An inactivating mutation in the histone deacetylase SIRT6 causes human perinatal lethality

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It has been well established that histone and DNA modifications are critical to maintaining the equilibrium between pluripotency and differentiation during early embryogenesis. Mutations in key regulators of DNA methylation have shown that the balance between gene regulation and function is critical during neural development in early years of life. However, there have been no identified cases linking epigenetic regulators to aberrant human development and fetal demise. Here, we demonstrate that a homozygous inactivating mutation in the histone deacetylase SIRT6 results in severe congenital anomalies and perinatal lethality in four affected fetuses. In vitro, the amino acid change at Asp63 to a histidine results in virtually complete loss of H3K9 deacetylase and demyristoylase functions. Functionally, SIRT6 D63H mouse embryonic stem cells (mESCs) fail to repress pluripotent gene expression, direct targets of SIRT6, and exhibit an even more severe phenotype than Sirt6-deficient ESCs when differentiated into embryoid bodies (EBs). When terminally differentiated toward cardiomyocyte lineage, D63H mutant mESCs maintain expression of pluripotent genes and fail to form functional cardiomyocyte foci. Lastly, human induced pluripotent stem cells (iPSCs) derived from D63H homozygous fetuses fail to differentiate into EBs, functional cardiomyocytes, and neural progenitor cells due to a failure to repress pluripotent genes. Altogether, our study described a germline mutation in SIRT6 as a cause for fetal demise, defining SIRT6 as a key factor in human development and identifying the first mutation in a chromatin factor behind a human syndrome of perinatal lethality.

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during early embryogenesis and fetal development [Naifee et al. 2008; Chen and Dent 2014]. Inactivating mutations in DNA methyltransferases [DNMTs] such as DNMT3a and MECP2 [Rett syndrome] are associated with severe congenital immunological, craniofacial, and neurodevelopmental disorders [Naifee et al. 2008]. Loss-of-function mutations in the chromatin regulator histone methyltransferase NSD1 result in Sotos syndrome, a disease characterized by physical overgrowth and macrocephaly [Douglas et al. 2003; Rayasam et al. 2003]. Inherited mutations in the class I histone deacetylase HDAC8 contribute to the pathologies observed in Cornelia de Lange syndrome, with the majority of children living into adulthood [Kaiser et al. 2014; Feng et al. 2015]. However, whether mutations in chromatin regulators contribute to embryonic or fetal demise in utero remains unknown.

The NAD+-dependent histone deacetylase SIRT6 is a chromatin-associated protein that targets H3K9ac and H3K56ac and is known to play roles in genomic stability, glucose homeostasis, and tumor suppression [Mostoslavsky et al. 2006; Michishita et al. 2009; Yang et al. 2009; Zhong et al. 2010; Sebastián et al. 2010; Toiber et al. 2013; Kugel et al. 2015, 2016]. Sirt6 knockout mice from the 129SvJ background are viable and born at Mendelian ratios; however, they are much smaller than wild-type littermates [Mostoslavsky et al. 2006]. At 2–3 wk of age, Sirt6 knockout mice demonstrate an overall failure to thrive, accompanied by loss of subcutaneous fat, osteopenia, and profound hypoglycemia, which eventually leads to their death at 4 wk of age [Mostoslavsky et al. 2006]. Interestingly, Sirt6 knockout mice in the C57/B6 background display a strain-dependent embryonic-lethal phenotype [J-P Etchegaray and R Mostoslavsky, unpubl.], suggesting a role for SIRT6 during embryonic development. Indeed, our previous studies have shown that SIRT6 directly represses expression of core pluripotent genes, which in turn controls embryonic stem cell (ESC) differentiation through Tet-mediated production of 5-hydroxymethylcytosine (5hmC) specifically on neuroectoderm genes [Etchegaray et al. 2015]. However, the role that SIRT6 plays in human embryonic and perinatal development is still unclear.

In this study, we identify a family presenting with four cases of perinatal loss consistent with a number of severe neurodevelopmental and cardiac anomalies. Through whole-exome sequencing (WES) and Sanger sequencing, we detected a single homozygous mutation in SIRT6 [c.187G>C; p.[Asp63His]] among family members in a manner consistent with an autosomal recessive trait (Fig. 1D). Both parents and one grandparent were tested and confirmed the presence of this variant in the four fetuses [Fig. 1C,D]. Both parents and one grandparent were tested and found to be heterozygous carriers [Fig. 1D]. The fifth pregnancy presented with a healthy neonate identified as a heterozygous carrier for SIRT6 [c.187G>C; p.[Asp63His]] [Fig. 1D]. The variant segregated among family members in a manner consistent with an autosomal recessive trait [Fig. 1D].

SIRT6 D63H mutation results in loss of deacetylase and demyristoylase activity

We first investigated the consequences of the Asp63His [D63H] mutation on SIRT6 protein localization and function. We generated the D63H SIRT6 mutation using site-directed mutagenesis in a GFP-tagged construct. Upon fluorescence imaging, D63H SIRT6 demonstrated nuclear localization similar to wild-type SIRT6 [Fig. 2A]. To gain further insight into the functional significance of the D63H mutation, we analyzed in silico the coocystal structure of SIRT6 bound to ADP-ribose and the H3K9 myristoylated peptide [Protein Data Bank [PDB] 3ZG6] [Fig. 2B; Jiang et al. 2013; Kugel et al. 2015]. Asp63 is located in the NAD+-binding pocket, forming hydrogen bonds with neighboring amino acids and thus providing structure to the NAD+-binding loop [Jiang et al. 2013]. Similar
to the cancer-associated Asp63 mutation to tyrosine (D63Y) (Kugel et al. 2015), the D63H mutation on SIRT6 should hinder NAD+ binding and catalysis (Fig. 2B). In vitro deacetylase assays using purified recombinant wild-type, D63Y, and D63H SIRT6 demonstrated that the mutations almost completely abolished the enzymatic activity of SIRT6 toward a H3K9ac peptide (Fig. 2C). Next, we wanted to determine whether SIRT6 D63H retains the previously reported demyristoylase activity (Jiang et al. 2013). Consistently, D63Y and D63H recombinant SIRT6 mutants displayed decreased demyristoylase activity when compared with wild-type SIRT6 (Fig. 2D). To test whether D63H mutation decreases SIRT6 affinity for NAD+, we measured steady-state rates of demyristoylation with increasing concentrations of NAD+. Steady-state kinetics revealed that the $k_{cat}/K_m$ value for D63H mutant is 688 times less than wild-type SIRT6 (Fig. 2E). The $K_m$ for NAD+ for both D63Y and D63H is increased 6.8-fold, suggesting a decreased affinity for NAD+ (Fig. 2E). Last, we performed site-directed mutagenesis and overexpressed doxycycline-inducible SIRT6 constructs in SIRT6 knockout mouse embryonic fibroblasts (MEFs) to determine whether the SIRT6 D63H mutant is able to deacetylate H3K9 and H3K56 in vivo. Notably, the D63H mutant was unable to deacetylate its targets, as clearly seen even on bulk chromatin (Fig. 2F). Together, these results indicate that a mutation of Asp63 to histidine inhibits the affinity of SIRT6 for NAD+ and therefore significantly reduces its catalytic activity both in vitro and on bulk chromatin in vivo.

