Effects of Adult or Perinatal Hormonal Environment on Ultradian Rhythms in Locomotor Activity of Laboratory LEW/Ztm Rats

FRANZISKA WOLLNIK
Central Animal Laboratory at the Medical School Hannover, D-3000 Hannover 61, F.R.G.

AND

KLAUS-DIETER DÖHLER
Department of Endocrinology at the Medical School Hannover, D-3000 Hannover 61, F.R.G.

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WOLLNIK, F. AND K.-D. DÖHLER. Effects of adult or perinatal hormonal environment on ultradian rhythms in locomotor activity of laboratory LEW/Ztm rats. PHYSIOL BEHAV 38(2) 229-240, 1986.—Four experiments were performed with male and female rats of the inbred strain LEW/Ztm maintained under a light-dark schedule of 12:12 hours. The animals were subject to castration (GOX) or ovariectomy (OVX), estradiol 17ß-implantation (E2-capsules), and perinatal hormonal treatments with testosterone propionate (TP) and an androgen antagonist (cyproterone acetate, CA). Results indicated a difference in the locomotor activity pattern between the two sexes as a result of the endogenous estradiol levels of the adult animals. The activity pattern of male LEW rats was characterized by ultradian rhythms of 4 and 4.8 hr periods. The female LEW rats, on the other hand, generally exhibited a clear circadian activity pattern and no ultradian activity rhythms. Following ovariectomy, each of the females showed distinct ultradian rhythms. These disappeared after E2-implantation. Castration of adult males had no effect on the ultradian activity pattern. Implantation of E2-capsules resulted in a marked decrease of the ultradian activity components. Perinatal treatment of the males with an androgen antagonist (CA) did not appear to effect ultradian rhythms during adulthood. Females treated perinatally with testosterone showed a significant increase in the ultradian activity components. This effect is assumed to be due to low estrogen levels in these animals during adulthood. Our study supports the assumption that ultradian rhythms are a result of changes in the phase relationships between several circadian oscillators. The synchrony of these oscillations seems to be facilitated by estradiol.

Ultradian rhythms  Locomotor activity  Gonadectomy  Estradiol  Testosterone
Brain differentiation  Laboratory rat

Several well-known phenomena of circadian rhythms in mammals can be explained by the model of the circadian system as a composite of multiple, highly ordered oscillators [40-42, 46] located in or close to the suprachiasmatic nuclei in the hypothalamus [47]. The origin of similar temporal variations on a smaller time scale, called ultradian or short-term rhythms, remains unclear [11]. Ultradian rhythms in locomotor activity are known to exist in laboratory rats [8, 9, 28-30, 49], however, attracting much less attention than circadian rhythms.

A recent study of LEW/Ztm rats has shown that ultradian rhythms of 4 and 4.8 hr periods are genetically fixed in this inbred strain [9]. The persistence of these ultradian rhythms following the disruption of circadian organization under continuous light supported the assumption that the activity rhythms of this strain are caused by an independent ultradian oscillator [9]. Alternatively, the persistence of ultradian rhythms can be explained by changes in the phase relationships between several circadian oscillators [46].

The ultradian rhythms of the inbred strain LEW/Ztm are sex-specific, being observed only in the males [56]. The question, therefore, arises as to whether the ultradian rhythms are the result of endocrine factors. Several studies have documented clear effects of gonadal hormones on the

Requests for reprints should be addressed to Dr. Franziska Wollnik, Northwestern University, Department of Neurobiology and Physiology, Evanston, IL 60201.
circadian system of mammals and birds [51]. Melatonin has been reported to shorten the freerunning period of activity and to induce continuous activity in the house sparrow [52]. Castration in mice resulted in a lengthening of the circadian running-wheel behavior, whereas continuous testosterone treatment reversed the effects of castration [12]. Testosterone, furthermore, modulated the duration of activity and induced “splitting” of the circadian locomotor activity pattern in the starling [24,25].

“Splitting” is defined as rapidly occurring transient dissociation of two oscillations which temporarily free-run with different circadian frequencies. In female hamsters and rats, exogenous estrogen shortened the free-running period of activity in a light dark cycle [1,36]. The phase and level of activity, therefore, appeared to vary with the estrous cycle [2,19]. A recent report [34] using female hamsters also indicated that low estrogen levels induce “splitting.” These various effects of gonadal steroids on the synchronization of the multi-oscillatory circadian system are of great interest lending an explanation to the observed sex-differences in the ultradian activity rhythms of the LEW strain.

