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MicroRNAs flip the switch for the production of hypothalamic GnRH before puberty

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Abstract

A sparse population of a few hundred primarily hypothalamic neurons forms the hub of a complex neuroglial network that controls reproduction in mammals by secreting the "master molecule", gonadotropin-releasing hormone (GnRH). Timely postnatal changes in GnRH expression are essential for puberty and adult fertility. Here, we report that a multilayered microRNA-operated switch with inbuilt feedback controls governs increased GnRH expression during the infantile-juvenile transition, and that impairing microRNA synthesis in GnRH neurons leads to hypogonadotropic hypogonadism and infertility in mice. Two essential components of this switch, miR-200 and miR-155, respectively regulate *Zeb1*, a repressor of *GnRH* transcriptional activators and *GnRH* itself, and *Cebpb*, an NO-mediated repressor of *GnRH* that acts both directly and through *Zeb1*, in GnRH neurons. This alteration in the delicate balance between inductive and repressive signals induces the normal GnRH-fuelled run-up to correct puberty initiation, and interfering with this process disrupts the neuroendocrine control of reproduction.

Introduction

The onset of puberty and the regulation of fertility in mammals are governed by a complex neural network, primarily in the hypothalamus, that converges onto gonadotropin-releasing hormone (GnRH)-producing neurons, the master regulators of gonadotropin secretion and postnatal gonadal growth and function ¹. The proper development of the GnRH system ², including timely changes in GnRH expression and signaling by this sparse population of a few hundred neurons ^{3,4}, is essential for sexual maturation and the normal functioning of the hypothalamic-pituitary-gonadal axis. However, despite the identification of several factors influencing GnRH production and release ⁵⁻⁹, the mechanisms regulating the clocklike precision of the pubertal activation process remain mysterious.

MicroRNAs (miRNAs) are short (~22-nucleotide) noncoding RNA molecules that silence gene expression posttranscriptionally, principally by binding to the 3' untranslated regions (3'UTR) of target mRNAs (Fig. S1). Mature miRNAs have been shown to be required for the normal differentiation and function of several different cell types ¹⁰⁻¹³ and, in particular, to regulate somatic growth and fertility at the level of peripheral organs ¹⁴⁻¹⁶. We therefore asked whether miRNAs could play a similarly critical role in the central neuroendocrine control of reproduction.

Here we show that a dramatic switch in miRNA expression patterns in infantile GnRH neurons inverts the balance between inductive and repressive signals, triggering an increase in hypothalamic GnRH expression and thereby controlling the crucial transition from the early infantile phase, when its levels are low, to the GnRH-fuelled run-up to puberty. These data raise the intriguing possibility that the miRNA-dependent epigenetic regulation of GnRH secretion could underlie the pathophysiology of human hypogonadotropic hypogonadism when a genetic cause is not evident, and hold therapeutic potential for disorders of puberty and infertility of hypothalamic origin.

Results

Loss of Dicer function in GnRH neurons causes postnatal defects leading to hypogonadotropic hypogonadism and infertility

To determine whether miRNAs play a role in the crucial postnatal increase in GnRH expression, we generated mice in which Dicer, an RNAse III endonuclease essential for miRNA biogenesis ¹⁷, was selectively inactivated in GnRH neurons. Animals harboring a floxed *Dicer* allele ¹² were crossed with those expressing Cre recombinase under the control of the endogenous GnRH promoter ¹⁸ (Fig. 1A), which is expressed in hypothalamic GnRH neurons but not in the gonads (Fig. S2A,B). The resulting Gnrh::Cre; DicerloxP/loxP mice were viable, born at Mendelian frequencies, and had body weights at adulthood indistinguishable from *Dicer*^{loxP/loxP} (wild-type) littermates (t₍₈₎=4.35, p=0.390, n=5 per group). All male and female mutant mice, however, exhibited severe hypogonadism (testicular weight: $t_{(13)}$ =24.4, p=0.0001, n=5-10 per group; uterine weight: $t_{(4.47)}$ =10.2, p=0.0005, n=5-6 per group) (Fig. 1B-E) and sterility (males: n=6; females: n= 6) (Fig. 1F). While some female Gnrh::Cre; Dicer mice underwent vaginal opening (5 of 7) (Fig. 1G), this external sign of sexual maturation occurred significantly later than in $Dicer^{JoxP/loxP}$ mice (t₍₁₄₎=-4.93, p=0.0002), and vaginal smears showed that they never reached puberty (n=7) (Fig. 1H) despite normal somatic growth (two-way ANOVA, $F_{(1.96, 11.96)} = 0.5741$, p=0.54, n=5 per group) (Fig. 1I). Similarly, male *Gnrh*::Cre; *Dicer*^{loxP/loxP} mice (n=5) did not exhibit balanopreputial separation, which occurred at 34.9 ± 1.8 days of age in *Dicer* littermates (n=11), or spermatozoa in the seminiferous tubules (Fig. 1C). Additionally, serum levels of the pituitary gonadotropins luteinizing hormone (LH) and follicle-stimulating hormone (FSH) were markedly reduced in adult mutants (Fig. S2C,D). LH levels could readily be increased by systemic GnRH injection, demonstrating that pituitary function in our *Gnrh*::Cre; *Dicer*^{loxP/loxP} mice was intact (Fig. S2E). However, LH levels were unaffected by the intracerebroventricular infusion of two potent stimulators of the GnRH system that trigger LH release in wild-type animals: kisspeptin, which acts through its receptor Gpr54, and NMDA, which bypasses the kisspeptidergic system to act directly on GnRH neurons 9,19 (Fig. S2F,G), suggesting that the hypogonadotropic hypogonadism observed was due primarily to a hypothalamic GnRH deficiency.

Following these physiological data, neuroanatomical analyses revealed a striking and complete absence of GnRH immunoreactivity in the hypothalamus of adult mutant mice (Fig. S3A, B), reminiscent of Kallmann syndrome in humans ², in which GnRH neurons, which originate in the olfactory placode, fail to migrate into the brain during development. To determine whether *Dicer* invalidation in mouse GnRH neurons similarly affected their migration, we examined them on embryonic day (E) 14.5, when half the migrating GnRH neurons are still in the nose, and at birth (postnatal day 0, P0), when GnRH neurons have completed their migration into the hypothalamic preoptic region, their principal site of residence, and sent projections to the median eminence, where they release their neurohormone into portal capillaries for delivery to the anterior pituitary ^{20,21}. Surprisingly, and unlike the case in Kallmann patients, GnRH neurons displayed no migratory defects and GnRH immunoreactivity was comparable in *Gnrh*::Cre; *Dicer*^{loxP/loxP} and *Dicer*^{loxP/loxP} littermates at P0 (t₍₄₎=0.27, p=0.8, n=3 per group) (Fig. S4). GnRH deficiency in mutant mice was thus not due to a developmental lack of GnRH neurons, but acquired postnatally.

The GnRH peptide content of the hypothalamus increases progressively between birth and puberty, accelerating markedly during the latter part of the infantile period (P7-P12) ⁷. We thus examined hypothalamic GnRH immunoreactivity at different postnatal ages and found that its disappearance in *Gnrh*::Cre; *Dicer*^{loxP/loxP} mice occurred gradually, starting during the infantile period (t₍₁₀₎=1.96, p=0.08, n=5-7 per group at P12) (Fig. 2A, Fig. S5) and becoming more rapid after weaning, during the juvenile period, i.e. P21 to ~P35 (t₍₁₁₎=8.53, p<0.0001, n=6-7 per group at P21 and t_(6.59)=10.98, p<0.0001, n=7-9 per group at P28) (Fig. 2A, Fig. S5). Unexpectedly, however, the generation of *Gnrh*::Cre; *Dicer*^{loxP/loxP}; td*Tomato*^{loxP/STOP} trigenic mice, in which the tracer tdTomato continues to be expressed in GnRH neurons even without ongoing GnRH promoter activity, revealed that in the absence of miRNAs, GnRH neurons did not die but simply lost GnRH immunoreactivity (Fig. 2B). Indeed, at P21, ~70% of GnRH neurons had lost GnRH peptide expression in *Gnrh*::Cre;

Dicer^{loxP/loxP}; td*Tomato*^{loxP/STOP} mice when compared to *Gnrh*::Cre; *Dicer*^{+/+}; td*Tomato*^{loxP/STOP} littermates. Importantly, the concomitant loss of GnRH immunoreactivity and green fluorescent protein (GFP) in *Gnrh*::Gfp; *Gnrh*::Cre; *Dicer*^{loxP/loxP} mice (Fig. S3C), in which GFP is expressed under the GnRH promoter (unlike tdTomato in the previous experiment), indicated that miRNAs probably did not regulate GnRH transcript stability and/or translation directly by binding to its 3'UTR, but rather affected genes necessary for GnRH transcription, such as promoter modulators.

