

Mapping the molecular and cellular complexity of cortical malformations

Esther Klingler, Fiona Francis, Denis Jabaudon, Silvia Cappello

► To cite this version:

Esther Klingler, Fiona Francis, Denis Jabaudon, Silvia Cappello. Mapping the molecular and cellular complexity of cortical malformations. Science, 2021, 371 (6527), pp.eaba4517. 10.1126/science.aba4517. hal-03425482

HAL Id: hal-03425482 https://hal.sorbonne-universite.fr/hal-03425482

Submitted on 10 Nov 2021

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

1 2	TITLE
$\frac{2}{3}$	Mapping the molecular and cellular complexity of cortical malformations
4	
5	
6	AUTHORS AND AFFILIATIONS
7	Esther Klingler ¹ , Fiona Francis ^{2,3,4} , Denis Jabaudon ^{1,5} * & Silvia Cappello ⁶ *
8	
9	¹ Department of Basic Neurosciences, University of Geneva, Geneva, Switzerland ; ² INSERM
10	U 1270, Paris, France ; ³ Sorbonne University, UMR-S 1270, F-75005 Paris, France ; ⁴ Institut
11	du Fer à Moulin, Paris, France ; ⁵ Clinic of Neurology, Geneva University Hospital, Geneva,
12	Switzerland ; ⁶ Max Planck Institute of Psychiatry, 80804 Munich, Germany.
13	* co-corresponding authors
14	
15	
16	

1 ABSTRACT

2 The cerebral cortex is an intricate structure underlying human features such as language and 3 cognition. Cortical functions rely on specialized neurons, which emerge during development 4 from complex molecular and cellular interactions. Neurodevelopmental disorders occur when 5 one or several of these steps are incorrectly executed. While a number of causal genes and 6 disease phenotypes have been identified, the sequence of events linking molecular disruption 7 to clinical expression mostly remains obscure. Here, focusing on human malformations of 8 cortical development, we illustrate how complex interactions at genetic, cellular and circuit 9 levels combinatorially contribute to diversity and variability in disease phenotypes. Through 10 select examples and an online resource, we propose that a multi-level assessment of disease 11 processes is key to identify points of vulnerability and develop novel therapeutic strategies.

1 MAIN TEXT

2 The cerebral cortex, or neocortex, is critical to key behavioral functions in mammals, including 3 language, sociability, and fine motor skills. This brain structure consists of several dozens of 4 specialized types of neurons organized across layers and areas, which are generated during 5 development by the divisions of diverse progenitor cells. Newborn neurons undergo sequential 6 molecular programs that drive their specific local and long-range circuit connectivity and adult 7 function (1). The complexity of corticogenesis is staggering and, while necessary for proper 8 cortical function to emerge, the myriads of molecular and cellular developmental processes 9 involved also provide multiple points of vulnerability for "malformations of cortical 10 development" (2), here termed "cortical malformations". These are usually first detected 11 through their clinical expression, including developmental delay with various combinations of 12 intellectual and motor disabilities, often associated with seizures. Despite the toll on patients 13 and their caregivers, only limited treatments exist and while a number of causal genes have 14 been identified, the sequence of events linking molecular disruption with clinical expression 15 mostly remains obscure.

16 Here, to interrogate cortical malformations and highlight potential points of intervention, 17 we first present basic principles of neocortical development and highlight vulnerable cellular 18 compartments and processes, with particular focus on neurogenesis and neuron migration. 19 Second, we present different "levels" of developmental processes, from genes and gene 20 products to cells, circuits, and clinical expression. Focusing on cortical malformations, we 21 illustrate how complex interactions within and across these levels may account for variable 22 disease patterns. Finally, we propose a framework integrating these different levels of 23 organization, towards better understanding and treatment of the disease, as well as an online 24 resource (http://genebrowser.unige.ch/humous/) to identify and compare genetic and cellular 25 developmental processes in mice, human embryos, and human organoids.

26 Neocortical development

During embryonic neocortical development, neurons are not born at the place where they will reside in adulthood, but instead migrate relatively long distances to reach their destination. Not only are cortical neurons born at different places, but distinct types of neurons are born at different times, over several days in mice (from embryonic day (E) 11 to 17 (*3*)) and several weeks in humans (from post-conception week (pcw) 8 to 24 (*4*)) (Fig. 1A). Generation of the cortex results from billions of cells undergoing four key sequential and partially overlapping

1 processes: (1) progenitor division and neurogenesis, (2) migration, (3) neuritogenesis (i.e. 2 extension of axon and dendrites) and (4) synaptogenesis (Fig. 1A-C). Cortical malformations 3 occur when one or several of these developmental events are improperly executed (Fig. 1B); 4 they typically manifest as macroscopic anatomical defects visible with brain imaging 5 techniques such as magnetic resonance imaging (MRI) (Fig. 2). Morphological manifestations 6 include microcephaly (decreased brain size), lissencephaly (disrupted cortical folding and 7 lamination), polymicrogyria (numerous small cortical folds), and heterotopias (abnormally 8 positioned cells in periventricular or subcortical regions) (5). In other cases, more subtle defects 9 are visible only in pathology specimens, e.g. in focal cortical dysplasias (abnormal cortical 10 lamination) (2).

11 From E8.5-E9.5 in mice and pcw4 in human (4), progenitors of the dorsal pallium, at the 12 rostral end of the neural tube, which initially form a single-layered neuroepithelium, undergo 13 self-replicating divisions to increase in numbers before generating apical radial glia (aRG, also 14 called apical progenitors) and forming an expanded germinal zone called the ventricular zone 15 (3). Around E11.5 in mice and pcw8 in human, aRGs start dividing asymmetrically to give rise 16 to their first neuronal progeny (Fig.1A) (4). Different types of excitatory cortical neurons are 17 then sequentially born between E11.5 and E17.5 in mice (pcw8-24 in human) and distribute to 18 form six radially-organized layers in an inside-out manner: deep layer neurons, which project 19 subcortically, are born first while superficial layer neurons, which project intracortically, are 20 born last (Fig. 1A) (1). Of note, inhibitory GABAergic neurons, which are not discussed in 21 detail in this review, are born in a distinct germinal zone, the ventral pallium, and migrate at 22 later timepoints into the cortex to form local circuits with glutamatergic neurons (6).

Radial cortical organization - The laminar organization of the adult neocortex tightly relies 23 24 on the radial polarity of progenitors and their daughter cells during development. Accordingly, 25 aRGs are highly polarized cells with a radial process extending to the pial surface, necessary 26 to guide the daughter neuron migration (Fig. 1A). Adherens junctions at the apical (*i.e.* towards 27 the ventricle) pole of aRGs are critical to this polarity, as they maintain a cohesive ventricular 28 zone and regulate the balance between proliferation and differentiation through cell-cell 29 interactions (3). Mutations of genes coding for adherens junction proteins, such as α/β -catenins and N-cadherin, indirectly disrupt cell divisions, leading to changes in cortical 30 size and folding, as well as neuronal heterotopias in mice and human (7-10). Similarly, 31 32 mutations in the genes coding for cell-cell contact proteins such as DCHS1 and FAT4 disrupt 33 aRG polarity and neuronal migration, resulting in human periventricular heterotopias (11).

Another notable determinant of aRG polarity is the primary cilium, a specialized organelle which transduces signals from ventricular cerebrospinal fluid (Fig. 1A) (3). Mutations in genes coding for cilium proteins ("ciliopathy" genes, *e.g. BBS1, BBS7, BBS10* and *TMEM216*) affect progenitor proliferation and newborn neuron migration in mice (12) and mutations in the microtubule-associated protein encoding gene *EML1*, which causes neuronal heterotopia in mice and humans, are likewise thought to act by destabilization of the primary cilium (13, 14).

