Hypothalamic Glial-to-Neuronal Signaling during Puberty: Influence of Alcohol

Vinod K. Srivastava, Jill K. Hiney and W. Les Dees *

Department of Veterinary Integrative Biosciences, College of Veterinary Medicine, Texas A&M University, College Station, TX 77843-4458, USA; E-Mails: vsrivastava@cvm.tamu.edu (V.K.S); jhiney@cvm.tamu.edu (J.K.H.)

* Author to whom correspondence should be addressed; E-Mail: ldees@cvm.tamu.edu;
Tel.: +1-979-8451430; Fax: +1-979-847-9038.

Received: 26 May 2011; in revised form: 27 June 2011 / Accepted: 12 July 2011 / Published: 14 July 2011

Abstract: Mammalian puberty requires complex interactions between glial and neuronal regulatory systems within the hypothalamus that results in the timely increase in the secretion of luteinizing hormone releasing hormone (LHRH). Assessing the molecules required for the development of coordinated communication networks between glia and LHRH neuron terminals in the basal hypothalamus, as well as identifying substances capable of affecting cell-cell communication are important. One such pathway involves growth factors of the epidermal growth factor (EGF) family that bind to specific erbB receptors. Activation of this receptor results in the release of prostaglandin-E2 (PGE2) from adjacent glial cells, which then acts on the nearby LHRH nerve terminals to elicit release of the peptide. Another pathway involves novel genes which synthesize adhesion/signaling proteins responsible for the structural integrity of bi-directional glial-neuronal communication. In this review, we will discuss the influence of these glial-neuronal communication pathways on the prepubertal LHRH secretory system, and furthermore, discuss the actions and interactions of alcohol on these two signaling processes.

Keywords: alcohol; puberty; transforming growth factor-α; glia; RPTPβ
1. Introduction

The hypothalamic region of the brain plays an important role in synchronizing events leading to the activation of the mammalian puberty. This process requires the interactive participation of both glial and neuronal regulatory circuitries that serve to control the secretion of luteinizing hormone-releasing hormone (LHRH) neurons [1,2]. The secretory activity of LHRH neurons is triggered by several transsynaptic inputs of both inhibitory and excitatory nature [1,3,4]. The decreased release of inhibitory neurotransmitters such as gamma amino butyric acid and the opioid peptides [5,6] as well as the increased release of excitatory neurotransmitters such as excitatory amino acids [7,8], transforming growth factor alpha (TGFα) [9], insulin-like growth factor-1 [10,11], and the kisspeptins [12] are capable of initiating the cascade of events leading to increased LHRH secretion at puberty. The glial cells within the medial basal hypothalamus (MBH) are key components of the central regulatory system that facilitate prepubertal LHRH release via pathways initiated by growth factors and cell adhesion molecules that act on LHRH neuron terminals to stimulate their secretory activity [13,14].

Growth factors of glial origin are important because they are intimately involved in glial-neuronal signaling processes by which the glial cells, through their close association with LHRH nerve terminals in the MBH, regulate LHRH secretion during puberty in rodents [13,15] and primates [16]. Growth factors such as basic fibroblast growth factor (bFGF) [17,18], transforming growth factor β (TGF-β) [19] and IGF-1 [10,20] have been shown to act directly on LHRH neurons to facilitate LHRH release. In contrast to these growth factors, members of the epidermal growth factor (EGF) family, including EGF itself, TGFα, and the neuregulins act indirectly on LHRH release through specific erbB receptors that have an extracellular ligand binding domain linked to a cytoplasmic domain containing tyrosine kinase activity [14,21,22]. While all of these peptides can influence LHRH release, a portion of this review will concentrate specifically on EGF/TGFα influences, since they have been shown to play a pivotal role in the glial control of neuronal LHRH secretion at the time of puberty.

