

Full Publication list

Articles:

Martins-Silva, T; Salatino-Oliveira, A; Genro, JP; Meyer, FDT.; Li, Y; Rohde, LA; **Hutz, MH**; Tovo-Rodrigues, L. 2021. Host genetics influences the relationship between the gut microbiome and psychiatric disorders. **Progress in Neuro-Psychopharmacology & Biological Psychiatry**. **106**, 110153 DOI 10.1016/j.pnpbp.2020.110153

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Botton, MR.; Viola, PP.; Meireles, MR.; Bruxel, EM.; Zuchinali, P; Bandinelli, E; Rohde, LE.; Leiria, Tiago LL.; Salamoni, JYY.; Garbin, AP.; **Hutz, MH.** 2020. Identification of environmental and genetic factors that influence warfarin time in therapeutic range **Genetics and Molecular Biology** **43**, e20190025

Silva, CA.; Fernandes, DCRO.; Braga, ACO.; Cavalcante, GC.; Sortica, VA.; **Hutz, MH.**; Leal, DFVB; Fernandes, MR.; Santana-da-Silva, MN.; Lopes Valente, SE.; Pastana, LF.; Pinto, PDC.; Costa, GE.; Ribeiro-dos-Santos, A.; Santos, S.; Santos, NPC. 2020. Investigation of genetic susceptibility to Mycobacterium tuberculosis (VDR and IL10 genes) in a population with a high level of substructure in the Brazilian Amazon region. **International Journal of Infectious Diseases**. **98**, 447-453

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