Role for the Na\(^+\)/K\(^+\) ATPase pump alpha 3 (ATP1A3) subunit in folding and lamination of the human neocortex

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Osmotic equilibrium and membrane potential in animal cells depend on concentration gradients of sodium (Na\(^+\)) and potassium (K\(^+\)) ions across the plasma membrane, a function that is catalyzed by the Na,K-ATPase alpha subunit. In vertebrates, four paralogous genes, \textit{ATP1A1-4}, encode distinct alpha subunit isoforms (\(a_1-a_4\)), three of which (\(a_1, a_2, a_3\)) are expressed in the brain, and two (\(a_1, a_3\)) predominantly in neurons. The \(a_3\) isoform, encoded by \textit{ATP1A3}, is critical to neuronal physiology, and a growing spectrum of neurological diseases are associated with \textit{ATP1A3} pathogenic variants, with ages of onset ranging from early childhood to adulthood. Here, we describe \textit{ATP1A3} variants encoding dysfunctional \(a_3\) subunits in children affected by polymicrogyria, a developmental malformation of the cerebral cortex characterized by abnormal folding and laminar organization. To gain cell-biological insights into the spatiotemporal dynamics of prenatal \textit{ATP1A3} expression, we established a transcriptional atlas of \textit{ATP1A3} expression during cortical development using mRNA \textit{in situ} hybridization and transcriptomic profiling of \(~125,000\) individual cells with single-cell RNA sequencing (Drop-Seq) from various areas of the midgestational human neocortex. We find that fetal expression of \textit{ATP1A3} is restricted to a subset of excitatory neurons carrying transcriptional signatures of neuronal activity and maturation characteristic of the developing subplate. Furthermore, by performing Drop-Seq on \(~52,000\) nuclei from four different areas of an infant human neocortex, we show that \textit{ATP1A3} expression persists throughout early postnatal development, not only within excitatory neurons across all cortical layers, but also and more predominantly in inhibitory neurons, with specific enrichment in fast-spiking basket cells. In addition, we show that \textit{ATP1A3} expression, both in fetal and postnatal neurons, tends to be higher in frontal cortical areas than in occipital areas, in a pattern consistent with the rostro-caudal maturation gradient of the human neocortex. Finally, we discover distinct
co-expression patterns linking catalytic α subunit isoforms \((ATP1A1,2,3)\) and auxiliary isoforms \((ATP1B1,2,3)\), suggesting the ATPase pump may form non-redundant, cell-type specific α-β combinations. Together, the importance of the developmental phenotypes and dynamic expression patterns of \(ATP1A3\) point to a key role for \(a3\) in the development and function of human cortex.

**Keywords**
Na,K-ATPase, \(ATP1A3\), Cortical Development, Cortical Malformation, Developmental Channelopathy, DYT12, EIEE, Subplate, Polymicrogyria
GLOSSARY

- **ATP1A3**: Na⁺,K⁺-ATPase alpha 3 subunit
- **PMG**: Polymicrogyria (an overfolded cerebral cortex)
- **MCD**: Malformation of cortical development
- **EN**: Excitatory neuron
- **IN**: Inhibitory neuron
- **NPC**: Neural progenitor cell
- **SP**: Subplate
- **Vm**: Resting membrane potential
- **WPC**: Weeks post conception
- **AHC**: Alternating hemiplegia of childhood
- **RDP**: Rapid-onset dystonia-parkinsonism
- **CNS**: Central Nervous System
INTRODUCTION

Generating a folded, 6-layered cerebral cortex relies on the integration of signals from a variety of cell types, coordinating schedules of cell proliferation, migration, differentiation, and maturation. These processes require proper maintenance of ionic gradients and cell membrane potential (Vm) and thus rely on the precise function of ion channels and pumps. Pathogenic mutations in genes encoding ion channels can cause malformations of cortical development (MCDs), collectively known as developmental channelopathies (Smith and Walsh, 2020). For example, pathogenic variants in prenatally expressed genes encoding glutamate (GRIN2B, GRIN1) and sodium channels (SCN3A) can lead to MCDs (Fry et al., 2018; Platzer et al., 2017; Smith et al., 2018), at least in part via gain-of-function, cell-autonomous effects in neurons and potentially other cell types (Smith et al., 2018). These developmental channelopathy variants in glutamate and sodium channels cause pathogenic increases in cationic flux into cells (Smith and Walsh, 2020), affecting Vm in neural progenitor cells and neurons, which has been shown to be sufficient to disrupt neurogenic and cortical lamination processes in experimental models (Hurni et al., 2017; Vitali et al., 2018).

Neuronal physiology, homeostasis, and signaling depend on the electrogenic activity of the Na,K-ATPase, an ionic pump powered by ATP hydrolysis which maintains Na\(^+\) and K\(^+\) gradients across the plasma membrane. The Na,K-ATPase complex contains a large catalytic subunit (α) and two auxiliary subunits (β, FXYD) (Holm et al., 2016). In vertebrates, four paralogous genes (ATPIA1-4) encode α subunits, of which the ATPIA3 isoform is predominantly expressed in the brain and is critical to the restitution of electrochemical gradients following neuronal excitation, among several other physiological functions (Kim et al., 2004; McGrail et al., 1991; Picton et al., 2017). Pathogenic variants in ATPIA3 have been increasingly associated with a spectrum of
neurological diseases, with phenotypes ranging broadly in age of onset, from the immediate postnatal period throughout adulthood (Brashear et al., 1993; Smedemark-Margulies et al., 2016), suggesting different vulnerabilities to ATP1A3 dysfunction during childhood brain maturation. These diseases include (in approximate order of onset in childhood development): early infantile epileptic encephalopathy (EIEE); alternating hemiplegia of childhood (ACH); cerebellar ataxia, areflexia, pes cavus, optic atrophy, and sensorineural hearing loss (CAPOS); childhood onset schizophrenia (COS); and rapid-onset dystonia-parkinsonism (RDP) (Brashear et al., 1993; Smedemark-Margulies et al., 2016). Variants in ATP1A3 associated with these postnatal diseases are generally categorized as heterozygous loss-of-function single nucleotide variants (Arystarkhova et al., 2019; Holm et al., 2016) which, among other deficits, have been shown to lead to depolarized Vm, such as in neurons from AHC patients (Simmons et al., 2018). During development, bioelectric changes can dramatically alter both intrinsic trajectories of individual cells (Levin et al., 2017) and cortical lamination more broadly (Hurni et al., 2017; Vitali et al., 2018), suggesting that bioelectric alterations in early brain development could in part contribute to subsequent ATP1A3-related neurological disorders.

