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Cortical neurons obtained from patient-derived iPSCs with GNAO1 p.G203R variant show altered differentiation and functional properties

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ABSTRACT

Pathogenic variants in the GNAO1 gene, encoding the alpha subunit of an inhibitory heterotrimeric guanine nucleotide-binding protein (Go) highly expressed in the mammalian brain, have been linked to encephalopathy characterized by different combinations of neurological symptoms, including developmental delay, hypotonia, epilepsy and hyperkinetic movement disorder with life-threatening paroxysmal exacerbations. Currently, there are only symptomatic treatments, and little is known about the pathophysiology of GNAO1-related disorders. Here, we report the characterization of a new in vitro model system based on patient-derived induced pluripotent stem cells (hiPSCs) carrying the recurrent p.G203R amino acid substitution in $G\alpha_0$, and a CRISPR-Cas9-genetically corrected isogenic control line. RNA-Seq analysis highlighted aberrant cell fate commitment in neuronal progenitor cells carrying the p.G203R pathogenic

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variant. Upon differentiation into cortical neurons, patients' cells showed reduced expression of early neural genes and increased expression of astrocyte markers, as well as premature and defective differentiation processes leading to aberrant formation of neuronal rosettes. Of note, comparable defects in gene expression and in the morphology of neural rosettes were observed in hiPSCs from an unrelated individual harboring the same *GNAO1* variant. Functional characterization showed lower basal intracellular free calcium concentration ($[Ca^{2+}]_i$), reduced frequency of spontaneous activity, and a smaller response to several neurotransmitters in 40- and 50-days differentiated p.G203R neurons compared to control cells. These findings suggest that the *GNAO1* pathogenic variant causes a neurodevelopmental phenotype characterized by aberrant differentiation of both neuronal and glial populations leading to a significant alteration of neuronal communication and signal transduction.

1. Introduction

The *GNAO1* gene, encoding the alpha-o1 subunit of guanine nucleotide-binding protein (G α o), has been associated with Developmental and Epileptic Encephalopathies (DEE), a clinically and genetically heterogeneous group of neurological disorders with onset during infancy or childhood [1]. Pathogenic variants in the *GNAO1* gene cause encephalopathy affecting psychomotor development with high clinical heterogeneity, including developmental delay, intellectual disability, early-onset hyperkinetic movement disorders with severe exacerbations, epileptic seizures, and prominent hypotonia. Such variants were first described in patients with early infantile epileptic encephalopathy 17 (EIEE17, OMIM #615473) [1]. Later, *GNAO1* variants were reported in patients affected by neurodevelopmental disorders with involuntary movements, with or without epilepsy (NEDIM, OMIM #617493) [2]. Approximately 50 pathogenic variants have been reported so far in ClinVar (www.ncbi.nlm.nih.gov/clinvar/). Most of them are missense variants in highly conserved residues distributed across the entire length of the protein. Codons 203, 209, and 246 represent pathogenic variant hotspots, occurring in about 50% of cases. A different class of variants leading to *GNAO1* haploinsufficiency has recently been associated with a relatively mild and delayed-onset dystonia phenotype ([3–5]). Recently, the examination of several patients with distinct *GNAO1* variants resulted in a new clinical severity score [6]. Interestingly, this study suggested that each variant is associated with a specific severity score and unique disease mechanisms. Despite the increasing number of children diagnosed with *GNAO1*-related disorders, no effective cure or treatments are available.

 $G\alpha$ o is the most abundant membrane protein in the mammalian central nervous system, playing major roles in neurodevelopment [7] and neurotransmission [8]. In the brain, Go controls the synthesis of cAMP, modulating inhibitory and stimulatory inputs to the adenylyl cyclase 5 (ADCY5) [9]. Go also mediates inhibitory signaling downstream of several neurotransmitters, including dopamine, adenosine, and GABA, through inhibition of calcium channels ([10,11]) or by preventing neurotransmitter release [12]. Go is ubiquitously expressed in the central nervous system, but it is highly enriched in the cerebral cortex, hippocampus, and striatum [13].

Little is known about the molecular mechanism that triggers the neurodevelopmental phenotype in patients carrying pathogenic *GNAO1* variants. Since a link between *GNAO1* variants and rare DEEs was reported for the first time a decade ago [1], several animal models have been raised to understand the pathophysiology of *GNAO1*-related disorders. Knockout models suggest that *GNAO1* plays a pivotal role both in the adult brain and during early neurogenesis. Indeed, its expression increases during axonogenesis ([14,15]), and knockout mice show impaired neurogenesis [16], while *Gnao1* deficient Drosophila models show defects during axon guidance [17]. *GNAO1* disease-linked pathogenic variants in mouse and worm models lead to a loss of function in G α -mediated signaling with or without a dominant negative effect ([18–20]).

Despite currently available animal models, which include mouse, Drosophila and *C. elegans* ([18–24]), greatly contributing to a deeper understanding of this pathology, species-specific differences (e.g. neonatal lethality in p.G203R/+ mice; [22] make it hard to translate findings from animal models to humans. Hence the necessity for more reliable models. In this regard, human induced pluripotent stem cells (hiPSCs) represent a valuable tool for disease modeling and drug screening since they can be derived directly from patients and then differentiated ideally into any cell type of interest. Moreover, gene editing allows for the introduction of specific nucleotide substitutions in existing iPSC lines from healthy individuals, generating pairs of mutated/wild-type isogenic lines. To the best of our knowledge, there is only one hiPSC-based study of *GNA01*-related disorders [25]. The authors showed abnormal structure and aberrant neurite outgrowth in *GNA01*-KO spheroids [25]. They assessed a defect in lamination and found few neural rosettes and the loss of *PAX6* expression compared to wild-type (WT) cells. They confirmed some of these findings in patient hiPSCs carrying the p. G203R substitution. However, a major limitation of this study is the lack of a proper isogenic *GNA01* WT control for the patient-derived line. Comparison of lines carrying a given pathogenic variant with their isogenic counterpart, indeed, is crucial to distinguish between phenotypes due to the presence of that variant from those originating from different genetic backgrounds ([26–28]).