7 *Cycling of progenitor cells* – The ability to undergo cell cycle is a central feature of progenitors. 8 A number of mutations can perturb this process, either by affecting cell divisions, or through 9 premature differentiation into neurons. For example, mutations of genes coding for proteins 10 controlling mitotic spindle organization (Fig. 1A) typically lead to premature differentiation 11 and / or aRG apoptosis, resulting in reduced neurogenesis, and manifesting in humans as 12 microcephaly with or without lissencephaly (e.g. ASPM, CENPJ, WDR62 (15)). Of note, both 13 premature differentiation into neurons as well as excessive self-replication of progenitors may 14 in principle both result in decreased neuronal output: in one case the progenitor pool is 15 prematurely depleted while in the other, progenitors fail to give rise to neurons. Divisions of 16 aRGs are initially rapid and then slow down as corticogenesis unfolds (from 8 hours at early 17 stages to 18 hours at the end of corticogenesis in mice, and from 23 to 54 hours in primates 18 (16, 17)). The slowing down of cell cycle as corticogenesis proceeds may per se confer new 19 properties to daughter cells as low-affinity DNA ligands such as transcription factors are more 20 likely to bind when target sites are available for longer times. Hence, disruption of cell cycle 21 not only affects the number of neurons being generated (typically decreasing neuronal output) 22 but also the relative proportions of daughter neuron types, since distinct types are sequentially 23 produced at different stages of corticogenesis (18).

Diversity of progenitor cells - aRGs not only generate neurons, but also a distinct type of 24 25 progenitor, termed intermediate progenitors (also called basal progenitors), which delaminate 26 from the ventricular zone and form a new germinal layer, the subventricular zone (Fig. 1A). 27 Intermediate progenitors are more neurogenic than aRGs, and act to boost neuronal production 28 (19). Consistent with such a function, human mutations affecting intermediate progenitors, such as mutations in EOMES (also known as TBR2), perturb gyri formation and result in 29 30 microcephaly, lissencephaly or polymicrogyria (20). In most gyrencephalic species (i.e. 31 species with a folded neocortex), including primates, additional progenitor types exist, 32 including basal radial glia (also called outer radial glia), which form a distinct proliferative 33 sub-compartment, the outer subventricular zone (Fig. 1A). Basal radial glia express distinctive

genes (*e.g. TMEM14B* (21)) and morphologies (*e.g.* many have a basal process but no apical
 process), and are thought to underlie the disproportionate expansion in superficial layers
 occurring during gyrification (22).

4 Neuronal migration - Following cell division, while undergoing a complex series of 5 morphological changes, postmitotic daughter neurons migrate radially along aRG basal 6 processes away from the ventricular zone to the cortical plate, where they settle in their 7 appropriate final positions. The radial migration of neurons relies on the extension of a leading 8 process, with the leading edge responding to environmental cues. The neuron's centrosome 9 moves within the leading process allowing somal translocation and locomotion (Fig. 1A) (23). 10 The proper execution of this migratory sequence is key to building the correct cortical 11 architecture and relies tightly on microtubule and actin cytoskeleton-associated proteins. 12 Accordingly, mutations in the genes coding for these proteins have dramatic effects on cortical 13 structure (5, 15), and account for a disproportionate fraction of cortical malformations, as will 14 be detailed in a later section.

15 Preplate splitting, which occurs at E11 in mouse (human pcw9-10) (24), is a critical event 16 during corticogenesis and evolution (Fig. 1A). The glycoprotein Reelin secreted by Cajal-17 Retzius cells present in the marginal zone plays an important role in these processes. Preplate 18 splitting defects consistently give rise to abnormal cortex development due to aberrant cortical 19 neuron position. Indeed, lamination defects in *Reeler* mice, mutant for the *Reln* gene, not only reflect abnormal neuronal migration, but also abnormal splitting of the preplate into a 20 21 superficial marginal zone (which becomes layer 1) and a transient deep layer subplate 22 (sometimes called layer 6b) (Fig. 1A). In humans, mutations in RELN cause autosomal 23 recessive lissencephaly (25). As discussed in a later section, despite the severe cortical 24 lamination defects in *Reeler* mouse mutants, long-range cortical input/output connectivity appears largely preserved (26, 27). Pomgnt2 mouse mutants, a model of "cobblestone" 25 26 lissencephaly caused by mutation in the POMGNT2 gene in human, also show preplate splitting 27 defects, since abnormal clusters of subplate neurons are observed in superficial layers (28). 28 The subplate itself plays an important role in the radial migration of the neurons which traverse 29 it and, later, in circuit assembly: subplate neurons make transient synaptic connections with 30 migrating neurons, which primes their radial migration (29), and also act as a scaffold for 31 thalamocortical and inhibitory circuit maturation (30, 31). Although mutations specifically 32 affecting this process have not been reported in humans, the subplate is usually visible in human fetal MRIs; in the future, it may be possible to reveal subtle subplate defects underlying
 more significant circuit dysfunction.

3 Building circuits - Neuritogenesis and synaptogenesis are critical for circuit assembly and a 4 bottleneck for cortical function. They have been discussed in detail in recent reviews (32) and 5 will only be briefly covered here. Axon-dendrite polarization starts soon after neurons are born, 6 when they still are migrating (33). Axon and dendrite growth relies on their growth cones which 7 are highly dynamic actin-rich protrusions sensitive to environmental guidance cues (Fig. 1A). 8 As is the case for migration, axonal guidance and extension thus heavily rely on cytoskeletal 9 function. Cortical malformations can be accompanied by anomalies in axonal tracts, most 10 notably in the form of atrophy or agenesis the corpus callosum, which connects both 11 hemispheres in human (34). Other anatomically less prominent axonal pathway defects are 12 likely to occur, and the use of MRI techniques such as diffusion tensor imaging (35) may in 13 the future further address this possibility. Synaptic dysfunction, including inhibitory synaptic 14 dysfunction, is thought to underlie many if not most neuropsychiatric disorders in humans (36), 15 and like migration and neuritogenesis, synaptogenesis involves a highly organized 16 cytoskeleton (Fig. 1A). Neuropsychiatric disorders are not generally associated with salient 17 MRI defects, yet some level of anatomical dysfunction likely exists. For example, patches of 18 cortical disorganization and white matter defects (particularly in the corpus callosum) have 19 been reported in patients with autism spectrum disorder (37, 38). Similarly, decreased neuronal 20 density have been reported in neuropathological specimens of patients with schizophrenia (39). 21 The extent to which these anatomical findings drive the clinical phenotype or are secondary to 22 another underlying process remains unclear.

23 Sources of complexity in cortical malformations

24 Cortical malformations have a broad range of clinical expression, including seizures, 25 intellectual disabilities, autistic features, failure to reach developmental milestones, and non-26 neurological co-morbidities (e.g. dysmorphic facial and musculoskeletal features, skin 27 abnormalities, cardiovascular defects) (5). These features alone often do not allow 28 identification of the cause of the disease, because genetically diverse defects lead to largely 29 overlapping clinical pictures, and impairment of single genes can lead to multiple clinical 30 defects. Thus, while an increasing number of causal genes is being identified and a diversity of 31 anatomical and clinical phenotypes have been recognized (15), the sequence of events linking 32 molecular disruption to final outcome remains mostly out of reach. In the following section, 33 we will highlight potential sources of variability which have precluded establishment of clear

1 causal relationships across genes and molecules, cell types and circuits, and clinical expression

2 (Fig. 3A).

