

# Agilent Cary 630 FTIR Spectrometer Supporting Organic Synthesis in Academic Teaching Labs

Application note

Academic

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## Introduction

As it has been for the past half century, IR spectroscopy is a fundamental analytical technique that undergraduate organic chemistry students must understand and have the opportunity to use. The demands of the undergraduate labs require that FTIR spectrometers possess certain design attributes associated with the multiuser environment. The spectrometer must be easy to use, rugged, reliable, compact and cost attractive.



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The recently introduced Cary 630 system (Figure 1) meets and exceeds these requirements and is an excellent choice for the undergraduate organic lab. In addition, it is equally at home in graduate level and research labs where organic chemists determine the identity and structure of synthesized compounds via a rapid IR analysis.



**Figure 1.** The Agilent Cary 630 FTIR spectrometer (all photographic elements shown in context)

## Agilent Cary 630 FTIR Spectrometer – features and benefits for academic organic synthesis applications in teaching and routine research support

### Overall System Design

- Most compact, superior performing FTIR available - ultracompact design preserves valuable bench top space
- Light weight allows system to be relocated, as needed
- Completely sealed optics and compact size enables use in standard fume hoods

- Broad range of sampling interfaces available, handling liquids, solids and gases
- Overall performance and ease of use advantages translate into virtually all analyses requiring less than a minute to execute

### Optomechanical Components

- Permanently aligned interferometer and optical system results in an extremely reliable, everyday “workhorse” system
- Interferometer and optics are exceedingly rugged – Cary 630 uses the same optomechanical components as those used in Agilent’s portable and handheld industrial systems, which require the highest level of robustness
- Large aperture optics and short internal optical paths provides class-leading performance

### Sampling Interfaces

- No-alignment, interchangeable sampling accessories allow students to understand and readily practice different experimental methods – transmission, ATR etc.
- Sampling interfaces available include: diamond ATR, diffuse reflectance, solid, liquid and gas transmission as well as Agilent’s innovative, exclusive DialPath technology for liquids analysis
- Diamond ATR sampling interface is ideal for analysis of reaction starting materials, reagents and product – impervious to scratching and highly chemically resistant (pH 1 -14)
- ATR sampling interface enables grab sample, neat reaction mixture analysis – no sample dilution necessary
- Powder press, which ensures good optical contact of solids with ATR crystal, cannot be overtightened, i.e. diamond window cannot be damaged by overpressure
- Sample interface RFID ensures that methods are matched with appropriate sampling technologies

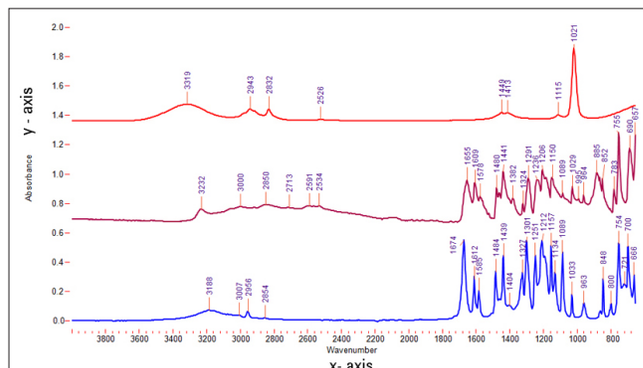
## Software and User Interface

- Highly visual, intuitive software allows data acquisition with essentially no training required.
- Real-time spectrum display works in conjunction with ATR powder press to ensure adequate contact between solid samples and diamond ATR crystal.
- Experimental results can be stored under individual student's name or other identifier
- Student can verify identity of starting materials, isolated intermediate compounds and final products by comparison with spectra in on-board IR spectral library
- For more advanced experiments or analysis, data is readily transferrable to commercially available data analysis package.
- On-board diagnostics ensure peak operating efficiency

