Follicle-Stimulating Hormone (FSH) and Luteinizing Hormone (LH) in the Urine of Prepubertal Children *

ARLEEN B. RIFKIND, HOWARD E. KULIN, AND GRIFF T. ROSS

(From the Endocrinology Section, National Institute of Child Health and Human Development, and the Endocrinology Branch, National Cancer Institute, National Institutes of Health, Bethesda, Maryland)

Abstract. Follicle-stimulating hormone (FSH) and luteinizing hormone (LH) have been measured by specific bioassays in pooled urine samples from prepubertal children, aged 2-6 yr, and from male adults. For children the mean urinary excretion of FSH was 2.2 U 2nd International Reference Preparation (2nd IRP) per liter and the mean urinary excretion of LH was 0.44 U 2nd IRP per liter. For adults the mean FSH excretion was 5.6 U 2nd IRP per liter and the mean LH excretion was 4.7 U 2nd IRP per liter. Our data show a 2.5-fold increase of FSH, a 10.7-fold increase of LH, and a consequent decrease in the FSH:LH ratio from 5 to 1 between childhood and adulthood. FSH and LH in urine from three patients with gonadal abnormalities have also been studied. The results from normal children, adults, and abnormal patients form a spectrum and reveal that sexual maturity is accompanied by a marked increase in the excretion of LH with relatively smaller increases in FSH.

Introduction

Several laboratories have reported the detection of gonadotropins in the urine of prepubertal children by bioassay (1-3). Those studies, however, report measurement of total gonadotropins, and very little information is available on the differential excretion of follicle-stimulating hormone (FSH) and luteinizing hormone (LH) in children. While Fitschen and Clayton (2) measured LH in a single pool of urine from six prepubertal boys under 6 yr, there have been no reported measurements of FSH in the urine of prepubertal children.

The present study was undertaken to delineate the pattern of gonadotropin excretion in prepubertal children. We have measured FSH and LH by means of specific bioassays in pooled urine samples from normal prepubertal children, aged 2-6 yr, and from adult males. The data show that an increase of FSH and LH and a decrease in the FSH/LH ratio are associated with sexual maturity.

Methods

Sources of urine

Children's urine was obtained from local private nursery schools. The children were not examined by the investigators but had had physical examinations during the school year and were considered healthy.

Nursery school A was attended by approximately 40 Negro children aged 3-6 yr, and nursery school B, by approximately 50 children aged 2-5 yr, most of whom were Caucasian.

Random urine specimens from the children were pooled in containers at the schools and were transported daily to the laboratory where they were stored at 4°C for 2 days to 2 wk until processed. Pools of urine from schools A and B were usually combined. A total of 480 liters was collected and assayed.

Laboratory personnel provided the normal adult male urine. Random specimens were pooled. A total of 426 liters was collected and assayed.

Urine was also collected and assayed from three patients with abnormalities that might be expected to be accompanied by alterations in gonadotropin excretion.
The first patient was a 44 yr old girl with moderately advanced idiopathic isosexual precocity evidenced by breast development, axillary and pubic hair, and adolescent type external genitalia, who had not yet had menarche. 11 24-hr urine specimens totalling 7.2 liters were collected from her and pooled. The second patient was a 61 yr old girl with more advanced idiopathic isosexual precocity who had had menarche at 6 yr 3 months. 7 24-hr specimens totalling 9.2 liters were collected from her and pooled. The third patient was a 23 yr old female with primary amenorrhea who had a normal chromosome complement and immature ovaries. 17 24-hr specimens totalling 29 liters were collected from her and pooled.

**Concentration and purification**

Batches of approximately 15 liters of urine were processed separately according to Albert's kaolin-acetone method for fraction A (4). The moist precipitate was lyophilized. Fraction B (5) was then prepared, usually on amounts of precipitate equivalent to 30 liters of urine, according to the following procedure. The dry precipitate was extracted in a centrifuge tube with freshly prepared 10% ammonium acetate in 70% ethanol (7 ml extractant for each 3 liter equivalent of precipitate) by vigorous shaking, followed by 20 min of mechanical stirring at 4°C. Centrifugation was carried out in a Sorvall RC2B centrifuge at 23,000 g for 15 min, and the supernatant was decanted and saved. The residue was extracted again in the same manner, the second supernatant was added to the first, and the residue was discarded. 2 volumes of freshly prepared 10% ammonium acetate in 100% ethanol were then added to the combined supernatants. A final precipitate was allowed to form at 4°C for periods ranging from 2 to 14 hr. It was then spun down at 23,000 g for 15 min and saved, and the supernatant was discarded. The precipitate was dissolved in 1 ml of distilled water for each liter equivalent of urine and frozen until used for bioassay.

