Klinefelter Syndrome

The Commonest Form of Hypogonadism, but Often Overlooked or Untreated

Eberhard Nieschlag

SUMMARY

Background: Klinefelter syndrome (KS) with the karyotype 47,XXY is one of the commonest types of congenital chromosomal disorder in males, with an incidence of 0.1% to 0.2% of newborn male infants. It causes hypogonadism and infertility. Until now, however, only about one-quarter of all persons with KS received the diagnosis during their lifetimes.

Methods: Selective review of the literature.

Results: KS is caused by aneuploidy of the sex chromosomes. Small, firm testes, the manifestations of androgen deficiency (sparse development of male-pattern body hair, greater than average height, lack of libido, erectile dysfunction) and, in more than 90% of affected men, azoospermia are its main features in adults. Affected boys may have verbalization difficulties and problems with learning and socialization. KS is often accompanied by other disturbances such as gynecomastia, varicose veins, thrombosis, osteoporosis, the metabolic syndrome, type 2 diabetes, bone fractures, epilepsy, and other neurological and mental disorders (4–6) (Table 1). This leads to a life expectancy 11.5 years below that of the male population as a whole (6).

As a result of this increased morbidity, Klinefelter syndrome patients require doctor and hospital treatment disproportionately often. However, a survey completed by 290 practising primary care physicians, internal medicine specialists, and urologists showed that two-thirds of primary care physicians and internal medicine specialists had had not knowingly seen any cases of Klinefelter syndrome in recent years, although their theoretical knowledge of it was good; urologists fared a little better regarding diagnosis of Klinefelter syndrome (7). On the basis of patient registries in Denmark, it is suspected that only 25% of all Klinefelter syndrome patients are diagnosed during their lifetimes (8). Nevertheless, suspected Klinefelter syndrome is easy to diagnose if physicians know the syndrome and thorough clinical examination is performed. Because knowledge of Klinefelter syndrome can only be derived from diagnosed cases, it is not known whether other cases remain undetected because they have a different range of symptoms. Furthermore, the known symptoms are not exclusive: Only one-quarter of patients in whom Klinefelter syndrome was suspected following clinical examination in a specialized facility actually showed a corresponding karyotype (9).

This article aims to attract greater medical attention to this important syndrome, in order to provide more patients with appropriate treatment. It is based on a selective search of the literature using a regular PubMed search over a 40-year period of clinical and scientific consideration of Klinefelter syndrome.

Pathophysiology

The disease pattern is caused by congenital aneuploidy of the sex chromosomes. 80% of patients have the Klinefelter syndrome 47,XXY was first described 70 years ago (1). With an incidence of 0.1% to 0.2% of male neonates (i.e. 1 to 2 per 1000), it is one of the commonest congenital chromosome disorders resulting in hypogonadism and genetically-determined infertility (2, 3).

Klinefelter syndrome is associated with a significantly higher morbidity rate compared to the male population as a whole. The main associated disorders are varicose veins, thrombosis, embolism, type 2 diabetes, bone fractures, epilepsy, and other neurological and mental disorders (4–6) (Table 1). This leads to a life expectancy 11.5 years below that of the male population as a whole (6).

Cite this as:
TABLE 1

<table>
<thead>
<tr>
<th>Comorbidities in Klinefelter syndrome: prevalence and mortality</th>
<th>Incidence %</th>
<th>Reference</th>
<th>Mortality (SMR*)</th>
<th>Reference</th>
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<tr>
<td>Gynecomastia</td>
<td>38</td>
<td>(2)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Breast cancer</td>
<td>0.3</td>
<td>(e7)</td>
<td>29 (e7)</td>
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</tr>
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<td>Thrombosis</td>
<td>4.7</td>
<td>(5)</td>
<td>8 (4)</td>
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<td>Pulmonary embolism</td>
<td>2.3</td>
<td>(5)</td>
<td>6 (4)</td>
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<tr>
<td>Metabolic syndrome</td>
<td>44</td>
<td>(14)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Type 2 diabetes</td>
<td>10</td>
<td>(14)</td>
<td>6 (4)</td>
<td></td>
</tr>
<tr>
<td>Osteopenia</td>
<td>40</td>
<td>(18)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Osteoporosis</td>
<td>10</td>
<td>(18)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Hip fracture</td>
<td>?</td>
<td>–</td>
<td>39 (4)</td>
<td></td>
</tr>
<tr>
<td>Maldescended testes</td>
<td>27</td>
<td>(2)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Mediastinal tumors</td>
<td>0.4</td>
<td>(5)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Epilepsy</td>
<td>5.5</td>
<td>(5)</td>
<td>7 (4)</td>
<td></td>
</tr>
<tr>
<td>Mental retardation</td>
<td>4.2</td>
<td>(5)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Delayed verbal development</td>
<td>40</td>
<td>(19)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Language disorder</td>
<td>70 to 80</td>
<td>(21)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Legasthenia</td>
<td>50 to 70</td>
<td>(21)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Learning difficulties</td>
<td>75</td>
<td>(19)</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

