysis of the brains of animals that died or that became moribund after transplantation. The remaining healthy mice were sacrificed 14-41 weeks after transplantation. Sections were stained with hematoxylin and eosin (HE). Thirty-six out of the 42 grafts ( $85.7 \%$ ) from "defective" clones contained non-neural lineage tissues, such as intestine-like
epithelial cells, cartilage or mesenchymal cells (Fig. 4D and Fig. S4A). In contrast, grafts from "good" clones largely consisted of neural tissues. Immunostaining confirmed grafts were positive for human NCAM (Fig. S4B). A qRT-PCR analysis of pre-transplanted cells from "defective" clones revealed higher expression levels of OCT3/4, suggesting that some undifferentiated cells still remained even after 29 days neural induction (Fig. 4E). We then depleted the TRA-1-60 cells on day 22 during neural induction, and transplanted cells on day 29. The TRA-1-60-depleted cells from "defective" clones resulted in significantly smaller grafts, which did not contain non-neural tissues (Fig. S4C).

We also observed that 14 out of the $63(22.2 \%)$ grafts from "good" clones, including those from hESCs, contained a non-neural component in the graft tissue after transplantation (Fig. 4D and Table S2), although these clones did not show high expression levels of OCT3/4 in the pre-transplantation samples (Fig. 4E). We referred to these clones as "type 2 defective" clones, which were distinct from "type 1 defective" clones that contained $O C T 3 / 4^{+}$undifferentiated cells in the pre-transplantation samples. We observed higher expression levels of SOX17 (an endoderm marker) and GSC (an endoderm and mesoderm marker) in the pre-transplantation samples of "type 2 defective" clones (Figs. 4F, G), demonstrating the presence of other lineages in these
pre-transplantation samples.

## Discussion

We identified two types of "defective" pluripotent stem cell lines in this study. The first type consisted of hiPSCs that retained a substantial number of undifferentiated cells after in vitro directed neural differentiation. Seven out of the 40 iPSCs (17.5\%) examined in this study fell into this category. In contrast, we did not observe such defects in any of the 10 hESCs . More clones should be analyzed to confirm that hESCs are free from this deficiency. Nevertheless, it is likely that "type 1 defectiveness" is more common in hiPSCs than in hESCs. The "type 1 defective" hiPSCs are accompanied by an aberrant epigenetic status. Among the 13 putative transcripts that were highly expressed in these defective clones, at least three were expressed from the LTR of endogenous retroviruses. Normally, these LTRs are silenced by various epigenetic modifications, including DNA methylation (22-24). In "type 1 -defective" iPSC clones, LTR locus in the three genes showed lower DNA methylation levels than in "good" clones and original somatic cells. Notably, the same regions were hypo-methylated in the nullipotent hECC line, 2102Ep 4D3, suggesting that the loss of DNA methylation in these LTR locus is correlated with the lower ability to differentiate. At present, the biological significance and relationship between activation of specific

LTRs and the defective phenotype is unclear. Recent reports showed that ERV may play roles in the establishment and maintenance of transcription network in pluripotent stem cells (25, 26). Furthermore, updated annotations revealed that one of differentially expressed probes in "type 1 defective" hiPSCs (A_19_P00325604) encoded large intergenic non-coding RNA regulator of reprogramming (Linc-ROR), which contained LTR7 in its 5' region. Linc-ROR is reported to have multiple roles in the induction and maintenance of pluripotency $(27,28)$. Future studies should be undertaken to clarify why these epigenetic abnormalities occur and how they are related to the defective differentiation.

Kim et al. (29) showed that there is an inverse correlation between the hsa-mir-371-373 expression and the efficiency of neural differentiation. They also showed that KLF4 may induce the expression of hsa-mir-371-373. In our analyses, the hsa-mir-371-373 cluster was highly expressed in all the seven "type 1 defective" hiPSC clones (Fig. S5A). However, the cluster was also highly expressed in many "good" clones. KLF4 was highly expressed in some "defective" clones (Fig. S5B), and four out of six retroviral "defective" clones failed to silence KLF4 retroviral transgenes (Fig. S5C). There was no correlation between the OCT3/4 transgene expression and "type 1 defectiveness" (Fig. S5D). Taken together, these findings indicate that high expression
levels of the hsa-mir-371-373 cluster, KLF4 and transgenes cannot function as absolute markers for "type 1 defectiveness".

We previously reported that the origin of mouse iPSCs was a major determinant of defectiveness in directed neural differentiation; mouse iPSCs from adult tail tip fibroblasts showed the highest incidence of resistance to differentiation (30). In the present study using human iPSCs, five out of seven "type 1 -defective" clones were derived from fibroblasts of donors of various ages, and six out of the seven clones were generated using retroviruses (Table 1). This may suggest that "type 1 defectiveness" is associated with fibroblast origin and retroviral induction. However in this study, most of the fibroblast-derived iPSCs were generated by retroviruses, and most of the non-fibroblast iPSCs were generated by non-retroviral methods. Future studies will need to be undertaken to determine whether the origin or the generation method (or both) has a significant impact on the frequency of "type 1 differentiation-defective" iPSCs.

The second type of "defective" group includes hiPSCs and hESCs that contained differentiated cells of non-neural lineages after in vitro directed differentiation into dopaminergic neurons. We have previously shown that the optimal conditions for hepatic differentiation are different for each clone (16). By optimizing the protocols, it
may be possible to induce complete neural differentiation to avoid "type 2 defective" clones. Alternatively, purification of neural cells using a cell sorter may work to avert "type 2 defective" clones.

Several studies have reported sets of genes of which DNA methylation status are different between hiPSCs and hESCs. We validated these CG-DMRs and found that many of them can distinguish some hESCs from hiPSCs (Group A in Fig. 1E). They are highly methylated in some hESCs, but not in hiPSCs or original somatic cells. Thus these CG-DMRs may represent epigenetic memories of somatic cells in iPSCs. However, we found a set of hESCs that showed low methylation status of these CG-DMRs, which were comparable to hiPSCs. We also found another set of the reported CG-DMRs that showed high methylation status in some hiPSCs, but not in original somatic cells or hESCs (Group C in Fig. 1E). These likely represent aberrant methylation associated with reprogramming. However, we also found many hiPSCs showed normal methylation patters of these CG-DMRs. A more recent study identified nine genes that can segregate hiPSCs from hESCs in DNA methylation and gene expression (31). However, we did not observe such a clear distinction in gene expression of these genes between our hiPSCs and hESCs (Fig. S6). Two of these genes, TCERG1L and FAM19A5 may distinguish some, but not all, hiPSCs from hESCs.

In our analyses, 35 hiPSCs had records of the donor's genetic background; 14 were derived from Caucasians and 21 were from Japanese subjects (Dataset S1). Thus, the similarity of some signatures between the Kyoto hESCs and our hiPSCs cannot be attributed to the racial or ethnic backgrounds of the donors. Another possible cause of the differences is the method used to establish the hESCs and the subsequent culture conditions. The Kyoto hESCs were generated on feeders consisting of a 1:1 mixture of MEFs and SL10 cells (17, 32), which were subcloned from STO cells. Most of our hiPSCs were established on SNL feeders, which were also derived from STO cells. A recent report showed that the feeders have profound effects on established hiPSCs (33). To confirm the importance of the culture conditions, more studies comparing $\mathrm{hESC} / \mathrm{hiPSCs}$ established under different conditions will be needed.

In conclusion, our results revealed that a subset of hiPSCs is defective in neural differentiation and marked with activation of endogenous retroviruses. We also confirmed that some hiPSCs are different from hESCs in molecular signatures, including CG-DMRs that have been previously reported. It remained to be determined whether these molecular signatures specific for some hiPSCs have functional consequences.

## Materials and Methods

The gene expression profiling was carried out using the SurePrint G3 human GE microarray (Agilent). Most of the data were analyzed using the Gene spring GX 11.5.1 software program (Agilent Technologies). Neural induction was performed as described previously (18). Detailed descriptions of methods in this article are available in SI Materials and Methods.

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## References

1. Thomson JA, et al. (1998) Embryonic stem cell lines derived from human blastocysts. Science 282(5391):1145-1147.
2. Takahashi K, et al. (2007) Induction of pluripotent stem cells from adult human fibroblasts by defined factors. Cell 131(5):861-872.
3. Yu J, et al. (2007) Induced pluripotent stem cell lines derived from human somatic cells. Science 318(5858):1917-1920.
4. Bock C, et al. (2011) Reference Maps of human ES and iPS cell variation enable high-throughput characterization of pluripotent cell lines. Cell 144(3):439-452.
5. Newman AM \& Cooper JB (2010) Lab-Specific Gene Expression Signatures in Pluripotent Stem Cells. Cell Stem Cell 7(2):258-262.
6. Guenther MG, et al. (2010) Chromatin structure and gene expression programs of human embryonic and induced pluripotent stem cells. Cell Stem Cell 7(2):249-257.
7. Chin MH, et al. (2009) Induced pluripotent stem cells and embryonic stem cells are distinguished by gene expression signatures. Cell Stem Cell 5(1):111-123.
8. Marchetto MC, et al. (2009) Transcriptional signature and memory retention of human-induced pluripotent stem cells. PLoS ONE 4(9):e7076.
9. Ghosh Z, et al. (2010) Persistent donor cell gene expression among human induced pluripotent stem cells contributes to differences with human embryonic stem cells. PLoS ONE 5(2): e 8975
10. Ohi Y, et al. (2011) Incomplete DNA methylation underlies a transcriptional memory of somatic cells in human iPS cells. Nat Cell Biol 13(5):541-549.
11. Deng J, et al. (2009) Targeted bisulfite sequencing reveals changes in DNA methylation associated with nuclear reprogramming. Nat Biotechnol 27(4):353-360.
12. Doi A, et al. (2009) Differential methylation of tissue- and cancer-specific CpG island shores distinguishes human induced pluripotent stem cells, embryonic stem cells and fibroblasts. Nat Genet 41(12):1350-1353.
13. Lister R, et al. (2011) Hotspots of aberrant epigenomic reprogramming in human induced pluripotent stem cells. Nature 471(7336):68-73.
14. Hu BY, et al. (2010) Neural differentiation of human induced pluripotent stem cells follows developmental principles but with variable potency. Proc Natl Acad Sci USA 107(9):4335-4340.
15. Yamanaka $S$ (2012) Induced pluripotent stem cells: past, present, and future. Cell Stem Cell 10(6):678-684.
16. Kajiwara M, et al. (2012) Donor-dependent variations in hepatic differentiation from human-induced pluripotent stem cells. Proc Natl Acad Sci $U S$ A 109(31):12538-12543.
17. Suemori H, et al. (2006) Efficient establishment of human embryonic stem cell lines and long-term maintenance with stable karyotype by enzymatic bulk passage. Biochem Biophys Res Commun 345(3):926-932.
18. Morizane A, Doi D, Kikuchi T, Nishimura K, \& Takahashi J (2011) Small-molecule inhibitors of bone morphogenic protein and activin/nodal signals promote highly efficient neural induction from human pluripotent stem cells. J Neurosci Res 89(2):117-126.
19. Kriks S, et al. (2011) Dopamine neurons derived from human ES cells efficiently engraft in animal models of Parkinson's disease. Nature 480(7378):547-551.
20. Kowalski PE, Freeman JD, \& Mager DL (1999) Intergenic splicing between a HERV-H endogenous retrovirus and two adjacent human genes. Genomics 57(3):371-379.
21. Bahrami AR, Matin MM, \& Andrews PW (2005) The CDK inhibitor p27 enhances neural differentiation in pluripotent NTERA2 human EC cells, but does not permit differentiation of 2102Ep nullipotent human EC cells. Mech Dev 122(9):1034-1042.
22. Rowe HM \& Trono D (2011) Dynamic control of endogenous retroviruses during development. Virology 411(2):273-287.
23. Hutnick LK, Huang X, Loo TC, Ma Z, \& Fan G (2010) Repression of retrotransposal elements in mouse embryonic stem cells is primarily mediated by a DNA methylation-independent mechanism. J Biol Chem 285(27):21082-21091.
24. Stoye JP (2012) Studies of endogenous retroviruses reveal a continuing evolutionary saga. Nat Rev Microbiol 10(6):395-406.
25. Macfarlan TS, et al. (2012) Embryonic stem cell potency fluctuates with endogenous retrovirus activity. Nature 487(7405):57-63.
26. Santoni FA, Guerra J, \& Luban J (2012) HERV-H RNA is abundant in human embryonic stem cells and a precise marker for pluripotency. Retrovirology 9:111.
27. Loewer S , et al. (2010) Large intergenic non-coding RNA-RoR modulates reprogramming of human induced pluripotent stem cells. Nat Genet 42(12):1113-1117.
28. Wang Y, et al. (2013) Endogenous miRNA sponge lincRNA-RoR regulates Oct4, Nanog, and Sox2 in human embryonic stem cell self-renewal. Dev Cell 25(1):69-80.
29. Kim H, et al. (2011) miR-371-3 expression predicts neural differentiation propensity in human pluripotent stem cells. Cell Stem Cell 8(6):695-706.
30. Miura K, et al. (2009) Variation in the safety of induced pluripotent stem cell lines. Nat Biotechnol 27(8):743-745.
31. Ruiz S, et al. (2012) Identification of a specific reprogramming-associated epigenetic signature in human induced pluripotent stem cells. Proc Natl Acad Sci USA 109(40):16196-16201.
32. Kawase E, et al. (1994) Strain difference in establishment of mouse embryonic stem (ES) cell lines. Int J Dev Biol 38(2):385-390.
33. Tomoda K, et al. (2012) Derivation conditions impact x-inactivation status in female human induced pluripotent stem cells. Cell Stem Cell 11(1):91-99.

## FIGURE LEGENDS

Fig. 1. hiPSCs and hESCs have overlapped variations in RNA expression and DNA

## methylation

(A)-(C) Scatter plots of the mRNA expression (A), miRNA expression (B) and DNA methylation (C) data comparing the average of 49 hiPSC lines ( y -axis) to the average of 10 hESC lines (x-axis) are shown. The RNA expression value is shown on a $\log 2$ scale. Green lines indicate two-fold differences in the RNA expression levels between the clones. Differentially expressed probes $(t$-test, FDR $<0.05)$ are shown in magenta. (D) The variations in the mRNA expression levels of 61 differentially expressed probes in hESCs (red) and hiPSCs (black) are shown. Probes are arranged in order of the absolute value of the fold change (FC) between hESCs and hiPSCs. (E) The DNA methylation profiles for CpGs contained in reported hES-hiPS DMRs and overlapping with our platform. Probes are arranged in order of the differences between the average DNA methylation level of hESCs and that of hiPSCs. The heat map represents the DNA methylation levels from completely methylated (= 1 , magenta) to unmethylated $(=0$, white) samples. (F)(G) The methylation status of the upstream region of PON3 (F) and TCERG1L (G) was examined by pyrosequencing.

Fig. 2. A differentiation-defective phenotype in a subset of hiPSC clones. (A) A schematic diagram of the SFEBq method used for neural differentiation. (B) Neural
induction was performed for two hESC and 21 hiPSC lines which were established from various origins by retroviral or episomal vector methods. On day 14 , we examined the proportion of PSA-NCAM-expressing cells by flow cytometry ( $\mathrm{n}=2$ ). (C) The proportions of PSA-NCAM- (white), OCT3/4- (grey) and TRA1-60- (black) positive cells 14 days after neural differentiation. (D) The proportions of OCT3/4 ${ }^{+}$cells on day 14 after neural differentiation are ranked in order of their maximum value. The numbers in parentheses show the number of trials.

Fig. 3 Activation of specific endogenous retroviral LTR7s in "defective" clones
(A) A scatter plot of the mRNA expression data comparing the average of 38 "good" clones ( y -axis) to the average of seven "defective" clones (x-axis). Green lines indicate five-fold differences in expression. A total of 19 differentially expressed probes are colored magenta. (B) The expression levels of LTR7-related genes (HHLA1, ABHD12B and C4orf51) were examined by microarray. (C) A schematic diagram of three LTR7-related genes. HHLA1 and OC90 are neighboring genes. Dots indicate microarray probes. Magenta dots show probes which are located after LTR7 regions, which were upregulated in "defective" clones. (D) A scatter plot of array probes which recognized LTR7-related genes and two other genes (EFR3A and KCNQ3) which are genes
neighboring HHLA1 and OC90, respectively. (E) The exon array of the ABHD12B and C4orf51 genes. The average levels of the normalized exon expression are shown. (F) The DNA methylation status of LTR7 and its neighboring regions of HHLA1, $A B H D 12 B$ and C4orf51 was examined by pyrosequencing. (n.s.; not significant, *; $\mathrm{p}<0.05$, **; p<0.01, Mann-Whitney's U-test)

Fig. 4 Transplantation of neural cells derived from hiPSCs and hESCs into mouse

## brains

(A) A schematic diagram of the SFEBq method used for DA neural differentiation. On day 29, the cells were transplanted into NOD/SCID mouse brains. (B) Magnetic resonance images of coronal sections of the grafted brains. The section surface of grafted cells indicated as the white shadow in right brain was measured as described in the lower panels. (C) A box-and-whisker plot of the surface sizes of graft sections 30 and 60 days after transplantation. The median, quartile and range are shown. $* \mathrm{t}$-test, $\mathrm{p}<0.05$. (D) The proportion of each kind of graft. Grafts were categorized according to their components as determined in HE sections, and were classified by the proportion of neural tissues by a microscopic observation. (E)-(G) The expression levels of the undifferentiated cell marker, OCT3/4 (E), the endoderm marker, SOX17 (F), and the
mesoderm and endoderm marker, GSC (G), in pre-transplantation cultures, undifferentiated hESC lines and somatic cells (HDF and DP) were examined by qRT-PCR. The colors of the dots were identical to the proportion of neural tissues in Fig.

4D.

Table 1. A summary of the iPSC clones used in this report

|  |  | Method used to generate clones |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Origin |  | Retrovirus | Episomal plasmid | Sendaivirus | Total |
|  | HDFs | $22(5,0)$ | $3(0,0)$ | $0(0,0)$ | $25(5,0)$ |
|  | DP cells | $1(0,0)$ | $2(1,1)$ | $0(0,0)$ | $3(1,1)$ |
|  | CB | $3(1,2)$ | $4(0,0)$ | $5(0,0)$ | $12(1,2)$ |
|  | PBMN | $0(0,0)$ | $5(0,1)$ | $4(0,0)$ | $9(0,1)$ |
|  | Total | $26(6,2)$ | $14(1,2)$ | $9(0,0)$ | $49(7,4)$ |

Total clone numbers with "type 1 defective" clone number and "type 2 defective" clone number in parentheses are shown.

