# QSAR model for boiling point (v1.0)





#### **ProtoPHYSCHEM**

ProtoPHYSCHEM is a computational (*in silico*) tool focused on the prediction of endpoints related with the physicochemical properties of chemical substances.

ProtoPHYSCHEM mainly includes, but is not limited to, endpoints used by REACH, a European Union regulation, adopted to improve the protection of human health and the environment from the risks that can be posed by chemicals, while enhancing the competitiveness of the EU chemicals industry.

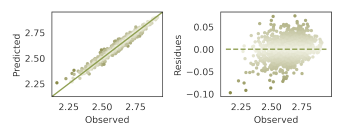
## **Endpoint**

## Physical-chemical properties: Boiling point.

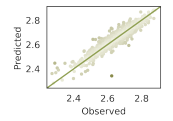
The boiling point is the temperature at which a substance's physical state changes from a liquid to a gas.

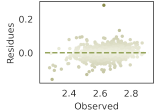
#### **Metrics**

### **Training set**









Parameters	Training	Validation
R <sup>2</sup> score	0.97	0.89
Mean absolute error (MAE)	0.01	0.02
Mean squared error (MSE)	0.00	0.00
Median absolute error	0.01	0.01
Explained variance	0.97	0.89

ProtoPHYSCHEM is part of



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



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