FT-IR Training Notebook: ATR

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November 14, 2019 (rev. 2.1)

Before you begin...

- Receive a user name and temporary password for Faces scheduling
- Identify your ENGR username and Password from Systems
 - □ If you don't have an ENGR account, send me the following:
 - Full name
 - Principal Investigator (PI)
 - UCR NetID
 - 🖵 email
- Coordinate a time with the lab manager for training
- Schedule a 1 hour block on Faces for your training

FT-IR Operation

- I. Pressure Tower Setup
- II. Initiate Software
- III. Collect Background
- IV. Sample Prep: Solids
- V. Sample Prep: Liquids
- VI. Collect Sample
- VII. Saving Data
- VIII. Peak Identification
- IX. Cleanup
- X. Library Search
- XI. Smart Transmission Accessory

I. Pressure Tower Setup – 1/2

- 1. To adjust the position of *Pressure Tower:*
 - Turn Knob counter-clockwise = raise Tower
 - Turn Knob clockwise = lower Tower



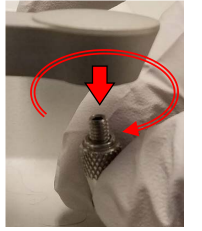
- 2. Inspect the *Pressure Tip* by moving *Tower Arm* to *Cleaning Position*
 - Move *Tower Arm* to the right until it stops





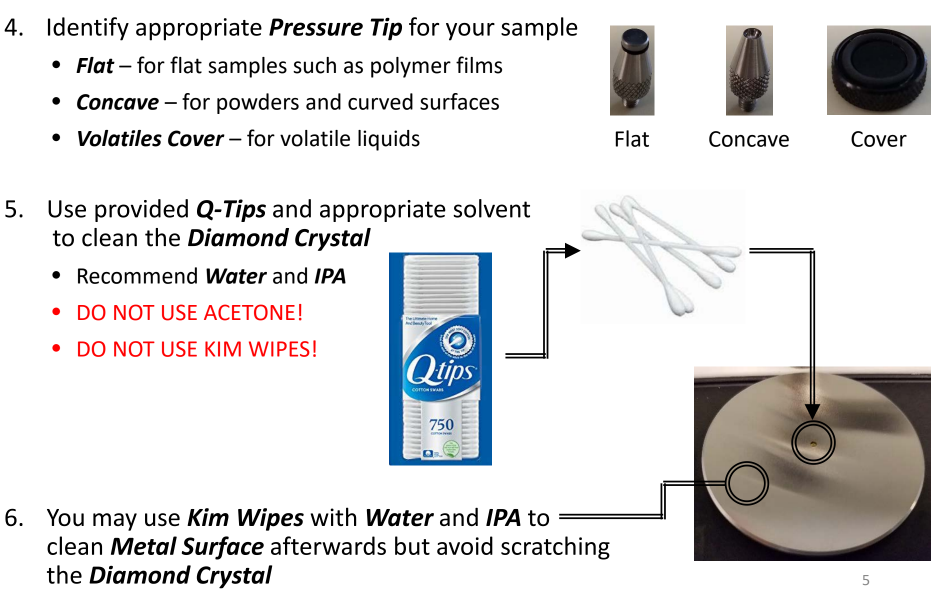
Cleaning Position

- 3. Clean the *Pressure Tip* (remove if necessary) with appropriate solvent
 - Recommend Water and IPA
 - DO NOT USE ACETONE!
- 4. To remove/install *Pressure Tip:*
 - Rotate *Tip clockwise = remove*
 - Rotate *Tip counter-clockwise = install*





I. Pressure Tower Setup – 2/2



II. Initiate Software – 1/10

- 1. Double left-click on the *OMNIC software icon* for FT-IR
- Ignore the Standards Expiration Warning and click OK
 Close the Thermo Scientific OMNIC Help popup window

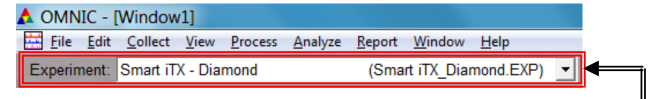
Contents Index Search

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Options

Print

4. The OMNIC main window will now appear



Confirm that *Smart iTX – Diamond (Smart iTX_Diamond.EXP)* appears in the Experiment window