**Bioassays**

**FSH.** FSH was measured by the ovarian augmentation reaction using a modification of the method of Steelman and Poiley (6). Holtzman immature female rats each weighing 40–50 g were used. Human Chorionic Gonadotropin,¹ 50 U/animal, was mixed and injected simultaneously with each dose of test material. Five animals were used for each test dose except when otherwise indicated. It was usually possible to discriminate between twofold dose intervals in the augmentation assay.

The 2nd International Reference Preparation for Human Menopausal Gonadotropin ² (2nd IRP) was used as a standard in all FSH assays. Significant responses in the linear portion of the dose response curve could be assured with doses of 2 and 4 U 2nd IRP.

The pooled adult urine extract was tested at three doses equivalent to 0.33, 0.67, and 1.33 liters of urine per animal to assure at least two responses in the linear portion of the dose response curve. The children's extract was tested at one dose equivalent to 1.5 or 2.0 liters of urine per animal, except for two assays (Nos. 11 and 13) in which two doses equivalent to 1 and 2 liters of urine were used.

Doses of extract equivalent to 0.75, 1.0, and 2.0 liters of urine were used for the patient with moderate isosexual precocity, advanced isosexual precocity, and primary amenorrhea, respectively. Four animals were used to test each abnormal child's extract.

There were saline and HCG injected controls in each assay. The test materials were diluted with normal saline so that each animal would receive its total dose in a volume of 4 cc. The total dose was delivered over 4 days with 1 cc injected the 1st day and 0.5 cc injected in the morning and afternoon of the 2nd, 3rd, and 4th days. Autopsy was performed on the morning of the 5th day, and both ovaries were removed and weighed on a torsion balance.

**LH.** The ventral prostate weight assay in hypophysectomized immature male rats as described by McArthur (7) was used to measure LH. Immature Sprague-Dawley rats weighing 40–50 g each were delivered 1 day after hypophysectomy at which time injections were begun. Injection and autopsy schedules and total volumes of injected material were the same as in the FSH assay. Unless otherwise indicated, five animals were used for each dose. In order to discriminate between doses in the ventral prostate assay, it was usually necessary to use fourfold dose intervals.

Humegon³ at dose levels of 0.8 and 3.2 U with five to seven animals per dose was used as a standard in all assays. The 2nd IRP was also used as a standard in assays No. 5 and 7–10. Doses of 1.2 and 4.8 U 2nd IRP assured consistent responses in the linear portion of the dose response curve. The data from those assays in which Humegon alone was used as a standard were converted to 2nd IRP by means of the factor 1 U Humegon = 1.5 U 2nd IRP, derived from the assays in which both Humegon and 2nd IRP were used.

Doses equivalent to 0.42 and 1.67 liters of male urine per animal consistently gave differential responses and were used for all assays of male pools. In assaying children's pools, it was necessary to use amounts of extract equivalent to 2.5 or 3 liters of urine to achieve consistently significant responses. In all assays one dose was tested with five animals except for assay No. 6 in which only three animals were used.

Doses equivalent to 0.75, 1.0, and 3.0 liters of urine were used for the patients with moderate isosexual precocity, advanced isosexual precocity, and primary amenorrhea.

of variance each of the Statistical evaluation was subjected to an analysis of variance using a Fortran IV computer program designed in this laboratory (8). F ratios to test for deviation from parallelism, indices of precision, and relative potencies with their 95% confidence limits, were evaluated.

Results

Table I shows the results of the LH assays for children and adults in terms of the 2nd IRP. All the adult pool assays had satisfactory indices of precision, and showed no significant deviation from parallelism with the standard. The mean value for LH in adult urine pools was 4.7 ± 0.6 se U 2nd IRP per liter.

The doses of children's urine extract used always gave responses at least two standard errors above control values, except in one assay (No. 5) in which the response was one standard error above control values. It was not possible to run two discriminating doses of the children's urine because of lethal animal toxicity at higher doses. Therefore, the data for LH in children should be interpreted with the reservation that technical limitation did not permit any evaluation of parallelism with the standard.

