Olfactory Stem Cells, a New Cellular Model for Studying Molecular Mechanisms Underlying Familial Dysautonomia

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Abstract

Background: Familial dysautonomia (FD) is a hereditary neuropathy caused by mutations in the IKBKAP gene, the most common of which results in variable tissue-specific mRNA splicing with skipping of exon 20. Defective splicing is especially severe in nervous tissue, leading to incomplete development and progressive degeneration of sensory and autonomic neurons. The specificity of neuron loss in FD is poorly understood due to the lack of an appropriate model system. To better understand and modelize the molecular mechanisms of IKBKAP mRNA splicing, we collected human olfactory ectomesenchymal stem cells (hOE-MSC) from FD patients. hOE-MSCs have a pluripotent ability to differentiate into various cell lineages, including neurons and glial cells.

Methodology/Principal Findings: We confirmed IKBKAP mRNA alternative splicing in FD hOE-MSCs and identified 2 novel spliced isoforms also present in control cells. We observed a significant lower expression of both IKBKAP transcript and IKAP/hELP1 protein in FD cells resulting from the degradation of the transcript isoform skipping exon 20. We localized IKAP/hELP1 in different cell compartments, including the nucleus, which supports multiple roles for that protein. We also investigated cellular pathways altered in FD, at the genome-wide level, and confirmed that cell migration and cytoskeleton reorganization were among the processes altered in FD. Indeed, FD hOE-MSCs exhibit impaired migration compared to control cells. Moreover, we showed that kinetin improved exon 20 inclusion and restores a normal level of IKAP/hELP1 in FD hOE-MSCs. Furthermore, we were able to modify the IKBKAP splicing ratio in FD hOE-MSCs, increasing or reducing the WT (exon 20 inclusion):MU (exon 20 skipping) ratio respectively, either by producing free-floating spheres, or by inducing cells into neural differentiation.

Conclusions/Significance: hOE-MSCs isolated from FD patients represent a new approach for modeling FD to better understand genetic expression and possible therapeutic approaches. This model could also be applied to other neurological genetic diseases.

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Introduction

Familial dysautonomia (FD, Riley-Day syndrome, hereditary sensory and autonomic neuropathy type III, MIM 223900) is an autosomal recessive genetic disorder that occurs in 1:3600 live births with a carrier frequency of 1 in 30 in the Ashkenazi Jewish population. The disease is characterized by incomplete development and the progressive depletion of autonomic and sensory neurons [1–3] resulting in variable symptoms including: insensitivity to pain, lack of overflow tearing, inappropriate blood pressure control manifested as orthostatic hypotension and episodic hypertension, poor oral coordination resulting in poor feeding and swallowing, and gastrointestinal dysmotility [4]. No cure is available for this disorder and treatment is aimed at controlling symptoms and avoiding complications.

FD is caused by mutations in the IKBKAP gene which encodes a protein termed IKAP/hELP1 [5,6]. The most prevalent mutation, is a splice mutation; the T-to-C transition in position 6 of the 5’ splice site (5’ss) of intron 20 (IVS20 +6T→C) of this gene. All FD cases have at least one copy of this mutation; >99.5% are homozygous [5–7]. This mutation leads to variable, tissue-specific skipping of exon 20 of IKBKAP mRNA, with the central and peripheral nervous system more prone to complete skipping than others tissues, which leads to reduced IKAP/hELP1 protein levels [8].
Although the exact function of the IKAP/hELP1 protein is not clearly understood, researchers have identified IKAP/hELP1 as the scaffold protein required to assemble a well conserved six-protein complex (ELP1-6) called the holo-Elongator complex that possess histone acetyltransferase activity directed against histone H3 and H4 in vitro [9]. IKAP/hElongator is recruited to the transcribed regions of some human genes essentially involved in actin cytoskeleton regulation and cell motility migration [10]. This role may underlie a cell motility deficiency in FD neurons because of impaired transcriptional elongation of some genes coding for proteins involved in cell migration. Indeed, one study found that mouse neurons defective in Elongator exhibit reduced levels of acetylated α-tubulin, causing defects in radial migration and branching of cortical projections neurons [11]. Another study showed that Caenorhabditis elegans Elongator complex is required for correct acetylation of microtubules and neuronal development [12].

IKAP/hELP1 protein is also involved in other cellular processes, including tRNA modifications [13–15], exocytosis [16], and zygotic paternal genome demethylation [17]. Recently, its homolog in fly (D-elp1) has also been suggested to be involved in RNA interference through a RNA-dependent RNA polymerase activity [18].

To better understand the molecular mechanisms leading to aberrant splicing of IKBKAP mRNA in FD, creation of model systems recapitulating the pathological development of neural cells is required. Because IKBKAP gene knock out causes embryonic lethality [19], an animal model that exhibits the major phenotypic characteristics observed in FD humans has not yet been established. However, a humanized IKBKAP transgenic mouse model for FD has been created [20], that reproduces the tissue-specific splicing of IKBKAP mRNA in nervous tissues. Such a model is a notable progress in the comprehension of this complex rare disease and offers a potential system for testing therapeutic agents. However, transgenic animals do not reproduce phenotypic features of FD as they maintain normal development. Alternatively, FD patient fibroblasts are an informative model of mRNA splicing regulation. However, a recent study suggests that IKAP/hELP1 expression is much higher in neurons compared to fibroblasts [21], and fibroblasts do not exhibit the same ratio of IKBKAP exon 20 including: exon 20 skipping transcripts (named WT:MU respectively for simplicity) as observed in nervous system-derived tissues [8]. This finding narrows the understanding of disease mechanisms in a neural context. Finally, generation of neural cells through the production of induced pluripotent stem (iPS) cell from FD fibroblasts has been recently established [22]. Neural cells derived from iPS cells have potential to be used for studies of neuropathologies [23]. However, the labor intensive reprogramming required to induce iPS cells erases the developmentally relevant epigenetic signature specific to the disease state. As a consequence, some important information may be lost impeding recreation of an accurate disease model. The demonstration that fibroblasts can be converted directly into neurons, without an initial reprogramming, as recently evidenced in mouse [24], is very attractive. Nevertheless, during their reprogramming, human iPS cells do not pass through the normal stages of embryonic development that human ES cells undergo. Although both stem cell types share a common transcriptional signature, a subset of genetic profiles found in human iPS cells suggests retention of transcriptional and epigenetic memory related to their tissue of origin, which can substantially affect their potential to differentiate into different cell types [25–27]. Thus, cells collected from primary sources that have been subjected to environmental signals appropriate for the pathological specificity of the targeted disease are likely important to mirror the biology of diseased human neural cells.