Figure. 1


## Figure. 2

A



## 




Figure. 3


## Figure. 4



## SI Material and Methods

## Cell culture

The hESCs and hiPSCs were maintained in Primate ES cell medium (ReproCELL) supplemented with $4 \mathrm{ng} / \mathrm{ml}$ of human recombinant basic fibroblast growth factor (bFGF, Wako) on SNL feeders (1-3). Human dermal fibroblasts (HDFs) were obtained from the Japanese Collection of Research Bioresources (JCRB) or were purchased from Cell Applications, Inc. Dental pulp (DP) cells were kindly provided by Dr. Ken-ichi Tezuka (Gifu University Graduate School of Medicine). HDFs and hECC lines were maintained in Dulbecco's modified Eagle medium (DMEM, Nacalai tesque) containing $10 \%$ fetal bovine serum (FBS, Thermo) and $0.5 \%$ Penicillin/Streptomycin (Pen/Strep, Life Technologies). DP cells were cultured in MSBGM medium (Lonza). CD34 ${ }^{+}$cord blood cells were obtained from the Stem Cell Resource Network in Japan (Banks at Miyagi, Tokyo, Kanagawa, Aichi, and Hyogo) through the RIKEN BioResource Center (Tsukuba, Ibaraki, Japan). The peripheral blood was harvested from healthy donors whose written informed consent was obtained in accordance with the institutional review board requirements. The mononuclear cells were then isolated by density gradient centrifugation with Ficoll-paque plus (GE healthcare).

## Generation of human iPSCs

The generation of hiPSCs from HDFs, DP cells and blood samples using a retroviral system or episomal vectors was performed as described previously (1, 4-6). TKCBV4-2, 5-6 and TKCB7-2 iPS cells were kindly provided by Drs. Koji Eto and Naoya Takayama (7).

During the generation of hiPSCs from blood using Sendai viral vectors, vectors encoding OCT3/4, SOX2, KLF4 and c-MYC (CytoTune-iPS, DNAVEC) were infected into $\mathrm{CD} 34^{+}$cells at a multiplicity of infection of 3 or 10 in $\alpha$ MEM medium supplemented with $10 \%$ FBS, $50 \mathrm{ng} / \mathrm{ml}$ IL-6, $50 \mathrm{ng} / \mathrm{ml} \mathrm{sIL-6R} 50 \mathrm{ng} /$,ml SCF, $10 \mathrm{ng} / \mathrm{ml}$ TPO, $20 \mathrm{ng} / \mathrm{ml}$ Flt3 ligand and $20 \mathrm{ng} / \mathrm{ml}$ IL-3. The next day, the infected cells were centrifuged to remove residual virus, plated onto 6-well plates covered with MEF feeder cells and cultured in Primate ES cell medium supplemented with $4 \mathrm{ng} / \mathrm{ml}$ of bFGF until colonies were formed. Sendai virus infection and the generation of iPSCs from $\alpha \beta T$ cells were carried out as described previously (8).

## RNA extraction

We lysed the cells at subconfluent density using the Trizol reagent (Life Technologies), and total RNA was purified by a standard protocol. The RNA
concentration and purity were determined through measurement of the A260/280 ratios with a Nanodrop instrument (Thermo Scientific). For microarrays, confirmation of the RNA quality was performed using the Agilent 2100 Bioanalyzer (Agilent Technologies).

## mRNA expression analysis

The gene expression profiling was carried out using the SurePrint G3 Human GE Microarray (Agilent) according to the manufacturer's protocol. The data were analyzed using the Gene spring GX 11.5.1 software program (Agilent Technologies). The data processing was performed as follows: (i) Threshold raw signals were set to 1.0 , (ii) Log base 2 transformation was performed, (iii) $75^{\text {th }}$ percentile normalization was chosen as the normalized algorithm (http://genespring-support.com/faq/normalization). The flag setting was performed as follows: feature is not positive and significant (not detected), not uniform (compromised), not above background (not detected), saturated (compromised) or is a population outlier (compromised). Control probes were removed and only the "detected" probes that were present in at least one sample in all hES/hiPS cell samples were used for the further analysis. The number of probes used in the analysis was 36,757 (Fig. 1A) and 36,083 (Fig. 3A).

## microRNA microarray analysis

The miRNA expression profiling was carried out using the Agilent Human miRNA microarray Rel 12.0 according to the manufacturer's protocol. The data were analyzed using the Gene spring GX 11.5.1 software program (Agilent Technologies) and data processing was performed in the same way as for the mRNA expression analysis, except that $90^{\text {th }}$ percentile normalization was chosen as the normalized algorithm. The number of probes used in the analysis was 476 (Fig. 1B).

## Genomic DNA extraction and bisulfite treatment

Genomic DNA extraction and purification from cultured cells was carried out using a Gentra Puregene kit (QIAGEN). Extracted DNA was quantitated by using the Nanodrop instrument, and the quality was assessed by gel electrophoresis. A total of 500 ng of genomic DNA was treated with bisulfite using the EZ DNA Methylation-Gold Kit (Zymo Research Corp., Irvine, CA) according to the manufacturer's protocol.

## DNA methylation analysis with a beads-array

Genome-wide DNA methylation profiling was performed using the Illumina Infinium Human Methylation27 BeadChip (Illumina). Bisulfite-converted DNA was
used, and the remaining assay steps were performed using the reagents supplied by Illumina and their specified conditions. The readout from the array was a $\beta$-value, which was defined as the ratio between the fluorescent signal from the methylated allele to the sum of both methylated and unmethylated alleles, and thus correlated with the level of DNA methylation. A $\beta$-value of 1.0 corresponds to complete methylation and 0 is equal to no DNA methylation. To exclude potential sources of technical bias, we only used CpG sites with detection $P$ values < 0.05 in at least 56 out of 59 samples. Normalization was not performed. The number of probes used in the analysis was 27,445 (Fig. 1E).

## Generation of heat maps

We used Microsoft Excel to visualize the values as heat maps. The color spectrum expands from green (lower value) to magenta (higher value) through black in the gene/miRNA expression analysis, and from white (hypomethylation) to magenta (hypermethylation) in the DNA methylation analysis.

## Bioinformatic analysis

A hierarchical clustering analysis was performed using the Gene Spring GX 11.5.1 software program.

## DNA methylation analysis with pyro- and clonal- sequencing

Pyrosequencing was carried out with primers designed using the Pyromark Assay Design Software program, ver. 2.0 (QIAGEN). The primer sequences are shown in

Table S1D. PCR was performed in a $25 \mu \mathrm{~L}$ reaction mixture containing 25 ng bisulfite-converted DNA, 1X Pyromark PCR Master Mix (QIAGEN), 1X Coral Load Concentrate and $0.2 \mu \mathrm{M}$ forward and 5 ' biotinylated reverse primers. The PCR conditions were 45 cycles of $95^{\circ} \mathrm{C}$ for $30 \mathrm{~s}, 56^{\circ} \mathrm{C}$ for 30 s and $72^{\circ} \mathrm{C}$ for 30 s. The PCR product was bound to streptavidin sepharose beads (GE Healthcare, Uppsala, Sweden), and was purified, washed, denatured and washed again. Then, $0.4 \mu \mathrm{M}$ of the sequencing primer was annealed to the purified PCR product. Pyrosequencing reactions were performed using the PSQ HS 96 Pyrosequencing System. The degree of methylation is shown as the percentage of methylated cytosines divided by the sum of methylated and unmethylated cytosines (percentage of 5 mC ). Bisulfite-clonal sequencing was performed as previously described (9). In Fig. 4F and Fig. S3C, we examined the DNA methylation status of LTR7 and its neighboring regions in "defective" clones with high expression levels of these genes $(\mathrm{n}=6$; TKCBV5-6, TIG118-4f1, TIG108-4f3, 1503-4f1, TIG107-3f1 and 451F3), "good" clones with low expression levels of these genes ( $\mathrm{n}=$

6; 703B1, 606B1, 201B7, H1, 253G1 and 454E2), two hECC lines (NTera2 and 2102Ep) and two somatic cell lines (HDF and DP).

## Exon arrays

We performed the exon array of the $A B H D 12 B$ and $C 4 o r f 51$ genes for three "defective" clones with high expression levels of these genes ( $\mathrm{n}=3$, TKCBV5-6, TIG108-4f3 and 451F3), three "good" clones with low expression levels of these genes ( $\mathrm{n}=3,201 \mathrm{~B} 7$, H1 and 253G1) and somatic cells (HDF and DP74). cDNA was generated with the WT expression kit (Ambion) per the manufacturer's instructions. The cDNA was fragmented and end-labeled with a GeneChip WT Terminal labeling kit (Affymetrix).

Approximately $5.5 \mu \mathrm{~g}$ of labeled DNA target was hybridized to the Affymetrix GeneChip Human Exon 1.0 ST Array at $45^{\circ} \mathrm{C}$ for 17 h , per the manufacturer's recommendations. Hybridized arrays were washed and stained on a GeneChip Fluidics Station 450 and scanned on a GCS3000 Scanner (Affymetrix). An exon array data analysis was performed using the Gene Spring GX11.5.1 software program employing ExonPLIER16 with core and extended probe sets. The exon probe sets were accepted if they had a DABG p-value $<0.05$ in at least one of the samples.

## Selection of microarray probes related to LTR7

The RepeatMasker database classifies sequences into subgroups by in silico analysis.

We used LTR7 sequences registered in Repeatmasker open 3.3.0 - Repeat Library 20110920, a database of human repetitive sequence (http://www.repeatmasker.org/species/homSap.html). According to this database, there are 3523 LTR7 elements, including LTR7A, 7B, 7C in the human genome. In order to extract microarray probes that were potentially affected by LTR7 elements, we first selected genes containing LTR7 elements in their gene bodies or in the regions 2 kb upstream from their transcription start sites. We then extracted microarray probes that were located between the each LTR7 and the 3' end of the corresponding gene body (The direction of LTR7 was not considered). As a result, we extracted 763 probes corresponding to 435 genes from the Agilent human G3 microarray as LTR7-related probes (Table S1C).

## Quantitative RT-PCR

To remove any potential contamination by genomic DNA, we treated purified RNA samples with a Turbo DNA free kit (Ambion). After DNase treatment, reverse
transcription was performed with the oligo $\mathrm{dT}_{20}$ primer using a ReverTra Ace- $\alpha$-kit (Toyobo). qRT-PCR was performed with SYBR Premix Ex Taq II (Takara), and samples were analyzed with the StepOne plus real-time PCR system (Applied Biosystems). The primer sequences are shown in Table S1D. The relative expression level was calculated by using plasmid DNA containing the PCR product (Fig 4 and Figs S5C,D).

## Neural induction

We performed the neural differentiation of human pluripotent stem cells with the quick method for serum-free embryoid body formation (SFEBq) as described previously (10). In brief, hESCs and hiPSCs treated with $10 \mu \mathrm{M} \mathrm{Y}-27632$ were dissociated into single cells and transferred at 9000 cells per well to 96 -well low cell adhesion plates (Lipidure-Coat Plate A-U96; NOF Corporation). The cells were cultured for 14 days in DFK5 medium consisting of DMEM/F-12 (Life Technologies), 5\% Knockout Serum Replacement (KSR, Life Technologies), 1\% MEM-non-essential amino acids (Life Technologies), 2 mM L-glutamine (Life Technologies), 0.1 mM 2-mercaptoethanol (Life Technologies) and $0.5 \%$ penicillin/streptomycin. We used DFK5 medium supplemented with $10 \mu \mathrm{M}$ Y-27632, $2 \mu \mathrm{M}$ dorsomorphin (SIGMA) and $10 \mu \mathrm{M}$ SB431542 (SIGMA) for the first four days.

Adhering neural differentiation of Dual SMAD inhibition was performed as described previously (11). Briefly, the cells were plated on matrigel-coated plate and after reaching to $90 \%$ confluency, they were cultured with 100 nM LDN193189 (Stemgent), $10 \mu \mathrm{M}$ SB431542 in $15 \%$ knockout serum replacement, 2 mM L-glutamine and $10 \mu \mathrm{M}$ $\beta$-mercaptoethanol-conteining D-MEM for 12 days.

For dopaminergic differentiation, we first transferred ESCs or iPSCs onto 96-well low cell adhesion plates with Y-27632, dorsomorphin and SB431542 in the same way as indicated for SFEBq. We supplemented the cultures with $100 \mathrm{ng} / \mathrm{ml}$ FGF8 (Peprotech) and $20 \mathrm{ng} / \mathrm{ml}$ WNT1 (Peprotech) from day five to 12, and with $200 \mathrm{ng} / \mathrm{ml}$ SHH (R\&D) from day eight to 12 . Twelve days after induction, aggregates were transferred onto 6-well plates coated with laminin (Becton-Dickinson) and Poly-L-ornithine (SIGMA) and were cultured with Neurobasal medium (Life Technologies) containing 2\% B27 supplement (Life Technologies), 2 mM L-glutamine and $0.5 \%$ penicillin/streptomycin. We added $200 \mathrm{ng} / \mathrm{ml}$ SHH to the medium from day 12 to 15 , and $1 \mathrm{ng} / \mathrm{ml}$ FGF20 and $12.5 \mathrm{ng} / \mathrm{ml}$ bFGF from day 15 to 22 . On day 22 , the cells were dissected into clumps and plated on new 6-well plates coated with laminin and Poly-L-ornithine, and were then cultured with Neurobasal medium supplemented with $2 \mathrm{ng} / \mathrm{ml}$ GDNF (R\&D), 20 $\mathrm{ng} / \mathrm{ml}$ BDNF (R\&D), 400 mM dbcAMP (SIGMA) and 200 mM Ascorbic Acid
(SIGMA) until day 29. TRA-1-60-positive cell labeling and depletion were performed on day 22 using an Anti-TRA-1-60 MicroBead kit and the autoMACS pro device (Miltenyi Biotech).

## Flow cytometric analysis

Neural aggregates were dissociated and processed for the flow cytometric analysis by a FACS Aria II instrument (Becton-Dickinson). To analyze the proportion of OCT3/4+ cells, the cells were fixed with $3.7 \%$ formaldehyde, permeabilized with $0.2 \%$ TritonX-100 and stained with the appropriate antibody. To count the number of PSA-NCAM ${ }^{+}$cells or TRA-1-60 ${ }^{+}$cells, cells were prepared without fixation. To eliminate the number of dead cells from the total cell population, we stained the cells with propidium iodide after labeling them with the anti-TRA-1-60 or anti-PSA-NCAM antibody, or with red fluorescent reactive dye from the LIVE/DEAD Fixable Dead Cell Stain Kits (Invitrogen) before fixing the cell suspension.

## Transplantation of ES/iPS cell-derived dopaminergic neuron cultures into the brains of NOD/SCID mice

To prepare samples for injection, we scraped and mechanically dissected cells by
gently pipetting them up and down a few times, suspended them in culture medium (1 x $10^{6}$ cells $/ \mu \mathrm{l}$ ) and injected $2 \mu \mathrm{l}$ of the cell suspension into the right striatum ( 2 mm lateral, 1 mm rostral to the bregma; depth, 3 mm from the dura) of NOD/SCID mice (6 weeks old, female) using a glass micropipette, as described previously (12).

## Magnetic resonance imaging

Graft imaging was performed with a MRmini SA instrument (DS Pharma Biomedical) by using a cylindrical slotted holder with a 20 mm radio frequency coil constructed for mice. T2-weighted images (repetition time $=2000 \mathrm{~ms}$, echo time $=69 \mathrm{~ms}$ ) were recorded. The brains were imaged coronally in a single section through the graft center by using an image matrix of $256 \times 128$, a field of view of $2 \times 4 \mathrm{~cm}^{2}$, and two excitations. Parametric images were generated by using the Sampler XP-NI software program (DS Pharma Biomedical). Graft section surfaces were measured by using the INTAGE Realia Professional imaging software program (CYBERNET).

## Immunostaining

Anti-NCAM (ERIC1) antibody (Santa Cruz) was used as a primary antibody and anti-Mouse Ig Biotin (Dako) was used as a secondary antibody for
immunocytechemistry.

## Statistical analysis

## Gene/miRNA expression

We conducted the $t$-test (variances assumed equal) for the normalized, filtered data and controlled the false discovery rate (FDR) at 0.05 using the Benjamini-Hochberg method to identify probes that differed significantly between hESCs and hiPSCs, or for neural differentiation "good" and "defective" clones.

## DNA methylation determined using the beads-array

We conducted Mann-Whitney's U-test on the filtered data controlling the FDR at 0.05 using the Benjamini-Hochberg method to identify probes that differed significantly between hESCs and hiPSCs.

Mann-Whitney's U-test was used to compare the quantitative methylation values between hESCs and hiPSCs or"defective" and "good" groups. Calculations were carried
out with the Statview software program.

## Graft size after transplantation

The $t$-test was used to compare the graft sizes derived from "defective" and "good" clones (Fig. 4C). In the case of comparisons between graft sizes between unsorted and depleted cell cultures, we performed the paired $t$-test.

1. Takahashi K, et al. (2007) Induction of pluripotent stem cells from adult human fibroblasts by defined factors. Cell 131(5):861-872.
2. Fujioka T, Yasuchika K, Nakamura Y, Nakatsuji N, \& Suemori H (2004) A simple and efficient cryopreservation method for primate embryonic stem cells. Int J Dev Biol 48(10):1149-1154.
3. Ohnuki M, Takahashi K, \& Yamanaka S (2009) Generation and characterization of human induced pluripotent stem cells. Curr Protoc Stem Cell Biol Chapter 4:Unit 4A 2.
4. Okita K, et al. (2011) A more efficient method to generate integration-free human iPS cells. Nat Methods 8(5):409-412.
5. Tamaoki N, et al. (2010) Dental Pulp Cells for Induced Pluripotent Stem Cell Banking. J Dent Res 89(8):773-778.
6. Okita K, et al. (2013) An efficient nonviral method to generate integration-free human-induced pluripotent stem cells from cord blood and peripheral blood cells. Stem Cells 31(3):458-466.
7. Kajiwara M, et al. (2012) Donor-dependent variations in hepatic differentiation from human-induced pluripotent stem cells. Proc Natl Acad Sci $U$ S A 109(31):12538-12543.
8. Seki T, et al. (2010) Generation of induced pluripotent stem cells from human terminally differentiated circulating T cells. Cell Stem Cell 7(1):11-14.
9. Takahashi K \& Yamanaka $S$ (2006) Induction of pluripotent stem cells from mouse embryonic and adult fibroblast cultures by defined factors. Cell 126(4):663-676.
10. Morizane A, Doi D, Kikuchi T, Nishimura K, \& Takahashi J (2011) Small-molecule inhibitors of bone morphogenic protein and activin/nodal signals promote highly
efficient neural induction from human pluripotent stem cells. J Neurosci Res 89(2):117-126.
11. Kriks S, et al. (2011) Dopamine neurons derived from human ES cells efficiently engraft in animal models of Parkinson's disease. Nature 480(7378):547-551.
12. Miura K, et al. (2009) Variation in the safety of induced pluripotent stem cell lines. Nat Biotechnol 27(8):743-745.

