

## Articole publicate de membrii disciplinei în reviste cotate ISI

1. Tripon F, Iancu M, Trifa A, Crauciuc GA, Bogliș A, Dima D, Lazar E, Banescu C. Modelling the Effects of MCM7 Variants, Somatic Mutations, and Clinical Features on Acute Myeloid Leukemia Susceptibility and Prognosis. *J Clin Med*. 2020;9(1):158. Published 2020 Jan 8. doi:10.3390/jcm9010158
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#### **Articole publicate IN EXTENSO, în reviste indexate BDI**

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