*SMR: standardized mortality rate, i.e. actual vs. predicted deaths (4)

Comorbidities in Klinefelter syndrome: prevalence and mortality

47,XXY karyotype. The remaining 20% have either mosaic 47,XXY/46,XY (i.e. different karyotypes in different cells), supernumerary X chromosome aneuploidy (48,XXX; 49,XXXXY), one or several additional Y chromosomes (e.g. 48,XXYY), or structurally abnormal additional X chromosomes (10, 11). These numerical chromosome abnormalities are the result of nondisjunction in maternal oogenesis in approximately two-thirds of cases, and in paternal spermatogenesis in the remaining third.

Although primordial germ cells (stem cells) are present in the testes of patients with Klinefelter syndrome, they degenerate unusually quickly, with the result that by puberty there are few or no remaining germ cells and very few remaining seminiferous tubules with complete spermatogenesis (12). The hyperplastic Leydig cells are unable to produce sufficient testosterone. This leads to testosterone deficiency, which can be modified by androgen receptor polymorphism and the severity of which varies between individual patients (13).

Ultimately almost all organ systems are associated with an elevated risk of morbidity and mortality in Klinefelter syndrome (4, 5) (Table 1). Gynecomastia, which is common in Klinefelter syndrome (occurring in 38% of our patients), is accompanied by a slightly higher incidence of breast cancer than in men with normal karyotype, who in total account for barely 1% of all breast cancers. In addition to increased frequency of breast cancer, mediastinal nonseminomatous germ cell tumors are also more common, most commonly between the ages of 15 and 30 years. Osteoporosis results in an increased incidence of bone fractures; femoral fractures are associated with a high mortality rate. Patients with Klinefelter syndrome suffer from vascular diseases, particularly varicose veins and thromboembolism, most commonly pulmonary embolisms. Central obesity with reduced glucose tolerance is often observed and can lead to metabolic syndrome and type 2 diabetes (14) (Figure 1). Epilepsy and other neurological and mental disorders occur significantly more frequently in Klinefelter syndrome patients.

Whether testosterone deficiency alone is responsible for this increased morbidity and mortality is doubtful, as a historical analysis of singers castrated before puberty, i.e. men with extremely low testosterone levels, showed the same life expectancy as for eugonadal singers (15). One risk factor may be the usually reduced arterial diameter of patients with Klinefelter syndrome, which leads to reduced perfusion of organ systems (16).

**Disease pattern**

Leading symptoms are signs of testosterone deficiency; very small, firm testes; and infertility (Box). Klinefelter syndrome patients can be reliably distinguished from eugonadal men by their small testes, which is always grounds for suspecting Klinefelter syndrome (Figure 2). They often display symmetrical gynecomastia with significantly palpable glands, and tall stature with long legs and a short trunk. However, unlike other forms of prepubertal-onset hypogonadism such as Kallman syndrome, in Klinefelter syndrome arm span does not exceed height. The severity of testosterone deficiency can vary greatly; thus the clinical phenotype can range from almost eugonadal to severe hypogonadism. While most patients pass through puberty with only mild symptoms and achieve normal penis size, approximately 70% of patients complain of falling libido and potency from the age of 25 years (17). Testosterone deficiency can also cause mild anemia. Beard growth and other secondary sexual hair growth are often sparse. Doctor’s certificates are often issued for lumbar pain and other musculoskeletal complaints resulting from the onset of osteopenia and osteoporosis (3, 18).

There are no symptoms that reliably indicate Klinefelter syndrome in prepubertal patients. Maldescended testes is present in one-quarter of cases, significantly more frequently than in boys in the population as a whole. Intellectual capacity may develop completely normally, but mild impairment of cognitive development is found in 4.2% of cases. Boys with Klinefelter syndrome may suffer from verbal and cognitive deficiencies (19) (Table 1) and often develop learning difficulties (75%). There may be attention deficit, and development of social skills is problematic. Klinefelter syndrome may suffer from verbal and cognitive deficiencies (19) (Table 1) and often develop learning difficulties (75%).
syndrome patients often achieve lower performance and professional status than their relatives (19–21). Evidence of learning disabilities has also been found in mice with Klinefelter syndrome, which means the phenomenon can be researched experimentally (22).

Many Klinefelter syndrome patients consult their doctors—and obtain a diagnosis—as a result of difficulties in becoming fathers. Azoospermia is present in more than 90% of cases, while in less than 10% of cases some or all sperm have reduced motility and morphology (2). Histological analysis of the testes reveals hyalinizing fibrosis of the seminiferous tubules, aspermatogenesis or focal spermatogenesis only, and hyperplasia of the Leydig cells. Spontaneous fatherhood in patients with Klinefelter syndrome is extremely rare.