## Supplemental figure legends.

Fig. S1. Gene expression patterns among Kyoto hESCs, other hESCs and hiPSCs and DNA methylation levels of previously reported ES-iPS DMRs in our cell lines
(A) Heat maps showing the expression levels of 15 probes that were differentially expressed between hESCs and hiPSCs (FDR < 0.05 and absolute FC > 3) and hsa-miR-142-3p and hsa-miR-886-3p in various cell lines. A hierarchical clustering analysis for 15 probes was performed using the Euclidean distance and average linkage algorithm. (B) A hierarchical clustering analysis of the global gene expression patterns in various cell lines was performed using the Euclidean distance and average linkage algorithm in the Gene Spring GX 11.5.1 software program.
(C) Some previously reported hES-iPS DMRs; A2BP1, IGF1R, ZNF184, POU3F4 and PTPRT were examined by pyrosequencing in $10 \mathrm{hESCS}, 49 \mathrm{hiPSCs}$ in our laboratory. Each CpG dinucleotide position was assayed in triplicate, and average values were plotted. Mann-Whitney's U-test was used to compare the quantitative methylation
values between hESCs and hiPSCs. (n.s.; not significant, *; $\mathrm{p}<0.05$, ** ; p<0.01)

Fig S2. Most of the cells can differentiate into neural cells although some clones retain undifferentiated cells after neural differentiation
(A) The expression levels of PAX6 (left panel) and MAP2 (right panel) in neurospheres of differentiated hESC line H9 and hiPSC lines (253G1, TIG108-4f3 and TKCBV5-6) were determined by quantitative RT-PCR. The expression levels in hESC line H9 before differentiation were set to 1 , and relative expression levels were presented in log scale. (B) A comparison of the proportion of TRA-1-60-positive cells after neural induction between the SFEBq method (white) and the adhesion culture method (black).

Fig. S3. Activation of LTR7 is not confined to HHLA1, ABHD12B and C4orf51; DNA hypomethylation exists in some, but not all, of LTR7s in "defective" hiPSC clones.
(A) The extraction of 763 probes corresponding to 435 genes as LTR7-related probes from the Agilent human G3 microarray (design ID 028004). (B) A comparison of the expression levels of 763 LTR7-related probes between "good" and "defective" clones. Magenta-colored genes (ABHD12B, HHLA1 and C4orf51) and yellow-colored genes (ARRB1, FAAH2 and TBC1D23) are differentially expressed between the "good" and "defective" clones (FDR $<0.05$ and FC $>5$ or FC $>2$,
respectively) and green-colored genes (DNMT3B, ABCA1 and APP) did not show any differences. (C) The DNA methylation status of LTR7 and its neighboring regions of ARRB1, FAAH2 and TBC1D23 were examined by pyro-sequencing and those of DNMT3B, ABCA1 and APP were examined by clonal-sequencing. Mann-Whitney's U-test was used to compare the quantitative methylation values between "defective" and "good" clones (n.s.; not significant, *; $\mathrm{p}<0.05$ ).

Fig. S4. The histology of grafts derived from hESCs and hiPSCs
(A) Transplanted mouse brains were fixed with formaldehyde, embedded in paraffin, sectioned, and stained with hematoxylin and eosin (HE). The ratios indicate neural cells and non-neural cells as determined by a microscopic observation. Scale bar, $500 \mu \mathrm{~m}$.
(B) HE-stained sections (left panel) and human NCAM-stained sections (right two panels) of mouse brains transplanted with 29 day-differentiated hESC/iPSCs. (C) The maximum surface size of graft sections 45 or 60 days after transplantation. Transplanted cells were prepared with or without depletion of TRA-1-60 ${ }^{+}$cells 22 days after differentiation. $*$ t-test, paired $\mathrm{p}<0.05$

Fig. S5 High expression levels of the hsa-mir-371-373 cluster, KLF4 and transgenes are not absolute markers for "type 1 defectiveness"

The expression levels of hsa-mir-371-373 (A) and KLF4 (B) were examined by a microarray analysis in seven "defective" clones, five "intermediate" clones, 38 "good" clones, two hECC lines (NTera2 and 2102Ep), six somatic cell lines (HDF, DP, CB1,CB2, PBMN1, PBMN2) and three cancer cell lines (HepG2, MCF7 and Jurkat). The total and retroviral transgene expression levels of KLF4 (C) and OCT3/4 (D) were measured by qPCR in 18 hiPS clones established by a retroviral method, the "defective" clone 451F3, which was generated using an episomal plasmid vector, four hESCs, two hECCs and two somatic cell lines (HDF, DP).

Fig. S6. previously reported iPS-specific aberrantly methylated genes' expressions

A heat map for 10 hESCs and 49 hiPSCs examined in our laboratory based on the gene expression levels of reported aberrantly methylated genes that can distinguish hiPSCs and hESCs. Of nine previously reported genes, probes for C22ORF34 were "not detected" in all the samples in our microarray platform, so we only evaluated the other 8 genes (8 probes, "detected" in at least two clones of our samples).

Figure.S1
A
mRNA expression

( $\log 2$ expression)


B



CpG1 CpG2


CpG1
CpG2



CpG4


Figure.S2
A


B

differentiated

Figure.S3

## A

LTR7(A), B, C $\quad \rightarrow 3523$ loci in whole human genome
$\rightarrow 658$ loci in gene bodies in Refseq genes


B


C


DNMT3B LTR7 (12 CpGs)


FAAH2 LTR7 (2 CpGs)


ABCA1 LTR7 (7 CpGs)


TBC1D23 LTR7 (3 CpGs)


APP LTR7 (6 CpGs)


Figure S4

(neural):(non-neural tissue)

B

(neural : non-neural tissue)

C

("defective" clones)
$\triangle$ 451F3
("good" clones)

- TIG108-4f3 (day60)
- TIG108-4f3 (day45)
- 1503-4f1
- TKCBV5-6
- TIG107-4f1

Figure S5


Figure. S6


Table S1A. List of CpGs reported as differentially methylated between hESCs and hiPSCs (only covered by the infinium platform)

|  | Chromosomal Coordinates (hg19 version) |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| cg number (infinium) | gene symbol | Chr. number | position of CpG (infinium) | start | end | reference | reported methylation status |
| cg23045073 | NPHS2 | chr1 | 179545458 | 179545343 | 179546543 | $\begin{gathered} \text { Doi et al., } \\ 2009 \end{gathered}$ | ES < iPS |
| cg13722123 | GRM1 | chr6 | 146350346 | 146349702 | 146350414 |  | ES > iPS |
| cg22411207 | MOS | chr8 | 57026301 | 57025648 | 57026328 |  | ES $>$ iPS |
| cg27205791 | COPZ1 | chr12 | 54719080 | 54718379 | 54719446 |  | ES $>\mathrm{iPS}$ |
| cg10316764 | TAOK2 | chr16 | 29984665 | 29984318 | 29984860 |  | ES < iPS |
| cg21756353 | DHPS | chr19 | 12792451 | 12791946 | 12792452 |  | ES > iPS |
| cg03257423 | ZNF228 | chr19 | 44860820 | 44860544 | 44862008 |  | ES > iPS |
| cg23850272 | ZNF228 | chr19 | 44861241 | 44860544 | 44862008 |  | ES $>\mathrm{iPS}$ |
| cg03000603 | DHX34 | chr19 | 47852595 | 47852502 | 47853505 |  | ES $>$ iPS |
| cg03020597 | SLITRK2 | chrX | 144898810 | 144897718 | 144898947 |  | ES < iPS |
| cg11108890 | VAMP5 | chr2 | 85811471 | 85811204 | 85812203 | Lister et al., 2010 | ES > iPS |
| cg25651505 | VAMP5 | chr2 | 85812023 | 85811204 | 85812203 |  | ES $>$ iPS |
| cg11024597 | ECRG4 | chr2 | 106681411 | 106681383 | 106682982 |  | $E S>\mathrm{iPS}$ |
| cg10885338 | ECRG4 | chr2 | 106682640 | 106681383 | 106682982 |  | ES $>$ iPS |
| cg17802847 | CFLAR | chr2 | 201980910 | 201980570 | 201982169 |  | ES $>$ iPS |
| cg27020690 | TERC | chr3 | 169482358 | 169482261 | 169483360 |  | ES $>$ iPS |
| cg01389761 | TERC | chr3 | 169482968 | 169482261 | 169483360 |  | $E S>\mathrm{iPS}$ |
| cg00936626 | PIGZ | chr3 | 196694856 | 196693558 | 196695157 |  | $E S>\mathrm{iPS}$ |
| cg10088985 | CXCL5 | chr4 | 74864313 | 74863700 | 74864699 |  | ES $>$ iPS |
| cg04263186 | TACR3 | chr4 | 104640489 | 104640415 | 104642014 |  | ES > iPS |
| cg05389335 | TACR3 | chr4 | 104641319 | 104640415 | 104642014 |  | ES $>$ iPS |
| cg25358289 | CD14 | chr5 | 140012728 | 140011238 | 140012837 |  | $E S>\mathrm{iPS}$ |
| cg11538128 | ZNF354C | chr5 | 178487481 | 178487016 | 178488015 |  | ES $>$ iPS |
| cg04488521 | ZNF354C | chr5 | 178487716 | 178487016 | 178488015 |  | ES $>$ iPS |
| cg08126211 | KAAG1 | chr6 | 24357720 | 24357674 | 24358773 |  | ES $>$ iPS |
| cg04515001 | DCDC2 | chr6 | 24358236 | 24357674 | 24358773 |  | $E S>\mathrm{iPS}$ |
| cg16306115 | DCDC2 | chr6 | 24358306 | 24357674 | 24358773 |  | ES $>\mathrm{iPS}$ |
| cg00463577 | C6orf150 | chr6 | 74161911 | 74160732 | 74162431 |  | ES $>$ iPS |
| cg09527362 | C6orf150 | chr6 | 74162142 | 74160732 | 74162431 |  | ES > iPS |
| cg08109815 | NMBR | chr6 | 142409831 | 142409560 | 142410659 |  | $E S>\mathrm{iPS}$ |
| cg17256157 | NMBR | chr6 | 142410100 | 142409560 | 142410659 |  | $E S>\mathrm{iPS}$ |
| cg07260592 | LPA | chr6 | 161100122 | 161099863 | 161100862 |  | $E S>\mathrm{iPS}$ |
| cg05158615 | NPY | chr7 | 24323559 | 24323459 | 24325058 |  | ES $>$ iPS |
| cg12614105 | NPY | chr7 | 24324435 | 24323459 | 24325058 |  | $E S>\mathrm{iPS}$ |
| cg10329418 | PON3 | chr7 | 95026181 | 95025448 | 95026447 |  | $\mathrm{ES}>\mathrm{iPS}$ |
| cg24750391 | PON3 | chr7 | 95026211 | 95025448 | 95026447 |  | ES $>$ iPS |
| cg26952662 | CTHRC1 | chr8 | 104383499 | 104383030 | 104384629 |  | ES $>$ iPS |
| cg19188612 | CTHRC1 | chr8 | 104384291 | 104383030 | 104384629 |  | ES > iPS |


| cg10303487 | DPYS | chr8 | 105479058 | 105478430 | 105479429 |  | $E S>i P S$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| cg20774846 | DPYS | chr8 | 105479420 | 105478430 | 105479429 |  | ES > iPS |
| cg10175795 | TCERG1L | chr10 | 133109194 | 133108510 | 133111409 |  | ES < iPS |
| cg03943081 | TCERG1L | chr10 | 133110646 | 133108510 | 133111409 |  | ES < iPS |
| cg08896945 | CALCB | chr11 | 15095068 | 15094797 | 15096396 |  | ES $>$ iPS |
| cg11784785 | SLC6A5 | chr11 | 20620089 | 20618097 | 20620296 |  | ES $>\mathrm{iPS}$ |
| cg15842276 | MTNR1B | chr11 | 92702648 | 92702225 | 92703224 |  | ES $>\mathrm{iPS}$ |
| cg20424530 | IRAK4 | chr12 | 44152509 | 44151863 | 44153462 |  | $E S>i P S$ |
| cg08992050 | IRAK4 | chr12 | 44152940 | 44151863 | 44153462 |  | ES $>$ iPS |
| cg20360244 | SLC35E3 | chr12 | 69140065 | 69139863 | 69140962 |  | $E S>\mathrm{iPS}$ |
| cg03469054 | KIAA1944 | chr12 | 130387861 | 130386777 | 130389576 |  | $E S<i P S$ |
| cg13234863 | KIAA1944 | chr12 | 130389138 | 130386777 | 130389576 |  | ES < iPS |
| cg23054883 | FZD10 | chr12 | 130647580 | 130643277 | 130649076 |  | ES < iPS |
| cg14912575 | C14orf162 | chr14 | 70038236 | 70038201 | 70039200 |  | ES > iPS |
| cg00815605 | ACOT2 | chr14 | 74035882 | 74035701 | 74037300 |  | $E S>\mathrm{iPS}$ |
| cg26780333 | ACOT4 | chr14 | 74058972 | 74058301 | 74059300 |  | $E S>\mathrm{iPS}$ |
| cg15309006 | LOC63928 | chr16 | 23766116 | 23765733 | 23766732 |  | $E S>\mathrm{iPS}$ |
| cg08085267 | C17orf57 | chr17 | 45401833 | 45401491 | 45402590 |  | ES $>\mathrm{iPS}$ |
| cg07177852 | CCDC68 | chr18 | 52626476 | 52625891 | 52626890 |  | ES $>\mathrm{iPS}$ |
| cg24673765 | HSPB6 | chr19 | 36247869 | 36246382 | 36248581 |  | ES $>$ iPS |
| cg15125472 | HSPB6 | chr19 | 36248077 | 36246382 | 36248581 |  | ES $>$ iPS |
| cg15925792 | MX1 | chr21 | 42798131 | 42797679 | 42799278 |  | $E S>\mathrm{iPS}$ |
| cg22152328 | MX1 | chr21 | 42798386 | 42797679 | 42799278 |  | ES $>$ iPS |
| cg00540544 | CSRP1 | chr1 | 201476297 | 201475983 | 201476336 | $\begin{gathered} \text { Ohi et al., } \\ 2011 \end{gathered}$ | $E S>\mathrm{iPS}$ |
| cg17780098 | CSRP1 | chr1 | 201476311 | 201475983 | 201476336 |  | ES $>\mathrm{iPS}$ |
| cg01626227 | TRIM4 | chr7 | 99517289 | 99516971 | 99517450 |  | ES $>\mathrm{iPS}$ |
| cg12927772 | C9orf64 | chr9 | 86571585 | 86571560 | 86571999 |  | ES $>$ iPS |
| cg23268677 | COMT (TXNRD2) | chr22 | 19929097 | 19929072 | 19929357 |  | $E S>\mathrm{iPS}$ |

Table S1B. List of probes which showed differentially expression between 38 "good" clones and 7 "defective" clones (FC>5, FDR<0.05)

| Putative transcript cluster | ProbeName | GeneSymbol | Description | Genomic Coordinates | Chromosome Strand_Avadis |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | A_23_P417821 | DMRTB1 | Homo sapiens DMRT-like family B with proline-rich C terminal, 1 (DMRTB1), mRNA [NM_033067] | $\begin{array}{\|c} \text { chr1:53933088- } \\ 53933147 \end{array}$ | + |
| 2 | A_33_P3236436 | C4orf51 | Homo sapiens chromosome 4 open reading frame 51 (C4orf51), mRNA [NM_001080531] | chr4:146653856- 146653915 | + |
| 3 | A_19_P00316694 | XLOC_005614 | BROAD lincRNAs version v2 (http://www.broadinstitute.org/genome_bio/human_lincrnas/) | $\begin{array}{\|c\|c\|} \hline \text { chr6:14281107- } \\ 14281048 \end{array}$ | - |
|  | A_19_P00320902 |  | lincRNA:chr6:14283301-14285685 reverse strand | $\begin{array}{\|c} \hline \text { chr6:14284343- } \\ 14284284 \end{array}$ | - |


|  | A_19_P00321571 |  | lincRNA:chr6:14283035-14285450 reverse strand | $\begin{gathered} \text { chr6:14285417- } \\ 14285358 \end{gathered}$ |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 4 | A_33_P3297020 | PSORS1C3 | Homo sapiens psoriasis susceptibility 1 candidate 3 (nonprotein coding) (PSORS1C3), non-coding RNA [NR_026816] | $\begin{gathered} \text { chr6:31141572- } \\ 31141513 \end{gathered}$ | - |
| 5 | A_32_P3955 | C7orf57 | Homo sapiens chromosome 7 open reading frame 57 (C7orf57), mRNA [NM_001100159] | $\begin{gathered} \text { chr7:48099967- } \\ 48100026 \end{gathered}$ | + |
| 6 | A_19_P00329841 |  | lincRNA:chr7:124873114-124899839 reverse strand | $\begin{gathered} \text { chr7:124873767- } \\ 124873708 \end{gathered}$ | - |
| 7 | A_19_P00331472 |  | lincRNA:chr8:129599518-129624118 reverse strand | $\begin{gathered} \text { chr8:129599638- } \\ 129599579 \end{gathered}$ | - |
| 8 | A_33_P3301050 | HHLA1 | Homo sapiens HERV-H LTR-associating 1 (HHLA1), mRNA [NM_001145095] | $\begin{gathered} \text { chr8:133073792- } \\ 133073733 \end{gathered}$ | - |
|  | A_19_P00325595 |  | lincRNA:chr8:133071643-133092468 reverse strand | $\begin{gathered} \text { chr8:133073545- } \\ 133073486 \end{gathered}$ | - |
|  | A_19_P00330523 |  | lincRNA:chr8:133071643-133092468 reverse strand | $\begin{array}{\|c\|} \text { chr8:133073455- } \\ 133073396 \end{array}$ | - |
|  | A_19_P00321436 | (HHLA1) | lincRNA:chr8:133073732-133075753 reverse strand | $\begin{array}{\|c\|} \text { chr8:133073792- } \\ 133073733 \end{array}$ | - |
|  | A_19_P00319204 | (HHLA1) | lincRNA:chr8:133076031-133093351 reverse strand | $\begin{array}{\|c\|} \hline \text { chr8:133076236- } \\ 133076177 \end{array}$ | - |
| 9 | A_19_P00327099 |  | lincRNA:chr8:138387843-138421643 reverse strand | $\begin{array}{\|c} \text { chr8:138395981- } \\ 138395922 \end{array}$ | - |
| 10 | A_23_P366035 | ABHD12B | Homo sapiens abhydrolase domain containing 12B (ABHD12B), transcript variant 1, mRNA [NM_001206673] | $\begin{array}{\|c} \text { chr14:51371212- } \\ 51371271 \end{array}$ | + |
| 11 | A_19_P00325604 |  | lincRNA:chr18:54721302-54731677 reverse strand | $\begin{gathered} \text { chr18:54722409- } \\ 54722350 \end{gathered}$ | - |
| 12 | A_23_P38959 | VAV1 | Homo sapiens vav 1 guanine nucleotide exchange factor (VAV1), mRNA [NM_005428] | $\begin{gathered} \text { chr19:6853995- } \\ 6854054 \end{gathered}$ | + |
| 13 | A_23_P432352 | CXorf61 | Homo sapiens chromosome $X$ open reading frame 61 (CXorf61), mRNA [NM_001017978] | $\begin{gathered} \text { chrX:115593025- } \\ 115592966 \end{gathered}$ | - |