Hide



A standard is expired

A standard you are using for Performance Verification or ValPro Qualification

II. Initiate Software – 2/10 MIC - [Withow1]

- 6. Select *Collect -> Experiment Setup* at the top window
- 7. Confirm that *Save interferograms* is *checked* =
 - Saving interferogram data lets you reprocess in case you want to restore the original data, even using a different background or changing parameter settings used
- 8. Confirm that *Save automatically* is *unchecked*
- 9. Set preferred *Background Handling* settings ———
 - Before every sample
 - After every sample
 - After 120 minutes (default)
 - Use specific file

Note: A new background will be requested if there is a change in resolution or data spacing of your sample spectrum!

| Collect | Bench Quality | Advanced Diagnostic | Canfigure |
|--|---|-----------------------|--|
| Estimated time for this collection: 00:00:23 No. of scans: 16 | | | Save automatically |
| | Resolution: 4. | | Base name: C:\My Documents\Omnic\autosave\Dongwei1908080 |
| | ata spacing: 1 Final format: A | | Background Handling O Collect background before every sample |
| | Correction: None Automatic atmospheric suppression Preview data collection Use transmittance data during preview | | Collect background after every sample Collect background after 120 minutes |
| | | | Collect background after 120 minutes Use specified background file: Browse |
| - | | its in collect window | Collect 64 scans for the background |
| ٨ | Min: 0.00 | Max: 2.00 | Experiment description: |
| Experiment title: Smart iTX - Diamond | | | Smart iTX Accessory with Diamond Crystal |

Experiment: Smart iTX - Diamond

II. Initiate Software – 3/10

10. Select desired No. of scans - recommend starting at 16 scans

- Increase to optimize desired spectrum signal/noise
- Recommend increments of powers of 4 (e.g. 16, 64, 256, 1024,...)
- 11. Select desired *Resolution value* recommend 8 or 4
 - Decrease value to increase spectrum resolution
 - Decreasing value too much may result in increased noise!
 Note: *Aperture* = *High resolution* if Resolution value is ≤ 2
- 12. Check Estimated time for collection
 - Time dependent on *No. of scans* and *Resolution*
- 13. Select desired Final format
 - % Transmittance
 - Absorbance (default)
 - Etc...

Note: Convert to other Y-axis units in *Process* menu

| ollect | Bench | Quality | Advanced | Diagnosti |
|-----------------------|-------------|-------------|----------------|-----------|
| Estin | nated tin | ne for this | s collection: | 00:00:23 |
| 1 | No. of sc | ans: 16 | | |
| | Resolu | tion: 4. | | • |
| D | lata spac | ing: 1.9 | 29 cm-1 | |
| | Final for | mat: Abs | sorbance | • |
| | Correct | tion: No | ne | - |
| Au | tomatic a | tmosphe | ric suppress | ion |
| Pre | eview dat | a collectio | on | |
| Us | e transm | ittance da | ata during pr | eview |
| Us | e fixed Y- | axis limits | s in collect w | vindow |
| 1 | Min: 0.0 | 0 | Max: 2 | .00 |
| Expe | riment ti | tle: | | |
| and the second second | t iTX - Dia | amond | | |

II. Initiate Software – 4/10

- 14. Select desired *Correction type* to *None*
- 15. Decide if *Automatic atmospheric suppression* is desired
 - Effects of water vapor and carbon dioxide will be automatically suppressed via quantitative model
- NOTE: Do NOT use this feature if atmospheric conditions change very slowly, only use if conditions change rapidly
- 16. Check *Preview data collection*
 - Views preliminary data before start of sample for verification
- 17. Decide if you want to preview data collection using % transmittance
 - May provide an improved preview of the data
- 18. Decide if fixed Y-axis limits will be used in the preview
 - Recommend using Min: -5% to Max: 105%

| Conect | Bench Qua | lity Advanced | Diagnos |
|--------|------------------|--------------------|----------|
| Estin | nated time for | this collection: | 00:00:23 |
| 1 | lo. of scans: | 16 | |
| | Resolution: | 4. | - |
| D | ata spacing: | 1.929 cm-1 | |
| | Final format: | Absorbance | • |
| | Correction: | None | • |
| Aut | omatic atmos | pheric suppress | ion |
| V Pre | eview data coll | ection | |
| Us | e transmittanc | e data during pr | eview |
| Us | e fixed Y-axis I | imits in collect w | indow |
| 1 | Vin: 0.00 | Max: 2 | 00 |
| Expe | riment title: | | |
| Smar | t iTX - Diamon | d | |
| | | | |

