



BÖLÜM 6

RENAL EMBRİYONAL NEOPLAZİLER

6.1 NEFROJENİK TÜMÖRLER

6.1.1 NEFROJENİK KALINTILAR

Yasemin YUYUCU KARABULUT¹

6.1.1.1. Tanım

- ▶ Nefrojenik kalıntılar, fetal böbrek gelişimi sırasında tamamen gerilemeyen ve nefroblastom ile ilişkili olabilen immatür renal dokulardır.

6.1.1.2. Köken/Patogenezi

- ▶ Prenatal böbrek gelişimi sürecinde diferansiyasyonun tamamlanamaması sonucunda oluşurlar (1). Özellikle Wilms tümörü gelişme riski taşırlar.

6.1.1.3. Epidemiyoloji

- ▶ Prematüre doğumlarda daha sık görülür.

6.1.1.4. Semptom ve Bulgular

- ▶ Çoğunlukla asemptomatik olup, rastlantısal olarak tespit edilir.
- ▶ Nadiren hipertansiyon ve hematüri görülebilir.

6.1.1.5. Radyoloji

- ▶ USG: Küçük, hipoekoik odaklar olarak izlenebilir.
- ▶ BT/MRG: Minimal kontrast tutan küçük nodüler lezyonlar şeklinde görülebilir.

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