Table S1C. List of LTR7-related probes from the Agilent human G3 microarray (design ID 28004)

| Probename | Genesymbl_in_Array | Genesymbl_in_refseq | chromosome | Probe_start | Probe_end |
| :---: | :---: | :---: | :---: | :---: | :---: |
| A_33_P3313258 | ATAD3B | ATAD3B | chr1 | 1417934 | 1417993 |
| A_33_P3396553 | LOC732419 | ATAD3B | chr1 | 1423229 | 1423288 |
| A_33_P3331588 | ATAD3B | ATAD3B | chr1 | 1431507 | 1431566 |
| A_33_P3385477 | ATAD3B | ATAD3B | chr1 | 1431522 | 1431581 |
| A_23_P103942 | DNAJC11 | DNAJC11 | chr1 | 6694205 | 6694234 |
| A_33_P3271387 | THAP3 | DNAJC11 | chr1 | 6695557 | 6695616 |
| A_33_P3241108 | DNAJC11 | DNAJC11 | chr1 | 6712909 | 6712968 |
| A_23_P126623 | PGD | PGD | chr1 | 10478911 | 10478970 |
| A_33_P3307960 | AADACL3 | AADACL3 | chr1 | 12785904 | 12785963 |
| A_33_P3307965 | AADACL3 | AADACL3 | chr1 | 12788667 | 12788726 |
| A_23_P34597 | CDA | CDA | chr1 | 20945069 | 20945128 |
| A_24_P353619 | ALPL | ALPL | chr1 | 21903084 | 21903884 |
| A_23_P104146 | ZMYM4 | ZMYM4 | chr1 | 35887010 | 35887069 |
| A_23_P85640 | INPP5B | INPP5B | chr1 | 38327872 | 38327931 |
| A_33_P3216694 | HIVEP3 | HIVEP3 | chr1 | 41975685 | 41975744 |
| A_23_P383118 | ZSWIM5 | ZSWIM5 | chr1 | 45482274 | 45482333 |
| A_23_P51660 | MUTYH | MUTYH | chr1 | 45794999 | 45795058 |
| A_23_P126057 | SCP2 | SCP2 | chr1 | 53516467 | 53516526 |
| A_24_P491397 | LDLRAD1 | LDLRAD1 | chr1 | 54474512 | 54474571 |
| A_33_P3332406 | LDLRAD1 | LDLRAD1 | chr1 | 54474691 | 54474750 |
| A_23_P23850 | DAB1 | DAB1 | chr1 | 57480639 | 57480698 |
| A_33_P3375334 | NOGENE | DAB1 | chr1 | 57536473 | 57536532 |
| A_32_P108655 | AK3L1 | AK4 | chr1 | 65692632 | 65692691 |
| A_32_P95067 | AK3L1 | AK4 | chr1 | 65694099 | 65694158 |
| A_23_P33093 | ST6GALNAC5 | ST6GALNAC5 | chr1 | 77516413 | 77516472 |
| A_33_P3315258 | CHD1L | CHD1L | chr1 | 146736128 | 146736187 |
| A_23_P45831 | CHD1L | CHD1L | chr1 | 146766111 | 146766170 |
| A_24_P336759 | MCL1 | MCL1 | chr1 | 150547622 | 150547681 |
| A_33_P3272952 | LOC100131311 | MCL1 | chr1 | 150552070 | 150552129 |
| A_23_P74309 | NOS1AP | NOS1AP | chr1 | 162337975 | 162338034 |
| A_19_P00813176 | NOGENE | LOC100506023 | chr1 | 173208439 | 173208498 |
| A_19_P00807358 | NOGENE | LOC100506023 | chr1 | 173222254 | 173222313 |
| A_19_P00813450 | NOGENE | LOC100506023 | chr1 | 173222232 | 173222291 |
| A_33_P3391120 | LOC646870 | LOC100506023 | chr1 | 173331503 | 173331562 |
| A_19_P00328574 | NOGENE | LOC100506023 | chr1 | 173382628 | 173382687 |
| A_19_P00811661 | NOGENE | LOC100506023 | chr1 | 173387004 | 173387063 |
| A_19_P00805950 | NOGENE | LOC100506023 | chr1 | 173386997 | 173387056 |
| A_19_P00317897 | NOGENE | LOC100506023 | chr1 | 173387179 | 173387238 |
| A_32_P52119 | NOGENE | LOC100506023 | chr1 | 173387368 | 173387427 |
| A_19_P00316156 | NOGENE | LOC100506023 | chr1 | 173387476 | 173387535 |
| A_19_P00328176 | NOGENE | LOC100506023 | chr1 | 173387703 | 173387762 |
| A_33_P3350758 | RASAL2 | RASAL2 | chr1 | 178442587 | 178442646 |
| A_23_P502747 | RASAL2 | RASAL2 | chr1 | 178443038 | 178443097 |


| A_32_P176594 | KIAA1614 | KIAA1614 | chr1 | 180913533 | 180913592 |
| :---: | :---: | :---: | :---: | :---: | :---: |
| A_23_P148990 | HMCN1 | HMCN1 | chr1 | 186159679 | 186159738 |
| A_23_P11685 | PLA2G4A | PLA2G4A | chr1 | 186957652 | 186957711 |
| A_33_P3311373 | LOC401980 | LOC401980 | chr1 | 202955580 | 202955639 |
| A_23_P149664 | TMEM183B | TMEM183B\|TMEM183A | chr1 | 202992196 | 202992255 |
| A_23_P256821 | CR1 | CR1 | chr1 | 207813016 | 207813075 |
| A_33_P3338634 | SYT14 | SYT14 | chr1 | 210334193 | 210334252 |
| A_24_P402415 | SYT14 | SYT14 | chr1 | 210335095 | 210335154 |
| A_33_P3221683 | C1orf227 | C1orf227 | chr1 | 213003515 | 213003574 |
| A_33_P3321919 | C1orf227 | C1orf227 | chr1 | 213009418 | 213009477 |
| A_33_P3212949 | USH2A | USH2A | chr1 | 215796333 | 215796392 |
| A_19_P00811854 | NOGENE | EPHX1 | chr1 | 226006283 | 226006342 |
| A_19_P00803678 | NOGENE | EPHX1 | chr1 | 226006301 | 226006360 |
| A_23_P34537 | EPHX1 | EPHX1 | chr1 | 226032883 | 226032942 |
| A_24_P62530 | RHOU | RHOU | chr1 | 228882099 | 228882158 |
| A_23_P62967 | DISC1 | TSNAX-DISC1\|DISC1 | chr1 | 232176696 | 232176755 |
| A_33_P3251727 | RYR2 | RYR2 | chr1 | 237996036 | 237996095 |
| A_23_P137797 | RYR2 | RYR2 | chr1 | 237996423 | 237996482 |
| A_23_P510 | PLD5 | PLD5 | chr1 | 242252899 | 242252958 |
| A_32_P8925 | C1orf100 | C1orf100 | chr1 | 244541935 | 244552305 |
| A_32_P103291 | SMYD3 | SMYD3 | chr1 | 245912797 | 245912856 |
| A_23_P51410 | SMYD3 | SMYD3 | chr1 | 246490544 | 246490603 |
| A_19_P00326008 | NOGENE | ZNF670-ZNF695 | chr1 | 247126279 | 247126338 |
| A_19_P00325181 | NOGENE | ZNF670-ZNF695 | chr1 | 247138953 | 247139012 |
| A_19_P00803131 | NOGENE | ZNF670-ZNF695 | chr1 | 247142823 | 247142882 |
| A_23_P35316 | ZNF695 | ZNF670-ZNF695 | chr1 | 247150712 | 247150771 |
| A_24_P254705 | ZNF695 | ZNF670-ZNF695 | chr1 | 247162725 | 247163255 |
| A_23_P74981 | ZNF670 | ZNF670-ZNF695\|ZNF670 | chr1 | 247200849 | 247200908 |
| A_23_P86751 | ADARB2 | ADARB2 | chr10 | 1228144 | 1228203 |
| A_33_P3282307 | ADARB2 | ADARB2 | chr10 | 1229183 | 1229242 |
| A_33_P3282305 | ADARB2 | ADARB2 | chr10 | 1246262 | 1246321 |
| A_33_P3244151 | NOGENE | ADARB2 | chr10 | 1259513 | 1259572 |
| A_33_P3416398 | LOC100129894 | ADARB2 | chr10 | 1334679 | 1334738 |
| A_33_P3641456 | NOGENE | ADARB2 | chr10 | 1379975 | 1380034 |
| A_24_P206604 | PFKFB3 | PFKFB3 | chr10 | 6266166 | 6268203 |
| A_33_P3223663 | NOGENE | PFKFB3 | chr10 | 6274345 | 6274404 |
| A_24_P261259 | PFKFB3 | PFKFB3 | chr10 | 6277062 | 6277121 |
| A_33_P3219939 | CUBN | CUBN | chr10 | 16865965 | 16866024 |
| A_33_P3366758 | ST8SIA6 | ST8SIA6 | chr10 | 17362676 | 17362735 |
| A_33_P3403963 | ST8SIA6 | ST8SIA6 | chr10 | 17363190 | 17363249 |
| A_23_P161352 | PTPLA | PTPLA | chr10 | 17645575 | 17645634 |
| A_33_P3235410 | PTPLA | PTPLA | chr10 | 17657511 | 17657570 |
| A_23_P61580 | NSUN6 | NSUN6 | chr10 | 18834724 | 18834783 |
| A_23_P161424 | PLXDC2 | PLXDC2 | chr10 | 20568719 | 20568778 |
| A_23_P300905 | CCNY | CCNY | chr10 | 35858729 | 35858788 |
| A_23_P338495 | 8-Mar | MARCH8 | chr10 | 45953754 | 45953813 |


| A_32_P163125 | SGMS1 | SGMS1 | chr10 | 52065560 | 52065619 |
| :---: | :---: | :---: | :---: | :---: | :---: |
| A_33_P3351087 | NOGENE | PRKG1 | chr10 | 53822566 | 53822622 |
| A_24_P250765 | PRKG1 | PRKG1 | chr10 | 54048737 | 54050026 |
| A_33_P3258244 | PCDH15 | PCDH15 | chr10 | 55562664 | 55562723 |
| A_33_P3258239 | PCDH15 | PCDH15 | chr10 | 55568452 | 55568511 |
| A_23_P161331 | PCDH15 | PCDH15 | chr10 | 55581652 | 55581711 |
| A_24_P111147 | PCDH15 | PCDH15 | chr10 | 55754659 | 55754718 |
| A_33_P3229412 | NRG3 | NRG3 | chr10 | 84745302 | 84745361 |
| A_33_P3229417 | NRG3 | NRG3 | chr10 | 84746874 | 84746933 |
| A_24_P527404 | BMPR1A | BMPR1A | chr10 | 88683628 | 88683687 |
| A_23_P1431 | BMPR1A | BMPR1A | chr10 | 88683898 | 88683957 |
| A_33_P3219256 | BMPR1A | BMPR1A | chr10 | 88684862 | 88684921 |
| A_33_P3283611 | IFIT3 | IFIT3 | chr10 | 91099762 | 91099821 |
| A_23_P500381 | HTR7 | HTR7 | chr10 | 92500784 | 92500843 |
| A_33_P3222788 | LOC100188947 | LOC100188947 | chr10 | 93067061 | 93067120 |
| A_23_P409489 | DNTT | DNTT | chr10 | 98098081 | 98098140 |
| A_33_P3220570 | UBTD1 | UBTD1 | chr10 | 99330900 | 99330959 |
| A_33_P3293524 | NEURL | NEURL | chr10 | 105352243 | 105352302 |
| A_23_P202034 | GUCY2GP | GUCY2GP | chr10 | 114074029 | 114074088 |
| A_24_P882914 | C10orf46 | C10orf46 | chr10 | 120441482 | 120441541 |
| A_33_P3377619 | C10orf46 | C10orf46 | chr10 | 120454654 | 120454713 |
| A_23_P86599 | DMBT1 | DMBT1 | chr10 | 124403152 | 124403211 |
| A_23_P364478 | FAM175B | FAM175B | chr10 | 126524866 | 126524925 |
| A_23_P87363 | ART1 | ART1 | chr11 | 3685463 | 3685523 |
| A_23_P124190 | TRIM34 | TRIM6-TRIM34\|TRIM34 | chr11 | 5655079 | 5655869 |
| A_24_P398323 | TRIM34 | TRIM34\|TRIM6-TRIM34 | chr11 | 5664680 | 5664739 |
| A_23_P139418 | GALNTL4 | GALNTL4 | chr11 | 11292608 | 11292667 |
| A_23_P150286 | PSMA1 | PSMA1 | chr11 | 14526569 | 14526628 |
| A_33_P3288384 | PSMA1 | PSMA1 | chr11 | 14535512 | 14535571 |
| A_24_P348806 | PLEKHA7 | PLEKHA7 | chr11 | 16809231 | 16809290 |
| A_24_P649282 | LUZP2 | LUZP2 | chr11 | 25104007 | 25104066 |
| A_32_P7316 | BDNF | BDNF-AS1 | chr11 | 27677013 | 27677072 |
| A_23_P127891 | BDNF | BDNF-AS1 | chr11 | 27679900 | 27679959 |
| A_33_P3323842 | BDNFOS | BDNF-AS1 | chr11 | 27699226 | 27699285 |
| A_24_P386622 | ARRB1 | ARRB1 | chr11 | 74977300 | 74978736 |
| A_23_P162165 | KCTD14 | NDUFC2-KCTD14 | chr11 | 77727492 | 77727551 |
| A_23_P363954 | THRSP | NDUFC2-KCTD14 | chr11 | 77775305 | 77775364 |
| A_24_P364236 | NDUFC2 | NDUFC2-KCTD14\|NDUFC2 | chr11 | 77779435 | 77779494 |
| A_33_P3336652 | NDUFC2 | NDUFC2-KCTD14\|NDUFC2 | chr11 | 77790643 | 77790702 |
| A_23_P47148 | NOX4 | NOX4 | chr11 | 89059826 | 89059885 |
| A_24_P169092 | MAML2 | MAML2 | chr11 | 95712375 | 95712434 |
| A_32_P41026 | SC5DL | SC5DL | chr11 | 121183280 | 121183339 |
| A_32_P158966 | KLRF1 | KLRF1 | chr12 | 9997324 | 9997383 |
| A_32_P720220 | C12orf36 | C12orf36 | chr12 | 13524582 | 13524641 |
| A_33_P3364089 | SLCO1B3 | SLCO1B3 | chr12 | 21036476 | 21036535 |
| A_24_P935986 | BCAT1 | BCAT1 | chr12 | 24964452 | 24964511 |


| A_24_P52921 | BCAT1 | BCAT1 | chr12 | 24989496 | 24995040 |
| :---: | :---: | :---: | :---: | :---: | :---: |
| A_23_P95231 | CASC1 | CASC1 | chr12 | 25261534 | 25261593 |
| A_33_P3378790 | CASC1 | CASC1 | chr12 | 25263063 | 25263122 |
| A_23_P150903 | FAR2 | FAR2 | chr12 | 29485623 | 29486586 |
| A_23_P98930 | C12orf35 | C12orf35 | chr12 | 32145764 | 32145823 |
| A_33_P3409090 | CNTN1 | CNTN1 | chr12 | 41352972 | 41353031 |
| A_23_P390700 | CNTN1 | CNTN1 | chr12 | 41414156 | 41414215 |
| A_24_P98914 | PFKM | PFKM | chr12 | 48539724 | 48539783 |
| A_23_P391443 | PPM1H | PPM1H | chr12 | 63038521 | 63038580 |
| A_23_P162300 | IRAK3 | IRAK3 | chr12 | 66642160 | 66642219 |
| A_23_P113382 | GRIP1 | GRIP1 | chr12 | 66742133 | 66742192 |
| A_33_P3252083 | GRIP1 | GRIP1 | chr12 | 66742800 | 66742859 |
| A_24_P129834 | TPH2 | TPH2 | chr12 | 72425334 | 72425393 |
| A_32_P326819 | KRR1 | GLIPR1 | chr12 | 75892026 | 75892085 |
| A_33_P3418125 | GLIPR1 | GLIPR1 | chr12 | 75895656 | 75895715 |
| A_33_P3258003 | ANKS1B | ANKS1B | chr12 | 99129338 | 99129397 |
| A_23_P356717 | ANKS1B | ANKS1B | chr12 | 99138178 | 99138237 |
| A_33_P3258004 | ANKS1B | ANKS1B | chr12 | 99145177 | 99145236 |
| A_32_P13113 | FAM71C | ANKS1B | chr12 | 100043693 | 100043752 |
| A_32_P189204 | GAS2L3 | GAS2L3 | chr12 | 101018594 | 101018653 |
| A_33_P3328289 | LOC100130902 | TXNRD1 | chr12 | 104680606 | 104680665 |
| A_33_P3328284 | NOGENE | TXNRD1 | chr12 | 104680832 | 104680891 |
| A_23_P65068 | EID3 | TXNRD1 | chr12 | 104698595 | 104698654 |
| A_33_P3351120 | TXNRD1 | TXNRD1 | chr12 | 104732949 | 104733008 |
| A_23_P348257 | NUAK1 | NUAK1 | chr12 | 106457546 | 106457605 |
| A_33_P3371889 | NUAK1 | NUAK1 | chr12 | 106461077 | 106461136 |
| A_23_P410312 | C12orf76 | C12orf76 | chr12 | 110479089 | 110479148 |
| A_23_P44643 | ANAPC7 | ANAPC7 | chr12 | 110813991 | 110815226 |
| A_23_P401361 | PITPNM2 | PITPNM2 | chr12 | 123468975 | 123469034 |
| A_33_P3295154 | NOGENE | PITPNM2 | chr12 | 123576390 | 123576449 |
| A_19_P00810888 | NOGENE | ZNF664-FAM101A | chr12 | 124573023 | 124573082 |
| A_33_P3342862 | FAM101A | ZNF664-FAM101A | chr12 | 124799228 | 124799287 |
| A_24_P288890 | FAM101A | ZNF664-FAM101A | chr12 | 124799557 | 124799616 |
| A_23_P308839 | TMEM132D | TMEM132D | chr12 | 129557350 | 129557409 |
| A_33_P3531979 | NOGENE | TMEM132D | chr12 | 129596347 | 129596406 |
| A_23_P344037 | CHFR | CHFR | chr12 | 133417712 | 133417771 |
| A_33_P3327956 | ZNF605 | ZNF605 | chr12 | 133498048 | 133498107 |
| A_33_P3352877 | SPG20 | SPG20 | chr13 | 36875853 | 36875912 |
| A_33_P3361741 | DNAJC15 | DNAJC15 | chr13 | 43683245 | 43683304 |
| A_24_P170774 | LRCH1 | LRCH1 | chr13 | 47317618 | 47317677 |
| A_33_P3333337 | LRCH1 | LRCH1 | chr13 | 47324726 | 47324785 |
| A_23_P117157 | SUCLA2 | SUCLA2 | chr13 | 48517388 | 48517447 |
| A_24_P330263 | EDNRB | EDNRB | chr13 | 78470568 | 78470627 |
| A_23_P204980 | UGGT2 | UGGT2 | chr13 | 96453925 | 96453984 |
| A_23_P205031 | COL4A2 | COL4A2 | chr13 | 111165261 | 111165320 |
| A_24_P161973 | ATP11A | ATP11A | chr13 | 113540918 | 113540977 |