**Figure 1.** WES identifies the homozygous c.187G > C; p.(Asp63His) SIRT6 mutation in four fetuses with severe congenital defects and perinatal lethality. (A) Description of clinical findings of fetuses with SIRT6 mutation. (B) Representative ultrasound images of abnormal cerebella, head circumferences, and frontal bossing observed in four affected fetuses. (C) Sanger sequencing confirms the SIRT6 c.187G > C missense mutation in affected fetal amniocytes corresponding to SIRT6 p.D63H. Protein sequence alignment of various species for the area around the D63 amino acid showing strong evolutionary conservation at this position. (D) Pedigree showing consanguinity and cosegregation of the SIRT6 p.D63H mutation. (na) Genotype not available; (black symbols) affected; (white symbols) unaffected; (?) unknown phenotype; (dots) carriers of mutation.

**Table 1.**

| General | Brain | Craniofacial | Heart |
|---------|-------|--------------|-------|
| Early Intratetral Growth Retardation | Cerebellar hypoplasia | Microcephaly | Septal Defects (VSD, ASD) |
| Generalized Edema | Microcephaly | Frontal Bossing | Cardiomegaly (dilated RA+LA) |
| Sex reversal (in males) | Brachycephaly | Valve Defects (Aortic valve stenosis, Tricuspid Valve insufficiency) |
| | | Absent Nasal Bone | |
| | | Short Palpebral Fissures | |
| | | Hypertelorism | |

**Figure 2.**

SIRT6 D63H mutant mESCs fail to form EBs and retain pluripotent gene expression

An overall down-regulation of the core pluripotent genes is essential for proper ESC differentiation (Young et al. 2011). We found previously that SIRT6 acts as a histone deacetylase to directly silence pluripotent genes, and SIRT6-deficient EB cells fail to repress pluripotent genes, significantly altering normal development of the three germ layers during differentiation (Etchegaray et al. 2015). To functionally test whether the SIRT6 D63H mutation is causing an aberrant developmental phenotype, we first measured the expression of pluripotent genes in
SIRT6 knockout mouse ESCs (mESCs) stably transduced with doxycycline-inducible SIRT6 wild-type and D63H constructs. Interestingly, quantitative RT–PCR (qRT–PCR) results revealed that D63H mESCs express significantly higher levels of the core pluripotent genes Oct4, Sox2, and Nanog compared with SIRT6 knockout and wild-type reconstituted mESCs (Fig. 3A). Additionally, we observed an increased presence of the pluripotent factor Oct4 on bulk chromatin in D63H mutant mESCs when compared with SIRT6 knockout mESCs (Fig. 3B). This result was accompanied by a failure to deacetylate H3K9, a SIRT6 target (Fig. 3B).

SIRT6 knockout ESCs cultured to form EBs are significantly smaller than their wild-type counterparts [Etchegaray et al. 2015]. Therefore, we tested the ability of D63H reconstituted SIRT6 knockout mESCs to differentiate in vitro into EBs. As demonstrated previously, expression of wild-type SIRT6 in SIRT6 knockout mESCs is sufficient to rescue the SIRT6 knockout EB phenotype [Etchegaray et al. 2015, Fig. 3C]. Interestingly, we observed that stable expression of SIRT6 D63H leads to an even more severe differentiation defect with a significantly smaller and decreased number of EBs compared with SIRT6 knockout ESCs [Fig. 3C]. This phenotype was associated with a significant derepression of pluripotent genes in the context of the mutant SIRT6 [Fig. 3D], increased binding of pluripotent factors to chromatin, and a failure to deacetylate H3K9 [Fig. 3E]. Such changes were significantly worse than in SIRT6 knockout cells, suggesting that the presence of the mutant protein could be biologically worse than the absence of the enzyme [see the Discussion below]. Interestingly, shRNA-mediated knockdown of Oct4 and SOX2 in SIRT6 knockout mESCs reconstituted with D63H SIRT6 [Supplemental Fig. S2B,C] resulted in a partial rescue of the EB defect observed in control shRNA-treated cells [Supplemental Fig. S2A]. Together, the data suggest that the D63H mutant SIRT6 severely impairs mESC differentiation due to a failure to suppress pluripotent genes.

The pluripotent factor Sox2 is responsible for promoting neuroectoderm differentiation [Wang et al. 2012]. Previously, our laboratory had demonstrated that SIRT6 knockout mESCs display a differentiation phenotype skewed toward the neuroectoderm lineage compared with wild-type counterparts [Etchegaray et al. 2015]. D63H EBs showed significantly elevated expression of SOX2 consistent with increases in neuroectoderm markers as measured by increased Nestin, Pax6, and Hoxa7.
compared with SIRT6 knockout EBs [Supplemental Fig. S1B], without compromising mesoderm and endoderm markers [Supplemental Fig. S1C]. Interestingly, combined knockdown of Oct4 and Sox2 was able to reverse the increase in neuroectoderm markers in SIRT6 D63H EBs [Supplemental Fig. S2D], together indicating that sustained SOX2 expression skewes D63H EBs further toward the neuroectoderm lineage.

