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General

How To Use This Manual

This manual describes the iTC_{200} operation, software and maintenance. Section 1 contains general descriptions of the system, its parts and tools, the software, how to perform a run, and the tutorials. These tutorials are recommended to every new user, to acquaint the user with the software and methods and to ensure that the instrument is working as expected. Section 2 describes the software in greater detail. Section 3 describes routine maintenance that users may need to perform, from basic cleaning, to periodic calibration checks, to removal and replacement of parts.

How to Get Help from MicroCal

The service department at MicroCal, LLC is available to help you during normal business hours, Monday through Friday from 8:00 AM to 5:00 PM (EST). The European office is available during normal business hours, Monday through Friday from 9:00 AM to 5:00 PM (GMT).

Service personnel may be contacted by e-mail, phone or fax, with preference being phone or e-mail. When e-mailing MicroCal for technical assistance, if possible, please attach a recent data file(s) (*.itc, raw ITC data file) that demonstrates the problem. Also, please include all details that may be relevant to the problem. For instances where the problem or question relates to post run data analysis, it is best to attach both the raw data file (*.itc) and the Origin project file (*.opj) generated during data analysis.

There are two general categories of problems associated with the iTC_{200} and its operation. The most extreme category is when a system is not working at all. Problems that prevent users from operating the iTC_{200} require immediate consultation with a MicroCal technician. Customers should not attempt to repair the iTC_{200} hardware or software unless instructed to do so by a MicroCal service representative.

The second, and less extreme general category of a problem is when an iTC_{200} instrument is functioning, but is not operating within its normal performance specifications. Large baseline drifting, non-repeatable control peaks (water/water) and/or an increase in short term noise level are examples of performance problems. These problems may be corrected by the operator in most cases. For these types of performance issues it is recommended that customers carry out the following minimum diagnostic steps prior to contacting MicroCal for service:

- 1. Thoroughly clean the cells. Do not assume they are clean; build-up or unexpected sample residue will cause problems. As a minimum, use the provided Cell Cleaning Apparatus to pass at least 50 ml of 10-15% detergent in water solution, followed by 50 ml of water, through the sample cell. Discretion may call for a more rigorous cleaning procedure.
- 2. Refill both the reference and sample cells with filtered degassed water. Thoroughly clean the syringe or use a different syringe. Load the syringe with water.
- 3. Go to the "Setup/Maintenance" window within the iTC₂₀₀ software and activate the extended data mode. While the software is in the extended data mode, the ITC data files will contain all available information produced by the iTC₂₀₀. This additional information is often helpful in the diagnosis of problems.
- 4. Carry out a minimum of 20, 1μL injections of water into water.

If, after completion of the 5 steps listed above, the ITC performance is not corrected, please contact the MicroCal service department for help. The water run, in extended data mode, should be provided to the MicroCal service technician for evaluation. Following the evaluation, a representative form the service department will contact you with comments and recommendations.

MicroCal, LLC Contact Information:

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Please forward corrections or suggestions concerning this manual to info@microcal.com

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Operational Safety

The points below are intended to enhance your safety awareness and to draw your attention to risks which only you, the operator, can prevent. While MicroCal works to ensure that the instrument is designed and tested to be as safe as possible, proper handling is also critical. The operators should be responsible people trained in basic laboratory protocol, and they should be familiar with the possible hazards before operating this instrument.

- Provide proper electrical power to the instrument. This should be 110 250 Volt, 50 60 Hertz alternating current, with a Ground Fault Circuit Interrupter (GFCI). Most power strips, including those provided by MicroCal, contain a GFCI. All power plugs and cords should be 3-prong, grounded cables or outlets.
- Replace fuses ONLY with 3.15 Amp 250 Volt Time Delay Fuses. Several spare fuses are provided with the original shipment.
- Repairs, alterations or modifications must only be carried out by specialist personnel, or with explicit directions from a MicroCal technician. Removal or modification of any cover or component could result in an unsafe or easily damaged instrument. The MicroCal service department will be happy to answer any questions and provide parts and service when necessary.
- A solution can become an electrical conductor when in contact with electricity. Use caution when using solutions near the instrument. If any liquid is spilled on or around the instrument, unplug the instrument immediately and wipe it up. Be careful not to overflow the small reservoirs around the cell access tubes. If there is any possibility that liquid may have leaked into the instrument case, contact MicroCal immediately. Do not plug the instrument into any electrical outlet until the problem is resolved.
- The operator should always follow proper laboratory procedures in handling and disposing of volatile or hazardous solutions.
- All solutions in the cells must be cooled down below 40 °C before removal. Any higher temperature may cause the syringe to break, and will increase the dangers of most hazardous solutions.
- Never allow liquid in the cells to freeze. The expansion of the liquid can distort the cells and rupture the most critical sensor, causing irreparable damage.
- In the iTC $_{200}$ instrument should always be moved in its normal operating orientation. Other orientations will subject delicate sensors inside the instrument to stress.
- The iTC₂₀₀ cells are constructed out of Tantalum or Hastalloy. **Strong acids and bases should be avoided**. Please refer to the accompanying material analysis booklet for further information
- This device is not designed to the Medical Devices Directive 93/42/EEC and should not be used for medical purposes and/or in the diagnosis of patients.

Instrument Specifications

Performance Specifications:

Operating Temperature Range: 2 - 80°C Titration Sensitivity Quotient: 8 nanocal * mL

Response Time: 10 seconds

Cell Design: 200µL, coin-shaped, non-reactive

Injection Syringe: 40μL, automated Smallest Injection Size: 0.1μL

Physical Specifications:

Cell Material: Tantalum or Hastelloy® Alloy C-276

Dimensions:

Cell: 6½ x 13 x 10½" Controller: 15½ x 16½ x7½" Monitor: 9 x 15 x 15½"

Weight (pounds)

Cell: 16 Controller: 25 Monitor: 12

Electrical Specifications:

All electrical specifications are for the cell only.

Electrical Ratings:

Voltage: 110 - 240 Volts Frequency: 50 - 60 Hz Power: 120 Watts

Fuses (2): 3.15A, 250V, Time Delay Output: Secondary/Data Connection Only

Protective Earth Terminals: Internal/external marked

Mode of Operation: Continuous

Classification: Class I

Atmospheric Specifications:

Operating:

Temperature: 10 - 40 °C

Humidity: 0 - 70% RH non-condensing Atmospheric Pressure: 700 – 1060 HPa

Storage (no liquid in cells):

Temperature: -40 - 70°C (no liquid in cells)

Humidity: 10 - 90% Atmospheric Pressure: 500 – 1060 HPa

Section 1: Introduction to the iTC_{200}

This section provides information about isothermal titration calorimetry and the basic parts and operation of the Isothermal Titration Calorimeter (ITC) instrument and software. At the end, it includes several suggested tutorial experiments.

1.1 The Basics of ITC

The iTC_{200} (Isothermal Titration Calorimeter, $200\mu L$ cell) unit directly measures heat evolved or absorbed in liquid samples as a result of mixing precise amounts of reactants. A spinning syringe is utilized for injecting and mixing of reactants. Spin rates are user selectable; the usual range is 500 to 1000 RPM. The normal temperature operating range is 1°C to 80°C. Wetted cell surfaces are Tantalum or Hastalloy, which are resistant to most solutions, however, strong acids and strong bases should be avoided.

Sample and reference cells are accessible for filling and cleaning through the top of the unit. The sample cell is on the left as one faces the front of the unit. A pair of identical coin shaped cells is enclosed within two shields; the inner shield is referred to as the jacket. Access stems travel from the top exterior of the instrument to the cells. Both the coin shaped cells and the access stems are completely filled with liquid during operation. This requires approximately $250~\mu L$ per cell even though the working volume of the cell is only $200~\mu L$.

Temperature differences between the reference cell and the sample cell are measured, calibrated to power units and displayed to the user as well as saved to disk. The data channel is referred to as the DP signal, or the differential power between the reference cell and the sample cell. This signal is sometimes thought of as the "feedback" power used to maintain temperature equilibrium. Calibration of this signal is obtained electrically by administering a known quantity of power through a resistive heater element located on the cell.

