Embryonic Development and Adult Regeneration of the Adrenal Gland

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The adrenal gland plays a pivotal role in an organism’s health span by controlling the endocrine system. Decades of research on the adrenal gland have provided multiscale insights into the development and maintenance of this essential organ. A particularly interesting finding is that founder stem/progenitor cells participate in adrenocortical development and enable the adult adrenal cortex to regenerate itself in response to hormonal stress and injury. Since major advances have been made in understanding the dynamics of the developmental process and the remarkable regenerative capacity of the adrenal gland, understanding the mechanisms underlying adrenal development, maintenance, and regeneration will be of interest to basic and clinical researchers. Here, we introduce the developmental processes of the adrenal gland and discuss current knowledge regarding stem/progenitor cells that regulate adrenal cortex remodeling and regeneration. This review will provide insights into the fascinating ongoing research on the development and regeneration of the adrenal cortex.

Keywords: Adrenal glands; Adrenal cortex; Developmental biology; Regeneration; Stem cells

INTRODUCTION

The adrenal gland is a bilateral endocrine organ consisting of the cortex and the medulla, which have different functions and developmental origins [1]. The adrenal cortex synthesizes steroid hormones from cholesterol through a series of biochemical metabolic pathways. The medulla produces epinephrine and norepinephrine as part of the sympathetic nervous system. The cortex develops from mesodermal tissue, whereas the medulla originates from the neuroectoderm [2-4]. The cortex is surrounded by mesenchymal cells that form the adrenal capsule. Underneath the capsule, there are three different cortical zones: the outermost zona glomerulosa (zG), the intermediate zona fasciculata (zF), and the innermost zona reticularis (zR) [1]. Although many animal models have provided extensive information on the development of the adrenal gland, the origin and regulation of the zonation of the human adrenal cortex remain to be elucidated.

Supported by pools of stem/progenitor cells, the adrenal gland has a high renewal capacity, which allows it to sustain physiological stress and meet hormonal demands [5-7]. In recent decades, exciting discoveries elucidating the identity, function, and fate of these stem/progenitor cell populations have emerged. Based on these findings, embryonic adrenal gland development and adult adrenal gland regeneration were found to share common features in terms of regulatory pathways and transcription factors. Understanding these shared features of adrenal development and regeneration will provide insights into the pathophysiological stress and meet hormonal demands [5-7]. In recent decades, exciting discoveries elucidating the identity, function, and fate of these stem/progenitor cell populations have emerged. Based on these findings, embryonic adrenal gland development and adult adrenal gland regeneration were found to share common features in terms of regulatory pathways and transcription factors. Understanding these shared features of adrenal development and regeneration will provide insights into the pathophysiological stress and meet hormonal demands [5-7]. In recent decades, exciting discoveries elucidating the identity, function, and fate of these stem/progenitor cell populations have emerged. Based on these findings, embryonic adrenal gland development and adult adrenal gland regeneration were found to share common features in terms of regulatory pathways and transcription factors. Understanding these shared features of adrenal development and regeneration will provide insights into the pathophysiological stress and meet hormonal demands [5-7]. In recent decades, exciting discoveries elucidating the identity, function, and fate of these stem/progenitor cell populations have emerged. Based on these findings, embryonic adrenal gland development and adult adrenal gland regeneration were found to share common features in terms of regulatory pathways and transcription factors. Understanding these shared features of adrenal development and regeneration will provide insights into the pathophysiological stress and meet hormonal demands [5-7]. In recent decades, exciting discoveries elucidating the identity, function, and fate of these stem/progenitor cell populations have emerged. Based on these findings, embryonic adrenal gland development and adult adrenal gland regeneration were found to share common features in terms of regulatory pathways and transcription factors. Understanding these shared features of adrenal development and regeneration will provide insights into the pathophysiological stress and meet hormonal demands [5-7]. In recent decades, exciting discoveries elucidating the identity, function, and fate of these stem/progenitor cell populations have emerged. Based on these findings, embryonic adrenal gland development and adult adrenal gland regeneration were found to share common features in terms of regulatory pathways and transcription factors. Understanding these shared features of adrenal development and regeneration will provide insights into the pathophysiological stress and meet hormonal demands [5-7]. In recent decades, exciting discoveries elucidating the identity, function, and fate of these stem/progenitor cell populations have emerged. Based on these findings, embryonic adrenal gland development and adult adrenal gland regeneration were found to share common features in terms of regulatory pathways and transcription factors. Understanding these shared features of adrenal development and regeneration will provide insights into the pathophysiological stress and meet hormonal demands [5-7].
DEVELOPMENT OF THE ADRENAL GLAND

