Rhythmic expression of circadian clock genes in the preovulatory ovarian follicles of the laying hen

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Abstract

The circadian clock is reported to play a role in the ovaries in a variety of vertebrate species, including the domestic hen. However, the ovary is an organ that changes daily, and the laying hen maintains a strict follicular hierarchy. The aim of this study was to examine the spatial-temporal expression of several known canonical clock genes in the granulosa and theca layers of six hierarchy follicles. We demonstrated that the granulosa cells (GCs) of the F1-F3 follicles harbored intrinsic oscillatory mechanisms in vivo. In addition, cultured granulosa cells (GCs) from F1 follicles exposed to luteinizing hormone (LH) synchronization displayed Per2 mRNA oscillations, whereas, the less mature GCs (F5 plus F6) displayed no circadian change in Per2 mRNA levels. Cultures containing follicle-stimulating hormone (FSH) combined with LH expressed levels of Per2 mRNA that were 2.5-fold higher than those in cultures with LH or FSH alone. These results show that there is spatial specificity in the localization of clock cells in hen preovulatory follicles. In addition, our results support the hypothesis that gonadotropins provide a cue for the development of the functional cellular clock in immature GCs.

Introduction

The circadian clock is a cell-autonomous system that coordinates physiology and metabolism to the correct time of the day [1, 2]. The endogenous timekeeper is based on intracellular transcriptional-translational feedback loops in which a few canonical clock gene products and genetic elements drive the rhythmic expression of downstream genes, thereby contributing to rhythmic physiology [3–5]. The circadian system is important for successful reproduction in vertebrates because it influences the follicle maturation and ovulation [6].

The avian central circadian organization consists of three separate oscillators located in the pineal gland, the retinae and a hypothalamic region, that is possibly equivalent to the mammalian SCN [7–9]. Prior studies have supported the intimate relationship between the circadian timing system and the hypothalamic-pituitary-gonadal (HPG) axis. Briefly, a signal from the
central clock is crucial for the initiation of the luteinizing hormone (LH) surge and subsequently, for ovulation. Recently, the ovarian circadian clock has been well documented in many mammalian and non-mammalian species, and its function has been investigated both in vivo and in vitro. In most known cases, the function of the ovarian clock is related to the timing of gene expression in mature granulosa cells (GCs), including genes related to steroidogenesis, gonadotropin responsiveness and ovulation [10].

Clock gene rhythms are possibly under the control of gonadotropins (follicle-stimulating hormone (FSH) and LH), acting to synchronize of follicular cell activities [11–14]. On the other hand, several candidate genes associated with the response of GCs to gonadotropins are known as clock-controlled genes (CCGs), such as the LH receptor (Lhcgr), steroidogenic enzymes (StAR), prostaglandin synthase (ptgs2) and gap junctions [15–17]. Obviously, there is a correlation and interaction between the presence of a functional clockwork and gonadotropin-induced follicular development. However, it is difficult to determine whether the activation of the ovarian clock is a cause or a result of follicle development and maturation.

The ovary is an ever-changing, multi-compartmental organ. In chickens, a strict follicular hierarchy is maintained, with 2 to approximately 6 hierarchical follicles (F1-F6) and a single follicle ovulated per day [12, 18–20]. In quails, Nakao found that the level of StAR mRNA exhibited a diurnal rhythm coincident with Per2 expression only in the largest follicle [21]. Therefore, gonadotropins and their associated cell signaling mechanisms combined with the ovarian clock might orchestrate a functional hierarchy of preovulatory follicles. To further understand the role of the ovarian clock during the maturation of follicles, a thorough analysis of clock gene expression in preovulatory follicles is needed to determine whether the rhythm is truly circadian and to reveal the detailed expression patterns in the specific components of different follicles [22].

