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Structural Modifications Introduced by NS2B Cofactor Binding to the NS3 Protease of the Kyasanur Forest Disease Virus		Kandagalla, Shivananda; Kumbar, Bhimanagoud; Novak, Jurica	
CYP3A4		Novak, Jurica	
Can Resveratrol Influence the Activity of 11 β - Hydroxysteroid Dehydrogenase Type 1? A Combined In Silico and In Vivo Study		Novak, Jurica; Tseilikman, Vadim E.; Tseilikman, Olga B.; Lazuko, Svetlana S.; Belyeva, Lyudmila E.; Rahmani, Azam; Fedotova, Julia	
The design of compounds with desirable properties – The anti-HIV case study		Novak, Jurica; Pathak, Prateek; Grishina, Maria A.; Potemkin, Vladimir A.	
Computational and experimental therapeutic efficacy analysis of andrographolide phospholipid complex self-assembled nanoparticles against Neuro2a cells		Mourya, Atul; Pingle, Purva; Babu, Chanti Katta; Veerabomma, Harithasree; Sainaga Jyothi, Vaskuri G.S; Novak, Jurica; Pathak, Prateek; Grishina, Maria; Verma, Amita; Kumar, Rahul; Singh, Pankaj Kumar; Khatri, Dharmendra Kumar; Singh, Shashi Bala; Madan, Jitender	
Structural modifications introduced by NS2B cofactor binding to the NS3 protease of the Kyasanur forest disease virus		Novak, Jurica	
Gaussian field-based 3D-QSAR and molecular simulation studies to design potent pyrimidine-sulfonamide hybrids as selective BRAFV600E inhibitors		Singh, A. K.; Novak, Jurica; Kumar, A.; Singh, H.; Thareja, S.; Pathak, Prateek; Grishina, M.; Verma, V.; Yadav, J. P.; Khalilullah, H.; Pathania, V.; Nandanwar, H.; Jaremko, M.; Emwas, A.-H.; Kumar, P.	
Računalna studija mehanizmom temeljenih ireverzibilnih inhibitora monoamin oksidaze B		Vrban, Lucija	
Analysis of ADAR protein during productive HSV-1 Infection		Echeta, Justina Oluchi	
Lijekovi protiv herpes simpleks virusa 1 i 2		Krišto, Ruža	
Withasomniferol C, a new potential SARS-CoV-2 main protease inhibitor from the Withania somnifera plant proposed by in silico approaches		Kandagalla, Shivananada; Rimac, Hrvoje; Krishnamoorthy, Gurushankar; Novak, Jurica; Grishina, Maria; Potemkin, Vladimir	
Molekulsko dinamičke simulacije retinala u otopini		Novak, Jurica	
Kiralne stacionarne faze		Novak, Jurica	

Exploring potential inhibitors against Kyasanur forest disease by utilizing molecular dynamics simulations and ensemble docking		Kandagalla, Shivananda; Novak, Jurica; Shekarappa, Sharath Belenahalli; Grishina, Maria A; Potemkin, Vladimir A; Kumbar, Bhimanagoud	
Dinamika modelnih bioloških sustava u osnovnom i pobuđenim elektronskim stanjima		Novak, Jurica	