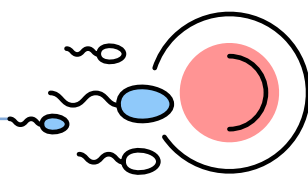


BÖLÜM 5

ERKEK ÜREME SİSTEMİ EMBRİYOLOJİSİ



Tuğba BAL TAŞTAN¹

GİRİŞ

GENİTAL SİSTEM GELİŞİMİ

Genital sistemin gelişimi, bir bireyin tüm cinsel farklılaşmasının bir aşamasıdır. Cinsiyetin belirlenmesi, döllenme sonucu, oositte mevcut olan X kromozomuna bir Y kromozomu veya bir X kromozomunun eklenmesi ile başlar. Bu aşama, cinsiyetin genetik olarak belirlenmesini temsil eder. Embriyonun genetik cinsiyeti döllenme sırasında kesin olmasına rağmen, embriyonun bütün fenotipik cinsiyeti ve gonadlar gelişimin yedinci haftasına kadar benzerdir ve **farklılaşmamış gonadlar** olarak tanımlanırlar. Mezodermal epitel, Embriyonik bağ dokusu ve primordial germ hücreleri testis ve overlerin (gonadlar) kaynağını oluşturmaktadır. 7. Hafadan önce, embriyo cinsiyetinin temel morfolojik göstergesi, dışında bulunan X kromozomunun inaktivasyonu sonucu oluşan cinsiyet kromatininin (**Barr cismi**) varlığı veya yokluğuudur. Cinsel gelişimin morfolojik olarak farklılaşmamış bu aşamasında, gametler vitellüs kesesinden gonadal primordiya göç eder.

Cinsiyetin fenotipik farklılaşmasının gonadlarla başladığı ve genital kanal sistemleri üzerindeki gonadal etkilerle ilerlediği kabul edilir. Dış genital organların

¹ Arş. Gör. Dr., Erzincan Binali Yıldırım Üniversitesi, Tıp Fakültesi, Histoloji ve Embriyoloji AD., tugba-bal07@gmail.com

- Bir birey sadece X kromozomu taşıyorsa bu durumda Turner' sendromu gelişir. Bazı bireyler çizgi gonad oluşumları ile dışı fenotipine sahip olurlar. Gerçek hermafroditizm veya psödohermafroditizm durumlarında çeşitli sebepler etkilidir. Testiküler feminizasyonda genetik olarak testosteron reseptörlerinin eksikliği mevcuttur. Bu durumda bireylerde dışı fenotipi izlenir. Genital kanalların majör anormallikleri ise çok nadir görülür.

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