3 Genetic and molecular dysfunction - Neurodevelopmental disorders can result from single 4 point mutations in critical genes (*i.e.* monogenic mutations; Fig. 3B), or from DNA sequence 5 variations at multiple loci that together increase disease probability. Convergent, divergent, or 6 mixed relationships between genetic defect(s) and disease phenotype(s) may occur (Fig. 3C-7 F), and penetrance of the genetic mutation may be incomplete and variable (*i.e.* the mutation 8 is not expressed at the clinical level (Fig. 3G)), together complexifying causal analyses. 9 Understanding this cross-level organization is critical in therapeutic terms: for example, 10 "hidden hubs" represent convergent points of pharmacological intervention across disease 11 phenotypes, while disorders with "hidden complexity" would require distinct treatments 12 despite similar clinical presentations (Fig. 3E, F). The contribution of monogenic causes varies 13 across conditions; in the case of lissencephaly for example, 20% of cases could not be 14 associated with a single genetic cause (40), which may reflect yet un-identified gene mutations, 15 mutations in non-coding regions (41), polygenic or multifactorial inheritance, modulated by 16 the influence of non-genetic factors (e.g. infectious or toxic), or somatic mosaicism (42). While 17 monogenic causes appear to predominate in cortical malformations, neuropsychiatric disorders 18 such as autism spectrum disorders and schizophrenia instead appear to mostly involve 19 polygenic processes including through chromosomal deletions (39). Of note, in chromosomal 20 deletions, disease-causing genes need not to be in the affected chromosomal region as local 21 DNA conformation changes may in principle epigenetically alter gene accessibility in other 22 parts of the genome. Although not discussed in detail here, some genetic mutations occur in a 23 clonal manner, *i.e.* do not affect the whole brain but only the progeny of some cells with 24 somatic mutations. Somatic mutations have been clearly established as a cause of focal cortical 25 dysplasia and hemi-megalencephaly (hypertrophy of a single hemisphere) (42), but whether 26 clonal processes are involved in other disorders, and in particular whether at least some 27 psychiatric disorders may result from mosaic synaptic dysfunction, is unknown (43). 28 Overlapping clinical pictures can emerge from mutations in genes which are not apparently 29 functionally correlated. For example, over 200 de novo variants have been identified in a cohort 30 of patients with periventricular heterotopia, suggesting that many genes will be associated with 31 this condition, with a high level of convergence in the final outcome (e.g. mismigration in the 32 case of heterotopias) (44).

1 Complexifying the situation, the aforementioned developmental processes (i.e. 2 neurogenesis, migration, axo- and synaptogenesis) often involve redundant molecular 3 mechanisms. For example, the cyclin-dependent kinase CDK5 (whose gene mutation is 4 associated with lissence phaly in human (45)) is critical for mitosis in mouse progenitors, but 5 also in their daughter neurons for migration and gyrification, synapse formation and even long-6 term plasticity in adults (Fig. 1C) (46, 47). Similarly, many, if not most, axon guidance 7 molecules also have earlier roles in neuron migration and even neuron generation, and/or in 8 synapse and spine formation and maturation (48-50). For example, the Ephrin receptor EphB2 9 both guides corpus callosum axons and stabilizes synaptic contacts (51, 52). Gene mutations 10 may therefore affect sequential cellular processes in parallel. Such versatility in protein 11 function likely contributes to the complexity and variability of disease processes.

12 Cytoskeletal perturbations – Given their involvement in multiple cellular processes, from 13 cytokinesis to migration and neuritogenesis, mutations in genes coding for cytoskeletal proteins 14 are particularly prevalent causes of cortical malformations. Strikingly however, quite different 15 anatomical defects can result from mutations in genes coding for apparently related proteins. 16 For example, in "tubulinopathies", mutations in a tubulin-encoding gene, TUBA1A, generally 17 decreases gyrification in humans, while disruption of another tubulin, TUBB3 often gives 18 polymicrogyria (53). Microtubule-related proteins can behave similarly, as mutations in the 19 gene coding for doublecortin (DCX, which stabilizes microtubules and regulates certain 20 molecular motors during neuronal migration (54)), or LIS1 (which regulates dynein and is 21 required for interkinetic migration of aRGs as well as neuronal migration (55)), variably result 22 in either subcortical heterotopias or lissencephaly in humans (5). In the case of DCX, which is 23 located on the X chromosome, mutations in women typically lead to heterotopia while in men 24 it is associated with lissencephaly, suggesting dose-dependent effects (56).

25 Post-transcriptional and post-translational dysregulations are also a cause of cortical 26 malformations. For example, abnormal alternative splicing can occur in the Filamin A actin-27 binding transcript (Flna), causing periventricular heterotopias in humans (57). Inhibition of a 28 neuronal "poison exon" by the Polypyrimidine Tract Binding Protein PTBP1 is necessary to 29 maintain neural progenitor identity. In the absence of PTBP1 function, de-repression of this 30 poison exon leads to decreased expression of FLNA and precocious neuronal differentiation 31 (Fig. 4A). Mutations of proteins modifying the conformation and activity of other proteins, 32 like kinases (e.g. PI₃K/AKT pathway (58)), glycosylases (59) and tubulin chaperones /

modifiers (*e.g.* Tubulin Folding Cofactor D (*TBCD*) (60)) are further post-translational causes
of cortical malformations.

Obviously, not all cortical malformations are the result of disrupted cytoskeletal function. Other classes of proteins, such as transcription factors, are also involved (*e.g.* ARX, PAX6, EOMES and FOXG1 whose gene mutations lead to severe cortical malformations in humans (Fig. 2) and mice (20, 61-63)). Such examples are, however, comparatively rare despite their critical function in orchestrating developmental genetic programs. One explanation for this apparent paradox is that many transcription factors may be "too" critical for corticogenesis, such that loss of their function is embryonically lethal.

10 Disrupted spatio-temporal choreographies - Normal gene expression is tightly regulated in 11 space and time during development. Hence, abnormal dynamic regulation of gene expression is likely a critical factor in the pathogenesis of cortical malformations. Supporting this 12 13 possibility, Flna mutant mice do not display neuronal migration defects, while acute RNAi-14 mediated knockdown of *Flna* in rats reproduces the periventricular heterotopias similar to those 15 observed in human patients (64). Also, constitutive Dcx mutant mice have a fairly normal 16 neocortex and behavior (65), while acute RNAi-mediated knockdown of Dcx causes 17 subcortical band heterotopia (66) (Fig. 4B). In principle, while dysfunction of early-onset genes 18 cascade onto a broad array of subsequent cellular processes and may lead to more severe 19 phenotypes, early-onset defects may also be more easily compensated for than defects affecting 20 late differentiation stages.

21 Spatial regulation of gene expression is also important, particularly in the cortex where 22 functionally specialized areas exist. In many cases, malformations are focal or regionalized 23 rather than diffuse, although what determines their location is usually unknown. Cortical 24 malformations often show antero-posterior gradients (40). For example, FLNA mutations typically cause frontal-predominant heterotopias (67), and GPR56 mutations frontal-25 26 predominant polymicrogyria (68). In contrast, CEP85L shows higher expression in posterior 27 cortical regions and mutations in this gene lead to posterior-predominant polymicrogyria (69). 28 Some affected genes, however, are ubiquitously expressed (e.g. GPR56 (70)); interactions with 29 locally expressed binding partners or specific developmental dynamics may here account for 30 the stereotypic location of the cortical defect. In some cases, such as FLNA, GPR56 or EOMES, 31 spatial restriction of gene expression in regions underlying gyri or sulci (71) could further 32 contribute to impairments in cortical folding. Finally, somatic mutations may also account for

the focal nature of a defect (*e.g.* PI₃K pathway (72)), although what determines area location
is often unknown (42).

3 Abnormal cell types – Cell types are the end-result of the dynamic spatio-temporal expression 4 of genes during development. Cortical malformations typically affect multiple types of neurons 5 and/or progenitors, which is in contrast with neurodegenerative diseases, where single cell-6 types (e.g. dopaminergic neurons in Parkinson disease) are predominantly affected. The 7 apparent lack of cell type specific impairments may reflect that differentiation processes are 8 mostly generic across neuron types, and that cell type identity mostly emerges from the 9 combinatorial interactions of multiple genes rather than single master regulators (73). In some 10 cases (mainly focal cortical dysplasia, autism spectrum disorder, schizophrenia and epilepsy), 11 specific defects have also been reported in inhibitory neurons, astrocytes and oligodendrocytes 12 (43, 74) (Fig. 4C). Failure in astrocyte-dependent removal of extracellular debris may lead to 13 a chronically pro-inflammatory environment, potentially increasing susceptibility to disease, 14 but the extent to which reported changes in non-neuronal cells are reactive to neuronal loss is, 15 however, often unclear (75). Of note, since astrocytes are born towards the end of neurogenesis 16 (likely from the progeny of aRGs), mutations that affect aRG divisions are probably associated 17 with a decreased number of astrocytes, although this has not been systematically assessed.