An infantile switch in miRNA species induces GnRH promoter activator expression in GnRH neurons *in vivo*

In order to tease apart the mechanism of miRNA action, we started by verifying the expression profile of the GnRH transcript in GnRH neurons isolated by FACS from Gnrh::Gfp mice wild-type for *Dicer* at two stages: P7, the beginning of the infantile period, and P12, when hypothalamic GnRH peptide content begins to increase in a functionally relevant manner in normal mice ⁷, whereas it begins to decrease in mice lacking *Dicer* even though the vast majority of their GnRH neurons (~80%) still express enough GFP to be detectable (Fig. S3). Real-time PCR analyses of GFP-expressing GnRH neurons isolated by fluorescence-activated cell sorting (FACS) from Gnrh::Gfp; Dicer of picer o S6) revealed that there was a 200% increase in GnRH mRNA levels between P7 and P12 in these *Dicer* wild-type mice ($t_{(5)}$ =-3.20, p=0.024, n=3-4 per group) (Fig 2D). This increase was accompanied by a significant upregulation of the mRNA levels of several promoter activators known to affect GnRH expression ^{22,23} (Fig. 2E), while transcripts of three known GnRH repressors ^{22,23} remained unchanged (Fig. 2F). However, in GnRH neurons lacking *Dicer* isolated from Gnrh::Gfp; Gnrh::Cre; DicerloxP/loxP trigenic mice at P12, GnRH mRNA expression was significantly lower than in neurons from Dicer wild-type Gnrh::Gfp; Dicer mice $(t_{(3,12)}=5.88, p=0.0098, n=4 per group)$ (Fig. 2G). In addition, there was a marked decrease in the expression of most GnRH promoter activators (Fig. 3A) and a striking 15-fold increase in the transcript of the GnRH promoter repressor Cebpb (Fig. 3B) in GnRH neurons in the absence of Dicer (Fig. 3A,B). *In silico* analysis of the 3'UTR sequence of these promoter modulators revealed the presence of putative binding sites for several miRNAs (p<0.01; Table S1), confirming that miRNAs could regulate the expression of these genes. Concordantly, we uncovered a dramatic inversion of the expression profile of these and other putative regulatory miRNAs in wild-type GnRH neurons between P7 and P12 (Fig. 3C), strongly suggesting that GnRH promoter activity during the infantile period is controlled by a switch operated by miRNAs.

The miR-200/Zeb1 network is involved in controlling hypothalamic GnRH expression during postnatal development

The decrease in the expression of GnRH promoter activators uncovered above suggests that in the absence of miRNAs, repressors of these activators might be reciprocally increased. To identify the putative molecular targets involved in this process, we looked for miRNAs differentially expressed between P7 and P12 (Fig. 3C) that were also specifically enriched in GnRH neurons. The results showed that members of the miRNA-200 family, including miR-141 and miR-429 (Fig. S7A), were among the most highly expressed miRNAs in infantile GnRH neurons when compared to non-GnRH-expressing cells (Fig. 3D and 3E). Within this family, the expression of miR-200a and miR-429, known to silence the mRNA for the transcriptional repressor Zeb1 outside the brain in pituitary gonadotropes, where it regulates LH expression ¹⁴, increased significantly between P7 and P12 (miR-200a, t₍₄₎=9.178, p=0.0008; miR-429, $t_{(5)}$ =3.931, p=0.0111, n=3-4 per group) (Fig 3F). To explore the possibility that the miR-200/Zeb1 network 24 (Fig. S7A) also controlled hypothalamic GnRH expression during postnatal development, we analyzed Zeb1 expression in the brain of mice that did or did not express miRNAs in GnRH neurons. Importantly, not only was Zeb1 mRNA expressed in infantile GnRH neurons, its expression decreased significantly between P7 and P12 in *Gnrh::*Gfp; *Dicer*^{+/+} mice ($t_{(6)}$ =4.86, p=0.0028, n=4 per group) (Fig. 4A). RT-PCR experiments in GnRH neurons isolated from Gnrh::Gfp; Gnrh::Cre; Dicer and Gnrh::Gfp; Dicer loxP/loxP littermates at P12 further showed that Zeb1 mRNA was significantly upregulated in mutant mice ($t_{(5)}$ =-3.38, p=0.0195, n=3-4 per group) (Fig. 4B). In silico sequence analysis revealed varying numbers of putative Zeb1-binding sites in the genes of all GnRH transcriptional activators under scrutiny (Fig. S7B), including Pou2f1, which in addition to activating the GnRH gene directly, could increase levels of miR-200 25, thereby indirectly influencing GnRH activation by repressing Zeb1. In addition, the GnRH gene itself was found to contain putative Zeb1-binding sites, three of which we validated as being functionally active using chromatin immunoprecipitation assays in an immortalized GnRHexpressing mouse cell line (Fig. 4C). Altogether these data support the idea that increased Zeb1 expression in the absence of miRNA biogenesis could mediate the downregulation of these activators and of GnRH in these neurons (Fig. 3A). To test this hypothesis, we infused a target-site blocker, TSB-200, into the brains of Gnrh::Gfp; Dicer+/+ mice to selectively hamper the binding of miR-200b/200c/429 to Zeb1 mRNA (Fig. S7C). TSBs are small, stabilized, freely diffusible locked nucleic acid (LNA) oligonucleotides, complementary to both the miRNA and the binding site being targeted (see for example Fig. S8C, D, E). As shown by fluorescence-tagged sequences (Fig. S8A), TSBs readily target the preoptic region and GnRH neurons 72h after their infusion into the lateral ventricle of *Gnrh::*Gfp; *Dicer**/+ pups. TSB-200 treatment significantly increased Zeb1 mRNA levels in GnRH neurons as compared to a scrambled sequence (t₍₉₎=-3.33, p=0.0087, n=5-6 per group) (Fig. 4D), but did not affect the transcripts of two other miR-200b/200c/429 target genes, Lpin1 26 (t₍₁₀₎=0.663, p=0.52, n=6 per group) and Maf²⁷ ($t_{(10)}$ =-0.192, p=0.85, n=6 per group) (Fig. 4D). This increase in Zeb1 expression was accompanied by a significant decrease in the induction of transcripts for two GnRH promoter activators that were upregulated in GnRH neurons at P12 compared to P7 (Fig. 2E), *Meis1* ($t_{(8)}$ =2.53, p=0.035, n=5 per group) and *Pou2f1* ($t_{(10)}$ =4.15, p=0.002, n=6 per group) (Fig. 4E), while transcripts for other known promoter activators, such as Aes, Otx2, Pbx1 and Pknox1, which did not significantly change between P7 and P12, were also unchanged by TSB-200 infusion. Interestingly, these genes also had fewer putative Zeb1binding sites than those whose expression was strongly modified by altered Zeb1 expression (Fig. S7B). As expected, GnRH transcription was also downregulated by the blockade of miR-200 binding to Zeb1 at P12 ($t_{(9)}$ =5.37, p=0.0004, n=4-7 per group) (Fig. 4F). This repressive effect on GnRH mRNA levels was apparently mediated by the inhibition of GnRH promoter activity, as shown by the dramatic decrease in Gfp reporter mRNA expression ($t_{(10)}$ =14.1, p<0.0001, n=6 per group) (Fig. 4G), suggesting that the miR-200/Zeb1 network is actively involved in the switch in the control of GnRH transcription in the infantile mouse hypothalamus (Fig. 4H).

