

# PhyTip<sup>®</sup> MEA Purification System Manual



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# Operating Instructions for PhyTip® MEA Personal Purification System

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## *Introduction*

In order to extend the benefits of PhyTip® columns for protein purification and enrichment, PhyNexus has developed the PhyTip® MEA Personal Purification System as the platform of choice for fully automated protein purification and enrichment. When used with the PhyNexus Operating Software, this system offers a range of flow rates that maximizes the purification and enrichment efficiency of the various affinity resins available for PhyTip® columns. In combination with the fully programmable positioning of the 12 channel system, and the ability to place samples, PhyTip® columns, wash stations and elution plates in any available position, the MEA Personal Purification System is an ideal automated system for micro scale protein purification.

These operating instructions are divided into the following sections:

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# Section A

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## 1. *Unpacking*

Each MEA system contains the following:

1. MEA Purification System
2. Two power supplies (one for MEA, one for 12 channel system). Use only the power supplies provided, see page 73 for ratings.
3. ME 12 channel system(s). The 12 Channel system(s) will be provided with attachment clamp already installed, ready to attach to the robotics tower.
4. Software CD
5. Communications cable
6. Consumables pack including PhyTip Evaporation Covers, Holders and 96-well plates
7. Literature, documents and Operating Instructions pack
8. Any optional cooling accessories ordered will ship with a third power supply. Use only the power supplies provided, see page 73 for ratings.

If these separate items are not in the original outer packing, please call PhyNexus Inc, at 408-267-7214.

Remove all contents from their individual packs and prepare to install the system.

## ***2. Setting up the system***

This section is divided into:

1. Initial location and set up of system parts
2. Installing the software
3. Powering up the system
4. Cleaning the system

### ***1. Initial location and set up of system parts***

#### **Quick guide:**

Once all contents have been removed from their individual packs, locate the MEA instrument and place it close to the computer to be used for operation. Remove the 12 channel system and attach to the robotics tower using the adaptor. Connect the communications cable between MEA COM1 and the computer. Connect the power supply to the MEA and separate power supply to the 12 channel system, and if installed, connect the power supply to the cooler accessory. Locate the Software CD and install in the computer.

#### **Detailed description:**

##### *Instrument placement.*

Remove the MEA system from its packing box and place on the specified lab bench. This must be close enough for two power supplies that are required to operate the MEA system (if the optional cooling accessory has been ordered, this will require a separate power supply). Enough vertical clearance must be given for the robot tower to move along the x axis (maximum height above the bench = 24”), and there must be enough space on the right hand side of the instrument to reach/press the red **Emergency Stop** button and the control switch for the cooling option. The surface for the MEA should be smooth and flat.

The full instrument dimensions are:

Width: 32”  
Depth: 14”, plus 6” to accommodate all necessary power supplies and cables.  
Height: 24”  
Weight: 55 lbs

##### *Installing the instrument.*

Each MEA is shipped with the ordered 12 channel ME system(s), either ME200+ (for 200+ PhyTip columns that operate with volumes up to 200 µL) or ME1000+ (for 1000+

PhyTip columns that operate with volumes up to 1000  $\mu\text{L}$ ) with the alignment clamp attached. Remove the ME system from its packing and install this unit onto the receptacle on the robot tower as follows:

1. The black alignment clamp has been factory installed on the 12 channel system. The LCD on the 12 channel system is at the front of the unit and the clamp has an alignment slot at the rear which will interface with an alignment bar on the moving Z-head on the robotic tower. The moving Z-head is a round bar with a single flat surface that extends forward of the robotic tower and the alignment bar that is parallel to the x axis of the system.
2. Locate the single attachment hole at the rear of the clamp and push the clamp onto Z-head bar.
3. With the alignment bar and alignment slot parallel to each other, push the clamp mechanism in fully until it snaps in place.
4. Turn the knurled screw on the side of the clamp to finally tighten the clamping mechanism in place.

To exchange the 12 channel system, simply loosen the knurled screw, ease the clamp from the alignment slot and pull away from the z-head bar. Repeat the first four steps with the alternate 12 channel system.

When replacing a 200+ channel system with a 1000+ 12 channel system, remove the front and rear retaining bars that are at positions 1 and 2. There are three allen key screws (two at the front and 1 at the rear). Unscrew the three screws and remove the retaining bars. The new height of positions 1 and 2 will be ready for the deep tip boxes used with the 1000+ 12 channel system.

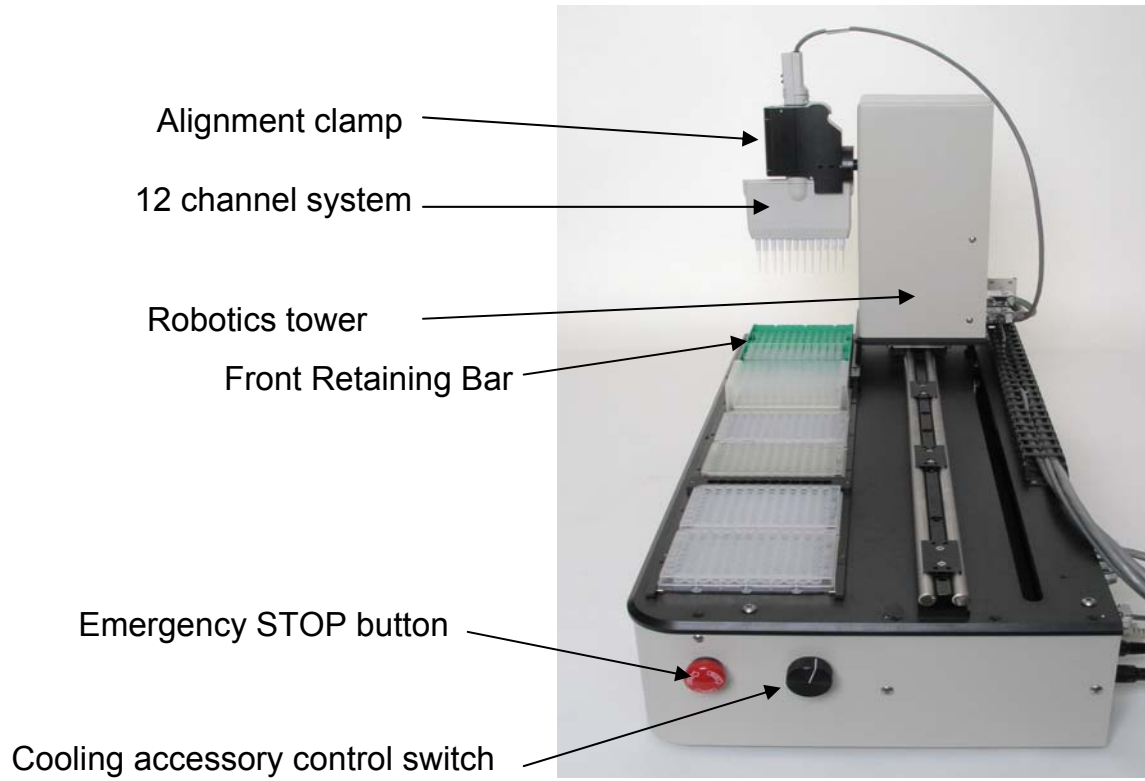
Take the communication cable that is attached to the 12 channel system and route it through the cable management channel on the robot track down to the communications port at the rear of the MEA (COM2). This is achieved by just pressing the cable into the track and snapping it in place. Using a flat head screw driver, attach the communications connector to the MEA.

Take the MEA power supply and attach to the supply port on the rear of the MEA (labeled "Power"). Take the 12 channel system power supply and attach to the 12 channel power supply port that is on the gray communication cable that runs from the top of the 12 channel system to the MEA. Connect all power supplies to the main voltage line.

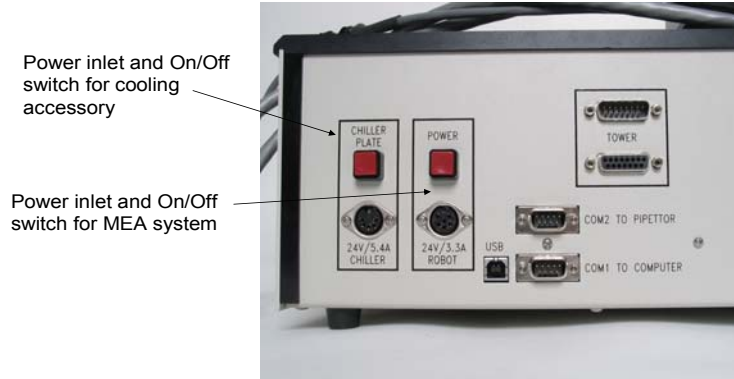
Take the separate communications cable and connect between the MEA port COM1 and a serial port of the computer to be used.

If the optional cooling accessory has been installed, simply connect the additional power supply at the rear of the instrument in the port labeled "Chiller Plate."

*Side view of MEA system.*



*Rear view of MEA system showing power supply inlets and cable attachment points*



## *2. Install the software in the PC*

The PhyNexus Operating Software is designed to operate with PCs running Windows XP, 2000, NT, and 98. With the computer turned on, insert the software disc into the CD or DVD drive of the system, double click on the setup icon, and follow the install instructions on the screen.

## *3. Powering up the system*

With all power supplies connected and the software installed in the computer, turn the system on by pressing the power supply switch at the rear of the MEA. A green “Power On” light will illuminate at the front of the instrument. If the green light does not illuminate, first check that all connects have been made and secondly make sure that the red emergency stop button has not been depressed. To do this, turn the emergency button in the direction for Reset (clockwise).

If the optional cooling accessory is installed, turn the main power on by pressing the power switch at the rear of the instrument and then turning the setting switch on the right of the instrument to the appropriate setting.

## *4. Cleaning the system.*

The MEA Personal Purification System can be cleaned by wiping with a damp cloth. Do not use solutions containing high concentrations of bleach or other oxidizing agents.

### 3. Using the PhyTip MEA Purification System

This portion of the manual is divided into the following sections:

1. Setting the instrument deck
2. Easy to Use Quick Guide Software Instructions
3. Easy To Use Quick Guide Operating Instructions
4. Detailed Software Information and Instructions

#### 1. Setting the instrument deck

The instrument deck can be configured in a number of different ways, but the standard configuration is as follows: On the left there are two positions for Tip boxes and to the right of these, there is enough space to hold 5 plates (standard SBS) and one reagent reservoir.

- Position 1. PhyTip Box
- Position 2. PhyTip Box
- Position 3. Plate position (Plate 1)
- Position 4. Plate position (Plate 2)
- Position 5. Plate position (Plate 3)
- Position 6. Reservoir position
- Position 7. Plate position (Plate 4)
- Position 8. Plate position (Plate 5)

If necessary, the positions of these plates can be changed by moving the attached pins and replacing them in the appropriate positions. This maintains perfect alignment of each plate in each position during MEA operation.

If the MEA has been installed with the PhyTip Cooling Accessory, the accessory will require its own power supply. When on, the cooling accessory will maintain the last two positions under the deck (Positions 7 & 8) at a temperature of approximately 4-6°C. This temperature can be modified by turning the control knob on the right side of the instrument. Position 1 represents the coldest setting and position 5 the warmest setting available as demonstrated below:

Position	Approximate Temperature (°C)
1	4
2	8
3	12
4	16
5	20

## **For a standard purification set up:**

### **Position 1:** PhyTip Columns – new/unused

For the MEA system, these should only be PhyTip columns with the appropriate part numbers for the 12 channel system that is being used, e.g. 200+ or 1000+, and the affinity resin that is required e.g. PTR 96-05-01 (Pack of 96 PhyTip columns for the 200+ system with Protein A as affinity resin).

### **Position 2:** Empty box for used PhyTip columns

Used PhyTip columns can be replaced in their original box or if required, in a separate empty box.

### **Position 3:** Extra plate position

This extra position can be used for a number of different applications e.g. extra wash step etc.

### **Position 4:** Wash Plate 1

Depending on the specific application, a number of different types of wash plates may be placed in this position, e.g. deep well 96-well plates. Care must be taken when choosing the type of deep well 96-well plate. When using the 1000+ system, PhyNexus recommends the use of 2 mL, square well plates (PhyNexus Part Number PCP 96-20-10).

### **Position 5:** Wash Plate 2

Depending on the specific application, a number of different types of wash plates may be placed in this position, e.g. deep well 96-well plates. Care must be taken when choosing the type of deep well 96-well plate. When using the 1000+ system, PhyNexus recommends the use of 2 mL, square well plates (PhyNexus Part Number PCP 96-20-10).

### **Position 6:** Reservoir Position

Depending on the application and specific affinity resin used, the Reservoir Position can be used as a holder for neutralization buffer. When used in this mode, the reservoir is filled with sufficient volume of neutralization buffer. Also Tip box position 2 should be used to hold standard Rainin LTS Pipette Tips. The system should then include the incorporation of a neutralization step using the Reservoir Position; use PhyNexus “reservoir inserts” (PhyNexus Part Number 96-60-00). The maximum total volume per well for each of the reservoir wells is approximately 300  $\mu$ L, therefore maximum recommended volume to be transferred (assuming 8 rows in a full 96-well plate) is 35  $\mu$ L.

**Position 7:** Sample Plate (samples to be purified)

Depending on the specific application, a number of different types of sample plates may be placed in this position, e.g. deep well 96-well plates. Care must be taken when choosing the type of deep well 96-well plate. When using the 1000+ system, PhyNexus recommends the use of 2 mL, square well plates (PhyNexus Part Number PCP 96-20-10).

**Position 8:** Elution Plate

PhyNexus recommends the use of the PhyTip Elution Plates (Part number PCP 96-00-10) for the final step of elution. These plates have v-bottoms which maximize the process of elution. It is also recommended to cover the plates, once the elution buffer has been added, with a PhyTip Plate Cover (Part number PCP 96-10-20) and PhyTip Plate Holder (Part number PCP 96-10-50).

With the cooling accessory attached, both positions 7 and 8 can be maintained at sub ambient temperatures.

## ***2. Easy to use Quick Guide Software Instructions***

The Auto controller Operating Software has been designed for simple to use automated methods that have been preprogrammed into the system. However, any combination of movements and positions can be built into a user defined method. To begin, PhyNexus recommends that the user start with the built-in methods and subsequently modify these to match the requirements of the application.

**Methods** manage the entire process of moving the robotics tower with 12 channel system attached to perform the Capture, Purify and Enrich process for all of the defined samples, **Protocols** manage the individual steps of the Capture, Purify and Enrich process. A method is a series of protocols and robotic movement/position instructions that fully automate the entire process of purifying a full plate of 96 samples. To load the software onto a PC, simply insert the CD into the CD or DVD drive of the PC and follow the installation instructions on the screen.

For simplest use of the system, run the preprogrammed methods for any of the given affinity resins. To do this, go to **File**, then **Open**. This will bring down a screen that shows either Method or Protocol, click on **Method** to show all of the stored Methods. Open the standard method IMAC; this will bring down the operational screen. Now go to **Method** and click on **Edit**; this will show all of the individual steps of the method.