In a typical experiment, the syringe containing a ligand is titrated (injected) into the cell containing a solution of macromolecule. An injection which results in the evolution of heat (exothermic) within the sample cell causes a negative change in the DP power, since the heat evolved chemically provides heat that the DP feedback is no longer required to provide. The opposite is true for endothermic reactions. Since the DP has units of power, the time integral of the peak yields a measurement of thermal energy, dH. This heat is released or absorbed in direct proportion to the amount of binding that occurs. When the macromolecule in the cell becomes saturated with added ligand, the heat signal diminishes until only the background heat of dilution is observed.

With the iTC_{200} system the entire experiment takes place under computer control. The user inputs the experimental parameters (temperature, number of injections, injection volumes) and the computer carries out the experiment. Origin software is then used to analyze the ITC data using fitting models to calculate reaction stoichiometry (n), binding constant (Ka), enthalpy (ΔH) and entropy (ΔS) .

1.2 Parts and Accessories

The main parts of the system are the cell unit and controller. The tower, on top of the iTC_{200} cell unit, guides the motion of the pipette, which holds the titration syringe. The pipette has three positions; cleaning, loading, and run positions. A 1 mL plastic syringe is used for loading the titration syringe, and a glass 0.5 mL syringe (Hamilton) is used to fill and rinse the sample and reference cells. A cell cleaning apparatus, consisting of a small plastic barrel and soft tubing, is used to run liquid through the sample cell, in combination with a small vacuum flask. The vacuum is provided by a ThermoVac unit, which is also used for sample preparation. Several tools are used for changing a syringe or a pipette tip. Also, a thin piece of wire is provided for vigorous cleaning of the syringe tip.

Complete System: ThermoVac unit; Cell; Controller; Pipette and Tower



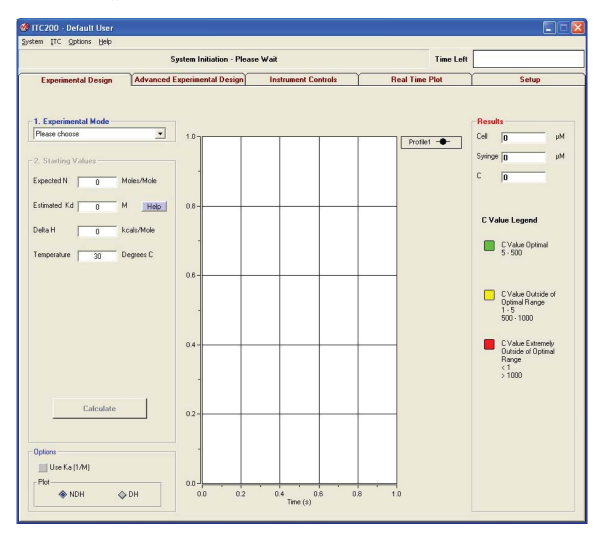






1.3 Software

In order for the system to initialize properly, all components must be powered up in the correct order. First, boot up the computer and log in to Windows. Once Windows has started, power the iTC_{200} by operating the switch at the rear of the unit. After several seconds, open the iTC_{200} software. If the option is selected, a real-time copy of Origin will open automatically, as well as the iTC_{200} control software.



The line just below the menus reads "System Initiation – Please Wait", which is the current status of the instrument. After a few seconds, the system will begin heating or cooling to the preset temperature. To the right of that, the "Time Left" box, during a run, will show the time left until the end of the run.

When the software is first started, the "Experimental Design" tab is selected; this contains the simple run controls. "Experimental Mode" can be "Highest Quality" or "Minimum Protein". The expected n, K_d , and Delta H and the desired run temperature will allow the software to

calculate the recommended concentrations for the cell and syringe, and set the run parameters. The "Advanced Experimental Design" tab contains more direct controls for the more advanced user. This tab should be very familiar to users of the VP line of instruments. The "Instrument Controls" tab allows the user to name the output files, choose post-run analysis options, and start and stop the run. The "Real Time Plot" tab shows the data currently being generated. The "Setup" tab contains various options and preferences.

Optional Feature:

The copy of Origin that is opened by iTC_{200} software, if desired, is for real-time data display. For analysis, the user should open a separate copy. At the top of the graph screen, the current status of the instrument is shown. Below that, the file name that the software is currently saving to is displayed. The graph displays DP in μ Cal/sec verses time in seconds. The "Main Display" box in the upper left corner shows the temperature of the cells, current DP, and DT. The lower left corner contains the graph view options: "Rescale To Show All" resizes the window to fit whatever data is currently displayed in it, "Auto-View 1" and "Auto-View 2" set the window to a certain scale, with the latest data point centered vertically, and "Saved View 1" and "Saved View 2" set the Y-axis to a specific range. All four of these views can be edited through the "Edit ranges" button.

1.4 Basics of Performing a Run

In order to perform a basic ITC titration experiment, the user must load the sample cell and the syringe, enter the desired parameters into the control software, and click "start". The reference cell should be filled with water or buffer, and may be left for several days.

Loading the Syringe:

To load the syringe, place a microcentrifuge tube in the tube holder. Be sure to push the tube to the bottom of the holder with the lid fitting into the slot provided (see image below). Be careful not to leave any part of the tube in the path of the syringe needle to prevent damage.



Disconnect the filling tube from the ThermoVac unit and connect the threaded end to the cell port if necessary (it should still be in place after cleaning and drying of the syringe). Insert the tip of the 1 mL plastic syringe into the end of the filling tubing. In the software, under the "Instrument Controls" tab, click the "Open Port" button to make sure that the fill port is open. Slide the pipette up, across, and down into the middle slot, so that the tip of the syringe rests in the microcentrifuge tube. Gently pull up on the plunger, until the sample has filled the titration syringe and is just overflowing into the tubing.







Click the "Close Port" button to close the fill port. Pull the plunger up approximately another 0.1mL to remove the excess liquid in the tubing, and disconnect the tubing from the fill port. Click the "Purge/Refill" button 2-3 times to move the pipette tip down and back up again, dislodging any air bubbles on the glass sides of the syringe.

Loading the Cell:

To load the sample cell, gently insert the glass Hamilton syringe into the (left) sample cell until it touches the bottom. Pull up on the plunger until bubbles are being pulled from the

cell, and there is no more liquid. Remove and empty the syringe. Clean the syringe if necessary. Pull approximately 300 μ L of sample into the syringe, and tap the syringe glass so that all air is at the top volume of the syringe. Do not allow air to be put into the cells. After removing the bubbles, insert the syringe into the cell and gently touch the bottom of the cell with the tip of the syringe needle. Raise the needle tip about 1 mm off the bottom of the cell, and hold it there until finished filling. Do not raise the syringe during the filling process. Slowly inject solution into the cell until it spills out the top of the cell stem. Finish the filling with several small abrupt spurts of solution to dislodge any bubbles in the cells. Finally, lift the tip of the syringe to the cell port (just below the visible portion of the cell port) and find the ledge that is formed where the cell stem meets the cell port. Place the syringe on the ledge at the top of the metal cell stem and remove the excess solution. If the reference cell needs refilling, follow the same procedure as for the sample cell.

If there is a saved run file for the current experiment, load the run parameters by clicking the "Load Run File" button at the top left of the "Advanced Experimental Design" tab in the iTC₂₀₀ software and select the run file. There are standard run files for most of the tutorial experiments. If no run file exists, enter parameters in the appropriate boxes in the "Experimental Design" tab (see sections 1.3 and 2.2 for more information). Once the parameters have been entered, click the "Start" button at the upper left corner of the "Instrument Controls" tab of the control software. Be sure to insert the syringe in to the sample cell before starting the experiment. The instrument will seek experimental temperature, equilibrate to that temperature, start the syringe stirring, wait until the DP signal is steady, and then start performing injections. The raw data will appear in the "Real Time Plot" tab.

Once a run has finished, the syringe and sample cell should be cleaned as soon as possible. See section 3.1.

1.5 Tutorial Experiments

The next two tutorial experiments are recommended for every new user before beginning analysis of their own samples. They are designed to provide standard results, which check that the instrument is working well and the user is comfortable with the procedures, before risking precious sample.

Water/Water Tutorial:

The first tutorial involves titrating water into water.