II. Initiate Software – 5/10

- 19. Select Bench tab
- 20. Confirm that the following are correct:
 - Source = IR
 - Accessory = Smart iTX
 - Window = Diamond
- 21. Select desired *Max* and *Min* range limit for your scans
 - Recommend using *Recommended range*
- 22. Select the *Gain* parameter
 - Electronically amplifies signal recommend Autogain
 - DO NOT set to Autogain if performing quantitative analysis
- 23. Select the desired Aperture
 - *High resolution* used with resolution at 2 or less for better stability and accuracy
 - *Medium resolution* recommended with resolution 4 for better Signal/Noise
- 24. Confirm the *Screen wheel* is set to *Open*

| Parameter | Value | |
|--------------------|-------------------|---|
| Sample compartment | Main | • |
| Detector | DTGS KBr | • |
| Beamsplitter | KBr | • |
| Source | IR | • |
| Accessory | Smart iTX | • |
| Window | Diamond | • |
| Recommended range | 4000 525 | |
| Max range limit | 4000 | |
| Min range limit | 400 | |
| Gain:8 | Autogain | • |
| Optical velocity | 0.4747 | • |
| Aperture | Medium resolution | • |
| Sample shuttle | | |
| Screen wheel | Open | + |

Bench Quality Advanced Diagnostic Configure Collect

II. Initiate Software – 6/10

25. Select Quality tab

Collect Bench Quality Advanced Diagnostic Configure

26. Determine if you want any spectral quality characteristics to be checked during

YOUR SCANS Select view:

Spectrum
Parameter
Background
Interferogram
All

- Spectrum checks quality of the spectrum scan
- *Parameter* checks the scan parameters
- **Background** checks the quality of the background scan
- *Interferogram* checks the raw interferogram signal
- All checks all the above characteristics
- 27. If you choose to check *Spectrum*...
 - Peaks present? checks for peaks and if sample is positioned correctly, recommend ON at 50%
 - Totally absorbing peaks checks for absorbing peaks, recommend ON at 50%
 - Fringes or channeling checks for back reflection inside sample, recommend ON at 50%
 - Derivative peaks checks for derivative-shaped peaks, recommend ON at 50%
 - Baseline error checks for baseline problems, recommend ON at 50%
 - **CO**₂ levels checks for CO₂ absorption, recommend **ON** at **50%**
 - H₂O levels checks for H₂O absorption, recommend ON at 50%

II. Initiate Software – 7/10

28. If you chose to check *Parameters*...

- **Spectral range** checks if spectral range is consistent for the hardware, recommend **ON**
- Apodization correct checks apodization type is appropriate, recommend ON
- *Resolution* checks if resolution is appropriate for the experiment, recommend *ON*

29. If you chose to check *Background*...

- Contamination peaks checks for contaminants, recommend ON at 50%
- Detector icing checks signs of detector icing, recommend NO
- **CO₂ levels** checks for CO₂ absorption, recommend **ON** at **50%**
- H₂O levels checks for H₂O absorption, recommend ON at 50%
- Background correct for accessory checks background spectrum, recommend ON at 50%
- 30. If you chose to check *Interferogram*...
 - *Peak amplitude within range* checks if amplitude is sufficient, recommend *ON*
 - Interferogram minimum = 0.20 and Interferogram maximum = 9.80
 - Minimum peak above noise checks if peak signal is above noise level, recommend ON
 - Peak Minimum = 10