During the past decade, evidence has accumulated suggesting LHRH secretory activity is also modulated by a specific glial-neuronal gene family which synthesizes adhesion/signaling proteins involved in the functional and structural integrity of bi-directional glial-neuronal communications. In this regard, various glial-neuronal adhesion genes have been identified within the hypothalamus which are not only involved in adhesive interactions, but also mediate intracellular signaling cascades that are critical for pubertal development [23,24]. The genes of this family include glial receptor protein tyrosine phosphatase-β (RPTPβ), neuronal contactin and contactin associated protein 1 (Caspr1). Once bound together, this family collectively contributes to glial-neuronal adhesiveness and to cell to cell communications [25–27]. The interaction between neuronal circuits and glial cells can be further influenced by metabolic signals, various environmental insults, and drugs of abuse. With regard to the latter, alcohol (ALC) is a drug of abuse that is known to alter hypothalamic functions that control reproduction. Chronic ALC exposure has been shown to cause depressed prepubertal LHRH secretion, and delayed pubertal development in both rodents [28–33] and primates [34,35]. This action of ALC is important since the onset of mammalian puberty is dependent upon an increase in the release of LHRH from the basal hypothalamus.
Recently, it has been suggested that some of the hypothalamic effects of ALC are due to ALC-induced interferences in glial-neuronal signaling networks involved in LHRH secretion. This review will mainly discuss two emerging glial-neuronal communication networks regarding their respective relationships to prepubertal LHRH secretion, and then address the actions and interactions of ALC on these glial-neuronal components during pubertal development.

2. EGF/TGFα Family of Growth Factors and Puberty

Growth factors of glial cell origin, acting via receptor tyrosine kinases, have been shown to be key components of the mechanism by which hypothalamic glial cells regulate LHRH neuronal function [36–38]. EGF and TGFα are peptides that signal through the erbB1 receptor. While both can stimulate LHRH release [9], TGFα has been shown to play a more pivotal role in the regulation of LHRH neuronal function during puberty in rodents [13,15] and primates [16]. TGFα mRNA and protein are highly expressed in glial cells and tanycytes in the MBH, their expressions increase significantly around the time of puberty [39], and pharmacological blockade of the erbB1 receptor [40] targeted to the MBH delays puberty [39]. Additionally, sexual maturation induced by hypothalamic lesions is associated with activation of TGFα expression in glial cells [41,42], and the effect of this lesion on puberty is blocked by using a selective inhibitor (RG-50864) of TGFα/EGF receptor tyrosine kinase activity [41]. Also, advanced puberty has been observed in transgenic mice overexpressing the TGFα gene [43] and in rats carrying grafted cells genetically engineered to secrete TGFα [44]. TGFα binds to and activates the erbB1/erbB2 receptor complex on adjacent glial cells in MBH. Activation of these receptors results in the production and release of prostaglandin-E2 (PGE2) from these glial cells, which then induces prepubertal LHRH secretion upon binding to specific receptors on nearby LHRH neuron terminals in the median eminence (ME) region of the MBH [22,37]. This is supported by the fact that the stimulatory effect of TGFα on PGE2 release was blocked by administering an erbB1 receptor antagonist (RG-50864) [37]. Several studies have shown that TGFα stimulates LHRH release via an indirect mechanism that involves a paracrine effect of this growth factor on glial cells in the release of LHRH. In this regard, the erbB1 receptors for TGFα have been shown immunohistochemically only in glial cells [15,16]. Furthermore, Ma et al. [37] have shown in vitro that the secretion of PGE2 from hypothalamic glial cells is increased after exposure to TGFα and that the conditioned medium of hypothalamic glial cells treated with TGFα is able to stimulate LHRH release from GT1 cells, which are immortalized LHRH secreting neurons. Moreover, in hypothalamic glial cells, PGE2 formation induced by TGFα and the stimulatory effect of the TGFα treated conditioned medium on LHRH release are shown to be prevented by the inhibition of erbB receptor tyrosine kinase activity or prostaglandin synthesis [37,45]. Collectively, these data strongly support the notion that TGFα acts indirectly in the functional control of neuronal networks governing mammalian puberty via hypothalamic glial-neuronal communications.
3. Effects of ALC on the TGF\textsubscript{α}/erbB1 Receptor/PGE\textsubscript{2} Pathway