Furthermore, it is also known that the brains of male and female rats differ functionally and also in the morphology of the suprachiasmatic nuclei [22,33]. These sex-related differences in brain function are established during an early critical stage of development, apparently by the masculinizing effect of androgens on the developing brain [3,14,21]. Preliminary reports with rats and hamsters indicate that perinatal exposure to androgens also permanently influences the organization of the circadian system [1,13,58].

The purpose of the present study was to examine the effects of perinatal and/or adult hormonal environment on the ultradian activity rhythm of the inbred rat strain LEW/Ztm. The experiments provide no evidence of perinatal hormonal treatment having an effect on the ultradian rhythms. It does seem certain, however, that the sex-related difference is caused by the dissimilar hormonal environments of the adult animals. Further findings indicate estradiol as being important in suppressing the ultradian rhythms in females as well as in males, thereby identifying an additional function of estradiol on the activity rhythms of rats.

GENERAL METHOD

Animals and Maintenance Conditions

The inbred rat strain LEW/Ztm has been found to exhibit genetically fixed [8,9] as well as sex-specific [56] ultradian activity rhythms. In addition to the limitation of ultradian rhythms to the males, it is likewise peculiar that the brief reproductive period of the females ends between 4 and 5 months of age (Döhler, unpublished observations). The strain itself is the result of strong inbreeding systems such as continuous brother-sister matings over at least twenty generations [17]. Individuals of this strain, therefore, provide a convenient material of genetically identical animals.

Both sexes were bred and raised in our laboratory under a light-dark cycle of LD 12:12 and controlled environmental conditions (room temperature 22±1°C, relative humidity 55±5%). The animals were free of all pathogens specified in the GV-SOLAS list [23]. During the course of the measurements, the animals were individually kept in polycarbonate cages (Makrolon Type III, 35×33×20 cm) on sterile wooden granules. A pelleted diet (Altromin 1324) and tap water from Makrolon bottles were available ad lib. The rooms were entered once a week for animal maintenance. The animals

FIG. 1. Analysis of artificial data by the spectral analysis [26,38] and the chi square periodogram [48]. Input data consisting of 1728 5-min values (=6 days) of the following functions were generated by a computer: (a) Cosine function of 25.5 hr period and an amplitude of 50 arbitrary units. A range of random numbers from +50 to −50 was added. (b) Saw tooth function of 24.0 hr period with an increase of the amplitude from 0 to 100 arbitrary units. A range of random numbers from +100 to −100 was added. (c) Cosine functions of PI=24.0 hr, P2=4.8 hr, and P3=4.0 hr period with amplitudes of A1=50, A2=40, and A3=30 arbitrary units. A range of random numbers from +180 to −180 was added. (d) Square wave functions of P1=24.0 hr, P2=4.8 hr, and P3=4.0 hr period with amplitudes of A1=60, A2=50, and A3=20 arbitrary units. A range of random numbers from +150 to −150 was added. (e) Cosine functions of P1=24.0 hr, P2=3.4 hr, and P3=3 hr periods with amplitudes of A1=60, A2=40, and A3=30 arbitrary units. A range of random numbers from +150 to −150 was added. (f) Square wave functions of P1=24.0 hr, P2=3.4 hr, and P3=3 hr periods with amplitudes of A1=60 and A2=30 arbitrary units. A range of random numbers of +150 to −150 was added.
were provided with new cages, food, and water every 3 weeks during long-term recordings.

**Apparatus and Data Analysis**

Locomotor activity was recorded with an electronic movement analyzer; which operating on a capacitant system [44] supplies a continuous signal proportional to the horizontal movement of the animal. The output was recorded at 10-sec intervals by a microcomputer (Apple II).

Five-min mean values were calculated from the 10-sec intervals. All further calculations were based on these 5-min averages. Due to the inconsistency in the body weights of the animals, the intensity of the output signals from the capacitant recording system varied. The measurements of locomotor activity were standardized by setting the overall mean value of each animal at 100%. Event records were printed at 5-min intervals over 48-hr time scale with each day being repeated for continuity (double plot). Analysis of ultradian rhythms in the presence of a circadian rhythm is a difficult problem. We therefore used two different mathematical techniques to examine the periodicity of the locomotor activity data. The results of the two methods were compared in order to check the reliability of each method. Statements about the periodicity of each animal were based on the application of both methods.

