

ABB MB3000 FT-IR Spectrometer Instructions

SAMPLE PREPARATION and SOFTWARE ACCESS:

- Obtain Spec. grade CH₂Cl₂ in the hood closest to the instrument. Prepare a concentrated solution of your analyte and apply to a salt plate. Salt plates can be found in a dessicator near the instrument.
- Access the FT-IR software by double clicking on the Horizon HB icon on the desktop.
- Enter the following information:

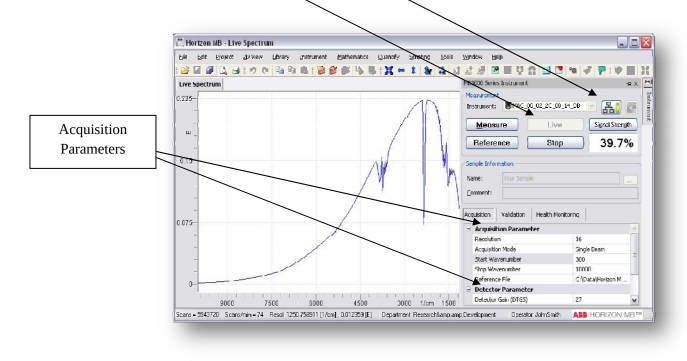
User Name: FTIR

Password: AJM328A.

• If asked, enter **chemistry** as the department name.

CHECKING SPECTROMETER COMMUNICATION:

- Click on **Acquisition** in the **Instrument** tab of the toolbar.
- Click on the **Connection** button in the Instrument panel.
- Click on **Live** and monitor the instrument signal strength.
- In the Acquisition Parameter window, set the Resolution according to application needs and adjust the Detector Gain value to adjust Signal Strength between 25% and 90%.





• Click on **Stop** to exit live mode.

SAVING THE DATA:

• Set the location to save data by clicking on the ... button:

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MB3000 Series	Instrument		-== :	н4 ×
Measurement Instrument:		00_02_2C_00_14_DE	· · A. C	Instrument
Measu	ire (Live	Signal Strength	
Referen	nce	Stop	39.7%	
Sample Inform	nation			
<u>N</u> ame:	Your Samp	ble		
Comment:				

Figure 2. Spectrum Acquisition Screen.

• Enter the **Sample Name** and make sure the **Path** and **Project** folder are defined as follows: Path: C:\Users\ABBBomem\Documents\horizon MB Data

Project: YOUR FOLDER'S NAME

ditional Sample Information			
Name Value			Configure
Comment			
to Save Options			
Path. C:\Users\ADDDor	mem\Documents\	Llonzon MB Data	
Save Data to File:	File Format.	ABB FTSW100 (*.apcctrum)	-
Save Data to Project:	Project:	Measured Data	
	Folder:	Uata	
her Settinus			
Append Counter to Sample	e Name:01		

Figure 3. Sample Information Editor.

• Check the Data to Project box and select the desired folder.

SPECTRUM ACQUISITION:

- Make sure the sample compartment is empty, and adjust the telescopic purge tubes as close to one another.
- The background will be collected in the single beam mode. The FT-IR will collect the sample spectrum as a ratio, subtracting out the background. In order to collect a background, set the acquisition mode to **Single Beam** and the desired number of **Scans** in the **Acquisition Parameter** window and click on the **Reference** button.
- In order to collect a spectrum, place the IR plate in the sample compartment and set the acquisition mode to **Absorbance** or **Transmittance** click on the **Measure** button.

ANALYSIS:

Baseline Correction:

- Select **Baseline Correction** from the **Mathematics** tab in the toolbar.
- Select the desired baseline correction by adjusting the red square boxes on either end of the correction line. Click on **Calculate** in the mathematics window.
- Close the baseline correction window.

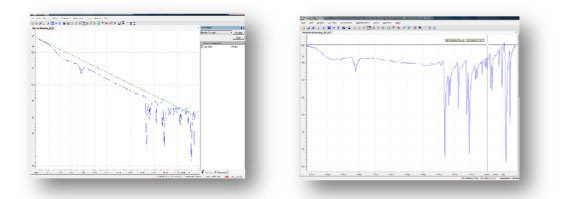


Figure 5. Baseline Correction.

Setting Trace Limits:

• Click on the axis to be modified and apply new limits. These will be reflected in the active window and print outs.

Labeling Peaks:

• Select **Peak Picking** from the **Mathematics** tab in the toolbar. A peak table will appear at underneath the spectrum.

- In order to add additional peaks, place the cursor just below the peak of interest and press the Ctrl key along the left mouse button.
- In order to delete unwanted peaks, just select the peak in the peak table and delete the corresponding line.
- Close the peak picking window.

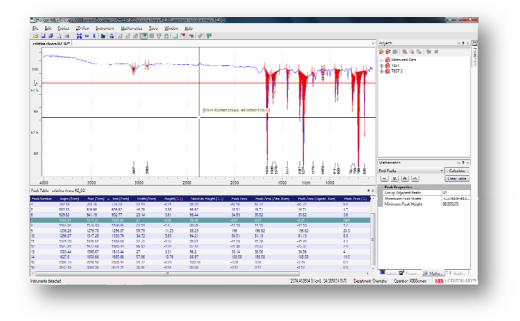


Figure 6. Peak Picking.

PRINTING:

- Select File then Print.
- Click **Set up** and select **Landscape** printing.
- Click on **Print** button.

FINISH:

- Remove the IR plate from the sample compartment and bring telescopic purge tubes close together.
- Clean the plate with CH₂Cl₂. **DO NOT USE WATER!!**
- Return plate to the dessicator.