Birds represent an excellent model to study the role of biological clocks in female reproduction for two reasons: ① The daily rhythms of ovulation-oviposition in adult female birds have been well described and are easy to monitor [23, 24]. ② The morphological characteristics and functional anatomical components of preovulatory follicles are well known, and the stage of follicular development is defined by the follicle size. Accordingly, the present study explored differences in spatial-temporal relationships among clock genes in GCs and theca cells (TCs). Moreover, on the basis of the present results, we examined the effects of gonadotropins on Per2 mRNA expression, providing preliminary support for the hypothesis that FSH provides a cue for the development of the functional cellular clock in immature GCs, whereas LH synchronizes the cellular clock of mature GCs.

Materials and methods

Ethics statement

All animal experiments were approved by The Committee on Experimental Animal Management of Sichuan Agriculture University, and carried out in strict accordance with the Regulations for the Administration of Affairs Concerning Experimental Animals of the State Council of the People’s Republic of China. The chickens involved in this study were sacrificed with as little pain as possible to reduce their suffering.

Animals and Zeitgeber Time (ZT)

Twenty-week-old Roman laying hens were housed individually in an experimental farm for poultry breeding at the Sichuan Agricultural University (Ya’an, China) and provided feed and water ad libitum. The hens were exposed to cycles of 16 h of light (L) and 8 h of darkness (D), with the light period beginning at 06:00 h local time. ZT0 was defined as the time at which the
lights came on. As described previously [25], adult birds (20–30 weeks of age) with regular clutches of at least 30 eggs were used for these experiments. Individuals or groups of chickens that were collected at specific times of the circadian day were sacrificed for RNA extraction. This procedure made it impossible to analyze the molecular rhythms within individuals over successive time points. Thus, we monitored individual daily laying patterns by assessing the oviposition time of each bird at 30-min intervals from ZT0 to ZT16. After 4 weeks of ovulatory cycle observations, 24 birds that had consistent laying sequences and oviposition times (between ZT2 and ZT2.5) were selected from among the 500 individuals for further experiments.

**Tissue collection and GC culture**

Selected hens were anesthetized with pentobarbital and sacrificed by decapitation at 2, 6, 10, 14, 18, and 22 hours after oviposition (n = 3 per time point). The experimental procedures were approved by the Animal Care Committee at our institute. After sacrifice, the ovary from each hen was immediately removed and placed into an ice-cold 0.9% NaCl solution. TCs and GCs were collected from large preovulatory follicles (F1-F6) according to previously described methods [26, 27]. Tissues were immediately snap frozen in liquid nitrogen and stored at -80˚C until RNA extraction.

For culture, hens were sacrificed 2 h after oviposition. Follicles were arranged according to size. GCs from F1 follicles and F5 plus F6 follicles (F5/F6) were studied separately. The isolated sheets were washed in pre-warmed Hanks’ balanced salt solution to remove the adherent yolk, cut into pieces and dissociated in Type IV collagenase (1.5 mg/mL, Sigma, St Louis, MO, USA) containing M199 medium (Gibco BRL Co.Ltd., USA). The viable dispersed cells were enumerated using the trypsin blue exclusion method and suspended in M199 medium containing 5% fetal bovine serum (FBS) and a 1% penicillin-streptomycin mixture. Cells were cultured at 37˚C in a humidified atmosphere of 95% air and 5% CO₂. Approximately 0.5 × 10⁶ GCs were seeded onto 12-well plates (Corning, NY, USA). After 24 h of incubation with a medium change at 6 h, the non-attached cells were removed by aspiration, and the adherent cells were washed three times with serum-free medium and used in all further incubations. Ovine LH (100 ng/mL), and recombinant human FSH (100 ng/mL) were dissolved in culture media.

