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| **C:\Users\hatfull lab\Desktop\2011 Teacher Workshop 6-9\Graphics and Logos\500px-UofPittsburgh_Seal.svg.png** | Phagehunting Program | C:\Users\hatfull lab\Desktop\2011 Teacher Workshop 6-9\Graphics and Logos\ForDarkBackgrounds-Logo.png |
| Macintosh HD:Users:Enoch:Desktop:New Protocols:Logos and Graphics:Workflow:PreparationPNG.png Macintosh HD:Users:Enoch:Desktop:New Protocols:Logos and Graphics:Workflow:IsolationPNG2.png Macintosh HD:Users:Enoch:Desktop:New Protocols:Logos and Graphics:Workflow:PurificationPNG.png C:\Users\Owner\Desktop\Protocol\picture\AmplificationWorkFlowgray.png Macintosh HD:Users:Enoch:Desktop:New Protocols:Logos and Graphics:Workflow:ExtractionPNG.png Macintosh HD:Users:Enoch:Desktop:New Protocols:Logos and Graphics:Workflow:CharacterizationPNG.png Macintosh HD:Users:Enoch:Desktop:New Protocols:Logos and Graphics:Workflow:SequencingPNG.png Macintosh HD:Users:Enoch:Desktop:New Protocols:Logos and Graphics:Workflow:AnnotationPNG.png Macintosh HD:Users:Enoch:Desktop:New Protocols:Logos and Graphics:Workflow:PhamerationPNG.png Macintosh HD:Users:Enoch:Desktop:New Protocols:Logos and Graphics:Workflow:FurtherDiscoveryPNG.png | | |

**BLASTable Mycobacteriophage Database**

**Purpose**: To have easy access to current known protein function through use of a DNA Master created protein database.

**Background**: When determining protein function, a variety of tools including BLASTP data, Phamerator maps, conserved domain identification, and HHpred are used. This is another tool to help. This database was created in DNA Master (procedure to follow) using 243 of the most recent mycobacteriophage genomes. This database was created December 2012.

**Overview:**  This protocol consists of three parts:

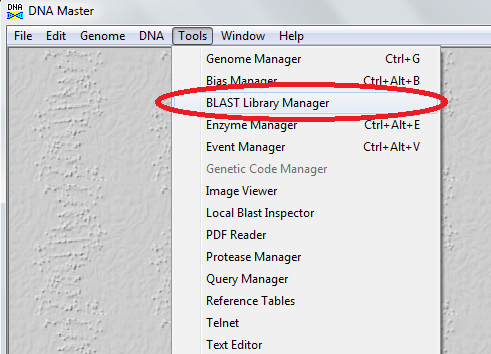
* Creation of local database (supplied, see addendum for procedure)
* Import of database
* Local Blast of database

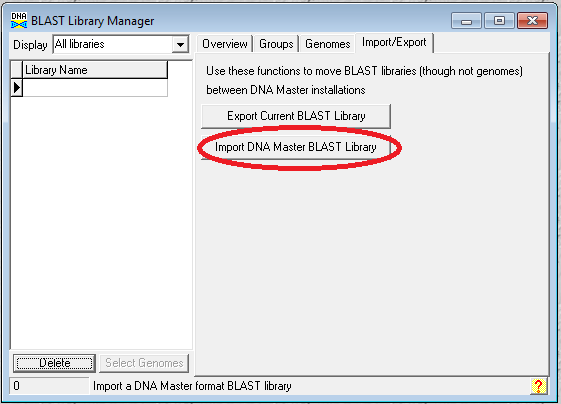
**Procedure:**

**Import of Database**

The library of completed mycobacteriophages with functions is currently a 3.41 MB file named Proteins\_243MP\_12.2012.dnamblib. This is a static file that will be periodically updated.

