Cardiac arrest

Eleshin Peter Femi
University of Lagos, Nigeria

In this discussion, the author will examine whether the active practice of Cardiac Arrest in the fundamental human body will be expressly dissolved. The purpose of this research is to know how far the Cardiac Arrest can be known to everybody; the author examines this through the study academic investigations into the subject, and is studying anecdotal evidence. The method used in this research is quantitative data with correlation analysis. The techniques used to collect the data are: Expression, Description and Presentation. The researchers developed three topics in order to know the reality of effectiveness of the Cardiac Arrest program in the conference. Furthermore, Sudden Cardiac Arrest (SCA) is an electrical problem with the heart that triggers a dangerously fast heart rhythm (ventricular fibrillation). This rapid, irregular heart rhythm causes the heart to quiver rather than contract or pump. When the heart stops pumping blood, oxygen cannot reach the body and brain. If not treated immediately, SCA can be fatal.

peterson12311@gmail.com

3D-QSAR and molecular docking study of ursolic acid derivatives used as phospholipase-A2 inhibitors in cardiotoxin venom

Preetama Singh¹ and Pallavi Somvanshi²
¹Sam Higginbottom Institute of Agriculture, Technology and Sciences, India
²TERI University, India

Snake bite is a serious medical problem especially in the tropical and subtropical countries including India. Snake Venom is the most dangerous and toxic protein of everything secreted from snakes poison glands. Phospholipase-A2 or PLA2 is the most affective and dangerous one. This PLA2 is the membrane protein responsible for releasing Arachidonic acid which causes serious inflammatory affect over the victim. Ursolic acid which is a natural occurring terpinoid generally found in fruits has the anti-functional properties against this PLA2. Ursolic acid (3β-hydroxy-urs-12-en-28-oic acid) isolated from many medicinal plants and has diverse pharmacologically important properties, including strong anti-inflammatory effect. QSAR models can also be used in designing new chemical entities (NCEs) and identify the most promising hit compounds along with reduced number of costly experiments at low cost. In this work, molecular modeling and QSAR studies of a set of 34 compounds, ursolic acid derivatives as phospholipase A2 antagonists were performed. QSAR results indicate the proposed kNN-MFA 3D-QSAR model gave good statical result in term of (q²=0.8956) and coefficient (pred_r²=0.8756) and provided most significant co-relation of steric field descriptors with the biological activities, therefore these model can be able to increase the activities of non-peptidic phospholipase A2 inhibitors from further modification of known ursolic acid derivatives.

psomvanshi@gmail.com