Ovarian ultrasound assessment in normal children, idiopathic precocious puberty, and during low dose pulsatile gonadotrophin releasing hormone treatment of hypogonadotropic hypogonadism

R STANHOPE, J ADAMS, H S JACOBS, AND C G D BROOK

Endocrine Unit, Middlesex Hospital, London

SUMMARY Ultrasound assessment of ovarian volume, follicular size, and uterine growth was undertaken in 40 normal premenarcheal girls aged 6 months to 14 years. Ovarian follicles were detected from 6 months of age and increased in size and number, so that after the age of 8·5 years there was a progressive increase in the incidence of ‘megalocystic’ appearance. Eight girls with idiopathic precocious puberty had large megalocystic ovaries which regressed on treatment with gonadotrophin releasing hormone analogue. These findings contrasted with those seen in two prepubertal patients with hypogonadotrophic hypogonadism in whom the natural sequence of ovarian development was greatly accelerated by pulsatile administration of native gonadotrophin releasing hormone.

The initiation and maintenance of human sexual development requires the pulsatile release of gonadotrophins from the pituitary in response to pulsatile gonadotrophin releasing hormone.¹ Treatment of hypogonadotropic hypogonadism with pulsatile gonadotrophin releasing hormone can restore reproductive function in both males² and females.³ The use of analogues that desensitise the pituitary response to endogenous gonadotrophin releasing hormone has been established as an effective treatment of idiopathic precocious puberty.⁴ Buserelin (6-D-Ser-(TBu)-GnRH-(1–9) ethylamide) is a superactive analogue that desensitises the pituitary by down regulation of the gonadotrope and thereby interferes with the secretion of gonadotrophins.⁵

Necropsy studies have shown that ovarian size increases gradually during childhood.⁶ Normal adolescent ovaries contain multiple follicular cysts⁷ whose development determines the rapid ovarian enlargement at puberty.⁸ Both ovarian and follicular size increase progressively from fetal life to puberty and there is histological evidence of a continued turnover of follicles throughout childhood.⁹ Ovaries containing multiple cysts have been shown by ultrasound in normal prepubertal girls¹⁰ and single or multiple follicular cysts have been reported in girls with precocious puberty.¹¹

We have used ultrasound as a non-invasive method of assessing ovarian size and structure as well as uterine changes in normal children. We have compared the ovarian morphological changes of normal childhood with those seen during the suppression of idiopathic precocious puberty and during induction of puberty in patients with hypogonadotropic hypogonadism.

Patients and methods

We studied 40 normal premenarcheal girls, eight with idiopathic precocious puberty, four of whom were treated with analogue, and two with delayed puberty treated by pulsatile administration of native gonadotrophin releasing hormone. All ultrasound scans were performed by JA using an Emisonic 4201 static scanner with a long focussed 3-5 MHz transducer and more recently an Aloka sector scanner with 3-5 mHz long focussed and 5-0 mHz transducers. Ovarian volume was calculated by measuring length, width, and depth and by assuming the shape to be ellipsoid.¹² Uterine size was estimated from measurements of maximum length and depth in sagittal section and expressed as cross sectional area.

Four of the patients with precocious puberty were treated with Buserelin (gonadotrophin releasing hormone analogue), given intranasally between 400 and 800 µg daily in divided doses. Native gonado-
trophin releasing hormone was administered subcutaneously in pulses using a miniature infusion pump: a dose of 15 µg was administered at 90 minute intervals throughout 24 hours.

Informed parental consent was given for normal control patients.

Results

Ovarian volume gradually increased during childhood from a mean of 1 cm³ at 2 years of age to a mean of 2 cm³ at 12 years (Fig. 1). A maximum follicular diameter of 7 mm was normal in early childhood (Fig. 2). We define a 'megalocystic' ovary as one containing more than six follicles greater than 4 mm in diameter present in each ovary (Fig. 3). There was a progressive increase in the proportion of normal girls with megalocystic ovaries over the age of 8-5 years (Fig. 1). Uterine size was unchanged from 6 months to 10 years, mean cross sectional area was 2 cm² (Fig. 4) and there was a rapid increase in uterine size after the age of 10 years associated with the onset of puberty.

![Fig. 1 Ovarian volume of 40 normal premenarcheal girls (circles) and eight girls with idiopathic precocious puberty (triangles). Megalocystic ovaries are represented as solid symbols, those not megalocystic as open symbols.](image)

![Fig. 2 Diameter of the largest ovarian follicle in either ovary of 40 normal premenarcheal girls (circles) and eight girls with idiopathic precocious puberty (triangles).](image)

![Fig. 3 Sagittal ultrasound image of a megalocystic ovary from a normal prepubertal 8.8 year old girl, seen through the bladder (B) which is the dark area at the top of the image. The white lines represent 1 cm markers. Arrows outline the margin of one ovary which contained eight 'cysts' greater than 4 mm in diameter. V indicates the ovarian vessels entering the superior pole of the ovary.](image)

![Fig. 4 Uterine cross sectional area of 40 normal premenarcheal girls (circles) and eight girls with idiopathic precocious puberty (triangles).](image)

At presentation, all eight girls with idiopathic precocious puberty had large megalocystic ovaries, greater than 3 cm³ in volume (Fig. 1), which approached or fell within the normal adult range. Maximum follicular diameter was within the normal range for age (Fig. 2). All eight had uterine cross sectional areas greater than 4 cm² (Fig. 4). Four of the eight patients were treated with intranasal gonadotrophin releasing hormone analogue which resulted in either arrest or regression of the clinical
puberty. The bladder (B).