II. Initiate Software – 8/10

31. Select **Advanced** tab

Collect Bench Quality Advanced

Diagnostic Configure

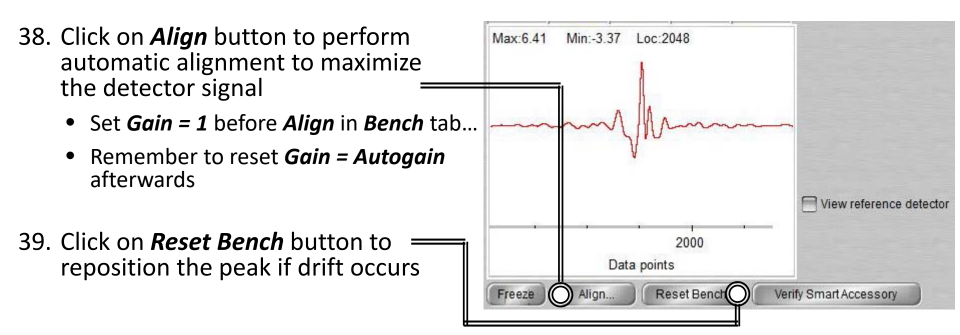
- 32. Confirm *Zero filling* is set to *None*
- 33. Confirm *Apodization* is set to *Happ-Genzel*
- 34. Confirm *Phase correction* is set to *Mertz*
- 35. Confirm that the following are checked:
 - Set sample spacing based on spectral range
 - Set filters based on velocity

| xperiment Setup - C:\My Docu Collect Bench Quality ^{Ad} | 500 B. C. B. | 25150 |
|---|--|-------|
| Zero filling: | None | - |
| Apodization: | Happ-Genzel | - |
| Phase correction: | Mertz | - |
| ☑ Set sample spacing ba | sed on spectral rang | le |
| Sample spacing: | 1.0 | - |
| Set filters based on velo | ocity | |
| Low pass filter: | 11000 | * |
| High pass filter: | 20 | - |
| Single-sided interferogr | am | |
| Reset bench at start of o | collection | |
| Start collection at extern | al trigger | |
| Help Open | Save Save | As) |

II. Initiate Software – 9/10

36. Select *Diagnostic* tab

- 37. Click on indicators to check spectrometer components
 - If the values are within the Acceptable Range, they will appear as a
 - If any values show X, contact the Lab Manager immediately!



Collect Bench Quality Advanced Diagnostic Configure

-Q-

II. Initiate Software – 10/10

40. Select *Configure* tab

Collect Bench Quality Advanced Diagnostic Configure

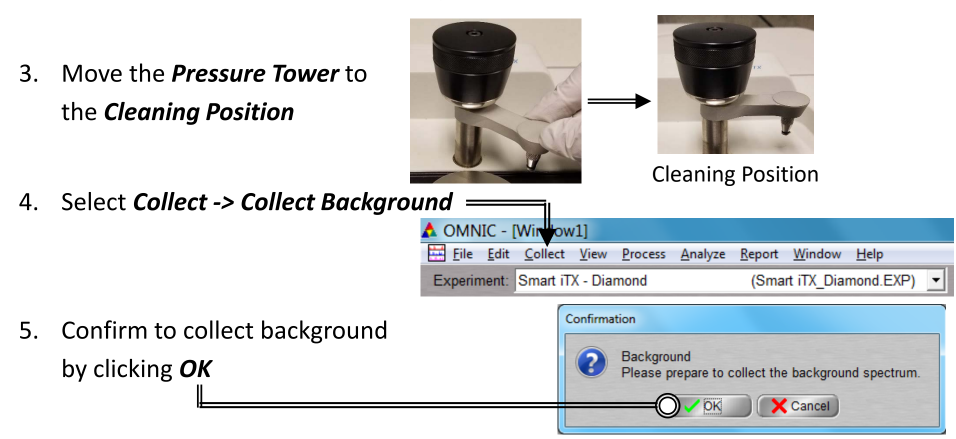
- 41. Confirm *Inactivity Rest mode* is checked
 - Confirm *Hours of inactivity* is set to "1" hour
- 42. Confirm *Daily Rest mode* is not checked

| V Inactivity Rest mode Hours of inactivity: 1 | |
|--|----------|
| Daily Rest mode | |
| Exit Rest mode: 5:30 AM | |
| Start Rest mode: 6:00 PM | |
| Rest days: Su Mo Tu We Th Fr Sa V IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII | |
| | OK Cance |
| Help Open Save O Save As | |