Here we report the characterization of a new hiPSC-based *in vitro* model of *GNAO1*-related disorders. We have generated cortical neurons from hiPSCs derived from a patient carrying the p.G203R amino acid change and performed molecular, cellular and functional studies at different time points during differentiation. Importantly, all experiments have been carried out in parallel with an isogenic *GNAO1* wild-type line generated by reverting the variant by CRISPR-Cas9-mediated gene editing. Compared to an isogenic control, *GNAO1* p.G203R neurons show altered expression of early and late key differentiation genes, altered timing of differentiation and an increased number of astrocytes, as well as functional impairment. In addition, the altered gene expression and differentiation of neuronal precursor cells was confirmed in a second hiPSC line derived from an unrelated individual harboring the *GNAO1* p.G203R

variant.

2. Results

2.1. RNA-seq analysis reveals impaired neuronal fate commitment in GNAO1 p.G203R neural progenitor cells

HiPSCs derived from an American patient and isogenic control hiPSCs generated by gene editing have been provided by Child's Cure Genetic Research (see Methods section). In this work, such lines are referred to as $GNAO1^{+/G203R}$ and $GNAO1^{+/G203G}$, respectively (Fig. 1A).

We first analyzed RNA-Seq data previously obtained from hiPSC-derived neural progenitor cells (NPCs; day 7 and day 11 of



Fig. 1. RNA-Seq analysis of neural progenitors **A.** Outline of the origin of the hiPSC lines used in this study. **B.** Heatmap showing the expression in GNAO1^{+/G203R} and GNAO1^{+/G203G} NPCs at day 7 and 11 of the leading edge genes identified via GSEA for the "cell fate commitment" Gene Ontology Biological Process category. Genes that were identified as leading edge ones only at day 7, only at day 11, or at both time points were split into three groups (Only_D7, Only D11, Common_D7-D11, respectively). Genes belonging to the "neuron fate commitment" or "neuron fate specification" categories were colored in purple. **C.** Heatmap showing the expression in GNAO1^{+/G203R} and GNAO1^{+/G203G} NPCs at day 7 and 11 of selected neuron progenitor- and reactive astrocyte-specific genes. The adjusted p-values displayed on the right pertain to the DGE analyses performed between GNAO1^{+/G203R} and GNAO1^{+/G203G} NPCs at day 7 (FDR D7) and day 11 (FDR D11). The expression values reported in both heatmaps correspond to row-scaled (Z-score), rlog-transformed count data.

Α



В







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Fig. 2. Cortical neuron differentiation and analysis at day 25. A. Schematic representation of the protocol used for cortical neuron differentiation of human iPSCs in conventional 2D cultures. NPC: neural progenitor cell; SB: SB431542 (TGF β type I receptor/ALK5 inhibitor); LDN: LDN193189 (BMP inhibitor); DAPT: Notch inhibitor; BDNF: Brain Derived Neurotrophic Factor; GDNF: Glial Derived Neurotrophic Factor; AA: ascorbic acid; cAMP: cyclic AMP. **B.** Phase contrast images of differentiating cells at day 25. **C.** Real-time qRT-PCR analysis of the expression of the indicated markers in differentiating cells. The graphs show the average and standard deviation (Student's *t*-test; paired; two tails; **p < 0.01, ***p < 0.001).

differentiation; n = 3 replicates from one differentiation batch). By performing differential gene expression (DGE) analysis, we detected 3451 downregulated genes and 3372 upregulated genes at day 7 and 4174 downregulated genes and 5081 upregulated genes at day 11 in GNAO1^{+/G203R} NPCs compared to GNAO1^{+/G203G} NPCs (adjusted *p*-value <0.01) (Supplementary Table S1). These results suggest that at early phases of neural induction the p.G203R variant has an extensive impact on gene expression, as also indicated by principal component analysis (PCA) (Supplementary Fig. S1A) showing that the transcriptional profile clearly segregates samples, exhibiting distinct patterns that can be attributed not only to their differentiation stage but also to their genotype. Such effects of the pathogenic variant on gene expression are particularly evident at day 11. We observed concordance between the sets of genes that exhibited deregulation at day 7 and day 11; however, the expression of several genes changed in opposite directions (Supplementary Fig. S1B). Gene Set Enrichment Analysis (GSEA) indicated that downregulated genes are enriched in those involved in cell fate commitment, many of which control neuronal differentiation (Fig. 1B). Among them, we noticed neural progenitor markers (PAX6, NES/NESTIN, EOMES/TBR2, CDH2, OTX1) and neuronal markers (CUX1), while astrocyte markers were mostly upregulated (Fig. 1C). In general, GSEA showed that deregulated genes are involved in many biological processes related to neural differentiation (Supplementary Fig. S2A). Specifically, we observed extensive downregulation of genes related to neurodevelopmental processes, such as "forebrain development", "neuron migration," "central nervous system", "neuron differentiation" and "hindbrain development". We also observed a downregulation of WNT pathway components [29], particularly on day 7, and to a somewhat lesser extent on day 11 (Fig. 1B; Supplementary Figs. S2B and C). Notably, this pathway regulates neurogenesis and gliogenesis in the cortex with a specific temporal gradient, and downregulation of WNT signaling has been shown to reduce the timing of neurogenesis and trigger precocious astrogenesis [30]. Further, $G\alpha o$ can act as a direct transducer of WNT-Frizzled signaling ([31,32]), reinforcing the importance of our findings that components of this pathway are downregulated in $GNAO1^{+/G203R}$ NPCs.

Collectively, this RNA-Seq data suggest that during early neurogenesis the *GNAO1* p.G203R pathogenic variant affects cell fate commitment, and in particular neuronal fate, causing the downregulation of genes implicated in neurodevelopment and, on the other hand, the upregulation of astrocyte markers.



