V. Klochkov ¹
P. Vassiliev ^{1,2,3}
D. Babkov ^{1,2}

DIRECTED SEARCH FOR NOVEL A-GLUCOSIDASE INHIBITORS USING STRUCTURE SIMILARITY WITH TESTED SUBSTANCES

¹ Chair of pharmacology;

² Research Institute of Pharmacology;

klochkovvladlen@gmail.com

Type 2 diabetes mellitus is one of the most fast-spreading socially-important diseases of our days, so researches worldwide are oriented for antidiabetic drug discovery. One of the approaches to hyperglycemia pharmacocorrection is the inhibition of α -glucosidase.

In this regard, the target of the present study is computer-aided search for novel α glucosidase inhibitors by method of structure similarity with tested substances. The prediction was carried outusing the database containing the information about structure and α -glucosidase inhibition activity of 183 newly tested substances. The data about their maximum inhibiting activity (Δ %) in 1 mM concentration were subjected to clusterization by k-means method, and 3 classes of activity were defined: high (19 subst.), moderate (40 subst.), low (74 subst.).

In silico activity prediction for 695 new compounds was made with the program TestSim 17.01.28 from IT Microcosm software complex [1]. This utility employs the method of similarity with standards, based on calculation of QL-modified Tanimoto similarity index T [2]. For each substance T values were calculated for all tested substances from database. The maximum value T_{max} was determined, with the indication of code and activity level of the most structurally similar tested substance.

A total of 15 compounds with predicted high value and $T_{max}>0.6$ were tested in vitro by method[3] in 1 mM concentration. The reference drug was acarbose.

According to the experimental results, out of 15 promising predicted substances 10 were found to possess high α -glucosidase inhibition activity. Five compounds were more active than reference drug and another five have the same activity as acarbose. So, the prediction accuracy of the α -glucosidase inhibitory properties by structure similarity with tested substances was defined as 66,7%.

As a result, we can conclude, that this method can be used for *in silico* prediction of new α -glucosidase inhibitors per their structural similarity with earlier tested substances. The accuracy was 66,7%, and five newly identified active compounds were selected for the further detailed pharmacological evaluation.

The work is funded by the Russian Science Foundation, 14-25-00139 project.

³ Laboratory of Molecular Modeling and Computer Searchfor Drug Substances, Volgograd State Medical University, Pavshikh Bortsov Square, 1, 400131 Volgograd, Russia

^{1.} Vassiliev P.M., Kochetkov A.N. IT Microcosm. State Registration Certificate for software program 2011618547 (Russian), 2011, 31 Oct 2011.

^{2.} Vassiliev P.M. et al. In: Target-oriented search for antidiabetic agents, VSMU, 2016: 126-181.

^{3.} Nair S.S. et al. European Journal of Experimental Biology, 2013, 3(1):128-132.