18 Depending on progenitor types and stage of corticogenesis, daughter neurons differ and the 19 aberrant generation of one of these progenitor types could in principle affect specific progenies. 20 For example, mutations affecting intermediate progenitor function could preferentially affect 21 superficial layers, since such progenitors become more and more prevalent as corticogenesis 22 unfolds and superficial layer neurons are born last. Recently, electrophysiological properties 23 of progenitors have been shown to contribute to the sequential generation of neuronal types 24 (76) and may thus contribute to cortical malformations. Accordingly, mutation in the sodium 25 channel Na_V1.3 (SCN3A) gene affects outer subventricular zone progenitors and causes focal 26 polymicrogyria in humans, potentially reflecting excess or altered superficial neuron 27 production (77).

Abnormal circuits – Although neuronal heterotopia likely affects circuit assembly, neuronal identity and basic circuit properties are often conserved despite abnormal neuronal position. In mice, ectopic neurons usually still express their proper laminar markers and display conserved long-range projections (27, 78, 79) (Fig. 4D). Similarly, L5-type neurons in L4 retain their connectivity, as do L4-type neurons in L2/3 (80, 81) and ectopic neurons still attract the proper subtypes of GABAergic interneurons with whom they make synaptic connections (as if they

1 were normally positioned in the cortex), suggesting that local connectivity might likewise be 2 preserved (82). How GABAergic interneuron migration is affected in microcephaly or other 3 cortical malformations has not been systematically examined (but see (83)), but disruption of 4 this process could provide a cellular/circuit explanation for the seizures seen in some patients. 5 In the *Reeler* mouse, although distinct layers are no longer visible, L4-type neurons still receive 6 proper innervation from the thalamus and somatotopic innervation of the cortex is relatively 7 preserved (26, 27) (Fig. 4D). Supporting spared circuit function, perception, learning and 8 memory in *Reeler* mutant mice are little affected by disrupted cortical lamination (27), although 9 more cortex-targeted behavioral tests would likely reveal some level of dysfunction. Together, 10 these results suggest that neuron identity emerges largely cell-autonomously, such that long-11 distance projections (and possibly microcircuits) may be relatively spared in neuronal 12 heterotopias.

13 Behavior and clinical features - As a consequence of the number and complexity of the 14 molecular and cellular processes involved in cortical disorders, the clinical manifestations are 15 heterogeneous, and include intellectual disabilities, epilepsy, and a broad spectrum of 16 neuropsychiatric manifestations. What determines this spectrum is unknown, but postnatal, 17 environmental factors and overall genetic background may contribute (84). Epilepsy often 18 occurs in the context of cortical malformations, but which factors determine seizure 19 susceptibility is unclear. In mouse models of cortical heterotopia, the ectopic and isotopic 20 cortices remain interconnected, which may contribute to abnormal neuronal activation and 21 seizures seen in human patients (85, 86). In some cases, abnormal electrical activity was 22 initiated within or adjacent to the heterotopia while in others, seizures were initiated from the 23 overlying cortex. This suggests either that aberrantly located neurons are capable of generating 24 dysfunctional circuits at a distance, or that the overlying homotopic cortex is abnormal (87). 25 Like other clinical features, intellectual functions are also variably affected, spanning from 26 essentially normal functions to severe disability, in which case seizures are often associated 27 (88). In the absence of additional biomarkers, this relatively generic array of symptoms and 28 signs, which have historically been the main criterion for disease classification, thus offer 29 limited opportunities for a mechanistic understanding of the disease process.

30 Towards a multi-level assessment of neurodevelopmental disorders

The examples above highlight several levels (*i.e.* genetic, molecular, cellular, circuit and behavioral) within and across which combinatorial interactions may occur during cortical malformations and preclude a causal understanding of the disease process (Figs. 3-4). Understanding the processes at play at each of these levels for individuals is key to provide
 patients and their families with prognostic indicators (including biomarkers and, more broadly,
 recognized "endophenotypes") and, in the longer term, therapeutic perspectives.

4 Towards these two aims, the use of animal models, including mouse, ferret and non-human 5 primates (the latter being gyrencephalic and thus closer to humans) is important, in particular, 6 at some point, to assess the therapeutic effect of approaches on behavior, including through insertion of human mutations via gene editing technologies (47). In vitro human models are 7 8 also emerging as a promising and feasible avenue: somatic cells can now be reprogrammed 9 into induced pluripotent stem cells (iPSCs) and serve as a basis for the generation of 3D 10 organoids and assembloids of the brain (89, 90). Organoids have been used to study a variety 11 of neurodevelopmental disorders including cortical malformations (e.g. CDK5RAP2 mutation 12 leading to cobblestone lissencephaly in humans) (91). Despite their limitations in terms of 13 reproducibility and restriction to early stages of corticogenesis (gyrification is for example 14 challenging to study in these models) (92), organoids are poised to become broadly used in 15 personalized medicine by giving access to the neurons of the patient under study.

16 Recent advances in single-cell transcriptomics now allow the dynamic developmental 17 expression of genes in emerging cell types to be assessed (93, 94) (Fig. 5A), potentially linking 18 cellular diversity and characteristics with circuits and behavioral repertoires. Comparison of 19 gene expression across brain development in mouse, monkey, human and primate- / human-20 derived brain organoids is particularly important to identify perturbed processes and select the 21 appropriate study model. As a first step in this direction, here we provide an online resource 22 (http://genebrowser.unige.ch/humous/) compiling transcriptional maps across development 23 and neuron differentiation for mouse embryos, human embryos and human-derived organoids 24 (94-96). As an example of an application of such a database, combinatorial analysis of arrays 25 of genes implicated in microcephaly, lissencephaly or polymicrogyria using this resource 26 confirms that microcephaly predominantly affects progenitor function, while lissencephaly and 27 polymicrogyria affect postmitotic neurons (Fig. 5B). In the latter two cases, human data 28 emphasizes that progenitors may also be affected, highlighting the value of trans-species 29 comparisons (see arrowheads in Fig. 5B).

Multi-omics approaches, in which transcriptomics are combined with genomics, epigenomics and proteomics will provide an opportunity to further unravel the cell type specific processes at play and distinguish core molecular / cellular disease processes from idiosyncratic or stochastic variability. Comparing the distribution of affected processes (at any 1 level of analysis) across patients will help to build a spectrum of abnormal states and predict 2 vulnerable processes (Fig. 5C). Integrative approaches including electrophysiological, 3 imaging, and clinical and biological data from patients, for example using state-of-the art 4 artificial intelligence algorithms (97, 98), may allow the bridging of DNA mutation(s) to gene 5 expression, cellular, anatomy and circuit consequences (Fig. 5D). This will be instrumental for 6 a physiopathogenic classification of diseases, an essential step for a more precise patient 7 stratification and for the design of personalized diagnostic and therapeutic tools. Although this 8 level of integration may seem futuristic, high-throughput techniques and analytical tools are 9 increasingly available, paving the road for such strategies in a realistically close future.

10

11

1 2

REFERENCES

- 3 1. D. Jabaudon, Fate and freedom in developing neocortical circuits. *Nature*4 *Communications*. 8, 16042–9 (2017).
- 5 2. M. Severino *et al.*, Definitions and classification of malformations of cortical
 6 development: practical guidelines. *Brain.* 32, 1123–21 (2020).
- 7 3. E. Taverna, M. Götz, W. B. Huttner, The cell biology of neurogenesis: toward an
 8 understanding of the development and evolution of the neocortex. *Annu. Rev. Cell Dev. Biol.*9 30, 465–502 (2014).
- J. C. Silbereis, S. Pochareddy, Y. Zhu, M. Li, N. Šestan, The cellular and molecular
 landscapes of the developing human central nervous system. *Neuron*. 89, 248–268 (2016).

R. Guerrini, W. B. Dobyns, Malformations of cortical development: clinical features and
genetic causes. *The Lancet Neurology*. 13, 710–726 (2014).

14 6. L. Lim, D. Mi, A. Llorca, O. Marín, Development and functional diversification of cortical
15 interneurons. *Neuron.* 100, 294–313 (2018).

16 7. W.-H. Lien, O. Klezovitch, T. E. Fernandez, J. Delrow, V. Vasioukhin, AlphaE-catenin
17 controls cerebral cortical size by regulating the hedgehog signaling pathway. *Science*. 311,
18 1609–1612 (2006).

