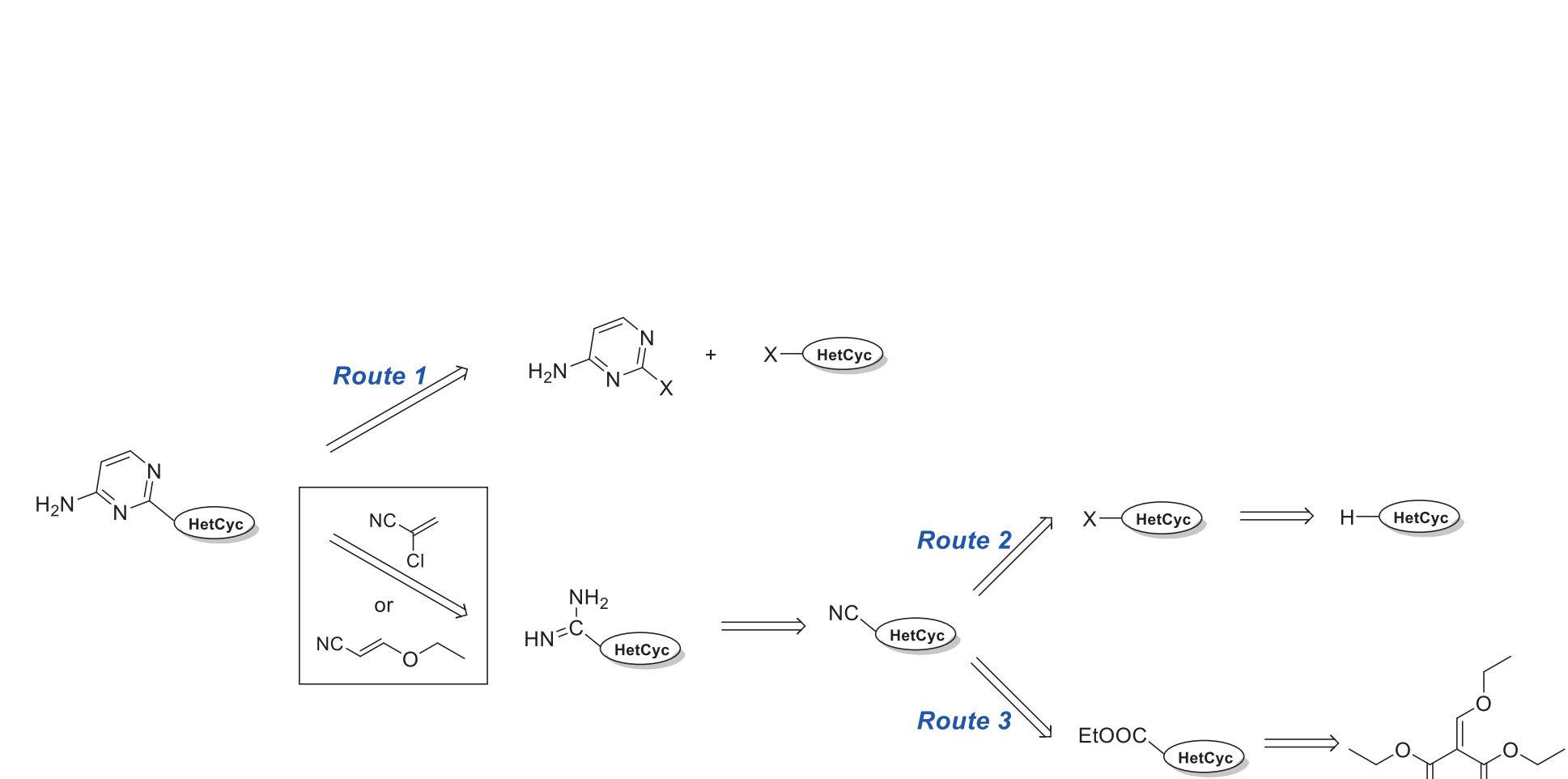


Efficient and Inexpensive Synthesis of A Functionalized Pyrimidine

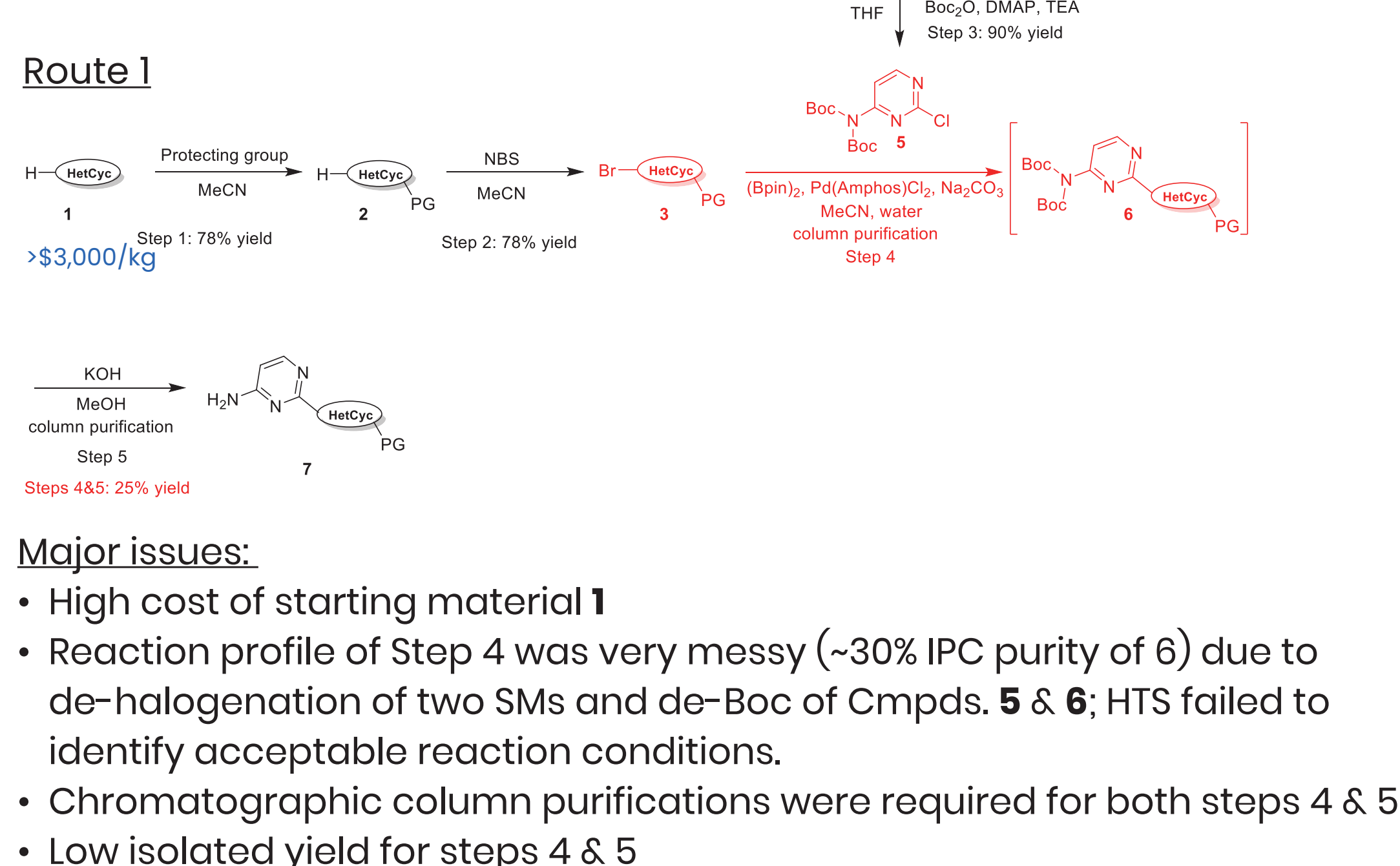
Yu Lu
Process R&D, WuXi STA



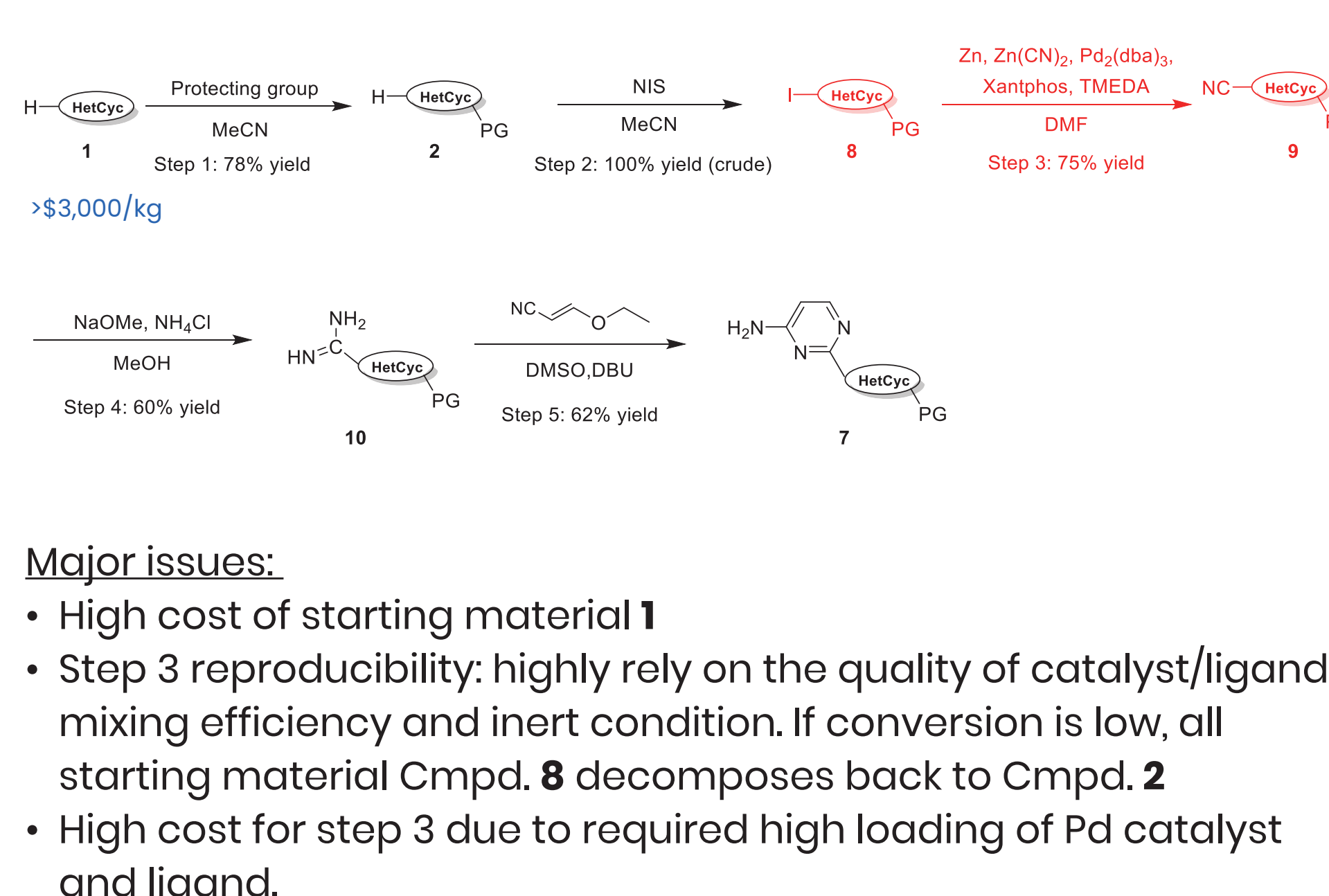
Retro-synthetic Analysis



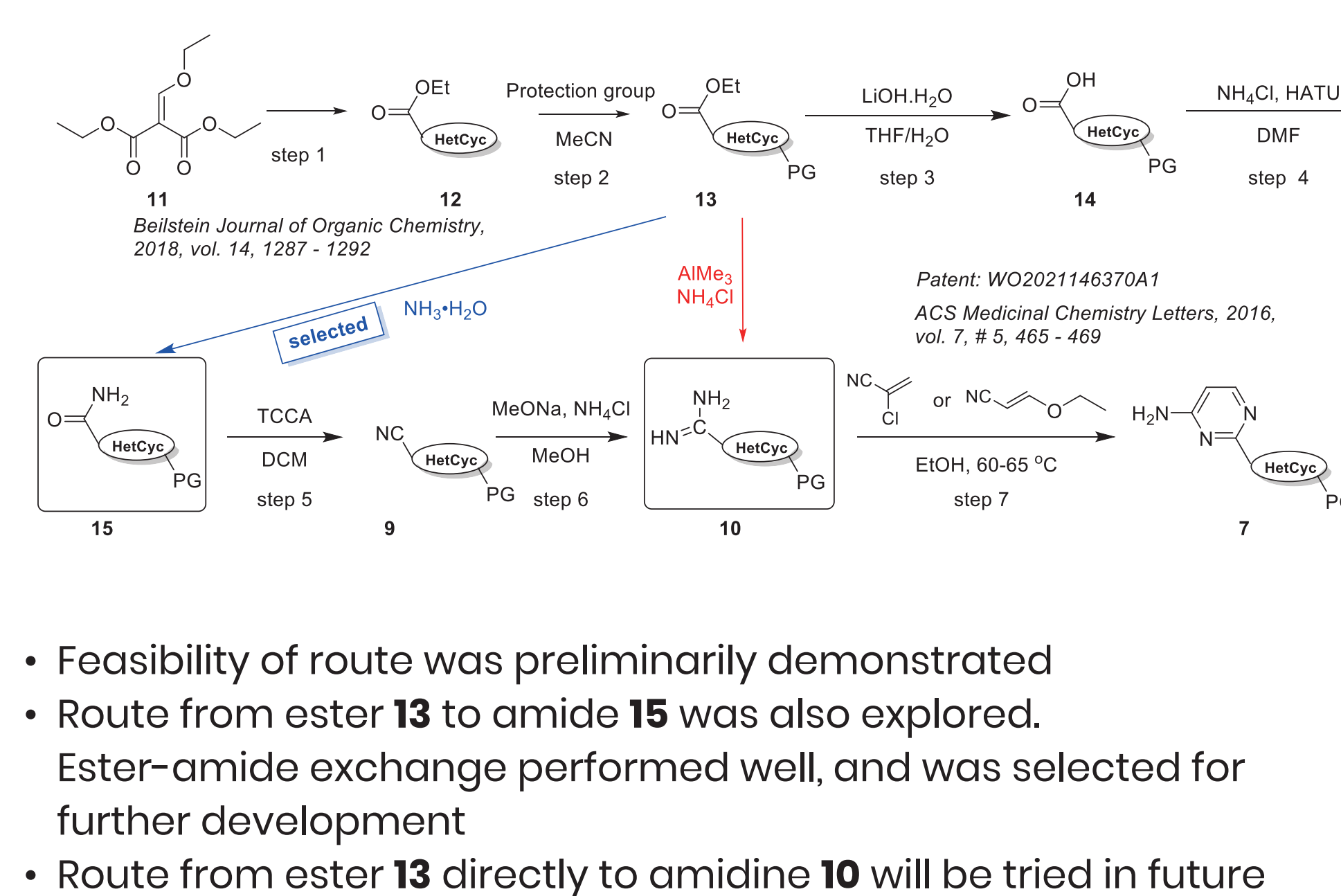