43. Click "*Save*" then "*OK*"

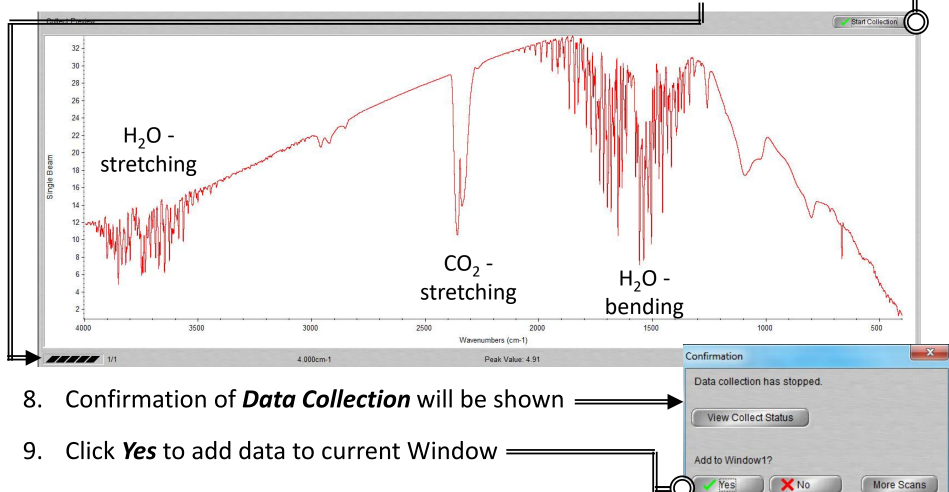
III. Collect Background – 1/2

- 1. It is critical that the *Crystal* is cleaned **BEFORE** *Background* is collected!
- 2. A single *Background* can be used to analyze multiple samples, but it is recommended to collect new *Background* at least every 2 hours



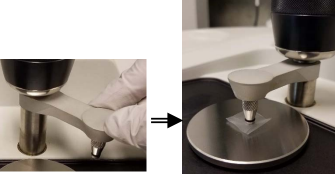
III. Collect Background – 2/2

- 6. Preview **Background Collection** then click **Start Collection** to begin =
- 7. The *Background Collection* will begin with the progress shown at the bottom



IV. Sample Prep: Solids – 1/1

- 1. For **Solid** and **Thin Films** use **Flat Tip** and for **Powder** use **Concave Tip...**
- 2. Ensure the *Flat* or *Concave Pressure Tip* is installed first
- 3. Place sample onto *Crystal*
- 4. Move the *Pressure Tip* into *Sampling Position*



Sampling Position

- 5. Lower the *Pressure Tower* to press the *Sample* against the *Crystal*
- 6. The *Pressure Tower Knob* will *Click* and *Freely Rotate* when the maximum pressure is reached

V. Sample Prep: Liquids – 1/1

- 1. For *Liquid*, *Paste*, or *Gel Sample*...
- 2. Move the *Pressure Tip* into *Cleaning Position* and dispense sample onto *Crystal*

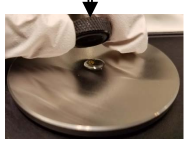




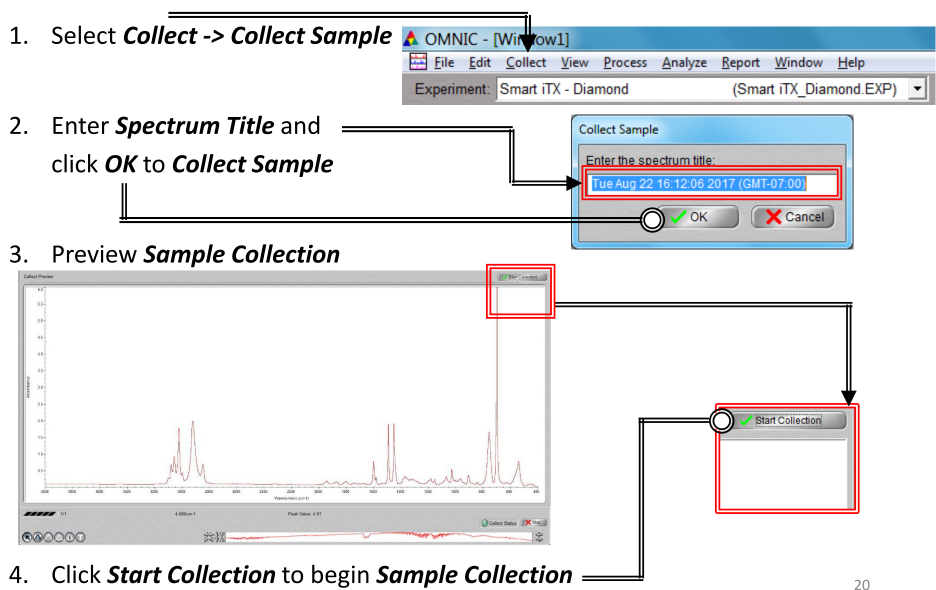
Cleaning Position

- 3. The sample should cover the *Crystal* but DO NOT OVERFILL or else the sample will run off the *Crystal Plate*
- 4. For *highly volatile samples*, place *Volatiles Cover* over sample to reduce of evaporation
 - Install *Flat Pressure Tip*, move into *Sampling Position*, and lower the *Pressure Tower* until the *Pressure Tower Knob Clicks* and *Freely Rotates* when the maximum pressure is reached



