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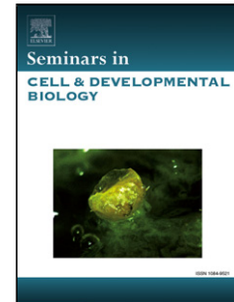
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# Conserved rules in embryonic development of cortical interneurons

By

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## Abstract

This review will focus on early aspects of cortical interneurons (cIN) development from specification to migration and final positioning in the human cerebral cortex. These mechanisms have been largely studied in the mouse model, which provides unique possibilities of genetic analysis, essential to dissect the molecular and cellular events involved in cortical development. An important goal here is to discuss the conservation and the potential divergence of these mechanisms, with a particular interest for the situation in the human embryo. We will thus cover recent works, but also revisit older studies in the light of recent data to better understand the developmental mechanisms underlying cIN differentiation in human. Because cIN are implicated in severe developmental disorders, understanding the molecular and cellular mechanisms controlling their differentiation might clarify some causes and potential therapeutic approaches to these important clinical conditions.

## Main Abbreviations:

CB, calbindin

cIN, cortical interneurons  
CGE, caudal ganglionic eminence  
CP, cortical plate  
CR, calretinin  
GABA, gamma-aminobutyric acid  
GFP, green fluorescent protein  
iSVZ, inner subventricular zone  
IZ, intermediate zone  
LGE, lateral ganglionic eminence  
MGE, medial ganglionic eminence  
MZ, marginal zone  
nNOS, neuronal nitric oxide synthase  
NPY, Neuropeptide Y  
oSVZ, outer subventricular zone  
POA, pre optic area  
PV, parvalbumin  
SP, subplate  
SST, somatostatin  
SVZ, subventricular zone  
VIP, vasoactive intestinal peptide  
VZ, ventricular zone

**Keywords:** human cortex, cortical interneurons, GABA, medial ganglionic eminence, caudal ganglionic eminence,

## Introduction

The unique cognitive abilities of the human brain most likely result from a relative expansion of regions responsible for associative and executive function in the cerebral cortex. During the course of evolution, the neocortex underwent a major expansion in its relative size and a dramatic complexification. Nevertheless, many aspects of the cortical organization and especially the underlying developmental processes, display a number of common and evolutionary conserved features in mammals. In all vertebrates the rostral-most part of the embryonic brain, called the telencephalon, is rapidly subdivided in two main regions, the dorsal pallium and ventral subpallium.

Cortical neurons can be subdivided in two major classes: the excitatory glutamatergic pyramidal projection neurons, and the inhibitory GABAergic interneurons. These two functional classes arise from spatially and molecularly segregated progenitors located respectively in the pallium and the subpallium. Importantly, the proper functions of the mature cerebral cortex require a balanced and coordinated action of these two major neuronal subtypes.

The embryonic development of the cerebral cortex relies on a highly precise orchestration of patterning events along multiple axes, and on a tight coordination of

cell proliferation, commitment, migration and differentiation. The incorrect formation of cortical organization and circuitry certainly leads to cognitive impairments and to psychiatric or neurological disorders. In particular, disturbing cortical inhibition results in severe clinical conditions, ranging from epilepsy to schizophrenia and autism disorders. This strongly suggests that the expansion of the cortex in mammals were rendered possible only by the co-evolution and expansion of inhibitory interneurons as well as excitatory neurons.

## 1. Diversity of the mature cIN

Locally projecting GABAergic interneurons, identified by Ramon y Cajal in 1899 as « short axon cells », represent a small percentage of the total neurons in the neocortex, ~20–30%, with the percentage varying substantially between cortical layers, areas and species (the remaining ~80–70% consisting in glutamatergic excitatory pyramidal neurons). In humans the percentage of cortical interneurons (cIN) is higher than in the mouse and can even reach 40–45% in the superficial cortical layers [1–3]. Ramon y Cajal already noticed a century ago that interneurons are more abundant and morphologically diverse in the primate neocortex (and particularly, in the human neocortex).

GABAergic cIN are inhibitory in the adult cortex and have a number of common features, including aspiny dendrites and local projections within the same cortical column or to adjacent columns in the cortex [4]. However they consist in a very heterogeneous population of neurons. The diverse subtypes are classified according to their location, morphology, target specificity, as well as electrophysiological and molecular characteristics. A number of excellent reviews highlight the trends in classification and nomenclature of GABAergic cIN [5–10]. The molecular criteria include different neurotransmitters and their synthesizing enzymes, neurotransmitter receptors, neuropeptides, calcium-binding proteins, transcription factors, structural proteins, connexins, ion channels and membrane transporters. The functional markers used to classify cIN are mainly the parvalbumin (PV), calbindin (CB), calretinin (CR), somatostatin (SST), vasoactive intestinal peptide (VIP), neuropeptide Y (NPY), reelin, neuronal nitric oxide synthase (nNOS)... Altogether, it has been suggest that the neocortex contains over 20 molecularly distinct classes of interneurons, and recent analyses of single cell transcriptomics further emphasized the cIN diversity [11–13]. The human and rodent cortices comprise equivalent classes of cIN although some morphological types seem specific to the human cortex [2]. cIN in rodents are presently reassembled into three main non-overlapping classes, each class expressing exclusively one out of the three following markers, PV, SST or the receptor to 5HT<sub>3a</sub> (5HT<sub>3a</sub>R) [9,14,15]. PV-expressing (PV(+)) cIN account for 40% and SST-expressing (SST(+)) cIN for 30% of the total number of GABAergic interneurons in the adult mouse cortex [15–17]. Similarly, PV and SST label two non-overlapping populations of GABAergic cIN in the adult human parietal cortex and in the somatosensory cortex of the young monkey [18]. But these two populations account for only 20% (PV(+)) and 25% (SST(+)) of the total number of interneurons in primates [18]. In primates, the relative decrease in PV and SST cIN is correlated with a strong increase in cIN expressing CR.

The attainment of subtype characteristics is progressively achieved during ontogenesis, after weeks of post-natal maturation. Dynamic changes in the expression of markers as well as specific schedules of development among species have hindered

the establishment of a complete and continuous overview of cIN development. The expression of PV is delayed during development, starting after birth in the mouse cortex, and by mid-term in the human cortex, together with SST and NPY. Certain calcium-binding proteins moreover change their cellular specificity during development. In the human embryo, CR antibodies label cortical plate neurons that resemble pyramidal neurons in addition to GABAergic neurons, and CB antibodies progressively label cortical pyramidal cells in addition to GABAergic cIN during cortical maturation [19-22].

During the last decades, fate-mapping approaches and genetics analyses in the mouse have demonstrated that the diversity of the functional markers expressed by the mature cIN largely depends on the ontogenic markers expressed by these cells at earlier stages [7,9,15]. Ontogenic markers are related to the spatial origin of the progenitors and to their date of birth. Maturing neurons can either express a temporal sequence of ontogenic markers (*Nkx2.1* and then *Lhx6* that is activated by *Nkx2.1*), or maintain forever the expression of an early marker that signs their ontogenic origin (*Coup-TFII*).

In contrast to the monkey cortex that presents very similar -although drastically shortened- sequence of prenatal development as compared to the human cortex, some embryonic structures that develop in the human cortex have no equivalent in the mouse cortex (Fig.1). However, two recent studies that analyzed in the human cortex and basal telencephalon, the distribution of ontogenic markers discovered in the mouse brain, have changed our understanding of cIN neurogenesis in humans [18,23,24].

