# Paths of analysis\*

# Synthia

February 13, 2024

# 1 Analysis parameters

Analysis type: Automatic Retrosynthesis

Rules: Expert-Coded Rules

Published Reactions: none selected

Filters: Exclude Diastereoselecitve reactions, Tunnels, FGI, FGI with protec-

tions

Max. paths returned: 50

Max. iterations: 2000

Commercial:

1. Max. molecular weight - 1000 g/mol

2. Max. price -  $1500 \$ /g

#### Published:

1. Max. molecular weight - 1000 g/mol

2. Popularity - 5

#### My Stockroom:

1. Max. molecular weight - 1000 g/mol

Shorter paths: no

Pathway linearity: COMBO

Protecting groups: BALANCED

<sup>\*</sup>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.

Reaction scoring formula: TUNNEL\_COEF\*FGI\_COEF\*STEP\*20+1000 000\*(FILTERS+CONFLICT+NON SELECTIVITY)+40\*PROTECT

Chemical scoring formula: SMALLER  $^{\circ}$  3,SMALLER  $^{\circ}$  1.5

Min. search width: 400

Max. reactions per product: 60

Strategies: none selected

FGI Coeff: 1

Tunnels Coeff: 1

JSON Parameters: {}

# 2 Paths

1 path found. Paths are sorted by score. Reactions are sorted in appearance order for each path.

# 2.1 Path 1

Score: 180.42



Figure 1: Outline of path 1

# 2.1.1 Knoevenagel condenstation

#### Substrates:

1. Cyanoacetic acid - available at Sigma-Aldrich

2. 2-Allyl-2-hydroxycyclohexanone

#### **Products:**

1. C=CCC1(O)CCCCC1=CC#N

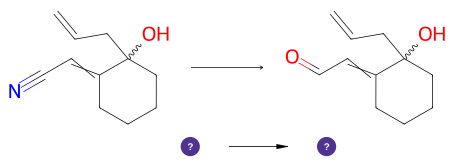
Typical conditions: NH4OAc.AcOH.heat

Protections: none

**Reference:** WO2005/44008 p.96 and 10.1002/hlca.201200162 and 10.1002/chem.200903053 and WO2005/44008 p.90 and US4012377 p.

Retrosynthesis ID: 32838

#### 2.1.2 Reduction of nitriles to aldehydes



#### Substrates:

1. C=CCC1(O)CCCCC1=CC#N

#### **Products:**

 $1. \ C{=}CCC1(O)CCCCC1{=}CC{=}O$ 

 ${\bf Typical\ conditions:}\ {\bf DIBALH.DCM}$ 

Protections: none

**Reference:** 10.1016/j.bmc.2006.01.061 and 10.1016/j.tet.2012.07.022 and 10.1016/j.bmcl.2009.01.075 and 10.1016/j.bmcl.2007.09.081 and 10.1021/jo000502v

Retrosynthesis ID: 31406

#### 2.1.3 Reduction of enones to saturated alcohols

#### Substrates:

 $1. \ C{=}CCC1(O)CCCCC1{=}CC{=}O$ 

#### **Products:**

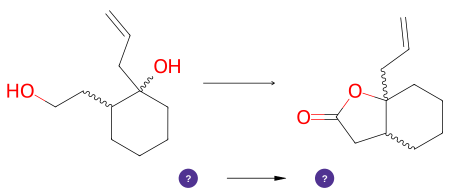
1. C=CCC1(O)CCCCC1CCO

 $\textbf{Typical conditions:} \ \ \text{NaBH4.transition.metal.salt.} (eg.Pd(OAc)2.or.CeCl3)$ 

Protections: none

Retrosynthesis ID: 15309

# 2.1.4 Oxidative lactonization of 1,4-diols



### Substrates:

1. C=CCC1(O)CCCCC1CCO

#### **Products:**

1. C=CCC12CCCC1CC(=O)O2

Typical conditions: Cp(st)RuCl(cod).tBuOK.acetone.30C

Protections: none

**Reference:** DOI: 10.1021/ol0706408

Retrosynthesis ID: 1501

#### 2.1.5Alkylation of Esters

#### Substrates:

1. Chlorallylene available at Sigma-Aldrich

 $2. \ C{=}CCC12CCCC1CC(=O)O2$ 

#### **Products:**

 $1. \ C=CCC1C(=O)OC2(CC=C)CCCCC12$ 

Typical conditions: base e.g. BuLi.THF

Protections: none

Reference: 10.1021/ja065404rand 10.1016/S0040-4020(01)88337-X and 10.1016/0040-4039(95)00562-Q and 10.1021/acs.orglett.6b01901 10.1021/jo00073a034 and 10.1021/ol2023278

Retrosynthesis ID: 31017152