Our aim is to understand what mechanisms drive IKBKAP mRNA splicing to the almost exclusive production of aberrant transcripts (MU) in neuronal cells. Here we demonstrate the potential of human olfactory ecto-mesenchymal stem cells (hOE-MSCs) to model this aspect in FD. Indeed, neurogenesis occurs throughout adult life in the olfactory mucosa, due to the presence of resident multipotent stem cells giving rise to olfactory neurons in vivo [28]. hOE-MSCs can be grown into neurospheres, that are multipotent, and differentiate in vitro into neurons, astrocytes, and oligodendrocytes as well as other cell types [29,30]. Isolated from patients, cultures of hOE-MSCs provide potential models for genetically determined neuropsychiatric diseases [31–33], and stand as an interesting human model to investigate gene networks and cellular pathways altered in disease like FD. For example, cell migration defects have been observed in cells lacking normal expression of IKAP/hELP1 [10,11,22,34], and we show here that FD hOE-MSCs exhibit impaired migration compared to control cells. Additionally, hOE-MSCs are an appropriate model for validating the potency of therapeutic agents such as kinetin, a cytokinin that has been shown to increase IKBKAP mRNA and protein expression in FD cell lines and in vivo models [29,22,35,36] as well as in leukocytes of healthy carriers of the FD mutation [37].

Results

FD hOE-MSCs express stem cell, glial and immature neuronal markers

To establish a human cellular model of FD, we collected 4 olfactory mucosa biopsies from homozygous patients for the IVS20+6T>C FD mutation. As previously demonstrated with control biopsies [30], after about 2 weeks of culture, the microscopic examination of the tissue crushed under a glass coverslip revealed stem cell proliferation (Figure 1A and 1B). After reaching confluency in a 4-well plate, the cells attached to the glass coverslip were further expanded by transfer into a 6-well plate (Figure 1C). Like control hOE-MSCs, we observed that FD hOE-MSCs could be cultured for long periods (at least 15 cycles of trypsin/EDTA treatment and expansion on larger plastic surface) with a doubling time of about 30–48 h. When subjected to immunostaining, all hOE-MSCs derived from control and FD biopsies express the neural stem cell–specific marker nestin (Figure 1D and 1E) and the immature neuronal marker β-III tubulin (Figure 1F and 1G) in the same proportions (Figure 1H and 1I). A comparatively low GFAP staining was observed in every hOE-MSCs (Figure 1J and 1K). In addition, cells were negative for a mature neuronal marker, MAP2 (Figure 1L and 1M). This analysis suggests that both control and FD hOE-MSCs display properties of neuralgial progenitor cells.

Expression of IKBKAP transcripts is dramatically reduced in FD hOE-MSCs

IKBKAP mRNA expression was investigated in cultures of 5 controls and 4 FD hOE-MSCs at early (P1, P2) and later cell passages (P5, P9). A semi quantitative RT-PCR analysis revealed that, while control hOE-MSCs expressed exclusively the WT mRNA transcript (Figure 2A, left panel), FD hOE-MSCs expressed the WT but also the MU transcript (Figure 2A, right panel). We also demonstrated that long time culture conditions and trypsin-EDTA mediated cell passages did not affect the IKBKAP gene expression pattern. In order to more accurately determine the level of expression of IKBKAP alternative transcripts, we designed primers, probes and plasmid calibrators to perform absolute quantification using quantitative real-time RT-PCR (RT-qPCR) on the same samples. Strikingly, WT transcripts were...
much less expressed in FD (5–8 fold), when compared to controls hOE-MSCs (Figure 2B). In addition, WT and MU transcripts were present in nearly equal amounts in FD hOE-MSCs (Figure 2B, right graph). Furthermore, the total amount of IKBKAP transcripts in FD (WT+MU) remains 3 to 5 times less abundant than WT in controls, which suggests a defect in IKBKAP transcription and/or mRNA stability. In FD cells, the differential expression of IKBKAP transcripts was also correlated to a reduced expression of IKAP/hELP1 protein in FD, when compared to controls, as revealed by western blot analysis (Figure 2C). Since MU transcripts contain a premature stop codon that may activate the nonsense-mediated mRNA decay (NMD) pathway, we wanted to confirm whether this pathway is responsible for the lower IKBKAP transcripts expression in FD cells. Thus, we tested cycloheximide, a protein synthesis inhibitor which also inhibits NMD. Indeed, FD cells preincubated for 6 h with cycloheximide exhibited a stabilization of the MU transcript as evidenced by semi-quantitative RT-PCR (Figure 2D, left panel). To accurately determine the level of WT and MU IKBKAP transcripts in these samples, absolute RT-qPCR analysis was performed (Figure 2D, right panel). The results clearly demonstrated that the WT:MU ratio decreases when mRNA surveillance is inhibited. Thus, a
large amount of IKBKAP MU transcripts is degraded through the NMD pathway resulting in much less IKBKAP transcripts and IKAP/hELP1 protein in FD compared to control cells.

Heterogeneous IKAP/hELP1 distribution in hOE-MSCs

Since the localization of IKAP/hELP1 remains controversial and is important to understand protein functions, we stained both control and FD hOE-MSCs with the monoclonal antibody directed against IKAP/hELP1 and previously used for detecting the protein by western blot analysis. In control cells, confocal imaging revealed a weak and diffuse signal with a dominant cytoplasmic staining within the perinuclear area. We could also detect the presence of IKAP/hELP1 in the nucleus of hOE-MSCs (Figure 3A–C). Significantly, FD hOE-MSCs exhibit a weaker anti-IKAP/hELP1 immunofluorescence staining compared to control cells, with a similar distribution of the staining (Figure 3D–F). Therefore, collectively, our results are in agreement with a wide distribution of IKAP/hELP1, including a much lower IKAP/hELP1 staining in FD hOE-MSCs, in agreement with RT-qPCR and western blot analysis.

Transcriptome analysis identified fifty dysregulated genes

It is widely accepted that culture conditions alone may exert effects on gene expression, resulting in experimental inconsistencies [38,39]. Thus, to investigate the involvement of candidate disease mechanisms in FD and to test whether differences in gene expression are stably imprinted in FD compared to control hOE-
MSCs, we explored the transcriptome of these cells at very early (P1, P2) and later (P5, P9) cell passages with the same samples used to quantify IKBKAP transcripts. Among the 8,780 cDNA represented on the microarray, 46 were significantly decreased and only 4 increased in FD hOE-MSCs, when compared to control hOE-MSCs (fold-change \( \leq 1.4 \)-fold; \( p \)-value \( \leq 6.10^{-3} \), Table 1 and Table S1), considering a false discovery rate (FDR) of 3% (Figure S1). Notably, the biological processes and the signaling pathways most significantly targeted by the effectors on our list were actin cytoskeleton organization, cell growth, and apoptosis (Table 1). More specifically, we identified 10 genes (Table 1 and Table S1) that also exhibited a significant dysregulated expression in previous microarray studies [10,22]. Interestingly, 2 genes, PMEPA1 and GSN (encoding TMEPAI and gelsolin, respectively), involved in cell growth and cytoskeleton organization, respectively, were dysregulated in both the IKBKAP RNAi and FD iPS cell studies.

In order to assess the robustness of our microarray analysis, RT-qPCR analysis was performed, on independent RNAs extracted from 4 control and 4 FD hOE-MSCs harvested at the second, fourth, and seventh cell passage. Since gene expression quantification using RT-qPCR requires a steady reference gene, we selected three genes frequently used for normalization of the data, ABL1, RPLP0, and HPRT1. We confirmed that PMEPA1 (Figure 4A), the most dysregulated gene on the microarray, and S100A16 (Figure 4B), were significantly underexpressed in FD samples. The expression pattern of these two candidate genes was essentially identical at all passages with the 3 reference genes, which demonstrates the validity and reliability of the array data.