Here, we present four individuals with novel de novo ATP1A3 variants resulting in polymicrogyria (PMG, the most common MCD (Jansen et al., 2016)), epilepsy, and global developmental delay, and consider a previously described individual case report of AHC with polymicrogyria (Gurrieri et al., 2016). The ATP1A3 variants we observed associated with PMG clustered in transmembrane regions 7 to 8 of the α3 protein. Using mRNA in situ analysis and single-cell RNA sequencing (Drop-Seq) in the human fetal cortex at mid-gestation, we show that ATP1A3 is restricted to a subset of deep-layer neurons located in the subplate. Moreover, using Drop-Seq in the human infant cortex, we show that postnatal expression of ATP1A3 at this stage
of development extends not only to all excitatory neuron subtypes (with cortical layer biases), but mostly predominates within cortical interneurons. The infant cortex also displays a rostral-to-caudal $ATPIA3$ gradient, with highest expression levels in the prefrontal cortex. Last, within individual cells, variable levels of ATPase $\alpha$ and $\beta$ subunit isoforms offers differential vulnerability between excitatory and inhibitory neurons, and may provide a partial explanation for the broad phenotypic range associated with $ATPIA3$ mutations.
RESULTS

Individuals with $ATP1A3$ variants display cortical malformation phenotypes

Magnetic resonance imaging (MRI) of four unrelated individuals with de novo $ATP1A3$ variants revealed a range of PMG severity, from extensive bilateral frontoparietal PMG to unilateral PMG (Figure 1A). Detailed descriptions of each case are included in supplemental text.

Case A: Bilateral Frontoparietal PMG (p.Arg901Met)

Case A, a female child of nonconsanguineous parents from Portugal, was born at full term and had a short postnatal hospital stay for respiratory distress and jaundice. Brain MRIs performed at 5 months and at 9 years of age revealed extensive bilateral perisylvian polymicrogyria involving the lateral temporal lobes, insulae, and posterior half of frontal and parietal lobes (Figure 1A). The posterior part of frontal and temporal cortex appears abnormally thick, with somewhat shallow sulci, giving a pachygyric appearance (bilateral and fairly symmetrical, but with a mild left to right predominance). The most anterior aspects of the frontal and parietal lobes, as well as the medial and inferior cerebral cortex, were spared. At 9 years she developed seizures, manifesting as focal impaired awareness, and was treated with oxcarbazepine. She was developmentally and cognitively delayed, able to sit alone and walk with a walker, point and make some signs but had no speech, and could hold a pencil but not write. Asymmetric spastic quadriparesis (left more significant than right) with hypertonia and athetoid movements of the hands and fingers were noted on examination, as were a high palate, thin upper lip and smooth philtrum. Her head circumference at 9 years was 51 cm (-0.73 standard deviations, SD), and she exhibited drooling and had feeding difficulties. Trio whole exome sequencing (WES) revealed a de novo $ATP1A3$ variant, c.2702G>T (p.Arg901Met), absent in both parents (Figure 1 & S1).
Case B: Extensive Bilateral PMG (c.2921+1G>A)

Case B, a male child of nonconsanguineous parents from the Philippines, was born at 38 weeks’ gestation. His head circumference at birth was 34 cm (-0.83 SD) and he developed seizures 12 hours after delivery. Brain MRI at 6 weeks revealed extensive bilateral polymicrogyria, and small, cyst-like areas were noted within the hippocampal heads and bodies. Video EEGs confirmed a severe epileptic encephalopathy resistant to essentially all antiepileptic drugs. EEGs at 2 months of age showed abundant electroclinical and electrographic seizures beginning predominantly from the right parasagittal region but also from the left and bilateral parasagittal regions. Visual and auditory evoked potential studies were normal at 4 and 5 months of age respectively. Evaluation at 14 months revealed microcephaly, with a head circumference of 41.5 cm (-4.63 SD), severe global developmental delay (never rolled over, sat up, or talked), intermittent nystagmus, significant axial and appendicular hypotonia, and hyporeflexia. Trio WES revealed a de novo ATP1A3 variant, c.2921+1G>A, disrupting a conserved splice site, absent in both parents (Figure 1A & S1).

Case C: Unilateral PMG (p.Leu924Pro)

Case C, a male child of nonconsanguineous parents from Brazil, was born at 38 weeks’ gestation by Cesarean section for fetal distress. He was hypotonic and noted to have a left clubfoot at birth and was admitted to neonatal intensive care when episodes of upward rolling of the eyes were noted on the first day of life. MRI revealed extensive cortical malformation of the right hemisphere with polymicrogyria comprising the frontal, parietal, and temporal right lobes and the insula, and a small focus of signal abnormality in the periventricular white matter, compatible with recent ischemic injury. EEG revealed the presence of electroclinical seizures and his neurological
exam at the time disclosed episodic dystonic posturing of the upper left extremity alternating with excessive movements of closing his left hand with thumb adduction. He was discharged on phenobarbital and levetiracetam. Upon neurological exam at 8 months of age, he had a head circumference of 42.5 cm (-1.75 SD), very poor eye contact, global hypotonia with present reflexes, absent head support, absence of rolling and of object gripping, and presence of babbling sounds. Trio WES revealed a de novo ATP1A3 variant, c.2771T>C (p.Leu924Pro), absent in both parents (Figure 1 & S1).

Case D: Extensive multifocal PMG (p.Gln851Arg)

Case D, a male child of nonconsanguineous parents of European descent from the United States, was born at 41 weeks by emergency Cesarean section; the pregnancy was complicated by fetal hydronephrosis. He developed electroclinical seizures and a neonatal encephalopathy requiring cooling and was discharged after 1 month. MRI soon after birth showed multifocal PMG involving both cerebral hemispheres, right greater than left, with the right hemisphere smaller than the left (Figure 1A). At 4 months of age he was admitted for congenital hip dysplasia surgery and noted to have frequent breath holding spells that worsened after the surgery. His varied seizure phenomenology included nystagmus, chin quivering, and hand shaking, which continue intermittently even through phenobarbital, levetiracetam, and oxcarbazepine combination therapy. At 15 and 30 months of age, head circumference was 42.7 cm (-3.5 SD) and 44.2 cm (-3.3 SD), respectively. Physical exam at 18 months demonstrated spastic hemiparesis involving primarily left arm, but including ipsilateral face, and central hypotonia. Other symptoms include feeding difficulties and persistent constipation. WES revealed a de novo ATP1A3 variant, c.2552A>G (p.Gln851Arg), absent in mother (father was not available for testing).
PMG-associated \textit{ATPIA3} mutations are predicted to disrupt function

\textit{ATPIA3} encodes the P-type Na,K-ATPase $\alpha$3 subunit (NM_152296), an integral membrane protein that relies on ATP hydrolysis to transport ions ($\text{Na}^+$ and $\text{K}^+$) via 10 membrane-inserted helices (TM1-TM10). This ATPase is functionally supported by the $\beta$ and FXYD subunits (Holm et al., 2016). \textit{ATPIA3} has several conserved domains across $\alpha$ isoform paralogs ($\alpha1$-$4$) and orthologous genes (Clausen et al., 2017), and is a highly constrained gene, with many fewer missense and loss of function variants seen in the general population than predicted (gnomAD Z score of 6.33 and pLI score of 1.0 respectively). Taken together with \textit{ATPIA3}'s known disease association with AHC, RDP, IEE, and CAPOS (Brashear et al., 1993), this suggests the newly identified variants in \textit{ATPIA3} are likely damaging (Table S1).