1. Get Proteins\_243MP\_12.2012.dnamblib from the PhagesDB website. Click on Resources🡪 Documents 🡪Annotation Resources. Save this file to a known location.
2. Go to Tools 🡪 BLAST Library Manager



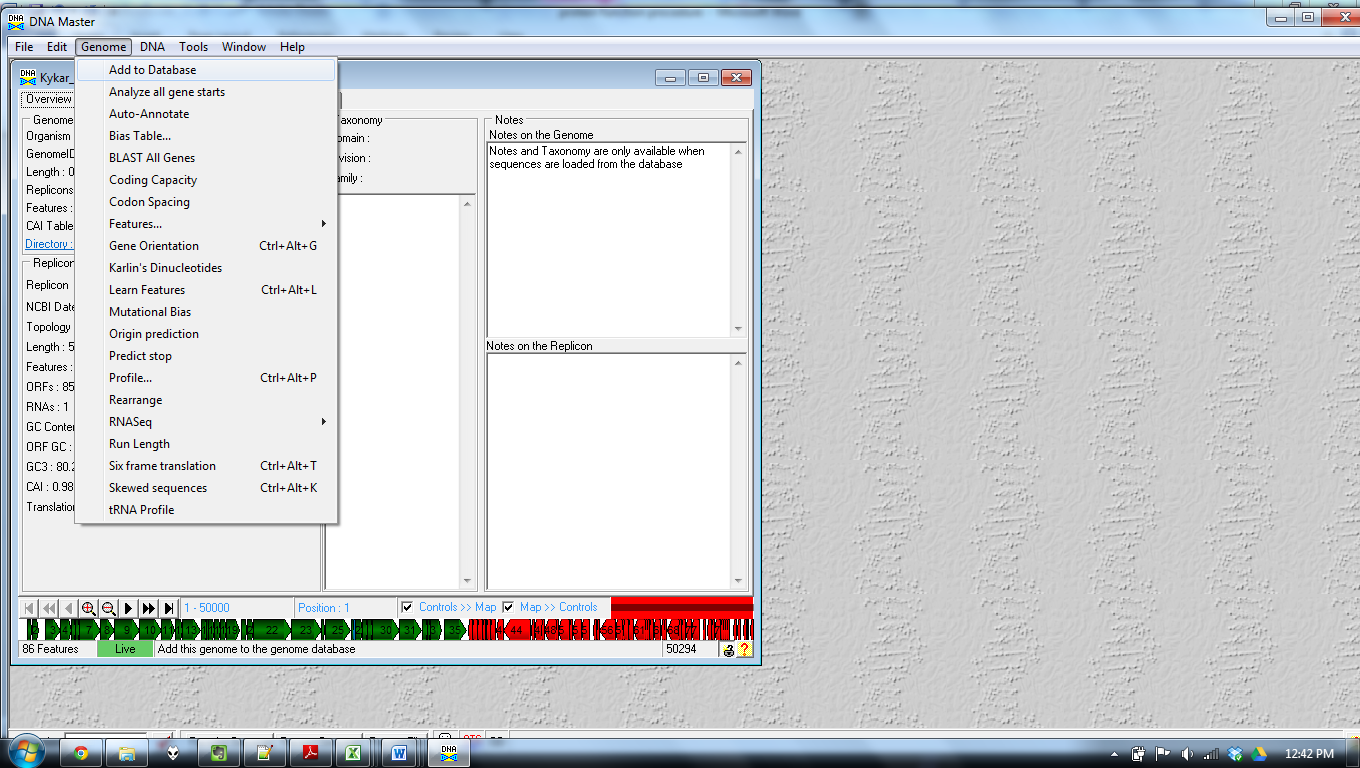


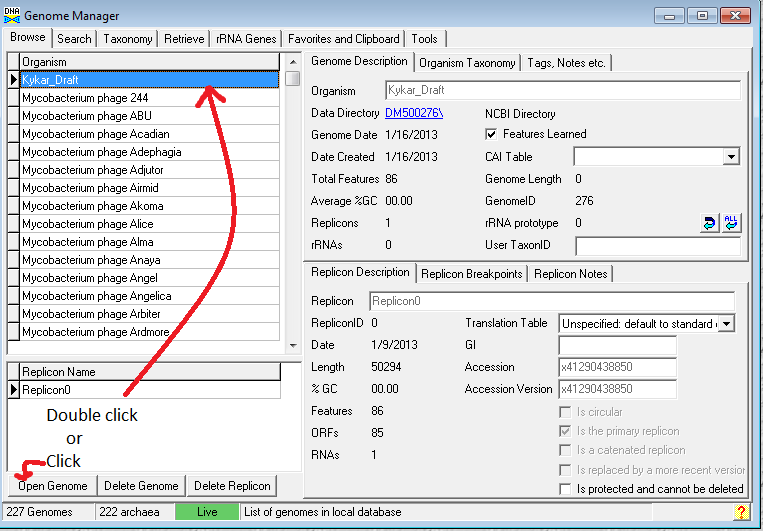
