Ugi Multicomponent Reaction

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Procedure

A. **N-(2,2-Dimethoxyethyl)-N-(2-oxo-2-(phenethylamino)ethyl)cyclohexane-carboxamide.** Into a 500 mL 3-necked, round-bottomed flask, equipped with a 6 cm Teflon blade overhead stirrer (Note 1), a 60-mL pressure equalizing dropping funnel holding a nitrogen inlet and a thermocouple (Figure 1), is added a mixture of **N-(phenethyl)formamide (7.46 g, 50 mmol)** and triethylamine (16.8 mL, 120 mmol 2.4 equiv) in dichloromethane (50 mL) (Note 2). The solution is cooled to approximately –10 °C using an ethanol-ice bath (Note 3).
Figure 1. Reaction set-up

Triphosgene (5.94 g, 20 mmol, 0.4 equiv) (Note 4) in dichloromethane (20 mL) is added dropwise to the stirring (500 rpm) mixture via the dropping funnel over a period of 45 min (Note 5), resulting in a dark red/brown solution. The reaction mixture is stirred at –10 °C for an additional 30 min (Note 6) (Figure 2).

Figure 2. Color changes as reaction progresses
A separate 50 mL round-bottomed flask, which is charged with a mixture of aminoacetaldehyde dimethyl acetal (5.52 g, 52.5 mmol, 1.05 equiv) and paraformaldehyde (1.50 g, 50 mmol, 1 equiv) in 50 mL methanol, is equipped with an air condenser and heated to 80 °C in an oil bath with stirring until a clear solution developed. After allowing the solution to cool to room temperature, cyclohexanecarboxylic acid (6.72 g, 52.5 mmol, 1.05 equiv) is added. The resulting solution is then added together with another 50 mL MeOH to the in situ formed isocyanide reaction mixture at –10 °C via cannula. The reaction mixture is left to warm to room temperature. The overhead mechanical stirrer and 60-mL pressure equalizing dropping funnel are replaced with red septa and a 2.5 cm Teflon stir bar (Figure 3).

Figure 3. Post-reagent addition set-up

After stirred at room temperature for 48 h (Note 7), the reaction mixture is transferred to a 1000 mL one-necked, round-bottomed flask, along with CH₂Cl₂ (100 mL) that is used to rinse the flask, and the solution is concentrated by rotary evaporation (40 °C, 7.5 mmHg) to remove the methanol. Then the mixture is redissolved in CH₂Cl₂ (50 mL) and transferred to a 250 mL separatory funnel, washed with water (3 x 50 mL), saturated NaHCO₃ (3 x 50 mL), and dried over MgSO₄ (10g). The drying agent is removed by filtration through a medium porosity 150 mL filter funnel (200 mmHg applied vacuum), and the resulting solution is collected into a 1000 mL round-bottomed flask, in which the solution is concentrated by rotary evaporation (40 °C, 7.5 mmHg) (Note 8).
The red/brown oil is purified by flash chromatography on silica using hexanes and ethyl acetate as the eluent (Note 9). The collected fractions are placed in a 1000-mL round-bottomed flask and then concentrated by rotary evaporation (40 °C, 7.5 mmHg) to yield a reddish-brown oil (picture above) which is then placed under high vacuum (45 °C, 0.2 mmHg) for 12 h (Note 10) to afford 7.46 g (46%, >99% purity) of \(N\)-(2,2-dimethoxyethyl)-\(N\)-(2-oxo-2-(phenethylamino)-ethyl)cyclohexanecarboxamide as a brown solid (Figure 4) (Notes 11 and 12).

**Notes**

1. The Teflon blade overhead stirrer was purchased from Arrow Engineering Mixing Products, model #1750; 115 VAC 60 Hz.
2. Reagents and solvents used in this preparation are commercially available and used without further purification, including solvents dichloromethane and methanol, which were purchased from Fisher Chemical. Triphosgene (98%) and aminoacetaldehyde dimethyl acetal (99%) from AK Scientific Inc., \(N\)-(phenethyl)formamide (97%), trimethylamine (≥99%), paraformaldehyde (95%) and cyclohexane carboxylic acid (≥98%) from Sigma Aldrich.
3. In a 1L Dewar cooling bath 275 g ice and 150 g ethanol were used to obtain a temperature of approximately –10 °C for the duration of the
reaction. Checkers utilized the thermocouple in the ice/ethanol bath to accurately measure for the duration of the reaction.

4. The solid reagent triphosgene is a less hazardous substitute for highly toxic gaseous phosgene, however should be handled very carefully. Triphosgene may be fatal if inhaled and causes burns by all exposure routes. This water-reactive substance liberates toxic gas upon exposure to water. Triphosgene is a lachrymator and can decompose violently at elevated temperatures. Triphosgene should be weighed out and the reaction should be performed in a well-ventilated fume hood.

5. Faster addition or temperatures above 0 °C will reduce the yield dramatically. Submitters noted that color is a good indicator of proper addition speed: slightly yellow color is good, orange towards brown indicates too fast addition of triphosgene.