**RNA extraction, reverse transcription and Real-Time PCR analysis**

Total RNA was extracted from the tissues using TRIzol reagent (Invitrogen, Carlsbad, CA, USA) according to the manufacturer’s instructions. For reverse transcription, cDNA was synthesized from 1 μg of total RNA with the PrimeScript® RT Master Mix (Perfect Real Time) (TaKaRa, Dalian, China) following the manufacturer’s protocol. All qPCR assays were performed on a Bio-Rad CFX96 Touch system (Bio-Rad, Hercules, CA, USA). Each reaction volume was 15 μl, containing 1 μl of cDNA, 0.5 μl of the reverse and forward primers (10 μM) for each gene, 5.5 μl of double-distilled H₂O, and 7.5 μl of the SsoFast Eva Green Supermix (Bio-Rad, Hercules, CA, USA). Triplette no-template controls containing DEPC-treated water were included in each run. All of the primers employed in this study were published previously [25]. Primer sequence details are provided in Supplementary S1 Table. Relative expression was determined using the 2^-ΔΔCT method. The quantity in each sample was measured relative to β-actin.

**Statistical analysis**

Data are presented as the mean ± SEM of at least three separate experiments. To detect the rhythmically expressed genes, we used 2 different algorithms. Cosinor analysis adjusts the data
to a cosinusoidal function and provides an objective test of whether the amplitude of the cosi-
nor curve differs from zero [28]. For this purpose, we fitted periodic sinusoidal functions to
the relative values for the expression of each gene across six time points (over a 24-h period)
using the formula \((t) = M + A \cos(2\pi t/24 + \sigma)\), where \((t)\) is the level of gene expression across
time, the mesor \(M\) is the middle value of the fitted cosine representing a rhythm-adjusted
mean, \(A\) is the amplitude of the oscillation in expression, \(t\) is the time in hours, and \(\sigma\) is the
acrophase (the peak time of the fitted cosine function). In addition to the performing the cosi-
nor analysis, we analyzed the data using CircWave software (R.A.Hut, Groningen, NL) [29].
CircWave uses a linear harmonic regression fit that describes the data by adding harmonics
to the principal wave function. To determine the number of harmonics to add, F-testing
was used for the primary fit and for each added harmonic, with a significance level of 0.001
adopted to reduce the chance of false positives. One-way ANOVA and Student’s \(t\)-test were
also performed, using SigmaPlot software (Ver. 12.0; Systat Software, San Jose CA) and Graph-
Pad Prism. Differences were considered significant at \(P < 0.05\).

Results

Circadian clock gene expression in preovulatory follicles in vivo

The mRNA of seven clock genes (\(Bmal1\), \(Bmal2\), \(Clock\), \(Per2\), \(Per3\), \(Cry1\) and \(Cry2\)) was
expressed at all time points in the preovulatory follicles of laying hens maintained under a 16:8
L: D schedule. The analysis of mRNA expression revealed that most of the genes showed diur-
nal changes, but not all of the clock gene transcripts were expressed rhythmically. The expres-
sion and rhythm characteristics of all clock genes in GCs and TCs are summarized in Table 1.
As shown in the table, clock genes that showed pronounced circadian rhythms in expression
were more common in GCs than in TCs, and the rhythm-adjusted 24-h means of the expres-
sion levels of most clock genes were higher in GCs than in TCs.

The normalized daily \(Bmal1\), \(Clock\), \(Per2\) and \(Cry2\) mRNA expression profiles are conven-
tionally used as circadian phase markers and are presented in Fig 1. The phase markers exhib-
itied rhythmic expression with the expected phase alignments in the GCs of the F1-F3 follicles.
For example, the \(Bmal1\) (positive element) and \(Per2\) (negative element) mRNA levels cycled in
antiphase (Fig 1). In contrast, the statistical analysis revealed that no or few clock genes showed
significant circadian variation in the TCs, and none of the expected phase alignments were
observed in any of the examined follicle TCs (S1 Fig).

Analysis of clock gene expression in cultured GCs

Preovulatory follicle GCs were divided into two groups (F1 and F5/F6). Expression levels of
the \(Lhcgr\) and \(STAR\) genes, as markers of GCs maturation, were significantly higher in F1 GCs
than in F5/F6 GCs (Fig 2A). When the cells were cultured in a serum-free medium for 2 days,
no or very weak oscillating mRNA expression was detected in all of the cultured cells. After
LH treatment, circadian oscillation of \(Per2\) expression was clearly observed in F1 GCs (Cir-
cWave, \(P < 0.001\)), whereas only time-dependent variations in mRNA levels were observed in
cultured F5/F6 GCs (Fig 2B).