Weight loss and mortality are thought to be criteria of significant toxicity in gonadotropin bio-

<table>
<thead>
<tr>
<th>Assay No.</th>
<th>Male adults</th>
<th>Children</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>IU 2nd IRP/liter</td>
<td>Bioassay design</td>
</tr>
<tr>
<td>11</td>
<td>6.7(5.5–8.3)§</td>
<td>2 + 2</td>
</tr>
<tr>
<td>12</td>
<td>1.1(0.86–1.4)</td>
<td>1 + 2</td>
</tr>
<tr>
<td>13</td>
<td>2.2(1.7–2.9)</td>
<td>1 + 2</td>
</tr>
<tr>
<td>14</td>
<td>1.8(1.4–2.2)</td>
<td>2 + 2</td>
</tr>
<tr>
<td>15</td>
<td>1.9(1.6–2.5)</td>
<td>1 + 2</td>
</tr>
<tr>
<td>Mean</td>
<td>5.6 ± 1.0 se</td>
<td>2.2 ± 0.4 se</td>
</tr>
</tbody>
</table>

* Abbreviations as in Table I.
† No. of doses of unknown plus No. of doses of standard.
§ Mean potency estimate with 95% confidence limits.
Differences (9). All our test rats gained weight and had mortality rates similar to the controls, thus showing no signs of major toxicity. With the high doses of children’s urine, however, the test animals occasionally evidenced lethargy and roughening of fur, suggesting some toxicity. That these changes did not affect the LH potency estimates has been shown by radioimmunoassay data from our laboratory 4 confirming LH potency estimates derived by bioassay from urine extracts of both children and adults.

Mean LH excretion for the children was 0.44 ± 0.07 × 10^-6 U 2nd IRP per liter of urine. The values for children and adults differ significantly (P < 0.001).

Table II shows the results of the FSH assays. Three adult pools were tested with a mean value of 5.6 ± 1.0 × 10^-6 U 2nd IRP per liter. All these assays also had satisfactory indices of precision and showed no significant deviation from parallelism with the standard.

Because of insufficient material, the children’s extract was tested at only one dose level in five assays. However, to assure that the children’s extract gave responses that were parallel and therefore comparable to the standard, two 2-point assays, Nos. 11 and 13, were run. In both these assays, the dose response curve of the children’s urine was satisfactorily parallel to the standard.

In order to test for marked variability between sources of children’s urine, several urine pools from nursery schools A and B were assayed separately for FSH. Three pools from nursery school A alone gave means of 2.2, 1.9, and 4.5 U 2nd IRP per liter. One pool from nursery school B gave a value of 2.5 U 2nd IRP per liter. These values were sufficiently similar to suggest that there were no variations in results caused by differences in sources.

The seven pools of children’s urine gave a mean value of 2.2 ± 0.4 × 10^-6 U 2nd IRP per liter. The values for children and adults are significantly different (P < 0.005).

Table III contrasts excretion of FSH and LH in children and adults. The results are listed in terms of units per liter and units per meter² per 24 hr. The estimated factor for converting excretion in terms of liters to meter² per 24 hr is similar in both children and adults, so that relative differences between children and adults remain the same when either units are used. Our data indicate that adult males excreted 2.5 times as much urinary FSH and 10.7 times as much urinary LH as prepubertal children, whether values are expressed per meter² per 24 hr or per liter of urine.

Table IV shows FSH/LH ratios in male adults,
children, and the abnormal patients. For adults, three values are listed. The first two represent pools of urine which were divided and aliquots of the same pool assayed for FSH and LH. The third represents the FSH/LH ratio calculated from the mean values for FSH and LH from all the assays of adult urine pools. The FSH/LH ratio found in adults was 1.2.

For the children, two values are listed. The first represents the ratio for one pool which was divided and aliquots assayed for FSH and LH. The second value is the ratio of means from all the FSH and LH assays. The ratio of 11.8 from the single pool is higher than the ratio of 5.0 for the means because the single pool gave a particularly low value for LH. Ratios between children and adults differ significantly ($P < 0.005$).

The ratio for the child with advanced isosexual precocity was 0.53. The ratio for the child with moderate isosexual precocity was 2.8. The ratio for the 23 yr old female with primary amenorrhea was 4.0.

**Discussion**

The main problems in measuring the separate gonadotropins in children's urine by bioassay have been the difficulty in obtaining sufficient material and the toxic effects of the large amounts of urine concentrates that are needed for a significant response. The use of Albert's fraction B purification method for urine extracts has enabled us to give doses sufficient to achieve responses without obvious toxicity.