6. The isocyanide intermediate Rf = 0.80 (1:1 Hexanes:EtOAc), checked by TLC EMD gel 60 F254 pre-coated plates (0.25 mm) (visualized with 254 nm UV lamp), appears nearly exclusively indicating consumption of the formamide (Figure 5).

Figure 5. TLC analysis of isocyanide formation

7. Not all reagents are fully consumed, but after 48 h no change in spot intensity, checked by TLC EMD gel 60 F254 pre-coated plates (0.25 mm) (visualized with 254 nm UV lamp), was observed.

Figure 6. TLC analysis of multi-component reaction
8. The bulk of solvent should be removed; however, leaving a trace of solvent (~0.5 g) is actually beneficial in allowing the product to crystallize. The crystallization process requires time (up to 2 weeks). The submitters report a yield of 9.05 g crude product. The submitters then triturate the crude crystals with 10 mL cold Et₂O, filtering through a P4 glass filter and washing with another 10 mL cold Et₂O. Yields after drying (0.5 mbar) 7.56 g (40%) pure N-(2,2-dimethoxyethyl)-N-(2-o xo-2-(phenethylamino)ethyl)cyclohexanecarb oxamide as pale yellow crystals (Figure 7).

Figure 7. Appearance of the Ugi product from submitters

9. The column chromatography was run as follows: A flash column with an outer diameter of 8 cm and capacity of 2000 mL was charged with silica (Silicycle Siliaflash P60 particle size 0.040–0.063 mm purchased from Silicycle; used as received) using a wet-pack method (316 g of silica in 700 mL of 66% hexanes in ethyl acetate). This gave a silica bed of 16 cm in height. The crude mixture was then dissolved in 15 mL of 66% hexanes in ethyl acetate and then loaded onto the silica using a pipette. Sand was then added to fill 2 cm above the silica. An eluent mixture of 66% hexanes in ethyl acetate (1300 mL) was used initially, followed by 50% ethyl acetate in hexanes (2000 mL) and 100% ethyl acetate (3000 mL). The flow rate was approximately 66 mL/min and 30-mL fractions were collected. Fractions 148–182 were then collected as the product fractions.

10. The checkers found that ethyl acetate was difficult to remove. Placing the purified reddish-brown oil under high vacuum (45 °C, 0.2 mmHg) for 12 h was required to remove the ethyl acetate.
11. \(N-(2,2\text{-Dimethoxyethyl})-N-(2\text{-oxo-2-(phenethylamino)ethyl})\) cyclohexanecarboxamide: \(\text{mp} = 83.4\text{--}85.2\) °C, \(R_f = 0.42\) (100\% EtOAc). \(^1\)H NMR (500 MHz, CDCl\(_3\)) \(\delta\): 1.16--1.30 (m, 3H), 1.37--1.52 (m, 2H), 1.56--1.71 (m, 2H), 1.71--1.82 (m, 2H), 2.24 (tt, \(J = 11.5, 3.4\) Hz, 0.5H), 2.58 (tt, \(J = 11.5, 3.4\) Hz, 0.5H), 2.80 (dt, \(J = 20.6, 7.2\) Hz, 2H), 3.33 (s, 3H), 3.37 (s, 3H), 3.42 (app t, \(J = 4.8\) Hz, 2H), 3.49 (q, \(J = 6.7\) Hz, 1H), 3.55 (q, \(J = 6.8\) Hz, 1H), 3.98 (d, \(J = 6.5\) Hz, 2H), 4.39 (t, \(J = 5.3\) Hz, 0.5H), 4.57 (t, \(J = 5.1\) Hz, 0.5H), 6.41--6.50 (m, 0.5H), 6.94--7.00 (m, 0.5H), 7.16--7.25 (m, 3H), 7.27--7.32 (m, 2H); \(^{13}\)C NMR (125 MHz, CDCl\(_3\)) \(\delta\): 25.7, 25.8, 25.9, 29.4, 29.5, 35.7, 35.8, 40.4, 40.6, 40.8, 41.2, 50.5, 51.6, 52.3, 54.2, 55.2, 55.6, 102.8, 103.6, 126.6, 126.8, 128.7, 128.8, 128.9, 138.7, 138.9, 169.4, 169.7, 178.0, 178.2; IR (film): 3301, 2929, 2854, 1627, 1544, 1451 cm\(^{-1}\). HRMS (ESI). \([M + H]^{+}\) calcd. for \(C_{21}H_{33}O_4N_2\): 377.2440. Found: 377.2408. The tertiary amide rotamers are clearly visible in the NMR spectrum and show a 1:1 ratio. The purity of the compound was determined using qNMR: 14.1 mg of the product are dissolved in 0.8 mL of CDCl\(_3\). 1,3,5-trimethoxybenzene, 10.2 mg (99\%, purchased from Alfa Aesar and used as received), is added as the internal standard. \(^1\)H NMR (500 MHz, CDCl\(_3\)) gave a product purity of 99.2\%.

12. A second reaction provided the product as a brown solid in 46\%.

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the procedures herein.