- M. Kadowaki *et al.*, N-cadherin mediates cortical organization in the mouse brain.
 Developmental Biology. 304, 22–33 (2007).
- 9. A. E. Schaffer *et al.*, Biallelic loss of human CTNNA2, encoding αN-catenin, leads to
 ARP2/3 complex overactivity and disordered cortical neuronal migration. *Nat Genet.* 50,
 1093–1101 (2018).
- 10. A. Winczewska-Wiktor *et al.*, A de novo CTNNB1 nonsense mutation associated with
 syndromic atypical hyperekplexia, microcephaly and intellectual disability: a case report. *BMC Neurology*, 1–6 (2016).
- 11. J. Klaus *et al.*, Altered neuronal migratory trajectories in human cerebral organoids derived
 from individuals with neuronal heterotopia. *Nat. Med.* 25, 561–568 (2019).

- 12. J. Guo *et al.*, Developmental disruptions underlying brain abnormalities in ciliopathies.
 Nature Communications. 6, 7857–13 (2015).
- 3 13. M. Kielar *et al.*, Mutations in Eml1 lead to ectopic progenitors and neuronal heterotopia
 4 in mouse and human. *Nat Neurosci.* 17, 923–933 (2014).
- 5 14. A. Uzquiano *et al.*, Mutations in the heterotopia gene Eml1/EML1 severely disrupt the
 6 formation of primary cilia. *CellReports.* 28, 1596–1611.e10 (2019).
- 7 15. D. M. Romero, N. Bahi-Buisson, F. Francis, Genetics and mechanisms leading to human
 8 cortical malformations. *Seminars in Cell and Developmental Biology*. 76, 33–75 (2018).
- 9 16. V. S. Caviness, T. Takahashi, Proliferative events in the cerebral ventricular zone. *Brain*10 *Dev.* 17, 159–163 (1995).
- 11 17. C. Dehay, H. Kennedy, Cell-cycle control and cortical development. *Nat Rev Neurosci.* 8,
 12 438–450 (2007).
- 18. L.-J. Pilaz *et al.*, Prolonged mitosis of neural progenitors alters cell fate in the developing
 brain. 89, 83–99 (2016).
- 15 19. A. Pontious, T. Kowalczyk, C. Englund, R. F. Hevner, Role of intermediate progenitor
 16 cells in cerebral cortex development. *Dev Neurosci.* 30, 24–32 (2008).
- 17 20. L. Baala *et al.*, Homozygous silencing of T-box transcription factor EOMES leads to
 18 microcephaly with polymicrogyria and corpus callosum agenesis. *Nat Genet.* 39, 454–456
 19 (2007).
- 20 21. J. Liu *et al.*, The primate-specific gene TMEM14B marks outer radial glia cells and
 21 promotes cortical expansion and folding. *Stem Cell.* 21, 635–649.e8 (2017).
- 22. J. H. Lui, D. V. Hansen, A. R. Kriegstein, Development and evolution of the human
 neocortex. *Cell.* 146, 18–36 (2011).
- 24 23. C. G. Silva, E. Peyre, L. Nguyen, Cell migration promotes dynamic cellular interactions
 25 to control cerebral cortex morphogenesis. *Nat Rev Neurosci.* 20, 318–329 (2019).
- 24. Z. Molnár *et al.*, Comparative aspects of cerebral cortical development. *European Journal*of *Neuroscience*. 23, 921–934 (2006).

S. E. Hong *et al.*, Autosomal recessive lissencephaly with cerebellar hypoplasia is
 associated with human RELN mutations. *Nat Genet.* 26, 93–96 (2000).

26. V. S. Caviness, Patterns of cell and fiber distribution in the neocortex of the reeler mutant
mouse. J. Comp. Neurol. 170, 435–447 (1976).

27. R. J. Wagener, C. David, S. Zhao, C. A. Haas, J. F. Staiger, The somatosensory cortex of
reeler mutant mice shows absent layering but intact formation and behavioral activation of
columnar somatotopic maps. *Journal of Neuroscience*. 30, 15700–15709 (2010).

- 8 28. N. Nakagawa, H. Yagi, K. Kato, H. Takematsu, S. Oka, Ectopic clustering of Cajal9 Retzius and subplate cells is an initial pathological feature in Pomgnt2-knockout mice, a model
 10 of dystroglycanopathy. *Sci Rep.* 5, 11163 (2015).
- 29. C. Ohtaka-Maruyama *et al.*, Synaptic transmission from subplate neurons controls radial
 migration of neocortical neurons. *Science*. 360, 313–317 (2018).
- 30. A. Hoerder Suabedissen, Z. Molnár, Development, evolution and pathology of neocortical
 subplate neurons. *Nat Rev Neurosci.* 16, 133–146 (2015).
- 15 31. P. O. Kanold, R. Deng, X. Meng, The integrative function of silent synapses on subplate
 16 neurons in cortical development and dysfunction. *Front. Neuroanat.* 13, 41 (2019).
- 17 32. C. R. Cadwell, A. Bhaduri, M. A. Mostajo-Radji, M. G. Keefe, T. J. Nowakowski,
 18 Development and arealization of the cerebral cortex. *Neuron*. 103, 980–1004 (2019).
- 33. C. Xu *et al.*, Radial glial cell-neuron interaction directs axon formation at the opposite side
 of the neuron from the contact site. *J. Neurosci.* 35, 14517–14532 (2015).
- 34. T. J. Edwards, E. H. Sherr, A. J. Barkovich, L. J. Richards, Clinical, genetic and imaging
 findings identify new causes for corpus callosum development syndromes. *Brain*. 137, 1579–
 1613 (2014).
- 35. M. Wahl, A. J. Barkovich, P. Mukherjee, Diffusion imaging and tractography of
 congenital brain malformations. *Pediatr Radiol.* 40, 59–67 (2010).
- 26 36. P. Monteiro, G. Feng, SHANK proteins: roles at the synapse and in autism spectrum
 27 disorder. *Nat Rev Neurosci*, 1–11 (2017).

- 37. R. Stoner *et al.*, Patches of disorganization in the neocortex of children with autism. N
 Engl J Med. 370, 1209–1219 (2014).
- 3 38. D. Dimond *et al.*, Reduced white matter fiber density in autism spectrum disorder. *Cereb.*4 *Cortex.* 29, 1778–1788 (2019).
- 5 39. M. Li *et al.*, Integrative functional genomic analysis of human brain development and 6 neuropsychiatric risks. *Science*. **362**, eaat7615–15 (2018).
- 40. N. Di Donato *et al.*, Analysis of 17 genes detects mutations in 81% of 811 patients with
 lissencephaly. *Genet. Med.* 20, 1354–1364 (2018).
- 9 41. E. Perenthaler, S. Yousefi, E. Niggl, T. S. Barakat, Beyond the Exome: The non-coding
 10 genome and enhancers in neurodevelopmental disorders and malformations of cortical
 11 development. *Front. Cell. Neurosci.* 13, 352 (2019).
- 42. A. M. D'Gama, C. A. Walsh, Somatic mosaicism and neurodevelopmental disease. *Nat Neurosci.* 21, 1504–1514 (2018).
- 14 43. D. Velmeshev *et al.*, Single-cell genomics identifies cell type-specific molecular changes
 15 in autism. *Science*. 364, 685–689 (2019).
- 44. E. L. Heinzen *et al.*, De novo and inherited private variants in MAP1B in periventricular
 nodular heterotopia. *PLoS Genet.* 14, e1007281 (2018).
- 45. D. Magen *et al.*, Autosomal recessive lissencephaly with cerebellar hypoplasia is
 associated with a loss-of-function mutation in CDK5. *Hum. Genet.* 134, 305–314 (2015).
- 46. A. H. Hawasli *et al.*, Cyclin-dependent kinase 5 governs learning and synaptic plasticity
 via control of NMDAR degradation. *Nat Neurosci.* 10, 880–886 (2007).
- 47. Y. Shinmyo *et al.*, Folding of the cerebral cortex requires CDK5 in upper-layer neurons in
 gyrencephalic mammals. *Cell Reports*. 20, 2131–2143 (2017).
- 48. H. Blockus, A. Chédotal, The multifaceted roles of Slits and Robos in cortical circuits:
 from proliferation to axon guidance and neurological diseases. *Current Opinion in Neurobiology*. 27, 82–88 (2014).

