

TA Instruments Standard Volume Isothermal Titration Calorimeter (ITC) Testing SOP

071724MB02

Sign in to the calendar if you have not already done so.

Prior to using the instrument, the instrument should be calibrated as detailed in the following section.

Note: The testing procedure can take up to three hours.

Testing Instrument Prior to Analysis and General Operation

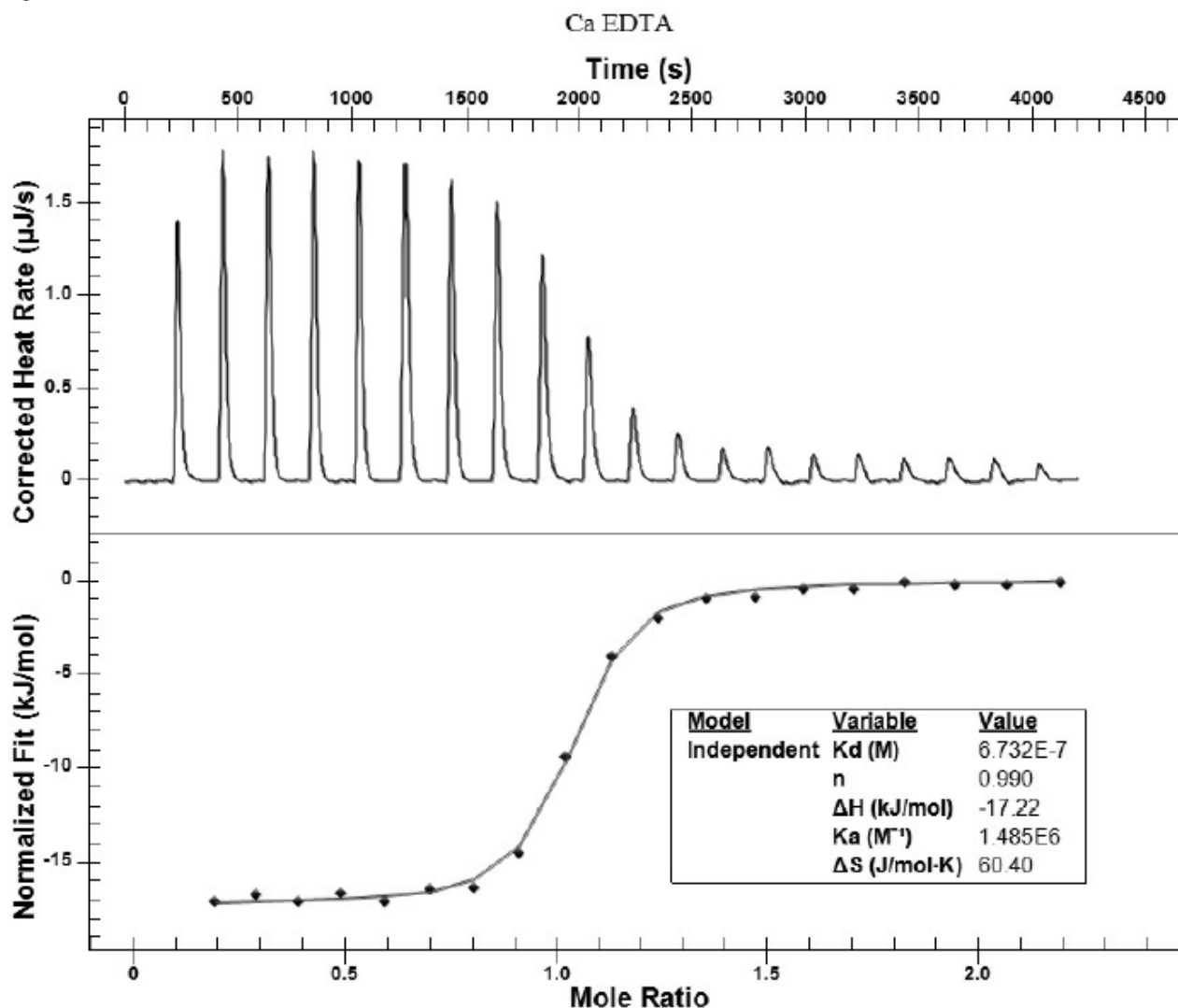
Software and CaCl_2 :EDTA Test Setup Procedure