| A_24_P144332 | NOGENE | C14orf167 | chr14 | 24408018 | 24408077 |
| :---: | :---: | :---: | :---: | :---: | :---: |
| A_32_P118250 | C14orf167 | C14orf167 | chr14 | 24408338 | 24408397 |
| A_23_P14351 | AKAP6 | AKAP6 | chr14 | 33301563 | 33301622 |
| A_33_P3408420 | MDGA2 | MDGA2 | chr14 | 47311008 | 47311067 |
| A_23_P366035 | ABHD12B | ABHD12B | chr14 | 51371212 | 51371271 |
| A_23_P205623 | DDHD1 | DDHD1 | chr14 | 53521264 | 53521323 |
| A_23_P140373 | FLVCR2 | FLVCR2 | chr14 | 76113756 | 76113815 |
| A_24_P150486 | SPTLC2 | SPTLC2 | chr14 | 77973992 | 77974051 |
| A_24_P42557 | TSHR | TSHR | chr14 | 81557464 | 81558910 |
| A_23_P88435 | FOXN3 | FOXN3 | chr14 | 89623539 | 89623598 |
| A_33_P3363710 | LOC100128075 | FOXN3 | chr14 | 89734789 | 89734848 |
| A_24_P306720 | LOC400236 | FOXN3 | chr14 | 89885621 | 89885680 |
| A_23_P48771 | C14orf159 | C14orf159 | chr14 | 91691358 | 91691417 |
| A_23_P106241 | TRIP11 | TRIP11 | chr14 | 92435892 | 92435951 |
| A_33_P3382271 | NOGENE | TRIP11 | chr14 | 92506805 | 92506864 |
| A_23_P432272 | KIAA1409 | KIAA1409 | chr14 | 94173274 | 94173333 |
| A_23_P65651 | WARS | WARS | chr14 | 100801041 | 100801100 |
| A_23_P76731 | RAGE | RAGE | chr14 | 102695289 | 102695348 |
| A_33_P3326989 | RAGE | RAGE | chr14 | 102698958 | 102699017 |
| A_33_P3326984 | RAGE | RAGE | chr14 | 102700128 | 102700187 |
| A_33_P3305958 | TECPR2 | TECPR2 | chr14 | 102900827 | 102900886 |
| A_32_P355396 | TECPR2 | TECPR2 | chr14 | 102968719 | 102968778 |
| A_33_P3277075 | GABRB3 | GABRB3 | chr15 | 26788857 | 26788916 |
| A_33_P3309206 | GABRB3 | GABRB3 | chr15 | 26869979 | 26870038 |
| A_23_P129133 | OCA2 | OCA2 | chr15 | 28000067 | 28000126 |
| A_32_P143000 | FAM189A1 | FAM189A1 | chr15 | 29412696 | 29412755 |
| A_24_P124973 | NDNL2 | FAM189A1 | chr15 | 29561202 | 29561261 |
| A_33_P3800734 | RYR3 | RYR3 | chr15 | 34158044 | 34158103 |
| A_19_P00328893 | NOGENE | LOC729082 | chr15 | 41584871 | 41584930 |
| A_24_P655268 | LOC729082 | LOC729082 | chr15 | 41591663 | 41591722 |
| A_33_P3215277 | TTBK2 | TTBK2 | chr15 | 43036550 | 43036609 |
| A_24_P652700 | CEP152 | CEP152 | chr15 | 49030699 | 49030758 |
| A_24_P333663 | MAPK6 | MAPK6 | chr15 | 52356907 | 52356966 |
| A_23_P3204 | MAPK6 | MAPK6 | chr15 | 52358055 | 52358114 |
| A_23_P346006 | CCPG1 | DYX1C1-CCPG1 | chr15 | 55648436 | 55648495 |
| A_32_P447001 | NOGENE | FLJ27352 | chr15 | 55710662 | 55710721 |
| A_23_P88559 | LIPC | LIPC | chr15 | 58861011 | 58861070 |
| A_23_P140475 | NOX5 | MIR548H4\|NOX5 | chr15 | 69348985 | 69349044 |
| A_19_P00321618 | NOGENE | MIR548H4 | chr15 | 69373246 | 69373305 |
| A_19_P00322896 | NOGENE | MIR548H4 | chr15 | 69373297 | 69373356 |
| A_19_P00319825 | NOGENE | MIR548H4 | chr15 | 69373355 | 69373414 |
| A_19_P00319776 | NOGENE | MIR548H4 | chr15 | 69373389 | 69373448 |
| A_19_P00322671 | NOGENE | MIR548H4 | chr15 | 69373449 | 69373508 |
| A_33_P3214229 | TMEM84 | MIR548H4 | chr15 | 69373563 | 69373622 |
| A_19_P00322669 | NOGENE | MIR548H4 | chr15 | 69383452 | 69383511 |
| A_19_P00321910 | NOGENE | MIR548H4 | chr15 | 69383675 | 69383734 |


| A_19_P00322662 | NOGENE | MIR548H4 | chr15 | 69387049 | 69387108 |
| :---: | :---: | :---: | :---: | :---: | :---: |
| A_23_P322632 | TMEM84 | MIR548H4 | chr15 | 69387230 | 69387289 |
| A_19_P00322879 | NOGENE | MIR548H4 | chr15 | 69392109 | 69392168 |
| A_23_P129169 | CYP11A1 | CYP11A1 | chr15 | 74631018 | 74631077 |
| A_23_P77304 | AP3B2 | AP3B2 | chr15 | 83328449 | 83328642 |
| A_33_P3572454 | LOC283692 | AP3B2 | chr15 | 83361512 | 83361571 |
| A_33_P3378081 | AGBL1 | AGBL1 | chr15 | 87531258 | 87531317 |
| A_23_P26184 | DET1 | DET1 | chr15 | 89055798 | 89055857 |
| A_33_P3229390 | NOGENE | DET1 | chr15 | 89061422 | 89061481 |
| A_24_P287691 | AP3S2 | C15orf38-AP3S2\|AP3S2 | chr15 | 90377719 | 90377778 |
| A_24_P943017 | LASS3 | LASS3 | chr15 | 100940872 | 100940931 |
| A_33_P3408305 | LASS3 | LASS3 | chr15 | 100942746 | 100942805 |
| A_24_P48162 | MPG | MPG | chr16 | 133183 | 133242 |
| A_23_P100326 | C16orf35 | MPG | chr16 | 135839 | 135898 |
| A_33_P3260209 | LOC100132944 | PMM2 | chr16 | 8941517 | 8941576 |
| A_23_P432360 | PMM2 | PMM2 | chr16 | 8942790 | 8942849 |
| A_23_P140928 | TMC7 | TMC7 | chr16 | 19074010 | 19074069 |
| A_23_P420281 | PRKCB | PRKCB | chr16 | 24226173 | 24226233 |
| A_33_P3294533 | PRKCB | PRKCB | chr16 | 24231495 | 24231554 |
| A_33_P3255314 | FLJ26245 | FLJ26245 | chr16 | 34989846 | 34989905 |
| A_23_P334123 | ITFG1 | ITFG1 | chr16 | 47192831 | 47192890 |
| A_33_P3316115 | LOC100127930 | ITFG1 | chr16 | 47196599 | 47196658 |
| A_24_P1773 | LONP2 | LONP2 | chr16 | 48386989 | 48387048 |
| A_33_P3367293 | MT1IP | MT1IP | chr16 | 56711498 | 56711557 |
| A_23_P140797 | CDH8 | CDH8 | chr16 | 61687697 | 61687756 |
| A_23_P100386 | IL34 | IL34 | chr16 | 70688562 | 70690551 |
| A_23_P418234 | PHLPP2 | PHLPP2 | chr16 | 71679160 | 71679219 |
| A_33_P3271810 | PHLPP2 | PHLPP2 | chr16 | 71686698 | 71686757 |
| A_23_P71972 | WWOX | WWOX | chr16 | 78458922 | 78466413 |
| A_23_P117992 | ATP2C2 | ATP2C2 | chr16 | 84497443 | 84497502 |
| A_33_P3253427 | LRRC37B2 | LRRC37BP1 | chr17 | 28935520 | 28935579 |
| A_33_P3564394 | SH3GL1P2 | LRRC37BP1 | chr17 | 28951611 | 28951670 |
| A_33_P3564399 | SH3GL1P2 | LRRC37BP1 | chr17 | 28952259 | 28952318 |
| A_32_P28402 | LRRC37B2 | LRRC37BP1 | chr17 | 28964201 | 28964260 |
| A_23_P389102 | MYO1D | MYO1D | chr17 | 30819832 | 30819891 |
| A_33_P3399566 | LOC100130931 | MYO1D | chr17 | 30903022 | 30903081 |
| A_24_P119609 | MYO1D | MYO1D | chr17 | 31092008 | 31092067 |
| A_33_P3221999 | GSDMB | GSDMB | chr17 | 38060850 | 38060909 |
| A_23_P73150 | TTC25 | TTC25 | chr17 | 40117524 | 40117583 |
| A_33_P3289113 | COX11 | TOM1L1 | chr17 | 53029465 | 53029524 |
| A_24_P73943 | COX11 | TOM1L1 | chr17 | 53038703 | 53038762 |
| A_23_P118493 | TOM1L1 | TOM1L1 | chr17 | 53038843 | 53038902 |
| A_23_P375566 | STXBP4 | STXBP4 | chr17 | 53111534 | 53111593 |
| A_32_P90483 | STXBP4 | STXBP4 | chr17 | 53240764 | 53240823 |
| A_33_P3414362 | USP32 | USP32 | chr17 | 58329743 | 58329802 |
| A_24_P410100 | NOGENE | USP32 | chr17 | 58356431 | 58356490 |


| A_19_P00811559 | NOGENE | LOC146880 | chr17 | 62758594 | 62758653 |
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| A_23_P66881 | RGS9 | RGS9 | chr17 | 63223600 | 63223659 |
| A_23_P84189 | PITPNC1 | PITPNC1 | chr17 | 65689234 | 65689293 |
| A_24_P255005 | NOGENE | LOC100499466 | chr17 | 66122851 | 66122910 |
| A_33_P3209404 | SH3GL1P3 | LOC100499466 | chr17 | 66130414 | 66130473 |
| A_33_P3314594 | RAB37 | RAB37 | chr17 | 72742734 | 72742793 |
| A_23_P414654 | RAB37 | RAB37 | chr17 | 72743222 | 72743281 |
| A_23_P38181 | GGA3 | GGA3 | chr17 | 73232920 | 73232979 |
| A_23_P26759 | CANT1 | CANT1 | chr17 | 76988694 | 76988753 |
| A_24_P116669 | CANT1 | CANT1 | chr17 | 76993163 | 76993222 |
| A_23_P327140 | RNF213 | LOC100294362 | chr17 | 78369857 | 78369916 |
| A_23_P66948 | FAM59A | FAM59A | chr18 | 29847581 | 29847640 |
| A_23_P170050 | RIT2 | RIT2 | chr18 | 40323249 | 40323308 |
| A_33_P3416588 | RIT2 | RIT2 | chr18 | 40503555 | 40503614 |
| A_24_P322354 | SKA1 | SKA1 | chr18 | 47919899 | 47919958 |
| A_33_P3401407 | hCG_33730 | LOC390858 | chr18 | 56718840 | 56718899 |
| A_33_P3706494 | LOC284294 | LOC284294 | chr18 | 62090743 | 62090802 |
| A_33_P3531970 | LOC643542 | LOC643542 | chr18 | 65566781 | 65566840 |
| A_23_P101208 | CYB5A | CYB5A | chr18 | 71920718 | 71920777 |
| A_33_P3311076 | CYB5A | CYB5A | chr18 | 71930598 | 71930657 |
| A_23_P131074 | THEG | THEG | chr19 | 362125 | 362184 |
| A_23_P39263 | ZNF57 | ZNF57 | chr19 | 2917775 | 2917834 |
| A_23_P360316 | FUT3 | FUT3 | chr19 | 5843092 | 5843151 |
| A_23_P101351 | ZNF426 | ZNF426 | chr19 | 9638979 | 9639038 |
| A_24_P935782 | ZNF121 | ZNF121 | chr19 | 9677285 | 9677344 |
| A_33_P3410925 | KLF1 | KLF1 | chr19 | 12995236 | 12995295 |
| A_32_P200238 | UCA1 | UCA1 | chr19 | 15946035 | 15946092 |
| A_23_P90419 | PBX4 | PBX4 | chr19 | 19672538 | 19672597 |
| A_32_P790361 | ZNF90 | ZNF90 | chr19 | 20228630 | 20228689 |
| A_33_P3474538 | ZNF90 | ZNF90 | chr19 | 20231608 | 20231667 |
| A_33_P3244574 | LOC100128675 | LOC100128675 | chr19 | 35550039 | 35550098 |
| A_23_P406782 | HPN | LOC100128675 | chr19 | 35556490 | 35556549 |
| A_23_P90444 | RBM42 | RBM42 | chr19 | 36128152 | 36128211 |
| A_33_P3403773 | ZNF569 | ZNF569 | chr19 | 37902079 | 37902138 |
| A_33_P3231602 | ZNF569 | ZNF569 | chr19 | 37902563 | 37902622 |
| A_23_P107994 | TMEM160 | TMEM160 | chr19 | 47549317 | 47549376 |
| A_23_P368779 | ZNF114 | ZNF114 | chr19 | 48790154 | 48790213 |
| A_33_P3327961 | ZNF615 | ZNF615 | chr19 | 52494748 | 52494807 |
| A_33_P3400424 | ZNF615 | ZNF615 | chr19 | 52496133 | 52496192 |
| A_33_P3345132 | ZNF578 | ZNF578 | chr19 | 53019319 | 53019378 |
| A_33_P3256868 | ZNF83 | ZNF83 | chr19 | 53116271 | 53116330 |
| A_23_P4962 | NLRP5 | NLRP5 | chr19 | 56569768 | 56572812 |
| A_23_P17287 | IAH1 | IAH1 | chr2 | 9628439 | 9628498 |
| A_23_P40049 | CAD | CAD | chr2 | 27465770 | 27465829 |
| A_23_P67847 | GALNT14 | GALNT14 | chr2 | 31133370 | 31133429 |
| A_23_P39718 | FEZ2 | FEZ2 | chr2 | 36779704 | 36779763 |


| A_33_P3311956 | FEZ2 | FEZ2 | chr2 | 36782827 | 36782886 |
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| A_33_P3401267 | FEZ2 | FEZ2 | chr2 | 36787931 | 36787990 |
| A_33_P3272352 | NOGENE | CEBPZ | chr2 | 37429079 | 37429138 |
| A_23_P119964 | CEBPZ | CEBPZ | chr2 | 37429942 | 37430001 |
| A_33_P3223648 | NOGENE | CEBPZ | chr2 | 37430462 | 37430521 |
| A_33_P3241489 | NOGENE | CEBPZ | chr2 | 37431823 | 37431882 |
| A_23_P108932 | RPL23AP32 | SPTBN1 | chr2 | 54756368 | 54756427 |
| A_33_P3258467 | SPTBN1 | SPTBN1 | chr2 | 54889385 | 54889444 |
| A_33_P3258472 | SPTBN1 | SPTBN1 | chr2 | 54898523 | 54898582 |
| A_23_P90565 | C2orf86 | WDPCP | chr2 | 63486522 | 63540419 |
| A_23_P253046 | UGP2 | UGP2 | chr2 | 64118483 | 64118542 |
| A_23_P56654 | MCEE | MCEE | chr2 | 71337089 | 71337148 |
| A_23_P5586 | MPHOSPH10 | MPHOSPH10 | chr2 | 71375141 | 71375200 |
| A_24_P148590 | TACR1 | TACR1 | chr2 | 75280843 | 75347719 |
| A_24_P666105 | LRRTM4 | LRRTM4 | chr2 | 76974947 | 76975006 |
| A_33_P3235204 | ELMOD3 | ELMOD3 | chr2 | 85617562 | 85617621 |
| A_33_P3297305 | ELMOD3 | ELMOD3 | chr2 | 85618030 | 85618089 |
| A_23_P154256 | ELMOD3 | ELMOD3 | chr2 | 85618691 | 85618750 |
| A_24_P945293 | VPS24 | VPS24\|RNF103-VPS24 | chr2 | 86730551 | 86730587 |
| A_24_P240065 | VPS24 | VPS24\|RNF103-VPS24 | chr2 | 86734691 | 86737514 |
| A_33_P3312489 | VWA3B | VWA3B | chr2 | 98908365 | 98908424 |
| A_32_P46603 | VWA3B | VWA3B | chr2 | 98920173 | 98920232 |
| A_23_P373464 | AFF3 | AFF3 | chr2 | 100163700 | 100163726 |
| A_23_P108761 | C2orf29 | C2orf29 | chr2 | 101885805 | 101885864 |
| A_32_P761797 | NCRNA00116 | NCRNA00116 | chr2 | 110970211 | 110970270 |
| A_33_P3402086 | NOGENE | MERTK | chr2 | 112766079 | 112766138 |
| A_33_P3402091 | MERTK | MERTK | chr2 | 112786885 | 112786944 |
| A_32_P86578 | LOC389023 | DPP10 | chr2 | 115901752 | 115901811 |
| A_33_P3211356 | DPP10 | DPP10 | chr2 | 116600660 | 116600719 |
| A_24_P140057 | DPP10 | DPP10 | chr2 | 116601306 | 116601365 |
| A_23_P373109 | CNTNAP5 | CNTNAP5 | chr2 | 125521661 | 125521720 |
| A_23_P56787 | CNTNAP5 | CNTNAP5 | chr2 | 125672642 | 125672701 |
| A_32_P328023 | WDR33 | WDR33 | chr2 | 128462034 | 128462093 |
| A_33_P3335840 | WDR33 | WDR33 | chr2 | 128492971 | 128493030 |
| A_24_P102920 | WDR33 | WDR33 | chr2 | 128521179 | 128521238 |
| A_33_P3335845 | WDR33 | WDR33 | chr2 | 128522116 | 128522175 |
| A_33_P3389779 | NOGENE | WDR33 | chr2 | 128568324 | 128568383 |
| A_23_P5342 | LRP1B | LRP1B | chr2 | 140989375 | 140989434 |
| A_33_P3244728 | LRP2 | LRP2 | chr2 | 169983769 | 169983828 |
| A_32_P116857 | PDE11A | PDE11A | chr2 | 178489045 | 178489104 |
| A_24_P43144 | PDE11A | PDE11A | chr2 | 178545554 | 178545613 |
| A_33_P3258772 | NOGENE | PDE11A | chr2 | 178587952 | 178588011 |
| A_33_P3348884 | CCDC141 | CCDC141 | chr2 | 179698900 | 179698959 |
| A_33_P3279004 | CCDC141 | CCDC141 | chr2 | 179700653 | 179700712 |
| A_33_P3495120 | LOC285026 | CCDC141 | chr2 | 179737753 | 179737812 |
| A_19_P00321823 | NOGENE | CCDC141 | chr2 | 179783570 | 179783629 |


