# Paths of analysis\*

## Synthia

March 3, 2022

## 1 Analysis parameters

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

<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.

## FGI Coeff: 0

JSON Parameters:  $\{\}$ 

## 2 Paths

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

## 2.1 Path 1

Score: 231.32

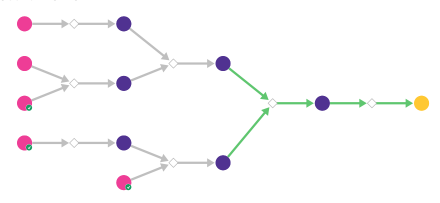


Figure 1: Outline of path 1

## 2.1.1 Reduction of nitro group

Substrates:

1. 1-Bromo-5-isopropoxy-2-methyl-4-nitrobenzene - available at Sigma-Aldrich

#### **Products:**

1. Cc1cc(N)c(OC(C)C)cc1Br

Typical conditions: Zn. aq NH4. EtOH //Zn.Hcl

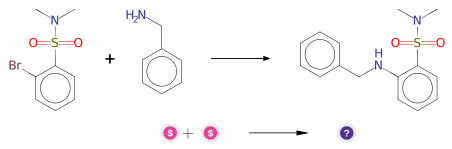
Protections: none

Yield: good

**Reference:** DOI: 10.1002/anie.201512005 and 10.1002/anie.201104681 and 10.3390/molecules17055497 and 10.3390/molecules19022655 and 10.1021/ol5033464 (SI,page 3) and 10.5012/bkcs.2013.34.4.1275

Retrosynthesis ID: 6145

## 2.1.2 Amination of aryl bromides



## Substrates:

1. 2-Bromo-N,N-dimethylbenzenesulphonamide 1g pack - Combi-Blocks

2. Benzylamine - available at Sigma-Aldrich

## **Products:**

1. CN(C)S(=O)(=O)c1ccccc1NCc1ccccc1

Typical conditions: Pd.ligand.base or CuI.ligand.base

Protections: none

100000101101

Yield: good

**Reference:** 10.1021/ja903049z and 10.1021/jo060945k and 10.1021/jo060190h and 10.1039/B923255A and 10.1021/jm8003625 and 10.1021/jo9006738

and 10:1000/ 20:000011 and 10:10:01/ Jhoodood and 10:10:01/ Journal

## 2.1.3 Synthesis of haloarenes via triflates

#### Substrates:

1. 5-chloro-uracil - Combi-Blocks

#### **Products:**

1. O=c1[nH]c(F)ncc1Cl

Typical conditions: 1.Tf2O 2. [Pd].MX

Protections: none
Yield: moderate

**Reference:** 10.1016/j.tetasy.2012.04.008 and WO2007/136577 (p46) and

10.1021/ol202098h and 10.1021/ol402859k and 10.1021/jacs.5b09308

Retrosynthesis ID: 23940

## 2.1.4 Photoredox Cross-Electrophile Coupling of Unactivated Alkyl Bromides

#### Substrates:

- 1. Cc1cc(N)c(OC(C)C)cc1Br
- 2. 2-Bromopropane available at Sigma-Aldrich

## **Products:**

1. Cc1cc(N)c(OC(C)C)cc1C(C)C

Typical conditions: [Ir]-photocat.[Ni]-cat.TTMSS.base.blue light

Protections: none

Yield: good

**Reference:** 10.1021/jacs.6b04818 and 10.1016/j.bbrc.2020.04.028 and 10.1021/ac-

smedchem lett. 8b00183

Retrosynthesis ID: 31016940

## 2.1.5 Amination of pyridones

## Substrates:

1. CN(C)S(=O)(=O)c1ccccc1NCc1ccccc1

2. O=c1[nH]c(F)ncc1Cl

## **Products:**

 $1. \ \mathrm{CN(C)S(=O)(=O)c1ccccc1N(Cc1ccccc1)c1nc(F)ncc1Cl}$ 

Typical conditions: 1.PCl5.2.amine

Protections: none
Yield: moderate

**Reference:** 10.1021/jm300780p AND 10.3390/molecules170910902 AND

10.1021/jm00392a017

## ${\bf 2.1.6}\quad {\bf Nucleophilic\ aromatic\ substitution}$

## Substrates:

- 1. Cc1cc(N)c(OC(C)C)cc1C(C)C
- $2. \ \ CN(C)S(=O)(=O)c1ccccc1N(Cc1ccccc1)c1nc(F)ncc1Cl$