Fig. 3. Immunostaining analysis at day 25 of differentiation A-B. Immunostaining analysis with the indicated primary antibodies and DAPI to label nuclei. PAX6 and NESTIN are neural progenitor markers; TBR2 is a neuronal precursor marker; TUJ1 and MAP2 are neuronal markers.

2.2. GNAO1 p.G203R hiPSCs show impaired neuronal differentiation

 $GNAO1^{+/G203R}$ hiPSCs, along with the isogenic $GNAO1^{+/G203G}$ control line, were differentiated into cortical neurons using a method previously described [33], based on dual SMAD inhibition followed by the inhibition of SHH signaling with cyclopamine to induce a neural cortical fate (Fig. 2A). This protocol recapitulates the steps of neural induction during embryonic development,



Fig. 4. Analysis of neuronal rosettes morphology in the independent $GNAO1^{+/G203R}$ #2 line **A.** Phase contrast images of $GNAO1^{+/G203R}$ #2 cells at the neural rosette stage, corresponding to day 30 of differentiation. Note that the in the protocol used for differentiation of $GNAO1^{+/G203R}$ #2 cells (as described in the Methods section) has a different timing from the one depicted in Fig. 2A. **B-D.** Immunostaining was performed using the specified primary antibodies along with DAPI for nuclear labeling. TBR1 is a marker of early-born neurons (B); and TUJ1 (B) and MAP2 (C) are markers for neurons; NESTIN serves as a marker for neural progenitors (C); β-catenin as a marker of apical polarization in neural rosettes (D). Number of individual cultures for experiments was 3.

including the formation of neural rosettes that mimic *in vitro* the forming neural tube. At day 25 of differentiation, when control cells showed the typical radial organization of neural progenitors in rosettes, GNAO1^{+//G203R}-derived cells did not acquire such morphology and showed a more advanced phenotype (Fig. 2B). Impaired timing of differentiation was confirmed by gene expression analysis



Day 40; n=3

100 µm

Fig. 5. Marker expression analysis at day 40 of differentiation A. Real-time qRT-PCR analysis of the expression of the indicated markers in differentiating cells. The graphs show the average and standard deviation (Student's *t*-test; paired; two tails; *p < 0.05, ***p < 0.001, n.s. nonsignificant). **B.** Immunostaining analysis with the indicated primary antibodies and DAPI to label nuclei.



С

GNAO1+/G203G

GNAO1+/G203R



Day 50; n=2

Fig. 6. Marker expression analysis at day 50 and 70 of differentiation A-B. Real-time qRT-PCR analysis of the expression of the indicated markers in differentiating cells at the indicated time points. The graphs show the average and standard deviation (Student's *t*-test; paired; two tails; *p < 0.05, $^{***}p < 0.001$, n.s. nonsignificant). C. Immunostaining analysis with the indicated primary antibodies and DAPI to label nuclei.

performed by qRT-PCR, showing altered levels of neural progenitor markers, such as *NESTIN*, *OTX2* and *PAX6* in patient-derived cells compared to the isogenic control (Fig. 2C), in agreement with RNA-Seq analysis at earlier time points (Fig. 1B). GNAO1^{+/G203R} cells also expressed lower levels of *TBR2* and *TBR1*, that in the developing neocortex are expressed in intermediate progenitor cells and postmitotic neurons, respectively [34], agreeing with the reduced numbers of TBR1-and TBR2-positive cells in the cortex of mice harboring pathogenic *GNAO1* variants [22]. These results were further confirmed by immunostaining analysis at the same time point, showing a reduced number of PAX6-, TBR2-and NESTIN-positive progenitors in patient-derived cells compared to the isogenic control (Fig. 3A and B).

Analysis of the neural rosettes generated from iPSCs collected from a Russian patient with the GNAO1^{+/G203R} variant (herafter GNAO1^{+/G203R}#2) showed similar disruption in the morphology of the neural rosettes. GNAO1^{+/G203R}#2 hiPSCs, along with the control cell line from a healthy donor, were differentiated into cortical neurons using a method previously described [35]. Neural rosette formation was examined on day 30 of differentiation in both control and GNAO1^{+/G203R}#2 cultures. The control cultures exhibited cellular rosettes formed by the accumulation of several tens to hundreds of cells, whereas the GNAO1^{+/G203R}#2 cultures showed clusters of only 10–20 cells (Fig. 4A and B). Immunocytochemical analysis of the neuronal markers TBR1, TUJ1 (Fig. 4B), and MAP2 (Fig. 4C) revealed the presence of positively stained cells at the edge of the neural rosettes in the control culture. In contrast, the number of these cells was significantly lower in the patient line-derived culture (Fig. 4B and C). Similarly, a marked reduction in NESTIN-positive progenitor cells was observed in the variant culture compared with the control (Fig. 4C). In addition, the variant-containing rosettes exhibited disrupted cytoarchitecture, a lack of cell polarization, and the absence of radial structure. Apical polarization of β -catenin was also impaired in the variant-containing culture (Fig. 4D).

Collectively, these data suggest an early impairment that affects *GNAO1* p.G203R neural progenitor cells. Importantly, morphologic abnormalities in neural rosettes were consistently observed in two independent patient-derived *GNAO1* p.G203R lines.



Fig. 7. Neurons carrying the p.G203R GNAO1 substitution exhibit lower values of basal $[Ca^{2+}]_i$ and a reduced spontaneous Ca^{2+} activity. **A.** representative spontaneous Ca^{2+} transients recorded in control cells. **B.** up, $[Ca^{2+}]_i$ basal values measured in individual cells. Please note the reduced basal $[Ca^{2+}]_i$ for p.G203R mutant neurons. Number of examined cells: 160, 159, 229 and 230, for p.G203G DIV40, p.G203R DIV40, p. G203G DIV50 and p.G203R DIV50, respectively. Bottom, percentage of neurons exhibiting spontaneous Ca^{2+} transients, same cells as upper panel. *, p < 0.001.