Hampering selective miR-155 binding to the GnRH repressor *Cebpb* activates *Zeb-1* expression and inhibits infantile GnRH promoter activity *in vivo*

In contrast to the downregulation of GnRH promoter activators in mice lacking Dicer in GnRH neurons, the mRNA for a GnRH repressor, Cebpb, which codes for the CAAT/enhancer binding protein β (C/EBP β), was found to be dramatically upregulated (see Fig. 3B above). We thus reasoned that the repression of C/EBPB expression by miRNAs would increase radically during the infantile period in wild-type mice. Using the miRWalk database, which provides validated information on miRNA-target interactions ²⁸, we found that among the miRNAs upregulated in GnRH neurons at P12 (Fig. 3C, 3D) was miR-155, whose seed region targets the 3'UTR of the Cebpb transcript (p<0.01, Table S2, Fig. S8B,C) 29. To determine whether the ten-fold increase in miR-155 expression between P7 and P12 (t₍₄₎=-4.85, p=0.0083, n=3 per group) (Fig. 5A) played an actual role in regulating C/EBPβ expression in GnRH neurons, we used TSB-155 to selectively hamper miR-155 binding to the Cebpb transcript in the hypothalamus of infantile mice (Fig. S8D, S8E). TSB-155 infusion into Gnrh::Gfp; Dicer+++ mice markedly increased the levels of Cebpb mRNA in GnRH neurons when compared to a scrambled sequence ($t_{(9)}$ =-3.41, p=0.0038, n=5-6 per group) (Fig. 5B), but did not change mRNA levels of Lpin1 ($t_{(10)}$ =0.601, p=0.56, n=6 per group) and Maf (t₍₁₀₎=0.406, p=0.69, n=6 per group) (Fig. 5B), which are also validated targets for miR-155 30,31. Importantly, this TSB-155-mediated upregulation of the Cebpb transcript was associated with a strong inhibition of both GnRH promoter activity, as indicated by Gfp mRNA levels ($t_{(10)}$ =14, p<0,0001, n=6 per group) (Fig. 5C) and GnRH mRNA levels ($t_{(8)}$ =3.57, p=0.0073, n=4-6 per group) (Fig. 5D). Surprisingly, we found, using ChIPBase, an integrated resource and platform for decoding transcription factor binding maps based on ChipSeq data ³², that the Zeb1 gene contained binding sites for C/EBPβ in six different regulatory regions ^{33,34} (Fig. S9) but no binding site for miR-155, suggesting that C/EBPβ could regulate GnRH mRNA levels not only directly but indirectly, by influencing Zeb1 expression. In agreement with this hypothesis, TSB-155 treatment, which selectively upregulates the Cebpb transcript, also increased Zeb1 mRNA expression in Dicer++ GnRH neurons (t₍₉₎=-2.89, p=0.0177, n=5-6 per group) (Fig. 5E). Furthermore, in silico and ChipSeq analysis revealed several binding sites for both C/EBPβ- and Zeb1 on the *Gpr54* gene (Fig. S7B and Fig. S9), suggesting that in addition to their direct and promoter-activator-mediated effects, C/EBPß and Zeb1 could repress GnRH gene transcription indirectly by blocking kisspeptinergic signaling through Gpr54, which has been shown to promote the nuclear translocation of the GnRH promoter activator Otx2 and the subsequent increase in GnRH transcription 35 (Fig. 5F). In support of this hypothesis, RT-PCR analysis revealed that in animals lacking Dicer in GnRH neurons, *Gpr54* mRNA levels were also strongly reduced ($t_{(6)}$ =4.22, p=0.0056, n=4 per group) (Fig. 5G), to a similar extent as that of other GnRH activators (Fig. 3A).

Blockade of infantile NO release partially rescues GnRH expression in GnRH neurons lacking functional Dicer

C/EBPβ is also known to mediate the nitric oxide (NO) -induced repression of GnRH gene expression ²². However, we have recently shown that the activation of neuronal NO synthase (nNOS) in neurons of the preoptic region, itself regulated by the actions of Kisspeptin on its receptor Gpr54 ³⁶, is required for sexual maturation ³⁷. Intriguingly, in wild-type mice in the present study, nNOS activity increased significantly in the preoptic region between P7 and P12, at a time when synaptic connectivity is being established in the GnRH neuronal network ^{38,39}, as shown by the increase in the phosphorylation of its Ser1412 activation site ⁴⁰ (p-

nNOS/NOS ratio, $t_{(6)}$ =-5.48, p=0.0015, n=4 per group) (Fig. 5H). To determine whether the marked decrease in GnRH expression in mice lacking *Dicer* could be related to this infantile increase in NO-C/EBPβ signaling, we next treated P12 Gnrh::Gfp; Gnrh::Cre; DicerloxP/loxP mice with the NOS inhibitor N-G-Nitro-L-arginine methyl ester (L-NAME; 50 mg/kg, i.p.) and assessed GnRH mRNA expression in FACS-isolated neurons 12h later. The reduced production of NO due to L-NAME treatment partially but significantly rescued GnRH transcript levels (t(8)=-2.40, p=0.043, n=6-7 per group) (Fig. 5I). In addition to the alleviation of the previously demonstrated repressive effect of C/EBP\$ on the GnRH promoter ²², this increase in GnRH expression could have resulted from the inhibition of the transcriptional activation of Zeb1 by C/EBPβ, as suggested by the 2.4-fold increase in transcripts for Pou2f1 (in L-NAME-injected vs. saline-injected mice; t(8)=-1.40, p=0.017, n=5-6 per group). This suggests that increased NO levels during this period crucial for subsequent GnRH neuronal function might actually participate in the multilayered transcriptional repression of the Gnrh gene in the absence of posttranscriptional silencing of C/EBPβ by miRNAs (Fig. 5J). In other words, not only do miRNAs control the switch in GnRH promoter activity in infantile GnRH neurons, and thus reproductive capacity, they do so through an intricate mechanism involving crosstalk between inductive and repressive pathways.

miR-200 family members bind to *Zeb-1* to control GnRH neuronal function throughout life

To assess the functional involvement of miR-200 family members and miR-155 in GnRH neurons in the onset of puberty, we performed a single bolus injection of the same TSB-200 and/or TSB-155 into the brains of female wild-type mice at P9, and monitored sexual maturation after weaning. Importantly, the use of fluorescently-labeled TSBs demonstrated that they could be detected in the brain up to 2 weeks after injection (Fig. S8A). Surprisingly, while growth (Fig S10) was similar in TSB- and scrambled-sequence-injected mice, TSB-injected mice exhibited a precocious onset of puberty when compared to control mice (Fig. 6A). This effect was most pronounced in TSB-200-infused mice, with 50% of the animals

displaying their first estrus by P32, i.e. more than 10 days before control mice. In line with these results, TSB-200-infused mice showed significantly elevated peripubertal afternoon surges of plasma LH at P38, while changes in LH levels induced by TSB-155 treatment did not reach statistical significance when compared to control mice, in which plasma LH levels were below the limit of detection (Fig.6B). The combination of TSB-200 and TSB-155 also increased LH to detectable levels, although this effect was minimal (Fig. 6B; F(3, 32)=3.475, p=0.0272, n=8-10 per group; TSB-200, p=0.0361; TSB-155, p=0.7541; TSB-200+TSB-155, p=0.9996) and resulted in earlier puberty onset by about 5 days when compared to control mice. In keeping with its lack of a significant effect on LH levels, treatment with TSB-155 alone failed to significantly change age at puberty (Fig. 6A; F(3, 35)=4.961, p=0.0057, n=9-10 per group; TSB-200, p=0.0019; TSB-155, p=0.073; TSB-200+TSB-155, p=0.0328). To verify whether this discrepancy in the impact of blocking either miR-200/429-Zeb1 binding or miR-155-Cebpb binding on GnRH promoter activity at P12 (Fig. 4F, 4G and Fig. 5C, 5D, respectively) and on subsequent sexual maturation (Fig 6A) was linked to differences in the specific cell types targeted by TSB-155 and TSB-200 in the hypothalamus, we analyzed the effects of TSB injection on the expression of Cebpb and Zeb1 in non-GnRH cells at P12. In agreement with the level of enrichment of miRNA expression in GnRH neurons compared to non-GnRH hypothalamic cells, which is about 50 fold higher for miR200/429 (Fig. 3E) than for miR-155 (Fig. S11A), Cebpb mRNA levels were significantly increased in non-GnRH cells of the preoptic region after TSB-155 infusion (Fig. S11B), while TSB-200 treatment altered neither Cebpb nor Zeb1 transcripts in these cells (Fig. S11B, S11C). Altogether, these data suggest that, in keeping with the complex feedback mechanisms at work (Fig. 7), selectively applying the brakes to GnRH production in an excessive manner (Fig. 4F, 4G) and then releasing them prematurely can trigger precocious puberty, possibly due to a rebound effect. To determine whether the miR-200/Zeb1 network could control GnRH function outside of this critical infantile time window leading up to puberty, we infused TSB-200 into the hypothalamic preoptic region of cycling adult female mice and followed their estrous cycles (Fig. 6C). While control mice infused with scrambled sequences did not exhibit any marked alteration of their estrous cycles, most mice treated with TSB-200 (5 out of 6) showed a slight prolongation in the number of days spent in diestrus (a basal stage when LH is released at nadir levels) and eventually reached proestrus (when the preovulatory surge of LH occurs), but displayed an incomplete cycle at least once during the first two weeks after injection (Fig. 6C). In line with this observation, LH levels measured during proestrus one week after injection showed a characteristic surge in control animals but not in TSB-200-treated mice (Fig. 6D; F(2, 14)=4.371, p=0.0335, n=5-6 per group; TSB-200 > 1 week, p=0.0262; TSB-200 > 2 weeks, p=0.6976). In 4 of the 6 animals treated with TSB-200, LH levels recovered and displayed normal preovulatory surges by the third week after TSB infusion (Fig. 6D).