| A_19_P00317490 | NOGENE | CCDC141 | chr2 | 179791240 | 179791299 |
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| A_19_P00317491 | NOGENE | CCDC141 | chr2 | 179791323 | 179791382 |
| A_19_P00325338 | NOGENE | CCDC141 | chr2 | 179815213 | 179815272 |
| A_19_P00328994 | NOGENE | CCDC141 | chr2 | 179828698 | 179828757 |
| A_19_P00812256 | NOGENE | CCDC141 | chr2 | 179866870 | 179866929 |
| A_19_P00811927 | NOGENE | CCDC141 | chr2 | 179866866 | 179866925 |
| A_19_P00316387 | NOGENE | CCDC141 | chr2 | 179914334 | 179914393 |
| A_32_P316136 | ZNF804A | ZNF804A | chr2 | 185803216 | 185803275 |
| A_23_P361049 | MYO1B | MYO1B | chr2 | 192289808 | 192289867 |
| A_23_P363878 | RFTN2 | RFTN2 | chr2 | 198436064 | 198436123 |
| A_23_P16817 | CLK1 | CLK1 | chr2 | 201718707 | 201719388 |
| A_33_P3211513 | CLK1 | CLK1 | chr2 | 201725962 | 201726021 |
| A_33_P3383283 | CASP10 | CASP10 | chr2 | 202086244 | 202086303 |
| A_33_P3255075 | CASP10 | CASP10 | chr2 | 202094051 | 202094110 |
| A_32_P199462 | C2orf80 | C2orf80 | chr2 | 209030194 | 209030253 |
| A_33_P3346966 | SPAG16 | SPAG16 | chr2 | 215013930 | 215013989 |
| A_33_P3367984 | ABCA12 | ABCA12 | chr2 | 215796277 | 215796336 |
| A_33_P3315134 | DIRC3 | DIRC3 | chr2 | 218148801 | 218148860 |
| A_23_P142574 | MOGAT1 | MOGAT1 | chr2 | 223574531 | 223574590 |
| A_23_P57110 | C20orf54 | C20orf54 | chr20 | 741010 | 741069 |
| A_23_P68511 | ANGPT4 | ANGPT4 | chr20 | 853663 | 853722 |
| A_33_P3482534 | LOC613266 | MACROD2 | chr20 | 15872981 | 15873038 |
| A_33_P3281033 | MACROD2 | MACROD2 | chr20 | 15967310 | 15967369 |
| A_32_P89352 | MACROD2 | MACROD2 | chr20 | 16033184 | 16033243 |
| A_23_P6119 | SEC23B | SEC23B | chr20 | 18526643 | 18529277 |
| A_24_P690924 | SEC23B | SEC23B | chr20 | 18531782 | 18534900 |
| A_33_P3344046 | RP11-218C14.6 | CSTT | chr20 | 23514881 | 23514940 |
| A_24_P194670 | RP11-218C14.6 | CSTT | chr20 | 23514944 | 23522368 |
| A_33_P3344044 | NOGENE | CSTT | chr20 | 23517548 | 23517607 |
| A_33_P3359508 | DNMT3B | DNMT3B | chr20 | 31384813 | 31384872 |
| A_23_P28953 | DNMT3B | DNMT3B | chr20 | 31397064 | 31397123 |
| A_23_P135576 | PTPRT | PTPRT | chr20 | 40701748 | 40701807 |
| A_23_P254181 | MGC5566 | LOC79015 | chr20 | 43285419 | 43285478 |
| A_23_P120442 | NCOA3 | NCOA3 | chr20 | 46283630 | 46283689 |
| A_24_P229536 | C21orf34 | C21orf34 | chr21 | 17909716 | 17979329 |
| A_33_P3312384 | C21orf34 | C21orf34 | chr21 | 17979483 | 17979542 |
| A_33_P3508822 | APP | APP | chr21 | 27253136 | 27253195 |
| A_33_P3296479 | APP | APP | chr21 | 27423347 | 27423406 |
| A_23_P109286 | GRIK1 | GRIK1 | chr21 | 30927563 | 30927622 |
| A_33_P3303557 | NOGENE | GRIK1 | chr21 | 30968929 | 30968988 |
| A_23_P413043 | C21orf41 | GRIK1 | chr21 | 30968994 | 30969053 |
| A_33_P3279362 | NCRNA00110 | GRIK1 | chr21 | 31121174 | 31121233 |
| A_33_P3353471 | NCRNA00110 | GRIK1-AS1 | chr21 | 31136183 | 31136242 |
| A_23_P211141 | DSCAM | DSCAM | chr21 | 41384961 | 41385020 |
| A_23_P154875 | BACE2 | BACE2 | chr21 | 42647696 | 42647755 |
| A_33_P3338341 | PRODH | PRODH | chr22 | 18900438 | 18900497 |


| A_33_P3239295 | PRODH | PRODH | chr22 | 18906965 | 18907024 |
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| A_33_P3239298 | NOGENE | PRODH | chr22 | 18909759 | 18909818 |
| A_33_P3235335 | MTMR3 | MTMR3 | chr22 | 30421915 | 30421974 |
| A_33_P3235330 | MTMR3 | MTMR3 | chr22 | 30426759 | 30426818 |
| A_23_P91697 | LARGE | LARGE | chr22 | 33669332 | 33669391 |
| A_23_P353149 | C22orf33 | C22orf33 | chr22 | 37387544 | 37387603 |
| A_33_P3349702 | LOC400927 | LOC400927 | chr22 | 38740670 | 38740729 |
| A_33_P3247372 | LOC400927 | LOC400927 | chr22 | 38755951 | 38756010 |
| A_33_P3262181 | APOBEC3F | APOBEC3F | chr22 | 39440368 | 39440427 |
| A_23_P357101 | APOBEC3F | APOBEC3F | chr22 | 39448588 | 39448647 |
| A_23_P143713 | APOBEC3G | APOBEC3G | chr22 | 39477481 | 39477540 |
| A_24_P340696 | SERHL2 | SERHL2 | chr22 | 42968453 | 42968512 |
| A_23_P120953 | SERHL2 | SERHL2 | chr22 | 42970262 | 42970321 |
| A_24_P242036 | RRP7B | SERHL2 | chr22 | 42970264 | 42970323 |
| A_23_P143987 | ATG7 | ATG7 | chr3 | 11468321 | 11468380 |
| A_23_P132595 | VGLL4 | ATG7 | chr3 | 11597959 | 11598018 |
| A_24_P944827 | ATG7 | ATG7 | chr3 | 11598763 | 11598822 |
| A_23_P18282 | DLEC1 | DLEC1 | chr3 | 38164126 | 38164185 |
| A_23_P319874 | C3orf23 | C3orf23 | chr3 | 44400419 | 44400478 |
| A_23_P373054 | C3orf23 | C3orf23 | chr3 | 44450680 | 44450739 |
| A_23_P250302 | CCR3 | CCR3 | chr3 | 46307828 | 46307887 |
| A_23_P155463 | LRRC2 | LRRC2 | chr3 | 46557386 | 46557445 |
| A_23_P334798 | LRRC2 | LRRC2 | chr3 | 46563082 | 46563141 |
| A_33_P3843873 | HESRG | ESRG | chr3 | 54666167 | 54666226 |
| A_33_P3373259 | CACNA2D3 | CACNA2D3 | chr3 | 54913050 | 54913109 |
| A_23_P40856 | LRTM1 | CACNA2D3 | chr3 | 54958694 | 54958753 |
| A_24_P402825 | CACNA2D3 | CACNA2D3 | chr3 | 55021774 | 55038842 |
| A_32_P14721 | DNAH12 | DNAH12 | chr3 | 57327797 | 57327856 |
| A_33_P3216601 | FHIT | FHIT | chr3 | 59737952 | 59738011 |
| A_33_P3300941 | NPCDR1 | FHIT | chr3 | 59957230 | 59957288 |
| A_33_P3245278 | PTPRG | PTPRG | chr3 | 62280514 | 62280573 |
| A_23_P80718 | SYNPR | SYNPR | chr3 | 63602485 | 63602544 |
| A_33_P3253089 | FAM19A1 | FAM19A1 | chr3 | 68587968 | 68588027 |
| A_32_P117693 | FAM19A1 | FAM19A1 | chr3 | 68594375 | 68594434 |
| A_23_P80503 | ROBO1 | ROBO1 | chr3 | 78649391 | 78649450 |
| A_23_P121082 | GBE1 | GBE1 | chr3 | 81538912 | 81538971 |
| A_23_P73114 | PROS1 | PROS1 | chr3 | 93592149 | 93592208 |
| A_23_P18342 | EPHA6 | EPHA6 | chr3 | 97367324 | 97367383 |
| A_33_P3408782 | EPHA6 | EPHA6 | chr3 | 97467486 | 97467545 |
| A_23_P212728 | TBC1D23 | TBC1D23 | chr3 | 100043741 | 100043800 |
| A_33_P3230319 | NOGENE | BBX | chr3 | 107513790 | 107513849 |
| A_33_P3369311 | BBX | BBX | chr3 | 107524368 | 107524427 |
| A_23_P121356 | BBX | BBX | chr3 | 107524534 | 107524593 |
| A_33_P3301499 | C3orf66 | C3orf66 | chr3 | 108903960 | 108904019 |
| A_33_P3358923 | BTLA | BTLA | chr3 | 112182938 | 112182997 |
| A_23_P212706 | ATG3 | ATG3 | chr3 | 112255356 | 112255415 |


| A_23_P92281 | GTPBP8 | GTPBP8 | chr3 | 112718398 | 112719742 |
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| A_23_P398189 | IGSF11 | IGSF11 | chr3 | 118619910 | 118619969 |
| A_23_P166633 | ITGB5 | ITGB5 | chr3 | 124482491 | 124482550 |
| A_23_P159316 | BFSP2 | BFSP2 | chr3 | 133193941 | 133194001 |
| A_32_P46214 | SLC9A9 | SLC9A9 | chr3 | 142986095 | 142986154 |
| A_33_P3313025 | NOGENE | SLC9A9 | chr3 | 143346956 | 143347015 |
| A_23_P40821 | HPS3 | CP | chr3 | 148890246 | 148890304 |
| A_33_P3296587 | CP | CP | chr3 | 148890649 | 148890708 |
| A_33_P3343196 | CP | CP | chr3 | 148890802 | 148890861 |
| A_33_P3424297 | SELT | SELT | chr3 | 150344826 | 150344885 |
| A_33_P3290667 | SELT | SELT | chr3 | 150344862 | 150344921 |
| A_33_P3305851 | SELT | SELT | chr3 | 150346997 | 150347056 |
| A_33_P3290672 | SELT | SELT | chr3 | 150348174 | 150348233 |
| A_33_P3332547 | SCHIP1 | SCHIP1\|IQCJ-SCHIP1 | chr3 | 159606672 | 159606731 |
| A_32_P62863 | SCHIP1 | SCHIP1\|IQCJ-SCHIP1 | chr3 | 159615048 | 159615107 |
| A_33_P3214690 | NLGN1 | NLGN1 | chr3 | 173998992 | 173999051 |
| A_23_P18123 | NLGN1 | NLGN1 | chr3 | 174000648 | 174000707 |
| A_33_P3273854 | NAALADL2 | NAALADL2 | chr3 | 175523323 | 175523382 |
| A_24_P85537 | MAP3K13 | MAP3K13 | chr3 | 185161379 | 185165590 |
| A_23_P58031 | MAP3K13 | MAP3K13 | chr3 | 185200168 | 185200227 |
| A_23_P250156 | IGF2BP2 | IGF2BP2 | chr3 | 185362047 | 185362106 |
| A_33_P3242973 | IGF2BP2 | IGF2BP2 | chr3 | 185363403 | 185364863 |
| A_33_P3324949 | C3orf65 | IGF2BP2 | chr3 | 185435895 | 185435954 |
| A_23_P155939 | ZNF595 | ZNF718\|ZNF595 | chr4 | 86842 | 86901 |
| A_23_P124438 | ZNF718 | ZNF718 | chr4 | 156150 | 156209 |
| A_33_P3274194 | KCNIP4 | KCNIP4 | chr4 | 20730270 | 20730329 |
| A_23_P18447 | PPARGC1A | PPARGC1A | chr4 | 23793937 | 23793996 |
| A_33_P3330952 | ATP8A1 | ATP8A1 | chr4 | 42410449 | 42410508 |
| A_23_P81131 | CORIN | CORIN | chr4 | 47596314 | 47596373 |
| A_23_P110345 | CHIC2 | CHIC2 | chr4 | 54876232 | 54876291 |
| A_24_P49687 | NOGENE | LPHN3 | chr4 | 62641374 | 62641433 |
| A_33_P3216746 | LPHN3 | LPHN3 | chr4 | 62775291 | 62775350 |
| A_23_P10980 | LPHN3 | LPHN3 | chr4 | 62937245 | 62937304 |
| A_33_P3366224 | NOGENE | SHROOM3 | chr4 | 77589366 | 77589425 |
| A_33_P3417328 | SHROOM3 | SHROOM3 | chr4 | 77637503 | 77637562 |
| A_33_P3417339 | SHROOM3 | SHROOM3 | chr4 | 77678028 | 77678087 |
| A_33_P3336700 | SHROOM3 | SHROOM3 | chr4 | 77704264 | 77704323 |
| A_23_P121695 | CXCL13 | CXCL13 | chr4 | 78532646 | 78532705 |
| A_23_P110412 | TMEM150C | TMEM150C | chr4 | 83405786 | 83405845 |
| A_33_P3242231 | NOGENE | TMEM150C | chr4 | 83412519 | 83412578 |
| A_33_P3851513 | LIN54 | LIN54 | chr4 | 83846981 | 83847040 |
| A_33_P3361067 | ABCG2 | ABCG2 | chr4 | 89011503 | 89011562 |
| A_33_P3297360 | NOGENE | FAM190A | chr4 | 92240185 | 92240244 |
| A_32_P306678 | FAM190A | FAM190A | chr4 | 92522638 | 92522697 |
| A_23_P32847 | GRID2 | GRID2 | chr4 | 94693252 | 94693311 |
| A_23_P255376 | CCDC109B | CCDC109B | chr4 | 110605745 | 110606408 |


| A_23_P7212 | CFI | CFI | chr4 | 110662064 | 110662123 |
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| A_24_P92472 | CFI | CFI | chr4 | 110670656 | 110670715 |
| A_23_P144458 | CAMK2D | CAMK2D | chr4 | 114381361 | 114386707 |
| A_33_P3268224 | SPATA5 | SPATA5 | chr4 | 124240537 | 124240596 |
| A_33_P3801085 | LOC641365 | LOC641365 | chr4 | 138948608 | 138948667 |
| A_33_P3214625 | INPP4B | INPP4B | chr4 | 142949208 | 142949267 |
| A_33_P3219115 | LOC100130178 | INPP4B | chr4 | 143238618 | 143238677 |
| A_33_P3236436 | C4orf51 | C4orf51 | chr4 | 146653856 | 146653915 |
| A_23_P320290 | ZNF827 | ZNF827 | chr4 | 146686254 | 146686313 |
| A_33_P3235891 | NOGENE | GALNTL6 | chr4 | 173908624 | 173908683 |
| A_33_P3320017 | GALNTL6 | GALNTL6 | chr4 | 173961168 | 173961227 |
| A_33_P3320022 | GALNTL6 | GALNTL6 | chr4 | 173961500 | 173961559 |
| A_24_P71904 | HPGD | HPGD | chr4 | 175411686 | 175411745 |
| A_23_P252236 | KLKB1 | KLKB1 | chr4 | 187178445 | 187178504 |
| A_23_P110624 | CTNND2 | CTNND2 | chr5 | 10973084 | 10973143 |
| A_19_P00327730 | NOGENE | TAG | chr5 | 12719497 | 12719556 |
| A_19_P00802660 | NOGENE | TAG | chr5 | 12734994 | 12735053 |
| A_33_P3826455 | TAG | TAG | chr5 | 12759692 | 12759751 |
| A_33_P3265941 | NOGENE | TAG | chr5 | 12794242 | 12794301 |
| A_19_P00316231 | NOGENE | TAG | chr5 | 12804384 | 12804443 |
| A_24_P7600 | FBXL7 | FBXL7 | chr5 | 15937236 | 15937295 |
| A_23_P144827 | FBXL7 | FBXL7 | chr5 | 15939565 | 15939624 |
| A_32_P10936 | CDH12 | CDH12 | chr5 | 21751312 | 21751368 |
| A_23_P92727 | RAI14 | RAI14 | chr5 | 34831952 | 34832011 |
| A_24_P40795 | NOGENE | NNT | chr5 | 43667002 | 43667061 |
| A_23_P70148 | NNT | NNT | chr5 | 43704546 | 43704605 |
| A_33_P3365995 | MAST4 | MAST4 | chr5 | 66255013 | 66255072 |
| A_23_P110571 | MAST4 | MAST4 | chr5 | 66462808 | 66462867 |
| A_23_P259521 | WDR41 | WDR41 | chr5 | 76728963 | 76729022 |
| A_33_P3359846 | NOGENE | SERINC5 | chr5 | 79439878 | 79439937 |
| A_23_P423457 | SERINC5 | SERINC5 | chr5 | 79473159 | 79473218 |
| A_24_P111912 | FAM172A | FAM172A | chr5 | 92953936 | 92953995 |
| A_23_P358564 | POU5F2 | FAM172A | chr5 | 93076020 | 93076079 |
| A_33_P3401058 | NOGENE | FAM172A | chr5 | 93198654 | 93198713 |
| A_24_P149124 | C5orf13 | C5orf13 | chr5 | 111065091 | 111065150 |
| A_33_P3391290 | C5orf13 | C5orf13 | chr5 | 111066368 | 111066427 |
| A_23_P346982 | DTWD2 | DTWD2 | chr5 | 118175647 | 118175706 |
| A_23_P144999 | RAPGEF6 | RAPGEF6 | chr5 | 130762075 | 130762134 |
| A_33_P3243519 | RAPGEF6 | RAPGEF6 | chr5 | 130766553 | 130766612 |
| A_33_P3243524 | RAPGEF6 | RAPGEF6 | chr5 | 130771644 | 130771703 |
| A_23_P110643 | CDKL3 | CDKL3 | chr5 | 133643886 | 133643945 |
| A_33_P3337632 | TRPC7 | TRPC7 | chr5 | 135549123 | 135549182 |
| A_24_P140569 | LRRTM2 | CTNNA1 | chr5 | 138205569 | 138205628 |
| A_33_P3251024 | LRRTM2 | CTNNA1 | chr5 | 138208699 | 138208758 |
| A_33_P3403418 | CTNNA1 | CTNNA1 | chr5 | 138266534 | 138266593 |
| A_23_P58647 | CTNNA1 | CTNNA1 | chr5 | 138270387 | 138270446 |


