

# SEQUENCHER®

## Tutorial for Windows and Macintosh

---

### Sequencher Connections

© 2016 Gene Codes Corporation

Gene Codes Corporation

T C A G E N E  
A G T C O D E S

Gene Codes Corporation  
775 Technology Drive, Ann Arbor, MI 48108 USA  
1.800.497.4939 (USA) +1.734.769.7249 (elsewhere)  
+1.734.769.7074 (fax)

[www.genecodes.com](http://www.genecodes.com) [ginfo@genecodes.com](mailto:ginfo@genecodes.com)

## Sequencher Connections

---

About Data Sources, Sessions, and Channels .....	3
Getting Started.....	3
Launching a Connections Session with BLAST .....	4
Viewing the Results .....	5
Changing Options for a BLAST Session .....	5
Launching a Connections Session with Primer-BLAST .....	7
Changing Options for your Primer-BLAST Session .....	8
Looking at the Results Using the Schematic.....	9
Saving Primers to your Project.....	10
Launching a Connections Session with MUSCLE .....	11
Adding a Second Connections Session with MUSCLE.....	13
The Session is Saved with your Project .....	15
The Session Expires from NCBI.....	16
Restored Results.....	17
Deleting Sessions .....	17
Conclusion .....	18

# Sequencher Connections

---

**Sequencher Connections** is a whole new way of performing multiple analyses on multiple sequences in parallel. This saves you time and gives you the opportunity to look at your data using different settings or parameters. **Sequencher Connections**, at the time of writing, allows you to use **BLAST**, **Primer-BLAST**, and **Local-BLAST** (if you have installed them) on individual sequences and **MUSCLE** on a group of related sequences. For these reasons we differentiate between individual and group or grouped data when talking about Connections. Not only can you view your **BLAST**, **Primer-BLAST**, **Local-BLAST**, or **MUSCLE** results at the time you performed the analysis, but you can also review your results later on since they are saved with your project. This tutorial covers working with sessions with **BLAST**, **Primer-BLAST**, and **MUSCLE**. For information on working with **Local-BLAST** in Connections, please refer to the **Sequencher Connections** chapter of the **Sequencher** User Manual or to the **Local-BLAST** tutorial.

## ABOUT DATA SOURCES, SESSIONS, AND CHANNELS

A data source in the context of **Sequencher Connections** is a **Sequencher** project containing reads and configs. A session is a coordinated set of data and analyses that operate on that data. A channel is an analysis such as **MUSCLE** or service such as **BLAST**. We will be using these terms throughout the tutorial.

## GETTING STARTED

In this tutorial, you will learn how to use **Sequencher Connections**, how to create a session, how to name your session and change its channel options. You will also learn how to review a session that you created previously.

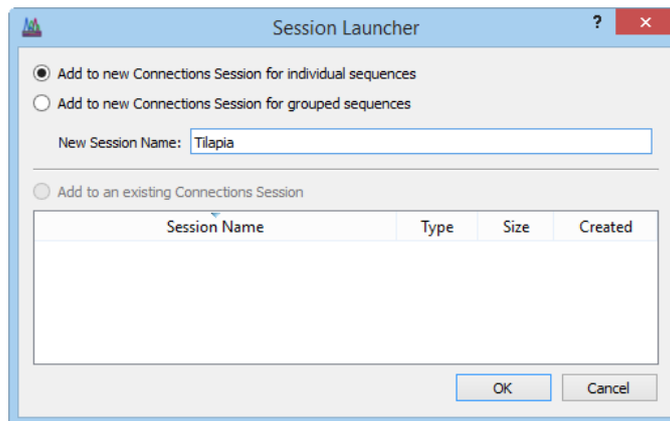
- Launch **Sequencher**.
- Go to the **File** menu and select **Import > Sequencher Project...**
- Navigate to the **Sample Data** folder inside the **Sequencher** application folder.
- Choose the **Sequencher Connections** project and select the **Open** button.
- Select **File > Save Project** and enter a name of **Sequencher Connections** and a location of your **Desktop** or **Documents** folder and select the **Save** button.
- Click on the **Label** column header in the **Project Window** to sort the project items by **Label**. This will make it easier to locate sequences later in the tutorial.

This project contains a set of fish mitochondrial sub-sequences. During the course of the tutorial, you will find out exactly what these sequences represent. These sequences will be analysed using **BLAST**. In the second part of the tutorial, you will analyse the sequences with **Primer-BLAST** and in part three, you will analyse the sequences with **MUSCLE**. There are also smaller sections on Restored Results and Deleting Sessions.

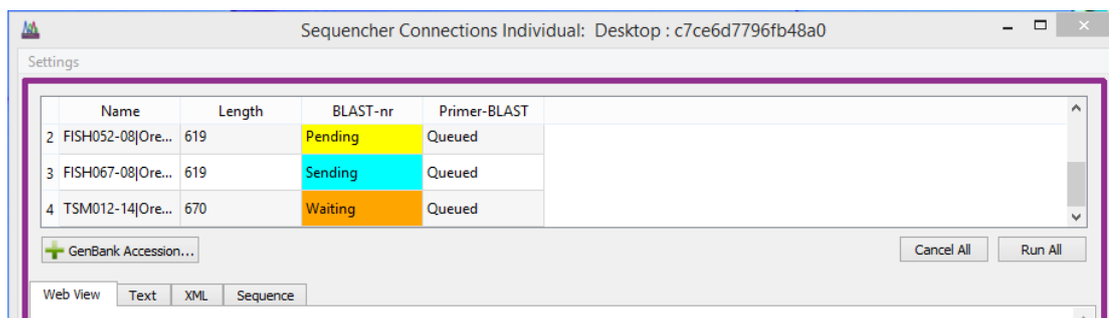
## LAUNCHING A CONNECTIONS SESSION WITH BLAST

In this session, you will use the default settings for **BLAST**. The sequences have been labeled using **Sequencher's Label** command to make it easier for you to choose the correct sequences in this tutorial.