PMG-associated \textit{ATPIA3} variants (p.Gln851Arg, p.Leu888Pro, p.Arg901Met, p.Leu924Pro, and c.2921+1G>A), occur at sites that are highly conserved among the $\alpha$ subunit paralogs (\textit{ATPIA1} – \textit{ATPIA4}) and invariant in other vertebrate \textit{ATPIA3} orthologs (Figures S1B).

\textit{In silico} pathogenicity predictions for p.Gln851Arg, p.Arg901Met, p.Leu924Pro, and c.2921+1G>A suggest these variants are likely deleterious and disease causing (Table S1). Two amino acid substitutions reported here, Leu924Pro and Gln851Arg, were previously associated with EIEE and AHC without MCD (Arystarkhova et al., 2019; Masoud et al., 2017), further supporting the wide spectrum of \textit{ATPIA3} phenotypes resulting from single amino acid variants (Table S3). Leu924Pro pathogenicity was also demonstrated to affect ATPase trafficking to the plasma membrane, categorized functionally as loss-of-function (LOF), or viewed as a toxic gain-of-function effect (Arystarkhova et al., 2019). Together, the conservation of nucleotides and absence of variants in population databases suggests these PMG \textit{ATPIA3} alleles are likely
pathogenic.

PMG-associated missense ATP1A3 variants (p.Gln851Arg, p.Leu888Pro, p.Arg901Met, p.Leu924Pro) cluster within TM7 to TM8 segments. The de novo ATP1A3 variant in Case A, NM_152296.4:c.2702G>T, p.Arg901Met (R901M), affects an arginine in the extracellular loop connecting transmembrane domains 7 and 8 (TM7-TM8; Figures 2A, S1A), in the same segment as a previously described variant (p.Leu888Pro) occurring in an AHC individual with PMG(Gurrieri et al., 2016). Arg901 and Leu888 amino acids are in close proximity to Glu899, Gln904, and Gln905, which interact with the Na,K-ATPase β subunit(Shinoda et al., 2009) (Figure 2B). Case C possesses a de novo ATP1A3 variant, NM_152296.4:c.2771T>C, (p.Leu924Pro), altering an amino acid in the TM7 helix just upstream of the loop (Figures 2B). Case D possesses a de novo ATP1A3 variant, NM_152296.4:c.2552A>G, p.Gln851Arg, altering an amino acid in TM7 (Figure 2B). Although the clustering of these missense variants may be seen as suggesting disruption of specific protein-protein interactions, on the other hand, the de novo ATP1A3 variant in Case B, NM_152296.4:c.2921+1G>A, abolishes the essential splice donor site of exon 21 (Figure S1), and similar ATP1A3 splice site variants are known to cause AHC(Viollet et al., 2015). Therefore, the PMG-associated variants may all represent LOF, with the more severe phenotype reflecting more complete LOF than for some other ATP1A3-associated conditions.

ATP1A3 mRNA expression in the human fetal neocortex

To investigate the transcriptional trajectory of ATP1A3 during human brain development underlying the prenatal phenotypes described in this study, we first contrasted the temporal expression patterns of ATP1A1-3 across brain regions, mining bulk neocortical transcriptome data from the Allen Brain Atlas(Jones et al., 2009), ranging from 12 weeks post conception (wpc) to adulthood (40 year-old) (Figure 3). We find that prenatal expression of ATP1A3 is higher than
that of its paralogs, and that ATP1A3 persists postnatally (Figure 3B). To gain further insights into the cell-biological mechanisms underlying the developmental cortical phenotypes described in this study, we characterized the patterns of ATP1A3 expression in the midgestational fetal human neocortex (wpc 19-21). This developmental age offers a uniquely informative window into corticogenesis as many and diverse biological processes – including neurogenesis, neuronal migration and differentiation, and cortical folding – co-occur at this age. At this stage, early-born excitatory neurons (ENs) have settled and started to differentiate within the innermost strata of the cortical plate (CP) (prospective layers 6a-5), while later-born ENs progressively migrate past their predecessors to occupy the most superficial aspect of the CP (prospective layers 4-2)(Miller et al., 2014). A large fraction of upper-layer ENs are still being generated in the germinal zones, while others – en route to the CP – are withheld within the subplate (SP), a transient developmental layer crucial for corticogenesis, where ENs are primed by thalamic input and form transient synaptic connections with one another and with resident SP neurons (prospective layer 6b)(Luhmann et al., 2018).

To investigate ATP1A3 expression in its tissue context, we performed RNA in situ hybridization (ISH) in a coronal cross-section of fetal neocortex at 19 wpc. ATP1A3 mRNA was virtually undetectable in the germinal zones (ventricular and subventricular zone, VZ and SVZ) but abundant in postmitotic neuronal layers, with highest expression in the inner CP and SP (Figure 3C & D). Multiplexed fluorescent ISH analysis of ATP1A3 with markers of two populations of cortical neural progenitors cells, radial glia and intermediate progenitor cells, confirmed ATP1A3 is not expressed in these dividing cells or their immediate progeny in either the VZ or the SVZ (Figure 3C). In addition, since ATP1A3-associated diseases display a range of cerebellar deficits (Brashear et al., 1993; Sweadner et al., 2016; Uchitel et al., 2019), we also
Smith et al., evaluated ATP1A3 expression in the fetal cerebellum (19 wpc), demonstrating an exclusive enrichment of ATP1A3 within the Purkinje cell layer (Figure 3E).

**Fetal neocortex single cell analysis demonstrates robust ATP1A3 enrichment to resident subplate neurons and rostral neocortical ATP1A3 enrichment**

Despite the high degree of compartmentalization of cell types into laminae, biological processes in the developmental neocortex are highly dynamic, and cellular compartments are heterogeneous in cell-type composition. Single-cell RNA sequencing enables large-scale, simultaneous comparisons of gene expression across thousands of individual cells, thus bridging functional data from human genetics, immunohistochemistry, and cell-type-specific biology (Macosko et al., 2015). To this end, we performed Drop-seq on 125,942 individual cells from a 21 wpc human neocortex (Figure 4A & S2A). Given the general trend of perisylvian to frontal cortex localization of the PMG phenotypes described in this study, we sought to investigate ATP1A3 expression across presumptive neocortical areal domains. To build our single-cell expression dataset, we collected cells from 11 adjacent sections spanning the dorso-medial and rostro-caudal axes of the neocortex, validating and refining the positional identity of these serial samples using the graded expression of cortical patterning genes as benchmarks (Figure 4B & S2)(Sun et al., 2005). Using unsupervised statistical methods (see materials and methods), cells were partitioned into four transcriptionally distinct clusters, corresponding to the major neural cell classes in the midgestational neocortex: excitatory neurons (EN), inhibitory neurons (IN), neural progenitor cells (NPCs), and glia (Figure 4C & S2C). We excluded from this analysis all non-neural cell types as our scope of inquiry was specifically regarding ATP1A3 dysfunction in neurons, and non-neural expression of ATP1A3 is limited (Figure 3B & S2). Within each of the
main neuronal cell classes we could further resolve transcriptionally distinct cell subtypes and states, which we interpreted by the distinctive enrichment of marker genes in each group (Figure S2C). Among ENs, consistent with the developmental landscape of mid-gestational corticogenesis described above, we could identify newborn and migratory upper-layer ENs at different stages of differentiation (clusters EN.0-3), and post-migratory deep-layer ENs expressing marker genes of maturing layer 6-5, including markers of resident SP neurons (EN-4, Figure 4C). Across cortical regions, ATP1A3 expression was virtually undetectable in the NPC clusters, low in INs and glia, more highly expressed in ENs (Figure 4C, S2C & D), and – among ENs – highest in the EN-4 cluster.