- 1. Fill both cells with degassed (5 min) distilled water, fill the titration syringe, and insert the syringe into the sample cell.
- 2. Click on the "Load Run File" button in the "Advanced Experimental Design" tab, choose "WATER.INJ", and click "Open". Name the run and click "Start". The instrument will seek the run temperature, equilibrate the system, and begin making injections. Each peak should be small, with all the peaks approximately the same shape.
- 3. Once the run has finished and the instrument is thermostatting again, open the ITC Calibrations project to analyze the data. It is located in the Origin directory, at C:\Origin70\ITCCalibrations.OPJ.
- 4. Click on the "Water Water Report" button. A window will pop up asking which files to analyze; select the file you just generated, click "Add Files", and click "OK".
- 5. Origin will analyze the data for several seconds. Once it has finished, it will ask you to save the results. Give the file a unique name. There should be a graph, with the water DP data, and a textbox. Zoom in using the magnifying glass over a plus sign icon on the top toolbar, so that you can read the textbox. To zoom in and read the text, select the magnifying glass over a plus sign icon on the top toolbar and click on the textbox.

```
Water-Water Report
VPITC11.06.834
12/5/2006 08:42:23
Raw DP Data
10uL injections
Water-Water Specifications:
Mean Energy/Injection less than 1.5 uCal
SD less than 0.15 uCal
Baseline Offsets Have Been Manipulated!
Wws1a, Mean Energy/Injection=-0.34863uCal, SD=0.08232 uCal
Results Within Specifications
```

Line 2 of the script is the instrument serial number, line 3 is the date and time that the report was generated, line 5 is the injection size, and lines 7 and 8 are instrument specifications. Origin manipulates the offsets so that multiple water runs can be displayed without overlapping. Line 10 gives the data file name and the mean and standard deviation of the energy per injection. The

last line will say "Results Within Specifications" or "Results Outside of Specifications". If the results are not within specifications, you may want to run the tutorial again.

CaCl₂/EDTA Tutorial:

This tutorial uses the standard sample kits provided with the instrument when it is shipped. Kits can also be obtained from MicroCal at any time.

- 1. Load the titration syringe with CaCl₂. The sample cell must be rinsed with EDTA in order to obtain an accurate fitted value for stoichiometry. Fill the sample cell with EDTA and allow the solution to remain in the cell. After 5 minutes, remove the EDTA solution from the cell. Load a fresh volume of EDTA in to the sample cell.
- 2. Set the run parameters. A set of parameters is provided with the sample.
- 3. Name the file and click "Start" to begin the experiment.
- 4. Once all injections have been completed, clean and dry the syringe. Use the cleaning apparatus to clean the sample cell. See section 2.9, "Origin Data Analysis Tutorial" for analysis of the results.

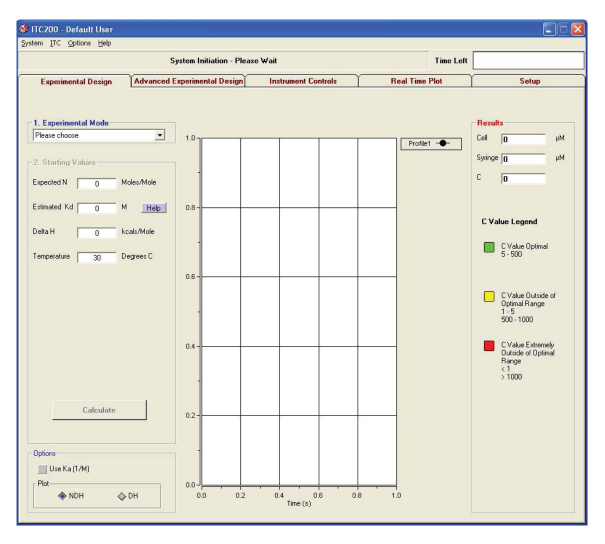
NOTE – Detailed instructions concerning sample loading and data collection using the standard CaCl₂ / EDTA kit are provided with each kit.

Section 2: iTC₂₀₀ Software and Origin

This section details the features of the iTC_{200} software. This software provides the user an interface to control the iTC_{200} instrument and perform an experiment. An optional copy of Origin can be used for plotting the data received from the iTC_{200} instrument. A separate copy of Origin (Origin for ITC) is used for analyzing data.

2.1 iTC₂₀₀ Software Overview

The iTC_{200} software contains drop down menus, a status bar, and several tabs. The drop down menus control global system functions. The status bar displays the current status of the instrument. Each tab contains controls for different areas of instrument control.



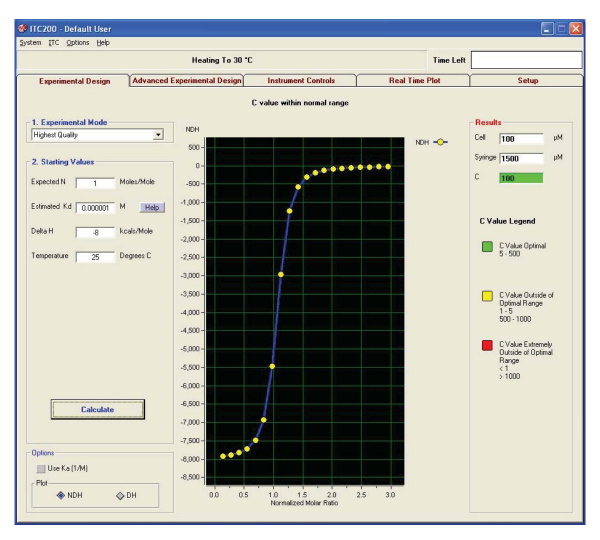
The status bar just below the menus reads "System Initiation – Please Wait", which is the current status of the instrument when it is first booting. After a few seconds, the system will begin heating or cooling to the preset temperature. Once the instrument reaches the set temperature, it will thermostat at that temperature. To the right of the status bar, the "Time Left" box. During an experiment this box will show the time left until the end of the run.

When the software is first started, by default, the "Experimental Design" tab is selected; this contains the simple run controls. "Experimental Mode" can be "Highest Quality" or "Minimum Protein". "Highest Quality" uses 20 injections and a c-value of 100. These parameters should produce data that is clear and easier to fit. "Minimum Protein" uses fewer

injections, 10, and a c-value of 5. The result of these parameters will be the use of the least amount of sample necessary for a successful titration. The expected n, K_d , and ΔH and the desired run temperature will allow the software to calculate the recommended concentrations for the cell and syringe, and set the run parameters based on mode chosen. If the user is unsure of the K_d for their system, clicking the "Help" button causes the software to prompt for the type of compound in cell and syringe. It will then make a guess as to the K_d . The user will still be required to choose values for ΔH and n.

The "Advanced Experimental Design" tab contains more direct controls for the more advanced user. This tab should be familiar to users of the VP line of instruments. The "Instrument Controls" tab allows the user to name the output files, choose post-run analysis options, and start and stop the run. The "Real Time Plot" tab shows the data currently being generated. The "Setup" tab contains various options and preferences.

2.2 Experimental Design



When the software is first started, by default, the "Experimental Design" tab is selected; this contains the simple run controls. "Experimental Mode" can be "Highest Quality" or "Minimum Protein". "Highest Quality" uses 20 injections and a c-value of 100. These parameters should produce data that is clear and easier to fit. "Minimum Protein" uses fewer injections, 10, and a c-value of 5. The result of these parameters will be the use of the least amount of sample necessary for a successful titration. The expected n, K_d , and ΔH and the desired run temperature will allow the software to calculate the recommended concentrations for the cell and syringe, and set the run parameters based on mode chosen. If the user is unsure of the K_d for their system, clicking the "Help" button causes the software to prompt for the type of compound in cell and syringe. It will then make a guess as to the K_d . The user will still be required to choose values for ΔH and n.

Click the "Calculate" button to calculate the results. The simulation window will update with a rough graph, and the "Results" column at the right of the screen will have values for the cell and syringe concentrations. The calculated C value is listed below; its background is color-coded. This value is equal to 5 in "Minimum Protein" mode, 100 in "Highest Quality" mode, and will vary if the user chooses to change cell concentration manually. The c-value predicts the shape or sigmoidicity of the curve. Optimal values for C are between 5 and 500 (green); values between 1 and 5, and 500 and 1000 should work but may not give the best result (yellow). C values less than 1 or greater than 1000 will probably not yield usable data (red). The user may adjust the two experimental concentrations by using the change buttons beside each concentration box. Once the cell concentration has been changed, clicking "Calculate" again will update the suggested syringe concentration relative to the new cell concentration.