- Select only the sequences labeled in blue (Tilapia). You should have selected 4 sequences.
- Go to the **Window** menu and select **Add to Connections Session...**
- Click on the radio button **Add to new Connections Session for individual sequences**.
- Type **Tilapia** in the **New Session Name** input field.



- Click on the **OK** button.
- A new **Sequencher Connections** session starts. You will see a window with a table pane at the top and a pane at the bottom containing a series of tabs.
- Right-click on the **BLAST-nr** column header and choose **Run on Each Sequence**.
- Watch the table as the colors of each cell in the **BLAST-nr** channel column change from white to orange to cyan to yellow and then to green and the status finally changes to **Done**. Once all the cells contain a green **Done**, your analyses are complete and you may review the results.

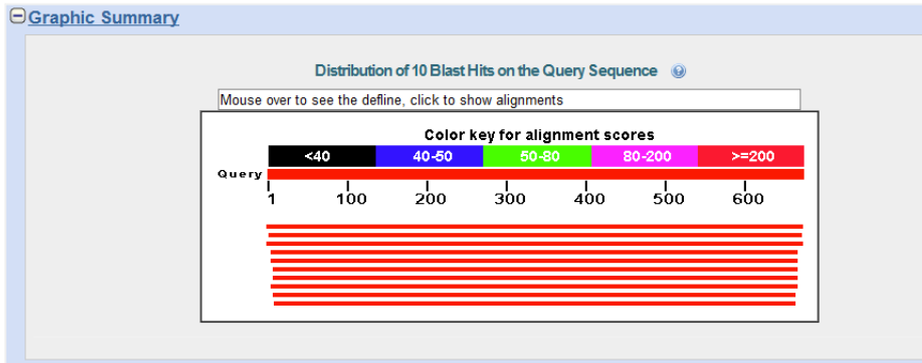


- If you get any red **Failed** statuses in the cells, try to rerun the blasts for those particular cells by right-clicking in the cells and selecting **Run**. If the status is still **Failed**, then check the **Options**.

## VIEWING THE RESULTS

Once the **BLAST** run has finished, clicking in a **Done** cell will change the view in the **Web View** tab to the familiar **BLAST** results page. You can scroll up and down, and click on links.

- Click on the **Done** cell in the BLAST-nr column for the sequence called FISH067-08 | Oreochromis in the Connections Session table.
- In the **Web View** tab, scroll down until you see the **Graphic Summary** representing the alignment of your query sequence and the alignments found by the **BLAST** algorithm.



- Click on the first red line. The web page jumps down to the alignment.

Download GenBank Graphics

Oreochromis niloticus voucher PNT-32 cytochrome oxidase subunit 1 (COI) gene, partial cds; mitochondrial

Sequence ID: [gb|KC789552.1](#) Length: 708 Number of Matches: 1

Range 1: 67 to 685 GenBank Graphics

Score	Expect	Identities	Gaps	Strand
1144 bits(619)	0.0	619/619(100%)	0/619(0%)	Plus/Plus

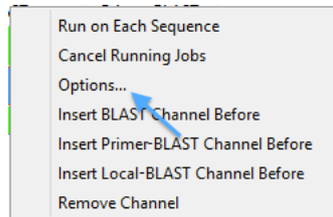
Query	Score	Subject	Score
1 GGAAGTGCCTAAGCCCTCTAATTCGGGAGCACTAAGCCAGCCCGGCTCTCTTCGGGA	60	67 GGAAGTGCCTAAGCCCTCTAATTCGGGAGCACTAAGCCAGCCCGGCTCTCTTCGGGA	126
61 GAGACCAAATCTAATAATGTAATTGTTACAGCACATGCTTTCGTAATAATTTTCITTATA	120	127 GAGACCAAATCTAATAATGTAATTGTTACAGCACATGCTTTCGTAATAATTTTCITTATA	186
121 GTAATACCAATTATGATTGGAGGCTTTGGAACTGACTAGTACCCCTCATGATTGGTGCA	180	187 GTAATACCAATTATGATTGGAGGCTTTGGAACTGACTAGTACCCCTCATGATTGGTGCA	246

- Click on the **Sequence ID** link. A new window opens displaying the **Feature Table** for the hit sequence to which your target has aligned. Notice that this is mitochondrial Cytochrome oxidase subunit 1.
- Now click on a different green **Done** cell and see if you get the same or similar results for the other sequences.

## CHANGING OPTIONS FOR A BLAST SESSION

**BLAST** offers a number of options for refining the scope of your search. Some of these options change values. One of the most important options is the E or expectation value. Another option you can change is the database to be searched. If you do not know much about the region you have just sequenced, you may want to cast the widest possible net by choosing the blastn algorithm rather than the megablast algorithm, which is the default setting. You can also ask **BLAST** to send you more than the default 10 results and descriptions as shown in this part of the tutorial below.

- Right-click in the header of the **BLAST-nr** channel.
- Choose **Insert BLAST Channel Before**.
- Right-click again in the header of the newly created **BLAST-nr** channel.
- Choose **Options...** from the context-sensitive menu.



