Klinefelter’s syndrome
- A genetic condition affecting 1 in 650 men
- Due to the presence of an extra X chromosome (47XXY)
- Chromosomal mosaicism (both 47XXY and 46XY cells) occurs in 10%
  - Usually have milder signs and symptoms, depending upon the level of mosaicism
- The most common cause of androgen deficiency
- Characterised by:
  - Impaired testosterone production (androgen deficiency)
  - Impaired spermatogenesis (azoospermia)
- Up to 70% of cases remain undiagnosed
- Classical features may be present (Figure 1), however, there is a wide spectrum of signs and symptoms
- Small firm testes <4mls is the only consistent feature
  - Men will benefit from life-long testosterone treatment

Clinical notes: penile development may be normal or at the lower end of the normal range.

The GP’s role
- The clinical presentation may be subtle and its diagnosis overlooked unless actively considered
- Most males are diagnosed prenatally, during puberty or in association with infertility, or androgen deficiency
- The GP’s role includes clinical assessment, laboratory investigation, treatment, referral and follow-up

Clinical notes: the low detection rate (~30%) of Klinefelter’s syndrome would be improved if testicular examination became a regular part of a male physical examination.

Diagnosis

Medical history
- Pubertal development (poor progression)
- Sexual function (low libido)
- Degree of virilisation
- Psychosocial (learning, schooling, behaviour)
- Infertility

Examination
Infancy:
- No hormonal features prior to puberty
- Undescended testes

Adolescence:
- Small firm testes (<4ml) characteristic from mid puberty
- Poor pubertal progression and facial, body and pubic hair relative to age
- Taller than average height
- Poor muscle development

Adult:
- Small firm testes (<4ml) Reduced facial, body and pubic hair
- Gynaecomastia
- Taller than average height
- Poor muscle development

Refer to Clinical Summary Guides 1-3

Testicular Volume
Assessment of testicular volume is essential
- Testicular volume is assessed using an orchidometer
- Normal testicular volume range:
  - childhood <3mls
  - puberty 4–14mls
  - adulthood 15–35mls
- Small firm testes <4mls is the only consistent feature of Klinefelter’s syndrome (Figure 2)

Clinical notes: the testes may start to develop in early puberty, but soon regress to <4mls by mid puberty.

Refer to Clinical Summary Guide 1

Investigations
- Two morning samples of serum total testosterone, taken on different mornings
- Total serum testosterone, low or low normal from mid puberty (normal range 8–27 nmol/L)
- Serum LH, elevated from mid puberty (normal range 1-8 IU/L)
- Serum FSH, elevated from mid puberty (normal range 1-8 IU/L)
- Karyotype (47XXY)
- Bone density study, DEXA (osteoporosis)
- Semen analysis if fertility is an issue (usually azoospermic)
- TFT (hypothyroidism)
- Fasting blood glucose (diabetes)

Other investigations: 10% mosaic 46XY/47XXY
### Testosterone replacement therapy (TRT)

- TRT is life-long and may be started from mid puberty although many boys initially virilise normally
- Gynaecomastia is an indication to start TRT
- Teenage boys usually start on a low dose and build to full adult dose as puberty progresses
- Even if measured T levels are normal, there is evidence that bone density is reduced in the presence of chronically raised LH levels, suggesting that TRT is indicated

**Clinical notes:** in adults, consult with a fertility specialist (if appropriate) to develop a plan for fertility prior to TRT, as TRT will suppress spermatogenesis.

### Treatment options

<table>
<thead>
<tr>
<th>Injections (IM)</th>
<th>Dosage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sustanon®, Primoteston®</td>
<td>250mg every 2–3 weeks</td>
</tr>
<tr>
<td>Reandron®</td>
<td>1000mg every 12 weeks</td>
</tr>
<tr>
<td></td>
<td>(range: 10–14 weeks):</td>
</tr>
<tr>
<td></td>
<td>loading dose 6 weeks</td>
</tr>
<tr>
<td></td>
<td>after initiating therapy</td>
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<tr>
<td></td>
<td>(e.g. 0, 6, 18, 30 weeks)</td>
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<table>
<thead>
<tr>
<th>Subcutaneous implants</th>
<th>Dosage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Organon</td>
<td>3–4 x 200mg pellets</td>
</tr>
<tr>
<td></td>
<td>every 4–6 months</td>
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</table>

<table>
<thead>
<tr>
<th>Transdermal patch</th>
<th>Dosage</th>
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</thead>
<tbody>
<tr>
<td>Androderm®</td>
<td>5mg applied nightly</td>
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<table>
<thead>
<tr>
<th>Transdermal gel</th>
<th>Dosage</th>
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</thead>
<tbody>
<tr>
<td>Testogel®</td>
<td>50mg/5g applied daily</td>
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<table>
<thead>
<tr>
<th>Transdermal cream</th>
<th>Dosage</th>
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</thead>
<tbody>
<tr>
<td>Andromen®, Andromen Forte®</td>
<td>2% or 5% (2–6cm)</td>
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<tr>
<td></td>
<td>applied daily</td>
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</tbody>
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<table>
<thead>
<tr>
<th>Oral undecanoate</th>
<th>Dosage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Andriol Testocaps®</td>
<td>160–240mg in 2–3 divided doses daily</td>
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</tbody>
</table>

### Other treatments

- Gynaecomastia, this may be transient, lasting one to three years
- Adequate testosterone replacement often results in complete resolution over 12 months
- Surgical removal, mastectomy (do not refer for early surgery, as it may resolve naturally or following TRT)

### Follow-up

**Monitoring TRT is essential**

**Prostate:**
- Men with Klinefelter’s syndrome are less likely to die from prostate cancer, and restoring testosterone levels to the normal range is likely only to return their risks to those of their eugonadal peers
- Subject to the same advice about screening for prostate cancer as their peers (DRE and PSA)

**Clinical notes:** exclusion of significant prostate pathology is essential for those aged >40 years at the commencement of therapy.

### Raising clinical awareness

Aside from cognitive and behavioural features, it is important to note that despite the following recognised disease associations with Klinefelter’s syndrome the absolute risk is low.

- **Tumours:** leukaemia, lymphoma, germ cell tumour, teratoma, breast cancer
- **Endocrine:** hypothyroidism, diabetes mellitus (Type 1 and 2, rare)
- **Cardiovascular:** venous ulcers, venous thromboembolic disease
- **Auto-immune:** systemic lupus erythematosus (SLE), coeliac disease
- **Tumours:** mediastinal, breast cancer

Some features of Klinefelter’s syndrome are specific to the syndrome (e.g. behavioural and cognitive) and some features relate to the androgen deficiency (e.g. osteoporosis).

### Learning and behaviour difficulties

The general intellectual ability of boys with Klinefelter’s syndrome is within the normal range. However, boys with Klinefelter’s syndrome may have:
- Difficulties with speech and reading
- Delayed motor development
- Reduced attention span
- Behavioural problems (particularly in adolescence)

Educational and allied health assistance may be required.

### Infertility

Infertility is a major implication of Klinefelter’s syndrome
- Most men are azoospermic
- In some men, sperm can be found in the ejaculate or by testicular biopsy
- Treatment options
  - Intracytoplasmic Sperm Injection (ICSI) - the risk of 47XXY offspring is low
  - Donor insemination
  - Counselling may be necessary

**Refer to Clinical Summary Guide 5**