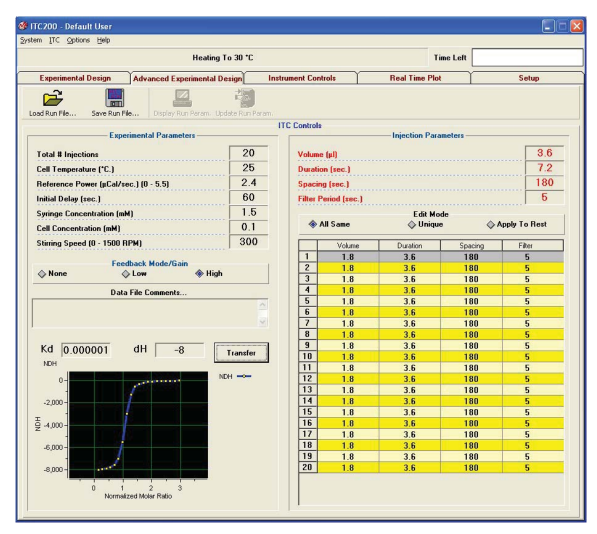
Any warnings will appear in the status bar near the top of the screen. These include "High heat warning", meaning that the estimated heat that will be generated is more than the instrument can detect.

It is highly recommended that the users look carefully at the projected curve and make sure that the shape and rough values are reasonable before proceeding.

A pair of options at the bottom of this tab allows users to work in K_d or K_a , and to choose whether to view the simulation plot using raw heat per injection (DH) or the heat normalized to the molar ratio (NDH).

2.3 Advanced Experimental Design

The advanced experimental design tab may be used in addition to the "Experimental Design" tab for operating the iTC_{200} instrument. Below is a description and suggested values for each of the run parameters displayed in this tab.



The top bar in this tab contains several buttons for opening, saving, and viewing run parameter files. The "Load Run File" button will display the file open dialog box, in which the user can select a file to load into the "Advanced Experimental Design" tab. Run parameters may be loaded from one of two types of files: a previous experiment's data file (*.itc) (the run parameters are extracted from the data file header), or a setup file (*.inj) which was previously saved using the "Save Run File" button.

"Save Run File" displays the file save dialog box which allows saving of the currently displayed run parameters. Once the parameters have been saved, they may be reloaded and reused in the future by selecting Load Run File as described above.

"Display Run Parameters" displays the current run parameters for the run in progress. This button is available only when the iTC $_{200}$ is in a non-idle state.

"Update Run Parameters" updates the current run parameters for the run in progress. When the ITC is in a non-idle state, this button must be clicked for the run parameter changes to take effect. This button is available only when the iTC $_{200}$ is in a non-idle state. Note: Only run parameters that have not yet been applied on the run in progress may be updated; all others will be ignored (i.e. after the stirring has commenced for an experiment, changes to the experimental stir speed will be ignored).

The "Total # Injections" (500 is the maximum) sets the number of injections for the titration (ITC) experiment. The multiple-injection method requires a minimum of 10-15 injections. There are no negative consequences (except the time spent) to titrating past the point where all sites are saturated, and these last few injections can be used for control heat information. The single injection method uses one single longer injection.

The "Cell Temperature" box is used to set the desired run temperature for the experiment. Most runs are performed between 25°C (room temperature) and 37°C (human body temperature); although the instrument's operating range is 2°C to 80°C.

Throughout an ITC experiment, a small constant amount of power (equal to the "Reference Power" entered) is continuously supplied to the offset heater of the reference cell. This causes the DP feedback system to become positive to supply compensating power to the sample cell that will equilibrate the temperatures. During an experiment the DP baseline will equilibrate near the value entered in the "Reference Power" box. The reference power setting is often referred to as the baseline setting. The best choice for the reference power setting will be determined by the anticipated size and direction of the titration peaks. Large exotherms will require a large reference power setting (ca. 5 μ Cal/sec.) and large endotherms will require a very small reference power setting (ca. 0.5 μ Cal/sec.). When working with an unknown sample, a reference power of 2-3 μ Cal/sec is best.

The "Initial Delay" refers to the time, in seconds, after the instrument has started a run and before the first injection. The standard parameter is 60 seconds; this is necessary to establish a baseline prior to the first injection.

The "Syringe Concentration" and "Cell Concentration" boxes are a place to enter the experimental concentrations. The "Experimental Design" tab will make suggestions of the proper concentrations if desired (see section 2.2), or refer to "Calculating Concentrations" later in this section for more information. If the "Experimental Design" tab was used, these values will already be filled in. These concentration values are stored in the data file header and used for ITC data analysis. These values are used only during data analysis and an entry is not required to perform an experiment.

Select the desired stirring rate from the drop down list (1000 RPM is recommended for most ITC experiments). If the solution in the sample cell contains suspended particles (e.g. agarose beads), then faster stirring may be necessary.

In the "Feedback Mode/Gain" section, the three available modes are "none", "low" or "high" (High is recommended for most ITC experiments). The feedback modes can be described as active ("low" or "high") or passive ("none"). High gain will provide the fastest response time, while passive mode ("none") will provide the highest sensitivity. Almost all ITC reactions will require using the high gain feedback mode. Monitoring of long, slow thermal process in the iTC $_{200}$ (i.e. kinetics, metabolic rates) might benefit from using the passive or low gain feedback modes.

The "Data File Comments" box allows the user to enter comments about the experiment to be saved in the header of the data file for future reference.

The simulation plot in this window is generated from the "Experimental Design" tab. Clicking "Transfer" pulls the information from that tab to the current one. In addition, the user may enter the K_d and dH here and change the plot from this tab.

The "Injection Parameters" group allows the user to define the volume, duration, spacing between injections, and the filter period for each injection. At the bottom of this group is a summary table that lists these parameters, ordered by injection number. The user may select an injection to edit by clicking on a row to highlight it; the parameters for this injection can be entered into the text boxes (see below). Then, depending on the "Edit Mode" selected, the user may edit the text boxes to effect all injections ("All Same"), only the currently highlighted injection ("Unique") or the currently highlighted injection and all the following injections ("Apply To Rest").

In the "Volume" box, the user may enter the volume of titrant (in μ L), to be injected from the pipette into the sample cell for the selected injection(s).

In the "Duration (sec.)" box, the user may enter the time, in seconds, that the instrument will take to inject the titrant into the sample cell. The iTC_{200} software will generally default this value to be twice the number entered in the "Volume" text box. For single injection method experiments, the standard duration is ten times the volume.

In the "Spacing (sec.)" box, the user may enter the time, in seconds, between the beginning of the selected injection and the beginning of the next injection (or end of the run). This spacing must allow enough time between the injections to allow the DT signal to return to the baseline after an injection peak deflection. Typical values for this parameter range from 90-180 seconds, depending on the size of the peak and the kinetics of the reaction. For Single Injection Method, the spacing should be 90 - 180 seconds greater than the duration (i.e. volume = $20 \,\mu$ l, duration = $200 \, \text{seconds}$, spacing = $300 \, \text{seconds}$).

The "Filter Period (sec.)" is the time period (in seconds) in which the data channel conversions are averaged to produce a single data point for plotting and saving. For fast reactions, a filter period of 5 seconds is sufficient to obtain enough data points for representation of the peak for accurate integration of the area. For monitoring of very long, slow thermal processes, the filter period may be increased accordingly to avoid accumulation of excess data points. Although there may be a practical limit for a data set size, only available disk space would limit the number of data points that the iTC₂₀₀ software is able to collect and save.

Calculating Concentrations:

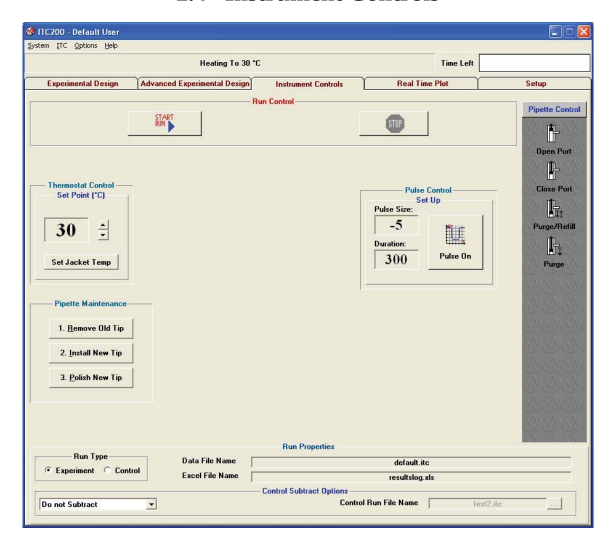
The easiest way to determine a good starting concentration for a system is to use the "Experimental Design" tab of the iTC_{200} software (see section 2.2). This section provides an alternative method for the slightly more advanced user.