- The **Channel Options** dialog opens. This dialog will contain a tab for each channel in your session.
- Click on **Optimize for** and change the algorithm to **Somewhat similar (blastn)**, clicking **OK** on the subsequent warning dialog.
- Change the word size to **11** to match the dialog you see below.
- Click on the **Default Graphic Color** picker, choose a new color for this channel (this will be visible in the Schematic), and click on the **OK** button.
- Change the number of alignments to **50** by choosing this value from the **Alignments** drop-down menu.
- Change the number of descriptions to **50** by choosing this value from the **Descriptions** drop-down menu.

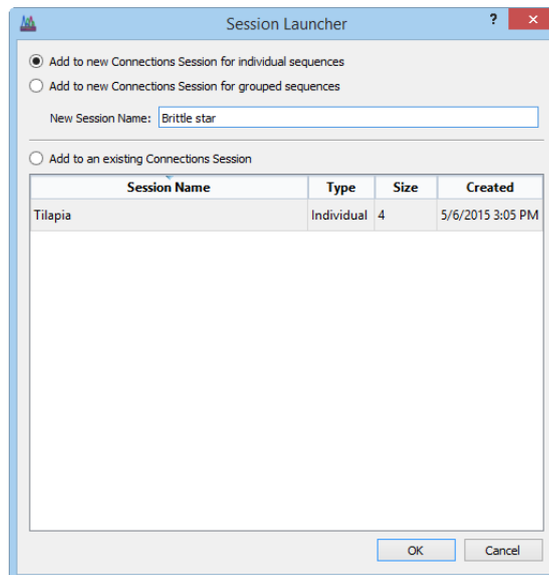
- Click the **OK** button to save these changes. This closes the **Channel Options** dialog.
- Right-click on this channel's column header and choose **Run on Each Sequence**.
- If you get any red **Failed** statuses in the cells, try to rerun the blasts for those particular cells by right-clicking in the cells and selecting **Run**.

You will see more results and descriptions with the above settings than in the first part of the tutorial. Note that, although you see more results, these are still all mitochondrial fish sequences related to the initial sequences you chose as your queries. It will not always be the case that the results are so highly related, this will depend on your query sequence as well as which database you chose to search and the settings you used.

## LAUNCHING A CONNECTIONS SESSION WITH PRIMER-BLAST

**Primer-BLAST** combines the capabilities of two resources into one. The first resource is the ability to harness the Primer 3 algorithm to predict primers and the second resource is the ability to use **BLAST** to check the specificity of those predictions. We will look at primer prediction only in this tutorial.

- Select the sequences labeled with magenta (Brittle star) in **Sequencher's Project Window**. You should have selected 4 sequences.
- Go to the **Window** menu and select **Add to Connections Session...**
- Click on the radio button **Add to new Connections Session for individual sequences**. Notice that your previous session is listed in the **Session Launcher** dialog.
- Type **Brittle star** into the **New Session Name** input field.



- Click on the **OK** button.
- A new **Sequencher Connections** session starts. You will see a window with a table pane at the top and a pane at the bottom containing a series of tabs.
- Right-click on the **Primer-BLAST** column header and choose **Run on Each Sequence**.
- Watch the table as the colors of each cell change from white to cyan to green and the status changes to **Done**. Once all the cells contain a green **Done**, your analyses are complete and you may review the results.
- If you get any red **Failed** statuses in the cells, try to rerun the blasts for those particular cells by right-clicking in the cells and selecting **Run**.
- Click on the **Done** cell for the sequence called ACAP127-12 | Ophiactis in the Primer-BLAST column. The Primer-BLAST results page loads in the **Web View** tab.

You can see your results displayed in this view with 10 sets of primer pairs in the display.

- Scroll down the page to view the detailed primer reports. You can see part of such a report in the image below.

## Detailed primer reports

Primer pair 1									
	Sequence (5'->3')	Template strand	Length	Start	Stop	Tm	GC%	Self complementarity	Self 3' complementarity
Forward primer	AAGAAAGAGGAGCCGGAACG	Plus	20	138	157	60.04	55.00	4.00	2.00
Reverse primer	ACAGTGGCATTTCGGTCCATT	Minus	20	335	316	59.96	50.00	3.00	1.00
Product length	198								

## CHANGING OPTIONS FOR YOUR PRIMER-BLAST SESSION

**Primer-BLAST** offers a number of options for refining the scope of your search for primers. Some of these options are specific to the primer calculations but you may also check on the specificity of your primers. Note that if you do check your primers specificity, the search will take longer.

- Right-click in the header of the **Primer-BLAST** channel.
- Choose **Insert Primer-BLAST Channel Before**.
- Right-click in the header of the new **Primer-BLAST** channel and choose **Options...** from the context-sensitive menu.
- The **Channel Options** dialog opens. This dialog opens at the **Primer-BLAST** channel options tab for the new channel but will also contain a tab for each channel in your session.
- Click on the **Default Graphic Color** picker and choose a new color for this channel (this will be visible in the Schematic) and select the **OK** button.
- Change the **PCR product size** minimum from **70** to **500**.
- Change the value of the **Number of primers to return** to **20**.
- Change the **Max%** for **GC Content** to **75**.