| A_32_P222695 | FLJ41603 | ARHGEF37 | chr5 | 149014156 | 149014215 |
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| A_33_P3245218 | ODZ2 | ODZ2 | chr5 | 167689754 | 167689813 |
| A_24_P299474 | ODZ2 | ODZ2 | chr5 | 167690063 | 167690122 |
| A_23_P58819 | RANBP17 | RANBP17 | chr5 | 170725916 | 170725975 |
| A_23_P321307 | ADAMTS2 | ADAMTS2 | chr5 | 178608104 | 178634520 |
| A_33_P3411204 | GCNT2 | GCNT2 | chr6 | 10627251 | 10627310 |
| A_24_P397489 | GCNT2 | GCNT2 | chr6 | 10628471 | 10628530 |
| A_24_P277657 | GMPR | GMPR | chr6 | 16290766 | 16290825 |
| A_33_P3212109 | DCDC2 | DCDC2 | chr6 | 24172180 | 24172239 |
| A_24_P166407 | HIST1H4B | HIST1H4B | chr6 | 26027149 | 26027208 |
| A_23_P259098 | ZSCAN16 | ZSCAN16 | chr6 | 28097263 | 28097322 |
| A_23_P42288 | C6orf27 | C6orf27 | chr6 | 31734142 | 31734279 |
| A_33_P3342420 | NOGENE | DNAH8 | chr6 | 38899007 | 38899066 |
| A_23_P145159 | DNAH8 | DNAH8 | chr6 | 38980339 | 38980398 |
| A_23_P145175 | ZNF318 | ZNF318 | chr6 | 43303992 | 43304051 |
| A_24_P204043 | ZNF318 | ZNF318 | chr6 | 43316128 | 43316187 |
| A_24_P204144 | NOGENE | ZNF318 | chr6 | 43331750 | 43331810 |
| A_23_P402187 | PKHD1 | PKHD1 | chr6 | 51480343 | 51480402 |
| A_23_P424617 | PKHD1 | PKHD1 | chr6 | 51586468 | 51586527 |
| A_33_P3387420 | PKHD1 | PKHD1 | chr6 | 51524767 | 51609241 |
| A_32_P480330 | EYS | EYS | chr6 | 64430448 | 64430507 |
| A_33_P3276833 | EYS | EYS | chr6 | 64430609 | 64430668 |
| A_19_P00803113 | NOGENE | EYS | chr6 | 64472319 | 64472378 |
| A_33_P3276808 | NOGENE | EYS | chr6 | 64472322 | 64472381 |
| A_19_P00321801 | NOGENE | EYS | chr6 | 64472463 | 64472522 |
| A_19_P00321802 | NOGENE | EYS | chr6 | 64487899 | 64487958 |
| A_19_P00316319 | NOGENE | EYS | chr6 | 64488013 | 64488072 |
| A_19_P00321803 | NOGENE | EYS | chr6 | 64488009 | 64488068 |
| A_33_P3276818 | EYS | EYS | chr6 | 65303024 | 65303083 |
| A_33_P3276828 | NOGENE | EYS | chr6 | 65532725 | 65532784 |
| A_33_P3276813 | EYS | EYS | chr6 | 65596591 | 65596650 |
| A_33_P3276805 | NOGENE | EYS | chr6 | 66005651 | 66005710 |
| A_24_P944973 | EYS | EYS | chr6 | 66044872 | 66044931 |
| A_33_P3359115 | LMBRD1 | LMBRD1 | chr6 | 70385779 | 70385838 |
| A_33_P3248799 | TRDN | TRDN | chr6 | 123869600 | 123869659 |
| A_23_P93524 | SAMD3 | SAMD3 | chr6 | 130465781 | 130465840 |
| A_23_P397937 | SAMD3 | SAMD3 | chr6 | 130505674 | 130505733 |
| A_23_P134113 | C6orf192 | C6orf192 | chr6 | 133090550 | 133090609 |
| A_33_P3321230 | C6orf192 | C6orf192 | chr6 | 133095381 | 133095440 |
| A_23_P134125 | MAP3K5 | MAP3K5 | chr6 | 136878349 | 136878408 |
| A_33_P3377130 | MAP3K5 | MAP3K5 | chr6 | 136901467 | 136901526 |
| A_33_P3246580 | KIAA1244 | KIAA1244 | chr6 | 138657434 | 138657493 |
| A_23_P393880 | KIAA1244 | KIAA1244 | chr6 | 138659313 | 138659372 |
| A_33_P3387951 | KIAA1244 | KIAA1244 | chr6 | 138662672 | 138662731 |
| A_33_P3416574 | AIG1 | AIG1 | chr6 | 143458067 | 143458126 |
| A_33_P3416563 | AIG1 | AIG1 | chr6 | 143654522 | 143654581 |


| A_23_P93431 | AIG1 | AIG1 | chr6 | 143661259 | 143661318 |
| :---: | :---: | :---: | :---: | :---: | :---: |
| A_33_P3412087 | C6orf97 | C6orf97 | chr6 | 151939801 | 151939860 |
| A_33_P3270489 | C6orf97 | C6orf97 | chr6 | 151942097 | 151942156 |
| A_23_P500861 | SYNE1 | SYNE1 | chr6 | 152443052 | 152443111 |
| A_33_P3335920 | SYNE1 | SYNE1 | chr6 | 152443528 | 152443587 |
| A_33_P3288919 | NOGENE | SYNE1 | chr6 | 152488859 | 152488918 |
| A_33_P3335915 | SYNE1 | SYNE1 | chr6 | 152748771 | 152748830 |
| A_33_P3335910 | SYNE1 | SYNE1 | chr6 | 152757076 | 152757134 |
| A_24_P58308 | NOGENE | ARID1B | chr6 | 157298027 | 157298086 |
| A_33_P3291877 | ARID1B | ARID1B | chr6 | 157469939 | 157469998 |
| A_33_P3291882 | NOGENE | ARID1B | chr6 | 157507679 | 157507738 |
| A_23_P70701 | ARID1B | ARID1B | chr6 | 157529022 | 157529081 |
| A_33_P3333078 | NOGENE | ARID1B | chr6 | 157531787 | 157531846 |
| A_23_P382045 | TULP4 | TULP4 | chr6 | 158932662 | 158932721 |
| A_33_P3312119 | LOC100130967 | C6orf99 | chr6 | 159331223 | 159331282 |
| A_23_P111395 | SLC22A2 | SLC22A2 | chr6 | 160638226 | 160638285 |
| A_23_P136077 | PARK2 | PARK2 | chr6 | 161769778 | 161769837 |
| A_23_P145134 | FGFR1OP | FGFR1OP | chr6 | 167453724 | 167453783 |
| A_32_P415151 | WDR27 | WDR27 | chr6 | 170033151 | 170034569 |
| A_23_P336992 | ZFAND2A | ZFAND2A | chr7 | 1195219 | 1197349 |
| A_33_P3251227 | C1GALT1 | C1GALT1 | chr7 | 7278507 | 7278566 |
| A_23_P252145 | C1GALT1 | C1GALT1 | chr7 | 7283565 | 7283624 |
| A_19_P00805702 | NOGENE | C1GALT1 | chr7 | 7288141 | 7288200 |
| A_24_P129277 | NOD1 | NOD1 | chr7 | 30464987 | 30465046 |
| A_24_P272967 | AVL9 | AVL9 | chr7 | 32598953 | 32599012 |
| A_23_P122947 | AVL9 | AVL9 | chr7 | 32619842 | 32620430 |
| A_23_P145718 | AOAH | AOAH | chr7 | 36561708 | 36570069 |
| A_23_P319027 | HECW1 | HECW1 | chr7 | 43602391 | 43602450 |
| A_23_P45087 | ZNF107 | ZNF107 | chr7 | 64171177 | 64171236 |
| A_23_P111517 | WBSCR17 | WBSCR17 | chr7 | 71177978 | 71178037 |
| A_23_P215484 | CCL26 | CCL26 | chr7 | 75399061 | 75401219 |
| A_23_P416894 | PION | PION | chr7 | 76940387 | 76940446 |
| A_33_P3280694 | PION | PION | chr7 | 76958871 | 76958930 |
| A_23_P21376 | MAGI2 | MAGI2 | chr7 | 77646565 | 77646624 |
| A_33_P3311992 | MAGI2 | MAGI2 | chr7 | 77649075 | 77649134 |
| A_33_P3312802 | RPL13AP17 | MAGI2 | chr7 | 77988714 | 77988773 |
| A_24_P123833 | SEMA3E | SEMA3E | chr7 | 82993805 | 82993864 |
| A_23_P59528 | ACN9 | ACN9 | chr7 | 96810787 | 96810846 |
| A_23_P358917 | CYP3A7 | CYP3A7 | chr7 | 99303037 | 99303096 |
| A_33_P3318117 | CYP3A7 | CYP3A7 | chr7 | 99303139 | 99303198 |
| A_33_P3264404 | LOC100132593 | CUX1 | chr7 | 101818665 | 101818724 |
| A_33_P3298785 | CUX1 | CUX1 | chr7 | 101892263 | 101892322 |
| A_24_P220454 | CUX1 | CUX1 | chr7 | 101893025 | 101893084 |
| A_23_P253375 | CUX1 | CUX1 | chr7 | 101926989 | 101927048 |
| A_23_P59637 | DOCK4 | DOCK4 | chr7 | 111366266 | 111366325 |
| A_33_P3253420 | NOGENE | DOCK4 | chr7 | 111611353 | 111611412 |


| A_33_P3354001 | CADPS2 | CADPS2 | chr7 | 121958491 | 121958550 |
| :---: | :---: | :---: | :---: | :---: | :---: |
| A_23_P42963 | RNF133 | CADPS2 | chr7 | 122338036 | 122338095 |
| A_24_P170234 | RNF148 | CADPS2 | chr7 | 122341976 | 122342035 |
| A_33_P3397418 | ZC3HAV1 | ZC3HAV1 | chr7 | 138728351 | 138728410 |
| A_33_P3298980 | NOGENE | ZC3HAV1 | chr7 | 138734016 | 138734075 |
| A_23_P20122 | ZC3HAV1 | ZC3HAV1 | chr7 | 138745443 | 138745502 |
| A_33_P3251801 | KLRG2 | KLRG2 | chr7 | 139138088 | 139138147 |
| A_33_P3306207 | KLRG2 | KLRG2 | chr7 | 139138858 | 139138917 |
| A_23_P309224 | AGK | AGK | chr7 | 141353184 | 141353243 |
| A_33_P3367196 | CNTNAP2 | CNTNAP2 | chr7 | 148117937 | 148117996 |
| A_33_P3264089 | ACTR3C | ACTR3C | chr7 | 149992342 | 149992401 |
| A_24_P132383 | GIMAP8 | GIMAP8 | chr7 | 150175990 | 150176049 |
| A_33_P3230244 | CSMD1 | CSMD1 | chr8 | 2792900 | 2792959 |
| A_33_P3230249 | CSMD1 | CSMD1 | chr8 | 2984937 | 2984996 |
| A_23_P31798 | NAT2 | NAT2 | chr8 | 18258142 | 18258201 |
| A_23_P94103 | SCARA5 | SCARA5 | chr8 | 27727875 | 27727934 |
| A_23_P215931 | LEPROTL1 | LEPROTL1 | chr8 | 29965452 | 29965511 |
| A_24_P306824 | MBOAT4 | LEPROTL1 | chr8 | 29989463 | 29989522 |
| A_33_P3252206 | LEPROTL1 | LEPROTL1 | chr8 | 29995141 | 29995200 |
| A_23_P72527 | ADAM18 | ADAM18 | chr8 | 39581412 | 39587461 |
| A_23_P83835 | SGK196 | SGK196 | chr8 | 42977869 | 42977928 |
| A_24_P4170 | CPA6 | CPA6 | chr8 | 68396078 | 68396957 |
| A_33_P3391778 | RALYL | RALYL | chr8 | 85441725 | 85441784 |
| A_33_P3316899 | RALYL | RALYL | chr8 | 85799861 | 85799920 |
| A_23_P388220 | RALYL | RALYL | chr8 | 85799966 | 85833142 |
| A_23_P371861 | CNBD1 | CNBD1 | chr8 | 88298829 | 88298888 |
| A_23_P252106 | RIPK2 | RIPK2 | chr8 | 90802345 | 90802404 |
| A_33_P3371219 | SDC2 | SDC2 | chr8 | 97623978 | 97624037 |
| A_24_P180680 | LAPTM4B | LAPTM4B | chr8 | 98864581 | 98864640 |
| A_24_P332816 | RIMS2 | RIMS2 | chr8 | 104928692 | 104928751 |
| A_23_P147786 | RIMS2 | RIMS2 | chr8 | 105263899 | 105263958 |
| A_24_P106794 | NUDCD1 | NUDCD1 | chr8 | 110253382 | 110253441 |
| A_23_P60166 | DEPDC6 | DEPTOR | chr8 | 121062582 | 121062641 |
| A_19_P00321436 | NOGENE | HHLA1 | chr8 | 133073733 | 133073792 |
| A_33_P3301050 | HHLA1 | HHLA1 | chr8 | 133073733 | 133073792 |
| A_19_P00319204 | NOGENE | HHLA1 | chr8 | 133076177 | 133076236 |
| A_19_P00330652 | NOGENE | HHLA1 | chr8 | 133088841 | 133088900 |
| A_19_P00321196 | NOGENE | HHLA1 | chr8 | 133090100 | 133090159 |
| A_19_P00319586 | NOGENE | HHLA1 | chr8 | 133092154 | 133092213 |
| A_23_P384023 | GLIS3 | GLIS3 | chr9 | 3828374 | 3829334 |
| A_33_P3326568 | GLIS3 | GLIS3 | chr9 | 3855481 | 3855540 |
| A_23_P312837 | C9orf70 | GLIS3 | chr9 | 3898837 | 3898896 |
| A_33_P3309551 | PTPRD | PTPRD | chr9 | 8314275 | 8314334 |
| A_24_P639665 | NOGENE | PTPRD | chr9 | 8861154 | 8861213 |
| A_32_P395879 | C9orf93 | C9orf93 | chr9 | 15920369 | 15971614 |
| A_33_P3608210 | LOC554202 | LOC554202 | chr9 | 21454716 | 21454775 |


| A_23_P302060 | IFNE | LOC554202 | chr9 | 21481102 | 21481161 |
| :---: | :---: | :---: | :---: | :---: | :---: |
| A_24_P255100 | C9orf135 | C9orf135 | chr9 | 72471488 | 72471547 |
| A_23_P32078 | SLC28A3 | SLC28A3 | chr9 | 86900326 | 86900385 |
| A_23_P43326 | SPTLC1 | SPTLC1 | chr9 | 94793717 | 94793776 |
| A_32_P48949 | C9orf129 | C9orf129 | chr9 | 96080485 | 96080544 |
| A_33_P3351175 | WNK2 | C9orf129 | chr9 | 96082754 | 96082813 |
| A_23_P157970 | INVS | INVS | chr9 | 103062898 | 103062957 |
| A_24_P235429 | ABCA1 | ABCA1 | chr9 | 107543704 | 107543763 |
| A_33_P3422897 | ABCA1 | ABCA1 | chr9 | 107624012 | 107624071 |
| A_23_P411806 | SLC44A1 | SLC44A1 | chr9 | 108136938 | 108136997 |
| A_23_P216630 | SLC44A1 | SLC44A1 | chr9 | 108153508 | 108153567 |
| A_23_P146486 | 1-Dec | DEC1 | chr9 | 118164718 | 118164777 |
| A_23_P216894 | MAPKAP1 | MAPKAP1 | chr9 | 128200564 | 128200623 |
| A_33_P3251538 | MAPKAP1 | MAPKAP1 | chr9 | 128305339 | 128305398 |
| A_33_P3397755 | MAPKAP1 | MAPKAP1 | chr9 | 128347923 | 128347982 |
| A_33_P3407937 | PLCXD1 | PLCXD1 | chrX | 208260 | 208319 |
| A_23_P61180 | PLCXD1 | PLCXD1 | chrX | 219727 | 219786 |
| A_23_P217704 | GYG2 | GYG2 | $\operatorname{chrX}$ | 2800115 | 2800174 |
| A_23_P254842 | HDHD1A | HDHD1 | $\operatorname{chrX}$ | 6967361 | 6967420 |
| A_23_P429950 | KAL1 | KAL1 | chr $X$ | 8497233 | 8497292 |
| A_24_P940275 | FRMPD4 | FRMPD4 | chrX | 12739959 | 12740018 |
| A_24_P672240 | FRMPD4 | FRMPD4 | chrX | 12742371 | 12742430 |
| A_33_P3780311 | LOC349408 | LOC349408 | chrX | 12921157 | 12921216 |
| A_23_P73837 | TLR8 | LOC349408 | $\operatorname{chrX}$ | 12940262 | 12940321 |
| A_33_P3224595 | OFD1 | OFD1 | chrX | 13769427 | 13769486 |
| A_24_P134653 | OFD1 | OFD1 | chrX | 13786879 | 13787218 |
| A_33_P3394178 | NHS | NHS | chrX | 17750175 | 17750234 |
| A_32_P126375 | NHS | NHS | chrX | 17753480 | 17753539 |
| A_33_P3351510 | IL1RAPL1 | IL1RAPL1 | chrX | 29973954 | 29974013 |
| A_24_P185854 | DMD | DMD | chrX | 31137349 | 31137408 |
| A_33_P3284763 | DMD | DMD | chrX | 31152220 | 31152279 |
| A_23_P321860 | DMD | DMD | chr X | 31196312 | 31196342 |
| A_33_P3297818 | NOGENE | DMD | chrX | 31199562 | 31199621 |
| A_24_P195724 | NOGENE | DMD | chrX | 32224776 | 32224837 |
| A_33_P3297813 | DMD | DMD | chrX | 32535057 | 32535116 |
| A_33_P3297808 | NOGENE | DMD | chrX | 32562406 | 32562465 |
| A_33_P3415087 | CLCN5 | CLCN5 | chrX | 49856988 | 49857047 |
| A_33_P3415092 | CLCN5 | CLCN5 | chrX | 49863833 | 49863892 |
| A_23_P113471 | FAAH2 | FAAH2 | chrX | 57515446 | 57515505 |
| A_24_P404840 | GJB1 | BCYRN1 | chrX | 70444287 | 70444346 |
| A_23_P137073 | ZMYM3 | BCYRN1 | chrX | 70459456 | 70459485 |
| A_24_P16815 | ZMYM3 | BCYRN1 | chrX | 70469422 | 70469481 |
| A_33_P3247753 | TSIX | TSIX | chrX | 73027005 | 73027064 |
| A_33_P3341686 | XIST | TSIX | chrX | 73040506 | 73040565 |
| A_19_P00331623 | NOGENE | TSIX | chrX | 73040566 | 73040625 |
| A_19_P00329511 | NOGENE | TSIX | chrX | 73040688 | 73040747 |


| A_19_P00323692 | NOGENE | TSIX | chrX | 73040697 | 73040756 |
| :---: | :---: | :---: | :---: | :---: | :---: |
| A_19_P00319151 | NOGENE | TSIX | chrX | 73040732 | 73040791 |
| A_19_P00321129 | NOGENE | TSIX | chrX | 73041907 | 73041966 |
| A_19_P00321917 | NOGENE | TSIX | chrX | 73043015 | 73043074 |
| A_19_P00802872 | NOGENE | TSIX | chrX | 73043696 | 73043755 |
| A_19_P00806762 | NOGENE | TSIX | chrX | 73043894 | 73043953 |
| A_19_P00332515 | NOGENE | TSIX | chrX | 73047086 | 73047145 |
| A_19_P00320438 | NOGENE | TSIX | chrX | 73048929 | 73048988 |
| A_19_P00316565 | NOGENE | TSIX | chrX | 73049001 | 73049060 |
| A_19_P00326132 | NOGENE | TSIX | chrX | 73049003 | 73049062 |
| A_33_P3405911 | TSIX | TSIX | chrX | 73049007 | 73049066 |
| A_19_P00319561 | NOGENE | MIR374AHG | chrX | 73289101 | 73289160 |
| A_19_P00320488 | NOGENE | MIR374AHG | chrX | 73289534 | 73289593 |
| A_33_P3294985 | NCRNA00183 | MIR374AHG | chrX | 73289703 | 73289762 |
| A_19_P00318107 | NOGENE | MIR374AHG | chrX | 73290680 | 73290739 |
| A_33_P3407235 | LOC100133180 | MIR374AHG | chrX | 73351716 | 73351775 |
| A_32_P464135 | DACH2 | DACH2 | chrX | 86087385 | 86087444 |
| A_33_P3363465 | NOGENE | PCDH11X | chrX | 91355188 | 91355247 |
| A_33_P3346448 | PCDH11X | PCDH11X | chrX | 91873836 | 91873895 |
| A_23_P254212 | RPA4 | DIAPH2 | chrX | 96140272 | 96140331 |
| A_23_P85004 | DIAPH2 | DIAPH2 | chrX | 96369895 | 96369954 |
| A_24_P144303 | LOC442459 | LOC442459 | chrX | 98716600 | 98716642 |
| A_33_P3318424 | NOGENE | BHLHB9 | chrX | 102000979 | 102001038 |
| A_23_P308954 | BHLHB9 | BHLHB9 | chrX | 102006979 | 102007038 |
| A_33_P3319176 | HTR2C | HTR2C | chrX | 114143005 | 114143064 |
| A_23_P433586 | HTR2C | HTR2C | chrX | 114143855 | 114143914 |
| A_33_P3281308 | LOC286467 | LOC286467 | chrX | 130836687 | 130836746 |
| A_19_P00325063 | NOGENE | LOC286467 | chrX | 130845660 | 130845719 |
| A_32_P137980 | PCDH11Y | PCDH11Y | chrY | 5610107 | 5610166 |
| A_23_P160017 | TTTY11 | TTTY11 | chrY | 8652207 | 8657062 |