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Fig. 8. The p.G203R GNAO1 substitution reduces the response of differentiated neurons to several neurotransmitters in terms of $[Ca^{2+}]_i$ elevations. A. left, percentage of neurons exhibiting Ca^{2+} transients induced by the application of glutamate (1 mM, 3 s); inset, representative Ca^{2+} transients elicited by glutamate administration (1 mM, 3 s), in a control (black) and GNAO1 p.G203R (red) iPSC-derived neuronal cells. Number of examined cells: 160, 159, 229 and 230, for p.G203G DIV40, p.G203R DIV40, p.G203G DIV50 and p.G203R DIV50, respectively. Right, amplitude of the $[Ca^{2+}]_i$ increase induced by glutamate application in the responding cells. **B.** left, percentage of neurons exhibiting Ca^{2+} transients induced by the application of GABA (100 μ M, 3 s). Same cells as Fig. 7A. Right, amplitude of the $[Ca^{2+}]_i$ increase induced by GABA application in the responding cells. **C.** left, percentage of neurons exhibiting Ca^{2+} transients induced by the application of baclofen (100 μ M, 3 s). Same cells as Fig. 7A. Right, amplitude of the $[Ca^{2+}]_i$ increase induced by baclofen application in the responding cells. **D.** left, percentage of neurons exhibiting Ca^{2+} transients induced by the application of $[Ca^{2+}]_i$ increase induced by glutamate application in the responding cells. **D.** left, percentage of neurons exhibiting Ca^{2+} transients induced by the application of ACh (100 μ M, 3 s). Same cells as Fig. 7A. Right, amplitude of the $[Ca^{2+}]_i$ increase induced by glucine application in the responding cells. **E.** left, percentage of neurons exhibiting Ca^{2+} transients induced by the application of ACh (100 μ M, 3 s). Same cells as Fig. 7A. Right, amplitude of the $[Ca^{2+}]_i$ increase induced by ACh application in the responding cells. **F.** left, percentage of neurons exhibiting Ca^{2+} transients induced by the application of ACh (100 μ M, 3 s). Same cells as Fig. 7A. Right, amplitude of the $[Ca^{2+}]_i$ increase induced by ACP application in the respondi

2.3. The GNAO1 p.G203R substitution alters neuronal maturation

At day 40, corresponding to a stage in which differentiated cells have entered the neuronal maturation stage, $GNAO1^{+/G203R}$ derived neurons continued to express lower levels of *TBR1*, which highlights a reduction in the presence of layer VI neurons (Fig. 5A and B). On the other hand, they tended to express higher levels of *GFAP*, an astrocyte marker, compared to the isogenic control, and failed to downregulate *PAX6* expression, suggesting an increased amount of astrocyte and progenitor cells (Fig. 5A and B). Furthermore, the expression levels of *FOXG1*, a telencephalic gene that plays an important role in neural differentiation and in balancing the excitatory/inhibitory network activity, were increased (Fig. 5A). At later time points (day 50 and day 70), we confirmed significant *TBR1* downregulation in variant-containing neural cultures, whereas we observed an increased trend of upregulation for *GFAP* (at both day 50 and day 70), *PAX6* (day 50, not tested at day 70) and *FOXG1* (only at day 50), despite statistical significance not always being reached (Fig. 6A–C).

Collectively, these findings indicate that the *GNAO1* p.G203R variant leads to altered gene expression as well as premature and defective differentiation processes affecting both neurons and astrocytes.

2.4. Spontaneous and evoked calcium transients were reduced in GNAO1 p.G203R cortical neurons

Functional characterization via calcium imaging was performed on days 40 and 50. As shown in Figs. 7 and 8, this analysis suggests impaired functional maturation in neuronal cultures obtained from patient-derived GNAO1^{+/G203R} iPSCs. At both time points, $GNAO1^{+/G203R}$ -derived neurons showed decreased levels of basal intracellular free calcium concentration ($[Ca^{2+}]_i$) and a reduced fraction of neurons with spontaneous activity (Fig. 7A and B). In line with this finding, calcium-dependent signaling was affected in $GNAO1^{+/G203R}$ neurons at day 50, which displayed lower levels of pGSK3 β at the inhibitory Ser9 residue compared to control neurons (Supplementary Fig. S3). We then tested the response of control and variant-containing neurons to different neurotransmitters or selective agonists. The highest response levels were obtained, both in control and GNAO1^{+//G203R} derived neurons, upon application of glutamate (Fig. 8A), but with a significant decrease in the percentage (at day 40) and in the amplitude of the responses in $GNAO1^{+/}$ G^{203R}-derived neurons (at both time points). The most striking difference was detected upon stimulation with GABA (Fig. 8B). While the vast majority of wild-type neurons responded to GABA stimulation at both time points, this effect was almost completely abolished by the pathogenic variant. Moreover, the amplitude of the $[Ca^{2+}]_i$ increase was significantly lower even in the few GNA01^{+/G203R} cells that were able to respond to GABA. We also tested baclofen, a specific agonist of GABA_B receptors (Fig. 8C). In this case, we observed a statistically significant difference only in the number of responding cells at day 50. Reduced activity did not correlate with a decrease in GABBR2 (GABA_B Receptor 2) and GRIA4 (Glutamate Ionotropic Receptor AMPA Type Subunit 2) mRNA levels (Fig. 5A), pointing to functional impairment rather than decreased expression of receptors. Reduced activity in response to other neurotransmitters, such as glycine, acetylcholine, and ATP was also observed in terms of the number of responding cells (Fig. 8D-F). These results suggest that the GNA01 p.G203R variant leads to a delay in neuronal maturation, which in turn causes a profound impairment of neuronal function.