It has been established that ALC acts within the hypothalamus to suppress the release of LHRH in both prepubertal and adult rats [46,47] and primates [35], and also causes delayed signs of pubertal maturation in both species [28,34]. Studies to discern the mechanism of this action of ALC to suppress LHRH release are important for understanding how this drug disrupts pubertal development. An important component of this ALC effect is PGE\textsubscript{2}, which plays a major role in the LHRH secretory process in prepubertal animals [48,49], and is known as a critical factor for glial-dependent regulation of LHRH release [21,37]. We showed previously [50] that acute ALC alters the EGF/TGF\textsubscript{α}-erbB1 receptor-COX (cyclooxygenase)-PGE\textsubscript{2} pathway by inhibiting the induction of COX, the rate limiting enzyme necessary for prostaglandin synthesis and lowers prepubertal PGE\textsubscript{2} secretion resulting in suppressed LHRH release [50,51]. Only recently have the mechanisms by which short-term ALC exposure affects the TGF\textsubscript{α}-erbB1 receptor-PGE\textsubscript{2} pathway been assessed with regard to glial-neuronal communications within the prepubertal hypothalamus [52]. That study has revealed that short-term ALC exposure for 4 and 6 days caused an increase in TGF\textsubscript{α} gene and protein expressions in prepubertal female rats. The gene expression of TGF\textsubscript{α} was increased markedly at 4 days (Figure 1). After 6 days of ALC exposure, the level of TGF\textsubscript{α} gene expression was still modestly but significantly elevated; however, the levels had declined markedly (not shown) as compared to 4 days of exposure. This effect paralleled an increase in TGF\textsubscript{α} protein expression at both 4 days (Figure 2A) and 6 days (Figure 2B). To determine if the elevated hypothalamic levels of TGF\textsubscript{α} protein were due to an inhibition of release, we assessed basal TGF\textsubscript{α} secretion from rat MBHs incubated in vitro following 6 days of ALC exposure in vivo. We determined that animals exposed to ALC had suppressed release of TGF\textsubscript{α} (Figure 3). Overall, these findings suggest that ALC exposure does not affect transcription or translation of TGF\textsubscript{α}, but is capable of interfering with the hypothalamic release of glial TGF\textsubscript{α}, resulting in suppressed prepubertal PGE\textsubscript{2} and LHRH secretion.

**Figure 1.** Effect of short-term ALC (ethanol) exposure on basal TGF\textsubscript{α} gene expression in the MBH of perpubertal female rats. Note that the ALC-treated animals showed an increase in the gene expression of TGF\textsubscript{α} compared with control animals. Values represent mean ± SEM. N = 12–14 animals per group; *** p < 0.001 versus control.
Figure 2. Effect of short-term ALC (ethanol) exposure on TGFα protein expression in the MBH of prepubertal female rats. Composite graphs that show the densitometric quantitation of the bands corresponding to TGFα protein. These data were normalized to the internal control β-actin protein, and the densitometric units represent the TGFα/β-actin ratio. Note that ALC caused an increase in TGFα protein expression on both 4 and 6 days compared with control animals. Values represent mean ± SEM. N = 7–8 per group. ** p < 0.01; *** p < 0.001 versus control.

Figure 3. Effect of short-term in vivo ALC (ethanol) exposure for on TGFα protein released in vitro from the MBH of prepubertal female rats. Note that TGFα release was decreased in ALC-treated animals compared with control animals. These data were normalized to the internal control β-actin protein, and the densitometric units represent TGFα/β-actin ratio. Values represent mean ± SEM. N = 14 for control, N = 9 for ALC, ** p < 0.01 versus control.

This study also showed that the erbB1 receptor, the principal receptor for TGFα was affected by ALC. Short-term ALC exposure for 4 and 6 days caused a marked decrease in the synthesis of the phosphorylated form of the erbB1 receptor at 4 days (Figure 4), with 6 days being almost identical (not shown), but did not elicit changes in erbB1 gene expression or the synthesis of total, non-phosphorylated erbB1 protein. It is possible that down regulation of erbB1 gene synthesis had not yet occurred because of this short-term duration of ALC exposure, but it does appear ALC affected the phosphorylation of the erbB1 protein. Interestingly, in this study ALC did not affect the synthesis of total and phosphorylated erbB2, the co-receptor necessary for activation of erbB1 signaling [38] (not
shown), which further demonstrates a specific effect of ALC on the erbB1 receptor. The mechanism of this action of ALC on erbB1, however, remains unclear. Since TGFα binding initiates the autophosphorylation of the erbB1 receptor [53,54], it is likely that the ALC-induced impairment of TGFα release (Figure 3) is a major contributing factor responsible for decreased erbB1 phosphorylation observed in the ALC-treated animals. However, we cannot rule out the possibility of a direct effect of ALC on the erbB1 autophosphorylation process that is independent of its effect to suppress TGFα secretion. In this regard, studies have demonstrated that chronic ALC exposure can disrupt phosphorylation of erbB1 by altering the receptor affinity and/or tyrosine kinase activity in rats [55,56].