FD OE-MSCs migration is altered compared to controls

To explore the functional consequence of a down-regulated expression of genes involved in cell migration in FD hOE-MSCs compared to control cells, we used the Boyden's chamber assay. After comparing the migration pattern of 3 control and 3 FD hOE-MSCs in serum medium and serum-free medium (ITS), we determined that FD cells invasion is significantly reduced compared to control cells both in serum and in ITS medium (Figure 5).

Confirmed down-expression of first and final IKBKAP exons in FD hOE-MSCs

Since we and others [22,40] did not detect IKBKAP among the significantly down-regulated transcripts in FD compared to control samples, we asked whether this discrepancy could be due to a lack of sensitivity of microarray compared to RT-qPCR. For this purpose, we decided to analyze IKBKAP levels of expression by investigating other exons distal from IKBKAP exon 20. By looking at the beginning of IKBKAP transcript, we identified a second event of alternative splicing. After amplifying transcripts from exon 1 to exon 5, we obtained 2 PCR products (Figure 6B, upper panel). The sequencing of the less abundant and shorter PCR product revealed the use of an alternative 3's within IKBKAP exon 2, which is shortened of 145 nt (Figure 6A, left schematic). Accordingly, the loss of the ATG start codon located within the 5't end of exon 2 can potentially induce the use of an alternative ATG start codon (in exon 4), resulting in the synthesis of a putative 114 amino acid-truncated IKAP/hELP1 protein (Figure 6D).
When investigating expression at the end of *IKBKP* coding sequence, again we observed a third alternative splicing event. The amplification from exon 33 to exon previously numbered exon 36 (and now called exon 37) revealed 2 products (Figure 6B, middle panel). The sequencing of the barely detectable and longer PCR product revealed the inclusion of an additional exon (Figure 6A, right schematic). This exon inclusion also induced a frameshift and resulted in a premature stop codon whose relative location may

### Table 1. Most dysregulated genes in FD hOE-MSCs are involved in seven over-represented cellular processes.

| Gene      | Clone ID | FC   | p-value  | Biological process                                      | Studies                  |
|-----------|----------|------|----------|---------------------------------------------------------|--------------------------|
| Actin cytoskeleton reorganization | p = 0.000275 |
| GSN       | 246170   | −1.91| 0.00124  | actin cytoskeleton reorganization                       | *1 *2                    |
| MYO9B     | 279085   | −1.70| 0.00144  | actin cytoskeleton remodeling                           |                          |
| MYPN      | 325601   | −1.52| 0.00131  | sarcomere organization through nebulin and \( \gamma \)-actinin interactions |                          |
| DSTN      | 149199   | −1.48| 0.00017  | actin filament depolymerization                         |                          |
| CORO2B    | 547561   | −1.44| 0.00079  | neuronal actin structure reorganization                 |                          |
| SLCA43R2  | 155467   | 2.42 | 0.00034  | adaptor of ion channels and receptors to the actin cytoskeleton | *1                      |
| Regulation of apoptosis | p = 0.00203 |
| CHEK2     | 1893020  | −2.20| 0.00007  | cell cycle arrest and apoptosis in response to DNA damage | *1                      |
| ZMAT3     | 525407   | −1.78| 0.00251  | positive regulation of p-53-mediated apoptosis           |                          |
| TNFSF10   | 713945   | −1.75| 0.00041  | induction of apoptosis by activation of caspase activity | *2                      |
| PARP3     | 436086   | −1.60| 0.00004  | positive regulation of apoptosis- maintenance of genomic stability |                          |
| Transport | p = 0.00224 |
| ABCG5     | 121977   | −1.59| 0.00089  | cholesterol transport in and out of the enterocytes     |                          |
| SLC35E1   | 487960   | −1.55| 0.00086  | monosaccharide transport                                |                          |
| SLC22A6   | 36482    | −1.52| 0.00050  | \( \gamma \)-ketoglutarate transmembrane transporter activity |                          |
| SFXN2     | 757192   | −1.45| 0.00021  | iron transport                                          |                          |
| Cell proliferation | p = 0.00552 |
| APOE      | 1870594  | −1.68| 0.00028  | cell proliferation-regulation of neurite extension       | *1                      |
| CD22      | 284220   | −1.58| 0.00032  | B cell proliferation                                    |                          |
| CD38      | 123264   | −1.54| 0.00074  | B cell proliferation                                    |                          |
| GBA       | 757264   | −1.53| 0.00012  | cell proliferation-ceramide metabolic process            |                          |
| SERINC2   | 149995   | 1.84 | 0.00014  | cell proliferation                                      |                          |
| Regulation of cell growth and cell cycle | p = 0.0091 |
| PMEPA1    | 366599   | −4.92| 0.00585  | EGF receptor signaling pathway - negative regulation of cell growth | *1 *2                   |
| STRBP     | 669157   | −1.75| 0.00007  | regulation of cell growth                               |                          |
| INO80B    | 323554   | −1.63| 0.00104  | growth induction and cell cycle arrest at the G1 phase   |                          |
| S100A16   | 739851   | −1.55| 0.00012  | regulation of cell cycle progression                    | *1                      |
| CDIPT     | 306047   | −1.49| 0.00088  | regulation of cell growth                               |                          |
| Nervous system process | p = 0.0302 |
| LRCH1     | 683580   | −3.62| 0.00080  | long-term memory and learning - signal transduction     |                          |
| KCNT2     | 38677    | −2.06| 0.00010  | synaptic transmission mediated by K (+) channels        |                          |
| NUMBL     | 1855110  | −1.61| 0.00118  | Notch signaling pathway inhibition - cerebral cortex morphogenesis |                          |
| DULLARD   | 346388   | −1.53| 0.00198  | Nuclear organization-negative regulation of BMP signaling |                          |
| Proteolysis | p = 0.0334 |
| FBXL15    | 166240   | −1.71| 0.00130  | ubiquitin-dependent protein catabolic process            |                          |
| WSB1      | 298983   | −1.61| 0.00123  | ubiquitination and proteasomal degradation of target proteins |                          |
| PCSK7     | 241130   | −1.53| 0.00155  | proteolysis- ubiquitous endoprotease activity           |                          |
| RNF115    | 471834   | −1.42| 0.00067  | proteolysis- vesicle-mediated transport - vesicle traffic |                          |
| MMP27     | 767086   | −1.35| 0.00067  | proteolysis of fibronectin, laminin, gelatins and/or collagens |                          |

Clone ID represents the number assigned to the original clones produced by the I.M.A.G.E Consortium. FC = Fold change, and p-values were calculated by SAM analysis as described in Methods. This list of genes was annotated with the ExplainTM System from Biobase. 7 majors processes are overrepresented in our list of genes, and, for each process, p-values were calculated and adjusted by the Bonferroni correction. The last column indicates the genes that were also found to be significantly dysregulated in 2 previous FD studies.

