A Visual Organic Chemistry Reaction: The Synthesis of 4-Amino-3-nitrobenzoic Acid Methyl Ester via Fischer Esterification

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ABSTRACT

The synthesis of 4-amino-3-nitrobenzoic acid methyl ester is a simple Fischer esterification reaction designed as an experiment for use in an introductory organic chemistry course. The compound was synthesized as a one-pot reaction within 30 mins to 16 hours, with one hour producing a workable yield. The bright yellow solid was purified using liquid-liquid extraction, and the extraction process is monitored by a marked color change. The product was then characterized by $^1$H NMR, $^{13}$C NMR, and thin-layer chromatography.

GRAPHICAL ABSTRACT

KEYWORDS

Ester, Synthesis, introductory chemistry, organic chemistry, hands-on learning, visual chemistry.

INTRODUCTION

Laboratories in undergraduate science incorporate visual, auditory, reading/writing, and kinesthetic aspects. Fundamental concepts that are difficult to appreciate otherwise, such as solubility and polarity, are demonstrated in laboratory sessions to enhance student learning outcomes. If students are able to visually monitor reactions and purification steps in an organic reaction, it helps to reinforce the main concepts learned. In introductory undergraduate chemistry laboratories, a typical first synthesis experiment would include known reactions such as aspirin synthesis. Aspirin synthesis is a simple and well-documented reaction with known hazards and risks. The synthesis applies itself to infrared
spectroscopy with the determination of the added carbonyl group. However, more commonly, we see that aspirin synthesis is now often performed in secondary education at high school, and thus there is a drop in the students’ engagement if repeated at a tertiary level.

Fischer esterification is an important topic included in most introductory chemistry courses, and examples of these reactions are abundant. The reaction is a reversible condensation reaction where a carboxylic acid reacts with an alcohol to produce an ester and water. The amount of water present in the reaction affects the equilibrium position and thus impact on the final yield of the product. Recent research into ester synthesis for undergraduate laboratories focuses on more transferable skills, including isolation of the final product and characterizing using 1H NMR, IR, gas chromatography or thin layer chromatography. This paper proposes the methylation reaction of 4-amino-3-nitrobenzoic acid for use in a typical 2-3 hour laboratory class. The reaction mixture is purified using a liquid-liquid extraction protocol that involves a salt-forming reaction. This extraction process is monitored visually with the carboxylic acid salt moving into the aqueous layer and the final ester retained in the ethyl acetate layer, allowing students to understand the practical approach of using liquid-liquid extraction. Once the extraction is complete, the ethyl acetate layer is filtered and concentrated in vacuo to yield a bright yellow solid. These are the key features that make this experiment a novel example for the teaching of chemistry. This laboratory experiment may be suited to being placed earlier in the curriculum, giving its colorimetric nature and ease, or used in a more advanced educational setting with the application of NMR analysis.
PEDAGOGICAL GOALS

Fundamental concepts of organic chemistry, such as solubility and polarity, are demonstrated in this experiment. Additionally, the pedagogical goals are:

- To introduce students to experimental organic synthesis techniques found in a traditional research laboratory
- To teach and consolidate understanding of the synthesis of ester compounds using Fischer esterification
- To demonstrate to students how to use thin-layer chromatography (TLC)
- To give students an opportunity to perform liquid-liquid extraction and understand the fundamentals of salt formation reactions
- To give practical skills to students on how to analyse and characterize organic compounds using IR, NMR, and melting point
- To provide experience to students on how to work safely in an organic chemistry laboratory

EXPERIMENTAL OVERVIEW

Scheme 1 – Synthesis of compound 2 (4-amino-3-nitrobenzoic acid methyl ester) from compound 1 (4-amino-3-nitrobenzoic acid) in neat methanol with catalytic sulfuric acid.
The process was completed in one laboratory period of 3 hours with students working in pairs. The NMR spectra and postlab questions were provided at the end of the laboratory session (see supplementary guides). The first part of this synthesis was the methylation of 4-amino-3-nitrobenzoic acid via Fischer esterification using methanol and concentrated sulphuric acid as a catalyst (Scheme 1). Students can monitor and confirm the formation of the product via TLC (with DCM as mobile phase, Figure 1) and visualize under UV light. Since the reaction is heated, in order to check the reaction progression, the reaction was allowed to cool slightly. Students monitored the reaction by TLC at 1 hour from commencement. After TLC, the reaction was quenched with saturated sodium bicarbonate solution and extracted into ethyl acetate, then filtered and dried to yield the final product. The final product’s melting point was determined and compared with literature.

![Figure 1: Silica gel thin-layer chromatography (TLC) with fluorescence indicator (f254) for visualization under UV light. S = starting compound (4-amino-3-nitrobenzoic acid), RXN = reaction mixture (monitored during the reaction), and P = product (4-amino-3-nitrobenzoic acid methyl ester).](image)

**HAZARDS**

The solvents used in this experiment, methanol and ethyl acetate are highly volatile, flammable, and of moderate toxicity. Dichloromethane is highly volatile, and excessive DCM could cause drowsiness and has moderate toxicity. 4-amino-3-nitrobenzoic acid and 4-amino-3-nitrobenzoic acid methyl ester (methyl 4-amino-3-nitrobenzoate) are irritants and may cause skin, eye, and respiratory irritation. Sulphuric acid is corrosive. The synthesis
step requires the heating of a flammable solvent. The reaction must be conducted in a fume extraction hood, and fire extinguishing equipment must be in place. Organic and aqueous waste should be disposed of correctly. Gloves, lab coat, and appropriate eye protection are required.