The "generalized harmonic spectral analysis" devised originally by Tukey [5] rests upon the mathematical principle that any finite sequence of discrete data taken at regular intervals can be fully and completely characterized as dis-

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**FIG. 2.** Left: Double plotted locomotor activity record of a female LEW rat (No. 4) under LD 12:12. The animal was ovariectomized (OVX) on day 21. Estradiol-17β implantation (E2-capsule) occurred on day 42. Surgery was performed during the light period of the lighting regime. A 3-4 hr interval of total rest is followed by a distinct peak of highly increased activity. Middle: Continuous spectral analysis: 6-day intervals were shifted over the total registration period in one day steps. For each of these 6-day intervals a spectral analysis similar to Fig. 3 was calculated according to Halberg and Panofsky [26,38]. The amplitudes of distinct spectral estimates, i.e., 24-hr (---), 4.8-hr ( ), and 4-hr (•) were appointed to the first day of the interval. Right: Chi square periodogram for distinct time intervals. The sloping line represents $p \leq 0.001$ ($\chi^2$) as derived according to Sokolove and Bushell [48]. For interpretation of the ultradian peaks see the Apparatus and Data Analysis section.
crete values taken at identical intervals from the sum of a
finite series of sine and cosine waves, known as the Fourier
components of the data.

We used a computational method described by Halberg
and Panofsky [26,38] for this type of analysis, whereby the
point of departure for the calculations was an autocorrelation
function of the original data. This method evaluates the
statistical significance of circadian rhythms as well as oscil-
lations having periods longer or shorter than circadian
rhythms. The only limitation is given by the fact that the
maximum meaningful resolution attainable is a spectral esti-
mate for each of T values of frequency (where T is the total
length of the data series). These components will differ from
each other in frequency by steps of I/T. Thus, spectral esti-
mates were attainable for only distinct periods (i.e., 24 hr,
19.2 hr, 16 hr, etc.). The amplitudes S' of distinct spectral
estimates are a measure of the extent to which oscillations of
various period lengths contribute to the total variability of a
time series. The statistical reliability given by the 95% con-
fidence limits could be improved by pooling the spectral esti-
mates of individuals; in this case animals of the same exper-
imental protocol. Differences between the amplitude of the
spectral estimates of a distinct period were tested statisti-
cally by t-tests.

In addition, we used the chi square periodogram to
determine the exact period length of ultradian rhythms. This
method is not bound by any implicit assumption about the
wave form of the dominant rhythmic components. The
calculation procedure developed by Sokolove and Bushell
[48] suggests a slightly different statistic than that origi-
nally recommended by Enright [16]. Here a simple test de-
termining the significance of a periodogram peak was pro-
posed and the sensitivity of the method was examined for
detecting periodicity in the presence of random fluctua-
tions and for resolving periodic components of similar am-
plitude, but with slightly different periods. The periodo-
gram has already been calibrated with a variety of artificial
as well as biological data. A broad description of several
deficiencies of the procedure, particularly when certain
kinds of non-stationarity are present, was given by Enright
[16]. Other calibrations of the periodogram, using several
different types of biological rhythm data, have been under-
taken [7,45] as well as comparisons of the periodogram with
the autocorrelation function and the power spectrum [6].

Because both methods, the spectral analysis and the
periodogram, were originally designed for the detection of
only circadian rhythms, we tested the usefulness of these
methods for the detection of ultradian rhythms with simu-
lated data. The input data were generated by computer and
contained random fluctuations as well as circadian and ul-
tradian rhythms of different period, amplitude and wave
form. Representative analyses are given in Fig. 1.

Although the number of data was limited (6 days), both
methods proved convenient for the analysis of our data.
Amplitudes of the spectral estimates and the periodogram
peaks were correlated to the "true" amplitude of the artificial
input data, although the wave form and the level of the
"random noise" also affected the amplitude. Two rhythms of
different amplitude having the same wave form and amplit-
ude resulted in spectral estimates of nearly the same amplit-
ude. One complication in the periodogram is evident when
analyzing circadian rhythms. The analysis cannot tell
whether the identified components are actually present, or
whether the periodogram peaks arise due to a component the
period of which is a submultiple (1/2, 1/3, etc.) of the apparent
values. An ultradian rhythm of 4.8 hr period will produce
periodogram peaks at 9.6 hr as well as at 14.4 hr and 19.2 hr.
Closer examination of the periodogram resolves this am-
biguity. If the periodogram contains a repeating pattern, the
"true" period length can be assigned to the appropriate
submultiple of the periods at which the repeating peaks oc-
cur.

Significant ultradian rhythms were not caused by
non-stationarities or asymmetry of a 24-hr rhythm in any of
the analyzed examples.