## 2. Origin, commitment and molecular determinants of cIN

A number of experimental approaches, such as tracing studies, in utero transplantation experiments, *in vitro* migration and differentiation assays, or gene knock-out in mice, have demonstrated the subpallial origin of cIN in rodents and the preponderant role of the medial ganglionic eminence (MGE) in producing cIN [25-30], review in [31-34].

### 2.1. The MGE and CGE are the main sources of cIN in rodents: spatial and temporal origin

The embryonic subpallium of mammals is subdivided into five major proliferative regions: three ganglionic eminences (GEs) initially defined by their location, medial (MGE), lateral (LGE) and caudal (CGE), the preoptic area (POA) and the septum anlage. All these anatomical subdivisions give rise to multiple neuronal types that differentiate locally or disperse tangentially before their terminal differentiation. The **MGE** gives rise to local projection neurons for the globus pallidus, the amygdala and the septum, and to migrating interneurons destined to the cortex, striatum or hippocampus, and also produces migrating and resident oligodendrocytes [35,36]. The **LGE** produces the local striatal projection neurons and interneurons migrating to the olfactory bulb. The **CGE** is considered as a caudal extension of the MGE ventrally, and a caudal extension of the LGE, dorsally. The CGE generates the local caudal striatal and pallidal neurons, as well as different subtypes of migrating interneurons destined for the cortex (caudal regions), hippocampus, amygdala and other limbic system nuclei [37].

Lineage tracing analyses in the mouse have revealed that cIN are produced in the MGE and the CGE and, to a lesser extent in the POA [27,30,38,39](Fig.2a). The MGE generates from E10.5 to E14.5 the large majority of all cIN (50%-70%), in particular the PV and SST populations (Fig.2B, 2C). The different subtypes of MGE-derived cIN are generated at different time point in a precise temporal sequence [36,40-43]. The CGE is the second region in the basal forebrain that contributes another large contingent of cIN (30%-40% of all cIN) identified by the expression of the 5HT3a receptor (5HT3aR) and additional markers such as CR, VIP or reelin. The CGE is responsible for producing the late-born cIN [16,40,42,44]. These neurons preferentially (75%) integrate the upper cortical layers independently of their birthdate (Fig.2A, 2B). A third region, the POA produces a small fraction of cIN, about 5%-10% of highly diverse subtypes with a large geographical distribution (Fig. 2B, 2E) [39,45].

## 2.2 Genetic control of the subpallium development

Here, we present a summary of the molecular mechanism that pattern the GEs and specify the fate of cIN generated in the mouse GE subdivisions (Fig.2B) and describe the expression pattern of same genes in the human telencephalon (Fig.3A, B).

Several families of transcription factors control the development of the subpallium and specify the fate of neurons originating in its neuroanatomical subdivisions. Some developmental genes are common determinants whereas others generate regional diversity.

### 2.2.1. cIN determinants with a widespread expression in the subpallium

Genes widely expressed within the three ganglionic eminences, especially the transcription factors *Ascl1* or *Dlx*, control the embryonic development of the subpallium and the generation and/or differentiation of GABAergic cells, including cIN. Mice lacking *Ascl1* or *Dlx1&2* in the GE lack 50% to 80% of cortical GABAergic neurons, confirming the origin of cortical GABAergic neurons in the ventral telencephalon [46-48] and demonstrating the existence of parallel and overlapping mechanisms of GE patterning [49,50].

*Ascl1* (also known as *Mash1*) is a proneural transcription factor strongly expressed in the subpallial ventricular zone (VZ) and subventricular zone (SVZ) (Fig.2B) required for the neurogenesis of GABAergic neurons [47,48,51,52] and for the generation of oligodendrocytes by restricting the number of *Dlx*-expressing progenitors [53,54]. In the mouse, *Ascl1* is expressed at very low level in pallial progenitors [47,55]. In primates, *ASCL1* shows a clear frontier of expression at the pallium/subpallium boundary, because *ASCL1*-expressing (*ASCL1*(+)) cells are numerous in the GE and sparse in the pallial VZ and SVZ (Fig.3B, Fig.4). According to Hansen et al. [23] and Ma et al. [18], *ASCL1*(+) cells located in the pallium co-express pallial markers, PAX6 or TBR2, but neither GAD65/67 nor GABA, suggesting that they very unlikely contribute to the GABAergic lineage. Because *Ascl1* is maintained in a proportion of cIN that reach the cortex [16], the pallial or subpallial origin of *Ascl1*(+) cells that later differentiate as GABAergic interneurons remains controversial and needs to be clarified.

*Dlx* homeobox genes (*Dlx1&2*, *Dlx5&6*) show a widespread expression throughout the subpallium [56,57] and are required for the differentiation of the three

GEs. They are also involved in the craniofacial development. In the subpallium, *Dlx1&2* expression commences as progenitors transit from the VZ to the SVZ and is extinguished after their phase of migration (Fig.2B) [58,59]. In the human brain, *DLX2* is strongly expressed in GE progenitors and more faintly in cells located in the cortex (Fig.3B, Fig.4A-C) [23,60]. The capacity of *DLX2* cells to proliferate in the human cortex is debated [23,61]. *Dlx1&2* promote the expression of glutamic acid decarboxylase (*Gad67*, also known as *Gad1*) and vesicular GABA transporter (*VGAT*) [62,63]. *Dlx* genes are thus essential for both the differentiation of GABAergic interneurons and the tangential migration of their precursors. *Dlx1* and *Dlx2* are functionally redundant and a 80% reduction in GABAergic cIN is uniquely observed in mice with *Dlx1&2* ablation [25,29,46,64]. *Dlx2* activates *Dlx5* gene expression [58]. Accordingly, *Dlx5&6* are expressed in more mature cells than *Dlx1&2* [65].

### 2.2.2. cIN determinants with a restricted expression in the subpallium

#### 2.2.2.1. Restricted to the MGE

*Nkx2.1 (Ttf1/Ttf1)* is a homeobox gene expressed early in the VZ and SVZ of the MGE and the POA, which controls the formation of the MGE and regulates the neurogenesis and differentiation of two main populations of MGE-derived GABAergic interneurons, the PV(+) and SST(+) cIN (Fig.2B) [27,40,41,66,67]. *Nkx2.1* expression is induced by sonic Hedgehog (SHH) in the ventral forebrain [68]. SHH expression and signaling level differ in the dorsal and ventro-medial parts of the MGE. PV(+) and SST(+) cIN seem specified by different levels of SHH. They are produced in different MGE regions, with SST(+) interneurons induced by higher level of SHH signaling in the dorsal MGE [17,69-71], Review in [72].

In the mouse, the genetic ablation of *Nkx2.1* results in 50% loss of GABAergic cIN, principally early migrating *Dlx2*-expressing (*Dlx2*(+)) cells, but spares *Dlx2*(+) cells that originate in the CGE. *Nkx2.1* seems to regulate the switch in the class of neurons generated at different time point and fated to different targets. For example, *Nkx2.1* is downregulated in GABAergic post-mitotic cells that migrate to the cortex, but maintained in GABAergic interneurons migrating to the striatum [73,74].