1. Click the Import/Export tab
2. Click the Import DNA Master BLAST Library button
3. Browse to the where the .dnamblib file is saved and select it.

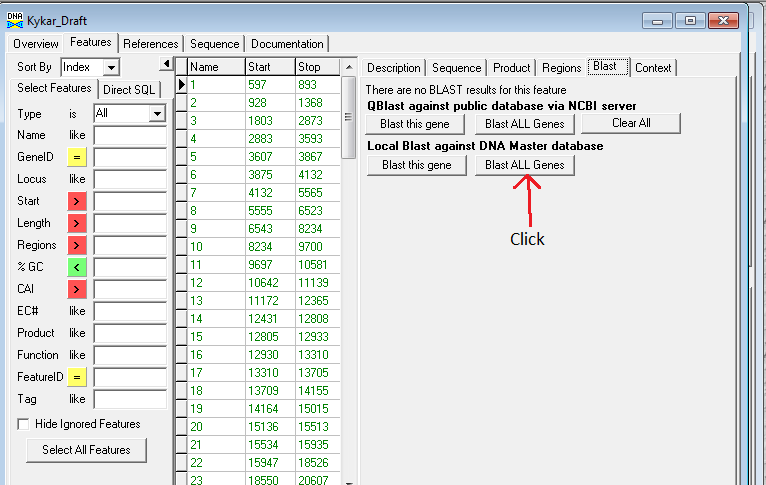
You have now installed the Blast Library of proteins of 243 genomes. The next step is to blast your genome against these.