The developmental program of the adrenal gland begins in the embryo and continues in the fetus and postnatal infant (Fig. 1). The adrenal cortex is established from the thickening of the mesoderm-derived coelomic epithelium between the urogenital ridge and the dorsal mesentery during 4 to 6 weeks post-conception in humans and at embryonic day 9 in mice [1,2,8]. Clusters of bipotential adrenal gonadal precursors, referred to as the adrenogonadal primordium (AGP), generate both the adrenal cortex and gonads. As the AGP grows along both sides, groups of cells separate from the coelomic epithelium and invade the mesenchymal layer in the intermediate mesoderm. The precursors next to the mesonephros migrate dorsolaterally to form the gonadal primordium, whereas the medial cells that abundantly express steroidogenic factor 1 (steroidogenic factor 1 [SF1], NR5A1, Ad4BP) migrate dorsomedially and converge to establish the adrenal primordium at the cranial pole of the mesonephros [9,10]. Thereafter, the cells of the adrenal primordium extensively proliferate and express 17α-hydroxylase while the adrenal gland undergoes vascularization [11,12].

During this stage, two dispersed distinct zones are detectable in the adrenal cortex. The fetal zone (FZ) appears in the inner

Fig. 1. The establishment of the adrenal gland. In the embryonic stage, following the growth of adrenogonadal primordium (AGP) on both sides, the adrenal progenitor population on the medial side of the AGP and the gonadal progenitor population on the lateral side of the AGP separate to form the adrenal primordium (AP) and the gonadal primordium (GP), respectively. From 6 weeks post-conception (wpc), the neural crest cells that later become the adrenal medulla invade the AP and the mesenchymal cells that become the capsule encapsulate them to establish the fetal adrenal gland. The enlarged fetal zone (FZ) is gradually replaced by the outer definitive zone (DZ). After birth, the FZ regresses through apoptosis and the adrenal cortex starts the zonation of the DZ into the zona glomerulosa (zG) and zona fasciculata (zF). Among the three major cortical zones, the zona reticularis (zR) is the last to develop. At around 6 to 8 years old, a period known as adrenarche, the zR is formed in the cortical-medullary boundary of the adrenal cortex. The production of adrenal androgens is clearly observed from this stage onwards. Unlike humans, the adrenal cortex in mice exhibits the X-zone as a transient cortical compartment at the cortical-medullary boundary. The regression of the X-zone is sexually dimorphic.
cortex with high levels of steroidogenic enzymes such as CYP17 and cholesterol side-chain cleavage enzyme (CYP11A), indicating that the fetal adrenal gland synthesizes dehydroepiandrosterone (DHEA) and its sulfated derivative (DHEAS). These steroids serve as sources of placenta-derived 17β-estradiol. Since the human placenta lacks P450c17 (17α-hydroxylase/17,20-lyase) and thus cannot synthesize estrogens de novo from cholesterol, estrogen formation from the DHEAS produced by FZ is essential for maintaining pregnancy [11,13]. Simultaneously, the definitive zone (DZ) in the outer cortex expresses relatively low levels of steroidogenic enzymes and emerges with FZ enlargement [13]. This rudimentary fetal adrenal gland is encapsulated by mesenchymal cells until 9 weeks of gestation and embryonic day 12.5 in humans and mice, respectively [3]. The translational zone (TZ) located in the intermediate region between the FZ and DZ is observed 14 weeks post-conception [11,14,15]. In the prenatal stage, the TZ produces cortisol via transient expression of 3β-hydroxysteroid dehydrogenase/Δ⁵-Δ⁴-isomerase (HSD3B2). The cortisol from the TZ is required for the development of various organ systems in utero. The negative regulation of the fetal hypothalamic-pituitary-adrenal (HPA) axis by this early cortisol synthesis plays a role in protecting normal female sexual development by inhibiting the production of adrenal androgens at 8 to 9 weeks post-conception [11].