Click on any one of the individual steps of the methods to show the options screen; this will be split into two sections. The top section deals with the operation of the 12 channel system (controller), while the bottom section deals with robotics movement of the autocontroller. Actions that are available are:

### **Top Section:**

*Execute:* executes a stored protocol, e.g. Capture

*Audible Alert:* introduces an audible alert at the end of an action

*Home (controller):* homes the 12 channel system to a zero volume

*Pause for Operator:* builds in an automatic pause in the method

*Comment:* allows operator to build in an automatic comment or prompt between actions

*Blowout:* causes a 12 channel system to blow out any air from the syringe

**Bottom Section:**

*Home Autocontroller:* returns the robotics tower and 12 channel system to registered Home position

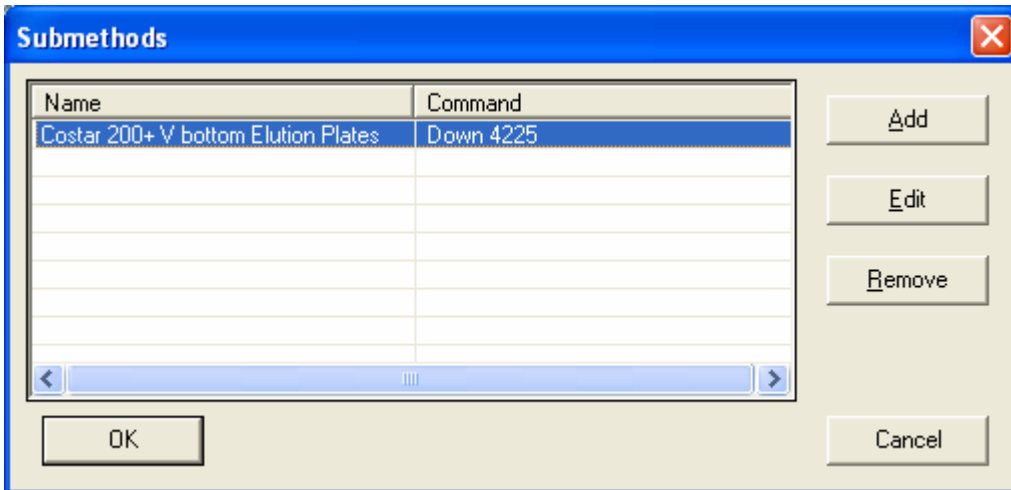
*Down:* moves the robotics tower with 12 channel system down to designated step position (0 is the highest position above the deck and 4725 is the position closest to the deck). A dialog box will appear where the number of steps can be entered.

*Up:* moves the robotics tower with 12 channel system up to the designated step position (0 is the highest position above the deck and 4725 is the position closest to the deck). A dialog box will appear where the data can be entered, if left unchecked the default is all the way up to the zero position. If data is entered, this information is only a delta, meaning that if 20 is entered the tower will move up 20 positions and NOT to location number 20.

*External Pump:* operates an external pump (when fitted) via the rear serial port, allowing for the automated addition of reagents.

*Rotate:* rotates the vertical 12 channel system (clockwise = 0 to + 335 and Counterclockwise = 0 to - 25)

*Sub method:* allows user to define a specific action and store this in memory e.g. “Costar 200+ V bottom plate” could reference the specific z height of a novel plate, once stored in memory, z numbers would not need to be recalled.



*Position:* moves the robotics tower with 12 channel system to the desired x-axis location on the deck. There are a total of 39,100 positions along the x-axis.

*Position/Row* moves the robotic tower with 12 channel system to the desired Position (1-8) and specific Row (1-8). This command assumes the use of plates that have 8 rows (e.g., standard 96 well plates, certain 48 well plates, etc.).

*Load Tips:* loads a row of 12 PhyTip columns onto the 12 channel system. This also incorporates the function of going down to the tips, loading the tips and moving them out from the tip box. This is set for the default PhyTip columns and standard Rainin LTS tips for neutralization.

*Eject:* ejects all 12 PhyTip columns from the 12 channel system

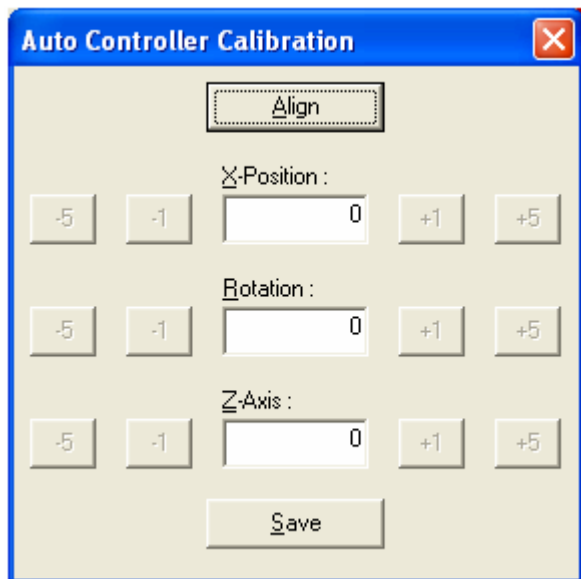
*XMOV:* Auto movement to specify position along x axis. CAUTION: This action does NOT move tower to zero position.

### 3. Easy to Use Quick Guide Operating Instructions

#### A. Calibrating the Instrument

Once installed in the correct position the instrument should be recalibrated in order to correctly zero the position of the robotic tower. It is also recommended that for each pipettor, (200 or 1200) calibration should be checked and stored.

To access the calibration software go to the general toolbar on the operating software and click on **Options**, this will display *Communications, Syringe, Calibration, Manual Positioning* and *SubMethods*. Click on **Syringe** and select either 200 or 1200 (for the 12 channel system that uses 200+ tips, choose 200: for 12 channel system that uses 1000+ tips, choose 1200). Next, click on **Calibration**, this will now display a screen with “Align” and three dialog boxes for *x-position, rotation and z-axis*.



Click align, and the 12 channel will move to the zero position on x-axis and then down to the normal tip loading height for the specific 12 channel system that is connected e.g. 200 or 1200. Adjustments can be made for x-axis, z-axis and rotate as follows:

For the x-axis, adjust the +1 and -1 settings to align the tip holders on the 12 channel system in the center of the tips

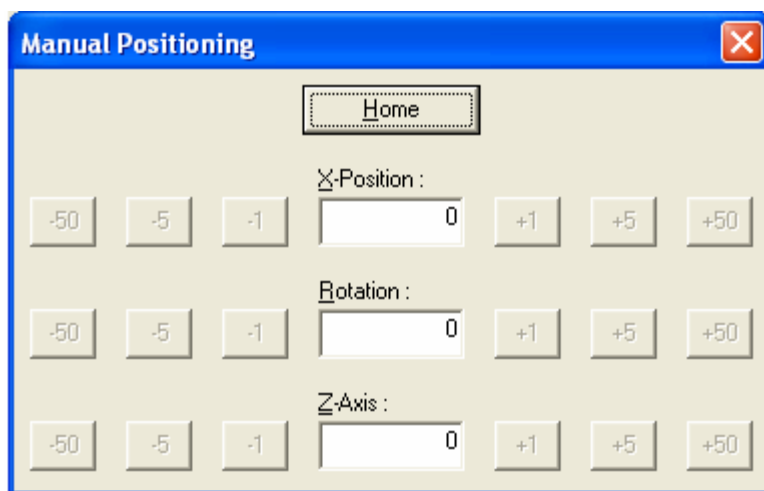
For the z-axis, adjust the +1 and -1 settings to bring the base of the tip holders on the 12 channel system to approximately 2 mm above the top of the tips.

For the rotation, adjust the +1 and -1 settings to bring the plane of the base of the tip holders on the 12 channel system parallel with the plane of the top of the tips.

Once complete, click Save. This will save the calibration settings for the specific 12 channel system. The software recognizes the difference between a 200+ system and 1200+ system and applies the saved settings accordingly.

## Manual Positioning

This allows the user to position the 12 channel system, anywhere on the deck. This is particularly useful when used in conjunction with Sub-method. Care must be taken when using this part of the software as the z-axis does not automatically return to a safe height above the deck, but instead moves the 12 channel system along the specific z-axis height that is seen on the screen.



### B. Instrument programming

Before programming a specific method note all of the specific steps that are likely to be carried out in the process that needs to be accomplished. Below is an example of some of the steps involved in a basic Method.

1. Prime the 12 channel system and home.
2. Move to Row 1 of Box 1 of PhyTip, move robot tower down and pick up row 1 of PhyTip columns.
3. Move robot tower up and the across to row 1 of sample plate at position 3.
4. Move tower down so that PhyTip columns enter the sample.
5. Begin preprogrammed Capture Step.
6. Upon completion of Capture step, bring tower up and move across to row 1 of wash plate 1 in plate position 4.
7. Move tower down so that PhyTip columns enter wash buffer 1.
8. Begin preprogrammed Wash 1 Step.
9. Upon completion of Wash step 1, bring tower up and move across to row 1 of wash plate 2 in plate position 5.
10. Move tower down so that PhyTip columns enter wash buffer 2.
11. Begin preprogrammed Wash 2 Step.
12. Upon completion of wash step 2, bring tower up and move across to row 1 of Enrichment Step in plate position 6.

13. Move tower down so that PhyTip columns enter Elution buffer.
14. Begin preprogrammed Enrichment step.
15. Upon completion of this step bring tower up and move back to row 1 of the original tip box in position 1. Move tower down and discard tips in row 1.
16. Repeat the process for row 2, etc.

The robotics tower is stepper motor driven and as such can be controlled by discrete steps in both the X and Z axis. In the Z axis, there are 4,725 steps and in the X axis, there are 39,100 steps. The robotics head can be driven to any of these steps by entering position numbers in the appropriate part of the software. Standard methods, built into the software, already have these positions preprogrammed.

To simplify this process the following standard and guidelines will help to speed up the process of programming:

***When using the 200+ 12 channel system:***

The simple programming of the software allows the user to move to any position and any row. However, to set the height of the tips, use these guidelines:

For z-axis heights in positions 3,4,5,7 & 8 it is assumed a PhyTip plate is used.

For z-axis heights:

**PhyTip columns**

	<b>Load</b>	<b>Eject</b>
Position 1	4210*	4100
Position 2	-	-
	<b>With cooler</b>	<b>Without cooler</b>
Position 3	4280	4280
Position 4	4280	4280
Position 5	4280	4280
Position 6	-	-
Position 7	4225	4280
Position 8	4225	4280

**Rainin LTS**

	<b>Load</b>	<b>Eject</b>
Position 1	-	-
Position 2	4210*	4100
	<b>With cooler</b>	<b>Without cooler</b>
Position 3	-	-
Position 4	-	-
Position 5	-	-
Position 6	4070	-
Position 7	4140	4195
Position 8	4140	4195

\* This is the initial z setting, then the 12 channel system moves down another 60 into the tips. This is for reference only. The system defaults to automatically load tips with the “Loadtips” command. The user should NOT program a down 4210 and follow this with a “Loadtips” command.

***When using the 1000+ 12 channel system:***

The simple programming of the software allows the user to move to any position and any row. However, to set the height of the tips, use these guidelines:

For z-axis height it is assumed that for positions 3, 4, 5 & 7, a PhyTip Deep Well Plate is used, and for position 8, a PhyTip Plate is used.

For z-axis heights:

**PhyTip columns**

	<b>Load</b>	<b>Eject</b>
Position 1	3600*	3500
Position 2	-	-
	<b>With cooler</b>	<b>Without cooler</b>
Position 3	2350	2350
Position 4	2350	2350
Position 5	2350	2350
Position 6	-	-
Position 7	2325	2380
Position 8	2325	2380

**Rainin LTS**

	<b>Load</b>	<b>Eject</b>
Position 1	-	-
Position 2	3600*	3500
	<b>With cooler</b>	<b>Without cooler</b>
Position 3	-	-
Position 4	-	-
Position 5	-	-
Position 6	2050	-
Position 7	2135	2135
Position 8	2190	2190

\* This is the initial z setting, then the 12 channel system moves down another 100 into the tips. This is for reference only. The system defaults to automatically load tips with the “Loadtips” command. The user should NOT program a down 4210 and follow this with a “Loadtips” command.

**B. Reagents**

**1. Preparing samples ready for Capture – Purify - Enrich™**

It is recommended that prior to the process of purification and enrichment by PhyTip columns, all samples be checked for particulates and clarity. If the final solution to be purified appears cloudy or contains particulates, it is recommended to spin the samples down one more time at 3,000 g for 5 minutes.

Samples can be placed in any position. However the default methods all have samples placed in position 3. With the cooling accessory (positions 7 & 8), it is possible to maintain both the pre-purified material and the purified sample at sub ambient temperatures.

## 2. Preparing wash buffer(s) ready for Purification process

Refer to the Product Insert that is included with each Affinity resin for details on each buffer used in the purification process. Some buffers are shipped as concentrates and will require dilution before use.

## 3. Prepare Elution Buffer ready for Enrichment process

Refer to the Product Insert that ships with each Affinity resin for details on the Elution Buffer used in the enrichment process. If using Protein A or Protein G affinity resins, it is recommended to add Neutralization Buffer at the end of the Enrichment step in order to return the sample to physiological pH. A separate Neutralization step should be incorporated into the automated purification process of the MEA (see standard methods). Once the elution buffer has been added to the elution plate, place the PhyTip Cover over the plate (see separate instructions) and place in the appropriate position. Place the PhyTip Holder over the plate in order to retain the plate in position during the final elution process.

## 4. Load the appropriate Method File

To load the appropriate Method File, go to **File** on the user screen. Enter **Open** and the choice of Protocol or Method will be shown. Enter **Method** and hit **OK**. This will take you to folders containing the preprogrammed method files available to run. Select the required file and press **Open**. The Method will now be available for the system to run. To verify the method click on **Method – Edit**, this displays the robotic steps and all of the linked protocols that constitute the given method. To run this method, press **Method – Execute**.

The PC is in constant communication with the MEA. Please do not run other programs while running the PhyNexus Operating Software.

## 5. Store/analyze samples

Once the final enrichment step has been completed, samples will be ready for storage/analysis.

## Standard Methods

### Example 1 - IMAC:

*Figure 1* below is an example of a standard method (without chiller) of Capture, Purify, Enrich using IMAC (Ni-NTA) PhyTip columns for one row of 12 x 200  $\mu$ L samples. For information on configuration of the instrument with and without chiller, see page 29.

Deck configuration for this example is:

Position 1, Row 1: IMAC 200+ PhyTip columns

Position 2, Row 1: Empty

Position 3, Row 1: Empty

Position 4, Row 1: Recommended volume = 200  $\mu$ L of Wash 1

Position 5, Row 1: Recommended volume = 200  $\mu$ L of Wash 2

Position 6, Row 1: Empty

Position 7, Row 1: Recommended volume = 200  $\mu$ L of Sample

Position 8, Row 1: 20  $\mu$ L Elution

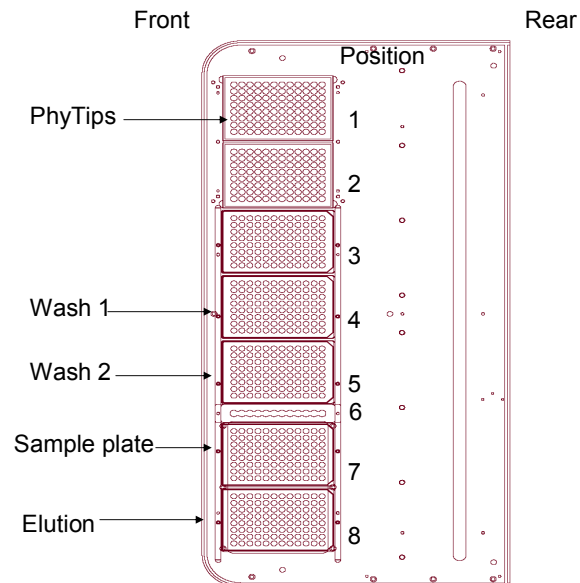


Figure 1 Standard Method for Capture, Purify & Enrich

Command	Parameters (if any)
Home Auto Controller	
Blowout	
Home Controller	
Execute Protocol	c:\PN200+Default IMAC Condition
Position	1,1
Loadtips	
Position	7,1
Down	4225
Execute Protocol	c:\PN200+Default IMAC Capture
Position	4,1
Down	4280
Execute Protocol	c:\PN200+Default IMAC Wash1
Position	5,1
Down	4280
Execute Protocol	c:\PN200+Default IMAC Wash2
Position	8,1
Down	4225
Execute Protocol	c:\PN200+Default IMAC Elution
Position	1,1
Down	4100
Eject	
Home Auto Controller	

This method shows the automated movement of the 12 channel system for 1 row of samples. In detail this can be broken down as follows:

### 1. Homing the system

Command	Parameters (if any)
Home Auto Controller	
Blowout	
Home Controller	
Execute Protocol	c:\PN200+Default IMAC Condition
Position/Row	1,1

**Home Auto Controller** moves the robotics tower to the Home position – full left position.

**Blowout** performs a complete blowout of the 12 channel system.

**Home Controller** performs a zeroing of the 12 channel system followed by a resetting of the system to zero.

**Execute** (c:\PN200+Default IMAC Condition) executes a stored protocol which in this case is a conditioning step. The 12 channel system withdraws air into the syringes before the PhyTip columns are attached. It does this to create an air buffer in the system that will later be used to push liquid out of the PhyTip columns. The volume of air will vary depending on the specific set of protocols being used.