*1 = Lee et al. 2009.  
*2 = Close et al. 2006.  
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lead to NMD of this new isoform (Figure 6D). We confirmed this exon 36 inclusion with specific primers (Figure 6B, lower panel).

Both new alternative splicing events we described were also observed in others cell types (fibroblasts, HeLa, peripheral mononuclear cells, data not shown) and we decided to focus on the two major splicing events, full exon 2 inclusion and exon 36 skipping. We derived the tools (plasmids, primers, probes) to perform absolute quantification of full-length exon 2 inclusion and exon 36 skipping by RT-qPCR on samples from 4 controls and 4 FD hOE-MSC cultures. Similar underexpression of IKBKAP transcripts (WT+MU) was observed in FD cells compared to control cells, regardless of the exon investigated (Figure 6C). Thus, these results confirmed a decreased level of IKBKAP transcripts (WT+MU) in FD cells.

In addition, we tested for the stability of exon 36-containing transcripts after cycloheximide treatment. Absolute RT-qPCR analysis revealed that the exon 36 skipping : exon 36 inclusion ratio decreases when NMD pathway is inhibited (Figure S2), suggesting that transcripts including exon 36 are degraded through NMD in FD, as well as in control OE-MSCs (data not shown).

Kinetin treatment corrects aberrant IKBKAP pre-mRNA splicing

Our next goal was to assess whether the production of both WT and MU IKBKAP mRNAs can be modulated in our model. In previous studies, one compound, kinetin (6-furfurylaminopurine) was found to correct IKBKAP splicing and increase IKAP/hELP1 production in FD cells [35]. We tested whether this drug could also modify the splicing defect of IKBKAP in FD hOE-MSC cells. For this purpose, we used increasing concentrations of kinetin (25 to 200 μM) on a FD hOE-MSCs culture for 72 h. As expected, after semi-quantitative RT-PCR, we observed a significant decrease of MU transcript compared to non-treated cells on agarose gel electrophoresis (Figure 7A). The level of IKBKAP mRNA splicing correction increased proportionally to the concentration of kinetin, and the MU transcript almost vanished at 100 μM. The dose-dependent action of kinetin on increasing WT:MU ratio was confirmed by RT-qPCR analysis (Figure 7B). A similar finding was observed when IKAP/hELP1 proteins were detected by western blot analysis (Figure 7C). Accordingly, when FD hOE-MSC were incubated with 100 μM kinetin for 24 h, we observed a major increase of anti-IKAP/hELP1 staining, in cytoplasmic as well as in nuclear areas (Figure 3G–I). However, the same kinetin treatment could not rescue the migration defect observed in FD hOE-MSCs with the Boyden’s chamber assay (Figure 5).

In order to determine how fast kinetin modulates IKBKAP mRNA splicing, we performed a time-course experiment with a constant concentration of 80 μM over 24 h. After performing semi-quantitative RT-PCR analysis, the first significant increase of WT:MU ratio was seen after 24 h of kinetin treatment (Figure 7D).
Figure 6. Expression and alternative splicing of IKBKAP mRNA at the extremities of the coding sequence. Two additional splicing events are described within IKBKAP gene. The first one represents the alternative use of a 3'ss for exon 2 and the second one concerns exon 36 skipping,
represented by a schematic (A). B, semi-quantitative RT-PCR illustrates relative amounts of both events on 2 control and 2 FD hOE-MSCs. C, RT-qPCR analysis performed at 2 different regions (exon 2 and exon 36), using 4 control and 4 FD samples, at the same cell passage (P7). Histograms represent the mean value of 4 samples, normalized with ABL1 gene. (*** P<0.001 using two-tailed Student’s test. IKAP/hELP1 truncated regions for all splicing events are represented by a schematic (D). Grey portions represent the conserved amino acids while black portions represent new amino acids resulting from a frame shift. Putative functional domains of the protein are indicated as well as the immunogenic region for the monoclonal antibody used in western blot and immunocytochemistry experiments.

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However, quantitative analysis by RT-qPCR revealed that kinetin significantly enhances the ratio 6 h after its addition to the culture (Figure 7E). Interestingly, time-response of kinetin was maximal at 24 h during treatment, but its effect on splicing lasted more than 6 h after the drug was washed out and WT transcript levels remained high compared to non-treated cells, at least 24 h after the wash out. Consistent results were observed for IKAP/hELP1 protein expression by western blot analysis, although a strong decrease of protein amount appeared after 24 h of wash out (Figure 7F).

Finally, we wanted to investigate kinetin activity along the IKBKAP transcript. Therefore, we compared the level of expression of IKBKAP transcripts by RT-qPCR to focus on different transcript levels in control cells, which likely excludes a potential modification of the WT:MU transcript ratio.

Commitment of FD OE-MSCs into neuronal and glial lineages leads to a more severe IKBKAP exon 20 skipping

FD hOE-MSCs were treated for 7 days to induce neuronal differentiation with a protocol previously used in hOE-MSC [41], which consists of adding retinoic acid, forskolin, and Sonic hedgehog in the medium (called rafnshh medium). Cells were first cultured in serum-free medium supplemented with N2 and B27 until they became adherent before being cultured in rafnshh (Figure 9B). The new culture medium induced a slight morphology change, as compared to the serum condition (Figure 9A). When hOE-MSCs were first cultured in rafnshh, they began to form long fine processes and neural-like cells (Figure 9C). After 7 days of treatment, a majority of cells adopted neuron-like morphologies (Figure 9D) and established a wide range of connections (Figure 9E, F and M). Using end-point PCR on 3 different FD cell cultures, we observed that IKBKAP mRNA splicing in rafnshh-treated cells was more prone to exon 20 skipping as compared to untreated cells (Figure 9G). This change can be quantified by RT-qPCR (Figure 9H). In contrast, we did not detect significant variations in exon 2 and exon 36 alternative splicing during neuronal differentiation (data not shown).

When assessing immunostaining on treated cells, we observed that rafnshh treatment increased the proportion of both GFAP (Figure 9I and J) and MAP2-expressing cells (Figure 9K and L). Double-labeling with β-III tubulin and nestin revealed a stronger expression of β-III tubulin compared to nestin during the differentiation process (Figure 9N–P). Collectively, these results indicate that rafnshh treatment influences the neural and glial lineage commitment. As a consequence, the splicing machinery in neuron or astrocyte-differentiated cells is impaired for IKBKAP exon 20 recognition.