**Figure 4.** Effect of short-term ALC (ethanol) exposure on phosphorylated erbB1 protein expressions in the MBH of prepubertal female rats. Note that the ALC-treated animals showed a marked decrease in phosphorylated erbB1 protein expression compared with controls. These data were normalized to the total, nonphosphorylated erbB1 protein, and the densitometric units represent the phosphorylated erbB1/total, non-phosphorylated erbB1 ratio. Values represent mean ± SEM. N = 8 per group. **p < 0.01 versus control.

Since it was expected that suppressed erbB1 phosphorylation would result in a downstream decrease in PGE2 release, animals were chronically exposed to ALC for 6 days and then their hypothalami removed and incubated in vitro in order to assess the amount of PGE2 released into the incubation medium. This revealed that ALC exposure suppressed the release of PGE2 (Figure 5), an action that was associated with the suppressed phosphorylation of the erbB1 receptor shown in Figure 4. These findings demonstrated for the first time the upstream inhibitory effects of ALC on glial TGFα/erbB1 pathways that control the production and secretion of PGE2 within the MBH; hence, providing a mechanism supporting previous reports showing that ALC is capable of suppressing PGE2 and subsequently, LHRH secretion [50,51,57,58].
Figure 5. Effect of *in vivo* ALC (ethanol) exposure on prostaglandin-E2 (PGE2) release *in vitro* from the MBH of prepubertal rats as determined by enzyme-linked immunoassay. Note that the ALC-treated animals showed a marked decrease in basal PGE2 release compared with control animals. Values represent mean ± SEM. \( N = 8 \) for control, \( N = 9 \) for ALC. ** \( p < 0.01 \) versus control.

Studies in recent years have identified upstream regulatory sites in the TGFα/erbB1/PGE2/LHRH-secretory pathway. Investigators have shown that the expression of the POU homeodomain gene, Oct2, increases in the MBH at puberty and that this increase was associated with the transactivation of TGFα gene expression [59]. Because IGF-1 plays an important role in control of prepubertal LHRH release, we investigated the possibility that IGF-1 may influence transcription of the Oct 2 gene and that this gene may be a target of ALC actions [60]. It was demonstrated that a single injection of IGF-1 to 25 day-old female rats caused increases in both Oct 2a and c gene transcripts in the MBH and furthermore, showed that an acute dose of ALC (3g/kg) did not alter basal expression of the gene transcripts, but blocked the IGF-1 induced expressions (Figure 6a and b). Similar effects were observed regarding Oct 2c in the POA, but not the Oct 2a transcript. Other investigators showed an inhibitory effect of chronic ALC exposure on Oct 2 proteins expressed in the entire rostro-caudal extent of the prepubertal hypothalamus [61].

Interestingly, ALC exposure has also been shown to affect the gene encoding thyroid transcription factor 1 (TTF1). TTF-1 is a member of the Nkx homeodomain gene family that increases in the hypothalamus at puberty and has the ability to activate LHRH neurons [38,62]. It was shown that TTF1 expression peaked at postnatal day 26–27 in controls but that ALC administration beginning at 24 days of age caused suppressed hypothalamic TTF1 protein expression between 25 and 27 days, which was followed by a significant increase by 30 days [61]; thus, those authors suggested that the ALC delayed the peak increase in prepubertal TTF1 protein expression. Recently, we observed that ALC exposure beginning on 27 days of age was associated with an increase in TTF1 gene expression in the MBH (Figure 7) at 31 days. This increase at 31 days did not occur in the POA, however, by day 33, both hypothalamic regions showed elevated TTF1 gene expressions compared with controls (not shown). While additional research is needed in this area, this observation supports the earlier report by Kim *et al.* [61], that ALC postponed the prepubertal increase in TTF1 synthesis.
Figure 6. Effect of acute ALC (ethanol) exposure on IGF-1 induced Oct 2a and 2c in the MBH of prepubertal female rats. Panels A and B depict densitometric quantitation corresponding to the Oct 2a and 2c transcripts. IGF-1 induced an increase in Oct2a and 2c mRNA over basal synthesis. Acute ALC exposure did not affect basal synthesis of Oct 2a and 2c mRNA but blocked the IGF-1 induced synthesis of both Oct 2a and 2c. N = 5–6/group. * p < 0.05.