**Diagnosis**

Suspected Klinefelter syndrome can be diagnosed as a result of thorough physical examination (paying attention to signs of testosterone deficiency and, in particular, testicle size) (Figure 2) or of chance observation during treatment of concomitant diseases (Table 1). Testicular volume is measured by ultrasound examination, which usually also shows testicular hypoechoegenicity. Smear testing can be used to look for Barr bodies in the buccal epithelium, corresponding to the inactive supernumerary X chromosome. Although no longer frequently used, this low-cost screening procedure has high specificity (95%) and adequate sensitivity (82%) (9). Diagnosis is finally confirmed cytogenetically, by karyotyping (Box).

Laboratory tests can reveal mild anemia as a result of testosterone deficiency. A marked increase in serum FSH levels is particularly characteristic of Klinefelter syndrome. Testosterone levels, which should always be determined in the morning due to fluctuations during the course of a day, may be either within normal range or abnormally low. For borderline cases in particular, it is recommended that sex hormone-binding globulin (SHBG) levels be determined and used with the total testosterone level to calculate free testosterone, which is a more sensitive indicator than total testosterone. Elevated LH values correlate with low testosterone but often remain elevated even after the testosterone level has risen to within normal range (Figure 3).

Bone density measurement by DXA (18), or inexpensively by phalanx osteosonography, is indicated in order to determine the extent and progression of any osteoporosis (3).

Semen analysis must be performed according to WHO guidelines. Azoospermia can only be reliably diagnosed following thorough examination of the sediment of centrifuged semen (23).

**Treatment**

In adults with Klinefelter syndrome, testosterone substitution should be used liberally, as this is generally thought to increase quality of life and prevent long-term effects. In hypogonadal men in general, rather
Testosterone deficiency should be treated only with natural testosterone in preparations that induce physiological serum levels. Of the wide variety of pharmaceutical forms available (3, 26), the commonest are intramuscular depot injections and transdermal gels. Complete substitution, which requires only four injections per year and achieves normal serum levels, involves intramuscular oil-based depot injections of 1000 mg testosterone undecanoate; however, this requires initial saturation, so the second injection is given six weeks after the first, and subsequent injections at 12-week intervals (27). This form of depot substitution has replaced the previously standard injections of 250 mg testosterone enanthate at two- to three-week intervals; the latter lead to major fluctuations in patients’ general condition as a result of fluctuating testosterone kinetics (26). There are various testosterone gels on the market, varying in testosterone concentration and composition. The higher a gel’s testosterone concentration, the smaller the quantity that must be applied once daily to the abdomen or upper arm to achieve normal serum levels (28, 29). Drug selection should take account of the patient’s preferences. In patients in whom endogenous production remains relatively high, however, low-dose, short-acting drugs should be preferred, possibly administered at prolonged intervals.

Essentially, testosterone therapy should be started when clinical symptoms so require and serum levels fall below the normal values of 12 mmol/L total testosterone and/or 250 pmol/L free testosterone (30, 31).
This occurs at very different ages in individual Klinefelter syndrome patients and thanks to regular annual checkups should not go unnoticed by physicians. In most patients it occurs between the ages of 20 and 30 years.

More recent research indicates that there are threshold values for various symptoms and groups of symptoms of testosterone deficiency; for example, loss of libido and drive occurs when levels drop below 15 nmol/L, while hot flashes and erectile dysfunction do not occur until levels fall below 8 nmol/L. Although these figures were obtained in patients with late-onset hypogonadism, they can be applied to other forms of hypogonadism (30, 32). In the future, pharmacokinetic issues that take account of androgen receptor polymorphism will also play a role (27).

Testosterone therapy must be accompanied by regular checkups that include inquiries into wellbeing, strength, and sexual function and examination of general condition, red blood cell count (excessively high doses lead to a risk of polycythemia), bone density, prostate, and PSA (3, 33) (Table 2). It must be considered lifelong substitution. In Klinefelter syndrome patients over the age of 50 years, guidelines on the treatment of late-onset hypogonadism should be followed (34); measurement of PSA levels and digital prostate examination are as important as for other men.

It is difficult to decide whether testosterone substitution should begin at puberty, as there are only a small number of studies on this question, and no controlled trials. Some researchers’ experience supports substitution as early as possible, as this achieves better physical, psychological, and scholastic development and social integration (35, 36). In individual cases it has been observed that criminal tendencies disappear in adolescent patients with Klinefelter syndrome receiving testosterone substitution (case studies in [35, 37]). Remaining endogenous production should not be suppressed, so intermittent treatment with short-acting testosterone preparations should be considered. Depot substitution would also suppress any remaining spermatogenesis, so later fertility must be taken into account even at this early age (38, 39). The decision must always be made on an individual basis until evidence-based recommendations can be given. Psychological support and speech therapy are also advisable.