BLAST-est BLAST-nr **Primer-BLAST**

**Primer-BLAST Search Tool** Name: Primer-BLAST

This applies to: **Individual Sequences** Default Graphic Color:

Primer Parameters

Server location:

PCR product size: Minimum:  Maximum:

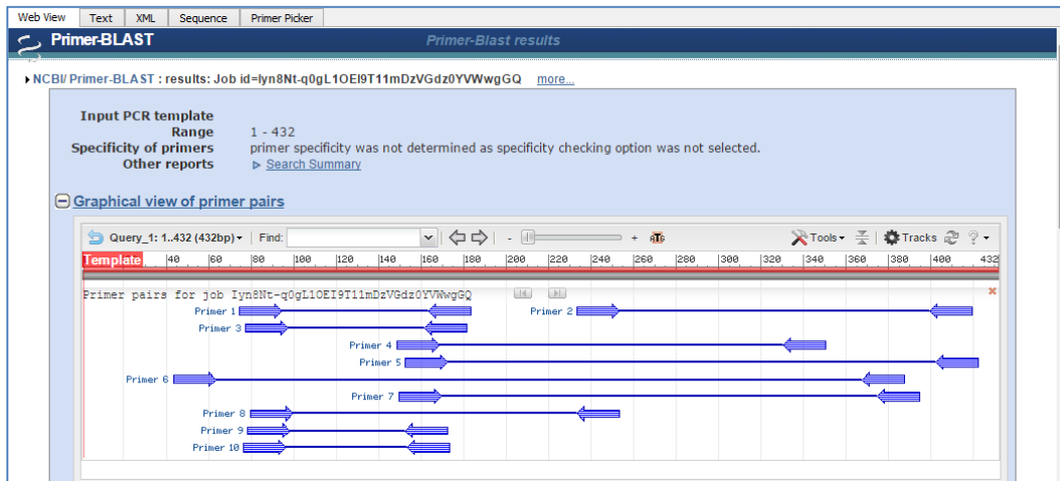
Number of primers to return:

Primer melting temperatures: Minimum:  Optimum:  Maximum:  Maximum Tm Difference:

GC Content: Min %:  Max %:  Clamp:

- Click on the **OK** button.
- Right-click on the new **Primer-BLAST** column header and choose **Run on Each Sequence**.
- Watch the table as the colors of each cell change from white to cyan to yellow to green and the status changes to **Done**. You should get a single red **Failed** status for one cell where **Primer-BLAST** failed to find any results within the minimum PCR product size which was the same as the length for this sequence.

- Click on one of the **Done** cells and click on the **Web View** tab.
- You can see your results displayed in a Graphical View with 10 sets of primer pairs in the display.



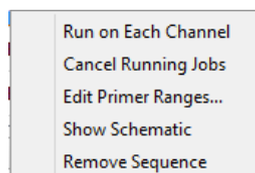
- Click on the **Text** tab. Now right-click in the tab view and choose **Save as Text...**, this saves a copy of the **Primer-BLAST** information to your chosen location.

Notice that if you insert the two sets of **Primer-BLAST** channels as neighbours, you can easily jump from one set of results to the other. In the next section, you will see how to view all the results for a single sequence using the **Schematic**.

## LOOKING AT THE RESULTS USING THE SCHEMATIC

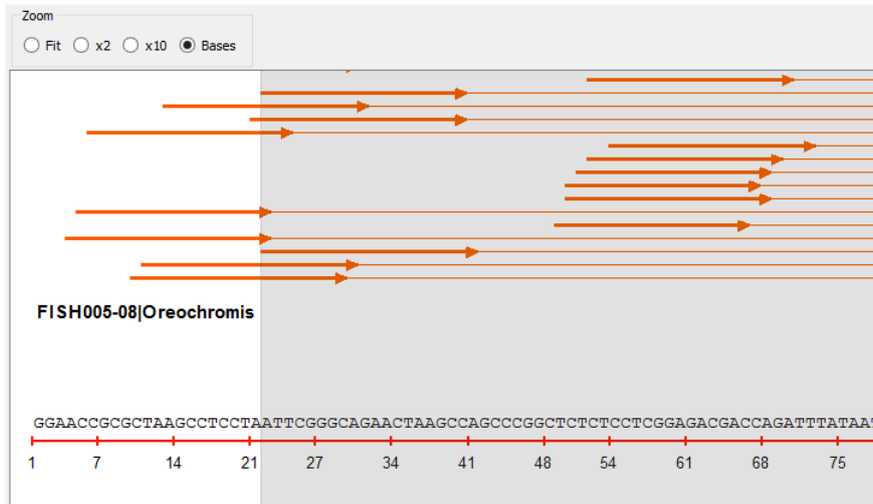
The **Schematic** enables you to combine several sets of different analyses into a single graphic. Thus for each sequence you are exploring, you can not only look at the results from several different **BLAST** searches but also combine that with the results of your **Primer-BLAST** analysis. This means that you can double check, with the aid of **BLAST**, that you are amplifying precisely the region you want and be assured of that in one single view instead of hopping about between different views or even different programs.

- Right-click on the sequence name in the second column of the session table and choose **Show Schematic**.



- A new dialog opens with the results of the all the analyses you have run on your data in one view.
- Place your cursor over one of the colored lines in the **Schematic**, note the contents of the tooltip that appears. It contains information on the hit sequence in a **BLAST** search.
- Scroll in the window until you see **Primer-BLAST** results. The arrows represent the primers and the thin lines connecting the arrows represent the insert. Place your cursor over one of the colored lines and note the contents of the tooltip. It contains information on the primer pair.

- Click on any line in the **Schematic** and see how it highlights a region corresponding to the range of the line you clicked on.
- Click on the radio button next to **Bases** and scroll to the bottom of the **Schematic**. You will see the bases contained within the range of the line you clicked on and their positions.