3. Discussion

In this study, we provide a side-by-side comparison of patient-derived hiPSCs and an isogenic control in which the p.G203R change has been reverted to WT (GNAO1^{+/G203G}). The highly conserved glycine 203 residue represents a pathogenic variant hotspot, associated with a mixed, severe phenotype characterized by epileptic seizures and movement disorders ([36,37]). The major clinical signs associated with the *GNAO1* p.G203R variant are neonatal/infantile-onset therapy-resistant epilepsy (generalized or focal), axial hypotonia, dystonia, and hyperkinetic movement disorders (chorea associated with paroxysmal events requiring intensive care). Together with glutamic acid 246 and arginine 209, this residue plays a crucial role in the allosteric regulation mediating $G\beta\gamma$ release [38]. The core molecular dysfunction for the three most common *GNAO1* pathogenic variants (p.G203R, p.R209C/H, and p.E246K) is the constitutive GTP-bound state of the mutant $G\alpha$ o proteins coupled to their inability to adopt the active conformation which allow proper binding to regulators and effectors [24]. Indeed, these three residues are located in the switch II or switch III regions, which are important for guanidine nucleotide-dependent regulation of downstream effectors.

In this work, the GNAO1^{+/G203R} hiPSC line from an American patient and the GNAO1^{+/G203G} isogenic control were differentiated

into cortical neurons using a method that recapitulates human corticogenesis. The comparison with the isogenic control shows that p. G203R is sufficient to impair the timeline of differentiation as early as at day 25, representing the onset of neuronal differentiation from neural progenitor cells. Similar impairment in the ability to form neural rosettes was consistently observed in a second $GNAO1^{+/}$ G^{203R} line derived from an unrelated Russian patient. This agreement confirms the importance of the hiPSCs *GNAO1* model in elucidating the pathogenesis of the disease and underscores the importance of further investigation of the role of this variant in the pathogenesis of *GNAO1*-related disorders. This neurodevelopmental phenotype is clinically relevant because, based on imaging, microcephaly, progressive and diffuse cerebral/cerebellar atrophy, hypomyelination/delayed myelination, and abnormalities of the basal ganglia (e.g., hypoplasia/atrophy of the caudate nucleus) or corpus callosum (thin corpus callosum) have been observed in the most severe cases harboring the p.G203R substitution [39].

Altered timing of proliferation and differentiation has been reported in many neurodevelopmental disease models, such as Fragile X syndrome ([40,41]), Tuberous Sclerosis Complex [42], and CDKL5 Deficiency Disorder [43]. The neurodevelopmental phenotype observed here is in line with previous observations from GNAO1 animal models. We have recently shown that there is a decrease in neural progenitor cells in two different mouse models of GNAO1-related disorders (p.C215Y and p.G203R), suggesting an impairment in neuronal differentiation or migration, specifically in the developing cerebral cortex [13]. We found enlargement of lateral ventricles and altered motor cortex thickness. We also reported a reduction of cell number in the total cortex, while no change was observed in the striatum and hippocampus. Accordingly, here we show that compared to the isogenic control line, human GNAO1 p.G203R differentiating neural progenitors are characterized by reduced expression of early neural genes, such as NESTIN, OTX2 and PAX6. Moreover, in line with the mouse model which has half of the number of *Tbr2*-positive cells in the ventricular and subventricular zones and has a lower cell density in layer VI of the cortex (tbr1-positive cells) [22], here we found decreased levels of both TBR1 and TBR2 transcripts in the iPSC-derived neurons. Altered expression of FOXG1, which was significantly upregulated at day 40 and shows the same trend at later time points, is intriguing. Indeed, variants in the FOXG1 gene have been linked to a rare form of epileptic encephalopathy (OMIM #613454) [44]. TBR1 is a master regulator of cortex development that plays a crucial role in the formation of neocortical layer VI, regulating both neuronal migration and synaptic development ([45,46]). Pathogenic variants in TBR1 gene are associated with cortical malformations, intellectual disability and autism spectrum disorder. Furthermore, it has been shown that TBR1 regulates neural stem cell fate by inhibiting astrocytes formation [47]. Interestingly, we report increased expression of GFAP, an intermediate progenitor marker expressed in astrocytes, suggesting that GNAO1 pathogenic variants could have an effect both on neuronal and glial populations. GFAP overexpression has been found both in neurodevelopmental and neurodegenerative diseases, such as Fragile X syndrome and Alzheimer's disease ([33,48-51]). Moreover, abnormal astrocytes proliferation linked to altered timing of neurogenesis and astrogenesis has been observed in RASopathies and Down Syndrome, further suggesting the role of astrocytes on the onset of neurodevelopmental diseases [52]. Increased GFAP expression is also considered as a hallmark of reactive astrocytes, which are astrocytes that underwent morphological and functional modification as a consequence of brain injury or disease [53]. Notably, since prolonged astrogliosis contributes to the epileptic phenotype [54] and astrocytes alteration has been observed in different DEE models ([55–57]), astrocytes are considered a target for therapeutic approaches in epilepsy [58]. Our present findings suggest that this might be extended to GNAO1-related disorders as well.

Our functional analysis supports the concept that the *GNAO1* p.G203R substitution leads to a lower degree of neuronal maturation. Control neurons exhibit sustained spontaneous $[Ca^{2+}]_i$ activity, which is significantly reduced in GNAO1^{+/G203R}-derived cells, in association with a reduction of the basal $[Ca^{2+}]_i$ level. As a probable consequence of the decrease in $[Ca^{2+}]_i$, we found lower levels of pGSK3 β (S9) in day 50 differentiated p.G203R neurons compared to control cells, which is in line with previous findings in neuronal cultures with knockout for the *GNAO1* gene or carrying the *GNAO1* p.G203R variant [25]. Considering the key role of Wnt-GSK3 signaling in the regulation of axon extension and branching [59], further studies are needed to explore the impact of aberrant GSK3 β phosphorylation in p.G203R neuronal morphogenesis. Furthermore, GNAO1^{+/G203R}-derived neurons are less responsive to several agonists, indicating a lower functional level of different crucial receptor systems. In particular, both GABA and glutamate induce a much lower response, suggesting a profound impairment of the neuronal function in Day 40 and Day 50 differentiated p. G203R neurons compared to control cells. Reduced sensitivity to neurotransmitters, and in particular GABA, is observed in several epileptic disorders [60]. The functional impairment observed in p.G203R neurons (loss of spontaneous Ca²⁺ transient activity, lower responsiveness to most neurotransmitters) strongly suggests a slower maturation of the molecular machinery necessary for proper synaptic connections, likely leading to dysfunctional cortical and subcortical circuitry, and consequently to the typical clinical signs, such as drug-resistant neonatal epilepsy and hyperkinetic movement disorders.