Alterations of \(Lhcgr\) and \(Per2\) expression upon LH and FSH stimulation

We investigated the expression of \(Lhcgr\) and \(Per2\) in cultured F1 and F5/F6 GCs. In the
F5/F6 group, the levels of \(Lhcgr\) mRNA were increased after treatment with FSH for 1 h
(1.6 ± 0.13-fold compared with the T1 control) and remained elevated and further increased
after 16 h of culture (5.5 ± 0.7-fold T16 control), whereas a significant increase in response to
Table 1. Circadian rhythm characteristics of clock genes in GCs and the TCs.

|    | Mesor | Amplitude | Acrophase(hh:mm) | P-value | Mesor | Amplitude | Acrophase(hh:mm) | P-value |
|----|-------|-----------|------------------|---------|-------|-----------|------------------|---------|
| F3 |       |           |                  |         |       |           |                  |         |
| Bmal1 | 4.85  | 1.41     | 13:08            | <0.05  | 0.32  | 0.07     | 12:10            | >0.05   |
| Bmal2 | 0.18  | 0.15     | 22:06            | >0.05  | 0.77  | 0.29     | 03:49            | <0.05   |
| Clock | 1.38  | 1.10     | 18:14            | <0.01  | 0.76  | 0.25     | 08:36            | >0.05   |
| Per2 | 0.01  | 0.06     | 10:25            | >0.05  | 0.02  | 0.01     | 01:18            | <0.05   |
| Cry1  | 0.70  | 0.18     | 06:24            | >0.05  | 0.17  | 0.03     | 01:48            | >0.05   |
| Cry2  | 2.38  | 1.09     | 06:40            | <0.01  | 0.58  | 0.09     | 03:40            | >0.05   |
| F4   |       |           |                  |         |       |           |                  |         |
| Bmal1 | 5.26  | 1.76     | 16:06            | >0.05  | 0.41  | 0.05     | 09:15            | <0.05   |
| Bmal2 | 0.94  | 1.64     | 18:41            | <0.05  | 1.05  | 0.26     | 03:46            | >0.05   |
| Clock | 1.14  | 1.45     | 06:05            | <0.05  | 0.81  | 0.11     | 06:53            | <0.05   |
| Per2  | 1.75  | 0.81     | 11:33            | <0.01  | 0.61  | 0.40     | 09:27            | <0.01   |
| Per3  | 0.14  | 0.05     | 13:05            | >0.05  | 0.05  | 0.05     | 13:27            | <0.01   |
| Cry1  | 1.01  | 0.73     | 15:36            | <0.05  | 0.22  | 0.07     | 17:45            | <0.05   |
| Cry2  | 3.91  | 2.24     | 18:42            | <0.01  | 0.71  | 0.04     | 17:13            | >0.05   |
| F5   |       |           |                  |         |       |           |                  |         |
| Bmal1 | 6.06  | 0.55     | 23:10            | >0.05  | 0.46  | 0.06     | 09:26            | >0.05   |
| Bmal2 | 0.42  | 0.11     | 19:24            | <0.05  | 1.01  | 0.16     | 05:02            | >0.05   |
| Clock | 0.67  | 0.06     | 06:05            | >0.05  | 0.72  | 0.06     | 19:28            | >0.05   |
| Per2  | 2.74  | 0.76     | 06:35            | <0.05  | 0.35  | 0.19     | 11:07            | <0.01   |
| Per3  | 0.18  | 0.02     | 16:39            | >0.05  | 0.02  | 0.00     | 13:16            | <0.05   |
| Cry1  | 1.37  | 0.53     | 16:42            | >0.05  | 0.18  | 0.02     | 17:55            | <0.05   |
| Cry2  | 5.14  | 0.59     | 15:16            | >0.05  | 0.63  | 0.01     | 00:30            | >0.05   |
| F6   |       |           |                  |         |       |           |                  |         |
| Bmal1 | 2.44  | 1.10     | 19:41            | <0.01  | 0.83  | 0.20     | 03:57            | <0.01   |
| Bmal2 | 0.62  | 0.23     | 14:08            | <0.05  | 1.33  | 0.27     | 03:31            | >0.05   |
| Clock | 0.72  | 0.50     | 18:32            | <0.001 | 1.37  | 0.14     | 05:40            | >0.05   |
| Per2  | 6.12  | 1.90     | 16:53            | <0.01  | 0.84  | 0.40     | 06:44            | <0.001  |
| Per3  | 0.42  | 0.12     | 16:37            | >0.05  | 0.06  | 0.02     | 10:00            | >0.05   |
| Cry1  | 0.37  | 0.09     | 17:21            | >0.05  | 0.37  | 0.07     | 13:24            | <0.05   |