**EXPERIMENT 1: EFFECT OF OVARIECTOMY AND ESTRADIOL
REPLACEMENT ON THE ACTIVITY PATTERN OF ADULT LEW/Ztm
FEMALES**

**Method**

Locomotor activity was measured under LD 12:12 in 6
adult female rats of the LEW/Ztm strain starting at approxi-
mately 80 days of age. As a control, locomotor activity of
these animals was recorded under these conditions for 20
days. On day 21, the animals were ovariectomized (OVX)
under anesthesia (100 mg/1 kg b.wt. Ketavet, Parke, Davis &
Company, München, + 0.05 ml Rompung Bayer Leverkusen,
IP) and were monitored for an additional 20 days. The
surgery took place in the experimental room during the light
period of the lighting regime.
On day 42, the animals were implanted with Silastic capsules (Dow-Corning, 1.57 mm i.d., 3.18 mm o.d.) filled to 15 mm with estradiol-17β (E2-capsule). Estradiol capsules of this size are reported to produce a relatively constant proestrus-like level of estradiol in ovariectomized Sprague-Dawley rats [32]. The capsules were incubated in water (21°C) for 30 min prior to implantation in order to minimize initial transitory hormone release. The E2-capsules were implanted subcutaneously in the lower dorsal thoracic region of the anesthetized rats.

The measurements of locomotor activity were continued for a total of 60 days. A processing failure caused the loss of data from one rat after day 22. Once activity recordings ceased, the animals were sacrificed with an overdose of anesthetic and were examined for the presence of the Silastic implant. The 15-mm estradiol Silastic capsules released 14.81±2.27 μg estradiol/day (means±SD). These values were determined by the difference in dry weight of the capsules before and after implantation. Vaginal smears were not taken during the experiment. Inherent handling would have disrupted the activity pattern.

**Results**

Figure 2 demonstrates the effects of ovariectomy and estradiol replacement on a representative female rat of the LEW/Ztm strain. The activity record is shown on the left side. A continuous spectral analysis appears in the middle of the figure. During the first 20 days under the LD 12:12 control period, the activity of the representative animal in Fig. 2 exhibited a clear 24-hr rhythm. The amplitude of the 24-hr spectral estimate ranged from $S'=300$ to $S'=500$ for the entire registration period. The amplitudes of the 4-hr and 4.8-hr spectral estimates were very low during the first 20 days. Shortly after ovariectomy (OVX), the activity pattern changed. At first, very short activity bouts were observed during the whole day. Within a few days post surgery, a stable pattern with 3-4 activity peaks occurred during the night phase, resulting in a marked increase of the amplitudes of the 4-hr and 4.8-hr spectral estimates. Implantation of an estradiol capsule on day 42 (E2-capsule) abolished the effect of ovariectomy. The activity record indicated no regular activity peaks during the night. The amplitudes of the 4-hr and
each animal by using the spectral analysis [26, 38] and the chi square treated with oil alone. Both animals were measured for 13 days under LD 12:12. The chi square periodogram of these individual data is shown below the activity record of each animal on the left: the spectral analysis on the right.

**TABLE I**

<table>
<thead>
<tr>
<th>Animal</th>
<th>Day 1-18</th>
<th>Day 30-41</th>
<th>Day 43-57</th>
<th>Day 64-81</th>
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<tbody>
<tr>
<td>1</td>
<td>4</td>
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<tr>
<td>7</td>
<td>4.8</td>
<td>4.8</td>
<td>4</td>
<td>4</td>
</tr>
</tbody>
</table>

Rhythmic components of distinct time intervals were derived for each animal by using the spectral analysis [26, 38] and the chi square periodogram [49]. Statements about the period of significant ultradian components were based on the application of both methods. Significant peaks in the periodogram were determined according to a $p<0.001$ level of significance. As shown previously, an ultradian rhythm with a period of 4 hours results not only in a peak at 4.0 hr, but also in peaks at 8.0, 12.0, 16.0 hr etc. This, likewise, holds for other ultradian rhythms. As a result, a series of multiple ultradian peaks was always interpreted as being caused by only one rhythmic component; i.e. the component of the shortest period. The 24-hr rhythm, which was present in all analyses, was not included in this table.

4.8-hr spectral estimates decreased to initial levels. As shown on the right side of Fig. 2, similar results were obtained with the chi square periodogram analysis of these data. Significant ultradian periodicities were observed during the second phase of the experiment only.

The effects of ovariectomy and estradiol replacement were observed in each of the 5 LEW/Ztm females studied. These results are shown in the pooled spectral analyses of Fig. 3. The pooled spectrum of the first experimental phase showed a single clear 24-hr component. After ovariectomy additional ultradian peaks of 4 hr ($S'=240\pm50$) and 4.8 hr ($S'=362\pm143$) periods appeared in the spectrum. The amplitudes of these ultradian components were significantly higher ($p<0.01$) than the corresponding values both before ovariectomy and after E2-replacement.