Discussion

The balance between inductive and repressive signals is a process essential for successful reproduction, not only in animals but in living organisms in general 41,42. Our data provide the first physiological and neuroanatomical evidence that miRNAs, which have been shown to regulate somatic growth, including the development of peripheral sexual organs 14-16, also play an essential role in the postnatal development of the hypothalamic neuroendocrine neurons controlling the onset of puberty and fertility in mammals, by modulating this balance in the GnRH gene network. During this infantile "critical period" lasting just a few days, when the first centrally-driven gonad-independent activation of the hypothalamic-pituitary-gonadal axis occurs, setting in motion, for example, the growth of the first wave of ovarian follicles that will ovulate at puberty in females and the development of the testes in males (see for review 43), our findings show that a switch in miRNA expression in turn flips a switch in a multilayered array of GnRH gene activators and repressors, permitting the sustained increase of the neurohormone required for normal sexual maturation. Interfering with the binding and functional effects of two key miRNA species, miR-200/429, which is not only upregulated during this critical period but selectively enhanced in GnRH neurons, and miR-155, which appears to act on other hypothalamic cell types as well, and mediates, for example, the effects of a concomitant release of NO upstream of GnRH neurons, blunts this

infantile increase in *GnRH* expression. In addition, this *in vivo* alteration of the miR-200/429-transcription factor micro-network leads to the disruption of normal puberty onset as well as normal estrous cyclicity in adulthood.

The fact that GnRH neurons form an extremely small population (~800 neurons in a mouse brain) and their scattered distribution in a continuum from the olfactory bulb to the hypothalamus, albeit with a strong concentration in the hypothalamic preoptic area, have made genetic and epigenetic studies in these neurons a veritable technical challenge until now. Our successful separation of GnRH from non-GnRH cells in the preoptic region of *GnRH*::Gfp mice has allowed us to determine miRNA expression profiles specific to this very limited population of neurons in the postnatal brain, i.e. once the processes of differentiation and the establishment of connectivity, including their projection over several millimeters to the hypothalamic median eminence, are complete. The difficulty of this approach is evident from the fact that other such studies, even those focusing on more abundant neuronal types, have not always succeeded ⁴⁴. Using this approach, we were not only able to study changes in miRNA expression in GnRH neurons at different stages of postnatal development, but even to analyze the effects of the selective invalidation of miRNA biogenesis and the acute manipulation of specific miRNA networks on the expression of transcription factors, GnRH promoter activity and GnRH expression in the postnatal brain *in vivo*.

Our findings show that miRNA processing in GnRH neurons is required to promote sexual development, but surprisingly, not GnRH neuronal migration *in vivo*. The specific role of miRNAs in sustaining the expression of GnRH is shown in *Gnrh*::Cre; *Dicer*^{loxP/loxP} mice, in which GnRH expression appears to be normal during the perinatal and the early infantile periods, and the process of sexual maturation commences normally and leads to vaginal opening in 80% of the individuals. However, these mice never achieve puberty because of the progressive decrease in GnRH promoter activity from the second week of postnatal life and its complete extinction during the juvenile period. Our data indicate that miRNA species whose expression is upregulated in GnRH neurons during this critical period of 7 to 12 days of age include miR-155 and members of the miR-200 family. Notably, the striking enrichment

of miR-200 family members during this period is restricted to GnRH neurons, indicating a highly specific function for these miRNA species in the regulation of GnRH expression. We have identified several genes targeted directly or indirectly by miR-155 and miR-200/429, whose repression (e.g. Cebpb and Zeb1) or activation (e.g. Pou2f1 or Meis1, which are both targets for Zeb1) is required for normal GnRH expression at 12 days of age. In particular miR-200/429 and its target Zeb1, which can directly repress the GnRH promoter, also seem to form a double-negative loop with the key GnRH promoter activator Pou2f1, while miR-155 counteracts NO/Cebpb-dependent GnRH repression. Nor can we rule out the possibility that some molecules act as both activators and repressors depending upon physiological conditions and cellular context, as we have seen in the case of Cebpb, a GnRH promoter repressor that nevertheless increases Zeb1 levels in GnRH neurons but not in non-GnRH hypothalamic cells (which do not express miR-200/429), or that some of these promoter modulators could exert a reciprocal regulatory effect on the miRNAs that control their expression, as is the case with Zeb1 and miR-200/429 ²⁴.

Surprisingly, the pattern of expression of various known GnRH promoter modulators is not the same at P12. While activators that are upregulated specifically in GnRH neurons in wild-type mice at P12 show reduced expression with the blockade of miR-200/429 binding (with the exception of Dlx5, which has no Zeb1 binding site and is thus probably regulated by other miRNA-effector pathways), the expression of one putative activator, Dlx1, is increased. Apart from a possible indirect effect, the fact that unlike Pou2f1 or Meis1, the expression of Dlx1 shows a downward trend in wild-type GnRH neurons at P12 could explain this seemingly paradoxical finding. The lack of effect of a miRNA-200/429 blockade on the expression of GnRH activator genes that contain no or few Zeb1-binding sites and that are not differentially expressed during the infantile-juvenile transition in GnRH neurons (but that could nevertheless play a role in baseline GnRH production or its regulation in other physiological contexts) further highlights the importance of the miR-200/Zeb1/Cebpb switch in the prepubertal increase in GnRH expression. Our findings show, in addition, that both miR-155 (by interfering with NO/Cebpb-mediated signaling by nNOS neurons, themselves activated

by Kisspeptin ³⁶) and miR-200/429 (through Zeb1) could also play a role in the intricate relationship between different cells of the hypothalamic GnRH network by affecting the expression of the Kisspeptin receptor Gpr54, and thus modulating the Kisspeptin-Gpr54-mediated activation of GnRH expression. NO itself, by being both essential for the initiation of puberty ³⁷ and participating in the repression of GnRH expression ²², could play a more complex role than suspected hithertofore. The Kisspetin/nNOS/GnRH neuronal microcircuit and miRNA-gene network in GnRH neurons (Figure 7) would be ideally poised to ignite rhythmic GnRH gene transcription, which has recently been associated to pulsatile GnRH secretion ^{45,46} in response to developmental and bodily cues during the infantile period in mice ⁴³, a developmental stage highly reminiscent of minipuberty in humans ⁴⁷.

Other feedback loops, possibly involving the expression of miR-155 in other cell types, could also exist, as suggested by the fact that simultaneously neutralizing both miR-200/429 binding to Zeb1 and miR-155 binding to Cebpb results in a considerably less striking effect on peripubertal LH levels than the blockade of miR-200/429 alone. Intriguingly, while blocking the binding of miR-200/429 to Zeb1 during this infantile-juvenile critical period dramatically reduces GnRH promoter activity at the cellular level, these animals go on to display precocious, not delayed, puberty. This counterintuitive finding could stem from a rebound effect, where the prolonged inhibition of GnRH expression (TSBs, once injected, remain at the injection site for around 2 weeks) results in a compensatory spike once the inhibition is relieved. Such a phenomenon has been demonstrated previously in human populations, where children whose GnRH system is blunted due to exposure to endocrine disruptors of agricultural origin undergo precocious puberty when they migrate to countries that do not use them ⁴⁸.

To conclude, our data are the first to indicate not only that miRNAs are active in the hypothalamic GnRH network, but that miRNA-200/429 and miRNA-155 are key components of a complex developmental switch that controls the activity of the GnRH promoter, as illustrated in the models presented in Figure 7 and Figure S12, and whose correct functioning is essential for the normal initiation of puberty and reproductive success in adulthood.

Moreover, this dramatic switch in gene expression appears to involve a multilayered and interconnected array of miRNAs and their target genes in GnRH neurons, with its own inbuilt feedback control mechanism, that alters the balance between inductive and repressive signals during the infantile critical period and thus leads to increased GnRH production. Deciphering the mechanisms through which miRNAs contribute to precocious or delayed puberty through their actions on the functioning of the GnRH neural network may thus not only provide novel insights into the epigenetic regulation of maturational processes, fascinating in itself, but pave the way to a better understanding of idiopathic infertility ^{49,50} and the elaboration of new diagnostic and therapeutic options in humans.