To approximate the concentration needed in the cell, estimate the K_d for the system and multiply by 10. With very tight binders, the concentration should be on the low side of this value; for weak binders, the concentration should be on the high side of this value. Multiply the cell concentration by n and by 10 - 15 for the syringe concentration. Again, for very tight binders, the concentration should be on the low side and for weak binders this value should be on the high side. The minimum and maximum concentrations that are commonly used in the iTC₂₀₀ are $3\mu M$ to $500\mu M$.

These are "rules of thumb" and not absolute. However, if K_d and n are unknown, a best guess should provide enough data to fine-tune the parameters. An analysis of the data from the run will show the average peak size, approximate n, and a better estimate of K_d . Either the concentrations or the injection sizes should be adjusted so that the large initial peaks are ~ 1 - 4 μ Cal/second in height. The peak height is roughly linearly proportional to each concentration and the injection size. The refined estimate of K_d allows for an adjustment of the cell and syringe concentrations.

As an example, assume we have an enzyme in the cell with a K_d of 0.000001. The concentration of enzyme should therefore be $10*K_d=0.00001$ M or $10~\mu M$. Since this is a reasonably average K_d , no further adjustment is required. Assuming a stoichiometry of 1, the concentration of inhibitor in the 40 μ l syringe should be 10 - 15-fold higher than the concentration of the enzyme in the cell, or $10~\mu M*10=100~\mu M$ to $10~\mu M*15=150~\mu M$.

2.4 Instrument Controls



The "Instrument Controls" tab contains the controls for direct operation of the instrument. At the top of this tab, the user can "Start Run", using whatever parameters are currently present in the "Experimental Design" or "Advanced Experimental Design" tabs. Before clicking this button, it is wise to check that all parameters are correct and that a valid, unique data file name has been entered. The software will double-check with the user before allowing any files to be overwritten. The "Stop" button, which is available only during a run, will abort the run immediately.

The "Thermostat Control" section allows for setting of the thermostat temperature, which will be maintained during the iTC_{200} thermostating (idle) state. Pre-thermostatting the iTC_{200} and samples at the run temperature will result in shorter equilibration times. Also, high temperature thermostatting during cell cleaning can improve the effects of the cleaning. Use the arrow buttons to raise or lower the temperature or click in the textbox and type in a new number. Click "Set Jacket Temp" to set the temperature.

The "Pipette Maintenance" section provides the software controls used for changing the pipette tip. Please see Section 3.?? for details on performing this maintenance.

The "Pulse Control" section allows for manually administering a DP calibration pulse. While this is not the most thorough method of checking the y-axis calibration, it is the quickest method. Pulses may be applied any time the DP signal is equilibrated and the resulting deflection used as a crude calibration assessment. If Origin for real-time plotting is enabled, Origin will calculate the error. If this is greater that 1%, please see Section 3.7 for a more thorough check of the DP calibration.

The "Pipette Control" panel at the right of this tab provides the controls for loading and cleaning the syringe. The "Open Port" button moves the plunger tip to above the fill port in the side of the syringe. "Close Port" moves the plunger tip down so that it blocks the fill port. "Purge/Refill" pushes the pipette tip all the way down and back up again, dislodging bubbles on the sides of the syringe. "Purge" moves the tip all the way down and leaves it there.

The "Run Properties" bar at the bottom of this tab lets the user choose output options. Postrun analysis will be carried out only if "Auto postrun data analysis" is selected under the "Options" menu. "Run Type" tells the computer whether this run is an experimental run or a control run. The two textboxes to the right of this allow the user to enter a data file name and an excel spreadsheet file name. In the bottom left textbox are the options for control heat subtraction. "Do not Subtract" tells the computer to do nothing. "Subtract Saturation Points" will cause the automated data analysis to shift the delta H data so that the final, blank heats are zero. "Subtract Straight Line" will fit a straight line to the data and subtract it. "Subtract Control File" will subtract the file named in the "Control Run File Name" box. Clicking the "..." button will open a pop-up in which the user can browse to the appropriate file. The file name can also be typed in directly.

States of Operation:

The iTC_{200} is a state driven instrument. The current state of the ITC (cell status) is displayed in both the control software and Origin, if it is enabled. There are five unique states that comprise an ITC run, thermostatting (idle), seeking temperature, prerun thermostat, final baseline equilibration and running. A description of each of these five states can be found below.

The thermostatting (idle) state is what is in effect when no run is being performed. The iTC_{200} will achieve and maintain the currently defined thermostat temperature. It will remain in its idle state until the operator starts a run by clicking the "Start Run" button.

When a run is started, the first operation that the iTC_{200} will perform is to heat or cool the cells and jacket to the entered starting temperature for the experimental run. This is accomplished when the cell temperature is within 0.1 degrees (of the entered temperature), and the DeltaT (temperature difference between cells and adiabatic jacket) is less than $0.001^{\circ}C$. To save equilibration time it is usually best to set the iTC_{200} thermostat temperature to the same as the starting temperature of the ensuing run, or slightly colder. When filling the cells with solution prior to a run, the sample being loaded should always be colder than the desired experimental temperature, in order to minimize the amount of time required to equilibrate the ITC. Loading solution that is hotter than the desired run temperature will cause for lengthy equilibrations and/or a run temperature that is slightly higher than requested.

Once the cells and jacket have reached the starting temperature, the ITC will enter the second prerun state, the prerun thermostat state. The purpose of this state is to maintain the starting temperature, for the requested period of time. This allows time for thermal gradients within the ITC cell thermal core to minimize before the experiment starts and will reduce their effect during the experiment. The default duration of the prerun thermostat state is 5 minutes.

After thermostatting, the ITC instrument will enter the final baseline equilibration or the stirring state. The syringe will start to rotate. The start of the stirring will cause the DP signal will deflect downward and then will increase, but will stabilize at a level slightly lower than the non-stirring baseline. The instrument will stir continuously for the remaining portion of the run.

Once the DP signal has stabilized after the start of the stirring, the instrument will enter the run state. The run commences with an initial delay to provide baseline data prior to the first injection. Then the injections will be executed according to the injection schedule. The instrument responds to an exothermic reaction by decreasing the feedback heat provided to the sample cell. That is, heat added to the sample cell by an exothermic reaction will result in a downward deflection in the baseline DP signal. This downward deflection is proportional to the energy not needed to maintain the temperature equilibrium between the sample and reference cells. An endothermic reaction will require more heat in the feedback loop to maintain the same temperature resulting in a positive deflection. The integrated area of this deflection is referenced to the baseline before the injection and after the reaction has completed, and a curve is fit to the points generated. In the single injection method, the shape of the peak from one injection is used to determine the characteristic curve.

2.5 Real Time Plot

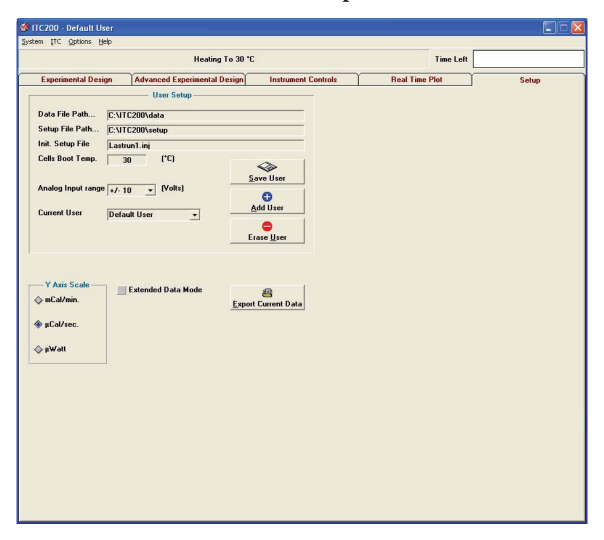


The "Real Time Plot" tab displays the current data. If no run has been started, the graph will be blank. The "Temp (°C)" box displays the current cell temperature. The "DP (μ Cal/s)" box displays the current DP. Technically, it is the reading from the sensor between the sample and reference cells. This data channel shows the changes in heat in the sample cell as events occur and is plotted in this graph.