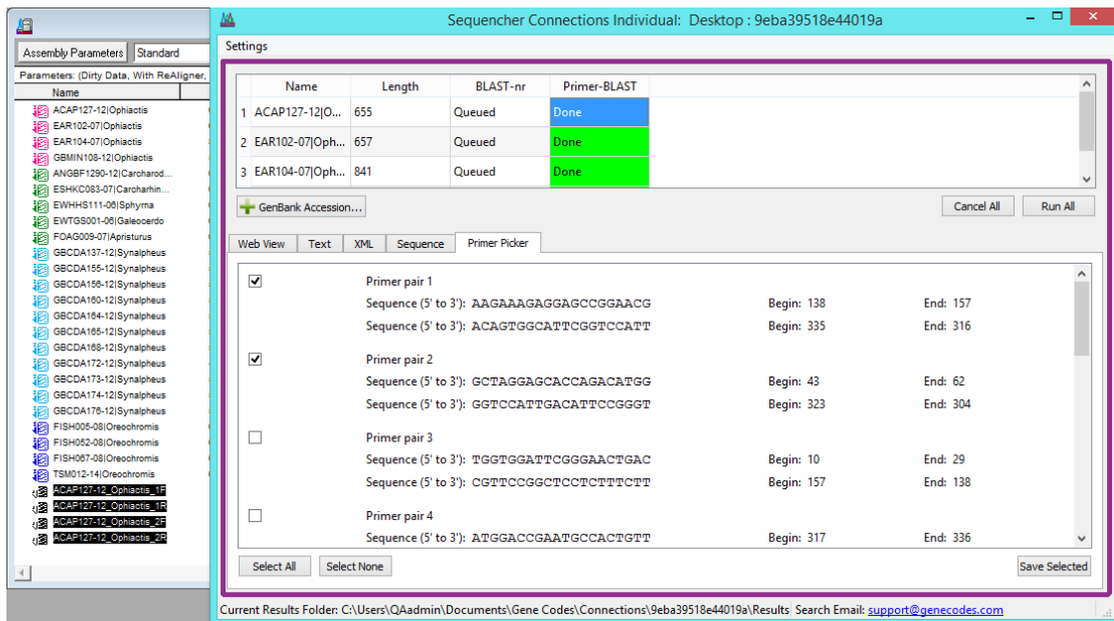


- In the bottom pane of the **Schematic** window, scroll down to view the table of results.
- Close the **Schematic** window.

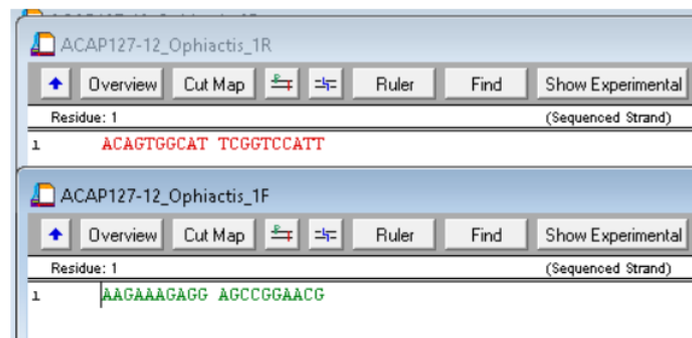
## SAVING PRIMERS TO YOUR PROJECT

Once you have reviewed the results using the **Schematic** and have decided which primers best suit your purposes, you will want to save the primers to your project. A new tab called the **Primer Picker** tab appears when you click in any cell containing a **Done** or **Done** status in a **Primer-BLAST** channel.

- Click in the **Done** cell for the sequence called **ACAP-12| Ophiactis**.
- Click on the **Primer Picker** tab and review the primers listed there.
- Click in the checkbox next each primer pair you want to save to your project.



- Click on the **Save Selected** button. The primers will appear in your project. Note that the primers contain the name of the parent sequence plus a number (for the primer pair) and either F or R for the direction of the primer.
- Double-click on the primers called **ACAP127-12\_Ophiactis\_1F** and **ACAP127-12\_Ophiactis\_1R** to open them in the **Sequence Editor**. Notice that the bases of ACAP127-12\_Ophiactis\_1F are colored green and ACAP127-12\_Ophiactis\_1R are colored red. This convention will help you to recognize the directionality of your primers simply by looking at the bases.



## LAUNCHING A CONNECTIONS SESSION WITH MUSCLE

In this session, you will use the default settings for **MUSCLE**. The sequences have been labeled using **Sequencher's Label** command to make it easier for you to choose the correct sequences in this tutorial.

- Select the sequences labeled in cyan (Snapping shrimp). You should have selected 11 sequences.
- Go to the **Window** menu and select **Add to Connections Session...**
- Click on the radio button **Add to new Connections Session for grouped sequences.**
- Type **Snapping shrimp** into the **New Session Name** input field.

Add to new Connections Session for individual sequences  
 Add to new Connections Session for grouped sequences  
 New Session Name:

- Click on the **OK** button.

A new window opens containing the grouped sequences session. You will notice that there is only one line in the table and that it is called Group 1. The name is already highlighted so that you can type in a new, more memorable name. The table gives you information on how many sequences are contained in the group.

	Name	Num Seqs	MUSCLE-1	MUSCLE-2
1	Group 1	11	Queued	Queued

- Type the name **Synalpheus** to replace the words **Group 1**.