**Position 1, Row 1** moves the robotics tower over position number 1 and row 1 (the tray of PhyTip columns).

## 2. Moving the PhyTip Columns and Capturing the Sample

Load tips	
Position	7,1
Down	4225
Execute Protocol	c:\PN200+Default IMAC Capture

**Load tips** moves the 12 channel system down towards the tips in row 1, then positions the 12 channel system at an angle to load and then secure all 12 PhyTip columns.

**Position 7, Row 1** automatically moves the robotics tower to the full extent (Position 0) and then moves it over to position 7, row 1 (the plate holding the sample to be purified).

**Down** moves the robotics head with PhyTip columns down into the sample solution in row 1 of position 7. This is a predefined position of 4225 (assumes use of PhyTip Plates with known dimensions) in readiness for the Capture step.

**Execute** (c:\PN200+Default IMAC Capture) executes the stored protocol that implements the Capture process with a given number of cycles, flow rates, volumes and pauses, e.g. in the standard method, this will be a volume of 170 µL for 2 cycles at a flow rate of 250 µL/min with 10 second pauses at the end of each step. Also note that at the end of the capture step, the last cycle expels 172 µL to make sure all of the sample solution is removed.

## 3. Wash and Elution Steps

Position	4,1
Down	4280
Execute Protocol	c:\PN200+Default IMAC Wash1
Position	5,1
Down	4280
Execute Protocol	c:\PN200+Default IMAC Wash2
Position	8,1
Down	4225
Execute Protocol	c:\PN200+Default IMAC Elution
Position	1,1
Down	4100
Eject	
Home Auto Controller	

**Position 4, Row 1** moves the robotics tower to the full extend position (Position 0), and then moves it over to position 4, row 1 ready for the first wash step.

**Down 4280** moves the robotics tower down to predefined position of 4280 (assumes use of PhyTip Plates with known dimensions) ready for the wash step.

**Execute** (c:\PN200+Default IMAC Wash 1) implements the stored protocol for the first wash step. This protocol, like Capture, will control the flow rate, volume, pause and number of cycles involved in this step.

**Position 5, Row 1** moves the robotic tower up to the fully extended zero position and then moves it to position 5 and row 1 in readiness for the next wash step.

**Down 4280** moves the robotics tower down to predefined position of 4280 (assumes use of PhyTip Plates with known dimensions) ready for the next wash step.

**Execute** (c:\PN200+Default IMAC Wash 2) implements the stored protocol for the second wash step. This protocol, like Wash 1, will control the flow rate, volume, pause and number of cycles involved in this step.

**Position 8, Row 1** moves the robotics tower to the full extend position (Position 0) and then moves it over to position 8, row 1, ready for the elution step.

**Down** moves the robotics tower down to predefined position of 4225 (assumes use of PhyTip Plates with known dimensions) ready for the elution step.

**Execute** (c:\PN200+Default IMAC Elution) implements the stored protocol for the elution step. This protocol, like Capture, will control the flow rate, volume, pause and number of cycles involved in this step. Note at the end of this step, the last expel step is greater volume than the other expel steps in order to maximize the total volume of elution buffer returned to the elution plate.

**Position 1, Row 1** moves the robotics tower to the full extend position (Position 0) and then moves it over to position 1, row 1, ready for the ejection of the PhyTip columns.

**Down 4100** moves the robotics tower down to predefined position of 4100 ready to eject the PhyTip columns back into their original position in row 1.

**Eject** is a command to remove all of the now used PhyTip columns.

**Home Auto Controller** takes the robotics tower back to its safe starting position and completes the purification of 1 row of samples.

If using this program as a template, just cut and paste the entire page and replace row 1 with row 2, etc.

When a method is running, each step of the process will display on the screen.

**Completed** steps will display **Green**, while steps **In Progress** will display **Blue**.

## Example 2 – Protein A:

Figure 2 below is an example of a modified method (without chiller) of Capture, Purify & Enrich for Protein A for one row of 12 x 200  $\mu$ L samples. For information on configuration of the instrument with and without chiller, see page 29.

Deck configuration for this example is as below.

Position 1, Row 1: Protein A 200+ PhyTip columns

Position 2, Row 1: Rainin LTS Tips

Position 3, Row 1: Empty

Position 4, Row 1: 200  $\mu$ L of Wash 1

Position 5, Row 1: 200  $\mu$ L of Wash 2

Position 6, Reservoir with Neutralization Buffer

Position 7, Row 1: 200  $\mu$ L of Sample

Position 8, Row 1: 20  $\mu$ L Elution

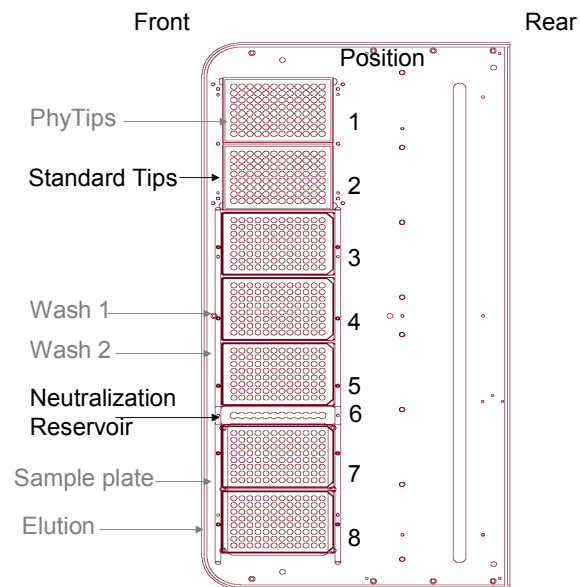


Figure 2 Standard method including addition of Neutralization step

Command	Parameters (if any)
Home Auto Controller	
Blowout	
Home Controller	
Execute Protocol	c:\PN200+Default Pro A Condition
Position	1,1
Load tips	
Position	7,1
Down	4225
Execute Protocol	c:\PN200+Default Pro A Capture
Position	4,1
Down	4280
Execute Protocol	c:\PN200+Default Pro A Wash1
Position	5,1
Down	4280
Execute Protocol	c:\PN200+Default Pro A Wash2
Position	8,1
Down	4225
Execute Protocol	c:\PN200+Default Pro A Elution
Position	1,1
Down	4100
Eject	
Eject	
Position	2,1
Load tips	
Execute Protocol	c:\PN200+ Condition 2
Position	6,1
Down	4070
Execute Protocol	c:\PN200+ Intake 5uL
Position	8,1
Down	4140
Execute Protocol	c:\PN200+ Expel 5uL
Position	2,1
Down	4100
Eject	
Home Auto Controller	

This method shows the automated movement of the 12 channel system for 1 row of samples. In detail, this can be broken down as follows:

## 1. Homing the system

Command	Parameters (if any)
Home Auto Controller	
Blowout	
Home Controller	
Execute Protocol	c:\PN200+Default Pro A Condition
Position/Row	1,1

**Home Auto Controller** moves the robotics tower to the Home position – full left position.

**Blowout** performs a complete blowout of the 12 channel system.

**Home Controller** performs a zeroing of the 12 channel system followed by a resetting of the system to zero.

**Execute** (c:\PN200+Default ProA Condition) executes a stored protocol which in this case is a conditioning step. The 12 channel system withdraws air into the syringes before the PhyTip columns are attached. It does this to create an air buffer in the system that will later be used to push liquid out of the PhyTip columns. The volume of air will vary depending on the specific set of protocols being used.

**Position 1, Row 1** moves the robotics tower over position number 1 and row 1 (The tray of PhyTip columns).

## 2. Moving the PhyTip Columns and Capturing the Sample

Load tips	
Position	7,1
Down	4225
Execute Protocol	c:\PN200+Default Pro A Capture

**Load tips** moves the 12 channel system down towards the tips in row 1 then moves the 12 channel system at an angle to load and secure all 12 PhyTip columns.

**Position 7, Row 1** automatically moves the robotics tower to the full extent (Position 0) and then moves it over to position 7, row 1 (the plate holding the sample to be purified).

**Down** moves the robotics head with PhyTip columns down into the sample solution in row 1 of position 7. This is a predefined position of 4225 (assumes use of PhyTip Plates with known dimensions) in readiness for the Capture step.

**Execute** (c:\PN200+Default ProA Capture) executes the stored protocol that implements the Capture process with a given number of cycles, flow rates, volumes and pauses e.g. in the standard method this will be a volume of 170  $\mu$ L for 2 cycles at a flow rate of 250  $\mu$ L/min with 10 second pauses at the end of each step. Also note that at the end of the capture step, the last cycle expels 172  $\mu$ L to make sure all of the sample solution is removed.

### 3. Wash and Elution Steps

Position	4,1
Down	4280
Execute Protocol	c:\PN200+Default Pro A Wash1
Position	5,1
Down	4280
Execute Protocol	c:\PN200+Default Pro A Wash2
Position	8,1
Down	4225
Execute Protocol	c:\PN200+Default Pro A Elution
Position	1,1
Down	4100
Eject	
Eject	

**Position 4, Row 1** moves the robotics tower to the full extend position (Position 0) and then moves it over to position 4, row 1 ready for the first wash step.

**Down 4280** moves the robotics tower down to predefined position of 4280 (assumes use of PhyTip Plates with known dimensions) ready for the wash step.

**Execute** (c:\PN200+Default Pro A Wash 1) implements the stored protocol for the first wash step. This protocol, like Capture, will control the flow rate, volume, pause and number of cycles involved in this step.

**Position 5, Row 1** moves the robotic tower up to the fully extended zero position and then moves it to position 5 and row 1 in readiness for the next wash step.

**Down 4280** moves the robotics tower down to predefined position of 4280 (assumes use of PhyTip Plates with known dimensions) ready for the next wash step.

**Execute** (c:\PN200+Default Pro A Wash 2) implements the stored protocol for the second wash step. This protocol, like Wash 1, will control the flow rate, volume, pause and number of cycles involved in this step.

**Position 8, Row 1** moves the robotics tower to the full extend position (Position 0) and then moves it over to position 8, row 1 ready for the elution step.

**Down** moves the robotics tower down to predefined position of 4225 (assumes use of PhyTip Plates with known dimensions) ready for the elution step.

**Execute** (c:\PN200+Default Pro A Elution) implements the stored protocol for the elution step. This protocol, like Capture, will control the flow rate, volume, pause and number of cycles involved in this step. Note at the end of this step, the last expel step is greater volume than the other expel steps in order to maximize the total volume of elution buffer returned to the elution plate.

**Position 1, Row 1** moves the robotics tower to the full extend position (Position 0) and then moves it over to position 1, row 1 ready for the ejection of the PhyTip columns.

**Down 4100** moves the robotics tower down to predefined position of 4100, ready to eject the PhyTip columns back into their original position in row 1.

**Eject** is a command to remove all of the now used PhyTip columns.

#### 4. Neutralization

Position	2,1
Load tips	
Execute Protocol	c:\PN200+ Condition 2
Position	6,1
Down	4070
Execute Protocol	c:\PN200+ Intake 5uL
Position	8,1
Down	4140
Execute Protocol	c:\PN200+ Expel 5uL
Position	2,1
Down	4100
Eject	
Home Auto Controller	

**Position 2, Row 1** moves the robotics tower over position number 2 and row 1 (The tray of Rainin LTS Tips).

**Load tips** moves the 12 channel head down to the pre assigned position to automatically load the tips onto the 12 channel system.

**Execute** (c:\PN200+ Condition 2) executes a stored protocol which in this case is a conditioning step. The 12 channel system withdraws air into the syringes before the LTS tips are attached. It does this to create an air buffer in the system that will later be used to push liquid out of the LTS tips. The volume of air will vary depending on the specific set of protocols being used.

**Position 6, Row 1** automatically moves the robotics tower to the full extent (Position 0) and then moves it over to position 6, (the Neutralization Buffer Reservoir).

**Down 4070** moves the robotics head with Tips down into the neutralization buffer. This is a predefined position of 4070 in readiness for removing the Neutralization buffer.

**Execute** (c:\PN200+Intake 5  $\mu$ L) executes the stored protocol that implements the removal of 5  $\mu$ L of neutralization buffer into the 12 Tips.

**Position 8, Row 1** automatically moves the robotics tower to the full extent (Position 0) and then moves it over to position 8, row 1 (the row that contains the purified protein).

**Down 4140** moves the robotics head with Tips with neutralization buffer down to the predefined position of 4140 in readiness for adding the Neutralization buffer.

**Execute** (c:\PN200+Expel 5  $\mu$ L) executes the stored protocol that implements the expelling of 5  $\mu$ L of neutralization buffer into row 1.

**Position 2, Row 1** automatically moves the robotics tower to the full extent (Position 0) and then moves it over to position 2, row 1 (the original tray for the Rainin Tips).

**Down 4100** moves the robotics head with Tips down to a predefined position of 4100 in readiness for removing the Tips from the 12 channel system.

**Eject** removes the tips into their original position.

**Home Auto Controller** takes the robotics tower back to its safe starting position and completes the purification of 1 row of samples.

If using this program as a template, just cut and paste the entire page and replace row 1 with row 2, etc.

### Example 3

#### *Optimization Method.*

The MEA can be used as a very rapid optimization platform to assess the optimum conditions for purifying protein samples. Below is an experimental design format as an example, with Ni-NTA resin:

#### **Capture**

Prepare the standard 96-well sample plate and program the MEA as follows:  
For the Capture protocol have Rows A, C, E & G have 2 Capture cycles, and Rows B, D, F & H have 4 Capture cycles.

		Column											
Row		1	2	3	4	5	6	7	8	9	10	11	12
A		2	2	2	2	2	2	2	2	2	2	2	2
B		4	4	4	4	4	4	4	4	4	4	4	4
C		2	2	2	2	2	2	2	2	2	2	2	2
D		4	4	4	4	4	4	4	4	4	4	4	4
E		2	2	2	2	2	2	2	2	2	2	2	2
F		4	4	4	4	4	4	4	4	4	4	4	4
G		2	2	2	2	2	2	2	2	2	2	2	2
H		4	4	4	4	4	4	4	4	4	4	4	4

#### **Purify**

The wash process for purification can be optimized when using Ni-NTA resin by varying the concentration of the Imidazole in the wash buffer.

In this example, rows A&B have 0 mM Imidazole in the wash buffer, rows C&D have 5 mM, E&F have 10 mM, and G&H have 20 mM Imidazole.

		Column											
Row		1	2	3	4	5	6	7	8	9	10	11	12
A		0	0	0	0	0	0	0	0	0	0	0	0
B		0	0	0	0	0	0	0	0	0	0	0	0
C		5	5	5	5	5	5	5	5	5	5	5	5
D		5	5	5	5	5	5	5	5	5	5	5	5
E		10	10	10	10	10	10	10	10	10	10	10	10
F		10	10	10	10	10	10	10	10	10	10	10	10
G		20	20	20	20	20	20	20	20	20	20	20	20
H		20	20	20	20	20	20	20	20	20	20	20	20

## Purification Step 2

This can be a repeat of Wash 1, using the exact same methodology.