Discussion

Deciphering the molecular basis of the tissue-specific pattern of IKBKAP mRNA splicing in FD nervous tissues is crucial for the comprehensive analysis of disease physiopathology. In this study, we aimed to recapitulate different aspects of IKBKAP gene expression using FD hOE-MSCs cultured with different conditions. While other human cellular models, such as fibroblasts or iPSCs cells, have been investigated to understand FD, we believe that hOE-MSCs hold a great promise to model the FD disease pathology. hOE-MSCs are easily obtained by a simple biopsy and can be maintained for an extended period of time and can be rapidly expanded in basic culture conditions without genetic manipulation. In addition, due to the origins of hOE-MSCs from a peripheral tissue, these cells are able to express neuroglial markers in vitro [42–47]. Thus, they constitute an efficient and simple method to derive neuronal cells in the original context of the genetic mutation studied.
In contrast to control cells, which constitutively include \textit{IKBKAP} exon 20, we confirmed the alternative splicing of that exon in FD cells (Figure 2A). Similar to neural precursors obtained from iPS cells [22], FD hOE-MSCs predominantly express the MU \textit{IKBKAP} transcript isoform (Figure 2A and B). Moreover, we demonstrated that FD cells exhibit notably lower \textit{IKBKAP} transcript levels (WT + MU), when compared to controls (Figures 2B, 6C and 7G). Such a difference is most likely explained by an extensive degradation of MU transcripts through the NMD pathway (Figure 2D), as was previously suggested [35]. However, NMD efficiency varies between cell types and individuals [48–50] and it is unclear how prevalent this mechanism of mRNA degradation is in the nervous system of FD individuals. In order to get a better insight into the actual contribution of NMD on the decay of \textit{IKBKAP} MU transcripts, it will be necessary to specifically block the NMD machinery.

Figure 7. Action of kinetin on \textit{IKBKAP} mRNA splicing in FD hOE-MSCs cells. A–C, hOE-MSCs were treated with increasing concentration of kinetin for 72 h. (i) Total RNAs were reverse transcribed and subjected to both semi-quantitative PCR (A) and absolute qPCR (B) of WT and MU \textit{IKBKAP} transcripts, (ii) total lysates were analyzed by western blot using a monoclonal mouse anti-IKAP/hELP1 antibody (C). D–F, kinetics of hOE-MSCs incubated for 24 h with 80 \textmu M kinetin which was then removed for the next 24 h. Total RNAs were reverse-transcribed and subjected to \textit{IKBKAP}-specific semi-quantitative PCR (D) and absolute RT-qPCR (E). Total lysates were analyzed by western blot (F). The level of WT and MU transcripts was normalized using \textit{ABL1} as a reference gene (B and E). G, Two controls (CTRL) and two FD hOE-MSCs treated or not with 80 \textmu M kinetin for 24h were analyzed by absolute RT-qPCR to determine the amount of \textit{IKBKAP} exon 2 inclusion, exon 20 inclusion, exon 20 skipping, and exon 36 exclusion after normalization with \textit{ABL1}. (* \(p<0.05\); *** \(p<0.001\) using two-tailed Student’s test). doi:10.1371/journal.pone.0015590.g007
So far, the proposed functions of IKAP/hELP1 are related to various cellular localizations. This has been the matter of a controversy because several studies have failed to detect IKAP/hELP1 in the nucleus or found it almost exclusively in the cytoplasm [11,21,34,51], which is difficult to reconcile with its suggested role in transcription elongation. As observed in most published studies, we observed that the immunolocalization of IKAP/hELP1 was mainly cytoplasmic within the perinuclear area. However, we also detected significant nuclear staining in agreement with other reported studies [9,52]. Altogether, our findings of IKAP/hELP1 distribution in hOE-MSCs support multiple roles for the protein within different subcellular compartments.

In order to establish a direct link between low levels of IKBKAP WT transcripts and decreased neuronal populations in FD patients, several groups have investigated transcriptome variations resulting from a reduced level of IKBKAP transcripts using microarray technology [10,22,34,40,53]. However, poor correlations were observed between these studies. Several reasons can explain these discrepancies. First, various cell types, used at different stages of development and differentiation, have been studied (brain tissue, fibroblasts, HeLa cells, HCT116 cells, iPS cells). The cells tested in the current study are likely at a stage between the iPS and iPS cell-derived neural crest precursors developed by the Studer group [22]. It is thus not surprising that most of our microarray results overlap with that of the iPS cell study. Second, a potential source of variability among transcriptome analyses derives from the technical manipulations employed to downregulate IKBKAP (presence of the FD mutation in its original context compared to WT IKBKAP knockdown using different interfering RNAs), resulting in differential residual IKBKAP/IKBKAP expression. Third, in the context of a rare disease, a small sample size may cast doubt upon the validity of drawn conclusions. To decrease statistical bias, we decided to increase the number of samples of our 5 control and 4 FD patients by collecting data from 4 different passages (P1, P2, P5 and P9) of each primary cell line. We hypothesized was that such a method would allow us to i) increase the statistical power of our analysis, and ii) explore the effects of culture conditions on gene expression. We anticipated that a strong gene dysregulation observed in microarray would be more significant if this expression is stably maintained at any cell passage. Most of the differentially expressed genes were found to have a modest (<2) fold-change (Table 1). Interestingly, like previous studies, we observed that a majority of genes were down-regulated in FD hOE-MSCs (Table 1, negative values), and only 4 genes were up-regulated (Table 1 and Table S1, positive values). This observation is in agreement with other studies [10,22,40] and may reflect a defect in transcription due to decreased Elongator activity, as previously proposed [10]. Importantly, 10 genes in our list (20%) appeared to be correlated with one or two previous investigations (Table 1 and Table S1, last column).

In one of the past studies, IKBKAP expression level can be downregulated by RNAi in control cells [10], where there is no production of MU transcripts. Thus, different studies share dysregulated genes in different contexts of either constitutive or alternative splicing of IKBKAP mRNA. This suggests that IKBKAP alternative splicing may not be the only pathological alteration in FD.

Similar to what was previously reported, our study revealed the downregulation of gelsolin (GSN), a protein involved in cell motility, that causes defects in cytoskeleton reorganization and cell migration in FD [10,11,22,34]. The most dysregulated gene observed in our study was TMEPAI (~4.92 fold change), encoding the TMEPAI protein which has recently been reported to be a direct target of the TGF-β signaling pathway and is involved in cell growth, cell differentiation, and apoptosis [54]. Due to its important cellular function and repeated reports of its dysregulation in FD cells, it would be very interesting to test TMEPAI in further studies. In agreement with previous studies correlating a decreased expression of IKAP/hELP1 with defects in cell migration [10,11,22,34], the Boyden’s chamber assay show that FD hOE-MSCs have decreased migration potential, compared to control cells (Figure 5).
Surprisingly, we did not find *IKBKAP* as a dysregulated gene in our microarray analysis. This result is all the more intriguing since this gene is expressed at much lower levels in FD hOE-MSCs, as shown by RT-qPCR in the exon 20 region (Figure 2B). However, previous analyses using microarrays also failed to detect *IKBKAP* as a down-regulated gene in FD cells [22,40]. To address the question of a possible PCR artifact or lowered microarray sensitivity and because the FD mutation is located in the middle region of the *IKBKAP* gene, we performed quantitative PCR at both ends of the *IKBKAP* gene.

Unexpectedly, we identified 2 new events of alternative splicing at both extremities of *IKBKAP* coding sequence (Figure 6A). However, these results are in agreement with EST sequences found in alternative splicing database such as ASD [55].

