# **QSAR model for CYP450 2C19 inhibitor** (v1.0)



### **ProtoADME**

ProtoADME is a computational (in silico) tool focused on the prediction of endpoints related with the ADME (Absortion, Distribution, Metabolism and Excretion) of chemical substances.

# **Endpoint**

#### Toxicokinetic: CYP450 2C19 inhibitor

The microsomal cytochrome P450 (CYP) family 4 monooxygenases are the major fatty acid omega-hydroxylases. These enzymes remove excess free fatty acids to prevent lipotoxicity, catabolize leukotrienes and prostanoids, and also produce bioactive metabolites from arachidonic acid omega-hydroxylation. In addition to endogenous substrates, recent evidence indicates that CYP4 monooxygenases can also metabolize xenobiotics, including therapeutic drugs. If a compound is a CYP inhibitor may decrease the metabolism of comedicated drugs.

#### **Metrics**

## **Training set**

QSAR predictions		
Non-inhibitor	Inhibitor	
124	19	
54	282	
	Non-inhibitor	

Validation set					
Experimental values	QSAR predictions				
	Non-inhibitor	Inhibitor			
Non-inhibitor	32	12			
Inhibitor	24	97			

Parameters	Training	Validation
Accuracy	0.85	0.78
Sensitivity / recall	0.84	0.80
Specificity	0.87	0.73
Precision	0.94	0.89
Negative predictive value	0.70	0.57
F-score	0.89	0.84
Matthews Correlation Coefficient	0.67	0.49
Critical Success Index	0.79	0.73
Area under the ROC	0.85	0.76



ProtoPRED platform allows the easy, fast and user-friendly prediction of different properties of chemical compounds, by proprietary (Q)SAR models.