These hormonal treatments did not appear to affect the 24-hr rhythm. Although the amplitude of the 24-hr spectral estimate seemed to be lower after ovariectomy ($S'=308\pm78$), these changes in the amplitude are insignificant. Although inspection of the ovaries after ovariectomy detected the presence of corpora lutea, no alterations in the activity onset that might depend upon the stage of estrous cycle were detectable during the first 18 days.

In conclusion, it is obvious from experiment 1 that ovariectomy and E2-capsule implantation have dramatic effects on the activity pattern of female LEW/Ztm rats. Following ovariectomy the females exhibited ultradian rhythms similar to those shown by untreated males of this strain. It seems likely that physiological levels of estradiol normally prevent ultradian rhythms in adult females.
**Experiment 2: Effect of Castration and Estradiol Implantation on the Ultradian Activity Pattern of Adult LEW/Ztm Males**

**Method**

Locomotor activity was measured under LD 12:12 in 7 adult male rats of the LEW/Ztm strain starting at approximately 80 days of age. All animals were recorded under these conditions for a period of 20 days. On day 21, five males (Nos. 1-5) were castrated (GOX) under anesthesia, two further males (Nos. 6 and 7) were sham operated. The animals were monitored for an extended period of 35 days to ensure elimination of testosterone from the plasma. Ultradian rhythms persisted throughout the whole testing period after castration. On day 56, 15 mm estradiol capsules (E2-capsule) were implanted in the lower thoracic region of five animals (Nos. 1, 2, 5-7). Two animals (Nos. 3 and 4) were sham operated. The measurements of locomotor activity continued until day 80.

**Results**

Figure 4 shows the effect of castration (GOX) and estradiol implantation (E2-capsule) on the activity pattern of one representative LEW/Ztm male (No. 2). During the first 20 days under LD 12:12 this animal exhibited a clear 24-hr rhythm. The amplitudes of the spectral estimates ranged between $S' = 400$ and $S' = 700$. The continuous spectral analysis showed additional significant ultradian rhythms of 4.8 hr, 4 hr, and 3.4 hr periods. In the periodogram, ultradian rhythms of 4.8 hr and 4 hr periods were detected. The ultradian activity pattern was not abolished by castration of the animal on day 21. The 24-hr component as well as the ultradian components persisted, but a 3-hr component became more prominent.

Implantation of an E2-capsule on day 56 had a dramatic effect on the circadian rhythms. Within a few days after implantation the amplitudes of the determined ultradian spectral estimates decreased to a non-significant level.

As shown on the right in Fig. 4, these effects of castration and E2-implantation were confirmed by the chi square periodogram analysis of these data. The periodogram from days 1-18 showed significant ultradian rhythms of 4 and 4.8 hr periods. The analysis of days 43-56 detected a main 3-hr rhythmic component. The periodogram from days 64-81 did not indicate any such significant ultradian components.

These effects of castration and E2-implantation were generally observed in all of the similarly treated animals (Table 1). No differences appeared in the activity patterns of the castrated and sham operated animals. On the other hand, a clear difference between the animals with E2-implantation and those without was obvious. Two animals (Nos. 3 and 4) which were sham operated on day 56 failed to show any reduction of ultradian rhythmicity during the experiment, whereas the animals with E2-implantation displayed no ultradian rhythms at all.

Although no statistical methods are available to secure the differences in the periods presented in Table 1, it can be concluded from experiment 2 that testosterone has no effect on the ultradian activity pattern of the LEW/Ztm males. The direct role of estradiol on the inhibition of ultradian rhythmicity in males of this strain, however, was confirmed.

**Experiment 3: Effect of Perinatal Treatment with an Androgen Antagonist on the Activity Pattern of Male LEW/Ztm Rats**

Androgens are known to effect several sexually differentiated behaviors of the adult rat. In addition, these steroid hormones have a permanent masculinizing effect on the brain during a critical stage of development which in the rat is around the day of birth [4, 14, 21]. Experiment 3 was designed to examine the effect of an androgen antagonist, cyproterone acetate, in preventing perinatal androgenization [4, 20, 37].