## Enrich

The final elution step can now be compared using two concentrations of Imidazole to remove the his-tagged protein, e.g. columns 1-6 are eluted with a 150 mM Imidazole buffer and columns 7-12 are eluted with a 250 mM Imidazole buffer.

		Column											
Row		1	2	3	4	5	6	7	8	9	10	11	12
A		150	150	150	150	150	150	250	250	250	250	250	250
B		150	150	150	150	150	150	250	250	250	250	250	250
C		150	150	150	150	150	150	250	250	250	250	250	250
D		150	150	150	150	150	150	250	250	250	250	250	250
E		150	150	150	150	150	150	250	250	250	250	250	250
F		150	150	150	150	150	150	250	250	250	250	250	250
G		150	150	150	150	150	150	250	250	250	250	250	250
H		150	150	150	150	150	150	250	250	250	250	250	250

This one experiment, which can be run in less than 2 hours, allows for a statistically significant number of comparisons of capture conditions, wash conditions and elution conditions.

## **Standard Methods:**

The following standard methods have been built into the standard operating software:

### **For 200+ PhyTip columns and 200 + 12 channel system**

*Without the Cooling Accessory:* Position 7 is for the sample and position 8 is for elution.

1. Protein A with included neutralization step
2. Protein G with included neutralization step
3. IMAC
4. Glutathione

*With the Cooling Accessory:* Position 7 is for the sample and position 8 is for elution.

1. Protein A with included neutralization step
2. Protein G with included neutralization step
3. IMAC
4. Glutathione

### **For 1000+ PhyTip columns and 1000 + 12 channel system**

*Without the Cooling Accessory:* Position 7 is for the sample and position 8 is for elution.

1. Protein A with included neutralization step
2. Protein G with included neutralization step
3. IMAC
4. Glutathione

*With the Cooling Accessory:* Position 7 is for the sample and position 8 is for elution.

1. Protein A with included neutralization step
2. Protein G with included neutralization step
3. IMAC
4. Glutathione

## Full x-axis and z-axis positions.

For the 200+ 12 channel system:

		x-axis #
<b>Position 1</b>	Row 1	0
<i>PhyTip columns</i>	Row 2	567
	Row 3	1134
	Row 4	1701
	Row 5	2268
	Row 6	2834
	Row 7	3401
	Row 8	3968

PhyTip columns z-axis:

Load = 4270. Eject = 4100

		x-axis #
<b>Position 2</b>	Row 1	5648
<i>PhyTip columns</i>	Row 2	6215
	Row 3	6782
	Row 4	7349
	Row 5	7916
	Row 6	8482
	Row 7	9049
	Row 8	9616

Rainin LTS Tips z-axis:

Load=4270. Eject = 4100

		x-axis #
<b>Position 3</b>	Row 1	11240
<i>Sample Plate</i>	Row 2	11807
<i>(PhyTip Plate)</i>	Row 3	12374
	Row 4	12941
	Row 5	13508
	Row 6	14074
	Row 7	14641
	Row 8	15208

PhyTip columns z-axis:

Sample Plate= 4280

		x-axis #
<b>Position 4</b>	Row 1	16723
<i>Wash 1</i>	Row 2	17290
<i>(PhyTip Plate)</i>	Row 3	17857
	Row 4	18424
	Row 5	18991
	Row 6	19558
	Row 7	20124
	Row 8	20691

PhyTip columns z-axis:

Wash Plate= 4280

		x-axis #
<b>Position 5</b>	Row 1	22206
<i>Wash 2</i>	Row 2	22773
<i>(PhyTip Plate)</i>	Row 3	23340
	Row 4	23907
	Row 5	24474
	Row 6	25041
	Row 7	25608
	Row 8	26175

PhyTip columns z-axis:

Wash Plate= 4280

		x-axis #
<b>Position 6</b>		
<i>Neutralization</i>		
<i>Reservoir</i>		27674

Rainin LTS Tips z-axis:

Neutralization= 4070

		x-axis #
<b>Position 7</b>	Row 1	29152
<i>Elution</i>	Row 2	29719
<i>(PhyTip Plate)</i>	Row 3	30286
	Row 4	30853
	Row 5	31420
	Row 6	31986
	Row 7	32553
	Row 8	33120

With Cooling System z-axis

PhyTip Elution = 4225

Rainin LTS = 4140

		x-axis #
<b>Position 8</b>	Row 1	34640
	Row 2	35207
	Row 3	35774
	Row 4	36341
	Row 5	36908
	Row 6	37474
	Row 7	38041
	Row 8	38608

With Cooling System z-axis

PhyTip Elution = 4225

Rainin LTS = 4140

Without Cooling System z-axis

PhyTip Elution = 4280

Rainin LTS = 4195

Without Cooling System z-axis

PhyTip Elution = 4280

Rainin LTS = 4195

For the 1000+ 12 channel system:

		x-axis #
<b>Position 1</b>	Row 1	0
<i>PhyTip columns</i>	Row 2	567
	Row 3	1134
	Row 4	1701
	Row 5	2268
	Row 6	2834
	Row 7	3401
	Row 8	3968

PhyTip columns z-axis:  
Load = 3600. Eject = 3500

		x-axis #
<b>Position 3</b>	Row 1	11240
<i>Sample Plate</i>	Row 2	11807
<i>(PhyTip Deep Well)</i>	Row 3	12374
	Row 4	12941
	Row 5	13508
	Row 6	14074
	Row 7	14641
	Row 8	15208

PhyTip columns z-axis:  
Sample Plate= 2310

		x-axis #
<b>Position 5</b>	Row 1	22206
<i>Wash 2</i>	Row 2	22773
<i>(PhyTip Deep Well)</i>	Row 3	23340
	Row 4	23907
	Row 5	24474
	Row 6	25041
	Row 7	25608
	Row 8	26175

PhyTip columns z-axis:  
Wash Plate= 2310

		x-axis #
<b>Position 7</b>	Row 1	29152
<i>Elution</i>	Row 2	29719
<i>(PhyTip Plate)</i>	Row 3	30286
	Row 4	30853
	Row 5	31420
	Row 6	31986
	Row 7	32553
	Row 8	33120

With Cooling System z-axis  
PhyTip Elution = 2310  
Rainin LTS = 2120

Without Cooling System z-axis  
PhyTip Elution = 2365  
Rainin LTS = 2175

		x-axis #
<b>Position 2</b>	Row 1	5648
<i>PhyTip columns</i>	Row 2	6215
	Row 3	6782
	Row 4	7349
	Row 5	7916
	Row 6	8482
	Row 7	9049
	Row 8	9616

Rainin LTS Tips z-axis:  
Load=3600. Eject = 3500

		x-axis #
<b>Position 4</b>	Row 1	16723
<i>Wash 1</i>	Row 2	17290
<i>(PhyTip Deep Well)</i>	Row 3	17857
	Row 4	18424
	Row 5	18991
	Row 6	19558
	Row 7	20124
	Row 8	20691

PhyTip columns z-axis:  
Wash Plate= 2310

		x-axis #
<b>Position 6</b>		
<i>Neutralization</i>		
<i>Reservoir</i>		27674

Rainin LTS Tips z-axis:  
Neutralization= 2050

		x-axis #
<b>Position 8</b>	Row 1	34640
	Row 2	35207
	Row 3	35774
	Row 4	36341
	Row 5	36908
	Row 6	37474
	Row 7	38041
	Row 8	38608

With Cooling System z-axis  
PhyTip Elution = 2310  
Rainin LTS = 2120

Without Cooling System z-axis  
PhyTip Elution = 2365  
Rainin LTS = 2175

## Protocols

Protocols define the flow rates, volume and pauses associated with the process of Capture-Purify-Enrich. There are a series of standard protocols that have been stored in the software, e.g.:

200+ ProA Condition

200+ ProA Capture

200+ ProA Wash 1

200+ ProA Wash 2

200+ ProA Elute

200+ ProA Condition 2

200+ ProA Intake 5  $\mu$ L

200+ ProA Expel 5  $\mu$ L

(These protocols represent a series of steps that would be used for the 200+ PhyTip columns containing Protein A affinity resin; there are others for 200+ Protein G, IMAC and Glutathione plus 1000+ Protein A, Protein G, IMAC and Glutathione).

The *standard methods* have been written to incorporate a series of the *standard protocols* so that the process of Capture-Purify-Enrich can simply be carried out without the need for further programming.

## 200 + Protocol

As an example of these protocols when using the standard 200+ Protein A Method, there are several “Execute” steps that reference the following steps:

200+ Pro A Condition

Volume (mL)	Rate (mL/min)	Times to do	Delay (sec)
0.01	1	1	0

**Condition:** This conditions the 12 channel system to prepare for the complete process of moving liquids over the resin bed in the PhyTip columns. The final process of elution will often require only 10  $\mu$ L of elution buffer to be moved over the resin bed and then expelled. In order to be sure that all of this liquid can be blown out, a sufficient reservoir of air must be added to the 12 channel system before the PhyTip columns are attached. In this case 10  $\mu$ L of air is moved into the 12 channel system at a rate of 1 mL/minute.

200+ Pro A Capture

Volume (mL)	Rate (mL/min)	Times to do	Delay (sec)
0.17	0.25	1	10
-0.17	0.25	1	10
0.17	0.25	1	10
-0.17	0.25	1	10
0.17	0.25	1	10
-0.17	0.25	1	10
0.17	0.25	1	10
-0.172	0.25	1	10

**Capture:** Passes the sample over the resin bed enabling the Protein A affinity resin to bind the protein of interest. In this case 170  $\mu\text{L}$  of sample is moved into the PhyTip (Intake) and 170  $\mu\text{L}$  is moved out (Expelled). Intake can be seen as the positive number and expel the negative number, also note that all expel steps are colored red. Note that there are two expel steps. Therefore, two cycles of Capture and that at the end of the second cycle 172  $\mu\text{L}$  is expelled from the PhyTip column. This extra volume is part of the original 10  $\mu\text{L}$  Condition step and helps to remove as much sample solution as possible prior to the Wash steps.

The Flow Rate is 250  $\mu\text{L}/\text{min}$  (expressed as 0.25 mL/min) with a delay of 10 seconds at the end of each step. This delay step of 10 seconds is built into the system to allow for the full equilibration of all PhyTip columns in all positions.

To modify any of the parameters, click on the parameter line and an “Edit Protocol Step” screen will display. This has several boxes that can be modified (see Section B, Detailed software Information): *Volume* (sets the desired volume), *Volume Units*: (references the volume, with mL or  $\mu\text{L}$  available), *Direction* (Expel or Intake), *Volume Prior to this Step* (indicates last volume input), *How Many Times* (number of times to repeat sequentially this specific step), *Rate* (the flow rate), *Rate units* (shows mL/hr, mL/min,  $\mu\text{L}/\text{hr}$ ,  $\mu\text{L}/\text{min}$ ) and *Delay* (in seconds).

200+ Pro A Wash 1

Volume (mL)	Rate (mL/min)	Times to do	Delay (sec)
0.17	0.5	1	10
-0.17	0.5	1	10
0.17	0.5	1	10
-0.17	0.5	1	10

**Wash 1:** This is the first of two washing (Purification) steps, here two cycles of moving 170  $\mu\text{L}$  of Wash Buffer 1 is accomplished at a flow rate of 500  $\mu\text{L}/\text{min}$  (expressed as 0.5 mL/min) with a delay of 10 seconds at the end of each step. Note that on the final expel step, the volume is the same as all the intake steps. At this first step, there is no need to expel any further volume since the next step is another wash step.

200+ Pro A Wash 2

Volume (mL)	Rate (mL/min)	Times to do	Delay (sec)
0.18	0.5	1	10
-0.18	0.5	1	10
0.18	0.5	1	10
-0.182	0.5	1	10

**Wash 2:** This is the second wash step, again two cycles at the same flow rate as Wash Step 1. Now however, there is a larger volume of wash buffer (180 µL) and at the final expel step there is an extra 2 µL of volume expelled from the PhyTip column (-182 µL).

200+ Pro A Elute

Volume (mL)	Rate (mL/min)	Times to do	Delay (sec)
0.05	0.5	1	10
-0.05	0.5	1	10
0.05	0.5	1	10
-0.05	0.5	1	10
0.05	0.5	1	10
-0.05	0.5	1	10
0.05	0.5	1	10
-0.05	0.5	1	10
0.05	0.5	1	10
-0.056	0.5	1	10

**Elute:** The final step in the purification process uses 5 cycles of rapid flow elution over the resin bed. In this case, although only 10 µL is being used as the Elution volume, the protocol is stored as 50 µL. This difference between actual physical volume and programmed volume allows for all of the Elution buffer to pass over the resin bed, thus maximizing the total yield of purified protein that can be obtained from the column. It is essential to “overdrive” this step (50µL to pick up 10µL of liquid) to overcome the surface tensions and pressures that are caused by the initial insertion of the tip, and bed resistance in the tip. The required balance is to pull up enough to get the fluid to move, but not too much such that the volume of fluid pulls an air bubble.

As a programming guideline, if a higher volume for elution is needed, add 40 to the final volume number to obtain the programmed elution volume, e.g. if 30 µL is the final elution volume required, program 30 + 40 = 70 µL for the final elution volume. **This guideline is valid for elution volumes less than 75 µL. For elution volumes greater or equal to 75 µL, simply program that same volume.**

Again note that the final step expels 56 µL to maximize the volume of Elution Buffer returned (Note: Initial Conditioning step has Intake volume of 10 µL. Capture has excess expel of 2 µL; Wash 2 has excess expel of 2 µL; and the final Elution step has an excess expel of 6 µL, thus balancing the Intake and Expel for the entire process).

200+ ProA Condition 2

Volume (mL)	Rate (mL/min)	Times to do	Delay (sec)
0.02	1	1	0

**Conditioning 2:** this step conditions the 12 channel system prior to adding the Rainin LTS Tips for the Neutralization Step. Air (20  $\mu$ L) is taken in at a flow rate of 1 mL/min.

200+ ProA Intake 5  $\mu$ L

Volume (mL)	Rate (mL/min)	Times to do	Delay (sec)
0.005	6	1	2

**Intake 5  $\mu$ L:** This step pulls 5  $\mu$ L of neutralization buffer into the Rainin LTS Tips at a flow rate of 6 mL/min.

200+ ProA Expel 5  $\mu$ L

Volume (mL)	Rate (mL/min)	Times to do	Delay (sec)
0.025	2	1	2

**Expel 5  $\mu$ L:** This step expels the 5  $\mu$ L of Neutralization Buffer from the Rainin LTS Tips into the specified row at a flow rate of 2 mL/min. The step requires 25  $\mu$ L of expel, the first 20  $\mu$ L is air and the remaining 5 is the neutralization buffer. Note the negative number indicating an expel step.

## 1000+ Protocol

Below is an example of the protocols for a 1000+ PhyTip column system using IMAC as the affinity resin:

1000+ IMAC Condition

Volume (mL)	Rate (mL/min)	Times to do	Delay (sec)
0.09	3	1	0

1000+ IMAC Capture

Volume (mL)	Rate (mL/min)	Times to do	Delay (sec)
0.45	0.25	1	20
-0.45	0.25	1	20
0.45	0.25	1	20
-0.45	0.25	1	20
0.45	0.25	1	20
-0.45	0.25	1	20
0.45	0.25	1	20
-0.47	0.25	1	20

1000+ IMAC Wash 1

Volume (mL)	Rate (mL/min)	Times to do	Delay (sec)
0.9	0.5	1	20
-0.9	0.5	1	20
0.9	0.5	1	20
-0.9	0.5	1	20

1000+ IMAC Wash 2

Volume (mL)	Rate (mL/min)	Times to do	Delay (sec)
0.5	0.5	1	20
-0.5	0.5	1	20
0.5	0.5	1	20
-0.52	0.5	1	20

1000+ IMAC Elute

Volume (mL)	Rate (mL/min)	Times to do	Delay (sec)
0.25	0.5	1	20
-0.25	0.5	1	20
0.25	0.5	1	20
-0.25	0.5	1	20
0.25	0.5	1	20
-0.25	0.5	1	20
0.25	0.5	1	20
-0.25	0.5	1	20
0.25	0.5	1	20
-0.3	0.5	1	20

Note: The initial conditioning step of 90  $\mu$ L and the subsequent combination of excess expels total 90  $\mu$ L. Also the final elution volume of 20  $\mu$ L requires an intake/expel of 250  $\mu$ L.