1. Turn on the instrument using the power switch located on the back if it is off.
2. The software for analysis is **ITCRUN**. The buret will need to be homed if the software has been started for the first time. Follow the onscreen instructions for this. The software will open and you will need to wait until the status has changed to **Idle**.
3. The cells of the instrument need to be flushed prior to analysis.
4. Using the tweezers located by the instrument, remove the reference needle plug and set it securely to the side.
5. Using the flushing needle and syringe, flush the reference cell at least 3 - 4 times with ~1.5 mL of degassed ultra-pure water. ultra-pure water can be degassed by sonication or by drawing a vacuum for ~15 minutes.
6. To flush the cell, dispense the syringe contents in and out of the cell the desired number of times. Discard the rinsing water.
7. Fill the reference cell with ~1.5 mL of degassed ultra-pure water. There should be no water visible once the flushing syringe has been removed. Remove some of the volume if there is. Replace the plug into the reference cell.
8. Using the flushing syringe, remove the contents of the sample cell.
9. Flush the sample cell with ~1.5 mL of the EDTA buffer at least twice. Discard the EDTA rinse.
10. Load ~1.5 mL of EDTA into the sample cell and allow this to soak within the cell for at least 5 minutes.
11. Following the 5 minutes, discard the EDTA that was in the cell and replace with ~1.5 mL of fresh EDTA for the actual test.
12. Locate the 250 μL buret syringe. Flush the syringe 2 times with EDTA, followed by one rinse of the calcium chloride solution. Discard the calcium chloride solution rinse and replace with fresh calcium chloride solution. You want to fill the syringe with 250 μL of the calcium chloride solution for the test.
13. Place the syringe in the black (aqueous) buret handle, and then insert the buret handle into the instrument.
14. Set the **Stirring Rate** to **350** and press the play icon to start the stirring.
15. Click the "Syringe Size" drop down menu and select the **250 μL syringe**.
16. The "Experiment Setup" window will now be adjusted. Click **Setup** and the "Setup Injections" window will open. For the Ca-EDTA titration, change the **number of injections to 25**, the **injection volume to 10 μL** , and the **injection interval to 300**. Click **Ok**.
17. Ensure the "Temperature Setpoint" is at **25**.
18. Under "Experiment Details" change the value of the syringe concentration to match the value of the calcium chloride solution (taken from the label) and the cell concentration to match the label of the EDTA buffer.
19. Under the "Equilibration" section, select **Auto Equilibrate**. **Small** should be selected for the "Expected Heats." This selection determines the statistics of the equilibration.
20. The timeout function should be unchecked.
21. Change the "Initial Baseline" to **300**.
22. The analysis can now begin. Click the green play icon at the top of the ITCRUN screen. Save your data and the analysis will begin. You can monitor the baseline as the instrument equilibrates by clicking the "Monitor" tab.

Analysis of the test data

1. Open NanoAnalyze from the desktop.
2. Add the titration file by clicking **File** followed by **Add File**. Click on the **Analysis** tab on the vertical toolbar to display the titration data.
3. Within this tab, you can subtract the baseline so long as it is satisfactory (resolved). Points can be added or removed as necessary, but should be done minimally and with caution to improve the baseline before subtracting it.
4. Click the **Area** tab on the **Analysis** tab toolbar to confirm the syringe and cell concentrations are correct. Likewise, ensure the sample cell volume is 950 μL . This value is set by TA Instruments at the factory.
5. Correct for the background heat by subtracting the average of the last 3 data points in the area tab under "Area correction." These values are the Q values within the data columns. It is extremely important that the sign of the value entered in to be subtracted matches that of the data that was averaged.
6. Click the **Modeling** tab. To import a model, click the asterisk located below the "Models" box.
7. Choose the "Independent" model and then click the **Select** button. The independent model calculates the equilibrium constant (k), enthalpy (H) and stoichiometry (n).

8. Anomalous data can be masked by clicking on the data point. The first data point should be masked due to diffusion of titrant during equilibration.
9. Fit the data using the green play icon.
10. The image below shows what the Ca-EDTA titration data should resemble once fitted.



11. Compare the values calculated from the fit to those of the table below.

Independent Fit Values	LV and SV
K _d (M)	3.90-8.99 x 10 ⁻⁷
ΔH (kJ/mol)	-17.5 ± 2
n (mol)	1.0 ± 0.1

Table 6. Acceptable values for Ca²⁺-EDTA Titration in MES buffer pH 6.0.

12. If your test values are within the acceptable values, you may clean the cells to begin your analysis. If they are not, contact Matt Burleson (Ap 346, mburleson@wcu.edu)

Acquiring and analyzing data

1. Data collection and analysis for an actual sample will follow the steps outlined above.
2. It is vital that the cells be filled with the same buffer (or as similar as possible).
3. If organics are to be used, use the organic buret handle instead.