**Method**

Newborn LEW/Ztm pups of several litters were allocated to separate groups according to their sex and hormonal treatment. Afterwards, each group was treated separately and members of one group always received identical injections. These precautions excluded the possibility of potential contamination via hormone leakage from litter mates. Six males were given 0.5 mg cyproterone acetate (Schering AG, Berlin) in 0.05 ml corn oil for the first 10 days after birth ($\alpha$-CA). Two males received pure 0.05 ml corn oil from day 1-10 ($\alpha$-oil). The number of control animals was limited to two, because applicable data from 7 control males in experiment 2 were available for direct comparison. All animals were weaned at 28 days of age and were maintained under LD 12:12 in treatment-segregated groups. From approximately 80 days of age on, the animals were individually housed and locomotor activity was recorded for 13 days under LD 12:12.
Results

Comparison of the activity patterns, the spectral analyses, and the chi square periodograms of two LEW/Ztm males is shown in Fig. 5. The LEW/Ztm male No. 4, treated with corn oil postnatally, showed a typical ultradian activity pattern of 4 hr and 4.8 hr period.

The activity pattern of the LEW/Ztm male No. 1, treated with cyproterone acetate postnatally, was also characterized by ultradian rhythms. Main rhythmic components of 4.8 and 3 hr periods were evident.

The individual spectral analyses were pooled and are shown in Fig. 6. Statistical comparisons of the 4.8-hr and 4-hr spectral estimates as well as the 24-hr spectral estimates indicated no significant differences in the amplitude of distinct spectral estimates between the untreated control males of experiment 2 (LEW 6), the cyproterone acetate-treated males (LEW 6-CA), and the oil-treated males (LEW 6-oil). The amplitudes of the 4.8 and 4-hr spectral estimates within both perinatally treated groups seemed to be somewhat lower than those found in the untreated LEW males. This effect may be due to considerable disturbances of the animals shortly after birth. Locomotor activity studies require conditions whereby the animals remain fully undisturbed. Therefore, treatment effectiveness on other behavioral endpoints was verified in separate groups of animals which received similar perinatal treatments (Döhler, unpublished results).

Perinatal treatment with cyproterone acetate was repeatedly shown to permanently feminize sexual behavior patterns and the mode of gonadotrophic hormone release in males [20,37]. Such treatment, furthermore, inhibits the masculinizing effect of exogenous testosterone in females [4]. Cyproterone acetate does not, however, have any effect on the ultradian activity patterns in the LEW/Ztm strain.

EXPERIMENT 4: EFFECT OF PERINATAL TREATMENT WITH TESTOSTERONE PROPIONATE ON THE ACTIVITY PATTERN OF LEW/Ztm FEMALES

Current views of sexual differentiation suggest that perinatal exposure to androgens masculinizes the brain and results in a permanent loss of ovarian and vaginal cyclicity [3, 14, 21]. Preliminary reports indicate that androgens are also responsible for sexual differentiation of the circadian system of the rat and hamster [1, 13, 58]. Experiment 4 examines the effect of postnatal testosterone treatment on differentiation of the activity pattern of female rats.

Method

Newborn LEW/Ztm pups were weaned and maintained under the conditions described in experiment 3. An initial group of 6 female animals was injected with 50 μg testosterone propionate (Serva Feinbiochemica, Heidelberg) in 0.05 ml corn oil during the first 6 days after birth (LEW 6-TP). Two females received 0.05 ml of pure corn oil (LEW 6-oil). Measurements of locomotor activity started at approximately 80 days of age and continued for 18 days.

A second group of 6 females was injected with 50 μg testosterone propionate in 0.05 ml corn oil on days 1–6. Activity recordings were performed with these animals for 18 days starting at approximately 80 days of age. On day 19 of
measurement these animals were ovariectomized under
anesthesia (LEW♀-TP + OVX) and were monitored for
another 22 days. The effectiveness of the perinatal testosterone
propionate treatment in this group was controlled by
inspection of the ovaries. The ovaries were weighed and the
presence or absence of corpora lutea was protocolled. Vagi­
nal smears were not taken during the experiments, as han­
dling would have disrupted the activity patterns.

Results

The activity patterns, the spectral analyses, and the chi
square periodograms of two female rats are compared in Fig.
7. The control female rat No. 4 was perinatally treated with
oil and showed the typical non-ultradian activity pattern of
dominating the activity pattern. The 4-hr and 4.8-hr spec­
tral estimates of the LEW♀-TP are marked with stars, because they
differ significantly (p<0.05) from the corresponding values of the
normal and the oil-treated females.

The persistence of the ultradian activity patterns in the
female LEW/Ztm rats. Female No. 1 which had been
perinatally treated with testosterone propionate (TP) exhib­
ted weak but significant ultradian rhythms of 4 and 4.8 hr
periods. The pooled spectral analyses obtained for the nor­
mal LEW/Ztm females of experiment 4, the testosterone
propionate-treated females (LEW♀-TP), and the oil-treated
females (LEW♀-oil) are shown in Fig. 8. Statistical analysis of the
data indicated no significant differences in the amplitudes of the 4-hr, 4.8-hr, and 24-hr spectral estimates be­tween the untreated (LEW♀) and the oil-treated females

(LEW♀-oil). Perinatal treatment with testosterone
propionate (LEW♀-TP) resulted in a significant increase
(p<0.05) of the 4.8-hr (S'=184±91) and 4-hr (S'=188±157)
spectral estimates with respect to both the untreated
(LEW♀) and the oil-treated control females (LEW♀-oil).