Route Scouting



Route 2



Route 3: constructing amidine **10** while avoiding expensive & problematic catalytic cyanation



Aspects for Process Development

Better control of heat and mass transfer
Avoid use and isolation of unstable, hazardous and energetic reagents, starting materials and intermediates

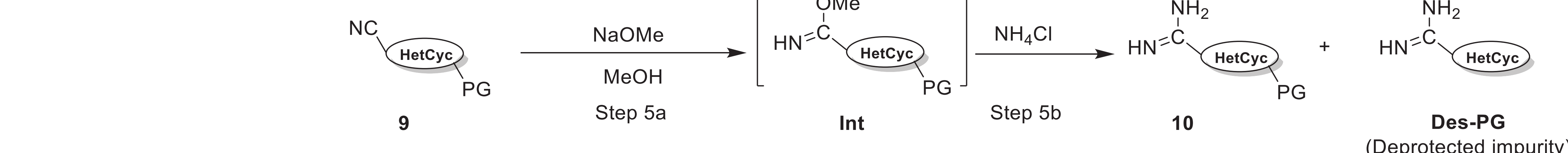
Consistent and reliable process performance despite different equipment and site, scale change and variations in process parameters
Supply of materials is stable, and quality is consistent
Stable intermediates for isolation, shipment and storage

Incorporation of green chemistry principles into synthetic route design
Selection of reagents and solvents based on atom economy and green principles economy and green principles cleaning cost, etc.