In humans, the localization and extension of the MGE has been identified recently in a KI67(+) rostral-medial domain of the GE thanks to the strong expression of *NKX2.1* (Fig.3B) [3,18,75]. The persistence of *NKX2.1* expression in cIN migrating through the cortex is debated. Two groups reported scattered *NKX2.1*-expressing (*NKX2.1*(+)) cells at GW24 and suggested that *NKX2.1* is down regulated in MGE cells after they reach the cortex, as in mice [18,23]. A third group [60] described *NKX2.1*(+) cells among cortical progenitors, and found a high percentage of KI67(+) and *PAX6*(+) cells (80%) in the *NKX2.1*(+) population (labeled “€” in Fig.4C,D). These authors reported nevertheless a low percentage of *NKX2.1*(+) cortical progenitors (3-5% of VZ cells and 5-9% of SVZ cells at GW15-GW22). *NKX2.1*(+) cells thereafter distributed in the intermediate zone (IZ) and subplate (SP). On the second half of the gestation, they distributed in all cortical layers and co-expressed PV or SST (Fig.4C,D) [76].

*Lhx6* is a transcription factor of the LIM homeodomain family directly activated by *Nkx2.1* and essential for the tangential migration of cIN precursors and for their differentiation in the cortex [27,77-79]. *Lhx6* is expressed in post-mitotic cIN lacking progenitor markers and is maintained in both PV(+) and SST(+) MGE-derived cIN

(Fig.2B) [27,42,57,78-81]. *LHX6* is also expressed in human MGE-derived cIN (Fig.3B) [23]. *Lhx6* is sufficient to rescue the production of cIN from *Nkx2.1*-/- subpallial cells, suggesting that the major function of *Nkx2.1* is to initiate the transcription of *Lhx6* in cIN progenitors [64]. *Lhx6* functions in part through promoting *Arx* and *Cxcr7* expression [82].

*Sox6* functions downstream *Lhx6* and is expressed in MGE-derived cIN from post-mitotic stage to adulthood. *Sox6* is required for normal PV(+) and SST(+) interneurons development, final positioning and maturation into the cortical layers (Fig.2B) [7,83]. In the developing mouse forebrain, *Sox6* is strongly expressed in MGE-derived cIN and, at lower level, in pallial VZ where it cooperates with *Ngn2* to repress *Ascl1* and maintain pallial identity and pallial/subpallial patterning [83]. The MGE in human strongly expresses *SOX6*. 80% of PV(+) cells and 90% of SST(+) cells co-express *SOX6* in the adult cortex, suggesting that they originate in the MGE (Fig.3B) [18]. A population of *SOX6*(+) pallial progenitors co-expresses *PAX6* but not GABA, and likely represents progenitors of cortical glutamatergic projection neurons (Fig.3A, Fig.4C).

#### 2.2.2.2.restricted to the CGE

CGE-derived cIN migrate later than MGE-derived cIN, do not require *Nkx2.1* function and differentiate as CR(+) and VIP(+) neurons [40,59,66]. The molecular signals involved in their fate determination are much less characterized but involve *Gsx2*, *CoupTFII*, and *Prox1* [17,84-86]. Gene expression profiles in CGE and LGE are largely overlapping, suggesting that the CGE is a caudal expansion of the LGE (Fig.2D) [56,57,87]. In the mouse, cIN generated outside of the *Nkx2.1*-expressing proliferative domains derive from CGE progenitors and can be traced through their expression of *Prox1* in post-mitotic cells [86,88]. *Prox1* has a crucial role in the acquisition of CGE-derived cIN properties and is maintained in the mature cortex (Fig.2B) [86].

All CGE-derived cIN express the serotonergic ionotropic receptor **5HT3aR** [89]. 5HT3aR is expressed in migrating and mature CGE- and POA-derived cIN that express neither PV nor SST, e.g. bipolar CR(+) (CALB2), VIP(+), CCK(+), reelin(+), and the multipolar NPY(+) cIN [14,89-92]. This receptor is presently used to define in the mouse a cell population that represents about 30% of cIN, targets the superficial cortical layers and comprises a variety of functional subpopulations (Fig.2B) [14,17]. Because of the lack of antibodies against 5HT3aR, these interneurons have been characterized in transgenic mouse lines (Lee et al, 2010, [91] and the receptor cannot be used to identify the non-PV and non-SST GABAergic interneurons in human and non-human primates.

*Coup-TFII* (*Nr2f2*) is an orphan nuclear receptor enriched in the CGE and required for the tangential migration of CGE-derived cIN in the mouse [84,93]. A small subset of *Coup-TFII*-expressing (*Coup-TFII*(+)) interneurons derives from the dorsal/caudal part of the MGE and co-expresses *Sox6* [83]. Therefore, *Coup-TFII* shows a minimum overlap with *Sox6* (Fig.2B).

In the human subpallium, *COUP-TFII* is expressed in a rostral-to-caudal increasing gradient (from LGE to CGE) identifying the CGE as a strongly labeled structure, with a high density of positive cells in the ventral part of the CGE (vCGE, Fig.3B) [75]. VZ and SVZ progenitors of the vCGE co-express KI67. *COUP-TFII*(+)

cells in the dorsal CGE and LGE (dCGE, dLGE, Fig.3B) are post-mitotic and co-express SP8 (see below). They could derive from the CGE and migrate toward the cortex [18,23]. *COUP-TFII* shows an abrupt decrease in expression at the pallium-subpallium boundary, but remains expressed at low level in the cortex [75]. The population of COUP-TFII(+) cortical cells is radially-oriented and co-expresses KI67 and PAX6 (Fig.4C,4D) [75]. This population very likely generates glutamatergic pyramidal neurons destined to cortical layers V and VI, which express COUP-TFII at low level [94].

Despite the small population of COUP-TFII(+) cells born in the MGE, *COUP-TFII* defines a third population of cIN that does not overlap with the PV(+) and SST(+) populations in the human and monkey cortex [18]. *COUP-TFII* is expressed in cIN that largely co-express the calcium-binding protein CR (80% of COUP-TFII(+) interneurons[94]), showing that CR, together with reelin, are two fairly good markers of the same population [18]. In the adult human cortex, interneurons expressing COUP-TFII are distributed in the supragranular layers and exhibit specific physiological properties [94].

*Sp8* (buttonhead-like zinc finger transcription factor) labels cIN derived from the dCGE (Fig.2B) and possibly from the dLGE [95]. *Sp8* is also strongly expressed in the dLGE, where it is required for neurogenesis, migration and survival of olfactory bulb GABAergic interneurons [96]. *Gsx2*, a downstream target of SHH, controls the differentiation of interneurons upstream *Sp8* [96-99]. In addition to regulating striatum development, *Gsx2* is particularly enriched in the LGE/CGE and contributes to the specification of NPY(+), bipolar CR(+) and VIP(+) CGE-derived cIN [42]. *Sp8*-expressing (*Sp8*(+)) cIN are born at later embryonic stages (peak at E15.5) than MGE-derived cIN. *Sp8* is moreover expressed at low level in the pallium where it controls the specification of cortical projection neurons [96,100].

