Molecular modelling studies of 2-substituted-3-benzyl-6-iodo quinazolin4(3H)-one analogs targeting for breast cancer activity

S. Rajasekaran and Zonunsiami

Department of Pharmaceutical Chemistry, Al-Ameen College of Pharmacy, Near Lalbagh Main Gate, Hosur Road, Bangalore-560027, India

E-mail: srajasekaran@gmail.com

ABSTRACT
Quinazoline moieties are existing in many pills such as Erlotinib, Gefatinib, Lapatinib, Afatinib, Raltitrexed, which are used clinically in the most cancers therapy. In the existing study, we have utilized crew quantitative structure–activity relationships (G-QSAR) for exploring the relationship between the buildings of a rising household of quinazoline derivatives and their breast most cancers activities. Group-quantitative shape exercise relationship (GQSAR) used to be carried out the use of the approach of more than one linear regression (MLR). Reasonable and superb G-QSAR fashions have been developed, in order to useful resource in similarly optimization and improvement of more recent anticancer agents. Further, the compounds had been subjected to molecular docking to apprehend the prolong of binding interplay between the ligands and the chosen protein (3HB5 and 4CQ0).

Keywords: Quinazolinone, Anticancer, Molecular properties, G-QSAR, Molecular docking

INTRODUCTION
Cancer is a main motive of loss of life in the international and accounted for 7.6 million deaths (around 13% of all deaths) in 2008. The international burden of most cancers rises in 2012 to 14 million new instances per year, a discern expected to upward shove to 22 million yearly inside the subsequent two decades. Over the equal period, most cancers deaths are anticipated to upward push from an estimated 8.2 million yearly to thirteen million per year. Globally, in 2012 the most frequent cancers recognized had been these of the lung (1.8 million cases, 13.0% of the total), breast (1.7 million, 11.9%), and massive bowel (1.4 million, 9.7%). The most frequent most cancers demise had been cancers of the lung (1.6 million, 19.4% of the total), liver (0.8 million, 9.1%), and belly (0.7 million, 8.8%). The breast is made up of glands known as lobules that can make milk and skinny tubes known as ducts that elevate the milk from the lobules to the nipple. Breast tissue additionally incorporates fats and connective tissue, lymph nodes, and blood vessels. Breast most cancers is a most cancers that develops from breast tissues [3]. Signs of breast most cancers can also consist of a lump in the breast, a alternate in breast shape, dimpling of the skin, fluid coming from the nipple, or a purple scaly patch of skin [4]. In these with far-off unfold of the disease, there may additionally be bone pain, swollen lymph nodes, shortness of breath, or yellow skin. Breast most cancers typically starts off evolved off in the internal lining of milk ducts or the lobules that furnish them with milk. Breast most cancers that develops in the lobules is regarded as lobular carcinoma, whilst that develops from the ducts is referred to as ductal carcinoma.

Quinazoline and their analogs are one of the most substantial scaffolds amongst bioactive compounds. There is an growing range of biochemical objectives for quinazoline compounds and has a large vary of a huge vary of organic things to do such as anticancer, antimicrobial, antimalarial, anti-inflammatory, antihypertensive, anti-diabetic, cholinesterase inhibition, dihydrofolate reductase inhibition, and kinase inhibitory activity. There are a number of quinazoline containing tablets on hand in the market (eg : Erlotinib, Gefitinib, Lapatinib, Afatinib, Raltitrexed, etc) for treating cancer.

DISCUSSION
As shown in Table 3 and Figure 2,3 and 4 the model 1,2,3 indicate that the descriptors SdssCcount, SsSHE-index, chi4pathCluster contributes positively for the biological activity while the descriptors T_2_N_1 and T_C_S_5 contribute negatively for the anticancer activity. The compound QSR_3, 13, 11 were found to have good binding affinity for 3HB5 and 4CQ0. It was interesting to note that the compound QSR_3 that had good binding affinity also showed high miLogP and large TPSA, however the molecular weight of the compound was above Lipinski rule, the hydrogen bond donor was 3, which is large for the compounds of the series.

CONCLUSION
A collection of 3-benzyl-6-iodo-2-mercaptio-3H-quinazolin-4-one analogs have been subjected for the molecular houses prediction by using molinspiration software program in order to locate the drug likeness. Then, team based totally qsar learn about used to be carried out the use of search engine algorithm, random and guide decision techniques for the division of the statistics units into education and take a look at sets. Three fashions had been generated by way of adopting the MLR approach with stepwise ahead backward and additionally simulated annealing. Statistically massive G-QSAR fashions had been generated. Among them the first mannequin has squared correlation coefficient (r2 ), go validated correlation coefficient (q2 ) and predictive correlation coefficient (pred r2 ) 0.81, 0.59 and 0.62; respectively The 2nd mannequin has 0.83,0.55 and 0.53; respectively. And additionally, the 1/3 mannequin has 0.82, 0.61 and 0.54 respectively. R1-SdssCcount, R1-SsSHE-index, R1-SssNHcount at R1 role are the three descriptors which are proven to expand the organic endeavor whilst the R1-T_2_N_1 and R1- T_C_S_5 are the two descriptors which are diminished organic activity. From the existing G-QSAR analysis, out of the generated three fashions any one can be used for predicting the undertaking of the newly designed compounds in discovering some greater powerful molecules. Furthermore, the
molecular docking find out about was once carried out in opposition to the 3HB5 and 4CQ0 protein for breast cancer, in which the compounds substituted with the acetamide, dihydrotriazinone, tetrahydropyridazinedione, 5-methyl dihydro-pyrazol-3-one, dimethyl pyrazole confirmed true docking in opposition to the 3HB5 protein with proper hydrogen bond interaction, $\pi-\pi$ and $\pi$-cation interplay whereas substitution with hydrazide, oxadiazolethione, pyrroldine Dione, methyl thiourea, phenyl thiourea, benzoic acid hydrazide, benzylidene hydrazine, phenyl thiazolidinone, N-formimidic ethyl ester, formamide, oxadiazole, thiadiazol and imidazole confirmed appropriate docking rating however with lesser interaction. The amino acids which are worried in the bond formation between the ligands and the protein are threonine 140, glycine 141, leucine 162, glutamine 163, glutamine 185 and cysteine 185. In case of 4CQ0 protein the compounds substituted with acetamide, benzoic acid hydrazide, benzylidene, 5-methyl dihydro-pyrazol-3-one, dimethyl pyrazole, formamide confirmed exact docking with interactions. However, the last compounds confirmed appropriate binding with bad hydrogen bond interplay and $\pi-\pi$ interaction. Finally, it is concluded that the work introduced right here will play an necessary position in grasp the relationship of physiochemical parameters with shape and organic activity. By reading the molecular property, G-QSAR mannequin and molecular docking one can pick out the appropriate substituent for in addition synthesizing bioactive compounds displaying most potency.

Bottom Note: This work is partly presented at International Conference on Genetics & Molecular Biology September on 24-25, 2020, Webinar.