Formula: C 23 H 18 O 10

MW: 454.39

MDL: MFCD02940844

TNP:

LogP: -2.02

LogS: -2.29

Acceptors: 10

Donors: 0

Rotation Bonds: 9

Chiral Centers: 0
$\mathrm{N}+\mathrm{O}: 10$

LIPINSKY: 4

IUPAC: 2-acetyloxy-4-(5,7-diacetyloxy-4-oxochromen-3-yl)phenyl acetate

Smiles: c1c(c2c(cc1OC(=O)C)occ(c2=O)c1cc(c(cc1)OC(C)=O)OC(C)=O)OC(C)=O