In humans, *SP8* is strongly expressed in the SVZ of the dLGE and dCGE (Fig. 3B). *SP8* is weakly expressed in the cortical VZ and SVZ (Fig.4C,D). Because *Gsx2* is not expressed in the human cortex, SP8(+) cortical progenitors unlikely generate interneurons. They moreover express PAX6 and likely generate excitatory projection neurons. At adult stage, in human and monkey, the vast majority of CR(+), VIP(+) and reelin(+) cIN express COUP-TFII and/or SP8, strongly suggesting that they are derived from the dLGE and CGE. These interneurons preferentially occupy the superficial cortical layers [101]. Altogether, SOX6(+), SP8(+) and COUP-TFII(+) cells account for nearly 90% of cIN [95].

### 2.2.2.3. restricted to the POA

The POA is ventral and immediately adjacent to the MGE. Progenitor cells in both regions express *Nkx2.1*. Cells in the POA moreover express specific markers (*Dbx1* and *Shh* in the VZ, *Nkx5.1* and *Nkx6-2* in the SVZ), which are not expressed in the MGE. Reciprocally, *Olig2*, which is highly expressed in the MGE, is virtually absent from the POA [57,67,102]. In mice, the POA contributes a minor but highly diverse pool of cIN expressing PV (50% of them in the somatosensory cortex), SST (25%) or reelin (15%), and for a very small number, CR, NOS or VIP (2-4%) [39,67].

POA derived cells first migrate in a rostro-medial direction through the septal region to invade the cortex at intermediate or rostral telencephalic level. Migratory routes are mostly through the marginal zone (MZ) and the subplate (SP) [67].

Finally, the possibility that a small fraction of the LGE (dLGE) produce some cIN is not entirely ruled out, neither in rodents, nor in humans [25,26,28-30,88,103-107].

### 3. The question of the pallial origin of cIN

The relative increase in CR(+) interneurons observed in the human cortex as compared with the rodent cortex (illustrated in Fig.3A, to compare with the upper panel in Fig.2B), was initially attributed to a local production of GABAergic interneurons within the cortex and considered as a specific feature of the primate cortex [61,76,108]. This initial assertion was further supported by human cases with severe holoprosencephaly lacking substantial parts of the GE and showing the persistence of CR(+) neurons in the cerebral cortex [110].

The capacity of cortical progenitors in the human cortex to generate GABAergic interneurons is presently a controversial issue. This proposal was initially supported by two observations: 1) GABAergic neurons in the human cortex co-express *ASCL1*, and 2) pallial progenitors in humans do express *ASCL1* and/or *DLX2*, two genes in the GABAergic lineage. As a consequence, it was proposed that *ASCL1*(+) pallial progenitors produce GABAergic neurons. We report below some arguments against.

Several studies in human showed that a large proportion of GABAergic cells (either GABA(+) or GAD65(+)) co-express *ASCL1* in the human pallial SVZ, subplate (SP) and cortical plate (CP) (horizontal arrows in Fig.4A,B) [61,109,111]. By this time, *Ascl1* and GABA (or GAD) co-expression had not been reported in the mouse cortex because cIN generated in the mouse subpallium down regulate *Ascl1* after reaching the cortex (but not *Dlx2*) [47,48,58,59]. The co-expression of GABA and *ASCL1* seemed thus unique to the human brain, and it was proposed that GABAergic neurons co-expressing *ASCL1* represent a pool of interneurons produced in the pallium by *ASCL1*(+) pallial progenitors. A later study performed in mice revealed however, that CGE-derived cIN maintain the expression of *Ascl1* as they migrate to the cortex as post-mitotic cells expressing Tuj1 or Map2 [16]. This study suggests that GABAergic neurons produced in the human subpallium can continue to express *ASCL1* after reaching the cortex and that *ASCL1*(+) GABA neurons present in the pallium can originate in the subpallium. As a consequence, two distinct cell lineages in the human telencephalon would express *ASCL1*: i) the cortical progenitors of excitatory neurons and ii) the CGE-derived precursors of cIN.

The pioneer study of Letinic et al. [61] reported the expression of *ASCL1* and *DLX2* in dividing cells in the human cortex at GW14-24. Pallial progenitors in the human cortex were identified by their location in the cortical VZ/SVZ and by markers of proliferation (KI67 or BrdU incorporation). Studies in the mouse showed accordingly that transcription factors regulating cIN neurogenesis are expressed in areas that largely overpass the ventral subpallium and often includes the dorsal pallium (Cf. §2.2.1 and §2.2.2.2 above). This is the case for *ASCL1* and also *SP8*, *COUP-TFII*, or *SOX6* (Fig.4C,D). Because these genes function in different genetic networks in their different areas of expression, they can both control the differentiation of GABAergic cells or oligodendrocytes in the subpallium and contribute to the differentiation of excitatory cortical cells in the pallium [83,96,100]. Therefore, the *ASCL1*(+) progenitors observed during the second trimester of gestation in the human cortex and from E64 to E88 in the monkey cortex

[61,108,109] cannot be attributed to the GABAergic lineage uniquely on the base of their expression of *ASCL1*.

A remaining question is whether transcription factors such as *DLX2* and *NKX2.1* that control the differentiation of cIN in the subpallium, can orient the fate of dorsal progenitors toward a novel dorsal GABAergic lineage in the human forebrain. There is no consensus among authors, neither on the expression level of *NKX2.1* in cortical progenitors, nor on the role of *NKX2.1* in human cortical neurogenesis (Fig.4C,D) [22,60,75,76]. Recent extensive analyses in the human cortex at GW10-14, revealed an almost null percentage of KI67(+) cells among the DLX2(+) cortical cells, a result confirmed by BrdU labeling experiments in cultured forebrain slices [23]. Ma et al. [18] did not find NKX2.1(+) cortical progenitors. In contrast, Radonjic et al. [76] described a small percentage of NKX2.1(+) cells among cortical progenitors at GW15-22, which could well contribute to a significant population of cIN because of the large size of the oSVZ at this stage [76]. In vitro, *NKX2.1* induces *LHX6* expression in pallial progenitors, indicating their commitment to the PV and SST lineages. SHH application on cortical progenitors increases the pool expressing NKX2.1, at the expense of CR(+) cells [119]. Since the human cortex is remarkable by the increased proportion of CR(+) cIN, the putative contribution of cortical cIN progenitors to the PV/SST lineage is somehow puzzling and suggests an important role of the SHH signaling pathway in regulating the neurogenesis of CR(+) cIN in the human cortex.

The suggestion of a pallial origin of cIN in the human cortex is further supported by the fact that 17% of the total number of CR(+) cells are co-labeled with KI67 antibodies in the oSVZ of the human cortex [22]. These dividing CR(+) cells either originate from the pallial precursors or come from the subpallium and undergo additional round of proliferation during their migration into the cortex [22,60]. Cells migrating from the GE to the cortex at early embryonic stages do not seem to divide ([118] and our own observations), although this possibility cannot be totally excluded. Indeed, it has been reported in the mouse that GAD65(+)CR(+) interneuron precursors derived from the CGE-LGE, can re-enter the cell cycle in the dorsal white matter after an initial phase of tangential migration [105]. Alternatively, the apparent pallial origin of certain olfactory interneurons [117] suggests that some cortical progenitors have the capacity to produce inhibitory neurons. And small numbers of mouse cIN are produced in the cortical SVZ after birth and completion of excitatory neuron production [116]. However, after an extensive search for the proliferative state of CR(+) interneurons in numerous cortical regions of human fetuses at GW10–14, Hansen et al. [23] concluded that virtually all cortical CR(+) and DLX2(+) inhibitory neuron precursors are non-proliferative in the human cortex.

