SUBFERTILITY - BIOCHEMICAL ASSESSMENT AND INITIAL INVESTIGATIONS

NOTES:

• SUBFERTILITY IS DEFINED AS FAILURE TO CONCEIVE FOR TWO YEARS.
  • The basic guidelines below are designed to allow initial classification of problems (typically in primary care) and appropriate specialist referral for further investigation.
  • Where possible, suggested cut-offs mirror those used in current laboratory reporting comments.

FEMALES

NB: Women with fertility problems are no more likely to have thyroid disease than the general population. Thyroid function testing should thus be confined to women with symptoms of thyroid disease (NICE Guideline).

A pre-requisite for logical investigation is knowledge of the frequency and regularity of menstrual cycles.

REGULAR MENSTRUAL CYCLE

Measure mid-luteal phase progesterone (e.g day 21 of 28 day cycle; day 28 of 35 day cycle): Sample should be taken 7 days before the expected period and interpreted when the next period has begun.
  • >30 nmol/L is consistent with ovulation and no further biochemical assessment is indicated
  • 10-30 nmol/L may require repeat with review of cycle timing
  • <30 nmol/L (confirmed) requires further investigation as below for anovulation/oligo-ovulation

Prolonged irregular menstrual cycles
Progesterone should be measured. Depending on the timing of the menstrual period, this may need to be conducted every 3rd or 4th day from day 21.

ANOVULATION AND OLIGO-OVULATION

Measure FSH, prolactin, oestradiol and testosterone (in early follicular phase days 2-7, if cycle history allows).

NB. Progesterone measurement has no role in the assessment of subfertility in females without a menstrual cycle.
Significant patterns include:

- FSH > 27 IU/L (usually with oestradiol <150 pmol/L) indicates primary ovarian failure on more than one occasion.
- FSH & LH <1.5 IU/L with low oestradiol (<100 pmol/L) suggests possible pituitary or hypothalamic disease. Anorexia and the effect of other medication should also be considered.
- Prolactin >1000 mU/L requires assessment for pituitary tumour after exclusion of macroprolactinaemia and drug-related causes. The laboratory will advise on this. Milder elevations (>700 but <1000 mU/L) should be repeated to eliminate a stress-related cause.
- Mildly raised testosterone (>3.7 nmol/L) suggests possible polycystic ovary syndrome
- A confirmed testosterone > 5 nmol/L requires further investigation: Consider 17- hydroxy progesterone (17-OHP), dehydroepiandrosterone sulphate (DHEAS) and androstenedione measurement

MALES

The key initial investigation is semen analysis performed according to a recommended protocol.

NORMAL SEMEN ANALYSIS

No further biochemical assessment required

AZOOSPERMIA OR OLIGOZOOSPERMIA

Measure FSH, LH, prolactin and testosterone (ideally early AM (8 – 10 am) sample)

Significant patterns include:

- Elevated FSH (>12 U/L)/LH (>9 U/L) suggests primary gonadal failure
- Reduced FSH (<1.5 U/L)/LH (<1.5 U/L) plus low testosterone (typically <8 nmol/L) suggests hypothalamic or pituitary disease
- Prolactin >1000 mU/L requires assessment for pituitary tumour after exclusion of macroprolactinaemia and drug-related causes. The laboratory will advise on this. Milder elevations (>700 but <1000 mU/L) should be repeated to eliminate stress-related cause

REFERENCES

2. ACP Best practice No 170, J Clin Pathol 2003;56:261-267