RESULTS AND DISCUSSION

Reaction Time

To establish a reasonable reaction time for an undergraduate laboratory, the reaction mixture was heated at reflux for 30 minutes, 1 hour, 2 hours, and 16 hours (yields detailed in Table 1). A reaction of 1 hour represented reasonable yields for isolation of the final compound and was an appropriate time for the undergraduate laboratory session.

Table 1 Correlation of Reaction Time and Percentage Yield

<table>
<thead>
<tr>
<th>Reaction Elapsed Time</th>
<th>Percentage Yield</th>
</tr>
</thead>
<tbody>
<tr>
<td>30 Minutes</td>
<td>20%</td>
</tr>
<tr>
<td>1 Hour</td>
<td>46%</td>
</tr>
<tr>
<td>2 Hours</td>
<td>50%</td>
</tr>
<tr>
<td>16 Hours</td>
<td>95%</td>
</tr>
</tbody>
</table>

Visual Reaction Workup

The workup procedure utilizes a salt-forming reaction along with polarity differences, and was visually monitored by the students. The starting material, compound 1, 4-amino-3-nitrobenzoic acid, contains a carboxylic acid, which reacts with sodium bicarbonate to form the corresponding sodium salt (shown below, Figure 2), and becomes soluble in the aqueous phase. The product, 4-amino-3-nitrobenzoic acid methyl ester, is not affected by the sodium bicarbonate and remains highly soluble in ethyl acetate and remains into the organic phase during the liquid-liquid extraction.

\[
\begin{align*}
\text{Aqueous (NaHCO}_3\text{) Soluble} & & \text{Ethyl acetate Soluble}
\end{align*}
\]
Once the reaction time was finished, the reaction was cooled, and a solution of saturated sodium bicarbonate was added to quench the reaction. From there, the organic phase was washed with the saturated sodium bicarbonate solution for a total of four times, or until the aqueous phase had become clear (Figure 3). Since the sodium carboxylate is also colored, its removal was monitored in this way. Students were thus able to monitor the purification progress of this reaction visually. Once the ethyl acetate layer was extracted, it was filtered through a silica plug and then concentrated in vacuo to yield the final product as a bright yellow solid.

Figure 3: Liquid-liquid extraction of 4-amino-3-nitrobenzoic acid methyl ester with ethyl acetate (top layer) and saturated sodium bicarbonate (bottom layer). Demonstrating the disappearing yellow colour of the bottom layer after successive washes with sodium bicarbonate.

Analysis and Elucidation of Structure

The final methyl ester compound, 4-amino-3-nitrobenzoic acid methyl ester was elucidated by both $^1$H NMR and $^{13}$C NMR, and the melting point determined. The melting point range was determined to be 188-191°C for the final product compared to 290-297°C for the starting 4-amino-3-nitrobenzoic acid, which is consistent with the literature. The $^1$H NMR spectra was provided to the students at the end of the laboratory session and showed five proton environments represented by five peaks. Due to the types of hydrogen environments within the compound, a low field benchtop NMR would be able to distinguishable them and thus elucidate the structure. When comparing the product to the starting material, there is a new singlet peak observed at 3.80 ppm representing the ester
methoxy CH$_3$ which was introduced in the reactant and not observed in the starting material (Figure 4). The other peaks at 7.04, 7.82, and 8.52 ppm elucidate the benzene ring with the doublet at 7.04 ($J = 8.95$ Hz) indicating a meta-proton, the doublet of doublets at 7.82 ppm ($J = 2.10, 8.95$ Hz) indicating an ortho-proton, and the final doublet at 8.52 ppm ($J = 2.01$ Hz) indicates the remaining proton. The broad peak at 7.97 ppm is indicative of an aromatic amine. On a 400 MHz NMR, the proton $J$ coupling values were determined, which corresponded to the expected values for an ortho- meta- and para- protons.

![Figure 4: 1H NMR of 4-amino-3-nitrobenzoic acid methyl ester at 400 MHz in DMSO-d$_6$.](image)

The carbon spectrum shows eight carbon peaks which correspond to the methoxy carbon at 51.9 ppm, and the aromatic region carbons at 116.1, 119.3, 128.0, 129.6, 134.7, and 148.8 ppm. The other significant peak is the carbonyl at 164.9 ppm (instructors notes).
CONCLUSION
In conclusion, this reaction is simple and cost-effective as a teaching reaction. The purification, liquid/liquid extraction process, provides a visual opportunity for students to track the salt-forming reaction and consolidate their understanding. Students can assign proton and carbon NMR or IR as well as determine the melting point and compare it with the starting 4-amino-3-nitrobenzoic acid. The experimental outcomes are consistent with existing curricula and offers a unique visual organic chemistry laboratory experience.

ASSOCIATED CONTENT
Supporting Information
The Supporting Information is available on the ACS Publications website at DOI: 10.1021/acs.jchemed.XXXXXXX. [ACS will fill this in]

[Instructor notes] including:

1. Chemical materials for experiments: reagent and solvent tables
2. Hazards and Safety Instructions
3. Instructor notes
4. Results for post lab questions (including the full assignment for \( ^1 \)H NMR and \( ^{13} \)C NMR spectra).

[Student laboratory handout] including:

1. Safety instructions
2. Experimental procedure
3. Postlab questions.

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REFERENCES