## **Products:**

 $1. \ Cc1cc(Nc2ncc(Cl)c(N(Cc3ccccc3)c3ccccc3S(=O)(=O)N(C)C)n2)c(OC(C)C)cc1C(C)Cnc(CC)$ 

Typical conditions: Solvent

Protections: none

Yield: good

Reference: 10.1002/9781118093559.ch4

Retrosynthesis ID: 49476

## 2.1.7 Debenzylation

## Substrates:

 $1. \ Cc1cc(Nc2ncc(Cl)c(N(Cc3ccccc3)c3ccccc3S(=O)(=O)N(C)C)n2)c(OC(C)C)cc1C(C)Cn2(CC)$ 

#### **Products:**

 $1. \ Cc1cc(Nc2ncc(C1)c(Nc3ccccc3S(=O)(=O)N(C)C)n2)c(OC(C)C)cc1C(C)C$ 

Typical conditions: H2. Pd/C or Pd(OH)2

Protections: none

Yield: good

**Reference:** DOI: 10.1002/1521-3773(20020603)41:11<1895::AID-ANIE1895>3.0.CO;2-3 and 10.1021/jo400589j and 10.1021/jm8012932 (SI,page S6) and 10.1080/00397911.2016.1261164

Retrosynthesis ID: 9995661

## 2.2 Path 2

Score: 331.78

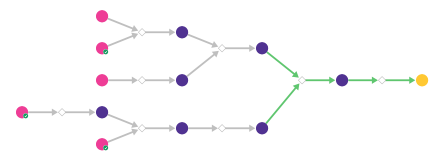


Figure 2: Outline of path 2

## 2.2.1 Reduction of nitro group

## Substrates:

 $\begin{array}{lll} \hbox{1. 1-Bromo-5-isopropoxy-2-methyl-4-nitrobenzene} & & available \ at \ Sigma-Aldrich \end{array}$ 

#### **Products:**

1. Cc1cc(N)c(OC(C)C)cc1Br

Typical conditions: Zn. aq NH4. EtOH //Zn.Hcl

Protections: none

Yield: good

**Reference:** DOI: 10.1002/anie.201512005 and 10.1002/anie.201104681 and 10.3390/molecules17055497 and 10.3390/molecules19022655 and 10.1021/ol5033464 (SI,page 3) and 10.5012/bkcs.2013.34.4.1275

Retrosynthesis ID: 6145

## 2.2.2 Photoredox Cross-Electrophile Coupling of Unactivated Alkyl Bromides

## Substrates:

1. Cc1cc(N)c(OC(C)C)cc1Br

2. Bromocyclopropane - available at Sigma-Aldrich

#### **Products:**

1. Cc1cc(N)c(OC(C)C)cc1C1CC1

 $\textbf{Typical conditions:} \ [Ir]-photocat.[Ni]-cat.TTMSS.base.blue \ light$ 

Protections: none

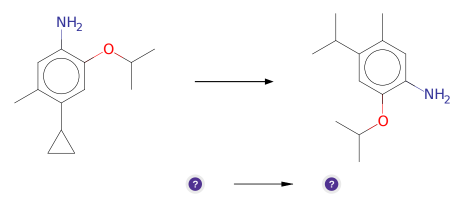
Yield: good

**Reference:** 10.1021/jacs.6b04818 and 10.1016/j.bbrc.2020.04.028 and 10.1021/ac-j.bbrc.2020.04.028 and 10.1021/ac-j.bbrc.2020.04.028