The cortisol from the cortex also induces the expression of the adrenaline-synthesizing enzyme phenylethanolamine-N-methyltransferase in chromaffin cells of the medulla and transcriptionally regulates the expression of chromaffin cell-specific genes and neuron-specific genes [16,17]. The sympathoadrenal primordium, which becomes the adrenal medulla, is induced by the migration of neural crest cells to the dorsal aorta [18]. Neural crest-derived sympathetic cells differentiate into tyrosine hydroxylase-expressing catecholaminergic neuronal progenitor cells in response to bone morphogenetic protein signaling from the surrounding niche [19]. These adrenal medulla precursors move along the nerves and the blood vessels to invade the adrenal primordium and form the adrenal medulla in the innermost region of the adrenal gland. After birth, the medullary islands consist of chromaffin cells within the cortex that coalesce to form a contiguous medulla [20].

**ADRENOCORTICAL ZONATION**

Postnatal adrenocortical zonation begins with the differentiation of the fetal adrenal cortex into three major cortical zones (Fig. 1). The DZ differentiates into the zG, zF, and zR. The zG produces aldosterone, which regulates sodium retention and the intravascular volume via the renin-angiotensin system. The zF produces glucocorticoids associated with glucose metabolism and the immune response, while the zR secretes adrenal androgens. Although the developmental origin and the interplay between the adult cortical zones and the FZs remain to be fully elucidated, considerable efforts in lineage tracing experiments have suggested that the adult cortical zones originate from stem/progenitor cells in the adrenal capsule and the sub-capsular population [6,21,22]. In the neonatal stage, the enlarged FZ during gestation rapidly remodels and regresses by apoptosis in the inner region of the FZ, the result of which is the rapid reduction of adrenal androgen secretion after 3 months of birth [23,24]. Subsequently, the zR gradually forms and begins to synthesize androgens again from around age 6 to 8, a period known as adrenarche in humans [25]. The role of adrenal androgens has not been clearly identified, but interestingly, their secretion profiles decrease with age. Current evidence is limited, but their function may be related to aging of the organism [26,27].

The adrenocortical zones differ among species [4]. In mice, the adrenal cortex has no recognizable zR. The mouse adrenal cortex has the X zone, a remnant of the fetal adrenal cortex, in contact with the adrenal medulla. The X zone begins to develop after birth and grows to occupy about 30% of the cortex. This zone degenerates in male mice at puberty and either at first pregnancy in females or spontaneously in old nulliparous females [28]. The function of the X-zone is thought to be involved in progesterone catabolism [29]. In the rat adrenal cortex, an undifferentiated region (zU) between the zG and zF is observed [30]. This zone contributes to adrenal cortical homeostasis and can be divided into two different zones, referred to as the outer zU and the inner zU [30]. The inner zU is assumed to be a transitional zone committed to steroidogenesis. Unlike conventional laboratory rodents, the adrenal cortex of spiny mice (Acomys cahirinus) contains the zR and secretes both cortisol and DHEA, similar to the human adrenal cortex, making it a captivating new model for research on adrenal cortex development and disease [31].