Figure 7. Effect of short-term ALC (ethanol) exposure on TTF1 mRNA expression from the MBH of prepubertal female rats. Note that TTF1 was increased in ALC-treated animals compared with control animals. Values represent mean ± SEM. N = 9 for control, N = 9 for ALC, * p < 0.05 versus control.

4. Adhesion/Signaling Genes Involved in Glial-Neuronal Communication during Puberty

There is growing evidence that glial cells can also regulate LHRH secretion by contributing to plastic arrangements associated with glial-neuronal adhesiveness [13]. Since adhesion/signaling molecules are abundant in the MBH, and have cell signaling actions [23,63], it has been suggested that they play a role in intracellular communication between glial cells and LHRH neuron terminals [64,65]. In this regard, several genes have been identified which synthesize adhesion/signaling proteins responsible for the structural integrity of bi-directional glial-neuronal communication. The synaptic cell adhesion molecule (SynCAM1) expressed by both glia and neuronal cells, promotes the glial-neuronal adhesiveness via homophylic, extracellular domain-mediated interactions required for synaptic assembly [13]. Similarly, another report using an in vitro adhesion assay shows that both hypothalamic glial cells and GT1-7 neuronal cells adhere to the extracellular domain of SynCAM1 and suggest the importance of this adhesion molecule in cell-cell communication between glial cells and LHRH neurons [64,65]. SynCAM1 is also associated with neuregulin activated erbB4 receptor, one of
the glial cell erbB receptors involved in LHRH secretion at puberty [64,66] and is expressed in hypothalamic glial cells. Specifically, erbB4 receptor forms a heterodimer complex with erbB2 and activation by neuregulins causes LHRH release via an action involving glial PGE2 release and facilitation of erbB-1 mediated signaling events [66,67]. Disruption of erbB4/-2 signaling results in delayed puberty and an inhibition of SynCam1 expression in hypothalamic glial cells [66,67]. Furthermore, a study has shown that transgenic mice with a double negative form of SynCam1 lacking the intracellular domain causes these animals to have delayed puberty [68]. These observations suggest that one of the mechanisms underlying erbB4 receptor facilitation of LHRH neuronal function involves SynCAM1 dependent signaling during pubertal development. The effect of ALC exposure on SynCAM1 gene and protein expression is under investigation.

Another example among the adhesion/signaling genes within the MBH is a three member family consisting of neuronal contactin associated protein-1 (Caspr1), a transmembrane protein that binds to contactin on the same neuronal cell membrane. The contactin portion of this Caspr1/contactin complex is bound by a glial transmembrane protein, receptor protein tyrosine phosphatase beta (RPTPβ); thus, forming the three member assembly that can contribute to glial-neuronal adhesiveness [25,26]. Contactin participates in axonal growth, synaptogenesis and neuroendocrine function, and contactin expression in hypothalamic secretory neurons has been shown to be altered in response to changes in glial-neuronal associations [27]. Caspr1 is proposed as a signaling molecule of contactin, which mediates its cell adhesion by binding to RPTPβ and activating intracellular signaling pathways in neurons [63]; thus, following binding, this three member assembly contributes to bidirectional cell-cell communications between glial and neuronal cells associated with hypothalamic neuroendocrine functions [27]. It is important to note that LHRH neurons express both contactin and Caspr1 [23]. It has been suggested that upon binding together this system not only provides adhesiveness between glial connections with LHRH terminals, but also regulates intracellular processes [64]. Interestingly, the intimate association between glial “end feet” and LHRH neuron terminals with the ME area is modified by the actions of different reproductive factors [69–71], in that changes in the contact between glia and neurons fluctuates depending on steroid milieu and stage of pubertal development [72]. Taken together, a concept is emerging that suggests that these glial-neuronal molecules facilitate hypothalamic neurosecretion via changes in the arrangement of glial-neuronal adhesion and signaling that could alter LHRH release and the pubertal process.