Horizontal lines represent 1 cm markers. The ovary was large (3.2×2.2 cm) and megalocystic. Breast development was Tanner stage III.

(b) Sagittal ovarian ultrasound image from the same girl after one year of gonadotrophin releasing hormone analogue treatment.

The scale of 1 cm markers is along the bottom of the image. The ovary (arrowed) was reduced in size (2.5×1.7 cm) and intra-ovarian 'cysts' were suppressed both in size and number. Breast development had regressed to Tanner stage II.

signs of their pubertal development. The change in pubertal status correlated with a decrease in ovarian size and regression of the megalocystic ovarian appearance (Fig. 5).

Case reports

Case 1. An 18 year old girl who presented with short stature and primary amenorrhoea, had been treated at the age of 17 years by transethmoidal resection of a craniopharyngioma. After the operation she had mild diabetes insipidus and adrenocorticotrophic and growth hormone deficiencies, for which she received hormone replacement treatment. Gonadotrophin responses to an intravenous diagnostic injection of gonadotrophin releasing hormone are shown in Fig. 6. Ovarian ultrasound showed small ovaries containing only a few small follicles, and a very small uterus. There was a gradual ovarian enlargement and increase in follicle numbers and size in response to pulsatile gonadotrophin releasing hormone treatment, so that by day 220 both ovaries were megalocystic.

Case 2. A 17 year old girl presented with no signs of puberty. Endocrine assessment showed no abnor-

mality except in the gonadotrophin responses to a diagnostic test of 100 μg intravenous gonadotrophin releasing hormone (Fig. 6). Ovarian ultrasound showed very small ovaries containing no follicles. Two weeks after beginning pulsatile treatment, there was a pronounced increase in the gonadotrophin response to a diagnostic test of intravenous gonadotrophin releasing hormone (Fig. 6). The pattern of growth of the ovary and uterus in response to treatment is shown in Fig. 7. After 8 weeks of treatment the ovaries became megalocystic. During a 10 day period of mechanical pump failure both ovaries and uterine size decreased,

Fig. 5(a) Sagittal ultrasound image of one ovary (arrowed) of a 2½ year old girl with idiopathic precocious puberty. The dark area at the top of the image is the bladder (B).

Fig. 6 Follicle stimulating hormone and luteinising hormone response, at 0, 20, and 60 minutes, to a diagnostic test of an intravenous injection of 100 μg native gonadotrophin releasing hormone.

In case 2 the test was repeated after two weeks of pulsatile gonadotrophin releasing hormone treatment.
although follicular size continued to increase. She had developed stage II breast development\textsuperscript{14} after three months of treatment.

**Discussion**

Although ultrasound imaging provides a non-invasive method of assessing ovarian and uterine changes, there have been few gynaecological ultrasound studies in normal childhood. Difficulty has been experienced in ovarian ultrasonic definition in infants and young children,\textsuperscript{15,16} even up to the age of 7 years.\textsuperscript{17} Our data on uterine and ovarian ultrasound assessment of normal girls is in agreement with more recent work.\textsuperscript{18,19} In addition to measuring ovarian size we have been able to assess intra-ovarian morphology using ultrasound in normal girls as young as 6 months of age.

We have found that pelvic ultrasound assessment is helpful in the diagnosis and classification of precocious puberty. The presence of a large uterus with a cross sectional area greater than 4 cm\textsuperscript{2} confirms this diagnosis. In addition to the uterine changes, the ovaries in idiopathic precocious puberty are large, approach adult size, and are always megalocystic. An ovarian volume of greater than 3 cm\textsuperscript{3} is outside the normal range and confirms a diagnosis of idiopathic precocious puberty. Treatment with gonadotrophin releasing hormone analogue is associated with regression of the megalocystic ovarian appearance.

In our experience of inducing puberty in patients with hypogonadotropic hypogonadism, changes occur in ovarian morphology and uterine size which closely resemble those seen in normal puberty. The megalocystic appearance is an ovarian response to pulsatile gonadotrophin secretion. We hypothesise that pulsatile secretion of gonadotrophin releasing hormone in low amplitude and long periodicity leads to the beginning of ovarian maturation early in childhood, before there are any clinical signs of puberty. A phase of megalocystic ovarian appearance occurs before the development of a dominant follicle. These changes can be reproduced in patients with hypogonadotropic hypogonadism treated with pulsatile gonadotrophin releasing hormone by varying pulse frequency and amplitude. The megalocystic ovarian appearance is dependent on pulsatile gonadotrophin releasing hormone secretion and can be suppressed with analogue.

Ultrasound imaging has a key role in the diagnosis and management of disorders of sexual maturation, reducing the requirement for more complex biochemical measurements. The information afforded by this non-invasive technique offers a new way of establishing the relation of the neuroendocrine changes with the more familiar clinical markers of sexual maturation.

We thank Dr Patrick Magill, (Senior Medical Advisor, Hoechst UK) for supplies of native gonadotrophin releasing hormone and analogue (Buserelin).

**References**


Correspondence to Dr C G D Brook, Middlesex Hospital, Mortimer Street, London W1A 9AA.

Received 28 August 1984
Ovarian ultrasound assessment in normal children, idiopathic precocious puberty, and during low dose pulsatile gonadotrophin releasing hormone treatment of hypogonadotrophic hypogonadism.

R Stanhope, J Adams, H S Jacobs and C G Brook

Arch Dis Child 1985 60: 116-119
doi: 10.1136/adc.60.2.116

Updated information and services can be found at:
http://adc.bmj.com/content/60/2/116

These include:

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/