**RENOVATION AND REMODELING OF ADRENOCORTICAL TISSUE**

In adult homeostasis, the adrenal cortex is responsible for the synthesis of steroid hormones through reciprocal hormonal regulation [1,4]. The adrenal cortex spontaneously replenishes dying cells that undergo rapid changes in response to hormonal
demands for steroid biosynthesis [6,7,22]. The adrenocortical stem cell population enables tissue renewal of the adrenal gland and helps to sustain physiological and homeostatic conditions despite constant cellular turnover throughout life (Fig. 2).

To identify bona fide stem cells and critical pathways involved in adrenocortical homeostasis and regeneration, specific lineage tracing and genetic ablation are currently under investigation. Different populations of adrenocortical progenitor cells have been uncovered in the adrenal capsular and sub-capsular region [3,5,7]. Similar to other organs, lineage progression in the adult adrenal cortex also proceeds from stem cells (no fate acquired, with self-renewal capacity) to progenitor cells (destiny acquired, with determinant gene activation), and then to steroidogenic cells (functional entities). This lineage progression in the adrenal cortex is controlled by the signaling response to both paracrine and endocrine signaling feedback loops, although the detailed mechanism remains to be clarified.

Two models of the adrenocortical renewal process have been proposed: the centripetal hypothesis for different adrenocortical zones, currently accepted [32,33] and the zone-specific hypothesis for lineage dependence, albeit old-fashioned [34]. In the centripetal migration model, undifferentiated progenitor cells in the capsular or sub-capsular region continually produce differentiated zG cells synthesizing mineralocorticoids. These cells migrate centripetally to the inner cortical zR/medulla boundary. In contrast, the zone-specific model claims that each different

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**Fig. 2.** Homeostasis, renewal, and regeneration of the adult adrenal cortex. To carry out the unique endocrine functions of the adrenal gland, three major compartments in the adrenal cortex, as well as the medulla, are controlled by external regulatory factors. Under the regulation of the renin-angiotensin system, the zona glomerulosa (zG) produces aldosterone to adjust the levels of sodium and potassium ions in plasma. In response to the circulating adrenocorticotropic hormone (ACTH) from the pituitary gland, the zona fasciculate (zF) and zona reticularis (zR) synthesize cortisol (corticosterone in mice) and adrenal androgens (dehydroepiandrosterone [DHEA] and dehydroepiandrosterone sulfate [DHEAS]), respectively. These steroids are governed by the hypothalamic-pituitary-adrenal (HPA) axis according to hormonal demands and external stress. The adrenocortical stem/progenitor cells in the capsule and sub-capsule regions of the adrenal cortex centripetally replenish senescent steroidogenic cells to maintain healthy steroidogenic cells in the cortical layer. The sonic hedgehog (SHH) and Wnt signaling pathway reciprocally regulate each type of signaling activity, and this mutual relationship is critical for maintaining proper functions of the adrenal cortex. ACTH-protein kinase A (PKA) signaling contributes to differentiating progenitors into steroidogenic cells in the zF. The adrenal cortical regeneration rates in males and females are similar to those of dihydrotestosterone (DHT)-treated females and gonadectomized (GDX) males, respectively, which explains the inhibitory effect of androgens on glioma-associated oncogene homolog 1+ (GLI1+) stem cell recruitment, and sexual dimorphism in the adrenal regeneration [36]. WT1, Wilms’ tumor 1; SF1, steroidogenic factor 1; GFP, green fluorescent protein; 20αHSD, 20α-hydroxysteroid dehydrogenase; CYB5, cytochrome B5; CC3, cleaved caspase 3.
zone is independently maintained by zone-specific progenitor cells, which can give rise to certain zonal cells. A series of prototype experiments have strengthened the centripetal migration hypothesis. The observation of complete adrenal cortex regeneration in an adrenal enucleation experiment, in which the inner content of the adrenal cortex was removed, leaving behind only the capsule and a layer of cells underneath the capsule, provided evidence that the capsular and/or subcapsular compartments in the adrenal cortex contain stem/progenitor cells for inner cortical regions [35]. Moreover, an intravital dye-labeling experiment performed using subcutaneous injections of trypan blue revealed that the initially dye-labeled capsule and zG cells were observed to move into the zF and eventually were detected in the zR. This finding suggests that newly synthesized cells are derived from the adrenal capsule [32].