The diverging conclusions discussed above should not hide that a major breakthrough was recently made by Hansen et al. [23], when they demonstrated a considerable increase in size of the primate CGE as compared with the CGE in rodents (illustrated in Fig.2E and Fig.3C). Other groups have confirmed this remarkable and novel observation [18,3]. Since the majority of CGE-derived cIN express CR, the dramatic increase in CR(+) cIN in the human cortex very likely results from the spectacular enlargement of the CGE. The number and diversity of GABAergic interneurons have increased concomitantly with the number of excitatory projection neurons so that a balance between excitatory and inhibitory activities was maintained in the human cortex [111,112]. In human and non-human primates, the

production of an increased number of excitatory projection neurons is related to the emergence of a new proliferative compartment, the outer subventricular zone (oSVZ). Such a neurogenic extension occurred thanks to developmental changes particularly visible around midterm in human, when both oSVZ and CGE are particularly developed (Fig.1). It has to be mentioned that the enlargement of both oSVZ and CGE in humans has no equivalent in rodents [113-115] and likely represents a derived feature of primates.

#### **4. A conserved subpallial origin of cIN in all gnathostomes, and conserved migratory properties.**

Even if authors disagree on the regional origin of cIN during the late phase of neurogenesis in primates, most of them agree that the GEs are the main sources of GABAergic neurons during the first trimester of gestation in humans and until E55-E65 in the *Cynomolgus* monkey [22,60,108,109,114]. Based on the continuity of temporal sequences of expression of GABA and of SOX6, COUP-TFII, SP8 transcription factors in the telencephalic vesicles, the studies of Ma et al. and Hansen et al. [18,23] show that the majority, if not all, GABAergic interneurons originate in the GEs in both human and non-human primates. These studies moreover reveal that developmental mechanisms responsible for the production of GABAergic interneurons are extremely well conserved in vertebrates [18,23].

The regionalization of the forebrain, with distinct domains marked by differential, specific gene expression patterns (or genoarchitecture), shows an impressive degree of conservation across all gnathostomes and might be dated back to more than 500 million years ago when the latest common ancestor of vertebrates arose [120-123].

Comparative analyses of orthologous genes expression in chick, turtle, frog, zebrafish or lamprey have shown that forebrain patterns are largely conserved among these species [122-127]. Thus, the basic developmental plan of the subpallium in the telencephalon has a deep evolutionary origin and the GE origin of cIN appears highly conserved in all Gnathostomes [128-130].

In mice as in other mammalian species examined so far, the capability to migrate long distances tangentially to the brain surface toward the dorsal pallium is a cardinal property of the GABAergic lineage born in the basal forebrain. The first experiments that evidenced the long distance migration of cINs from the subpallium to the developing cortical plate, consisted of vital dye injections in organotypic slices of mouse embryos. Precursors of cIN labeled in the subpallium migrated long distances in the organotypic cortical slices and reached the cortex after one or two days in culture [25,26,103]. Analyses of the long distance migration of cIN precursors strongly benefited from the development of transgenic mouse strains that express fluorescent markers in cIN [131,132]. Live cell imaging of cIN expressing fluorescent proteins (such as GFP) in slices of embryonic cortex helped to identify the migratory routes of cIN within the cortical wall [133,134]. Immature cIN first enter the cortical wall by a tangential route located either superficially in the marginal zone (MZ) or deep in the SVZ/lower IZ. After the CP differentiates, an additional tangential route forms in the SP. cIN then reorient radially and move to the CP where they settle and differentiate (Fig.2D,2E). A proportion of cIN follows oblique trajectories rather than tangential ones, and the radial re-orientation can be directed toward the VZ rather

than the CP. In various mammalian species examined so far (rat [135,136], opossums [137], cynomolgus monkey [109]) as well as in the human brain [22,61,138,139] GABAergic cIN migrate tangentially from the subpallium to the pallium via the MZ and SVZ/IZ routes (Fig.4), as described in the mouse. A tangential mode of migration is already present in infra mammals, in the chicken, turtle, sharks and lamprey where MGE cells migrate from the subpallium to the dorsal pallium along more or less well defined pathways [126,127,140].

The intrinsic migratory ability of subpallium-derived GABAergic cIN was tested by interspecies transplantations. Ex vivo interspecies transplantations showed that turtle MGE cells transplanted into the mouse MGE migrated tangentially and entered the mouse neocortex, and vice versa [127], suggesting that the ability of MGE cells to enter the neocortical primordium had already been established in the most recent common ancestor of amniotes. In utero interspecies transplantation of chicken, turtle or marmoset MGE cells into the mouse MGE revealed that, in contrast to mouse and marmoset MGE cells, a majority of chicken and turtle MGE cells fail to enter the cortical plate and remain in the SVZ/IZ [130]. Nevertheless, 29 days after transplantation the chicken MGE cells had differentiated into GABAergic cIN within the mouse neocortex and expressed either PV or SST. Therefore, the intrinsic ability of MGE cells to generate both PV(+) and SST(+) cells is conserved in all amniotes.

## 5. Migration routes of cIN in the human brain

### 5.1 Migration routes toward the cortex

Dye injections in forebrain slices from human fetuses at GW15 [61] label a significant number of GE cells that migrated to the cortical VZ/SVZ (Fig.4f in [61]). This experimental approach is highly challenging in human, for ethical and technical reasons [18,23], notably because of the length, duration and complexity of cIN trajectories in the human cortex. Therefore, analyses in humans are restricted to small portions of cIN trajectories [139]. The analysis of cIN migratory pathways in the developing human brain thus essentially relies on neuroanatomical observations on fixed tissue using markers stably expressed in cIN, both spatially and temporally. GABA, GAD, SOX6, SP8 or COUP-TFII are stable and reliable markers of cIN (however not exclusive to cIN). Whereas SOX6, SP8 and COUP-TFII antibodies stain nuclei, GABA and GAD distribute in the whole cell including thin protrusions. GABA immunostaining reveals the morphology of cIN precisely enough to identify their polarity and their direction of migration. GABA immunostaining moreover reflects the activity of the glutamic acid decarboxylases (GAD), whose expression is activated by DLX proteins [65]. The two isoforms 65 and 67 (also known as Gad1) are co-expressed in almost all GABAergic neurons in the mouse, monkey and human adult brain[21,65,109].

Before genetic studies established that GABA and GAD are reliable ontogenic markers, careful and detailed descriptions of the distribution and morphology of cells expressing GABA and/or GAD had been performed in the developing cortex of both rodents and primates [109,141-146]. Authors took advantage of the polarized morphology of GABA neurons that exhibited growth cones at the end of long leading processes, for identifying the direction of movement of GABAergic neurons in the

developing cortex. By correlating the putative directions of movement with the changes in distribution of the GABA neurons, authors inferred from analyses on fixed tissue that pioneer GABA cells have a lateral to medial direction of movement in the developing cortical wall. However, complex morphological changes associated with trajectories reorientation and transient polarity reversals are major limitations of these analyses on fixed tissue. The rodent subplate is a typical decision region where cIN exhibit complex morphologies, which had not been associated with trajectories reorientations in early studies [141,142]. In addition, the radial re-orientation of cIN away from their tangential routes of migration progressively hampers tangential streams of cIN in fixed sections and prevents analyses of the late stages of migration. Because GABA cells were found in two transient structures of the developing cortex, the cortical subplate and the marginal zone, it was initially proposed that GABAergic cells are pioneer cells that disappear with these transient structures.