Materials and Methods

Animals

All C57Bl/6J mice were housed under specific pathogen-free conditions in a temperature-controlled room (21-22°C) with a 12h light/dark cycle and *ad libitum* access to food and water. *Gnrh:*:Cre (Tg(*Gnrh1::*Cre)1Dlc), *Dicel*-LoxP, and *Gnrh::*Gfp mice were a generous gift of Dr. Catherine Dulac (Howard Hughes Medical Institute, Cambridge MA) ¹⁸, Dr. Brian Harfe (University of Florida, FL) ¹² and Dr. Daniel J. Spergel (Section of Endocrinology, Department of Medicine, University of Chicago, IL) ⁵¹, respectively. *tdTomato*loxP/STOP mice (B6.Cg-*Gt*(*ROSA*)26Sortm9(CAG-tdTomato)Hze/J) were purchased from the Jackson laboratory (Maine, USA). Mice were genotyped by PCR using primers listed in table S3. Animal studies were approved by the Institutional Ethics Committees for the Care and Use of Experimental Animals of the Universities of Lille and Cordoba; all experiments were performed in accordance with the guidelines for animal use specified by the European Union Council Directive of September 22, 2010 (2010/63/EU).

Physiological measurements

Fertility index. Male and female fertility indices were calculated from the number of litters per females during a 120-day-long mating.

Puberty onset. Weaned female mice were checked daily for vaginal opening. After vaginal opening, vaginal smears were performed daily and analyzed under an inverted microscope to identify the specific day of the estrous cycle. Weaned males were checked daily for balanopreputial separation.

Hormone level measurements. Protocols and doses for in vivo testing of the LH response to GnRH, kisspeptin-10 and NMDA were as described in detail elsewhere ⁵². Serum LH and FSH levels were measured using radioimmunoassay kits supplied by the National Institutes of Health (Dr. A. F. Parlow, National Hormone and Peptide Program, Torrance, CA). Rat LH-I-10 and FSH-I-9 were labeled with ¹²⁵I using Iodo-gen tubes, following the instructions of the manufacturer (Pierce, Rockford, IL). Hormone concentrations were determined using reference preparations of LH-RP-3 and FSH-RP-2 as standards. Intra- and inter-assay coefficients of variation were <8 and 10% for LH and <6 and 9% for FSH, respectively. The sensitivity of the assay was 3.75 pg/tube for LH and 20 pg/tube for FSH. The accuracy of hormone measurements was confirmed by the assessment of rodent serum samples of known concentration (external controls).

Ovarian histology and quantitative analysis

Ovaries and testes were collected from 3-month-old *Dicer*^{LoxP/LoxP} and *Gnrh::*Cre; *Dicer*^{LoxP/LoxP} mice, immersion-fixed in 4% PFA solution and stored at 4°C. Paraffin-embedded ovaries and testes were sectioned at a thickness of 5 µm (histology facility, University of Lille 2, France) and stained with hematoxylin-eosin.

Tissue preparation

For immunohistochemical analysis, embryos were obtained after cervical dislocation from timed-pregnant *Gnrh::*Cre; *Dicer*^{LoxP/+} crossed with *Gnrh::*Cre; *Dicer*^{LoxP/LoxP} male mice. Embryos were washed thoroughly in cold 0.1 m PBS, fixed in fixative solution [4%]

paraformaldehyde (PFA), 0.2% picric acid in 0.1 m PBS, pH 7.4] for 6–8 h at 4°C and cryoprotected in 20% sucrose overnight at 4°C. The following day, embryos were embedded in OCT embedding medium (Tissue-Tek), frozen on dry ice, and stored at -80°C until sectioning. Postnatal (P0 to P28) and adult mice (3–5 months old) were anesthetized with 50-100 mg/kg of Ketamine-HCl and 5-10mg/kg Xylazine-HCl and perfused transcardially with 2-10 ml of saline, followed by 10-100 ml of 4% PFA, pH7.4. Brains were collected, postfixed in the same fixative for 2 h at 4°C, embedded in OCT embedding medium (Tissue-Tek), frozen on dry ice, and stored at -80°C until cryosectioning.

Immunohistochemistry

Tissues were cryosectioned (Leica cryostat) at 16μm for embryos and pre-weaning postnatal mice, and at 35μm (free-floating sections) for post-weaning and adult brains. Immunohistochemistry was performed as previously reported ^{53,54}, using Alexa-Fluor 488-(1:400) and Cy3-conjugated (1:800) secondary antibodies (Invitrogen). Fluorescent specimens were mounted using 1,4-diazabicyclo[2.2.2]octane (Sigma-Aldrich). The primary antisera used were as follows: rabbit anti-GnRH (1:3000), a generous gift from Prof. G. Tramu (Centre Nationale de la Recherche Scientifique, URA 339, Université Bordeaux I, Talence, France) ⁵⁵.

Isolation of hypothalamic GnRH neurons using Fluorescence-Activated Cell Sorting

The preoptic regions of *Gnrh::*Gfp; *Dicer*^{LoxP/LoxP} and *Gnrh::*Gfp; *Gnrh::*Cre; *Dicer*^{LoxP/LoxP} mice were microdissected and enzymatically dissociated using a Papain Dissociation System (Worthington, Lakewood, NJ) to obtain single-cell suspensions. FACS was performed using an EPICS ALTRA Cell Sorter Cytometer device (BD Bioscience). The sort decision was based on measurements of GFP fluorescence (excitation: 488nm, 50 mW; detection: GFP bandpass 530/30 nm, autofluorescence bandpass 695/40nm) by comparing cell suspensions from *Gnrh::*Gfp and wild-type animals, as indicated in figure S5. For each animal, 400 to 800

GFP-positive cells were sorted directly into 10µl extraction buffer: 0.1% Triton® X-100 (Sigma-Aldrich) and 0.4 U/µl RNaseOUT™ (Life Technologies).

Quantitative RT-PCR analyses

software (Applied Biosystems).

For gene expression analyses, mRNAs obtained from FACS-sorted GnRH neurons were reverse transcribed using SuperScript® III Reverse Transcriptase (Life Technologies) and a linear preamplification step was performed using the TaqMan® PreAmp Master Mix Kit protocol (P/N 4366128, Applied Biosystems). Real-time PCR was carried out on Applied Biosystems 7900HT Fast Real-Time PCR System using exon-boundary-specific TagMan® Gene Expression Assays (Applied Biosystems): Aes (Aes-Mm00507847 m1), Cebpb (Cebpb-Mm00843434_s1), *Dlx1*(Dlx1-Mm00438424_m1), *Dlx5* (Dlx5-Mm00438430_m1), (Gfp-Mr03989638_mr), (Gnrh1-Mm01315605_m1), Gfp Gnrh1 Gpr54 (Gpr54-Mm00475046_m1), Lpin1 ((Lpin1-Mm00550511_m1), Maf (Maf-Mm02581355_s1), Meis1 (Meis1-Mm00487664 m1), Msx1 (Msx1-Mm00440330 m1), Otx2 (Otx2-Mm00446859 m1), Pbx1 (Pbx1-Mm04207617_m1), Pknox1 (Pknox1-Mm00479320_m1), Pou2f1 (Pou2f1-Mm00448332_m1), Tle4 (Tle4-Mm01195172_m1), Zeb1 (Zeb1-Mm00495564_m1). Control housekeeping genes: r18S (18S-Hs99999901_s1); ACTB (Actb-Mm00607939_s1). MicroRNA expression analyses were performed using stem-loop RT-PCR based TaqMan Rodent MicroRNA Arrays (Applied Biosystems). Briefly, miRNAs obtained from FACS-sorted GnRH neurons were reverse transcribed using the TaqMan miRNA Reverse Transcription Kit (Applied Biosystems) in combination with the stem-loop Megaplex primer pool sets A and B according to the manufacturer's instructions. A linear preamplification step was performed

recommended cycling conditions.