In recent, genetic lineage-tracing experiments have provided more direct evidence for the centripetal migration hypothesis. Lineage tracing experiments to monitor the cell fates of sonic hedgehog (SHH)- and glioma-associated oncogene homolog 1 (GLI1)-expressing progenitor cells showed radial stripes that appeared to migrate through the zG into the zF [7]. CYP11B2 cell-specific lineage tracing demonstrated that differentiated adrenocortical cells carry out adrenocortical zonation through direct lineage conversion of zG cells into zF cells. The lineage conversion of zG cells into zF cells also contributes to adrenal cortex regeneration following by dexamethasone-induced damage [6]. Taken together, these two findings reveal that through centripetal migration, SHH+ cells, which are descendants of GLI1+ cells, are major contributors to the replenishment of adrenocortical cells in adult cortical cell renewal and regeneration.

REGULATORY FACTORS INVOLVED IN THE DEVELOPMENT AND RENEWAL OF THE ADRENA L CORTEX

The adrenal capsule is composed of SF1+ mesenchymal cells, which are necessary for maintaining sub-capsular SF1+ progenitor cells (Fig. 2) [36]. In response to hormonal requirements, SF1+ capsular stem cells generate sub-capsular SF1+ progenitors that differentiate into steroidogenic cells within the zG and zF. Several regulatory factors that are important for the centripetal replenishment of the adrenal cortex have been found. Since the adult renewal and regeneration of the adrenal cortex recapitulate the developmental lineage determination in the embryo, signaling molecules that participate in embryonic adrenal development are often key regulators of adult adrenal cortex homeostasis.

Hedgehog pathway
SHH, a hedgehog family ligand, is secreted by subcapsular cells that express SF1, but not steroidogenic enzymes such as CYP11B1 and CYP11B2 [7,37,38]. The related transcriptional effector, GLI1, is upregulated by the activation of the SHH pathway, and the GLI1 gene thus serves as a marker of hedgehog signaling activation [39]. Capsular cells that express GLI1 in response to SHH from subcapsular cells migrate centripetally into the cortex and differentiate into steroidogenic cells with a distinct zone-specific marker of the zG (CYP11B2) or zF (CYP11B1). GLI1+ progenitor cells are also responsible for differentiation into steroidogenic lineage cells during the cortical development in embryo, fetus, and newborn mice [7]. Moreover, it has been reported that the SHH pathway plays a critical role in adult adrenal regeneration. Following dexamethasone-induced atrophy, constitutively activating SHH signaling in capsular cells enhances adrenal cortex regeneration, whereas pharmacological inhibition of Shh signaling reduces the regeneration capacity. Mechanistically, the constitutive activation of SHH signaling is correlated with Wnt signaling activity, a pathway that is activated in the cortex during the regeneration process [40]. Taken together, these studies show not only the potential role of SHH-expressing cells as a stem/progenitor population for adrenal cortex homeostasis, but also the necessity of SHH signaling for adrenocortical regeneration.

WNT pathway
The canonical Wnt signaling pathway is responsible for both the embryonic establishment and homeostatic maintenance of the adrenal cortex. Mouse models of Wnt signaling ablation in steroidogenic cells have shown that Wnt signaling is critical for adrenocortical maintenance in both embryonic and adult mouse adrenal glands [21]. Among the Wnt signaling ligands, Wnt4 is produced by cells located in the subcapsular/zG region in mice, and it appears to be involved in adrenocortical homeostasis via the canonical Wnt signaling pathway. Homozygous loss of Wnt4 in mice results in postnatal adrenal defects, suggesting that Wnt4 plays an important role in neonatal zonation and steroidogenesis [41]. In humans, a homozygous mutation in WNT4 was found to result in severe developmental defects, including adrenal hypoplasia [42].