SIRT6 D63H mutant mESCs fail to differentiate into functional cardiomyocyte foci

SIRT6 in the adult heart has been shown to protect against concentric cardiac hypertrophy [Sundaresan et al. 2012]. However, the role of SIRT6 in cardiac development and morphogenesis still remains to be investigated. Analysis of SIRT6 knockout mESC-derived cardiomyocytes revealed decreased expression of cardiomyogenesis markers, a phenotype associated with irregular beating patterns and a decreased number of beating foci when compared with wild-type SIRT6 cardiomyocytes (data not shown). Since a number of cardiac anomalies were identified in D63H homozygote fetuses [Fig. 1A], we tested the capacity of D63H mutant mESCs to differentiate into functional cardiomyocytes. To address this question, we used an EB-directed differentiation protocol in mESCs to aid in mesoderm layer specification (Lee et al. 2011). From day 0 to day 4, mESCs were cultured under hanging drop conditions as EBs. Subsequently, from day 4 to day 25, cells were placed in EGM-2 [endothelial cell type] medium containing growth factors essential for normal cardiac development [Fig. 4A, top panel; Lee et al. 2011]. On day 25, wild-type SIRT6 differentiated cardiomyocytes morphologically resembled cardiomyocyte foci and exhibited regular spontaneous contraction patterns [Supplemental Fig. S3A, Supplemental Movies S1–S4]. SIRT6 knockout ESCs, on the other hand, formed less differentiated cardiomyocyte foci, a phenotype rescued with the re-
expression of wild-type SIRT6 but not with expression of the mutant D63H protein [Supplemental Fig. S3A; Supplemental Movies S1–S4]. This severe phenotype was accompanied by a dramatic derepression of Oct4 and increased acetylation of the SIRT6 target H3K56 when the mutant protein was expressed [Fig. 4A, bottom panel]. Immunofluorescence staining for the cardiac differentiation marker cardiac troponin T (cTnT) demonstrated intense staining in SIRT6 wild-type cardiomyocytes when compared with SIRT6 knockout, which exhibited weak staining and aberrant morphology [Fig. 4B]. cTnT staining was rescued when wild-type SIRT6 was reconstituted in SIRT6 knockout cardiomyocytes [Fig. 4B]. However, D63H-expressing mouse cardiomyocytes failed to stain for cTnT, morphologically remaining as EBs [Fig. 4B]. These data indicate that expression of SIRT6 D63H suppresses cardiomyocyte differentiation, likely through failure to suppress Oct4 expression.

Next, we sought to characterize molecular signatures of SIRT6 D63H mESC-derived cardiomyocytes. SIRT6 wild-type and SIRT6 knockout cardiomyocytes reconstituted with wild-type SIRT6 exhibit expression of cardiac sarcomeric genes αMHC, βMHC, and cTnT, indicating fully differentiated foci [Fig. 4C, Lian et al. 2012]. Interestingly, SIRT6 D63H-expressing cardiomyocytes expressed significantly lower levels of αMHC and βMHC when compared with SIRT6 knockout cardiomyocytes [Fig. 4C], consistent with a failure to differentiate. The transcription factor TBX5 is an important regulator of cardiomyocyte development, and decreased levels are associated with ventral septal defects (VSDs) and atrial septal defects (ASDs) (Horb and Thomsen 1999; Posch et al. 2010). Tbx5 expression was decreased in SIRT6 knockout cardiomyocytes compared with wild-type counterparts [Fig. 4C]. However, Tbx5 expression was further compromised in D63H-expressing cardiomyocytes compared with wild-type counterparts [Fig. 4C]. These data indicate that expression of SIRT6 D63H suppresses cardiomyocyte differentiation, likely through failure to suppress Oct4 expression.

Figure 4. D63H mutant mESCs fail to differentiate into functional cardiomyocyte foci and retain pluripotent gene expression. (A, top panel) Schematic showing the differentiation from mESCs into cardiomyocytes. (Bottom panel) Western blot analysis on bulk chromatin for Oct4 and Ac-H3K56 in wild-type versus SIRT6 knockout ESC-derived cardiomyocytes with doxycycline-inducible overexpression of wild-type SIRT6 and SIRT6 D63H. (B) Immunofluorescence staining for cTnT in mESC-derived cardiomyocytes (20×). (C) Cardiac marker expression in wild-type versus SIRT6 knockout differentiated cardiomyocytes with doxycycline-inducible overexpression of wild-type SIRT6 and SIRT6 D63H assessed by qRT–PCR analysis. Data are expressed relative to knockout cardiomyocytes. (D) Spontaneous contractile foci were quantified on day 21 as the average number of foci from five clones.
significantly reduced FBN1 levels when compared with SIRT6 knockout cells [Fig. 4C], in line with the ventricular and cardiac valve defects observed in the D63H homozygous fetuses [Fig. 4C]. Thus, molecular characterization of SIRT6 D63H mESCs indicates a failure to differentiate and express cardiac morphogenesis markers, likely due to a failure to repress pluripotent gene expression.

To test the functional significance of the D63H mutation on cardiomyocyte differentiation, we quantified the number of spontaneously contractile foci in SIRT6 D63H cardiomyocytes. SIRT6 D63H cardiomyocytes formed significantly less spontaneous contractile foci, exhibiting irregular beating patterns [Supplemental Fig. S2B] compared with SIRT6 wild-type and wild-type reconstituted SIRT6 knockout cardiomyocytes [Fig. 4D]. This was consistent with gene expression data demonstrating significantly decreased levels of the α1-adrenergic receptor a1c (Fig. 4C; Jensen et al. 2014). Collectively, SIRT6 D63H mESCs exhibit a robust failure to differentiate into cardiomyocyte foci and display irregular beating patterns.

D63H homozygous iPSCs fail to differentiate into EBs and retain pluripotent gene expression

Mouse-derived ESCs expressing SIRT6 D63H displayed aberrant pluripotent gene expression and failure to differentiate into EBs and functional cardiomyocytes [Figs. 3, 4]. To determine whether the endogenous expression of the human homozygous D63H mutation is sufficient to cause differentiation defects, we generated iPSCs from amniocytes isolated from two D63H homozygous fetuses (one female gestational age 20 wk and one male gestational age 27 wk) as well as peripheral blood mononuclear cells (PBMCs) from the heterozygous mother and viable sibling [Supplemental Fig. S4A]. Immunofluorescence staining for pluripotency surface markers SSEA-4 and TRA-1-81 confirmed the undifferentiated state of multiple generated iPSC clones [Supplemental Fig. S4A]. Interestingly, undifferentiated iPSCs derived from homozygous amniocytes did not exclusively retain pluripotent factors on bulk chromatin [Supplemental Fig. S4B] or display increased pluripotent gene expression [Supplemental Fig. S4C]. To test whether the presence of the homozygous D63H mutation prevents these cells from undergoing proper differentiation, we enabled spontaneous differentiation by modulating Wnt signaling [Fig. 6A, top panel; Lian et al. 2012, 2013; Aguilar et al. 2015; Rajasingh et al. 2015]. Day 3 after Wnt activation resulted in retained expression of pluripotent genes and significantly lower levels of cTnT in the D63H homozygous cells [Fig. 6B, top panel]. Further differentiation (day 7) exacerbated the phenotype, where both βMHC and cTnT were severely reduced in the mutant iPSC cardiomyocytes [Fig. 6B, bottom panel], consistent with delayed differentiation of the overall population as measured by cTnT-FITC+ staining [Supplemental Fig. S5A]. Notably, Wnt-mediated differentiation of D63H homozygous iPSC cardiomyocytes resulted in a failure to fully differentiate, and the cells crashed prior to completion on day 16 with only a few cells remaining, and, from those, <50% of cells stained positive for cTnT-FITC+ [Fig. 6A, bottom panel]. Genome-wide RNA sequencing (RNA-seq) analysis confirmed the lack of suppression of pluripotent genes [Supplemental Fig. S4C]. In addition, gene ontology analysis identified an altered gene expression signature with defects in HOX genes, Wnt signaling, aorta and cardiac morphogenesis, and cardiomyopathy [Supplemental Fig. S7C]. Interestingly, many of these aberrantly expressed genes are critical for cardiac development [TBX20, NKKX2-3, MYOC, NKKX2-5, GATA4, FBN1, Wnt1, and Wnt3A], with deregulation of these genes contributing to cardiomyopathy, proper valve formation, and cardiac septation [Supplemental Fig. S7D; Brand et al. 2003; Perrino and Rockman 2006; Pepe et al. 2014].