The EN-4 cluster was composed of cells sourcing from all cortical regions, and was marked by enriched expression of SP-specific genes (CRYM and LPL, Figure 4C), as well as by the expression of genes associated with neuronal maturation (e.g. axonal genes NEFL/NEFM and synaptic genes SEMA3E). This is consistent with the notion that the earliest-born neurons (a subset of which will persist in L6b in the adult cortex) settle and mature early in the developing SP (Hoerder-Suabedissen and Molnár, 2013). Within this cluster, ATP1A3 appeared to be differentially expressed across cortical region, with highest expression levels found in cells from frontal samples, intermediate levels in cells from medial samples, and lowest levels in cells sampled from more caudal cortical regions (Figure 4C & S2A). In agreement with this, expression of ATP1A3 in the EN-4 cluster was found to be enriched in neurons co-expressing SP markers and frontal cortex markers, but much less so in neurons expressing caudal cortex markers (Figure 4D, S2 & S3).

Cortical development proceeds along a rostral-to-caudal maturation gradient, with asynchronous growth patterns accounting for the differential expansion of frontal and caudal
cortical regions (Rakic, 1974). This implies that, at any given time, neurons in frontal regions are on average more mature than their caudal counterparts. To investigate whether the observed \( ATP1A3 \) enrichment in the EN-4 cluster might reflect a differential maturation state of EN neurons along the rostrocaudal axis, we performed pseudotime analysis of \( ATP1A3 \) across all ENs. To do this, we used the Monocle3 algorithm, an established analytical tool allowing single-cell transcripomes to be ordered and connected along a branched topology based on the differential expression of variable genes (Cao et al., 2019). By selecting as an “origin” cell, this algorithm allows projection of a “pseudo-differentiation” trajectory onto the network topology (Camp et al., 2015). This analysis revealed a positive correlation between \( ATP1A3 \) expression and EN differentiation, suggesting \( ATP1A3 \) may be required during neuronal differentiation, as neurons transition to a mature state, most readily observed in resident subplate neurons (Figure 4E & F).

Finally, to identify potential isoform-specific and cell-type specific vulnerabilities to \( ATP1A1-3 \) variants during corticogenesis, we compared the expression patterns of the \( ATP1A1-3 \) paralogs across cell clusters (Figure S3A). We find that expression of \( ATP1A2 \) is restricted to NPCs (and specifically to radial glia), and is virtually undetectable in neurons – in an almost mutually-exclusive pattern to \( ATP1A3 \). \( ATP1A1 \) expression more closely mimicked \( ATP1A3 \) expression, albeit its expression in NPCs was higher, and partially overlapped \( ATP1A2 \) (Figure S3A). Accordingly, pairwise correlation analysis of the three isoforms across clusters revealed that expression of \( ATP1A1 \) and \( ATP1A3 \) correlates more strongly with the EN-4-SP cluster, while expression of \( ATP1A1 \) and \( ATP1A2 \) correlates more highly with NPC clusters (Figure S3B). Across all cells, pairwise correlation analysis indicated a positive correlation, though moderate, between \( ATP1A1 \) and \( ATP1A3 \), but weak-to-no correlation between \( ATP1A1 \) and \( ATP1A2 \), and between \( ATP1A2 \) and \( ATP1A3 \) (Figure S3C). Together, this analysis suggests that neurons...
transitioning within the fetal SP may be particularly susceptible to \textit{ATP1A3} variants, with potential broad consequences for cortical lamination. Moreover, the single cell variable expression of ATPase isoforms could underlie compensation or vulnerability mechanisms for disease haploinsufficiency or differential penetrance within specific cell populations.

\textbf{ATP1A3 enrichment in parvalbumin interneurons and excitatory neurons in infant neocortex}

Given that \textit{ATP1A3} expression persists in the postnatal neocortex (Figure 3B), and that several known \textit{ATP1A3} mutations show postnatal onset of symptoms (Brashear et al., 1993; Smedemark-Margulies et al., 2016), we profiled \textit{ATP1A3} expression in the infant neocortex. Using Drop-seq, we profiled 51,878 nuclei from a ~8-month-old human cortex, sampled from prefrontal, temporal, occipital, and parietal lobes (Figure 5A). Unsupervised hierarchical clustering sorted nuclear transcriptomes into 3 major classes – ENs, INs, and glia – subdivided into 24 transcriptionally distinct cell classes (Figure 5A & S4). Within the EN clade, clusters stratified largely by layer and neuronal subtype, rather than cortical area, with several clusters enriched in gene markers of deep layer 5 and 6 (Figure 5B, See supplemental S4 for full clading). Within the IN clade, cells stratified by developmental origin (Caudal or Medial Ganglionic Eminence) and major “cardinal types”, according to previous studies (Mayer et al., 2018). In the infant neocortex, like in the fetal neocortex, \textit{ATP1A3} was also enriched in neurons compared to glia, but unlike in the fetal dataset, \textit{ATP1A3} was more highly expressed in INs than in ENs (Figure 5B & S5). Among neuronal clusters, and across cortical regions, \textit{ATP1A3} expression was fairly variable, with lowest expression in V1/occipital cortex cells, and wide range of expression levels within deep and upper layer EN clusters (Figure 5B). No specific enrichment was found in L6b ENs (i.e. the postnatal
remnant of the fetal SP layer), suggesting that fetal enrichment of ATP1A3 in the SP is transient, with maturation and migration of cells continuing to populate the upper layers. Of note, the lack of specific enrichment in postnatal SP may reflect the fact that, unlike the fetal dataset, the postnatal dataset was built from nuclear RNA rather than whole cell RNA, so that some transcripts may be under-detected, primarily those localized primarily in the somato-dendritic compartment.