Interesting evidence regarding the regulation of Wnt signaling in both the establishment of adrenocortical zonation and in the maintenance of the capsular stem/progenitor cell niche has recently been reported. SF1+ capsular stem cells secrete R-spondin 3 (Rspo3) to induce Wnt/β-catenin signaling activation in
the underlying steroidogenic compartment, and impaired expression of Rspo3 in capsular progenitors reduces Shh and Wnt4 expression in zG cells [43]. These results indicate that reciprocal inter-compartmental signaling maintains the zG characteristics and adrenal homeostasis through a cooperative mechanism.

**ACTH/PKA signaling**

The production of cortisol and androgen in humans (corticosterone in mice) by adrenocortical steroidogenic cells is governed by the HPA axis. Adrenocorticotropic hormone (ACTH) from the pituitary gland stimulates cortisol production from the adrenal zF by activating cAMP-dependent protein kinase A (PKA) signaling [44,45]. The complex of melanocortin 2 receptor (MC2R) and its accessory protein (MRAP) on the surface of adrenocortical cells plays an essential role in mediating the action of pituitary ACTH and subsequent steroidogenesis [46,47]. MRAP deficiency was found to lead to neonatal lethality in mice, and the administration of exogenous glucocorticoids rescued the lethality. However, the MRAP-deficient mice rescued by glucocorticoid treatment showed impaired steroidogenesis due to an accumulation of progenitor cells from activated SHH and Wnt signaling [48]. These results explain why ACTH signaling via PKA induces the differentiation of adrenocortical stem/progenitor cells into steroidogenic cells of the zF.

**Regulation of SF1 transcriptional activity**

SF1 is a major transcription factor involved in adrenal development and function. Forced expression of SF1 in adipose-derived and bone marrow-derived mesenchymal cells [49-52], as well as in mouse embryonic stem cells [51,53], induces their differentiation into steroidogenic cells. Moreover, transgenic mice that expressed SF1 under the control of fetal adrenal enhancer showed increased adrenal size and the formation of ectopic adrenal tissue in the thorax [54]. These results indicate that the forced expression of SF1 causes the switch of nonadrenal lineage cells to the adrenal cell fate. The activation of SF1 expression and modulation of SF1-dependent transcription have been found to play an important role in these phenomena. Both odd-skipped related 1 [55] and Sal-like 1 (SALL1) [56] deletion cause reduced expression of Wilms’ tumor protein 1 (WT1). WT1 binds to the promoter region within the SF1 gene and cooperates with the coactivator Cbp/p300-interacting transactivator 2. This complex strongly induces SF1 expression, which is required for adrenal primordium development [57]. Furthermore, to adjust to the differentiation to steroidogenic cells, SF1 self-inhibits its transcriptional activity by activating DAX1, the repressor of SF1 [58]. These findings indicate that fine control of SF1 transcriptional activity is required for proper adrenal development.

**CONCLUSIONS**

The signaling pathways that regulate the formation of the adrenal cortex in the embryo are shared in the regeneration process that occurs in response to hormonal demand and surgical damage. Several animal experiments have shown that the regulatory genes involved in adrenal cortical stem/progenitor cell identity, fate, and differentiation during development are also necessary during the adult adrenal regeneration process. However, the regulatory network in the human adrenal gland remains to be validated. Thus, the unraveling of the undefined gene regulatory network between human adrenal cortical development and regeneration will be a topic of major interest.

Despite the remaining challenges, human organoids and single-cell RNA sequencing techniques based on epigenetic modulation have provided great potential in translational studies [59,60]. Technological advances in both approaches may elucidate aspects of human biology relevant for research on the adrenal glands and pathological processes therein. As the establishment of many organoid systems has gained insights from the understanding of organogenesis, current and future knowledge on adrenal development and regeneration will lay the foundation for the development of human adrenal organoid systems.

**CONFLICTS OF INTEREST**

No potential conflict of interest relevant to this article was reported.

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