As a programming guideline, if a higher volume for elution is needed, add 230 to the final volume number to obtain the programmed elution volume, e.g. if 30  $\mu$ L is the final elution volume required, program  $30 + 230 = 260$   $\mu$ L for the final elution volume. **This guideline is valid for elution volumes less than 250  $\mu$ L. For elution volumes greater or equal to 250  $\mu$ L, simply program that same volume.**

## Manual methods programming

The MEA software not only includes standard Methods and Protocols that can be simply modified, but also a feature that allows the user to calculate and program specific positions on the deck. The standard windows screen shows five basic headings:

File – Protocol – Method – Options – Help.

**File** is the standard New/Open/Save/Save as found in most Windows applications.

**Protocol** contains Execute/Edit. Execute begins the loaded Protocol and Edit allows the user to Edit the loaded Protocol.

**Method** contains Execute/Verify/Edit. Execute begins the loaded Method, Verify checks that the Method can be operated and Edit allows the user to Edit the loaded Method.

**Options** contain Communications/Syringe. Communications sets the COM port for the system and Syringe verifies the 12 channel system type (200+ or 1000+).

**Help** shows the About information regarding the system.

To access the ability to specify x-z positioning of the 12 channel system press Shift – Alt, this will add two new functions to the Options field:

**Simulation Mode:** allows the user to run a method on the software only, disconnecting the functions of the MEA. This can be used to verify/check that all of the software programming is compatible without running the MEA.

**Terminal:** allows the user to move the 12 channel system, via a series of typed commands to any position on the deck. This function allows the user to verify specific positions on the MEA and to design new methods that are different to the stored methods.

Specific position commands can be entered in the window and once the “send” command is hit, the robotics tower with 12 channel system will move to that specific position.

Commands are entered in the window to the right of the “send” command button. Once the command has been entered, press ‘send.’ This will initiate the command and in the screen box above this window there will be a confirmation that the command has been implemented e.g. Response OK.

## *Commands*

**Home:** moves the robotics tower and 12 channel system back to starting home position (x= 0 and z=0)

**Down #:** (where # = any number between 0 and 4275) moves the 12 channel system down to a specific point. It is recommended to place two Tips that will be used in the process on positions 1 and 12 of the 12 channel system in order to position the base of the tip correctly. Note that when the 12 channel system is in the full up position (0), then to move the system down, just type in the desired position number e.g. 2500. However, if the 12 channel system is not at the zero position e.g. 2500 and you want to move the system down a further 100 stepper motor points, type in Down 100 and this moves the system to position 2600.

**Up:** moves the 12 channel system back to zero position, fully up.

**X pos - #:** (where # = any number between 0 and 39,100) moves the robotics tower and 12 channel system to a specific point on the x- axis along the deck. It is recommended to place two Tips that will be used in the process on positions 1 and 12 of the 12 channel system in order to position the tip correctly.

# Section B

## 1. Introduction

PhyNexus auto controller software is a graphical user interface application which is intended to allow the user to perform the following tasks:

1. Open, verify, and execute files which perform intake and expel operations.
2. Edit and save protocol files.
3. Open, verify, and execute method files which execute protocol files and other commands, such as instructions to the auto controller.
4. Edit and save method files.
5. Perform manual intake, expel, and blowout / home operations.

The auto controller software is intended for use on 32-bit Microsoft Windows platforms (Windows '95, '98, Me, NT, 2000, and XP) and communicates with the Rainin automated pipettor / controller via the Auto Controller.

### 1.1 Conventions

The auto controller software will afterwards be referred to as the *app*.

Menu and submenu options will be referred to using the -> notation. For example, the Open option under the File menu is referred to as File->Open.

### 1.2 Definitions

A protocol is a series of steps which command the controller to intake or expel liquid. Each step of a protocol specifies a volume (positive to intake, negative to expel), a rate, and a number of times to repeat. An optional post-step delay may also be specified.

A method is a series of steps which group protocol(s) together and also provide some user-interface functionality. The following actions can be steps of a method:

1. Execute (specified by filename).
2. Audible alert (beep).
3. Pause (display message, and wait for user input to either continue method or abort).
4. Blow out contents.
5. Reset to home position.
6. Comment (solely for documentation of the method file – not seen by the operator).
7. Auto controller instructions:

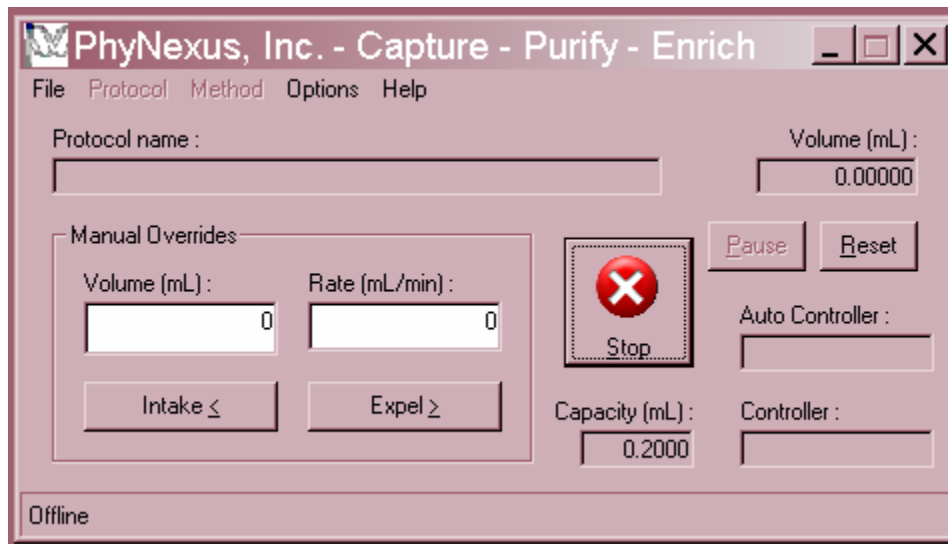
- a. Reset all axes to home position.
- b. Position above tray at specified step.
- c. Move controller down towards tray by specified number of steps.
- d. Load tips onto controller.
- e. Move controller to retracted position (performed automatically by tray command [b]).
- f. Remove tips from controller.
- g. Turn external controller on / off.

### 1.3 Compatibility

The auto controller (MEA system) is assumed to be connected in between the host computer and the controller (the 12 channel system). No other system or options are compatible with the MEA system.

## 2. How to open and execute a protocol

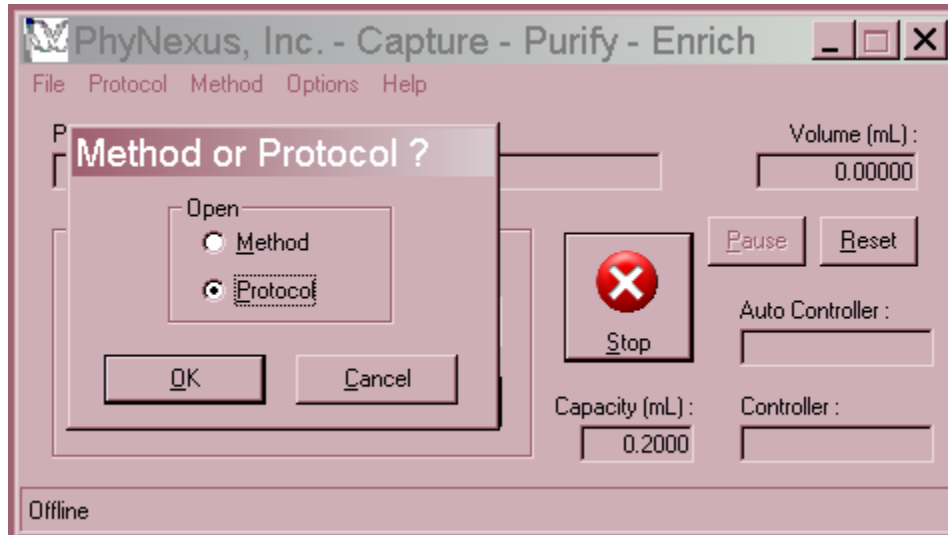
The following window appears when the **app** first runs:



The *status bar* (lower portion of the screen) shows the **app** status at any time. Initially the **app** is not communicating with the controller, so the status bar shows the message "Offline" when it starts.

## 2.1 Open protocol file

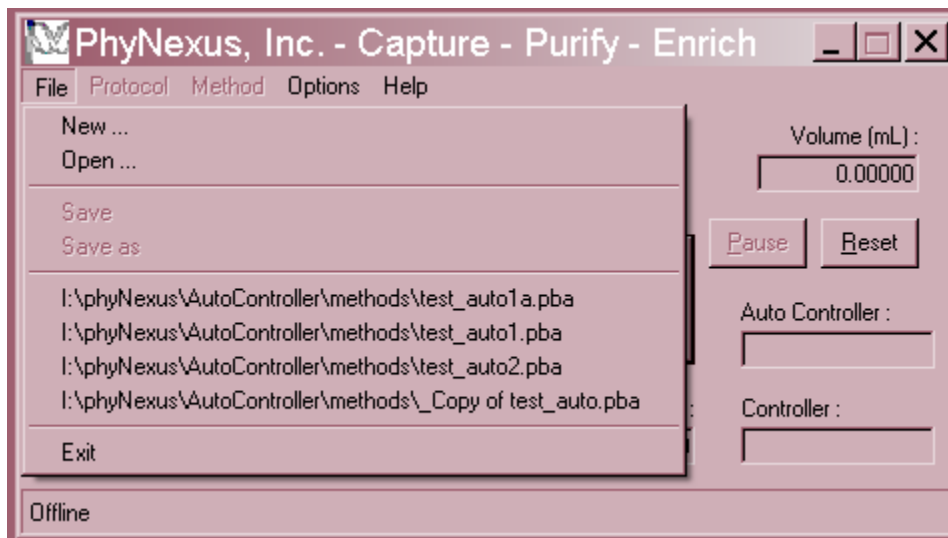
Use the File->Open menu option and select Protocol to browse for a protocol file, as shown below:



Protocol files are assumed to have a .psc file extension. If the protocol file was opened successfully, then it may be verified, executed and/or edited.

If the **app** displays an error message when you try to open the protocol file, then it may have been corrupted, e.g. via disk error or hand-editing, or the file selected is simply not a protocol.

The last four successfully-opened files are added to an **MRU** (most-recently-used) list in the file menu, as shown below:



When the filename is selected from the MRU, the **app** attempts to open the file. If the file cannot be opened or no longer exists, then it is removed from the MRU.

## 2.2 Execute

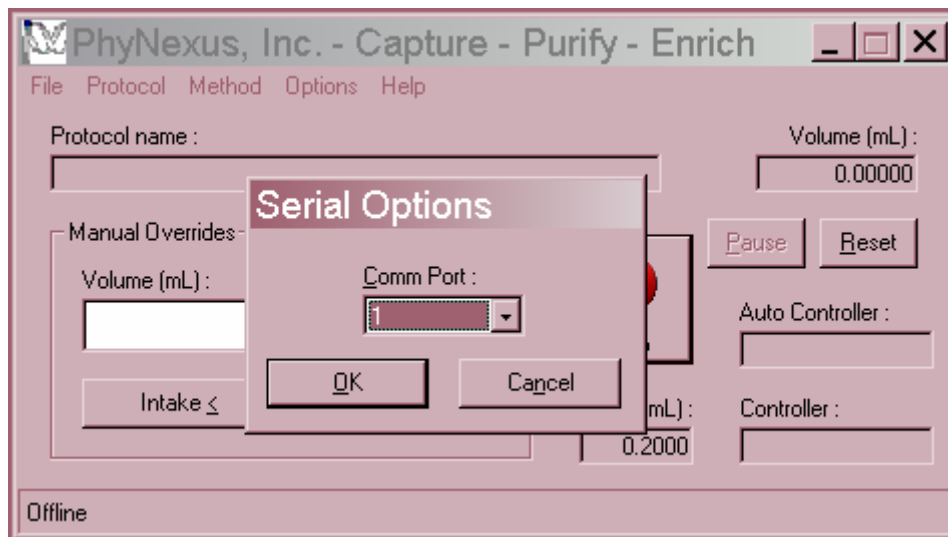
Before executing the protocol, make sure of the following:

1. All PhyTip columns, sample plates, wash buffers, etc., are in place.
2. **App's** serial parameters are set to match those of the controller.
3. The capacity of the PhyTip columns matches the volume required to be processed.

After these items have been checked, the protocol can be executed.

### 2.2.1 Check serial port

The **app's** serial port setting may be adjusted by selecting the Options->Serial menu, which pops up a dialog as follows:



Make sure that the port setting corresponds to the one on the PC that is connected to the auto controller. Initially, the **app** uses port 1, but remembers the last setting used (See *registry settings* below).

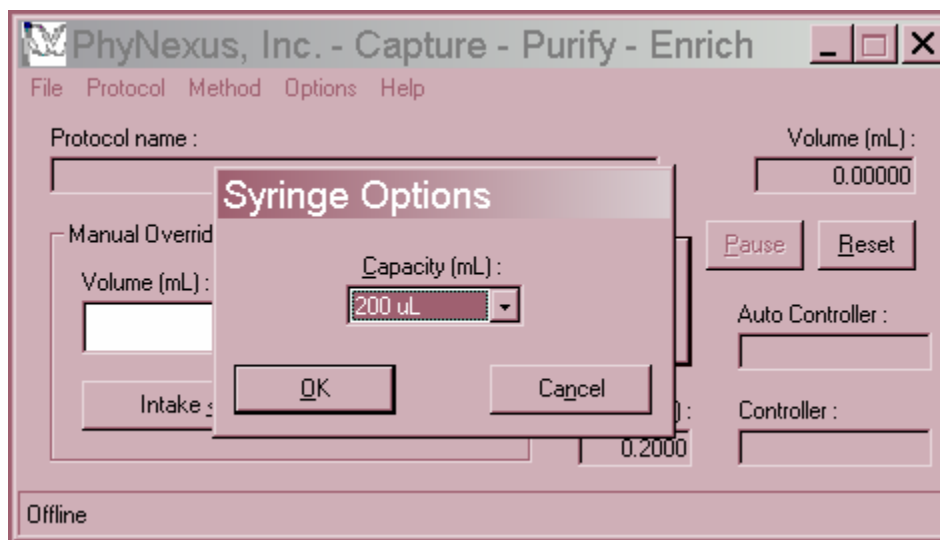
#### *Note:*

- Settings for the comm port can only be changed when the **app** is not communicating with the auto controller (not executing a protocol, method or

manual override command). Pressing 'Stop' terminates communication, and allows these settings to be changed (see below).