D63H homozygous iPSCs display delayed NPC differentiation

As described above, EBs derived from SIRT6 D63H homozygous iPSCs failed to express the early CNS intermediate filament protein Nestin [Supplemental Fig. S4D]. In order to directly test whether reprogrammed SIRT6 D63H
homozygous iPSCs properly differentiate into CNS precursors, we performed directed monolayer differentiation into NPCs (Fig. 6C). By day 9 of differentiation, heterozygous iPSCs readily expressed Nestin while suppressing Oct4, SOX2, and Nanog expression (Fig. 6C,D, top panel). However, iPSC-derived NPCs generated from SIRT6 D63H homozygous fetuses, while expressing Nestin, retained pluripotent gene expression (Fig. 6C,D, top panel).
Furthermore, even late stage (day 15) D63H homozygous iPSCs–NPCs retained Oct4 and Nanog expression (Fig. 6D, bottom panel). This was confirmed by RNA-seq analysis (Supplemental Fig. S4C), and, consistent with the severe neural phenotypes observed in the fetuses, we also found aberrant expression of HOX and POU genes, limb morphogenesis, Wnt signaling, and neural differentiation pathways (Supplemental Fig. S7A). Interestingly, many of these aberrantly expressed genes are critical for CNS development [POU4F1, FOXG1, SHH, NEUROD1, GBX2, MSX1, and SOX10], and deregulation of these genes has been associated with microcephaly and cerebellar hypoplasia (Supplemental Fig. S7B; Nanni et al. 1999; Ramos et al. 2004; Kortüm et al. 2011; Szulc et al. 2013).
Together, directed differentiation of D63H homozygous iPSCs resulted in delayed NPC generation characterized by sustained pluripotent gene expression, which could explain the neurodevelopmental defects observed in the fetuses.

Discussion

As many as one-third of cases of late stage fetal loss remain unexplained, and idiopathic stillbirth is now the most common contributor to human perinatal mortality (Robson and Leader 2010; Bukowski et al. 2014; Lamont et al. 2015). Here we identify, for the first time, a family with four cases of perinatal lethality carrying a naturally occurring homozygous loss-of-function mutation in the histone deacetylase SIRT6. This mutation resulted in severe brain and heart developmental defects. When tested in culture, we observed a failure of mESCs and iPSCs carrying the homozygous mutant D63H SIRT6 to undergo proper differentiation (when challenged to differentiate into EBs, cardiomyocytes, and NPCs) due to sustained pluripotent gene expression. Importantly, ectopic expression of the human SIRT6 D63H mutant in SIRT6 knockout mESCs recapitulates the severe differentiation phenotype observed in D63H homozygous human iPSCs, thereby confirming a conserved role for SIRT6 in mouse and human pluripotency and development. In previous studies, we found that SIRT6 knockout mESCs fail to properly differentiate due to an inability to repress pluripotent genes [Etchegaray et al. 2015]. Interestingly, the presence of the mutant D63H SIRT6 presents a more severe ESC differentiation phenotype than the one observed in the absence of SIRT6. It is plausible that the presence of the enzymatically inactive SIRT6 D63H on chromatin may impede compensatory mechanisms from other histone deacetylases from removing H3K9 and H3K56 marks. Alternatively, the SIRT6 D63H protein may exhibit a slower turnover than the wild-type protein, altering the balance between protein synthesis and degradation and disrupting histone acetylation in a more severe manner than when SIRT6 is completely absent. Of note, we observed a severe defect in the ability of the mutant iPSCs to differentiate into human EBs in vitro, yet D63H homozygous fetuses are miscarried relatively late in gestation. It is possible that in vivo, there is some redundancy with other chromatin factors that manage to keep early embryos differentiating and viable, and the phenotype manifests only in later developmental stages. Such a mechanism may be lost in the EB in vitro setting, where the differentiation to the three germ layers is driven at a much faster pace. Another explanation is that maternal contribution from the wild-type copy of SIRT6 may be weakening the developmental consequences of the D63H homozygous mutant fetuses in vivo. In this case, the embryos exhibit a milder defect when compared with the in vitro EBs due to compensation from the maternal wild-type SIRT6. For instance, Dnmt1o (Howell et al. 2001) and Dnmt3l (Bourc’his et al. 2001) are other epigenetic modifiers that exhibit similar maternal contribution in vertebrates, demonstrating late developmental phenotypes in vivo.