Figure 9. WT:MU ratio is decreased in differentiated hOE-MSCs. A, Phase contrast microscopy of FD hOE-MSCs cultured for 48 h in either serum (A), or N2B27 conditions (B), or with the rafnshh cocktail including retinoic acid, forskolin and Sonic hedgehog (C), or rafnshh for 7 days (D). Details of connections established between cells and extensive cellular arborization after 7 days in rafnshh condition are shown in E and F. G, agarose gel electrophoresis of semi-quantitative RT-PCR products showing *IKBKAP* WT and MU transcripts of 3 different FD patients before and after rafnshh treatment. H, histograms represent the mean level of *IKBKAP* transcripts expression normalized with *ABL1* gene expression for 3 FD patients, after RT-qPCR analysis. (*P<0.05, using two-tailed Student’s test). FD hOE-MSCs treated for 7 days in rafnshh were fixed and stained for GFAP (I, one cell with ramified neuritic process is magnified in J), MAP2 expression (K and L). M–P, double labelling of rafnshh-treated cells with anti-β III tubulin and anti-nestin antibodies. Scale bars represent either 50 μm (A–D, I, K, M–P) or 25 μm (E–F, J, L). doi:10.1371/journal.pone.0015590.g009
revealed that the alternative use of a 3’ss (downstream of the ATG start codon) leads to a shorter exon 2 which can potentially induce the use of an alternative ATG start codon (in exon 4), resulting in the synthesis of an N-terminal truncated IKAP/hELP1 protein. In addition, we detected the presence of intronic sequences at the end of IKBKAP gene leading to a supplementary exon in the mRNA, named “exon 36”. This exon inclusion also induced a frameshift and resulted in a premature stop codon whose relative location likely led to NMD of this new isoform, as observed by stabilization of the transcript after cycloheximide treatment (Figure 52).

IKAP/hELP1 plays the role of a scaffold protein in Elongator complex assembly and the C-terminus half of IKAP/hELP1 is responsible for this function [34]. It has also been shown that IKAP/hELP1 contains five WD-like repeats domains in the N-terminal part that may play a role for protein-protein interactions [56]. When comparing the different protein isoforms resulting from the 3 alternative splicing events we described (Figure 6D), only the isoform resulting from exon 20 skipping seems to lack a functional domain and may play a pathological role during FD progression. However, the protein domains of IKAP/hELP1 important for Elongator integrity have not been precisely mapped and other roles for IKAP/hELP1 have been proposed outside of the Elongator complex. Thus, it is not clear whether the new IKBKAP isoforms we described, may have functional roles. Future investigations with specific reagents (antibodies) will be required to address this issue.

Nevertheless, we consistently detected a lower expression of IKBKAP gene (including the full length exon 2 transcript and the transcripts skipping exon 36) in FD hOE-MSCs (Figure 6C), as determined when investigating the exon 20 region. Thus, the relatively stable expression of IKBKAP observed in microarray analysis may be due to a weak expression that is masked within the noise signals. Furthermore, during the analysis of our microarray data and those of previous studies [10,22,40] we noticed that a high fraction of genes were expressed at background levels. This points to the limitation of using microarray technology to establish the whole genome expression pattern. We expect that new technologies such as RNA deep sequencing will rival PCR sensitivity and specificity in the near future.

The model of hOE-MSCs from FD patients has also been very useful to test compounds, such as kinetin, that can correct the defective splicing process. As reported in the other cell types tested, we confirmed that kinetin corrects splicing in a dose-dependent manner in FD hOE-MSCs (Figure 7A-C). This suggests that kinetin activity is not cell-type specific. Although the mechanism by which kinetin modulate splicing remains poorly understood [36,57], it is unlikely that kinetin acts directly on the general transcription machinery as the level of IKBKAP transcripts was not significantly modulated by kinetin in control hOE-MSCs (Figure 7G). This effect of kinetin has also been previously observed in control iPS cells [22]. Time-course experiments of kinetin treatment revealed that the drug acts quite rapidly on correcting IKBKAP mRNA splicing and enhancing IKAP/hELP1 synthesis, but its effects last only a short time after removal (Figure 7E and F). This information provides new perspectives in the strategy of kinetin delivery to FD patients. First, kinetin as an FD treatment would potentially decrease deleterious consequences of the mutation at the protein level. In addition, drug efficacy may be achieved if adequate levels of kinetin is maintained over a long period of time. However, as observed for FD iPS cells [22], kinetin did not improve cell migration in FD hOE-MSCs (Figure 5), suggesting incomplete phenotype complementation.

Using the hOE-MSCs model, we were also able to modulate the expression of IKBKAP WT and MU transcripts, by exposing the cells to different culture conditions to simulate variations in alternative splicing occurring during development and differentiation. hOE-MSCs form free floating spheres in approximately 7 days, when cultured with EGF and bFGF in serum-free medium. It is known that in sphere conditions, cells can form a niche prevent differentiation and ensure self-renewal. The cell populations contained in hOE-MSC-derived spheres are not well-known. Some reports indicate that they include a heterogeneous mixture of stem cells and neuronal progenitors [29,43,44,47,58]. However, immunostainings of nestin and β-III tubulin show no significant differences. Interestingly, PCR analysis demonstrated that spheres express higher levels of WT IKBKAP transcript compared to hOE-MSCs in serum and express very few levels of MU transcript. However, when the cells were transferred back to culture conditions with serum, the enhanced IKBKAP exon 20 inclusion was not maintained (Figure 8G and H). FD hOE-MSCs that were cultured in serum-free conditions without forming spheres did not exhibit significant changes in IKBKAP isoforms, suggesting that there is subpopulation of cells within the spheres that can promote IKBKAP exon 20 inclusion. These results indicate that when FD cells are turned back into a more “primitive” developmental stage, IKBKAP aberrant splicing is corrected, as was described during the fibroblast to iPS cell reprogramming process [22]. Accordingly, commitment into a more differentiated neuronal state would alter IKBKAP exon 20 inclusion. Therefore, we differentiated FD hOE-MSCs, using a previously established protocol which included retinoic acid (RA), forskolin (FN), and Sonic hedgehog (Shh) in the culture medium [44]. In these conditions, we observed that differentiated cells express the highest levels of MU IKBKAP transcript (Figure 9G and H). This result correlates with the specific low WT:MU IKBKAP isoform ratio in nervous tissues [8], and suggests that stem cells engaged in a neuronal lineage with appropriate culture conditions can rapidly switch their IKBKAP WT:MU transcript ratio.

Previous studies have shown that i) IKBKAP exon 20 is poorly defined in a “healthy” context, due to the presence of a weak 3’ss and exonic splicing silencers, and ii) the FD mutation exacerbates the environment leading to alternative exon 20 inclusion in FD tissues [59,60]. We propose that some transcription/splicing factors involved in IKBKAP exon 20 recognition are also downregulated in a tissue-specific manner. This would explain why the pattern of IKBKAP alternative mRNA splicing is more aberrant in the nervous system. Interestingly, Lee and colleagues determined that the neuron-specific splicing factor NOVA1 [61] was underexpressed in FD versus control iPS cell-derived neural crest precursors [22]. The new model described in this study will allow us further test whether candidate splicing factors may be involved in the tissue-specific regulation of IKBKAP mRNA alternative splicing.

Materials and Methods

Ethics Statement

All control and FD participants gave informed and written consents (provided by the parents for the children) and biopsies were obtained under a protocol, which was approved by the local ethical committees in New York (Institutional Review Board of the New York University School of Medicine) and Marseille (Comité Consultatif de Protection des Personnes dans la Recherche Biomédicale Marseille 2).