**Local Blast of Database**

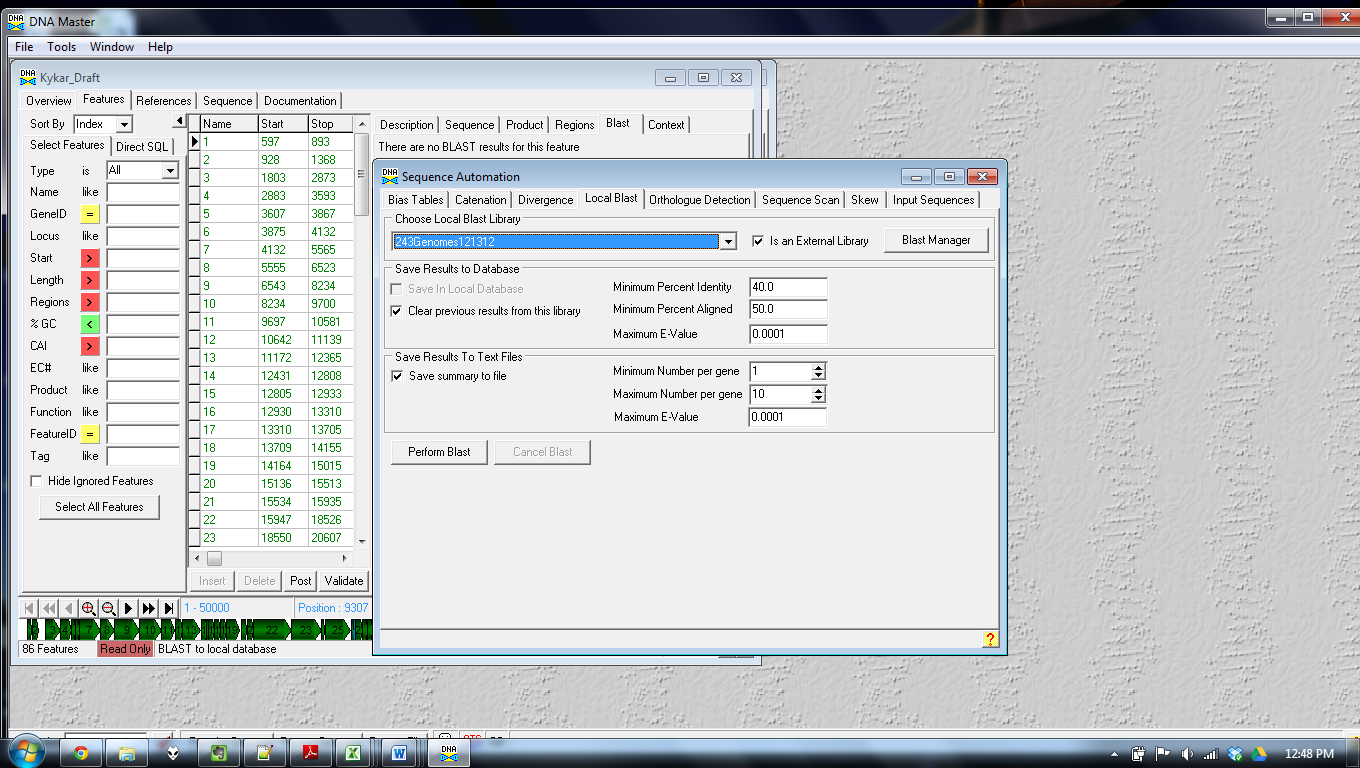
1. The genome must be added to the Genome Manager. To do this, open your genome in DNA Master. Click on Main menu🡪Genome🡪Add to Database. You can now close your genome file.



1. Now the sequence must be opened from the Genome Manager.
2. To open your file in Genome Manager: click on Tools🡪 Genome Manager
3. Double click on your genome to open the file. To blast this genome, it must be in the Genome Manager.
4. Now in this genome, go to the Feature tables and find a gene that has no blast data.
5. Click on the Blast tab and the following window appears.