A number of mutations in DNA methylation and chromatin factors have been associated with neurodevelopmental syndromes with affected individuals living into adulthood [Douglas et al. 2003; Nafee et al. 2008; Jakovecski and Akbarian 2012; Kaiser et al. 2014; Feng et al. 2015; Vallianatos and Iwase 2015]. Although severe neurological phenotypes are observed in patients with chromatin-associated mutations in NSD1 (Sotos syndrome), EHMT1 (Kleefstra syndrome), and KDM5C (X-linked mental retardation), these mutations rarely result in perinatal lethality, thus pointing to possible compensation for the enzymatic activities of these histone modifiers during development [Douglas et al. 2003; Iwase et al. 2007; Nafee et al. 2008]. For example, it has been suggested that the mild phenotype observed in KDM5C [JARID1C]-deficient mice is due to compensation by other Jumonji domain-containing histone demethylases [Iwase et al. 2007; Klose et al. 2007; Vallianatos and Iwase 2015]. In contrast, our data indicate a key role for SIRT6 as a critical enzyme required for proper cell lineage differentiation and organogenesis throughout fetal development. A loss-of-function mutation results in a completely penetrant perinatal lethal phenotype, likely due to an inability to repress early embryonic genes. The main features observed in these fetuses were severe heart and neurological defects, a phenotype that we mimicked in vitro when forcing mutant iPSCs derived from amniocytes to differentiate into neural progenitors and cardiomyocytes. Previous studies have shown critical roles for SIRT6 in modulating glucose homeostasis, with SIRT6-deficient mice in certain strains dying early after birth due to severe hypoglycemia [Mostoslavsky et al. 2006; Zhong et al. 2010]. Unfortunately, we were unable to measure glucose in these fetuses before their demise; therefore, it remains to be determined whether a glucose imbalance also contributed to the lethal phenotype. Although we detected an increase in LDHA and PDK1 at the protein level in the context of the homozygous SIRT6 mutation [iPSCs], one fetus (fetus #3) failed to derepress these genes to a lesser extent than the other fetus [fetus #1] [Supplemental Fig. S6A,B]. Notably, the glucose transporter GLUT1 (SLC2A1) was actually down-regulated in D63H homozygous fetus-derived EBs [Supplemental Fig. S6B], suggesting some compensatory mechanism. Despite this increase in LDHA and PDK1 expression, treatment with the glycolytic inhibitor dichloroacetate (DCA) did not rescue the severe EB phenotype [Supplemental Fig. S6C], as we observed for previous SIRT6-related phenotypes [Zhong et al. 2010; Sebastián et al. 2012]. Additionally, genome-wide RNA-seq expression analysis demonstrated that iPSCs, NPCs, and cardiomyocytes derived from D63H homozygous fetuses do not contain elevated levels of glycolytic genes and ribosome biogenesis genes [Supplemental Fig. S6D]. These results confirm that SIRT6-dependent metabolic changes cannot account for the developmental defects seen in the D63H homozygous iPSCs, but rather the inability of SIRT6 D63H fetuses to suppress pluripotent gene expression and cell death.
expression is likely the dominant molecular mechanism contributing to perinatal loss.

In separate studies, SIRT6 has been shown to prevent genomic instability and maintain pericentromeric silencing, inhibiting cellular senescence [Michishita et al. 2009, Mao et al. 2011; Toiber et al. 2013; Tasselli et al. 2016; Van Meter et al. 2016]. However, cytogenetic analysis of fetal amniocytes indicated no evidence of copy number variations or structural aneuploidy [data not shown]. Although D63H-overexpressing mESC-derived EBs fail to reduce DNA damage signaling in the context of the SIRT6 knock-out [Supplemental Fig. S6E], the D63H homozygous patient-derived iPSCs and EBs do not exhibit increased cleaved caspase and γ-H2AX [both signs of DNA damage] compared with the phenotypically normal heterozygous iPSCs [Supplemental Fig. S6F]. This suggests that these cells somehow are able to compensate for the absence of SIRT6 function in order to sustain genomic stability, further confirming that DNA damage is not causal in the developmental defects observed in D63H homozygous fetuses.

Although SIRT6 has been shown to protect against cardiac hypertrophy in the adult (Sundaresan et al. 2012), our studies identify a novel role for this chromatin modifier in regulating the balance between pluripotency and cardiac lineage specification during development. Contrary to results seen in adult hearts, SIRT6 D63H-expressing cells exhibit inconsistent IGF signaling, with increased p-AKT and decreased p-ERK levels [data not shown]. Additionally, iPSC-derived cardiomyocytes from D63H homozygous fetuses fail to exhibit elevated IGF signaling compared with cardiomyocytes derived from the heterozygous mother, as measured by RNA-seq [Supplemental Fig. S5B], suggesting that, unlike the adult, IGF signaling may not drive the phenotypes that we observed during development. However, RNA-seq analysis demonstrates aberrant expression of important downstream mediators of cardiac specification in the D63H homozygous fetuses (for example, GATA4 and Wnt3a) [Supplemental Fig. S7D]. Future studies will investigate the functional role of these genes in cardiac differentiation in the context of the homozygous SIRT6 D63H mutation. Whether they are direct targets of SIRT6 or function downstream from the pluripotent genes or the Tet enzymes [Etchegaray et al. 2015] remains to be determined.

Patients with de novo mutations in H3K4me regulators (MLL2, KDM6A, and CHD7) display diverse congenital heart defects commonly accompanied by neurodevelopmental and growth abnormalities [Zaidi et al. 2013]. H3K4me2 is an activating epigenetic mark associated with both promoters and enhancers involved in transcriptional activation of key developmental genes [Iwase et al. 2007; Klose et al. 2007; Vallianatos and Iwase 2015]. Previously, our laboratory demonstrated that SIRT6 knock-out mESCs contain elevated Tet1 levels and enriched 5hmC within exons encoding neuroectoderm genes [Etchegaray et al. 2015]. Importantly, these regions were also enriched for H3K4me2, and the up-regulation of coregulated genes was rescued upon Tet inhibition [Etchegaray et al. 2015]. Interestingly, D63H-expressing mouse cardiomyocytes retained Tet2 on chromatin [data not shown]. In the future, it will be interesting to explore the link between SIRT6 deacetylation and H3K4me during cardiac differentiation. Whether such a link represents an example of coordination of two related marks [histone code] or whether SIRT6 may directly deacetylate nonchromatin targets, including lysine methyltransferases and demethylases, remains to be explored.

In summary, our work identified a novel human syndrome defined by a naturally occurring loss-of-function mutation in the chromatin enzyme SIRT6, associated with a fully penetrant human perinatal lethal phenotype. We demonstrated a failure to suppress pluripotent gene expression in homozygous SIRT6 D63H iPSCs, resulting in an inability of the cells to undergo proper differentiation into EBs, cardiomyocytes, and NPCs. Together, our data implicate SIRT6 chromatin remodeling activity as a critical modulator of human development, regulating the transition from pluripotency to differentiated states.

Materials and methods

Patient amniocyte/PBMC collection and sequencing

Peripheral blood was collected from the mother, father, and grandmother. Amniotic fluid was collected from amniocentesis during all five pregnancies. Written consent was obtained in accordance with guidelines of the Academic Medical Centre [AMC], Netherlands Institutional Review Board. DNA was extracted from blood or amniotic fluid cells using standard methods. WES was performed on child #4 and both parents. Exonic targets were enriched with the SeqCap EZ human exome library version 3.0 kit [NimbleGen]. The resulting libraries were sequenced on a HiSeq 2500 version 4 chemistry (Illumina) according to the manufacturer’s recommendations for paired-end 125-base-pair [bp] reads. Alignment of sequence reads to the human reference genome [hg19] was done using BWAMEM 0.7.5 [http://www.bio-bwa.sourceforge.net], and variants were called using the GATK3.3 software package [http://www.broadinstitute.org/gatk]. Annotation and filtering of variants was done using Cartagenia Bench Lab NGS [Agilent]. Variants with a frequency of >1% or homozygously present in public [ESP, ExAC, dbSNP, and 1KG] and/or in-house databases were excluded. Evaluation of the homozygous variants revealed possible causal variants in five genes [ARHgef17, EGLAM, RBM4N, RAF1, and SIRT6]. Subsequent analysis of the three affected siblings using Sanger sequencing showed that only the SIRT6 variant [c.187G>C p.[Asp63His]] segregated with the disorder.

