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Eğitim Bilgileri

Tıpta Uzmanlık, İstanbul Üniversitesi, Tıp Fakültesi, Türkiye 1984 - 1989

Yaptığı Tezler

Tıpta Uzmanlık, Asfiksili term yenidoğanlarda böbrek fonksiyonlarının incelenmesi, İstanbul Üniversitesi, İstanbul Tıp Fakültesi, 1989

Araştırma Alanları

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Akademik Unvanlar / Görevler

Prof.Dr., Marmara Üniversitesi, Tıp Fakültesi, Dahili Tıp Bilimleri Bölümü, 2004 - Devam Ediyor

Verdiği Dersler

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