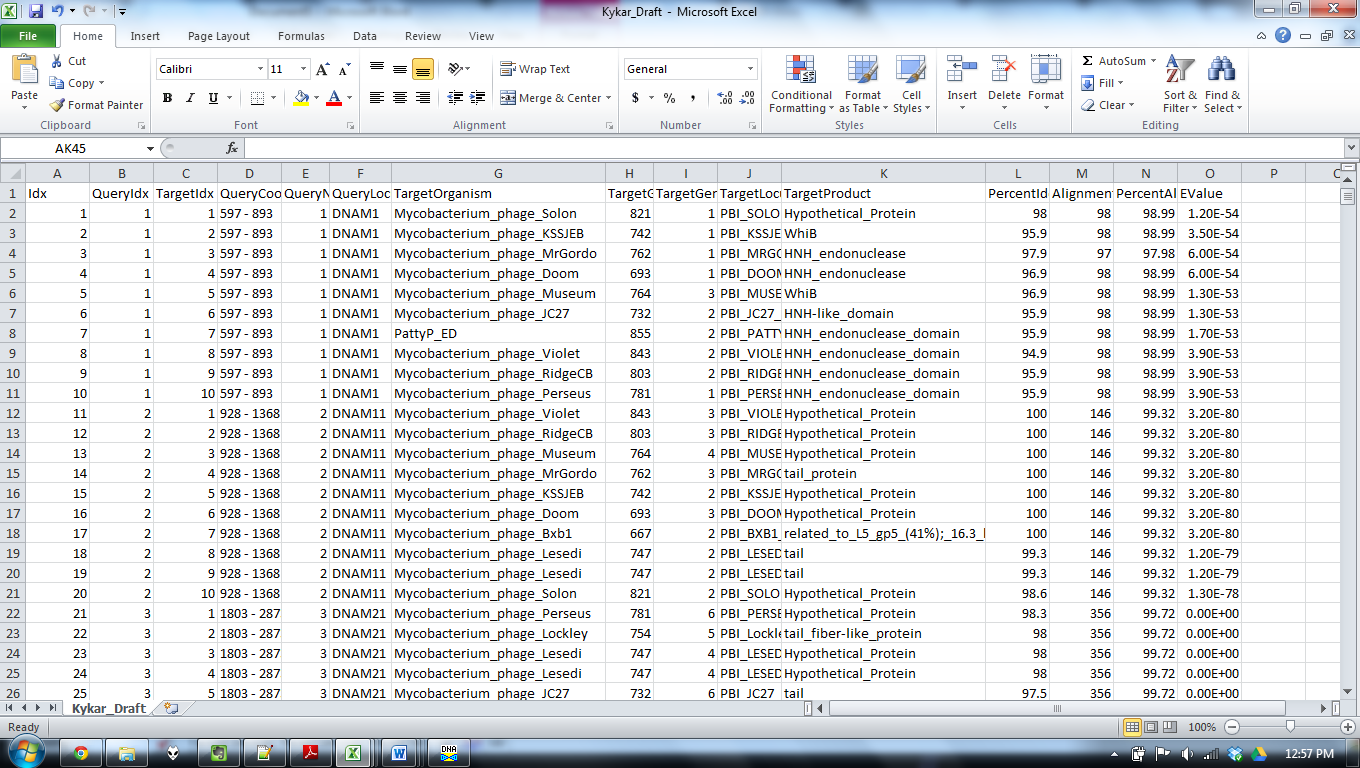


1. Find Local Blast against DNA Master database and select the button Blast ALL Genes.
2. The Sequence Automation window pops up. Choose the following features.



* + Choose the Local Blast library: 243Genomes121312.
  + Under Save Results to Database: Use default settings.
  + Under Save Results to Text Files: Checkmark the "Save summary to file" option. Use default settings for the remaining features.
  + Click the Perform Blast button and choose where to save the output file.