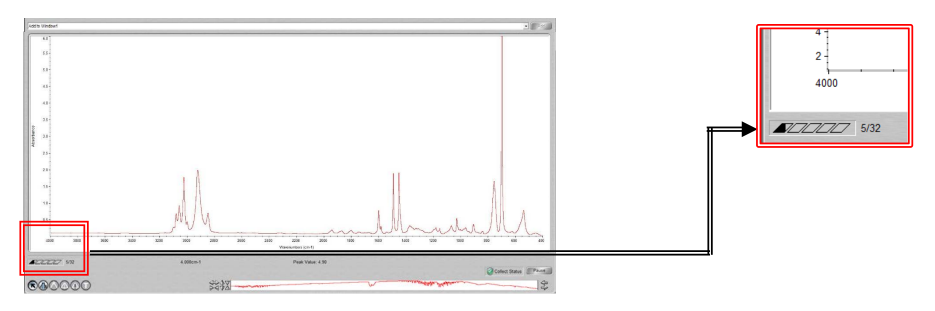


VI. Collect Sample – 1/2



VI. Collect Sample – 2/2

5. The Sample Collection will begin with the progress shown at the bottom

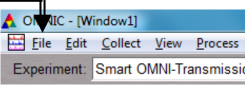


- 6. Confirmation of *Data Collection* will be shown
- 7. Click **Yes** to add data to current Window

| Data collection has stoppe | ed. |
|----------------------------|-----|
| View Collect Status | |
| Add to Window1? | |

VII. Saving Data – 1/1

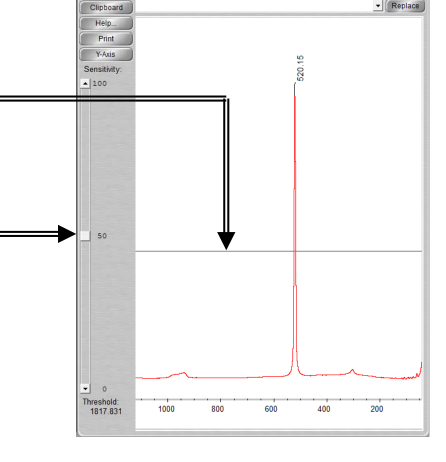
- 1. Specific spectra can be selected using the 🕟 selection tool at the bottom of window and clicking on it or selecting from dropdown box 🗊 No spectra selected
- 2. Multiple spectra can be selected/deselected by holding down the *Ctrl* key and clicking spectra
- Click *File -> Save* to save a spectrum (e.g. default is SPA) using the current filename
- Click *File -> Save As* to save a spectrum into another file type (e.g. CSV or TIFF)
- Click *File -> Save Group* to save more than one spectrum as a group in one file having file extension .SPG to open later



VIII. Peak Identification – 1/1

1. Click on "*Find Pks*" button at the top _____

- Click the spectrum window to adjust the *Threshold* position on where peaks are to be considered ===
- 3. Adjust the *Sensitivity* button to separate peaks from noise _____