The relative ovarian weights of the testosterone
propionate-treated females were significantly lower than
those of untreated females and none of them demonstrated
corpora lutea. This observation confirmed the existence of
permanent anovulatory sterility due to defeminization of the
gonadotropin release pattern.

Female rats are normally characterized by cyclic
hypophyseal gonadotropin secretion during sexual maturity.
This can be indirectly demonstrated by the presence of
corpora lutea in the ovaries. Perinatal treatment with testos­
terone propionate results in tonic gonadotropin release, lead­
ing to permanent sterility of the females [114] for review) and
in lower production of estradiol [3].

The persistence of the ultradian activity patterns in the
testosterone-treated females may either be the result of a
masculinized brain or the result of low estrogen levels in
adulthood. This hypothesis was verified by a further experi­
ment in which female rats were not only perinatally an­
drogenized with testosterone propionate but were also
ovariectomized as adults. The results of this treatment are
shown in Fig. 9. During the first 18 days of measurement, the females showed increased amplitudes for the 4.8-hr \(S' = 203 \pm 32\) and 4-hr \(S' = 157 \pm 34\) spectral estimates which did not differ significantly from those of the LEW ?-TP females in Fig. 8.

Ovariectomy resulted in a further increase of the ultradian spectral estimates, whereby alteration of the 4-hr spectral estimate \(S' = 262 \pm 76\) was statistically significant \((p \leq 0.05)\). This supports the suggestion that expression and non-expression of ultradian activity patterns in this strain is directly related to the circulating levels of estradiol rather than to an organizational effect of testosterone.

**GENERAL DISCUSSION**

Gonadal hormones have two distinct effects on the circadian system. The presence of androgens during the perinatal period has a permanent organizational effect on the circadian system of hamsters [13, 33, 57, 58] and rats [1, 22], as well as on several other physiological and behavioral variables [3, 14, 18, 20, 21]. The results of the present study provide no evidence for a perinatal determination of the ultradian activity pattern by testosterone. Perinatal treatment of male LEW/Ztm rats with cyproterone acetate, an androgen antagonist [4, 20, 37], did not eliminate the ultradian activity pattern of the adult animals (experiment 3). Stimulation of the ultradian activity patterns in female rats when perinatally exposed to testosterone propionate (experiment 4) apparently contradicts the cyproterone acetate experiments. These contradicting observations can be explained by the fact that female rats perinatally treated with testosterone propionate display lower production levels of estradiol.

This leads to a second and more transient type of hormonal effect of circulating gonadal hormones in the adult animal. In male rodents, testosterone is known to increase the amplitude and to shorten the period of circadian activity rhythms [12]. Castration of adult LEW/Ztm males did not prevent ultradian rhythmicity (experiment 2) but seemed to effect the uniformity of the activity pattern. As shown in Table 1, non-castrated males (day 1–18) normally exhibited one or two ultradian components of 4 and 4.8 hr periods. Castration of these males (days 30–41 and 43–57) resulted in stimulation of rhythmic components with shorter periods (e.g., 2.6, 3, and 3.4 hr). Since these experiments were conducted under LD 12:12 entrainment, they do not allow for any clarification of the effect of testosterone on the endogenous ultradian rhythms.

Nevertheless, the results support previous observations which pointed out the correlation between the deterioration of precision in entrained hamsters and photoperiodic castration (i.e., testicular atrophy) during prolonged exposure to a short light-dark cycle [15]. Furthermore, gonadically testosterone was reported to restore running onset precision in hamsters [35]. Studies involving castration and testosterone replacement of male LEW/Ztm rats under continuous light conditions, where the activity rhythms are not entrained to any lighting regime and therefore display their spontaneous periods, are needed to investigate the testosterone dependent changes in the activity pattern of this strain.

The results of this study demonstrate the effect of estradiol in the prevention of ultradian activity rhythms in laboratory rats of the LEW/Ztm strain. Estradiol released continuously from long-term subcutaneous implants generally prevented ultradian rhythms in ovariectomized females as well as in castrated and non-castrated males. As a result, this effect of estradiol can be added to the list of effects caused by this steroid on a biological rhythm system. The present results support the hypothesis that the ultradian activity pattern of this strain may be the result of a hormonally based change in the phase relationship between several circadian oscillators.