(Continued)
FSH treatment was not observed in the F1 follicle GCs. However, the levels of Lhcgr mRNA were markedly lower in F5/F6 follicles GCs than in F1 follicles regardless of FSH-treat (Fig 3A). As shown in Fig 3B, time-dependent variation in the mRNA levels of Per2 was observed in the F1 GCs but not F5/F6 GCs. Both FSH and LH stimulation significantly increased the levels of Per2 mRNA in F1 GCs compared with the no-treatment controls (P < 0.05); however, FSH treatment did not affect the levels of Per2 mRNA expression in F5/F6 GCs (P > 0.05). Treatment with FSH plus LH induced a significant increase in Per2 expression in the cultured cells from both the F5/F6 and F1 groups compared with the controls (3.6 ± 0.28-fold and 5.2 ± 0.78-folds, respectively).

Discussion

Ovarian preovulatory follicles also known as hierarchical follicles (F1-F6) are selected from the small yellow follicles (SYF) and are destined to ovulate; the F1 follicle will be the next to ovulate. Oviposition is followed 15–45 min later by ovulation, after which each follicle moves up one position in the hierarchy. The GCs of different follicles are at different levels of differentiation, and steroidogenic ability differs among the six preovulatory follicles.

In the present study, we showed the cyclical gene expression patterns of several canonical clock genes in ovarian preovulatory follicles during the ovulation-oviposition cycle (~24 h) of commercial hens under farming light-dark conditions. Typically, the expression of a positive element (Bmal1 and Clock) in an antiphase relationship to the negative elements (Pers and Crys) in any given tissue is considered evidence of a functional clock [30]. In a previous study, Nakao et al. [21] found a robust change over 24 h in Per2 and Per3 gene expression only in mature F1 follicles based on the Fisher’s least significant difference (LSD) post hoc test. In our study, the data were analyzed using CircWave 1.4 software [29] to test the daily rhythmicity of gene expression. We found that several of the clock genes exhibited rhythmic expression (P < 0.001), with the expected phase alignments in the GCs of the F1, F2 and F3 follicles (Fig 1). Although several clock genes also appeared to change over 24 h in other tissues (e.g., in the GCs of F6 follicles), no significant rhythmic pattern or anticipated antiphase relationship was observed (Fig 1). Our results provide a thorough and precise analysis of clock gene expression in preovulatory follicles and reveal tissue-specific variations in phase and amplitude. These findings are in agreement with those of previous studies suggesting that the specificity of clock gene oscillations may be fine-tuned to functional compartmentalization of the preovulatory follicles.