Find Pks

IX. Cleanup – 1/1

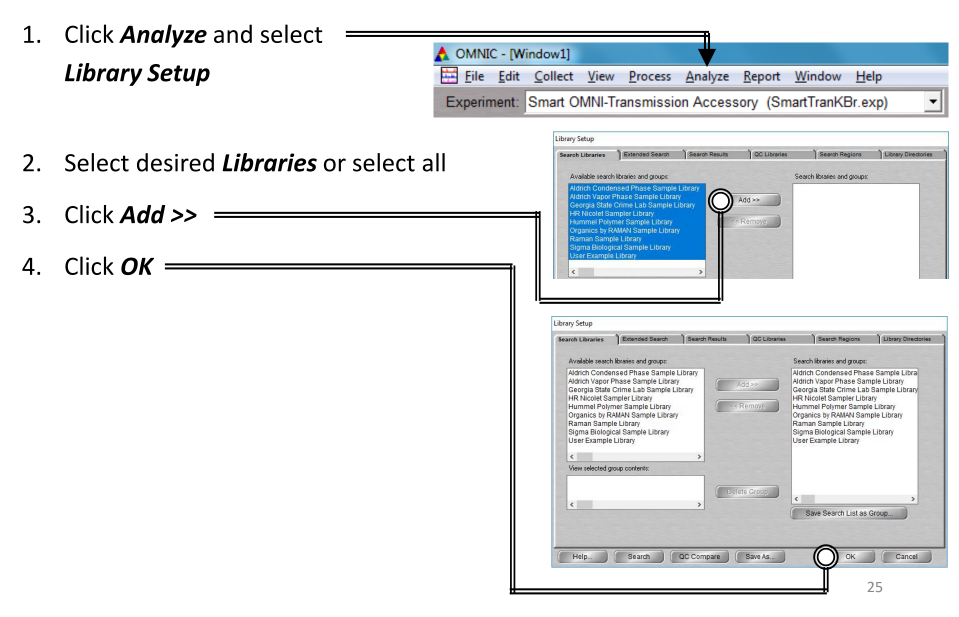
- 1. Remove *Sample* from the *Crystal* without scratching the *Crystal*
- 2. Use provided *Q-tips* and appropriate solvent to clean the *Crystal*
 - Recommend Water and IPA
 - DO NOT USE ACETONE!
 - DO NOT USE KIM WIPES!
- 3. Clean the *Pressure Tip* (remove if necessary) and *Metal Surface* with appropriate solvent and *Kim Wipes*
 - Recommend Water and IPA
 - DO NOT USE ACETONE!



- 4. Click on *File -> Exit* to shut down the software
- 5. Log off of your ENGR account

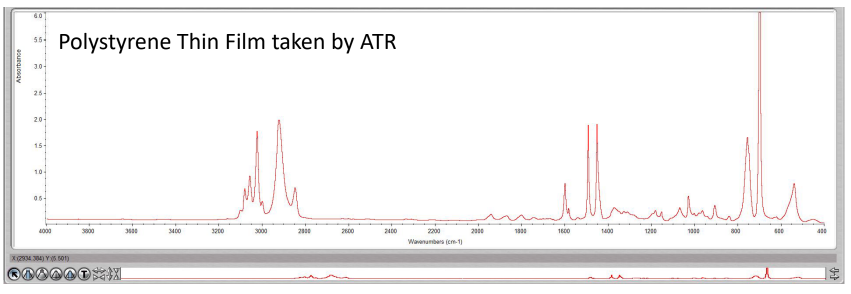


X. Library Search – 1/5



X. Library Search – 2/5

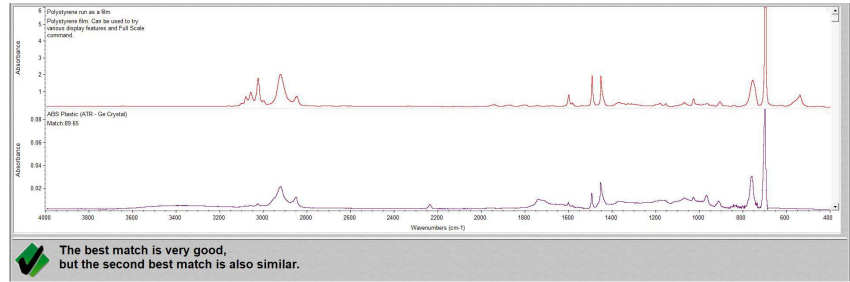
5. Select the desired spectra you wish to search for a library match