Purification of hOE-MSCs

Human nasal mucosae were obtained from biopsies of 4 FD patients (3 females and 1 male aged 12–16 years) at the
Dysautonomia Treatment and Evaluation Center, New York. All four FD patients were homozygous for the splicing mutation. Biopsies form 5 healthy controls (3 females and 2 males, aged 19–39 years) were collected by the ENT Department in Marseille (Hôpital Nord, France). Biopsies were harvested as previously described [30,46] to obtain a cell culture of hOE-MSCs. The cells were continuously cultured in DMEM/HAM’S F12 (Gibco) supplemented with 10% fetal bovine serum (FBS) and 50 μg/ml gentamicin (Gibco) and trypsinized once a week with 0.05% trypsin-EDTA (Gibco) at 60–80% confluence. Cycloheximide (Sigma) was used to inhibit protein synthesis. For RNA extraction, and 24-well plates for immunostaining), in a concentration ranging from 25 to 200 μM for various incubation times, as specified in the text.

Generation of spheres and cell differentiation

Cells were plated at 15,000 cells/cm² into 6-well plates pretreated with poly-L-lysine (5 μg/cm², Sigma) in a serum-free medium of DMEM supplemented with insulin-transferrin-selenium (ITS, 1g/l insulin, 0.55 g/l transferrin, 0.67 mg/l sodium selenite; Gibco), epidermal growth factor (EGF, 50 ng/ml, R&D system) and basic fibroblast growth factor-2 (bFGF, 50 ng/ml, R&D system). Half of the medium was changed every 2 days. Multipotent spheres were obtained after 1 week and harvested by aspiration of the culture medium and centrifugation (5 min, 300g). The pellet, containing approximately 10⁶ cells, was resuspended in 0.5 ml 2× Laemml buffer (0.5 M Tris pH 6.8, 4.4 ml Glycerol, 20% SDS, 1% Bromophenol Blue, 0.5 ml β-mercaptoethanol). 50 μl of cell lysates were separated on 6.5% SDS-polyacrylamide gel electrophoresis and transferred to a nitrocellulose membrane (Amersham Biosciences). After blocking with 5% nonfat milk in PBS, 0.1%, tween 20 (PBST) buffer, blots were probed for 1 h at room temperature with a mouse monoclonal anti-IKBKAP antibody (1:5,000, BD Biosciences, clone 33) in PBST; followed by incubation with horseradish peroxidase-conjugated goat anti-mouse IgG (1:15,000, Jackson Immunoresearch) for 45 min at room temperature. For IKAP/hELP1 staining, primary antibody was incubation at 4°C. Vessel size was set to 58 nm in x and y and to 162 nm in z.

Western blot analysis

Cells were harvested by trypsinization and centrifugation (5 min, 300g). The pellet, containing approximately 10⁶ cells, was resuspended in 0.5 ml 2× Laemml buffer (0.5 M Tris pH 6.8, 4.4 ml Glycerol, 20% SDS, 1% Bromophenol Blue, 0.5 ml β-mercaptoethanol). 50 μl of cell lysates were separated on 6.5% SDS-polyacrylamide gel electrophoresis and transferred to a nitrocellulose membrane (Amersham Biosciences). After blocking with 5% nonfat milk in PBS, 0.1%, tween 20 (PBST) buffer, blots were probed for 1 h at room temperature with a mouse monoclonal anti-IKBKAP antibody (1:5,000, BD Biosciences, clone 33) in PBST; followed by incubation with horseradish peroxidase-conjugated goat anti-mouse IgG (1:15,000, Jackson Immunoresearch) for 45 min at room temperature. As a control, the membrane was also probed for β-actin (1:5,000, Sigma, clone D6C10). Proteins were visualized by chemiluminescent detection using the ECL detection kit (Enhanced Chemiluminescence, Amersham) and films were digitized and analyzed using the Bio 1D software.

RNA Isolation and semi-quantitative reverse transcription-polymerase chain reaction analysis

Total RNA was isolated using the RNaseasy Mini Kit (Qiagen) with DNase treatment on the column according to manufacturer’s recommendation. Total RNA was subjected to reverse transcription-RT using the High-Capacity cDNA Archive Kit (Applied Biosystems). End-point polymerase chain reaction (PCR) analysis was performed using the Go-Taq polymerase system (Promega) and IKBKAP-specific primers listed in Table 2. PCR products were separated on a 1.7% agarose gel by electrophoresis in 1× TBE buffer (Tris 0.89 M, boric acid 0.89 M and EDTA 0.02 M). DNA was visualized under UV light after ethidium bromide incorporation and documented using BioVision Camera.

Plasmid calibrators

A fragment of WT IKBKAP cDNA, containing exon 19-exon 21 and the 16 first nt of intron 22, was cloned into pcDNA 3.1 TOPO vector (Invitrogen) and named IKBKAP cal. Similarly, a piece of MU IKBKAP cDNA, containing exon 19-exon 21 and the 19 first nt of exon 22, was cloned into pcDNA 3.1 TOPO vector and named IKBKAP skipEx22cal. A piece of WT IKBKAP cDNA, containing the last 103 nt of exon 35-exon37-exon38 first 90 nt, was cloned into KpnI-XbaI cloning sites of pcDNA 3.1 TOPO vector and named IKBKAP skipEx32cal. A piece of WT IKBKAP cDNA, containing the last 30 nt of exon1-exon2-exon3 first 110 nt, was cloned into KpnI-XbaI cloning sites of pcDNA 3.1 TOPO vector and named IKBKAP fullEx32cal. For ABL1, the last 37 nt of exon 2 and first 102 nt of exon 3 were amplified from ABL1 cDNA, cloned into KpnI-XbaI cloning sites of pcDNA 3.1 TOPO vector and named ABL1 cal. All plasmid calibrators were linearized with XbaI and serially diluted in a

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solution of Tris 10 mM, EDTA 1 mM pH 8, containing 20 ng/ml of E. coli 16S and 23S rRNA (Roche).

Real-time PCR assay

The PCR reactions were performed in triplicate in a final volume of 25 µl, including 300 nM primers, 200 nM TaqMan® probe, 12.5 µl of TaqMan® universal PCR master mix (Applied Biosystems) and 5 µl of either cDNA or plasmid calibrator in an AB Prism 7900 HT thermocycler with 50 cycles and the protocol recommended by the manufacturer. For relative quantification and microarray results validation, we selected primer sets and probes, matching sequences present in the IMAGE human cDNA clones of the nylon microarrays with those displayed on the web portal of Applied Biosystems. The assay IDs were the following: Hs00375306_m1 (PMEPA1) and Hs00293488_m1 (S100A16) for the dysregulated genes in FD, and Hs01003267_m1 (HPRT1) and Hs00293488_m1 (RPLP0) for reference genes used to normalize the data. We also used previously validated primers and probe for ABL1 as a third reference gene [62]. Results were calculated using the 2(-ΔΔCT) method [63]. For absolute quantification, IKBKAP primers and hydrolysis probes (FAM TAMRA) were designed using the Primer 3 software and are listed in Table 2. Serial dilutions of plasmid calibrators (10^6, 10^5, 10^4, 10^3, 10^2 copies in 5 µl) were prepared and used to construct the standard curves. The number of IKBKAP and ABL1 transcripts was extrapolated automatically by the Sequence Detection System v2.2.2 software (Applied Biosystems).