5. Effects of ALC on Glial-Neuronal Adhesion and Signaling

In addition to ALC affecting neuronal inputs involved in prepubertal LHRH secretion, there has been some recent attention given to the actions of this drug on the contactin-Caspr1-RPTPβ signaling system as it relates to glial-LHRH neuronal interactions [73]. In this regard, short-term ALC exposure caused a marked decrease in the basal expression of the RPTPβ gene (Figure 8A), but did not affect the expression of either contactin (Figure 8B) or Caspr1 (Figure 8C). Similarly, ALC caused suppressed levels of the RPTPβ protein (Figure 9A), with the expressions of both contactin (Figure 9B) and Caspr1 (Figure 9C) proteins being unaltered. The ALC-induced decrease in glial RPTPβ gene expression in this study indicates a reduced amount of peptide available for binding to the contactin-Caspr1 complex on LHRH neurons. As a result, decreased adhesiveness between the glia
and LHRH neurons in the MBH following ALC exposure would be expected, an action that could affect neuronal functions associated with pubertal development. The binding of glial RPTPβ to the contactin/Caspr1 complex on neuron terminals initiates the cell adhesion and subsequently, activates neuronal intracellular signaling pathways [63]. The precise contribution of contactin-Caspr1 signaling to LHRH secretion remains to be determined, but previous studies have suggested that a function of the RPTPβ-contactin-Caspr1 family is to facilitate neurosecretion through changes in glial-neuronal signaling [23,27]. The cell adhesiveness component also promotes a more secure proximity for glial-derived secretions to bind to their receptors on the LHRH nerve terminals. Whatever the mechanism, the fact that ALC alters the synthesis of prepubertal glial RPTPβ in the hypothalamus, which is required for binding to the neuronal contactin/Caspr1 complex, suggests diminished glial-neuronal adhesiveness and thus, altered facilitation of LHRH release by the products secreted from the neighboring glial cells.

**Figure 8.** Effect of short-term ALC (ethanol) exposure on the gene expressions of basal RPTPβ (A), Contactin (B) and Caspr1(C) in the MBH of prepubertal female rats. Note that ALC caused a marked decrease in the gene expression of basal RPTPβ compared with control animals. Values represent mean ± SEM. N = 12–13 per group. *** p < 0.001 versus control.

**Figure 9.** Effect of short-term ALC (ethanol) exposure on the protein expressions of basal RPTPβ (A), contactin (B) and Caspr1(C) in the MBH of prepubertal rats. Note that the protein expression of RPTPβ was markedly decreased in ALC exposed animal; however, the protein expressions of contactin and Caspr1 were unaltered. These data were normalized to the internal control β-actin protein, and the densitometric units represent the respective specific protein/β -actin ratio. Values represent mean ± SEM. N = 10 for control, N = 6 for ALC. *** p < 0.001 versus control.
6. Actions and Interactions of IGF-1 and ALC on Glial-Neuronal Adhesion and Signaling:

IGF-1 plays a critical role in the pubertal process. Initially, it was shown that the peptide is capable of inducing release of LHRH from the prepubertal hypothalamus in vitro [10], then later demonstrated that IGF-1 acts within the hypothalamus to accelerate the timing of female puberty [11]. Subsequently, it was shown that premature elevation of serum IGF-1 advanced first ovulation in rhesus monkeys [74]. While IGF-1 is produced by neurons and glia in the MBH, the majority of IGF-1 present in this region during puberty is derived from peripheral sources, such as liver [11,75,76]. IGF-1 binds to type 1 IGF receptors (IGF-1R) in different tissues including the brain, where the greatest concentration of IGF-1R has been observed in the ME of both rats [77] and primates [78]. Recently, DiVall et al. [79] showed that transgenic mice lacking the IGF-1R on their LHRH neurons have delayed puberty, further supporting the importance of this peptide for pubertal development. IGF-1 is also capable of acting at both glial and neuronal levels involved in neuroendocrine events within the hypothalamus in association with prepubertal LHRH secretion [47]. These findings suggest that IGF-1 may play a role in regulating the expression of adhesion and signaling molecules. Interestingly, we showed that IGF-1 induced an increase in the expression of the RPTPß gene in prepubertal female rat hypothalamus (Figure 10); however, it did not affect the gene expression of either contactin or Caspr1 [73]. Glial RPTPß is expressed abundantly within the MBH [80]. The knowledge that increased circulating levels of IGF-1 at puberty can cross the blood brain barrier and enter the MBH region [11], and that the circulating peptide is taken up by hypothalamic glia in the MBH [81], suggests that the IGF-1-induced expression of RPTPß in the MBH may facilitate neurosecretion via changes in glial-neuronal adhesive signaling.