Gene and miRNA expression data were analyzed using SDS 2.4.1 and Data Assist 3.0.1

using the TaqMan® PreAmp Master Mix Kit protocol (P/N 4366128, Applied Biosystems) and

quantitative real-time PCR were performed using TaqMan Low-Density Arrays (Applied

Biosystems) on an Applied Biosystems 7900HT thermocycler using the manufacturer's

Zeb1 chromatin immunoprecipitation (ChIP) assays

Immortalized GnRH neurons (GN11 cells) were cultured in DMEM + 10% FBS (Life Technology). Prior to experimental assays, cells were washed twice in PBS and incubated overnight in serum-free medium. The next morning, 10% FBS was added for 2h. Chromatin isolation was performed using the ChIP-IT Express kit (Active Motif) following the manufacturer's instructions. After DNA shearing by sonication (9 min total, 20s pulses at 70% power followed by a 40s pause; on ice), protein-bound DNA was immunoprecipitated at 4°C overnight using 5 µg of a polyclonal goat anti-rabbit Zeb1 antibody (E20; Santa Cruz Biotechnology; SC-10572) or control goat IgGs (Santa Cruz Biotechnology; sc-2028). ChIP-DNA fragments were purified using a QIAquick PCR purification kit (Qiagen), and analyzed by qPCR on a CFX96 thermal cycler (Biorad) using GoTaq® qPCR Master Mix (Promega) and the primers indicated in Table S4.

Western blot analyses

The preoptic area of the hypothalamus was dissected from each animal using Wecker scissors (Moria, France) under a binocular magnifying glass, and protein extracted in 200 μl of lysis buffer (25 mM Tris, pH 7.4, β-glycerophosphate, 1.5 mM EGTA, 0.5 mM EDTA, 1 mM sodium pyrophosphate, 1 mM sodium orthovanadate, 10 μg/ml leupeptin and pepstatin A, 10 μg/ml apoprotinin, 100 μg/ml PMSF, and 1% Triton-X100) by trituration of the fragments through 22 and 26 gauge needles in succession. Tissue lysates were cleared by centrifugation at 14000 rpm for 15 min at 4°C. Protein content was determined using the Bradford method (Bio-Rad, Hercules, CA) and equal amounts of protein were mixed with 4X sample buffer (Invitrogen). Samples were boiled for 5 min and stored at -80°C until use. Samples were reboiled for 5 min after thawing and electrophoresed for 75 min at 150 V in 7% Tris-acetate, or for 50 min at 200 V in precast 4–12% MES SDS-polyacrylamide gels according to the protocol supplied with the NuPAGE system (Invitrogen). After size fractionation, the proteins were transferred onto 0.2 μm pore-size polyvinylidene difluoride

membranes (LC2002; Invitrogen) in the blot module of the NuPAGE system (Invitrogen) for 75 min at room temperature. Membranes were blocked for 1 h in blocking buffer [TBS with 0.05% Tween 20 (TBST) and 5% nonfat milk] at room temperature, and incubated overnight at 4°C with the appropriate primary antibody diluted in blocking buffer (rabbit polyclonal anti-RFP, 600-401-379, 1:1000, Rockland antibodies & assays; rabbit polyclonal anti-nNOS, sc-8309, 1:500, Santa Cruz technologies; rabbit polyclonal anti-Ser¹⁴¹² phospho-nNOS, PA1-032; 1:1000, Affinity BioReagents; goat polyclonal anti-actin, sc-1616, 1:1000). Membranes were washed four times with TBST the following day before being exposed to HRP-conjugated secondary antibodies diluted in blocking buffer for 1 h at room temperature. Immunoreactions were visualized using the ECL detection kit (NEL101; PerkinElmer, Boston, MA). Immunoblots were scanned using a desktop scanner (Epson Expression 1680 PRO) and Adobe Photoshop, and band intensities were determined using ImageJ software (NIH, Bethesda).

Target site blockers (TSBs)

To selectively hamper the silencing activity of miR-155 and miR-200b/200c/429 on *Cebpb* and *Zeb1* respectively in vivo, we used custom-designed target sites blockers with phosphorothioate backbone modifications from Exiqon (miRCURY LNATM microRNA Target Site Blockers, in vivo ready). TSB sequences are designed with a large arm that covers the miRNA binding site and a short arm outside the miRNA seed to ensure target specificity. In detail, one sequence was generated to protect the unique miR-155-binding site in the *Cebpb* 3'UTR (TSB-155, Fig. S7). Five sequences were generated to protect the five different miR-200b/200c/429-binding sites in the *Zeb1* 3'UTR (Fig. S7) and mixed at equimolar ratios (TSB-200).

Stereotactic brain infusions of TSBs

Gnrh::Gfp mice were placed in a stereotaxic frame (Kopf® Instruments, California) under anesthesia (isoflurane), and a burr hole was drilled 1 mm lateral to the bregma, according to

a mouse brain atlas (Paxinos and Franklin, 2004). A 10 μ l Hamilton syringe was slowly inserted into the bottom of the left lateral ventricle (3.5 mm deep relative to the dura), and 1 μ l of the different treatment solutions (TSB-200, 50 μ M, n = 6; TSB-155, 50 μ M, n = 6) or vehicle (PBS, pH7.4, n = 6) was injected using an infusion pump (KD Scientific, Holliston, MA) over 4 min. Animals were subjected to intracranial surgery at P9 and killed at P12.

Identification of putative Zeb1-binding sites

To identify putative Zeb1- and C/EBPβ-binding sites in the promoter regions of target genes, 1.5 Kb of the genomic sequence upstream of the transcription initiation site of GnRH, Gpr54 and the GnRH promoter activators under study, were analyzed using ALGEN PROMO 3.0 software (http://alggen.lsi.upc.es/cgi-bin/promo_v3/promo/promoinit.cgi?dirDB=TF_8.3) and jaspar database (http://jaspar.genereg.net/).

Statistics

All analyses were performed using Prism 5 (GraphPad Software) and assessed for normality (Shapiro-Wilk test) and variance, when appropriate. Sample sizes were chosen according to standard practice in the field. Data were compared using an unpaired two-tailed Student's t test, a one-way ANOVA for multiple comparisons or a two-way repeated-measures ANOVA. A Tukey's *post hoc* test was performed when appropriate. The significance level was set at p < 0.05. Data are indicated as means \pm SEM. The number of biologically independent experiments, p values and degrees of freedom are indicated either in the main text or in the figure legends.

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The supplemental information includes 11 figures and 3 tables.

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Figures

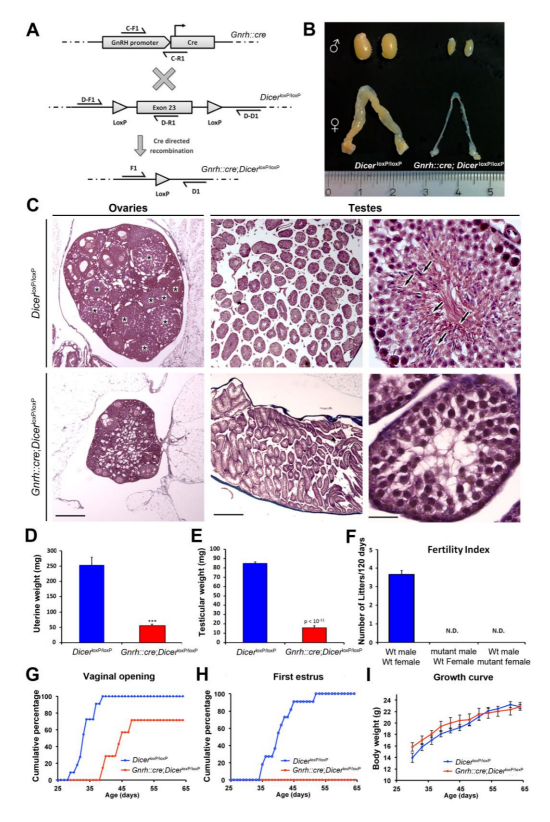


Figure 1. Lack of miRNA biogenesis in GnRH neurons leads to hypogonadotropic hypogonadism and infertility. (A) Genetic strategy to invalidate *Dicer* expression specifically in GnRH-expressing cells in mice. (B) Smaller testes and ovaries and thread-like

uteri in mutant mice. (**C**) Hematoxylin-eosin-stained ovarian and testicular sections of mutant *Gnrh*::Cre; *Dicer*^{loxP/loxP} mice and their wild-type *Dicer*^{loxP/loxP} littermates. Asterisks show corpora lutea and arrows, spermatozoa. Scale bars: 500 μm (left and middle panels) and 30 μm (right panels). (**D** to **F**) Both male and female *Gnrh*::Cre; *Dicer*^{loxP/loxP} mice exhibit hypogonadism and are infertile. N.D.: not detected. (***: p< 0.001, ****: p<0.0001). Values shown are mean ± SEM. (**G** to **J**) While some *Gnrh*::Cre; *Dicer*^{loxP/loxP} female mice undergo vaginal opening during postnatal development (**G**), they never reach puberty (**H**) despite normal somatic growth (**I**).