The hiPSC-based model described here will be used as an *in vitro* platform for translational studies. Both the neurodevelopmental phenotype and the functional impairment represent possible readouts to evaluate the effects of promising new drug candidates, such as caffeine and zinc ([19,24]), and new gene therapy approaches, such as those based on recombinant adeno-associated virus (rAAV)-mediated delivery of genetic constructs encoding WT *GNAO1*, as well as short hairpin RNA (shRNA) to suppress the endogenous gene [61]. A major limitation of the present work, however, lies in the absence of additional patient-derived hiPSC lines. Future investigations will be imperative to determine whether the phenotypes observed in p.G203R cells are replicated in the presence of different *GNAO1* pathogenic variants.

In summary, our study solidifies the understanding of GNAO1 encephalopathy as a neurodevelopmental disorder, shedding light on the specific molecular and functional disruptions caused by the p.G203R variant. Our findings illuminate critical aspects of impaired neurogenesis, altered neuronal differentiation, and dysfunctional responses in affected neural cells. While this research delves into the early stages of neurodevelopment, further investigations into the variant's impact on fully matured neurons are imperative. Additionally, exploring a broader spectrum of *GNAO1* pathogenic variants associated with diverse clinical features will enhance our comprehension of this complex disorder. These avenues of study not only promise to deepen our understanding of *GNAO1*-related

neurological conditions but also pave the way for targeted interventions and therapeutic strategies.

4. Materials and Methods

4.1. Reprogramming of patient-derived hiPSCs and generation of the gene edited GNAO1^{+/G203G} control line

GNAO1^{+/G203R} iPSCs derived from an American patient and the isogenic control line were kindly provided by the nonprofit organization Child's Cure Genetic Research (Fremont, CA, USA). Briefly, fibroblasts were derived at the Stanford Children's Hospital (CA, USA) from a skin punch biopsy of a GNAO1^{+/G203R} patient. The patient (male, born May 2017, 2 years old at the time of biopsy) began to exhibit seizures at the age of 6 weeks and epilepsy was controlled with anti-epileptic drugs. Overtime seizures transformed to infantile at the age 4 months and he was treated with adrenocorticotropic hormone (ACTH) and vigabatrin. Seizures have been in remission until the age of 6 years old. Patient exhibited global development delay, weak muscle tone, movement disorders (chorea and dystonia that were more pronounced with fever) and limited communication skills through visual eye gaze device. Reprogramming to iPSCs, followed by standard characterization, was performed by Applied StemCell Inc. The isogenic control line was generated using CRISPR/Cas9-mediated genome editing by the Genome Engineering & Stem Cell Center (GESC@MGI) at Washington University in St. Louis (https://GeneEditing.wustl.edu).

 $GNAO1^{+/G203R}$ #2 hiPSCs were generated by reprogramming of skin fibroblasts from a patient from Moscow with GNAO1 encephalopathy carrying the c.607 G > A (p.Gly203Arg) variant, as described in Ref. [61], which were kindly provided by Dr. Bardina (Institute of Gene Biology, Russian Academy of Sciences, Moscow) upon the proper written patient consent. The patient was born in October 2016 (male, 1 year old at the time of biopsy). From day 7 after birth, epileptic seizures were noted in the form of bilateral tonic seizures. Remission of seizures was achieved at 5 months of age upon treatment with levetiracetam and vigabatrin. In addition to seizures, the patient presented with a complex set of symptoms, including psychomotor developmental delay, choreiform hyperkinesia, and dystonic hyperkinesia. To generate control fibroblasts, a skin biopsy was obtained from a healthy 26-year-old female donor after informed consent. The biopsy was cut into small pieces and cultured under the coverslip in DMEM (PanEco, Russia) with 10% FBS (Hyclone, USA), 2 mM Glutamine (PanEco), 1X penicillin-streptomycin (PanEco). Medium was changed once in three days. After seven days, fibroblasts were passaged with 0.25% Trypsin (Gibco). Human skin fibroblasts from the patient and the healthy donor were reprogrammed using the protocol described in Shuvalolva et al., 2020 [62].