Across INs clusters, expression of ATP1A3 was highest in cells sampled from the frontal lobe (prefrontal BA10), and more strikingly so within Parvalbumin (PV+) INs (Figure 5B & S4C). This differential expression pattern raised the hypothesis that ATP1A3 may be required dynamically across neuronal types, cell-biological processes, and across developmental ages. To gain insights into the nature of the cell-biological signature associated with ATP1A3 expression, we compared the top-100 genes correlating with ATP1A3 in EN and IN clusters. Interestingly, we found that 50/100 ATP1A3-correlating genes were shared across cell types, suggesting that ATP1A3 partakes in similar transcriptional programs, and thus potentially similar function, in different neuronal types. In both EN and IN, among the top genes correlating with ATP1A3, was FXDY6, the FXYD subunit of the ATPase complex (Figure 5D). We did find, however, that the ATPase β subunit beta subunit ATP1B1 correlated with ATP1A3 only in ENs, and ATP1B2 in INs, suggesting that, at least in part, ATP1A3 may form cell-type specific ATPase complexes in different neuronal classes (Figure 5D). Gene Ontology Enrichment analysis of the top 100-genes most strongly correlated with ATP1A3 in both ENs and INs revealed significant over-representation of gene products associated with transmembrane ionic transport and the somatodendritic compartment, including synaptic vesicles (Table S2). This is consistent with the established role of ATP1A3 in synaptic transmission and membrane potential, and suggests
ATP1A3 may partake in a shared transcriptional network for which ATP1A3 variants may perturb aspects of neuronal physiology central to many cell types.

Given the association of ATP1A3 pathological variants with phenotypes affecting higher-order cognitive functions with no clear counterpart in animal models, it was of interest to investigate the degree to which ATP1A3 enrichment in INs may be conserved among humans, non-human primates, and rodents. We took advantage of a recently published single-cell RNAseq dataset profiling gene expression in homologous IN types in humans, non-human primates (macaques and marmosets) and mice (Krienen et al., 2019). We found that relative ATP1A3 expression levels across IN types, both in the neocortex and in the striatum, are largely conserved across species, with highest expression in the fast spiking PV+ INs (Figure S6). This suggests that the observed differential expression of ATP1A3 across neuronal types is genetically encoded and evolutionarily conserved.
DISCUSSION

Mutations in \textit{ATP1A3}, the gene encoding the Na,K-ATPase \(\alpha_3\) catalytic subunit, have been associated with a wide spectrum of neurological diseases affecting childhood development, ranging in onset age from early infancy to late adolescence into adulthood. Here, we describe \textit{ATP1A3} pathogenic variants in individuals presenting with PMG, a severe prenatal malformation of the cerebral cortex characterized by abnormal gyrification and laminar organization. These case studies extend the known phenotypic range of \textit{ATP1A3}-related diseases, and provide new insights into the molecular pathology of PMG and PMG-associated developmental channelopathies. Furthermore, by compiling a single-cell atlas of \textit{ATP1A3} expression in the fetal and early postnatal neocortex, we revealed cell-type-specific \textit{ATP1A3} expression in the fetal and infant human brain, establishing a spatiotemporal roadmap for future genotype-phenotype associations studies, and studies of \textit{ATP1A3} function in the subplate and cortical interneurons. \textit{ATP1A3} associated conditions comprise a broad spectrum of overlapping phenotypes, including the blending of encephalopathies with more paroxysmal phenotypes, with a complex genotype-phenotype correlation. Further confounding the genotype to phenotype relationship, identical amino acid substitutions can impart variable presentations (\textit{Table S3}); for example, Leu924Pro and Gln851Arg reported here have been previously associated with EIEE and AHC without MCDs(Arstarkhova et al., 2019; Masoud et al., 2017), but with postnatal microcephaly (Leu924Pro), suggesting pathogenic susceptibility differs between individuals. Differential penetrance between individuals may reflect \(\alpha_3\) ATPase expression levels, competition between disease and wild-type \(\alpha_3\) subunits for utilization within Na/K-ATPases(Arstarkhova et al., 2019), and as described in this study, variable expression of \(\alpha_3\) (and coexpression with other \(\alpha\) and \(\beta\) subunits) within different cell populations (\textit{Figure 4D, S3 & S5}). Individuals with heterozygous
ATP1A3 variants presented here with PMG also exhibited a range of phenotype severity, from unilateral to extensive bilateral PMG, variable degrees of developmental delay, postnatal microcephaly, ataxia-like phenotypes and EIEE. ATP1A3’s wide phenotypic spectrum is similar to known glutamate and sodium channel “developmental channelopathies”, in which a subset of cases present with EIEE and developmental delay, both with and without MCDs and postnatal microcephaly (Smith and Walsh, 2020). Taken together, ATP1A3 PMG-associated variants likely represent a continuum of LOF features, with the more severe phenotype reflecting more complete LOF than for some other ATP1A3-associated conditions. In a future opportunity to examine cortical tissue of PMG associated with ATP1A3 variants, it would be useful to document cortical architecture, including whether gaps occur in the pia mater and glia limitans. This anatomical defect permits continuity between the molecular zones of adjacent microgyri for synaptic excitatory short-circuitry leading to epilepsy, as with other genetic mutations in PMG (Jansen et al., 2016).

Functional analysis of ATP1A3 variants associated with postnatal diseases are largely described as LOF; including decreased membrane localization in Leu924Pro, causing toxic effects within cells (Arystarkhova et al., 2019). LOF mechanisms described for ATP1A3 disease-causing variants include changes in Na⁺ affinity, impaired conformational changes, voltage-dependence shifts, and disruption of protein expression (Arystarkhova et al., 2019; Heinzen et al., 2014; M. Li et al., 2015). Since the Na,K-ATPase exports Na⁺, and elevated Na⁺ can be deleterious to cellular processes (Azarias et al., 2013; Toustrup-Jensen et al., 2014), α3 dysfunction that results in a shift in Na⁺ affinity could trigger a known developmental pathophysiology associated with increased intracellular Na⁺ concentration, leading to PMG (Smith et al., 2018; Smith and Walsh, 2020; Zaman et al., 2020). While some ATP1A1 variants have been described to increase inward leak of
Na’ (Azizan et al., 2013; Kaneko et al., 2014), similar pathogenic gain of function effects have not been described in ATPIA3 related diseases. In addition to acute ATPase pump dysfunction, PMG pathophysiology could also reflect an accelerated activation of toxic downstream mechanisms that underlie postnatal ATPIA3-related disorders causing neurodegeneration or cell death, including postnatal microcephaly (Paciorkowski et al., 2015), regional neuronal loss (Oblak et al., 2014), and cerebellar hypoplasia (Sweedner et al., 2016; Uchitel et al., 2019). Moreover, two of the PMG-causing ATPIA3 variants we describe are located in the TM7-TM8 extracellular segment of the α3 that interacts with the β-subunit (Shinoda et al., 2009) and Leu924Pro demonstrates β subunit associated LOF membrane localization properties (Arystarkhova et al., 2019), suggesting the final formation of α-β complex could potentially underlie the pathophysiology. The β-subunit supports localization of the α-subunit to the plasma membrane and modulates affinity of Na+ and K+ for the ATPase (Blanco et al., 1998). Moreover, α- and β- subunits can trigger several intracellular signaling pathways associated with MCDs, including MAPK and AKT signaling via phosphoinositide 3-kinase (PI3K) and EGFR pathways, resulting in changes in cell polarity, growth, motility and gene expression (Z. Li and Langhans, 2015).