Material cost: solvents, reagents, catalyst, etc.
Labor cost: operators, analysts, etc.
Production cost: production cycle time, equipment, instruments, waste treatment, cleaning cost, etc.

Zhang, Tony Y. *Chemical Reviews*, 2008, 106(7), 2583-2595.

Step 5



Equivalents of reagents and reaction temperature

Entry	NaOMe (equiv.)	MeOH (Vol)	Step 5a T (°C)	Step 5a t (h)	Step 5b IPC (g/ml)	NH ₄ Cl (equiv.)	Step 5b T (°C)	Step 5b t (h)	Step 5b IPC (g/ml)	Des-PG impurity
1	1.25	20	40-50	18	18.3 / 87.8	3.5	40-50	19	14.9 / 83.5 / 0	0%
2	4.5	20	40-50	20	12.2 / 89.0	3.5	40-50	19	13.1 / 83.6 / 0	0.5%
3	1.25	20	40-50	2	15.0 / 84.7	2.5	20-30	20	12.7 / 85 / 78.7 / 0.8	1.3%

- Step 5a is reversible; conversion could not be completed;
- Ratio of NaOMe/NH₄Cl is important; highly excess NH₄Cl led to des-PG impurity growth in Step 5b.

- Add NaOMe and NH₄Cl alternately to drive conversion of step 5a while mitigating PG- deprotection.

Reagent	sol.	T (°C)	T (h)	9 / Int / 10	Des-PG impurity
NaOMe: 1.25 eq	MeOH	40-50	2	12.6 / 83.5 / 0	0%
+NH ₄ Cl: 2.5 eq	10V	20-30	18	12.0 / 2.6 / 80.7	0.5%
+NaOMe: 2.0 eq	MeOH	40-50	2	4.5 / 21.7 / 69.6	0.4%
+NH ₄ Cl: 2.5 eq	10V	20-30	18	4.0 / 0.3 / 90.0	1.3%

- Conversion of step 5a is improved, and des-PG impurity is suppressed

From:

1.0 eq SM
2.5 eq NaOMe, 3.5 eq NH₄Cl
20V MeOH
45°C

Filter, concentrate filtrate, and slurry with EtOAc

Charge NaOMe and NH₄Cl alternately to improve conversion
Reduce reaction volume

Isolate free base instead of salt through base wash and crystallization
Yield is improved from 60% to 75%

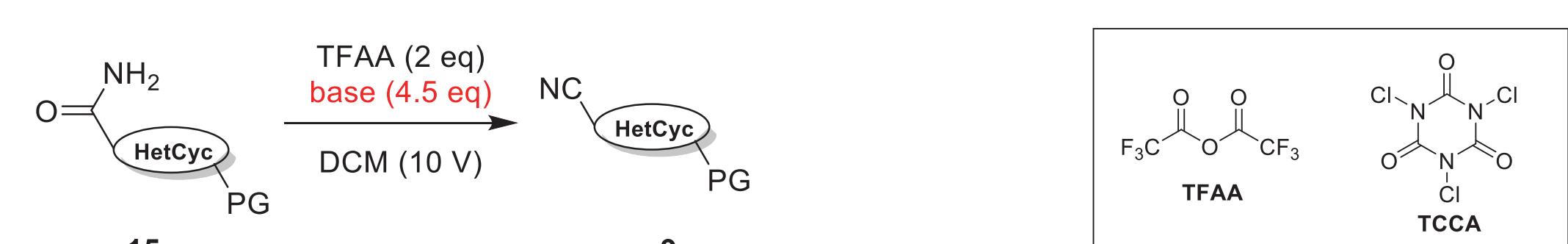
Demo result: 100 g scale, 75.0% isolated yield, 99.9% purity, 98.2% assay

To:

1.0 eq SM
12x20 eq NaOMe, 25x25 eq NH₄Cl
10V MeOH
40-50 °C for step 5a
20-30 °C for step 5b

Filter, concentrate filtrate, dilute with DCM, wash with aqueous Na₂CO₃ solution, extract with DCM, concentrate and switch with EtOAc. Filter and dry.