4.1.1. RNA-sequencing and bioinformatic analysis

FASTO files from paired-end RNA-Seq analysis performed using the Illumina® Stranded Total RNA Prep kit with Ribo-Zero Plus treatment on GNAO1^{+/G203R} and GNAO1^{+/G203G} hiPSC-derived NPCs at day 7, generated at Rarebase PBC, and day 11, generated as previously described in Whye et al. (2023) (basic protocol 1 and support protocol 3) [63]. For each time point, 3 replicates from one differentiation batch were used. Raw RNA-Seq datasets are available at the NCBI Sequence Read Archive (SRA) with the accession number PRJNA991628. Forward and reverse reads were aligned to a database of rRNA sequences using Bowtie 2 v2.4.4 [64] with -nofw and -nofw parameters, respectively; only read pairs in which both mates did not map to these abundant molecules were retained for downstream analyses. STAR v2.7.10b [65] with default mapping parameters was used to align reads to the human GRCh38 genome, providing the GENCODE 42 [66] GTF file to generate a splice junction database and to calculate gene-level read counts with the -quantMode TranscriptomeSAM GeneCounts option. Read counts reported in the fourth column of ReadsPerGene.out.tab files were combined into a count matrix using a custom Python script. This matrix was used to calculate normalized expression values, expressed as FPKM (Fragments Per Kilobase of exon per Million mapped reads), using the "fpkm" function from the DESeq2 v1.34.0 R package [67]. We retained only protein-coding or lncRNA genes with FPKM >0.5 in at least three samples for subsequent analysis. Prior to performing PCA using the DESeq2 "plotPCA" function, we applied regularized-log (rlog) transformation to the count data. To identify genes which are differentially express (DEGs) between $GNAO1^{+/G203R}$ and $GNAO1^{+/G203G}$ samples at both time points, Wald test with independent filtering was performed using the "DESeq" function. Apeglm method [68] was employed for log2(fold change) shrinkage. The FDR-adjusted p-value threshold to select DEGs was set to 0.01. Additionally, in both contrasts, we only selected genes that had an average FPKM greater than 0.5 in at least one of the two conditions being compared. The intersections between the DEGs identified in the two comparisons were visualized using the UpSetR v1.4.0 R package [69]. Gene expression heatmaps were generated with the ComplexHeatmap v2.10 R package [70] using row-scaled (Z-score), rlog-transformed count data. The WebGestaltR v0.4.4 R package [71] was employed to perform Gene Set Enrichment Analysis (GSEA), using shrunken log2(fold change) values as a ranking metric.

4.2. HiPSC culture and differentiation

 $GNAO1^{+/G203R}$ iPSCs were cultured in Nutristem-XF (Biological Industries) supplemented with 0.1% Penicillin-Streptomycin (Thermo Fisher Scientific) onto Biolaminin 521 LN (Biolamina) functionalized plates. The culture medium was refreshed every day and cells were passaged every 4–5 days using 1 mg/mL Dispase II (Thermo Fisher Scientific). Cells were routinely tested for mycoplasma contamination. $GNAO1^{+/G203R}$ hiPSCs were differentiated into cortical neurons as previously described [33].

 $GNAO1^{+/G2O3R}$ #2 hiPSCs were cultured in mTeSRTM1 medium (Stem Cell Technologies) on Matrigel (Corning)-coated plates following the manufacturer's guidelines. Every day, the culture medium was renewed, and cells were passaged every 4–5 days up to 70–85% confluency utilizing 0.5 mM EDTA (Thermo Fisher Scientific). On the first day post-subculturing, cells were cultured in medium supplemented with 5 μ M of the ROCK inhibitor StemMACSTM Y27632 (Miltenyi Biotec). The differentiation of hiPSCs into midbrain neurons was performed as previously described [35]. The differentiation of $GNAO1^{+/G203R}$ #2 hiPSCs into mature human neurons comprised three main stages [1]: hiPSC differentiation into neuronal progenitor cells (NPCs) [2]; expansion and maturation of NPCs; and [3] differentiation of NPCs into mature neurons. In brief, hiPSCs were differentiated into NPCs using the commercially available PSC Neural Induction Medium (Thermo Fisher Scientific) following the manufacturer's guidelines. This was considered as day 0 (D0). After 10 days of induction to NPCs, cells were detached from the substrate with 0.5 mM EDTA and were plated at density of 250,000–400,000 cells per cm² in Petri dishes or multi-well plates coated with Matrigel. Further cultivation took place in the medium for expansion and maturation of NPCs consisting of advanced DMEM/F12 (Thermo Fisher Scientific), Neurobasal Medium (Thermo Fisher Scientific), 2% Neural Induction Supplement (GIBCO). The ROCK inhibitor Y27632 5 μ M was used for every seeding overnight with the following replace to the medium without ROCK inhibitor. After 18 days the differentiation protocol for maturation into neurons was initiated. Neural precursors were cultured and frozen until the third passage. We reseeded neural precursors to a density of 300,000–400,000 cells per cm². NPCs were seeded into wells coated with Matrigel and cultured in the medium containing DMEM/F12 (Thermo Fisher Scientific), 1X GlutaMAX (Thermo Fisher Scientific), 2% B27 (Thermo Fisher Scientific), 1X Penicilin-Streptomicin (Thermo Fisher Scientific), BDNF 20 ng/mL (GenScript), GDNF 20 ng/mL (GenScript), 200 μ M ascorbic acid (Sigma Aldrich), 4 μ M forskolin (Sigma Aldrich) for 2 weeks with subculturing during the first week. The ROCK inhibitor was used every passage overnight with the following replacement to the medium without the ROCK inhibitor.

4.2.1. Real-time PCR

RNA was extraction and Real-time RT-PCR was performed as described in Brighi et al., 2021 [33]. A complete list of primers, including the housekeeping control gene *ATP50*, is provided in Supplementary material (Supplementary Table S2).

4.2.2. Immunostaining

GNAO1^{+/G203R} differentiating cells were immunostained as described in Brighi et al., 2021 [33] using the following primary antibodies: mouse anti-PAX6 (sc81649 Santa Cruz Biotechnology, 1:50), mouse anti-GFAP (MAB360 Merck Life Sciences, 1:500), chicken anti-MAP2 (ab5392 Abcam, 1:2000), rabbit anti-β-TUBULIN III (TUJ1) (T2200 Merck Life Sciences, 1:2000), Rabbit Anti-Nestin (MA5-32272 Thermo Fisher Scientific, 1:100), Rabbit Anti-TBR2 (ab15894 Millipore,1:50). AlexaFluor secondary antibodies (Thermo Fisher Scientific) were used at the concentration of 1:250, and DAPI (Merck Life Sciences) was used to stain nuclei. An inverted Olympus iX73 microscope equipped with an X-Light V3 spinning disc head (Crest Optics), a LDI laser illuminator (89 North), using 405 nm, 470 nm, 555 nm and 640 nm wavelengths, depending on the fluorophores, a Prime BSI sCMOS camera (Photometrics) and MetaMorph software (Molecular Devices) with a 20x objective (Olympus), was used for image acquisition.