On the same line one would expect a 4-5 day modulation in the activity pattern of sexually mature LEW-females, not only with respect to the onset and level of activity as previously found in rats [2, 19] and hamsters [36], but also in the appearance of ultradian rhythms. Unfortunately, our data provide no such evidence of an estrous correlated modulation of the activity pattern. This is most likely caused by the circadian system [44] used for measurement of locomotor activity, whereby all behavioral movements of the animal including feeding and grooming were registered.

In studies performed by Albers et al. [2] and by Gentry and Wade [18], no mention of ultradian rhythmicity was specifically made. Nevertheless, visual inspection of their running wheel data from female rats shows an obvious bimodal activity pattern on the days of di-, met-, and proestrous and an almost constant plateau of activity on the day of estrous.

Additional data have recently been presented which support the hypothesis that estradiol may uniformly strengthen the coupling between constituent oscillators. Circadian activity rhythms of hamsters exposed to constant illumination can be dissociated or split into components that free-run with different periods. Moin [34] has demonstrated that the administration of estradiol prevents splitting of the activity rhythm of ovariectomized female hamsters. Estradiol, likewise, inhibits other forms of abnormal rhythmicity classified by the presence of high or low frequency oscillations without obvious “splitting” [34].

In contrast to the hamster, where the circadian system is sexually differentiated and males do not apparently respond to exogenous estradiol [58], the present data did not show any sex-dependent effects of estradiol on ultradian rhythms in the LEW/Ztm rat. This is in agreement with data from a previous study [1] in which perinatal androgenization did not eliminate estrogen sensitivity in the circadian system of the rat. According to the structural model of the circadian pacemaker [40–43], “splitting” develops when mutual interactions between constituent oscillators cease as a result of a differential effect of light on the period of each oscillator. Daan et al. [12] proposed that hormones influence the length of circadian rhythms by strengthening or weakening the synchronization between the different oscillators which together govern circadian periodicity. Interpretation of the present data in the context of the coupling model presented above, suggests that estrogen suppresses the occurrence of circadian rhythms by uniformly strengthening synchrony between oscillators. Furthermore, the present data provide strong support for a multi-oscillator system with more than two oscillators regulating the activity rhythm in this special inbred strain of rats (i.e., three 24-hr oscillations which are 4.8 hr out of phase). Hohmann [27] has presented evidence for more than two free-running oscillators in the day-active animal *Tupaia belangeri*. The complex running patterns of arctic red-backed voles also suggest more than two oscillations regulating running rhythmicity [50].

It is presently unknown what aspects of neural organization are responsible for the effect of estradiol on the internal clock. Various authors have pointed out that there are a variety of direct and indirect ways steroid hormones could
influence the circadian system [12, 24, 25, 34–36, 51, 52, 57].
Evidence that the SCN does not accumulate high concentrations
of radiolabeled estrogen in either rats or hamsters
[31,39] suggests that other areas of steroid uptake connected
to the SCN (e.g., the preoptic-anterior hypothalamic area;
[10,55]) may be involved in steroid effects on circadian
rhythms. In addition, due to hormonal changes which are
known to occur in response to alterations in circulating
steroid levels, the effects of steroids on circadian rhythmicity
may be mediated by other hormones (i.e., pituitary or
hypothalamic hormones).

One of the most consistent effects of estrogens is the
dramatic increase in various activities [10, 18, 19, 53–55]. In
addition, steroid hormones also influence food intake [55],
body weight [54] and a variety of autonomic functions in
rodents [53]. Such observations support the alternative
hypothesis that steroid hormones alter circadian properties
by changing the general physiological state of an organism [51].

The described method of measuring locomotor activity
[44] records overall integrated activity and does not
differentiate between locomotion, feeding-related behavior and
other types of activity.

Although the effect of estrogens on food intake is well
known, it does not explain the observed effects of estrogens
on ultradian activity. Estrogens stimulate food intake signifi-
cantly more in control female rats than in males or
androgenized females [18]. In our study, estrogens inhibited
ultradian activity in both sexes equally. Our data correspond
well with data on running wheel activity, where males and
females responded equally to the effects of estrogen or
progestosterone [1,19]. Since the observed sex-differences in
the activity pattern of the LEW/Ztm strain are not deter-
mined by perinatal androgenization, and since exogenous
estrogen has the same effect in males and females, it is most
likely that the observed sex-differences in the activity pattern
of the LEW/Ztm strain reflect differences in locomotor
behavior alone. Further investigations of this strain in the
running wheel and measurements of feeding patterns and
food intake are necessary to answer this question satisfac-
torily.

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