### 2.2.2 Check capacity setting

The capacity setting may be adjusted via the Options->Syringe menu, which pops up the following dialog:



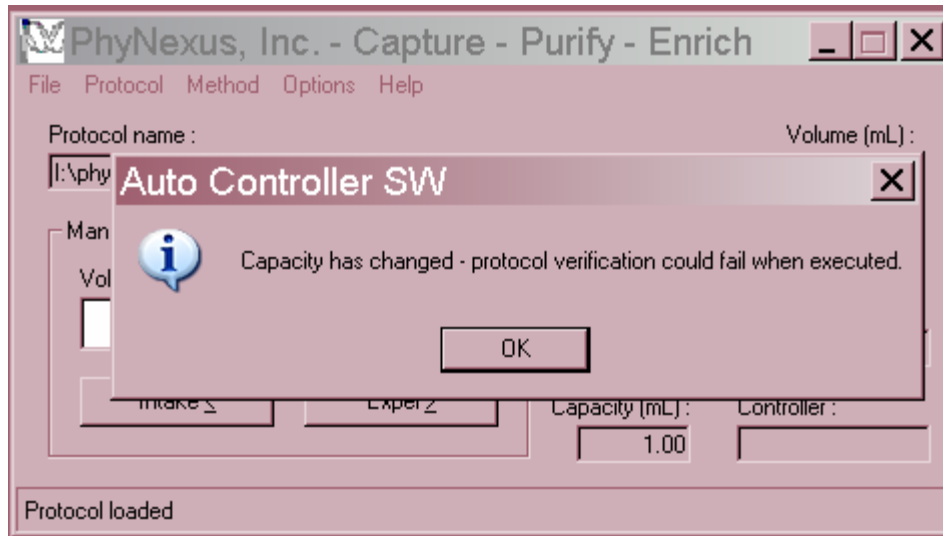
The setting shown above is for the 200  $\mu\text{L}$  capacity. The **app** is initially set up for 200  $\mu\text{L}$ , but remembers the last settings used (See *Registry Settings* below).

Also note that the current capacity is displayed on the main window of the **app**.

The capacity setting can only be changed when the **app** is not communicating with the auto controller (not executing a protocol, method or manual override command). Pressing 'Stop' also stops communication so that these settings can be changed.

#### 2.2.2.1 Capacity and verification

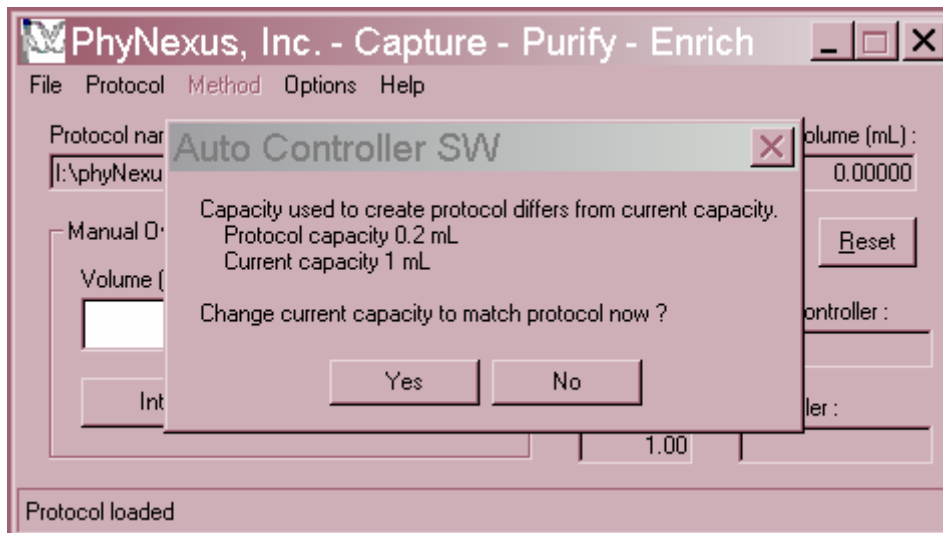
If the capacity setting is changed while a protocol is open, the user is prompted as follows:



This message indicates that the protocol may not execute properly with the new capacity setting.

### 2.2.2.2 Capacity and protocols

If a protocol specifies a different capacity than the current **app** settings, the user is prompted as follows when the protocol file is opened:



Answer “yes” to set the **app**’s capacity setting to match the protocol, or “no” to do so later.

**Note:** Answering “no” may prevent the protocol from executing properly.

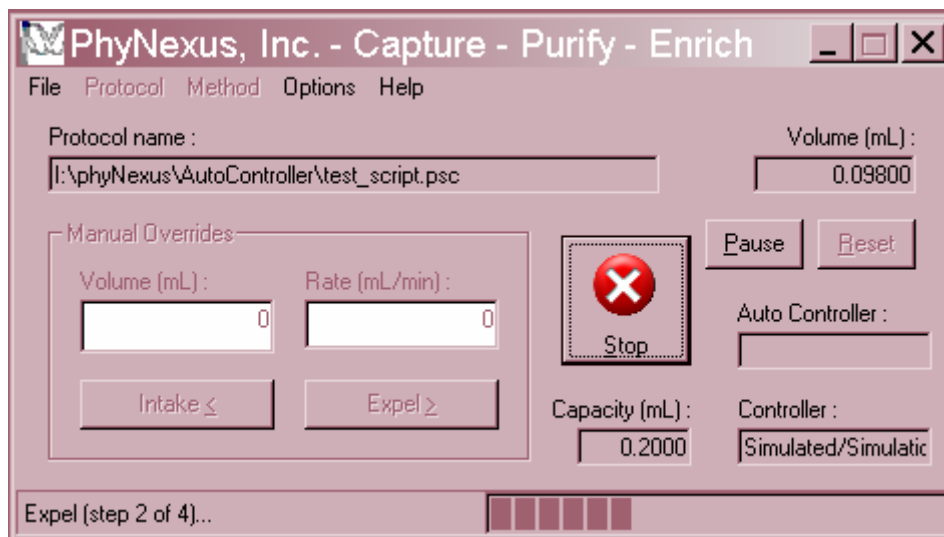
### 2.2.3 Start protocol execution

Once the protocol has been opened, and the controller has been prepared for protocol execution, select the Protocol->Execute option. The **app** will establish communication with the controller (go online), get the controller's software version and capacity, disable the controller's keyboard / speaker / LCD display, and get the current volume. The protocol will then be automatically **verified**, as noted in section 2.3 below, using the controller's current volume. Then the **app** will instruct the controller to execute the intake or expel steps contained in the protocol.

The **app** displays the following during protocol execution:

1. Status bar displays the protocol step and step type (intake/expel).
2. **Progress bar** (to the right of the status bar text) shows progress in total execution time, as calculated by the volume and rate of each step.
3. **Volume** displays the current volume in the syringe, as reported by the controller after the most-recently-executed step.
4. **Controller** shows the firmware version of the controller.
5. **Auto Controller** shows the firmware version of the auto controller.

An example is as follows:



When the protocol is completed, the status bar will display "*Protocol execution complete*" and the Volume indicator will show how much volume is in the syringe at the end of the protocol.

The protocol is automatically terminated if an error occurs during execution, and the error is shown in the status bar. See “Possible errors and corrective action” below (section 2.2.3.2).

### 2.2.3.1 Stopping protocol execution

If the protocol needs to be stopped before it’s complete, then press the Stop button. Pressing stop terminates the **app’s** execution of the protocol, but the controller will continue executing the last command issued by the controller until it has been completed.

### 2.2.3.2 Possible errors and corrective action

If a *Port in use* error results when the protocol is executed, then some other application is using the serial port selected (See section 2.2.1 above). Possibly a modem is plugged in and is being used, or the HyperTerminal app (which is found on most versions of Windows). To fix this, select a different serial port from the Serial Options dialog and try again, or find the application which is using the serial port, and shut it down.

The error *No response from controller (offline)* indicates that the serial port selected is not actually connected to the controller. Select a different serial port and try again. Similarly, *No response from auto controller (offline)* indicates that the serial port selected is not actually connected to the auto controller.

*Controller error* (command not echoed, bad checksum, invalid parameter or command) indicates a problem communicating with the controller over the serial cable, or an incompatible version of controller firmware. Check that the serial cable is in good condition, and that the controller firmware version is supported by the app (see section 1 above).

## 2.3 Verify protocol

Verification is performed automatically by the **app** prior to executing a protocol, but may also be invoked manually. In this process, the **app** checks that the following rules are followed during the protocol:

1. Volume never falls below zero.
2. Volume never rises above the capacity (see also section 2.2.2.1).
3. Flow rates are within limits for the capacity (see Appendix D).

When verification is performed automatically just prior to protocol execution, the initial volume is assumed to be that reported by the controller.

A protocol may be manually verified using the Protocol->Verify menu option. In this case, whatever current volume is presently reported by the controller is assumed to be the initial volume for verification purposes. If the **app** has not yet gone online, then the initial volume is assumed to be zero.

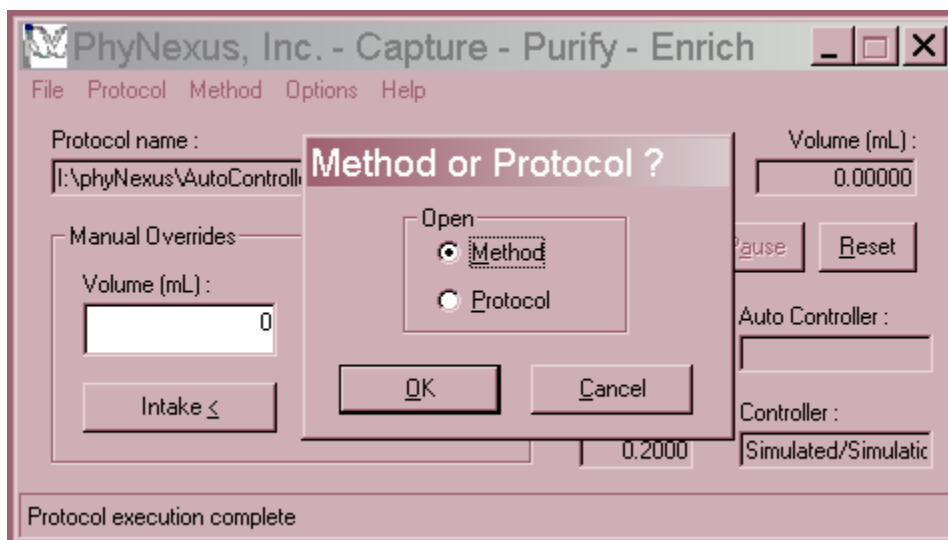
If protocol verification fails, then an error message appears showing the line in the protocol at which the error occurred.

### 3. How to open and execute a method

Handling of methods is similar to that of protocols.

#### 3.1 Open Method

Use the File->Open menu option and select Method to browse for a method file, as shown below :

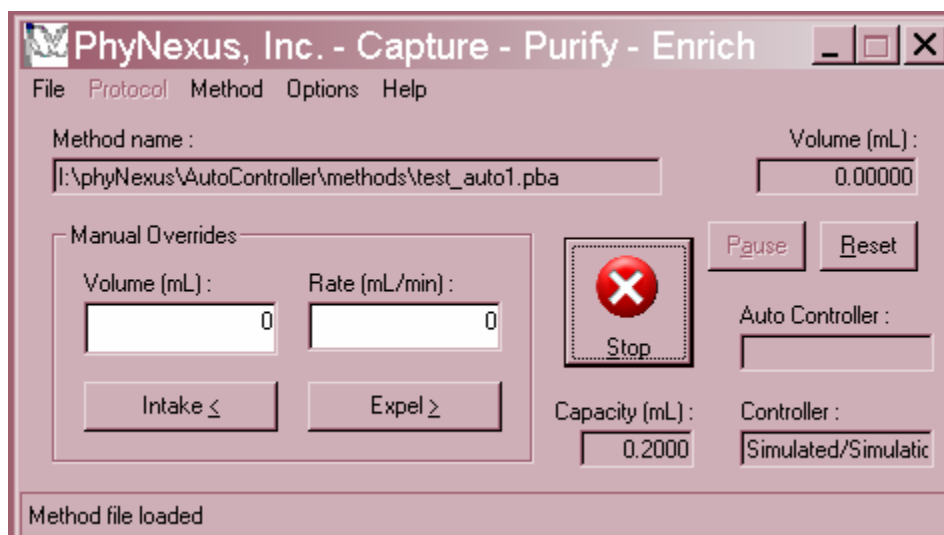


Method files are assumed to have a .pba file extension. If the method file was opened successfully, then it may be executed and/or edited.

If the **app** displays an error message when you try to open the method file, then it may have been corrupted, e.g. via disk error or hand-editing, or the file selected is simply not a method.

The **MRU** (most-recently-used) list in the file menu also handles method files.

After opening the method file, the display will indicate the currently-open method, as shown below:



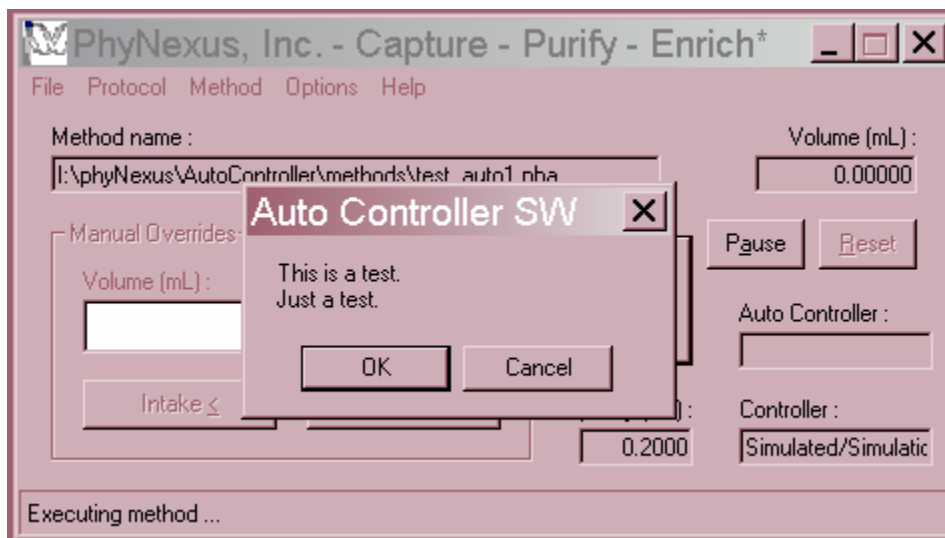
### 3.2 Execute method

Before executing the method, make sure of the following: (see section 2.2 above)

1. 12 channel system is correctly mounted on the controller.
2. **App's** serial parameters are set to match those of auto controller.
3. **App's** capacity setting matches the syringe capacity.

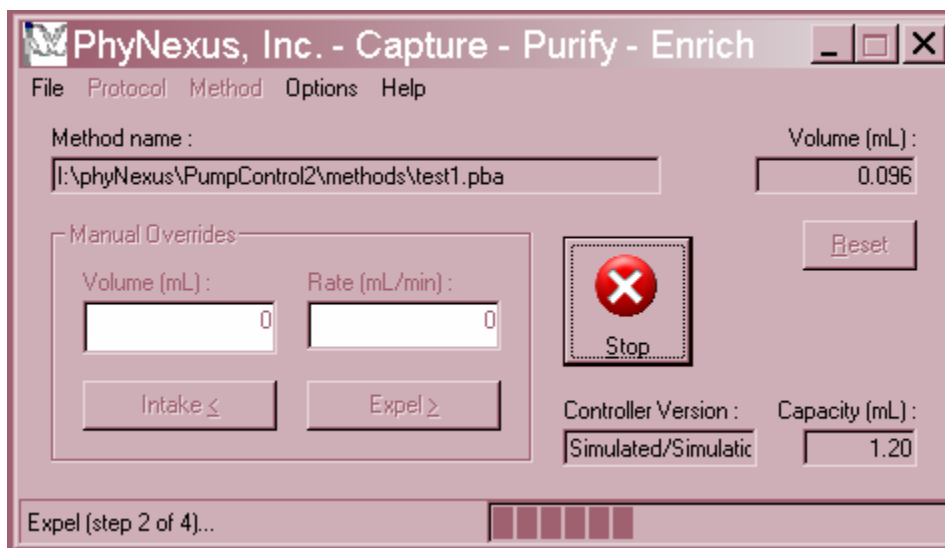
To execute the currently-open method, select the Method->Execute menu option. The **app** will then go online with the controller, much as it would while executing a protocol.