smedchem lett. 8b00183

Retrosynthesis ID: 31016940

## 2.2.3 Hydrogenolysis of cyclopropane



## Substrates:

1. Cc1cc(N)c(OC(C)C)cc1C1CC1

## **Products:**

1. Cc1cc(N)c(OC(C)C)cc1C(C)C

Typical conditions: H2.Pd/C

Protections: none

Yield: good

**Reference:** 10.1021/jo00264a013 AND 10.1021/ja00272a046 AND

10.1021/jm950677a

## 2.2.4 Balz-Schiemann Reaction

#### Substrates:

1. 4,5-Dichloropyrimidin-2-amine - Combi-Blocks

#### **Products:**

1. Fc1ncc(Cl)c(Cl)n1

Typical conditions: NaNO2.HF-pyridine.-25 to  $0\mathrm{C}$ 

Protections: none
Yield: moderate

**Reference:** 10.1021/jm100432w and 10.1021/jm5008177 and 10.1021/ol401540k and 10.1021/jm401551n (main text and SI, page S8) and 10.1515/chempap-2016-0033 and 10.1021/jm0311442 and 10.1016/j.jfluchem.2007.03.012 and 10.1021/jo00185a023

Retrosynthesis ID: 29906

## 2.2.5 Chan-Lam Coupling



#### Substrates:

1. 2-(N,N-DIMETHYLSULPHAMOYL)BENZENEBORONIC ACID Combi-Blocks

2. Benzhydrazide - available at Sigma-Aldrich

## **Products:**

1. CN(C)S(=O)(=O)c1ccccc1NNC(=O)c1ccccc1

Typical conditions: Cu(Oac)2.dioxane

Protections: none

Yield: good

**Reference:** 10.1039/C4OB02343A **Retrosynthesis ID:** 31015967

## 2.2.6 Nucleophilic aromatic substitution

#### Substrates:

1. Fc1ncc(Cl)c(Cl)n1

 $2. \ \mathrm{CN(C)S(=O)(=O)c1ccccc1} \\ \mathrm{NNC(=O)c1ccccc1}$ 

## **Products:**

1. CN(C)S(=O)(=O)c1ccccc1N(NC(=O)c1ccccc1)c1nc(F)ncc1Cl

Typical conditions: EtOH.Heat or DMF.K2CO3

Protections: none

Yield: good

 $\textbf{Reference:} \quad 10.1016/0040\text{-}4020(95)00966\text{-}3 \quad \text{and} \quad 10.1002/\text{jhet}.5570220420 \quad \text{and} \quad 10.1002/\text{jhet}.55702020420 \quad \text{and} \quad 10.1002/\text{jhet}.55702020420 \quad \text{and} \quad 10.1$ 

10.1021/jo402481t

Retrosynthesis ID: 29647

## 2.2.7 Nucleophilic aromatic substitution

## Substrates:

1. Cc1cc(N)c(OC(C)C)cc1C(C)C

 $2. \ \mathrm{CN(C)S(=O)(=O)c1ccccc1N(NC(=O)c1ccccc1)c1nc(F)ncc1Cl} \\$ 

## Products:

Typical conditions: Solvent

Protections: none

.. .

Yield: good

**Reference:** 10.1002/9781118093559.ch4

## ${\bf 2.2.8} \quad {\bf Reduction \ of \ phenylhydrazines}$

## Substrates:

 $1. \ \ Cc1cc(Nc2ncc(Cl)c(N(NC(=O)c3ccccc3)c3ccccc3S(=O)(=O)N(C)C)n2)c(OC(C)C)cc1C(C)Cnc(CC)$ 

## Products:

 $1. \ Cc1cc(Nc2ncc(Cl)c(Nc3ccccc3S(=O)(=O)N(C)C)n2)c(OC(C)C)cc1C(C)C)\\$ 

 $\textbf{Typical conditions:} \ \, \text{RaNi.H2.MeOH}$ 

Protections: none
Yield: moderate

**Reference:** DOI: 10.1021/ol501424f