- 49. W. D. Andrews, M. Barber, J. G. Parnavelas, Slit-Robo interactions during cortical
 development. *Journal of Anatomy*. 211, 188–198 (2007).
- 50. E. Arbeille *et al.*, Cerebrospinal fluid-derived Semaphorin3B orients neuroepithelial cell
 divisions in the apicobasal axis. *Nature Communications*, 1–14 (2019).
- 5 51. M. B. Dalva *et al.*, EphB receptors interact with NMDA receptors and regulate excitatory
 synapse formation. *Cell.* 103, 945–956 (2000).
- 52. M. S. Kayser, M. J. Nolt, M. B. Dalva, EphB receptors couple dendritic filopodia motility
 to synapse formation. *Neuron.* 59, 56–69 (2008).
- 9 53. N. Bahi-Buisson, M. Cavallin, Tubulinopathies overview (available at 10 https://www.ncbi.nlm.nih.gov/books/).
- 54. F. Francis *et al.*, Doublecortin is a developmentally regulated, microtubule-associated
 protein expressed in migrating and differentiating neurons. *Neuron.* 23, 247–256 (1999).
- 13 55. J.-W. Tsai, K. H. Bremner, R. B. Vallee, Dual subcellular roles for LIS1 and dynein in
 radial neuronal migration in live brain tissue. *Nat Neurosci.* 10, 970–979 (2007).
- 15 56. R. Guerrini, T. Filippi, Neuronal migration disorders, genetics, and epileptogenesis. J.
 16 *Child Neurol.* 20, 287–299 (2005).
- 57. X. Zhang *et al.*, Cell-type-specific alternative splicing governs cell fate in the developing
 cerebral cortex. *Cell.* 166, 1147–1162.e15 (2016).
- 19 58. L. A. Jansen *et al.*, PI3K/AKT pathway mutations cause a spectrum of brain malformations
 20 from megalencephaly to focal cortical dysplasia. *Brain.* 138, 1613–1628 (2015).
- 59. A. R. Nickolls, C. G. Bönnemann, The roles of dystroglycan in the nervous system:
 insights from animal models of muscular dystrophy. *Dis Model Mech.* 11 (2018),
 doi:10.1242/dmm.035931.
- 60. E. Flex *et al.*, Biallelic mutations in TBCD, encoding the tubulin folding cofactor D,
 perturb microtubule dynamics and cause early-onset encephalopathy. *Am. J. Hum. Genet.* 99,
 962–973 (2016).

- M. Kato *et al.*, Mutations of ARX are associated with striking pleiotropy and consistent
 genotype-phenotype correlation. *Hum. Mutat.* 23, 147–159 (2004).
- 3 62. L. K. Davis *et al.*, Pax6 3' deletion results in aniridia, autism and mental retardation. *Hum.*4 *Genet.* 123, 371–378 (2008).
- 63. M. Pringsheim *et al.*, Structural brain anomalies in patients with FOXG1 syndrome and in
 Foxg1+/- mice. *Ann Clin Transl Neurol.* 6, 655–668 (2019).
- 64. A. Carabalona *et al.*, A glial origin for periventricular nodular heterotopia caused by
 impaired expression of Filamin-A. *Human Molecular Genetics*. 21, 1004–1017 (2012).
- 9 65. J. C. Corbo *et al.*, Doublecortin is required in mice for lamination of the hippocampus but
 10 not the neocortex. *Journal of Neuroscience*. 22, 7548–7557 (2002).
- 66. S. Sahu *et al.*, Spontaneous epileptiform activity in a rat model of bilateral subcortical band
 heterotopia. *Epilepsia*. 60, 337–348 (2019).
- 13 67. E. Parrini *et al.*, Periventricular heterotopia: phenotypic heterogeneity and correlation with
 Filamin A mutations. *Brain.* 129, 1892–1906 (2006).
- 15 68. X. Piao *et al.*, G protein-coupled receptor-dependent development of human frontal cortex.
 16 *Science*. 303, 2033–2036 (2004).
- 17 69. A. Kodani et al., Posterior neocortex-specific regulation of neuronal migration by CEP85L
- 18 identifies maternal centriole-dependent activation of CDK5. *Neuron.* **106**, 246–255 (2020).
- 70. B.-I. Bae *et al.*, Evolutionarily dynamic alternative splicing of GPR56 regulates regional
 cerebral cortical patterning. *Science*. 343, 764–768 (2014).
- 71. C. De Juan Romero, C. Bruder, U. Tomasello, J. M. Sanz Anquela, V. Borrell, Discrete
 domains of gene expression in germinal layers distinguish the development of gyrencephaly.
- 23 The EMBO Journal. 34, 1859–1874 (2015).
- 72. G. M. Mirzaa *et al.*, Characterisation of mutations of the phosphoinositide-3-kinase
 regulatory subunit, PIK3R2, in perisylvian polymicrogyria: a next-generation sequencing
 study. *The Lancet Neurology*. 14, 1182–1195 (2015).

- 73. L. Telley, D. Jabaudon, A mixed model of neuronal diversity. *Nature*. 555, 452–454
 (2018).
- 74. X. Jin *et al.*, In vivo Perturb-Seq reveals neuronal and glial abnormalities associated with
 Autism risk genes. *BioRxiv.* 10, 2233–37 (2019).
- 5 75. M. Kielbinski, K. Gzielo, Z. Soltys, Review: Roles for astrocytes in epilepsy: insights from
 6 malformations of cortical development. *Neuropathol Appl Neurobiol.* 42, 593–606 (2016).
- 7 76. I. Vitali *et al.*, Progenitor hyperpolarization regulates the sequential generation of neuronal
 8 subtypes in the developing neocortex. *Cell.* 174, 1264–1276.e15 (2018).
- 9 77. R. S. Smith *et al.*, Sodium channel SCN3A (NaV1.3) regulation of human cerebral cortical
 10 folding and oral motor development. *Neuron*. 99, 905–913.e7 (2018).
- 78. F. Schottler, D. Couture, A. Rao, H. Kahn, K. S. Lee, Subcortical connections of
 normotopic and heterotopic neurons in sensory and motor cortices of the tish mutant rat. *J. Comp. Neurol.* 395, 29–42 (1998).
- 79. A. Croquelois *et al.*, Characterization of the HeCo mutant mouse: a new model of
 subcortical band heterotopia associated with seizures and behavioral deficits. *Cereb. Cortex.*16 19, 563–575 (2009).
- 17 80. A. De la Rossa *et al.*, In vivo reprogramming of circuit connectivity in postmitotic
 18 neocortical neurons. *Nat Neurosci.* 16, 193–200 (2013).
- 19 81. E. Klingler, *et al.*, A translaminar genetic logic for the circuit identity of intracortically20 projecting neurons. *Curr Biol.*, **29**, 332–339 (2019).
- 82. S. Lodato *et al.*, Excitatory projection neuron subtypes control the distribution of local
 inhibitory interneurons in the cerebral cortex. *Neuron*. 69, 763–779 (2011).
- 83. M. Thom, L. Martinian, J. G. Parnavelas, S. M. Sisodiya, Distribution of cortical
 interneurons in grey matter heterotopia in patients with epilepsy. *Epilepsia*. 45, 916–923
 (2004).
- 84. S. S. Jeste, D. H. Geschwind, Disentangling the heterogeneity of autism spectrum disorder
 through genetic findings. *Nat Rev Neurol.* 10, 74–81 (2014).