Expression and purification of recombinant human SIRT6

His-tagged wild-type and mutant SIRT6 proteins were overexpressed in the BL21(DE3) Escherichia coli strain as described previously [Feldman et al. 2015]. Cells were harvested by centrifugation and stored at −80°C. SIRT6 wild-type and mutant proteins were purified by nickel affinity resin chromatography as reported previously [Feldman et al. 2015]. Protein concentrations were determined by the Bradford reagent assay.

HPLC deacetylation assay

Peptides corresponding to residues 4–17 of histone H3 [acetyl: AcKQTARKAcSTGGKAPRWW-NH2, and myristoyl: Ac-KQT ARKmyrSTGGKAPRWW-NH2] were synthesized as described.
previously (Feldman et al. 2013). Decacylation reactions were analy-
zed by reversed-phase high-performance liquid chromatogra-
phy on a Kinetex C18 column (100 Å, 100 mm × 4.6 mm, 2.6
µm, Phenomenex) by monitoring the formation of the decacylated
product at 214 nm (Feldman et al. 2013).

Steady-state kinetics analyses
Steady-state rates were measured by varying NAD⁺ [5–640 µM] in
the presence of 0.5 µM wild-type, D63Y, and D63H SIRT6 in
20 mM sodium phosphate (pH 7.5) at 37°C. Initial velocities
were determined, and data were fitted to the Michaelis-Menten
equation. Deacetylase and demyristoylase activities were mea-
sured in the presence of 50 mM corresponding substrate, 0.5
mM NAD⁺, and 2 µM wild-type or mutant SIRT6. Reactions
were quenched at <15% substrate depletion by addition of 2% TFA
final concentration.

Generation of human SIRT6 constructs and virus production
Generation of human eGFP and pRetroX-Tight-Pur SIRT6 con-
structs have been described previously (Kugel et al. 2015). Site-di-
rected mutagenesis of wild-type SIRT6 was performed using the
QuickChange Lightning site-directed mutagenesis kit (Strata
gene). pLVX-Tet-On was obtained from Clontech. Viral particles
containing the above-mentioned plasmids were synthesized us-
ing retroviral packaging plasmids pCL-ECO (Addgene) and
cMV-VSV-G (Addgene).

Subcellular localization of SIRT6 constructs
293T cells were transfected using Trans-IT 293 [Mirus] with empty
t vector, pEGFP-SIRT6, or the SIRT6 mutant constructs. Twenty-
four hours after transfection, cells were trypsinized and seeded
onto eight-well chamber slides and allowed to adhere overnight.
Cells were then fixed using 2% paraformaldehyde in 1% phos-
phate-buffered saline (PBS) solution and permeabilized with 0.1%
Triton X-100, and nuclei were stained using 4,6-diamidino-2-
phenylindole (DAPI). Images were taken using a fluorescent microscope.

Mouse-derived cell lines and tissue culture conditions
SIRT6 knockout primary MEFs were generated from 13.5-d-old
embryos as described (Mostoslavsky et al. 2006). Cells were cul-
tured at 37°C under 5% CO₂ in high-glucose Dulbecco’s modified
Eagle’s medium [DMEM] supplemented with 10% fetal bovine
serum (FBS), 1% 100 U/mL penicillin/streptomycin [Invitrogen],
2 mM L-glutamine, 0.1 mM NEAA, 1 mM sodium pyruvate, and
20 mM HEPES. 293T cells were cultured in high-glucose DMEM
supplemented with 10% FBS and 1% 100 U/mL penicillin/strep-
томycin [Invitrogen]. SIRT6 knockout MEFs were infected by in-
cubating with virus and 10 µg/mL polybrene. Forty-eight hours
later, cells were selected in 2.5 µg/mL puromycin on puromy-
cin-resistant γ-irradiated feeders. Clones were picked and
screened by qRT–PCR for knockdown of Oct4 and Sox2 expres-
sion. Prior to experiments, clones were removed from feeders
and grown on gelatin-coated plates.

Generation of Oct4 and Sox2 knockdown ESC lines
Oct4 and Sox2 shRNA constructs were used as described previ-
ously [Etchegary et al. 2015]. Briefly, ESCs were infected by in-
cubating with virus and 10 µg/mL polybrene. Forty-eight hours
later, cells were selected in 2.5 µg/mL puromycin on puromy-
cin-resistant γ-irradiated feeders. Clones were picked and
screened by qRT–PCR for knockdown of Oct4 and Sox2 expres-
sion. Prior to experiments, clones were removed from feeders
and grown on gelatin-coated plates.

Differentiation of mESCs into cardiomyocytes
SIRT6 wild-type and knockout ESCs were infected and treated
with doxycycline as described above. Differentiation toward the
cardiomyocyte lineage was carried out using the standard hang-
drop method with EBs collected on day 4 and plated onto ad-
herent plates in EGM-2 bullet kit medium [Lonza]. Spontaneous
contractile foci within differentiating EBs were detected as early
as day 15 and quantified as the average number of foci per five
clones. The medium was subsequently changed until day 21,
and cells were collected for bulk chromatin and RNA isolation
and fixed for immunofluorescence staining for CTnT.

