Paths of analysis* Analysis 6

Synthia

March 3, 2022

Analysis parameters 1

Analysis type: Automatic Retrosynthesis

Rules: none selected

Filters: FGI, FGI with protections

Max. paths returned: 5

Max. iterations: 300

Commercial:

1. Max. molecular weight - 1000 g/mol

2. Max. price - 1000 \$/g

Published:

1. Max. molecular weight - 1000 g/mol

2. Popularity - 10

My Stockroom:

1. Max. molecular weight - 1000 g/mol

Reaction scoring formula: TUNNEL COEF*FGI COEF*STEP*20+1000 000*(CONFLICT+NON SELECTIVITY+FILTERS+PROTECT)

Chemical scoring formula: SMALLER^ 3,SMALLER^ 1.5

Min. search width: 400

Max. reactions per product: 60

Strategies: none selected

^{*}The results stated herein were generated using the proprietary platform owned and maintained by Grzybowski Scientific Inventions, Inc., a subsidiary of Merck KGaA, Darmstadt Germany. The results are provided on an as is basis, and shall be used solely in connection with the rights afforded in the license agreement and for no other purpose.

FGI Coeff: 0

JSON Parameters: {}

2 Paths

 $1~\mathrm{path}$ found. Paths are sorted by score. Reactions are sorted in appearance order for each path.

2.1 Path 1

Score: 77.94

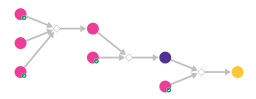


Figure 1: Outline of path 1

2.1.1 Synthesis of N-arylamides from arenediazonium salts

$$\begin{array}{c} & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & &$$

Substrates:

- 1. Vitamin Bx available at Sigma-Aldrich
- 2. 2-Chloronicotinamide Combi-Blocks
- 3. Calcium nitrite solution available at Sigma-Aldrich

Products:

1. 4-(2-chloro-nicotinoylamino)-benzoic acid - Vitas-MLaboratory

Typical conditions: 1) HCl.NaNO2 2) CuI.TBAI.N,N'-dimethylethane-1,2-

diamine.K2CO3.DMSO.110C

Protections: none
Yield: moderate

Reference: DOI: 10.1055/s-0034-1378556

Retrosynthesis ID: 1922

2.1.2 Amide coupling

Substrates:

1. Cyclopentylamine - available at Sigma-Aldrich

 $2. \ \, 4\hbox{-}(2\hbox{-chloro-nicotinoylamino})\hbox{-benzoic acid -} \qquad \textit{Vitas-MLaboratory}$

Products:

 $1. \ O = C(NC1CCCC1)c1ccc(NC(=O)c2cccnc2Cl)cc1$

Typical conditions: DCC.DCM or EDC.DCM or SOCl2.DCM

Protections: none

Yield: good

Reference: 10.1021/cr100048w 10.1039/B701677H and and 10.1039/C5RA24527C and 10.3727/000000006783981206 and 10.1021/np060007f and 10.1021/jo00012a058 and 10.1016/j.bmcl.2007.08.03710.1039/C0OB00355Gand $10.1021/\mathrm{j}m500031w$ (p.3056)and 10.1016/j.tet.2011.03.046

Retrosynthesis ID: 10087

${\bf 2.1.3}\quad {\bf Nucleophilic\ aromatic\ substitution}$

Substrates:

1. O=C(NC1CCCC1)c1ccc(NC(=O)c2cccnc2Cl)cc1

2. 4-Picolylamine - available at Sigma-Aldrich

Products:

1. O=C(NC1CCCC1)c1ccc(NC(=O)c2cccnc2NCc2ccncc2)cc1

Typical conditions: solvent. Heating or pressure

Protections: none

Yield: good

Reference: 10.1021/jm00040a009 or 10.1111/bph.12233 or 10.1246/cl.1987.1187

Retrosynthesis ID: 5003