When a Pause method command is executed, the **app** displays the message specified in the command, and stops to wait for operator intervention, as shown below:



The operator can press **OK** to proceed with method execution, or **Cancel** to abort the method.

The appearance of the **app** while executing a protocol within a method is very similar to that when simply executing a protocol standalone – the only difference is that the method name is shown instead of the protocol name, as shown here:



Similarly, the Blowout and Home commands cause the status bar to be updated, and Volume to be zeroed.

See section 2.2.3 above for information on stopping execution, and possible errors that may occur, since this information applies equally to methods as well as protocols.

## 4. Manual Commands

The controller can be instructed to perform individual intake and expel steps when the **app** isn't running a protocol. The following procedure should be used:

1. Verify that the 12 channel system is mounted in the controller correctly, as above.
2. Verify that the **app's** serial parameters match those of the controller, as above.
3. Verify that the **app's** capacity setting matches that of the syringe, as above.
4. Enter the volume and rate in the text boxes under the *Manual Overrides* frame, and then press the *Intake* or *Expel* button.

This will cause the **app** to go online with the controller and execute the intake or expel step. Status will be shown in the status bar, and the Volume indicator will be updated when the operation is completed, if successful.

See section 2.2.3.2 above for troubleshooting.

### 4.1 Reset volume

The Reset button may be used to manually blow out the syringe contents, and reset its volume to the "home" position (zero).

### 4.2 Stopping manual commands

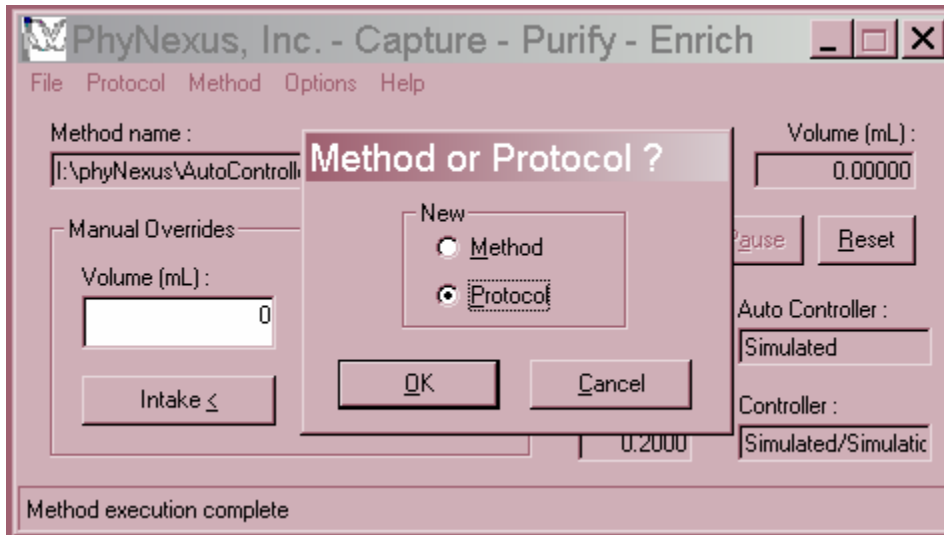
Manual intake/expel operations may be aborted using the Stop button. As with aborting protocol or method execution, pressing stop will only prevent the **app** from issuing further commands to the controller, but the controller will finish executing the last command it was given. When this last command is complete, the **app** will display the current volume as last reported by the controller.

**Note:** Communications and Syringe settings can only be changed when the **app** is not communicating with the controller (not executing a protocol, method or manual override command). Pressing 'Stop' terminates communication, and allows these settings to be changed.

## 5. Protocol editing

### 5.1 How to create a new protocol

To create a new protocol, select File->New and select Protocol, as shown below:



Then press OK, which will pop up the editor dialog:



Press the **Add** button to insert new steps. New steps may be edited after they have been added (see below).

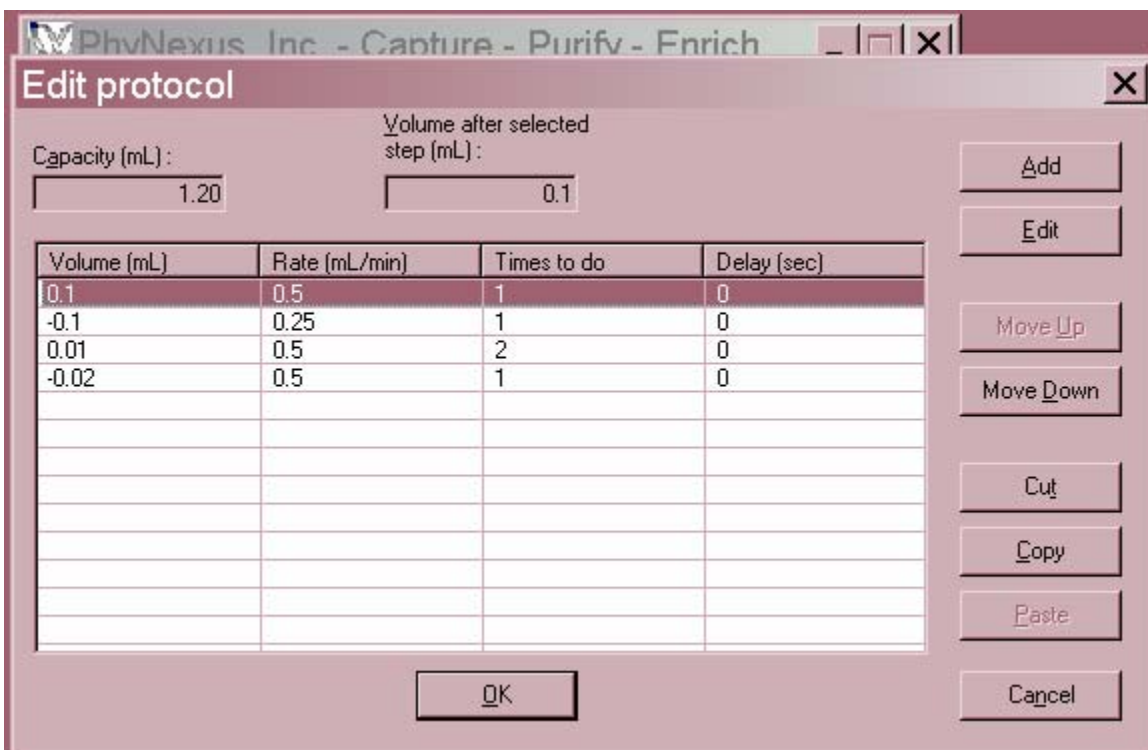
Steps may be re-ordered, either individually or in groups, by selecting the steps to be re-ordered, and using the **Move Up** and **Move Down** buttons.

A single step or a group of steps may be cut, copied, or re-inserted, using the **Cut**, **Copy**, and **Paste** buttons – these work as most Windows users would expect, and the actions of the buttons may also be invoked by pressing <Ctrl-X>, <Ctrl-C>, and <Ctrl-V>, respectively.

When a single step is selected (highlighted using the mouse or keyboard), the **Volume after selected step** indicator shows the volume in the syringe after the current step has been executed. The volume at the beginning of the protocol is assumed to be the value in the **Initial volume** edit box. The user may change the Initial volume, as desired.

## 5.2 How to edit an existing protocol

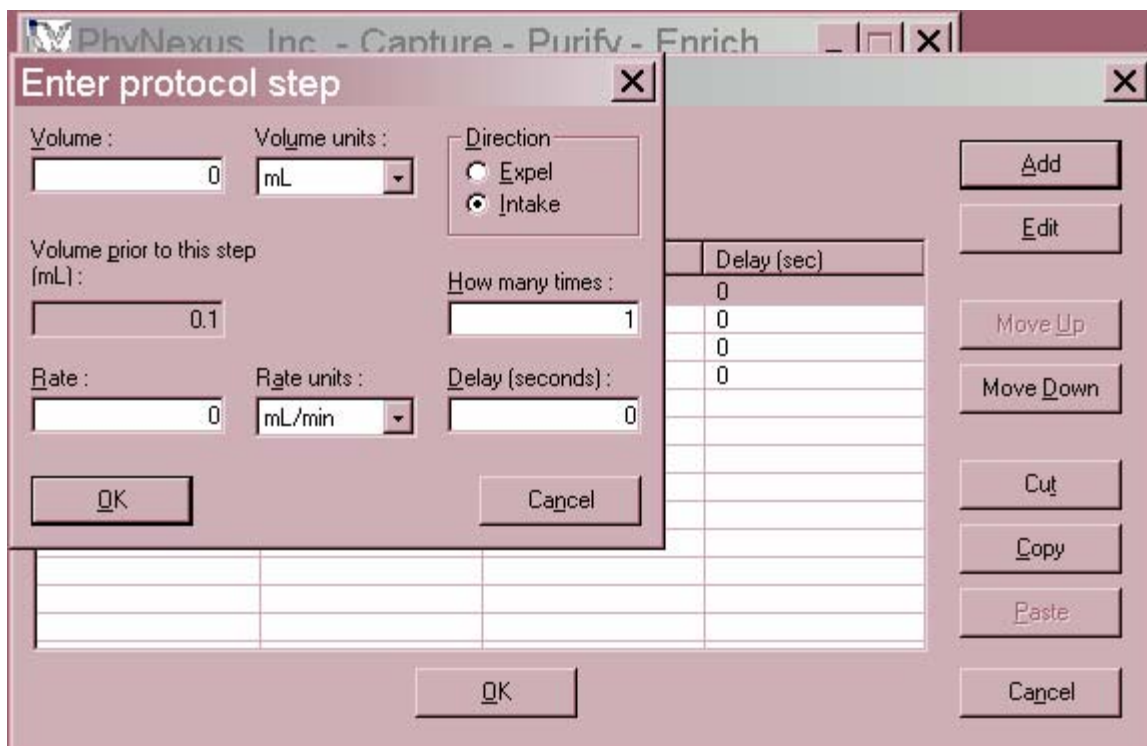
To edit a protocol, select Protocol->Edit to pop up the editor dialog, as described above. An example is shown below:



See below for a description of creating and editing steps.

### 5.3 Create protocol step

To create a new protocol step, press the Add button – this pops up the following dialog:



**Note:** If no protocol step was selected in the editor dialog, then the new step is placed at the end of the protocol. If any protocol steps were selected, then the new step is placed at the end of the selected steps. The *Move Up* or *Move Down* button may be used to move the new step, if desired.

Make sure to enter the correct values and units for the volume and rate, select the direction (intake/expel), the number of times to execute the step, and the post-step delay, if any. Delay times are rounded to the nearest millisecond. If a step is to be repeated, then the post-step delay is also repeated when executed.

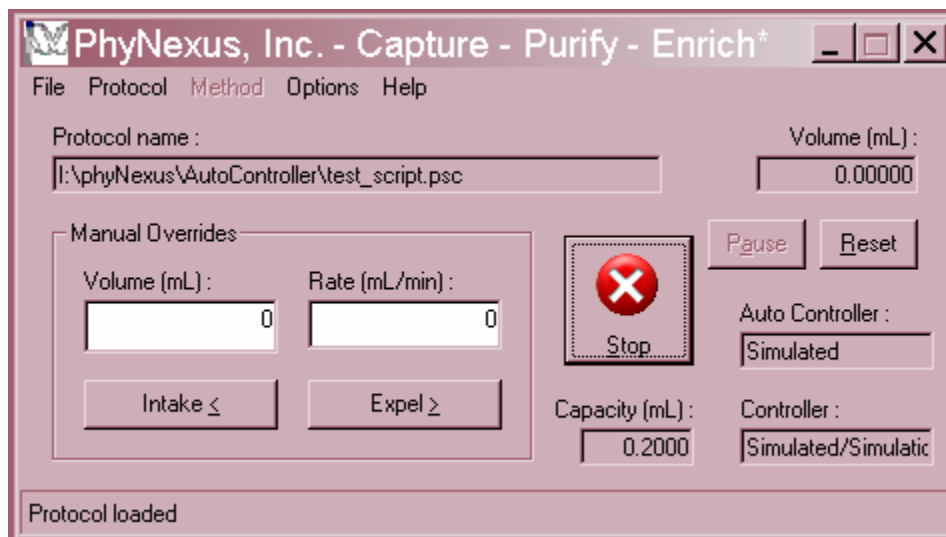
The ***Volume prior to this step*** indicator shows the calculated volume prior to the step being added. This dialog will not allow the volume to exceed the capacity, and also checks the rate against the capacity (see Appendix C).

## 5.4 Edit protocol step

To edit an existing protocol step, select the step in the editor window, and either double-click on the step, or press the **Edit** button. The protocol step dialog described above is then shown.

## 5.5 Save protocol to file

After a user presses **OK** in the editor window, the protocol is assumed to have been changed, and the app's caption indicates this via an asterisk next to the title, as shown below:



The asterisk means that there are changes to the protocol that have not been saved. Select File->Save to save the changes back to the previous filename, or File->Save As to save changes to a new filename.

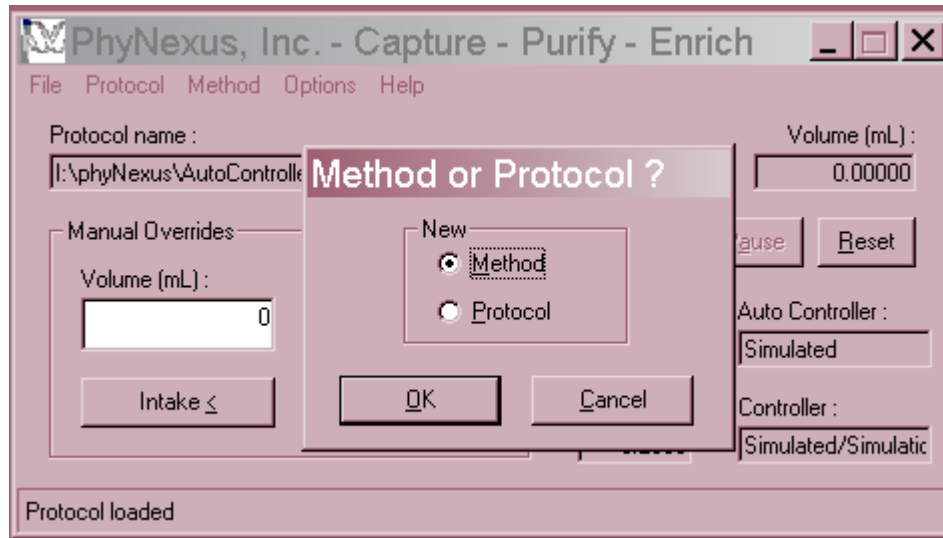
Note that the **app's** current capacity setting is also saved to the protocol file as a comment (see Appendix B).

## 6. Method editing

The steps involved in creating and editing a method are similar to that for a protocol.

### 6.1 How to create a new method

To create a new method, select File->New and select Method, as shown below:



Then press OK, which will pop up the editor dialog:





### 6.3 Create method step

To create a new method step, press the Add button.

**Note:** If no method step was selected in the editor window, then the new step is placed at the end of the method. If any steps were selected, then the new step is placed at the end of the selected steps. The *Move Up* or *Move Down* button may be used to move the new step, if desired.

Select the type of method step desired – this will cause the entry field(s) at the bottom of the dialog to change appearance as needed. Enter the parameter needed, as follows:

- Execute:* executes a stored protocol, e.g. Capture
- Audible Alert:* introduces an audible alert at the end of an action
- Home (controller):* homes the 12 channel system to a zero volume
- Pause for Operator:* builds in an automatic pause in the method
- Comment:* allows operator to build in an automatic comment or prompt between actions
- Blowout:* causes a 12 channel system to blow out any air from the syringe

The following method steps are auto controller instructions:

- Home Autocontroller:* returns the robotics tower and 12 channel system to registered Home position
- Down:* moves the robotics tower with 12 channel system down to designated step position (0 is the highest position above the deck and 4725 is the position closest to the deck).
- Up:* moves the robotics tower with 12 channel system up to the designated step position (0 is the highest position above the deck and 4725 is the position closest to the deck).
- External Pump:* operates an external pump (when fitted) via the rear serial port, allowing for the automated addition of reagents.
- Rotate:* rotates the vertical 12 channel system.
- Position:* moves the robotics tower with 12 channel system to the desired

x-axis location on the deck. There are a total of 39,100 positions along the x-axis.