The three buttons at the lower left control the graph resolution; all three resize the y axis relative to the last data point plotted. "Zoom ± 0.05 " alters the vertical range to the last data point plus or minus $0.05\mu Cal/s$. "Zoom ± 0.5 " alters the range to the last data point plus or minus $0.5\mu Cal/s$. "Show All" rescales the y axis so that all the data points are visible.

The "Plot Idle Data" box, when checked, will display the current DP readings on the graph even though no run is being performed. It is frequently helpful for troubleshooting.

2.6 Setup



The "Setup" tab allows customization for each user. This is often most helpful when different users have different default data file paths, so that each user's files are stored in a different place in the computer.

The "Data File Path" box indicates the path (excluding file name) where iTC_{200} data files will be saved. The path should be located on a local hard drive and not on any floppy or CD-ROM drive. This setting may only be changed while the iTC_{200} is in the idle state, thermostatting. To change the current data file path, double click on the existing path text and a dialog box will open allowing the user to change the path. The desired data directory must already exist in order to select it from the dialog box. In order to be read into Origin for data analysis the path cannot contain any spaces.

The "Setup File Path" box indicates the path (excluding file name) where iTC_{200} setup files will be saved and loaded. The specified path should be located on a local hard drive and not on any floppy or CD-ROM drives. To change the current setup file path, double click on the

existing path text and a dialog box will open allowing you to change the path. The desired setup directory must already exist in order to select it from the dialog box.

The "Init. Setup File" box specifies the initial setup file which will be loaded into the experimental control window, when it is first opened. Specifying the initial setup file as LastRun1.inj will result in the last run parameters that were executed to be loaded at startup.

The "Cells Boot Temperature" box sets the temperature that the instrument will heat or cool to and then thermostat at.

The "Analog Input Range" selector determines the maximum range of measurable DC voltages for all iTC $_{200}$ data channels. The default range of +/- 10 volts is adequate for almost all ITC applications and provides for the greatest sensitivity. Please consult a MicroCal engineer prior to making any changes.

The "Save/Add/Erase User" buttons provide a convenient way of maintaining unique startup parameters and placing data files in different folders for multiple users. To enable additional users, select the Add User button and enter a user name in the text box and click OK. "Save User" will save the current settings for the current user. "Erase User" will remove a user.

The Y-axis group allows for changing of the current y-axis (DP) units. Available y-axis units are mCal/min., μ Cal/sec. and μ Watts. By default, y-axis units will be displayed in μ Cal/sec. Changing y-axis units will only change the power units for 'local' data displayed in real-time. All data files will be saved with the default units of μ Cal/sec

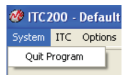
The "Extended Data Mode" button will tell the software to save all data the iTC_{200} instrument produces to the data file. This provides MicroCal engineers with extra data for troubleshooting.

The "Export Current Data" button will bring up a "save as" window to prompt the user for a filename, and then save whatever is currently displayed in the real-time window to that file. By default, it will be a *.dat file.

2.7 Menu Options

The main drop down menus at the top of the iTC_{200} software will provide access to some of the less frequently used functions of the application.

The "System \Rightarrow Quit Program" menu option prompts the user to confirm that they want to terminate the application. Responding with yes will terminate the program and any run in progress. All ITC run data will be saved to disk. Approximately a minute or so after program termination, the power to the iTC₂₀₀ will also be shut down. Power to the ITC will not be restored until the application is run again.



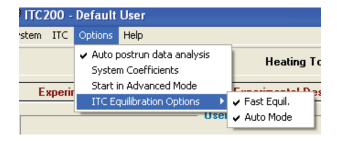
The "ITC => Print/Save As Text Run Parameters" menu option allows users to print or save to a file the run parameters that are currently loaded in the iTC₂₀₀ software. While run parameters are always included as part of the ITC data file header, the "Print/Save As Text Run Parameters" menu option provides the user with a nicely formatted listing of run parameters, ideal for data presentation or general record keeping.



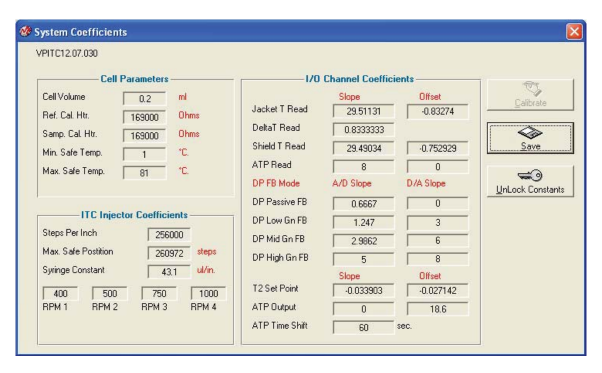
The "Pipette Tools" menu option opens a popup window for adjusting syringe calibrations.

The "Start ITC Calibration Run" menu item is used to check the accuracy of the instrument's calibrations. See sections 3.6 and 3.7 for more information.

The "Options" menu allows for several user settings. "Auto postrun data analysis" tells the computer to automatically analyse the data after each run. The exact procedures are controlled from the bottom of the "Instrument Controls" tab.



"System Coefficients" opens a popup window with the most critical of the software calibration constants. This is often the easiest place to find the instrument serial number, which is in the upper left corner. None of these numbers should be changed without direct instruction from a MicroCal technician.



"Start in Advanced Mode" tells the software to open with the "Advanced Experimental Design" tab selected by default.

The "ITC Equilibration Options" affect the equilibration before a run starts and are generally only used for troubleshooting. When the "Fast Equil." option is disabled, the system will go through an additional prerun state, a non-stirring equilibration period before stirring is started. This can be useful when stirring-related noise problems are suspected. When the "Auto Mode" option is selected the ITC experiment will proceed through the equilibration states (Prestirring and Final Baseline) and complete the experiment without any further interaction from the user. With this option disabled, the prerun equilibration will not progress from one state to another until the user double clicks on the DP data box.

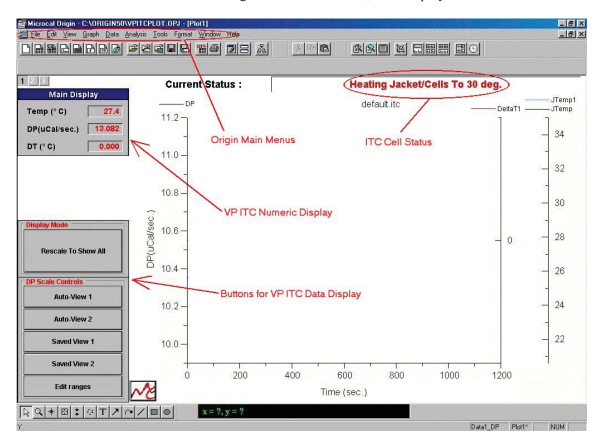


The "Help => Contact Info" menu option provides contact information for MicroCal support. The "About" menu option provides specific details about this version of software.

2.8 Origin Real-Time Display

This section describes the functionality of the optional copy of Origin for real-time display. When the software is opened, it will open the Origin TM project window VPITCPLOT.OPJ for real time data display. This project of Origin is dedicated to data display only, and should not be used for data analysis. Users should open a separate copy of Origin for ITC to perform data analysis.

Pictured below is the main Origin window for ITC₂₀₀ data display.



The ITC cell status, the ITC $_{200}$ numeric display and the buttons for ITC $_{200}$ data tools (as indicated above) have been added for user convenience in viewing data generated by the ITC $_{200}$. Following is a description of each of these sets of tools.