There are very few reports on LH excretion in children with which to compare our data. McArthur et al. (10), using kaolin-acetone extracts (fraction A) and the ventral prostate assay, were able to detect urinary LH activity in only 3 out of 12 children, aged 4–11 yr. Fitschen and Clayton (2), using fraction B and the ovarian ascorbic acid depletion assay for LH, found $0.89 \mu g$ NIH-LH-S1/24 hr in a single pool of urine from six boys under 6 yr. By converting this value to 2nd IRP $^5$

$^5$ Because the 2nd IRP has not been used as a standard in most published studies of urinary FSH and LH, in order to compare our results with those of other investigators conversion factors have been used. To express NIH-FSH-S1 and NIH-LH-S1 standards in terms of the 2nd IRP, Rosenberg and Lewis' (11) factors, 1 mg NIH-FSH-S1 = 25 U 2nd IRP and 1 mg NIH-LH-S1 = a value of 0.45 U 2nd IRP/24 hr is obtained.

FSH has not been measured previously in the urine of children under 10 yr.

Our results for FSH and LH in adult males agree well with other published values on male urine pools (5, 12, 13) when the reported values have been converted to 2nd IRP (11).

Urinary excretion of both FSH and LH appears to undergo an increase as sexual maturity is reached. However, our data suggest that whereas FSH is present in the urine in easily detectable amounts before puberty and increases only about 2.5-fold by adulthood, LH is barely detectable in the urine of younger children and increases approximately 10.7-fold in adults. Several investigators (10, 14), have speculated, on the basis of indirect evidence, that there would be more FSH than LH in the urine of prepubertal children. The greater increase of LH associated with puberty is responsible for a shift in FSH/LH ratios from prepubertal values of about 5.0 to adult values of about 1.2.

Though information derived from a few individuals may be misleading, the data from the pathologic cases which have been studied here fit well into the observed pattern of decreasing FSH/LH ratios associated with sexual maturity. Whether the FSH/LH ratio or a specific amount of FSH and/or LH is a critical factor in the initiation of puberty cannot be answered by this study. The failure of sexual development in the 23 yr old patient with primary amenorrhea who had both a childhood type of FSH/LH ratio and childhood levels of FSH and LH does not help in discriminating among these possibilities. The ratio of 0.53 in the child with advanced isosexual precocity was closest to the adult ratio. The ratio of 2.8 in the patient with moderate isosexual precocity who had not had the onset of menarche was intermediate between the ratios for children and adults. The development of puberty in both these children in whom LH, but not FSH, had increased significantly above childhood levels suggests that physiologic changes may be associated with increments in LH alone.

There are no data with which to compare our 62.5 U 2nd IRP, were used. It must be emphasized that conversion factors applicable in one laboratory cannot be used by another with any certainty of accuracy.
FSH/LH ratios in children. However, our ratio of 1.2 in males agrees well with the results of other investigators (5, 12, 13, 15, 16) who have reported FSH/LH ratios in male urine ranging from 1.1 to 2.5. As Rosenberg and Solod (15) have pointed out, individual variability of FSH/LH ratios may be masked by pooled samples.

Differences in FSH/LH ratios between men, postmenopausal women, and eunuchs are still a matter of some controversy. Whereas most investigators (12, 13, 15–19) found that FSH/LH ratios in urine pools from men and postmenopausal women were similar, others (20–21) report differences. Eunuchs, however, have uniformly been found to have urinary FSH/LH ratios 2.0 to 6 times higher than either normal men or postmenopausal women (13, 16, 17). Although prepubertal children excrete smaller amounts of gonadotropin than eunuchs they exhibit the same high FSH/LH ratio.

We cannot be certain that the differences in urinary gonadotropin excretion between children and adults reported here are accurate reflections of differences in plasma levels. The resolution of this problem awaits the perfection of techniques for measurement of FSH and LH in low titer plasma.

While it is possible that the observed changes in gonadotropin excretion are causal factors in sexual development they may be mere accompaniments of that process and other factors may be crucial. Thus, consideration should be given to explanations of sexual maturation in terms of possible differences in chemical structure between gonadotropins of children and adults, or the development of circadian peaks of gonadotropin release by the pituitary, or as yet undetermined differences in gonad sensitivity between children and adults. Though significant changes in gonadotropin excretion between childhood and adulthood have been demonstrated in this study, the causes of puberty are still unknown.

Acknowledgments

We are grateful to Mrs. C. Bowden and the staff and children of Green Hill Day Nursery and Mrs. A. Trainor and the staff and children of the Bethesda Community School for their cooperation in the difficult job of collecting urine, to Mr. James Brice for his excellent technical assistance, and to Dr. John Gart for his help in the statistical evaluation.

References