Human iPSC generation and immunostaining
Frozen amniocytes were thawed and maintained on 0.1% gelatin-
coated six-well plates using amniocyte medium. The medium
contained Alpha-MEM [Invitrogen] supplemented with 10% FBS
[Hyclone], 1× GlutaMax [Invitrogen], 1× nonessential amino acids
[Invitrogen], and 0.1 mg/mL Primocin [InvivoGen]. Two days be-
tween transduction, amniocytes were plated onto 0.1% gelatin-
coated six-well plates to obtain 2 × 10⁵ and 5 × 10⁴ cells on the
day of transduction, which represents 50%–80% confluency. To trans-
duce the cells, CytoTune 2.0 Sendai reprogramming kit [Invitro-
gen] was thawed and prepared according to the manufacturer’s proto-
col. Amniocytes were infected with each virus at multiplici-
ty of infection [KOS:hc-Myc:hKlf4 = 5:5:3] using amniocyte me-
dium with 5 µg/mL polybrene [MilliporeSigma]. The viruses
were washed the following day, and the infected cells were cul-
tured for six more days by changing the amniocyte medium every
other day. At day 7 after transduction, the cells were passaged and
plated onto human ESC [hESC]-qualified Matrigel-coated [Cor-
ing] six-well plates at ∼1.0 × 10⁴ to 8.5 × 10⁴ cells per well. From

mESC cultures, formation, and immunostaining of EBs
mESCs derived from Sirt6 knockout and wild-type 129 mouse
strain were maintained on γ-irradiated MEFs in knockout
DMEM [Gibco] containing 15% ES-qualified FBS, 0.1 mM
each) nonessential amino acids, 2 mM L-glutamine, 0.1 mM β-
mercaptoethanol, and 50 U/mL penicillin/streptomycin [Invitro-
gen] and supplemented with leukemia-inhibiting factor. SIRT6
knockout and wild-type mESCs were infected by incubating
with virus and 10 µg/mL polybrene. Forty-eight hours later, cells
were selected in 2.5 µg/mL puromycin, and five clones were se-
picted for use in various experiments. For all experiments involv-
ing doxycycline-inducible SIRT6 knockout ESCs, cells were
-treated with 1 µg/mL doxycycline for 7 d prior to collection and
differentiation. For all experiments described, cells were trypsi-
nized and plated for 30 min on standard tissue culture dishes to
remove feeder cells before floating ESCs were collected and re-
plated on gelatin-coated dishes or wells before differentiation to-
ward EBs. ESCs were dissociated with trypsin [day 0] and cultured
at a density of 5000 cells per milliliter in Iscove’s modified Dul-
becco’s medium [IMDM] with 15% fetal calf serum [Atlanta Bio-
logicals], 2 mM L-glutamine [Gibco], 200 µg/mL transferrin
[Roche], 0.5 mM ascorbic acid [Sigma], and 4.5 × 10⁻⁴ M monothio-
glycerol [Sigma]. Differentiation was carried out using the stan-
dard hanging drop method, and EBs were collected from days 6
to 12. EB size was quantified using ImageJ 64 software as an aver-
age of five independent clones in a 2.5x field. EB number was
quantified as the number of EBs in a 2.5x field and was an average
of five independent clones.

Generation of Oct4 and Sox2 knockdown ESC lines
Oct4 and Sox2 shRNA constructs were used as described previ-
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cubating with virus and 10 µg/mL polybrene. Forty-eight hours
later, cells were selected in 2.5 µg/mL puromycin on puromy-
cin-resistant γ-irradiated feeders. Clones were picked and
screened by qRT–PCR for knockdown of Oct4 and Sox2 expres-
sion. Prior to experiments, clones were removed from feeders
and grown on gelatin-coated plates.

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Frozen amniocytes were thawed and maintained on 0.1% gelatin-
coated six-well plates using amniocyte medium. The medium
contained Alpha-MEM [Invitrogen] supplemented with 10% FBS
[Hyclone], 1× GlutaMax [Invitrogen], 1× nonessential amino acids
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tween transduction, amniocytes were plated onto 0.1% gelatin-
coated six-well plates to obtain 2 × 10⁵ and 5 × 10⁴ cells on the
day of transduction, which represents 50%–80% confluency. To trans-
duce the cells, CytoTune 2.0 Sendai reprogramming kit [Invitro-
gen] was thawed and prepared according to the manufacturer’s proto-
col. Amniocytes were infected with each virus at multiplici-
ty of infection [KOS:hc-Myc:hKlf4 = 5:5:3] using amniocyte me-
dium with 5 µg/mL polybrene [MilliporeSigma]. The viruses
were washed the following day, and the infected cells were cul-
tured for six more days by changing the amniocyte medium every
other day. At day 7 after transduction, the cells were passaged and
plated onto human ESC [hESC]-qualified Matrigel-coated [Cor-
ing] six-well plates at ∼1.0 × 10⁴ to 8.5 × 10⁴ cells per well. From
day 8 after transduction, the cells were fed with complete ReproTeSR (Stem Cell Technologies). The medium was changed daily. Once human iPSC colonies appeared, clonal human iPSCs were picked and expanded using mTeSR (Stem Cell Technologies).

Frozen PBMCs were thawed and expanded for 9 d using Expansum medium containing QBSF-60 stem cell medium [Quality Biological] supplemented with 50 µg/mL ascorbic acid (Millipore Sigma), 50 ng/mL SCF (R&D Systems), 10 ng/mL IL-3 (R&D Systems), 2 µg/mL EPO (R&D Systems), 40 ng/mL IGF-1 (R&D Systems), 1 µM dexamethasone (MilliporeSigma), and 0.1 mg/mL Primocin (InvivoGen). Expanded PBMCs were transduced with CytoTune 2.0 Sendai reprogramming kit (Invitrogen) by spinoculation at 2250 rpm for 90 min at room temperature. After 1 d of incubation, the viruses were washed, and the cells were plated onto hESC-qualified Matrigel-coated (Corning) six-well plates at 8.0 × 10^4 to 2 × 10^5 cells per well using Expansion medium. At day 3 after transduction, the infected cells started being fed with ReproTeSR (Stem Cell Technologies). Once human iPSC colonies appeared, clonal human iPSCs were picked and expanded using mTeSR (Stem Cell Technologies).

All generated human iPSCs were fixed with 4% paraformaldehyde/PBS and immunofluorescence-stained with mouse anti-human SSEA-4, mouse anti-human TRA-1-81, and mouse anti-human TRA-1-60 (EMD Millipore) followed by secondary antibody, Alexa fluor 488-conjugated goat anti-mouse immunoglobulin M [ThermoFisher Scientific], and Alexa fluor 488-conjugated goat anti-mouse immunoglobulin G [ThermoFisher Scientific]. All human iPSC differentiation experiments were performed using two iPSC clones from each genotype.

**Human EB formation**

Human iPSC lines were mechanically dissociated using Stem-Pro EZPassage (Invitrogen) to make clumps of a consistent size. Human EBs were generated in suspension culture over a low-speed shaker for 10 d. Culturing conditions included medium containing DMEM/F12, 20% knockout serum replacement, 1 mM L-glutamine, 100 µM M MEM nonessential amino acids, and 0.1 mM β-mercaptoethanol in the absence of FGF-2.