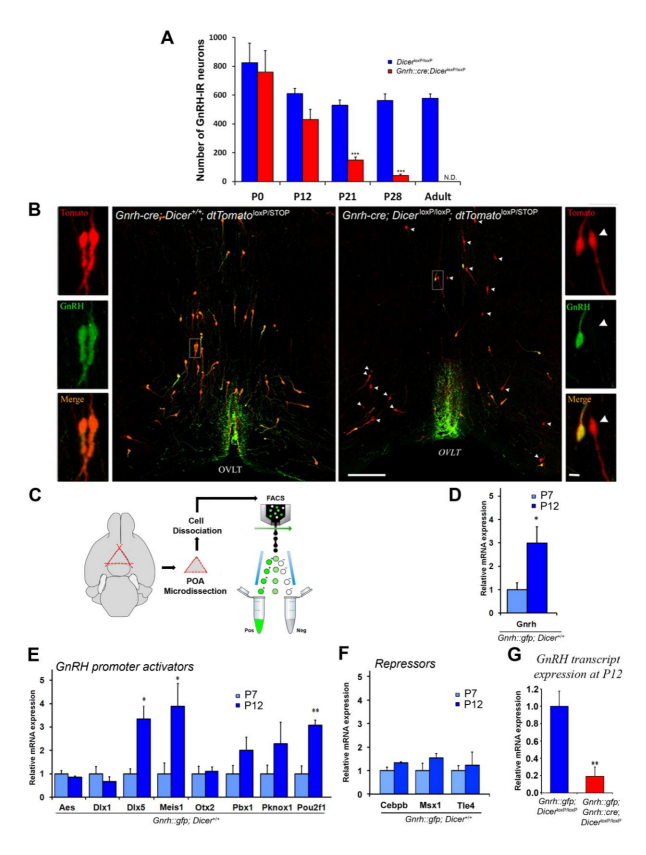


Figure 2. A switch in the control of GnRH gene expression is operated by miRNAs during postnatal developmental. (A) Gradual disappearance of GnRH-immunoreactive (GnRH-IR) neurons in *Gnrh*::Cre; *Dicer*^{loxP/loxP} during postnatal development. (***: p<0.001).

P0, postnatal day 0; N.D., not detected. (**B**) Neuronal lineage tracing shows normal distribution of GnRH neurons in P21 juvenile mice with GnRH deficiency (Gnrh::Cre; $Dicer^{JoxP/IoxP}$; $tdTomato^{JoxP/STOP}$), when compared to control littermates ($Dicer^{JoxP/IoxP}$; $tdTomato^{JoxP/STOP}$). Arrowheads show GnRH-immunonegative Tomato-positive neurons. OVLT, organum vasculosum of the lamina terminalis. Scale bars, 200 μ m; 10 μ m (inset). (**C**) GnRH-GFP neuron isolation by FACS from the preoptic region of Gnrh::Gfp mice. (**D-H**) Real-time PCR analysis of expression levels of GnRH mRNA (**D**, **G**), GnRH promoter activators (DIx5, $t_{(5)}$ =-3.62, p=0.015; Meis1, $t_{(5)}$ =-3.67, p=0.014; Pou211, $t_{(5)}$ =-6.14, p=0.0017; n=3-4 per group) (**E**) and repressors (**F**) in FACS-sorted GnRH-GFP neurons. ((*: p<0.05; **: p<0.01). Values are expressed relative to P7 or wild-type values, as appropriate, set at 1. Values shown are mean ± SEM.

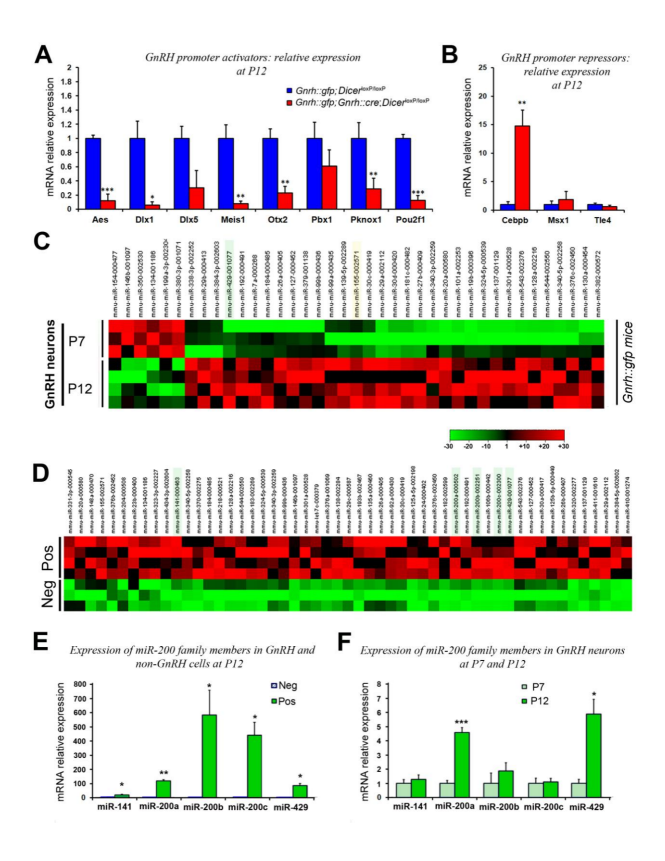


Figure 3. MiRNAs regulate the expression of GnRH promoter modulators in infantile GnRH neurons. (A and B) RT-PCR analysis of the expression of GnRH promoter activators (A) and repressors (B) in FACS-sorted GnRH-GFP neurons from mice selectively lacking

Dicer in GnRH neurons (red bars) or wild-type for Dicer (blue bars) (all values are expressed relative to wild-type values, set at 1). Aes, $t_{(5)}$ =9.58, p=0.0002; Dlx1, $t_{(5)}$ =3.78, p=0.032; Meis1, $t_{(3.17)}$ =4.71, p=0.018; Otx2, $t_{(5)}$ =4.34, p=0.0074; Pou2f1, $t_{(5)}$ =10, p=0.0002; n=3-4 per group. Pknox1, $t_{(6)}$ =2.63, p=0.039; Cebpb, $t_{(3.17)}$ =-4.79, p=0.0173, n=4 per group (*: p<0.05; **: p<0.01; ***: p<0.001). (C-D) miRNAs whose expression changes between P7 and P12 (C) and that are specifically enriched in GnRH neurons at P12 (D). Pos: GFP-expressing neurons; Neg: GFP-negative cells. (E and F) RT-PCR analysis of the expression of miR-200 family members in FACS-sorted cells from preoptic explants isolated from Gnrh::Gfp mice wild-type for Dicer either at P12 (E; Pos vs. Neg: miR-141, $t_{(5)}$ =3.370, p=0.0199; miR-200a, $t_{(4)}$ =13.57, p=0.0002; miR-200b, $t_{(5)}$ =2.775, p=0.0391; miR-200c, $t_{(5)}$ =4.016, p=0.0102; miR-429, $t_{(5)}$ =4.778, p=0.005; n=3-4 per group) or at P7 and P12 (F; P7 vs. P12: miR-141, $t_{(5)}$ =0.6652, p=0.5354; miR-200a, $t_{(4)}$ =9.178, p=0.0008; miR-200b, $t_{(5)}$ =0.9572, p=0.3824; miR-200c, $t_{(5)}$ =0.2604, p=0.805; miR-429, $t_{(5)}$ =3.931, p=0.0111; n= 3-4 per group). Values shown are mean ± SEM.