During embryonic brain development, bioelectric cellular properties are required for the propagation of key organizing signals, directly implicating ion channels and pumps in a variety of morphogenetic processes (Levin et al., 2017). For instance, in mice and ferrets, experimental modulations in membrane conductance and Na+-dependent excitation have been linked to neuronal fate specification and directed migration (Hurni et al., 2017; LoTurco et al., 1995; Smith et al., 2018; Vitali et al., 2018). The fetal SP represents a hub for the spatiotemporal integration and transduction of signals from many cellular sources. Early electrical activity patterns in the SP accompany the dynamic formation of transient synaptic networks between resident SP neurons...
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502 and migratory ENs and INs en route to the CP, ascending radial glia processes, and ingressing thalamic fibers\textsuperscript{,23}. Our finding that (i) *ATP1A3*-expressing cells are located in the developing SP (ISH, Fig 3D), and (ii) co-express SP-specific marker genes (DropSeq, Fig 4 and S2), is therefore consistent with a role for *ATP1A3* in regulating electric activity within this compartment. This is consistent with the notion that human SP neurons possess large Na\textsuperscript{+} currents with prolonged depolarized states\textsuperscript{,23}(Moore et al., 2009), and therefore require a stout mechanism to recover Na\textsuperscript{+} and K\textsuperscript{+} electrochemical gradients following excitation, a known key function of the \( \alpha_3 \) ATPase in postnatal neurons\textsuperscript{,24}(Kim et al., 2004; Picton et al., 2017; Vaillend et al., 2002). In addition, the finding that *ATP1A3*-expressing cells are also found in the inner aspect of the CP (ISH, Fig 2C), express L6-5 EN markers (DropSeq, Fig4C), together with the rostral-to-caudal gradient of *ATP1A3* expression and positive correlation with neuronal pseudo-age, (Monocle, Fig 4F), all are consistent with a role of *ATP1A3* in the maturation of early-born deep-layer ENs.

The observed differential prenatal expression of Na\textsuperscript{+},K\textsuperscript{+},ATPase isoforms parallels the enrichment pattern reported for specific glutamate and sodium channel subtypes implicated in developmental channelopathies. These glutamate and sodium channel diseases maintain robust RNA expression of developmental subtypes; specific channel subtypes are utilized for developmental electrical functions (*SCN3A, GRIN2B*), whereas others are used for adult electrical functions (*SCN1A, GRIN2A*)\textsuperscript{,25}(Smith and Walsh, 2020). The \( \alpha_3 \) isoform may have a similar specialized function in fetal brain neurons compared to \( \alpha_1 \) isoforms. For example, Na\textsuperscript{+} affinity is lower in \( \alpha_3 \) than \( \alpha_1 \) isoforms\textsuperscript{,25}(Azarias et al., 2013; Dobretsov and Stimers, 2005), enabling \( \alpha_3 \) containing ATPase pumps to respond to larger Na\textsuperscript{+} influxes. Lower Na\textsuperscript{+} affinity provides a wider range to buffer aberrant Na\textsuperscript{+} flux, therefore could provide an evolutionarily advantageous neuroprotective effect in cells during development. Known developmental channelopathy genes

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also maintain a robust cell type and subtype expression profiles as ATP1A3; for example, SCN3A and GRIN2B are enriched in excitatory neurons and some progenitor types (Bagasrawala et al., 2017; Smith et al., 2018; Smith and Walsh, 2020), before transitioning to mostly GRIN2A-containing glutamate receptors and SCN1A in the postnatal cortex (Bagasrawala et al., 2017; Smith and Walsh, 2020). Intriguingly, the observed sc-RNA-seq ATP1A2 enrichment in NPCs could offer a pathological basis for ATP1A2 variants associated with fetal or early life demise, including microcephaly (Chatron et al., 2019; Monteiro et al., 2019).

In the postnatal cortex, the observed ATP1A3 enrichment in interneurons likely supports physiological processes necessary for maintaining inhibitory tone of cortical circuits (Z. Li and Langhans, 2015), a proposed mechanism underling EIEE, or perhaps AHC. We were surprised to find that even after 90 million years of divergence, the human and rodent postnatal cortex possess similar single-cell distribution of ATP1A3, most notably in PV+ INs, underscoring the utility of mice as a model for studying excitatory/inhibitory balance (Bøttger et al., 2011), especially in the context of PV IN loss relating to postnatal brain disorders (Hunanyan et al., 2018). On the other hand, while ATP1A3 expression in INs appears to be conserved, differences in circuits and connectivity, and how interneurons shape gene expression differentially could offer pathological differences between species. For example, the observed layer specific rostral enrichment of ATP1A3 could contribute an early susceptibility for ATP1A3 variants associated childhood onset schizophrenia (Chaumette et al., 2018; Smedemark-Margulies et al., 2016). Intriguingly, the identified ATP1A3 enriched cell types in the pre- and postnatal cortex (subplate and PV+ neurons), and Purkinje cells of the cerebellum, are all characterized by their robust firing properties (Moore et al., 2009; Tremblay et al., 2016). Within these cell types with dynamic physiological needs, the low affinity α3 would help maintain ionic gradients following excitation and suggests neurons with
higher neurophysiological demands would be most susceptible to Na⁺,K⁺,ATPase dysfunction. Additionally, even within the cortical pyramidal neuron population, differential Na⁺,K⁺,ATPase activity levels exist (Anderson et al., 2010). Therefore, ATP1A3 PMG-associated variants are likely LOF (affecting levels of Na⁺ accumulation in cells), with severe phenotypes reflecting more complete LOF effects and toxicity within more vulnerable cell types.
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AUTHOR CONTRIBUTIONS

C.A.W., A.J.B., R.S.S., S.K.A., F.P.V., E.M., L.F.S. and L.B. collected and evaluated clinical data; J.E.N. coordinated clinical data; R.S.S. performed and analyzed tissue experiments; M.F. and R.S.S. performed cortical dissections; M.G., C.D.M., N.R. and M.F. generated Drop-Seq libraries; M.F. analyzed Drop-Seq data; R.S.S., M.F., J.E.N., S.K.A. and C.A.W. wrote the manuscript, with contributions from all co-authors. We thank Kathleen Sweadner for productive discussion. The authors declare no competing financial interests.
MAIN FIGURES

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Single-cell analysis of ATP1A3 expression in the infant human neocortex
Figure 1. *ATPIA3* variants associated with cerebral cortex malformations
Figure 1. *ATP1A3* variants associated with cerebral cortex malformations

