



Formula: C<sub>48</sub>H<sub>52</sub>N<sub>2</sub>O<sub>10</sub>

MW: 816.95



LogP: -1.64

LogS: -1.84

Acceptors: 10

Donors: 6

Rotation Bonds: 21

Chiral Centers: 0

N+O: 12

LIPINSKY: 2

IUPAC: 8-[(1E)-4-(3-methoxyphenoxy)-2-azabut-1-enyl]-2-{8-[(1E)-4-(3-methoxyphenoxy)-2-azabut-1-enyl]-1,6,7-trihydroxy-3-methyl-5-(methylethyl)(2-naphthyl)}-3-methyl-5-(methylethyl)naphthalene-1,6,7-triol

Smiles:

c1(c2c(c3c(/C=NCCOc4cc(ccc4)OC)c(O)c(c(c3cc2C)C(C)C)O)O)c(c2c(/C=NCCOc3cccc(c3)OC)c(O)c(c(c2cc1C)C(C)C)O)O