Buttons for Data Display:

This group of buttons allows you to rescale the y-axis by simply clicking on a single button. "Rescale To Show All" shows all the data currently on the plot. "Auto-View 1" will rescale the y-axis so that the last DP data point plotted will be centered on a full scale determined by the entry in the "Full Scale-Auto View 1" text box in the "Edit Ranges" box (see description below). By default, it is $0.1\mu\text{Cal/sec}$. "Auto-View 2" rescales the y-axis to a second scale centered on the last DP data point plotted. By default, it is $1\mu\text{Cal/sec}$. "Saved View 1" rescales the y-axis to a specific range, specified

"Saved View 1" rescales the y-axis to a specific range, specified in the "Edit Range" box. By default, the range is -1 to 1μCal/sec. "Saved View 2" rescales the y-axis to a second specific range. By default, the range is -10 to 10μCal/sec.

"Edit Ranges" opens a window that contains parameters to be used with the aforementioned buttons. In addition to the "Auto View" and "Saved View" entries, which are described above, the "X Axis Options" drop down list box controls the automatic rescaling of the x-axis when the data exceeds the X Axis (time) display range.

"Disabled": Takes no action. Current data may be plotted off screen.

"Rescale": Extends the maximum X Axis setting by 25%, resulting in a larger display range.

"Scroll": Extends the minimum and maximum X Axis setting by 25%, preserving the same range of the display, but always displaying the most recently plotted

Display Mode

Rescale To Show All

DP Scale Controls

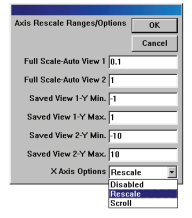
Auto-View 1

Auto-View 2

Saved View 1

Saved View 2

Edit ranges



data. Click "OK" to exit and save any changes or "Cancel" to exit without saving any changes.

The ITC "Main Display" shows the same data channels as the iTC $_{200}$ software: cell temperature, DP, and DT. The status bar shows the current state of the instrument and is the same as the iTC $_{200}$ software status bar.

2.9 Origin Data Analysis Tutorial

- 1. Open a copy of Origin for ITC data analysis. Click on the "Read Data..." button at the top left of the window.
- 2. In the new window, select your CaCl/EDTA file and click "Open". Origin will load the data into a worksheet and calculate and display the integrated heat for each peak (delta H).
- 3. If you did not enter the concentrations earlier, click on the "Concentration..." at the top left of the window. In the popup window, enter the concentrations and click "OK".
- 4. Click on the "Remove Bad Data..." button just below the concentration button. Click on the first data point and press enter to remove the point. This first point is always smaller than the succeeding several points, and must be removed for a better fit. Many users will make a smaller first injection to minimize sample usage on this first injection.
- 5. Click on the "One Set of Sites..." button at the top of the "Model Fitting" section. A new window will pop up, and a red curve will appear in the graph window.
- 6. Click on the "100 Iter." (100 iterations) button until the Chi-squared is as low as it will go. The red curve will come closer and closer to the curve formed by the delta H points. The dialog box will indicate when the Chi-squared is no longer being reduced. This means that the fit is as good as the computer can make it.

Example dialog box text:

Click "Done". A textbox will appear on the graph window.

7. Select Origin menu ITC => Final Figure. A new layout will be created, with the raw data in the top graph and the delta H data in the bottom graph.

Section 3: Instrument Maintenance

This section provides the user with information on the proper maintenance of the instrument to ensure proper function. The tools are pictured and labeled below.





Syringe Cleaning Wire

3.1 Cleaning The Cells And Syringe

Cell Cleaning:

The iTC $_{200}$ uses fixed in place cells in order to provide maximum sensitivity and stability. These cells must be cleaned routinely to maintain the high performance of the instrument. Dirty cells will contribute greatly to cell filling problems, repeatability problems and possibly misinterpretation of data. Inadequate cleaning is the cause of many problems experienced with the iTC $_{200}$.

The reference cell generally requires no special cleaning; rinsing and refilling every week or so is sufficient. The mildest method of cleaning the sample cell is simply to rinse the cell with buffer several times before loading the sample. This should be done whenever the necessary buffer is available.

Every day or two, or after any sample that precipitates, the sample cell should be cleaned with detergent using the Thermovac. Insert the long needle into the Sample cell and push down carefully until the o-ring has sealed, as shown below. Immerse the end of the plastic tubing from the upper tube of the Cell cleaning Apparatus into a beaker of 50-100 ml of detergent cleaning solution. Connect the tubing to the top of the waste flask. Make sure that the other connection on the flask is connected to the ThermoVac and that the ThermoVac cover is seated.









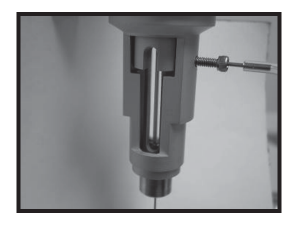
Turn on the ThermoVac vacuum pump. The vacuum will pull the detergent solution from the beaker, through the cell and into the waste flask. Do not allow the waste flask to become full enough for liquid to be sucked into the ThermoVac's vacuum pump, as this may cause damage.

Once sufficient detergent solution has passed through the cell, remove the plastic tubing from the solution and placed into another beaker containing 100 ml or more of water for rinsing. After rinsing with water, remove the plastic tubing from the rinse water and allow time for the vacuum to drain the fluid out of the plastic tubing, then remove the cleaning apparatus from the cell and remove the remaining water from the cell using a long needle syringe.

If this cleaning method is insufficient, fill the cell with \sim 5% LiquiNox and set the instrument to thermostat at 70°C for several hours. Be careful to cool the cells before removing the LiquiNox, as hot liquid may shatter the loading syringe.

Syringe Cleaning:

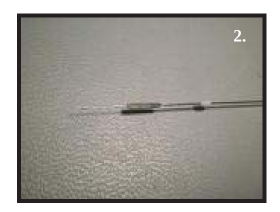
Conceptually, cleaning the syringe is similar to using the ThermoVac to clean the sample cell; the vacuum pulls liquid from a beaker, through the syringe, and into the waste beaker in the ThermoVac chamber. With the syringe in the filling hole, attach the filling tubing. You will need to line up the hole in the pipette with the hole at the top of the syringe. Spin the syringe assembly to find the hole, and then turn it so that the hole is turned "out", toward the outer filling port. The screw on the end of the tubing will align itself as it is screwed in (see image below).



Attach the end of the tubing to the port on top of the waste flask and make sure that the waste flask is attached to the ThermoVac. Move the pipette so that it is in the cleaning hole, furthest from the cell. Push down gently until it seals in place. Turn on the ThermoVac's vacuum. Hold the end of the tubing in a beaker of ~5% Liqui-Nox for several seconds. Make sure that liquid is being pulled into the syringe and through the tubing to the ThermoVac. Hold the end of the tubing in a beaker of laboratory-grade methanol for several seconds, and leave the end open to the air for several minutes to dry the syringe. Once it is dry, turn off the vacuum and disconnect the tubing at the ThermoVac. Move the pipette to the filling hole to fill the syringe.

If clogging of the syringe is suspected, it should be cleaned by running the thin wire through it. This must be done very carefully. First, remove the syringe from the pipette (see section 3.2). As always when handling the syringe, be careful not to bend the needle. The wire should always be inserted through the glass first, both to ensure that the clog is fully removed from the syringe and to decrease the likelihood of bending the syringe. It may be difficult to insert the end of the wire from the glass bore into the metal needle; good light and a magnifying glass will help. Continue to insert the wire until it emerges from the tip of the needle (see images 1 and 2). Carefully pull the wire back through the needle and glass bore.

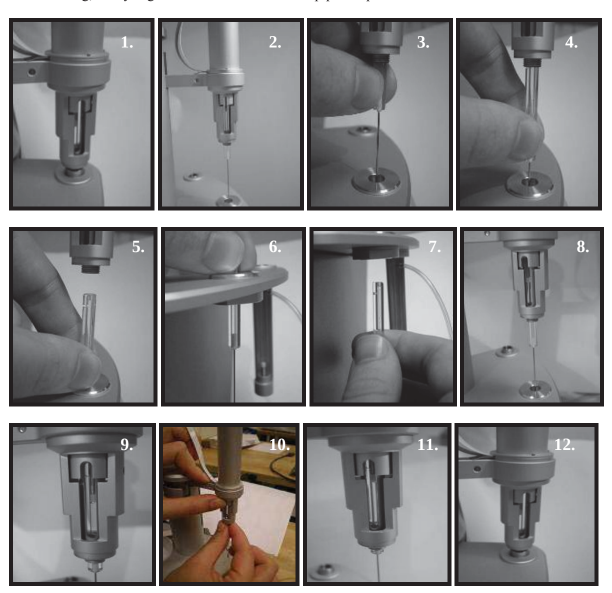




3.2 Changing Injection Syringes

Clean and dry the syringe before removal. Remove sample tube form loading hole and disconnect the cleaning apparatus. Be very careful whenever the syringe is not secured in the pipette, as the needle bends extremely easily. See images 1-7 below for illustrated removal.