The fact that IGF-1 is important for LHRH release at puberty [10,11] and the observation that ALC can alter prepubertal IGF-1 signaling [29] suggests that ALC could have a detrimental impact on pubertal development by interfering with glial neuronal signaling networks. In support of this, it has been shown recently [73] that ALC blocked the ability of IGF-1 to induce prepubertal RPTPß gene expression (Figure 10). The mechanism by which ALC influenced the IGF-1 induced expression of RPTPß remains to be determined; however, this effect could be due to an action of ALC at the level of the IGF-1R and/or to an altered post-receptor event. Interestingly, ALC administration does not alter IGF-1R gene or protein expression [29], although we cannot rule out that it may affect mechanisms regulating receptor function. This suggests that ALC may act on a pathway component downstream from the IGF-1R. In this regard, chronic ALC administration was shown to suppress the basal protein expression of phosphorylated Akt, a transduction signal activated by IGF-1 [82]. Acutely, ALC was shown to block the IGF-1 induced expression of other genes involved in the pubertal process [60,83], and that this occurs by blocking the peptides ability to induce phosphorylation of Akt [83]. Chronic exposure to ALC has been shown to decrease circulating levels of IGF-1 in prepubertal rats [29] and rhesus monkeys [34]; actions associated with altered pubertal development. Based on these collective results, it is suggested that the decreased prepubertal levels of circulating IGF-1 available to the MBH, as well as altered post-receptor transduction signals, may contribute to the ability of ALC to cause the suppression in RPTPß gene expression.
Figure 10. Effect of acute ALC exposure on basal and IGF-1 stimulated gene expressions of RPTPβ in the MBH of prepubertal female rats. Note that IGF-1 increased the gene expression of RPTPβ compared with the basal level of control. Importantly, the basal expression of the RPTPβ gene was not altered by ALC alone, but the IGF-1 induced expression of RPTPβ was blocked by the presence of ALC. Values represent mean ± SEM. N = 8–10 per group. ** p < 0.01 versus control and ALC + IGF-1.

7. Conclusions

It is becoming increasingly clear that glial-neuronal interactions play an important role in prepubertal LHRH secretion and the subsequent acquisition of female pubertal development. In this review, we have mainly discussed the actions of glial-neuronal communication networks at puberty and how these actions are influenced by ALC, a drug of abuse known to alter pubertal development. A summary of these cellular communications and the sites of ALC actions are shown in Figure 11. One of these communication systems is the TGFα-erbB1 receptor signaling system. We have made reference to research indicating that glial TGFα activates the erbB1/erbB2 receptor complex on adjacent glia in the MBH. This activation causes a cascade of events leading to the increased synthesis and release of PGE2, which then binds to its receptor on nearby LHRH neuron terminals causing stimulated release of the peptide. Additionally, evidence was presented showing that ALC is capable of interfering with hypothalamic glial to glial signaling involved with prepubetal PGE2 synthesis/release. The other communication network discussed in detail is the contactin-Caspr1 complex that binds to glial RPTPβ. After binding together, the newly formed three member family provides adhesiveness and promotes signaling between glia and LHRH nerve terminals in the MBH. Interestingly, we provided evidence that ALC exposure can interfere with this glial-neuronal communication family at puberty by suppressing the synthesis of glial RPTPβ. Additionally, we discussed how IGF-1 can influence both PGE2 and RPTPβ signaling. Overall, this review further indicates that glial-neuronal communications are important for LHRH secretion at puberty, and that ALC is capable of altering these cell-cell interactions. There are obviously other glial-neuronal systems not discussed here that deserve further investigation into their respective roles in neuroendocrine secretion at puberty, as well as determining whether they are affected by endocrine disruptive substances that may alter their functions.
Figure 11. Schematic drawing showing glial-LHRH neuronal associations and sites of ALC effects on RPTPβ in the median eminence of juvenile female rats. For clarity, details of other downstream pathways in this region are not shown. ALC, alcohol; BBB, blood brain barrier; Caspr1, contactin associated protein-1; EGF, epidermal growth factor; erbB1, EGF/TGF receptor; ER, estrogen receptor; IGF-1, insulin-like growth factor-1; IGFR, Insulin-like growth factor-1 receptor; LHRH, luteinizing hormone releasing hormone; PGE2, prostaglandin-E2; PGFR, prostaglandin-E2 receptor; RPTPβ, receptor protein tyrosine phosphatase β; TGF, transforming growth factor [73].

Acknowledgements

This work was supported by the NIH grant AA07216 (to WLD). The authors declare no conflict of interest.

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