# SEQUENCHER®

## **Tutorial for Windows and Macintosh**

# **De Novo Sequence Assembly with Velvet**

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# De Novo Sequence Assembly with Velvet

| About File Formats   | 3 |
|--|---|
| Getting Started  | 3 |
| Aligning Your Data with Velvet                               | 4 |
| Aligning Your Data with a Different Hash Length              | 5 |
| Using the Advanced (Edit) Options                            | 6 |
| Using the Advanced (Edit) Options To Capture Unaligned Reads | 7 |
| Conclusion   |   |

## De Novo Sequence Assembly with Velvet

Velvet is a de novo assembler, like Sequencher's default algorithm "Assemble Automatically". If you do not have a reference sequence or the reference sequence you do have is too evolutionarily distant to be useful, then you need to use de novo assembly. You can use this type of assembly whether you are working with a new organism, genome, chromosome, or region of DNA. Velvet itself consists of two programs, velveth, which analyzes the reads, and velvetg, which uses that analysis to perform the assembly.

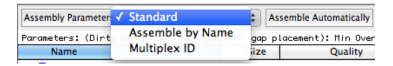
#### **ABOUT FILE FORMATS**

As with GSNAP reference-guided alignment, you can use reads in FastA or FastQ format. In single-end de novo assembly, all reads in a single file are compared to each other and contigs are built. In paired-end de novo assembly, reads from two files are used along with the paired information (distance between pairs) to construct longer scaffolds. As this is an NGS assembly, you don't need any reads to be imported into Sequencher; instead you tell Sequencher where to find the files it needs.

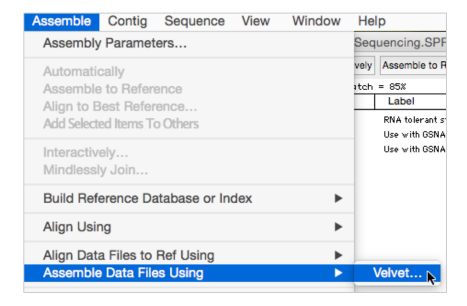
#### **GETTING STARTED**

In this tutorial, you will use one of the Next-Gen algorithms in Sequencher to align your reads. We provide a sample set of data for you to use with this tutorial.

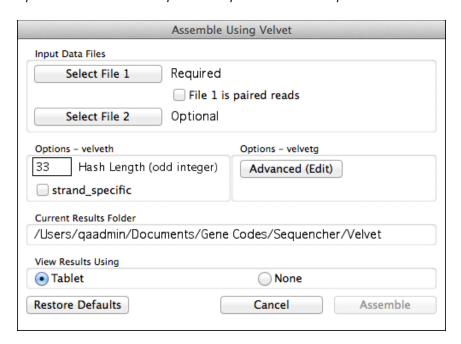
- Launch Sequencher.
- Set the Assembly Mode to Standard.



Go to the Assemble menu and select Assemble Data Files Using>Velvet...



The **Assemble Using Velvet** dialog appears. This is the launch point for your de novo assembly. You use this dialog to choose the data files that you are going to work with. You will be working with paired-end data so you will need to select two files. You may also want to decide whether you will view your results immediately or not.



- Click on the Select File 1 button.
- Navigate to the Sample Data folder inside the Sequencher application folder, then to the NGS Data folder.
- Choose read 1.fq and click on the Open button.
- Click on the Select File 2 button.
- The file picker will automatically take you to the last location you chose which, conveniently, is where the second file you need is located.
- Choose read2.fq and click on the Open button.

#### ALIGNING YOUR DATA WITH VELVET

The **Hash Length** is probably the single most important parameter when working with Velvet. It sets the minimum read length for consideration in an assembly. Setting the value higher will mean that fewer reads are used by Velvet and the assembly will occur more quickly, but the downside is that coverage may be reduced and may affect the overall accuracy.

In this tutorial, you will be looking at the effect of changing the **Hash Length** on the same data set.

Change the value in the Hash Length input field to 21.



- Click on the None radio button in the View Results Using group box.
- Click on the Assemble button.

Note how many results are returned to Sequencher's Project Window. Each result is the consensus of a contig created by Velvet (which you can view using the Tablet genome viewer). Your run may not return exactly the same results as seen in the image below.



- Click on any sequence.
- Go to the Contig menu and select Show NGS Data Using>Tablet.
- When you have finished browsing with Tablet, quit the program.
- Before moving to the next section, ensure that no data is selected in the Project Window by going to the Select menu and choosing Select None.