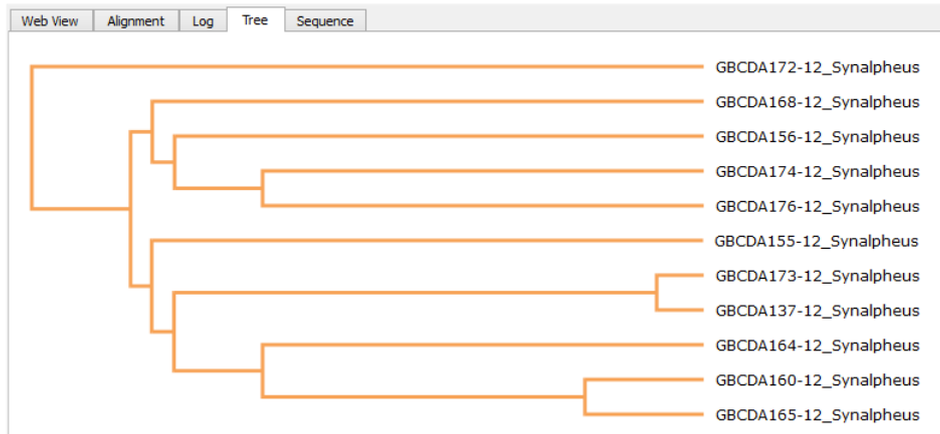
	Name	Num Seqs	MUSCLE-1	MUSCLE-2
1	Synalpheus	11	Queued	Queued

- Right-click on the MUSCLE-1 column header and choose **Run on Each Group**.
- The colors and text in the MUSCLE-1 column change in response to the status of the analysis.
- When the color in the cell is green and the text has changed to **Done**, click in the cell.
- The contents of the first tab change from the Connections landing page to a display of the aligned sequences.

```

Web View | Alignment | Log | Tree | Sequence
GBCDA172-12_Synalpheus -----CGGAATAATcTCTC
GBCDA168-12_Synalpheus -----TCaC
GBCDA156-12_Synalpheus -----TCCTAATtccCGnGCcITCGGtATAATcTCTC
GBCDA174-12_Synalpheus -----TtCTAATcCTcCGgGCTITCGGAATAATcTCTC
GBCDA176-12_Synalpheus -----TCCTAATTCIcCGaGCTITCGGAATAATTCcC
GBCDA155-12_Synalpheus -----TctTAATctTACCcGCcITtGGAAATAATTCTC
GBCDA173-12_Synalpheus -----TttTAATTTCInCCgGCTITCGGAATgATTTC
GBCDA137-12_Synalpheus gattcctgggacaccctgaagttataTttTAATTTCInCCgGCTITCGGAATgATTTC
GBCDA164-12_Synalpheus -----CCTAATtTACCgGCTITCGGtATAATTGgC
GBCDA160-12_Synalpheus -----tGGtATAATTTCaC
GBCDA165-12_Synalpheus -----TttTAATtTACCgGgITtGGtATAATTTCaC
  
```

- Click on the **Tree** tab.



- You now see a phylogram of the members of your data set.
- Right-click in the page and change the tree from rectangular to circular.
- Leave the session window open.

## ADDING A SECOND CONNECTIONS SESSION WITH MUSCLE

When you add sequences to **Sequencher Connections**, you can either add them to a new session or an existing session. If you are working with a set of related sequences you might want to add those to an existing group session so you can compare and contrast your results. The sequences are added as a new group and appear in the **Sequencher Connections** table as an additional row.

- Select all the sequences in the project - green (Sharks), magenta (Brittle stars), cyan (Snapping shrimp), and blue (Tilapia). You should have selected 24 sequences.
- Go to the **Window** menu and select **Add to Connections Session...**
- Click on the radio button **Add to an existing Connections Session.**
- Choose the **Snapping shrimp** group.

Open an existing Connections window

Session Name	Type	Size	Created
Snapping shrimp	Individual	1	5/6/2015 3:19 PM
Brittle star	Individual	4	5/6/2015 3:19 PM
Tilapia	Individual	4	5/6/2015 3:05 PM

- Click on the **OK** button.
- Notice that a new group has been added to the open session. Leave this name as **Group 2**.

	Name	Num Seqs	MUSCLE-1	MUSCLE-2
1	Synalpheus	11	Done	Queued
2	Group 2	24	Queued	Queued

- Right-click on the **MUSCLE-2** column header and choose **Run on Each Group**.
- When the color in both cells is green and the text has changed to **Done**, click in the **Group 2** cell in the **MUSCLE-2** channel. In the **Web View**, you will see the alignment for all the sequences.