Table S1D. List of primer sequences

|  | Figures |  |  | sequence | others |
| :---: | :---: | :---: | :---: | :---: | :---: |
| pyro sequencing | Fig.1F | hPON3 | forward primer | GGGAATGGAGGGGAGTTTTAGTTTAGAG | cg24750391 |
|  |  |  | reverse primer | CCTATCTTTTCCTTCTTTTCTCCTAATAT (5'-biotinylated) |  |
|  |  |  | sequence primer | TGTTTTATTTAGGAGTGTGTTG |  |
|  | Fig. 1G | hTCERG1L | forward primer | GGGTGTTTGGTTTGAAGTT (5'-biotinylated) | cg03943081 |
|  |  |  | reverse primer | AATAATCCTACCCCACCCAAAAAATATC |  |
|  |  |  | sequence primer | CCACCTACCTAATACCTT |  |
|  | Fig. 3F | $\begin{aligned} & \text { hABHD12B } \\ & \text { 5'LTR-1 } \end{aligned}$ | forward primer | TGTGTATTAATGTATGGTTAATTTTGGTAA |  |
|  |  |  | reverse primer | CAAACCATCTAAACAAATACCTACAA (5'biotinylated) |  |
|  |  |  | sequence primer | GTTGTTTTTTATGTAGTGTTT |  |
|  |  | hABHD12B 5'LTR-2 | forward primer | TGTGTATTAATGTATGGTTAATTTTGGTAA |  |
|  |  |  | reverse primer | CAAACCATCTAAACAAATACCTACAA (5'biotinylated) |  |
|  |  |  | sequence primer | TTAGGTTTTTGAGTTTAAGTTAA |  |
|  |  | hABHD12B | forward primer | AAGTTTGTTTGGTGGTTTTTTTATATAGA (5'biotinylated) |  |




Table S2. Information about transplantation, graft size and compoments

|  | experiment No. | clone name | passage No. | qPCR (pretransplantation) | Graft size (Day30) | Graft size (Day60) | graft |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |  | neural | nonneural |  |
| defective (42) | \#0 | TKCBV5-6 | p99 | N.D. | N.D. | N.D. | 0 | 10 | A |
|  | \#1-1 | TKCBV5-6 | p101 | $\checkmark$ | N.D. | 8.552 | 1 | 9 | A |
|  | \#1-2 | TKCBV5-6 | p101 | $\checkmark$ | N.D. | 9.66 | 2 | 8 | A |
|  | \#3-1 | TKCBV5-6 | p82 | $\checkmark$ | 4.939 | 10.15 | 9 | 1 | D |
|  | \#3-2 | TKCBV5-6 | p82 | $\checkmark$ | 4.787 | 16.46 | 6 | 4 | B |
|  | \#3-3 | TKCBV5-6 | p82 | $\checkmark$ | 8.556 | 20.68 | 8 | 2 | C |
|  | \#0 | TIG118-4f1 | p43 | N.D. | N.D. | 20.35 | 9 | 1 | D |
|  | \#1-1 | TIG118-4f1 | p45 | $\checkmark$ | N.D. | 4.888 | 10 | 0 | E |
|  | \#1-2 | TIG118-4f1 | p45 | $\checkmark$ | N.D. | 38.71 | 8 | 2 | C |
|  | \#3-1 | TIG118-4f1 | p53 | $\checkmark$ | 1.056 | 22.66 | 3 | 7 | A |
|  | \#3-2 | TIG118-4f1 | p53 | $\checkmark$ | 3.492 | 16.62 | 8 | 2 | C |
|  | \#3-3 | TIG118-4f1 | p53 | $\checkmark$ | 9.194 | 28.85 | 9 | 1 | D |
|  | \#0 | TIG108-4f3 | p36 | N.D. | N.D. | N.D. | 7 | 3 | C |
|  | \#1-1 | TIG108-4f3 | p30 | $\checkmark$ | N.D. | Died | 5 | 5 | B |
|  | \#1-2 | TIG108-4f3 | p30 | $\checkmark$ | N.D. | Died | 5 | 5 | B |
|  | \#3-1 | TIG108-4f3 | p17 | $\checkmark$ | 10.86 | Died | 6 | 4 | B |
|  | \#3-3 | TIG108-4f3 | p17 | $\checkmark$ | 17.88 | Died | 5 | 5 | B |
|  | \#4-1 | TIG108-4f3 | p18 | $\checkmark$ | 6.601 | Died | 5 | 5 | B |
|  | \#4-2 | TIG108-4f3 | p18 | $\checkmark$ | 6.889 | 16.71 | 6 | 4 | B |
|  | \#4-3 | TIG108-4f3 | p18 | $\checkmark$ | 14.82 | Died | 7 | 3 | C |
|  | \#9-1 | 1503-4f1-1 | p54 | $\checkmark$ | 3.152 | 12.87 | 9 | 1 | D |
|  | \#9-1 | 1503-4f1-2 | p54 | $\checkmark$ | 0.783 | 12.328 |  |  | N.D. |
|  | \#0 | TIG107-3f1 | p48 | N.D. | N.D. | N.D. | 5 | 5 | B |
|  | \#1-1 | TIG107-3f1 | p56 | $\checkmark$ | N.D. | Died | 7 | 3 | C |
|  | \#1-2 | TIG107-3f1 | p56 | $\checkmark$ | N.D. | Died | 5 | 5 | B |
|  | \#2-1 | TIG107-3f1 | p56 | $\checkmark$ | 21.05 | Died | 9 | 1 | D |
|  | \#2-2 | TIG107-3f1 | p56 | $\checkmark$ | 11.03 | Died | 9 | 1 | D |
|  | \#2-3 | TIG107-3f1 | p56 | $\checkmark$ | 21.25 | Died | 9 | 1 | D |
|  | \#0 | 451F3 | p75 | N.D. | N.D. | 22.81 | 9 | 1 | D |
|  | \#1-1 | 451F3 | p77 | $\checkmark$ | N.D. | 11.43 | 10 | 0 | E |
|  | \#1-2 | 451F3 | p77 | $\checkmark$ | N.D. | 13.14 | 10 | 0 | E |
|  | \#2-1 | 451F3 | p65 | $\checkmark$ | 2.61 | 7.556 | 10 | 0 | [ |
|  | \#2-2 | 451F3 | p65 | $\checkmark$ | 2.937 | 9.967 | 8 | 2 | C |
|  | \#2-3 | 451F3 | p65 | $\checkmark$ | 8.2 | 34.77 | 7 | 3 | C |
|  | \#7-1 | 451F3 | p63 | $\checkmark$ | 4.852 | 11.82 | 10 | 0 | E |
|  | \#7-2 | 451F3 | p63 | $\checkmark$ | 2.441 | 6.26 | 9 | 1 | D |
|  | \#7-3 | 451F3 | p63 | $\checkmark$ | 2.99 | 1.46 | 1 | 9 | A |


| neural tissue <br> containing |  |
| :---: | :---: |
| $0-4$ | A |
| $5-6$ | B |
| $7-8$ | C |
| 9 | D |
| 10 | E |
| not determined | N.D. |


|  | \#2-1 | TIG107-4f1 | p24 | $\checkmark$ | 3.608 | 14.17 | 6 | 4 | B |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | \#2-2 | TIG107-4f1 | p24 | $\checkmark$ | 3.801 | 23.31 | 7 | 3 | C |
|  | \#2-3 | TIG107-4f1 | p24 | $\checkmark$ | 4.889 | 24.22 | 6 | 4 | B |
|  | \#9-1 | TIG107-4f1-1 | p24 | $\checkmark$ | 2.076 | 12.07 | 5 | 5 | B |
|  | \#9-1 | TIG107-441-2 | p24 | $\checkmark$ | 3.715 | 10.598 | 6 | 4 | B |
|  | \#8-2 | 604A1 | p32 | $\checkmark$ | N.D. | N.D. |  |  | N.D. |
|  | \#8-3 | 604A1 | p32 | $\checkmark$ | 2.411 | 7.894 | 10 | 0 | E |
|  | \#8-1 | 610B1 | p24 | $\checkmark$ | 1.196 | 4.279 | 10 | 0 | E |
|  | \#8-2 | 610B1 | p24 | $\checkmark$ | 1.906 | 3.741 | 10 | 0 | E |
| intermediate (9) | \#8-3 | 610B1 | p24 | $\checkmark$ | 0.7776 | 1.968 | 10 | 0 | E |
|  | \#0 | khES1 | p79 | N.D. | N.D. | 12.32 | 10 | 0 | E |
|  | \#7-1 | khES1 | p72 | $\checkmark$ | 4.879 | 9.16 | 10 | 0 | E |
|  | \#7-2 | khES1 | p72 | $\checkmark$ | 4.65 | 8.917 | 10 | 0 | E |
|  | \#7-3 | khES1 | p72 | $\checkmark$ | 2.66 | 6.951 | 8 | 2 | C |
| good (63) | \#8-1 | 604B1 | p24 | $\checkmark$ | 2.924 | 5.083 | 10 | 0 | E |
|  | \#8-2 | 604B1 | p24 | $\checkmark$ | 2.131 | 5.082 | 10 | 0 | E |
|  | \#8-3 | 604B1 | p24 | $\checkmark$ | 1.852 | 8.705 | 10 | 0 | E |
|  | \#7-1 | TKCBV4-2 | p77 | $\checkmark$ | 0.862 | 9.654 | 0 | 10 | A |
|  | \#7-2 | TKCBV4-2 | p77 | $\checkmark$ | 0.193 | Died | 0 | 10 | A |
|  | \#7-3 | TKCBV4-2 | p77 | $\checkmark$ | 1.992 | 6.908 | 0 | 10 | A |
|  | \#7-1 | TKCB7-2 | p41 | $\checkmark$ | 2.751 | 12.776 | 10 | 0 | E |
|  | \#7-2 | TKCB7-2 | p41 | $\checkmark$ | 1.666 | 2.939 | 10 | 0 | E |
|  | \#7-3 | TKCB7-2 | p41 | $\checkmark$ | 0.76 | Died | 6 | 4 | B |
|  | \#0 | H9 | p47 | N.D. | N.D. | N.D. | 10 | 0 | E |
|  | \#2-1 | H9 | p50 | $\checkmark$ | 1.338 | 3.081 | 10 | 0 | E |
|  | \#2-2 | H9 | p50 | $\checkmark$ | N.D. | N.D. | 10 | 0 | E |
|  | \#2-3 | H9 | p50 | $\checkmark$ | 4.403 | 13.42 | 10 | 0 | E |
|  | \#4-1 | H9 | p16 | $\checkmark$ | 3.047 | 10.49 | 9 | 1 | D |
|  | \#4-2 | H9 | p16 | $\checkmark$ | 5.178 | 15.91 | 9 | 1 | D |
|  | \#4-3 | H9 | p16 | $\checkmark$ | 1.625 | 5.94 | 5 | 5 | B |
|  | \#7-1 | 606A1 | p27 | $\checkmark$ | 2.359 | 2.903 | 10 | 0 | E |
|  | \#7-2 | 606A1 | p27 | $\checkmark$ | N.D. | 5.734 | 10 | 0 | E |
|  | \#7-3 | 606A1 | p27 | $\checkmark$ | 5.241 | 16.14 | 10 | 0 | E |
|  | \#8-2 | 606B1 | p23 | $\checkmark$ | 2.184 | 6.033 | 10 | 0 | E |
|  | \#8-3 | 606B1 | p23 | $\checkmark$ | 1.655 | 1.646 | 10 | 0 | E |
|  | \#3-1 | 253G4 | p27 | $\checkmark$ | 1.19 | 1.814 | 10 | 0 | E |
|  | \#3-2 | 253G4 | p27 | $\checkmark$ | 0.5012 | 1.252 | 10 | 0 | E |
|  | \#3-3 | 253G4 | p27 | $\checkmark$ | 1.219 | 1.839 | 10 | 0 | E |
|  | \#2-1 | 201B7 | p25 | $\checkmark$ | 0.8161 | 1.747 | 10 | 0 | E |
|  | \#2-2 | 201B7 | p25 | $\checkmark$ | 1.462 | 2.485 | 10 | 0 | E |
|  | \#2-3 | 201B7 | p25 | $\checkmark$ | 2.052 | 2.568 | 10 | 0 | E |
|  | \#0 | khES3 | p76 | N.D. | N.D. | 2.405 | 10 | 0 | E |
|  | \#1-1 | khES3 | p78 | $\checkmark$ | N.D. | 1.177 | 10 | 0 | E |

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| \#1-2 | khES3 | p78 | $\checkmark$ | N.D. | 0.5812 | 10 | 0 |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| \#3-1 | khES3 | p76 | $\checkmark$ | 1.105 | 6.036 | 10 | 0 | E |
| \#3-3 | khES3 | p76 | $\checkmark$ | 1.321 | 2.151 | 10 | 0 | L |
| \#8-2 | 604A3 | p25 | $\checkmark$ | 2.528 | 4.077 | 10 | 0 | E |
| \#8-3 | 604A3 | p25 | $\checkmark$ | 1.611 | 2.911 | 10 | 0 |  |
| \#7-1 | 409B2 | p24 | $\checkmark$ | 0.5389 | 1.034 | 10 | 0 |  |
| \#7-2 | 409B2 | p24 | $\checkmark$ | 0.541 | 1.093 | 10 | 0 | E |
| \#7-3 | 409B2 | p24 | $\checkmark$ | 0.627 | 0.2324 | 10 | 0 |  |
| \#8-1 | H1 | p56 | $\checkmark$ | 2.652 | 9.702 | 10 | 0 | E |
| \#8-2 | H1 | p56 | $\checkmark$ | 3.089 | 8.957 | 10 | 0 |  |
| \#8-3 | H1 | p56 | $\checkmark$ | 2.124 | 23.46 | 9 | 1 | D |
| \#9-1 | H1-1 | p62 | $\checkmark$ | 2.954 | 8.729 | 10 | 0 | E |
| \#9-1 | H1-2 | p62 | $\checkmark$ | 2.367 | 13.34 |  |  | N.D. |
| \#7-1 | 585A1 | p24 | $\checkmark$ | 2.627 | 5.63 | 9 | 1 | D |
| \#7-2 | 585A1 | p24 | $\checkmark$ | 4.3 | 17.33 | 9 | 1 | D |
| \#7-3 | 585A1 | p24 | $\checkmark$ | 3.623 | 5.309 | 6 | 4 | B |
| \#8-1 | 585A1 | p25 | $\checkmark$ | 4.18 | 9.747 | 10 | 0 | E |
| \#8-2 | 585A1 | p25 | $\checkmark$ | 2.07 | 6.665 | 10 | 0 | E |
| \#8-3 | 585A1 | p25 | $\checkmark$ | 2.562 | 6.714 | 10 | 0 | E |
| \#7-1 | 414C2 | p23 | $\checkmark$ | 1.03 | 2.919 | 10 | 0 |  |
| \#7-2 | 414C2 | p23 | $\checkmark$ | 1.21 | 2.441 | 10 | 0 | E |
| \#7-3 | 414C2 | p23 | $\checkmark$ | 1.229 | 2.498 |  |  | N.D. |
| \#2-1 | 253G1 | p25 | $\checkmark$ | 0.8773 | 3.505 | 10 | 0 |  |
| \#2-2 | 253G1 | p25 | $\checkmark$ | 0.9547 | 1.308 | 10 | 0 | E |
| \#2-3 | 253G1 | p25 | $\checkmark$ | 3.066 | 4.442 | 10 | 0 | E |
| \#8-1 | 610A2 | p25 | $\checkmark$ | 1.47 | 3.751 | 10 | 0 | E |
| \#8-2 | 610A2 | p25 | $\checkmark$ | 1.963 | 3.279 | 10 | 0 | E |
| \#8-3 | 610A2 | p25 | $\checkmark$ | 2.143 | 3.452 | 10 | 0 | E |
| \#7-1 | 454E2 | p23 | $\checkmark$ | 3.126 | N.D. | 9 | 1 | D |
| \#7-2 | 454E2 | p23 | $\checkmark$ | 1.241 | 4.443 | 1 | 9 | A |
| \#7-3 | 454E2 | p23 | $\checkmark$ | 1.051 | 2.35 | 1 | 9 | A |
| \#3-1 | 201B6 | p26 | $\checkmark$ | 0.7227 | 1.992 | 10 | 0 | E |
| \#3-2 | 201B6 | p26 | $\checkmark$ | 0.5638 | 1.034 | 10 | 0 |  |
| \#3-3 | 201B6 | p26 | $\checkmark$ | 0.3843 | Died |  |  | N.D. |
| \#8-1 | medium |  |  | 0 | 0 |  |  |  |
| \#8-2 | medium |  |  | 0 | 0 |  |  |  |
| \#8-3 | medium |  |  | 0 | 0 |  |  |  |
| \#9-1 | PBS |  |  | 0 | 0 |  |  |  |