In primates, the large size of structures and the long duration of developmental phases make the identification of streams of GABAergic cells easier than in rodents. GABAergic cells oriented tangentially (i.e. parallel) to the brain surface are organized in continuous and dense streams between the subpallium and pallium. A large stream of GABAergic cells was first described between the GE and the cortex in the rostral part of the telencephalic vesicles in human [22,60,61]. GABA and GAD65 immunostaining labeled similar migratory stream in the rostral telencephalon of the Cynomolgus Monkey [109]. In the human brain, Ma et al. [18] and Hansen et al. [23] have recently identified large streams of cells that express subpallial genes and extend between the GE and the developing cortex. Transcription factors expressed either in the MGE, or in the ventral and dorsal parts of the CGE label distinct migratory streams and identify different routes of migration between the GE and the cortex. Cells in streams usually do not express proliferation markers. Two main routes of migration have been described. NKX2.1(+) cells materialize the beginning of the route between the MGE and the cortex, which extend through the LGE. SOX6, a stable marker of MGE-derived cells, identifies the end of the same route that crosses the LGE and reaches the lateral and anterior cortex. A second route of migration is identified by large and continuous streams of cells expressing COUP-TFII or SP8, which extend between the CGE and occipital cortical regions [18,23,75]. As in the mouse, interneurons from the CGE migrate caudally toward the hippocampus and the caudal cerebral cortex [44] under *Coup-TFII* control [84]. A study from Clowry and coll. in humans [75], shows different cortical distribution of cIN generated in distinct parts of the GE, strengthening previous observations of Ma et al [18], who found different ratios of CR and PV/SST cIN in the frontal and caudal cortex.

Doublecortin (CDX) is another marker expressed in young migrating neurons in the human cortex [147], which reveals interneuron morphology. DCX recently helped to identify the directions of movement of a population of GABAergic neurons present at birth in the human frontal cortex. These neurons moreover express transcription factors identifying their ventral telencephalic origin, in particular SP8, COUP-TFII, NKX2.1, or LHX6 [139].

## 5.2 Migration routes within the cortex

### 5.2.1 Embryonic and fetal stages

In the human brain, the first GABAergic cells are observed at the surface of the cortical anlage at very early stage (GW6.5-7) when the cortical wall only comprises the proliferative neuroepithelium (VZ) and a superficial layer of post-mitotic cells, the

plexiform layer (PPL, Fig.1) [21,138]. At this stage, GABAergic cells distribute in a latero-medial gradient. Most of them exhibit either bipolar or unipolar morphologies suggestive of migrating cells. They often extend tangentially to the brain surface one long process ending with a growth cone. Very few radially oriented GABA(+) cells are present in the VZ [60]. Radially oriented GABA(+) cells could either originate locally or, more likely, colonize the VZ from the superficial PPL using a ventricle-directed migration as described in organotypic cortical slices from rodents [148,149]. Accordingly, some radial DLX2-expressing cells that co-express MAP2, a neuronal marker, are also observed in the VZ at same stage [60]. These neurons could migrate from the PPL/MZ that contains tangentially oriented cells co-expressing DLX2 and MAP2 (Fig.4A). Young DLX2(+) neurons likely migrate from the GE to the cortex as observed in the mouse.

As development proceeds, GABAergic cells become more numerous in the human cerebral wall and distribute between the MZ and the VZ in the developing CP (Fig. 4B). Regional differences in cell density are no longer observed at GW11. GABA(+) cells extend horizontal/tangential or oblique processes, and structures resembling growth cones are observed at the tip of the processes. From GW11, another population of tangentially oriented GABAergic cells appears in the lower part of the IZ and in the SVZ [138,143]. Some horizontally oriented GABA(+) cells distribute along the ventricle. GABAergic cells in the cortical VZ and SVZ express markers of young neurons (TUJ1, MAP2) showing that they are post-mitotic immature neurons extending long processes parallel to the surface of the ventricle.

This early distribution and laminar location of GABAergic cells corresponds well to the distribution in rodents and nonhuman primates [150,151]. In the CP, GABAergic cells exhibit preponderant radial orientation suggesting that they use both radial and tangential migration to disperse in the developing cortex [138].

From GW14 to midterm [22,143] (Fig.4C), the density of GABAergic cells increases in the SVZ/IZ, SP and MZ. Authors report variable orientations of process, but agree on a large proportion of tangentially oriented cells in the VZ, IZ and oSVZ. In the IZ and SP, horizontal processes intersect radially orientated GABAergic cells. At these stage, the distribution, morphology and orientation of CR(+) cells is reminiscent of GABAergic cells (Fig.4C) [22].

During the second half of the gestation (Fig.4D), the peak of GABA immunoreactivity moves superficially in the developing CP, where the proportion of multipolar cells increases in comparison with bipolar GABAergic cells [143].

In the Cynomolgus Monkey, a similar sequence of development is observed for GAD65-expressing (GAD65(+)) cells [109]. A large stream of tangentially oriented GAD65(+) cells distributes in the lower IZ/upper SVZ of all cortical regions at E55 (equivalent of GW13/14 in human). Smaller streams of GAD65(+) cells are also observed in the MZ and SP. At a stage corresponding to midterm in human, GAD65(+) are numerous in the VZ/SVZ and GAD65(+) cells lacking the horizontal orientation observed at early stages, distribute in the whole thickness of the cortical wall.

### 5.2.2 Post-natal migration of cIN

Neuronal migration is considered to be largely complete before birth, except for specific population of GABAergic interneurons that contributes to the olfactory bulb and, in a limited extent, to the anterior forebrain [152]. Throughout life, neural stem cells in different domains of the SVZ of the adult rodent brain generate several subtypes of interneurons that travel from the walls of the lateral ventricle, along the rostral migratory stream (RMS) and add to circuits into the olfactory bulb [153,154]. In humans, a sub-population of young SVZ-derived neurons migrate along a ventral route branching off the RMS, the medial migratory stream (MMS) and settle in the medial prefrontal cortex [154]. At birth in human neonates, although many SP8(+) and/or COUP-TFII(+) cells already reached the superficial cortical layers, some of them are still present in the cortical SVZ [18], suggesting that their migration to the cortex continues during infancy [139,154]. Post-mortem analyses in infant human brains have recently established that a large population of young neurons still migrates at birth in the frontal lobe of the cortex [139]. This population will integrate the frontal cortex during the first months of post-natal life. These young neurons express DCX and GABA, but neither progenitors nor pallial markers. Most of them express subpallial markers showing that they likely originate in the CGE, MGE, and possibly LGE. They concentrate at birth in the anterior horn of the lateral ventricle, where they distribute along the ventricle and blood vessels in an arc-shaped structure. Infection of tissue with GFP-expressing adenovirus reveals that these cells migrate using a saltatory mode of progression identical to the mode of migration of GABAergic neurons in the embryonic cortex [133,155,156]. Near the ventricle, cell trajectories are predominantly tangential to the brain surface. Within the cortical parenchyma, young neurons reorient their trajectories perpendicular to the cortical layers [139].