Discussion

The Ugi multicomponent reaction of amines, oxo components,
isocyanides and inorganic or organic acids leads to different scaffold types
dependent on the nature of the acid component. For example carboxylic
acids yield α-amino acylamides (Table 1). The Ugi multicomponent reaction
is gaining increasing attention due to the rapid and convergent assembly of
functional structures based on four classes of widely available starting
material classes. In addition, our recent work on generating the isonitrile in situ
is making the Ugi reaction even more attractive, as the isolation of the
infamous isonitrile is circumvented in this methodology. While several
approaches towards α-amino acylamides are possible, the Ugi approach is
faster and moreover impresses by a very large substrate scope. Here the
Ugi intermediate to the schistosomiasis drug praziquantel (PQZ, 2-
(cyclohexanecarbonyl)-2,3,6,7-tetrahydro-1H-piazino[2,1-a]isoquinolin-4(11bH)-one) is prepared to exemplify the usefulness of this methodology. It
comprises a one-pot synthesis from the readily available bulk starting
materials, 2-phenylethyl formamide, formaldehyde, cyclohexanecarboxylic
acid and aminoacetaldehyde dimethylacetal. N-Phenethylformamide reacts
with triphosgene, which in turn reacts with paraformaldehyde, aminoacetaldehyde dimethylacetal, and cyclohexylcarboxylic acid in an Ugi
four component reaction to the advanced praziquantel precursor in 46% yield. Many similar derivatives have been synthesized accordingly (Table
1).
Table 1. Examples of Ugi products and their yields

The yields described are calculated from the combined yields over the two subsequent Ugi and Pictet-Spengler reaction steps, as the crude Ugi intermediate was directly used in the Pictet-Spengler reaction.

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*The yields described are calculated from the combined yields over the two subsequent Ugi and Pictet-Spengler reaction steps, as the crude Ugi intermediate was directly used in the Pictet-Spengler reaction.*
References

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Appendix

Chemical Abstracts Nomenclature (Registry Number)

N-(Phenethyl)formamide (23069-99-0)
Triphosgene (32315-10-9)
Triethylamine (121-44-8)
Paraformaldehyde; (30525-89-4)
Aminoacetaldehyde dimethyl acetal; (22483-09-6)
Cyclohexane carboxylic acid; (98-89-5)
N-(2,2-Dimethoxyethyl)-N-(2-oxo-2-(Phenethylamino)ethyl)cyclohexanecarboxamide; (90142-13-5)
Alexander Dömling studied chemistry & biology at the Technical University Munich. After performing his Ph.D. under the supervision of Ivar Ugi he spent his postdoctoral year at the Scripps Research Institute in the group of Barry Sharpless under a prestigious Feodor Lynen research fellowship from the Alexander von Humboldt society. He is founder of several biotech companies, including Morphochem, Telesis and SMIO. In 2004 he performed his habilitation in chemistry at the Technical University of Munich. Since 2006 he was Professor at the University of Pittsburgh in the Department of Pharmacy and Chemistry and in 2011 he became Chair of Drug Design at the University of Groningen. He is author of more than 200 papers, reviews, book contributions and patents. His research interest focuses on novel aspects of multicomponent reaction chemistries and its applications to drug design.

André Boltjes was born in the Netherlands. He received his Bachelor’s degree in 2007 from Hanze University and started working at the University of Groningen, developing dopamine agonists, HAT inhibitors, doxorubicin prodrugs and performing various contract synthesis projects. In 2011 he joined Dömling’s group as a technician in University of Groningen.
Haixia Liu was born in Tianjin, China. She received her undergraduate chemistry degree at Nanjing University in 2002. Then, she moved to Shanghai, China to pursue her doctoral studies with Dr. Zhu-Jun Yao at the Shanghai Institute of Organic Chemistry, where she studied chemical biology with particular focus on mimetics of Annonaceous acetogenin. In January 2008, she came to the U.S. to start her post-doctoral work with Dr. Alexander Dömling at the University of Pittsburgh to study multi-component reactions to build small-molecule libraries. She then continued her post-doctoral work when she came to UCLA in January 2009, where she is now working on syntheses of ghrelin O-acyltransferase (GOAT) inhibitors. Currently she is working for Roche/Shanghai.

Haiping Cao was born in Zhejiang, China. She received her BA degree in 2002 from Fudan University, China. After five years of studies, she obtained her PhD in organic chemistry from Shanghai Institute of Organic chemistry, China under the supervision of Professor Qingyun Chen in 2007. Her dissertation focused on new methodology to synthesize fluorinated compounds. In 2009 she joined Professor Alexander Dömling’s group as a postdoctoral associate at the University of Pittsburgh.

Emma L. Baker-Tripp received her B.A. in chemistry from Grinnell College in 2013. She is currently pursuing her Ph.D. in organic chemistry in the laboratory of Professor Neil K. Garg at the University of California, Los Angeles. Her graduate research has focused on the total synthesis of welwitindolinone alkaloids and the development of new C-C and C-heteroatom bond forming reactions of amides mediated by nickel catalysis.
N-(2,2-dimethoxyethyl)-N-(2-oxo-2-(phenethylamino)ethyl)cyclohexanecarboxamide

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