- 85. R. Mai *et al.*, A neuropathological, stereo-EEG, and MRI study of subcortical band
 heterotopia. *Neurology*. 60, 1834–1838 (2003).
- 3 86. J. A. Christodoulou *et al.*, Integration of gray matter nodules into functional cortical
 4 circuits in periventricular heterotopia. *Epilepsy Behav.* 29, 400–406 (2013).
- 5 87. S. V. Kothare *et al.*, Seizure onset from periventricular nodular heterotopias: depth6 electrode study. *Neurology*. 51, 1723–1727 (1998).
- 88. P. Chiurazzi, F. Pirozzi, Advances in understanding genetic basis of intellectual
 disability. *F1000Res.* 5 (2016), doi:10.12688/f1000research.7134.1.
- 89. E. Di Lullo, A. R. Kriegstein, The use of brain organoids to investigate neural development
 and disease. *Nat Rev Neurosci.* 18, 573–584 (2017).
- 90. S. P. Paşca, The rise of three-dimensional human brain cultures. *Nature*. 553, 437–445
 (2018).
- 13 91. M. A. Lancaster *et al.*, Cerebral organoids model human brain development and
 14 microcephaly. *Nature*. 501, 373–379 (2013).
- 92. S. Velasco *et al.*, Individual brain organoids reproducibly form cell diversity of the human
 cerebral cortex. *Nature*. 570, 523-527 (2019).
- 17 93. J. A. Farrell *et al.*, Single-cell reconstruction of developmental trajectories during
 18 zebrafish embryogenesis. *Science*. 360, eaar3131 (2018).
- 19 94. L. Telley *et al.*, Temporal patterning of apical progenitors and their daughter neurons in
 20 the developing neocortex. *Science*. 364, eaav2522–9 (2019).
- 95. T. J. Nowakowski *et al.*, Spatiotemporal gene expression trajectories reveal developmental
 hierarchies of the human cortex. *Science*. 358, 1318–1323 (2017).
- 96. A. A. Pollen *et al.*, Establishing cerebral organoids as models of human-specific brain
 evolution. *Cell.* 176, 743–756 (2019).
- 97. A. Esteva *et al.*, Dermatologist-level classification of skin cancer with deep neural
 networks. *Nature*, 1–12 (2017).

- 1 98. N. Schwalbe, B. Wahl, Artificial intelligence and the future of global health. *The Lancet*.
- **395**, 1579–1586 (2020).

1 ACKNOWLEDGMENTS

2 We thank members of the Jabaudon, Cappello and Francis laboratories for insightful 3 discussions and suggestions, Natalia Baumann and Quentin LoGiudice for their support with 4 the bioinformatic analyses, Julien Prados for setting the up 5 http://genebrowser.unige.ch/humous/ website and Sergi Roig for help in figure design. We 6 thank Stéphanie Baulac and Camilla Bellone for comments on this review and Nadia Bahi-7 Buisson and the NeuroMIG consortium for fruitful interactions and for providing gene-disorder 8 information. We thank colleagues whose work we were not able to cite due to space constraints 9 for their understanding. SC, DJ and FF laboratories take part in the NeuroMIG COST action 10 (CA16118).

Fundings: This work is supported by ERA-Net E-Rare (HETEROMICS ERARE 18-049) and
national funding agencies.

13 Author contributions: D.J., E.K., F.F. & S.C. conceived and designed the presented work.

D.J., E.K. & S.C. wrote the manuscript, with the help of F.F. E.K. performed the data analysesand designed the figures.

16 **Competing interests:** The authors declare no competing interests.

17 Data and materials availability: In Fig. 5, data from mouse embryos are from (94); data

18 from human embryos and human-derived organoids are from (95, 96).

19

1 FIGURE LEGENDS

2 Fig. 1: Key steps of corticogenesis and their susceptibility to disease. (A) Summary of 3 embryonic corticogenesis. One neuron can be followed from its birth at the ventricular zone, 4 to its migration toward the cortical plate, and then its maturation (arrow). The role of 5 cytoskeletal processes at each of these steps is highlighted on the right. (B) Literature-based 6 summary of knowledge on the involvement of these developmental steps in specific 7 neurodevelopmental disorders. The numbers of publications were found using the text mining 8 EasyPubMed R package, which automatically identifies publications containing given the 9 combinations of keywords indicated (e.g. "microcephaly" and "proliferation"). These numbers 10 were then normalized for number of publications for each of the developmental steps and each 11 of the disorders. The dendrogram represents unbiased clustering of the diseases based on this 12 data. (C) Overlapping cellular processes and pleiotropic molecular mechanisms involving 13 CDK5 across development and in the adult brain. aRG, apical radial glia; bRG, basal radial 14 glia; CP, cortical plate; E, embryonic day; IN, interneuron; IPC, intermediate progenitor cell; 15 N, excitatory neuron; pcw; post-conception week; SVZ, subventricular zone; VZ, ventricular 16 zone; WM, white matter.

17

18 Fig. 2: Cellular mechanisms at play in human cortical malformations. Cells with abnormal 19 developmental trajectories are in pink. Reduced progenitor proliferation (sometimes associated 20 with abnormal neuron migration) leads to microcephaly. Abnormal neuron position can lead 21 to agyria and lissencephaly (reduced migration and/or progenitor misposition), polymicrogyria 22 (increased migration and/or excess of basal radial glia), cobblestone lissencephaly (excessive 23 migration), periventricular heterotopia (neurons stay close to the ventricle) or subcortical band 24 heterotopia (neurons stay below the cortex). In focal cortical dysplasia, clonal mutations affect subpopulations of mature cells, which become dysmorphic (balloon cells, dysmorphic 25 26 neurons). Genes mentioned in this review are listed (bold: genes with published human-derived 27 brain organoid models, *: gene studied in the ferret, green: genes coding for transcription 28 factors). Pink arrowheads on MRIs highlight the malformations. MRI illustrations are from 29 (15) and (2) for polymicrogyria and focal cortical dysplasia (Reprinted from Semin Cell Dev 30 Biol vol 76, D. M. Romero, N. Bahi-Buisson & F. Francis, Genetics and mechanisms leading 31 to human cortical malformations, pages 33-75, 2018, with permission from Elsevier; 32 Definitions and classification of malformations of cortical development: practical guidelines; 33 M. Severino et al., Brain 2020; awaa174, with permission from Oxford University Press).

1

2 Fig. 3: From gene mutations to diseases: levels of complexity. (A) Levels of organization 3 during corticogenesis, from DNA (*i.e.* genes, represented by letters) to RNA and proteins (*i.e.* 4 gene expression), to cells, circuit and anatomy, and phenotype. Each circle represents a given 5 feature of that level (e.g. a gene, a protein, a circuit). Interactions within levels are linked 6 through complex relationships (dashed lines) to states at other levels (black lines). (B-G) 7 Examples of abnormal feature relationships across levels in disease (highlighted in pink). In 8 **B-D**, linear (monogenic), convergent (polygenic) and divergent relationships between genetic 9 defects and phenotype(s) are illustrated. (E) Hidden complexity: note that from a gene-10 phenotype point of view, this relationship appears linear. (F) Hidden hub: different genes may 11 lead to distinct phenotypes *via* hidden feature hubs influencing several downstream pathways. 12 (G) Incomplete penetrance: not all genetic features are expressed at other levels. In A, source 13 images for human brain section and phenotype are from https://msu.edi/~brains/brains/ and 14 ©biorender.com, respectively.

15

16 Fig. 4: Spatiotemporal and cell type regulations of gene expression and malformations. 17 (A) Alternative splicing of FLNA gene regulates its role in proliferation versus neuron 18 differentiation of cortical progenitors (top). Des-inhibition of the FLNA poison exon leads to 19 precocious neuron differentiation and periventricular heterotopia in humans (bottom, pink 20 arrowheads). (**B**) *DCX* mutation in human is associated with subcortical band heterotopia (pink 21 arrowheads). In mice, constitutive Dcx loss of function does not cause major cortical defects, 22 while acute uni/bilateral loss of function induces large subcortical heterotopia. Heterotopia are 23 highlighted in pink. KO, knock-out. (C) The numbers of publications containing the indicated 24 combinations of keywords (cell types and neurodevelopmental diseases) were identified with 25 the EasyPubMed R package as described in Fig. 1. (D) Basic connectivity rules in the mouse 26 somatosensory cortex. This connectivity appears largely conserved when neurons are 27 mispositioned, like in the "scrambled" cortex of Reeler mice or in case of heterotopia. ASD, 28 autism spectrum disorder; FCD, focal cortical dysplasia; HET, heterotopia; L, layer; LIS, 29 lissencephaly; MIC, microcephaly; PMG, polymicrogyria; SZ., schizophrenia. In B, source 30 image for mouse is from Cbiorender.com.