#### ALIGNING YOUR DATA WITH A DIFFERENT HASH LENGTH

- Go to the Assemble menu and select Assemble Data Files Using>Velvet...
- Click on the Select File 1 button.
- Navigate to the Sample Data folder inside the Sequencher application folder, then to the NGS Data folder.
- Choose **read 1.fq** and click on the **Open** button.
- Click on the **Select File 2** button.
- The file picker will automatically take you to the last location you chose which conveniently is where the second file you need is located.
- Choose read2.fq and click on the Open button.
- Change the value in the Hash Length input field to 29.
- Click on the None radio button in the View Results Using group box.
- Click on the Assemble button.

| NODE_259_length_1747_cov_2.568403            | 1775 BPs |
|--|----------|
| <b>↓</b> NODE_309_length_1552_cov_2.275773   | 1580 BPs |
| <b>∔</b> NODE_148_length_1550_cov_2.201936   | 1578 BPs |
| <b>∔</b> 2 NODE_308_length_1516_cov_2.184037 | 1544 BPs |
| <b>↓</b> NODE_276_length_1478_cov_2.102842   | 1506 BPs |

There are many consensus sequences in the Project Window. In the above image, the longest sequence's overall length is 1,775 bases. (Your results may vary from this.) Another thing to note is that, included in the name of each consensus, is the coverage of its original coverage, which has decreased. So in this case, increasing the **Hash Length** has not improved the result.

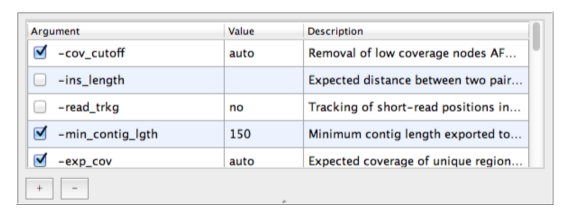
#### **USING THE ADVANCED (EDIT) OPTIONS**

Another way you can control the outcome of the assembly is to use the Advanced Parameters.

- Go to the Assemble menu and select Assemble Data Files Using>Velvet...
- Click on the Advanced (Edit) button.

The **Velvet Advanced Options** dialog opens. The dialog consists of two parts, a table with 3 columns and a **Current Parameters** preview window. You can read more about these parameters in the Velvet manual at this link. http://www.ebi.ac.uk/~zerbino/velvet/Manual.pdf.

You click in a checkbox (in the first column) to enable or disable a parameter. If you are used to using Velvet from the command line, this is the equivalent of choosing whether or not to include this argument on the command line. You click in the Value field (in the second column) to change the parameter's value. If you are uncertain about a specific parameter, then read its description (in the third column) first for more information.



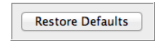
If there is a set of parameters you are not interested in, select the parameter you wish to remove and click on the — (dash) button. If there is a parameter you wish to add, click on the + (plus) button and enter the relevant information.



- Change the value for the parameter called min contig lath to 100.
- Scroll down the table to the parameter called **min\_pair\_count** and enable it by clicking on its check box.
- Click on the OK button.
- Click on the Select File 1 button.
- Navigate to the Sample Data folder inside the Sequencher application folder, then to the NGS Data folder.
- Choose read 1.fg and click on the Open button.
- Click on the Select File 2 button.
- The file picker will automatically take you to the last location you chose which conveniently is where the second file you need is located.
- Choose read2.fq and click on the Open button.
- Click on the **Assemble** button.
- The consensus sequences appear in the Project Window. Select one of the newly created consensus sequences so it is highlighted.
- Go to the Contig menu and select Show NGS Data Using>Tablet.

If you find that your results are not getting better, then you may need to revert the parameters to their original values.

Click on the Restore Defaults button on the Velvet Advanced Options dialog.



Note how the parameters in the **Current Parameters** preview window change back to the initial values you saw at the beginning of this tutorial.

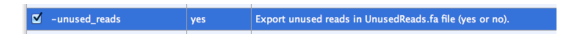
### USING THE ADVANCED (EDIT) OPTIONS TO CAPTURE UNALIGNED READS

If you wish to capture any unaligned reads you will need to use an Advanced setting.

- Go to the **Assemble** menu and select **Assemble Data Files Using>Velvet...**
- Click on the Advanced (Edit) button.

The **Velvet Advanced Options** dialog opens. The dialog consists of two parts, a table with 3 columns and a **Current Parameters** preview window.

Ensure the value for the parameter called unused reads is set to yes.



- Set any other arguments you wish to use or change.
- Click on the OK button.



- Click on the Select File 1 button.
- Navigate to the Sample Data folder inside the Sequencher application folder, then to the NGS Data folder
- Choose read 1.fq and click on the Open button.
- Click on the **Select File 2** button.
- The file picker will automatically take you to the last location you chose which conveniently is where the second file you need is located.
- Choose read2.fq and click on the Open button.
- Click on the Assemble button.
- The consensus sequences appear in the Project Window. Click anywhere in the Project Window to deselect all items.
- Select one of the newly created consensus sequences so it is highlighted. Right-click and select the Open
  External Data Folder menu item.

You will see a file called UnusedReads.fa that contains the unaligned reads in the appropriate Run folder.

#### CONCLUSION

In this tutorial, you have learned how to use the Velvet de novo assembler. You have learned how to use the advanced settings to refine its parameters in order to improve your results. You have also seen how using Sequencher's internal assembly algorithms can refine your results even further.

For more information on using Sequencher, this tutorial and others are a good place to start. You can also read the manual or consult our website by visiting http://www.genecodes.com.