Step 4



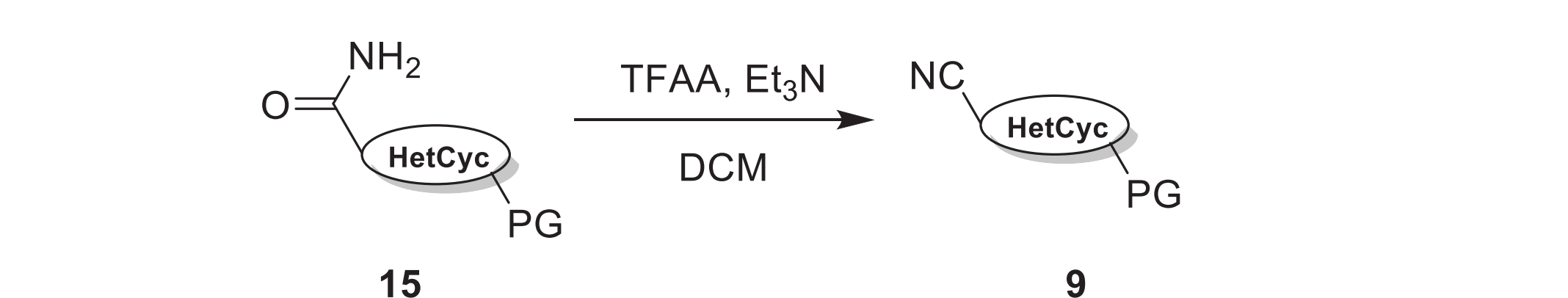
Screen reagent (TFAA>TCCA) and temp. (no significant effect)
Reduce reagent equiv. and solvent vol.

Et₃N works best among screened organic bases

base effect on reaction conversion

Entry	Reagents	Solvent	Temp (°C)	Time (h)	IPC (HPLC area%) Cmpd. 15 / Cmpd. 9
1	DPEA: 3.0eq TCCA: 1.0eq	DCM (10 v)	0-10	20	34.3 / 9.8
2	DPEA: 4.5eq TFAA: 2.0eq	DCM (10 v)	20-30	18	39 / 75.1
3	Et ₃ N: 4.5eq TFAA: 2.0eq	DCM (10 v)	20-30	2	0/90.3
4	Et ₃ N: 4.5eq TFAA: 2.0eq	DCM (10 v)	0-10	2	2.8/89.6
5	TFAA: 5.0eq TFAA: 1.5eq	DCM (5 v)	0-10	2	15/94.5

TFAA/Et₃N was selected due to better conversion and reaction impurity profile



From: 1.0eq SM, 3.0eq DIPEA, 1.0eq TCCA, 10V DMF, 0-25 °C, 14hrs

To: 1.0eq SM, TEA: 3.5eq, TFAA: 1.5eq, 5V DCM, 0-10 °C, 17hrs

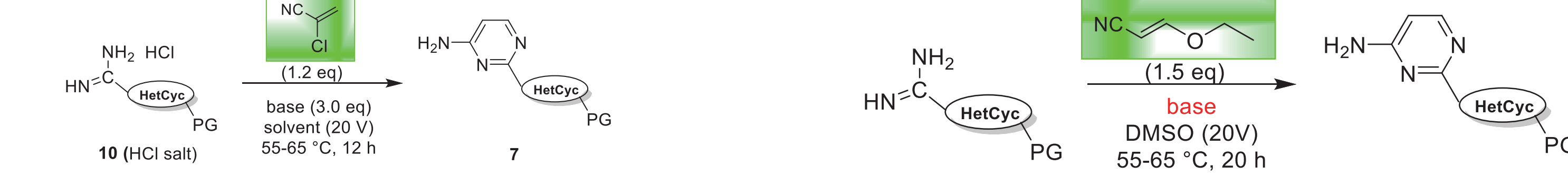
Replace TCCA with TFAA to avoid allergic reagent and improve reaction profile
Replace solvent and reduce volume

Develop extraction and crystallization process to purify product
Yield is improved from 48% to 86%

Quench with water, extract with DCM, concentrate and crystallize with DCM/MTBE. Filter and dry