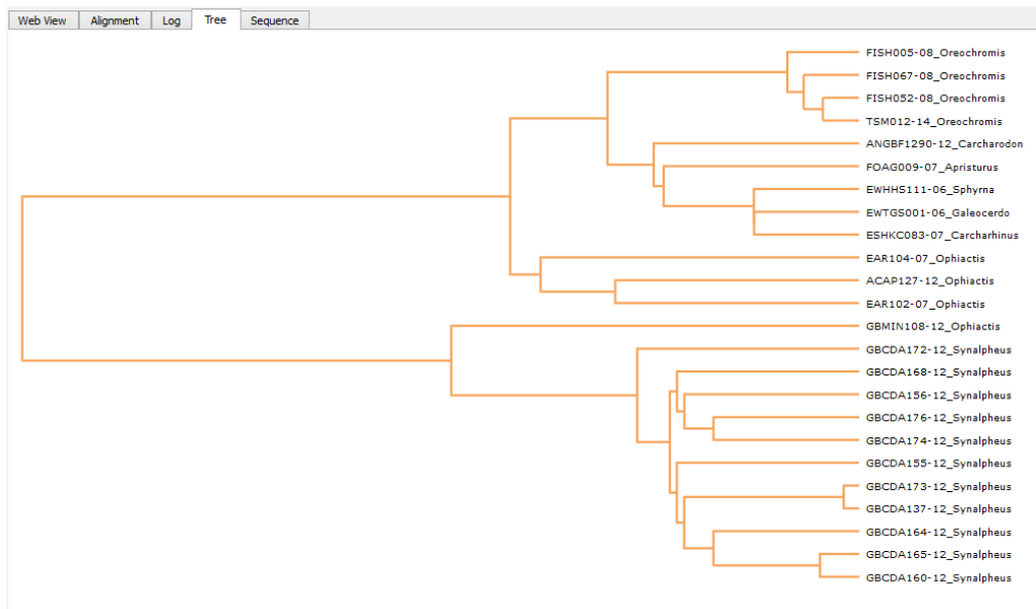
Web View	Alignment	Log	Tree	Sequence
FISH005-08_Oreochromis	-----	-----	-----	-----G
FISH067-08_Oreochromis	-----	-----	-----	-----G
FISH052-08_Oreochromis	-----	-----	-----	-----G
TSM012-14_Oreochromis	-----	-----	-----	-----G
ANGBF1290-12_Carcharodon	-----	-----	-----	-----G
FOAG009-07_Apristurus	-----	-----	-----	-----G
ENHHS111-06_Sphyrna	-----	-----	-----	-----G
EWTGS001-06_Galeocerdo	-----	-----	-----	-----G
ESHKC083-07_Carcharhinus	-----	-----	-----	-----G
EAR104-07_Ophiactis	-----	-----	-----	-----G
ACAP127-12_Ophiactis	-----	-----	-----	-----G
GBMIN108-12_Ophiactis	-----	-----	-----	-----G
GBCDA172-12_Synalpheus	-----	-----	-----	-----G
GBCDA168-12_Synalpheus	-----	-----	-----	-----G
GBCDA156-12_Synalpheus	-----	-----	-----	-----G
GBCDA176-12_Synalpheus	-----	-----	-----	-----G
GBCDA174-12_Synalpheus	-----	-----	-----	-----G
GBCDA155-12_Synalpheus	-----	-----	-----	-----G
GBCDA173-12_Synalpheus	-----	-----	-----	-----G
GBCDA137-12_Synalpheus	-----	-----	-----	-----G
GBCDA164-12_Synalpheus	-----	-----	-----	-----G
GBCDA165-12_Synalpheus	-----	-----	-----	-----G
GBCDA160-12_Synalpheus	-----	-----	-----	-----G

In this example, you can see that the Synalpheus portion of the alignment is much more fragmented, there are more gaps as a result of adding in the additional sequences, and fewer matching bases.

- Now switch to the **Alignment** tab. Here you scroll right and left to view the alignment rather than down and up with the results in the **Web View** tab.

Web View	Alignment	Log	Tree	Sequence
FISH005-08_Oreochromis	-----	-----	-----	-----G
FISH067-08_Oreochromis	-----	-----	-----	-----G
FISH052-08_Oreochromis	-----	-----	-----	-----G
TSM012-14_Oreochromis	-----	-----	-----	-----G
ANGBF1290-12_Carcharodon	-----	-----	-----	-----G
FOAG009-07_Apristurus	-----	-----	-----	-----G
ENHHS111-06_Sphyrna	-----	-----	-----	-----G
EWTGS001-06_Galeocerdo	-----	-----	-----	-----G
ESHKC083-07_Carcharhinus	-----	-----	-----	-----G
EAR104-07_Ophiactis	-----	-----	-----	-----G
ACAP127-12_Ophiactis	-----	-----	-----	-----G
GBMIN108-12_Ophiactis	-----	-----	-----	-----G
GBCDA172-12_Synalpheus	-----	-----	-----	-----G
GBCDA168-12_Synalpheus	-----	-----	-----	-----G
GBCDA156-12_Synalpheus	-----	-----	-----	-----G
GBCDA176-12_Synalpheus	-----	-----	-----	-----G
GBCDA174-12_Synalpheus	-----	-----	-----	-----G
GBCDA155-12_Synalpheus	-----	-----	-----	-----G
GBCDA173-12_Synalpheus	-----	-----	-----	-----G
GBCDA137-12_Synalpheus	-----	-----	-----	-----G
GBCDA164-12_Synalpheus	-----	-----	-----	-----G
GBCDA165-12_Synalpheus	-----	-----	-----	-----G
GBCDA160-12_Synalpheus	-----	-----	-----	-----G

- Now switch to the **Tree** tab.



- Now click in the **Done** cell in the **MUSCLE-2** Synalpheus group. In the **Tree** tab, you will see the tree for these sequences. In this way, you can visually compare the results of several sets of alignments quickly.

In this example, you added both sets of data to the same session, this makes sense for closely related data. If your data is less closely related, you can add it to a new **Sequencher Connections** session for grouped sequences instead.

## THE SESSION IS SAVED WITH YOUR PROJECT

Your session is saved with your project and can be reviewed next time you open your project. You can test this with the following steps:

- Close the project.
- You should be prompted to save. Go ahead and **Save**.
- Exit/Quit Sequencher.

The next time you open the **Sequencher Connections** project, you will be able to review the results from this tutorial as follows.

- Launch **Sequencher**.
- Go to the **File** menu and select **File > Open Project...**
- Navigate to the location you saved your project to earlier.
- Choose the **Sequencher Connections** project and select the **Open** button.
- Go to the **Window** menu and choose **Open Existing Connections Session...**
- **Save** the project if prompted to do so.