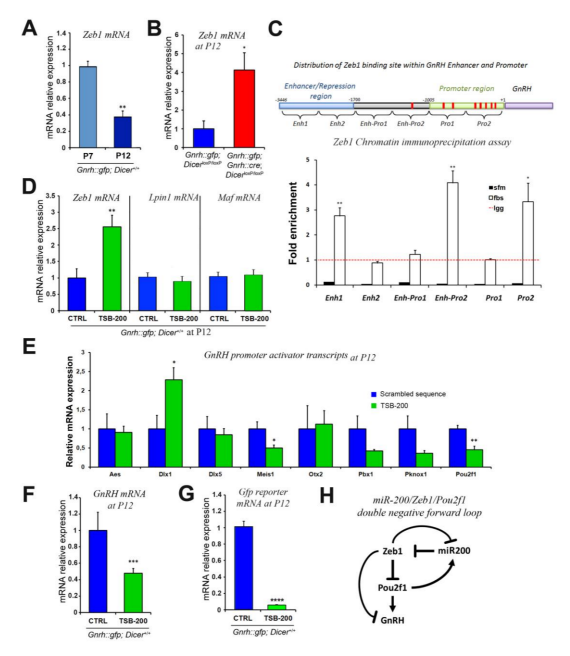


Figure 4. Knocking down the ability of miR-200 to repress *Zeb1* expression in the brain alters infantile GnRH promoter activity and *Gnrh* mRNA expression. (A and B) RT-PCR analysis of the expression of *Zeb1* in GnRH-GFP neurons isolated from *Gnrh::*Gfp; *Dicer*^{+/+} mice at P7 and P12 (A) or from mice in which *Dicer* expression is invalidated or not in GnRH neurons (B). (C) Diagram showing the distribution of putative *Zeb1* binding sites on the *Gnrh* gene (upper panel) and their validation using a *Zeb1* chromatin immunoprecipitation assay in an immortalized mouse cell line secreting GnRH and cultured in the presence or absence of fetal bovine serum (fbs). Values are expressed relative to the immunoprecipitation of chromatin containing the GnRH promoter region with irrelevant IgG species, arbitrarily set at

1 (red dotted line). Serum-free medium (sfm) vs. fbs: Enh1, $t_{(6)}$ =4.756, p=0.0031; Enh2, $t_{(6)}$ =2.320, p=0.0595; Enh-Pro1, $t_{(6)}$ =0.9063, p=0.3997; Enh-Pro2, $t_{(6)}$ =5.462, p=0.0016; Pro1, $t_{(6)}$ =0.2104, p=0.8403; Pro2, $t_{(6)}$ =2.467, p=0.0486, n=4 per group. (**D-G**) Effect of the i.c.v. injection of TSB-200 (green bar) or a scrambled sequence (CTRL, blue bar) on the expression of Zeb1, Expression = Express

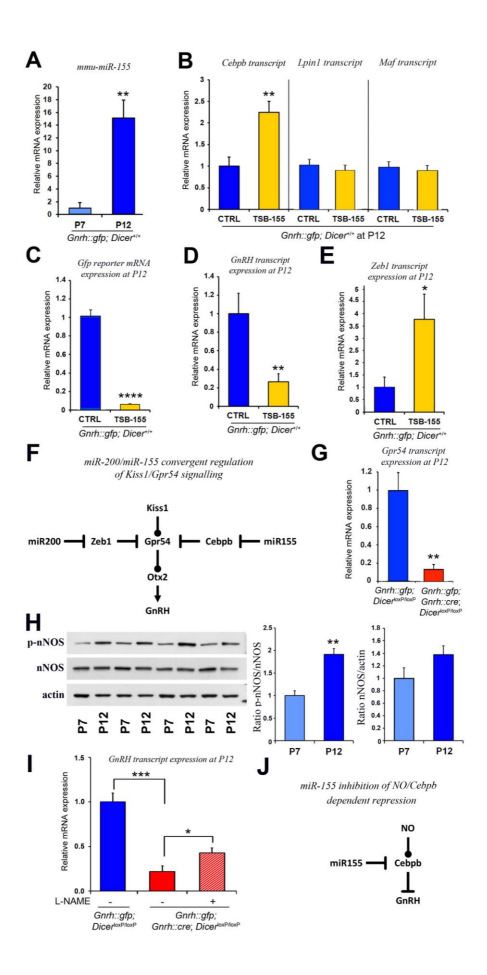


Figure 5. miR-155 modulates Cebpb, Gnrh and Zeb1 mRNA expression in infantile GnRH neurons. (A) RT-PCR analysis of the expression of miR-155 in FACS-isolated GnRH-GFP neurons between P7 and P12 (**: p<0.01). (B-E) Effect of the i.c.v. injection of TSB-155 (yellow bar) or a scrambled sequence (Ctrl, blue bar) on the expression of Cebpb, Lpin1 and Maf (B), GnRH promoter activity (C), and GnRH (D) and Zeb1 (E) transcripts in GnRH-GFP neurons of infantile mice. (F) Putative miRNA-gene network illustrating the convergent contributions of miR200 and miR155, and their respective target genes Zeb1 and Cebpb, in the infantile control of GnRH expression via the repression of the kisspeptin/Gpr54/Otx2signaling pathway (see also Fig. S7A and S9). (G) RT-PCR analysis of expression levels of Gpr54 mRNA in FACS-sorted GnRH-GFP neurons (**p<0.01). (H) Western blot (upper panel) and quantitative (bottom panel) comparison of phosphorylated and total nNOS protein expression in the preoptic region of wild-type mice between P7 and P12. Actin was used as a loading control. Values are expressed relative to P7, set at 1. (I) L-NAME treatment partially rescues GnRH mRNA expression in P12 mice harboring a Dicer deficiency in GnRH neurons. (J) Schematic representation of a miRNA-gene network potentially regulating GnRH expression in the infantile period, and its crosstalk with nitrergic neurotransmission. (*: p<0.05; ****: p<0.0001, $t_{(10)}$ =6.66, n=6 per group). Values are expressed relative to untreated wild-type values, set at 1. Values shown are mean \pm SEM.

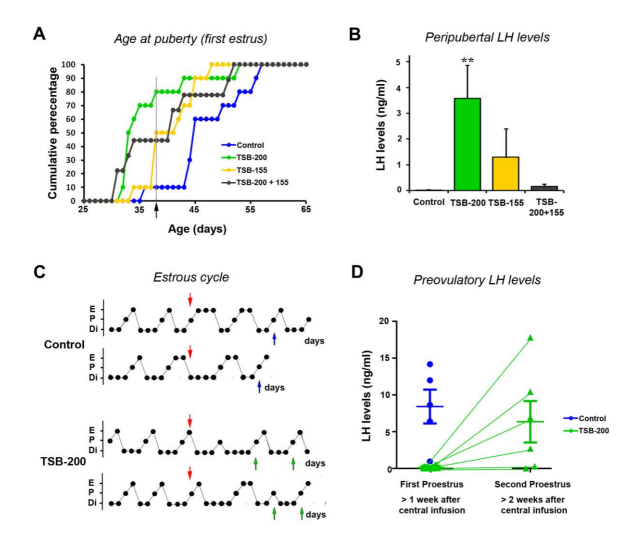


Figure 6. miR-200 binding to *Zeb1* modulates the function of GnRH neurons both in infantile and adult mice. (A) Knocking down the ability of miR-200 and/or miR-155 to repress *Zeb1* and *Cebpb* expression, respectively, during the infantile period markedly accelerates puberty. (B) Circulating levels of LH in P38 peripubertal mice (arrow in A), in which TSB-200, TSB-155, TSB-200+155 or scrambled control sequences were infused into the brain during the infantile period (**: p<0.01). (C) Estrous cyclicity before and after the infusion of TSB-200 into the preoptic region of adult mice (red arrows). Upward arrows show the time of blood collection in control- (blue) and TSB-200-treated (green) mice. (D) LH levels in blood samples collected during proestrus as indicated in C. Values shown are means ± SEM.

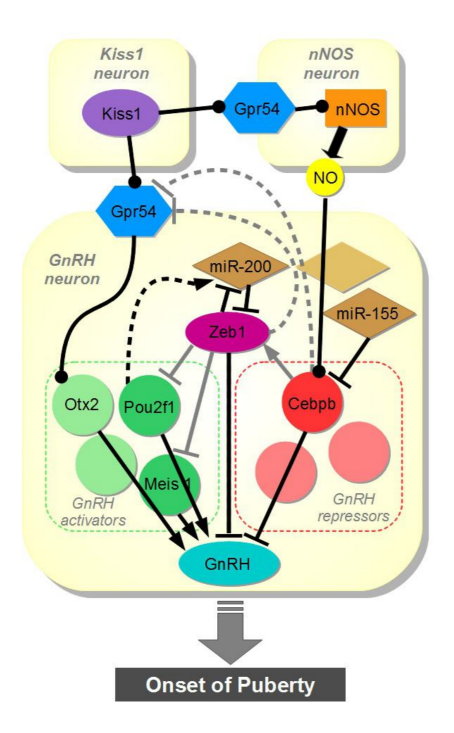


Figure 7. Neuronal microcircuit and miRNA-gene network in hypothalamic GnRH neurons regulating the onset of puberty. Schematic diagram illustrating the potential contribution of the miR-155 and miR-200 families and their target genes to the increase in

GnRH gene expression during the infantile period of postnatal development and how these events may be intertwined with the integration of postmigratory GnRH neurons into the neural network responsible for regulating the timely onset of puberty (solid lines: demonstrated/published; grey lines: ChipSeq data; dotted lines: predictive *in silico* analyses).