1. The output file is a .csv file that can be opened in Excel.



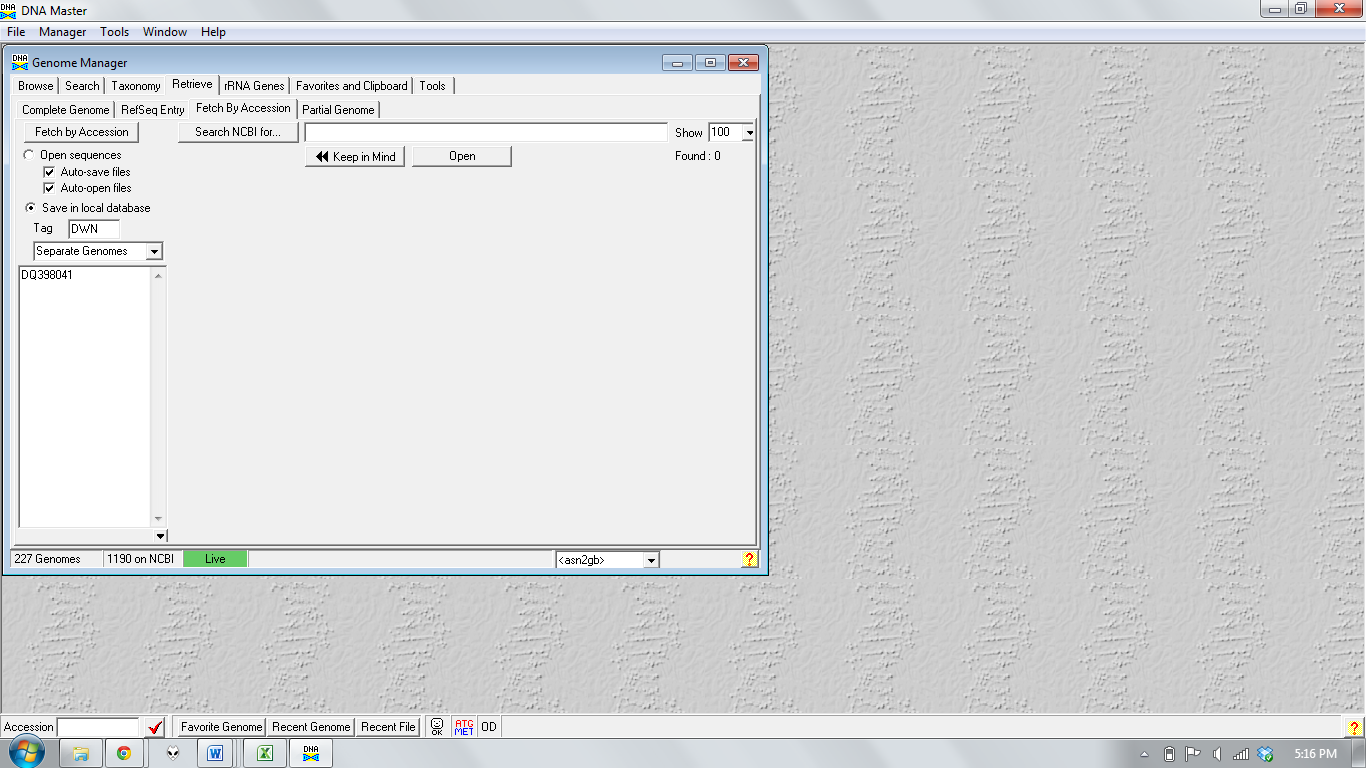
Notes:

* This is a static database created December 2012 with 243 mycobacteriophage genomes.
* At present, you can only retrieve the best 10 hits.
* Functions are present in this database as called by Phage Hunters. Information obtained should still undergo evaluation.