GNAO1^{+/G203R}#2 cells at the stage of neuronal rosettes (30 day of differentiation) were fixed in 4% paraformaldehyde solution (Merck Life Sciences) for 15 min at room temperature and washed thrice with 1X PBS (Thermo Fisher Scientific). Fixed cells were permeabilized for 1 min using ice-cold PBS supplemented with 0.1% Triton X-100, blocked for 30 min with PBS supplemented with 1% BSA. Cells were then incubated overnight at 4 °C with primary antibodies at the following dilutions: rabbit anti-MAP2 (ab254264 Abcam, 1:2000), mouse anti- β TUBULIN III (TUJ1) (T8660 Sigma-Aldrich, 1:500), mouse Anti-Nestin (ab6142 Abcam, 1:250), rabbit anti-TBR1 (ab183032 Abcam,1:200), mouse anti- β -Catenin (571–781 BD Transduction Laboratories, 1:200). The day after, the primary antibody solution was washed out and the cells were incubated with secondary antibodies (Alexa Fluor 488 or Alexa Fluor 594 conjugates, Jackson ImmunoResearch, 1:250) for 2 h at room temperature in the dark. DAPI (Merck Life Sciences) was used for nucleus staining. Images were acquired with an LSM710 confocal microscope.

4.3. Western blotting

Differentiating cells were lysed with RIPA Buffer with freshly added protease and phosphatase inhibitors (Thermo Fisher Scientific), and kept on ice for 30 min. Afterwards they were centrifuged at $12.000 \times g$ (20 min, 4 °C), the pellet was discarded and the supernatant transferred into new tubes, and stored at -80 °C. Protein content was determined using the BCA protein Assay (Thermo Fisher Scientific). Protein extracts (20 µg) were resuspended in Novex Tris-Glycine SDS Sample Buffer (Thermo Fisher Scientific) containing Sample Reducing Agent, heated for 5 min at 85 °C, and separated on precast Novex Tris-Glycine Mini Gels. Proteins were transferred using iBlotTM 2 on a PVDF membrane, which was then blocked in 5% nonfat dry milk (Biorad Laboratories) in T-TBS for 1h at RT. Membranes were then incubated O/N with anti-Phospho-Gsk3 β (Ser9 - D85E12) rabbit monoclonal antibody (1:1.000, cat#5558, Cell Signaling), anti-GSK-3 β (3D10) mouse monoclonal antibody (1:1.000, cat#9832, Cell Signaling), and anti-GAPDH (6C5) mouse monoclonal antibody (1:500, cat# sc-32233, Santa Cruz Biotechnologies) in T-TBS with 5% BSA. After incubation, membranes were washed three times in T-TBS for 10 min at RT, and incubated for 1h with anti-mouse horseradish peroxidase-conjugated antibodies (1:10.000, cat#115-035-003, Jackson Immunoresearch Laboratories) at RT. Reactivity was detected using Western Bright ECL spray device (cat#K-12049-D50, Advansta, Aurogene) and Fluorchem E (ProteinSimple). Densitometric analysis was performed using the ImageJ software (http://rsb.info.nih.gov/ij/).

4.4. Calcium imaging

Changes in free intracellular Ca²⁺ concentration $[Ca^{2+}]_i$ were quantified by time-resolved digital fluorescence microscopy using the Ca²⁺ indicator Fura-2 (excitation 340 nm and 380 nm, emission 510 nm). The changes of $[Ca^{2+}]_i$ were expressed as R=F340/F380. iPSCs-derived neuronal cells were incubated with the cell-permeant Fura-2 acetoxylmethylester (2 μ M; Molecular probes, Life

technologies) for 1 h at 37 °C in culture medium. Ca^{2+} responses were elicited by administering the neurotransmitter or the agonist dissolved in normal external solution for 3 s. Cells were continuously perfused during the experiment. $[Ca^{2+}]_i$ variations were obtained separately from each individual cell, using the MetaFluor 7.0 software (Molecular Devices, USA).

Amplitude values were expressed as means \pm S.D. and analyzed using one-way ANOVA test. When necessary, the non-parametric Dunn's one-way ANOVA on ranks was used. In case of significance, all pairwise multiple comparison procedure was used (Holm-Sidak, or Dunn's method for non-parametric tests). Responsive cells data were expressed as % and analyzed using chi-square test. The minimum power of statistical tests was set at 0.8. The significance for all tests was set at p < 0.05.

Data availability

The RNA-Seq data from $GNAO1^{+/G203G}$ and $GNAO1^{+/G203R}$ NPCs at day 7 and day 11 are available at the NCBI Sequence Read Archive (SRA) with the accession number PRJNA991628.

CRediT authorship contribution statement

Maria Cristina Benedetti: Writing – review & editing, Methodology, Investigation, Formal analysis, Conceptualization. Tiziano D'Andrea: Writing – review & editing, Methodology, Investigation, Formal analysis. Alessio Colantoni: Writing – review & editing, Methodology, Formal analysis. Denis Silachev: Investigation, Formal analysis. Valeria de Turris: Methodology. Zaira Boussadia: Methodology. Valentina A. Babenko: Methodology. Egor A. Volovikov: Methodology. Lilia Belikova: Methodology. Alexandra N. Bogomazova: Methodology. Rita Pepponi: Methodology. Dosh Whye: Methodology. Elizabeth D. Buttermore: Writing – review & editing, Supervision. Gian Gaetano Tartaglia: Supervision. Maria A. Lagarkova: Supervision. Vladimir L. Katanaev: Writing – review & editing, Supervision, Conceptualization. Ilya Musayev: Resources. Simone Martinelli: Writing – review & editing, Supervision, Conceptualization. Sergio Fucile: Writing – review & editing, Supervision, Conceptualization. Alessandro Rosa: Writing – review & editing, Writing – original draft, Supervision, Project administration, Conceptualization.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.heliyon.2024.e26656.

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