## Application example: Syntheses of esters of o-hydroxybenzoic acid

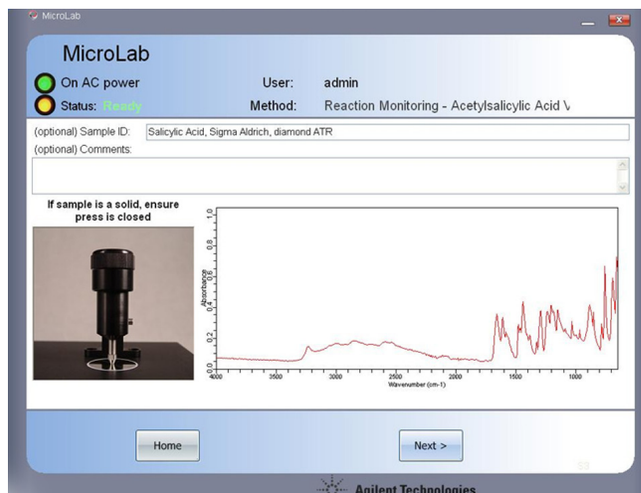
A classic undergraduate organic chemistry experiment, which is also ideal for FTIR analysis, is the synthesis of o-hydroxybenzoic acid esters, i.e. the condensation reaction of salicylic acid with methanol to form methyl salicylate (oil of wintergreen), and the esterification of salicylic acid with acetic anhydride to form acetylsalicylic acid (aspirin). This lab synthesis is particularly useful as a teaching tool because the synthesis is associated with product isolation, purification and the analysis of final product for purity and yield. Melting point, FTIR analysis and proton NMR are often employed in support of this teaching experience, so the student receives an understanding of how synergistic analysis provides a better overall understanding of chemistry. With respect to FTIR analysis, these reactions allow the student to measure solids, liquids and slurries using a variety of IR sampling methods.

In the preparation of methyl salicylate, the starting materials and final product have highly characteristic infrared spectra (Figure 2). The diamond ATR sampling



**Figure 2.** The IR spectra of methanol (red), salicylic acid (maroon), and methyl salicylate product (blue).

interface is ideal for the measurement of these compounds, which are solids and liquids. For methanol reagent, the student simply puts a drop of the liquid on the diamond sensor and initiates analysis. A high quality spectrum only requires 2 seconds to collect and the system's rapid analysis capability is useful when volatile solvents or reagents are analyzed. For the solid salicylic acid starting material, a few grains are placed on the diamond sensor; the powder press is rotated downwards until enough contact is made so that a spectrum appears in the real-time spectrum display (Figure 3). After the reaction is complete, a drop of the oily product is placed on the diamond sensor



**Figure 3.** Powder press ensures that solid sample is in contact with diamond ATR crystal to obtain high quality IR spectrum. Powder press design eliminates overtightening. Instantaneously refreshed spectra obtained in real-time analysis window shows that solid sample is in contact with the Diamond ATR crystal.



## Synthesis of o-hydroxybenzoic acid esters – Use of infrared spectroscopy and analysis of spectra

If the mixture that is formed from the reaction of salicylic acid and methanol is dried down and analyzed without purification, the students will observe the methyl CH stretch absorbance band at  $2956\text{ cm}^{-1}$  ( see Figure 2) for methyl salicylate, along with bands from the unreacted salicylic acid starting material. Bands that the students should watch for is the loss of the salicylic acid broad OH---O stretch absorbance in the region from  $2800\text{-}2200\text{ cm}^{-1}$  and the OH---O wag absorbance at  $885\text{ cm}^{-1}$ . If the reaction mixture is not dried down, but rather further heated on a hot plate at  $100^{\circ}\text{C}$ , methyl salicylate will condense on the sides of the vial. This clear oily condensate can be collected with a swab or plastic pipet for measurement on the diamond crystal for IR analysis. Very little of the liquid is necessary, the 1.5 mm diameter diamond only needs a thin film on its surface, to produce an excellent infrared spectrum of the product.

The carbonyl region of the IR spectra of salicylic acid, methyl salicylate, and acetylsalicylic acid makes for an interesting study on how IR can be used to characterize hydrogen bonding influences on carbonyl

group frequencies. Hydrogen bonding lowers the carbonyl group frequency in the IR spectrum and under normal conditions, the carboxylic acid group forms a H-bonded dimer at  $1710\text{-}1700\text{ cm}^{-1}$ . The carbonyl frequency can shift as high as  $1750\text{ cm}^{-1}$  when analyzed in the gaseous state (no H-bonding is possible) or in very dilute solution in a non-polar solvent. The carboxylic acid carbonyl band (COOH) for salicylic acid (Figure 4) is lower in frequency ( $1655\text{ cm}^{-1}$ ) than that in acetylsalicylic acid ( $1678\text{ cm}^{-1}$ ). This is due to the slight very weak hydrogen bonding influence from the phenolic OH group present in the salicylic acid and not in the aspirin, which is enough to disturb the dimer found in normal non-influenced aromatic COOH. Benzoic acid which has no phenolic OH group has the carboxylic acid dimer absorbance at  $1678\text{ cm}^{-1}$ , nearly identical to that in acetylsalicylic acid. Interestingly, the methyl ester in methyl salicylate (Figure 2) is at the same frequency ( $1678\text{ cm}^{-1}$ ) as the carboxylic acid groups. This is due to the slight hydrogen bonding from the phenolic OH group, which is either intramolecular, intermolecular, or a mix of both H-bonded states, since the normal aromatic methyl ester without a phenolic OH group should be much higher at  $\sim 1730\text{ cm}^{-1}$ , such as in orthophthalates.

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