Slide the pipette to the cleaning position and insert it firmly into the hole. Turn the inner metal syringe holder to the right several full rotations. This will unscrew the metal ring that holds the syringe in the pipette. One the ring is loose, move the pipette to the empty loading hole. Use the manual pipette controls to lower the pipette tip about 0.3 inches. Without the bottom ring, the syringe will move down with the pipette tip.

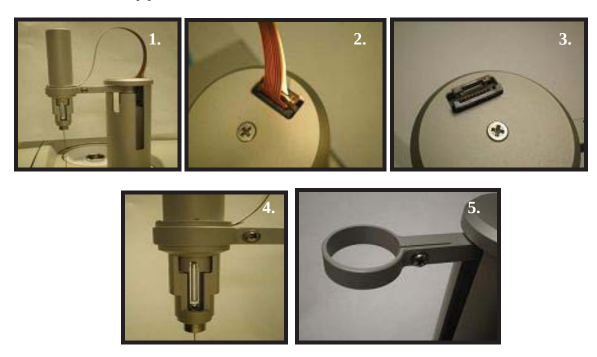


Enough of the syringe bore should now be exposed to hold it firmly and pull straight down, pulling the syringe out of the pipette and down through the filling hole. The soft-grip tweezers can be used to help grip the syringe without damaging it. Carefully put the syringe in a place where it will not roll or be damaged.

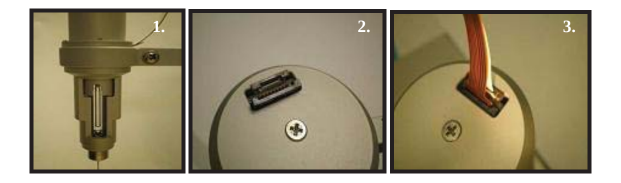
See images 7-12 for illustrated syringe replacement. Gently push the syringe up through the filling hole and into the pipette. It will come to a stop with about 4mm of syringe glass exposed below the metal. Carefully push up on the glass while spinning the metal syringe holder slowly. Once the notch in the syringe aligns with the notch in the holder, the syringe will slide up approximately another 2mm. Move the pipette to the cleaning hole and turn the inner metal syringe holder to the left until the bottom ring is secure.

3.3 Removing Pipette

The pipette is held by the arm and connected to the tower by two small cables. Detach the two cables from the top of the tower. First pull up on the tiny tabs on each side of the locking pieces. The cables should slide out of their sockets. Loosen the screw on the side of the arm to remove the pipette.

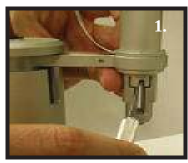


To reinsert the pipette, place the pipette in the socket on the arm and tighten the screw. Insert the cables into their slots, and press the locking pieces back down to hold them. The white cable goes in the slot to the rear of the instrument, and the orange/brown cable goes into the front slot.



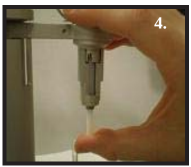
3.4 Changing Pipette Tips

See the instructions above to remove the syringe from the pipette. The pipette should remain connected to the tower. In the "Instrument Controls" tab, click the "Remove Old Tip" button to position the tip. Use the blade of the X-acto knife provided with your instrument to make a diagonal cut on the side of the white Teflon tip (image 1), and use the small-tipped tweezers to pull the tip off (image 2). Click the "Install New Tip" button to position the tip. Use the tip pusher tool, with the new tip in the slot hole-side up, and firmly press the new tip into place (image 3-5). Rotate the tool to ensure that the tip is firmly seated. Remove the tip pusher tool and inspect the tip to be sure it is seated properly. Reinstall the syringe as described above to finish.







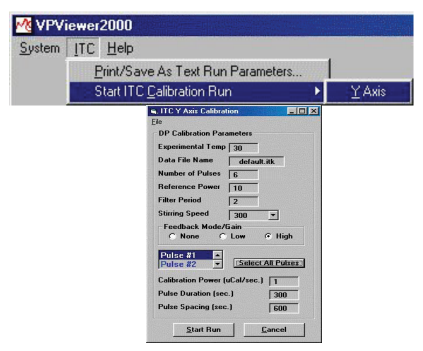




3.5 Y-axis Calibration Check

It is recommended that the y-axis calibration be checked every few months to ensure accurate data. The automatic calibration check routine will send a series of pulses to the cell heaters, dissipating a known power. The offset in the DP as a result of this power is analyzed in comparison to the correct DP offset.

Make sure the cells are clean, and fill both cells with degassed, distilled water. Load the titration syringe with water and insert it into the sample cell. It is recommended that Origin for real-time data be enabled.

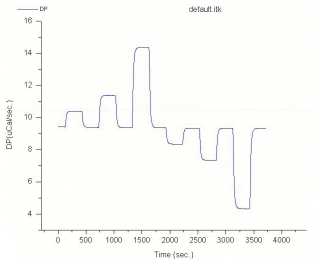


To begin the y-axis calibration check procedure select iTC_{200} software menu "ITC => Start ITC Calibration Run => Y Axis Check". Once the menu has been selected, the Calibration Pulse Setup Window will appear. This window allows the calibration pulses to be modified. Individual pulse parameters are entered by first selecting a pulse or multiple pulses, then entering the desired parameter value into the pulse parameter boxes (Calibration Power, Pulse Duration and Pulse spacing). Users are encouraged to simply use the default y-axis calibration parameters. After the run and pulse parameters are entered, clicking on the "Start Run" button will start the run. The ITC will equilibrate in the same manner as it would during a titration experiment.

If creating customized calibration parameters, users must be aware of the DP range limits when setting reference power and pulse sizes. The reference power must be high enough to allow all pulses without hitting saturation, and if a pulse size is too small, it can show abnormally high error.

After the final equilibration phase has completed, the initial delay will begin and the pulses will be applied as entered. As each pulse completes, Origin will analyze the pulse region and determine the deflection of the baseline as well as the energy (area) of the pulse. The requested power and energy will also be displayed as will a percent error for both power and energy. The reported error in deflection or energy should be less than 1%. If the error is reported as higher than 1%, please contact MicroCal.

For a more rigorous analysis, once the calibration is done and the system is thermostatting again, open the ITC Calibrations project. Click on the "Y-Axis Calibration (DP,



uCal/sec)" button. Origin will ask for the DP check file. Select the data file just created and click "Open". The computer will think for a few moments. If any of the pulses are out of specifications, a pop-up will inform you. It will ask you to save.

Origin will show four graphs. The upper left graph holds the raw data. The lower two graphs show the energy and power of each pulse. The upper right graph displays the percent error for the energy and power of each pulse. Right-click at the upper right portion of the graph and select "Go To Window" from the menu. Check the sizes of the errors. If any of the errors is greater than 1%, please contact MicroCal.

3.6 Temperature Calibration Check

It is recommended that the ITC temperature calibration be checked approximately once every 12 months. The measured temperatures should be within \pm 0.2 degrees of the set temperatures in the iTC₂₀₀ software. This procedure requires a Control Company digital meter and submersible probe or equivalent. The Control Company meter is model #9612; the probe is model #4021.

Make sure the cells are clean, and load them with degassed, distilled water. Residue in the cells or empty cells will yield unpredictable results. Insert the probe into the sample cell and turn it on

- 1. Use the "Temperature" control box under the "Instrument Controls" tab in the iTC_{200} software to set the temperature to 30°C.
- 2. Set the Plot Idle Data flag so that the current data is displayed.
- 3. When the instrument is thermostatting at the target temperature, the DP has settled at a negative, stable value, and the DT is 0, use the data reader tool and the display from the meter to take temperature data points, as close together as possible.
- 4. Repeat for the second check point at 70°C. If the error for each is not less than 0.2°C, please contact MicroCal for assistance.

It is recommended that users keep a log with the date of the calibration check and the software and meter readings for each temperature point.