On the basis of our present understanding of the spatial specificity in the localization of clock cells in the hierarchal follicles, we cultured GCs from F1 and F5/F6 follicles synchronized with LH. LH is a potent synchronization factor in vitro as described in previous studies [16, 31–33]. Our results showed that Per2 oscillations were generated by LH synchronization only in GCs from F1 follicles; the clock genes of less mature GCs prepared from F5/F6 follicles displayed no circadian pattern. The present findings are consistent with previous reports of much more active cellular clock in matured GCs than in less mature GCs [16, 21, 32]. LH plays

| Table 1. (Continued) |
|-----------------------|
| **Cry2** | Mesor | Amplitude | Acrophase(hh:mm) | P-value | Mesor | Amplitude | Acrophase(hh:mm) | P-value |
|-----------------------|
| 6.84 | 3.05 | 16:42 | <0.001 | 1.17 | 0.09 | 02:03 | >0.05 |

The acrophases and amplitudes were calculated according to the non-linear regression fit of a cosine function. CircWave was used to detect a significant rhythm, and P < 0.001 was considered statistically significant.

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Fig 1. Daily expression patterns of selected clock genes in GCs of various preovulatory follicles under 16L:8D. Each data point represents the mRNA amount of the corresponding clock gene normalized to β-actin expressed as the mean ± SEM (n = 3). The white areas of the bars in the bottom of the figure indicate the light period, and the black areas indicate the dark period. ZT24 values are a duplicate of ZT0 shown for clarity. Genes were identified as rhythmically expressed in unison by CircWave, P < 0.001 was considered to indicate significant rhythmic expression (solid lines). The dotted lines indicate no rhythmic expression.

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various roles through combination with its receptor (Lhcgr). Lhcgr mRNA is first present in the granulosa layer of the smallest preovulatory follicles and increases with follicular maturation [34, 35]. Thus, it is possible that the elevation of Per2 oscillations in F1 GCs may be due to the increased transcription of Lhcgr. In this study, the Lhcgr mRNA level was statistically significantly higher in cultured GCs from F1 follicles than in those from F5/F6 follicles (Fig 2A). LH-stimulated Per2 gene expression may be dependent on high levels of Lhcgr expression.

FSH treatment upregulated the expression of Lhcgr mRNA in GCs from F5/F6 follicles (Fig 3A). Of the core clock genes, Pers (Per1-Per3) are the most responsive to various inducers in vivo or in vitro. However, FSH treatment had no effect on the expression of Per2 in F5/F6 GCs. In contrast, a slight but significant increase in Per2 mRNA levels was observed after LH treatment. Interestingly, FSH plus LH treatment was most effective in increasing the expression of Per2 mRNA (Fig 3B). We failed to observe oscillatory expression of Per2 in FSH-treated GCs by LH synchronization.

Based on work performed by the present authors and others, one hypothesis is that FSH can induce the sustained increase of Lhcgr, which is accompanied with follicular development, and alter the responsiveness of follicles to LH as a result. The high-responsiveness follicles may be synchronized by the pulsatile LH surge, leading to a gradual activation of the functional clock in follicles. Furthermore, the follicular clock system maximizes the expression of LH receptors preceding the arrival of the LH surge. Our data, consistent with several previous reports, showed that there was spatial specificity in the localization of clock cells in the
preovulatory follicles. Our results support the hypothesis that FSH provides a cue for the development of the functional cellular clock in immature GCs, and that LH synchronizes the cellular clock of mature GCs. This work contributes to our understanding of the physiological significance of rhythmic clock gene expression in ovarian follicles.

Supporting information

S1 Fig. Daily expression patterns of selected clock genes in TCs of various preovulatory follicles under 16L:8D. Each data point represents the mRNA amount of the corresponding clock gene normalized to β-actin expressed as the mean ± SEM (n = 3). The white areas of the bars in the bottom of the figure indicate the light period, and the black areas indicate the dark period. ZT24 values are a duplicate of ZT0 shown for clarity. Genes were identified as rhythmically expressed in unison by CircWave, \( P < 0.001 \) was considered to indicate significant rhythmic expression (solid lines). The dotted lines indicate no rhythmic expression.

(TIF)

S1 Table. Real-time PCR primer sequences for the clock genes.

(DOCX)

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Author Contributions

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