Demo result: 120 g scale, 86.1% isolated yield, 99.9% purity, 99.7% assay

Step 6



pyrimidine ring formation w/ 2-chloroprop-2-enitrile: Patent: WO2020146570A1

Tried 2-chloroprop-2-enitrile as in the patent literature: low to moderate conversion

base & solvent screening

base screening

Freebase of **10** was used

HCl salt of **10** was used

Use of 3-ethoxyacrylonitrile provided higher conversion in general
Use of **10** as freebase gave better results and easier handling in Step 5
DBU performed best among screened inorganic & organic bases

Further reduction of reagent equiv. with DBU (2 eq) in DMSO (5V) at 55-65 °C for 18h:

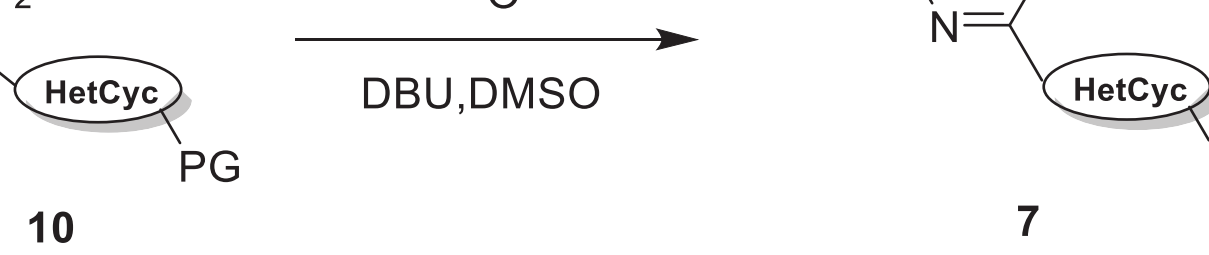
3-ethoxyacrylonitrile (equiv.)	IPC (HPLC area%) Cmpd. 10 / Cmpd. 7
11	1.0 / 90.5
12	0.6 / 91.9
13	0.5 / 92.4

DCM was selected. Reaction solvent DMSO can be removed by water wash.

Crystallization study of Cmpd. 7 to determine crystallization solvent system for purity upgrade and material recovery

Solvent system	Input purity	Purity of wet cake	Loss in mother liquor
THF	87%	99.8%	11%
MeCN	87%	99.5%	20%
DCM/n-heptane (4/16, v/v)	87%	91.0%	Less than 1%
DCM/EtOAc (4/16, v/v)	87%	99.6%	11%
IPA/n-heptane (5/15, v/v)	87%	99.7%	19%
IPA/water (5/15, v/v)	87%	97.2%	More than 40%

DCM/EtOAc was selected to avoid solvent switch with acceptable impurity purging efficiency.



From: 1.0eq SM, 1.0eq 2-chloroprop-2-enitrile, 10V EtOH, 60 °C, 3hrs

To: 1.0eq SM, 1.2eq 3-ethoxyacrylonitrile, 2eq DBU, 5V DMSO, 55-65 °C, 22hrs

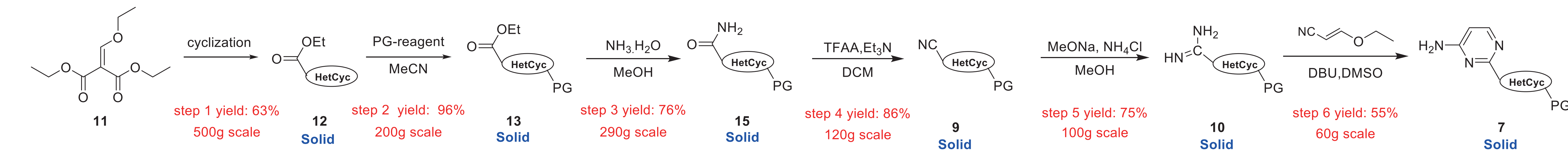
Screen reagent, replace 2-chloroprop-2-enitrile with 3-ethoxyacrylonitrile; Reduce reaction volume

Isolate free base instead of salt through base wash and crystallization
Yield is significantly improved from 14% to 55%

Dilute with DCM, wash with water, extract with DCM, concentrate, and crystallize with DCM and EtOAc. Filter and dry.

Demo result: 60 g scale, 55% isolated yield, 100% purity, 99.1% assay

Route 3 after process optimization



- Inexpensive and readily available starting materials
- No transition-metal catalyzed chemistry; good robustness in lab scale-up
- Process is ready for kg scale-up