(A) MRI images of Case A, B, C and D showing the cortical malformation, polymicrogyria (PMG). White arrows denote gross location of affected brain regions. The Control image comes from an unaffected 11-year-old. Scale bar (1cm) (B) Pedigrees with novel *de novo* pathogenic single nucleotide variants in *ATP1A3*: Case A (missense, p.Arg901Met) with bilateral frontoparietal PMG, Case B (splice donor site, c.2921+1G>A) with extensive bilateral PMG, Case C (missense, p.Leu924Pro) with unilateral PMG, and Case D (missense, p.Gly851Arg) with extensive bilateral PMG, more pronounced in the right hemisphere. Individual features: Genotype (-/+, *de novo* heterozygous), PMG, infantile seizures, developmental delay, and postnatal microcephaly shown in pedigree summary. Square, male; circle, female; black and/or gray shading, affected individual. See supplemental text for genetic validation and comprehensive clinical phenotyping.
Figure 2. Schematic of ATP1A3 variants associated with cerebral cortex malformations and other ATP1A3-related diseases
Figure 2. Schematic of \textit{ATP1A3} variants associated with cerebral cortex malformations and other \textit{ATP1A3} related diseases

(A) Overview of the $\alpha$ subunit (green) of the Na,K-ATPase with novel PMG alleles mapped (colored amino acids) and previous case report of PMG associated allele (Leu888Pro)(Gurrieri et al., 2016). Red, p.Leu888Pro (L888P); Orange, p.Arg901Met (R901M); Purple, p.Leu924Pro (L924P); Blue, p.Gln851Arg (Q851R). Image generated with PyMOL using Protein Databank 2ZXE(Shinoda et al., 2009). (B) Transmembrane topology schematic of Na,K-ATPase for visualization of PMG causing variants, including $\alpha$, $\beta$, and FXYD subunits. We also denote a variant enriched region TM7-TM8, where $\beta$ subunits interact with the $\alpha3$ subunit in the extracellular TM7-TM8 segment Glu899, Gln904, and Gln905(Shinoda et al., 2009). (C) Overview of genotype-phenotype correlation of \textit{ATP1A3} associated disease single nucleotide variants, phenotypes were grouped by primary symptomatic presentation: including early infantile epileptic encephalopathy (EIEE); alternating hemiplegia of childhood (ACH); cerebellar ataxia, areflexia, pes cavus, optic atrophy, and sensorineural hearing loss (CAPOS); rapid-onset dystonia-parkinsonism (RDP); and childhood onset schizophrenia (COS), and/or a cortical malformation syndrome (MCD+), cerebellar ataxia (CA). Alleles from several sources including(Arstarkhova et al., 2019; Brashear et al., 1993; Holm et al., 2016; Swoadner et al., 2019). See supplemental Figure S7 and Table S3 for complete allele to protein topology breakdown.
Figure 3. *ATP1A3* is differentially enriched to the human fetal period and localized to subplate and deep-layer neurons
Figure 3. *ATP1A3* is differentially enriched to the human fetal period and localized to subplate and deep-layer neurons

(A) Chromosome locations for CNS enriched Na\(^+\),K\(^+\)-ATPase alpha isoforms, *ATP1A1* and *ATP1A2* (Chr1), and *ATP1A3* (Chr19). (B) RNA transcriptome analysis of bulk brain regions revealed *ATP1A3* transcripts are high during fetal gestational and persist postnatally. CNS expressed paralogous *ATP1A1* and *ATP1A2* paralogs show relative expression pattern changes during development, while *ATP1A4* is not expressed significantly in the brain. Raw transcriptome data for (B) from Allen Brain Atlas, presented as log\(_2\) RPKM (reads per kilobase per million) values and a polynomial fit to average across time points (Jones et al., 2009). (C) Left, *ATP1A3* mRNA chromogenic *in situ* hybridization performed on 19 weeks post conception (pcw) coronal brain sections demonstrate highest *ATP1A3* expression in the human cortical plate (CP), and subplate region (SP). Scale bar left, 500µm. Right, Corresponding fluorescence imaging of 19 wpc fetal brain with cell type specific markers for intermediate progenitors (TBR2) and neural progenitors (vimentin, VIM) demonstrates *ATP1A3* transcripts are not present within cells of the sub-ventricular zone (SVZ) and ventricular zone (VZ). Arrows indicate high expressing neurons in deep cortical layers and subplate region. (D) Magnified image from (C) depicting high *ATP1A3* expressing subplate layer cells. (E) Left, Schematic of developing sagittal fetal cerebellum tissue section at 19 wpc demonstrates enrichment of *ATP1A3* in Purkinje cell layer. (i) *ATP1A3* mRNA fluorescence in situ at 19 wpc demonstrates highest expression in the Purkinje cell layer, colocalizing with Purkinje cell marker calbindin (CALB1, green). (ii), Zoomed fluorescence image, including labeling of external granule layer. Scale bar left, 50µm. DAPI stain for nuclei in blue. Scale bar 25µm. VZ, ventricular zone; SVZ, subventricular zone; IMZ, intermediate zone; SP, subplate; CP, cortical plate; PCL, purkinje cell layer; EGL, external granule layer.
Figure 4. Single cell *ATP1A3* expression atlas in the developing human cerebral cortex
**Fig 4. Single cell ATP1A3 expression atlas in the developing human cerebral cortex**

*(A)* **Left:** Schematic showing a 21 post conception weeks (pcw) forebrain and location of regions sampled (for anatomical detail see fig. S2A). **Top:** dorsolateral view. **Bottom:** sagittal view. **Right:** Uniform Manifold Approximation and Projection (UMAP) of 125,943 single cells profiled by DropSeq. Each dot represents a cell, color-coded by sample of origin. Key: blue-to-red: frontal-to-caudal. Occ: Occipital; Par: Parietal; Temp: Temporal; Ins: Insula; Orb: Orbital; Str: Striatum; PFC: Prefrontal Cortex; V1: Primary Visual Cortex; Hp: Hippocampus; CGE: Caudal Ganglionic Eminence; MGE: Medial Ganglionic Eminence **(B)** Graded expression of areal marker genes *LMO3, LMO4* and *NR2F1* (COUP-TF1) across samples ordered caudal-to-rostral (x-axis). Gene expression (y-axis) was normalized by cell (divided by total UMIs/cell x 100k), then the mean value for each gene was aggregated by sample. Notice the opposite gradient of *LMO3* and *NR2F1* expression, and the U-shaped pattern of LMO4 expression along the rostro-caudal axis. See fig. S2B for additional areal markers. **(C) Left:** Histogram showing relative expression of *ATP1A3* (x-axis) across cell clusters (y-axis). Notice enriched expression of *ATP1A3* in cluster EN.4-SP, containing subplate (SP) EN neurons (See Fig. SC and H for SP markers enriched in this cluster). Mean *ATP1A3* expression was aggregated by cluster, then rescaled from 0 to 1. Cell clusters are color-coded by cell type (blue: excitatory neurons (EN); gold: glia; blue: inhibitory neurons (IN); red: neural progenitor cells (NPC)). See fig.SB for cluster markers and assignments. **Right:** Dot-plot showing *ATP1A3* expression across clusters split by areas of origin (x-axis; sample 1-11, ordered from caudal to rostral). Color scale codes for mean expression by group; size of the dots codes for percentage of cells expressing *ATP1A3* in each group. **(D) Top,** Violin-Boxplot showing differential expression of *ATP1A3* (y-axis) in the EN.4-SP cluster across three main cortical partitions (x-axis; Caudal: samples 1-3 (occipital cortex); Medial: samples 4-8 (including parietal cortex).
and temporal cortex, and subcortical structures); Frontal: samples 9-11 (frontal cortex). Dots represent individual cells. Violins show probability density distributions; boxes show interquartile ranges; notches show confidence intervals around the median (horizontal lines). Outliers were trimmed. Mean expression of $ATP1A3$ was aggregated by region, then log10-transformed. Notice the highest expression of $ATP1A3$ in cells sampled from frontal cortex, and lowest expression in cells sampled from caudal cortex. Bottom: Nissl-stained frontal coronal section of a 21 pcw forebrain highlighting anatomical position of SP. Source: Atlas of the Developing Human Brain, BrainSpan (www.brainspan.org), Reference Atlas. (E) $ATP1A3$ expression across cells in the EN clusters (color-coded in blue in C, left) ordered by pseudotime. Dots represent individual cells. Y-axis: log-transformed, scaled expression of $ATP1A3$ by cell. X-axis: pseudotime score (color-coded) for each cell, calculated using the Monocle3 algorithm. Trend-line shows the increase of $ATP1A3$ expression as function of pseudotime, calculated by fitting a quasipoisson model to the data. (F) UMAP of all profiled cells color-coded by cluster (see C, left). The EN.4-SP cluster is highlighted. (G) UMAP of cells in the EN.4-SP cluster (highlighted in F) color-coded by rostral, medial and caudal anatomical partition of origin (see also D, top). Notice that cells segregate by origin. (H) Relative expression of areal marker genes in the EN.4-SP cluster (see G). Relative gene expression is coded by color and size. Notice that $ATP1A3$ is enriched in cells expressing SP markers ($CRYM$, $WNT7B$, $HS3ST4$) and frontal cortex markers ($GRP$, $SEMA3E$), but not caudal cortex markers (NPY, LPL, NEFL).
Figure 5. Single-cell analysis of \textit{ATP1A3} expression in the infant human neocortex
Figure 5. Single-cell analysis of ATP1A3 expression in the infant human neocortex