Microarray analysis and normalization

RNA integrity was assessed using an Agilent 2100 Bioanalyzer (Palo Alto, CA). Samples with an RNA integrity number (RIN)<9 were excluded from the analysis (the samples concerned were C2P5, C3P5, and FD2P5). Gene expression analyses were carried out with cDNA Nylon microarrays containing 8,780 spotted cDNA clones and radioactive detection as previously described [64], with 5 µg of RNA reverse transcribed (oligo-dT priming) in presence of [α-33P]dCTP (Amersham Pharmacia Biotech). Details about microarray construction, clones list, probes preparations, hybridizations and washes have been previously described [65]. After image acquisition, signal intensities were quantified using BZScan software (http://tagc.univ-mrs.fr/bioinformatics/bzscan, [66]). A specific R library that uses the ‘S4’ system of formal classes and methods was used to process and normalize nylon microarray data [67]. Quantile normalization was applied to vector probe data (V) and complex probe data (C), to correct for global intensity and dispersion. Correction by the vector signal was made for each spot signal by calculating a C:V ratio before log transformation (base 2). No background correction or overshining correction was used. All data are MIAME compliant and have been loaded into ArrayExpress database (http://www.ebi.ac.uk/microarray-as/ae/) under accession number E-MTAB-281.

Statistical and gene ontology analysis

Significant Analysis of Microarray (SAM version 1.13; Stanford University) was applied to determine significant differential gene expression using the Multieperiment viewer (MEV) program. The data were analyzed using a two-class unpaired response type, which compared control samples versus FD samples. SAM calculated a significant score for each gene based on the gene expression change relative to the standard deviation of repeated values for that gene. We used 100 permutations and a

| Table 2. Sequence of primers used for end-point and TaqMan real-time PCR. |
|-----------------------------------------------|
| Primer Probe | Sequence | T (°C)* | Amplicon size (bp) | Splicing events |
|-----------------------------------------------|
| **End-point PCR** |
| hIKBKAP ex17-18F | TCAATAGACATTGAGGGTG | 55 | 446 (WT) | ex20 incl/skip |
| hIKBKAP ex22R | ATGATTCAAGAATCTATCTG | 57 | 372 (MU) | |
| hIKBKAP ex1F | CGCAGGCTAATCTTTGG | 60 | 485 | alt 3’ss for ex2 |
| hIKBKAP ex4-5R | TCAGGTTCTGTAAGCATCTG | 59 | 449 | ex36 incl/skip |
| hIKBKAP ex33-34F | TCCAGTAGATCAACGAGATC | 59 | 449 | ex36 incl/skip |
| hIKBKAP ex37R | GCTGATAAATGCGCTGATAC | 59 | 414 | ex36 incl |
| hIKBKAP ex35-36R | TGGGACCTAGAACCCTGT | 59 | 414 | ex36 incl |
| **Real-time PCR** |
| hELP1 ex19F | GGTTCCAGGATTGTCACTGGT | 60 | 133 | ex20 incl |
| hELP1 ex20-21R | ACATAAGGGTGGGCAACCCCTCC | 60 | 133 | ex20 incl |
| P-WTELP1 ex20R | AAGCAAGGGCTGGTATGAGAACA | 60 | 133 | ex20 incl |
| hELP1 ex19-21F | GGCACAAAGGCTTTATACACACTTAA | 60 | 133 | ex20 incl |
| hELP1 ex21-22R | CCACCTTGCCAGAACCCTTACT | 60 | 133 | ex20 incl |
| P-MUELP1 ex21F | AGAGGCAATTAGGCTGAGAAACGC | 60 | 133 | ex20 incl |
| hELP1 ex2F | CCGGAGTTCTAGGGCTGCT | 60 | 133 | ex20 incl |
| hELP1 ex2-3R | TTACCTCTCTGGAGACAGGGTCTAC | 60 | 133 | ex20 incl |
| P-WTELP1 ex2F | TCCGACTGAACGGGGACGTT | 60 | 133 | ex20 incl |
| hELP1 ex35-37F | TCAGTACCCGGTTCTAGGTT | 60 | 133 | ex20 incl |
| hELP1 ex38R | GGTTCTCCTGGTATGCTTGTG | 60 | 133 | ex20 incl |
| P-WTELP1 ex37-38R | AAGCTAGCATCTCAAGAACCAGAACCC | 60 | 133 | ex20 incl |

*Annealing temperature. doi:10.1371/journal.pone.0015590.t002
false discovery rate (FDR) of 3%. A total of 50 genes appearing in the heat map generation were called as significant with a p-value < 0.006. For gene ontology analysis, we generated a set of human protein associated with the gene appearing as significant with the SAM test, by using the BioKnowledge® Library (BKL) Retrieval™ search tool [http://www.biobase-international.com/]. This set of proteins was analyzed for overrepresentation of Gene Ontology (GO) Biological Process (BP) terms.

**Boydner chamber-based cell migration assay**

hOE-MSCs were detached by trypsin/EDTA, counted and seeded into the upper chamber of transwell polystyrene terephthalate filter membranes with 8 µm diameter pores (BD Biosciences), at a density of 3 × 10^4 cells/well, in a final volume of 200 µl serum or serum free culture medium, with or without 100 µM kinetin. Cells were allowed to migrate through the membrane filter for 24h at 37°C, 5% CO2. Cells migrating through the membrane pore and invading the underside surface of the membrane were fixed with 4% paraformaldehyde. Non-migratory cells on the upper membrane surface were removed with a cotton swab, and nuclei were stained with 0.5 µg/mL DNA intercalant Hoechst #33258. For quantitative assessment, the number of stained migrating cells was counted with imageJ software on 10 random fields per membrane filter at x20 magnification.

**Supporting Information**

**Figure S1** 50 genes are differentially expressed between control and FD hOE-MSCs. Heatmap representation of overexpressed (red) and underexpressed (green) genes in 5 controls and 4 FD OE-MSCs at passage 1, 2, 3, and 9. Normalized signal intensities were treated with the SAM software to highlight the most differentially expressed genes, with a FDR set at 3% (TIF).

**Figure S2 IKBKAP exon 36 inclusion increases after cycloheximide treatment.** NMD pathway was blocked by the translation inhibitor cycloheximide and results in an elevated expression of exon 36-including transcripts in 2 FD OE-MSC cultures (FD3 and FD4) as determined by absolute RT-qPCR. (* P<0.05). (TIF)

**Table S1** Dysregulated genes involved in other processes. (DOC)

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**Author Contributions**

Conceived and designed the experiments: NB FF FBA ECI. Performed the experiments: NB BL AB OS CFT ECI. Analyzed the data: NB ECI. Contributed reagents/materials/analysis tools: JG MK CN FF BFA. Wrote the paper: NB ECI.

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