These novel data, by showing that a pool of young post mitotic neurons are stored in tangential migratory streams after birth, suggest that cIN can be integrated postnatally in perhaps fully functional circuits. This opens new avenues for understanding the clinical consequences of perinatal strokes and experience-dependent functional maturation of the cortex.

## 6. Conclusion

cIN are implicated in a wide range of neuropsychiatric disorders as epilepsy, bipolar disorders, attention deficit hyperactivity disorder (ADHD), schizophrenia and autism. The heterogeneity of cIN and their complex and long-lasting phase of migration could make them particularly sensitive to a wide variety of intrinsic (genetic) and extrinsic (environmental, functional, ...) factors that can alter their development and lead to defective cortical circuits. For example, dysfunction or cell death of specific types of GABAergic neurons has been described as a hallmark of various psychiatric and neurological disorders, such as schizophrenia and epilepsy. Several studies associate dysfunction of PV(+) interneurons in frontal cortex with schizophrenia [157-160]. Neocortical interneurons in the upper layers are believed to be involved in the physiopathology of these psychiatric disorders [161]. For example, the specific removal of *Lhx6* and *Sox6* from MGE-derived cIN leads to abnormal PV and SST expression related to mispositioning of basket and Martinotti cIN. As a

consequence, this results in a severe epileptic encephalopathy [162]. An imbalance between excitation and inhibition in the cerebral cortex has also been suggested as a possible etiology of autism [163]. Interestingly, *DLX1 & 2* polymorphism is associated with autism [164]. Beside genetic mutations, intracerebral hemorrhage observed in the cortical SVZ in premature babies and neonates [165] may have serious consequences on later brain development. Hemorrhage can destroy a large number of interneurons and oligodendrocytes destined for the cerebral cortex and still located in the tangential migratory streams [166]. By depleting neonate or preterm brains in cIN, intracerebral hemorrhage could be implicated in schizophrenia and in neuropsychiatric disorders resulting from alteration in the excitation/inhibition balance [167]. The recent study of Paredes et al. [139], by opening a new and previously unexpected chapter of the biology of interneurons in the human cortex, give us a glimpse of a potential cell-based therapy in the treatment of neurological disorders, even if any plan to correct brain activity should be based on a stronger knowledge on the normal and pathological human brain development. Although caution should be the rule with mouse–human extrapolations [3], it remains that the power of genetic studies in mouse models is considerable and served well our increasing knowledge of the human brain development. Studies in the mouse are at the origin of the recent highlight on a large territory in the human basal forebrain that expresses CGE specific genes and likely contributes the numerically and functionally important population of CR-expressing cIN in humans. Reciprocally, incoming observations in humans should provide valuable insight to further understand the mouse model and the regional differences in the migration and maturation of cortical interneurons.

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## Caption

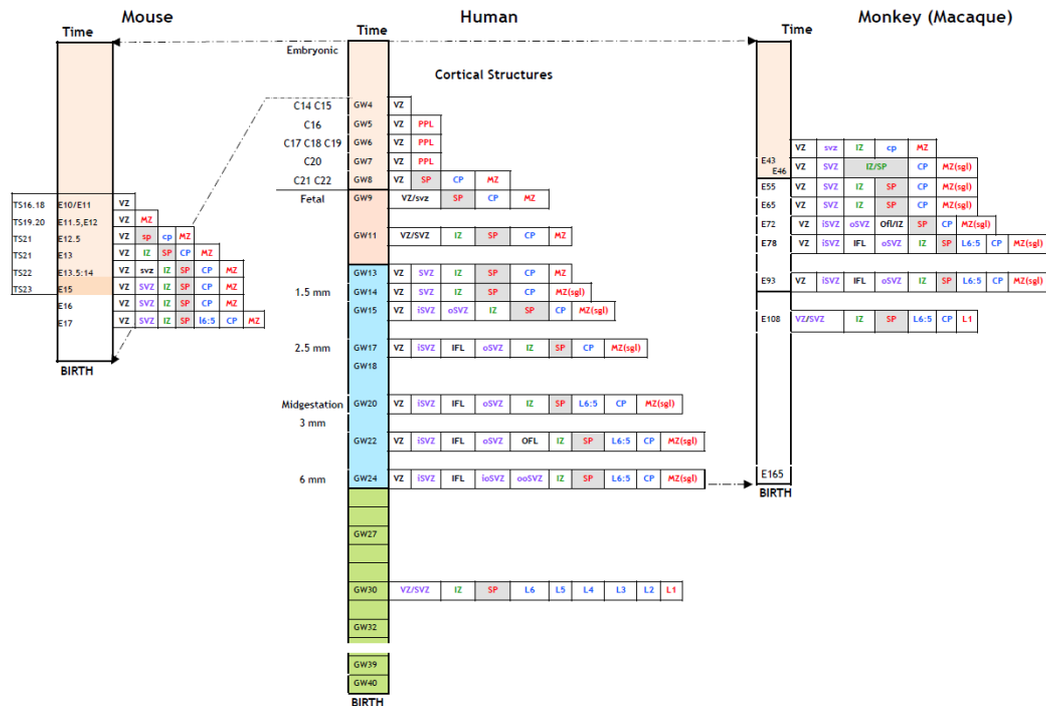
**Figure 1: Timetable of the development of embryonic cortical structures in human (middle): comparison with mouse (left) and monkey (right).**

The panel in the middle shows the development of embryonic cortical structures in the human brain during the first (orange) and second (blue) trimesters of the gestation. Time is counted in gestation weeks (GW, linear scale). During the last (green) trimester of the gestation, cortical layers develop at the expense of subjacent embryonic structures. The embryonic stages defined by Carnegie are indicated on the left side of the timetable. (After [18,20-22,60,61,75,108,114,138,143,169-171])

The panel on the right shows the sequence of development of embryonic cortical structures observed in the monkey brain until mid-gestation. The developmental sequence observed during the 100 first days of the gestation resembles the sequence observed during the first semester of the gestation in human. Time is counted in embryonic days (E, linear scale) (After [109,113])

The panel on the left recalls the sequence of development of embryonic cortical structures in the developing mouse brain. Because of the shortened duration of cortical development in the mouse, the time scale has been expanded. Embryonic structures in the mouse and human cortex derive after the embryonic (E) days 15/16 in mice and the gestation week (GW) 13 in humans

VZ, ventricular zone; SVZ, subventricular zone; iSVZ, inner subventricular zone; IFL, inner fiber layer; oSVZ, outer subventricular zone; OFL, outer fiber layer; ioSVZ, inner oSVZ; ooSVZ, outer oSVZ; IZ, intermediate zone; SP, subplate; CP, cortical plate; MZ, marginal zone; PP, pre plate; PPL, plexiform layer; LI,2,..., layer 1,2,...; sgl, subpial granular layer.