Testosterone substitution has almost no effect on gynecomastia, and antiestrogens and aromatase inhibitors are only beneficial in isolated cases. However, no controlled trials are available. If gynecomastia is cosmetically problematic, mastectomy should be performed by surgeons with experience in this area.

Treatment for infertility has changed drastically in the last 15 years, although spontaneous pregnancy in partners of Klinefelter syndrome patients remains extremely rare. Also, to date only a few pregnancies have been reported from sperm taken from the semen of Klinefelter syndrome patients via ICSI (intracytoplasmic sperm injection). However, pregnancy is achieved more frequently using TESE (testicular sperm extraction) followed by ICSI treatment (2, 40). Individual seminiferous tubules may contain sperm that can be obtained by biopsy and introduced directly into the egg cell (ICSI). This achieves a higher success rate if microTESE, which in recent years has become established, is used in preference to conventional TESE, as shown by comparative reviews from 2004 (2) and 2011 (40). Pregnancy and live birth were achieved in approximately half the Klinefelter syndrome patients in whom TESE was attempted and whose data have been published to date (40).

Some groups have reported positive outcomes following hCG and/or antiestrogen pretreatment (38, e1). However, these trials were not controlled, and it remains questionable whether such pretreatment really has an effect, as controlled, double-blind trials in other types of male fertility disorders have shown no positive effects (e2). There is also evidence that sperm are more likely to be found in the testes of younger Klinefelter syndrome patients than those of older patients (39, e3, e4). Cryopreservation of testicular biopsy material of pubertal boys with Klinefelter syndrome is therefore being considered (e5).

In any event, if patients with Klinefelter syndrome wish to have children the initial step is to interrupt testosterone substitution, as testosterone suppresses residual spermatogenesis. In patients already receiving testosterone substitution, it must be interrupted for at least three to six months to allow remaining spermatogenesis to recover. The few spermatogonia produced in the testes of patients with Klinefelter syndrome seem to have a 46,XY chromosome set, unlike somatic cells (11, e6). This explains why most children of fathers with Klinefelter syndrome conceived in this way to date have normal karyotypes (2, 11, 40). However, it should be explained to parents that embryos and children have an increased risk of aneuploidy, not only

### TABLE 2

<table>
<thead>
<tr>
<th>Clinical parameters</th>
<th>Laboratory parameters</th>
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<tbody>
<tr>
<td>General wellbeing</td>
<td>Serum testosterone levels</td>
</tr>
<tr>
<td>Drive and mood</td>
<td>(Serum LH and FSH)</td>
</tr>
<tr>
<td>Libido and sexual activity (erections, ejaculations, intercourse)</td>
<td>RBC/hemoglobin/Hc/lipids/fibrinolysis</td>
</tr>
<tr>
<td>Body weight/BMI</td>
<td>Ejaculate volume</td>
</tr>
<tr>
<td>Bodily proportions</td>
<td>Prostate size and consistency</td>
</tr>
<tr>
<td>Hair growth</td>
<td>PSA (over 50 years)</td>
</tr>
<tr>
<td>Sebum production</td>
<td>Bone density</td>
</tr>
<tr>
<td>Voice</td>
<td></td>
</tr>
</tbody>
</table>

*Modified according to (33)

LH: luteinizing hormone; FSH: follicle-stimulating hormone; PSA: prostate-specific antigen; Hc: hematocrit
of sex chromosomes but also of chromosomes 18 and 21. Invasive genetic prenatal testing is therefore recommended.

Overall, treatment of infertility in patients with Klinefelter syndrome remains only partly successful, and counseling provided for couples should not raise their hopes too high.

Conflict of interest statement
Prof. Nieschlag has received fees for a Springer textbook on andrology and a textbook on testosterone from Cambridge University Press. He has also received reimbursement of conference participation fees and travel and accommodation expenses from the Association of German Internal Medicine Specialists (BDI, Berufsverband der Deutschen Internisten).

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REFERENCES


KEY MESSAGES

- Klinefelter syndrome is one of the commonest forms of chromosome aneuploidy and combinom form most common form of hypogonadism and genetically-determined infertility.

- Although diagnosis is easy (small testicular volume), Klinefelter syndrome still often remains overlooked or untreated.

- Children and adolescents with Klinefelter syndrome can suffer from impaired verbal development, learning difficulties, and problems in the development of social skills.

- Klinefelter syndrome is associated with a high comorbidity rate and increased mortality.

- Testosterone substitution and prevention of comorbid disorders are the most important elements of treatment. Nowadays some patients can become fathers via TESE/CSI.


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For eReferences please refer to: www.aerzteblatt-international.de/ref2013
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