6. Click Analyze and select Search...
 or click Search icon

X. Library Search – 3/5

7. The top matches will be shown (below) your acquired spectra (top)



8. Click View Match List and select either Overlay or Stack view

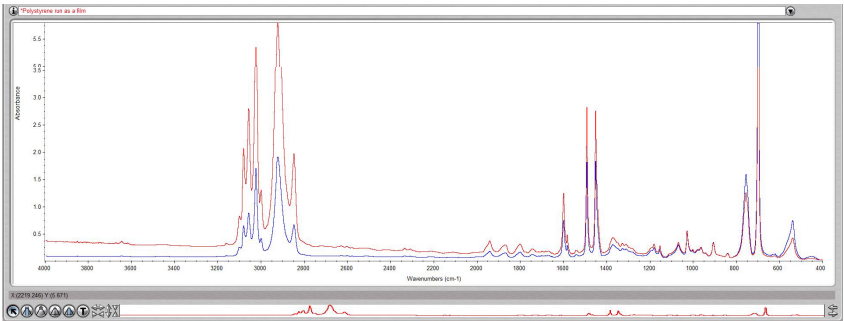


9. Perform *ATR Correction* to achieve better match results

| .0. Click Process > Other Corrections and select ATR | Other Corrections |
|---|----------------------|
| | Select a correction: |
| Monte - [Window1] | ATR |
| <u>File Edit Collect View Process Analyze Report Window Help</u> Experiment: Smart OMNI-Transmission Accessory (SmartTranKBr.exp) | OK Cancel |

X. Library Search – 4/5

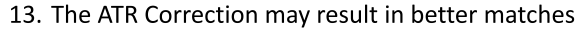
11. The ATR Corrected spectra will be created and marked with a *

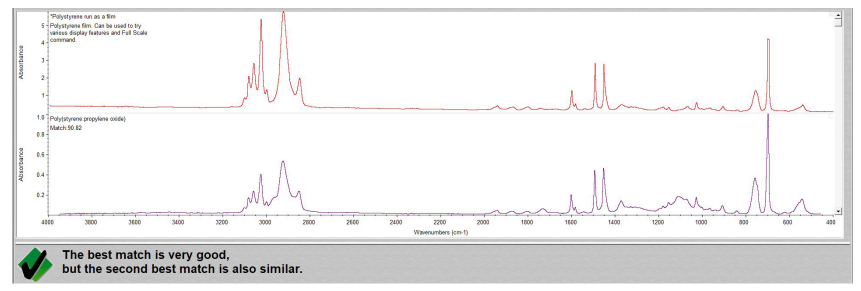


12. Click **Analyze** and select **Search...** or click **Search** icon Search

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X. Library Search – 5/5



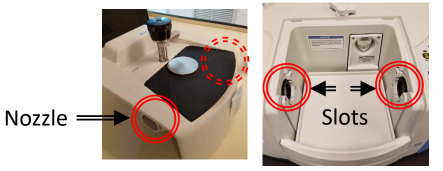


14. If a Match does not result, you will have to find matching spectra online instead

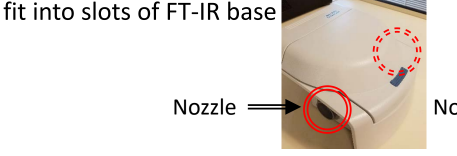
XI. Smart Transmission Accessory – 1/3

- The Smart ATR Accessory is the default accessory installed 1.
- 2. Please contact the Lab Manager if you need to use the *Smart Transmission Accessory* for Transmission FT-IR measurements
- The *Smart ATR Accessory* contains mirrored optics that need = 3. to be carefully taken care to avoid damage and contamination
- Both **Smart ATR Accessory** and **Smart Transmission Accessory** have nozzles to 4.





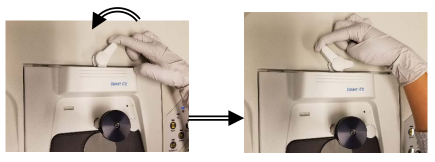


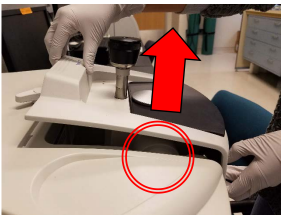




XI. Smart Transmission Accessory – 2/3

- 5. To remove the *Smart ATR Accessory,* move the lock to the *Unlocked* position
- Carefully remove *Smart ATR Accessory* by gently pulling upward and position nozzles out of slots
- 7. Carefully place aside and KEEP AWAY FROM CONTAMINANTS!
- 8. Carefully insert the *Smart Transmission Accessory* by gently aligning the nozzles into the slots







XI. Smart Transmission Accessory – 3/3

 Once firmly seated into the FT-IR base, move the lock to *Locked* position



10. Remember to remove *Smart Transmission Accessory* and reinsert the *Smart ATR Accessory* before leaving...