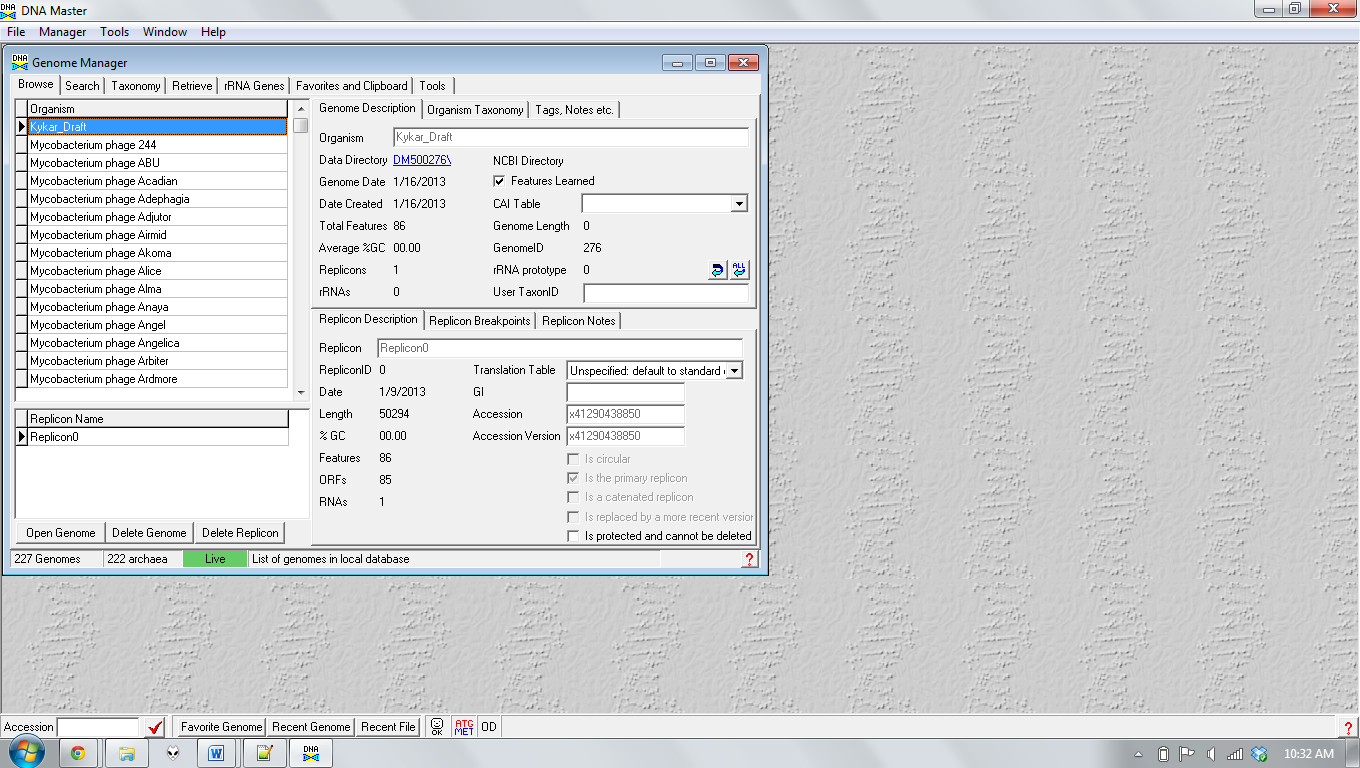
**Addendum:**

**Creation of Local Database**

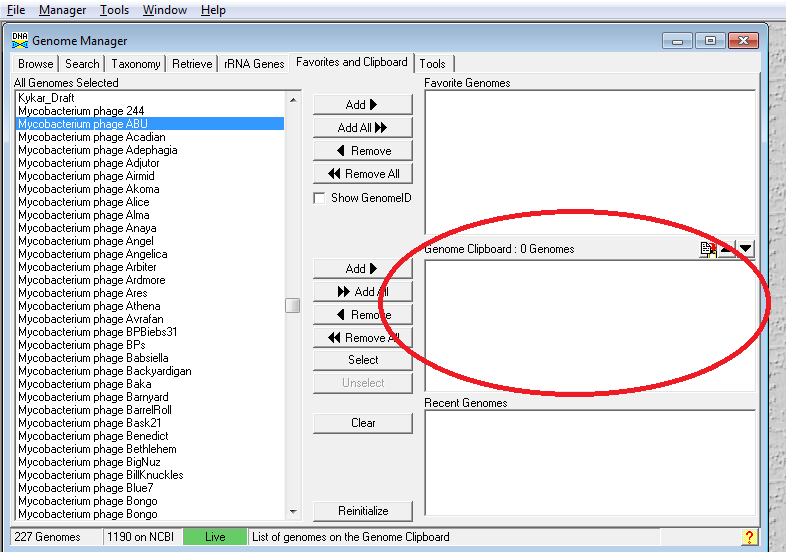
1. To create a local database, go to Tools🡪Genome Manager.
2. Select the genomes you wish to place in the library by importing desired genomes into your version of DNA Master Genome Manager. Any genome with a DNA Master file or one found in GenBank can be added. To add a DNA Master file, see first step of Local Blast of Database. To add a file from GenBank, go to the Retrieve tab in Genome Manager. From the submenu there, select the Fetch by Accession tab. If you know the accession number(s), place that number(s) in the window on the left side of the window. Once your list is complete, click Save in local database.