1 Fig. 5: Multimodal assessment of neurodevelopmental disorders. (A) Identification of 2 developmental gene dynamics in specific cell types through single-cell RNA sequencing (top). 3 Illustration from the online resource http://genebrowser.unige.ch/humous/ comparing data 4 from mouse, human embryos, and human derived- brain organoids (bottom). (B) Average 5 expression of genes associated with microcephaly (MIC), lissencephaly (LIS) and polymicrogyria (PMG) in mouse, human embryos, and human derived- brain organoids. 6 7 Arrowheads show human-specific patterns. (C) Schematic representation of the probabilistic 8 distribution of affected processes in sample disorders. Pink cubes define abnormal states. (**D**) 9 Hypothetical patient classification through integration of multiple datasets. Here patients were 10 clustered into 2 groups based on the integration of several features, from genes to clinical 11 expression. ATAC-seq (Assay for Transposase-Accessible Chromatin with high-throughput 12 sequencing) allows to identify chromatin accessible regions. Mass-spectrometry and co-13 immunoprecipitation allow to identified proteins based on their molecular weight and interaction(s) between proteins, respectively. In vitro and animal model data bring 14 complementary information about the cellular effect(s) of a gene mutation. aRG, apical radial 15 16 glia; bRG, basal radial glia; CSF, cerebrospinal fluid; fMRI, functional magnetic resonance 17 imaging; iN, immature neurons; IPC, intermediate progenitor cells; MCD, malformations of 18 cortical development; mN mature neurons. In B, data from mouse embryos are from (94); data 19 from human embryos and human-derived organoids are from (95, 96). The 2-dimensional 20 landscapes of gene expression across time and differentiation were performed as described in 21 (94). In B, source images for mouse and human are from ©biorender.com.



Fig. 1: Key steps of corticogenesis and their susceptibility to disease. (A) Summary of embryonic corticogenesis. One neuron can be followed from its birth at the ventricular zone, to its migration toward the cortical plate, and then its maturation (arrow). The role of cytoskeletal processes at each of these steps is highlighted on the right. (B) Literature-based summary of knowledge on the involvement of these developmental steps in specific neurodevelopmental disorders. The numbers of publications were found using the text mining EasyPubMed R package, which automatically identifies publications containing given the combinations of keywords indicated (e.g. "microcephaly" and "proliferation"). These numbers were then normalized for number of publications for each of the developmental steps and each of the disorders. The dendrogram represents unbiased clustering of the diseases based on this data. (C) Overlapping cellular processes and pleiotropic molecular mechanisms involving CDK5 across development and in the adult brain. aRG, apical radial glia; bRG, basal radial glia; CP, cortical plate; E, embryonic day; IN, interneuron; IPC, intermediate progenitor cell; N, excitatory neuron; pcw; post-conception week; SVZ, subventricular zone; VZ, ventricular zone; WM, white matter.



Fig. 2: Cellular mechanisms at play in human cortical malformations. Cells with abnormal developmental trajectories are in pink. Reduced progenitor proliferation (sometimes associated with abnormal neuron migration) leads to microcephaly. Abnormal neuron position can lead to agyria and lissencephaly (reduced migration and/or progenitor misposition), polymicrogyria (increased migration and/or excess of basal radial glia), cobblestone lissencephaly (excessive migration), periventricular heterotopia (neurons stay close to the ventricle) or subcortical band heterotopia (neurons stay below the cortex). In focal cortical dysplasia, clonal mutations affect subpopulations of mature cells, which become dysmorphic (balloon cells, dysmorphic neurons). Genes mentioned in this review are listed (bold: genes with published human-derived brain organoid models, *: gene studied in the ferret, green: genes coding for transcription factors). Pink arrowheads on MRIs highlight the malformations. MRI illustrations are from (15) and (2) for polymicrogyria and focal cortical dysplasia (Reprinted from Semin Cell Dev Biol vol 76, D. M. Romero, N. Bahi-Buisson & F. Francis, Genetics and mechanisms leading to human cortical malformations, pages 33-75, 2018, with permission from Elsevier; Definitions and classification of malformations of cortical development: practical guidelines; M. Severino et al., Brain 2020; awaa174, with permission from Oxford University Press).



Fig. 3: From gene mutations to diseases: levels of complexity. (A) Levels of organization during corticogenesis, from DNA (i.e. genes, represented by letters) to RNA and proteins (i.e. gene expression), to cells, circuit and anatomy, and phenotype. Each circle represents a given feature of that level (e.g. a gene, a protein, a circuit). Interactions within levels are linked through complex relationships (dashed lines) to states at other levels (black lines). (B-G) Examples of abnormal feature relationships across levels in disease (highlighted in pink). In B-D, linear (monogenic), convergent (polygenic) and divergent relationships between genetic defects and phenotype(s) are illustrated. (E) Hidden complexity: note that from a gene-phenotype point of view, this relationship appears linear. (F) Hidden hub: different genes may lead to distinct phenotypes via hidden feature hubs influencing several downstream pathways. (G) Incomplete penetrance: not all genetic features are expressed at other levels. In A, source images for human brain section and phenotype are from https://msu.edi/~brains/brains/ and ©biorender.com, respectively.



Fig. 4: Spatiotemporal and cell type regulations of gene expression and malformations. (A) Alternative splicing of *FLNA* gene regulates its role in proliferation *versus* neuron differentiation of cortical progenitors (top). Des-inhibition of the *FLNA* poison exon leads to precocious neuron differentiation and periventricular heterotopia in humans (bottom, pink arrowheads). **(B)** *DCX* mutation in human is associated with subcortical band heterotopia (pink arrowheads). **(B)** *DCX* mutation induces large subcortical heterotopia. Heterotopia are highlighted in pink. KO, knock-out. **(C)** The numbers of publications containing the indicated combinations of keywords (cell types and neurodevelopmental diseases) were identified with the EasyPubMed R package as described in Fig. 1. **(D)** Basic connectivity rules in the mouse somatosensory cortex. This connectivity appears largely conserved when neurons are mispositioned, like in the "scrambled" cortex of *Reeler* mice or in case of heterotopia. ASD, autism spectrum disorder; FCD, focal cortical dysplasia; HET, heterotopia; L, layer; LIS, lissencephaly; MIC, microcephaly; PMG, polymicrogyria; SZ., schizophrenia. In B, source image for mouse is from ©biorender.com.



Fig. 5: Multimodal assessment of neurodevelopmental disorders. (A) Identification of developmental gene dynamics in specific cell types through single-cell RNA sequencing (top). Illustration from the online resource http://genebrowser.unige.ch/humous/ comparing data from mouse, human embryos, and human derived- brain organoids (bottom). (B) Average expression of genes associated with microcephaly (MIC), lissencephaly (LIS) and polymicrogyria (PMG) in mouse, human embryos, and human derived- brain organoids. Arrowheads show human-specific patterns. (C) Schematic representation of the probabilistic distribution of affected processes in sample disorders. Pink cubes define abnormal states. (D) Hypothetical patient classification through integration of multiple datasets. Here patients were clustered into 2 groups based on the integration of several features, from genes to clinical expression. ATAC-seq (Assay for Transposase-Accessible Chromatin with high-throughput sequencing) allows to identify chromatin accessible regions. Mass-spectrometry and co-immunoprecipitation allow to identified proteins based on their molecular weight and interaction(s) between proteins, respectively. In vitro and animal model data bring complementary information about the cellular effect(s) of a gene mutation. aRG, apical radial glia; bRG, basal radial glia; CSF, cerebrospinal fluid; fMRI, functional magnetic resonance imaging; iN, immature neurons; IPC, intermediate progenitor cells; MCD, malformations of cortical development; mN mature neurons. In B, data from mouse embryos are from (94); data from human embryos and human-derived organoids are from (95, 96). The 2-dimensional landscapes of gene expression across time and differentiation were performed as described in (94). In B, source images for mouse and human are from ©biorender.com.