(A) Left, Schematic showing an 8-month-old neocortex and location of regions sampled. BA: Brodmann area; PFC: Prefrontal Cortex; V1: Primary Visual Cortex. Right, T-distributed Stochastic Neighbor Embedding (tSNE) of 51,878 single nuclei profiled by DropSeq. Each dot represents a nucleus, color-coded by cluster (left), cell type (middle) and origin sample (right). EN: excitatory neurons; IN: inhibitory neurons. See C for cluster and cell-class assignments. (B) Top, dendrogram summarizing hierarchical clustering of data. Clusters are color-coded as in A, left). See fig.S4 for cluster markers and assignments. Middle, bar graph showing relative expression of ATP1A3 (y-axis) across cell clusters (x-axis). Mean expression was aggregated by cluster, then rescaled from 0 to 1 across all clusters. Data are color-coded by cell type (as in A, middle – blue: EN; gold: IN; brown: glia). Bottom, Dot-plot showing ATP1A3 expression across the IN and EN clusters (x-axis) split by areas of origin (y-axis). Color scale codes for mean expression by group; size of the dots codes for percentage of cells expressing ATP1A3 in each group. Right, tSNE plots showing specific enrichment of cortical marker genes including genes associated with early activity in the developing cortex (MEFC2, SLC17A7 and SLC17A6 (VGLU1 and VGLU2), and MCD associated gene TUBB2A. (D) Venn diagram showing intersection between the top-100 genes correlating with ATP1A3 in the EN cluster (blue) and IN cluster (gold). 47/100 genes are shared between the two groups. Top 50 genes correlating with ATP1A3 in the EN and IN clusters are shown. Dots indicate Pearson’s R for each gene in the EN cluster (top; blue) and IN cluster (bottom; gold). Genes found in both groups are indicated in red, and the respective correlation coefficients are connected by red lines.
REFERENCES


Smith et al.,


Monteiro, F.P., Curry, C.J., Hevner, R., Elliott, S., Fisher, J.H., Turcay, J., Dobyns, W.B., Costa,
cause hydrops fetalis, microcephaly, arthrogryposis and extensive cortical malformations.
Eur J Med Genet 63, 103624. doi:10.1016/j.ejmg.2019.01.014

excitability of early neurons in the human cerebral cortex during the second trimester of

onset dystonia-parkinsonism associated with the I758S mutation of the ATP1A3 gene: a
doi:10.1007/s00401-014-1279-x

S., Gunter, S.A., Vasta, V., Zhang, Q., Tran, T., Liu, Y.B., Ozelius, L.J., Brashear, A.,
Sweadner, K.J., Dobyns, W.B., Hahn, S., 2015. Novel mutations in ATP1A3 associated with
catastrophic early life epilepsy, episodic prolonged apnea, and postnatal microcephaly.
Epilepsia 56, 422–430. doi:10.1111/epi.12914

Mediate Activity-Dependent Changes in Mammalian Motor Networks. J. Neurosci. 37, 906–

Platzer, K., Yuan, H., Schütz, H., Winschel, A., Chen, W., Hu, C., Kusumoto, H., Heyne, H.O.,
Helbig, K.L., Tang, S., Willing, M.C., Tinkle, B.T., Adams, D.J., Depienne, C., Keren, B.,
Mignot, C., Frengen, E., Stromme, P., Biskup, S., Döcker, D., Strom, T.M., Mefford, H.C.,
Myers, C.T., Muir, A.M., LaCroix, A., Sadleir, L., Scheffer, I.E., Brilstra, E., van Haelst,
Goldberg, E.M., De Bie, I., Fox, S., Major, P., Jones, J.R., Zackai, E.H., Abou Jamra, R.,
Rolf, S., Leventer, R.J., Lawson, J.A., Roscioli, T., Jansen, F.E., Ranza, E., Korff, C.M.,
Leheşjoki, A.-E., Courage, C., Linnankivi, T., Smith, D.R., Stanley, C., Mintz, M.,
McKnight, D., Decker, A., Tan, W.-H., Tarnopolsky, M.A., Brady, L.I., Wolff, M., Dondit,
L., Pedro, H.F., Parisotto, S.E., Jones, K.L., Patel, A.D., Franz, D.N., Vanzo, R., Marco, E.,
GRIN2B encephalopathy: novel findings on phenotype, variant clustering, functional
doi:10.1136/jmedgenet-2016-104509

Rakic, P., 1974. Neurons in rhesus monkey visual cortex: systematic relation between time of

Shinoda, T., Ogawa, H., Cornelius, F., Toyoshima, C., 2009. Crystal structure of the sodium-
potassium pump at 2.4 A resolution. Nature 459, 446–450. doi:10.1038/nature07939

Simmons, C.Q., Thompson, C.H., Cawthon, B.E., Westlake, G., Swoboda, K.J., Kiskinis, E.,
doi:10.1016/j.nbd.2018.03.009
Smith et al.,