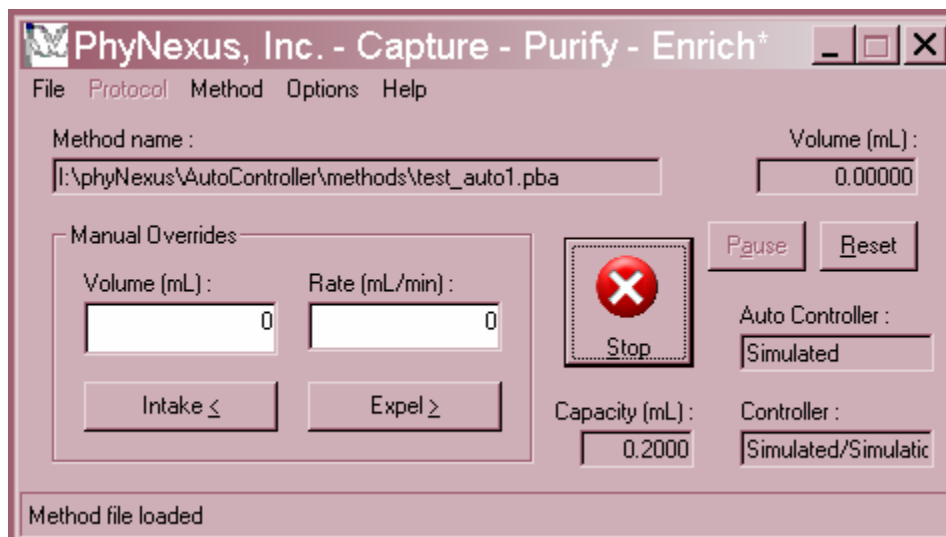
- Position/Row* moves the robotic tower with 12 channel system to the desired Position (1-8) and specific Row (1-8). This command assumes the use of plates that have 8 rows (e.g., standard 96 well plates, certain 48 well plates, etc.).
- Load:* loads a row of 12 PhyTip columns onto the 12 channel system
- Eject:* ejects all 12 PhyTip columns from the 12 channel system

## 6.4 Edit method step

To edit an existing method step, select the step in the editor window, and either double-click on the step, or press the **Edit** button. The method step dialog described above is then shown.

## 6.5 Save method to file

After a user presses **OK** in the editor window, the method is assumed to have been changed, and the **app's** caption indicates this via an asterisk next to the title, as shown below:



The asterisk means that there are changes to the method that have not been saved. Select File->Save to save the changes back to the previous filename, or File->Save As to save changes to a new filename.

## Appendix A : Protocol File Format

Protocols are stored as simple text files, and may be edited with Notepad, if desired (however, this may result in a protocol which cannot be opened and/or verified by the app). The **app** registers the file extension .psc for PhyNexus protocol files, so that clicking on a .psc file opens the protocol in the **app** (if the **app** isn't already running).

Each line of the file represents one step of the protocol, and is a comma-delimited list of the following fields :

1. Volume, in mL - positive for intake, negative for expel.
2. Rate, in mL per minute.
3. Number of times to do the step - optional.
4. Post-step delay, in seconds (**app** rounds this to nearest millisecond) – optional.

For example, the following line indicates an expel step of 20  $\mu$ L, with a rate of 5  $\mu$ L per minute, done once:

```
-0.02,0.005
```

The following line indicates an intake step of 1.5 mL, with a rate of 10 mL per minute, done three times :

```
1.5,10,3
```

This line indicates an intake step of 0.5 mL, with a rate of 2 mL per minute, with a post-step delay of 2 seconds :

```
0.5,2,1,2
```

The line below indicates an intake step of 1.5 milliliters, with a rate of 10 milliliters per minute, done three times, each with a post-step delay of 1.5 seconds (if a step is to be repeated, then the post-step delay is also repeated when executed):

```
1.5,10,3,1.5
```

Any line which begins with # is treated as a comment and ignored. The **app** stores only one comment line to a file, to indicate the capacity in milliliters. An example is as follows:

```
# vol =1
```

This indicates that when the file was saved, the **app**'s capacity setting was 1 mL.

## Appendix B: Method File Format

Methods are also stored as text files. The **app** registers the file extension .pba for PhyNexus method files. Each line of a method file represents a step of the method, and may take the following forms:

1. Execute specified protocol file :

```
execute <protocol file path>
```

For example, the following line indicates that the protocol file c:\protocols\test1.psc should be executed:

```
execute c:\protocols\test1.psc
```

Note that file paths may be absolute (as above) or relative. Relative paths are relative to the **app** executable file. For example, if the **app** resides at c:\phyNexus\pumpControl2.exe and the path protocols\test1.psc is specified, then the protocol file c:\phyNexus\protocols\test1.psc will be executed.

2. Audible alert:

```
alert
```

3. Pause, show message, and wait for operator intervention:

```
pause <message>
```

The message is shown with OK and Cancel buttons. OK continues the method, and Cancel terminates it. New lines may be embedded in the message by inserting the “\n” escape sequence. For example, the following line pops up a message box containing the lines “Hello” and “World”.

```
pause Hello\nWorld
```

4. Comment (ignored) :

```
#<comment>
```

Comments are ignored by the **app** when the method is executed, but can be useful for documenting methods.

5. Home:

```
home
```

6. Blowout:

## blowout

The following method steps are provided as instructions to the auto controller:

1. Home:

auto\_home

2. Move tray to specified step:

tray <step>

The step must be between 0 and 39100.

3. Move down towards tray specified number of steps:

down <number of steps>

The number of steps must be between than 0 and 4725.

4. Load tips:

load

5. Move up to retracted position:

up

6. Eject tips:

eject

7. Turn external pump on/off:

external\_pump <on | off>

### Appendix C: Flow rate limits

Flow rates are limited to a range which is dependent on capacity, as indicated by the following table:

Capacity	Minimum flow rate	Maximum flow rate
200 $\mu\text{L}$	20 $\mu\text{L}/\text{min}$	17.1 mL/min
1 mL	20 $\mu\text{L}/\text{min}$	85.7 mL/min
1.2 mL	20 $\mu\text{L}/\text{min}$	102.9 mL/min

## Appendix D: Flow rate implementation

The 12 channel controller does not actually implement continuous flow rates, but operates at ten discrete motor speeds, each of which is computed as follows:

$$\text{flow rate [mL/min]} = 60 \times \text{syringe capacity [mL]} / \text{full-scale duration [sec]}$$

where the motor speed determines full-scale duration, as per the following table: (only the 200 and 1000  $\mu\text{L}$  capacities are shown):

Motor speed	Full-scale duration [sec]	Flow rate for 200 $\mu\text{L}$ capacity [mL/min]	Flow rate for 1000 $\mu\text{L}$ capacity [mL/min]
1	5.3	2.3	13.6
2	4.5	2.7	16.0
3	3.8	3.2	18.9
4	3.2	3.8	22.5
5	2.8	4.3	25.7
6	2.4	5.0	30.0
7	1.9	6.3	37.9
8	1.5	8.0	48.0
9	1.0	12.0	72.0
10	0.7	17.1	102.9

The motor speed used to implement a particular flow rate is usually arrived at by interpolating between the motor speeds. For example, a flow rate of 13 mL/min for the 200  $\mu\text{L}$  capacity would use motor speed 9 because the midpoint between speeds 9 and 10 is 14.6 mL/min.

For flow rates below motor speed 1, the controller is commanded to perform *move / wait* cycles in which small intake or expel volume increments are made, interspersed by delays of several hundred to several thousand milliseconds. These cycles are used when the flow rate falls below 2, 10, or 12 mL/min, for the 200 and 1000  $\mu\text{L}$  capacities, respectively. Motor speed 1 is used to perform the cycles.

The volume increments for move/wait cycles are as follows:

Flow rate range [ $\mu\text{L}/\text{min}$ ]	Volume increment [ $\mu\text{L}$ ]
20 – 100	0.5 for 200 $\mu\text{L}$ capacity, 1 otherwise
100 – 200	1
200 – 400	2
400 – 800	5
800 – 1500	10
1500 – 2000	15
2000 – 4000	20
4000 – 8000	40
8000 – 12000	50

The move/wait cycle delay is computed as follows:

$$\text{delay [msec]} = 60000 \times \text{volume increment [mL]} / \text{desired flow rate [mL/min]}$$

The number of cycles is simply the desired volume divided by the volume increment, rounded up so that at least one cycle is performed.

For example, suppose a 100  $\mu\text{L}$  intake step needs to be made at 300  $\mu\text{L}/\text{min}$ . In this case 50 cycles will be performed, each withdrawing 2  $\mu\text{L}$  at motor speed 1, with a 400 msec wait until the next cycle.

## Appendix E: Registry Settings

The **app** uses the Windows registry for storing settings from session to session. The registry key for the app is as follows, and found under HKEY\_CURRENT\_USER, so that different users get different settings:

Software\VB and VBA Program Settings\PhyNexus PumpControl 2

Settings are stored under the following subkeys:

<b>Settings</b>	<b>Subkey name</b>
Serial communications	Serial
Syringe settings	Syringe
MRU list	MRU
Browse directory (protocol and method)	General

## Cleaning Instructions

To clean the exterior of this instrument, complete the following steps:

- a. Turn off and unplug the instrument
- b. Wipe the instrument exterior surfaces only, using a towel dampened with laboratory grade cleaning agent
- c. Repeat step b, using a towel dampened with clean water
- d. Repeat step c with a second water wash step
- e. Dry the instrument exterior using a dry towel

**Warning** – do not allow any liquid to enter the instrument, or come in contact with any electrical components. The instrument must be thoroughly dry before you reconnect the power or turn the instrument on.

## Safety Notices

**Warning** – When cleaning instrument, do not allow any liquid to enter the instrument, or come in contact with any electrical components. The instrument must be thoroughly dry before you reconnect the power or turn the instrument on.

### Caution

#### *Power Cord Set Requirements*

The power cord set supplied with your instrument meets the requirements of the country where you purchased the instrument. If you use the instrument in another country, you must use a power cord set that meets the requirements of that country.

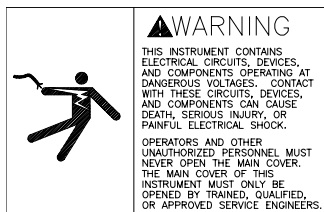
To reduce the risk of fire hazard and electrical shock, do not expose the unit to rain or humidity. To reduce the risk of electrical shock, do not open or remove the instrument casing. All maintenance is to be performed by an authorized PhyNexus service provider.

### Warning

This equipment is designed for connection to a grounded (earthed) outlet. The grounded type plug is an important safety feature. To reduce the risk of electrical shock or damage to the instrument, do not disable this feature.

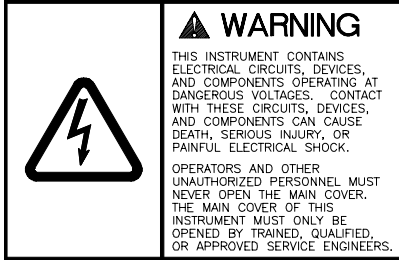
## Environmental

Operating Temperature:	4° to 30°C
Relative Humidity	0% to 95%
Altitude	0 to 10000 Ft



**⚠ Avertissement**

TOUT CONTACT AVEC LES HAUTES TENSIONS PEUT ENTRAINER LA MORT OU DES BLESSURES SÉVÈRES. CE PANNEAU NE DOIT ÊTRE ENLEVÉ QUE PAR UN RÉPARATEUR QUALIFIÉ.



**⚠ Avertissement**

TOUT CONTACT AVEC LES HAUTES TENSIONS PEUT ENTRAINER LA MORT OU DES BLESSURES SÉVÈRES. CE PANNEAU NE DOIT ÊTRE ENLEVÉ QUE PAR UN RÉPARATEUR QUALIFIÉ.



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**⚠ AVERTISSEMENT**  
 TOUT CONTACT AVEC LES HAUTES TENSIONS PEUT ENTRAINER LA MORT OU DES BLESSURES SÉVÈRES. CE

**⚠ WARNING**  
 HIGH LEAKAGE CURRENT -  
 ENSURE PROPER GROUNDING

**⚠ AVERTISSEMENT**  
 COURANT DE FUITE ÉLEVÉ — FOURNIR UNE MISE À LA TERRE EFFICACE.

**WARNING**  
 If the MEA is used in a manner not specified by PhyNexus Inc, the protection provided the equipment may be impaired.

MEA Power Supply Input: AC Voltage and Current 100-240V ~ 47-63Hz 1.9A	MEA Power Supply Output: DC Voltage and Current  ==== 24V 3.33A	Pipettor Power Supply Input: AC Voltage and Current 100-240V ~ 47-63Hz 0.5A Output: 5VDC, 3.0A
---	--	--

Chiller Power Supply Output: DC Voltage and Current  ==== 24V 5.4 A	Chiller Power Supply Input: AC Voltage and Current 100-240V ~ 47-63Hz 1.9A
--	---

**WARNING**  
 Pollution Degree - 1

## **MEA Personal Purification System Warranty**

The MEA Personal Purification System has a warranty period of one (1) year from the date of purchase. If the system should fail during the warranty period, please contact your local PhyNexus office for further instructions and information.

### **Limited Warranty and Limitation of Liability**

Seller (PhyNexus) warrants the merchandise to conform to specifications. Under no circumstances shall Seller be responsible for alleged nonconformities with respect to any merchandise which has been used for purposes or in any manner for which it was not intended, or any merchandise which has been customized or modified without Seller's prior written consent, or damaged or misused. As Buyer's exclusive remedy in the event of breach of warranty, Seller shall repair or replace, as its option, any nonconforming merchandise or parts thereof for a period of one (1) year after delivery. All claims must be made in writing to the Seller. Any claims not made within the period specified above shall be deemed waived and released.

THE PROVISIONS OF THE FOREGOING WARRANTIES ARE IN LIEU OF ANY OTHER WARRANTY, WHETHER EXPRESS OR IMPLIED, WRITTEN OR ORAL (INCLUDING ANY WARRANTY OF MERCHANTABILITY, FITNESS FOR A PARTICULAR PURPOSE, TITLE OR NON-INFRINGEMENT). SELLER'S LIABILITY ARISING OUT OF THE MANUFACTURE, SALE OR SUPPLYING OF A PRODUCT OR ITS USE OR ITS DISPOSITION, WHETHER BASED UPON WARRANTY CONTRACT, TORT OR OTHERWISE, SHALL NOT FOR ANY REASONS EXCEED THE AGGREGATE PURCHASE PRICE PAID BY BUYER FOR SUCH PRODUCT. IN NO EVENT SHALL SELLER BE LIABLE TO BUYER OR ANY OTHER PERSON OR ENTITY FOR SPECIAL, INCIDENTAL, CONSEQUENTIAL OR EXEMPLARY DAMAGES (INCLUDING, BUT NOT LIMITED TO, DAMAGES FOR LOSS OF PROFITS, LOSS OF DATA OR LOSS OF USE) ARISING OUT OF THE MANUFACTURE, SALE, SUPPLY, USE, MARKETING, RESALE OR OPERATION OF THE MERCHANDISE, EVEN IF SELLER HAS BEEN ADVISED OF THE POSSIBILITY OF SUCH DAMAGES OR LOSSES.

PhyNexus, Inc.

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