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Hakemli Kongre / Sempozyum Bildiri Kitaplarında Yer Alan Yayınlar

- **A novel missense variant in the homogentisate 1,2-dioxygenase (HGD) gene in a patient with clinical symptoms of alkaptonuria.**
KAZIMLI U., DOĞAN M. E. , BAYSAL K., ÖZKUL Y., ŞENEL S., DÜNDAR M.
13th Balkan Congress of Human Genetics, Edirne, Türkiye, 17 - 20 Nisan 2019, cilt.22, ss.166
- **Clinical characteristics of pulmonary arterial hypertension associated with congenital heart disease: baseline results from the Turkish congenital heart disease-associated pulmonary arterial hypertension (THALES) registry.**
Sagin Saylam G., Kucukoglu S. N. , Alehan D., Kula S., Kaymaz C., Akcevin A., Cicek S., KAYA M. G. , Karademir S., Varan B., et al.
46th Annual Meeting of the Association for European Paediatric and Congenital Cardiology, İstanbul, Türkiye, 23 - 26 Mayıs 2012, ss.1