Basic Cleaning and Shutting down the instrument

1. Once your analysis is complete, the cells will be flushed with degassed ultra-pure water. This is a basic cleaning. More rigorous cleaning methods are detailed later.
2. Return the cells to 25 °C. **Rinsing room temperature fluids through a hot cell can damage sensors.**
3. Close the software.
4. Flush the reference cell at least 5 times with degassed ultra-pure water. Fill the cell with fresh degassed ultra-pure water after the last rinse, and replace the reference cell plug.
5. The sample cell will be flushed using the cleaning tool which shown below.



6. Ensure the instrument temperature has adjusted to near ambient before starting the cleaning process.
7. Remove the buret assembly and syringe from the top opening of the Nano ITC, and withdraw the cell contents using the filling syringe.
8. Carefully lower the shaft into the cell opening.
9. Connect the shorter rubber tubing to the side port of the cleaning tool. Place the free end of this tube in a beaker of clean, degassed ultra-pure water.
10. Connect the longer rubber tubing to the top port of the cleaning tool and the other end to the vacuum flask used for cleaning.
11. The connected tubing should resemble that shown below.



12. Apply a vacuum to draw the water through the system and flush the cell.
13. Refill the flushed cell with ~1.5 mL of fresh, degassed ultra-pure water.
14. Clean the titration syringe using a 0.5 % Contrad solution by back filling the syringe with a pipette and pushing out the liquid with the plunger. Repeat this 5x and follow with 10x of water. Use extreme caution when handling the titration syringe, especially if trying to clean the outside of the stir paddle. Clean the outside of the syringe by using a squirt bottle to rinse and then just gently brush a kimwipe to absorb any remaining liquid.
15. Contact Matt Burleson to perform a water-water titration to ensure the instrument is ready for the next user.
16. The instrument may be turned off by flipping the switch on the back.

Preparation of Solutions for Ca/EDTA Titration

Buffer matching is key for ITC experiments. Prepare the EDTA and CaCl₂ solutions using the same MES buffer.

Solutions Needed	Details
Sodium Hydroxide	5 - 10 N, prepared and stored in plastic containers
MES Buffer	10 mM MES, pH 6.0
EDTA	0.15 mM EDTA in 10 mM MES, pH 6.0
Calcium Chloride	1 mM CaCl ₂ in 10 mM MES, pH 6.0

10 mM MES, pH 6.0 (MW of MES: 195.2 g/mol, MW of MES Free Acid Monohydrate: 213.2 g/mol) = 2.132 g/L in ultra-pure water. Adjust pH using the sodium hydroxide solution above. Degas this solution before moving on.

0.15 mM EDTA in 10 mM MES (MW of 292.24 g/mol, MW of EDTA disodium salt dihydrate: 372.24) = 0.0056 g in 100-mL of 10 mM MES from above.

1 mM CaCl₂ in 10 mM MES (MW: 110.98 g/mol) = 0.0111 g in 100-mL of 10 mM MES from above. **Calcium chloride should be stored in a desiccator when not in use.**

Be sure to complete the ITC Solution Logbook EACH time a solution is made.

Rigorous Cleaning methods

10% v/v Contrad Solution

1. Discard any content in the sample cell and fill it with ~1.2 mL of 10% (v/v) Contrad solution. Set the temperature of the sample cell to 40 °C for ~1 hour. After 1 hour, return the temperature of the sample cell to 25 °C or to the temperature of the experiment.
2. Set up the rinsing apparatus as shown previously under Basic Cleaning and flush 100-mL of 10% (v/v) Contrad solution.
3. Empty the cell contents and rinse with 4-L of ultra-pure water.

Harsh Cleaning Method

1. Close the ITC software.
2. Load a mixture of 10% Contrad, 20% MeOH and 1 N NaOH into the sample cell.
3. Using the loading syringe, pipette up and down to agitate the solution.
4. Remove solution from cell.
5. Load 4N NaOH into the sample cell and cover the cell with a Kimwipe, but keep buret handle out.
6. Set the temperature to 65 °C for 30-60 min.
7. Cool the cell back to 25 °C. **Rinsing room temperature fluids through a hot cell can damage sensors.**
8. Rinse with at least 1-L of ultra-pure water.
9. Load a 50% formic acid solution into the sample cell.
10. Cover the cell with a kimwipe, but keep buret handle out.
11. Set temperature to 65 °C for 30-60 min.
12. Cool the cell back to 25 °C. **Rinsing room temperature fluids through a hot cell can damage sensors.**
13. Rinse with 4-L of ultra-pure water.