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Naslov	URL	Autori	Naslov izvornika
Genotype-phenotype correlation in contactin-associated protein-like 2 (CNTNAP-2) developmental disorder		D'Onofrio, Gianluca; Accogli, Andrea; Severino, Mariasavina; Caliskan, Haluk; Kokotović, Tomislav; Blažeković, Antonela; Gotovac Jerčić, Kristina; Markovic, Silvana; Žigman, Tamara; Goran, Krnjak; Barišić, Nina; Duranovic, Vlasta; Ban, Ana; Borovečki, Fran; Petković Ramadža, Danijela; Barić, Ivo; Fazeli, Walid; Herkenrath, Peter; Marini, Carla; Vittorini, Roberta; Gowda, Vykuntaraju; Bouman, Arjan; Rocca, Clarissa; Alkhawaja, Issam Azmi; Murtaza, Bibi Nazia; Rehman, Malik Mujaddad Ur; Al Alam, Chadi; Nader, Gisele; Mancardi, Maria Margherita; Giacomini, Thea; Srivastava, Siddharth; Alvi, Javeria Raza; Tomoum, Hoda; Matricardi, Sara; Iacomino, Michele; Riva, Antonella; Scala, Marcello; Madia, Francesca; Pistorio, Angela; Salpietro, Vincenzo; Minetti, Carlo; Rivière, Jean-Baptiste; Srouf, Myriam; Efthymiou, Stephanie; Maroofian, Reza; Houlden, Henry; Vernes, Sonja Catherine; Zara, Federico; Striano, Pasquale; Nagy, Vanja	
SNCA 3' UTR Genetic Variants in Patients with Parkinson's Disease		Blažeković, Antonela; Gotovac Jerčić, Kristina; Borovečki, Fran	
A first report of a rare TP53 variant associated with Li-Fraumeni syndrome manifesting as invasive breast cancer and malignant solitary fibrous tumor		Prejac, Juraj; Dedić Plavetić, Natalija; Gotovac Jerčić, Kristina; Borovečki, Fran	
Clinical and pathohistological characteristics of Alport spectrum disorder caused by COL4A4 mutation c.193-2A>C: a case series		Šenjug, Petar; Nikuševa Martić, Tamara; Šenjug Perica, Marija; Oroz, Maja; Horaček, Matija; Gotovac Jerčić, Kristina; Galešić, Krešimir; Galešić Ljubanović, Danica	
Congenital factor XI deficiency caused by a novel F11 missense variant: a case report		Gotovac Jerčić, Kristina; Blažeković, Antonela; Hančević, Mireia; Bilić, Ervina; Borovečki, Fran	

The influence of dopamine-beta-hydroxylase and catechol O-methyltransferase gene polymorphism on the efficacy of insulin detemir therapy in patients with type 2 diabetes mellitus		Božek, Tomislav; Blažeković, Antonela; Nikolac Perkovic, Matea; Gotovac Jerčić, Kristina; Šustar, Aleksandra; Smirčić- Duvnjak, Lea; Outeiro, Tiago; Pivac, Nela; Borovečki, Fran	
Complex intrachromosomal rearrangement in 1q leading to 1q32.2 microdeletion: a potential role of SRGAP2 in the gyrification of cerebral cortex		Rinčić, Martina; Radoš, Milan; Krsnik, Željka; Gotovac, Kristina; Borovečki, Fran; Liehr, Thomas; Brečević, Lukrecija	
Genetic changes observed in a case of adult pilocytic astrocytoma revealed by array CGH analysis		Pećina-Šlaus, Nives; Gotovac, Kristina; Kafka, Anja; Tomas, Davor; Borovečki, Fran	
Aberrant expression of shared master-key genes contributes to the immunopathogenesis in patients with juvenile spondyloarthritis		Lamot, Lovro; Borovečki, Fran; Tambić Bukovac, Lana; Vidović, Mandica; Perica, Marija; Gotovac, Kristina; Harjaček, Miroslav	
Molekularni učinci proteina alfa sinukleina u jezgri: vezanje na molekulu DNA i regulacija transkripcije		Gotovac, Kristina	
Pilot study of the association between the HLA region and testicular carcinoma among Croatian patients		Gotovac, Kristina; Grubić, Zorana; Kaštelan, Željko; Štingl, Katarina; Kuliš, Tomislav; Krhen, Ivan; Hudolin, Tvrtko; Kaštelan, Maja; Žunec, Renata	