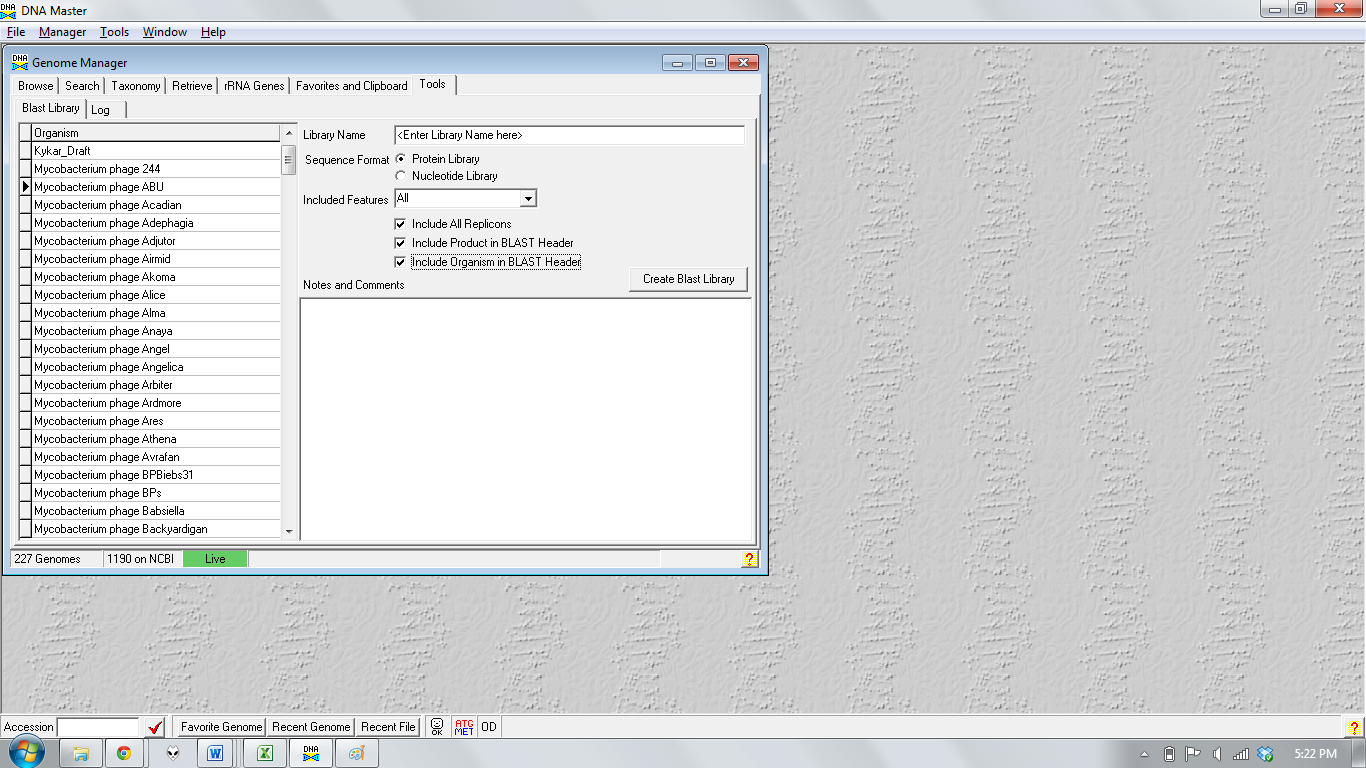
1. Click on the Browse tab. This will populate your Genome Manager with as many genomes as you choose.



1. Go to Favorites and Clipboard tab and add all genomes of interest to the clipboard (middle window).



1. Go to the Tools tab🡪 Blast Library
2. Enter a name of the library, choose a protein library, specify that the Products and Organisms by included in the BLAST header.



1. Create the library
2. This will create a .dnamblib file with the name you chose in step 5.
3. Proceed to Import of Database.

djs/mpp January, 2013