**Figure 2: Ontogenic and functional classification of the cortical interneurons (cIN) in the mouse cortex.**

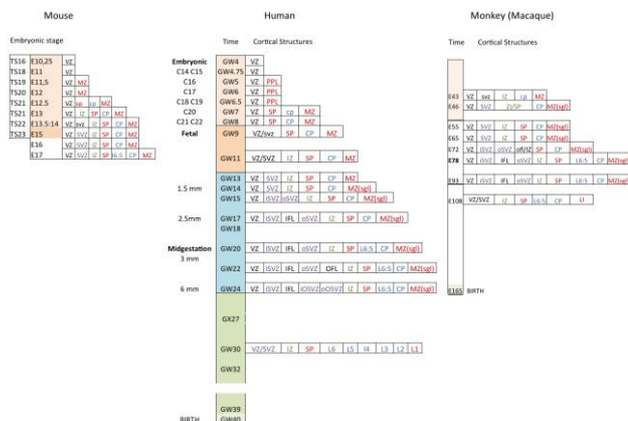
A. Distribution of cortical interneurons (cIN) within the 6 layers of the cortex according to their embryonic origin MGE (orange) or CGE (blue) (adapted from [168])

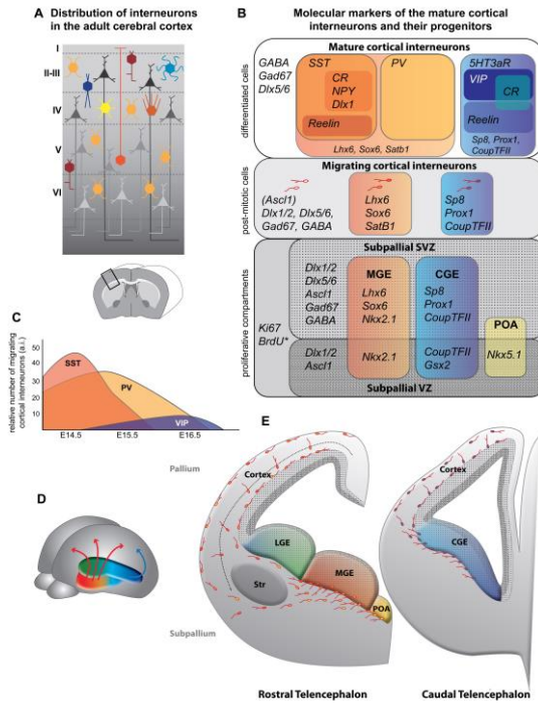
B. Up. Diversity of differentiated cIN. cIN express one of the main three non-overlapping markers: somatostatin (SST, dark orange), parvalbumin (PV, light orange), and the ionotropic serotonin receptor 5HT3a (5HT3aR, blue). Further subdivisions within each molecular group are revealed by the expression of other markers, as well as morphological, electrophysiological, and synaptic properties not indicated. The different subtypes have different embryonic origins indicated below.

Bottom. Mouse subtypes of cIN are generated from distinct proliferative regions within the subpallium. Boxes indicate the specific and common markers in the ventricular zone (VZ, dark grey) and subventricular zone (SVZ, medium grey) of the medial ganglionic eminence (MGE, orange) and the caudal ganglionic eminence (CGE, blue). Each region produces specific cIN. Markers expressed in the different subpopulations of migrating cIN are indicated in the upper boxes

C. Diagram showing the temporal origin of three subtypes of cIN. Early generated cIN originate from the MGE and give rise to SST+ (pink) or PV+ (orange) cIN. Late generated cIN derive from the CGE and produce VIP+ (blue) (adapted from [162])

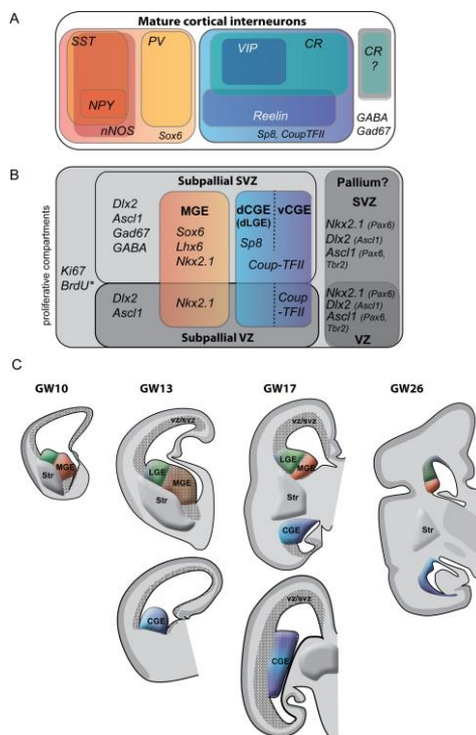
D. & E. Schemes showing the anatomical origin of cINs generated in the MGE and CGE, and their major routes of migration (D. adapted from [34]). E. Trajectories of cIN generated in the MGE (red), CGE (blue) and POA (yellow) on rostral (left) and caudal (right) coronal sections. VZ in dark grey, SVZ in light grey, LGE in green.





**Figure 3: Ontogenic and functional classification of the cortical interneurons (cIN) in the human cortex**

- A. Boxes illustrate the subpopulations of cortical interneurons cIN in the human cortex identified by their expression of functional and ontogenic markers (modified from [18], abbreviation as in Fig.2). A contingent of calretinin-expressing (CR+) cIN whose pallial origin is debated is indicated on the right.
- B. Boxes summarize the origin of cIN subtypes in human (after [18,23,75]). cIN are generated from distinct proliferative regions in the medial ganglionic eminence (MGE, orange) and the caudal ganglionic eminence (CGE, blue). The dorsal and ventral parts of the CGE are considered separately. Markers expressed by the putative pallial progenitors of cIN are listed in the grey box (after [22,60,61,76,108]). Ventricular zone, VZ, dark grey; subventricular zone, SVZ, medium grey.
- C. Schematic representation of frontal sections of the left telencephalic vesicle at gestation weeks (GW) 10,13,17 and 26 in human (after [172]), showing the location and relative size of the lateral ganglionic eminence (LGE, green), MGE (red) and CGE (blue). Rostral level is shown at each stage; caudal level (bottom) is shown only at GW13 and GW17, during the period of maximal CGE expansion. VZ and SVZ in patterned grey.



**Figure 4: Cortical localization at different gestational ages of cells labeled with interneuron markers (GABA, CR) or transcription factors expressed in subpallial and pallial domains (Ascl1, Dlx2, Nkx2.1, Sox6, Sp8, Coup-TFII)**

The figure summarizes results of 8 studies [18,21,22,60,61,114,138,143] that describe the distribution and orientation of GABAergic cIN at different developmental stages in the human cortex.

The distribution of CR+ cells summarizes results published in [21,22,114]. The cortical distribution at same stages of cells expressing transcription factors of the GABAergic lineage is also indicated on the figure (summarizes results published in [18,22,23,60,61,75,76,108,114,170]).

For each marker, the color intensity is proportional to the density of labeled cells. The expression of transcription factors in VZ and/or SVZ cortical progenitors, attested by the co-expression of transcription factors and either KI67 or Sox2, is figured by vertical stripes. Arrows indicate the tangential versus radial orientation of cells expressing GABA, CR, or neuronal markers (TUJ1). In transcription factor columns, arrows indicate the alignment of immunostained post-mitotic cells. The figure summarizes data in good agreement with each other, except the column “Nkx2.1” that illustrates results published in [60,76,108,170] in contradiction with results published in ([18,23,75] ), as indicated by the symbol “€”.

