To Be or Not To Be … Male

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Introduction:
Pats. with Klinefelter Syndrome (KS) have elevated morbidity and mortality due to several reasons. Yet, there is no connection between KS and male-to-female (MTF) sex change.

Case Report
A 74-year old man was sent for endocrine work-up prior to surgery due to gynecomastia. The patient had lived as a MTF transgenger for many years. He was told not to qualify for a transgender surgical approach when he was 55 years old. He took estradiol substitution for several years, but by age 60 after developing recurrent severe pulmonary embolism he stopped substitution. After that he lived “asexual” and bilateral, non-painful gynecomastia developed (or became disturbing). Examination revealed small testes (Picture 1), bilateral Gynecomastia (Tanner IV, Picture 2) and signs of chronic venous insufficiency. The biochemical analyses showed hypergonadotropic hypogonadism with otherwise normal values. The molecular analyses revealed a 47XXY Karyotype. Osteodensitometry showed low peak bone mass (Picture 3). We started topical testosterone replacement and calcium/vitamin D3 substitution.

Discussion:
KS affects about 1 in 660 men and is the most frequent chromosomal aberration in males, but remains often undetected with only about 25% of patients receiving the correct diagnosis. Age at diagnosis is around 35 yrs. The phenotype is thought to be linked to non-inactivated genes from the extra X-chromosome, but alternative mechanisms are possible. The excess morbidity and mortality (expected lifespan reduced by 1.5-2 yr) may be explained by endocrine dysfunction and diseases of the cardiovascular/respiratory systems and malignancies. Besides there are some socioeconomic differences to the healthy population (shorter education, lower income etc.) also an increased risk for criminal activity was found (crime rate for sexual abuse and arson). Only few studies have examined the association of KS and transsexuality. The prevalence of chromosomal abnormalities and disorders of sexual development (DSD) in MIF individuals varies between 1.11’000 to 1.45’000. In a retrospective study in 83 MIF only 1 was found with primary hypogonadism (0.5%) and normal karyotype (Figure 1), one was found to be KF. The study by Fisher et al. showed besides other findings a higher rate of Gender Dysphoria in KS, which was mediated by the presence of autistic traits. Yet, no significant increase in KS was reported in previous studies looking at MIF transgender populations. Still, whether or not KS is associated with MIF has to our knowledge not systematically been investigated.

Conclusion
In MIF Individuals further work-up towards KS should be envisaged if there is additional clinical suspicion – such as clues from patient history, clinical examination (small testes) or lab work (hypergonadotropic hypogonadism) as found in our case.

References
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