Add to new Connections Session for individual sequences  
 Add to new Connections Session for grouped sequences  
 New Session Name:

Open an existing Connections window

Session Name	Type	Size	Created
Snapping shrimp	Individual	1	5/6/2015 3:19 PM
Brittle star	Individual	4	5/6/2015 3:19 PM
Tilapia	Individual	4	5/6/2015 3:05 PM

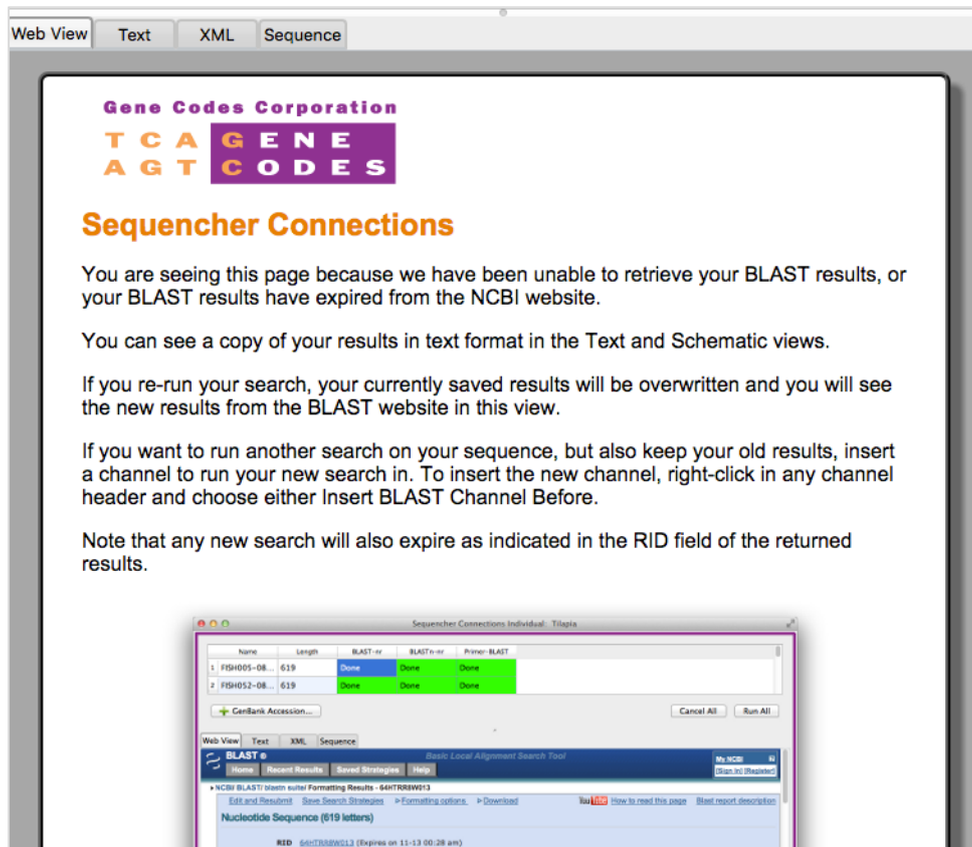
- Choose the **Snapping shrimp** Grouped session by clicking on it. Notice that it shows 2 groups are associated with this session.
- Click on the **OK** button.

The session opens and the saved results from the last time you used this project are now available for review.

### THE SESSION EXPIRES FROM NCBI BUT NOT FROM SEQUENCHER

If more than 36 hours have passed since you ran a **BLAST** or **Primer-BLAST** analysis, the results will have expired from the NCBI website. When you click on a Done status cell, **Sequencher** will display a different view in the **Web View** tab informing you that the results have expired. However, since **Sequencher Connections** has saved your results, you will still be able to view them.

- Click on the **Text** tab to view the text version of your search results.
- Right-click on the sequence name in the second column of the session table and choose **Show Schematic** to view the graphic view of your results.



Note that if you rerun any of the existing results and the runs complete, you will lose the originals. If some time has passed since you last ran a search and you suspect new data may have been added to the databases you searched against, add a new channel and just search using that channel, then use the **Schematic** view to compare the results.

## RESTORED RESULTS

If your sessions have earlier search results associated with them, those results are restored when those sessions are reopened. Any previous green **Done** statuses will have gray **Done** statuses.

Likewise, if your sessions have earlier search results associated with them and you choose to run another query, then cancel that request before the query is complete, your earlier saved results are also restored.

## DELETING SESSIONS

If you have accidentally added sessions or no longer need sessions, you can remove them from your project.

- Go to the **Window** menu and choose **Delete Existing Connections Session...**
- In the **Delete Session(s)** dialog, select the session or sessions you wish to delete and click on the **Delete Selected Session(s)** button. You will be prompted to **Cancel** or **Continue**.

If you select **Continue** and you have any open and/or running sessions, your running sessions will be stopped, closed, and removed from your project.

If you select **Continue** and you have no open and/or running sessions, your selected sessions will simply be removed from your project.

If you select **Cancel**, any running or open sessions will remain so and will not be removed from your project.

## **CONCLUSION**

In this tutorial, you have worked with **Sequencher Connections**. You have learned how to add data to a session, how to give your session a meaningful name, and how to change session options. You have used **BLAST**, **Primer-BLAST**, and **MUSCLE**. You have learned how to view your results using the **Schematic** view and how to view your **MUSCLE** results as phylograms. You have also learned that your results are saved with your project and can be reviewed at a later time. You have also learned how to save primers from a **Primer-BLAST** session to your project. Finally, you have learned that, although results expire from the NCBI website, you can always view your results in the **Text** and **Schematic** views.