For rescue experiments, human iPSCs were infected by incubating three separate clones with either pRetro empty vector or pRetro-S6 wild-type virus, 10 µg/mL polybrene, and 10 µM Rock inhibitor. Forty-eight hours later, cells were selected with 2.5 µg/mL puromycin. After puromycin selection, cells were treated with 1 µg/mL doxycycline for 6 d prior to differentiation into EBs. For glycolysis inhibition, iPSCs were treated with either dH2O or 5 mM dichloroacetate (DCA), and, 24 h later, cells were mechanically dissociated into EBs as described above.

**Gel electrophoresis and Western blotting**

Chromatin extraction and Western blot analysis were performed as described previously [Sebastián et al. 2012]. Briefly, the cell pellet was resuspended in lysis buffer containing 10 mM HEPES [pH 7.4], 10 mM KCl, 0.05% NP-40 supplemented with a protease inhibitor cocktail (Complete EDTA-free, Roche Applied Science), 5 µM TSA, 5 mM sodium butyrate, 1 mM DTT, 1 mM PMSF, 50 mM NaF, 0.2 mM sodium orthovanadate, and phosphatase inhibitors [phosphatase inhibitor cocktail sets I and II, Calbiochem] and incubated for 20 min on ice. The lysate was then centrifuged at 14,000 rpm for 10 min at 4°C. The supernatant was removed (cytosolic fraction), and the pellet (nuclei) was acid-extracted using 0.2 N HCl and incubated for 20 min on ice. The lysate was then centrifuged at 14,000 rpm for 10 min at 4°C. The supernatant containing acid-solu-
in 1× PBS). Pellets were then spun down and incubated in primary antibody [anti-cTnT-FITC (Abcam, ab105439) or IgG isotype control] in 100 µL of Tween-20 buffer at a 1:50 dilution for 30 min at room temperature using an end-over-end rotator. Pellets were washed twice with Tween-20 buffer and then resuspended in FACS buffer for flow cytometry analysis.

**Directed monolayer iPSC differentiation into NPCs**

Human NPC differentiation from monolayer iPSCs was accomplished using the STEMdiff neural system according to the manufacturer’s instructions (Stem Cell Technologies). Briefly, on day 0, human iPSCs maintained in mTESR1 medium on Matrigel-coated plates were dissociated and plated in STEMdiff neural induction medium + 10 µM Y-27632. The medium was changed, and cells were split (as required per the manufacturer’s instructions) until day 6. On day 6, the medium was changed to STEMdiff neural progenitor medium until the completion of differentiation. Time points were collected every 3 d for RNA and immunofluorescence staining.

**Immunofluorescence staining of mouse and human differentiated cells**

In brief, cells were fixed with 4% formalin for 20 min at room temperature. Fixed cells were washed with PBS-glycine [130 mM NaCl, 7 mM Na2HPO4, 100 mM glycine] three times for 10 min. Wells were then blocked using immunofluorescence buffer [130 mM NaCl, 7 mM Na2PO4, 7.7 mM NaN3, 0.1% bovine serum, 0.2% Triton X-100, 0.05% Tween 20] plus 2% goat serum for 1 h. Cells were incubated with primary antibodies diluted in blocking buffer overnight at 4°C on a shaker. After three 20-min washes with immunofluorescence buffer, cells were incubated with anti-mouse or anti-rabbit secondary antibodies coupled with Alexa fluor dyes [Invitrogen] diluted in immunofluorescence buffer supplemented with 10% goat serum for 1 h at room temperature. After incubation with secondary antibodies, cells were washed three times with immunofluorescence buffer and then incubated with 0.5 ng/mL DAPI [Sigma-Aldrich] for 10 min at room temperature. Cells were washed again for 5 min with immunofluorescence buffer prior to imaging using a Nikon Eclipse Ti-E microscope. Ten images were taken per genotype. The antibodies used were as follows: anti-Sox2 (Abcam, ab92494), anti-Nestin (Abcam, ab18102), and anti-cTnT (Abcam, ab8295).

**RNA isolation and library construction**

Total RNA was isolated using Trizol-LS from three biological replicates for each condition. Library construction was performed using NEBNext poly[A] mRNA magnetic isolation module (New England Biolabs, E7490) and NEBNext Ultra Directional RNA library preparation kit for Illumina (New England Biolabs, E7420). The total RNA input amount for the poly[A] kit was 150 ng total. After poly[A] selection, all material was taken through library construction. Fifteen cycles of PCR were performed for PCR amplification. After library constructions, the samples were validated using a 2200 Tapestation system and high-sensitivity D1000 ScreenTape kit. Libraries were quantified using the Kapa Biosystems library quantification kit (catalog no. KK4828) and the Bio-Rad CFX96 instrument. Each lane of sequencing was pooled into a 21-plex (21 samples per lane) with unique barcodes over four total lanes. Pooled libraries are also quantified using the Kapa Biosystems library quantification kit (catalog no. KK4828) and the Bio-Rad CFX96 instrument. These pools are then denatured to 16 pM with 1% phiX and sequenced on an Illumina HiSeq 2000 instrument. The sequencing length read was paired-end 50 bp.

**Methods used for data analysis**

STAR aligner (Dobin et al. 2013) was used to map sequencing reads to the mouse reference transcriptome (mm9 assembly). Read counts over transcripts were calculated using HTSeq version 0.6.0 (Anders et al. 2015) based on a current Ensembl annotation file for NCBI37/mm9 assembly. Differential expression analysis was performed using EdgeR (Robinson et al. 2010); genes were classified as differentially expressed based on the cutoffs of at least twofold change and false discovery rate <0.05. Analysis of enriched functional categories among detected genes was performed using DAVID (Subramanian et al. 2005).

**Statistics**

For steady-state deacetylation/demystroylation assays, real-time RT-PCR analysis, and EB size and number and spontaneous contractile focus quantification, significance was analyzed using two-tailed Student's t-test. A P-value of <0.05 was considered statistically significant.

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**Author contributions**

C.M.F. performed and designed most of the experiments, analyzed the data, and wrote the manuscript. M.C. and R.I.S. performed bioinformatic analysis. M.A., A.V.P., E.P., A.G., S.v.K., V.M.C., M.M.A.M.M., L.K., and M.C.v.M. identified the family, collected materials from the patients, and performed the clinical analysis and sequencing of the individuals. M.C.v.M also helped design the study and provided conceptual input and discussions to write the manuscript. S.P. and G.M. derived human iPSCs from the patients and helped with some of the iPSC experiments. M.A.K. and J.M.D. performed the in vitro assays with recombinant SIRT6. J.P.E. assisted with ESC experiments and provided advice with experimental design. R.M. supervised and designed the study, interpreted the results, and wrote the manuscript with C.M.F.

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